## PA Criteria

Prior Authorization Group

Drug Names

ACITRETIN ACITRETIN

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D, prevention of non-

melanoma skin cancers in high risk individuals.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

ACTIMMUNE ACTIMMUNE

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D, mycosis fungoides,

Sezary syndrome, atopic dermatitis.

**Exclusion Criteria** 

**Required Medical Information** 

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

ADAGEN ADAGEN

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Plan Year

Other Criteria

Prior Authorization Group ADEMPAS
Drug Names ADEMPAS

Covered Uses All FDA-approved indications not otherwise excluded from Part D.

Exclusion Criteria

Required Medical Information

For pulmonary arterial hypertension (PAH) (WHO Group 1): PAH was confirmed by right heart catheterization. For chronic thromboembolic pulmonary hypertension (CTEPH) (WHO Group 4): Patient has persistent or recurrent CTEPH after pulmonary endarterectomy (PEA), OR patient has inoperable CTEPH with the diagnosis confirmed by right heart catheterization AND by computed tomography (CT), magnetic resonance imaging (MRI), or pulmonary angiography. For new starts only (excluding recurrent/persistent CTEPH after PEA): 1) pretreatment mean pulmonary arterial pressure is greater than or equal to 25 mmHg, AND 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, AND 3) pretreatment pulmonary vascular resistance is greater than 3 Wood units.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Plan Year

Prior Authorization Group
Drug Names
Covered Uses

AFINITOR

AFINITOR, AFINITOR DISPERZ

All FDA-approved indications not otherwise excluded from Part D, classical Hodgkin lymphoma, thymomas and thymic carcinomas, Waldenstrom's macroglobulinemia/lymphoplasmacytic lymphoma, soft tissue sarcoma subtypes: perivascular epithelioid cell tumors (PEComa), angiomyolipoma, lymphangioleiomyomatosis, osteosarcoma.

Exclusion Criteria
Required Medical Information

Breast cancer: 1) The patient has recurrent or metastatic hormone receptor positive. HER2 negative disease, AND 2) Afinitor will be used in combination with exemestane, AND 3) The patient's disease a) has progressed while on or within 12 months of nonsteroidal aromatase inhibitor therapy, OR b) was previously treated with tamoxifen. Renal cell carcinoma: 1) The disease is relapsed or unresectable, AND 2) For disease that is of non- clear cell histology, Afinitor will be used as first-line systemic therapy AND Afinitor will be used as a single agent, AND 3) For disease that is of predominantly clear cell histology, Afinitor will be used as a single agent or in combination with Lenvima AND disease has progressed on prior anti-angiogenic therapy (e.g. sunitinib). Classical Hodgkin lymphoma: 1) Afinitor will be used as a single agent, AND 2) Patient meets ONE of the following: a) The disease is relapsed or refractory, OR b) Afinitor will be used as palliative therapy. Thymomas and Thymic carcinomas: 1) The disease has progressed on a platinum-based chemotherapy regimen, AND 2) Afinitor will be used as a single agent. Soft tissue sarcoma: 1) The patient has one of the following subtypes of STS: a) Perivascular epithelioid cell tumors (PEComa), or b) Angiomyolipoma, or c) Lymphangioleiomyomatosis, AND 2) Afinitor will be used as a single agent. Osteosarcoma: Afinitor will be used in combination with sorafenib [Nexavar]. Subependymal giant cell astrocytoma associated with tuberous sclerosis complex (TSC): The patient is not a candidate for curative surgical resection. Renal angiomyolipoma associated with TSC: The patient does not require immediate surgery.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Plan Year

**Drug Names** 

**ALDURAZYME ALDURAZYME** 

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Diagnosis of mucopolysaccharidosis I was confirmed by an enzyme assay

demonstrating a deficiency of alpha-L-iduronidase enzyme activity or by genetic testing. Patients with Scheie syndrome must have moderate to severe symptoms.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

**ALECENSA** 

**ALECENSA** 

All FDA-approved indications not otherwise excluded from Part D.

For NSCLC, patient meets all of the following: 1) Tumor is ALK-positive, and 2)

Disease is recurrent or metastatic.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

**ALIQOPA** 

**ALIQOPA** 

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

Drug Names
Covered Uses

Exclusion Criteria

**Required Medical Information** 

**ALOSETRON** 

ALOSETRON HYDROCHLORIDE

All FDA-approved indications not otherwise excluded from Part D.

1) The requested drug is being prescribed for a biological female or a person that self-identifies as a female with a diagnosis of severe diarrhea-predominant irritable bowel syndrome (IBS) AND 2) Chronic IBS symptoms lasting at least 6 months AND 3) Gastrointestinal tract abnormalities have been ruled out AND 4) Inadequate response to conventional therapy.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

Drug Names Covered Uses

**Exclusion Criteria** 

**Required Medical Information** 

ALPHA1-PROTEINASE INHIBITOR

ARALAST NP, GLASSIA, PROLASTIN-C, ZEMAIRA

All FDA-approved indications not otherwise excluded from Part D.

Patients must have clinically evident emphysema. Patients must have a pretreatment serum alpha1-proteinase inhibitor level less than 11 micromol/L (80 mg/dl by radial immunodiffusion or 50 mg/dl by nephelometry). Patients must have a pretreatment post-bronchodilation forced expiratory volume in 1 second (FEV1) greater than or equal to 25 percent and less than or equal to 80 percent of predicted.

Age Restrictions

Prescriber Restrictions

Coverage Duration
Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

Coverage Duration

Other Criteria

ALUNBRIG ALUNBRIG

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

**Drug Names** 

**AMPYRA AMPYRA** 

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

For new starts: Prior to initiating therapy, patient demonstrates sustained walking impairment. For continuation of therapy: Patient must have experienced an improvement in walking speed or other objective measure of walking ability since starting Ampyra.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

ANADROL ANADROL-50

All FDA-approved indications not otherwise excluded from Part D. Cachexia associated

with AIDS (HIV-wasting) or due to chronic disease, Fanconi's anemia.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

6 months

**Prior Authorization Group** 

**Drug Names** 

**APOKYN APOKYN** 

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Plan Year

Other Criteria

**Drug Names Covered Uses**  **ARANESP** 

ARANESP ALBUMIN FREE

All FDA-approved indications not otherwise excluded from Part D, anemia due to

myelodysplastic syndromes (MDS).

**Exclusion Criteria** 

**Required Medical Information** 

Patients receiving chemotherapy with curative intent. Patients with myeloid cancer. For all uses: Pretreatment (no erythropoietin treatment in previous month) hemoglobin

(Hgb) is less than 10 g/dL.

Age Restrictions **Prescriber Restrictions Coverage Duration** Other Criteria

16 weeks

Coverage under Part D will be denied if coverage is available under Part A or Part B as the medication is prescribed and dispensed or administered for the individual (eg, used for treatment of anemia for a patient with chronic renal failure who is undergoing dialysis, or furnished from physician's supply incident to a physician service). Requirements regarding Hgb values exclude values due to a recent transfusion. For reauthorizations (patient received erythropoietin in previous month): 1) For all uses, there is an increase in Hgb of at least 1 g/dL after at least 12 weeks of therapy, 2) For all uses except anemia due to myelosuppressive cancer chemotherapy: current Hgb is less than or equal to 12 g/dL, and 3) For anemia due to myelosuppressive cancer chemotherapy: current Hgb is less than 11 g/dL.

**Prior Authorization Group** 

**Drug Names Covered Uses**  ARCAI YST **ARCALYST** 

All FDA-approved indications not otherwise excluded from Part D. Prevention of gout

flares in patients initiating or continuing urate-lowering therapy.

**Exclusion Criteria** Required Medical Information

For prevention of gout flares in members initiating or continuing urate-lowering therapy (i.e., allopurinol or febuxostat) (new starts): all of the following criteria must be met: 1) serum uric acid concentration greater than or equal to 445 micromol/L (7.5 mg/dL) prior to initiating Arcalyst, 2) two or more gout flares within the previous 12 months, 3) inadequate response, intolerance or contraindication to maximum tolerated doses of non-steroidal anti-inflammatory drugs and colchicine, and 4) concurrent use with uratelowering therapy (i.e., allopurinol or febuxostat). For prevention of gout flares in members initiating or continuing urate-lowering therapy (i.e., allopurinol or febuxostat) (continuation): 1) Member must have achieved or maintain a clinical benefit (i.e., a fewer number of gout attacks or fewer flare days) compared to baseline and 2) have continued use of urate-lowering therapy concurrently with Arcalyst.

Age Restrictions **Prescriber Restrictions Coverage Duration** 

Other Criteria

For prevention of gout flares: 4 months. Other: Plan Year

Abbreviation: CAPS = Cryopyrin-Associated Periodic Syndromes.

CAPS: 12 years of age or older. Gout: 18 years of age or older.

**Drug Names** 

ARMODAFINIL ARMODAFINIL

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

1) Diagnosis is narcolepsy confirmed by sleep lab evaluation OR 2) Diagnosis is

obstructive sleep apnea (OSA) confirmed by polysomnography OR 3) Diagnosis is Shift

Work Disorder (SWD).

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

Required Medical Information

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

AUSTEDO AUSTEDO

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

Prior Authorization Group
Drug Names
Covered Uses

AVASTIN AVASTIN

All FDA-approved indications not otherwise excluded from Part D, breast cancer, central nervous system (CNS) tumor types: adult intracranial and spinal ependymoma and anaplastic gliomas, endometrial cancer, ovarian malignant sex cord-stromal tumors, soft tissue sarcoma subtypes: angiosarcoma, solitary fibrous tumor, and hemangiopericytoma, malignant pleural mesothelioma, choroidal neovascularization associated with: ocular histoplasmosis, pathologic myopia, angioid streaks, inflammatory conditions, or of idiopathic etiology, neovascular (wet) age-related macular degeneration including polypoidal choroidopathy and retinal angiomatous proliferation subtypes, macular edema due to retinal vein occlusion, diabetic macular edema, ocular neovascularization of the choroid, retina, or iris associated with proliferative diabetic retinopathy, neovascular glaucoma, retinopathy of prematurity, and proliferative diabetic retinopathy (as adjunct prior to vitrectomy).

Exclusion Criteria
Required Medical Information

Colorectal cancer: The disease is unresectable advanced or metastatic. Non-squamous non-small cell lung cancer: the requested drug will be used as first-line therapy, subsequent therapy, or continuation maintenance therapy (ie, continuation of the requested drug as first-line therapy beyond 4-6 cycles in the absence of disease progression).

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Plan Year

Coverage under Part D will be denied if coverage is available under Part A or Part B as the medication is prescribed and dispensed or administered for the individual.

## Prior Authorization Group Drug Names

B VS. D

ABELCET, ABRAXANE, ACETYLCYSTEINE, ACYCLOVIR SODIUM, ADRIAMYCIN, ADRUCIL, ALBUTEROL SULFATE, ALIMTA, AMBISOME, AMINOSYN, AMINOSYN 7%/ELECTROLYTES. AMINOSYN 8.5%/ELECTROLYTE. AMINOSYN II. AMINOSYN II 8.5%/ELECTROL. AMINOSYN M. AMINOSYN-HBC. AMINOSYN-PF. AMINOSYN-PF 7%, AMINOSYN-RF, AMPHOTERICIN B, APREPITANT, ARRANON, ARZERRA, ASTAGRAF XL, ATGAM, AZACITIDINE, AZASAN, AZATHIOPRINE, BENDEKA, BLEOMYCIN SULFATE, BROVANA, BUDESONIDE, BUSULFAN, CALCITONIN-SALMON, CALCITRIOL, CAMPTOSAR, CARBOPLATIN, CESAMET, CISPLATIN, CLADRIBINE, CLINIMIX 2.75%/DEXTROSE 5, CLINIMIX 4.25%/DEXTROSE 1, CLINIMIX 4.25%/DEXTROSE 2, CLINIMIX 4.25%/DEXTROSE 5, CLINIMIX 5%/DEXTROSE 15%, CLINIMIX 5%/DEXTROSE 20%, CLINIMIX 5%/DEXTROSE 25%, CLINIMIX E 2.75%/DEXTROSE, CLINIMIX E 4.25%/DEXTROSE, CLINIMIX E 5%/DEXTROSE 15, CLINIMIX E 5%/DEXTROSE 20, CLINIMIX E 5%/DEXTROSE 25. CLINISOL SF 15%, CLOFARABINE, COSMEGEN, CROMOLYN SODIUM. CYCLOPHOSPHAMIDE, CYCLOSPORINE, CYCLOSPORINE MODIFIED, CYTARABINE AQUEOUS, DACARBAZINE, DACTINOMYCIN, DECITABINE, DEPO-MEDROL, DEPO-PROVERA, DEXRAZOXANE, DIPHTHERIA/TETANUS TOXOID, DOCETAXEL, DOXERCALCIFEROL, DOXORUBICIN HCL, DOXORUBICIN HCL LIPOSOME, DRONABINOL, DUOPA, ELIGARD, ELITEK, EMEND, ENGERIX-B. ENVARSUS XR, EPIRUBICIN HCL, ERBITUX, ETOPOPHOS, ETOPOSIDE, FASLODEX, FIRMAGON, FLUDARABINE PHOSPHATE, FLUOROURACIL. FREAMINE HBC 6.9%, FREAMINE III, GAMASTAN S/D, GANCICLOVIR, GEMCITABINE, GEMCITABINE HCL, GENGRAF, GRANISETRON HCL, HALAVEN, HECTOROL, HEPARIN SODIUM, HEPATAMINE, HUMULIN R U-500 (CONCENTR, HYDROMORPHONE HCL, HYDROXYPROGESTERONE CAPRO, IBANDRONATE SODIUM, IFEX, IFOSFAMIDE, INTRALIPID, INTRON A, IPRATROPIUM BROMIDE, IPRATROPIUM BROMIDE/ALBUT, IRINOTECAN, IXEMPRA KIT, KADCYLA, KEPIVANCE, LEUCOVORIN CALCIUM, LEVALBUTEROL, LEVALBUTEROL HCL. LEVOCARNITINE, LEVOLEUCOVORIN, LIDOCAINE HCL, MEDROL, MELPHALAN HYDROCHLORIDE, MESNA, METHOTREXATE SODIUM, METHYLPREDNISOLONE, METHYLPREDNISOLONE ACETAT, METHYLPREDNISOLONE SODIUM, MIACALCIN, MILLIPRED, MITOMYCIN, MITOXANTRONE HCL, MORPHINE SULFATE, MUSTARGEN, MYCOPHENOLATE MOFETIL, MYCOPHENOLIC ACID DR, NEBUPENT, NEPHRAMINE, NIPENT, NULOJIX, NUTRILIPID, ONDANSETRON HCL, ONDANSETRON ODT, ONIVYDE, OXALIPLATIN, PACLITAXEL, PAMIDRONATE DISODIUM, PARICALCITOL, PERFOROMIST, PLENAMINE, PREDNISOLONE, PREDNISOLONE SODIUM PHOSP, PREDNISONE, PREDNISONE INTENSOL, PREMASOL, PROCALAMINE. PROGRAF, PROSOL, RAPAMUNE, RAYOS, RECOMBIVAX HB, SANDIMMUNE, SENSIPAR, SIROLIMUS, SMOFLIPID, SOLU-MEDROL, SYNDROS, TACROLIMUS. TAXOTERE, TENIVAC, TETANUS/DIPHTHERIA TOXOID, THIOTEPA. THYMOGLOBULIN, TOPOSAR, TOPOTECAN HCL, TORISEL, TPN

ELECTROLYTES, TRAVASOL, TREANDA, TREXALL, TRISENOX, TROPHAMINE, VARUBI, VECTIBIX, VINBLASTINE SULFATE, VINCASAR PFS, VINCRISTINE SULFATE, VINORELBINE TARTRATE, XATMEP, ZANOSAR, ZOLEDRONIC ACID. ZOMETA. ZORTRESS. ZUPLENZ

**Covered Uses** 

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.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

Prescriber Restrictions

**Coverage Duration** 

Other Criteria

N/A

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

**BANZEL** 

**BANZEL** 

All FDA-approved indications not otherwise excluded from Part D.

1 year of age or older.

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

**BELBUCA** 

**BELBUCA** 

All FDA-approved indications not otherwise excluded from Part D.

1) The requested drug is being prescribed for pain associated with cancer, a terminal condition, or pain being managed through hospice or palliative care OR 2) The requested drug is being prescribed for pain severe enough to require daily, around-theclock, long-term treatment in a patient who has been taking an opioid AND 3) The patient can safely take the requested dose based on their history of opioid use AND 4) The patient has been evaluated and will be monitored for the development of opioid

use disorder

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Drug Names** 

**BELEODAQ** BELEODAQ

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D, adult T-cell leukemia/lymphoma, mycosis fungoides/sezary syndrome, and primary cutaneous CD30+ T cell lymphoproliferative disorders: relapsed or refractory cutaneous anaplastic large cell lymphoma.

**Exclusion Criteria** 

**Required Medical Information** 

For ATLL: patient must be a non-responder to first-line therapy and belinostat is used

for acute disease or lymphoma.

Age Restrictions

Prescriber Restrictions **Coverage Duration** 

Plan Year

Other Criteria

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** Required Medical Information All FDA-approved indications not otherwise excluded from Part D.

Severe active lupus nephritis. Severe active central nervous system lupus.

Patient has been diagnosed with active, autoantibody-positive systemic lupus erythematosus (SLE). Patient is currently receiving standard therapy for SLE (e.g., corticosteroids, azathioprine, leflunomide, methotrexate, mycophenolate mofetil, hydroxychloroquine, non-steroidal anti-inflammatory drugs) OR patient is not currently

receiving standard therapy for SLE because patient tried and had an inadequate

response or intolerance to standard therapy.

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**BESPONSA** 

**BENLYSTA** 

**BENLYSTA** 

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**BESPONSA** All FDA-approved indications not otherwise excluded from Part D.

**Required Medical Information** For relapsed or refractory B-cell precursor acute lymphoblastic leukemia: The tumor is

CD22-positive as confirmed by testing or analysis to identify the CD22 protein on the surface of the B-cell.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Plan Year

Other Criteria

**Drug Names** 

BETASERON BETASERON

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Have a relapsing form of multiple sclerosis (MS) (e.g., relapsing-remitting MS,

progressive-relapsing MS, or secondary progressive MS with relapses) OR first clinical

episode of MS.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

Drug Names

**BEXAROTENE** 

BEXAROTENE, TARGRETIN

**Covered Uses**All FDA-approved indications not otherwise excluded from Part D, mycosis fungoides,

Sezary syndrome (capsules only), primary cutaneous CD30-positive T-cell

lymphoproliferative disorder types: primary cutaneous anaplastic large cell lymphoma

(capsules only) and lymphomatoid papulosis (capsules only), adult T-cell

leukemia/lymphoma (gel only), primary cutaneous B-cell lymphoma types: primary cutaneous marginal zone lymphoma (gel only) and primary cutaneous follicle center

lymphoma (gel only).

**Exclusion Criteria** 

**Required Medical Information** 

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Drug Names** 

**TRACLEER** 

**Covered Uses** 

**BOSENTAN** 

**Exclusion Criteria Required Medical Information** 

Pulmonary arterial hypertension (WHO Group 1) was confirmed by right heart

All FDA-approved indications not otherwise excluded from Part D.

catheterization. For new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than or equal to 25 mmHg, 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, and 3) pretreatment pulmonary vascular

resistance is greater than 3 Wood units.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

**BOSULIF BOSULIF** 

All FDA-approved indications not otherwise excluded from Part D.

Diagnosis of chronic myelogenous leukemia (CML) was confirmed by detection of the

Philadelphia chromosome or BCR-ABL gene. For CML, patient meets any of the following: 1) Patient has chronic phase, accelerated phase, or blast phase CML and meets one of the following: a) has not received prior therapy with a tyrosine kinase inhibitor (TKI) (e.g., imatinib, dasatinib, nilotinib, ponatinib), b) experienced intolerance or toxicity to a prior TKI, or c) experienced resistance to a prior TKI and is negative for

T315I mutation, OR 2) Patient received a hematopoietic stem cell transplant.

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

18 years of age or older

Prior Authorization Group BOTOX
Drug Names BOTOX

**Covered Uses**All FDA-approved indications not otherwise excluded from Part D, excessive salivation

secondary to advanced Parkinson's disease, hemifacial spasm.

**Exclusion Criteria** Cosmetic use.

**Required Medical Information** For chronic migraine prophylaxis, initial treatment: patient experiences at least 15

headache days per month, and patient had an inadequate response to at least 4 weeks of oral migraine preventative therapy. For chronic migraine prophylaxis, continuation of treatment (after 2 injection cycles): More headache-free days per month since starting therapy. For urinary incontinence in a patient with a neurologic condition (eg, spinal cord injury, multiple sclerosis) or with overactive bladder: patient had an inadequate

response to or is intolerant of an anticholinergic medication.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** Chronic migraine, initial tx: 6 months, renewal: plan year. Plan Year for all other

indications.

Other Criteria

Prior Authorization Group BRIVIACT Drug Names BRIVIACT

Covered Uses All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

BUPRENORPHINE
BUPRENORPHINE HCL

All FDA-approved indications not otherwise excluded from Part D.

1) The drug is being prescribed for the treatment of opioid dependence AND 2) If the patient is pregnant or breastfeeding and being prescribed buprenorphine for induction therapy and/or subsequent maintenance therapy for opioid dependence treatment OR 3) If buprenorphine is being prescribed for induction therapy for transition from opioid use to opioid dependence treatment OR 4) If buprenorphine is being prescribed for maintenance therapy for opioid dependence treatment in a patient who is intolerant to naloxone

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Induction 3 months, Maintenance Plan Year, Pregnancy/Breastfeeding Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

Required Medical Information

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

**BUPRENORPHINE-NALOXONE** 

BUNAVAIL, BUPRENORPHINE HCL/NALOXON, SUBOXONE, ZUBSOLV

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

Coverage Duration

Other Criteria

CABOMETYX

**CABOMETYX** 

All FDA-approved indications not otherwise excluded from Part D.

The disease expresses clear cell histology and is advanced or metastatic.

Plan year

Prior Authorization Group CALQUENCE
Drug Names CALQUENCE

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** For mantle cell lymphoma: Patient has received at least one prior therapy.

Age Restrictions

Prescriber Restrictions

Coverage Duration
Other Criteria

Plan year

Prior Authorization GroupCAPRELSADrug NamesCAPRELSA

Covered Uses All FDA-approved indications not otherwise excluded from Part D, non-small lung

cancer, papillary, follicular, or Hurthle cell thyroid cancer

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

For lung cancer, the disease expresses an RET gene rearrangement

Plan Year

**CARBAGLU** 

**Prior Authorization Group** 

**Drug Names** CARBAGLU

Covered Uses All FDA-approved indications not otherwise excluded from Part D, methylmalonic

acidemia, propionic acidemia.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

Coverage Duration

**Other Criteria** 

Diagnosis of NAGS deficiency was confirmed by enzymatic or genetic testing.

Plan Year

**Prior Authorization Group** CAYSTON **Drug Names** CAYSTON

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** The diagnosis of cystic fibrosis is confirmed by appropriate diagnostic or genetic

testing. Pseudomonas aeruginosa is present in airway cultures.

Age Restrictions

**Prescriber Restrictions Coverage Duration** 

Other Criteria

Plan Year

**CERDELGA** 

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses Exclusion Criteria** 

**Required Medical Information** 

**CERDELGA** 

All FDA-approved indications not otherwise excluded from Part D.

Diagnosis of Gaucher disease was confirmed by an enzyme assay demonstrating a

deficiency of beta-glucocerebrosidase enzyme activity or by genetic testing. The patient's CYP2D6 metabolizer status has been established using an FDA-cleared test. The patient is a CYP2D6 extensive metabolizer, an intermediate metabolizer, or a poor

metabolizer.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

18 years of age or older

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**CEREZYME CEREZYME** 

All FDA-approved indications not otherwise excluded from Part D, type 3 Gaucher

disease.

**Exclusion Criteria** 

**Required Medical Information** 

Diagnosis of Gaucher disease was confirmed by an enzyme assay demonstrating a

deficiency of beta-glucocerebrosidase enzyme activity or by genetic testing.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**CHANTIX Drug Names** 

**Covered Uses** 

CHANTIX, CHANTIX CONTINUING MONTH, CHANTIX STARTING MONTH PA

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

6 months

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

**CINQAIR CINQAIR** 

All FDA-approved indications not otherwise excluded from Part D.

For initial therapy: 1)Patient has baseline eosinophil count of at least 400 cells per

microliter, and 2) Patient has a history of severe asthma attacks (exacerbations) despite current treatment with both of the following medications at optimized doses: a)

inhaled corticosteroid AND b) additional controller (long acting beta2-agonist,

leukotriene modifier, or sustained-release theophylline). For continuation therapy:

Asthma control has improved on Cingair treatment, demonstrated by a reduction in the

frequency and/or severity of symptoms and exacerbations.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

18 years of age or older

Plan Year

Drug Names Covered Uses Exclusion Criteria

All FDA-approved indications not otherwise excluded from Part D.

**Required Medical Information** 

Diagnostic laboratory testing for HAE has been performed (eg, C4, C1 inhibitor functional, and C1 inhibitor antigenic protein levels). For patients with HAE with C1 inhibitor deficiency, C1 inhibitor antigenic protein level and/or C1 inhibitor functional level is below the lower limit of normal as defined by the laboratory performing the test. For patients with HAE with normal C1 inhibitor, other causes of angioedema have been ruled out (eg, drug-induced) and EITHER 1) Patient tested positive for the F12 gene mutation OR 2) Patient has a family history of angioedema and the angioedema was refractory to a trial of antihistamine (eg, levocetirizine) for at least one month.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Plan Year

**CINRYZE** 

**CINRYZE** 

Prior Authorization Group
Drug Names

Covered Uses Exclusion Criteria

Required Medical Information

CLORAZEPATE

**CLORAZEPATE DIPOTASSIUM** 

All FDA-approved indications not otherwise excluded from Part D.

1) For the management of anxiety disorders or for the short-term relief of the symptoms of anxiety, the requested drug is being used with a selective serotonin reuptake inhibitor (SSRI) or serotonin-norepinephrine reuptake inhibitor (SNRI) until the antidepressant becomes effective for the symptoms of anxiety OR the patient has experienced an inadequate treatment response, intolerance or contraindication to a selective serotonin reuptake inhibitor (SSRI) or a serotonin-norepinephrine reuptake inhibitor (SNRI) OR 2) For adjunctive therapy in the management of partial seizures OR 3) Symptomatic relief in acute alcohol withdrawal AND 4) The benefit of therapy with the prescribed medication outweighs the potential risk in a patient 65 years of age or older.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Anxiety Disorders-4 Months, All other Diagnoses-Plan Year

This Prior Authorization requirement only applies to patients 65 years of age or older. (The American Geriatrics Society identifies the use of this medication as potentially inappropriate in older adults, meaning it is best avoided, prescribed at reduced dosage, or used with caution or carefully monitored.)

Prior Authorization GroupCLOZAPINE ODTDrug NamesCLOZAPINE ODT

Covered Uses All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** The patient is unwilling or unable to take tablets or capsules orally or is at high risk for

non-compliance.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

Prior Authorization Group COMETRIQ
Drug Names COMETRIQ

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

Coverage Duration

Other Criteria

Plan Year

Prior Authorization Group COTELLIC Drug Names COTELLIC

Covered Uses All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

**Age Restrictions** 

**Prescriber Restrictions** 

Coverage Duration

Other Criteria

Plan Year

**Drug Names** 

**CRINONE CRINONE** 

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D. progesterone

supplementation during a confirmed pregnancy.

**Exclusion Criteria** 

**Required Medical Information** 

1) Treatment of secondary amenorrhea OR 2) As a progesterone supplementation during a confirmed pregnancy AND 3) Not being prescribed to promote fertility.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

**CYSTAGON CYSTAGON** 

All FDA-approved indications not otherwise excluded from Part D.

Diagnosis of nephropathic cystinosis was confirmed by the presence of increased

cystine concentration in leukocytes or by genetic testing.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**CYSTARAN** 

**CYSTARAN** 

All FDA-approved indications not otherwise excluded from Part D.

**Required Medical Information** Diagnosis of cystinosis was confirmed by the presence of increased cystine

concentration in leukocytes or by DNA testing. The patient has corneal cystine crystal

accumulation.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

Drug Names

DAKLINZA DAKLINZA

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D, chronic hepatitis C

genotype 2 or 4 infection.

**Exclusion Criteria** 

**Required Medical Information** 

Chronic hepatitis C infection confirmed by presence of HCV RNA in serum prior to starting treatment. Planned treatment regimen, genotype, prior treatment history, presence or absence of cirrhosis (compensated or decompensated [Child Turcotte Pugh class B or C]), presence or absence of HIV coinfection, presence or absence of resistance-associated variants where applicable, liver transplantation status if applicable. Coverage conditions and specific durations of approval will be based on current AASLD treatment guidelines.

Age Restrictions

**Prescriber Restrictions** 

Coverage Duration
Other Criteria

Criteria will be applied consistent with current AASLD-IDSA guidance For HCV/HIV coinfection, patient meets criteria for requested regimen.

**Prior Authorization Group** 

**Drug Names** 

Covered Uses

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

**DEFERASIROX** 

EXJADE, JADENU, JADENU SPRINKLE

All FDA-approved indications not otherwise excluded from Part D.

For chronic iron overload due to blood transfusions: pretreatment serum ferritin level is

greater than 1000 mcg/L.

Plan Year

Drug Names
Covered Uses
Exclusion Criteria
Required Medical Information

DIAZEPAM

DIAZEPAM, DIAZEPAM INTENSOL

All FDA-approved indications not otherwise excluded from Part D.

1) For the management of anxiety disorders or for the short-term relief of the symptoms of anxiety, the requested drug is being used with a selective serotonin reuptake inhibitor (SSRI) or serotonin-norepinephrine reuptake inhibitor (SNRI) until the antidepressant becomes effective for the symptoms of anxiety OR the patient has experienced an inadequate treatment response, intolerance or contraindication to a selective serotonin reuptake inhibitor (SSRI) or a serotonin-norepinephrine reuptake inhibitor (SNRI) OR 2) For symptomatic relief in acute alcohol withdrawal OR 3) For use as an adjunct for the relief of skeletal muscle spasms OR 4) For adjunctive therapy in the treatment of convulsive disorders AND 5) The benefit of therapy with the prescribed medication outweighs the potential risk in a patient 65 years of age or older.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Anxiety Disorders-4 Months, All other Diagnoses-Plan Year

This Prior Authorization requirement only applies to patients 65 years of age or older. (The American Geriatrics Society identifies the use of this medication as potentially inappropriate in older adults, meaning it is best avoided, prescribed at reduced dosage, or used with caution or carefully monitored.)

**Prior Authorization Group** 

Drug Names
Covered Uses
Exclusion Criteria

DICLOFENAC GEL 1% DICLOFENAC SODIUM

All FDA-approved indications not otherwise excluded from Part D.

Required Medical Information

1) The patient has osteoarthritis pain in joints susceptible to topical treatment such as feet, ankles, knees, hands, wrist, and elbow. AND 2) Treatment with the requested drug is necessary due to intolerance or a contraindication to oral nonsteroidal anti-inflammatory (NSAID) drugs.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Plan Year

**Prior Authorization Group** EGRIFTA **Drug Names** EGRIFTA

Covered Uses All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** Use for weight loss.

**Required Medical Information** Patient has a diagnosis of HIV infection and lipodystrophy and is receiving anti-

retroviral therapy. For patients who have received at least 6 months of Egrifta therapy: patient has demonstrated clear clinical improvement from baseline that is supported by

a waist circumference or CT scan.

Age Restrictions

Prescriber Restrictions Infectious disease specialist, endocrinologist

Coverage Duration

Other Criteria

6 months

Prior Authorization Group ELAPRASE
Drug Names ELAPRASE

Covered Uses All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** Diagnosis of mucopolysaccharidosis II was confirmed by an enzyme assay

demonstrating a deficiency of iduronate 2-sulfatase enzyme activity or by genetic

testing.

**Age Restrictions** 

**Prescriber Restrictions** 

Coverage Duration

Other Criteria

Plan Year

Prior Authorization Group ELELYSO

**Drug Names** ELELYSO **Covered Uses** All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** Diagnosis of Gaucher disease was confirmed by an enzyme assay demonstrating a

deficiency of beta-glucocerebrosidase enzyme activity or by genetic testing.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

Drug Names
Covered Uses

Exclusion Criteria

**Required Medical Information** 

EMSAM EMSAM

All FDA-approved indications not otherwise excluded from Part D.

1) Patient experienced an inadequate treatment response, intolerance, or

contraindication to any one of the following antidepressants: bupropion, trazodone, mirtazapine, serotonin norepinephrine reuptake inhibitors (SNRIs (e.g., venlafaxine)), selective serotonin reuptake inhibitors (SSRIs (e.g., citalopram, fluoxetine, fluvoxamine,

paroxetine, sertraline)), tricyclic or tetracyclic antidepressants (e.g., amitriptyline,

nortriptyline) OR 2) Patient is unable to swallow oral formulations.

Age Restrictions

**Prescriber Restrictions** 

Coverage Duration
Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

Covered Uses

**Exclusion Criteria** 

**Required Medical Information** 

ENTYVIO ENTYVIO

18 years of age or older.

All FDA-approved indications not otherwise excluded from Part D.

For moderately to severely active Crohn's disease (CD), patient must have an inadequate response, intolerance, or contraindication to at least one conventional therapy option (e.g., corticosteroids, sulfasalazine, azathioprine, 6-mercaptopurine) OR a tumor necrosis factor (TNF) inhibitor for CD (e.g., adalimumab). For moderately to severely active ulcerative colitis (UC), patient must have an inadequate response, intolerance, or contraindication to at least one conventional therapy option (e.g., oral aminosalicylates, corticosteroids, azathioprine, 6-mercaptopurine) OR a tumor necrosis factor (TNF) inhibitor for UC (e.g., adalimumab).

18 years of age or older

Age Restrictions

Prescriber Restrictions
Coverage Duration

Other Criteria

Plan Year

Prior Authorization GroupEPCLUSADrug NamesEPCLUSA

**Covered Uses**All FDA-approved indications not otherwise excluded from Part D. **Exclusion Criteria** 

Required Medical Information Chronic hepatitis C infection confirmed by presence of HCV RNA in the serum prior to

starting treatment. Planned treatment regimen, genotype, prior treatment history, presence or absence of cirrhosis (compensated or decompensated [CTP class B or C]), presence or absence of HIV coinfection, presence or absence of resistance-associated variants where applicable, liver transplantation status if applicable.

Coverage conditions and specific durations of approval will be based on current AASLD

treatment guidelines.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Criteria will be applied consistent with current AASLD-IDSA guidance.

Prior Authorization Group
Drug Names
Covered Uses

EPO

EPOGEN, PROCRIT

All FDA-approved indications not otherwise excluded from Part D, anemia due to myelodysplastic syndromes (MDS), anemia in congestive heart failure (CHF), anemia in rheumatoid arthritis (RA), anemia due to hepatitis C treatment (ribavirin in combination with either interferon alfa or peginterferon alfa), anemia in primary myelofibrosis, post-polycythemia vera myelofibrosis, and post-essential thrombocythemia myelofibrosis.

Exclusion Criteria
Required Medical Information

Patients receiving chemotherapy with curative intent. Patients with myeloid cancer. For all uses except surgery: Pretreatment (no erythropoietin treatment in previous month) hemoglobin (Hgb) is less than 10 g/dL (less than 9 g/dL for anemia in CHF only). Additional requirements for primary myelofibrosis (MF), post-polycythemia vera MF, post-essential thrombocythemia MF: 1) Patient has symptomatic anemia and 2) For initial therapy, pretreatment serum erythropoietin level is less than 500 mU/mL. For surgery: 1) Patient is scheduled for elective, noncardiac, nonvascular surgery and 2) Pretreatment Hgb is greater than 10 but not more than 13 g/dL.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

## 16 weeks

Coverage under Part D will be denied if coverage is available under Part A or Part B as the medication is prescribed and dispensed or administered for the individual (e.g., used for treatment of anemia for a patient with chronic renal failure who is undergoing dialysis, or furnished from physician's supply incident to a physician service). Coverage includes use in anemia in patients whose religious beliefs forbid blood transfusions. Requirements regarding Hgb values exclude values due to a recent transfusion. For reauthorizations (patient received erythropoietin in previous month): 1) For all uses except surgery, there is an increase in Hgb of at least 1 g/dL after at least 12 weeks of therapy, 2) For anemia in chronic kidney disease, MDS, CHF, RA, HIV, hepatitis C treatment, primary MF, post-polycythemia vera MF, post-essential thrombocythemia MF, or patients whose religious beliefs forbid blood transfusions: current Hgb is less than or equal to 12 g/dL, and 3) For anemia due to myelosuppressive cancer chemotherapy: current Hgb is less than 11 g/dL.

**Prior Authorization Group** ERIVEDGE

**Drug Names** ERIVEDGE

Covered Uses All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

Coverage Duration
Other Criteria

Plan Year

**Prior Authorization Group** 

Drug Names

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

ESBRIET ESBRIET

All FDA-approved indications not otherwise excluded from Part D.

Initial Review Only: The patient does not have a known etiology for interstitial lung disease and meets one of the following: 1) a high-resolution computed tomography (HRCT) study of the chest or surgical lung biopsy reveals the usual interstitial pneumonia (UIP) pattern, or 2) HRCT study of the chest reveals a possible UIP pattern and the diagnosis is supported either by surgical lung biopsy or by a multidisciplinary discussion between at least a radiologist and pulmonologist who are experienced in idiopathic pulmonary fibrosis if surgical lung biopsy has not been conducted. For initial

and continuation: Esbriet will not be used in combination with Ofev.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Drug Names** 

EUCRISA EUCRISA

**Covered Uses** 

EUCKISA

Exclusion Criteria

Required Medical Information

All FDA-approved indications not otherwise excluded from Part D.

1) If the requested drug is being prescribed for use on sensitive skin areas (e.g., face, body skin folds, genital area, armpit, or around the eyes), the patient experienced an inadequate treatment response, intolerance, or contraindication to a topical calcineurin inhibitor OR 2) If the requested drug is being prescribed for use on non-sensitive (or remaining) skin areas, the patient experienced an inadequate treatment response, intolerance, or contraindication to a medium or higher potency topical corticosteroid or a topical calcineurin inhibitor.

Age Restrictions

2 years of age or older.

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

FABRAZYME FABRAZYME

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Diagnosis of Fabry disease was confirmed by an enzyme assay demonstrating a deficiency of alpha-galactosidase enzyme activity or by genetic testing, or the patient is an obligate female carrier with a first degree male relative diagnosed with Fabry disease.

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Plan Year

Other Criteria

**Prior Authorization Group** 

**Drug Names** 

FARYDAK

Drug Humos

**FARYDAK** 

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Plan Year

Other Criteria

**Drug Names Covered Uses**  **FASENRA** 

**FASENRA** 

**Exclusion Criteria** 

**Required Medical Information** 

All FDA-approved indications not otherwise excluded from Part D.

For new start:1) Member has baseline blood eosinophil count of at least 150 cells per microliter. 2) Member has a history of severe asthma despite current treatment with both of the following medications at optimized doses: a) Inhaled corticosteroid and b) Additional controller (long acting beta2-agonist, leukotriene modifier, or sustained release theophylline). For continuation of therapy: Asthma control has improved on Fasenra treatment, as demonstrated by a reduction in the frequency and/or severity of symptoms and exacerbations or a reduction in the daily maintenance oral corticosteroid dose

**Age Restrictions** 

**Prescriber Restrictions Coverage Duration** Other Criteria

12 years of age or older

Plan Year

**Prior Authorization Group** 

**Drug Names Covered Uses Exclusion Criteria Required Medical Information**  FENTANYL PATCH

**FENTANYL** 

All FDA-approved indications not otherwise excluded from Part D.

1) The requested drug is being prescribed for pain associated with cancer, a terminal condition, or pain being managed through hospice or palliative care OR 2) The requested drug is being prescribed for pain severe enough to require daily, around-theclock, long-term treatment in a patient who has been taking an opioid AND 3) The patient can safely take the requested dose based on their history of opioid use AND 4) The patient has been evaluated and will be monitored for the development of opioid

Age Restrictions **Prescriber Restrictions Coverage Duration** 

Other Criteria

Plan Year

use disorder

**Drug Names** 

FERRIPROX FERRIPROX

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

Exclusion Criteria

Exclusion Criteria

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

Covered Uses

**FILGRASTIM** 

GRANIX. NEUPOGEN

All FDA-approved indications not otherwise excluded from Part D, treatment of chemotherapy-induced febrile neutropenia (FN), following chemotherapy for acute lymphocytic leukemia (ALL), leukemic relapse following allogeneic stem cell transplantation, myelodysplastic syndromes (MDS), agranulocytosis, aplastic anemia,

HIV-related neutropenia.

**Exclusion Criteria** 

Required Medical Information

For prophylaxis of myelosuppressive chemotherapy-induced FN patients must meet all of the following: 1) Patient has a non-myeloid cancer, 2) Patient is currently receiving or will be receiving treatment with myelosuppressive anti-cancer therapy. For treatment of myelosuppressive chemotherapy-induced FN patients must meet all of the following: 1) Patient has a non-myeloid cancer, 2) Patient is currently receiving or has received treatment with myelosuppressive anti-cancer therapy. For the treatment of anemia in MDS patients must meet all of the following: 1) Patient has symptomatic anemia, 2) The requested G-CSF product will be used in combination with epoetin or darbepoetin, 3) Patient has MDS with a low or intermediate-1 risk stratification, 4) The serum erythropoietin level is less than, or equal to, 500 mU/ml. For neutropenia in MDS: 1) Member is neutropenic, 2) Patient experiences recurrent or resistant infections.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

6 months

**Prior Authorization Group** FIRAZYR **Drug Names** FIRAZYR

All FDA-approved indications not otherwise excluded from Part D, Angiotensin-converting enzyme inhibitor (ACEI)-induced angioedema.

Exclusion Criteria
Required Medical Information

**Covered Uses** 

Firazyr is being requested for the treatment of acute angioedema attacks. For hereditary angioedema (HAE), 1) Diagnostic laboratory testing for HAE has been performed (eg, C4, C1 inhibitor functional, and C1 inhibitor antigenic protein levels), 2) For patients with HAE with C1 inhibitor deficiency, C1 inhibitor antigenic protein level and/or C1 inhibitor functional level is below the lower limit of normal as defined by the laboratory performing the test and 3) For patients with HAE with normal C1 inhibitor, other causes of angioedema have been ruled out (eg, drug induced) and EITHER a) Patient tested positive for the F12 gene mutation OR b) Patient has a family history of angioedema and the angioedema was refractory to a trial of antihistamine (eg, levocetirizine) for at least one month.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

18 years of age or older

Plan Year

Drug Names
Covered Uses

All FDA-approved indications not otherwise excluded from Part D.

Exclusion Criteria

**Required Medical Information** 

For postmenopausal osteoporosis: patient has ONE of the following (1. or 2.): 1) A history of fragility fractures, OR 2) A pre-treatment T-score of less than or equal to -2.5 or osteopenia with a high pre-treatment FRAX fracture probability and patient has ANY of the following: a) Indicators for higher fracture risk (e.g., advanced age, frailty, glucocorticoid therapy, very low T-scores, or increased fall risk), OR b) Patient has failed prior treatment with or is intolerant to a previous osteoporosis therapy (i.e., oral bisphosphonates or injectable antiresorptive agents). For primary or hypogonadal osteoporosis in men: patient has a) a history of osteoporotic vertebral or hip fracture OR b) a pre-treatment T-score of less than or equal to -2.5 OR c) osteopenia with a high pre-treatment FRAX fracture probability. For glucocorticoid-induced osteoporosis: a) patient has had an oral bisphosphonate trial of at least 1-year duration unless patient has a contraindication or intolerance to an oral bisphosphonate, AND Patient has a) a history of fragility fracture, OR b) a pre-treatment T-score of less than or equal to -2.5, OR c) osteopenia with a high pre-treatment FRAX fracture probability.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

24 months (lifetime)

Patient has high FRAX fracture probability if the 10 year probability is either greater than or equal to 20% for any major osteoporotic fracture or greater than or equal to 3% for hip fracture

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

Required Medical Information

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

FYCOMPA FYCOMPA

**FORTEO** 

**FORTEO** 

... \_\_ .

All FDA-approved indications not otherwise excluded from Part D.

12 years of age or older.

Plan Year

**GATTEX Prior Authorization Group** 

**Drug Names GATTEX** 

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** For initial therapy: Patient was dependent on parenteral support for at least 12 months.

For continuation: Requirement for parenteral support has decreased from baseline

while on Gattex therapy.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Plan Year

**Prior Authorization Group** 

**Drug Names** 

Other Criteria

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

Other Criteria

**GILENYA GILENYA** 

All FDA-approved indications not otherwise excluded from Part D.

Have a relapsing form of multiple sclerosis (MS) (e.g., relapsing-remitting MS,

progressive-relapsing MS, or secondary progressive MS with relapses).

**Prescriber Restrictions Coverage Duration** 

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**GILOTRIF GILOTRIF** 

All FDA-approved indications not otherwise excluded from Part D.

**Required Medical Information** For non-small cell lung cancer (NSCLC), patient meets either of the following: 1)

> Patient has metastatic squamous NSCLC that progressed after platinum-based chemotherapy, OR 2) Patient had EGFR mutation testing and is positive for a known

sensitizing EGFR mutation (e.g., EGFR exon 19 deletion or exon 21 (L858R)

substitution mutation), AND Gilotrif is prescribed for treatment of recurrent or metastatic

disease.

**Age Restrictions** 

Other Criteria

**Prescriber Restrictions** 

**Coverage Duration** 

Plan Year

**Drug Names Covered Uses**  **GLATIRAMER** 

COPAXONE, GLATIRAMER ACETATE, GLATOPA

All FDA-approved indications not otherwise excluded from Part D. first clinical episode

of MS.

**Exclusion Criteria** 

**Required Medical Information** 

Have a relapsing form of multiple sclerosis (MS) (e.g., relapsing-remitting MS,

progressive-relapsing MS, or secondary progressive MS with relapses) OR first clinical

episode of MS.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

**GOCOVRI** 

**GOCOVRI** 

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

**GONADOTROPIN** 

CHORIONIC GONADOTROPIN, NOVAREL, PREGNYL W/DILUENT BENZYL

All FDA-approved indications not otherwise excluded from Part D.

Patient is female.

Plan Year

Prior Authorization Group Drug Names

**GROWTH HORMONE** 

GENOTROPIN, GENOTROPIN MINIQUICK, HUMATROPE, HUMATROPE COMBO PACK, NORDITROPIN FLEXPRO, NUTROPIN AQ NUSPIN 10, NUTROPIN AQ NUSPIN 20, NUTROPIN AQ NUSPIN 5, OMNITROPE, SAIZEN, SAIZEN CLICK.EASY, ZOMACTON

Covered Uses
Exclusion Criteria
Required Medical Information

All medically accepted indications not otherwise excluded from Part D. Pediatric patients with closed epiphyses (except in patients with PWS).

Pediatric GHD: Younger than 2.5 yrs old, when applicable: Pretreatment (pre-tx) height (ht) more than 2 SD below mean and slow growth velocity. Two and a half vrs old or older: a) Pre-tx 1-year ht velocity more than 2 SD below mean OR b) Pre-tx ht more than 2 SD below mean and 1-year ht velocity more than 1 SD below mean. Pediatric GHD: Failed 2 stimulation tests (peak below 10 ng/mL) prior to starting treatment, OR Pituitary/CNS disorder (eg., genetic defects, CNS tumors, congenital structural abnormalities) and pre-tx IGF-1 more than 2 SD below mean. OR Patient is a neonate or was diagnosed with GHD as a neonate. TS: 1) Confirmed by karyotyping AND 2) Pre-treatment height is less than the 5th percentile for age. SGA:1) Birth weight (wt) below 2500g at gestational age (GA) more than 37 weeks OR birth wt or length below 3rd percentile for GA or at least 2 SD below mean for GA, AND 2) Did not manifest catch-up growth by age 2. Adult GHD: 1) Failed 2 stimulation tests (peak below 5 ng/mL) prior to starting tx, OR 2) Structural abnormality of the hypothalamus/pituitary AND 3 or more pituitary hormone deficiencies, OR 3) Childhood-onset GHD with congenital (genetic or structural) abnormality of the hypothalamus/pituitary/CNS, OR 4) Low pre-tx IGF-1 and failed 1 stimulation test (peak below 5 ng/mL) prior to starting tx. SGA: 2 years of age or older

Age Restrictions
Prescriber Restrictions

Endocrinologist, Pediatric Endocrinologist, Pediatric nephrologist, Infectious disease specialist, Gastroenterologist/Nutritional support specialist, geneticist.

Coverage Duration
Other Criteria

Plan Year

**Drug Names** 

HAEGARDA **Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

All FDA-approved indications not otherwise excluded from Part D.

This medication is being used for the prevention of acute angioedema attacks. Patient has hereditary angioedema (HAE) with C1 inhibitor deficiency confirmed by laboratory testing OR patient has hereditary angioedema with normal C1 inhibitor confirmed by laboratory testing. For patients with HAE with normal C1 inhibitor, EITHER 1) Patient tested positive for the F12 gene mutation OR 2) Patient has a family history of angioedema and the angioedema was refractory to a trial of antihistamine for at least one month

Age Restrictions **Prescriber Restrictions Coverage Duration** 

Other Criteria

Plan Year

HAEGARDA

**Prior Authorization Group** 

**Drug Names Covered Uses** 

**Exclusion Criteria** 

Required Medical Information

**HARVONI HARVONI** 

All FDA-approved indications not otherwise excluded from Part D.

Chronic hepatitis C infection confirmed by presence of HCV RNA in the serum prior to starting treatment. Planned treatment regimen, genotype, prior treatment history, presence or absence of cirrhosis (compensated or decompensated [Child Turcotte Pugh class B or C]), presence or absence of HIV coinfection, presence or absence of resistance-associated variants where applicable, liver transplantation status if applicable. Coverage conditions and specific durations of approval will be based on current AASLD treatment guidelines.

Age Restrictions **Prescriber Restrictions** 

**Coverage Duration** 

Criteria applied consistent with current AASLD-IDSA guidance.Reminder for 8wk option

if appropriate.

Other Criteria Harvoni will not be used with other drugs containing sofosbuvir, including Sovaldi.

Drug Names

HERCEPTIN HERCEPTIN

Covered Uses All FDA-approved indications not otherwise excluded from Part D, neoadjuvant

treatment for HER2-positive breast cancer, recurrent HER2-positive breast cancer, leptomeningeal metastases from HER2-positive breast cancer, HER2-positive

esophageal and esophagogastric junction cancer.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Neoadjuvant therapy for breast cancer: 6 months. Other: Plan Year.

Coverage under Part D will be denied if coverage is available under Part A or Part B as

the medication is prescribed and dispensed or administered for the individual.

**Prior Authorization Group** 

Drug Names

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

**HETLIOZ** 

HETLIOZ
All FDA-approved indications not otherwise excluded from Part D.

For initial therapy and continuation of Hetlioz therapy: 1) diagnosis of Non-24 Hour Sleep-Wake Disorder, and, 2) diagnosis of total blindness in both eyes (e.g.,

nonfunctioning retinas), and 3) unable to perceive light in both eyes. For patients currently on Hetlioz therapy, must meet at least one of the following: 1) increased total

nighttime sleep or 2) decreased daytime nap duration.

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Initiation: 3 Months, Renewal: Plan Year

Prior Authorization Group Drug Names

HIGH RISK MEDICATION

ALORA, CYPROHEPTADINE HCL, DIGITEK, DIGOX, DIGOXIN, DISOPYRAMIDE PHOSPHATE, ESTRADIOL, FYAVOLV, GUANFACINE ER, JINTELI, LANOXIN,

MEGESTROL ACETATE, MENOSTAR, MINIVELLE, NORETHINDRONE ACETATE/ETH, NORPACE CR, SCOPOLAMINE, TRANSDERM-SCOP

All FDA-approved indications not otherwise excluded from Part D.

Covered Uses

Exclusion Criteria

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

This Prior Authorization requirement only applies to patients 65 years of age or older. (The American Geriatrics Society identifies the use of this medication as potentially inappropriate in older adults, meaning it is best avoided, prescribed at reduced dosage, or used with caution or carefully monitored.) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient.

**Prior Authorization Group** 

**Drug Names** 

Covered Uses
Exclusion Criteria

**Required Medical Information** 

Age Restrictions

Prescriber Restrictions
Coverage Duration

Other Criteria

HRM-ANTICONVULSANTS

PHENOBARBITAL, PHENOBARBITAL SODIUM

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

This Prior Authorization requirement only applies to patients 65 years of age or older. (The American Geriatrics Society identifies the use of this medication as potentially inappropriate in older adults, meaning it is best avoided, prescribed at reduced dosage, or used with caution or carefully monitored.) 1) Two non-HRM alternative drugs carbamazepine, lamotrigine, levetiracetam, topiramate, or valproic acid have not been tried. AND 2) The patient has a contraindication to two non-HRM alternative drugs carbamazepine, lamotrigine, levetiracetam, topiramate, or valproic acid AND 3) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient OR 4) Two non-HRM alternative drugs carbamazepine, lamotrigine, levetiracetam, topiramate, or valproic acid have been tried. AND 5) The patient experienced an inadequate treatment response OR intolerance to two non-HRM alternative drugs carbamazepine, lamotrigine, levetiracetam, topiramate, or valproic acid AND 6) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient.

**Drug Names** 

HRM-ANTIDEPRESSANTS TCA

AMITRIPTYLINE HCL, DOXEPIN HCL, IMIPRAMINE HCL, IMIPRAMINE PAMOATE,

TRIMIPRAMINE MALEATE

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D, Neuropathic pain for amitriptyline or imipramine.

Exclusion Criteria
Required Medical Information
Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Plan Year

This Prior Authorization requirement only applies to patients 65 years of age or older. (The American Geriatrics Society identifies the use of this medication as potentially inappropriate in older adults, meaning it is best avoided, prescribed at reduced dosage. or used with caution or carefully monitored.) Depression: 1) Two non-HRM alternative drugs SSRIs (citalopram, escitalopram, fluoxetine, or sertraline), SNRIs (duloxetine, venlafaxine, or venlafaxine ER), bupropion, mirtazapine, or trazodone have been tried. AND 2) The patient experienced an inadequate treatment response OR intolerance to two non-HRM alternative drugs SSRIs (citalogram, escitalogram, fluoxetine, or sertraline), SNRIs (duloxetine, venlafaxine, or venlafaxine ER), bupropion, mirtazapine, or trazodone AND 3) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient. Neuropathic pain for amitriptyline or imipramine: 1) Two non-HRM alternative drugs duloxetine, gabapentin, pregabalin, or lidocaine patch have been tried. AND 2) The patient experienced an inadequate treatment response OR intolerance to two non-HRM alternative drugs duloxetine, gabapentin, pregabalin, or lidocaine patch AND 3) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient.

**Drug Names** 

Covered Uses
Exclusion Criteria

Required Medical Information

Age Restrictions

**Prescriber Restrictions** 

Coverage Duration

**Other Criteria** 

HRM-ANTIPARKINSON

BENZTROPINE MESYLATE, TRIHEXYPHENIDYL HCL

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

This Prior Authorization requirement only applies to patients 65 years of age or older. (The American Geriatrics Society identifies the use of this medication as potentially inappropriate in older adults, meaning it is best avoided, prescribed at reduced dosage, or used with caution or carefully monitored.) EPS: 1) One non-HRM alternative drug amantadine has not been tried. AND 2) The patient has a contraindication to one non-HRM alternative drug amantadine AND 3) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient. OR 4) One non-HRM alternative drug amantadine has been tried. AND 5) The patient experienced an inadequate treatment response OR intolerance to one non-HRM alternative drug amantadine AND 6) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient. Parkinson's: 1) Two non-HRM drugs amantadine, carbidopa/levodopa, pramipexole, or ropinirole have been tried. AND 2) The patient experienced an inadequate treatment response OR intolerance to two non-HRM drugs amantadine, carbidopa/levodopa, pramipexole, or ropinirole AND 3) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient.

**Prior Authorization Group** 

Drug Names
Covered Uses

Exclusion Criteria

**Required Medical Information** 

Age Restrictions

Prescriber Restrictions

Coverage Duration
Other Criteria

HRM-ANTIPSYCHOTICS
THIORIDAZINE HCL

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

This Prior Authorization requirement only applies to patients 65 years of age or older. (The American Geriatrics Society identifies the use of this medication as potentially inappropriate in older adults, meaning it is best avoided, prescribed at reduced dosage, or used with caution or carefully monitored.) 1) Two non-HRM alternative drugs aripiprazole, asenapine, iloperidone, lurasidone, quetiapine, risperidone, or ziprasidone have been tried. AND 2) The patient experienced an inadequate treatment response OR intolerance to two non-HRM alternative drugs aripiprazole, asenapine, iloperidone, lurasidone, quetiapine, risperidone, or ziprasidone. AND 3) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient.

Drug Names

Covered Uses

Exclusion Criteria

Required Medical Information

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

HRM-CLOMIPRAMINE CLOMIPRAMINE HCL

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

This Prior Authorization requirement only applies to patients 65 years of age or older. (The American Geriatrics Society identifies the use of this medication as potentially inappropriate in older adults, meaning it is best avoided, prescribed at reduced dosage, or used with caution or carefully monitored.) 1) Two non-HRM alternative drugs escitalopram, fluoxetine, fluvoxamine, sertraline, venlafaxine or venlafaxine ER have been tried. AND 2) The patient experienced an inadequate treatment response OR intolerance to two non-HRM alternative drugs escitalopram, fluoxetine, fluvoxamine, sertraline, venlafaxine or venlafaxine ER AND 3) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient.

Prior Authorization Group
Drug Names
Covered Uses
Exclusion Criteria
Required Medical Information
Age Restrictions
Prescriber Restrictions
Coverage Duration

Other Criteria

HRM-HYDROXYZINE
HYDROXYZINE HCL, HYDROXYZINE PAMOATE
All FDA-approved indications not otherwise excluded from Part D.

Plan Year

This Prior Authorization requirement only applies to patients 65 years of age or older. (The American Geriatrics Society identifies the use of this medication as potentially inappropriate in older adults, meaning it is best avoided, prescribed at reduced dosage. or used with caution or carefully monitored.) For pruritus 1) A non-HRM alternative drug levocetirizine has not been tried. AND 2) The patient has a contraindication to a non-HRM alternative drug levocetirizine AND 3) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient. OR 4) A non-HRM alternative drug levocetirizine has been tried. AND 5) The patient experienced an inadequate treatment response OR intolerance to a non-HRM alternative drug levocetirizine AND 6) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient. For anxiety 1) Two non-HRM alternative drugs buspirone, duloxetine, escitalopram, sertraline, or venlafaxine ER have been tried. AND 2) The patient experienced an inadequate treatment response OR intolerance to two non-HRM alternative drugs buspirone, duloxetine, escitalopram, sertraline, or venlafaxine ER AND 3) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient.

Drug Names

Covered Uses
Exclusion Criteria

**Required Medical Information** 

Age Restrictions

Prescriber Restrictions
Coverage Duration

Other Criteria

HRM-HYDROXYZINE INJ HYDROXYZINE HCL

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

This Prior Authorization requirement only applies to patients 65 years of age or older. (The American Geriatrics Society identifies the use of this medication as potentially inappropriate in older adults, meaning it is best avoided, prescribed at reduced dosage. or used with caution or carefully monitored.) Alcohol Withdrawal Syndrome:1) One non-HRM alternative drug clorazepate or lorazepam have not been tried AND 2) The patient has a contraindication to one non-HRM alternative drug clorazepate or lorazepam AND 3) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient OR 4) One non-HRM alternative drug clorazepate or lorazepam have been tried AND 5) The patient experienced an inadequate treatment response OR intolerance to one non-HRM alternative drug clorazepate or lorazepam AND 6) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient Anxiety: 1) Two non-HRM alternative drugs buspirone, duloxetine, escitalopram. sertraline or venlafaxine ER have been tried AND 2) The patient experienced an inadequate treatment response OR intolerance to two non-HRM alternative drugs buspirone, duloxetine, escitalopram, sertraline or venlafaxine ER AND 3) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient OR 4) If being requested for nausea/vomiting, prescriber must acknowledge that medication benefits outweigh potential risks for this patient.

**Prior Authorization Group** 

Drug Names
Covered Uses

**Exclusion Criteria** 

**Required Medical Information** 

**Age Restrictions** 

Prescriber Restrictions
Coverage Duration
Other Criteria

HRM-HYPNOTICS ZOLPIDEM TARTRATE

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

This Prior Authorization requirement only applies to patients 65 years of age or older. (The American Geriatrics Society identifies the use of this medication as potentially inappropriate in older adults, meaning it is best avoided, prescribed at reduced dosage, or used with caution or carefully monitored.) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient. APPLIES TO GREATER THAN CUMULATIVE 90 DAYS OF THERAPY PER YEAR.

Prior Authorization Group Drug Names

HRM-NITROFURANTOIN
NITROFURANTOIN MACROCRYST, NITROFURANTOIN
MONOHYDRAT

Covered Uses
Exclusion Criteria
Required Medical Information
Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

This Prior Authorization requirement only applies to patients 65 years of age or older. (The American Geriatrics Society identifies the use of this medication as potentially inappropriate in older adults, meaning it is best avoided, prescribed at reduced dosage, or used with caution or carefully monitored.) 1) Two non-HRM alternative drugs cephalexin, ciprofloxacin, levofloxacin, sulfamethoxazole/trimethoprim, or trimethoprim have not been tried. AND 2) The patient has a contraindication to two non-HRM alternative drugs cephalexin, ciprofloxacin, levofloxacin,

sulfamethoxazole/trimethoprim, or trimethoprim AND 3) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient. OR 4) Two non-HRM alternative drugs cephalexin, ciprofloxacin, levofloxacin,

sulfamethoxazole/trimethoprim, or trimethoprim have been tried. AND 5) The patient experienced an inadequate treatment response OR intolerance to two non-HRM alternative drugs cephalexin, ciprofloxacin, levofloxacin,

sulfamethoxazole/trimethoprim, or trimethoprim) AND 6) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient. APPLIES TO GREATER THAN CUMULATIVE 90 DAYS OF THERAPY PER YEAR.

**Drug Names** 

HRM-PROMETHAZINE

PHENADOZ, PHENERGAN, PROMETHAZINE HCL, PROMETHAZINE HCL PLAIN,

**PROMETHEGAN** 

Covered Uses
Exclusion Criteria

**Required Medical Information** 

Age Restrictions

Prescriber Restrictions
Coverage Duration

Other Criteria

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

This Prior Authorization requirement only applies to patients 65 years of age or older. (The American Geriatrics Society identifies the use of this medication as potentially inappropriate in older adults, meaning it is best avoided, prescribed at reduced dosage, or used with caution or carefully monitored.) Rhinitis: 1) One non-HRM alternative drug levocetirizine, azelastine nasal, fluticasone nasal, or flunisolide nasal has been tried. AND 2) The patient experienced an inadequate treatment response OR intolerance to one non-HRM alternative drug levocetirizine, azelastine nasal, fluticasone nasal, or flunisolide nasal AND 3) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient Urticaria: 1) One non-HRM alternative drug levocetirizine has not been tried. AND 2) The patient has a contraindication to one non-HRM alternative drug levocetirizine AND 3) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient. OR 4) One non-HRM alternative drug levocetirizine has been tried. AND 5) The patient experienced an inadequate treatment response OR intolerance to one non-HRM alternative drug levocetirizine AND 6) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient OR 7) The drug is being requested for antiemetic therapy in postoperative patients or motion sickness AND 8) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient.

**Prior Authorization Group** 

Drug Names
Covered Uses

Exclusion Criteria

**Required Medical Information** 

Age Restrictions

Prescriber Restrictions
Coverage Duration

Other Criteria

HRM-SKELETAL MUSCLE RELAXANTS

CYCLOBENZAPRINE HCL

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

This Prior Authorization requirement only applies to patients 65 years of age or older. (The American Geriatrics Society identifies the use of this medication as potentially inappropriate in older adults, meaning it is best avoided, prescribed at reduced dosage, or used with caution or carefully monitored.) Prescriber must acknowledge that medication benefits outweigh potential risks for this patient.

Prior Authorization Group Drug Names

**Covered Uses** 

Exclusion Criteria
Required Medical Information

HUMIRA

HUMIRA, HUMIRA PEDIATRIC CROHNS D, HUMIRA PEN, HUMIRA PEN-CROHNS DISEASE, HUMIRA PEN-PSORIASIS STAR

All FDA-approved indications not otherwise excluded from Part D, axial spondyloarthritis.

For moderately to severely active rheumatoid arthritis (new starts only): Patient meets at least one of the following: 1) Inadequate response, intolerance or contraindication to methotrexate (MTX), OR 2) Inadequate response or intolerance to a prior biologic disease-modifying antirheumatic drug (DMARD) or a targeted synthetic DMARD (e.g., tofacitinib). For moderately to severely active polyarticular juvenile idiopathic arthritis (new starts only): Patient meets ANY of the following: 1) Inadequate response. intolerance or contraindication to MTX, OR 2) Inadequate response or intolerance to a prior biologic DMARD. For active ankylosing spondylitis and axial spondyloarthritis (new starts only): Inadequate response to a non-steroidal anti-inflammatory drug (NSAID) trial at maximum recommended or tolerated dose OR intolerance or contraindication to NSAIDs. For moderate to severe chronic plaque psoriasis (new starts only): 1) At least 5% of body surface area (BSA) is affected OR crucial body areas (e.g., feet, hands, face, neck, groin, intertriginous areas) are affected at the time of diagnosis, AND 2) Patient meets any of the following: a) Inadequate response or intolerance to either phototherapy (e.g., UVB, PUVA) or pharmacologic treatment with methotrexate, cyclosporine, or acitretin, b) Pharmacologic treatment with methotrexate, cyclosporine, or acitretin is contraindicated, c) Patient has severe psoriasis that warrants a biologic DMARD as first-line therapy. For moderately to severely active Crohn's disease (new starts only): 1) Inadequate response to at least one conventional therapy (e.g., corticosteroids, sulfasalazine, azathioprine, mesalamine), OR 2) Intolerance or contraindication to conventional therapy. For moderately to severely active ulcerative colitis (new starts only): 1) Inadequate response to at least one immunosuppressant therapy (e.g., corticosteroids, azathioprine, mercaptopurine), OR 2) Intolerance or contraindication to immunosuppressant therapy.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Plan Year

**Drug Names** 

HYPNOTIC BENZODIAZEPINES **TEMAZEPAM** 

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

This Prior Authorization requirement only applies to patients 65 years of age or older. (The American Geriatrics Society identifies the use of this medication as potentially inappropriate in older adults, meaning it is best avoided, prescribed at reduced dosage. or used with caution or carefully monitored.) 1) One non-HRM alternative drug Silenor (3mg or 6mg) or trazodone has not been tried. AND 2) The patient has a contraindication to two non-HRM alternative drugs Silenor (3mg or 6mg) and trazodone. AND 3) Prescriber must acknowledge that medication benefits outweigh potential risk in a patient 65 years of age or older. OR 4) One non-HRM alternative drug Silenor (3mg or 6mg) or trazodone has been tried. AND 5) The patient experienced an inadequate treatment response OR intolerance to one non-HRM alternative drug Silenor (3mg or 6mg) or trazodone. AND 6) Prescriber must acknowledge that medication benefits outweigh potential risk in a patient 65 years of age or older. APPLIES TO GREATER THAN CUMULATIVE 90 DAYS OF THERAPY PER YEAR.

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**IBRANCE IBRANCE** 

All FDA-approved indications not otherwise excluded from Part D, well-

differentiated/dedifferentiated retroperitoneal liposarcoma.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

Drug Names

ICLUSIG ICLUSIG

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

For chronic myeloid leukemia (CML) or Philadelphia chromosome positive acute lymphoblastic leukemia (Ph+ ALL), diagnosis was confirmed by detection of the

Philadelphia chromosome or BCR-ABL gene.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

18 years of age or older

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

Required Medical Information

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

IDHIFA

**IDHIFA** 

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

**Prior Authorization Group** 

**Drug Names** 

IMATINIB

Covered Uses

IMATINIB MESYLATE

All FDA-approved indications not otherwise excluded from Part D, Philadelphia chromosome positive (Ph+) lymphoblastic lymphoma, desmoid tumors, pigmented villonodular synovitis/tenosynovial giant cell tumor (PVNS/TGCT), chordoma, and

melanoma.

**Exclusion Criteria** 

**Required Medical Information** 

For chronic myeloid leukemia (CML) or Philadelphia chromosome positive acute lymphoblastic leukemia (Ph+ ALL)/lymphoblastic lymphoma, diagnosis was confirmed

by detection of the Philadelphia chromosome or BCR-ABL gene. For CML, patient did

not fail (excluding failure due to intolerance) prior therapy with a tyrosine kinase inhibitor (eg, dasatinib, nilotinib, bosutinib, ponatinib). For melanoma, c-Kit mutation is

positive.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

Drug Names

IMBRUVICA IMBRUVICA

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D, lymphoplasmacytic

lymphoma.

**Exclusion Criteria** 

Required Medical Information

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

INCRELEX INCRELEX

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

Closed epiphyses.

**Required Medical Information** 

Must meet all of the following prior to beginning Increlex therapy (new starts only): 1) height 3 or more standard deviations below the mean for children of the same age and gender AND 2) basal IGF-1 level 3 or more standard deviations below the mean for children of the same age and gender AND 3) stimulation test showing a normal or elevated growth hormone level. For renewal, patient is experiencing improvement AND

the current IGF-1 level is normal for age and gender.

Age Restrictions

Prescriber Restrictions Coverage Duration Other Criteria Endocrinologist Plan year

**Prior Authorization Group** 

**Drug Names** 

INLYTA INLYTA

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D, papillary, Hurthle

cell, or follicular thyroid carcinoma.

**Exclusion Criteria** 

**Required Medical Information** 

For renal cell carcinoma: the disease is relapsed or unresectable. For thyroid

carcinoma: 1) the disease has papillary, Hurthle cell, or follicular histology, and 2) the

disease is unresectable or metastatic

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Drug Names** 

**IRESSA IRESSA** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

For non-small cell lung cancer (NSCLC): 1) Patient had EGFR mutation testing and is positive for a known sensitizing EGFR mutation (e.g., EGFR exon 19 deletion OR exon

21 (L858R) substitution mutation), AND 2) Iressa is prescribed for treatment of

All FDA-approved indications not otherwise excluded from Part D.

recurrent or metastatic disease.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names Covered Uses**  ISOTRETINOIN

AMNESTEEM, CLARAVIS, MYORISAN, ZENATANE

All FDA-approved indications not otherwise excluded from Part D, refractory acne. severe refractory rosacea, neuroblastoma, cutaneous T-cell lymphoma (CTCL) (e.g., mycosis fungoides, Sezary syndrome), reduction of the development of skin cancer (squamous cell cancers) in high risk patients, transient acantholytic dermatosis (Grover Disease), keratosis follicularis (Darier Disease), lamellar ichthyosis, pityriasis rubra

pilaris.

Exclusion Criteria

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**ITRACONAZOLE ITRACONAZOLE** 

All FDA-approved indications not otherwise excluded from Part D, Coccidioidomycosis,

Cryptococcosis, Microsporidiosis, Penicilliosis, Sporotrichosis, Pitvriasis

versicolor/Tinea versicolor, Tinea corporis/Tinea cruris, Tinea manuum/Tinea pedis.

**Exclusion Criteria** 

**Required Medical Information** 

If for the treatment of onychomycosis due to tinea, the diagnosis has been confirmed

by a fungal diagnostic test.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

6 months

Prior Authorization Group Drug Names

IVIG

BIVIGAM, CARIMUNE NANOFILTERED, FLEBOGAMMA DIF, GAMMAGARD LIQUID, GAMMAGARD S/D IGA LESS TH, GAMMAKED, GAMMAPLEX, GAMUNEX-C,

OCTAGAM, PRIVIGEN

**Covered Uses**All FDA-approved indications not otherwise excluded from Part D, primary

immunodeficiency, chronic inflammatory demyelinating polyneuropathy, multifocal motor neuropathy, dermatomyositis, polymyositis, Guillain-Barre syndrome (GBS), myasthenia gravis, Lambert-Eaton myasthenic syndrome, Kawasaki syndrome, idiopathic thrombocytopenic purpura, pure red cell aplasia (PRCA), fetal/neonatal alloimmune thrombocytopenia, Stiff-person syndrome, and prophylaxis of bacterial infections in B-cell chronic lymphocytic leukemia (CLL), bone marrow/hematopoietic stem cell transplant (BMT/HSCT) recipients, and pediatric HIV infection.

Exclusion Criteria
Required Medical Information

For CLL: serum IgG less than 500 mg/dL OR a history of recurrent bacterial infections. For BMT/HSCT: IVIG is requested within the first 100 days post-transplant OR serum IgG less than 400 mg/dL. For pediatric HIV infection: 1) Serum IgG less than 400 mg/dL, OR 2) History of recurrent bacterial infections, patient is not able to take combination antiretroviral therapy, and antibiotic prophylaxis was not effective. For dermatomyositis and polymyositis: at least one standard first-line treatment (corticosteroids or immunosuppressants) has been tried but was unsuccessful or not tolerated OR patient is unable to receive standard therapy because of a contraindication or other clinical reason. For GBS: physical mobility must be severely affected such that the patient requires an aid to walk AND IVIG therapy must be initiated within 2 weeks of symptom onset. For myasthenia gravis: IVIG is requested for worsening weakness, acute exacerbation or use in preparation for surgery. PRCA is secondary to parvovirus B19 infection. For Stiff-person syndrome: inadequate response or intolerance to at least one first-line therapy such as a benzodiazepine (eg, diazepam) and/or baclofen unless contraindicated.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

For pediatric HIV infection: age 12 years or younger.

Plan Year

Coverage under Part D will be denied if coverage is available under Part A or Part B as the medication is prescribed and dispensed or administered for the individual.

Prior Authorization GroupJAKAFIDrug NamesJAKAFI

Covered Uses All FDA-approved indications not otherwise excluded from Part D, low-risk,

intermediate-risk, accelerated phase, or blast phase myelofibrosis.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

Prior Authorization Group
Drug Names
Covered Uses
Exclusion Criteria
Required Medical Information

JUXTAPID JUXTAPID

All FDA-approved indications not otherwise excluded from Part D.

For initiation of therapy: 1) Patient has a diagnosis of homozygous familial hypercholesterolemia (HoFH) confirmed by genetic analysis or clinical criteria (see Other Criteria), AND 2) Prior to initiation of treatment with Juxtapid, patient is/was receiving a combination lipid-lowering regimen consisting of at least 2 of the following treatment options: high-intensity statin (eg, atorvastatin, rosuvastatin), fibrate (eg, fenofibrate, fenofibric acid, gemfibrozil), bile acid sequestrant (eg, cholestyramine, colesevelam, colestipol), ezetimibe, or niacin, at maximally tolerated doses or at the maximum doses approved by the FDA, AND 3) Prior to initiation of treatment with Juxtapid, patient is/was experiencing an inadequate response to such combination regimen as demonstrated by treated LDL-C greater than 160 mg/dL or 100 mg/dl if the patient has coronary heart disease or other atherosclerotic cardiovascular disease. OR diabetes, OR a family history of very early coronary heart disease (less than 45 years of age in men and less than 55 years of age in women), OR current smoker, OR two or more coronary heart disease risk factors, OR lipoprotein(a) levels of 50 mg/dl or greater. For renewal of therapy: 1) Patient meets all initial criteria AND 2) Has responded to therapy as demonstrated by a reduction in LDL-C.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Lipid specialist, cardiometabolic specialist, cardiologist, or endocrinologist Plan Year

Diagnosis of HoFH must be confirmed by one of the following: 1) Genetic diagnosis: Mutations in both alleles at LDL receptor, ApoB, PCSK9 or LDL receptor adaptor protein/ARH gene locus, or 2) Clinical diagnosis: Untreated LDL-C greater than 500 mg/dL or unknown untreated LDL-C with treated LDL-C greater than 300 mg/dL plus one of the following: a) Tendon or cutaneous xanthomas at age 10 or younger, or b) Diagnosis of definite FH by genetic analysis, Simon-Broome Diagnostic Criteria or Dutch Lipid Clinic Network Criteria in both parents, or c) Evidence of FH in both parents with a history including any of the following: Total cholesterol greater than or equal to 310 mg/dL, premature ASCVD [before 55 years in men and 60 years in women]. tendon xanthoma, or sudden premature cardiac death. Diagnosis of definite FH must be confirmed by one of the following: 1) Genetic diagnosis: An LDL-receptor mutation, familial defective apo B-100, or a PCSK9 gain-of-function mutation, or 2) Simon-Broome Diagnostic Criteria for definite FH: Total cholesterol greater than 290 mg/dL or LDL-C greater than 190 mg/dL, plus tendon xanthoma in patient, first-degree (parent, sibling or child) or second-degree relative (grandparent, uncle or aunt), or 3) Dutch Lipid Clinic Network Criteria for definite FH: Total score greater than 8 points.

**KALYDECO Prior Authorization Group Drug Names KALYDECO** 

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** The patient has a diagnosis of cystic fibrosis. The patient has one mutation in the cystic

fibrosis transmembrane conductance regulator (CFTR) gene that is responsive to

ivacaftor potentiation based on clinical and/or in vitro assay data. Granules: 2 years of age or older, Tablets: 6 years of age or older

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria Kalydeco will not be used in combination with Orkambi.

Plan Year

**Prior Authorization Group** 

**Drug Names Covered Uses** 

**Exclusion Criteria** 

Required Medical Information

Age Restrictions **Prescriber Restrictions Coverage Duration** 

Other Criteria

**KETOCONAZOLE** 

**KETOCONAZOLE** 

All FDA-approved indications not otherwise excluded from Part D, Cushing's syndrome.

Acute or chronic liver disease. Current use with dofetilide, guinidine, pimozide, cisapride, methadone, disopyramide, dronedarone, ranolazine, ergot alkaloids,

alprazolam or simvastatin.

1) Patient has one of the following diagnoses: blastomycosis, coccidioidomycosis,

histoplasmosis, chromomycosis, or paracoccidioidomycosis, OR 2) The requested drug is being prescribed for a patient with Cushing's syndrome who cannot tolerate surgery

or surgery has not been curative.

6 months

**Drug Names** 

**KEVEYIS KEVEYIS** 

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Severe pulmonary disease, hepatic insufficiency, concomitant use of high-dose aspirin.

For primary HYPOkalemic periodic paralysis: 1) The diagnosis was supported by

genetic test results. OR 2) patient has a family history of primary hypokalemic periodic

paralysis, OR 3) patient's attacks are associated with hypokalemia AND both

Andersen-Tawil syndrome and thyrotoxic periodic paralysis have been ruled out. For primary HYPERkalemic periodic paralysis: 1) The diagnosis was supported by genetic test results, OR 2) patient has a family history of primary hyperkalemic periodic paralysis, OR 3) patient's attacks are associated with hyperkalemia AND Andersen-

Tawil syndrome has been ruled out.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Initial: 2 months Continuation: 12 months.

Keveyis is used as maintenance therapy to prevent attacks. For continuation of

therapy, patient is demonstrating a response to Keveyis therapy as demonstrated by a

decrease in the number of attacks.

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

**KEYTRUDA** 

**KEYTRUDA** 

All FDA-approved indications not otherwise excluded from Part D.

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**KISQALI** 

Plan Year

Plan year

KISQALI, KISQALI FEMARA 200 DOSE, KISQALI FEMARA 400 DOSE, KISQALI

FEMARA 600 DOSE

**Exclusion Criteria** 

**Required Medical Information** 

**Age Restrictions** 

Prescriber Restrictions

**Coverage Duration** 

Other Criteria

All FDA-approved indications not otherwise excluded from Part D.

Drug Names

KORLYM KORLYM

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

**KUVAN** 

KUVAN

All FDA-approved indications not otherwise excluded from Part D.

For patients who have not yet received a therapeutic trial of Kuvan, the patient's

pretreatment, including before dietary management, phenylalanine level is greater than 6 mg/dL (360 micromol/L). For patients for whom this is the first treatment after a therapeutic trial of Kuvan: a) The patient must have experienced a reduction in blood phenylalanine level from baseline OR b) the patient has demonstrated an improvement

in neuropsychiatric symptoms.

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Initial: 2 months. Continuation of treatment: Plan Year.

Prior Authorization Group
Drug Names
Covered Uses
Exclusion Criteria
Required Medical Information

KYNAMRO KYNAMRO

All FDA-approved indications not otherwise excluded from Part D.

For initiation of therapy: 1) Patient has a diagnosis of homozygous familial hypercholesterolemia (HoFH) confirmed by genetic analysis or clinical criteria (see Other Criteria), AND 2) Prior to initiation of treatment with Kynamro, patient is/was receiving a combination lipid-lowering regimen consisting of at least 2 of the following treatment options: high-intensity statin (eg, atorvastatin, rosuvastatin), fibrate (eg, fenofibrate, fenofibric acid, gemfibrozil), bile acid sequestrant (eg, cholestyramine, colesevelam, colestipol), ezetimibe, or niacin, at maximally tolerated doses or at the maximum doses approved by the FDA, AND 3) Prior to initiation of treatment with Kynamro, patient is/was experiencing an inadequate response to such combination regimen, as demonstrated by treated LDL-C greater than 160 mg/dL or 100 mg/dl if the patient has coronary heart disease or other atherosclerotic cardiovascular disease. OR diabetes, OR a family history of very early coronary heart disease (less than 45 years of age in men and less than 55 years of age in women), OR current smoker, OR two or more coronary heart disease risk factors, OR lipoprotein(a) levels of 50 mg/dl or greater. For renewal of therapy, 1) Patient meets all initial criteria AND 2) Has responded to therapy as demonstrated by a reduction in LDL-C.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Lipid specialist, cardiometabolic specialist, cardiologist, or endocrinologist Plan Year

Diagnosis of HoFH must be confirmed by one of the following: 1) Genetic diagnosis: Mutations in both alleles at LDL receptor, ApoB, PCSK9 or LDL receptor adaptor protein/ARH gene locus, or 2) Clinical diagnosis: Untreated LDL-C greater than 500 mg/dL or unknown untreated LDL-C with treated LDL-C greater than 300 mg/dL plus one of the following: a) Tendon or cutaneous xanthomas at age 10 or younger, or b) Diagnosis of definite FH by genetic analysis, Simon-Broome Diagnostic Criteria or Dutch Lipid Clinic Network Criteria in both parents, or c) Evidence of FH in both parents with a history including any of the following: Total cholesterol greater than or equal to 310 mg/dL, premature ASCVD [before 55 years in men and 60 years in women]. tendon xanthoma, sudden premature cardiac death. Diagnosis of definite FH must be confirmed by one of the following: 1) Genetic diagnosis: An LDL-receptor mutation, familial defective apo B-100, or a PCSK9 gain-of-function mutation, or 2) Simon-Broome Diagnostic Criteria for definite FH: Total cholesterol greater than 290 mg/dL or LDL-C greater than 190 mg/dL, plus tendon xanthoma in patient, first-degree (parent, sibling or child) or second-degree relative (grandparent, uncle or aunt), or 3) Dutch Lipid Clinic Network Criteria for definite FH: Total score greater than 8 points.

**Drug Names** 

LENVIMA

LENVIMA 10 MG DAILY DOSE, LENVIMA 14 MG DAILY DOSE, LENVIMA 18 MG DAILY DOSE, LENVIMA 20 MG DAILY DOSE, LENVIMA 24 MG DAILY DOSE.

LENVIMA 8 MG DAILY DOSE

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria Required Medical Information** 

For differentiated thyroid cancer: 1) histologic subtype is papillary, follicular, or Hurthle cell, AND 2) disease is iodine-refractory. For renal cell carcinoma which meets all of the following: 1) Patient has relapsed or advanced disease, 2) Lenvima will be used in combination with everolimus, 3) For disease that is of predominantly clear cell histology. Lenvima will be used as subsequent therapy for disease that has progressed on prior anti-angiogenic therapy (e.g., bevacizumab, sunitinib, sorafenib).

Age Restrictions

**Prescriber Restrictions Coverage Duration** 

Other Criteria

Plan Year

**LETAIRIS** 

**LETAIRIS** 

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** Pulmonary arterial hypertension (WHO Group 1) was confirmed by right heart

> catheterization. For new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than or equal to 25 mmHg, 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, and 3) pretreatment pulmonary vascular

resistance is greater than 3 Wood units.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

Prior Authorization Group
Drug Names

**Covered Uses** 

LEUKINE LEUKINE

All FDA-approved indications not otherwise excluded from Part D, prevention and treatment of chemotherapy-induced febrile neutropenia (FN), following chemotherapy for acute lymphocytic leukemia (ALL) or acute myeloid leukemia (AML), neutropenia in myelodysplastic syndromes (MDS), agranulocytosis, aplastic anemia, HIV-related neutropenia.

**Exclusion Criteria** 

**Required Medical Information** 

For prophylaxis of myelosuppressive chemotherapy-induced FN the patient must meet all of the following: 1) Patient has a non-myeloid cancer, 2) Patient is currently receiving or will be receiving treatment with myelosuppressive anti-cancer therapy. For treatment of myelosuppressive chemotherapy-induced FN the patient must meet all of the following: 1) Patient has a non-myeloid cancer, 2) Patient is currently receiving or has received treatment with myelosuppressive anti-cancer therapy. For MDS: Patient has neutropenia and experiences recurrent or resistant infections.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

6 months

**Prior Authorization Group** 

**Drug Names Covered Uses** 

LIDOCAINE PATCHES

LIDOCAINE

All FDA-approved indications not otherwise excluded from Part D, pain associated with diabetic neuropathy, pain associated with cancer-related neuropathy (including treatment-related neuropathy [e.g. neuropathy associated with radiation treatment or chemotherapy]).

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Drug Names** 

LONSURF LONSURF

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

For colorectal cancer: The disease is unresectable advanced or metastatic. Patient has

progressed on treatment with EITHER a) FOLFOXIRI (fluorouracil, leucovorin,

oxaliplatin, and irinotecan) regimen OR b) irinotecan- AND oxaliplatin-based regimens.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

LUMIZYME

LUMIZYME

All FDA-approved indications not otherwise excluded from Part D.

Diagnosis of Pompe disease was confirmed by an enzyme assay demonstrating a

deficiency of acid alpha-glucosidase (GAA) enzyme activity or by genetic testing.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

Required Medical Information

LUPANETA

LUPANETA PACK

All FDA-approved indications not otherwise excluded from Part D.

For retreatment patient must have had a recurrence of symptoms (one-time

retreatment course allowed).

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

18 years of age or older

6 months

**Drug Names** 

LUPRON

LEUPROLIDE ACETATE, LUPRON DEPOT (1-MONTH), LUPRON DEPOT (3-

MONTH), LUPRON DEPOT (4-MONTH), LUPRON DEPOT-PED (1-MONTH, LUPRON

DEPOT-PED (3-MONTH

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D, in combination with growth hormone for children with growth failure and advancing puberty (leuprolide acetate only), breast cancer (3.75 mg only), malignant sex cord-stromal tumors (3.75 mg and 11.25 mg), epithelial ovarian cancer/fallopian tube cancer/primary peritoneal cancer (3.75 mg only), preoperative use for uterine leiomyomata (3.75 mg and 11.25

mg).

Exclusion Criteria
Required Medical Information

For central precocious puberty (CPP), patients not currently receiving therapy must meet all of the following criteria: 1) Diagnosis of CPP confirmed by: a) A pubertal response to a gonadotropin releasing hormone (GnRH) agonist test OR a pubertal level of a third generation luteinizing hormone (LH) assay AND b) Assessment of bone age versus chronological age, and 2) The onset of secondary sexual characteristics occurred prior to 8 years of age for female patients OR prior to 9 years of age for male patients. For endometriosis retreatment patient must meet all of the following: 1) Patient has had a recurrence of symptoms, and 2) Patient will be receiving add-back therapy (eg, norethindrone). For uterine fibroids patient must meet one of the following: 1) Diagnosis of anemia (eg, hematocrit less than or equal to 30 percent and/or hemoglobin less than or equal to 10g/dL), OR 2) Lupron Depot will be used in the preoperative setting to facilitate surgery. For epithelial ovarian cancer/fallopian tube cancer/primary peritoneal cancer: Lupron (3.75mg only) will be used as a single agent AND disease is persistent or recurrent. For breast cancer (3.75mg only), patient must be premenopausal with hormone receptor positive disease.

Age Restrictions

CPP: Less than 12 years old if female and less than 13 years old if male. Endometriosis, fibroids, breast cancer, stromal tumors, epithelial ovarian/fallopian tube/primary peritoneal cancer: 18 years of age or older.

Prescriber Restrictions
Coverage Duration

Fibroids: 3 months (mo), max 6 mo total. Endometriosis: 6 mo, max 12 mo total.

Others: Plan Year

Other Criteria

**Prior Authorization Group** LYNPARZA

**Drug Names** LYNPARZA

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** For HER2-negative, recurrent or metastatic breast cancer patient must meet both of the

following criteria: 1) patient has a deleterious or suspected deleterious germline BRCA mutation, and 2) patient has received prior treatment with chemotherapy or endocrine

therapy.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**MAVYRET MAVYRET** 

All FDA-approved indications not otherwise excluded from Part D.

Decompensated cirrhosis/moderate or severe hepatic impairment (Child Turcotte Pugh

class B or C)

Chronic hepatitis C infection confirmed by presence of HCV RNA in the serum prior to Required Medical Information

starting treatment. Planned treatment regimen, genotype, prior treatment history, presence or absence of cirrhosis (compensated or decompensated [CTP class B or C1), presence or absence of HIV coinfection, presence or absence of resistanceassociated variants where applicable, liver transplantation status if applicable.

Coverage conditions and specific durations of approval will be based on current AASLD

treatment guidelines.

Age Restrictions

Prescriber Restrictions

**Coverage Duration** 8-16 weeks per package insert or Criteria will be applied consistent w/ current AASLD-

IDSA guidance

Other Criteria

MEGESTROL **Prior Authorization Group** 

MEGESTROL ACETATE **Drug Names** 

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Drug Names** 

**MEKINIST MEKINIST** 

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D, non-small cell lung

cancer (NSCLC) with BRAF V600E mutation.

**Exclusion Criteria** 

**Required Medical Information** 

For unresectable or metastatic melanoma, the tumor is positive for BRAF V600E or V600K mutation and Mekinist is used as a single agent or in combination with dabrafenib. For NSCLC, tumor is positive for BRAF V600E mutation and Mekinist is used in combination with dabrafenib.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**MEMANTINE** 

MEMANTINE HCL, MEMANTINE HYDROCHLORIDE, NAMENDA XR, NAMENDA XR

TITRATION PACK

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

All FDA-approved indications not otherwise excluded from Part D.

The drug is being prescribed for the treatment of moderate to severe dementia of the

Alzheimer's type. [Note: Common indicators of moderate to severe disease include MMSE scores of less than or equal to 20 and/or when ADLs are significantly impacted.]

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

This edit only applies to patients less than 30 years of age.

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

**MODAFINIL MODAFINIL** 

All FDA-approved indications not otherwise excluded from Part D.

1) Diagnosis is narcolepsy confirmed by sleep lab evaluation OR 2) Diagnosis is

obstructive sleep apnea (OSA) confirmed by polysomnography OR 3) Diagnosis is shift

work disorder (SWD).

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Drug Names** 

**MOZOBIL MOZOBIL** 

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

6 months

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**MYLOTARG MYLOTARG** 

All FDA-approved indications not otherwise excluded from Part D, acute promyelocytic

leukemia (APL).

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**NAGLAZYME** 

**NAGLAZYME** 

All FDA-approved indications not otherwise excluded from Part D.

Diagnosis of mucopolysaccharidosis VI disease was confirmed by an enzyme assay **Required Medical Information** 

demonstrating a deficiency of N-acetylgalactosamine 4-sulfatase (arylsulfatase B)

enzyme activity or by genetic testing.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Drug Names** 

**NATPARA NATPARA** 

**Covered Uses Exclusion Criteria**  All FDA-approved indications not otherwise excluded from Part D. Acute postsurgical hypoparathyroidism (within 6 months of surgery).

Hypoparathyroidism due to calcium-sensing receptor mutations.

**Required Medical Information** 

Natpara is prescribed to control hypocalcemia associated with hypoparathyroidism. Total serum calcium levels are inadequately controlled despite treatment with calcitriol.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

**Prescriber Restrictions** 

**Coverage Duration** 

**NERLYNX** 

**NERLYNX** 

All FDA-approved indications not otherwise excluded from Part D.

The patient has early stage HER2-positive breast cancer. Nerlynx is initiated within two

years after completing adjuvant trastuzumab based therapy.

Age Restrictions

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**NEULASTA** 

NEULASTA. NEULASTA ONPRO KIT

All FDA-approved indications not otherwise excluded from Part D. mobilization of

peripheral blood progenitor cells prior to autologous transplantation.

**Exclusion Criteria** 

**Required Medical Information** 

For prophylaxis of myelosuppressive chemotherapy-induced febrile neutropenia the patient must meet all of the following: 1) Patient has a non-myeloid cancer, 2) Patient is

currently receiving or will be receiving treatment with myelosuppressive anti-cancer

therapy

Age Restrictions

Other Criteria

**Prescriber Restrictions** 

**Coverage Duration** 

6 months

Drug Names

NEXAVAR NEXAVAR

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D, osteosarcoma, soft tissue sarcoma subtypes: angiosarcoma, desmoid tumors (aggressive fibromatosis), gastrointestinal stromal tumor (GIST), medullary thyroid carcinoma, acute myeloid leukemia.

**Exclusion Criteria** 

**Required Medical Information** 

For renal cell carcinoma: the patient has relapsed or unresectable disease. For follicular, papillary, or Hurthle cell thyroid carcinoma: the disease is unresectable or metastatic. For medullary thyroid carcinoma: the patient has progressive or metastatic disease. For gastrointestinal stromal tumor: the disease has progressed after treatment with imatinib, sunitinib, or regorafenib. For acute myeloid leukemia: 1) the disease is relapsed or refractory, and 2) the patient has FLT3-ITD mutation-positive disease.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

Drug Names

**Covered Uses** 

NINLARO NINLARO

All FDA-approved indications not otherwise excluded from Part D, multiple myeloma in

combination with dexamethasone and lenalidomide, relapsed, refractory, or

progressive multiple myeloma in combination with dexamethasone

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

**Other Criteria** 

Plan Year

Drug Names
Covered Uses

Exclusion Criteria

**Required Medical Information** 

NORTHERA NORTHERA

All FDA-approved indications not otherwise excluded from Part D.

Prior to initial therapy, patient has a persistent, consistent decrease in systolic blood pressure of at least 20 mmHg OR decrease in diastolic blood pressure of at least 10 mmHg within 3 minutes of standing. Northera will be used for patients with neurogenic orthostatic hypotension associated with one of the following diagnoses: 1) Primary autonomic failure due to Parkinson's disease, multiple system atrophy, or pure autonomic failure, OR 2) Dopamine beta hydroxylase deficiency, OR 3) Non-diabetic autonomic neuropathy

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

3 months

Patients currently on Northera must experience a sustained decrease in dizziness to continue on therapy.

Prior Authorization Group
Drug Names
Covered Uses
Exclusion Criteria
Required Medical Information

NUCALA NUCALA

All FDA-approved indications not otherwise excluded from Part D.

For initial therapy for severe asthma with an eosinophilic phenotype: 1) Patient has baseline eosinophil count of at least 150 cells per microliter, and 2) Patient has a history of severe asthma attacks (exacerbations) despite current treatment with both of the following medications at optimized doses: a) inhaled corticosteroid and b) additional controller (long acting beta2-agonist, leukotriene modifier, or sustained-release theophylline). For continuation therapy for severe asthma with an eosinophilic phenotype: Asthma control has improved on Nucala treatment, demonstrated by a reduction in the frequency and/or severity of symptoms and exacerbations or a reduction in the daily maintenance oral corticosteroid dose. For initial therapy for eosinophilic granulomatosis with polyangiitis (EGPA): Patient has a history or the presence of an eosinophil count of more than 1000 cells per microliter or a blood eosinophil level of greater than 10 percent. For continuation of therapy for EGPA: Patient has a beneficial response to treatment with Nucala, demonstrated by any of the following: 1) a reduction in the frequency of relapses, 2) a reduction in the daily oral corticosteroid dose, or 3) no active vasculitis.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Asthma: 12 years of age or older, EGPA: 18 years of age or older

Plan Year

**Drug Names** 

NUEDEXTA NUEDEXTA

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

Covered Uses

**Exclusion Criteria** 

**Required Medical Information** 

NUPLAZID NUPLAZID

All FDA-approved indications not otherwise excluded from Part D.

The diagnosis of Parkinson's disease was made prior to the onset of psychotic

symptoms.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

Covered Uses

OCTREOTIDE

OCTREOTIDE ACETATE, SANDOSTATIN LAR DEPOT

All FDA-approved indications not otherwise excluded from Part D, meningiomas, thymomas and thymic carcinomas, and neuroendocrine tumors (NETs) of the

gastrointestinal (GI) tract, thymus, lung, pancreas and adrenal gland.

**Exclusion Criteria** 

Required Medical Information

For acromegaly: 1) Patient has a high pretreatment insulin-like growth factor-1 (IGF-1)

level for age and/or gender, and 2) Patient had an inadequate or partial response to surgery or radiotherapy OR there is a clinical reason for why the patient has not had surgery or radiotherapy. For NETs of the GI tract, thymus, and lung: patient has metastatic or unresectable disease. For adrenal gland NETs: patient has non-adrenocorticotropic hormone (non-ACTH) dependent Cushing's syndrome. For meningiomas: patient has unresectable disease. For thymomas and thymic

carcinomas: patient has progressed on at least one prior chemotherapy regimen.

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Plan Year

Other Criteria For acromegaly continuation of therapy: patient's IGF-1 level has decreased or

normalized since initiation of therapy.

**ODOMZO Prior Authorization Group Drug Names ODOMZO** 

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group OFEV OFEV Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

All FDA-approved indications not otherwise excluded from Part D.

Initial Review Only: The patient does not have a known etiology for interstitial lung disease and meets one of the following: 1) a high-resolution computed tomography (HRCT) study of the chest or surgical lung biopsy reveals the usual interstitial pneumonia (UIP) pattern, or 2) HRCT study of the chest reveals a possible UIP pattern and the diagnosis is supported either by surgical lung biopsy or by a multidisciplinary discussion between at least a radiologist and pulmonologist who are experienced in idiopathic pulmonary fibrosis if surgical lung biopsy has not been conducted. For initial and continuation: Ofey will not be used in combination with Esbriet

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

ONFI **Prior Authorization Group** 

**ONFI Drug Names** 

**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** 

Plan Year

Other Criteria

**Drug Names** 

ONMEL ONMEL

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

All FDA-approved indications not otherwise excluded from Part D.

Exclusion Criteria

**Required Medical Information** 

Treatment of onychomycosis of the toenail due to Trichophyton that has been

confirmed by a fungal diagnostic test.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

3 months

**Prior Authorization Group** 

**Drug Names** 

OPSUMIT OPSUMIT

**Covered Uses** 

Exclusion Criteria

**Required Medical Information** 

Pulmonary arterial hypertension (WHO Group 1) was confirmed by right heart catheterization. For new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than or equal to 25 mmHg, 2) pretreatment pulmonary capillary wedge

pressure is less than or equal to 15 mmHg, and 3) pretreatment pulmonary vascular

resistance is greater than 3 Wood units.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

Drug Names Covered Uses

**Exclusion Criteria** 

Required Medical Information

ORAL-INTRANASAL FENTANYL

ABSTRAL, FENTANYL CITRATE ORAL TRA, FENTORA, LAZANDA, SUBSYS

All FDA-approved indications not otherwise excluded from Part D.

1) The patient has CANCER related pain AND 2) The ICD diagnosis code provided supports the CANCER RELATED diagnosis [Note: For drug coverage approval, ICD diagnosis code provided MUST support the CANCER RELATED diagnosis.] AND 3) The drug is being prescribed for the management of breakthrough pain in a CANCER patient who is currently receiving around-the-clock opioid therapy for underlying CANCER pain AND 4) The patient can safely take the requested dose based on their current opioid use history.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

Prior Authorization Group ORALAIR
Drug Names ORALAIR

Covered Uses All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** Prescribed as immunotherapy for grass pollen-induced allergic rhinitis confirmed by

positive skin test or in vitro testing for pollen-specific IgE antibodies for any of the five grass species contained in this product. (Sweet Vernal, Orchard, Perennial Rye,

Timothy, Kentucky Blue Grass).

Age Restrictions 10 through 65 years of age

Prescriber Restrictions Allergist/Immunologist or in consultation with an Allergist/Immunologist.

Coverage Duration 6 months
Other Criteria

Prior Authorization Group ORENITRAM
Drug Names ORENITRAM

**Covered Uses**All FDA-approved indications not otherwise excluded from Part D.

Required Medical Information Pulmonary arterial hypertension (WHO Group 1) was confirmed by right heart

catheterization. For new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than or equal to 25 mmHg, 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, and 3) pretreatment pulmonary vascular

resistance is greater than 3 Wood units.

Age Restrictions

**Exclusion Criteria** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

Prior Authorization Group ORFADIN

**Drug Names** ORFADIN

Exclusion Criteria

Covered Uses All FDA-approved indications not otherwise excluded from Part D.

Required Medical Information Diagnosis of hereditary tyrosinemia type 1 is confirmed by one of the following: 1)

biochemical testing (e.g., detection of succinylacetone in urine) or 2) DNA testing

(mutation analysis).

Age Restrictions

**Prescriber Restrictions** 

Coverage Duration

Other Criteria

Plan Year

Prior Authorization GroupORKAMBIDrug NamesORKAMBI

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** The patient is positive for the F508del mutation on both alleles of the cystic fibrosis

transmembrane conductance regulator (CFTR) gene.

Age Restrictions 6 years of age or older

**Prescriber Restrictions** 

Coverage Duration Plan Year

Other Criteria Orkambi will not be used in combination with Kalydeco.

Prior Authorization Group OXANDROLONE
Drug Names OXANDROLONE

Covered Uses All FDA-approved indications not otherwise excluded from Part D, Cachexia associated

with AIDS (HIV-wasting) or due to chronic disease or to enhance growth in patients

with Turner's Syndrome.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

Coverage Duration

Other Criteria

6 months

Drug Names
Covered Uses

**PEGASYS** 

PEGASYS, PEGASYS PROCLICK

All FDA-approved indications not otherwise excluded from Part D, myeloproliferative neoplasm (primary myelofibrosis and post-polycythemia vera or post-essential

thrombocytopenia myelofibrosis).

**Exclusion Criteria** 

**Required Medical Information** 

For chronic hepatitis C (CHC): CHC infection confirmed by presence of HCV RNA in serum prior to starting treatment. Planned treatment regimen, genotype, prior treatment history, presence or absence of cirrhosis (compensated or decompensated [Child Turcotte Pugh class B or C]), presence or absence of HIV coinfection, presence or absence of resistance-associated variants where applicable, liver transplantation status if applicable. Coverage conditions and specific durations of approval will be based on current AASLD-IDSA treatment guidelines.

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

HCV=Criteria will be applied consistent with current AASLD-IDSA guidance. HBV=48

wks.Other=Plan Yr

Other Criteria

**Prior Authorization Group** 

**Drug Names** 

Covered Uses

PERJETA

**PERJETA** 

All FDA-approved indications not otherwise excluded from Part D, adjuvant therapy for

HER2-positive breast cancer, recurrent HER2-positive breast cancer.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

**Other Criteria** 

Neoadjuvant therapy: 6 months. Other: Plan Year

**Prior Authorization Group** 

**Drug Names** 

Covered Uses

**Exclusion Criteria** 

**Required Medical Information** 

**PHENYLBUTYRATE** 

BUPHENYL. SODIUM PHENYLBUTYRATE

All FDA-approved indications not otherwise excluded from Part D.

n Diagnosis of urea cycle disorder (UCD) was confirmed by enzymatic, biochemical or

genetic testing. Requested drug will be used for chronic management of UCD.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

Prior Authorization GroupPOMALYSTDrug NamesPOMALYST

Covered Uses All FDA-approved indications not otherwise excluded from Part D, systemic light chain

amyloidosis

**Exclusion Criteria** 

**Required Medical Information** Multiple myeloma: The patient has previously received at least two prior therapies for

multiple myeloma, including an immunomodulatory agent (ie, thalidomide,

lenalidomide) AND a proteasome inhibitor (ie, bortezomib, carfilzomib, ixazomib).

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

Prior Authorization Group
Drug Names
Covered Uses
Exclusion Criteria
Required Medical Information

PRALUENT PRALUENT

All FDA-approved indications not otherwise excluded from Part D.

Member must have one of the following conditions (new starts and continuation): 1) Prior clinical atherosclerotic cardiovascular disease (ASCVD) or cardiovascular event (see Other Criteria) OR, 2) Heterozygous familial hypercholesterolemia (HeFH): Diagnosis of FH (See Other Criteria). For new starts: For members with prior clinical ASCVD or cardiovascular event, at least one of the following requirements is met: 1) Current low density lipoprotein (LDL-C) level 70 mg/dL or greater after treatment with a high-intensity statin (eg. atorvastatin, rosuvastatin), 2) Current LDL-C level 70 mg/dL or greater with intolerance to a high-intensity statin AND is taking a maximally tolerated dose of any statin, 3) Current LDL-C level 70 mg/dL or greater with contraindication to statin (see Other Criteria) OR intolerance to any dose of two statins. For members with HeFH, at least one of the following requirements is met: 1) With ASCVD: See requirements for members with prior ASCVD above, 2) Current LDL-C level 100 mg/dL or greater after treatment with a high-intensity statin (eg, atorvastatin, rosuvastatin), 3) Current LDL-C level 100 mg/dL or greater with intolerance to a high-intensity statin AND is taking a maximally tolerated dose of any statin, 4) Current LDL-C level 100 mg/dL or greater with contraindication to statin OR intolerance to any dose of two statins. For continuation: Response to therapy as demonstrated by a reduction in LDL-C.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Plan Year

Prior clinical atherosclerotic cardiovascular disease (ASCVD) or cardiovascular event is defined as: acute coronary syndromes, myocardial infarction, stable or unstable angina, coronary or other arterial revascularization procedure [eg, PTCA, CABG], stroke of presumed atherosclerotic origin, transient ischemic attack [TIA], non-cardiac peripheral arterial disease of presumed atherosclerotic origin, or obstructive coronary artery disease [defined as fifty percent or greater stenosis on cardiac computed tomography angiogram or catheterization]. Diagnosis of FH must be confirmed by one of the following: 1) Genetic confirmation: An LDL-receptor mutation, familial defective apo B-100, or a PCSK9 gain-of-function mutation, 2) Simon-Broome Diagnostic Criteria for FH: Simon-Broome Diagnostic Criteria for FH: Total cholesterol greater than 290 mg/dL or LDL-C greater than 190 mg/dL in patients over 16 years of age or total cholesterol greater than 260 mg/dl or LDL-C greater than 155 mg/dl in patients less than 16 years of age and one of the following: a)Tendon xanthomas in the patient, first (parent, sibling or child) or second degree relative (grandparent, uncle or aunt), b)Family history of myocardial infarction in a first degree relative before the age of 60 or in a second degree relative before the age of 50, c) Total cholesterol greater than 290 mg/dl in an adult first or second degree relative, d) Total cholesterol greater than 260 mg/dl in a child, brother, or sister aged younger than 16 years. 3) Dutch Lipid Clinic Network

Criteria for definite or probable FH: Total score greater than 5 points.

Prior Authorization GroupPROCYSBIDrug NamesPROCYSBI

Covered Uses All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** Diagnosis of nephropathic cystinosis was confirmed by the presence of increased

cystine concentration in leukocytes or by genetic testing. Patient has tried and

experienced intolerance to prior therapy with Cystagon.

Age Restrictions 1 year of age or older

Prescriber Restrictions

Plan Year

Coverage Duration
Other Criteria

Prior Authorization Group
Drug Names
Covered Uses
Exclusion Criteria
Required Medical Information

PROMACTA PROMACTA

All FDA-approved indications not otherwise excluded from Part D.

For chronic or persistent immune thrombocytopenia (ITP): 1) For new starts: a) patient has had an inadequate response or is intolerant to corticosteroids, immunoglobulins or splenectomy, AND b) untransfused platelet count at time of diagnosis is less than 30,000/mcL OR 30,000-50,000/mcL with symptomatic bleeding or risk factor(s) for bleeding. 2) For continuation of therapy, platelet (plt) count response to Promacta: a) current plt count is 50,000-200,000/mcL OR b) current plt count is less than 50,000/mcL and sufficient to avoid clinically important bleeding OR c) current plt count is less than 50,000/mcL and patient has not received a maximal dose of Promacta for at least 4 weeks OR d) current plt count is greater than 200,000/mcL and dosing will be adjusted to a plt count sufficient to avoid clinically important bleeding. For thrombocytopenia associated with chronic hepatitis C: 1) For new starts: a) Promacta is used for initiation and maintenance of interferon-based therapy, AND b) untransfused platelet count at time of diagnosis is less than 75,000/mcL. 2) For continuation of therapy: patient is receiving interferon-based therapy. For severe aplastic anemia (AA): 1) For new starts: a) patient has had an inadequate response to immunosuppressive therapy, AND b) untransfused platelet count at time of diagnosis is less than or equal to 30,000/mcL. 2) For continuation of therapy, plt count response to Promacta: 1) current plt count is 50,000-200,000/mcL OR b) current plt count is less than 50,000/mcL and patient has not received appropriately titrated therapy for at least 16 weeks OR c) current plt count is less than 50.000/mcL and patient is transfusion-independent OR d) current plt count is greater than 200,000/mcL and dosing will be adjusted to achieve and maintain an appropriate target plt count.

Age Restrictions
Prescriber Restrictions
Coverage Duration

Other Criteria

HCV:6mo, INITIAL:ITP/AA-6mo, REAUTH:1)ITP/AA APR-Plan Yr, 2)ITP IPR-3mo, 3)AA IPR-16wks

APR:adequate platelet response (greater than or equal to 50k/mcL), IPR:inadequate platelet response (less than 50k/mcL)

**Drug Names** 

**PULMOZYME PULMOZYME** 

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Diagnosis of cystic fibrosis was confirmed by appropriate diagnostic or genetic testing.

Plan Year

Coverage under Part D will be denied if coverage is available under Part A or Part B as

the medication is prescribed and dispensed or administered for the individual.

**Prior Authorization Group** 

**Drug Names** 

**QUININE SULFATE QUININE SULFATE** 

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D, Babesiosis,

uncomplicated Plasmodium vivax malaria.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

1 month

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

**RAVICTI RAVICTI** 

All FDA-approved indications not otherwise excluded from Part D.

Diagnosis of urea cycle disorder (UCD) was confirmed by enzymatic, biochemical or

genetic testing. Ravicti will be used for chronic management of UCD. Patient has experienced intolerance to prior Buphenyl therapy OR patient has not tried Buphenyl because of a comorbid condition that prohibits a trial due to its sodium content (e.g.,

heart failure, hypertension, renal impairment, edema).

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

2 months of age or older

Plan Year

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

REGRANEX REGRANEX

All FDA-approved indications not otherwise excluded from Part D.

For the treatment of lower extremity diabetic neuropathic ulcers that extend into the

subcutaneous tissue or beyond and have an adequate blood supply

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

20 weeks

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

**RELISTOR INJ RELISTOR** 

All FDA-approved indications not otherwise excluded from Part D.

1) The requested drug is being prescribed for opioid-induced constipation in an adult patient with advanced illness or pain caused by active cancer who requires opioid dosage escalation for palliative care OR 2) The requested drug is being prescribed for opioid-induced constipation in an adult patient with chronic non-cancer pain, including chronic pain related to prior cancer or its treatment who does not require frequent (e.g., weekly) opioid dosage escalation AND 3) The patient is unable to tolerate oral medications OR 4) An oral drug indicated for opioid-induced constipation in an adult patient with chronic non-cancer pain has been tried. (Note: Examples are Amitiza or Movantik) AND 5) The patient experienced an inadequate treatment response or intolerance to an oral drug indicated for opioid-induced constipation in an adult patient with chronic non-cancer pain. (Note: Examples are Amitiza or Movantik) OR 6) The patient has a contraindication to an oral drug indicated for opioid-induced constipation in an adult patient with chronic non-cancer pain (Note: Examples are Amitiza or Movantik).

Age Restrictions **Prescriber Restrictions Coverage Duration** Other Criteria

4 months

Prior Authorization GroupRELISTOR TABDrug NamesRELISTOR

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

REMICADE REMICADE

All FDA-approved indications not otherwise excluded from Part D, axial spondyloarthritis, Behcet's syndrome, granulomatosis with polyangiitis (Wegener's granulomatosis), hidradenitis suppurativa, juvenile idiopathic arthritis, pyoderma gangrenosum, sarcoidosis, Takayasu's arteritis, uveitis.

Exclusion Criteria
Required Medical Information

For moderately to severely active Crohn's disease (new starts only): 1) Patient has fistulizing disease, OR 2) Inadequate response or intolerance to a self-injectable tumor necrosis factor (TNF) inhibitor (e.g., adalimumab). For moderately to severely active ulcerative colitis (new starts only): 1) Inadequate response to at least one conventional therapy (e.g., corticosteroids, sulfasalazine, azathioprine, mesalamine), OR 2) Intolerance or contraindication to conventional therapy. For moderately to severely active rheumatoid arthritis (new starts only): 1) Will be used in combination with methotrexate (MTX) or leflunomide OR patient has intolerance or contraindication to MTX or leflunomide, AND 2) Inadequate response or intolerance to a self-injectable TNF inhibitor (e.g., adalimumab) or a targeted synthetic disease-modifying antirheumatic drug (DMARD) (e.g., tofacitinib). For active ankylosing spondylitis and axial spondyloarthritis (new starts only): Inadequate response to a non-steroidal antiinflammatory drug (NSAID) trial at maximum recommended or tolerated dose OR intolerance or contraindication to NSAIDs. For moderate to severe chronic plaque psoriasis (new starts only): 1) At least 5% of body surface area (BSA) is affected OR crucial body areas (e.g., feet, hands, face, neck, groin, intertriginous areas) are affected at time of diagnosis, AND 2) Inadequate response or intolerance to a selfinjectable TNF inhibitor (e.g., adalimumab). For juvenile idiopathic arthritis (new starts only): Inadequate response or intolerance to a self-injectable TNF inhibitor (e.g., adalimumab). For hidradenitis suppurativa (new starts only): patient has severe, refractory disease. For uveitis (new starts only): Patient has experienced an inadequate response or intolerance or has a contraindication to a trial of immunosuppressive therapy for uveitis (e.g., methotrexate, azathioprine, or mycophenolate mofetil).

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Plan Year

REVLIMID REVLIMID

All FDA-approved indications not otherwise excluded from Part D, systemic light chain amyloidosis, classical Hodgkin lymphoma, myelofibrosis-associated anemia, non-Hodgkin's lymphoma with the following subtypes: chronic lymphocytic leukemia/small lymphocytic lymphoma, AIDS-related diffuse large B-cell lymphoma, primary effusion lymphoma, lymphoma associated with Castleman's disease, diffuse large B-cell lymphoma, follicular lymphoma, nongastric/gastric MALT lymphoma, primary cutaneous B-cell lymphoma, splenic marginal zone lymphoma, multicentric Castleman's disease, adult T-cell leukemia/lymphoma, mycosis fungoides/Sezary syndrome, angioimmunoblastic T-cell lymphoma, peripheral T-cell lymphoma not otherwise specified, enteropathy-associated T-cell lymphoma, primary cutaneous anaplastic large cell lymphoma

Exclusion Criteria
Required Medical Information

Myelodysplastic syndrome (MDS): Low- to intermediate-1 risk MDS with symptomatic anemia

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Plan Year

RITUXAN RITUXAN

All FDA-approved indications not otherwise excluded from Part D, primary CNS lymphoma, leptomeningeal metastases from lymphomas, Hodgkin's lymphoma (lymphocyte-predominant), non-Hodgkin's lymphoma subtypes [marginal zone lymphomas (splenic, MALT), Mantle cell lymphoma, Burkitt lymphoma, AIDS-related B-cell lymphoma, relapsed/refractory hairy cell leukemia, small lymphocytic lymphoma (SLL), post-transplant lymphoproliferative disorder (PTLD), primary cutaneous B-cell lymphoma, lymphoblastic lymphoma, Castleman's disease], acute lymphoblastic leukemia (ALL), autoimmune hemolytic anemia, chronic graft-versus-host disease (GVHD), refractory immune or idiopathic thrombocytopenic purpura (ITP), Waldenstrom's macroglobulinemia, lymphoplasmacytic lymphoma, Sjogren syndrome, thrombotic thrombocytopenic purpura, prevention of Epstein-Barr virus (EBV)-related PTLD, multiple sclerosis.

## Exclusion Criteria Required Medical Information

For moderately to severely active rheumatoid arthritis (new starts only): 1) Rituxan is used in combination with methotrexate (MTX) unless MTX is contraindicated or not tolerated, AND 2) Patient has an inadequate response, intolerance or contraindication to a self-injectable tumor necrosis factor (TNF) inhibitor (e.g., adalimumab) or a targeted synthetic disease-modifying antirheumatic drug (DMARD) (e.g., tofacitinib). Hematologic malignancies must be CD20-positive. For Burkitt lymphoma and ALL, Rituxan is used as a component of a chemotherapy regimen. For Wegener's Granulomatosis (WG) and Microscopic Polyangiitis (MPA), Rituxan will be used in combination with glucocorticoids. For multiple sclerosis: 1) Patient has a diagnosis of relapsing remitting multiple sclerosis, AND 2) Patient has had an inadequate response to two or more disease-modifying drugs indicated for multiple sclerosis despite adequate duration of treatment.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Plan Year

**Drug Names** 

RITUXAN HYCELA RITUXAN HYCELA

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Malignancies must be CD20 positive. Patient must receive at least one full dose of a rituximab product by intravenous infusion without experiencing severe adverse

reactions.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

**RUBRACA RUBRACA** 

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

RUCONEST RUCONEST

All FDA-approved indications not otherwise excluded from Part D.

This medication is being used for the treatment of acute angioedema attacks. Diagnostic laboratory testing for HAE has been performed (eg, C4, C1 inhibitor

functional, and C1 inhibitor antigenic protein levels). For patients with HAE with C1 inhibitor deficiency, C1 inhibitor antigenic protein level and/or C1 inhibitor functional level is below the lower limit of normal as defined by the laboratory performing the test. For patients with HAE with normal C1 inhibitor, other causes of angioedema have been ruled out (eg, drug induced) and EITHER 1) Patient tested positive for the F12 gene mutation OR 2) Patient has a family history of angioedema and the angioedema was

refractory to a trial of antihistamine (eq. levocetirizine) for at least one month.

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

Drug Names

RYDAPT RYDAPT

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

For newly diagnosed FLT3 mutation-positive AML, Rydapt is/was used in combination

with standard cytarabine with daunorubicin or idarubicin induction followed by

cytarabine consolidation chemotherapy.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

18 years of age or older

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

**SABRIL** 

SABRIL, VIGABATRIN

All FDA-approved indications not otherwise excluded from Part D.

For infantile spasms (IS): The requested drug is used as a single agent in the treatment of IS. For complex partial seizures (CPS): 1) patient had an inadequate response to at least 2 alternative therapies for CPS (e.g., carbamazepine, phenytoin, levetiracetam, topiramate, oxcarbazepine or lamotrigine), AND 2) The requested drug is used as

adjunctive therapy.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

Drug Names

SAMSCA SAMSCA

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

Underlying liver disease.

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

0 5 "

**Coverage Duration** 

30 days

Other Criteria

Samsca therapy was initiated (or re-initiated) in the hospital.

Prior Authorization Group SEROSTIM
Drug Names SEROSTIM

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** Active malignancy.

**Required Medical Information** Serostim is used in combination with antiretroviral therapy. Patient has had a

suboptimal response to at least 1 other therapy for wasting or cachexia (e.g., megestrol, dronabinol, cyproheptadine, or testosterone therapy if hypogonadal) OR patient has contraindication or intolerance to alternative therapies. For initial approval, patient must have a body mass index (BMI) less than 18.5 kg per meter squared. The patient must also have experienced unintentional weight loss greater than 5 percent of body weight in the previous 6 months. For continuation of therapy, patient must have demonstrated a response to therapy with Serostim (i.e., BMI has increased or

stabilized).

Age Restrictions

**Prescriber Restrictions** Infectious disease specialist

Coverage Duration 12 weeks

Other Criteria

Prior Authorization Group SIGNIFOR

**Drug Names**SIGNIFOR **Covered Uses**All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** Patient has had pituitary surgery that was not curative or the patient is not a candidate

for surgery. For continuation of therapy, patient must show a clinically meaningful reduction in 24-hour urinary free cortisol levels and/or improvement in signs or

symptoms of the disease.

Age Restrictions
Prescriber Restrictions

Coverage Duration
Other Criteria

Plan Year

Prior Authorization GroupSIGNIFOR LARDrug NamesSIGNIFOR LAR

Covered Uses All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

Required Medical Information Patient meets both of the following criteria: 1) Patient has high pretreatment insulin-like

growth factor-1 (IGF-1) level for age and/or gender, and 2) Patient had an inadequate or partial response to surgery OR there is a clinical reason for why the patient has not

had surgery.

Age Restrictions

**Prescriber Restrictions** 

Coverage Duration Plan Year

Other Criteria For continuation of therapy: patient's IGF-1 level has decreased or normalized since

initiation of therapy.

**Prior Authorization Group** SILDENAFIL

**Drug Names** REVATIO, SILDENAFIL

Covered Uses All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** Pulmonary arterial hypertension (WHO Group 1) was confirmed by right heart

catheterization. For new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than or equal to 25 mmHg, 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, and 3) pretreatment pulmonary vascular

resistance is greater than 3 Wood units.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**SIRTURO** 

**Prior Authorization Group** 

**Drug Names** SIRTURO

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** The requested drug is being prescribed for the treatment of latent infection due to

Mycobacterium tuberculosis, drug-sensitive tuberculosis, extra-pulmonary tuberculosis,

or infection caused by the non-tuberculous mycobacteria

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

Coverage Duration

Other Criteria

6 months

**Drug Names Covered Uses**  SOMATULINE DEPOT SOMATULINE DEPOT

All FDA-approved indications not otherwise excluded from Part D. neuroendocrine tumors (NETs) of the gastrointestinal (GI) tract, thymus, lung, pancreas, and adrenal

gland.

**Exclusion Criteria** 

**Required Medical Information** 

For acromegaly: 1) Patient has a high pretreatment insulin-like growth factor-1 (IGF-1) level for age and/or gender, and 2) Patient had an inadequate or partial response to surgery or radiotherapy OR there is a clinical reason for why the patient has not had surgery or radiotherapy. For NETs of the GI tract, thymus, and lung: patient has metastatic or unresectable disease. For adrenal gland NETs: patient has nonadrenocorticotropic hormone (non-ACTH) dependent Cushing's syndrome.

Age Restrictions

**Prescriber Restrictions Coverage Duration** 

Other Criteria

Plan Year

**SOMAVERT** 

**SOMAVERT** 

For acromegaly continuation of therapy: patient's IGF-1 level has decreased or

All FDA-approved indications not otherwise excluded from Part D.

for why the patient has not had surgery or radiotherapy.

normalized since initiation of therapy.

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions **Prescriber Restrictions Coverage Duration** 

Other Criteria

Plan Year

For continuation of therapy: patient's IGF-1 level has decreased or normalized since

Patient meets both of the following criteria: 1) Patient has a high pretreatment insulin-

inadequate or partial response to surgery or radiotherapy OR there is a clinical reason

like growth factor-1 (IGF-1) level for age and/or gender, and 2) Patient had an

initiation of therapy.

Prior Authorization Group
Drug Names
Covered Uses
Exclusion Criteria
Required Medical Information

SOVALDI SOVALDI

All FDA-approved indications not otherwise excluded from Part D.

Chronic hepatitis C infection confirmed by presence of HCV RNA in serum prior to treatment. Planned treatment regimen, genotype, prior treatment history, presence or absence of cirrhosis (compensated or decompensated [Child Turcotte Pugh class B or C]), presence or absence of HIV coinfection, presence or absence of resistance-associated variants (eg, NS3 Q80K polymorphism) where applicable, liver transplantation status if applicable. For patients with genotype 1, 2, 3, or 4 infection and hepatocellular carcinoma awaiting liver transplantation: must meet MILAN criteria. Coverage conditions and specific durations of approval will be based on current AASLD treatment guidelines.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Criteria will be applied consistent with current AASLD-IDSA guidance For HCV/HIV coinfection, patient meets criteria for requested regimen. For patients prescribed a treatment regimen that includes Olysio, no prior treatment failure with an HCV protease inhibitor (eg, telaprevir, simeprevir, boceprevir, paritaprevir) despite adequate dosing and duration of therapy. MILAN criteria defined as: 1) tumor size 5 cm or less in diameter in pts with single hepatocellular carcinoma OR 3 tumor nodules or less, each 3 cm or less in diameter in pts with multiple tumors, and 2) no extrahepatic manifestations of the cancer or evidence of vascular invasion of tumor.

**Drug Names** 

**SPRYCEL SPRYCEL** 

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D. gastrointestinal

stromal tumor (GIST).

**Exclusion Criteria** 

**Required Medical Information** 

For CML or ALL, diagnosis was confirmed by detection of the Philadelphia chromosome or BCR-ABL gene. For CML, 1) patient has received a hematopoietic stem cell transplant, OR 2) Patient has accelerated or blast phase CML, OR 3) For chronic phase CML, patient has one of the following a) patient is 21 years of age or younger, or b) high or intermediate risk for disease progression, or c) low risk for disease progression and has experienced resistance, intolerance or toxicity to imatinib or an alternative tyrosine kinase inhibitor. If patient experienced resistance to imatinib or an alternative tyrosine kinase inhibitor for CML, patient is negative for T315I mutation. For GIST, patient must have PDGFRA D842V mutation and disease progression on imatinib, sunitinib, or regorafenib.

Age Restrictions **Prescriber Restrictions Coverage Duration** 

Plan Year

**Prior Authorization Group** 

**Drug Names** 

Other Criteria

**Covered Uses** 

Other Criteria

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions **Prescriber Restrictions Coverage Duration** 

**STIVARGA** STIVARGA

All FDA-approved indications not otherwise excluded from Part D.

For colorectal cancer: The disease is unresectable advanced or metastatic. The patient has progressed on treatment with EITHER a) FOLFOXIRI (fluorouracil, leucovorin,

oxaliplatin, and irinotecan) regimen OR b) irinotecan- AND oxaliplatin-based regimens.

Plan Year

Drug Names

SUTENT SUTENT

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D, thyroid carcinoma (follicular, papillary, Hurthle cell, or medullary), angiosarcoma, solitary fibrous tumor, hemangiopericytoma, chordoma (bone cancer), lung neuroendocrine tumor, thymic

carcinoma.

**Exclusion Criteria** 

**Required Medical Information** 

For renal cell carcinoma: Either 1) The disease is relapsed or unresectable OR 2) the patient is at high risk of disease recurrence following nephrectomy. For gastrointestinal stromal tumor: the patient experienced disease progression on imatinib or was intolerant to imatinib. For follicular, papillary, or Hurthle cell thyroid carcinoma: the disease is unresectable or metastatic. For medullary thyroid carcinoma: the patient has progressive or metastatic disease. For thymic carcinoma: the disease has progressed on a platinum-based chemotherapy regimen.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

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**Required Medical Information** 

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

SYLATRON
All FDA-app

**SYLATRON** 

All FDA-approved indications not otherwise excluded from Part D, myelofibrosis

er Criteria

**Prior Authorization Group** 

**Drug Names** 

Covered Uses

**Exclusion Criteria** 

Required Medical Information

**SYMLIN** 

Plan Year

SYMLINPEN 120, SYMLINPEN 60

All FDA-approved indications not otherwise excluded from Part D.

1) The patient has a diagnosis of type 1 or type 2 diabetes mellitus AND 2) The patient

is currently receiving optimal mealtime insulin therapy AND 3) The patient has

experienced an inadequate treatment response to insulin.

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

Prior Authorization Group SYNRIBO Drug Names SYNRIBO

Covered Uses All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** For chronic myeloid leukemia (CML), the patient has experienced resistance, toxicity or

intolerance to prior therapy with at least two tyrosine kinase inhibitors (TKIs) (eg,

imatinib, dasatinib, nilotinib, bosutinib, ponatinib).

Age Restrictions

Prescriber Restrictions

Coverage Duration
Other Criteria

Plan Year

**Prior Authorization Group** 

Drug Names

Covered Uses

**Exclusion Criteria** 

**Required Medical Information** 

TADALAFIL (PAH)

**ADCIRCA** 

All FDA-approved indications not otherwise excluded from Part D.

Pulmonary arterial hypertension (WHO Group 1) was confirmed by right heart

catheterization. For new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than or equal to 25 mmHg, 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, and 3) pretreatment pulmonary vascular

resistance is greater than 3 Wood units.

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

Drug Names

**Covered Uses** 

TAFINLAR TAFINLAR

All FDA-approved indications not otherwise excluded from Part D, non-small cell lung

cancer (NSCLC).

**Exclusion Criteria** 

**Required Medical Information** 

For unresectable or metastatic melanoma, the tumor is positive for BRAF V600E or

V600K mutation, and Tafinlar will be used as a single agent or in combination with trametinib. For NSCLC, the tumor is positive for the BRAF V600E mutation and Tafinlar

will be used as a single agent or in combination with trametinib.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Drug Names** 

TAGRISSO TAGRISSO

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

Exclusion Criteria

Required Medical Information

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

TARCEVA TARCEVA

All FDA-approved indications not otherwise excluded from Part D, chordoma, renal cell

carcinoma (RCC).

**Exclusion Criteria** 

**Required Medical Information** 

For non-small cell lung cancer (NSCLC), patient meets any of the following: 1) Tarceva is used as first-line therapy (EGFR mutation discovered prior to first-line chemotherapy or during first-line chemotherapy) or as subsequent therapy following disease progression on first-line therapy with erlotinib AND the patient has recurrent or metastatic NSCLC with a known sensitizing EGFR mutation (e.g., EGFR exon 19 deletion or exon 21 (L858R) substitution mutation) confirmed by EGFR mutation testing, OR 2) Tarceva is used for metastatic NSCLC as maintenance therapy or as second or greater line treatment after progression following at least one prior chemotherapy regimen in patients with an EGFR exon 19 deletion or exon 21 (L858R) substitution mutation confirmed by EGFR mutation testing. For pancreatic cancer, Tarceva is prescribed in combination with gemcitabine for locally advanced unresectable or metastatic pancreatic cancer. For chordoma, Tarceva is prescribed for recurrent disease. For RCC, Tarceva is prescribed for relapsed or unresectable stage IV disease with non-clear cell histology.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Plan Year

**Drug Names** 

**TASIGNA TASIGNA** 

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D. Philadelphia

chromosome positive acute lymphoblastic leukemia (Ph+ ALL), gastrointestinal stromal

tumor (GIST).

**Exclusion Criteria** 

**Required Medical Information** For CML or ALL, diagnosis was confirmed by detection of the Philadelphia

chromosome or BCR-ABL gene. For CML, 1) patient has received a hematopoietic stem cell transplant, OR 2) Patient has accelerated or blast phase CML, OR 3) For chronic phase CML, patient has one of the following a) high or intermediate risk for disease progression, or b) low risk for disease progression and has experienced resistance, intolerance or toxicity to imatinib or an alternative tyrosine kinase inhibitor. If patient experienced resistance to imatinib or an alternative tyrosine kinase inhibitor for CML, patient is negative for T315I mutation. For GIST, patient must have progressed

on imatinib, sunitinib or regorafenib.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

**TAZORAC** 

TAZAROTENE, TAZORAC

All FDA-approved indications not otherwise excluded from Part D.

For patients being treated for plague psoriasis, the requested drug must be applied to

less than 20 percent of the patient's body surface area.

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

**TECENTRIQ TECENTRIQ** 

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

TESTOSTERONE CYPIONATE INJ TESTOSTERONE CYPIONATE

All FDA-approved indications not otherwise excluded from Part D, Gender Dysphoria in Female-to-Male transgender patients

Exclusion Criteria
Required Medical Information

1) Request is for continuation of testosterone therapy and requested drug is being prescribed for hypogonadism in a male patient or a patient that self-identifies as male who has had a confirmed low testosterone level according to current practice guidelines or your standard male lab reference values before starting testosterone therapy OR 2) Request is not for continuation of testosterone therapy and requested drug is being prescribed for hypogonadism in a male patient or a patient that self-identifies as male who has at least two confirmed low testosterone levels according to current practice guidelines or your standard male lab reference values OR 3) Requested drug is being prescribed for female-to-male gender reassignment in a patient who is 12 years of age or older and able to make an informed, mature decision to engage in therapy

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

12 years of age or older (applies to gender reassignment only)

Plan Year

Prior Authorization Group
Drug Names
Covered Uses
Exclusion Criteria
Required Medical Information

TESTOSTERONE ENANTHATE INJ TESTOSTERONE ENANTHATE

All FDA-approved indications not otherwise excluded from Part D.

1) Requested drug is being prescribed for inoperable metastatic breast cancer in a patient who is 1 to 5 years postmenopausal and who has had an incomplete response to other therapy for metastatic breast cancer OR 2) Requested drug is being prescribed for a pre-menopausal patient with breast cancer who has benefited from oophorectomy and is considered to have a hormone-responsive tumor OR 3) Request is for continuation of testosterone therapy and requested drug is being prescribed for hypogonadism in a male patient or a patient that self-identifies as male who had a confirmed low testosterone level according to current practice guidelines or your standard male lab reference values before starting testosterone therapy OR 4) Request is not for continuation of testosterone therapy and requested drug is being prescribed for hypogonadism in a male patient or a patient that self-identifies as male who has at least two confirmed low testosterone levels according to current practice guidelines or your standard male lab reference values OR 5) Requested drug is being prescribed for delayed puberty in a male patient.

Age Restrictions
Prescriber Restrictions
Coverage Duration
Other Criteria

Plan Year

Prior Authorization Group Drug Names

Covered Uses

Exclusion Criteria
Required Medical Information
Age Restrictions
Prescriber Restrictions

Other Criteria

**Coverage Duration** 

TETRABENAZINE TETRABENAZINE

All FDA-approved indications not otherwise excluded from Part D, chronic tics, tardive dyskinesia, hemiballismus, chorea not associated with Huntington's disease.

Plan Year

**Drug Names** 

THALOMID THALOMID

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D, myelofibrosis-related

anemia, systemic light chain amyloidosis, Waldenstrom's

macroglobulinemia/lymphoplasmacytic lymphoma, recurrent aphthous stomatitis, recurrent HIV-associated aphthous ulcers, cachexia, HIV-associated diarrhea, Kaposi's sarcoma, Behcet's syndrome, chronic graft-versus-host disease, Crohn's disease,

multicentric Castleman's disease.

**Exclusion Criteria** 

**Required Medical Information** 

Cachexia: Cachexia must be due to cancer or HIV-infection. Kaposi's sarcoma: The

patient has HIV infection.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

**Other Criteria** 

Plan Year

**Prior Authorization Group** 

**Drug Names** 

TOBI PODHALER

TOBI INHALER

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D, non-cystic fibrosis

bronchiectasis.

**Exclusion Criteria** 

**Required Medical Information** 

The patient has a diagnosis of cystic fibrosis that is confirmed by appropriate diagnostic or genetic testing OR the patient has a diagnosis of non-cystic fibrosis bronchiectasis. Pseudomonas aeruginosa is present in the patient's airway cultures OR the patient has

a history of pseudomonas aeruginosa infection or colonization in the airways.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

**Other Criteria** 

Plan Year

Drug Names
Covered Uses

**TOBRAMYCIN** 

BETHKIS, TOBRAMYCIN

All FDA-approved indications not otherwise excluded from Part D, non-cystic fibrosis

bronchiectasis.

**Exclusion Criteria** 

**Required Medical Information** 

The patient has a diagnosis of cystic fibrosis that is confirmed by appropriate diagnostic or genetic testing OR the patient has a diagnosis of non-cystic fibrosis bronchiectasis. Pseudomonas aeruginosa is present in the patient's airway cultures OR the patient has a history of pseudomonas aeruginosa infection or colonization in the airways.

Age Restrictions

Prescriber Restrictions
Coverage Duration

Other Criteria

Plan Year

Coverage under Part D will be denied if coverage is available under Part A or Part B as

the medication is prescribed and dispensed or administered for the individual.

**Prior Authorization Group** 

**Drug Names** 

**TOPICAL LIDOCAINE** 

GLYDO, LIDOCAINE, LIDOCAINE HCL, LIDOCAINE HCL JELLY,

All FDA-approved indications not otherwise excluded from Part D.

LIDOCAINE/PRILOCAINE, SYNERA

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

3 months

1) If being used as part of a compounded product, all active ingredients in the

compounded product are FDA approved for topical use. 2) Coverage under Part D will

be denied if coverage is available under Part A or Part B as the medication is

prescribed and dispensed or administered for the individual.

**Drug Names** 

**TOPICAL TESTOSTERONES** 

ANDRODERM, ANDROGEL, ANDROGEL PUMP, AXIRON, STRIANT, TESTIM,

TESTOSTERONE, TESTOSTERONE PUMP, VOGELXO

**Covered Uses Exclusion Criteria** 

**Required Medical Information** 

All FDA-approved indications not otherwise excluded from Part D.

1) Request is for continuation of testosterone therapy and requested drug is being prescribed for hypogonadism in a male patient or a patient that self-identifies as male who had a confirmed low testosterone level according to current practice guidelines or your standard male lab reference values before starting testosterone therapy OR 2) Request is not for continuation of testosterone therapy and requested drug is being prescribed for hypogonadism in a male patient or a patient that self-identifies as male who has at least two confirmed low testosterone levels according to current practice guidelines or your standard male lab reference values.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

TOPICAL TRETINOIN

AVITA, RETIN-A MICRO, RETIN-A MICRO PUMP, TRETINOIN, TRETINOIN

All FDA-approved indications not otherwise excluded from Part D.

**MICROSPHERE** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

**TRELSTAR** 

TRELSTAR MIXJECT

All FDA-approved indications not otherwise excluded from Part D.

Plan Year

**Drug Names** 

TREPROSTINIL INJ
REMODULIN

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Pulmonary arterial hypertension (WHO Group 1) was confirmed by right heart catheterization. For new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than or equal to 25 mmHg, 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, and 3) pretreatment pulmonary vascular

resistance is greater than 3 Wood units.

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

Coverage under Part D will be denied if coverage is available under Part A or Part B as

the medication is prescribed and dispensed or administered for the individual.

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

TYKERB TYKERB

All FDA-approved indications not otherwise excluded from Part D, metastatic CNS

lesions from HER2-positive breast cancer.

**Exclusion Criteria** 

**Required Medical Information** 

For HER2-positive breast cancer, the requested drug will be used in combination with:

1) aromatase inhibitor (e.g., anastrozole, letrozole, exemestane), or 2) capecitabine, or

3) trastuzumab.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** TYMLOS

Drug Names TYMLOS

Covered Uses All FDA-approved indications not otherwise excluded from Part D.

Exclusion Criteria

**Required Medical Information** Patient has ONE of the following (1. or 2.): 1) Member has a history of fragility

fractures, OR 2) Member has a pre-treatment T-score of less than or equal to -2.5 and meets ANY of the following criteria (a or b): a) Member has indicators of higher fracture risk (e.g., advanced age, frailty, glucocorticoid use, very low T-scores, or increased fall

risk), OR b) Member has failed prior treatment with or is intolerant to previous

osteoporosis therapy (i.e., oral bisphosphonates or injectable antiresorptive agents)

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 24 months lifetime total for parathyroid hormone analogs (e.g., abaloparatide or

teriparatide)

Other Criteria

Prior Authorization Group TYSABRI
Drug Names TYSABRI

Covered Uses All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** For Crohn's disease (CD), patient must have an inadequate response, intolerance or

contraindication to one conventional CD therapy (e.g., corticosteroid, azathioprine, mesalamine) AND one tumor necrosis factor (TNF) inhibitor (e.g., adalimumab).

Age Restrictions

**Prescriber Restrictions** 

Coverage Duration

Other Criteria

Plan Year

**TYVASO Prior Authorization Group Drug Names TYVASO** 

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** Pulmonary arterial hypertension (WHO Group 1) was confirmed by right heart

> catheterization. For new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than or equal to 25 mmHg, 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, and 3) pretreatment pulmonary vascular

resistance is greater than 3 Wood units.

Age Restrictions

**Prescriber Restrictions Coverage Duration** 

Other Criteria Coverage under Part D will be denied if coverage is available under Part A or Part B as

the medication is prescribed and dispensed or administered for the individual.

**Prior Authorization Group** 

**UPTRAVI Drug Names UPTRAVI** 

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** Pulmonary arterial hypertension (WHO Group 1) was confirmed by right heart

> catheterization. For new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than or equal to 25 mmHg, 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, and 3) pretreatment pulmonary vascular

resistance is greater than 3 Wood units.

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**VALCHLOR** 

Plan Year

**Prior Authorization Group** 

**Drug Names VALCHLOR** 

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D, adult T-cell

leukemia/lymphoma, primary cutaneous marginal zone lymphoma, primary cutaneous

follicle center lymphoma, lymphomatoid papulosis.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Adult T-cell leukemia/lymphoma: The disease is chronic or smoldering.

Plan Year

Drug Names

VELCADE

Covered Uses

BORTEZOMIB, VELCADE

All FDA-approved indications not otherwise excluded from Part D, systemic light chain

amyloidosis, Waldenstrom's macroglobulinemia/lymphoplasmacytic lymphoma,

multicentric Castleman's disease.

**Exclusion Criteria** 

**Required Medical Information** 

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

**Duration** Plan Year

Other Criteria Coverage under Part D will be denied if coverage is available under Part A or Part B as

the medication is prescribed and dispensed or administered for the individual.

**Prior Authorization Group** 

**Drug Names** 

VENCLEXTA

VENCLEXTA, VENCLEXTA STARTING PACK

Covered Uses All FDA-approved indications not otherwise excluded from Part D, small lymphocytic

lymphoma.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

Exclusion Criteria

Exclusion ontena

**Required Medical Information** 

VENTAVIS

**VENTAVIS** 

All FDA-approved indications not otherwise excluded from Part D.

Pulmonary arterial hypertension (WHO Group 1) was confirmed by right heart

catheterization. For new starts only: 1) pretreatment mean pulmonary arterial pressure is greater than or equal to 25 mmHg, 2) pretreatment pulmonary capillary wedge pressure is less than or equal to 15 mmHg, and 3) pretreatment pulmonary vascular

resistance is greater than 3 Wood units.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Plan Year

Other Criteria Coverage under Part D will be denied if coverage is available under Part A or Part B as

the medication is prescribed and dispensed or administered for the individual.

Prior Authorization Group VERSACLOZ Drug Names VERSACLOZ

Covered Uses All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** The patient is unwilling or unable to take tablets or capsules orally or is at high risk for

non-compliance.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

Prior Authorization Group VERZENIO

Drug Names VERZENIO

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

Coverage Duration

Other Criteria

Plan Year

Prior Authorization Group VIBERZI Drug Names VIBERZI

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

**Age Restrictions** 

**Prescriber Restrictions** 

Coverage Duration Plan Year

Other Criteria

**Drug Names** 

**VIMIZIM** VIMIZIM

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

Diagnosis of mucopolysaccharidosis IVA disease was confirmed by an enzyme assay demonstrating a deficiency of N-acetylgalactosamine 6-sulfatase enzyme activity or by

genetic testing.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**VOSEVI** VOSEVI

All FDA-approved indications not otherwise excluded from Part D.

Decompensated cirrhosis/moderate or severe hepatic impairment (Child Turcotte Pugh

class B or C)

**Required Medical Information** 

Chronic hepatitis C infection confirmed by presence of HCV RNA in the serum prior to starting treatment. Planned treatment regimen, genotype, prior treatment history, presence or absence of cirrhosis (compensated or decompensated [CTP class B or C]), presence or absence of HIV coinfection, presence or absence of resistanceassociated variants where applicable, liver transplantation status if applicable.

Coverage conditions and specific durations of approval will be based on current AASLD

treatment guidelines.

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

12 weeks or Criteria will be applied consistent with current AASLD-IDSA guidance.

Drug Names
Covered Uses

VOTRIENT VOTRIENT

All FDA-approved indications not otherwise excluded from Part D,

dermatofibrosarcoma protuberans, thyroid carcinoma (follicular, papillary, Hurthle cell,

or medullary), uterine sarcoma.

**Exclusion Criteria** 

**Required Medical Information** 

For renal cell carcinoma: the disease is relapsed or unresectable. For soft tissue sarcoma (STS): 1) the patient does not have an adipocytic soft tissue sarcoma, and 2) The patient has one of the following subtypes of STS: a) gastrointestinal stromal tumor (GIST), b) angiosarcoma, c) pleomorphic rhabdomyosarcoma, d) retroperitoneal/intra-abdominal sarcoma, or e) extremity/superficial trunk sarcoma. For follicular, papillary, or Hurthle cell thyroid carcinoma: the disease is unresectable or metastatic. For medullary thyroid carcinoma: the patient has progressive or metastatic disease. For dermatofibrosarcoma protuberans: the disease is metastatic.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

VPRIV VPRIV

All FDA-approved indications not otherwise excluded from Part D.

Diagnosis of Gaucher disease was confirmed by an enzyme assay demonstrating a

deficiency of beta-glucocerebrosidase enzyme activity or by genetic testing.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Drug Names** 

VRAYLAR VRAYLAR

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** 

The patient experienced an inadequate treatment response, intolerance, or

contraindication to one of the following: Latuda, aripiprazole, olanzapine, paliperidone,

quetiapine, risperidone, or ziprasidone.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

XALKORI XALKORI

All FDA-approved indications not otherwise excluded from Part D, non-small cell lung cancer (NSCLC) with high-level MET amplification or MET exon 14 skipping mutation.

inflammatory myofibroblastic tumors (IMT).

**Exclusion Criteria** 

**Required Medical Information** 

For (ALK)-positive NSCLC, patient has recurrent or metastatic disease. For ROS1-positive NSCLC, patient has recurrent or metastatic disease. For NSCLC with high-level MET amplification or MET exon 14 skipping mutation, patient has recurrent or metastatic disease. For IMT, the tumor is ALK-positive and Xalkori is being used as a

single agent.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** XELJANZ

**Drug Names** XELJANZ, XELJANZ XR

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

**Required Medical Information** For moderately to severely active rheumatoid arthritis (new starts only): Patient meets

at least one of the following criteria: 1) Inadequate response, intolerance or

contraindication to methotrexate (MTX), or 2) Inadequate response or intolerance to a prior biologic disease-modifying antirheumatic drug (DMARD) (e.g., adalimumab). For active psoriatic arthritis (new starts only): Patient meets BOTH of the following criteria:

1) Inadequate response to methotrexate (MTX) or other nonbiologic disease-modifying antirheumatic drugs (DMARDs) (e.g., leflunomide, sulfasalazine, etc.) OR a prior biologic DMARD (e.g., adalimumab), and 2) Xeljanz/Xeljanz XR is used in combination

with a nonbiologic DMARD (e.g., methotrexate, leflunomide, sulfasalazine, etc.)

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

Drug Names

**Covered Uses** 

Exclusion Criteria

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

XEOMIN

XEOMIN

All FDA-approved indications not otherwise excluded from Part D.

Cosmetic use.

Plan Year

Prior Authorization Group

**Drug Names** 

Covered Uses

Exclusion Criteria

**Required Medical Information** 

XGEVA XGEVA

All FDA-approved indications not otherwise excluded from Part D.

For hypercalcemia of malignancy, condition is refractory to intravenous (IV)

bisphosphonate therapy (eg. zoledronic acid, pamidronate).

Age Restrictions

**Prescriber Restrictions** 

Coverage Duration Plan Year

Other Criteria Coverage under Part D will be denied if coverage is available under Part A or Part B as

the medication is prescribed and dispensed or administered for the individual.

Drug Names

All FDA-approved indications not otherwise excluded from Part D.

Covered Uses
Exclusion Criteria

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Reduction in risk of overt HE recurrence-6 Months, IBS-D-Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

XOLAIR XOLAIR

**XIFAXAN** 

**XIFAXAN** 

All FDA-approved indications not otherwise excluded from Part D.

For allergic asthma initial therapy: 1)Patient has positive skin test (or blood test) to at least 1 perennial aeroallergen, 2)Patient has baseline IgE level greater than or equal to 30 IU/mL, 3) Patient has inadequate asthma control despite current treatment with both of the following medications at optimized doses: a) Inhaled corticosteroid b) Additional controller (long acting beta2-agonist, leukotriene modifier, or sustained-release theophylline) unless patient has an intolerance or contraindication to such therapies. For allergic asthma continuation therapy only: Patient's asthma control has improved on Xolair treatment since initiation of therapy. Chronic idiopathic urticaria (CIU) initial therapy: 1) Patient has been evaluated for other causes of urticaria, including bradykinin-related angioedema and IL-1-associated urticarial syndromes (autoinflammatory disorders, urticarial vasculitis) 2) Patient has experienced a spontaneous onset of wheals, angioedema, or both, for at least 6 weeks. For CIU continuation therapy: Patient has experienced a response (e.g., improved symptoms) since initiation

of therapy.

Age Restrictions
Prescriber Restrictions

Coverage Duration
Other Criteria

For CIU: 12 years of age or older. For allergic asthma: 6 years of age or older.

Allergic asthma: Plan Year. CIU initial: 6 months. CIU continuation: Plan Year.

**Drug Names** 

**XTANDI XTANDI** 

**Covered Uses** 

**Exclusion Criteria** 

All FDA-approved indications not otherwise excluded from Part D.

**Required Medical Information** 

For non-castration-resistant disease, Xtandi will be used in combination with androgen deprivation therapy to: 1) enhance the effectiveness of radiation therapy, 2) supplement androgen deprivation therapy if the patient experienced inadequate testosterone suppression, OR 3) prevent androgen flare in androgen deprivation therapy naive patients who are at risk of developing symptoms.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

Required Medical Information

**XYREM XYREM** 

All FDA-approved indications not otherwise excluded from Part D.

1) The drug is being prescribed for the treatment of excessive daytime sleepiness in a patient with narcolepsy without cataplexy AND 2) The patient experienced an inadequate treatment response or intolerance to a CNS stimulant drug and a CNS promoting wakefulness drug OR 3) the patient has a contraindication to a CNS stimulant drug or a CNS wakefulness promoting drug (NOTE: Examples of a CNS stimulant drug are amphetamine, dextroamphetamine, or methylphenidate. Example of a CNS wakefulness promoting drug is armodafinil. Coverage of armodafinil or amphetamines or methylphenidates may require prior authorization). OR 4) The drug is being prescribed for the treatment of cataplexy in a patient with narcolepsy

Age Restrictions **Prescriber Restrictions Coverage Duration** Other Criteria

Plan Year

If the request is for the continuation of Xyrem, the patient experienced a decrease in daytime sleepiness with narcolepsy or a decrease in cataplexy episodes with narcolepsy.

YERVOY **Drug Names** YERVOY

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D. CNS metastases

from primary tumor (melanoma), small cell lung cancer

**Exclusion Criteria** 

**Required Medical Information** For CNS metastases from primary tumor (melanoma), member must meet all of the

> following: 1) Yervoy was active against the primary tumor (melanoma) AND 2) the disease is recurrent. For small cell lung cancer, Yervoy will be used on combination

with nivolumab

Age Restrictions

Prescriber Restrictions

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

Required Medical Information

Age Restrictions

Prescriber Restrictions

**Coverage Duration** 

Other Criteria

**ZAVESCA** 

**ZAVESCA** 

All FDA-approved indications not otherwise excluded from Part D.

All FDA-approved indications not otherwise excluded from Part D.

Diagnosis of Gaucher disease was confirmed by an enzyme assay demonstrating a

deficiency of beta-glucocerebrosidase enzyme activity or by genetic testing.

18 years of age or older

Plan Year

**ZEJULA** 

**ZEJULA** 

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

Required Medical Information

Age Restrictions

**Prescriber Restrictions** 

Plan Year **Coverage Duration** 

Other Criteria Treatment is being started or was started no later than 8 weeks after the most recent

platinum-based chemotherapy.

Drug Names Covered Uses ZELBORAF ZELBORAF

All FDA-approved uses not otherwise excluded from Part D, melanoma with BRAF

V600K mutation, non-small cell lung cancer (NSCLC) with BRAF V600E mutation, and

hairy cell leukemia.

**Exclusion Criteria** 

**Required Medical Information** 

For unresectable or metastatic melanoma, the tumor is positive for either BRAF V600E or V600K mutation, and Zelboraf is used as a single agent or in combination with cobimetinib. For NSCLC, the tumor is positive for the BRAF V600E mutation. For refractory hairy cell leukemia, Zelboraf will be used as a single agent for disease progression after non-response to purine analog therapy.

Age Restrictions

**Prescriber Restrictions** 

Coverage Duration
Other Criteria

Plan Year

**Prior Authorization Group** 

Drug Names

Covered Uses

**Exclusion Criteria** 

ZEPATIER ZEPATIER

All FDA-approved indications not otherwise excluded from Part D.

Decompensated cirrhosis/moderate or severe hepatic impairment (Child Turcotte Pugh

class B or C). Liver transplant recipient or awaiting liver transplantation

Required Medical Information

Chronic hepatitis C infection confirmed by presence of HCV RNA in the serum prior to starting treatment. Planned treatment regimen, genotype, prior treatment history, presence or absence of cirrhosis (compensated or decompensated [CTP class B or C]), presence or absence of HIV coinfection, presence or absence of resistance-associated variants (eg, NS5A polymorphisms) where applicable, liver transplantation status if applicable. Coverage conditions and specific durations of approval will be based on current AASLD-IDSA treatment guidelines.

Age Restrictions
Prescriber Restrictions
Coverage Duration

Other Criteria

Criteria will be applied consistent with current AASLD-IDSA guidance.

**ZOLINZA Prior Authorization Group Drug Names ZOLINZA** 

**Covered Uses** All FDA-approved indications not otherwise excluded from Part D, mycosis fungoides,

Sezary syndrome.

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

**ZORBTIVE** 

**ZORBTIVE** 

All FDA-approved indications not otherwise excluded from Part D.

Gastroenterologist or nutritional support specialist

Plan Year

**Prior Authorization Group** 

**Covered Uses** 

**Exclusion Criteria** 

**Required Medical Information** 

**ZURAMPIC ZURAMPIC Drug Names** 

All FDA-approved indications not otherwise excluded from Part D.

The requested drug is being used in combination with a xanthine oxidase inhibitor (i.e.,

allopurinol or febuxostat).

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

Prior Authorization Group
Drug Names

**Covered Uses** 

ZYDELIG ZYDELIG

All FDA-approved indications not otherwise excluded from Part D, relapsed or refractory chronic lymphocytic leukemia (CLL)/ small lymphocytic lymphoma (SLL) as a single agent or in combination with rituximab, refractory, relapsed or progressive follicular lymphoma, primary cutaneous B-cell lymphoma [primary cutaneous marginal

zone lymphoma and follicle center lymphoma], and marginal zone lymphomas [gastric mucosa associated lymphoid tissue (MALT) lymphoma, non-gastric MALT lymphoma.

and splenic marginal zone lymphoma].

**Exclusion Criteria** 

Required Medical Information

For relapsed or refractory CLL/SLL, Zydelig is used as a single agent or in combination with rituximab. For gastric mucosa associated lymphoid tissue (MALT) lymphoma, the disease is recurrent or progressive. For non-gastric MALT and Splenic marginal zone lymphomas, the disease is refractory or progressive.

Age Restrictions

Prescriber Restrictions
Coverage Duration

Other Criteria

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

ZYKADIA ZYKADIA

All FDA-approved indications not otherwise excluded from Part D, anaplastic

lymphoma kinase (ALK)-positive inflammatory myofibroblastic tumor.

**Exclusion Criteria** 

**Required Medical Information** 

For NSCLC, patient meets all of the following: 1) Tumor is ALK-positive, and 2)

Disease is recurrent or metastatic, and 3) Zykadia is prescribed as a single agent. For ALK-positive inflammatory myofibroblastic tumor: Zykadia is prescribed as a single

agent.

**Age Restrictions** 

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year

**Drug Names** 

ZYPREXA RELPREVV ZYPREXA RELPREVV

**Covered Uses** 

All FDA-approved indications not otherwise excluded from Part D.

**Exclusion Criteria** 

Exclusion Cinteria

**Required Medical Information** 

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Tolerability with oral olanzapine has been established.

Plan Year

**Prior Authorization Group** 

**Drug Names** 

**Covered Uses** 

**ZYTIGA - PENDING CMS REVIEW** 

ZYTIGA

All FDA-approved indications not otherwise excluded from Part D and newly diagnosed

metastatic or high-risk locally advanced prostate cancer.

**Exclusion Criteria** 

**Required Medical Information** 

For metastatic castration-resistant prostate cancer: 1) Patient has been previously treated with Xtandi unless the patient has a contraindication to Xtandi therapy and 2) Zytiga will be used in combination with prednisone. For metastatic or locally advanced prostate cancer: 1) Zytiga will be used in combination with prednisone and concurrent androgen-deprivation therapy. Androgen deprivation therapy is not required in patients who have had bilateral orchiectomy, 2) Disease is newly diagnosed and metastatic, node-positive, high-risk locally advanced, or was previously treated with radical surgery

or radiotherapy and is now relapsing with high risk features.

Age Restrictions

**Prescriber Restrictions** 

**Coverage Duration** 

Other Criteria

Plan Year