

ABALOPARATIDE

Products Affected

- Tymlos

| PA Criteria | Criteria Details |
|------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | ONE OF THE FOLLOWING: (1) HIGH RISK FOR FRACTURES DEFINED AS ONE OF THE FOLLOWING: HISTORY OF OSTEOPOROTIC (I.E., FRAGILITY, LOW TRAUMA) FRACTURE(S). 2 OR MORE RISK FACTORS FOR FRACTURE (E.G., HISTORY OF MULTIPLE RECENT LOW TRAUMA FRACTURES, BMD T-SCORE LESS THAN OR EQUAL TO -2.5, CORTICOSTEROID USE, OR USE OF GNRH ANALOGS SUCH AS NAFARELIN, ETC.). NO PRIOR TREATMENT FOR OSTEOPOROSIS AND FRAX SCORE OF AT LEAST 20% FOR ANY MAJOR FRACTURE OR OF AT LEAST 3% FOR HIP FRACTURE. (2) UNABLE TO USE ORAL THERAPY (I.E., UPPER GASTROINTESTINAL PROBLEMS UNABLE TO TOLERATE ORAL MEDICATION, LOWER GASTROINTESTINAL PROBLEMS UNABLE TO ABSORB ORAL MEDICATIONS, TROUBLE REMEMBERING TO TAKE ORAL MEDICATIONS OR COORDINATING AN ORAL BISPHOSPHONATE WITH OTHER ORAL MEDICATIONS OR THEIR DAILY ROUTINE). (3) ADEQUATE TRIAL OF, INTOLERANCE TO, OR A CONTRAINDICATION TO BISPHOSPHONATES (E.G., ALENDRONATE, RISEDRONATE, IBANDRONATE). |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

ABATACEPT IV

Products Affected

- Orencia (with maltose)

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A RHEUMATOLOGIST |
| Coverage Duration | INITIAL: RA: 6 MOS. JIA: 4 MOS. PSA: 12 MOS. RENEWAL: 12 MOS ALL INDICATIONS |
| Other Criteria | INITIAL: RHEUMATOID ARTHRITIS (RA): PREVIOUS TRIAL OF HUMIRA AND ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG) SUCH AS METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE. JUVENILE IDIOPATHIC ARTHRITIS (JIA): PREVIOUS TRIAL OF HUMIRA AND ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG) SUCH AS METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE. |

ABATACEPT SQ

Products Affected

- Orenzia
- Orenzia ClickJect

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A RHEUMATOLOGIST. |
| Coverage Duration | INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS. |
| Other Criteria | INITIAL: RHEUMATOID ARTHRITIS (RA): PREVIOUS TRIAL OF HUMIRA AND ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG) SUCH AS METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE. JUVENILE IDIOPATHIC ARTHRITIS (JIA): PREVIOUS TRIAL OF HUMIRA AND ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG) SUCH AS METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE. |

ABEMACICLIB

Products Affected

- Verzenio

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | TRIAL OF OR CONTRAINDICATION TO IBRANCE (PALBOCICLIB) WHEN REQUEST IS FOR COMBINATION THERAPY WITH FULVESTRANT FOR HORMONE RECEPTOR (HR)-POSITIVE, HER2-NEGATIVE ADVANCED OR METASTATIC BREAST CANCER. |

ABIRATERONE

Products Affected

- Zytiga

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

ACALABRUTINIB

Products Affected

- Calquence

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

ADALIMUMAB

Products Affected

- Humira
- Humira Pediatric Crohn's Start
- Humira Pen
- Humira Pen Crohn's-UC-HS Start
- Humira Pen Psoriasis-Uveitis

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | INITIAL: POLYARTICULAR JUVENILE IDIOPATHIC ARTHRITIS: CURRENT WEIGHT. PLAQUE PSORIASIS: MODERATE TO SEVERE PLAQUE PSORIASIS INVOLVING GREATER THAN OR EQUAL TO 5% BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, OR GENITAL AREA. RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | RHEUMATOID ARTHRITIS, POLYARTICULAR JUVENILE IDIOPATHIC ARTHRITIS, ANKYLOSING SPONDYLITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A RHEUMATOLOGIST. PSORIATIC ARTHRITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST. PSORIASIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST. CROHN'S DISEASE/ULCERATIVE COLITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A GASTROENTEROLOGIST. |
| Coverage Duration | INITIAL:RA:6 MO PSA/AS:4 MO PJIA:5 MO PSO/CD/UC/HS:3 MO UVEITIS:6 MO RENEWAL:12 MO ALL INDICATIONS |

| PA Criteria | Criteria Details |
|-----------------------|--|
| Other Criteria | <p>INITIAL: RHEUMATOID ARTHRITIS (RA): PREVIOUS TRIAL OF ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG) SUCH AS METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE. POLYARTICULAR JUVENILE IDIOPATHIC ARTHRITIS (PJIA): PREVIOUS TRIAL OF ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG) SUCH AS METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE. PSORIATIC ARTHRITIS (PSA): PREVIOUS TRIAL OF ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG) SUCH AS METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE. ANKYLOSING SPONDYLITIS: TRIAL OF FORMULARY AGENTS NOT REQUIRED. PLAQUE PSORIASIS (PSO): PREVIOUS TRIAL OF ONE OF THE FOLLOWING CONVENTIONAL THERAPIES SUCH AS PUVA (PHOTOTHERAPY ULTRAVIOLET LIGHT A), UVB (ULTRAVIOLET LIGHT B), TOPICAL CORTICOSTEROIDS, CALCIPOTRIENE, ACITRETIN, METHOTREXATE, OR CYCLOSPORINE. CROHN'S DISEASE (CD): PREVIOUS TRIAL OF ONE CONVENTIONAL AGENT SUCH AS A CORTICOSTEROID (I.E., BUDESONIDE, METHYLPREDNISOLONE), AZATHIOPRINE, MERCAPTOPURINE, METHOTREXATE, OR MESALAMINE. ULCERATIVE COLITIS (UC): PREVIOUS TRIAL OF ONE CONVENTIONAL AGENT SUCH AS A CORTICOSTEROID (I.E., BUDESONIDE, METHYLPREDNISOLONE), AZATHIOPRINE, MERCAPTOPURINE, METHOTREXATE, OR MESALAMINE.</p> |

AFATINIB DIMALEATE

Products Affected

- Gilotrif

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

ALECTINIB

Products Affected

- Alecensa

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

ALEMTUZUMAB - LEMTRADA

Products Affected

- Lemtrada

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | INITIAL: 1 MONTH. RENEWAL: 12 MONTHS. |
| Other Criteria | TRIAL WITH TWO OF THE FOLLOWING AGENTS FOR MULTIPLE SCLEROSIS: AUBAGIO, AVONEX, GILENYA, PLEGRIDY, REBIF, TECFIDERA, OR GLATIRAMER. RENEWAL REQUESTS FOR ALEMTUZUMAB REQUIRE THAT AT LEAST 12 MONTHS HAVE ELAPSED SINCE RECEIVING THE FIRST COURSE OF LEMTRADA. PATIENTS ARE LIMITED TO TWO COURSES OF THERAPY WITH LEMTRADA WITHIN A LIFETIME. |

ALIROCUMAB

Products Affected

- Praluent Pen

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | 18 YEARS OF AGE AND OLDER. |
| Prescriber Restrictions | CARDIOLOGIST, ENDOCRINOLOGIST OR LIPIDOLOGIST |
| Coverage Duration | INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS |

| PA Criteria | Criteria Details |
|-----------------------|--|
| Other Criteria | <p>MUST HAVE A LDL CHOLESTEROL LEVEL GREATER THAN 100MG/DL WHILE ON MAXIMAL DRUG TREATMENT FOR THE PAST 2 MONTHS AND ONE OF THE FOLLOWING DIAGNOSES: (1) HETEROZYGOUS FAMILIAL HYPERCHOLESTEROLEMIA (HEFH) DETERMINED BY SIMON BROOME DIAGNOSTIC CRITERIA FOR HEFH OR A SCORE OF 6 OR GREATER ON THE DUTCH LIPID NETWORK CRITERIA FOR HEFH OR (2) HISTORY OF ATHEROSCLEROTIC CARDIOVASCULAR DISEASE (ASCVD) AS DOCUMENTED BY PHYSICIAN ATTESTATION. PATIENT MUST NOT HAVE CONCURRENT USE OF REPATHA OR OTHER PCSK9 AGENT. INITIAL THERAPY: FOR STATIN TOLERANT PATIENTS: MUST HAVE TAKEN ATORVASTATIN OR ROSUVASTATIN FOR THE PAST 2 MONTHS. FOR STATIN INTOLERANT PATIENTS: DOCUMENTATION OF STATIN INTOLERANCE BY ONE OF THE FOLLOWING: (1) PHYSICIAN ATTESTATION, (2) PATIENT HAS TRIED ROSUVASTATIN, ATORVASTATIN, OR STATIN THERAPY AT ANY DOSE AND HAS EXPERIENCED SKELETAL-MUSCLE RELATED SYMPTOMS (E.G., MYOPATHY). PATIENTS WITH CONTRAINDICATIONS TO STATINS INCLUDING ACTIVE DECOMPENSATED LIVER DISEASE, NURSING FEMALE, PREGNANCY OR PLANS TO BECOME PREGNANT OR HYPERSENSITIVITY REACTIONS WILL BE APPROVED FOR PRALUENT THERAPY WITHOUT REQUIREMENT OF DOCUMENTATION OF STATIN INTOLERANCE. RENEWAL CRITERIA: RECEIVING PRIOR PRALUENT THERAPY FOR THE PAST 6 MONTHS AND NO CLAIMS FOR REPATHA, JUXTAPID, OR KYNAMRO SINCE PRALUENT APPROVAL.</p> |

AMANTADINE

Products Affected

- Gocovri oral capsule, extended release 24hr
137 mg, 68.5 mg

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

ANAKINRA

Products Affected

- Kineret

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | RHEUMATOID ARTHRITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A RHEUMATOLOGIST. |
| Coverage Duration | INITIAL: RA: 6 MONTHS NOMID/CAPS: 12 MONTHS. RENEWAL: 12 MONTHS. |
| Other Criteria | INITIAL: RHEUMATOID ARTHRITIS (RA): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING PREFERRED AGENTS: ORENCIA, XELJANZ, CIMZIA, OR ACTEMRA. |

APREMILAST

Products Affected

- Otezla
- Otezla Starter

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | INITIAL: MODERATE TO SEVERE PLAQUE PSORIASIS INVOLVING GREATER THAN OR EQUAL TO 5% OF BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, OR GENITAL AREA. RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | PSORIATIC ARTHRITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST. PSORIASIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST |
| Coverage Duration | INITIAL: PSORIATIC ARTHRITIS: 4 MONTHS. PSORIASIS: 5 MONTHS. RENEWAL: 12 MONTHS |
| Other Criteria | INITIAL: PSORIATIC ARTHRITIS (PSA): PREVIOUS TRIAL OF HUMIRA AND ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG) SUCH AS METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE. PLAQUE PSORIASIS (PSO): PREVIOUS TRIAL OF HUMIRA AND ONE CONVENTIONAL THERAPY SUCH AS PUVA (PHOTOTHERAPY ULTRAVIOLET LIGHT A), UVB (ULTRAVIOLET LIGHT B), TOPICAL CORTICOSTEROIDS, CALCIPOTRIENE, ACITRETIN, METHOTREXATE, OR CYCLOSPORINE. |

ASFOTASE

Products Affected

- Strensiq

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | NON-SPECIFIC ALKALINE PHOSPHATASE (TNSALP) (ALPL) GENE MUTATION, SERUM ALKALINE PHOSPHATASE (ALP) LEVEL, SERUM PYRIDOXAL-5'-PHOSPHATE (PLP) LEVELS, URINE PHOSPHOETHANOLAMINE (PEA) LEVEL, RADIOGRAPHIC EVIDENCE OF HYPOPHOSPHATASIA (HPP) |
| Age Restrictions | PERINATAL/INFANTILE-ONSET HYPOPHOSPHATASIA (HPP): 6 MONTHS OF AGE OR YOUNGER AT HYPOPHOSPHATASIA (HPP) ONSET. JUVENILE-ONSET HYPOPHOSPHATASIA (HPP): 18 YEARS OF AGE OR YOUNGER AT HYPOPHOSPHATASIA (HPP) ONSET. |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH AN ENDOCRINOLOGIST, A GENETICIST, OR A METABOLIC SPECIALIST. |
| Coverage Duration | INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS. |

| PA Criteria | Criteria Details |
|-----------------------|---|
| Other Criteria | <p>INITIAL: FOR PATIENTS WITH PERINATAL/INFANTILE-ONSET HYPOPHOSPHATASIA (HPP), ALL OF THE FOLLOWING CRITERIA MUST BE MET: POSITIVE FOR A TISSUE NON-SPECIFIC ALKALINE PHOSPHATASE (TNSALP) (ALPL) GENE MUTATION AS CONFIRMED BY GENETIC TESTING OR MEETS AT LEAST TWO OF THE FOLLOWING CRITERIA: 1.) SERUM ALKALINE PHOSPHATASE (ALP) LEVEL BELOW THAT OF NORMAL RANGE FOR PATIENT AGE 2.) SERUM PYRIDOXAL-5'-PHOSPHATE (PLP) LEVELS ELEVATED AND PATIENT HAS NOT RECEIVED VITAMIN B6 SUPPLEMENTATION IN THE PREVIOUS WEEK 3.) URINE PHOSPHOETHANOLAMINE (PEA) LEVEL ABOVE THAT OF NORMAL RANGE FOR PATIENT AGE 4.) RADIOGRAPHIC EVIDENCE OF HYPOPHOSPHATASIA (HPP) (E.G., FLARED AND FRAYED METAPHYSES, OSTEOPENIA, WIDENED GROWTH PLATES, AREAS OF RADIOLUCENCY OR SCLEROSIS) 5.) PRESENCE OF TWO OR MORE OF THE FOLLOWING: RACHITIC CHEST DEFORMITY, CRANIOSYNOSTOSIS (PREMATURE CLOSURE OF SKULL BONES), DELAY IN SKELETAL GROWTH RESULTING IN DELAY OF MOTOR DEVELOPMENT, HISTORY OF VITAMIN B6 DEPENDENT SEIZURES, NEPHROCALCINOSIS, OR HISTORY OF ELEVATED SERUM CALCIUM. HISTORY OR PRESENCE OF NON-TRAUMATIC POSTNATAL FRACTURE AND DELAYED FRACTURE HEALING. FOR PATIENTS WITH JUVENILE-ONSET HYPOPHOSPHATASIA (HPP), ALL OF THE FOLLOWING CRITERIA MUST BE MET: POSITIVE FOR A TISSUE NON-SPECIFIC ALKALINE PHOSPHATASE (TNSALP) (ALPL) GENE MUTATION AS CONFIRMED BY GENETIC TESTING OR MEETS AT LEAST TWO OF THE FOLLOWING CRITERIA: 1.) SERUM ALKALINE PHOSPHATASE (ALP) LEVEL BELOW THAT OF NORMAL RANGE FOR PATIENT AGE 2.) SERUM PYRIDOXAL-5'-PHOSPHATE (PLP) LEVELS ELEVATED AND PATIENT HAS NOT RECEIVED VITAMIN B6 SUPPLEMENTATION IN THE PREVIOUS WEEK 3.)URINE PHOSPHOETHANOLAMINE (PEA) LEVEL ABOVE THAT OF NORMAL RANGE FOR PATIENT AGE 4.)RADIOGRAPHIC EVIDENCE OF HYPOPHOSPHATASIA (HPP) (E.G., FLARED AND FRAYED METAPHYSES, OSTEOPENIA, OSTEOMALACIA, WIDENED GROWTH PLATES, AREAS OF RADIOLUCENCY OR SCLEROSIS) 5.)PRESENCE OF TWO OR MORE OF THE FOLLOWING:RACHITIC DEFORMITIES (RACHITIC CHEST,</p> |

| PA Criteria | Criteria Details |
|-------------|---|
| | <p>BOWED LEGS, KNOCK-KNEES),PREMATURE LOSS OF PRIMARY TEETH PRIOR TO 5 YEARS OF AGE, DELAY IN SKELETAL GROWTH RESULTING IN DELAY OF MOTOR DEVELOPMENT, OR HISTORY OR PRESENCE OF NON-TRAUMATIC FRACTURES OR DELAYED FRACTURE HEALING. STRENSIQ WILL NOT BE APPROVED FOR THE FOLLOWING PATIENTS: PATIENTS CURRENTLY RECEIVING TREATMENT WITH A BISPHOSPHONATE [E.G., BONIVA (IBANDRONATE), FOSAMAX (ALENDRONATE), ACTONEL (RISEDRONATE)], PATIENTS WITH SERUM CALCIUM OR PHOSPHATE LEVELS BELOW THE NORMAL RANGE, PATIENTS WITH A TREATABLE FORM OF RICKETS. RENEWAL: PATIENT HAS EXPERIENCED AN IMPROVEMENT IN THE SKELETAL CHARACTERISTICS OF HYPOPHOSPHATASIA (HPP) (E.G., IMPROVEMENT OF THE IRREGULARITY OF THE PROVISIONAL ZONE OF CALCIFICATION, PHYSEAL WIDENING, METAPHYSEAL FLARING, RADIOLUCENCIES, PATCHY OSTEOSCLEROSIS, RATIO OF MID-DIAPHYSEAL CORTEX TO BONE THICKNESS, GRACILE BONES, BONE FORMATION AND FRACTURES.</p> |

ASPARAGINASE

Products Affected

- Oncaspar

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 3 MONTHS |
| Other Criteria | |

ATEZOLIZUMAB

Products Affected

- Tecentriq

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

AVELUMAB

Products Affected

- Bavencio

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

AXITINIB

Products Affected

- Inlyta oral tablet 1 mg, 5 mg

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | TRIAL OF AT LEAST ONE SYSTEMIC THERAPY FOR THE TREATMENT OF RCC SUCH AS NEXAVAR (SORAFENIB), TORISEL (TEMSIROLIMUS), SUTENT (SUNITINIB), VOTRIENT (PAZOPANIB), OR AVASTIN (BEVACIZUMAB) IN COMBINATION WITH INTERFERON. |

BECAPLERMIN

Products Affected

- Regranex

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | NON-DIABETIC ULCERS, NEOPLASM AT APPLICATION SITE, PRESSURE OR VENOUS STASIS ULCERS AND ULCERS THAT DO NOT EXTEND THROUGH THE DERMIS. |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | VASCULAR SURGEON, PODIATRIST, ENDOCRINOLOGIST, PHYSICIAN PRACTICING IN A SPECIALTY WOUND CLINIC OR INFECTIOUS DISEASE SPECIALIST. |
| Coverage Duration | 3 MONTHS |
| Other Criteria | |

BEDAQUILINE FUMARATE

Products Affected

- Sirturo

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | 18 YEARS OF AGE AND OLDER. |
| Prescriber Restrictions | |
| Coverage Duration | 24 WEEKS |
| Other Criteria | SIRTURO USED IN COMBINATION WITH AT LEAST 3 OTHER ANTIBIOTICS FOR THE TREATMENT OF PULMONARY MULTI-DRUG RESISTANT TUBERCULOSIS. |

BELIMUMAB

Products Affected

- Benlysta intravenous
- Benlysta subcutaneous

| PA Criteria | Criteria Details |
|------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | AUTOANTIBODY POSITIVE LUPUS TEST. |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS |
| Other Criteria | INITIAL: MEMBER IS CURRENTLY TAKING CORTICOSTEROIDS, ANTIMALARIALS, NSAIDS, OR IMMUNOSUPPRESSIVE AGENTS. NO APPROVAL FOR DIAGNOSIS OF SEVERE ACTIVE LUPUS NEPHRITIS, SEVERE CENTRAL NERVOUS SYSTEM LUPUS OR CONCURRENT USE OF BIOLOGIC AGENTS OR INTRAVENOUS CYCLOPHOSPHAMIDE. RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |

BELINOSTAT

Products Affected

- Beleodaq

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

BENDAMUSTINE

Products Affected

- Bendeka

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

BENRALIZUMAB

Products Affected

- Fasenra

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

BEVACIZUMAB

Products Affected

- Avastin

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

BEXAROTENE

Products Affected

- bexarotene
- Targretin topical

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

BLINATUMOMAB

Products Affected

- Blincyto intravenous kit

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | INITIAL: 3 MONTHS. RENEWAL: 12 MONTHS |
| Other Criteria | INITIAL: DIAGNOSIS OF PHILADELPHIA CHROMOSOME-NEGATIVE (PH-) B-CELL PRECURSOR ACUTE LYMPHOBLASTIC LEUKEMIA IN A PATIENT WHO HAS PREVIOUSLY TRIED CHEMOTHERAPY BUT HAS RELAPSED OR IS REFRACTORY TO TREATMENT. INITIAL APPROVAL IS FOR 2 CYCLES, MAY APPROVE FOR 1 ADDITIONAL CYCLE DUE TO TREATMENT INTERRUPTION FOR DOSE MODIFICATION. RENEWAL IS APPROVED FOR PATIENTS WHO HAVE ACHIEVED COMPLETE REMISSION WITH OR WITHOUT PARTIAL HEMATOLOGICAL RECOVERY OF PERIPHERAL BLOOD COUNTS AFTER 2 CYCLES OF TREATMENT. RENEWAL IS NOT APPROVED FOR PATIENTS WHO RECEIVED AN ALLOGENEIC HEMATOPOIETIC STEM-CELL TRANSPLANT. |

BORTEZOMIB

Products Affected

- Velcade

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

BOSUTINIB

Products Affected

- Bosulif oral tablet 100 mg, 400 mg, 500 mg

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | CML: BCR-ABL MUTATIONAL ANALYSIS CONFIRMING THAT BOTH T315I AND V299L MUTATIONS ARE NOT PRESENT. |

BOTULINUM NEUROTOXIN

Products Affected

- Botox injection recon soln 100 unit, 200 unit unit/2 mL, 2,500 unit/0.5 mL, 5,000 unit/mL
- Myobloc intramuscular solution 10,000

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | COSMETIC DIAGNOSIS: WRINKLES. |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | MIGRAINE HEADACHE: TRIAL OF TWO OF THE FOLLOWING: BETA BLOCKERS, TRICYCLIC ANTIDEPRESSANTS, OR VALPROIC ACID. OVERACTIVE BLADDER: TRIAL OF OR CONTRAINDICATION TO THE USE OF ONE ANTICHOLINERGIC MEDICATION SUCH AS ORAL OXYBUTYNIN, ORAL OXYBUTYNIN ER, TOLTERODINE, TOLTERODINE ER, TOVIAZ, TROSPIUM, OR TROSPIUM ER. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION. |

BRIGATINIB

Products Affected

- Alunbrig

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

BRODALUMAB

Products Affected

- Siliq

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | INITIAL: MODERATE TO SEVERE PLAQUE PSORIASIS INVOLVING GREATER THAN OR EQUAL TO 5% OF BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, OR GENITAL AREA. RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST. |
| Coverage Duration | INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS |
| Other Criteria | INITIAL: PLAQUE PSORIASIS (PSO): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING: COSENTYX OR OTEZLA. PATIENT HAS BEEN COUNSELED ON AND EXPRESSES UNDERSTANDING OF THE RISK OF SUICIDAL IDEATION AND BEHAVIOR. RENEWAL: PATIENT HAS NOT DEVELOPED OR REPORTED WORSENING DEPRESSIVE SYMPTOMS OR SUICIDAL IDEATION AND BEHAVIORS WHILE ON TREATMENT WITH SILIQ. |

C1 ESTERASE INHIBITOR

Products Affected

- Cinryze
- Haegarda

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | HEMATOLOGIST, IMMUNOLOGIST |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

CABOZANTINIB

Products Affected

- Cometriq

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

CABOZANTINIB S-MALATE - CABOMETYX

Products Affected

- Cabometyx oral tablet 20 mg, 40 mg, 60 mg

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | PATIENT HAS RECEIVED PRIOR ANTIANGIOGENIC THERAPY (E.G., SUTENT [SUNITINIB], VOTRIENT [PAZOPANIB], INLYTA [AXITINIB], NEXAVAR [SORAFENIB]) |

CANAKINUMAB

Products Affected

- Ilaris (PF)

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | CAPS: 4 YEARS AND OLDER. SJIA: 2 YEARS AND OLDER. |
| Prescriber Restrictions | PRESCRIBED OR SUPERVISED BY RHEUMATOLOGIST, DERMATOLOGIST, OR AN IMMUNOLOGIST. |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

CANNABINOIDS

Products Affected

- dronabinol

| PA Criteria | Criteria Details |
|------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 6 MONTHS |
| Other Criteria | B VS D COVERAGE CONSIDERATION. PART D COVERAGE CONSIDERATION FOR A DIAGNOSIS OF NAUSEA AND VOMITING ASSOCIATED WITH CANCER CHEMOTHERAPY REQUIRES A TRIAL OF OR CONTRAINDICATION TO CONVENTIONAL ANTIEMETIC THERAPIES SUCH AS ONDANSETRON, STEROIDS INDICATED FOR EMESIS OR EMEND. NO ADDITIONAL REQUIREMENTS FOR A DIAGNOSIS OF ANOREXIA ASSOCIATED WITH WEIGHT LOSS IN PATIENTS WITH AIDS. |

CANNABINOIDS ORAL SOLUTION

Products Affected

- Syndros

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 6 MONTHS |
| Other Criteria | B VS D COVERAGE CONSIDERATION. PART D COVERAGE CONSIDERATION FOR A DIAGNOSIS OF NAUSEA AND VOMITING ASSOCIATED WITH CANCER CHEMOTHERAPY REQUIRES A TRIAL OF OR CONTRAINDICATION TO FORMULARY DRONABINOL CAPSULES AND ONE CONVENTIONAL ANTIEMETIC THERAPY SUCH AS ONDANSETRON, STEROIDS INDICATED FOR EMESIS OR APREPITANT. PART D COVERAGE CONSIDERATION FOR A DIAGNOSIS OF ANOREXIA ASSOCIATED WITH WEIGHT LOSS IN PATIENTS WITH AIDS REQUIRES A TRIAL OF OR CONTRAINDICATION TO FORMULARY DRONABINOL CAPSULES. |

CARFILZOMIB

Products Affected

- Kyprolis

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

CERITINIB

Products Affected

- Zykadia

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | POSITIVE FOR ANAPLASTIC LYMPHOMA KINASE (ALK) FUSION ONCOGENE. |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

CERTOLIZUMAB PEGOL

Products Affected

- Cimzia
- Cimzia Powder for Reconst

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | RHEUMATOID ARTHRITIS/ANKYLOSING SPONDYLITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A RHEUMATOLOGIST. PSORIATIC ARTHRITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST. CROHN'S DISEASE: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A GASTROENTEROLOGIST. |
| Coverage Duration | INITIAL: RA: 6 MONTHS. PSA/AS: 4 MONTHS. CD: 12 MONTHS. RENEWAL: 12 MONTHS FOR ALL DIAGNOSES. |
| Other Criteria | INITIAL: RHEUMATOID ARTHRITIS (RA): PREVIOUS TRIAL OF HUMIRA AND ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG) SUCH AS METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE. PSORIATIC ARTHRITIS (PSA): PREVIOUS TRIAL OF HUMIRA AND ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG) SUCH AS METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE. ANKYLOSING SPONDYLITIS: PREVIOUS TRIAL OF HUMIRA. CROHN'S DISEASE (CD): PREVIOUS TRIAL OF HUMIRA AND ONE CONVENTIONAL AGENT SUCH AS A CORTICOSTEROID (I.E., BUDESONIDE, METHYLPREDNISOLONE), AZATHIOPRINE, MERCAPTOPYRINE, METHOTREXATE, OR MESALAMINE. |

CLOBAZAM

Products Affected

- Onfi oral suspension
- Onfi oral tablet 10 mg, 20 mg

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | 2 YEARS OF AGE OR OLDER |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | TRIAL OF LAMOTRIGINE OR TOPIRAMATE. REQUESTS FOR ORAL SUSPENSION APPROVABLE IF PATIENT IS UNABLE TO SWALLOW OR IS UNDER THE AGE OF 5 YEARS. |

COBIMETINIB FUMARATE

Products Affected

- Cotellic

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

COPANLISIB DI-HCL

Products Affected

- Aliqopa

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

CORTICOTROPIN

Products Affected

- Acthar H.P.

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | INFANTILE SPASMS AND MULTIPLE SCLEROSIS: 28 DAYS. OTHER FDA APPROVED INDICATIONS:12 MONTHS. |
| Other Criteria | ALL FDA APPROVED INDICATIONS EXCEPT INFANTILE SPASMS REQUIRE A TRIAL OF OR CONTRAINDICATION TO IV CORTICOSTEROIDS. NOT APPROVED FOR DIAGNOSTIC PURPOSES. |

CRIZOTINIB

Products Affected

- Xalkori

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

DABRAFENIB MESYLATE

Products Affected

- Tafinlar

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

DACLATASVIR

Products Affected

- Daklinza

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD-IDSA GUIDANCE AND ADDITIONAL CONSIDERATION FOR COVERAGE CONSISTENT WITH FDA LABELING. |
| Exclusion Criteria | |
| Required Medical Information | HCV RNA LEVEL WITHIN PAST 6 MONTHS. |
| Age Restrictions | |
| Prescriber Restrictions | GASTROENTEROLOGIST, INFECTIOUS DISEASE SPECIALIST, PHYSICIAN SPECIALIZING IN THE TREATMENT OF HEPATITIS (HEPATOLOGIST), OR A SPECIALLY TRAINED GROUP SUCH AS ECHO (EXTENSION FOR COMMUNITY HEALTHCARE OUTCOMES) MODEL. |
| Coverage Duration | CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE. |

| PA Criteria | Criteria Details |
|-----------------------|--|
| Other Criteria | <p>CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE. 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. NO APPROVALS FOR CONCURRENT USE WITH ANY OF THESE (CONTRAINDICATED OR NOT RECOMMENDED BY THE MANUFACTURER) MEDICATIONS: AMIODARONE, CARBAMAZEPINE, PHENYTOIN, OR RIFAMPIN. APPROVAL FOR INTERFERON INELIGIBLE PATIENTS - INTERFERON INELIGIBILITY INCLUDES CONCURRENT DIAGNOSIS OF AUTOIMMUNE HEPATITIS OR OTHER AUTOIMMUNE DISORDER, A KNOWN HYPERSENSITIVITY REACTION (SUCH AS URTICARIA, ANGIOEDEMA, BRONCHOCONSTRICTION AND ANAPHYLAXIS TO ALPHA INTERFERONS, PEG, OR ANY COMPONENT OF PEGINTERFERON), DOCUMENTED DEPRESSION, DECOMPENSATED HEPATIC DISEASE: A BASELINE NEUTROPHIL COUNT BELOW 1,500 PER MICROLITER, A BASELINE PLATELET COUNT BELOW 90,000, OR A BASELINE HEMOGLOBIN BELOW 10G/DL THAT HAS NOT RESPONDED TO TREATMENT.</p> |

DACLIZUMAB

Products Affected

- Zinbryta

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | PRE-EXISTING HEPATIC DISEASE OR IMPAIRMENT, INCLUDING: ACTIVE HEPATITIS B AND C, AUTOIMMUNE HEPATITIS OR OTHER AUTOIMMUNE CONDITIONS INVOLVING THE LIVER, BASELINE ALT AND AST GREATER THAN OR EQUAL TO 2 TIMES UPPER LIMIT OF NORMAL (ULN). |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | INITIAL: 3 MONTHS. RENEWAL: 12 MONTHS. |
| Other Criteria | PREVIOUS TRIAL OF TWO OF THE FOLLOWING PREFERRED AGENTS FOR MULTIPLE SCLEROSIS, SUCH AS AUBAGIO, AVONEX, GILENYA, PLEGRIDY, REBIF, TECFIDERA, OR FORMULARY GLATIRAMER ACETATE. RENEWAL: REQUESTS FOR DACLIZUMAB WILL NOT BE APPROVED FOR THE FOLLOWING PATIENTS: PATIENT WITH AUTOIMMUNE HEPATITIS OR HEPATIC INJURY. |

DALFAMPRIDINE

Products Affected

- Ampyra

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | WALKING DISABILITY SUCH AS MILD TO MODERATE BILATERAL LOWER EXTREMITY WEAKNESS OR UNILATERAL WEAKNESS PLUS LOWER EXTREMITY OR TRUNCAL ATAXIA. |
| Age Restrictions | |
| Prescriber Restrictions | NEUROLOGIST |
| Coverage Duration | INITIAL: 3 MONTHS. RENEWAL: 12 MONTHS |
| Other Criteria | RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT IN WALKING ABILITY. |

DARATUMUMAB

Products Affected

- Darzalex

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

DASATINIB

Products Affected

- Sprycel oral tablet 100 mg, 140 mg, 20 mg, 50 mg, 70 mg, 80 mg

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | PREVIOUSLY-TREATED CHRONIC MYELOID LEUKEMIA (CML) REQUIRES BCR-ABL MUTATIONAL ANALYSIS NEGATIVE FOR THE FOLLOWING MUTATIONS: T315I, V299L, T315A, F317L/V/I/C. |

DEFERASIROX

Products Affected

- Exjade
- Jadenu
- Jadenu Sprinkle

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A HEMATOLOGIST OR HEMATOLOGIST/ONCOLOGIST |
| Coverage Duration | INITIAL: 6 MONTHS RENEWAL: 12 MONTHS |
| Other Criteria | CHRONIC IRON OVERLOAD DUE TO BLOOD TRANSFUSIONS INITIAL: SERUM FERRITIN LEVEL CONSISTENTLY ABOVE 1000 MCG/L. RENEWAL: SERUM FERRITIN LEVEL CONSISTENTLY ABOVE 500 MCG/L. NON-TRANSFUSION DEPENDENT THALASSEMIA (NTDT) INITIAL: SERUM FERRITIN LEVEL CONSISTENTLY ABOVE 300 MCG/L AND LIVER IRON CONCENTRATION (LIC) OF 5 MG FE/G DRY WEIGHT OR GREATER. RENEWAL: SERUM FERRITIN LEVEL CONSISTENTLY ABOVE THAN 300 MCG/L AND LIC OF 3 MG FE/G DRY WEIGHT OR GREATER |

DEFERIPRONE

Products Affected

- Ferriprox

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A HEMATOLOGIST OR HEMATOLOGIST/ONCOLOGIST |
| Coverage Duration | INITIAL: 6 MONTHS RENEWAL: 12 MONTHS |
| Other Criteria | INITIAL CRITERIA: REQUIRES TRIAL OF EXJADE, JADENU, OR GENERIC DEFEROXAMINE AND ONE OF THE FOLLOWING CRITERIA 1) PHYSICIAN ATTESTATION THAT PATIENT IS EXPERIENCING INTOLERABLE TOXICITIES, CLINICALLY SIGNIFICANT ADVERSE EFFECTS, OR CONTRAINDICATION TO THESE THERAPIES OR 2) INADEQUATE CHELATION DEFINED BY ONE OF THE FOLLOWING: A) SERUM FERRITIN LEVEL CONSISTENTLY ABOVE 2500 MCG/L OR B) EVIDENCE OF CARDIAC IRON ACCUMULATION. RENEWAL: SERUM FERRITIN LEVELS MUST BE CONSISTENTLY ABOVE 500MCG/L |

DEFEROXAMINE

Products Affected

- deferoxamine

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | AT LEAST 3 YEARS OF AGE OR OLDER |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A HEMATOLOGIST OR HEMATOLOGIST/ONCOLOGIST |
| Coverage Duration | INITIAL: 6 MONTHS RENEWAL: 12 MONTHS |
| Other Criteria | INITIAL: SERUM FERRITIN LEVEL CONSISTENTLY ABOVE 1000MCG/L RENEWAL: SERUM FERRITIN LEVELS MUST BE CONSISTENTLY ABOVE 500MCG/L |

DEFLAZACORT

Products Affected

- Emflaza oral suspension mg
- Emflaza oral tablet 18 mg, 30 mg, 36 mg, 6

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | REPORTED INTOLERANCE TO PREDNISONE OR PREDNISOLONE IS A REPORTED INTOLERANCE IN THE EMFLAZA PRESCRIBING INFORMATION AS AN ADVERSE EVENT OF EMFLAZA |
| Required Medical Information | PHYSICIAN ATTESTATION OF GENETIC TESTING CONFIRMING DMD DIAGNOSIS. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A NEUROLOGIST SPECIALIZING IN THE TREATMENT OF DMD. |
| Coverage Duration | INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS |

| PA Criteria | Criteria Details |
|-----------------------|--|
| Other Criteria | <p>INITIAL CRITERIA: REQUIRE TRIAL OF PREDNISON OR PREDNISOLONE FOR AT LEAST 6 MONTHS. REQUEST DUE TO LACK OF EFFICACY OF PREDNISON OR PREDNISOLONE AND ALL OF THE FOLLOWING CRITERIA ARE MET: 1)PATIENT IS NOT IN STAGE 1 (PRE-SYMPTOMATIC PHASE) 2) STEROID MYOPATHY HAS BEEN RULED OUT 3) PHYSICIAN ATTESTATION OF DETERIORATION IN AMBULATION, FUNCTIONAL STATUS, OR PULMONARY FUNCTION CONSISTENT WITH ADVANCING DISEASE USING STANDARD MEASURES [SUCH AS 6-MINUTE WALK DISTANCE (6MWD), TIME TO ASCEND/DESCEND 4 STAIRS, RISE FROM FLOOR TIME (GOWER'S MANEUVER), 10-METER RUN/WALK TIME, OR NORTH STAR AMBULATORY ASSESSMENT (NSAA), PHYSICIAN GLOBAL ASSESSMENTS (PGA), PULMONARY FUNCTION (FVC, PFTS), UPPER LIMB STRENGTH (PROPELLING A WHEELCHAIR 30 FEET)]. REQUEST DUE TO ADVERSE EFFECTS OF PREDNISON OR PREDNISOLONE THAT ARE NOT LISTED IN THE PRESCRIBING INFORMATION OF EMFLAZA REQUIRE PHYSICIAN ATTESTATION OF LITERATURE SUPPORTING EMFLAZA MITIGATES NAMED ADVERSE CONSEQUENCE.</p> <p>RENEWAL: APPROVAL FOR PATIENTS CURRENTLY AMBULATORY REQUIRES PHYSICIAN ATTESTATION OF FUNCTION, STABILIZATION, OR IMPROVEMENT IN STANDARD MEASURES SINCE TREATMENT WITH EMFLAZA THAT ARE BEING MONITORED, TRACKED, AND DOCUMENTED CONSISTENTLY. APPROVAL FOR PATIENTS CURRENTLY NON-AMBULATORY REQUIRES PHYSICIAN ATTESTATION OF MAINTENANCE OR LESS THAN EXPECTED DECLINE IN PULMONARY FUNCTION AND/OR UPPER LIMB STRENGTH ASSESSED BY STANDARD MEASURES (SUCH AS PULMONARY FUNCTION [FVC, PFTS], UPPER LIMB STRENGTH MEASURES [PROPELLING A WHEELCHAIR 30 FEET], PHYSICIAN GLOBAL ASSESSMENTS [PGA]) SINCE TREATMENT WITH EMFLAZA THAT ARE BEING MONITORED, TRACKED, AND DOCUMENTED CONSISTENTLY.</p> |

DELAFLOXACIN

Products Affected

- Baxdela oral

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | ANIMAL OR HUMAN BITE, NECROTIZING FASCIITIS, DIABETIC FOOT INFECTION, DECUBITIS ULCER FORMATION, MYONECROSIS OR ECTHYMA GANGRENOSUM |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | ONE MONTH |
| Other Criteria | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH AN INFECTIOUS DISEASE SPECIALIST OR ABSSSI ORGANISM ANTIMICROBIAL SUSCEPTIBILITY TESTING SHOWS SUSCEPTIBILITY TO DELAFLOXACIN AND RESISTANCE TO ONE PREFERRED FORMULARY STANDARD OF CARE AGENT OR IF SENSITIVITY RESULTS ARE UNAVAILABLE: TRIAL OF OR CONTRAINDICATION TO ONE OF THE FOLLOWING PREFERRED FORMULARY AGENTS: A PENICILLIN, A FLUOROQUINOLONE, A CEPHALOSPORIN, OR A GRAM POSITIVE TARGETING ANTIBIOTIC |

DENOSUMAB-XGEVA

Products Affected

- Xgeva

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | DIAGNOSIS OF MULTIPLE MYELOMA |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

DESIRUDIN

Products Affected

- Iprivask

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 1 MONTH |
| Other Criteria | |

DEUTETRABENAZINE

Products Affected

- Austedo oral tablet 12 mg, 6 mg, 9 mg

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

DICHLORPHENAMIDE

Products Affected

- Keveyis

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | HEPATIC INSUFFICIENCY, PULMONARY OBSTRUCTION, OR A HEALTH CONDITION THAT WARRANTS CONCURRENT USE OF HIGH-DOSE ASPIRIN |
| Required Medical Information | |
| Age Restrictions | 18 YEARS AND OLDER |
| Prescriber Restrictions | |
| Coverage Duration | INITIAL: 2 MONTHS RENEWAL: 12 MONTHS |
| Other Criteria | RENEWAL REQUIRES THE PATIENT EXPERIENCED AT LEAST TWO FEWER ATTACKS PER WEEK FROM THEIR BASELINE |

DICLOFENAC EPOLAMINE

Products Affected

- Flector

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION. |

DICLOFENAC TOPICAL

Products Affected

- diclofenac sodium topical gel 3 % pump
- Pennsaid topical solution in metered-dose

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

DIMETHYL FUMARATE

Products Affected

- Tecfidera oral capsule, delayed release(DR/EC) 120 mg, 120 mg (14)- 240 mg (46), 240 mg

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

DINUTUXIMAB

Products Affected

- Unituxin

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

DROXIDOPA

Products Affected

- Northera

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | BLOOD PRESSURE READINGS WHILE THE PATIENT IS SITTING AND ALSO WITHIN 3 MINUTES OF STANDING FROM A SUPINE (LYING FACE UP) POSITION AT BASELINE AND RENEWAL. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A NEUROLOGIST OR CARDIOLOGIST. |
| Coverage Duration | INITIAL: 3 MONTHS RENEWAL: 12 MONTHS |
| Other Criteria | INITIAL: DIAGNOSIS OF ORTHOSTATIC HYPOTENSION AS DOCUMENTED BY 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. RENEWAL: PATIENT HAD AN INCREASE IN SYSTOLIC BLOOD PRESSURE FROM BASELINE OF AT LEAST 10 MMHG UPON STANDING FROM A SUPINE (LYING FACE UP) POSITION. |

DUPILUMAB

Products Affected

- Dupixent

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | INITIAL: PATIENT HAS MINIMUM BODY SURFACE AREA (BSA) INVOLVEMENT OF AT LEAST 10%, ECZEMA AREA AND SEVERITY INDEX (EASI) SCORE OF AT LEAST 16, OR PHYSICIAN GLOBAL ASSESSMENT (PGA) SCORE OF AT LEAST 3. RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST, ALLERGIST OR IMMUNOLOGIST |
| Coverage Duration | INITIAL: 6 MONTHS. RENEWAL:12 MONTHS |
| Other Criteria | INITIAL: PREVIOUS TRIAL OF OR CONTRAINDICATION TO TWO OF THE FOLLOWING: TOPICAL CORTICOSTEROIDS, TOPICAL CALCINEURIN INHIBITORS [E.G., ELIDEL (PIMECROLIMUS), GENERIC TACROLIMUS OINTMENT], OR TOPICAL PDE4 INHIBITOR [E.G., EUCRISA (CRISABOROLE)]. |

DURVALUMAB

Products Affected

- Imfinzi

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

EDARAVONE

Products Affected

- Radicava

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

ELBASVIR/GRAZOPREVIR

Products Affected

- Zepatier

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD-IDSA GUIDANCE AND ADDITIONAL CONSIDERATION FOR COVERAGE CONSISTENT WITH FDA LABELING. |
| Exclusion Criteria | MODERATE OR SEVERE LIVER IMPAIRMENT (CHILD PUGH B OR C) |
| Required Medical Information | HCV RNA LEVEL WITHIN PAST 6 MONTHS. FOR GENOTYPE 1A -TESTING FOR NS5A RESISTANCE-ASSOCIATED POLYMORPHISMS. |
| Age Restrictions | |
| Prescriber Restrictions | GASTROENTEROLOGIST, INFECTIOUS DISEASE SPECIALIST, PHYSICIAN SPECIALIZING IN THE TREATMENT OF HEPATITIS (HEPATOLOGIST), OR A SPECIALLY TRAINED GROUP SUCH AS ECHO (EXTENSION FOR COMMUNITY HEALTHCARE OUTCOMES) MODEL. |
| Coverage Duration | CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE. |

| PA Criteria | Criteria Details |
|-----------------------|--|
| Other Criteria | <p>CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE. 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. NO CONCURRENT USE OF SOVALDI AND ANY OF THE FOLLOWING AGENTS: PHENYTOIN, CARBAMAZEPINE, RIFAMPIN, EFAVIRENZ, ATAZANAVIR, DARUNAVIR, LOPINAVIR, SAQUINAVIR, TIPRANAVIR, CYCLOSPORINE, NAFCILLIN, KETOCONAZOLE, MODAFINIL, BOSENTAN, ETRAVIRINE, ELVITEGRAVIR/COBICISTAT/EMTRICITABINE/TENOFOVIR, ATORVASTATIN AT DOSES GREATER THAN 20MG PER DAY OR ROSUVASTATIN AT DOSES GREATER THAN 10MG PER DAY.</p> |

ELIGLUSTAT TARTRATE

Products Affected

- Cerdelga

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

ELOSULFASE ALFA

Products Affected

- Vimizim

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | LIFETIME OF MEMBERSHIP IN PLAN. |
| Other Criteria | |

ELOTUZUMAB

Products Affected

- Empliciti

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

ELTROMBOPAG

Products Affected

- Promacta

| PA Criteria | Criteria Details |
|------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | ITP:INITIAL: 2MO.RENEW:AFTER RESPONSE:12MO, INADEQUATE DOSE:2MO.HCV:12MO.SEVERE APLASTIC ANEMIA:12MO |
| Other Criteria | CHRONIC IMMUNE (IDIOPATHIC) THROMBOCYTOPENIA PURPURA (ITP): INITIAL: TRIAL OF OR CONTRAINDICATION TO CORTICOSTEROIDS, IMMUNOGLOBULINS, OR AN INSUFFICIENT RESPONSE TO SPLENECTOMY. ITP: RENEWAL: PATIENT HAS A CLINICAL RESPONSE AS DEFINED BY AN INCREASE IN PLATELET COUNT OF GREATER THAN OR EQUAL TO 50 X10 ⁹ /L (GREATER THAN OR EQUAL TO 50,000 PER UL) AT THE MAX DOSE OF 75MG PER DAY FOR 4 WEEKS. HEPATITIS C: CONCURRENT INTERFERON THERAPY. |

ENASIDENIB

Products Affected

- Idhifa

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

ENDOTHELIN RECEPTOR ANTAGONISTS

Products Affected

- Letairis
- Opsumit
- Tracleer oral tablet
- Tracleer oral tablet for suspension

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | DOCUMENTED CONFIRMATORY PULMONARY ARTERIAL HYPERTENSION (PAH) DIAGNOSIS BASED ON RIGHT HEART CATHETERIZATION. PATIENT HAS NYHA-WHO FUNCTIONAL CLASS II-IV SYMPTOMS. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST |
| Coverage Duration | INITIAL AND RENEWAL: 12 MONTHS |
| Other Criteria | INITIAL: MEAN PULMONARY ARTERY PRESSURE (PAP) OF AT LEAST 25 MMHG OR GREATER, PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 3 WOOD UNITS. LETAIRIS: PATIENT DOES NOT HAVE IDIOPATHIC PULMONARY FIBROSIS (IPF) TRACLEER: PATIENT DOES NOT HAVE ELEVATED LIVER ENZYMES (ALT, AST) MORE THAN 3 TIMES UPPER LIMIT OF NORMAL (ULN) OR INCREASES IN BILIRUBIN BY 2 OR MORE TIMES ULN. PATIENT IS NOT CONCURRENTLY TAKING CYCLOSPORINE A OR GLYBURIDE. RENEWAL: PATIENT SHOW IMPROVEMENT FROM BASELINE IN THE 6-MINUTE WALK DISTANCE OR PATIENT HAS A STABLE 6-MINUTE WALK DISTANCE WITH A STABLE/IMPROVED WHO FUNCTIONAL CLASS. |

ENZALUTAMIDE

Products Affected

- Xtandi

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | TRIAL OF OR CONTRAINDICATION TO ZYTIGA (ABIRATERONE ACETATE) IS ALSO REQUIRED IN PATIENTS WHO DO NOT HAVE A CONTRAINDICATION OR INTOLERANCE TO PREDNISONE. |

EPOPROSTENOL IV

Products Affected

- epoprostenol (glycine)

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | COVERED UNDER LOCAL COVERAGE POLICY OF APPLICABLE MEDICARE DMERC. |
| Required Medical Information | FORMULARY DRUG ADMINISTERED IN A LONG TERM CARE FACILITY TO A PATIENT WHOSE PART A COVERAGE HAS EXPIRED OR FORMULARY DRUG NOT ADMINISTERED VIA AN IMPLANTABLE PUMP OR AN EXTERNAL PUMP OR DRUG ADMINISTERED VIA AN IMPLANTABLE PUMP/AN EXTERNAL PUMP. DOCUMENTED CONFIRMATORY PULMONARY ARTERIAL HYPERTENSION (PAH) DIAGNOSIS BASED ON RIGHT HEART CATHETERIZATION. PATIENT HAS NYHA-WHO FUNCTIONAL CLASS III-IV SYMPTOMS. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST. |
| Coverage Duration | INITIAL AND RENEWAL: 12 MONTHS |
| Other Criteria | INITIAL: MEAN PULMONARY ARTERY PRESSURE (PAP) OF AT LEAST 25 MMHG OR GREATER, PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 3 WOOD UNITS. RENEWAL: PATIENT HAS SHOWN IMPROVEMENT FROM BASELINE IN THE 6-MINUTE WALK DISTANCE OR PATIENT HAS A STABLE 6-MINUTE WALK DISTANCE WITH A STABLE/ IMPROVED WHO FUNCTIONAL CLASS. |

ERLOTINIB

Products Affected

- Tarceva oral tablet 100 mg, 150 mg, 25 mg

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

ERYTHROPOIESIS STIMULATING AGENTS - EPOETIN ALFA

Products Affected

- EPOGEN 10,000 UNITS/ML VIAL SDV, P/F, OUTER
- Epogen injection solution 2,000 unit/mL, 20,000 unit/2 mL, 20,000 unit/mL, 3,000 unit/mL, 4,000 unit/mL
- Procrit 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 |
|--------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. ADDITIONAL OFF LABEL ANEMIA IN HEPATITIS C BEING TREATED IN COMBINATION WITH RIBAVIRIN AND INTERFERON ALFA OR PEGINTERFERON ALFA. |
| Exclusion Criteria | |

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Required Medical Information | <p>INITIAL: CHRONIC RENAL FAILURE (CRF) AND ANEMIA RELATED TO ZIDOVUDINE THERAPY REQUIRES A HEMOGLOBIN LEVEL OF LESS THAN 10G/DL. CANCER CHEMOTHERAPY REQUIRES A HEMOGLOBIN LEVEL OF LESS THAN 10G/DL ANEMIA DUE TO CONCURRENT HEPATITIS C TREATMENT WITH RIBAVIRIN PLUS INTERFERON ALFA/PEGINTERFERON ALFA REQUIRES A HEMOGLOBIN LEVEL LESS THAN 10G/DL AND RIBAVIRIN DOSE REDUCTION (UNLESS CONTRAINDICATED).ELECTIVE NON-CARDIAC OR NON-VASCULAR SURGERY REQUIRES A HEMOGLOBIN LEVEL LESS THAN 13G/DL. RENEWAL: CHRONIC RENAL FAILURE REQUIRES THAT THE PATIENT MEETS ONE OF THE FOLLOWING: IF THE PATIENT IS CURRENTLY RECEIVING DIALYSIS TREATMENT: 1) HEMOGLOBIN LEVEL OF LESS THAN 11G/DL OR 2) HEMOGLOBIN LEVEL THAT HAS REACHED 11G/DL AND DOSE REDUCTION/INTERRUPTION IS REQUIRED TO REDUCE THE NEED FOR BLOOD TRANSFUSIONS. IF THE PATIENT IS NOT RECEIVING DIALYSIS TREATMENT: 1) HEMOGLOBIN LEVEL OF LESS THAN 10G/DL OR 2) HEMOGLOBIN LEVEL THAT HAS REACHED 10G/DL AND DOSE REDUCTION/INTERRUPTION IS REQUIRED TO REDUCE THE NEED FOR BLOOD TRANSFUSIONS. ANEMIA DUE TO CONCURRENT HEPATITIS C TREATMENT WITH RIBAVIRIN PLUS INTERFERON ALFA/PEGINTERFERON ALFA, OR ANEMIA DUE TO ZIDOVUDINE THERAPY REQUIRES HEMOGLOBIN LEVELS BETWEEN 10G/DL AND 12G/DL. ANEMIA DUE TO EFFECT OF CONCOMITANTLY ADMINISTERED CANCER CHEMOTHERAPY REQUIRES A HEMOGLOBIN LEVEL OF LESS THAN 10 G/DL OR THAT THE HEMOGLOBIN LEVEL DOES NOT EXCEED A LEVEL NEEDED TO AVOID RBC TRANSFUSION.</p> |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | <p>ANEMIA FROM MYELOSUPPRESSIVE CHEMO/CKD WITHOUT DIALYSIS/ZIDOVUDINE:12 MONTHS.SURGERY:1 MO.HCV:6 MOS.</p> |

| PA Criteria | Criteria Details |
|-----------------------|--|
| Other Criteria | ALL INDICATIONS: TRIAL OF PROCRIT. PART D MEMBER RECEIVING DIALYSIS OR IDENTIFIED AS A PART D END STAGE RENAL DISEASE MEMBER: PAYS UNDER PART B. |

ERYTHROPOIESIS STIMULATING AGENTS - MIRCERA

Products Affected

- Mircera injection syringe 100 mcg/0.3 mL, 200 mcg/0.3 mL, 50 mcg/0.3 mL, 75 mcg/0.3 mL

| PA Criteria | Criteria Details |
|------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | INITIAL: CHRONIC RENAL FAILURE REQUIRES HEMOGLOBIN LEVELS LESS THAN 10G/DL RENEWAL: CHRONIC RENAL FAILURE REQUIRES THAT THE PATIENT MEETS ONE OF THE FOLLOWING: IF THE PATIENT IS CURRENTLY RECEIVING DIALYSIS TREATMENT: 1) HEMOGLOBIN LEVEL OF LESS THAN 11G/DL OR 2) HEMOGLOBIN LEVEL THAT HAS REACHED 11G/DL AND DOSE REDUCTION/INTERRUPTION IS REQUIRED TO REDUCE THE NEED FOR BLOOD TRANSFUSIONS. IF THE PATIENT IS NOT RECEIVING DIALYSIS TREATMENT: 1) HEMOGLOBIN LEVEL OF LESS THAN 10G/DL OR 2) HEMOGLOBIN LEVEL THAT HAS REACHED 10G/DL AND DOSE REDUCTION/INTERRUPTION IS REQUIRED TO REDUCE THE NEED FOR BLOOD TRANSFUSIONS. |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | ANEMIA DUE TO CKD WITH OR WITHOUT DIALYSIS: 12 MONTHS. |
| Other Criteria | TRIAL OF PROCRIT. PART D MEMBER RECEIVING DIALYSIS OR IDENTIFIED AS A PART D END STAGE RENAL DISEASE MEMBER: PAYS UNDER PART B. |

ETANERCEPT

Products Affected

- Enbrel
- Enbrel SureClick

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | INITIAL: PLAQUE PSORIASIS: MODERATE TO SEVERE PLAQUE PSORIASIS INVOLVING AT LEAST 5% BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, OR GENITAL AREA. RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | RHEUMATOID ARTHRITIS, ANKYLOSING SPONDYLITIS, PSORIATIC ARTHRITIS: 18 YEARS OR OLDER |
| Prescriber Restrictions | RHEUMATOID ARTHRITIS, POLYARTICULAR JUVENILE IDIOPATHIC ARTHRITIS, ANKYLOSING SPONDYLITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A RHEUMATOLOGIST. PSORIATIC ARTHRITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST. PLAQUE PSORIASIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST. |
| Coverage Duration | INITIAL: RA: 6 MONTHS. PJIA: 3 MONTHS. PSA/AS/PSO: 4 MONTHS. RENEWAL: 12 MONTHS FOR ALL DIAGNOSES |

| PA Criteria | Criteria Details |
|-----------------------|---|
| Other Criteria | <p>INITIAL: RHEUMATOID ARTHRITIS (RA): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING PREFERRED AGENTS: ORENCIA, XELJANZ, CIMZIA, OR ACTEMRA. POLYARTICULAR JUVENILE IDIOPATHIC ARTHRITIS (PJIA): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING PREFERRED AGENTS: ORENCIA OR ACTEMRA. PSORIATIC ARTHRITIS (PSA): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING PREFERRED AGENTS: CIMZIA, OTEZLA, OR COSENTYX. ANKYLOSING SPONDYLITIS (AS): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING PREFERRED AGENTS: CIMZIA OR COSENTYX. PLAQUE PSORIASIS (PSO): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING PREFERRED AGENTS: COSENTYX OR OTEZLA.</p> |

ETEPLIRSEN

Products Affected

- Exondys 51

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | PHYSICIAN ATTESTATION OF GENETIC TESTING CONFIRMING THAT MUTATION IN DUCHENNE MUSCULAR DYSTROPHY (DMD) GENE IS AMENABLE TO EXON 51 SKIPPING. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A NEUROLOGIST. |
| Coverage Duration | INITIAL: 24 WEEKS. RENEWAL: 12 MONTHS. |
| Other Criteria | INITIAL CRITERIA: PATIENT IS AMBULATORY AND IS CURRENTLY RECEIVING TREATMENT WITH OR HAS A CONTRAINDICATION TO CORTICOSTEROIDS. RENEWAL CRITERIA: PATIENT HAS 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) DURING THE PAST 24 WEEKS. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D. |

EVEROLIMUS

Products Affected

- Afinitor Disperz 7.5 mg
- Afinitor oral tablet 10 mg, 2.5 mg, 5 mg,

| PA Criteria | Criteria Details |
|------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | ADVANCED RENAL CELL CARCINOMA (RCC): TRIAL OF OR CONTRAINDICATION TO SUTENT OR NEXAVAR. |

EVOLOCUMAB

Products Affected

- Repatha Pushtronex
- Repatha SureClick
- Repatha Syringe

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | HEFH OR ASCVD: 18 YEARS OF AGE AND OLDER. |
| Prescriber Restrictions | CARDIOLOGIST, ENDOCRINOLOGIST OR LIPIDOLOGIST |
| Coverage Duration | INITIAL: 6 MONTHS RENEWAL: 12 MONTHS |

| PA Criteria | Criteria Details |
|-----------------------|--|
| Other Criteria | <p>FOR HETEROZYGOUS FAMILIAL HYPERCHOLESTEROLEMIA (HEFH) OR ATHEROSCLEROTIC CARDIOVASCULAR DISEASE (ASCVD): MUST HAVE LDL LEVEL GREATER THAN 100MG/DL ON MAXIMAL DRUG TREATMENT (MDT) FOR AT LEAST 2 MONTHS WITHIN THE PAST 2 MONTHS AND ONE OF THE FOLLOWING: (1) HEFH DETERMINED BY SIMON BROOME DIAGNOSTIC (SBD) CRITERIA OR A SCORE OF 6 OR GREATER ON THE DUTCH LIPID NETWORK (DLN) CRITERIA OR (2) ASCVD AS SUBSTANTIATED BY PHYSICIAN ATTESTATION.</p> <p>HOMOZYGOUS FAMILIAL HYPERCHOLESTEROLEMIA (HOFH): LDL LEVEL GREATER THAN 100MG/DL ON MDT FOR AT LEAST 2 MONTHS WITHIN THE PAST 2 MONTHS AND HOFH DETERMINED BY ONE OF THE FOLLOWING: 1) SBD CRITERIA, 2) A SCORE OF 8 OR GREATER ON THE DLN CRITERIA, OR 3) A 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. NO CONCURRENT USE OF OTHER PCSK9 INHIBITORS. INITIAL THERAPY: FOR STATIN TOLERANT PATIENTS: MUST HAVE TRIED MAXIMALLY TOLERATED DOSE OF HIGH INTENSITY STATIN SUCH AS ATORVASTATIN OR ROSUVASTATIN. FOR STATIN INTOLERANT PATIENTS WITH HEFH OR ASCVD: ONE OF THE FOLLOWING MUST BE MET: PHYSICIAN ATTESTATION OF STATIN INTOLERANCE (INCLUDING BUT NOT LIMITED TO MYOPATHY), OR PATIENT HAS TRIED ROSUVASTATIN OR ATORVASTATIN AT ANY DOSE. PATIENTS WITH CONTRAINDICATIONS TO STATINS INCLUDING ACTIVE DECOMPENSATED LIVER DISEASE, NURSING FEMALE, PREGNANCY OR PLANS TO BECOME PREGNANT OR HYPERSENSITIVITY REACTIONS WILL BE APPROVED FOR REPATHA THERAPY WITHOUT DOCUMENTED STATIN INTOLERANCE. FOR STATIN INTOLERANT PATIENTS WITH HOFH: MUST BE ON MAX LIPID-LOWERING THERAPY INCLUDING ONE OF THE FOLLOWING: NIACIN, BILE ACID SEQUESTRANT, LOMITAPIDE OR MIPOMERSEN. QUALIFIERS MUST PROVIDE DOCUMENTATION OF STATIN INTOLERANCE TO ONE OF THE FOLLOWING: A HIGH INTENSITY STATIN (ROSUVASTATIN OR ATORVASTATIN) OR OTHER STATIN THERAPY AT ANY DOSE. STATIN INTOLERANT PATIENTS</p> |

| PA Criteria | Criteria Details |
|-------------|--|
| | <p>MUST BE ON MAXIMAL LIPID-LOWERING MEDICATION (NON-STATIN THERAPY) FOR AT LEAST 2 MONTHS WITHIN THE PAST 2 MONTHS WITH DOCUMENTATION OF STATIN INTOLERANCE TO ATORVASTATIN OR ROSUVASTATIN OR STATIN THERAPY AT ANY DOSE. DOCUMENTATION OF STATIN INTOLERANCE INCLUDES: (1) PHYSICIAN ATTESTATION, OR (2) PATIENT HAS TRIED ROSUVASTATIN OR ATORVASTATIN AND HAS EXPERIENCED SKELETAL MUSCLE RELATED EVENTS (E.G. MYOPATHY). RENEWAL CRITERIA: RECEIVING PRIOR REPATHA THERAPY FOR AT LEAST 6 MONTHS AND NOT ON CONCURRENT THERAPY WITH OTHER PCSK9 INHIBITORS, MIPOMERSEN, OR LOMITAPIDE.</p> |

FENTANYL NASAL SPRAY

Products Affected

- Lazanda

| PA Criteria | Criteria Details |
|------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 6 MONTHS |
| Other Criteria | <p>CANCER: CURRENTLY ON A MAINTENANCE DOSE OF CONTROLLED-RELEASE OPIOID PAIN MEDICATION (SUCH AS MORPHINE SULFATE ER, OXYCODONE ER, OR FENTANYL). EITHER A TRIAL OR CONTRAINDICATION TO AT LEAST ONE (1) IMMEDIATE-RELEASE ORAL OPIOID PAIN AGENT (SUCH AS MORPHINE SULFATE IR, OXYCODONE/ASPIRIN, OXYCODONE/ACETAMINOPHEN, CODEINE/ACETAMINOPHEN, HYDROMORPHONE, OR MEPERIDINE) OR MEMBER HAS DIFFICULTY SWALLOWING TABLETS/CAPSULES AND TRIAL OR CONTRAINDICATION TO GENERIC FENTANYL CITRATE LOZENGE. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION.</p> |

FENTANYL TRANSMUCOSAL AGENTS - FENTANYL CITRATE

Products Affected

- fentanyl citrate

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 6 MONTHS |
| Other Criteria | CANCER: CURRENTLY ON A MAINTENANCE DOSE OF CONTROLLED-RELEASE OPIOID PAIN MEDICATION (SUCH AS MORPHINE SULFATE ER, OXYCODONE ER, OR FENTANYL). EITHER A TRIAL OR CONTRAINDICATION TO AT LEAST ONE (1) IMMEDIATE-RELEASE ORAL OPIOID PAIN AGENT (SUCH AS MORPHINE SULFATE IR, OXYCODONE/ASPIRIN, OXYCODONE/ACETAMINOPHEN, CODEINE/ACETAMINOPHEN, HYDROMORPHONE, OR MEPERIDINE) OR MEMBER HAS DIFFICULTY SWALLOWING TABLETS/CAPSULES. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION. |

FINGOLIMOD

Products Affected

- Gilenya

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

GEFITINIB

Products Affected

- Iressa

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

GEMTUZUMAB OZOGAMICIN

Products Affected

- Mylotarg

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

GLATIRAMER ACETATE

Products Affected

- Copaxone subcutaneous syringe 40 mg/mL
- Glatopa mg/mL, 40 mg/mL
- glatiramer subcutaneous syringe 20

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

GLECAPREVIR/PIBRENTASVIR

Products Affected

- Mavyret

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD-IDSA GUIDANCE. |
| Exclusion Criteria | MODERATE OR SEVERE HEPATIC IMPAIRMENT (CHILD PUGH B OR C) |
| Required Medical Information | HCV RNA LEVEL WITHIN PAST 6 MONTHS |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH: GASTROENTEROLOGIST, INFECTIOUS DISEASE SPECIALIST, PHYSICIAN SPECIALIZING IN THE TREATMENT OF HEPATITIS (HEPATOLOGIST), OR A SPECIALLY TRAINED GROUP SUCH AS ECHO (EXTENSION FOR COMMUNITY HEALTHCARE OUTCOMES) MODEL. |
| Coverage Duration | CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE. |

| PA Criteria | Criteria Details |
|-----------------------|--|
| Other Criteria | <p>CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE. 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. PATIENT IS NOT CONCURRENTLY TAKING ANY OF THE FOLLOWING MEDICATIONS NOT RECOMMENDED OR CONTRAINDICATED BY THE MANUFACTURER: CARBAMAZEPINE, RIFAMPIN, ETHINYL ESTRADIOL-CONTAINING MEDICATION, ATAZANAVIR, DARUNAVIR, LOPINAVIR, RITONAVIR, EFAVIRENZ, ATORVASTATIN, LOVASTATIN, SIMVASTATIN, ROSUVASTATIN AT DOSES GREATER THAN 10MG, OR CYCLOSPORINE AT DOSES GREATER THAN 100MG PER DAY. PATIENT MUST NOT HAVE PRIOR FAILURE OF A DAA REGIMEN WITH NS5A INHIBITOR AND HCV PROTEASE INHIBITOR.</p> |

GLYCEROL PHENYL BUTYRATE

Products Affected

- Ravicti

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | TRIAL OF OR CONTRAINDICATION TO SODIUM PHENYL BUTYRATE (BUPHENYL). |

GOLIMUMAB IV

Products Affected

- Simponi ARIA

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | RHEUMATOID ARTHRITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A RHEUMATOLOGIST |
| Coverage Duration | INITIAL: 6 MONTHS, RENEWAL: 12 MONTHS |
| Other Criteria | INITIAL: PREVIOUS TRIAL OF HUMIRA AND ONE OF THE FOLLOWING PREFERRED AGENTS: ORENCIA, XELJANZ, CIMZIA, OR ACTEMRA. |

GOLIMUMAB SQ

Products Affected

- Simponi

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | RHEUMATOID ARTHRITIS, ANKYLOSING SPONDYLITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A RHEUMATOLOGIST. PSORIATIC ARTHRITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST. ULCERATIVE COLITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A GASTROENTEROLOGIST. |
| Coverage Duration | INITIAL: RA: 6 MONTHS. PSA/AS: 4 MONTHS. UC: 12 MONTHS. RENEWAL: 12 MONTHS FOR ALL DIAGNOSES. |
| Other Criteria | INITIAL: RHEUMATOID ARTHRITIS (RA): PREVIOUS TRIAL OF HUMIRA AND ONE OF THE FOLLOWING PREFERRED AGENTS: ORENCIA, XELJANZ, CIMZIA OR ACTEMRA. PSORIATIC ARTHRITIS (PSA): PREVIOUS TRIAL OF HUMIRA AND ONE OF THE FOLLOWING PREFERRED AGENTS: CIMZIA, OTEZLA, OR COSENTYX. ANKYLOSING SPONDYLITIS (AS): PREVIOUS TRIAL OF HUMIRA AND ONE OF THE FOLLOWING PREFERRED AGENTS: CIMZIA OR COSENTYX. ULCERATIVE COLITIS (UC): PREVIOUS TRIAL OF HUMIRA AND ONE OF THE FOLLOWING CONVENTIONAL AGENTS SUCH AS A CORTICOSTEROID (I.E., BUDESONIDE, METHYLPREDINSONE), AZATHIOPRINE, MERCAPTOPYRINE, METHOTREXATE, OR MESALAMINE. |

GUSELKUMAB

Products Affected

- Tremfya

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | INITIAL: MODERATE TO SEVERE PLAQUE PSORIASIS INVOLVING GREATER THAN OR EQUAL TO 5% OF BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, OR GENITAL AREA. RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST. |
| Coverage Duration | INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS |
| Other Criteria | INITIAL: PLAQUE PSORIASIS (PSO): PREVIOUS TRIAL WITH HUMIRA FOLLOWED BY ONE OF THE FOLLOWING: COSENTYX OR OTEZLA. |

HYDROXYPROGESTERONE CAPROATE- DELALUTIN GENERIC

Products Affected

- hydroxyprogesterone caproate

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| 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. |

IBRUTINIB

Products Affected

- Imbruvica

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

IBUPROFEN-FAMOTIDINE

Products Affected

- Duexis

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| 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, NIZATIDINE, OR RANITIDINE, AND TRIAL OF GENERIC, FEDERAL LEGEND IBUPROFEN. |

IDELALISIB

Products Affected

- Zydelig

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

IMATINIB MESYLATE

Products Affected

- imatinib oral tablet 100 mg, 400 mg

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | ALL DIAGNOSES: 12 MONTHS. ADJUVANT GASTROINTESTINAL STROMAL TUMOR (GIST) TREATMENT: 36 MONTHS. |
| Other Criteria | PATIENTS WITH PREVIOUSLY-TREATED CML REQUIRE A BCR-ABL MUTATIONAL ANALYSIS CONFIRMING THAT THE PATIENT IS NEGATIVE FOR THE FOLLOWING MUTATIONS: T315I, V299L, F317L/V/I/C, Y253H, E255K/V, F359V/C/I. |

IMIQUIMOD - ALDARA

Products Affected

- imiquimod

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | ACTINIC KERATOSIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST. SUPERFICIAL BASAL CELL CARCINOMA: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST OR AN ONCOLOGIST. |
| Coverage Duration | 4 MONTHS |
| Other Criteria | EXTERNAL GENITAL WARTS: TRIAL OF PODOFILOX (CONDYLOX) 0.5% TOPICAL SOLUTION. ACTINIC KERATOSIS BRAND DRUG REQUEST: TRIAL OF GENERIC IMIQUIMOD 5% CREAM. SUPERFICIAL BASAL CELL CARCINOMA: LESS THAN 2CM IN SIZE AND NOT ON THE FACE. |

INFLIXIMAB

Products Affected

- Remicade

| PA Criteria | Criteria Details |
|------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | INITIAL: PLAQUE PSORIASIS: MODERATE TO SEVERE PLAQUE PSORIASIS INVOLVING GREATER THAN OR EQUAL TO 5 % BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, OR GENITAL AREA. RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | RHEUMATOID ARTHRITIS, ANKYLOSING SPONDYLITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A RHEUMATOLOGIST. PSORIATIC ARTHRITIS PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST. PSORIASIS PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST. CROHN'S DISEASE/ULCERATIVE COLITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A GASTROENTEROLOGIST. |
| Coverage Duration | INITIAL: CD/UC: 8 MO. RA: 6 MO. PSA/AS/PSO: 4 MO. RENEWAL FOR ALL INDICATIONS: 12 MO. |

| PA Criteria | Criteria Details |
|-----------------------|---|
| Other Criteria | <p>INITIAL: RHEUMATOID ARTHRITIS (RA): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING: ORENCIA, XELJANZ, CIMZIA, OR ACTEMRA. PSORIATRIC ARTHRITIS (PSA): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING: CIMZIA, OTEZLA, OR COSENTYX. ANKYLOSING SPONDYLITIS (AS): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY CIMZIA OR COSENTYX. PLAQUE PSORIASIS (PSO): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING: COSENTYX OR OTEZLA. CROHN'S DISEASE (CD): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY CIMZIA. ULCERATIVE COLITIS (UC): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING: SIMPONI OR ENTYVIO.</p> |

INFLIXIMAB-ABDA

Products Affected

- Renflexis

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | INITIAL: PLAQUE PSORIASIS: MODERATE TO SEVERE PLAQUE PSORIASIS INVOLVING GREATER THAN OR EQUAL TO 5% BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, OR GENITAL AREA. RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | RHEUMATOID ARTHRITIS, ANKYLOSING SPONDYLITIS: PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. PSORIATIC ARTHRITIS PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST. PSORIASIS PRESCRIBED BY OR IN CONSULTATION WITH A DERMATOLOGIST. CROHN'S DISEASE/ULCERATIVE COLITIS: GASTROENTEROLOGIST. |
| Coverage Duration | INITIAL: CD/UC: 8 MOS. RA: 6 MOS. PSA/AS/PSO: 4 MOS. RENEWAL FOR ALL INDICATIONS: 12 MOS. |

| PA Criteria | Criteria Details |
|-----------------------|---|
| Other Criteria | <p>INITIAL: RHEUMATOID ARTHRITIS (RA): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING: ORENCIA, XELJANZ, CIMZIA, OR ACTEMRA. PSORIATRIC ARTHRITIS (PSA): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING: CIMZIA, OTEZLA, OR COSENTYX. ANKYLOSING SPONDYLITIS (AS): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY CIMZIA OR COSENTYX. PLAQUE PSORIASIS (PSO): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING: COSENTYX OR OTEZLA. CROHN'S DISEASE (CD): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY CIMZIA. ULCERATIVE COLITIS (UC): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING: SIMPONI OR ENTYVIO.</p> |

INFLIXIMAB-DYYB

Products Affected

- Inflectra

| PA Criteria | Criteria Details |
|------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | INITIAL: PLAQUE PSORIASIS: SEVERE PLAQUE PSORIASIS INVOLVING GREATER THAN OR EQUAL TO 5 PERCENT BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, OR GENITAL AREA. RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | RHEUMATOID ARTHRITIS, ANKYLOSING SPONDYLITIS: PRESCRIBED BY GIVEN OR IN CONSULTATION WITH A RHEUMATOLOGIST. PSORIATIC ARTHRITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST OR RHEUMATOLOGIST. PSORIASIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST. CROHN'S DISEASE/ULCERATIVE COLITIS: GASTROENTEROLOGIST. |
| Coverage Duration | INITIAL: CD/UC: 8 MOS. RA: 6 MOS. PSA/AS/PSO: 4 MOS. RENEWAL FOR ALL INDICATIONS: 12 MOS. |

| PA Criteria | Criteria Details |
|-----------------------|--|
| Other Criteria | <p>INITIAL: RHEUMATOID ARTHRITIS (RA): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING: ORENCIA, XELJANZ, CIMZIA, OR ACTEMRA. PSORIATIC ARTHRITIS (PSA): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING: CIMZIA, OTEZLA, OR COSENTYX. ANKYLOSING SPONDYLITIS (AS): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING: CIMZIA OR COSENTYX. PLAQUE PSORIASIS (PSO): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING: COSENTYX OR OTEZLA. CROHN'S DISEASE (CD): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY CIMZIA. ULCERATIVE COLITIS (UC): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING: SIMPONI OR ENTYVIO.</p> |

INOTUZUMAB OZOGAMICIN

Products Affected

- Besponsa

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

INTERFERON ALFA-2B

Products Affected

- Intron A injection

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | HEPATITIS C: GASTROENTEROLOGIST, INFECTIOUS DISEASE SPECIALIST, PHYSICIAN SPECIALIZING IN THE TREATMENT OF HEPATITIS (E.G. HEPATOLOGIST). NO REQUIREMENT FOR OTHER FDA APPROVED INDICATIONS. |
| Coverage Duration | 6 MONTHS |
| Other Criteria | LIMITED TO 1 YEAR OF THERAPY EXCEPT 18 MONTHS FOR FOLLICULAR LYMPHOMA. HEPATITIS C GENOTYPE 1, 2, 3, 4, 5, OR 6: REQUIRES A TRIAL OF OR CONTRAINDICATION TO PEGINTERFERON ALFA-2A OR PEGINTERFERON ALFA-2B USED IN COMBINATION WITH RIBAVIRIN UNLESS CONTRAINDICATED. |

INTERFERONS FOR MS-AVONEX, PLEGRIDY, REBIF

Products Affected

- Avonex (with albumin)
- Avonex intramuscular pen injector kit
- Avonex intramuscular syringe kit
- Plegridy
- Rebif (with albumin)
- Rebif Rebidose
- Rebif Titration Pack

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

INTERFERONS FOR MS-BETASERON, EXTAVIA

Products Affected

- Betaseron subcutaneous kit
- Extavia subcutaneous kit

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | TRIAL WITH TWO OF THE FOLLOWING AGENTS FOR MULTIPLE SCLEROSIS: AUBAGIO, AVONEX, GILENYA, PLEGRIDY, REBIF, TECFIDERA, AND GLATIRAMER |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

IPILIMUMAB

Products Affected

- Yervoy

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | INITIAL: UNRESECTABLE/METASTATIC MELANOMA: 3 MO ADJVNT MELANOMA: 6 MO RENEWAL: ADJVNT MELANOMA: 6 MO |
| Other Criteria | RENEWAL FOR ADJUVANT CUTANEOUS MELANOMA: NO EVIDENCE OF DISEASE RECURRENCE (DEFINED AS THE APPEARANCE OF ONE OR MORE NEW MELANOMA LESIONS: LOCAL, REGIONAL OR DISTANT METASTASIS). |

IVABRADINE

Products Affected

- Corlanor

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | PATIENT MUST HAVE NEW YORK HEART ASSOCIATION (NYHA) CLASS II TO IV HEART FAILURE |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A CARDIOLOGIST. |
| Coverage Duration | INITIAL AND RENEWAL: 12 MONTHS. |
| Other Criteria | INITIAL: APPROVAL REQUIRES THE PATIENT DOES NOT HAVE A DEMAND PACEMAKER SET TO A RATE OF 60 BEATS PER MINUTE OR GREATER. PATIENT IS CURRENTLY RECEIVING TREATMENT WITH OR HAS AN INTOLERANCE TO A FORMULARY BETA BLOCKER SUCH AS METOPROLOL SUCCINATE, BISOPROLOL, OR CARVEDILOL. RENEWAL: APPROVAL REQUIRES DIAGNOSIS OF HEART FAILURE AND PATIENT MUST BE IN SINUS RHYTHM. |

IVACAFTOR

Products Affected

- Kalydeco

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | HOMOZYGOUS FOR F508DEL MUTATION IN CFTR GENE. |
| Required Medical Information | CONFIRMED MUTATION IN CFTR GENE ACCEPTABLE FOR THE TREATMENT OF CYSTIC FIBROSIS. |
| Age Restrictions | 6 YEARS OF AGE OR OLDER. |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

IVACAFTOR - GRANULE PACKETS

Products Affected

- Kalydeco

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | F508DEL MUTATION IN CFTR GENE. |
| Required Medical Information | CONFIRMED MUTATION IN CFTR GENE ACCEPTABLE FOR THE TREATMENT OF CYSTIC FIBROSIS. PATIENT WEIGHT. |
| Age Restrictions | 2 YEARS OF AGE TO 5 YEARS OF AGE |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

IXAZOMIB

Products Affected

- Ninlaro

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

IXEKIZUMAB

Products Affected

- Taltz Autoinjector
- Taltz Syringe

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | PLAQUE PSORIASIS (PSO): MODERATE TO SEVERE PLAQUE PSORIASIS INVOLVING GREATER THAN OR EQUAL TO 5PERCENT BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, OR GENITAL AREA. RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | PLAQUE PSORIASIS (PSO): PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST |
| Coverage Duration | PLAQUE PSORIASIS (PSO): INITIAL: 4 MONTHS. RENEWAL: 12 MONTHS |
| Other Criteria | INITIAL: PLAQUE PSORIASIS (PSO): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING PREFERRED AGENTS: COSENTYX OR OTEZLA. |

LEDIPASVIR-SOFOSBUVIR

Products Affected

- Harvoni

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD-IDSA GUIDANCE AND ADDITIONAL CONSIDERATION FOR COVERAGE CONSISTENT WITH FDA LABELING. |
| Exclusion Criteria | |
| Required Medical Information | HCV RNA LEVEL WITHIN PAST 6 MONTHS. |
| Age Restrictions | |
| Prescriber Restrictions | GASTROENTEROLOGIST, INFECTIOUS DISEASE SPECIALIST, PHYSICIAN SPECIALIZING IN THE TREATMENT OF HEPATITIS (HEPATOLOGIST), OR A SPECIALLY TRAINED GROUP SUCH AS ECHO (EXTENSION FOR COMMUNITY HEALTHCARE OUTCOMES) MODEL. |
| Coverage Duration | CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE. |
| Other Criteria | CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE. PATIENT IS NOT CONCURRENTLY TAKING ANY OF THE FOLLOWING: CARBAMAZEPINE, PHENYTOIN, PHENOBARBITAL, OXCARBAZEPINE, RIFAMPIN, RIFABUTIN, RIFAPENTINE, ROSUVASTATIN, SIMEPREVIR, SOFOSBUVIR (AS A SINGLE AGENT), STRIBILD (ELVITEGRAVIR/COBICISTAT/EMTRICITABINE /TENOFVIR), OR TIPRANA VIR/RITONAVIR. |

LENALIDOMIDE

Products Affected

- Revlimid

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

LENVATINIB MESYLATE

Products Affected

- Lenvima

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

LETERMОВIR

Products Affected

- Prevymis intravenous solution 240 mg/12 mL, 480 mg/24 mL
- Prevymis oral

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 4 MONTHS |
| Other Criteria | |

L-GLUTAMINE

Products Affected

- Endari

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR IN CONSULTATION WITH A HEMATOLOGIST |
| Coverage Duration | 12 MONTHS |
| Other Criteria | INITIAL CRITERIA FOR ADULTS (18 YEARS OR OLDER): PHYSICIAN ATTESTATION OF ONE OF THE FOLLOWING: (1) AT LEAST 3 SICKLE CELL CRISES IN THE PAST YEAR OR (2) SICKLE-CELL ASSOCIATED SYMPTOMS WHICH ARE INTERFERING WITH ACTIVITIES OF DAILY LIVING OR (3) HISTORY OF OR HAS RECURRENT ACUTE CHEST SYNDROME (ACS). RENEWAL FOR ALL PATIENTS: PHYSICIAN ATTESTATION PATIENT HAS MAINTAINED OR EXPERIENCED REDUCTION IN ACUTE COMPLICATIONS OF SICKLE CELL DISEASE. |

LIDOCAINE

Products Affected

- lidocaine topical adhesive patch,medicated
- lidocaine topical ointment

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. ADDITIONAL COVERAGE FOR DIABETIC NEUROPATHY WILL BE CONSIDERED FOR REQUESTS FOR LIDOCAINE TOPICAL PATCHES. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | PATCH: 12 MONTHS. OINTMENT: 3 MONTHS. |
| Other Criteria | THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION. |

LIFITEGRAST OPHTHALMIC

Products Affected

- Xiidra

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH AN OPHTHALMOLOGIST, OPTOMETRIST OR RHEUMATOLOGIST. |
| Coverage Duration | 12 MONTHS |
| Other Criteria | PREVIOUS TRIAL OF OR CONTRAINDICATION TO CYCLOSPORINE OPHTHALMIC (RESTASIS). |

LOMITAPIDE

Products Affected

- Juxtapid oral capsule 10 mg, 20 mg, 30 mg, 40 mg, 5 mg, 60 mg

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | LDL CHOLESTEROL LEVEL, LDL RECEPTOR STATUS |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST, ENDOCRINOLOGIST OR LIPIDOLOGIST. |
| Coverage Duration | INITIAL: 7 MONTHS RENEWAL: 6 MONTHS |

| PA Criteria | Criteria Details |
|-----------------------|--|
| Other Criteria | <p>DIAGNOSIS CONFIRMED BY ONE OF THE FOLLOWING: SIMON BROOME DIAGNOSTIC CRITERIA (DEFINITE), (E.G. GENETIC TESTING CONSISTENT WITH HOFH AND PRETREATMENT BASELINE LDL CHOLESTEROL IS GREATER THAN 190 MG/DL), CASCADE SCREENING, DUTCH LIPID NETWORK CRITERIA WITH A SCORE OF AT LEAST 6, OR HISTORY OF UNTREATED CHOLESTEROL GREATER THAN 500MG/DL (OR TREATED CHOLESTEROL GREATER THAN 300MG/DL) AND CUTANEOUS XANTHOMA BEFORE 10 YEARS OF AGE. LOMITAPIDE WILL NOT BE APPROVED FOR PATIENTS CONCURRENTLY USING ANY OF THE FOLLOWING STRONG OR MODERATE CYP3A4 MEDICATIONS: CLARITHROMYCIN, CONIVAPTAN, INDINAVIR, ITRACONAZOLE, KETOCONAZOLE, LOPINAVIR/RITONAVIR, MIBEFRADIL, NEFAZODONE, NELFINAVIR, POSACONAZOLE, RITONAVIR, SAQUINAVIR, TELITHROMYCIN, TIPRANAVIR/RITONAVIR, VORICONAZOLE, AMPRENAVIR, APREPITANT, ATAZANAVIR, CIPROFLOXACIN, CRIZOTINIB, DARUNAVIR/RITONAVIR, DILTIAZEM, ERYTHROMYCIN, FLUCONAZOLE, FOSAMPRENAVIR, IMATINIB, OR VERAPAMIL. INITIAL: LDL CHOLESTEROL LEVEL OF AT LEAST 160MG/DL WHILE ON LIPID-LOWERING THERAPY PRIOR TO INITIATING LOMITAPIDE. PREVIOUS TRIAL OF A PCSK9 INHIBITOR (E.G. ALIROCUMAB OR EVOLOCUMAB), UNLESS THE PATIENT HAS NON-FUNCTIONING LDL RECEPTORS. PREVIOUS TRIAL OF ROSUVASTATIN OR ATORVASTATIN, UNLESS THE PATIENT HAS AN ABSOLUTE CONTRAINDICATION TO STATIN THERAPY (E.G. ACTIVE, DECOMPENSATED LIVER DISEASE, NURSING FEMALE, PREGNANCY OR PLANS TO BECOME PREGNANT, HYPERSENSITIVITY REACTION). STATIN-TOLERANT PATIENTS MUST BE TAKING ATORVASTATIN OR ROSUVASTATIN FOR THE PAST 2 MONTHS PRIOR TO STARTING LOMITAPIDE. LOMITAPIDE MUST BE USED IN COMBINATION WITH ATORVASTATIN OR ROSUVASTATIN. IF THE PATIENT HAS PREVIOUSLY TRIED ATORVASTATIN OR ROSUVASTATIN, LOMITAPIDE MUST BE USED IN COMBINATION WITH ANOTHER STATIN OR FORMULARY LDL-LOWERING AGENT (E.G. BILE ACID SEQUESTRANT, GEMFIBROZIL OR OTHER FIBRATE, EZETIMIBE, OR NIACIN). STATIN-INTOLERANT PATIENTS REQUIRE EITHER PHYSICIAN ATTESTATION OF</p> |

| PA Criteria | Criteria Details |
|-------------|---|
| | <p>STATIN INTOLERANCE OR HISTORY OF SKELETAL-MUSCLE RELATED SYMPTOMS (E.G., MYOPATHY) DUE TO A PREVIOUS TRIAL OF STATINS (E.G. ROSUVASTATIN OR ATORVASTATIN). FOR STATIN-INTOLERANT PATIENTS, LOMITAPIDE MUST BE USED IN COMBINATION WITH ONE OF THE FOLLOWING FORMULARY LIPID-LOWERING TREATMENTS: EZETIMIBE, FENOFIBRATE, NIACIN, OR A BILE ACID SEQUESTRANT (E.G. CHOLESTYRAMINE, COLESTIPOL, COLESEVELAM). RENEWAL: PATIENT HAS RECEIVED AT LEAST 6 MONTHS OF THERAPY WITH LOMITAPIDE IN COMBINATION WITH ANOTHER AND LIPID-LOWERING AGENT.</p> |

LUMACAFITOR-IVACAFITOR

Products Affected

- Orkambi

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | CONFIRMED MUTATION IN CFTR GENE ACCEPTABLE FOR THE TREATMENT OF CYSTIC FIBROSIS. BASELINE FEV1. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A PULMONOLOGIST OR CYSTIC FIBROSIS EXPERT. |
| Coverage Duration | INITIAL: 6 MONTHS. RENEWAL 12 MONTHS. |
| Other Criteria | RENEWAL: MAINTAINED OR IMPROVEMENT IN FEV1 OR REDUCTION IN NUMBER OF PULMONARY EXACERBATIONS OR IMPROVEMENT IN BODY MASS INDEX (BMI). |

MEPOLIZUMAB

Products Affected

- Nucala

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | CONCURRENT USE OF XOLAIR |
| Required Medical Information | BLOOD EOSINOPHIL LEVEL GREATER THAN OR EQUAL TO 150 CELLS/MCL WITHIN THE LAST 6 WEEKS OR GREATER THAN OR EQUAL TO 300 CELLS/MCL WITHIN THE LAST 12 MONTHS |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR IN CONSULTATION WITH A PHYSICIAN SPECIALIZING IN PULMONARY MEDICINE, AN ALLERGIST OR AN IMMUNOLOGIST. |
| Coverage Duration | INITIAL 24 WEEKS. RENEWAL 12 MONTHS |
| Other Criteria | INITIAL THERAPY: PATIENT CURRENTLY TREATED WITH A MAXIMALLY TOLERATED DOSE OF INHALED CORTICOSTEROIDS AND AT LEAST ONE OTHER MAINTENANCE MEDICATION WHICH INCLUDES ANY OF THE FOLLOWING: LONG-ACTING INHALED BETA2-AGONIST, LONG-ACTING MUSCARINIC ANTAGONIST, A LEUKOTRIENE RECEPTOR ANTAGONIST, THEOPHYLLINE, OR ORAL CORTICOSTEROID. RENEWAL REQUIRES DOCUMENTATION THAT THE PATIENT HAS EXPERIENCED IMPROVEMENT IN ASTHMA EXACERBATIONS FROM BASELINE (PHYSICIAN ATTESTATION) AND A REDUCTION IN ORAL CORTICOSTEROID DOSE (IF THE PATIENT WAS ON A MAINTENANCE REGIMEN OF ORAL CORTICOSTEROIDS AT THE INITIATION OF TREATMENT). |

METHYLNALTREXONE

Products Affected

- Relistor subcutaneous solution
- Relistor subcutaneous syringe

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | ADVANCED ILLNESS: OPIOID-INDUCED CONSTIPATION. |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 6 MONTHS FOR PATIENTS RECEIVING PALLIATIVE CARE, 12 MONTHS FOR PATIENTS WITH CHRONIC, NON-CANCER PAIN. |
| Other Criteria | ADVANCED ILLNESS: PATIENT IS RECEIVING PALLIATIVE CARE. CHRONIC NON-CANCER PAIN: PATIENT HAS BEEN TAKING OPIOIDS FOR AT LEAST 4 WEEKS AND HAD A PREVIOUS TRIAL OF OR CONTRAINDICATION TO NALOXEGOL (MOVANTIK). |

METHYLNALTREXONE ORAL

Products Affected

- Relistor oral

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | PATIENT HAS BEEN TAKING OPIOIDS FOR AT LEAST 4 WEEKS AND HAD A PREVIOUS TRIAL OF OR CONTRAINDICATION TO NALOXEGOL (MOVANTIK). |

MIDOSTAURIN

Products Affected

- Rydapt

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | ACUTE MYELOID LEUKEMIA: 6 MONTHS. ADVANCED SYSTEMIC MASTOCYTOSIS: 12 MONTHS |
| Other Criteria | |

MIFEPRISTONE

Products Affected

- Korlym

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

MILTEFOSINE

Products Affected

- Impavido

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

MIPOMERSEN

Products Affected

- Kynamro

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | LDL CHOLESTEROL LEVEL, LDL RECEPTOR STATUS |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A CARDIOLOGIST, ENDOCRINOLOGIST OR LIPIDOLOGIST |
| Coverage Duration | INITIAL: 7 MONTHS RENEWAL 12 MONTHS |

| PA Criteria | Criteria Details |
|-----------------------|---|
| Other Criteria | <p>DIAGNOSIS OF HOMOZYGOUS FAMILIAL HYPERCHOLESTEROLEMIA AS DETERMINED BY ONE OF THE FOLLOWING CRITERIA: SIMON BROOME DIAGNOSTIC CRITERIA (DEFINITE) [EXAMPLE: GENETIC TESTING CONSISTENT WITH HOFH AND PRETREATMENT BASELINE LDL CHOLESTEROL IS GREATER THAN 190 MG/DL], CASCADE SCREENING, DUTCH LIPID NETWORK CRITERIA WITH A SCORE AT LEAST 6, OR HISTORY OF UNTREATED CHOLESTEROL GREATER THAN 500MG/DL (OR TREATED GREATER THAN 300MG/DL) AND CUTANEOUS XANTHOMA BEFORE AGE 10. INITIAL CRITERIA: CURRENT LDL CHOLESTEROL LEVEL IS AT LEAST 160MG/DL. PATIENT DOES NOT HAVE ANY OF THE FOLLOWING CONTRAINDICATIONS TO KYNAMRO (MIPOMERSEN): MODERATE OR SEVERE HEPATIC IMPAIRMENT OR ACTIVE LIVER DISEASE, INCLUDING UNEXPLAINED PERSISTENT ELEVATIONS OF SERUM TRANSAMINASES. PREVIOUS TRIAL OF A PCSK9 INHIBITOR (SUCH AS ALIROCUMAB OR EVOLOCUMAB) UNLESS THE PATIENT HAS NON-FUNCTIONING LDL RECEPTORS. PREVIOUS TRIAL WITH ONE OF THE FOLLOWING STATINS: ROSUVASTATIN OR ATORVASTATIN. PATIENTS WITH ABSOLUTE CONTRAINDICATION TO STATIN THERAPY (ACTIVE, DECOMPENSATED LIVER DISEASE, NURSING FEMALE, PREGNANCY OR PLANS TO BECOME PREGNANT, HYPERSENSITIVITY REACTION) WILL BE APPROVED FOR THERAPY WITHOUT REQUIREMENT OF A TRIAL WITH A STATIN. STATIN-TOLERANT PATIENTS: PRIOR TO (KYNAMRO), PATIENT MUST HAVE BEEN TAKING ONE OF THE FOLLOWING: ATORVASTATIN OR ROSUVASTATIN, FOR AT LEAST 2 MONTHS WITHIN THE PAST 2 MONTHS. FOR STATIN-INTOLERANT PATIENTS: DOCUMENTATION OF STATIN INTOLERANCE WHICH INCLUDES THE FOLLOWING: PHYSICIAN ATTESTATION OR PATIENT HAS TRIED ROSUVASTATIN OR ATORVASTATIN AND HAS EXPERIENCED SKELETAL-MUSCLE RELATED SYMPTOMS (E.G., MYOPATHY). UNLESS CONTRAINDICATED, PATIENT MUST BE ON CONCURRENT THERAPY WITH ONE OF THE FOLLOWING LIPID-LOWERING TREATMENTS (SUCH AS A STATIN [SIMVASTATIN, ATORVASTATIN], EZETIMIBE, FENOFIBRATE, NIACIN, OR BILE ACID SEQUESTRANT [CHOLESTYRAMINE, COLESTIPOL, COLESEVELAM]). RENEWAL CRITERIA:</p> |

| PA Criteria | Criteria Details |
|--------------------|--|
| | PATIENT HAS RECEIVED THERAPY FOR AT LEAST 6 MONTHS AND MUST ALSO BE TAKING KYNAMRO IN COMBINATION WITH ANOTHER LIPID-LOWERING AGENT. |

NAPROXEN- ESOMEPRAZOLE

Products Affected

- Vimovo

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | TRIAL OF ONE OF THE FOLLOWING GENERIC, FEDERAL LEGEND PROTON PUMP INHIBITORS: OMEPRAZOLE, LANSOPRAZOLE, OR PANTOPRAZOLE AND A TRIAL OF GENERIC, FEDERAL LEGEND NAPROXEN. |

NATALIZUMAB

Products Affected

- Tysabri

| PA Criteria | Criteria Details |
|------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | CROHN'S DISEASE: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A GASTROENTEROLOGIST. |
| Coverage Duration | MULTIPLE SCLEROSIS: 12 MONTHS. CROHN'S DISEASE: INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS |
| Other Criteria | MULTIPLE SCLEROSIS INITIAL CRITERIA: PREVIOUS TRIAL OF TWO OF THE FOLLOWING PREFERRED AGENTS FOR MULTIPLE SCLEROSIS: GLATIRAMER, REBIF, AVONEX, PLEGRIDY, TECFIDERA, GILENYA, OR AUBAGIO. CROHN'S DISEASE INITIAL CRITERIA: PREVIOUS TRIAL OF HUMIRA AND CIMZIA. CROHN'S DISEASE RENEWAL CRITERIA: PATIENT HAS RECEIVED AT LEAST 12 MONTHS OF THERAPY WITH TYSABRI WITH PHYSICIAN ATTESTATION THAT THE PATIENT HAS NOT REQUIRED MORE THAN 3 MONTHS OF CORTICOSTEROID USE WITHIN THE PAST 12 MONTHS TO CONTROL THEIR CROHN'S DISEASE WHILE ON TYSABRI, OR PATIENT HAS ONLY RECEIVED 6 MONTHS OF THERAPY WITH TYSABRI WITH PHYSICIAN ATTESTATION THAT THE PATIENT HAS TAPERED OFF CORTICOSTEROIDS DURING THE FIRST 24 WEEKS OF TYSABRI THERAPY. |

NECITUMUMAB

Products Affected

- Portrazza

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

NERATINIB MALEATE

Products Affected

- Nerlynx

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | EARLY-STAGE TUMOR (STAGE I-III) AND TUMOR IS HORMONE-RECEPTOR POSITIVE AND THE MEDICATION IS BEING REQUESTED WITHIN 2 YEARS OF COMPLETING THE LAST TRASTUZUMAB DOSE |

NILOTINIB

Products Affected

- Tasigna

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | PREVIOUSLY TREATED CML REQUIRES BCR-ABL MUTATIONAL ANALYSIS NEGATIVE FOR THE FOLLOWING MUTATIONS: T315I, Y253H, E255K/V, AND F359V/C/I. |

NINTEDANIB

Products Affected

- Ofev

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | NOT APPROVED FOR PATIENTS WITH 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, AND CANCER. NOT APPROVED IF PATIENT DOES NOT HAVE A PREDICTED FORCED VITAL CAPACITY (FVC) OF AT LEAST 50 PERCENT OR HAS NOT OBTAINED LIVER FUNCTION TESTS |
| Required Medical Information | 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. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A PULMONOLOGIST |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

NIRAPARIB TOSYLATE

Products Affected

- Zejula

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

NITISINONE

Products Affected

- Orfadin

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | DIAGNOSIS OF HEREDITARY TYROSINEMIA TYPE 1 AS CONFIRMED BY ELEVATED URINARY OR PLASMA SUCCINYLACETONE LEVELS OR A MUTATION IN THE FUMARYLACETOACETATE HYDROLASE GENE. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A PRESCRIBER SPECIALIZING IN INHERITED METABOLIC DISEASES. |
| Coverage Duration | INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS |
| Other Criteria | ORFADIN SUSPENSION: TRIAL OF ORFADIN CAPSULES. RENEWAL: THE PATIENT'S URINARY OR PLASMA SUCCINYLACETONE LEVELS HAVE DECREASED FROM BASELINE WHILE ON TREATMENT WITH NITISINONE. |

NIVOLUMAB

Products Affected

- Opdivo intravenous solution 100 mg/10 mL, 40 mg/4 mL

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | MELANOMA: OPDIVO IS NOT APPROVED FOR COMBINATION THERAPY WITH TAFINLAR, MEKINIST (TRAMETINIB), COTELLIC (COBIMETINIB), OR ZELBORAF. |

OBETICHOLIC ACID

Products Affected

- Ocaliva

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | PATIENTS WITH COMPLETE BILIARY OBSTRUCTION. |
| Required Medical Information | DIAGNOSIS OF PRIMARY BILIARY CHOLANGITIS AS CONFIRMED BY AT LEAST TWO OF THE FOLLOWING CRITERIA: AN ALKALINE PHOSPHATASE LEVEL OF AT LEAST 1.5 TIMES THE UPPER LIMIT OF NORMAL (ULN), THE PRESENCE OF ANTIMITOCHONDRIAL ANTIBODIES AT A TITER OF 1:40 OR HIGHER, HISTOLOGIC EVIDENCE OF NON-SUPPURATIVE DESTRUCTIVE CHOLANGITIS AND DESTRUCTION OF INTERLOBULAR BILE DUCTS. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A GASTROENTEROLOGIST OR HEPATOLOGIST. |
| Coverage Duration | 12 MONTHS |
| Other Criteria | INITIAL: USED IN COMBINATION WITH URSODEOXYCHOLIC ACID (E.G., URSODIOL, URSO 250, URSO FORTE) IN ADULTS WITH AN INADEQUATE RESPONSE TO URSODEOXYCHOLIC ACID AT A DOSAGE OF 13-15 MG/KG/DAY FOR AT LEAST 1 YEAR, OR AS MONOTHERAPY IN ADULTS UNABLE TO TOLERATE URSODEOXYCHOLIC ACID. RENEWAL: PATIENT'S ALKALINE PHOSPHATASE LEVELS HAVE DECREASED BY AT LEAST 15% FROM BASELINE WHILE ON TREATMENT WITH OBETICHOLIC ACID. |

OBINUTUZUMAB

Products Affected

- Gazyva

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 6 MONTHS |
| Other Criteria | |

OCRELIZUMAB

Products Affected

- Ocrevus

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | RELAPSING FORM OF MULTIPLE SCLEROSIS: TRIAL OF TWO OF THE FOLLOWING AGENTS FOR MULTIPLE SCLEROSIS: AUBAGIO, AVONEX, GILENYA, PLEGRIDY, REBIF, TECFIDERA, OR GLATIRAMER. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D. |

OLAPARIB

Products Affected

- Lynparza oral capsule
- Lynparza oral tablet

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

OLARATUMAB

Products Affected

- Lartruvo

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | THIS DRUG MAY BE COVERED UNDER MEDICARE PART B OR D DEPENDING UPON THE CIRCUMSTANCES. INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE DETERMINATION. |

OMACETAXINE

Products Affected

- Synribo

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | INDUCTION: 3 MONTHS. POST INDUCTION OR RENEWAL: 3 TO 12 MONTHS |
| Other Criteria | CML INDUCTION THERAPY: TRIAL OF OR CONTRAINDICATION TO AT LEAST TWO OF THE FOLLOWING AGENTS: GLEEVEC, SPRYCEL, TASIGNA, BOSULIF OR ICLUSIG. APPROVAL FOR POST-INDUCTION THERAPY DURATION WILL DEPEND ON THE PATIENT'S HEMATOLOGIC RESPONSE, DEFINED AS AN ABSOLUTE NEUTROPHIL COUNT (ANC) GREATER THAN OR EQUAL TO $1.5 \times 10^9/L$, PLATELETS GREATER THAN OR EQUAL TO $100 \times 10^9/L$ WITHOUT BLOOD BLASTS OR THE PATIENT HAS BONE MARROW BLASTS AT LESS THAN 5 PERCENT. APPROVAL IS FOR 12 MONTHS IF HEMATOLOGIC RESPONSE IS MET. IF NOT MET, APPROVAL IS FOR 3 MONTHS. |

OMALIZUMAB

Products Affected

- Xolair

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | INITIAL CRITERIA FOR ASTHMA: PATIENT MEETS THE CRITERIA OF MODERATE TO SEVERE ASTHMA, POSITIVE SKIN PRICK OR RAST TEST, FEV1 LESS THAN 80%, DEMONSTRATED INADEQUATELY CONTROLLED SYMPTOMS ON INHALED CORTICOSTEROIDS AND SECOND ASTHMA CONTROLLER, BASELINE IGE SERUM LEVEL GREATER THAN OR EQUAL TO 30IU/ML. RENEWAL CRITERIA FOR ASTHMA: PHYSICIAN ATTESTATION OF IMPROVEMENT IN ASTHMA EXCERBATIONS FROM BASELINE AND A REDUCTION IN ORAL OR INHALED CORTICOSTEROID USE. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A SPECIALIST IN ALLERGY, PULMONARY MEDICINE, DERMATOLOGY OR IMMUNOLOGY. |
| Coverage Duration | ASTHMA: 12 MONTHS. CHRONIC IDIOPATHIC URTICARIA: 6 MONTHS. |
| Other Criteria | FOR CHRONIC IDIOPATHIC URTICARIA: PREVIOUS TRIAL OF OR CONTRAINDICATION TO A MAXIMALLY TOLERATED DOSE OF AN H1 ANTI-HISTAMINE (SUCH AS CLARINEX OR XYZAL) AND PATIENT STILL EXPERIENCES HIVES ON MOST DAYS OF THE WEEK. |

OMBITASVIR-PARITAPREVIR-RITONAVIR

Products Affected

- Technivie

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD-IDSA GUIDANCE AND ADDITIONAL CONSIDERATION FOR COVERAGE CONSISTENT WITH FDA LABELING. |
| Exclusion Criteria | DECOMPENSATED CIRRHOSIS, MODERATE OR SEVERE LIVER IMPAIRMENT (CHILD-PUGH B OR C). |
| Required Medical Information | HCV RNA LEVEL WITHIN PAST 6 MONTHS. |
| Age Restrictions | |
| Prescriber Restrictions | GASTROENTEROLOGIST, INFECTIOUS DISEASE SPECIALIST, PHYSICIAN SPECIALIZING IN THE TREATMENT OF HEPATITIS (HEPATOLOGIST), OR A SPECIALLY TRAINED GROUP SUCH AS ECHO (EXTENSION FOR COMMUNITY HEALTHCARE OUTCOMES) MODEL. |
| Coverage Duration | CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE. |

| PA Criteria | Criteria Details |
|-----------------------|---|
| Other Criteria | <p>CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE. 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. MUST BE USED CONCURRENTLY WITH RIBAVIRIN UNLESS PATIENT IS TREATMENT NAIVE AND HAS CONTRAINDICATION TO RIBAVIRIN. PATIENT IS NOT CONCURRENTLY TAKING ANY OF THE FOLLOWING (CONTRAINDICATED OR NOT RECOMMENDED BY THE MANUFACTURER): ALFUZOSIN, CARBAMAZEPINE, PHENYTOIN, PHENOBARBITAL, RIFAMPIN, ERGOTAMINE, DIHYDROERGOTAMINE, ERGONOVINE, METHYLERGONOVINE, ETHINYL ESTRADIOL CONTAINING MEDICATIONS (SUCH AS COMBINED ORAL CONTRACEPTIVES, NUVARING, ORTHO EVRA OR XULANE TRANSDERMAL PATCH SYSTEM), LOVASTATIN, SIMVASTATIN, PIMOZIDE, EFAVIRENZ (ATRIPLA, SUSTIVA), REVATIO (SILDENAFIL DOSE OF 20MG AND/OR DOSED THREE TIMES DAILY FOR PAH), TRIAZOLAM, ORAL MIDAZOLAM, LOPINAVIR/RITONAVIR, RILPIVIRINE, SALMETEROL.</p> |

OMBITASVIR-PARITAPREVIR-RITONAVIR-DASABUVIR

Products Affected

- Viekira Pak
- Viekira XR

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD-IDSA GUIDANCE AND ADDITIONAL CONSIDERATION FOR COVERAGE CONSISTENT WITH FDA LABELING. |
| Exclusion Criteria | DECOMPENSATED CIRRHOSIS, MODERATE OR SEVERE LIVER IMPAIRMENT (CHILD-PUGH B OR C). |
| Required Medical Information | HCV RNA LEVEL WITHIN PAST 6 MONTHS. |
| Age Restrictions | |
| Prescriber Restrictions | GASTROENTEROLOGIST, INFECTIOUS DISEASE SPECIALIST, PHYSICIAN SPECIALIZING IN THE TREATMENT OF HEPATITIS (HEPATOLOGIST), OR A SPECIALLY TRAINED GROUP SUCH AS ECHO (EXTENSION FOR COMMUNITY HEALTHCARE OUTCOMES) MODEL. |
| Coverage Duration | CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE. |

| PA Criteria | Criteria Details |
|-----------------------|---|
| Other Criteria | <p>CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE. 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. PATIENT IS NOT CONCURRENTLY TAKING ANY OF THE FOLLOWING: ALFUZOSIN, CARBAMAZEPINE, PHENYTOIN, PHENOBARBITAL, GEMFIBROZIL, RIFAMPIN, ERGOTAMINE, DIHYDROERGOTAMINE, ERGONOVINE, METHYLERGONOVINE, ETHINYL ESTRADIOL CONTAINING MEDICATIONS (SUCH AS COMBINED ORAL CONTRACEPTIVES, NUVARING, ORTHO EVRA OR XULANE TRANSDERMAL PATCH SYSTEM), ST. JOHN'S WORT, LOVASTATIN, SIMVASTATIN, PIMOZIDE, EFAVIRENZ, REVATIO, TRIAZOLAM, ORAL MIDAZOLAM, DARUNAVIR/RITONAVIR, LOPINAVIR/RITONAVIR, RILPIVIRINE, SALMETEROL.</p> |

OSIMERTINIB

Products Affected

- Tagrisso

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | CONCURRENT THERAPY WITH AN EGFR TYROSINE KINASE-INHIBITOR |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

OXYMETHOLONE

Products Affected

- Anadrol-50

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | CARCINOMA OF THE PROSTATE OR BREAST IN MALE PATIENTS, CARCINOMA OF THE BREAST IN FEMALES WITH HYPERCALCEMIA, WOMEN WHO ARE OR MAY BECOME PREGNANT, NEPHROSIS OR THE NEPHROTIC PHASE OF NEPHRITIS, HYPERSENSITIVITY TO THE DRUG AND SEVERE HEPATIC DYSFUNCTION. |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

PALBOCICLIB

Products Affected

- Ibrance

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | PREVIOUS TRIAL OF OR CONTRAINDICATION TO THE PREFERRED FORMULARY ALTERNATIVE RIBOCICLIB (KISQALI). |

PALIVIZUMAB

Products Affected

- Synagis

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | GESTATIONAL AGE |
| Age Restrictions | LESS THAN 24 MONTHS OF AGE. |
| Prescriber Restrictions | |
| Coverage Duration | 1 MONTH TO 5 MONTHS. SEE OTHER CRITERIA FOR MORE INFORMATION. |
| Other Criteria | CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT RECOMMENDATIONS FROM THE AMERICAN ACADEMY OF PEDIATRICS FOR PALIVIZUMAB PROPHYLAXIS FOR RESPIRATORY SYNCYTIAL VIRUS INFECTIONS. INITIAL: APPROVAL WILL BE FOR AT LEAST 1 MONTH AND NO GREATER THAN 5 MONTHS DEPENDENT UPON REMAINING LENGTH OF RESPIRATORY SYNCYTIAL VIRUS (RSV) SEASON. RENEWAL: ADDITIONAL 1 MONTH OF TREATMENT FOR CARDIOPULMONARY BYPASS SURGERY DURING RSV PROPHYLAXIS SEASON. |

PANITUMUMAB

Products Affected

- Vectibix

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

PANOBINOSTAT

Products Affected

- Farydak

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | RENEWAL: PATIENT HAS TOLERATED THE FIRST 8 CYCLES OF THERAPY WITHOUT UNRESOLVED SEVERE OR MEDICALLY SIGNIFICANT TOXICITY. |

PARATHYROID HORMONE

Products Affected

- Natpara

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

PAZOPANIB

Products Affected

- Votrient

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

PDE5 INHIBITORS FOR PULMONARY ARTERIAL HYPERTENSION

Products Affected

- Adcirca
- sildenafil (antihypertensive) oral

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | PATIENT CANNOT CONCURRENTLY OR INTERMITTENTLY BE TAKING ORAL ERECTILE DYSFUNCTION AGENTS (E.G. CIALIS, VIAGRA), ANY ORGANIC NITRATES IN ANY FORM, OR GUANYLATE CYCLASE (GC) STIMULATORS (ADEMPAS). |
| Required Medical Information | DOCUMENTED CONFIRMATORY PULMONARY ARTERIAL HYPERTENSION (PAH) DIAGNOSIS BASED ON RIGHT HEART CATHETERIZATION. PATIENT HAS NYHA-WHO FUNCTIONAL CLASS II-IV SYMPTOMS |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST |
| Coverage Duration | INITIAL AND RENEWAL: 12 MONTHS |
| Other Criteria | INITIAL: MEAN PULMONARY ARTERY PRESSURE (PAP) OF AT LEAST 25 MMHG OR GREATER, PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 3 WOOD UNITS. REQUEST FOR ADCIRCA REQUIRE TRIAL OR CONTRAINDICATION TO REVATIO. RENEWAL: PATIENT SHOWS IMPROVEMENT FROM BASELINE IN THE 6-MINUTE WALK DISTANCE OR PATIENT HAS A STABLE 6-MINUTE WALK DISTANCE WITH A STABLE/ IMPROVED WHO FUNCTIONAL CLASS. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION. |

PDE5 INHIBITORS FOR PULMONARY ARTERIAL HYPERTENSION - IV

Products Affected

- sildenafil (antihypertensive) intravenous

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | PATIENT CANNOT CONCURRENTLY OR INTERMITTENTLY BE TAKING ORAL ERECTILE DYSFUNCTION AGENTS (E.G. CIALIS, VIAGRA), ANY ORGANIC NITRATES IN ANY FORM, OR GUANYLATE CYCLASE (GC) STIMULATORS (ADEMPAS). |
| Required Medical Information | DOCUMENTED CONFIRMATORY PULMONARY ARTERIAL HYPERTENSION (PAH) DIAGNOSIS BASED ON RIGHT HEART CATHETERIZATION. PATIENT HAS NYHA-WHO FUNCTIONAL CLASS II-IV SYMPTOMS |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST |
| Coverage Duration | INITIAL AND RENEWAL: 12 MONTHS |
| Other Criteria | INITIAL: MEAN PULMONARY ARTERY PRESSURE (PAP) OF AT LEAST 25 MMHG OR GREATER, PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 3 WOOD UNITS. RENEWAL: PATIENT SHOWS IMPROVEMENT FROM BASELINE IN THE 6-MINUTE WALK DISTANCE OR PATIENT HAS A STABLE 6-MINUTE WALK DISTANCE WITH A STABLE/ IMPROVED WHO FUNCTIONAL CLASS. |

PEG-INTERFERON ALFA-2B-SYLATRON

Products Affected

- Sylatron

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | OVERALL DURATION OF THERAPY LIMITED TO 5 YEARS. |

PEMBROLIZUMAB

Products Affected

- Keytruda intravenous recon soln
- Keytruda intravenous solution

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

PENICILLAMINE

Products Affected

- Cuprimine

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | RHEUMATOID ARTHRITIS: HISTORY OR OTHER EVIDENCE OF RENAL INSUFFICIENCY |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | WILSON'S DISEASE: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A HEPATOLOGIST. CYSTINURIA: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A NEPHROLOGIST. RHEUMATOID ARTHRITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A RHEUMATOLOGIST. |
| Coverage Duration | 12 MONTHS |
| Other Criteria | WILSON'S DISEASE: GENETIC TESTING FOR ATP7B MUTATIONS. CYSTINURIA: DIAGNOSIS REQUIRES THE PRESENCE OF NEPHROLITHIASIS AND 1 OR MORE OF THE FOLLOWING: STONE ANALYSIS SHOWING PRESENCE OF CYSTEINE, IDENTIFICATION OF PATHOGNOMONIC HEXAGONAL CYSTINE CRYSTALS ON URINALYSIS, POSITIVE FAMILY HISTORY OF CYSTINURIA WITH POSITIVE CYANIDE-NITROPRUSSIDE SCREEN. REQUESTS FOR CUPRIMINE FOR THE TREATMENT OF WILSONS DISEASE, CYSTINURIA, AND RHEUMATOID ARTHRITIS REQUIRE A PREVIOUS TRIAL OF OR CONTRAINDICATION TO DEPEN. |

PENICILLAMINE-DEPEN

Products Affected

- Depen Titratabs

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | RHEUMATOID ARTHRITIS: HISTORY OR OTHER EVIDENCE OF RENAL INSUFFICIENCY |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | WILSON'S DISEASE: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A HEPATOLOGIST. CYSTINURIA: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A NEPHROLOGIST. RHEUMATOID ARTHRITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A RHEUMATOLOGIST. |
| Coverage Duration | 12 MONTHS |
| Other Criteria | WILSON'S DISEASE: GENETIC TESTING FOR ATP7B MUTATIONS. CYSTINURIA: DIAGNOSIS REQUIRES THE PRESENCE OF NEPHROLITHIASIS AND 1 OR MORE OF THE FOLLOWING: STONE ANALYSIS SHOWING PRESENCE OF CYSTEINE, IDENTIFICATION OF PATHOGNOMONIC HEXAGONAL CYSTINE CRYSTALS ON URINALYSIS, POSITIVE FAMILY HISTORY OF CYSTINURIA WITH POSITIVE CYANIDE-NITROPRUSSIDE SCREEN. |

PERTUZUMAB

Products Affected

- Perjeta

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | INITIAL: 4 MONTHS. RENEWAL: 12 MONTHS |
| Other Criteria | |

PIMAVANSERIN

Products Affected

- Nuplazid

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | 18 YEARS OR OLDER |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A NEUROLOGIST, GERIATRICIAN, OR A BEHAVIORAL HEALTH SPECIALIST (SUCH AS A PSYCHIATRIST). |
| Coverage Duration | INITIAL 12 MONTHS. RENEWAL 12 MONTHS. |
| Other Criteria | RENEWAL REQUIRES THAT THE PATIENT HAS EXPERIENCED AN IMPROVEMENT IN PSYCHOSIS SYMPTOMS FROM BASELINE AND DEMONSTRATES A CONTINUED NEED FOR TREATMENT. |

PIRFENIDONE

Products Affected

- Esbriet oral capsule
- Esbriet oral tablet 267 mg, 801 mg

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | PATIENTS WITH 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, AND CANCER). NOT APPROVED IF THE PATIENT HAS NOT OBTAINED LIVER FUNCTION TESTS. |
| Required Medical Information | PATIENT WITH USUAL INTERSTITIAL PNEUMONIA (UIP) PATTERN AS EVIDENCED BY HIGH-RESOLUTION COMPUTED TOMOGRAPHY (HRCT) ALONE OR VIA A COMBINATION OF SURGICAL LUNG BIOPSY AND HRCT |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A PULMONOLOGIST |
| Coverage Duration | 12 MONTHS |
| Other Criteria | PATIENT HAS A PREDICTED FORCED VITAL CAPACITY (FVC) OF AT LEAST 50%. |

POMALIDOMIDE

Products Affected

- Pomalyst

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

PONATINIB

Products Affected

- Iclusig oral tablet 15 mg, 45 mg

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

PRAMLINTIDE

Products Affected

- SymlinPen 120
- SymlinPen 60

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | TYPE I OR TYPE II DIABETES: REQUIRING INSULIN OR CONTINUOUS INSULIN INFUSION (INSULIN PUMP) FOR GLYCEMIC CONTROL |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

PYRIMETHAMINE

Products Affected

- Daraprim

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D, ADDITIONAL CONSIDERATION FOR CHRONIC MAINTENANCE THERAPY FOR TOXOPLASMOSIS AND TOXOPLASMOSIS PROPHYLAXIS. |
| Exclusion Criteria | |
| Required Medical Information | MALARIA: PLASMODIA SUSCEPTIBLE TESTING. TOXOPLASMOSIS:CD4 LEVEL |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | ACUTE MALARIA AND CHEMOPROPHYLAXIS: INITIAL: 3 MONTHS. RENEWAL: 12 MONTHS. SEE OTHER CRITERIA FIELD |

| PA Criteria | Criteria Details |
|-----------------------|---|
| Other Criteria | <p>INITIAL: ACUTE MALARIA TREATMENT AND MALARIA CHEMOPROPHYLAXIS REQUIRES THAT THE PATIENT HAS MALARIA SUSCEPTIBLE TO PYRIMETHAMINE AND A PREVIOUS TRIAL OF PLAQUENIL (HYDROXYCHLOROQUINE SULFATE) AND MALARONE (ATOVAQUONE/PROGUANIL) (UNLESS THESE REGIMENS ARE RESISTANT IN THE SPECIFIC REGION AS INDICATED BY REGIONAL PLASMODIA SUSCEPTIBILITY). PRIMARY PROPHYLAXIS OF TOXOPLASMOSIS IN PATIENTS WITH HIV REQUIRES PREVIOUS TRIAL OF OR CONTRAINDICATION TO BACTRIM (SMX/TMP). RENEWAL: CONTINUATION OF TREATMENT FOLLOWING ACUTE MALARIA REQUIRES PREVIOUS INFECTION WITH MALARIA SUSCEPTIBLE TO PYRIMETHAMINE WITH SUBSEQUENT CLINICAL CURE (ELIMINATION OF MALARIA SYMPTOMS DEFINED AS CHILLS, FEVER, SWEATS, GENERAL MALAISE) FOLLOWED BY SYMPTOMS OF RELAPSE. CONTINUATION OF MALARIA CHEMOPROPHYLAXIS REQUIRES THE PATIENT WILL BE TRAVELING TO OR RESIDING IN AN AREA WHERE PLASMODIA SUSCEPTIBLE TO PYRIMETHAMINE EXISTS (MALARIA MUST BE SENSITIVE TO PYRIMETHAMINE).CONTINUED TREATMENT OF TOXOPLASMOSIS REQUIRES 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 CURRENT ANTI-RETROVIRAL THERAPY IF HIV POSITIVE. CONTINUATION OF PRIMARY PROPHYLAXIS FOR TOXOPLASMOSIS WITH HIV REQUIRES CD4 COUNT LESS THAN 200 CELLS/MM3 AND CURRENT ANTI RETROVIRAL THERAPY. TOXOPLASMOSIS: INITIAL: 8 WEEKS. RENEWAL: 6 MONTHS. PRIMARY PROPHYLAXIS OF TOXOPLASMOSIS: INITIAL AND RENEWAL IS 12 MONTHS.</p> |

QUININE SULFATE

Products Affected

- quinine sulfate

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

RAMUCIRUMAB

Products Affected

- Cyramza

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

REGORAFENIB

Products Affected

- Stivarga

| PA Criteria | Criteria Details |
|------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | FOR COLORECTAL CANCER: TRIAL OF OR CONTRAINDICATION TO AN ANTI-VEGF THERAPY SUCH AS AVASTIN OR ZALTRAP AND A FLUOROPYRIMIDINE-, OXALIPLATIN- AND IRINOTECAN-BASED CHEMOTHERAPY SUCH AS FOLFOX, FOLFOXIRI, FOLFIRI, CAPEOX, INFUSIONAL 5-FU/LV OR CAPECITABINE. IF APPLICABLE, A TRIAL OF OR CONTRAINDICATION TO AN ANTI-EGFR THERAPY SUCH AS ERBITUX OR VECTIBIX IS ALSO REQUIRED FOR KRAS WILD TYPE COLORECTAL CANCER. FOR GIST, A TRIAL OF OR CONTRAINDICATION TO GLEEVEC AND SUTENT IS REQUIRED. |

RESLIZUMAB

Products Affected

- Cinqair

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | CONCURRENT USE OF XOLAIR |
| Required Medical Information | BLOOD EOSINOPHIL LEVEL GREATER THAN OR EQUAL TO 400 CELLS/MCL WITHIN THE LAST 6 MONTHS |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A PHYSICIAN SPECIALIZING IN PULMONARY MEDICINE |
| Coverage Duration | INITIAL 24 WEEKS. RENEWAL 12 MONTHS |
| Other Criteria | INITIAL THERAPY: PATIENT CURRENTLY TREATED WITH A MAXIMALLY TOLERATED DOSE OF INHALED CORTICOSTEROIDS. RENEWAL REQUIRES DOCUMENTATION THAT THE PATIENT HAS EXPERIENCED AT LEAST A 25 PERCENT REDUCTION IN ASTHMA EXACERBATIONS (FOR EXAMPLE: HOSPITALIZATIONS, URGENT OR EMERGENT CARE VISITS, USE OF RESCUE MEDICATIONS, ETC.) FROM BASELINE. |

RIBOCICLIB

Products Affected

- Kisqali mg/day(200 mg x 2)-2.5 mg, 600
- Kisqali Femara Co-Pack oral tablet 200 mg/day(200 mg x 3)-2.5 mg
mg/day(200 mg x 1)-2.5 mg, 400

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

RIFAXIMIN

Products Affected

- Xifaxan oral tablet 200 mg, 550 mg

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | TRAVELERS' DIARRHEA:1 FILL IN 1MONTH.HEPATIC ENCEPHALOPATHY:12 MO.IBS-D:INITIAL:12 WKS.RENEWAL:12 MO |
| Other Criteria | FOR RIFAXIMIN 550 MG TABLETS ONLY: INITIAL: HEPATIC ENCEPHALOPATHY (HE): PREVIOUS TRIAL OF OR CONTRAINDICATION TO LACTULOSE OR CONCURRENT LACTULOSE THERAPY. IRRITABLE BOWEL SYNDROME WITH DIARRHEA (IBS-D): PREVIOUS TRIAL OF OR CONTRAINDICATION TO DICYCLOMINE. RENEWAL FOR IBS-D REQUIRES THAT AT LEAST 10 WEEKS HAVE PASSED SINCE THE LAST TREATMENT COURSE OF RIFAXIMIN AND PHYSICIAN ATTESTATION OF IMPROVEMENT. |

RIOCIGUAT

Products Affected

- Adempas

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | INITIAL FOR PAH: PATIENT IS NOT CONCURRENTLY TAKING NITRATES OR NITRIC OXIDE DONORS (E.G. AMYL NITRATE), PHOSPHODIESTERASE INHIBITORS (E.G. SILDENAFIL, TADALAFIL, OR VARDENAFIL), OR NON-SPECIFIC PDE INHIBITORS (E.G. DIPYRIDAMOLE, THEOPHYLLINE). INITIAL FOR CTEPH: PATIENT IS NOT A CANDIDATE FOR SURGERY OR HAS INOPERABLE CTEPH. PERSISTENT OR RECURRENT DISEASE AFTER SURGICAL TREATMENT. PATIENT IS NOT CONCURRENTLY OR INTERMITTENTLY TAKING NITRATES, NITRIC OXIDE DONORS OR ANY PDE INHIBITORS (E.G. VIAGRA, CIALIS, DIPYRIDAMOLE). |
| Required Medical Information | DOCUMENTED CONFIRMATORY PULMONARY ARTERIAL HYPERTENSION (PAH) DIAGNOSIS BASED ON RIGHT HEART CATHETERIZATION. PATIENT HAS NYHA-WHO FUNCTIONAL CLASS II-IV SYMPTOMS. DIAGNOSIS OF PERSISTENT/RECURRENT CHRONIC THROMBOEMBOLIC PULMONARY HYPERTENSION (CTEPH) WHO GROUP 4. PATIENT HAS NYHA-WHO FUNCTIONAL CLASS II-IV SYMPTOMS. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST. |
| Coverage Duration | INITIAL AND RENEWAL: 12 MONTHS |

| PA Criteria | Criteria Details |
|-----------------------|---|
| Other Criteria | <p>INITIAL FOR PAH: MEAN PULMONARY ARTERY PRESSURE (PAP) OF AT LEAST 25 MMHG OR GREATER, PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 3 WOOD UNITS. PREVIOUS TRIAL OF OR CONTRAINDICATION TO A PHOSPHODIESTERASE-5 (PDE-5) INHIBITOR, SUCH AS REVATIO OR ADCIRCA.</p> <p>RENEWAL FOR PAH AND CTEPH: PATIENT SHOW IMPROVEMENT FROM BASELINE IN THE 6-MINUTE WALK DISTANCE OR PATIENT HAS A STABLE 6-MINUTE WALK DISTANCE WITH A STABLE/ IMPROVED WHO FUNCTIONAL CLASS.</p> |

RITUXIMAB

Products Affected

- Rituxan

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | RHEUMATOID ARTHRITIS: PRESCRIBED BY OR IN CONSULTATION WITH A RHEUMATOLOGIST. NHL, CLL: ONCOLOGIST. |
| Coverage Duration | RA: INITIAL: 6 MO. RENEWAL: 12 MONTHS. NHL: 1 YEAR. CLL: 6 MO. WG, MPA: 3 MONTH. |
| Other Criteria | INITIAL: RHEUMATOID ARTHRITIS: PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING PREFERRED AGENTS: ORENCIA, XELJANZ, CIMZIA OR ACTEMRA. |

RITUXIMAB SQ

Products Affected

- Rituxan Hycela

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | THE PATIENT HAS RECEIVED OR WILL RECEIVE AT LEAST ONE FULL DOSE OF A RITUXIMAB PRODUCT BY INTRAVENOUS INFUSION PRIOR TO INITIATION OF RITUXIMAB AND HYALURONIDASE. |

ROMIPLOSTIM

Products Affected

- Nplate

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | INITIAL: ADEQUATE RESPONSE TO CORTICOSTEROIDS, IMMUNOGLOBULINS, OR SPLENECTOMY. RENEWAL: NO CLINICAL RESPONSE AS DEFINED BY AN INCREASE IN PLATELET COUNT OF GREATER THAN OR EQUAL TO $50 \times 10^9/L$ AT THE MAX DOSE OF 10 MCG/KG PER DAY FOR 4 WEEKS. |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | INITL: 2 MO., RENEW: IF NO RESPONSE AFTER INITIAL APPROVAL: 1 MO. AT MAX DOSE. IF RESPONSE: 12 MO. |
| Other Criteria | |

RUCAPARIB

Products Affected

- Rubraca

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

RUXOLITINIB

Products Affected

- Jakafi

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | RENEWAL: IMPROVEMENT OR MAINTENANCE OF SYMPTOM IMPROVEMENT SUCH AS A 50% OR GREATER REDUCTION IN TOTAL SYMPTOM SCORE ON THE MODIFIED MYELOFIBROSIS SYMPTOM ASSESSMENT FORM (MFSAF) V2.0 OR 50% OR GREATER REDUCTION IN PALPABLE SPLEEN LENGTH, OR REDUCTION OF 35% OR GREATER FROM BASELINE SPLEEN VOLUME AFTER 6 MONTHS OF THERAPY. |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS |
| Other Criteria | |

SAFINAMIDE MESYLATE

Products Affected

- Xadago

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

SARILUMAB

Products Affected

- Kevzara

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A RHEUMATOLOGIST. |
| Coverage Duration | INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS |
| Other Criteria | INITIAL: PREVIOUS TRIAL OF HUMIRA AND ONE OF THE FOLLOWING PREFERRED AGENTS: ACTEMRA, CIMZIA, ORENCIA, OR XELJANZ. |

SEBELIPASE ALFA

Products Affected

- Kanuma

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | BLOOD TEST OR DRIED BLOOD SPOT TEST INDICATING LOW OR ABSENT LYSOSOMAL ACID LIPASE DEFICIENCY (LAL) ENZYME ACTIVITY, OR A GENETIC TEST INDICATING THE PRESENCE OF ALTERED LIPA GENE(S). |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH AN ENDOCRINOLOGIST, HEPATOLOGIST, GASTROENTEROLOGIST, MEDICAL GENETICIST, LIPIDOLOGIST, OR A METABOLIC SPECIALIST. |
| Coverage Duration | LAL INITIAL 6 OR 12 MONTHS, SEE OTHER CRITERIA. RENEWAL: 12 MONTHS |

| PA Criteria | Criteria Details |
|-----------------------|--|
| Other Criteria | <p>INITIAL: DIAGNOSIS OF LYSOSOMAL ACID LIPASE (LAL) DEFICIENCY, AS CONFIRMED BY THE PRESENCE OF CLINICAL FEATURES (E.G., HEPATOMEGALY, ELEVATED SERUM TRANSAMINASES, DYSLIPIDEMIA, SPLENOMEGALY) PLUS ANY OF THE FOLLOWING: A BLOOD TEST INDICATING LOW OR ABSENT LEVELS OF LAL ENZYME ACTIVITY, A DRIED BLOOD SPOT TEST INDICATING LOW OR ABSENT LAL ENZYME ACTIVITY, OR A GENETIC TEST INDICATING THE BI-ALLELIC PRESENCE OF ALTERED LIPA GENE(S).</p> <p>RENEWAL:DIAGNOSIS OF LYSOSOMAL ACID LIPASE (LAL) DEFICIENCY PRESENTING AFTER THE FIRST 6 MONTHS OF LIFE AND NOT CONSIDERED RAPIDLY PROGRESSIVE REQUIRES DOCUMENTED IMPROVEMENT IN ANY ONE OF THE FOLLOWING CLINICAL PARAMETERS ASSOCIATED WITH LYSOSOMAL ACID LIPASE (LAL) DEFICIENCY DURING THE PAST 6 MONTHS: A RELATIVE REDUCTION FROM BASELINE IN ANY ONE OF THE FOLLOWING LIPID LEVELS (LDL-C, NON-HDL-C, OR TRIGLYCERIDES), NORMALIZATION OF ASPARTATE AMINOTRANSFERASE (AST) BASED ON AGE- AND GENDER-SPECIFIC NORMAL RANGES, A DECREASE IN LIVER FAT CONTENT COMPARED TO BASELINE ASSESSED BY ABDOMINAL IMAGING (E.G., MULTI-ECHO GRADIENT ECHO [MEGE] MRI). DIAGNOSIS OF RAPIDLY PROGRESSIVE LYSOSOMAL ACID LIPASE (LAL) DEFICIENCY PRESENTING WITHIN THE FIRST 6 MONTHS OF LIFE: 12 MONTHS. A DIAGNOSIS OF LYSOSOMAL ACID LIPASE (LAL) DEFICIENCY PRESENTING AFTER THE FIRST 6 MONTHS OF LIFE AND NOT CONSIDERED RAPIDLY PROGRESSIVE: INITIAL: 6 MONTHS</p> |

SECUKINUMAB

Products Affected

- Cosentyx (2 Syringes)
- Cosentyx Pen (2 Pens)

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | PLAQUE PSORIASIS (PSO): MODERATE TO SEVERE PLAQUE PSORIASIS INVOLVING GREATER THAN OR EQUAL TO 5 PERCENT BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, OR GENITAL AREA. RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | PLAQUE PSORIASIS (PSO): PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A DERMATOLOGIST. PSORIATIC ARTHRITIS (PSA): PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A RHEUMATOLOGIST OR A DERMATOLOGIST. ANKYLOSING SPONDYLITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A RHEUMATOLOGIST. |
| Coverage Duration | INITIAL: 4 MONTHS. RENEWAL: 12 MONTHS FOR ALL DIAGNOSES |
| Other Criteria | INITIAL: PLAQUE PSORIASIS (PSO): PREVIOUS TRIAL OF HUMIRA AND ONE CONVENTIONAL THERAPY SUCH AS PUVA (PHOTOTHERAPY ULTRAVIOLET LIGHT A), UVB (ULTRAVIOLET LIGHT B), TOPICAL CORTICOSTEROIDS, CALCIPOTRIENE, ACITRETIN, METHOTREXATE, OR CYCLOSPORINE. PSORIATIC ARTHRITIS (PSA): PREVIOUS TRIAL OF HUMIRA AND ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG) SUCH AS METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE. ANKYLOSING SPONDYLITIS (AS): PREVIOUS TRIAL OF HUMIRA. |

SELEXIPAG

Products Affected

- Upravi oral tablet 1,000 mcg, 1,200 mcg, 1,400 mcg, 1,600 mcg, 200 mcg, 400 mcg, 600 mcg, 800 mcg
- Upravi oral tablets, dose pack

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | DOCUMENTED CONFIRMATORY PULMONARY ARTERIAL HYPERTENSION (PAH) DIAGNOSIS BASED ON RIGHT HEART CATHETERIZATION. PATIENT HAS NYHA-WHO FUNCTIONAL CLASS II-IV SYMPTOMS. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST |
| Coverage Duration | INITIAL AND RENEWAL: 12 MONTHS |
| Other Criteria | INITIAL: MEAN PULMONARY ARTERY PRESSURE (PAP) OF AT LEAST 25 MMHG OR GREATER, PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 3 WOOD UNITS. RENEWAL: PATIENT SHOWS IMPROVEMENT FROM BASELINE IN THE 6-MINUTE WALK DISTANCE OR PATIENT HAS A STABLE 6-MINUTE WALK DISTANCE WITH A STABLE/ IMPROVED WHO FUNCTIONAL CLASS. |

SILTUXIMAB

Products Affected

- Sylvant

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

SIMEPREVIR

Products Affected

- Olysio

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD-IDSA GUIDANCE AND ADDITIONAL CONSIDERATION FOR COVERAGE CONSISTENT WITH FDA LABELING. |
| Exclusion Criteria | |
| Required Medical Information | HCV RNA LEVEL WITHIN PAST 6 MONTHS. FOR ALL GENOTYPE 1A: NS3 80K POLYMORPHISM LAB TEST AT BASELINE. |
| Age Restrictions | |
| Prescriber Restrictions | GASTROENTEROLOGIST, INFECTIOUS DISEASE SPECIALIST, PHYSICIAN SPECIALIZING IN THE TREATMENT OF HEPATITIS (FOR EXAMPLE HEPATOLOGIST), OR A SPECIALLY TRAINED GROUP SUCH AS ECHO (EXTENSION FOR COMMUNITY HEALTHCARE OUTCOMES) MODEL. |
| Coverage Duration | CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD-IDSA GUIDANCE. |

| PA Criteria | Criteria Details |
|-----------------------|--|
| Other Criteria | <p>CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD-IDSa GUIDANCE. 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. PATIENT MUST NOT BE TAKING ANY OF THE FOLLOWING INTERACTING MEDICATIONS: CARBAMAZEPINE, PHENYTOIN, PHENOBARBITAL, OXCARBAZEPINE, RIFAMPIN, RIFABUTIN, RIFAPENTINE, ERYTHROMYCIN (DOES NOT INCLUDE TOPICAL FORMULATIONS), CLARITHROMYCIN, TELITHROMYCIN, ITRACONAZOLE, KETOCONAZOLE, POSACONAZOLE, FLUCONAZOLE (DOES NOT INCLUDE TOPICAL FORMULATIONS), VORICONAZOLE, DEXAMETHASONE, CISAPRIDE, CYCLOSPORINE, ROSUVASTATIN DOSE ABOVE 10MG, ATORVASTATIN DOSE ABOVE 40MG, OR ANY OF THE FOLLOWING HIV MEDICATIONS: COBICISTAT-CONTAINING MEDS (E.G., STRIBILD), ANY HIV PROTEASE INHIBITOR (ATAZANAVIR, FOSAMPRENAVIR, LOPINAVIR, INDINAVIR, NELFINAVIR, SAQUINAVIR, OR TIPRANAVIR) RITONAVIR, DARUNAVIR/RITONAVIR, DELAVIRDINE, ETRAVIRINE, NEVIRAPINE, EFAVIRENZ). PATIENT MUST ALSO NOT BE TAKING AMIODARONE IF ON COMBINATION REGIMEN OF SOVALDI AND OLYSIO.</p> |

SOFOSBUVIR

Products Affected

- Sovaldi

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD-IDSa GUIDANCE AND ADDITIONAL CONSIDERATION FOR COVERAGE CONSISTENT WITH FDA LABELING. |
| Exclusion Criteria | PATIENT WITH END STAGE RENAL DISEASE OR REQUIRES DIALYSIS. |
| Required Medical Information | FOR ALL GENOTYPE 1 PATIENTS USING OLYSIO AND SOVALDI AND HAVE GENOTYPE 1A: NS3 80K POLYMORPHISM LAB TEST AT BASELINE. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A GASTROENTEROLOGIST, INFECTIOUS DISEASE SPECIALIST, PHYSICIAN SPECIALIZING IN THE TREATMENT OF HEPATITIS (HEPATOLOGIST), OR A SPECIALLY TRAINED GROUP SUCH AS ECHO (EXTENSION FOR COMMUNITY HEALTHCARE OUTCOMES) MODEL |
| Coverage Duration | CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSa GUIDANCE |

| PA Criteria | Criteria Details |
|-----------------------|--|
| Other Criteria | <p>CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE. 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. FOR PATIENTS ON SOVALDI PLUS DAKLINZA REGIMENS THERE WILL BE NO APPROVALS FOR CONCURRENT USE OF ANY OF THESE (CONTRAINDICATED OR NOT RECOMMENDED BY THE MANUFACTURER) MEDICATIONS: AMIODARONE, CARBAMAZEPINE, PHENYTOIN, OR RIFAMPIN. REQUESTS FOR SOVALDI IN COMBINATION WITH DAKLINZA OR OLYSIO WILL REQUIRE THAT THE PATIENT ALSO MEETS ALL CRITERIA FOR THE RESPECTIVE AGENT USED (DAKLINZA OR OLYSIO).</p> |

SOFOSBUVIR/VELPATASVIR

Products Affected

- Epclusa

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD-IDSA GUIDANCE. |
| Exclusion Criteria | |
| Required Medical Information | HCV RNA LEVEL. |
| Age Restrictions | 18 YEARS OF AGE AND OLDER. |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH: GASTROENTEROLOGIST, INFECTIOUS DISEASE SPECIALIST, PHYSICIAN SPECIALIZING IN THE TREATMENT OF HEPATITIS (HEPATOLOGIST), OR A SPECIALLY TRAINED GROUP SUCH AS ECHO (EXTENSION FOR COMMUNITY HEALTHCARE OUTCOMES) MODEL. |
| Coverage Duration | CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE. |
| Other Criteria | CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE. HCV RNA LEVEL WITHIN PAST 6 MONTHS. PATIENT IS NOT CONCURRENTLY TAKING ANY OF THE FOLLOWING MEDICATIONS NOT RECOMMENDED BY THE MANUFACTURER: AMIODARONE, CARBAMAZEPINE, PHENYTOIN, PHENOBARBITAL, OXCARBAZEPINE, RIFAMPIN, RIFABUTIN, RIFAPENTINE, HIV REGIMEN THAT CONTAINS EFAVIRENZ, ROSUVASTATIN AT DOSES ABOVE 10MG, TIPRANAVIR/RITONAVIR OR TOPOTECAN. PATIENT MUST NOT HAVE SEVERE RENAL IMPAIRMENT, ESRD OR ON HEMODIALYSIS. RIBAVIRIN USE REQUIRED FOR PATIENTS WITH DECOMPENSATED CIRRHOSIS. |

SOFOSBUVIR/VELPATASVIR/VOXILAPREVIR

Products Affected

- Vosevi

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD-IDSA GUIDANCE. |
| Exclusion Criteria | SEVERE RENAL IMPAIRMENT, ESRD OR ON HEMODIALYSIS. MODERATE OR SEVERE HEPATIC IMPAIRMENT (CHILD-PUGH B OR C). |
| Required Medical Information | HCV RNA LEVEL WITHIN PAST 6 MONTHS |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH: GASTROENTEROLOGIST, INFECTIOUS DISEASE SPECIALIST, PHYSICIAN SPECIALIZING IN THE TREATMENT OF HEPATITIS (HEPATOLOGIST), OR A SPECIALLY TRAINED GROUP SUCH AS ECHO (EXTENSION FOR COMMUNITY HEALTHCARE OUTCOMES) MODEL. |
| Coverage Duration | CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE. |
| Other Criteria | CRITERIA WILL BE APPLIED CONSISTENT WITH CURRENT AASLD/IDSA GUIDANCE. PATIENT IS NOT CONCURRENTLY TAKING ANY OF THE FOLLOWING MEDICATIONS NOT RECOMMENDED BY THE MANUFACTURER: 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 OR TIPRANA VIR/RITONAVIR. |

SOMATROPIN - GROWTH HORMONE

Products Affected

- Humatrope
- Omnitrope
- Saizen
- Saizen click.easy
- Zomacton

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | ATHLETIC ENHANCEMENT, ANTI-AGING PURPOSES, GROWTH FAILURE WITH CLOSED EPIPHYSES. |
| Required Medical Information | INDUCTION - PATIENT'S HEIGHT AT LEAST 2 STANDARD DEVIATIONS (SD) BELOW THE MEAN HEIGHT FOR NORMAL CHILDREN OF THE SAME AGE AND GENDER. RENEWAL: GROWTH VELOCITY AND/OR TARGET HEIGHT. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH: ENDOCRINOLOGIST. |
| Coverage Duration | 12 MONTHS |
| Other Criteria | INITIAL: PREVIOUS TRIAL OF PREFERRED FORMULARY ALTERNATIVES NORDITROPIN AND GENOTROPIN PER FDA INDICATION. RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION. |

SOMATROPIN - SEROSTIM

Products Affected

- Serostim subcutaneous recon soln 4 mg, 5 mg, 6 mg

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | ATHLETIC ENHANCEMENT, ANTI-AGING PURPOSES |
| Required Medical Information | HIV/WASTING: MEETS CRITERIA OF WEIGHT LOSS: 10% UNINTENTIONAL WEIGHT LOSS OVER 12 MONTHS, OR 7.5% OVER 6 MONTHS, OR 5% BODY CELL MASS (BCM) LOSS WITHIN 6 MONTHS, OR A BCM LESS THAN 35% (MEN) OF TOTAL BODY WEIGHT AND A BODY MASS INDEX (BMI) LESS THAN 27 KG PER METER SQUARED, OR BCM LESS THAN 23% (WOMEN) OF TOTAL BODY WEIGHT AND A BODY MASS INDEX (BMI) LESS THAN 27 KG PER METER SQUARED, OR BMI LESS THAN 20 KG PER METER SQUARED. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A GASTROENTEROLOGIST, NUTRITIONAL SUPPORT SPECIALIST (SBS), OR INFECTIOUS DISEASE SPECIALIST |
| Coverage Duration | 3 MONTHS |
| Other Criteria | HIV/WASTING: CURRENTLY ON ANTIRETROVIRAL THERAPY. IF CURRENTLY ON GROWTH HORMONE, PATIENT HAS SHOWN CLINICAL BENEFIT IN MUSCLE MASS AND WEIGHT OR IF NOT ON GROWTH HORMONE, PATIENT HAS HAD INADEQUATE RESPONSE TO PREVIOUS THERAPY. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION. |

SOMATROPIN - ZORBTIVE

Products Affected

- Zorbtive

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | ATHLETIC ENHANCEMENT, ANTI-AGING PURPOSES |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A GASTROENTEROLOGIST |
| Coverage Duration | SHORT BOWEL: 4 WEEKS ONCE |
| Other Criteria | THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION. |

SOMATROPIN-NORDITROPIN AND GENOTROPIN

Products Affected

- Genotropin
- Genotropin MiniQuick
- Norditropin FlexPro

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | ATHLETIC ENHANCEMENT, ANTI-AGING PURPOSES, GROWTH FAILURE WITH CLOSED EPIPHYSES. |
| Required Medical Information | INDUCTION - PATIENT'S HEIGHT AT LEAST 2 STANDARD DEVIATIONS (SD) BELOW THE MEAN HEIGHT FOR NORMAL CHILDREN OF THE SAME AGE AND GENDER. RENEWAL: GROWTH VELOCITY AND/OR TARGET HEIGHT. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH: ENDOCRINOLOGIST. |
| Coverage Duration | 12 MONTHS |
| Other Criteria | RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT (I.E. INCREASED HEIGHT OR INCREASED GROWTH VELOCITY). THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION. |

SOMATROPIN-NUTROPIN AND NUTROPIN AQ

Products Affected

- Nutropin AQ Nuspin

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | ATHLETIC ENHANCEMENT, ANTI-AGING PURPOSES, GROWTH FAILURE DUE TO CKD IF PATIENT HAS HAD A RENAL TRANSPLANT, OR GROWTH FAILURE WITH CLOSED EPIPHYSES. |
| Required Medical Information | INDUCTION - PATIENT'S HEIGHT AT LEAST 2 STANDARD DEVIATIONS (SD) BELOW THE MEAN HEIGHT FOR NORMAL CHILDREN OF THE SAME AGE AND GENDER. RENEWAL: GROWTH VELOCITY AND/OR TARGET HEIGHT. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH: ENDOCRINOLOGIST. FOR GROWTH HORMONE FAILURE DUE TO CRI: NEPHROLOGIST. |
| Coverage Duration | INITIAL AND RENEWAL: 12 MONTHS |
| Other Criteria | ALL DIAGNOSES EXCEPT FOR CHRONIC KIDNEY DISEASE (CKD): INITIAL: PREVIOUS TRIAL OF PREFERRED FORMULARY ALTERNATIVES NORDITROPIN AND GENOTROPIN PER FDA INDICATION. RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT (I.E. INCREASED HEIGHT OR INCREASED GROWTH VELOCITY). FOR GROWTH FAILURE SECONDARY TO CKD: PATIENT HAS NOT RECEIVED A RENAL TRANSPLANT. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION. |

SONIDEGIB

Products Affected

- Odomzo

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | BASELINE SERUM CREATINE KINASE (CK) AND SERUM CREATININE LEVELS |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

SORAFENIB TOSYLATE

Products Affected

- Nexavar

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

SUNITINIB MALATE

Products Affected

- Sutent

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | GASTROINTESTINAL STROMAL TUMORS (GIST): TRIAL OF OR CONTRAINDICATION TO GLEEVEC. |

TADALAFIL

Products Affected

- Cialis oral tablet 2.5 mg, 5 mg

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | ERECTILE DYSFUNCTION WITHOUT DIAGNOSIS OF BENIGN PROSTATIC HYPERPLASIA. |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | TRIAL OF ONE FORMULARY ALPHA BLOCKER SUCH AS DOXAZOSIN, TERAZOSIN, TAMSULOSIN OR ALFUZOSIN) AND ONE FORMULARY 5-ALPHA-REDUCTASE (SUCH AS FINASTERIDE OR DUTASTERIDE). APPLIES TO 2.5MG AND 5MG STRENGTHS ONLY. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION. |

TALIMOGENE

Products Affected

- Imlygic injection suspension 10exp6 (1 million) PFU/mL, 10exp8 (100 million) PFU/mL

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | HISTORY OF PRIMARY OR ACQUIRED IMMUNODEFICIENT STATES, LEUKEMIA, LYMPHOMA, OR AIDS. PATIENT IS NOT CURRENTLY RECEIVING IMMUNOSUPPRESSIVE THERAPY. |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | IMLYGIC TO BE INJECTED INTO CUTANEOUS, SUBCUTANEOUS, AND OR NODAL LESIONS THAT ARE VISIBLE, PALPABLE, OR DETECTABLE BY ULTRASOUND GUIDANCE. NO CONCURRENT USE WITH PEMBROLIZUMAB, NIVOLUMAB, IPILIMUMAB, DABRAFENIB, TRAMETINIB, VEMURAFENIB, INTERLEUKIN-2, INTERFERON, DACARBAZINE, TEMOZOLOMIDE, PACLITAXEL, CARBOPLATIN, IMATINIB, MELPHALAN, IMIQUIMOD, OR RADIATION THERAPY. |

TASIMELTEON

Products Affected

- Hetlioz

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

TEDUGLUTIDE

Products Affected

- Gattex 30-Vial

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | 18 YEARS OF AGE AND OLDER |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | PATIENT IS DEPENDENT ON INTRAVENOUS PARENTERAL NUTRITION DEFINED AS REQUIRING PARENTERAL NUTRITION AT LEAST THREE TIMES PER WEEK. |

TELOTRISTAT

Products Affected

- Xermelo

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

TEMOZOLOMIDE

Products Affected

- Temodar intravenous

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

TERIFLUNOMIDE

Products Affected

- Aubagio

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

TERIPARATIDE

Products Affected

- Forteo

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | 24 MONTHS OR MORE OF ANABOLIC THERAPY. |
| Required Medical Information | ONE OF THE FOLLOWING: (1) HIGH RISK FOR FRACTURES DEFINED AS ONE OF THE FOLLOWING: A) HISTORY OF OSTEOPOROTIC (I.E., FRAGILITY, LOW TRAUMA) FRACTURE(S). B) 2 OR MORE RISK FACTORS FOR FRACTURE (E.G., HISTORY OF MULTIPLE RECENT LOW TRAUMA FRACTURES, BMD T-SCORE LESS THAN OR EQUAL TO -2.5, CORTICOSTEROID USE, OR USE OF GNRH ANALOGS SUCH AS NAFARELIN, ETC.). C) NO PRIOR TREATMENT FOR OSTEOPOROSIS AND FRAX SCORE OF AT LEAST 20% FOR ANY MAJOR FRACTURE OR OF AT LEAST 3% FOR HIP FRACTURE. (2) UNABLE TO USE ORAL THERAPY (I.E., UPPER GASTROINTESTINAL PROBLEMS UNABLE TO TOLERATE ORAL MEDICATION, LOWER GASTROINTESTINAL PROBLEMS UNABLE TO ABSORB ORAL MEDICATIONS, TROUBLE REMEMBERING TO TAKE ORAL MEDICATIONS OR COORDINATING AN ORAL BISPHOSPHONATE WITH OTHER ORAL MEDICATIONS OR THEIR DAILY ROUTINE). (3) ADEQUATE TRIAL OF, INTOLERANCE TO, OR A CONTRAINDICATION TO BISPHOSPHONATES (E.G., ALENDRONATE, RISEDRONATE, IBANDRONATE). |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

TESTOSTERONE

Products Affected

- Androderm (1.62 mg/2.5 gram)
- AndroGel transdermal gel in metered-dose pump 20.25 mg/1.25 gram (1.62 %)
- AndroGel transdermal gel in packet 1.62 % (20.25 mg/1.25 gram), 1.62 % (40.5 mg/2.5 gram)
- testosterone cypionate
- testosterone enanthate
- testosterone transdermal gel in packet

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. ADDITIONAL CONSIDERATION FOR GENDER DYSPHORIA. |
| Exclusion Criteria | |
| Required Medical Information | MALE HYPOGONADISM CONFIRMED BY EITHER: 1) LAB CONFIRMED TOTAL SERUM TESTOSTERONE LEVEL OF LESS THAN 300 NG/DL OR 2) A LOW TOTAL SERUM TESTOSTERONE LEVEL AS INDICATED BY A LAB RESULT WITH A REFERENCE RANGE OBTAINED WITHIN 90 DAYS, OR 3) A FREE SERUM TESTOSTERONE LEVEL OF LESS THAN 5 PG/ML. |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | LIFETIME OF MEMBERSHIP IN PLAN |
| Other Criteria | |

TETRABENAZINE

Products Affected

- tetrabenazine

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | NEUROLOGIST |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

THALIDOMIDE

Products Affected

- Thalomid

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

TOCILIZUMAB IV

Products Affected

- Actemra

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | RENEWAL FOR RA, PJIA, OR SJIA: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | MODERATE TO SEVERE RHEUMATOID ARTHRITIS (RA)/POLYARTICULAR JUVENILE IDIOPATHIC ARTHRITIS (PJIA)/ SYSTEMIC JUVENILE IDIOPATHIC ARTHRITIS (SJIA): PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A RHEUMATOLOGIST |
| Coverage Duration | INITIAL: RA: 6 MONTHS. PJIA: 5 MOS. SJIA: 12 MOS. CRS: 1 MO. RENEWAL: 12 MOS FOR RA, PJIA, OR SJIA |
| Other Criteria | INITIAL: MODERATE TO SEVERE RA AND PJIA: PREVIOUS TRIAL OF HUMIRA AND ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG) AGENT SUCH AS METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE. INITIAL SJIA: PREVIOUS TRIAL WITH ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG) AGENT SUCH AS METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE. |

TOCILIZUMAB SQ

Products Affected

- Actemra

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | RA RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | RHEUMATOID ARTHRITIS (RA): PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A RHEUMATOLOGIST. |
| Coverage Duration | RA INITIAL: 6 MONTHS. RA RENEWAL: 12 MONTHS. GIANT CELL ARTERITIS: 12 MONTHS |
| Other Criteria | RA INITIAL : PREVIOUS TRIAL OF HUMIRA AND ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG) AGENT SUCH AS METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE. |

TOFACITINIB

Products Affected

- Xeljanz
- Xeljanz XR

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | RENEWAL: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | RHEUMATOID ARTHRITIS: PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A RHEUMATOLOGIST. |
| Coverage Duration | RA: INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS. |
| Other Criteria | INITIAL: RHEUMATOID ARTHRITIS: PREVIOUS TRIAL OF HUMIRA AND ONE DMARD (DISEASE-MODIFYING ANTIRHEUMATIC DRUG) SUCH AS METHOTREXATE, LEFLUNOMIDE, HYDROXYCHLOROQUINE, OR SULFASALAZINE. |

TOPICAL TRETINOIN

Products Affected

- tretinoin topical cream
- tretinoin topical gel 0.01 %, 0.025 %

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | WRINKLES, PHOTOAGING, MELASMA. |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

TRABECTEDIN

Products Affected

- Yondelis

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

TRAMETINIB DIMETHYL SULFOXIDE

Products Affected

- Mekinist oral tablet 0.5 mg, 2 mg

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

TRASTUZUMAB

Products Affected

- Herceptin

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | BREAST CANCER, METASTATIC BREAST CANCER, GASTRIC CANCER: HER2 POSITIVE |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | B VS D COVERAGE CONSIDERATION. |

TREPROSTINIL DIOLAMINE

Products Affected

- Orenitram

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | PATIENT DOES NOT HAVE SEVERE HEPATIC IMPAIRMENT. |
| Required Medical Information | DOCUMENTED CONFIRMATORY PULMONARY ARTERIAL HYPERTENSION (PAH) DIAGNOSIS BASED ON RIGHT HEART CATHETERIZATION. PATIENT HAS NYHA-WHO FUNCTIONAL CLASS II-IV SYMPTOMS. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST. |
| Coverage Duration | INITIAL AND RENEWAL: 12 MONTHS |
| Other Criteria | INITIAL: MEAN PULMONARY ARTERY PRESSURE (PAP) OF AT LEAST 25 MMHG OR GREATER, PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 3 WOOD UNITS. PREVIOUS OR CURRENT TREATMENT WITH ONE OF THE FOLLOWING AGENTS: A FORMULARY PHOSPHODIESTERASE-5 INHIBITOR (E.G., SILDENAFIL [GENERIC FOR REVATIO] OR ADCIRCA [TADALAFIL]) OR AN ENDOTHELIN RECEPTOR ANTAGONIST (E.G., TRACLEER [BOSENTAN], LETAIRIS [AMBRISENTAN], OR OPSUMIT [MACITENTAN]). TRIAL OF A FORMULARY PHOSPHODIESTERASE-5 INHIBITOR OR ENDOTHELIN RECEPTOR ANTAGONIST IS NOT REQUIRED IF THE PATIENT WAS PREVIOUSLY STABLE ON ORENITRAM. RENEWAL: PATIENT SHOWS IMPROVEMENT FROM BASELINE IN THE 6-MINUTE WALK DISTANCE OR PATIENT HAS A STABLE 6-MINUTE WALK DISTANCE WITH A STABLE/ IMPROVED WHO FUNCTIONAL CLASS. |

TREPROSTINIL INHALED

Products Affected

- Tyvaso

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | DOCUMENTED CONFIRMATORY PULMONARY ARTERIAL HYPERTENSION (PAH) DIAGNOSIS BASED ON RIGHT HEART CATHETERIZATION. PATIENT HAS NYHA-WHO FUNCTIONAL CLASS III-IV SYMPTOMS. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST |
| Coverage Duration | INITIAL AND RENEWAL: 12 MONTHS |
| Other Criteria | THIS DRUG MAYBE COVERED UNDER MEDICARE PART B OR D DEPENDING UPON THE CIRCUMSTANCES. INFORMATION MAY NEED TO BE SUBMITTED DESCRIBING THE USE AND SETTING OF THE DRUG TO MAKE THE DETERMINATION. NEBULIZER THERAPY IS COVERED UNDER PART B FOR PATIENTS WHO ARE USING THE MEDICATION VIA A NEBULIZER IN THEIR OWN HOME. THOSE WHO ARE NOT USING IT IN THEIR HOME WILL BE COVERED UNDER PART D. INITIAL: MEAN PULMONARY ARTERY PRESSURE (PAP) OF AT LEAST 25 MMHG OR GREATER, PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 3 WOOD UNITS. RENEWAL: PATIENT SHOW IMPROVEMENT FROM BASELINE IN THE 6-MINUTE WALK DISTANCE OR PATIENT HAS A STABLE 6-MINUTE WALK DISTANCE WITH A STABLE/ IMPROVED WHO FUNCTIONAL CLASS. |

TREPROSTINIL SODIUM INJECTABLE

Products Affected

- Remodulin

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | COVERED UNDER LOCAL COVERAGE POLICY OF APPLICABLE MEDICARE DMERC. |
| Required Medical Information | FORMULARY DRUG ADMINISTERED IN A LONG TERM CARE FACILITY TO A PATIENT WHOSE PART A COVERAGE HAS EXPIRED OR FORMULARY DRUG NOT ADMINISTERED VIA AN IMPLANTABLE PUMP OR AN EXTERNAL PUMP OR DRUG ADMINISTERED VIA AN IMPLANTABLE PUMP/AN EXTERNAL PUMP. DOCUMENTED CONFIRMATORY PULMONARY ARTERIAL HYPERTENSION (PAH) DIAGNOSIS BASED ON RIGHT HEART CATHETERIZATION. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR IN CONSULTATION WITH A CARDIOLOGIST OR PULMONOLOGIST |
| Coverage Duration | INITIAL AND RENEWAL: 12 MONTHS |

| PA Criteria | Criteria Details |
|-----------------------|---|
| Other Criteria | <p>INITIAL: MEAN PULMONARY ARTERY PRESSURE (PAP) OF AT LEAST 25 MMHG OR GREATER, PULMONARY CAPILLARY WEDGE PRESSURE (PCWP) OF 15 MMHG OR LESS, PULMONARY VASCULAR RESISTANCE (PVR) GREATER THAN 3 WOOD UNITS. CONTINUATION OF CURRENT REMODULIN THERAPY: PATIENT MUST HAVE NYHA/WHO FC II-IV SYMPTOMS. NEW REQUESTS FOR REMODULIN THERAPY: PATIENT MUST HAVE NYHA/WHO FC III-IV SYMPTOMS. NEW REQUESTS FOR REMODULIN THERAPY FOR PATIENTS WITH NYHA/WHO FC II SYMPTOMS REQUIRES A TRIAL OF OR CONTRAINDICATION TO A PHOSPHODIESTERASE-5 INHIBITOR (PDE-5) (E.G., REVATIO, ADCIRCA) OR AN ENDOTHELIN RECEPTOR ANTAGONIST (ERA) (E.G., LETAIRIS, OPSUMIT, TRACLEER). RENEWAL: PATIENT SHOW IMPROVEMENT FROM BASELINE IN THE 6-MINUTE WALK DISTANCE OR PATIENT HAS A STABLE 6-MINUTE WALK DISTANCE WITH A STABLE/ IMPROVED WHO FUNCTIONAL CLASS.</p> |

TRIENTINE

Products Affected

- Syprine

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | KNOWN FAMILY HISTORY OF WILSON'S DISEASE OR PHYSICAL EXAMINATION CONSISTENT WITH WILSON'S DISEASE. PLASMA COPPER-PROTEIN CERULOPLASMIN LESS THAN 20 MG/DL. LIVER BIOPSY POSITIVE FOR AN ABNORMALLY HIGH CONCENTRATION OF COPPER (GREATER THAN 250 MCG/G DRY WEIGHT) OR THE PRESENCE OF KAYSER-FLEISCHER RINGS. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A HEPATOLOGIST. |
| Coverage Duration | 12 MONTHS |
| Other Criteria | PREVIOUS TRIAL OF OR CONTRAINDICATION TO PENICILLAMINE (DEPEN). |

TRIFLURIDINE/TIPIRACIL

Products Affected

- Lonsurf oral tablet 15-6.14 mg, 20-8.19 mg

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

URIDINE TRIACETATE

Products Affected

- Xuriden

| PA Criteria | Criteria Details |
|------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | INITIAL: DIAGNOSIS CONFIRMED BY 1) GENETIC MUTATION OF URIDINE MONOPHOSPHATE SYNTHASE (UMPS) GENE AND 2) ELEVATED URINE OROTIC ACID PER AGE-SPECIFIC REFERENCE RANGE. RENEWAL: IMPROVEMENT FROM BASELINE OR STABILIZATION OF AGE DEPENDENT HEMATOLOGIC PARAMETERS (E.G., NEUTROPHIL COUNT, NEUTROPHIL PERCENT, WBC COUNT, MEAN CORPUSCULAR VOLUME) |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | INITIAL: 6 MONTHS. RENEWAL: 12 MONTHS |
| Other Criteria | |

USTEKINUMAB

Products Affected

- Stelara subcutaneous syringe

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | INITIAL: PLAQUE PSORIASIS: MODERATE TO SEVERE PLAQUE PSORIASIS INVOLVING GREATER THAN OR EQUAL TO 5% BODY SURFACE AREA OR PSORIATIC LESIONS AFFECTING THE HANDS, FEET, OR GENITAL AREA. RENEWAL FOR PSORIATIC ARTHRITIS OR PLAQUE PSORIASIS: PHYSICIAN ATTESTATION OF IMPROVEMENT. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH: PSORIATIC ARTHRITIS: DERMATOLOGIST OR RHEUMATOLOGIST. PLAQUE PSORIASIS: DERMATOLOGIST. CROHN'S DISEASE: GASTROENTEROLOGIST. |
| Coverage Duration | INITIAL: PSA, PSO, CD: 4 MONTHS. CD WITH PREVIOUS DOSE IV: 2 MONTHS. RENEW ALL: 12 MO |
| Other Criteria | INITIAL: PSORIATIC ARTHRITIS (PSA): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING PREFERRED AGENTS: CIMZIA, OTEZLA, OR COSENTYX. PLAQUE PSORIASIS (PSO): PREVIOUS TRIAL OF HUMIRA FOLLOWED BY ONE OF THE FOLLOWING PREFERRED AGENTS: COSENTYX OR OTEZLA. CROHN'S DISEASE (CD): PREVIOUS TRIAL OF OR CONTRAINDICATION TO PREFERRED TNF INHIBITORS: HUMIRA FOLLOWED BY CIMZIA. |

USTEKINUMAB IV

Products Affected

- Stelara intravenous

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A GASTROENTEROLOGIST. |
| Coverage Duration | 2 MONTHS |
| Other Criteria | PREVIOUS TRIAL OF OR CONTRAINDICATION TO THE PREFERRED TNF INHIBITORS: HUMIRA FOLLOWED BY CIMZIA. THIS DRUG ALSO REQUIRES PAYMENT DETERMINATION AND MAY BE COVERED UNDER MEDICARE PART B OR D. |

VALBENZAZINE TOSYLATE

Products Affected

- Ingrezza oral capsule 40 mg, 80 mg

| PA Criteria | Criteria Details |
|-------------------------------------|---|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | PATIENT HAS BEEN STABLE ON ANTIPSYCHOTIC MEDICATIONS OR METOCLOPRAMIDE PER PHYSICIAN ATTESTATION. |
| Age Restrictions | |
| Prescriber Restrictions | PRESCRIBED BY OR GIVEN IN CONSULTATION WITH A NEUROLOGIST, PSYCHIATRIST, OR MOVEMENT DISORDER SPECIALIST. |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

VANDETANIB

Products Affected

- Caprelsa oral tablet 100 mg, 300 mg

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

VEMURAFENIB

Products Affected

- Zelboraf

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | BRAFV600E MUTATION |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

VENETOCLAX

Products Affected

- Venclexta oral tablet 10 mg, 100 mg, 50 mg
- Venclexta Starting Pack

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

VINCRIStINE SULFATE LIPOSOMAL

Products Affected

- Marqibo

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| 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. |

VISMODEGIB

Products Affected

- Erivedge

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

ZIV-AFLIBERCEPT

Products Affected

- Zaltrap

| PA Criteria | Criteria Details |
|-------------------------------------|--|
| Covered Uses | ALL FDA APPROVED INDICATIONS NOT OTHERWISE EXCLUDED FROM PART D. |
| Exclusion Criteria | |
| Required Medical Information | |
| Age Restrictions | |
| Prescriber Restrictions | |
| Coverage Duration | 12 MONTHS |
| Other Criteria | |

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