2020 PDP Basic Prior Authorization Document

Abraxane

Products Affected

• ABRAXANE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For relapsed or refractory melanoma, individual is using as a single agent and Eastern Cooperative Oncology Group (ECOG) performance status of 0-2 following at least one prior therapy. For persistent or recurrent ovarian cancer (epithelial ovarian cancer, fallopian tube cancer, or primary peritoneal cancer) OR Individual is using for the treatment of persistent or recurrent ovarian cancer (epithelial ovarian cancer, fallopian tube cancer, or primary peritoneal cancer) in an individual with confirmed taxane (solvent- based paclitaxel or docetaxel) hypersensitivity. For recurrent, metastatic or high-risk uterine/endometrial cancer in individual with confirmed taxane (solvent-based paclitaxel or docetaxel) hypersensitivity OR individual using for treatment of solid tumors where treatment with taxane is medically appropriate and the individual has confirmed taxane (solvent-based paclitaxel or docetaxel) hypersensitivity. For NSCLC, individual has current ECOG performance status of 0-2 OR individual is suing for using for NSCLC with confirmed taxane (solvent-based paclitaxel or docetaxel) hypersensitivity.
Indications	All Medically-accepted Indications.
Off Label Uses	

Abstral

Products Affected

• ABSTRAL

PA Criteria	Criteria Details
Exclusion Criteria	Treatment of acute or postoperative pain OR treatment of migraine headache pain OR treatment of non-cancer related breakthrough pain.
Required Medical Information	
Age Restrictions	Individual is 18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a diagnosis of active cancer (such as, metastatic or locally invasive cancer for which the individual is currently seeking treatment) with breakthrough cancer pain (provide diagnosis) AND has tried and failed fentanyl citrate lozenge (generic Actiq) AND Individual is already receiving opioid therapy and is TOLERANT to opioid therapy as defined as receiving around the clock medicine consisting of one of the following: At least 60mg morphine per day, OR At least 25mcg/hr transdermal fentanyl/hour, OR At least 30mg of oxycodone daily, OR At least 8mg of oral hydromorphone daily, OR At least 25mg of oral oxymorphone daily, OR An equianalgesic
	dose of another opioid for a week or longer. Individual will also continue around the clock opioids when taking Abstral (fentanyl) for cancer related breakthrough pain.
Indications	continue around the clock opioids when taking Abstral (fentanyl)
Indications Off Label Uses	continue around the clock opioids when taking Abstral (fentanyl) for cancer related breakthrough pain.

Y0114_20_114077_I_012

Actemra

Products Affected

- ACTEMRA
- ACTEMRA ACTPEN

PA Criteria	Criteria Details
Exclusion Criteria	Tuberculosis, or invasive fungal infections or other active serious infections, or history if recurrent infections. Individuals who have not had a tuberculin skin test or Center for Disease Control recommended equivalent to evaluate for latent tuberculosis prior to initiating Actemra (tocilizumab). Using Actemra in combination with other TNF antagonists, IL-1R antagonists, janus kinase inhibitor, anti-cd20 monoclonal antibodies or selective co-stimulation modulators. At initiation of therapy, absolute neutrophil count (ANC) below 2000/mm3, platelet count below 100,000/mm3, or ALT or AST above 1.5 times the upper limit of normal.
Required Medical Information	
Age Restrictions	individual is 18 years of age or older, except for the diagnosis of JIA, PJIA. For JIA, PJIA patient is 2 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For rheumatoid arthritis (RA), Individual has had an inadequate response to ONE non-biological or biologic disease modifying anti- rheumatic drug (DMARD) such as methotrexate (MTX) or a tumor necrosis factor (TNF) antagonist drug AND individual has had a trial and inadequate response or intolerance to: Humira(adalimumab) OR Enbrel(etanercept). For Systemic Juvenile Idiopathic Arthritis (SJIA), agent is being used to reduce signs/symptoms or induce/maintain clinical response. Individual has failed to respond to, is tolerant of, or has a medical contraindication to ONE corticosteroid or nonsteroidal anti-inflammatory drug (NSAID). For Polyarticular Juvenile Idiopathic Arthritis (PJIA), Individual has had inadequate response to, is intolerant of, or has a medical contraindication to ONE Conventional therapy [non-biologic DMARD (such as methotrexate)] AND individual has had a trial and inadequate response or intolerance to: Humira(adalimumab) OR Enbrel(etanercept). For any of the above indications, if the TNF agent (Enbrel(etanercept)/Humira(adalimumab)) tried and failed are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Actemra (tocilizumab) or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis OR mbr has either concomitant clinical condition: (a) Demyelinating disease or (b) Heart failure with documented left ventricular dysfunction. For Multicentric Castleman Disease (MCD), agent is being used as a single agent for tx of relapsed/refractory or progressive MCD. Individual is HIV (human immunodeficiency virus) and HHV-8 (human herpes-8) negative. And individual has no concurrent lymphoma. For Giant Cell Arteritis, agent used in combination with a tapering course of corticosteroids (such as, prednisone) OR being used as a single agent after discontinuing corticosteroids. For chronic Antibody-
	mediated renal transplant rejection with the following are met (Choi 2017): mbr has chronic active antibody-mediated rejection plus donor-specific antibodies and transplant glomerulopathy AND has failed to respond to IVIG plus rituximab therapy with or without plasma exchange.
Indications	All Medically-accepted Indications.
Y0114_20_114077	7 I 012 December 2020

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

Actimmune

Products Affected

• ACTIMMUNE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Actiq

Products Affected

• ACTIQ

PA Criteria	Criteria Details
Exclusion Criteria	Treatment of acute or postoperative pain OR treatment of migraine headache pain OR treatment of non-cancer related breakthrough pain
Required Medical Information	
Age Restrictions	Individual is 16 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a diagnosis of active cancer (such as, metastatic or locally invasive cancer for which the individual is currently seeking treatment) with breakthrough cancer pain (provide diagnosis) AND has tried and failed fentanyl citrate lozenge (generic Actiq) AND Individual is already receiving opioid therapy and is TOLERANT to opioid therapy as defined as receiving around the clock medicine consisting of one of the following: At least 60mg morphine per day, OR At least 25mcg/hr transdermal fentanyl/hour, OR At least 30mg of oxycodone daily, OR At least 8mg of oral hydromorphone daily, OR At least 25mg of oral oxymorphone daily, OR An equianalgesic dose of another opioid for a week or longer. Individual will also continue around the clock opioids when taking Actiq (fentanyl).
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Adcetris

Products Affected

• ADCETRIS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 Year

PA Criteria	Criteria Details
Other Criteria	For diagnosis of Hodgkin lymphoma with either one of the following: For relapsed or refractory disease in a single line of therapy or in combination with bendamustine OR as consolidation therapy after autologous stem cell transplant for ind at high risk of relapse or progression with any of the following: a) Primary refractory Hodgkin lymphoma or b) Relapsed Hodgkin lymphoma with an initial remission duration of less than 12 months or c) Extranodal involvement at the start of pre-transplantation salvage chemotherapy OR as maintenance therapy for 1 yr following high dose therapy and autologous stem cell rescue for relapsed or refractory dx those who are brentuximab vedotin naive and have Deauville score less than 5. For CD30+ non-Hodgkins Lymphoma with either one of the following: cutaneous anaplastic large cell lymphoma OR cutaneous T-cell lymphoma, including mycosis fungoides/Sezary syndrome which is relapsed, refractory or as first line therapy for advanced disease presentation (for example, follicultropic, large cell transformation or extracutaneous disease) OR previously untreated peripheral T-cell lymphoma in combination with cyclophosphamide, doxorubicin and prednisone OR relapsed or refractory disease after at least one prior multi-agent chemotherapy regimen for treatment of ANY if the following: a) systemic anaplastic large cell lymphoma b) T-cell lymphoma estic argue. Stat is symptomatic or characterized by extensive cutaneous lesions OR As a single-agent for adult T-cell leukemia/lymphoma after high dose therapy and autologous stem cell rescue OR Adjuvant systemic therapy and autologous stem cell rescue OR Adjuvant systemic therapy for breast implant-associated anaplastic large cell lymphoma for either of the following: a) residual, localized disease (confined to capsule/implant/breast) following partial excision or capsuletomy OR b) extended disease (stage II-IV).
Indications	All Medically-accepted Indications.
Off Label Uses	

Adcirca

Products Affected

- ADCIRCA
- ALYQ
- tadalafil (pah)

PA Criteria	Criteria Details
Exclusion Criteria	Use in combination with phosphodiesterase-5 (PDE5) inhibitors [such as but not limited to, Cialis (tadalafil)] or use in combination with organic nitrates [such as but not limited to, isosorbide mono/dinitrate or nitroglycerin] or guanylate cyclase stimulators [such as but not limited to, Adempas (riociguat)]. For the treatment of benign prostatic hypertension or erectile dysfunction. Diagnosis of severe hepatic impairment (Child-Pugh Class C), pulmonary veno-occlusive disease (PVOD), severe renal impairment (creatinine clearance less than or equal to 30 mL/min) or on dialysis. Individual has a known degenerative retinal disorder (such as but not limited to, retinitis pigmentosa).
Required Medical Information	Catheterization-proven diagnosis of Pulmonary Arterial Hypertension World Health Organization (WHO) Group 1. Individual has WHO functional class II- IV symptoms.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Addyi

Products Affected

• ADDYI

PA Criteria	Criteria Details
Exclusion Criteria	Use in individuals who cannot abstain from alcohol use. Treatment of HSDD in postmenopausal women OR men. Use for enhancement of sexual performance. Use in individuals with hepatic impairment OR utilizing moderate (such as but not limited to atazanavir, ciprofloxacin, diltiazem, erythromycin, fluconazole, fosamprenavir, verapamil, grapefruit juice) or strong CYP3A4 inhibitors (examples include, but not limited to ketoconazole, itraconazole, posaconazole, clarithromycin, nefazodone, ritonavir, saquinavir, nelfinavir, indinavir, boceprevir, telaprevir, telithromycin, conivaptan) OR concomitant use with CYP3A4 inducers (such as but not limited to, carbamazepine, phenobarbital, phenytoin, rifabutin, rifampin, rifapentine, St. John's Wort).
Required Medical Information	
Age Restrictions	18 years old
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	Individual has Acquired, Generalized Hypoactive Sexual Desire Disorder (HSDD)/Acquired Female Sexual Interest Arousal Disorder (FSIAD) for at least 24 weeks characterized by low sexual desire that causes marked distress or interpersonal difficulty AND it is confirmed that the diagnosis of HSDD/FSIAD is not caused by any of the following: i) A co-existing psychiatric condition, OR ii) A co- existing medical condition that could contribute to sexual dysfunction, OR iii) Problems within a relationship, OR iv) Major life stressor (such as, loss of income, death of a family member), OR v) Effects of a medication or other drug substance.
Indications	All Medically-accepted Indications.
Off Label Uses	

Adempas

Products Affected

• ADEMPAS

PA Criteria	Criteria Details
Exclusion Criteria	Use in combination with nitrates (such as but not limited to, nitroglycerin) or nitric oxide donors (such as but not limited to, amyl nitrite) in any form OR Use in combination with phosphodiesterase (PDE) inhibitors [such as, PDE-5 inhibitors (sildenafil, tadalafil, vardenafil) or nonspecific PDE inhibitors (dipyridamole, theophylline)]. Individual has a diagnosis of severe hepatic impairment (Child-Pugh class C). Individual is on dialysis or has creatinine clearance less than 15 ml/min. Individual has a diagnosis of pulmonary veno-occlusive disease (PVOD), or pulmonary hypertension associated with idiopathic interstitial pneumonias (PH-IIP).
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	Individual has catheterization-proven diagnosis of pulmonary arterial hypertension (PAH), WHO Group 1 AND has WHO functional class II-IV symptoms. Or individual has catheterization- proven diagnosis of chronic thromboembolic pulmonary hypertension (CTEPH), WHO Group 4 AND Individual has WHO functional class II-IV symptoms AND using for one of the following: Persistent or recurrent pulmonary hypertension after at least 180 days following surgical treatment with pulmonary endarterectomey OR Inoperable (via pulmonary endarterectomey) CTEPH.
Indications	All Medically-accepted Indications.
Off Label Uses	

AFINITOR

Products Affected

- AFINITOR
- AFINITOR DISPERZ
- everolimus oral tablet 2.5 mg, 5 mg, 7.5 mg

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Products Affected

 AFREZZA INHALATION POWDER 12 UNIT, 4 & 8 & 12 UNIT, 4 UNIT, 8 UNIT, 90 X 4 UNIT & 90X8 UNIT, 90 X 8 UNIT & 90X12 UNIT

PA Criteria	Criteria Details
Exclusion Criteria	Individuals with a diagnosis of chronic lung disease, such as asthma or chronic obstructive pulmonary disease. Individuals who smoke cigarettes or who recently (within 6 months) quit smoking. Using as a treatment for diabetic ketoacidosis.
Required Medical Information	Individual has had a physical examination including detailed medical history to identify potential lung disease.
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a diagnosis of diabetes mellitus and using for one of the following: 1. For type 1 diabetes, individual will be using concurrently with long-acting insulin. OR 2. For type 2 diabetes, individual has inadequate control, intolerance, or contraindication to at least 2 oral anti-diabetic medications.
Indications	All Medically-accepted Indications.
Off Label Uses	

Ajovy

Products Affected

• AJOVY

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Initial: 3 months, Maintenance: 1 Year
Other Criteria	For Initial requests: (I) Individual has a diagnosis of one of the following: (a) Episodic migraine defined as at least 4 and fewer than 15 migraine days per month and fewer than 15 headache days per month on average during the previous 3 month period OR (b) Chronic migraine defined as headache occurring on 15 or more days per month for more than 3 months, which on at least 8 days per month, has features of a migraine headache (ICHD-3) AND (II) Individual is using for migraine prophylaxis. For Renewal requests: Individual has a reduction in the overall number of migraine days or reduction in number of severe migraine days per month AND Individual has obtained clinical benefit deemed significant by individual or prescriber.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Aldurazyme

Products Affected

ALDURAZYME

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Diagnosis is confirmed by either of the following (Clarke 2016, Lehman 2011): (a) Documented (written or verbal attestation) deficiency in alpha-L-iduronidase enzyme activity as measured in fibroblasts or leukocytes or (b) Documented (written or verbal attestation) alpha-L-iduronidase gene mutation.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For dx of mucopolysaccharidosis I (MPS I) AND Individual has one of the following forms of MPS I: (1) Hurler OR (2) Hurler-Scheie OR (3) Scheie with moderate to severe symptoms manifestations including any of the following: (a) Cardiac valve abnormalities (such as aortic or mitral valve regurgitation, with or without insufficiency or stenosis) or (b) Corneal clouding, open-angle glaucoma, and retinal degeneration, progressive or (c) Craniofacial or growth retardation or (d) Frequent, moderate to severe upper respiratory infections or (e) Hepatosplenomegaly or (f) Hernias (such as hiatal, inguinal, or umbilical) or (g) Neurological symptoms resulting from cervical instability or cervical spinal cord compression or (h) Skeletal and joint involvement, progressive (such as, arthropathy, back pain, joint stiffness, lumbar spondylolisthesis, lumbar spinal compression, osteopenia, or osteoporosis).

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Alecensa

Products Affected

ALECENSA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Alimta

Products Affected

• ALIMTA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Aliqopa

Products Affected

ALIQOPA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	Initial 1 year. Continuation 6 months.
Other Criteria	For initial use in the treatment of follicular lymphoma, Individual has received at least two prior systemic therapies and have not had previous treatment with another PI3-kinase inhibitor previously (for example, idelalisib [Zydelig]). For continued use, there is objective evidence of continuing clinical benefit (for example, complete response, partial response, or stable disease) verified at least every 6 months that is objectively measured.
Indications	All Medically-accepted Indications.
Off Label Uses	

Aloxi

Products Affected

- ALOXI
- palonosetron hcl

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	6 months
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Alpha1-Proteinase Inhibitor

Products Affected

- ARALAST NP
- GLASSIA
- PROLASTIN-C
- ZEMAIRA

PA Criteria	Criteria Details
Exclusion Criteria	Alpha 1 proteinase inhibitors may not be approved for individuals with IgA antibodies.
Required Medical Information	Confirmed alpha-1 antitrypsin level is less than or equal to 11micro-mol/L (approximately equivalent to 80mg/dL measured by radial immunodiffusion or 57 mg/dL measured by nephelometry) (ATS/ERS 2003, Stoller 2017). Individual has clinically evident emphysema and one of the following: Moderate airflow obstruction is evidenced by forced expiratory volume (FEV1) of 30-65 percent of predicted value, prior to initiation of therapy OR a rapid decline in lung function as measured by a change in FEV1 greater than 120 ml/year.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Alunbrig

Products Affected

- ALUNBRIG ORAL TABLET 180 MG, 30 MG, 90 MG
- ALÚNBRIG ORAL TABLET THERAPY PACK

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Amphetamine Line

Products Affected

- ADZENYS ER
- ADZENYS XR-ODT
- amphetamine er
- DYANAVEL XR

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	Individual is 6 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Amphetamine Salts

Products Affected

- amphetamine-dextroamphet er
- amphetamine-dextroamphetamine oral tablet 10 mg, 12.5 mg, 15 mg, 20 mg, 30 mg, 5 mg, 7.5 mg

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	For dx ADHD, 3 years of age or older for immediate release. For Narcolepsy, 6 years of age or older for immediate release
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Amphetamine Salts - B

Products Affected

 ADDERALL ORAL TABLET 10 MG, 12.5 MG, 15 MG, 20 MG, 30 MG, 5 MG, 7.5 MG

ADDERALL XR

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	For dx ADHD, 3 years of age or older for immediate release, 6 years of age or older for extended-release. For Narcolepsy, 6 years of age or older for immediate release
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

MYDAYIS

Ampyra

Products Affected

- AMPYRA
- dalfampridine er

PA Criteria	Criteria Details
Exclusion Criteria	Individual has a history of seizures, OR moderate or severe renal impairment (defined as creatinine clearance less than or equal to 50 mL/min)
Required Medical Information	For renewal, individual achieved and sustained clinically significant improvement in ambulation related functional status.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Initial approval 12 weeks, renewal 1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Anadrol 50

Products Affected

• ANADROL-50

PA Criteria	Criteria Details
Exclusion Criteria	Anadrol 50 may not be used to not replace other supportive measures for anemia such as transfusion, correction of iron, folic acid, B12 or pyridoxine deficiency, antibacterial therapy, or the appropriate use of corticosteroids. Using to enhance athletic ability. Individual has a diagnosis of Carcinoma of the prostate or breast in male individuals or Carcinoma of the breast in females with Hypercalcemia. Individual has a diagnosis of nephrosis (nephrotic phase of nephritis). Individual has a diagnosis of severe hepatic dysfunction.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	6 months
Other Criteria	Individual has a diagnosis of a deficient red cell production- associated anemia, such as but not limited to: acquired aplastic anemia, congenital aplastic anemia, myelofibrosis, or myelotoxic drug-associated hypoplastic anemia.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Apokyn

Products Affected

- APOKYN
- KYNMOBI

PA Criteria	Criteria Details
Exclusion Criteria	Erectile Dysfunction (ED) use
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual is using in conjunction with an antiemetic (excluding 5ht3 antagonist agents).
Indications	All Medically-accepted Indications.
Off Label Uses	

Aranesp

Products Affected

• ARANESP (ALBUMIN FREE)

PA Criteria	Criteria Details
Exclusion Criteria	Sudden loss of response with severe anemia and low reticulocyte count. Anemia in cancer patients receiving myelosuppressive chemotherapy and anemia can be managed by transfusion. Continued use when the hemoglobin level exceeds 11.0 g/dL unless otherwise specified in the criteria. Individuals with uncontrolled HTN. Use beyond 12 weeks in the absence of response in individuals with chronic kidney disease. Use beyond 8 weeks in the absence of response of response or if transfusions are still required in individuals with myelosuppressive chemotherapy agents known to produce anemia. Continued use beyond 6 weeks after therapy with myelosuppressive chemotherapy agents known to produce anemia is completed.
Required Medical Information	Hemoglobin (Hgb) levels are less than 10 g/dL, prior to initiation of therapy (unless otherwise specified) AND the individual iron status reveals, transferrin saturation at least 20% or ferritin at least 80 ng/mL or bone marrow demonstrates adequate iron stores AND For individuals with hypertension. For MDS, endogenous erythropoietin level is less than 500 mU/ml. For tx of anemia due to myelosuppressive chemotherapy, chemo is planned for a minimum of 2 months and the dx is non-myeloid cancer and anticipated outcome is not cure. For anemia associated with CKD ON dialysis use is to achieve and maintain hgb levels within the range of 10.0 to 11.0 g/dL. For anemia associated with CKD NOT ON dialysis use is to achieve and maintain hgb levels of 10.0g/dL.
Age Restrictions	

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Prescriber Restrictions	
Coverage Duration	8wk.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Arcalyst

Products Affected

• ARCALYST

PA Criteria	Criteria Details
Exclusion Criteria	Use in combination with other IL-1 inhibitors, JAK inhibitors, or other biologic drugs (such as IL-6 inhibitors, TNF antagonists, or selective co-stimulation modulators). Tuberculosis, other active serious infections or a history of recurrent infections. Individual has not had a tuberculin skin test (TST) or Centers for Disease Control (CDC)-recommended equivalent to evaluate for latent tuberculosis prior to initiating treatment with rilonacept.
Required Medical Information	
Age Restrictions	Individual is 12 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Arzerra

Products Affected

• ARZERRA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

AUBAGIO

Products Affected

• AUBAGIO

PA Criteria	Criteria Details
Exclusion Criteria	Concurrent use with other immunomodulatory agents (such as Gilenya, tecfidera, Tysabri, Copaxone, Extavia, Plegridy, Rebif, Avonex or Betaseron). Individual has an active acute or chronic infection at the initiation of therapy or has not had a tuberculin skin test (TST) or Centers for Disease Control (CDC)-recommended equivalent to evaluate for latent tuberculosis prior to initiation of therapy.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has been on Aubagio in the past 180 days OR individual has tried therapy with ONE of the following agents: Avonex (interferon beta-1a) OR Plegridy (interferon beta-1a) OR Betaseron (interferon beta-1b) OR Tecfidera (dimethyl fumarate) OR Copaxone/Glatopa (glatiramer).
Indications	All Medically-accepted Indications.
Off Label Uses	

Auryxia

Products Affected

• AURYXIA

PA Criteria	Criteria Details
Exclusion Criteria	Individual has a diagnosis of an iron overload syndrome (for example, hemochromatosis) or has a diagnosis of iron deficiency anemia associated with chronic kidney disease (CKD) stages 3, 4, or 5 and is not on dialysis [Not Part D].
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Austedo

Products Affected

• AUSTEDO

PA Criteria	Criteria Details
Exclusion Criteria	Individual is suicidal or has untreated or inadequately treated depression. Individual has hepatic impairment. Individual is currently utilizing monoamine oxidase inhibitors (MAOIs), reserpine or tetrabenazine.
Required Medical Information	
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For initial requests, Individual has a diagnosis of chorea associated with Huntington's disease. Has a diagnosis of Tardive dyskinesia confirmed by the following DSM-5 AND (a.) At least 60 days of stable (drug, dose) medication exposure (either typical or first generation antipsychotic agents [such as, chlorpromazine, haloperidol, fluphenazine], atypical or second-generation antipsychotic agents [such as, clozapine, risperidone, olanzapine, quetiapine, aripiprazole], or certain dopamine receptor-blocking drugs used in treatment of nausea and gastroparesis [such as, prochlorperazine, promethazine, metoclopramide]) and (b.) Presence of involuntary athetoid or choreiform movements lasting at least 30 days. For continuation requests, Individual has experienced an improvement in symptoms deemed to be clinically significant by the provider (verbal attestation).
Indications	All Medically-accepted Indications.
Y0114_20_114077	_I_012 December 2020

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

AVASTIN

Products Affected

- AVASTIN
- MVASI
- ZIRABEV

	-
PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Ayvakit

Products Affected

• AYVAKIT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Confirmed test results (written or verbal) for individual with a platelet-derived growth factor receptor alpha (PDGFRA) exon 18 mutation, including D842V mutation.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Bafiertam

Products Affected

• BAFIERTAM

PA Criteria	Criteria Details
Exclusion Criteria	Concurrent use with other MS disease modifying agents (including Tecfidera, Vumerity).
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Balversa

Products Affected

 BALVERSA ORAL TABLET 3 MG, 4 MG, 5 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has confirmed (written or verbal attestation) disease susceptible to genetic alterations.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Products Affected

- BANZEL ORAL SUSPENSION
- BANZEL ORAL TABLET 200 MG, 400 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	1 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Baraclude

Products Affected

- BARACLUDE
- entecavir

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has a diagnosis of Chronic Hepatitis B virus (HBV) infection and has not been previously treated with lamivudine (AASLD 2016) OR is using as prophylaxis for hepatitis B reactivation in the setting of immune suppression (AGA 2015) or in combination with hepatitis C direct-acting antiviral therapy (AASLD 2017) OR Individual is a solid organ transplant recipient and using as prophylaxis for hepatitis B reactivation post (AASLD 2018).
Age Restrictions	2 years of age and older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Bavencio

Products Affected

BAVENCIO

PA Criteria	Criteria Details
Exclusion Criteria	Receiving treatment with another PD-1 agent (for example, Opdivo (nivolumab) or Keytruda (pembrolizumab)). Receiving therapy for an autoimmune disease or chronic condition requiring treatment with systemic immunosuppressant.
Required Medical Information	Current Eastern Cooperative Oncology Group (ECOG) performance status of 0-2 for metastatic merkel cell carcinoma and locally advanced or metastatic urothelial carcinoma.
Age Restrictions	12 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For metastatic merkel cell carcinoma, Bavencio is used when individual has not received treatment with another PD-1 (programed death receptor -1) agent (for example, Opdivo or Keytruda) and is not receiving treatment with a systemic immunosuppressant. For locally advanced or metastatic urothelial carcinoma, Bavencio is used as a single agent and individual has not received treatment with another PD-1 agent (for example, Opdivo or Keytruda) and is not receiving therapy for an autoimmune disease or chronic condition requiring treatment with a systemic immunosuppressant and individual meets ONE of the following criteria: has demonstrated disease progression on or after platinum-containing chemotherapy or has demonstrated disease progression within 12 months of receiving neoadjuvant or adjuvant treatment with platinum-containing chemotherapy.
Indications	All Medically-accepted Indications.
Y0114_20_114077	_I_012 December 2020

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

Beleodaq

Products Affected

• BELEODAQ

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Benlysta

Products Affected

• BENLYSTA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For initial treatment, individual has a Clinical diagnosis of SLE per the American College of Rheumatology [ACR] criteria AND Unequivocally positive ANA (anti-nuclear antibody) titer greater than or equal to 1:80 or anti-dsDNA (double stranded DNA antibody) greater than or equal to 30 IU/mL AND SLE is active as documented by a SELENA-SLEDAI score greater than or equal to 6 while on current treatment regimen AND There is no evidence of severe renal disease (proteinuria greater than 6 gm/day, serum creatinine greater than 2.5 mg/dl, or requiring renal dialysis) AND There is no evidence of active central nervous system lupus (e.g. psychosis and seizures) AND SLE remains active while on corticosteroid, antimalarials, and/or immunosuppressants for at least the last 30 days. For continuation of therapy, individual has a clinical diagnosis of SLE per the American College of Rheumatology [ACR] criteria AND documentation of previous improvement in disease activity following treatment with belimumab indicating a therapeutic response AND there is no evidence of severe renal disease AND there is no evidence of active central nervous system lupus.
Age Restrictions	Individual is 18 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Beovu

Products Affected

• BEOVU

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Berinert

Products Affected

• BERINERT

PA Criteria	Criteria Details
Exclusion Criteria	Prophylaxis for HAE attacks.
Required Medical Information	HAE to be confirmed by a C4 level below the lower limit of normal (as defined by the laboratory performing the test) and either a C1 inhibitor antigenic level below the lower limit of normal (as defined by the lab performing test) or a C1 inhibitor functional level below the lower limit of normal (as defined by the lab performing the test).
Age Restrictions	Individual is 5 years or older.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a history of moderate or severe attacks (for example, airway swelling, severe abdominal pain, facial swelling, nausea and vomiting, painful facial distortion) and using Berinert for acute HAE attacks.
Indications	All Medically-accepted Indications.
Off Label Uses	

Blincyto

Products Affected

• BLINCYTO

	1
PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 Year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Bosulif

Products Affected

 BOSULIF ORAL TABLET 100 MG, 400 MG, 500 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Confirmed (written or verbal attestation) BCR-ABL1 positive chronic phase disease. Individual has any of the following confirmed mutations (written or verbal attestation): E255K/V, F317L/V/I/C, F359V/C/I, T315A or Y253H.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has chronic myelogenous leukemia (CML) AND has newly-diagnosed Philadelphia-positive (Ph+) or confirmed(written or verbal attestation) BCR-ABL1 positive chronic phase disease OR is using in combination as primary treatment in lymphoid blast phase or myeloid blast phase disease OR using in post-allogenic HCT therapy for those with prior accelerated or blast phase disease with complete cytogenic response.
Indications	All Medically-accepted Indications.
Off Label Uses	

Botox-Myobloc-Dysport

Products Affected

- BOTOX
- DYSPORT
- MYOBLOC
- XEOMIN

PA Criteria	Criteria Details
Exclusion Criteria	Botulinum toxin is considered cosmetic as a treatment of skin wrinkles or other cosmetic indications and is not approvable.
Required Medical Information	For Cervical Dystonia (spasmodic torticollis) of mod or greater severity when all the following are met: Individual is requesting initial tx AND HX of recurrent clonic and/or tonic involuntary contractions of 1 or more of the following muscles: sternocleidomastoid, splenius, trapezius and/or posterior cervical muscles AND sustained head tilt and/or abnormal posturing with limited range of motion in the neck AND duration of the condition is greater than 6 months. Subsequent injections for the tx of cervical dystonia of mod or greater severity when all the following is met: Individual is requesting subsequent injections AND there is a response to initial tx documented in medical records. An initial 6 month trial For prevention of chronic migraine, PT must have migraine on 15 or more days/month for more than 3 months which on at least 8 days per month has features of a migraine HA (ICHD- 3) AND Individual has had trial of/inadequate response/intolerance to 2 agents for migraine prophylaxis (at least 1 agent in any 2 of the following classes) or has contraindication to all of the following meds (AAN/AHA 2012/2015, Level A and B evidence, ICSI 2013, high quality evidence): 1 of these antidepressants: amitriptyline, venlafaxine OR 1 of these antiepileptics: valproate sod, divalproex sod, topiramate, gabapentin. Continuing tx medically nec when individual has completed an initial 6 month trial and has a reduction in overall number of migraine days or reduction in number of severe migraine days/month AND individual has obtained clinical benefit deemed significant by individual or prescriber.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year, chronic migraine 6 months
Other Criteria	Treatment of primary hyperhidrosis. Treatment of secondary hyperhidrosis. Treatment of significant drooling in patients who are unable to tolerate scopolamine. Treatment neurogenic overactive bladder (also referred to as detrusor overactivity or detrusor sphincter dyssynergia) that is inadequately controlled with anticholinergic therapy. Treatment of idiopathic overactive bladder in adults who are unresponsive to or intolerant of a trial of anticholinergic therapy. Treatment of Hirschsprung disease and associated functional obstruction caused by the inability of the internal anal sphincter to relax after prior surgical treatment.
Indications	All Medically-accepted Indications.
Off Label Uses	

Braftovi

Products Affected

 BRAFTOVI ORAL CAPSULE 50 MG, 75 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Written or verbal attestation has been provided for either BRAF V600E or V600K genetic mutation.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Briviact

Products Affected

- BRIVIACT INTRAVENOUS
- BRIVIACT ORAL SOLUTION
- BRIVIACT ORAL TABLET 10 MG, 100 MG, 25 MG, 50 MG, 75 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Brukinsa

Products Affected

• BRUKINSA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has no prior BTK inhibitor usage.
Age Restrictions	18 years of age and older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Buphenyl

Products Affected

- BUPHENYL
- sodium phenylbutyrate

PA Criteria	Criteria Details
Exclusion Criteria	Management of acute hyperammonemia
Required Medical Information	Using as adjunctive therapy for chronic management of hyperammonemia
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Cabometyx

Products Affected

• CABOMETYX

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Calquence

Products Affected

CALQUENCE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Caplyta

Products Affected

• CAPLYTA

PA Criteria	Criteria Details
Exclusion Criteria	Individual is currently using a CYP3A4 inducer (such as carbamazepine, phenytoin, rifampin, St. John's wort, bosentan, efavirenz, etravirine, modafinil, nafcillin, aprepitant, armodafinil, pioglitazone, prednisone) and cannot discontinue the medication OR Individual is using a moderate (such as amprenavir, ciprofloxacin, cyclosporine, diltiazem, erythromycin, fluconazole, fluvoxamine, verapamil) or strong (such as, clarithromycin, grapefruit juice, itraconazole, voriconazole, nefazodone, ritonavir, nelfinavir) CYP3A4 inhibitor and cannot discontinue the medication OR Individual has moderate (Child-Pugh Class B) or severe hepatic impairment (Child-Pugh Class C).
Required Medical Information	
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Caprelsa

Products Affected

CAPRELSA ORAL TABLET 100 MG, 300 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Carbaglu

Products Affected

• CARBAGLU

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Cayston

Products Affected

CAYSTON

PA Criteria	Criteria Details
Exclusion Criteria	Individual has a forced expiratory volume in 1 second (FEV1) of less than 25% or greater than 75% of predicted
Required Medical Information	
Age Restrictions	7 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Celebrex

Products Affected

- CELEBREX
- celecoxib

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has failed to respond to, is intolerant of, or has a medical contraindication to ONE prescription Non-Steroidal Anti-Inflammatory Drugs (NSAID) or salicylates
Indications	All Medically-accepted Indications.
Off Label Uses	

Cequa

Products Affected

• CEQUA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has an abnormal result or response to one or more of the following dry eye disease diagnostic/assessment methods (AAO, 2018): 1)Tear break-up time (less than 10 seconds) or 2) Ocular surface dye staining using fluorescein, rose bengal, or lissamine green dyes or 3) Schirmer test (aqueous tear production of less than or equal to 10 mm of strip wetting in 5 minutes) or 4) Fluorescein clearance test/tear function index or 5) Tear osmolarity (indicating tear film instability) or 6) Tear lactoferrin concentrations in the lacrimal gland (decreased) or 7) Matrix metalloproteinase-9 (MMP-9) test.
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual using for moderate to severe dry eye AND has had a trial and inadequate response or intolerance to Xiidra OR has a known hypersensitivity to any ingredient to Xiidra (preferred agent) which is not also present in the requested non-preferred agent
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Cerdelga

Products Affected

• CERDELGA

PA Criteria	Criteria Details
Exclusion Criteria	Use of glucosylceramide synthase inhibitor in conjunction with one another or in conjunction with a substrate reduction therapy (SRT) agent or enzyme replacement therapy (ERT) agent. Use in an ultra- rapid metabolizers of CYP2D6. Individual has moderate or severe renal impairment or end-stage renal disease (ESRD). Individual has mild, moderate or severe hepatic impairment or cirrhosis OR individual has pre-existing cardiac disease or long QT syndrome.
Required Medical Information	Presence of type 1 Gaucher disease is confirmed by either of the following: Glucocerebrosidase activity in the white blood cells or skin fibroblasts less than or equal to 30% of normal activity, OR genotype testing indicating mutation of two alleles of the glucocerebrosidase genome. And patient has clinically significant manifestations of Type 1 gauchers disease including any of the following: (A) skeletal disease (demonstrated by ANY of the following: avascular necrosis, erlenmeyer flask deformity, lytic disease, marrow infiltration, osteopenia, osteosclerosis, pathological fracture, joint deterioration or replacement) OR (B) individual presents with at least 2 of the following: clinically significant hepatomegaly as confirmed by medical imaging [such as but not limited to, volumetric magnetic resonance imaging (MRI)], clinically significant splenomegaly as confirmed by medical imaging [such as but not limited to, volumetric (MRI)], hgb less than or equal to 11.5 grams per dl for females or 12.5 grams per dl for males or 1 gram per dl below lower limit for normal for age and sex, platelet counts less than or equal to 120,000 mm3 OR (C) individual is a CYP2D6 extensive metabolizer (EM), intermediate metabolizer (IM), or poor metabolizer (PM) as confirmed by a FDA-approved genotype test.
Age Restrictions	Individual is 18 years of age or older.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Chantix

Products Affected

- CHANTIX CONTINUING MONTH PAK
- CHANTIX ORAL TABLET 0.5 MG, 1 MG
- CHANTIX STARTING MONTH PAK

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	3 months
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Chenodal

Products Affected

CHENODAL

PA Criteria	Criteria Details
Exclusion Criteria	Individual has calcified (radiopaque) or radiolucent bile pigment stones OR has preexisting hepatic impairment OR has known hepatocyte dysfunction or bile ductal abnormalities (such as but not limited to intrahepatic cholestasis, primary biliary cirrhosis, or sclerosing cholangitis) OR has a gallbladder confirmed as nonvisualizing after two consecutive single doses of dye OR has gallstone complications or compelling reasons for gallbladder surgery (such as but not limited to unremitting acute cholecystitis, cholangitis, biliary obstruction, gallstone pancreatitis, or biliary gastrointestinal fistula).
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	2 years. Requests to continue therapy beyond 24 months (2 years) should not be approved.
Other Criteria	Individual is using for gallstone dissolution AND has a well- opacifying gallbladder with radiolucent stones AND has an increased surgical risk due to systemic disease or advanced age. For continuation, Repeat imaging studies show partial dissolution of gallstone(s).
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Cholbam

Products Affected

• CHOLBAM

PA Criteria	Criteria Details
Exclusion Criteria	Used to treat extrahepatic manifestations (such as but not limited to neurologic symptoms) of SED-associated-BASDs or PDs.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Initial therapy: 3 months. Continuation therapy: 1 year
Other Criteria	For initial therapy: (A) Diagnosis of bile acid synthesis disorders (BASDs) due to single enzyme defects (SEDs) including but not limited to 3 beta-hydroxy-delta 5-C27-steroid oxidotrductase defects OR (B) Diagnosis of peroximal disorders (PDs) including but not limited to Zellweger spectrum disorders AND (C) Individual has one of the following: (a) Manifestations of liver disease (for example, jaundice, hepatomegaly) (b) steatorrhea (c) Complications from decreased fat soluble vitamin (such as but not limited to vitamin D and K) absorption (for example, rickets, hypocalcemia, bleeding). For maintenance therapy: Meets the initial request criteria AND has had a clinical improvement (symptoms, lab values) in liver function and/or cholestasis AND has not developed a complete biliary obstruction.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Chorionic Gonadotropin

Products Affected

- chorionic gonadotropin
- NOVAREL
- PREGNYL

PA Criteria	Criteria Details
Exclusion Criteria	Use in the following: Infertility treatments (Including use with IVF, ART), Obesity, Weight loss, Stimulation of spermatogenesis in males, Treatment of anovulation in females with infertility, Ovulation induction in females.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	6 months
Other Criteria	Individual is using for Pre-pubertal cryptorchidism not caused by anatomical obstruction in males OR Hypogonadotropic hypogonadism from pituitary deficiency in males.
Indications	All Medically-accepted Indications.
Off Label Uses	

Cialis BPH

Products Affected

- CIALIS ORAL TABLET 2.5 MG, 5 MG
- tadalafil oral tablet 2.5 mg, 5 mg

PA Criteria	Criteria Details
Exclusion Criteria	Erectile dysfunction. Currently on nitrate therapy.
Required Medical Information	Individual has a diagnosis of benign prostatic hyperplasia (BPH) [with or without ED]
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Cimzia

Products Affected

- CIMZIA
- CIMZIA PREFILLED
- CIMZIA STARTER KIT

PA Criteria	Criteria Details
Exclusion Criteria	Tuberculosis, invasive fungal infection, other active serious infections, or a history of recurrent infections prior to initiating Cimzia (certolizumab pegol). Individuals who have not had a tuberculin skin test or Center for Disease Control recommended equivalent to evaluate for latent tuberculosis prior to initiating Cimzia (certolizumab pegol). Using Cimzia in combination with other TNF antagonists, non-TNF immunomodulatory drugs: abatacept, anakinra, natalizumab, tofacitinib or rituximab.
Required Medical Information	Individual has chronic moderate to severe (that is, extensive or disabling) plaque psoriasis with either of the following: Individual has greater than 5% of body surface area with plaque psoriasis OR less than or equal to 5% of body surface area with plaque psoriasis involving sensitive areas or areas that would significantly impact daily function (such as palms, soles of feet, head/neck, or genitalia).
Age Restrictions	Individual is 18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For moderate to severe Crohn's Disease, Individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional therapy (e.g. 5-ASA products, systemic corticosteroids, or immunosuppressants) AND Individual has had trial and inadequate response or is intolerant to Humira (adalimumab). For moderate to severe Rheumatoid Arthritis, Individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE nonbiologic DMARD AND Individual has had a trial and an inadequate response or is intolerant to Humira (adalimumab) OR Enbrel (etanercept). For moderate to severe Psoriatic Arthritis, Individual has had an inadequate response or is intolerant to Humira (adalimumab) OR Enbrel (etanercept). For moderate to severe Psoriatic Arthritis, Individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional therapy (such as nonbiologic DMARDs) AND has had a trial and an inadequate response or is intolerant to Humira (adalimumab) OR Enbrel (etanercept). For moderate to severe Ankylosing Spondylitis (AS), Individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional therapy (such as NSAIDs or non-biologic DMARDs) AND has had a trial and an inadequate response or is intolerant to Humira (adalimumab) OR Enbrel (etanercept). For plaque psoriasis individual has had an inadequate response or is intolerant of, or has a actiretin, cyclosporine, or methotrexate) AND has had a trial of and inadequate response or intolerance to Humira (adalimumab) OR Enbrel (etanercept). For any of the above indications, if the TNF agent (Enbrel/Humira) tried and failed are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Cimzia or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infectio
	non-radiographic axial spondyloarthritis, individual has had an inadequate response to, or has a contraindication to conventional therapy [such as NSAID or nonbiologic such as sulfasalazine)].
Indications	All Medically-accepted Indications.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

Cinqair

Products Affected

• CINQAIR

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has blood eosinophil counts (in the absence of other potential causes of eosinophilia, including hypereosinophilic syndromes, neoplastic disease and known or suspected parasitic infection) greater than or equal to 400 cells/microliter at initiation of therapy. The individual has pretreatment forced expiratory volume in one second (FEV1) less than 80% predicted AND FEV1 reversibility of at least 12% and 200mL after albuterol administration.
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For initial requests, individual must have a diagnosis of eosinophilic asthma AND individual has had a 3 month trial and inadequate response or intolerance to combination controller therapy (high dose inhaled corticosteroids plus long acting beta2-agonist, leukotriene modifiers, theophylline or oral corticosteroids) (ERS/ATS 2013) AND individual has experienced 2 or more asthma exacerbations in the prior 12 months requiring use of a systemic corticosteroid or temporary increase in the individual's usual maintenance dosage of oral corticosteroids (ERS/ATS 2013). For Maintenance Therapy: Treatment has resulted in clinical improvement as confirmed one or more of the following: i) Decreased utilization of rescue medications OR ii) A decreased frequency of exacerbation (defined as worsening of asthma that requires increase in inhaled corticosteroid dose or treatment with systemic corticosteroid) OR iii) An increase in percent predicted FEV1 from pretreatment baseline OR iv) A reduction in reported asthma-related symptoms, such as, but not limited to wheezing, shortness of breath, coughing, fatigue, sleep disturbance or asthmatic symptoms upon awakening
Indications	All Medically-accepted Indications.
Off Label Uses	

Cinryze

Products Affected

• CINRYZE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	HAE to be confirmed by a C4 level below the lower limit of normal (as defined by the laboratory performing the test) and ANY of the following: 1. C1 inhibitor antigenic level below the lower limit of normal (as defined by the lab performing test). 2. C1 inhibitor functional level below the lower limit of normal (as defined by the lab performing the test) Or 3. The presence of a known HAE-causing C1-INH mutation.
Age Restrictions	6 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a history of moderate or severe attacks and is using Cinryze as prophylaxis for short-term use prior to surgery, dental procedures or intubation OR for long-term prophylaxis and member has failed, or is intolerant to, or has contraindication to 17-alpha- alkylated androgens or antifibrinolytic agents.
Indications	All Medically-accepted Indications.
Off Label Uses	

Cometriq

Products Affected

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

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Copaxone

Products Affected

COPAXONE SUBCUTANEOUS SOLUTION
 PREFILLED SYRINGE 20 MG/ML, 40
 MG/ML

prefilled syringe 20 mg/ml, 40 mg/ml

- glatopa subcutaneous solution prefilled syringe 20 mg/ml, 40 mg/ml
- glatiramer acetate subcutaneous solution

PA Criteria	Criteria Details
Exclusion Criteria	Individual with primary progressive MS (PPMS). Individual with non-active secondary progressive MS (SPMS). Concurrent use with other MS Disease modifying agents (such as, Aubagio, Gilenya, Tecfidera, Tysabri, Lemtrada, Ocrevus, Copaxone/Glatopa, Extavia, Rebif, Avonex, Plegridy or Betaseron)
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Copiktra

Products Affected

• COPIKTRA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Corlanor

Products Affected

- CORLANOR ORAL SOLUTION
- CORLANOR ORAL TABLET

DA Critoria	Criteria Details
PA Criteria	
Exclusion Criteria	Individual has a heart rate maintained exclusively by a pacemaker. Individual has severe hypotension (blood pressure less than 90/50 mmgHg). Individual has severe hepatic impairment (Child-Pugh Class C).
Required Medical Information	
Age Restrictions	For NYHA Class II, II, or IV due to DCM, age 6 months or older.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	(A) Individual is using for the treatment of New York Heart Association (NYHA) class II, III or IV heart failure symptoms AND has a left ventricular ejection fraction less than or equal to 35% AND will be utilizing in combination with a beta-blocker OR has a contraindication or intolerance to beta-blocker therapy AND is in normal sinus rhythm AND If initiating treatment with Corlanor, individual has a resting heart rate greater than or equal to 70 beats per minute. OR (B) Individual is using for the treatment of New York Heart Association (NYHA) class II, III, or IV heart failure symptoms due to dilated cardiomyopathy AND has a left ventricular ejection fraction less than or equal to 45% AND is in normal sinus rhythm AND If initiating treatment with Corlanor, individual has an elevated resting heart rate.
Indications	All Medically-accepted Indications.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

Cosentyx

Products Affected

- COSENTYX
- COSENTYX (300 MG DOSE)
- COSENTYX SENSOREADY (300 MG)
- COSENTYX SENSOREADY PEN

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has moderate to severe plaque psoriasis with either of the following: Individual has greater than 5% of body surface area with plaque psoriasis OR Less than or equal to 5% of body surface area with plaque psoriasis involving sensitive areas or areas that would significantly impact daily function (such as palms, soles of feet, head/neck, or genitalia) OR Individual is using for the treatment of non-radiographic axial spondyloarthritis (nr-axSpA) with objective signs of inflammation.
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For moderate to severe Ankylosing Spondylitis (AS), individual had an inadequate response to/intolerant of or has a medical contraindication to ONE conventional therapy [such as, NSAIDs or nonbiologic DMARDS such as sulfasalazine or a tumor necrosis factor (TNF) antagonist] AND Individual has had a trial of/inadequate response or is intolerant to Enbrel (etanercept) OR Humira (adalimumab). For moderate to severe plaque psoriasis (Ps), individual had an inadequate response to/intolerant of or has contraindication to phototherapy or other systemic therapy (such as acitretin, cyclosporine or methotrexate) AND Individual has had a trial of/inadequate response or is intolerant to Enbrel (etanercept) OR Humira (adalimumab). For moderate to severe Psoriatic Arthritis (PsA), individual has had an inadequate response to/intolerant of or has a medical contraindication to conventional therapy [nonbiologic DMARDs (such as methotrexate, sulfasalazine or leflunomide)] or TNF antagonist (AAD 2011) AND Individual has had a trial of and an inadequate response or is intolerant to Enbrel (etanercept) OR Humira (adalimumab). For any of the above indications, if the TNF agent (Enbrel(etanercept)/Humira (adalimumab)) tried and failed are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Cosentyx or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis OR individual has either concomitant clinical condition: (a) Demyelinating disease or (b) Heart failure with documented left ventricular dysfunction, Cosentyx may be allowed without trial of preferred TNF agents (Enbrel/Humira).
Indications	All Medically-accepted Indications.
Off Label Uses	

Cotellic

Products Affected

COTELLIC

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Diagnosis of unresectable or metastatic melanoma with confirmed (written or verbal attestation) BRAF V600E or V600K mutation.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual is using Cotellic (cobimetinib) in combination with Zelboraf (vemurafenib).
Indications	All Medically-accepted Indications.
Off Label Uses	

Cresemba

Products Affected

• CRESEMBA

PA Criteria	Criteria Details
Exclusion Criteria	Individual has diagnosis or history of familial short QT syndrome. Use in combination with strong CYP3A4 inhibitors (such as but not limited to ketoconazole) OR strong CYP3A4 inducers (such as but not limited to rifampin).
Required Medical Information	
Age Restrictions	18 years and older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual initiated treatment in an inpatient setting and requires continued treatment of invasive aspergilliosis or mucormycosis in an outpatient setting. For invasive aspergillosis individual has an inadequate response/intolerance to or contraindication to voriconazole or liposomal amphotericin B (ATS 2011, IDSA 2008). For invasive mucormycosis individual has had an inadequate response/intolerance to or contraindication to amphotericin B (ATS 2001).
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Crinone

Products Affected

CRINONE

PA Criteria	Criteria Details
Exclusion Criteria	Progesterone supplementation or replacement as part of an Assisted Reproductive Technology ("ART") treatment for infertile women with progesterone deficiency, Progesterone supplementation/deficiency.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Cyramza

Products Affected

• CYRAMZA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For urothelial cancer, an Eastern Cooperative Oncology Group (ECOG) performance status of 0 or 1.
Age Restrictions	For urothelial cancer, 18 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 Year
Other Criteria	For locally advanced, unresectable or metastatic urothelial cancer originating from bladder, urethra, ureter or renal pelvis and using in combination with docetaxel AND disease has progressed after platinum-containing chemotherapy (cisplatin or carboplatin) AND individual has received treatment with no more than one immune checkpoint inhibitor (such as, atezolizumab, avelumab, durvalumab, nivolumab or pembrolizumab) AND has received treatment with no more than one prior systemic chemotherapy regimen in the relapsed or metastatic setting AND individual has not received prior systemic taxane therapy in any setting (neoadjuvant, adjuvant or metastatic).
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

D.H.E Inj

Products Affected

- D.H.E. 45
- *dihydroergotamine mesylate injection*

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For migraine attacks with aura in individuals meeting the following International Headache Society (IHS) diagnostic criteria (must meet criteria A-D): A) At least 2 or more headache attacks AND B) Aura consisting of at least 1 of the following fully reversible aura sx: 1. visual symptoms (such as, flickering lights, spots or lines) OR 2. Sensory symptoms (for example, pins and needles, numbness) OR 3. Speech and/or language (for example, aphasia) OR 4. Motor (for example, weakness) OR 5. Brainstem (for example, ataxia or vertigo) OR 6. Retinal (for example, blindness) AND C) At least 3 of the following characteristics: a) At least 1 aura sx develops gradually over 5 or more minutes or b) 2 or more aura sx occur in succession or c) Each individual aura lasts 5 to 60 minutes or d) At least 1 aura sx is unilateral or e) At least 1 aura sx is positive (scintillations and pins and needles are examples of positive sx of aura) or f) The aura is accompanied or followed within 60 minutes, by headache AND D) Individual's headache is not attributed to another headache disorder (for example, transient ischemic attack). For migraine attacks without aura in adults meeting the following IHS diagnostic criteria: 1) At least 5 or more headache attacks AND 2) Headaches lasting 4-72 hrs (untreated or unsuccessfully treated) AND 3) Headache has at least 2 or more of the following: i) Unilateral location ii) Pulsating quality iii)Moderate or severe pain intensity iv) Aggravation by or causing avoidance of routine physical activity (such as, walking or climbing stairs) AND 4) Individual's headache is accompanied by 1 or more of the following: i) Nausea, vomiting or both ii) Photophobia or phonophobia AND 5) Individual's headache is not attributed to another headache disorder (for example, transient ischemic attack). For cluster headache episodes in adults meeting the following IHS diagnostic criteria: A) At least 5 or more attacks B) Severe or very severe unilateral orbital, supraorbital and/or temporal pain lasting 15 to 180 minute

PA Criteria	Criteria Details
	One or more of the following sx or signs, ipsilateral to the headache: (i) conjunctival injection and/or lacrimation (ii) nasal congestion and/or rhinorrhea (iii) eyelid edema (iv) forehead and facial sweating (v) forehead and facial flushing (vi) miosis and ptosis OR 2. A sense of restlessness or agitation AND D) Attacks have a frequency from one every other day to 8 per day for more than half of the time the disorder is active AND E) Individual's headache is not attributed to another headache disorder. DHE may also be may be approved: For Status migrainosus or rebound withdrawal type of headaches OR As alternative to narcotic therapy for severe migraine or cluster headaches OR individual is unresponsive to prior use of triptans for severe migraine or cluster headache.
Indications	All Medically-accepted Indications.
Off Label Uses	

DAKLINZA

Products Affected

• DAKLINZA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Documentation is provided for a diagnosis of chronic hepatitis C (CHC) infection, which includes genotype and a positive HCV RNA result (AASLD/IDSA 2017, CDC 2013) AND Individual has received baseline evaluation for liver fibrosis to guide appropriate therapy AND Individual does not have a short life expectancy (less than 12 months owing to non-liver related comorbid conditions) that cannot be remediated by treating HCV, by transplantation or other directed therapy (AASLD/IDSA 2016).
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	Criteria will be applied consistent with current AASLD/IDSA guidance.

PA Criteria	Criteria Details
Other Criteria	Criteria will be applied consistent with current AASLD/IDSA guidance. For GT 1, Individual has had a prior trial and inadequate response to Harvoni (sofosbuvir/ledipasvir) OR individual is currently on and completing a course of therapy with the requested regimen OR Individual has a documented hypersensitivity, as manifested by a severe allergic reaction, to any ingredient in Harvoni (ledipasvir/sofosbuvir) which is not also in Daklinza OR Individual is concurrently using an agent that cannot be substituted with another agent or temporarily discontinued and is contraindicated or not recommended for concomitant use with Harvoni (sofosbuvir/ledipasvir). For GT 4, individual has had a prior trial and inadequate response to Harvoni (sofosbuvir/ledipasvir) or Epclusa (sofosbuvir/velpatasvir). OR individual is currently on and completing a course of therapy with the requested regimen OR Individual has a documented hypersensitivity, as manifested by a severe allergic reaction, to any ingredient in Harvoni(sofosbuvir/ledipasvir) or Epclusa (sofosbuvir/velpatasvir) which is not also in Daklinza OR Individual is concurrently using an agent that cannot be substituted with another agent or temporarily discontinued and is contraindicated or not recommended for concomitant use with Harvoni (sofosbuvir/ledipasvir) or Epclusa (sofosbuvir/velpatasvir).
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Daliresp

Products Affected

• DALIRESP

PA Criteria	Criteria Details
Exclusion Criteria	Individual is using to treat acute bronchospasm OR moderate (Child-Pugh B) or severe (Child-Pugh C) hepatic impairment OR using concomitantly with strong cytochrome P450 enzyme inducer (such as but not limited to phenobarbital, carbamazepine or phenytoin)
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual is currently or will be concomitantly using with a long- acting bronchodilator.
Indications	All Medically-accepted Indications.
Off Label Uses	

Darzalex

Products Affected

• DARZALEX FASPRO

mg/20ml

100 MG/5ML*darzalex intravenous solution 400*

DARZALEX INTRAVENOUS SOLUTION

PA Criteria	Criteria Details
Exclusion Criteria	Has received treatment with daratumumab or another anti-CD38 agent
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Daurismo

Products Affected

 DAURISMO ORAL TABLET 100 MG, 25 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	Individual is 75 years old or older OR has comorbidities that preclude use of intensive induction chemotherapy.
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Desoxyn

Products Affected

- DESOXYN
- methamphetamine hcl

PA Criteria	Criteria Details
Exclusion Criteria	Adjunct treatment of exogenous obesity/weight loss.
Required Medical Information	
Age Restrictions	Individual is 6 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a diagnosis of attention deficit hyperactivity disorder (ADHD) AND has had an appropriate trial of one of the following: (a) methylphenidate containing agent OR (b) amphetamine containing agent (such as, amphetamine/dextroamphetamine, lisdexamfetamine, or dextroamphetamine).
Indications	All Medically-accepted Indications.
Off Label Uses	

Dificid

Products Affected

• DIFICID

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has had a trial and inadequate response or intolerance to or has a contraindication to a course of oral vancomycin.
Age Restrictions	18 years of age or older.
Prescriber Restrictions	
Coverage Duration	30 days
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Doptelet

Products Affected

• DOPTELET

PA Criteria	Criteria Details
Exclusion Criteria	Doptelet should not be administered to individuals with chronic liver disease in an attempt to normalize platelet counts.
Required Medical Information	
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Doxil

Products Affected

- DOXIL
- doxorubicin hcl liposomal

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Products Affected

- alogliptin benzoate oral tablet 12.5 mg, 25 mg, 6.25 mg
- alogliptin-metformin hcl ٠
- alogliptin-pioglitazone oral tablet 12.5-15 mg, 12.5-30 mg, 12.5-45 mg, 25-15 mg, • ONGLYZA ORAL TABLET 2.5 MG, 5 MG 25-30 mg, 25-45 mg
- KAZANO
- KOMBIGLYZE XR ORAL TABLET

EXTENDED RELEASE 24 HOUR 2.5-1000 MG, 5-1000 MG, 5-500 MG

- NESINA ORAL TABLET 12.5 MG, 25 MG, 6.25 MG
- OSENI ORAL TABLET 12.5-15 MG, 12.5-30 MG, 12.5-45 MG, 25-15 MG, 25-30 MG, 25-45 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has had a trial and inadequate response or intolerance to metformin OR Individual has a contraindication to metformin therapy [such as but not limited to, renal insufficiency (eGFR less than 45 mL/min/1.73 m2)] AND Individual has had a trial and inadequate response or intolerance to ONE of the following: Januvia (sitagliptin), Tradjenta (linagliptin), Janumet (sitagliptin/metformin), Janumet XR (sitagliptin/metformin), Jentadueto (linagliptin/metformin), OR Jentadueto XR (linagliptin/metformin).
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Duavee

Products Affected

• DUAVEE

PA Criteria	Criteria Details
Exclusion Criteria	May not be approved for the following: (1) If individual had a hysterectomy (2) Individual has undiagnosed abnormal uterine bleeding (3) Individual has known, suspected, or past history of breast cancer (4) Individual has known or suspected estrogen- dependent neoplasia (5) Individual has active or past history of venous thromboembolism (6) Individual has active or past history of arterial thromboembolism (7) Individual has known hepatic impairment or disease OR (8) Individual has known protein C, protein S, or antithrombin deficiency or other known thrombophilic disorders.
Required Medical Information	
Age Restrictions	Age 18 through age 75
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual is using for ONE of the following: Treatment of moderate to severe vasomotor symptoms associated with menopause OR Individual is using for prevention of postmenopausal osteoporosis AND is using solely for prevention of osteoporosis and has had a trial of and inadequate response or intolerance or has a contraindication to non-estrogen agents for osteoporosis.
Indications	All Medically-accepted Indications.
Off Label Uses	
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Y0114_20_114077_I_012

Duexis

Products Affected

• DUEXIS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual has had a trial and inadequate response or intolerance to one (1) oral generic prescription Non-Steroidal Anti-Inflammatory Drugs (NSAID) AND has had a trial and inadequate response or intolerance to one (1) of the following (Lanza 2009): (a) Preferred proton pump inhibitor (PPI) OR (b) Generic misoprostol OR (c) Generic histamine-2 receptor antagonist (H2RA) AND Individual has had an adequate response (pain relief and appropriate gastro protection) with a trial of ibuprofen and famotidine used at the same time AND Documentation has been provided for why the combination agent is clinically necessary and not for convenience.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Duobrii

Products Affected

• DUOBRII

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual has a diagnosis of plaque psoriasis AND Documentation (verbal or written) has been provided for why the combination agent is clinically necessary and not for convenience.
Indications	All Medically-accepted Indications.
Off Label Uses	

Duopa

Products Affected

• DUOPA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For advanced Parkinsons disease with complicated motor fluctuations that have not been adequately controlled with optimal medical therapy with any TWO of the following: Oral levodopa- carbidopa, a Dopamine agonist [such as, but limited to Apokyn (apomorphine), Mirapex (pramipexole), Requip (ropinirole) and Neupro (rotigotine)], a catechol-0-methyl transferase (COMT) inhibitor [such as, but not limited to Comtan (entacapone) and Tasmar (tolcapone)], or a monoamine oxidase B (MAO)-B inhibitor [such as, but not limited to Eldepryl (selegiline), and Azilect (rasagiline)].
Indications	All Medically-accepted Indications.
Off Label Uses	

Dupixent

Products Affected

DUPIXENT SUBCUTANEOUS SOLUTION
 PEN-INJECTOR

300 MG/2ML

 DUPIXENT SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 200 MG/1.14ML,

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	A)Chronic atopic dermatitis that has been present for 3 years or more AND B) failure of topical pharmacological therapy as indicated by one or more of the following: 1) Daily treatment of topical corticosteroids of medium to higher potency for the maximum treatment period indicated in the product prescribing information has failed to achieve and maintain remission of low or mild disease activity state OR 2) topical calcineurin inhibitors (for example, Elidel, Protopic) if topical corticosteroids are not indicated for the maximum treatment period indicated in the product prescribing information has failed to achieve and maintain remission of low or mild disease activity state OR 3) Topical treatment is medically inadvisable as defined by treatments which have side effects or safety concerns which outweigh potential treatment benefits as evidenced by any of the following: Intolerance to treatment, hypersensitivity reactions, significant skin atrophy or systemic effects AND C) One of the following: Phototherapy (UVB or PUVA) has failed to achieve and maintain remission of low or mild disease activity state or is contraindicated OR systemic treatment (for example, immunosuppressants) has failed to achieve and maintain remission of low or mild disease activity state or is contraindicated.
Indications	All Medically-accepted Indications.
Off Label Uses	

Duragesic Patch

Products Affected

- DURAGESIC-100
- DURAGESIC-12
- DURAGESIC-25
- DURAGESIC-50

• DURAGESIC-75	
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• fentanyl

PA Criteria	Criteria Details
Exclusion Criteria	Individual is requesting or using as an as-needed analgesic OR Individual has one of the following conditions: (a) Significant respiratory depression OR (b) Acute or severe bronchial asthma or hypercarbia OR (c) Known or suspected paralytic ileus.
Required Medical Information	
Age Restrictions	Individual is 2 years of age or older
Prescriber Restrictions	
Coverage Duration	Initial 3 months. Maintenance 6 months. Cancer Pain/Terminal Dx or Palliative Care 1 Year.

PA Criteria	Criteria Details
Other Criteria	For initial use, Individual has one of the following: (a) Diagnosis of cancer related pain and/or is actively undergoing cancer therapy (provide cancer diagnosis) OR (b) Diagnosis of terminal illness and is receiving palliative/end-of-life care (provide terminal diagnosis) OR Individual has pain severe enough to require daily, around-the-clock, long term opioid treatment (provide diagnosis) AND Individual has one of the following: (a) An inadequate response to ONE alternative treatment options, such as but not limited to non-opioid analgesics and immediate-release opioids OR (b) Alternative treatment options would otherwise be inadequate to provide sufficient management of pain OR (c) Individual has contraindications to non-opioid analgesics (such as NSAID use in individuals with active ulcer condition/gastrointestinal bleeding, renal failure) AND Individual is not opioid naïve as noted by the following: (a) Individual is currently using a short-acting opioid analgesic to another long-acting opioid analgesic OR (c) already receiving at least 60 mg/day of oral morphine, 30 mg/day of oral oxycodone, 8 mg/day of oral hydromorphone, or an equianalgesic dose of another opioid. For continued use, Attestation (verbal or written) that long-acting opioid therapy has provided meaningful improvement in pain and/or function compared to baseline AND prescriber has consulted with individual regarding risks of opioid therapy AND clear treatment goals have been defined and outlined as part of overall pain.
Indications	All Medically-accepted Indications.
Off Label Uses	

Egrifta

Products Affected

- EGRIFTA
- EGRIFTA SV

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For initial requests, individual has a body mass index (BMI) is greater than 20 kg/m2 AND waist circumference and a waist-to-hip ratio of one of the following (Falutz 2010): (a) For males, waist circumference greater than or equal to 95cm and waist-to-hip ratio greater than or equal to 0.94 OR (b) For females, waist circumference greater than or equal to 94cm and waist-to-hip ratio greater than or equal to 0.88 AND fasting blood glucose (FBG) is less than 150 mg/dL (8.33 mmol/L) AND no history of type 1 diabetes or insulin-treated type 2 diabetes AND no active malignancy (e.g., a potential cancer which is being evaluated or a diagnosed cancer which is being treated). For continuation, individual must demonstrate there is a clear response in reduction of visceral adipose tissue measured by waist circumference or computed tomography (CT) scan.
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	Initial 6 months, renewal 1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Elaprase

Products Affected

• ELAPRASE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Documented deficiency in iduronate 2-sulfatase enzyme activity as measured in fibroblasts or leukocytes combined with normal enzyme activity level of another sulfatase OR Documented pathologic iduronate 2-sulfatase gene mutation.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has symptoms attributable to MPS II such as: (a) Developmental delay or cognitive impairment or (b) Frequent infections or (c) Hearing loss or (d) Hepatosplenomegaly or (e) Hernias or (f) Impaired respiratory function or (g) Joint pain or (h) Skeletal deformities or (i) Sleep apnea or (j) Valvular heart disease.
Indications	All Medically-accepted Indications.
Off Label Uses	

Elidel

Products Affected

- ELIDEL
- pimecrolimus

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	Individual is 2 years of age and older.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual had a trial of and inadequate response or intolerance to one topical prescription strength corticosteroid.
Indications	All Medically-accepted Indications.
Off Label Uses	

ELIGARD_GNRH

Products Affected

• ELIGARD SUBCUTANEOUS KIT 22.5 MG, 30 MG, 45 MG, 7.5 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For Prostate cancer: Clinically localized disease with intermediate (T2b to T2c cancer, Gleason score of 7/Gleason grade group 2-3, or prostate specific antigen (PSA) value of 10-20 ng/mL) OR higher risk of recurrence as neoadjuvant therapy with radiation therapy or cryosurgery OR Other advanced, recurrent, or metastatic disease.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Elitek

Products Affected

• ELITEK

PA Criteria	Criteria Details
Exclusion Criteria	Individual has a diagnosis of glucose-6-phosphate dehydrogenase (G6PD) deficiency.
Required Medical Information	Individual is receiving treatment in a setting appropriate for providing necessary monitoring and supportive care for tumor lysis syndrome AND Individual has not received a course of Elitek therapy in the past.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	30 days
Other Criteria	Individual has been diagnosed with leukemia, lymphoma or other hematologic malignancy with risk factors for tumor lysis syndrome AND Individual is receiving chemotherapy.
Indications	All Medically-accepted Indications.
Off Label Uses	

Elzonris

Products Affected

• ELZONRIS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has a current Eastern Cooperative Oncology Group (ECOG) status of 0-1.
Age Restrictions	2 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Emflaza

Products Affected

• EMFLAZA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	One of the following: (1) Documentation has been provided for excessive weight-gain with prednisone (increase of greater than 0.5 Z score from prior growth curve expectations [American Academy of Pediatrics/CDC Weight for Age Growth Chart, Z-score data files, CDC, Weight-for-age charts, 2 to 20 years, selected weight z-scores in kilograms, by sex and age]) AND Weight gain is likely to be a direct result of prednisone use. Or (2) Documentation has been provided regarding the presence of clinically significant neuropsychiatric side effects while on prednisone (such as but not limited to aggression) AND Neuropsychiatric side effects are likely to be the direct result of prednisone use.
Age Restrictions	Individual is 5 years of age or older
Prescriber Restrictions	
Coverage Duration	6 months

PA Criteria	Criteria Details
Other Criteria	For initial treatment of Duchenne Muscular Dystrophy (DMD) AND Individual has had a 6 month trial of oral prednisone (AAN 2016, DrugPoints B, IIa). Request for continuation of therapy when one of the following: (1) when approved due to excess weight gain with prednisone, individual has experienced a return to baseline growth curve expectations or remained on the same growth curve that was in effect when Emflaza was initiated (American Academy of Pediatrics/CDC Weight for Age Growth Chart, Z-score data files, CDC, Weight-for-age charts, 2 to 20 years, selected weight z- scores in kilograms, by sex and age) Or (2) When approved due to neuropsychiatric side effects while on prednisone, individual has shown improvement in neuropsychiatric symptoms.
Indications	All Medically-accepted Indications.
Off Label Uses	

Empliciti

Products Affected

• EMPLICITI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Emsam

Products Affected

• EMSAM

PA Criteria	Criteria Details
Exclusion Criteria	Individuals with pheochromocytoma OR Individual is currently taking one of the following: (1) Selective serotonin reuptake inhibitors (SSRIs) (for example, fluoxetine) OR (2) Serotonin and norepinephrine reuptake inhibitors (SNRIs) (for example, venlafaxine) OR (3) Tricyclic antidepressants (clomipramine or imipramine) OR (4) Opiate analgesics (meperidine, tramadol, methadone, pentazocine) OR (5) Dextromethorphan OR (6) Carbamazepine.
Required Medical Information	Individual has no known contraindications to the use of a monoamine oxidase inhibitor (MAOI).
Age Restrictions	Individual is 18 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Enbrel

Products Affected

- ENBREL MINI
- ENBREL SUBCUTANEOUS SOLUTION PREFILLED SYRINGE 25 MG/0.5ML, 50 • ENBREL SURECLICK MG/ML
- ENBREL SUBCUTANEOUS SOLUTION RECONSTITUTED

PA Criteria	Criteria Details
Exclusion Criteria	Entanercept used in combination with other TNF antagonist JAK inhibitors, other biologic drugs (such as, abatacept, anakinra, vedolizumab), or cyclophosphamides. Tuberculosis, other active serious infections or history of recurrent infections. Individual who have not had a tuberculin skin test or a CDC-recommended equivalent to evaluate for latent tuberculosis prior to initiating etanercept.
Required Medical Information	For chronic moderate to severe plaque psoriasis with either of the following: Individual has greater than 5% of body surface area with plaque psoriasis OR Less than or equal to 5% of body surface area with plaque psoriasis involving sensitive areas or areas that would significantly impact daily function (such as palms, soles of feet, head/neck, or genitalia).
Age Restrictions	Individual is 18 years of age or older, except for the diagnosis of JIA and plaque psoriasis. For JIA individual is 2 years of age or older. For plaque psoriasis, 4 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year except for Initial high dose tx chronic plaque psoriasis 12 wk

PA Criteria	Criteria Details
Other Criteria	For moderate to severe Ankylosing Spondylitis, individual has had an inadequate response to, is intolerant of, or has a contraindication to ONE conventional therapies: (such as NSAIDs or nonbiologic DMARDs) (ACR 2015). For Moderate to severe Chronic Plaque Psoriasis, individual has had inadequate response to, is intolerant of, or has a medical contraindication to phototherapy OR ONE other systemic therapies (such as, methotrexate, acitretin, or cyclosporine). For moderate to severe Rheumatoid Arthritis, individual has had an inadequate response to, is intolerant of, or has a contraindication to ONE conventional therapy [nonbiologic DMARDs (such as methotrexate, sulfasalazine, leflunomide or hydroxychloroquine)] (ACR 2015). For moderate to severe Polyarticular JIA, individual has had an inadequate response to, is intolerant of, or has a contraindication to ONE conventional therapy [nonbiologic DMARDs such as methotrexate] (ACR 2011). For moderate to severe Psoriatic Arthritis, individual has had an inadequate response to, is intolerant of, or has a contraindication to ONE conventional therapy [nonbiologic DMARDs (such as methotrexate, sulfasalazine or leflunomide)] (AAD 2011).
Indications	All Medically-accepted Indications.
Off Label Uses	

Enhertu

Products Affected

• ENHERTU

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has unresectable or metastatic HER2-positive (HER2+) breast cancer confirmed (written or verbal) by either Immunohistochemistry (IHC) is 3+ OR In situ hybridization (ISH) positive
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual is using Enhertu as monotherapy.
Indications	All Medically-accepted Indications.
Off Label Uses	

Entresto

Products Affected

• ENTRESTO

PA Criteria	Criteria Details
Exclusion Criteria	Individual is pregnant/ wishing to become pregnant OR breastfeeding OR has a history of angioedema related to previous ACE inhibitor or ARB therapy OR has severe hepatic impairment (Child-Pugh C). OR Individual will be utilizing an angiotensin- converting enzyme (ACE) inhibitor OR angiotensin receptor blocker (ARB) in combination with Entresto (sacubitril/valsartan). Individual will be utilizing Entresto (sacubitril/valsartan) in combination with Tekturna (aliskiren)/Tekturna HCT (aliskiren/hydrochlorothiazide) and has a diagnosis of diabetes or renal impairment (eGFR less than 60 mL/min/1.73 m2).
Required Medical Information	Individual has a left ventricular ejection fraction less than or equal to 40%.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Entyvio

Products Affected

• ENTYVIO

PA Criteria	Criteria Details
Exclusion Criteria	Entyvio used in combination with other biologic drugs (such as TNF antagonists or natalizumab or non-TNF antagonists immunomodulatory drugs, such as or Tysabri (natalizumab)). Active, severe infections, or a history of recurrent infections.
Required Medical Information	
Age Restrictions	Individual is 6 years or older.
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For UC or CD, individual has had an inadequate response to, is intolerant of, or has a contraindication to conventional therapy (such as 5-aminosalicylic acid products, systemic corticosteroids or immunosuppressants) or a tumor necrosis factor (TNF) antagonists AND Individual has had a trial and an inadequate response or is intolerant to Humira (adalimumab) OR the TNF agent (Humira (adalimumab)) tried and failed is not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Entyvio or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis OR individual has either concomitant clinical condition: (a) Demyelinating disease or (b) Heart failure with documented left ventricular dysfunction, (c) Malignancy [such as, but not limited to, solid or hematologic cancers excluding superficial skin cancers (such as basal and squamous cell], or (d) tuberculosis infection. Entyvio may be allowed without trial of preferred TNF agents (Humira).
Indications	All Medically-accepted Indications.
Off Label Uses	

Epclusa

Products Affected

- EPCLUSA
- sofosbuvir-velpatasvir

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Documentation is provided for a diagnosis of chronic hepatitis C (CHC) infection, which includes genotype and positive HCV RNA result (AASLD/IDSA 2017, CDC 2013) AND Individual has received baseline evaluation for liver fibrosis to guide appropriate therapy AND Individual does not have a short life expectancy (less than 12 months owing to non-liver related comorbid conditions) that cannot be remediated by treating HCV, by transplantation or other directed therapy (AASLD/IDSA 2016).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Criteria will be applied consistent with current AASLD/IDSA guidance.
Other Criteria	Criteria will be applied consistent with current AASLD/IDSA guidance.
Indications	All Medically-accepted Indications.
Off Label Uses	

Epidiolex

Products Affected

• EPIDIOLEX

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Epogen and Procrit

Products Affected

- EPOGEN
- PROCRIT
- RETACRIT

PA Criteria	Criteria Details
Exclusion Criteria	Anemia in patients due to other factors such as iron deficiency, folate deficiency or B12 deficiency, hemolysis, gastrointestinal bleeding, other active or occult bleeding, or underlying hematologic diseases (such as sickle cell anemia, thalassemia, and porphyria). Sudden loss of response with severe anemia and low reticulocyte count. Treatment of anemia in patients with cancer not treated by chemotherapy known to produce anemia. Treatment of in any indication not listed in criteria including anemia of prematurity. Anemia in cancer patients receiving myelosuppressive chemotherapy when the anticipated outcome is cure. Anemia in cancer patients receiving concomitant myelosuppressive chemotherapy unless receiving concomitant myelosuppressive chemotherapy in whom the anemia can be managed by transfusion. Continued use when the hemoglobin level exceeds 11.0 g/dL unless otherwise specified in the criteria. Use beyond 12 weeks in the absence of response in individuals with myelodysplastic syndrome (MDS). Use beyond 8-9 weeks in the absence of response or if transfusions are still required in individuals with metastatic, non-myeloid cancer being treated with myelosuppressive chemotherapy agents known to produce anemia. Continued use beyond 6 weeks after therapy with myelosuppressive chemotherapy known to produce anemia is completed. Pre-operative use for patients who are willing to donate autologous blood.

Y0114_20_114077_I_012

Criteria Details
Hemoglobin (Hgb) levels are less than 10.0 g/dL, prior to initiation of therapy (unless otherwise specified) AND prior to initiation of therapy, (baseline) evaluation of the individual iron status reveals, transferrin saturation at least 20% or ferritin at least 80 ng/mL or bone marrow demonstrates adequate iron stores AND For individuals with uncontrolled hypertension, blood pressure is adequately controlled before initiation of therapy and closely monitored and controlled during therapy. For MDS, endogenous erythropoietin level is less than 500 mU/ml. For anemia related to zidovudine in HIV-infected patients when the dose of zidovudine is less than or equal to 4200 mg per week, endogenous erythropoietin level is less than or equal 500 mU/ml. Reduction of Allogeneic Blood Transfusion in Pre-Operative Surgery Patients: Patient's hgb is greater than 10.0 and less than or equal to 13.0 g/dL, individual is scheduled to undergo elective, noncardiac, nonvascular surgery, individual is at high risk for perioperative transfusions with significant, anticipated blood loss, individual is unable or unwilling to donate autologous blood, Antithrombotic prophylaxis has been considered. For tx of anemia due to myelosuppressive chemotherapy known to produce anemia when the following are met: chemo is planned for a minimum of 2 months and the dx is non-myeloid cancer and the anticipated outcome is not cure. For anemia associated with CKD ON dialysis use is to achieve and maintain hgb levels within the range of 10.0 to 11.0 g/dL. For anemia associated with CKD NOT ON dialysis use is to achieve and maintain hgb levels of 10.0g/dL.
8wk.
For Hepatitis C, patient is concomitantly treated with combination of ribavirin and interferon alfa, or ribavirin and peginterferon alfa. Myelosuppressive drugs known to produce anemia in individuals with a diagnosis of chronic inflammatory disease. Allogeneic bone marrow transplantation.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Eraxis

Products Affected

• ERAXIS

	1
PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Erbitux

Products Affected

• ERBITUX

PA Criteria	Criteria Details
Exclusion Criteria	Erbitux is used in combination with other anti-VEGF agents (e.g., bevacizumab). Erbitux is used in more than one line of therapy.
Required Medical Information	For stage IV, kras wild type colon, rectal, colorectal, small bowel, appendix, or anal adenocarcinoma when used as a single agent or as part of combination therapy. For squamous cell carcinoma of the Head and/or Neck cancer, Erbitux is used in combination with radiation therapy, for the treatment of locally or regionally advanced disease. Or as a single agent for the treatment of patients with recurrent or metastatic disease for whom prior platinum-based therapy has failed. Or in combination with platinum-based therapy with 5-FU (fluorouracil) as first-line treatment for individuals with recurrent locoregional disease or metastatic SCCHN. OR as a single agent or in combination therapy with or without radiation therapy for any of the following indications, unresectable locoregional recurrence or second primary in individuals who have received prior radiation therapy OR resectable locoregional recurrence in individuals who have not received prior radiation therapy OR distant metastases.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Erivedge

Products Affected

• ERIVEDGE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Erleada

Products Affected

• ERLEADA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Erwinase

Products Affected

• ERWINAZE

PA Criteria	Criteria Details
Exclusion Criteria	History of serious thrombosis with prior L-asparaginase therapy. History of pancreatitis with prior L-asparaginase therapy. History of serious hemorrhagic events with prior L-asparaginase therapy.
Required Medical Information	Individual is using Erwinase as a component of a multi-agent chemotherapeutic regimen AND is using for Acute lymphoblastic lymphoma or acute lymphocytic leukemia (ALL) or Extranodal natural killer T-cell lymphoma, nasal type (ENKL). Individual has developed a documented systemic allergic reaction or anaphylaxis to prior treatment with Oncaspar (Pegaspargase).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Esbriet

Products Affected

- ESBRIET ORAL CAPSULE
- ESBRIET ORAL TABLET 267 MG, 801 MG

PA Criteria	Criteria Details
Exclusion Criteria	Individuals using in combination with Ofev (nintedanib). Individuals with end-stage renal disease (ESRD). Individuals with severe hepatic impairment (child pugh class C) or end-stage liver disease.
Required Medical Information	Diagnosis of idiopathic pulmonary fibrosis (IPF) is confirmed by: Exclusion of other known causes of interstitial lung disease (ILD) such as domestic and occupational environmental exposures, connective tissue disease, and drug toxicity AND High resolution computed tomography (HRCT) with or without surgical lung biopsy. Individual has documented pulmonary function tests within prior 60 days with a Forced Vital Capacity (% FVC) greater than or equal to 50%.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Ethyol

Products Affected

• ETHYOL

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Evekeo

Products Affected

- amphetamine sulfate oral tablet 10 mg, 5 mg
- EVEKEO ODT ORAL TABLET DISPERSIBLE 10 MG, 15 MG, 20 MG, 5

MG

• EVEKEO ORAL TABLET 10 MG, 5 MG

PA Criteria	Criteria Details
Exclusion Criteria	Individual is using for exogenous obesity/weight loss [Exclusion from Part D].
Required Medical Information	
Age Restrictions	3 years of age or older for attention deficit hyperactivity disorder (ADHD). 6 years of age or older for narcolepsy.
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Evenity

Products Affected

• EVENITY

PA Criteria	Criteria Details
Exclusion Criteria	Individual is using Evenity (romosozumab-aqqg) in combination with any of the following: Prolia (denosumab), Bisphosphonates, Evista (raloxifene), Miacalcin/Fortical (calcitonin nasal spray), Reclast (zoledronic acid), Forteo (teriparatide), Tymlos (abaloparatide).
Required Medical Information	Osteoporosis is defined as a BMD T-Score in the spine, femoral neck, total hip or distal 1/3 of the radius of less than or equal to - 2.5 as compared to young adult reference population OR a Clinical diagnosis based on a history of a low trauma fracture (fragility fracture) at high risk for fracture.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.

PA Criteria	Criteria Details
Other Criteria	Individual meets one of the following: (A) Individual has been refractory to a trial of an oral bisphosphonate therapy OR (B) Intolerance or contraindications to oral bisphosphonate as defined by having at least one of the following: 1. Hypersensitivity to TWO oral bisphosphonates (one of which must be alendronate) 2. Inability to sit or stand upright for at least 30 minutes, 3. Pre- existing gastrointestinal disorders (Barrett's esophagus, hypersecretory disorders, delayed esophageal emptying, etc.). 4. Uncorrected hypocalcemia. 5. Severe renal insufficiency as defined by creatinine clearance less than 35 mL/min for alendronate agents or creatinine clearance less than 30 mL/min for risedronate and ibandronate. Individual has utilized Evenity (romosozumab-aqqg) for a total duration of less than 12 months in their lifetime.
Indications	All Medically-accepted Indications.
Off Label Uses	

Evrysdi

Products Affected

• EVRYSDI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Diagnosis of spinal muscular atrophy (SMA) by documentation (written or verbal attestation) of either: (A) Spinal Muscular Atrophy (SMA) diagnostic test results confirming 0 copies of SMN1 OR (B) Molecular genetic testing of 5q SMA for any of the following: (1) Homozygous gene deletion OR (2) Homozygous conversion mutation OR (3) Compound heterozygote.
Age Restrictions	Individual is 2 months of age or older.
Prescriber Restrictions	
Coverage Duration	Initial and Continuation 6 months.

PA Criteria	Criteria Details
Other Criteria	Initial requests, individual has documentation (written or verbal attestation) of SMA-associated signs and symptoms AND (1) has documentation (written or verbal attestation) of genetic testing confirming 2 copies of SMN2 (NCT02913482) AND has documentation of symptom onset before 3 months of age (NCT02913482). OR (2) Individual is 2 years of age or older AND is non-ambulant as defined by being unable to walk unassisted for greater than or equal to 10m (NCT02908685). AND Individual does not require use of invasive ventilation or tracheostomy as a result of advanced SMA disease. For continuation, when initial therapy was determined to meet the above criteria AND individual has documentation (written or verbal attestation) of clinically significant improvement in spinal muscular atrophy-associated signs and symptoms (i.e., progression, stabilization, or decreased decline in motor function) compared to the predicted natural history trajectory of disease AND does not require use of invasive ventilation or tracheostomy as a result of advanced SMA disease. For INITIAL use following treatment with Zolgensma (onasemnogene abeparvovec-xioi): individual meets initial criteria above AND individual has experienced a decline in clinical status (for example, loss of motor milestone) since receipt of gene therapy.
Indications	All Medically-accepted Indications.
Off Label Uses	

Exjade

Products Affected

- deferasirox
- EXJADE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	Individual is 2 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Exondys 51

Products Affected

• EXONDYS 51

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has a confirmed genetic mutation of the DMD gene that is amenable to exon 51 skipping
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For initial use in the treatment of Duchenne muscular dystrophy (DMD), individual is ambulatory (with or without needing an assistive device, such as a cane or walker). For continued therapy following each 12 month period post initiation of therapy, the initial therapy has been met AND the individual remains ambulatory (with or without needing an assistive device, such as a cane or walker).
Indications	All Medically-accepted Indications.
Off Label Uses	

Eylea

Products Affected

• EYLEA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Fabrazyme

Products Affected

• FABRAZYME

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Diagnosis of Fabry disease is confirmed with either of the following: (a) Documentation of complete deficiency or less than 5% of mean normal alpha-galactosidase A enzyme activity in leukocytes, dried blood spots or serum (plasma) analysis or (b) Documented galactosidase alpha gene mutation by gene sequencing.
Age Restrictions	Individual is 8 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has one or more symptoms or physical findings attributable to Fabry disease (ACMG), such as, but not limited to: (a) Burning pain in the extremities (Acroparesthesias) or (b) Cutaneous vascular lesions (Angiokeratomas) or (c) Corneal verticillata (whorls) or (d) Decreased sweating (anhidrosis or hypohidrosis) or (e) Personal or family history of exercise, heat, or cold intolerance or (f) Personal or family history of kidney failure.
Indications	All Medically-accepted Indications.
Off Label Uses	

Farydak

Products Affected

 FARYDAK ORAL CAPSULE 10 MG, 15 MG, 20 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Fasenra

- FASENRA
- FASENRA PEN

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Evidence of asthma is demonstrated by the following (NAEPP, 2008): (a) pretreatment forced expiratory volume in 1 second (FEV1) less than 80% predicted AND (b) FEV1 reversibility of at least 12% and 200 milliliters after albuterol administration.
Age Restrictions	12 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year.

PA Criteria	Criteria Details
Other Criteria	For Initial use, individual has had a 3 month trial and inadequate response or intolerance to combination controller therapy (high dose inhaled corticosteroids plus long acting beta2 -agonists, leukotriene modifiers, theophylline or oral corticosteroids) (ERS/ATS, 2013) AND has experienced two or more asthma exacerbations in the prior 12 months requiring use of a systemic corticosteroid or temporary increase in the individual's usual maintenance dosage of oral corticosteroids (ERS/ATS, 2013) AND has a blood eosinophil count (in the absence of other potential causes of eosinophilia, including hypereosinophilic syndromes, neoplastic disease, and known or suspected parasitic infection) greater than or equal to 300 cells/microliter (300 cells/mm3) at initiation of therapy. For Continuation use, treatment has resulted in clinical improvement as confirmed by one or more of the following: (a) decreased utilization of rescue medications OR (b) decreased frequency of exacerbations (defined as worsening of asthma that requires an increase in inhaled corticosteroid dose or treatment with systemic corticosteroids) OR (c) increase in percent predicted FEV1 from pretreatment baseline OR (d) reduction in reported asthma-related symptoms, such as asthmatic symptoms upon awakening, coughing, fatigue, shortness of breath, sleep disturbance, or wheezing.
Indications	All Medically-accepted Indications.
Off Label Uses	

Faslodex

Products Affected

- FASLODEX
- fulvestrant

DA Cuitaria	
PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Fentora

Products Affected

- fentanyl citrate buccal
- FENTÓRA

PA Criteria	Criteria Details
Exclusion Criteria	Using for treatment of acute or postoperative pain OR migraine headache pain OR non-cancer related pain.
Required Medical Information	
Age Restrictions	Individual is 18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a diagnosis of active cancer (such as, metastatic or locally invasive cancer for which the individual is currently seeking treatment) with breakthrough cancer pain (provide diagnosis) AND has tried and failed fentanyl citrate lozenge (generic Actiq) AND Individual is already receiving opioid therapy and is TOLERANT to opioid therapy as defined as receiving around the clock medicine consisting of one of the following: At least 60mg morphine per day, OR At least 25mcg/hr transdermal fentanyl/hour, OR At least 30mg of oxycodone daily, OR At least 8mg of oral hydromorphone daily, OR At least 25mg of oral oxymorphone daily, OR An equianalgesic dose of another opioid for a week or longer. Individual will also continue around the clock opioids when taking fentanyl citrate for cancer related breakthrough pain.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Ferriprox

Products Affected

- deferiprone
- FERRIPROX
- FERRIPROX TWICE-A-DAY

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Fetzima

- FETZIMA ORAL CAPSULE EXTENDED RELEASE 24 HOUR 120 MG, 20 MG, 40 MG, 80 MG
- FETZIMA TITRATION

PA Criteria	Criteria Details
Exclusion Criteria	May not be approved for treatment of fibromyalgia
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For MDD, individual has had a trial of TWO of the following: Desvenlafaxine ER, desvenlafaxine Fumerate ER, fluoxetine, fluvoxamine, escitalopram, citalopram, paroxetine, sertraline, mirtazapine, immediate-release venlafaxine, extended-release venlafaxine or bupropion.
Indications	All Medically-accepted Indications.
Off Label Uses	

Fintepla

Products Affected

• FINTEPLA

PA Criteria	Criteria Details
Exclusion Criteria	Individual is using for weight loss/reduction OR using within 14 days of taking a monoamine oxidase inhibitor (MAOI).
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual has a diagnosis of seizures associated with Dravet Syndrome AND has responded inadequately to two previous antiepileptic drugs (Lagae 2019, Wirrell 2017, Ziobro 2018).
Indications	All Medically-accepted Indications.
Off Label Uses	

Firazyr

- FIRAZYR
- *icatibant acetate*

PA Criteria	Criteria Details
Exclusion Criteria	Prophylaxis for HAE attacks.
Required Medical Information	HAE to be confirmed by a C4 level below the lower limit of normal (as defined by the laboratory performing the test) and either a C1 inhibitor antigenic level below the lower limit of normal (as defined by the lab performing test) or a C1 inhibitor functional level below the lower limit of normal (as defined by the lab performing the test).
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a history of moderate or severe attacks (for example, airway swelling, severe abdominal pain, facial swelling, nausea and vomiting, painful facial distortion) and using Icatibant for acute HAE attacks.
Indications	All Medically-accepted Indications.
Off Label Uses	

Firdapse

- FIRDAPSE
- RUZURGI

PA Criteria	Criteria Details
Exclusion Criteria	History of seizures or Using in combination with compounded form of 3,4 diaminopyridine
Required Medical Information	Diagnosis is confirmed by one of the following: Presence of anti- P/Q type voltage-gated calcium channel (VGCC) antibodies or Characteristic electrodiagnostic findings.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For initial requests, individual has diagnosis of Lambert Eaton myasthenic syndrome. Continued treatment with Firdapse may be approved when there is objective evidence that the individual achieved and sustained meaningful improvement in muscle strength.
Indications	All Medically-accepted Indications.
Off Label Uses	

Firmagon

- FIRMAGON (240 MG DOSE)
- FIRMAGON SUBCUTANEOUS SOLUTION RECONSTITUTED 120 MG, 80 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For Prostate cancer: Clinically localized disease with intermediate (T2b to T2c cancer, Gleason score of 7/Gleason grade group 2-3, or prostate specific antigen (PSA) value of 10-20 ng/mL) OR higher risk of recurrence as neoadjuvant therapy with radiation therapy or cryosurgery OR Used for progressive castration-naïve disease or for castration-recurrent disease OR Other advanced, recurrent, or metastatic disease.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Flector Patch

- diclofenac epolamine
- FLECTOR
- LICART

PA Criteria	Criteria Details
Exclusion Criteria	Individual is using for perioperative pain in the setting of coronary artery bypass graft (CABG) surgery. Individual is using on non- intact or damaged skin resulting from any etiology, including exudative dermatitis, eczema, infection lesions, burns or wounds.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	3 months
Other Criteria	Individual is using for the treatment of acute pain from one of the following: (a) Minor strain OR (b) Sprain OR (c) Contusion.
Indications	All Medically-accepted Indications.
Off Label Uses	

Forteo

- FORTEO
- teriparatide (recombinant)

PA Criteria	Criteria Details
Exclusion Criteria	Individual is using Forteo (teriparatide) in combination with any of the following: Prolia (denosumab), Bisphosphonates, Evista (raloxifene), Miacalcin/Fortical (calcitonin nasal spray), Recalst (zolendronic acid), or Tymlos (abalaoparatide).
Required Medical Information	Osteoporosis is defined as a BMD T-Score in the spine, femoral neck, total hip or distal 1/3 of the radius of less than or equal to - 2.5 as compared to young adult reference population OR a Clinical diagnosis based on a history of a low trauma fracture (fragility fracture) at high risk for fracture OR associated with sustained systemic glucocorticoid therapy (daily dosage equivalent to 5mg or greater of prednisone for at least 3 months) at high risk for fracture.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	2 years. Requests to continue therapy beyond 24 months (2 years) should not be approved.

PA Criteria	Criteria Details
Other Criteria	Individual meets one of the following: (A) Individual has been refractory to a trial of an oral bisphosphonate therapy OR (B) Intolerance or contraindications to oral bisphosphonate as defined by having at least one of the following: 1. Hypersensitivity to TWO oral bisphosphonates (one of which must be alendronate) 2. Inability to sit or stand upright for at least 30 minutes, 3. Pre- existing gastrointestinal disorders (Barrett's esophagus, hypersecretory disorders, delayed esophageal emptying, etc.). 4. Uncorrected hypocalcemia. 5. Severe renal insufficiency as defined by creatinine clearance less than 35 mL/min for alendronate agents or creatinine clearance less than 30 mL/min for risedronate and ibandronate.
Indications	All Medically-accepted Indications.
Off Label Uses	

Galafold

Products Affected

• GALAFOLD

PA Criteria	Criteria Details
Exclusion Criteria	Individual has severe renal impairment or end-stage renal disease.
Required Medical Information	Individual has a diagnosis of Fabry disease as confirmed with either Documentation (written or verbal attestation is acceptable) of complete deficiency or less than 5% of mean normal alpha- galactosidase A (a-Gal A) enzyme activity in leukocytes, dried blood spots or serum (plasma) analysis OR Documented (written or verbal attestation is acceptable) galactosidase alpha (GLA) gene mutation by gene sequencing. Individual has an amendable GLA gene variant based on the human embryonic kidney-293 assay.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has one or more symptoms or physical findings attributable to Fabry disease (ACMG), such as but not limited to: (a) Burning pain in the extremities (acroparesthesias), or (b) Cutaneous vascular lesions (angiokeratomas), or (c) Corneal verticillata (whorls), or (d) Decreased sweating (anhidrosis or hypohidrosis), or (e) Personal or family history of exercise, heat, or cold intolerance, or (f) Personal or family history of kidney failure.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

GamaSTAN

Products Affected

- GAMASTAN
- GAMASTAN S/D

PA Criteria	Criteria Details
Exclusion Criteria	Individual with isolated immunoglobulin A (IgA) deficiency. Individual with severe thrombocytopenia or any coagulation disorder that would contraindicate intramuscular injections. Prophylaxis of viral hepatitis type B. Routine post-exposure prophylaxis or treatment of rubella, poliomyelitis, mumps, or varicella. Allergy or asthma in individuals who have normal levels of immunoglobulin. Treatment to prevent recurrent spontaneous abortion in pregnant women with a history of recurrent spontaneous abortion (ASRM 2012).
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	Pre-Exposure of HAV, mbr will get IM inj prior to exposure AND mbr has no clinical manifestations of hepatitis A AND is unvaccinated (CDC 2007/2015) along with one of the following: unable to receive HAV vaccine (such as, contraindication to or unavailability of the vaccine) OR mbr is considered high-risk (such as but not limited to, travel to an endemic area, older adults, immunocompromised, or diagnosis of chronic liver disease) and will receive a simultaneous dose of HAV vaccine unless contraindicated. Post-Exposure of HAV, mbr will get IM inj within 2 weeks of exposure AND mbr has no clinical manifestations of hepatitis A AND is unvaccinated (CDC 2007/2015) along with one of the following: unable to receive HAV vaccine (such as, contraindication to or unavailability of the vaccine) OR mbr is considered high-risk (such as but not limited to, immunocompromised, diagnosis of chronic liver disease, or vaccine contraindication). For post exposure prophylaxis of rubeola, must be given within 6 days of exposure and not concomitantly with a vaccine containing the measles virus AND eligible exposed, non-immune individuals will receive a vaccine containing the measles virus greater than or equal to 6 months after receiving intramuscular immune globulin (CDC 2013) AND used in mbr considered at risk for severe disease and complications: infants or previously unvaccinated and ineligible to receive a vaccine contraindication or an initial exposure greater than 72 hours) or no evidence of measles immunity in particular pregnant woman or severely immunocompromised individuals. For post-exposure prophylaxis of varicella infection in susceptible individuals (such as, immunocompromised) AND varicella-zoster immune globulin (human) (VZIG) and immune globulin intravenous (IGIV) are not available. For post-exposure prophylaxis administered within 72 hours of exposue to a confirmed case of rubella to modify to suppress symptoms (label, CDC 2001) AND
	mbr is in the early stages (first trimester) of pregnancy, and will not consider terminating the pregnancy under any circumstance.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Gamifant

Products Affected

• GAMIFANT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has a diagnosis of active primary active primary hemophagocytic lymphohistiocytosis (HLH) as confirmed (written or verbal attestation is acceptable) by one of the following: 1) A genetic mutation known to cause HLH or 2) A family history consistent with primary HLH or 3) Individual meets five of the following criteria: Fever, Splenomegaly, Cytopenias affecting 2 of 3 lineages in the peripheral blood (HGB less than 9g/dL (or less than 10g/dL in infants), platelets less than 100 x 10 9/L, neutrophils less than 1 x 10 9/L), Hypertriglyceridemia (fasting TG greater than or equal to 265mg/dL) and/or hypofibrinogenemia (fibrinogen less than or equal to 1.5g/L), Hemophagocytosis in bone marrow, spleen, or lymph nodes with no evidence of malignancy, Low or absent NK-cell activity, Ferritin greater than or equal to 500 mcg/L, Soluble CD25 greater than or equal to 2400U/mL
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual is using in combination with dexamethasone AND has had an inadequate response to, is intolerant of or has a contraindication to conventional therapy (such as etoposide, dexamethasone or cyclosporine).
Indications	All Medically-accepted Indications.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

Gattex

Products Affected

• GATTEX

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For diagnosis of Short Bowel Syndrome (SBS) individual has been dependent on parenteral nutrition/intravenous (PN/IV) support, For at least 12 months AND requires PN at least 3 times per week.
Indications	All Medically-accepted Indications.
Off Label Uses	

Gauchers

Products Affected

- CEREZYME
- ELELYSO
- VPRIV

PA Criteria	Criteria Details
Exclusion Criteria	Use of enzyme replacement therapy (ERT) agents in conjunction with one another or in conjunction with a substrate reduction therapy (SRT) agent. Use of ERT agents for the treatment of type 2 gaucher disease.
Required Medical Information	Type 1 Gaucher is confirmed by either (Weinreb 2004, Wang 2011): Deficiency in Glucocerebrosidase activity as measured in white blood cells or skin fibroblasts OR genotype tests indicating mutation of two alleles of the glucocerebrosidase genome. And indiv has clinically significant manifestations of gauchers (Andersson 2005, Weinreb 2004) including for type 1,3: [Adults] skeletal disease (such as but not limited to avascular necrosis, Erlenmeyer flask deformity, osteopenia or pathological fracture) OR presents with 2 or more of the following: clinically significant hepatomegaly/splenomegaly, hgb at least 1 gm/dl below lower limit for normal for age and sex, platelet counts less than or equal to 120,000 mm3. [Children] clinical manifestations such as but not limited to hepatomegaly, splenomegaly, anemia, thrombocytopenia, skeletal disease or growth failure (Andersson 2005) OR Type 3 gauchers is confirmed by genotype testing indicating mutation of 2 alleles of the glucocerebrosidase genome (Kaplan 2013, Wang 2011) And has clinically significant manifestations of gauchers listed above in type 1 AND Neurological findings are consistent with type 3 gaucher disease based on neurological evaluation including brain imaging[MRI or CT and EEG].
Age Restrictions	
Prescriber Restrictions	

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Gavreto

Products Affected

• GAVRETO

PA Criteria	Criteria Details
Exclusion Criteria	Pending CMS Review
Required Medical Information	Pending CMS Review
Age Restrictions	Pending CMS Review
Prescriber Restrictions	Pending CMS Review
Coverage Duration	Pending CMS Review
Other Criteria	Pending CMS Review
Indications	Pending CMS Review
Off Label Uses	Pending CMS Review

Gazyva

Products Affected

• GAZYVA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 Year
Other Criteria	Gazyva may be approved for treatment of chronic lymphocytic leukemia/small lymphocytic lymphoma for any of the following: a) first-line in individuals without del (17p) mutation when used in combination with chlorambucil or bendamustine OR as first-line single agent in individuals who are frail or with del (17p) mutation OR as a single agent for the treatment of relapsed/refractory disease without del (17p) mutation. Approved for the treatment of follicular lymphoma when used as a component of ONE of the following combination therapies regimens and as monotherapy for up to 24 months or until disease progression, following the listed combination therapy regimens: cyclophosphamide, doxorubicin, vincristine and prednisone (CHOP regimen) or cyclophosphamide, vincristine, and prednisone (CVP regimen) or bendamustine.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Gilenya

Products Affected

• GILENYA

PA Criteria	Criteria Details
Exclusion Criteria	Concurrent use with other MS disease modifying agents (such as, Aubagio, Tecfidera, Tysabri, Ocrevus, Copaxone/Glotopa, Extavia, Rebif, Avonex, Plegridy, Betaseron). Individual has history or presence of Mobitz Type II second- or third-degree atrioventricular (AV) block or sick sinus syndrome, unless individual has a functioning pacemaker. Individual has a baseline QTc interval greater than or equal to 500 ms. Individual is being treated with Class Ia (such as quinidine, procainamide, or disopyramide) or Class III [such as amiodarone, Multaq (dronedarone), Tikosyn (dofetilide), or sotalol] anti-arrhythmic drugs. Individual has had a recent (within the past 6 months) occurrence of one of the following: Myocardial infarction, Unstable angina, Stroke, Transient ischemic attack (TIA), Decompensated heart failure requiring hospitalization, Class III/IV heart failure or individual has an active acute or chronic infection at the initiation of therapy.
Required Medical Information	I. Individual has had a trial and inadequate response or intolerance to one of the following: Avonex (interferon beta-1a), Plegridy (interferon beta-1-a), Betaseron (interferon beta-1b), Tecfidera (dimethyl fumarate), Copaxone/Glatopa (glatiramer) OR II. Individual has high disease activity despite treatment with a disease modifying drug (Aubagio, Avonex, Plegridy, Rebif, Betaseron, Extavia, Copaxone/Glatopa, Tecfidera) defined as the following: At least 1 relapse in the previous year while on therapy AND At least 9 T2-hyperintense lesions in cranial MRI OR At least 1 Gadolinium-enhancing lesion. OR III. Individual is treatment naive (no previous history of use of disease modifying drugs such as Aubagio, Avonex, Plegridy, Rebif, Betaseron, Extavia, Copaxone/Glatopa, Tecfidera) AND IV. Individual has rapidly evolving severe relapsing multiple sclerosis defined as the following: Two or more disabling relapses in 1 year AND One or more Gadolinium-enhancing lesions on brain MRI.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Gilotrif

Products Affected

• GILOTRIF

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Test results confirmed for individuals with metastatic non-small cell lung cancer (NSCLC) with non-resistant epidermal growth factor receptor (EGFR) mutation.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Givlaari

Products Affected

• GIVLAARI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has acute hepatic porphyria and confirmation (written or verbal) of one of the following subtypes (APF 2010-2019): Acute intermittent porphyria (AIP) OR Hereditary coproporphyria (HCP) OR Variegate porphyria (VP) OR ALA dehydratase-deficiency porphyria (ADP) AND documentation (written or verbal) of elevated urinary or plasma porphobiligen or delta-aminolevulinic acid within the past year (NCT03338816) AND individual has active symptomatic disease with at least 2 documented (written or verbal) porphyria attacks within the last 6 months (NCT03338816).
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	Initial: 6 months and continuance: 1 year
Other Criteria	For Continuation Therapy: Individual has experienced a clinical response to therapy (for example, a reduction in the number of porphyria attacks) AND does not have severe or clinically significant transaminase elevations defined as alanine aminotransferase (ALT) greater than 5 times the upper limit of normal (Balwani 2019).
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Gleevec

- GLEEVEC ORAL TABLET 100 MG, 400 MG
- imatinib mesylate oral tablet 100 mg, 400 mg

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Gleostine

Products Affected

• GLEOSTINE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Granix

Products Affected

• GRANIX

PA Criteria	Criteria Details
Exclusion Criteria	Use as prophylaxis for FN during concomitant chemotherapy and radiation therapy. Continued use if no response is seen within 28-42 days (individuals who have failed to respond within this time frame are considered non-responders).
Required Medical Information	Primary prophylaxis of FN in Individual that has a risk of developing FN is greater than or equal to 10% and less than of 20% based on chemotherapy regimens and individuals have any of the following risk factors for FN: Individual age greater than 65 years, Poor performance status, Bone marrow involvement by tumor producing cytopenias, persistent neutropenia (ANC less than 1500mm3), poor renal function (GFR less than 60mL/min), liver dysfunction, recent surgery and/or presence of open wounds. Individual has not received prophylactic therapy with granulocyte colongy stimulating factor AND has a high-risk for infection associated complications as demonstrated by any of the following: Expected prolonged (greater than 10 day) and profound (less than 0.1 x 10to the power of 9/L) neutropenia, Age greater than 65 years, Pneumonia or other clinically documented infections, Hypotension and multi organ dysfunction (sepsis syndrome), Invasive fungal infection, prior episode of febrile neutropenia or Hospitalized at the time of the development of fever.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Other Criteria	Individual has trial and inadequate response to Zarxio (filgrastim- sndz). Individual is using to mobilize progenitor cells into peripheral blood for collection by leukapheresis as adjunct to peripheral blood/hematopoietic stem cell transplantation (PBSCT/PHSCT).
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Grastek

Products Affected

• GRASTEK

PA Criteria	Criteria Details
Exclusion Criteria	Severe, unstable or uncontrolled asthma. History of any severe systemic allergic reaction. History of eosinophilic esophagitis. Or, individual is receiving concomitant therapy with other allergen immunotherapy products.
Required Medical Information	For grass pollen induced allergic rhinitis, individual has a confirmed (verbal or written attestation) positive skin test OR positive in vitro testing for pollen-specific IgE antibodies for Timothy grass or cross- reactive grass pollens. Individual has had a trial of, and inadequate symptom control or intolerance to (1) nasal steroid and (1) non- sedating antihistamine AND individual has a confirmed (verbal or written attestation) prescription for an auto-injectable epinephrine product.
Age Restrictions	Individual is between the ages of 5 years and 65 years old.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Treatment is initiated at least 12 weeks before the expected onset of grass pollen season and is continued throughout the season.
Indications	All Medically-accepted Indications.
Off Label Uses	

Haegarda

Products Affected

• HAEGARDA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Hereditary angioedema (HAE) is confirmed by a C4 level below the lower limit of normal as defined by the laboratory performing the test AND ANY of the following (a, b, or c): (a) C1 inhibitor (C1-INH) antigenic level below the lower limit of normal as defined by the laboratory performing the test or (b) C1-INH functional level below the lower limit of normal as defined by the laboratory performing the test or (c) Presence of a known HAE-causing C1-INH mutation.
Age Restrictions	12 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a history of moderate or severe attacks and is using Haegarda as prophylaxis for short-term use prior to surgery, dental procedures or intubation OR for long-term prophylaxis and individual has failed, or is intolerant to, or has contraindication to 17-alpha-alkylated androgens or antifibrinolytic agents.
Indications	All Medically-accepted Indications.
Off Label Uses	

Halaven

Products Affected

• HALAVEN

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Halven is used as a single agent and in a single line of therapy for recurrent or metastatic breast cancer. Member has previously received at least two chemotherapeutic regimens for locally recurrent or metastatic disease. Individual is using in combination with trastuzumab (Herceptin) in the treatment of locally recurrent or metastatic HER2+ breast cancer with: (a) Symptomatic visceral disease OR (b) Either hormone receptor-negative disease or hormone receptor-positive and endocrine refractory disease. For soft tissue sarcoma, agent is used as a single agent in a single line of therapy and has previously received at least 2 chemotherapeutic regimens for locally recurrent or metastatic disease.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Harvoni

Products Affected

- HARVONI
- ledipasvir-sofosbuvir

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Documentation is provided for a diagnosis of chronic Hepatitis C virus (CHC) infection, which includes Genotype and a positive HCV RNA result (AASLD/IDSA 2017, CDC 2013) AND Individual has received baseline evaluation for liver fibrosis to guide appropriate therapy AND Individual does not have a short life expectancy (less than 12 months owing to non-liver related comorbid conditions) that cannot be remediated by treating HCV, by transplantation or other directed therapy (AASLD/IDSA 2016).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Criteria will be applied consistent with current AASLD/IDSA guidance.
Other Criteria	Criteria will be applied consistent with current AASLD/IDSA guidance.
Indications	All Medically-accepted Indications.
Off Label Uses	

Hepsera

Products Affected

- adefovir dipivoxil
- HEPSERA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	12 years of age and older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has had a previous trial and inadequate response or intolerance to or has a contraindication to an alternative antiviral agent with a higher genetic barrier to resistance for Hepatitis B [such as entecavir or tenofovir] (AASLD 2016).
Indications	All Medically-accepted Indications.
Off Label Uses	

Hetlioz

Products Affected

• HETLIOZ

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Horizant

Products Affected

 HORIZANT ORAL TABLET EXTENDED RELEASE 300 MG, 600 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For diagnosis post herpetic neuralgia (PHN), individual has had a trial of immediate release gabapentin. For diagnosis restless leg syndrome (RLS) individual has has had a trial of or contraindication/intolerance to either pramipexole OR Ropinirole.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

HP Acthar

Products Affected

• ACTHAR

PA Criteria	Criteria Details
Exclusion Criteria	All other uses except those listed under Other Criteria section.
Required Medical Information	
Age Restrictions	For West Syndrome, infant or child less than 2 years of age.
Prescriber Restrictions	
Coverage Duration	6 month
Other Criteria	Individual is using for infantile spasm (West Syndrome).
Indications	Some FDA-approved Indications Only.
Off Label Uses	

HRM Age

Products Affected

- amitriptyline hcl
- amoxapine
- ANAFRANIL
- chlordiazepoxide-amitriptyline
- clomipramine hcl
- desipramine hcl
- doxepin hcl oral capsule
- doxepin hcl oral concentrate
- imipramine hcl
- imipramine pamoate
- NEMBUTAL
- NORPRAMIN

- nortriptyline hcl
- PAMELOR
- pentobarbital sodium
- perphenazine-amitriptyline
- phenobarbital oral elixir
- phenobarbital oral solution
- phenobarbital oral tablet 100 mg, 15 mg, 16.2 mg, 30 mg, 32.4 mg, 60 mg, 64.8 mg, 97.2 mg
- phenobarbital sodium
- protriptyline hcl
- TOFRANIL

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Prescriber must acknowledge this is a high risk medication (HRM) (as identified by American Geriatric Society) for individuals greater than 65 and medication benefits outweigh potential risk for this individual.
Age Restrictions	Individuals that are 64 years of age or younger are NOT subject to the prior authorization requirements. Prior Authorization applies to individuals that are 65 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

HRM Age AU

Products Affected

- ACTIVELLA
- ALLZITAL
- ALORA
- AMABELZ
- AMBIEN •
- AMBIEN CR
- AMRIX
- AMYTAL SODIUM •
- ANGELIQ
- ARMOUR THYROID
- ASCOMP-CODEINE
- benztropine mesylate oral
- BIJUVA
- BONJESTA
- BUPAP
- butalbital-acetaminophen ٠
- butalbital-apap-caff-cod
- butalbital-apap-caffeine •
- butalbital-asa-caff-codeine
- butalbital-asa-caffeine
- butalbital-aspirin-caffeine
- BUTISOL SODIUM ٠
- carbinoxamine maleate ٠
- carisoprodol ٠
- carisoprodol-aspirin
- carisoprodol-aspirin-codeine
- chlordiazepoxide-clidinium
- chlorzoxazone oral tablet 250 mg, 500 ٠ mg
- CHLORZOXAZONE ORAL TABLET 375 MG, 750 MG
- clemastine fumarate
- CLIMARA
- CLIMARA PRO
- COMBIPATCH
- cyclobenzaprine hcl
- cyclobenzaprine hcl er
- cyproheptadine hcl ٠
- DEMEROL
- dexchlorpheniramine maleate Y0114_20_114077_I_012

- DICLEGIS •
- digitek oral tablet 250 mcg
- digox oral tablet 250 mcg
- digoxin injection
- digoxin oral tablet 250 mcg
- diphenhydramine hcl oral •
- dipyridamole oral
- disopyramide phosphate
- DIVIGEL
- DOTTI
- doxepin hcl oral tablet ٠
- doxylamine-pyridoxine
- EDLUAR
- ELESTRIN
- ergoloid mesylates
- ESGIC
- ESTRACE ORAL
- estradiol oral
- · estradiol transdermal patch twice weekly
- estradiol transdermal patch weekly
- estradiol-norethindrone acet
- ESTROGEL
- EVAMIST
- FEMHRT LOW DOSE
- FEXMID
- FIORICET
- FIORICET/CODEINE
- FIORINAL
- FIORINAL/CODEINE #3
- fyavolv
- glyburide micronized oral tablet 1.5 mg, • 3 mg, 6 mg
- glyburide oral tablet 1.25 mg, 2.5 mg, 5 та
- glyburide-metformin oral tablet 1.25-250 ma, 2.5-500 mg, 5-500 mg
- GLYNASE ORAL TABLET 1.5 MG, 3 MG, 6 MG
- quanfacine hcl
- hydroxyzine hcl

- hydroxyzine pamoate
- INDOCIN ORAL ٠
- indomethacin
- indomethacin er ٠
- INTERMEZZO
- *jinteli*
- KARBINAL ER
- ketorolac tromethamine injection
- ketorolac tromethamine intramuscular
- ketorolac tromethamine oral •
- LANOXIN INJECTION
- LANOXIN ORAL TABLET 187.5 MCG, 250 PROCARDIA MCG
- levothyroxine-liothyronine
- LIBRAX ٠
- LOPREEZA
- LORZONE
- MEGACE ES
- megestrol acetate oral suspension 625 mg/5ml
- MENEST
- MENOSTAR
- meperidine hcl injection ٠
- meperidine hcl oral solution
- meperidine hcl oral tablet ٠
- meprobamate ٠
- METAXALL
- metaxalone ٠
- methocarbamol ٠
- methyldopa
- methyldopa-hydrochlorothiazide ٠
- MIMVEY
- MIMVEY LO
- MINIVELLE
- MOTOFEN
- nifedipine •
- norethindrone-eth estradiol ٠
- norgesic forte
- NORPACE ٠
- NORPACE CR
- np thyroid ٠
- orphenadrine citrate
- orphenadrine citrate er
- Y0114_20_114077_I_012

- orphenadrine-asa-caffeine
- orphenadrine-aspirin-caffeine
- ORPHENGESIC FORTE
- pentazocine-naloxone hcl
- PHENADOZ
- PHENERGAN
- PHRENILIN FORTE
- PREFEST
- PREMARIN ORAL
- PREMPHASE
- PREMPRO
- promethazine hcl
- promethazine vc plain
- promethazine-phenylephrine
- PROMETHEGAN
- propantheline bromide
- ROBAXIN
- ROBAXIN-750
- RYCLORA
- RYVENT
- SECONAL
- SILENOR
- SKELAXIN
- SOMA
- TENCON
- thyroid oral tablet 120 mg, 15 mg, 30 mg, 60 mg, 90 mg
- THYROID ORAL TABLET 65 MG
- trihexyphenidyl hcl
- VANADOM
- VANATOL LO
- VANATOL S
- VISTARIL
- VIVELLE-DOT
- VTOL LQ
- ZEBUTAL
- zolpidem tartrate
- zolpidem tartrate er
- ZOLPIMIST

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Prescriber must acknowledge this is a high risk medication (HRM) (as identified by American Geriatric Society) for individuals greater than 65 and medication benefits outweigh potential risk for this individual.
Age Restrictions	Individuals that are 64 years of age or younger are NOT subject to the prior authorization requirements. Prior Authorization applies to individuals that are 65 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Human Growth Hormone

Products Affected

- NORDITROPIN FLEXPRO
- OMNITROPE

PA Criteria	Criteria Details
Exclusion Criteria	Use of GH tx when applicable criteria have not been met may not be approved for, but not limited to, the following: Renal transplant, Anabolic tx, except for AIDS, provided to counteract acute or chronic catabolic illness (e.g. surgery, trauma, cancer,chronic hemodialysis) producing catabolic (protein wasting) changes in both adult and ped. Anabolic tx to enhance body mass or strength for professional, recreational or social reasons, Constitutional delay of growth and development. Cystic fibrosis, GH Tx in combo with GnRH agonist (Lupron-leuprolide) as a Tx of precocious puberty. Hypophosphatemic rickets, osteogenesis imperfecta, osteoporosis, Short Stature associated with GH insensitivity (Laron Syndrome). Therapy in older adults with normally occurring decrease in GH, who are not congenitally GH deficient and who have no evidence of organic pituitary dz (this is referred to as age-related GH deficiency). Tx of CHF, burn patients, fibromyalgia, glucocorticoid- induced growth failure. Tx of HIV lipodystrophy (fat redistribution syndrome), also referred to as altered body habitus (e.g. buffalo hump), associated with antiviral therapy in HIV-infected patients. Tx of intrauterine growth restriction (IUGR)or Russell-Silver Syndrome that does not result in SGA. Tx of obesity. Other etiologies of SS where GH has not been shown to be associated with an increase in final height, including but not limited to achondrodoplasia and other skeletal dysplasias. GH tx used for reconstruction should not continue when BA =16 yr (M), or = 14 yr (F) is reached OR Epiphyseal fusion has occurred OR Mid- parenteral height is achieved. Child over 12: an X-ray report with evidence that epiphyses have closed or SMR of 3 or more.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Required Medical Information	Initial Requests, For idiopathic GHD, has signs/sym of GHD, GV 2 SD below age-appropriate mean or ht 2.25 SD below age- appropriate mean and A subnormal (SubNL) response (less than10ng/ml)to 2GH stim tests OR Neonates with hypoglycemia and clinical/hormone evidence of hypopit and low GH (less than 10ng/ml)OR 20ther pit hormone deficiencies and low IGF-1 OR mbr has cranial irradiation with low IGF1 and normal thyroid tests.Child born SGA(birth wt or length 2 or more SD below the mean for gest age),Child fails to manifest catch up growth by age 2 yr(ht 2 or more SD below the mean for age,gender)AND Other causes for SS have been ruled out.Transitioning adolescent: completed linear growth(growth rate of less than 2cm/yr) AND either of the following: A)GH tx has been stopped at least a month and GHD reconfirmed by: 1)idiopathic isolated GHD(SubNL response to 2 GH stim tests OR SubNL response (GH conc of less than10 ng/mL)to 1 provocative test and low IGF-I/IGFBP-3) OR 2)multiple pit hormone deficiency,(SubNL response to 1 provocative GH test and/or low IGF-I/IGFBP-3)or 3) with cranial irradiation, low IGF with normal thyroid OR B) any of the following:known genetic mutation assoc with def GH production/secretion or Hypothalamic-pit tumor/structural defect or 3 other pit hormone deficiencies.Adult GHD confirmed/reconfirmed:SubNL response in adults to 2 GH stim tests (GH conc of less than or equal to 5ng/ml when using insulin induced hypoglycemia OR GH conc of less than or equal to 4.1ng/ml when using arginine)OR SubNL response to 1 stim test for adults with hypothalamic or pit dz and 1 pit hormone deficits OR 3 other pit hormone deficiencies.Initial requests for therapy in child: Reconstructive GH tx may be approved if either mean ht is at least 2.25 but less than 2.55D below the mean for age, gender and GV is less than the 10th percentile over 1yr OR mean ht is at least 2.55D below the mean for age, gender for conditions known responsive to GH.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Other Criteria	Continuation therapy in child (including reconstructive tx) when following are met: individ evaluated AND growth rate remains above 2.5cm/year (does not apply to children with prior documented hypopituitarism)(Grimberg2016). GH may be approved for Adolescents with childhood onset GHD who have completed linear growth. Treatment with GH in other populations approved when: Individual has AIDS wasting syndrome, defined as greater than 10% of baseline wt loss that cannot be explained by a concurrent illness other than HIV infection AND is being tx with antiviral therapy AND continues tx until above definition is no longer met OR individual dx with short bowel syndrome AND is receiving specialized nutritional support.
Indications	All Medically-accepted Indications.
Off Label Uses	

Humira

Products Affected

- HUMIRA PEDIATRIC CROHNS START SUBCUTANEOUS PREFILLED SYRINGE KIT 40 MG/0.8ML, 40 MG/0.8ML (6 PACK), 80 MG/0.8ML, 80 MG/0.8ML & 40MG/0.4ML
- HUMIRA PEN
- HUMIRA PEN-CD/UC/HS STARTER SUBCUTANEOUS PEN-INJECTOR KIT 40

MG/0.8ML, 80 MG/0.8ML

- HUMIRA PEN-PS/UV/ADOL HS START SUBCUTANEOUS PEN-INJECTOR KIT 40 MG/0.8ML, 80 MG/0.8ML & 40MG/0.4ML
- HUMIRA SUBCUTANEOUS PREFILLED SYRINGE KIT 10 MG/0.1ML, 10 MG/0.2ML, 20 MG/0.2ML, 20 MG/0.4ML, 40 MG/0.4ML, 40 MG/0.8ML

PA Criteria	Criteria Details
Exclusion Criteria	Using adalimumab in combination with other TNF agents, JAK inhiitors, or other biologic drugs (such as, Abatacept, anakinra or vedolizumab). Tuberculosis or other active serious infections or a history of recurrent infections. Individuals who have not had a tuberculin skin test or CDC-recommended equivalent to evaluate for latent tuberculosis prior to initiating adalimumab.
Required Medical Information	For Chronic moderate to severe plaque psoriasis with either of the following: Patient has greater than 5% of body surface area with plaque psoriasis OR Less than or equal to 5% of body surface area with plaque psoriasis involving sensitive areas or areas that significantly impact daily function (such as palms, soles of feet, head/neck, or genitalia) AND agent is used to reduce signs/symptoms OR to induce/maintain clinical response.
Age Restrictions	Patient is 18 years of age or older for all indications except JIA, non-infectious Uveitis, Hidradenitis Suppurativa (HS) and Crohns disease. Patient must be at least 2 years old for JIA and non- infectious uveitis. Patient must be at least 6 years of age for Crohns disease. Patient must be at least 12 years old for HS.
Prescriber Restrictions	
Coverage Duration	1 year

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Other Criteria	For moderate to severe RA, individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional therapy [nonbologic DMARDs (such as methotrexate, sulfasalazine, leflunomide, or hydroxylchloroquine)] (ACR 2015). For moderate to severe Psoriatic Arthritis, individual has had an inadequate response to, is intolerant of, has medical contraindication to ONE conventional therapy [non-biologic DMARDs (such as methotrexate, sulfasalazine or leflunomide)]. For moderate to severe Ankylosing Spondylitis (AS), individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional therapy (such as NSAIDs or nonbiologic DMARDs (such as sulfasalazine). For moderate to severe Crohn's disease, individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional therapy (such as NSAIDs or nonbiologic DMARDs (such as sulfasalazine). For moderate to severe Crohn's disease, individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional therapy (e.g. s-ASA products, sulfasalazine, systemic corticosteroids, or immunosuppressants). For chronic moderate to severe plaque psoriasis, individual has had an inadequate response to, is intolerant of, or has a medical contraindication to phototherapy or ONE other systemic therapy (e.g. methotrexate, acitretin, or cyclosporine). For moderate to severe Polyarticular Juvenile Idiopathic Arthritis (PJIA), individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional therapy [nonbologic DMARDs (such as methotrexate)] (ACR 2011). For moderate to severe Ulcerative Colitis (UC), individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional therapy [nonbologic DMARDs (such as metical contraindication to ONE conventional therapy [nonbologic DMARDs (such as metical contraindication to ONE conventio

PA Criteria	Criteria Details
	progressive, treatment-refractory Sarcoidosis (Sweiss 2014), individual has had an inadequate response to, is intolerant of or has a contraindication to systemic corticosteroids AND nonbiologic DMARDs (such as methotrexate or azathioprine). For moderate to severe Hidradenitis Suppurativa (Hurley stage II or Hurley stage III disease) AND has had an inadequate response to, is intolerant of, or has a medical contraindication to conventional therapy (such as oral antibiotics).
Indications	All Medically-accepted Indications.
Off Label Uses	

Humulin U500

Products Affected

- HUMULIN R U-500 (CONCENTRATED)
- HUMULIN R U-500 KWIKPEN

PA Criteria	Criteria Details
Exclusion Criteria	For use as a continuous subcutaneous infusion.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual has a diagnosis of diabetes mellitus AND requires more than 200 units of insulin per day.
Indications	All Medically-accepted Indications.
Off Label Uses	

Ibrance

Products Affected

• IBRANCE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Iclusig

Products Affected

• ICLUSIG ORAL TABLET 15 MG, 45 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Idhifa

Products Affected

• IDHIFA ORAL TABLET 100 MG, 50 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has confirmed (written or verbal attestation) isocitratedehydrogenase-2 (IDH2) mutation.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Ilaris

Products Affected

• ILARIS

PA Criteria	Criteria Details
Exclusion Criteria	Tuberculosis, invasive fungal infection, other active serious infection, or a history of recurrent infection. Individual who have not had a tuberculin skin test or a CDC-recommended equivalent to evaluate for latent tuberculosis prior to initiating Ilaris (canakinumab). Using Ilaris in combination with other biologic disease-modifying antirheumatic drugs (DMARDs), such as, tumor necrosis factor (TNF) antagonists, IL-1R antagonists, Janus kinase inhibitors (for example, tofacitinib citrate), selective co-stimulation modulators or IL-6 receptor antagonists.
Required Medical Information	
Age Restrictions	For cryopyrin-associated periodic syndromes age 4 years and older and for systemic juvenile idiopathic arthritis 2 years of age and older.
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	FFor SIJA, individual has had an inadequate response to, intolerant of, or has a medical contraindication to ONE corticosteroids or nonsteroidal anti-inflammatory drugs (NSAIDs) AND may be used alone or in combination with corticosteroids, methotrexate or NSAIDs. For FMF, individual has active type 1 FMF disease with genetic confirmation of the diagnosis (MEFV gene exon 10 mutation) and confirmed recurrent, active disease (that is, at least one flare per month) and has failed to respond to, or is intolerant of colchicine therapy. For HIDS/MKD, individual has HIDS with genetic confirmation of the diagnosis by deoxyribonucleic acid (DNA) analysis or enzymatic studies (that is, mutations in the MVK gene or markedly reduced mevalonate kinase activity) and confirmed prior history of greater than or equal to three febrile acute flares within a 6-month period when not receiving prophylactic treatment. For TRAPS, genetic confirmation of the diagnosis (TNFRSF1A gene mutation) and has chronic or recurrent disease activity defined as six flares in a 12-month period.
Indications	All Medically-accepted Indications.
Off Label Uses	

Ilumya

Products Affected

• ILUMYA

PA Criteria	Criteria Details
Exclusion Criteria	Using in combination with other biologic drugs or phototherapy. Tuberculosis, other active serious infections, or a history of recurrent infections. Individual has not had a tuberculin skin test or a Centers for Disease Control and Prevention-recommended equivalent test to evaluate for latent tuberculosis prior to initiating Ilumya.
Required Medical Information	Diagnosis of moderate to severe plaque psoriasis with either of the following: Patient has greater than 5% of body surface area with plaque psoriasis OR Less than or equal to 5% of body surface area with plaque psoriasis involving sensitive areas or areas that would significantly impact daily function (such as palms, soles of feet, head/neck, or genitalia).
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year.

PA Criteria	Criteria Details
Other Criteria	For of chronic plaque psoriasis, individual has had an inadequate response to, is intolerant of, or has a medical contraindication to phototherapy or any ONE systemic therapy (for example acitretin, methotrexate, cyclosporine) AND individual has had a trial of and an inadequate response or is intolerant to Humira (adalimumab) OR Enbrel (etanercept). For any of the above indications, if the TNF agent (Enbrel/Humira) tried and failed are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Ilumya or (b) Individuals age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis OR individual has either concomitant clinical condition: (a) Demyelinating disease or (b) Heart failure with documented left ventricular dysfunction, Ilumya may be allowed without trial of preferred TNF agents (Enbrel/Humira).
Indications	All Medically-accepted Indications.
Off Label Uses	

Imbruvica

Products Affected

- IMBRUVICA ORAL CAPSULE 140 MG, 70 MG
- IMBRUVICA ORAL TABLET 140 MG, 280 MG, 420 MG, 560 MG

DA Cuitoria	Criteria Detaile
PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Imfinzi

Products Affected

• IMFINZI

PA Criteria	Criteria Details
Exclusion Criteria	History of immunodeficiency or severe autoimmune disease. Requires systemic immunosuppression, active immune-medicated disease, severe or life-threatening infections or untreated central nervous system (CNS) metastases. Has received treatment with another anti-PD-1 or anit-PD-L1 agent.
Required Medical Information	For locally advanced or metastatic urothelial carcinoma, Inoperable or metastatic transitional-cell urothelial carcinoma histologically or cytologically confirmed. For locally advanced, unresectable non- small cell lung cancer, histologically or cytologically confirmed stage III and current Eastern Cooperative Oncology Group performance status 0-2.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has inoperable or metastatic urothelial carcinoma AND Either the disease has progressed during or following platinum- containing therapy OR disease has progressed within 12 months of neoadjuvant or adjuvant treatment with platinum-containing therapy. For locally advanced, unresectable non-small cell lung cancer (NSCLC), disease has not progressed after definitive chemoradioation and disease has progressed or individual has reached a maximum of 12 months of treatment.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Imlygic

Products Affected

• IMLYGIC

PA Criteria	Criteria Details
Exclusion Criteria	Individual is immunocompromised. Individual is pregnant.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has diagnosis of unresectable melanoma AND is using as intralesional treatment for one of the following: a) Stage III disease with clinical or satellite/in-transit metastases b) Local satellite recurrence of disease c) in-transit recurrence of disease.
Indications	All Medically-accepted Indications.
Off Label Uses	

Increlex

Products Affected

• INCRELEX

PA Criteria	Criteria Details
Exclusion Criteria	Individual has suspected or known malignancy. Individual has closed. Individual has of secondary IGF-1 deficiency (for example, due to GH deficiency, untreated malnutrition, untreated hypothyroidism).
Required Medical Information	For initial treatment of growth failure associated with severe primary IGF-1 deficiency as defined by: Height standard deviation score of less than or equal to -3.0 AND Basal IGF-1 standard deviation score of less than or equal to -3.0 AND normal or elevated growth hormone levels (greater than 10ng/mL on standard GH stimulation tests) are present OR GH gene deletion who have development of neutralizing antibodies to GH.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For Continuation of treatment with Increlex (mecasermin), Growth velocity is greater than or equal to 2cm (greater than equal to 2.0 cm) total growth in 1 year AND Final adult height has not been reached.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Ingrezza

Products Affected

- INGREZZA ORAL CAPSULE 40 MG, 80
 MG
- INGREZZA ORAL CAPSULE THERAPY PACK

PA Criteria	Criteria Details
Exclusion Criteria	Has congenital long QT-syndrome or arrhythmia associated with a prolonged QT interval. Individual is currently using a strong CYP 3A4 inducer (examples, rifampin, carbamazepine, phenytoin, St. John's wort). Individual is currently using a monoamine oxidase inhibitor (MAOI) (examples, isocarboxazid, phenelzine, selegiline)
Required Medical Information	Tardive dyskinesia confirmed by the following (DSM-5): A) At least 60 days of stable (drug, dose) medication exposure (either typical or first generation antipsychotic agents [such as, chlorpromazine, haloperidol, fluphenazine], atypical or second-generation antipsychotic agents [such as, clozapine, risperidone, olanzapine, quetiapine, aripiprazole], or certain dopamine receptor-blocking used in treatment of nausea and gastroparesis [such as prochlorperazine, promethazine, metoclopramide] AND B) Presence of involuntary athetoid or choreiform movements lasting at least 30 days.
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Requests for continuation of therapy may be approved for individuals who meet the following criteria: Individual has experienced an improvement in symptoms deemed to be clinically significant by the provider.
Indications	All Medically-accepted Indications.
Off Label Uses	
V0114 20 114077	L 012 December 2020

Y0114_20_114077_I_012

Inlyta

Products Affected

• INLYTA ORAL TABLET 1 MG, 5 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Inqovi

Products Affected

• INQOVI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Inrebic

Products Affected

• INREBIC

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	Individual is 18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Interferons for MS

- AVONEX
- AVONEX PEN
- AVONEX PREFILLED
- BETASERON

r	
PA Criteria	Criteria Details
Exclusion Criteria	Individuals with primary progressive MS. Individuals with secondary progressive MS without relapsing disease. Concurrent use with other MS disease modifying agents (such as Aubagio, Gilenya, tecfidera, tysabri, Lemtrada, Ocrevus, Copaxone/Glatopa, Extavoa. Rebif, Avonex, Plegridy or Betaseron).
Required Medical Information	Individual has experienced a first clinical episode and has MRI features consistent with multiple sclerosis OR Individual has a diagnosis of relapsing multiple sclerosis (RMS) OR Individual has secondary progressive MS (SPMS) with a history of superimposed relapses (AHFS).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Intuniv

Products Affected

- guanfacine hcl er INTUNIV

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual is using for Attention Deficit Hyperactivity Disorder/Attention Deficit Disorder (ADHD/ADD).
Age Restrictions	Individual is 6 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Istodax

- ISTODAX (OVERFILL)
- romidepsin

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Isturisa

Products Affected

 ISTURISA ORAL TABLET 1 MG, 10 MG, 5 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

ITRACONAZOLE

- itraconazole oral capsule
- SPORANOX ORAL CAPSULE
- SPORANOX PULSEPAK
- tolsura

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	For fingernail/toenail onychomycosis 12 weeks. For all other indications 1 year.
Other Criteria	For second-line non-onychomycosis indications include tinea infections (including, but limited to tinea versicolor, tinea cruris, tinea corporis, tinea pedis, tinea manuum, and tinea capitis) where the individual has received at least one prior topical therapy: clotrimazole, ketoconazole, econazole, or nystatin.
Indications	All Medically-accepted Indications.
Off Label Uses	

Products Affected

- GAMUNEX-C
- OCTAGAM

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Hyperimmunoglobulinemia E synd when dx is evidenced by high level of serum IgE. Autoimmune mucocutaneous blistering dx when mbr had inadeq response/intolerance/contraindication to other tx such as corticosteroids,immunosuppressants. For autoimmune neutropenia, active INFECT is excluded as cause. For tx of 1 neurologic DZ: A) Lambert-Eaton myasthenic syndrome when having muscle weakness AND dx confirmed w/characteristic electrodiagnostic (ED) finding using nerve conduction tests, repetitive nerve stimulation (RNS), exercise testing or single fiber electromyography (SFE) or presence of antibodies (AB) directed against voltage-gated Ca channels B) Myasthenia Gravis (MG) and dx confirmed by presence of AB against the acetylcholine receptor or muscle specific tyrosine kinase or characteristic ED findings using RNS or SFE AND using for worsening sx or exacerbation or short-term therapy as immunosuppressive tx is taking effect or MAINT therapy of MG when mbr had inadeq response/intolerance/contraindication to other tx such as steroids, immunosuppressants C) Chronic Inflammatory Demyelinating Polyneuropathy (CIDP), as INIT trial up to 12wks when muscle weakness or sensory dysfx is caused by neuropathy in more than 1 limb and nerve conduction studies or diagnostic criteria confirm evidence of demyelinating neuropathy and other polyneuropathies. For cont use of CIDP, clinically sig improvement in neurological sx on exam and cont need is shown by attempts on annual basis to titrate dose or interval of therapy result in sx worsening. As INIT exam(up to 12wks), clinical presentation w/electrodiagnostic test confirm MMN. For MMN cont use, mbr had improvement in strength and fx and need shown by attempts annually to titrate dose or interval therapy results in worse sx.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Other Criteria	Tx of primary humoral immunprimaryodeficiency (PI) when hx of recurrent sinopulmonary infection (SI) req ABX tx AND lack of/inadeq response to immunization AND no evidence of renal (nephrotic synd) and GI as causes of hypogammaglobulinemia (HGG) AND INIT pre-tx total serum IgG below the lower limit of age adj lab ref range or more than 2SD below adj mean. hyperimmunoadj mean AND hx of recurrent SI requiring ABX therapy AND lack of/inadeq response to immunization OR Use for ONE: A) B-cell CLL w/ hx of recurrent bacterial or active INFECT not responding to antimicrobial therapy and HGG w/ total IgG less than 500mg/dL B)Multiple myeloma with hx of recurrent bacterial or clinically severe INFECT and HGG with total IgG less than 500mg/dL C) HIV infected children to prevent opportunistic bacterial infection w/HGG (IgG less than 400mg/dL) or recurrent INFECT D) PARVO B19 chronic INFECT and severe anemia assoc w/bone marrow suppression OR using in context of transplant for ONE: 1) hematopoietic stem cell transplant 2) Solid organ transplantation including prior desensitization for transplantation for suppression of panel reactive anti-HLA antibody (AB) in ppl with high panel reactive AB (PRA) levels to human leukocyte antigens OR Transplant recipients at risk of CMV 3) Transplant recipients experiencing AB-mediated rejection w/ donor-specific AB OR for tx of ONE autoimmune DZ: A) ITP w/either active bleed or platelet count less than 30,000 mcL B) Fetal alloimmune TCP or fetal blood sample shows TCP C) Isoimmune hemolytic dx of newborn, tx of severe hyperbilirubinemia D) Dermatomyositis (DMM) or polymyositis when mbr had inadeq response/intolerance/contraindication to other tx,e.g., corticosteroids, immunosuppressive agents AND X confirmed having at least 4 sx: weak trunk/proximal extremities, high serum CK or aldolase levels, unexplainable muscle pain, electromyography

PA Criteria	Criteria Details
	findings, anti-Jo-1 AB, arthralgia/arthritis w/out joint destruction, sign of systemic inflamm, e.g.,fever/elevated C-reactive protein/high SED rate or inflamm myositis seen on muscle biopsy AND using for DMM and skin lesions present. For 1 MISC DX: post- exposure prophylaxis to stop measles, give in 6dys of exposure (not w/VACC having measles virus), eligible/exposed/non-immune mbr will get a VACC w/measles virus greater than/equal to 8 mth after Ig admin and used in mbrs at risk of severe dx/complications and no evidence of measles immunity in PREG or severely immunocompromised ppl OR for Kawasaki Dz tx initiated w/in 10dys of onset and tx for more than 5dys.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Ixempra

Products Affected

• IXEMPRA KIT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Jadenu

- deferasirox
- deferasirox granules
- JADENU
- JADENU SPRINKLE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	For dx non-transfusion-dependent thalassemia (NTDT) syndrome, 10 years of age or older. For dx of chronic iron overload, 2 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Jakafi

Products Affected

 JAKAFI ORAL TABLET 10 MG, 15 MG, 20 MG, 25 MG, 5 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Jetrea

Products Affected

• JETREA

PA Criteria	Criteria Details
Exclusion Criteria	"Individual has any of the following: Proliferative diabetic retinopathy, Neovascular age-related macular degeneration, Retinal vascular occlusion, Aphakia, High myopia (more than ?8 diopters), Uncontrolled glaucoma, Macular hole greater than 400 ?m in diameter, Vitreous opacification, Lenticular or zonular instability, History of retinal detachment in either eye, Prior vitrectomy in the affected eye, Prior laser photocoagulation of the macula in the affected eye, Prior treatment with ocular surgery, intravitreal injection or retinal laser photocoagulation in the previous 3 months."
Required Medical Information	Posterior segment optical coherence tomography (OCT) demonstrates all of the following: there is vitreous adhesion within 6-mm of the fovea (center of macula) AND elevation of the posterior vitreous cortex (outer layer of the vitreous). Individual has best-corrected visual acuity of 20/25 or worse in the eye to be treated with ocriplasmin.
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Jevtana

Products Affected

• JEVTANA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For hormone-refractory metastatic prostate cancer, individual has a Eastern Cooperative Oncology Group (ECOG) performance status is 0-2.
Indications	All Medically-accepted Indications.
Off Label Uses	

Juxtapid

Products Affected

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

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has a clinical diagnosis of homozygous familial hypercholesterolemia (HoFH), based on the presence of the following (Cuchel 2014, Singh 2015): (A) Genetic confirmation of two (2) mutant alleles at the LDL receptor, apoB, PCSK9 or ARH adaptor protein (LDLRAP1) gene locus OR (B) One of the following: 1. an untreated LDL-C concentration greater than 500 mg/dL (13 mmol/L) OR 2. Treated LDL-C greater than or equal to 300 mg/dL (7.76 mmol/L) AND one of the following: (i) cutaneous or tendonous xanthoma before age of 10 years OR (ii) untreated LDL- C levels consistent with heterozygous familial hypercholesterolemia in both parents (greater than 190 mg/dL).
Age Restrictions	18 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has had an adequate trial and titration of Repatha and achieved suboptimal lipid lowering response.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Jynarque

Products Affected

- JYNARQUE ORAL TABLET
- JYNARQUE ORAL TABLET THERAPY PACK

PA Criteria	Criteria Details
Exclusion Criteria	Individual has an uncorrected abnormal blood sodium level or urinary outflow obstruction OR unable to sense or appropriately respond to thirst OR has dx of hypovolemia OR is anuric OR has underlying significant liver disease (not including uncomplicated polycystic liver disease) OR will be concurrently utilizing a strong CYP3A inhibitor (such as clarithromycin, ketoconazole, itraconazole, ritonavir, indinavir, nelfinavir, saquinavir or nefazodone)
Required Medical Information	
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Kadcyla

Products Affected

• KADCYLA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Tumor(s) have been evaluated with an assay validated to predict HER2 positive (HER2+) protein overexpression. Individuals are considered HER2 positive (HER2+) as documented by one of the following, immunohistochemistry (IHC) 3+ or In Situ hybridization (ISH) positive by any of the following: Single probe average HER2 copy number greater than or equal to 6.0 signals/cell OR Dual- probe HER2/CEP 17 ratio greater than or equal to 2.0 OR Dual- probe HER2/CEP 17 ratio less than 2.0 with an average HER2 copy number greater than or equal to 6.0 signals/cell.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For metastatic breast cancer, individual has previously received trastuzumab and a taxane, separately or in combination. AND has either received prior therapy for metastatic disease OR developed disease recurrence during or within six (6) months of completing adjuvant therapy. Kadcyla is only used in a single line of therapy.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Kalbitor

Products Affected

• KALBITOR

PA Criteria	Criteria Details
Exclusion Criteria	Prophylaxis for HAE attacks.
Required Medical Information	HAE to be confirmed by a C4 level below the lower limit of normal (as defined by the laboratory performing the test) and either a C1 inhibitor antigenic level below the lower limit of normal (as defined by the lab performing test) or a C1 inhibitor functional level below the lower limit of normal (as defined by the lab performing the test).
Age Restrictions	12 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a history of moderate or severe attacks (for example, airway swelling, severe abdominal pain, facial swelling, nausea and vomiting, painful facial distortion) and using Kalbitor for acute HAE attacks.
Indications	All Medically-accepted Indications.
Off Label Uses	

Kalydeco

- KALYDECO ORAL PACKET
- KALYDECO ORAL TABLET

PA Criteria	Criteria Details
Exclusion Criteria	Using Kalydeco (ivacaftor) monotherapy, without concurrent use of lumacaftor or tezacaftor, for the F508del mutation in the CFTR gene.
Required Medical Information	Individual has a diagnosis of cystic fibrosis (CF). Individual has confirmed (verbal or written attestation) mutation positive result in the cystic fibrosis membrane conductance regulator (CFTR) gene.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Kanuma

Products Affected

• KANUMA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Diagnosis has been confirmed by one of the following: (1) A dried blood spot (DBS) test demonstrating deficient lysosomal acid lipase (LAL) activity or (2) Documented molecular genetic test revealing mutations in the lipase A, lysosomal acid type (LIPA) gene.
Age Restrictions	For diagnosis of lysosomal acid lipase deficiency (LAL-D) disorder [also known as Wolman disease (WD)], individual is equal to or less than 4 year of age. For diagnosis of LAL-D disorder [also known as cholesteryl ester storage disease (CESD)], individual is greater than 4 years of age.
Prescriber Restrictions	
Coverage Duration	Initial request: 6 months, Maintenance: 1 year
Other Criteria	FFor diagnosis of LAL-D disorder [also known as cholesteryl ester storage disease (CESD)], Individual has a baseline alanine aminotransferase (ALT) level greater than or equal to 1.5 times the upper limit of normal (ULN). Maintenance therapy requests for Kanuma (sebelipase alfa) may be approved if the following criteria are met: Individual has had clinical improvement in symptoms and/or lab values.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Keveyis

Products Affected

• KEVEYIS

PA Criteria	Criteria Details
Exclusion Criteria	Individual has a diagnosis of hepatic insufficiency OR severe pulmonary obstruction OR a known hypersensitivity to sulfonamides OR individual is concurrently using high-dose aspirin.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Initial 3 months, renewal 1 year
Other Criteria	For initial therapy, individual experiences greater than or equal to one episode of muscle weakness per week. For continuation therapy individual has provided written or verbal attestation that the individual has achieved and sustained clinically significant improvement in the number of episodes of muscle weakness experienced per week.
Indications	All Medically-accepted Indications.
Off Label Uses	

Kevzara

Products Affected

• KEVZARA

PA Criteria	Criteria Details
Exclusion Criteria	Using in combination with other biologic disease modifying anti- rheumatic drugs such as anti-CD20 monoclonal antibodies, IL-1R antagonists, selective co-stimulation modulators, or tumor necrosis factor antagonists. At initiation of therapy, absolute neutrophil count less than 2000/mm3 , platelet count less than 150,000/mm3 , alanine aminotransferase or aspartate aminotransferase greater than 1.5 times the upper limits of normal. Tuberculosis, invasive fungal infection, or other active serious infections or a history of recurrent infections. Individual has not had a tuberculin skin test or Centers for Disease Control and Prevention-recommended equivalent to evaluate for latent tuberculosis prior to initiating sarilumab.
Required Medical Information	
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For Rheumatoid Arthritis, individual has had an inadequate response to, is intolerant of, or has a contraindication to ONE conventional therapy [nonbiologic DMARDs (such as methotrexate)] AND individual has had an inadequate response or is intolerant to Humira OR Enbrel. For any of the above indications, if the TNF agent (Enbrel/Humira) tried and failed are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Kevzara or (b) Individuals age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis OR individual has either concomitant clinical condition: (a) Demyelinating disease or (b) Heart failure with documented left ventricular dysfunction, Kevzara may be allowed without trial of preferred TNF agents (Enbrel/Humira).
Indications	All Medically-accepted Indications.
Off Label Uses	

Keytruda

Products Affected

• KEYTRUDA

PA Criteria	Criteria Details
Exclusion Criteria	Previous treatment with another programmed death receptor-1 (PD-1) blocking antibody agent. OR Presence of human immunodeficiency virus (HIV) infection, hepatitis B virus infection, or hepatitis C virus infection. Receiving therapy for an autoimmune disease or chronic condition requiring treatment with an immunosuppressant.
Required Medical Information	Current ECOG (Eastern Cooperative Oncology Group) performance status of 0-2.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For melanoma, used as single agent and tx is 1st line in untreated dz or 2nd line in dz progression while receiving or since completing most recent therapy. For adv melanoma w/lymph node, resected high risk stage III. For colorectal cancer, used as single agent, primary tx as single agent for unresectable metachronous metastases (defective mismatch repair/high microsatellite instability [dMMR/MSIH] only) and previous adjuvant FOLFOX or Cape OX w/in past 12mon or subsequent therapy as single agent if nivolumab or pembrolizumab not previously given following oxaliplatin-irinotecan and fluropyrimidine based therapy or oxaliplatin-irinotecan. For adv/metastatic NSCLC, used as 1st line, single agent, cytologically confirmed stage III or IV, tumor expresses PD_L1 gene on at least 1% or grtr of tumor cells, no sensitizing epidermal growth factor receptor (EGFR) mutations or anaplastic lymphoma kinase (ALK) translocations. For 1st line adv/metastatic nonsquamous NSCLC, used in combo w/pemetrexed and carboplatin, cytologically confirmed stage IIIb or IV, no sensitizing EFGR mutation or ALK translocations. For 1st line metastatic squamous NSCLC, used in combo with carboplatin and paclitaxel or nab-paclitaxel and cytologically confirmed stage IV and has not undergone prev systemic tx for dz. For CONT/MAINT of recurrent/metastatic nonsquamous NSCLC, used as single agent, given 1st line as part of pembrolizumab/carboplatin/paclitaxel regimen, achieved tumor response or stable dz after initial cytotoxic therapy. For CONT/MAINT therapy of recurrent/metastatic Squamous NSCLC, Used 2nd line, single agent, given 1st line as part of pembrolizumab/carboplatin/paclitaxel regimen, achieved dz progression rafter platinum-containing chemo, ALK or EGFR genomic tumor aberrations prisent and dz progression on FDA approved therapy for aberrations prior to

PA Criteria	Criteria Details
	receiving pembrolizumab. For small cell lung cancer, used as single agent and demonstrated dz relapse w/in 6mon after complete or partial response or stable dz with initial tx or primary progressive dz and not received another PD-1 agent. For cHL, except for those with lymphocyte-predominant HL, or who have relapsed after 3 or more prior lines of therapy. For Merkel-cell carcinoma (MCC), used as single agent, Presence of metastatic or recurrent locoregional MCC determined to be not amendable to definitive surgery or radiation therapy. For unresectable or metastatic solid tumors (dMMR/MSIH only), used as single agent. For hepatocellular carcinoma, used as single agent, demonstrated dz progression or intolerance on or after tx w/an approved 1st line agent.
Indications	All Medically-accepted Indications.
Off Label Uses	

Kineret

Products Affected

• KINERET

PA Criteria	Criteria Details
Exclusion Criteria	Individual is using Kineret in combination with other tumor necrosis factor (TNF) antagonists. Tuberculosis or other active serious infections or a history of recurrent infections. Individual has not had a tuberculin skin test (TST) or Centers for Disease Control (CDC) Prevention-recommended equivalent to evaluate for latent tuberculosis prior to initiating Kineret. In combination with Xeljanz (tofacitinib) or with NONTNF immunomodulatory drugs [such as but not limited to Actemra (tocilizumab) or Orencia (abacept)].
Required Medical Information	
Age Restrictions	For RA, individual is 18 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For RA, Individual has had an inadequate response to is intolerant of or has a contraindication to at least ONE conventional therapy [disease modifying anti-rheumatic agent (DMARD)] AND Individual has had a trial and an inadequate response to or intolerance to Humira (adalimumab) OR Enbrel (etanercept) or the TNF agent (Humira(adalimumab)/Enbrel(etanercept)) tried and failed are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with kineret or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis OR the individual has either concomitant clinical condition: 1) Demyelinating disease or 2) Heart failure with documented left ventricular dysfunction. Kineret may be allowed without trial of preferred TNF agents (Enbrel/Humira).
Indications	All Medically-accepted Indications.
Off Label Uses	

Kisqali

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

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Korlym

Products Affected

• KORLYM

PA Criteria	Criteria Details
Exclusion Criteria	History of unexplained vaginal bleeding. Current endometrial hyperplasia with atypia or endometrial carcinoma. Diagnosis of severe hepatic impairment (Child Pugh Class C). Concomitant use with any of the following: (1) Long term systemic corticosteroids for serious medical conditions or illnesses OR (2) Simvastatin or lovastatin OR (3) CYP3A substrates with narrow therapeutic ranges (such as but not limited to cyclosporine, fentanyl, sirolimus, tacrolimus) OR (4) Agents or co-morbid conditions which prolong the QT interval
Required Medical Information	
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Initial therapy: Individual is not a candidate for surgery that is expected to correct the cause of endogenous Cushing's Syndrome OR disease persists or recurs following surgery intended to correct the cause of endogenous Cushing's Syndrome. For continuation of therapy: Individual continues to meet the initial request approval criteria AND has experienced an improvement in or stabilization of glucose control as assessed by fasting serum glucose test, oral glucose tolerance test or hemoglobin A1c test.
Indications	All Medically-accepted Indications.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

Koselugo

Products Affected

• KOSELUGO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	2 years old or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Krystexxa

Products Affected

• KRYSTEXXA

PA Criteria	Criteria Details
Exclusion Criteria	Individual has asymptomatic hyperuricemia. Individual has a known glucose-6-phosphate dehydrogenase (G6PD) deficiency.
Required Medical Information	Individual has 1 or more of the following: 3 or more gout flares in the previous 18 months OR 1 or more tophus OR History of chronic gouty arthropathy, defined clinically or radiographically as joint damage due to gout. Individual has a baseline serum uric acid of 6 mg/dL or greater prior to initiating pegloticase.
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has chronic, treatment-refractory and has failed to respond to, is intolerant of, or has a medical contraindication to ONE of the following conventional therapies: A xanthine oxidase inhibitor (allopurinol or febuxostat) or combination therapy of 1 xanthine oxidase inhibitor given with a uricosuric agent (for example, probenecid).
Indications	All Medically-accepted Indications.
Off Label Uses	

Kuvan

- KUVAN
- sapropterin dihydrochloride

PA Criteria	Criteria Details
FACILLEIIa	
Exclusion Criteria	If blood phenylalanine levels do not decrease from baseline at a dose of 10mg/kg/day administered for up to one month. The dose may be increased up to 20mg/kg/day. Individuals are non-responders if phenylalanine levels do not decrease after 1 month and tx should be discontinued
Required Medical Information	For initial requests, Individual has Dx of Hyperphenylalaninemia (HPA) due to tetrahydrobiopterin-(BH4) responsive PKU. For continued use, individual is showing signs of continuing improvement as evidenced by blood phenylalanine levels.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Initial 8 weeks, 1 year for continuation
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

KYNAMRO

Products Affected

• KYNAMRO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has a diagnosis of homozygous familial hypercholesterolemia (HoFH), based on the presence of the following (Cuchel 2014, Singh 2015): (A) Genetic confirmation of two (2) mutant alleles at the LDL receptor, apoB, PCSK9 or ARH adaptor protein (LDLRAP1) gene locus OR (B) One of the following: 1. an untreated LDL-C concentration greater than 500 mg/dL (13 mmol/L) OR 2. Treated LDL-C greater than or equal to 300 mg/dL (7.76 mmol/L) AND one of the following: (i) cutaneous or tendonous xanthoma before age of 10 years OR (ii) untreated LDL- C levels consistent with heterozygous familial hypercholesterolemia in both parents (greater than 190 mg/dL).
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has had an adequate trial and titration of Repatha and achieved suboptimal lipid lowering response.
Indications	All Medically-accepted Indications.
Off Label Uses	

Kyprolis

Products Affected

KYPROLIS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 Year
Other Criteria	For multiple myeloma (for primary treatment) and being used in combination with lenalidomide plus dexamethasone. For the treatment of Waldenstrom's macroglbulinemia when the following criteria are met: (a) Used as a primary agent, in combination with rituximab and dexamethasone OR (b) Used for relapsed disease when the primary therapy of carfilzomib, rituximab and dexamethasone was given and relapse is greater than 12 months after therapy.
Indications	All Medically-accepted Indications.
Off Label Uses	

Lartruvo

Products Affected

• LARTRUVO

PA Criteria	Criteria Details	
Exclusion Criteria		
Required Medical Information		
Age Restrictions	Individual is 18 years of age or older	
Prescriber Restrictions		
Coverage Duration	1 year	
Other Criteria	Individual has a histologically confirmed diagnosis tissue sarcoma (locally advanced or metastatic) n treated with an anthracycline and Olaratumab is u combination with doxorubicin and Radiotherapy of curative treatment option and Individual's current Cooperative Oncology Group (ECOG) performance Individual has a histologically confirmed diagnosis tissue sarcoma (locally advanced or metastatic) n treated with an anthracycline and Olaratumab is u combination with doxorubicin and after at least 8 doxorubicin or earlier discontinuation of doxorubic and then if so chosen, continuing olaratumab as n the absence of unacceptable toxicities until diseas and Radiotherapy or surgery is not a treatment of Individual's current Eastern Cooperative Oncology performance status is 0-2.	ot previously used in or surgery is not a : Eastern e status is 0-2. s of late stage soft ot previously used in cycles with cin due to toxicity, monotherapy in se progression ption and
Indications	All Medically-accepted Indications.	
Y0114_20_114077	7_I_012 Decer	nber 2020

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

Lazanda

Products Affected

• LAZANDA

PA Criteria	Criteria Details
Exclusion Criteria	Treatment of acute or postoperative pain, migraine headache pain OR non-cancer related breakthrough pain
Required Medical Information	
Age Restrictions	Individual is 18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a diagnosis of active cancer (such as, metastatic or locally invasive cancer for which the individual is currently seeking treatment) with breakthrough cancer pain (provide diagnosis) AND has tried and failed fentanyl citrate lozenge (generic Actiq) AND Individual is already receiving opioid therapy and is TOLERANT to opioid therapy as defined as receiving around the clock medicine consisting of one of the following: At least 60mg morphine per day, OR At least 25mcg/hr transdermal fentanyl/hour, OR At least 30mg of oxycodone daily, OR At least 8mg of oral hydromorphone daily, OR At least 25mg of oral oxymorphone daily, OR An equianalgesic dose of another opioid for a week or longer. Individual will also continue around the clock opioids when taking Lazanda (fentanyl) for cancer related breakthrough pain.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Lemtrada

Products Affected

• LEMTRADA

PA Criteria	Criteria Details
Exclusion Criteria	Individual with primary progressive MS. Individual with secondary progressive MS. Concurrent use with other MS disease modifying agents (such as Aubagio, Gilenya, Tecfidera, Tysabri, Ocrevus, Copaxone/Glatopa, Extavia, Rebif, Avonex, Plegridy, Betaseron).
Required Medical Information	Individual is Human immunodeficiency virus (HIV) negative.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has received prior treatment with at least two alternative drug therapies indicated for the treatment of MS (for example, interferons, glatiramer) and failed to achieve an adequate response.
Indications	All Medically-accepted Indications.
Off Label Uses	

Lenvima

Products Affected

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

 - LENVIMA (8 MG DAILY DOSE)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Letairis

Products Affected

- ambrisentan
- LETAIRIS

PA Criteria	Criteria Details
Exclusion Criteria	Individual has idiopathic pulmonary fibrosis (IPF). Individual has a diagnosis of moderate (Child-Pugh Class B) or severe (Child-Pugh Class C) hepatic impairment. Individual is initiating therapy and has a diagnosis of clinically significant anemia/severe anemia. Using in combination with other endothelin receptor antagonist (ERA) agents, such as but not limited to Opsumit (macitentan) or Tracleer (bosentan).
Required Medical Information	Individual has catheterization-proven diagnosis of pulmonary arterial hypertension (PAH) (WHO Group 1) AND individual has WHO Functional Class II-IV symptoms.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Leukine

Products Affected

• LEUKINE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individuals who are at high risk for infection-associated complications demonstrated by any of the following: Expected prolonged (greater than 10 day) and profound (less than 0.1 x 10to the power of 9/L) neutropenia, Age greater than 65 years, Pneumonia or other clinically documented infection, Hypotension and multi organ dysfunction (sepsis syndrome), Invasive fungal infection, Prior episode of febrile neutropenia, Hospitalized at the time of the development of fever. Primary prophylaxis of FN in patients who have a risk of FN of 20% or greater based on chemotherapy regimens. OR when the risk of developing FN is greater than or equal to 10% and less than 20% based on chemotherapy in patients who have other patient specific risk factors for FN including any of the following: Individual age greater than 65 years, Poor performance status (ECOG status 3-4), Bone marrow involvement by tumor producing cytopenias, Persistent neutropenia (ANC less than 1500mm3), poor renal function (GFR less than 60mL/min), liver dysfunction, Recent surgery and/or presence of open wounds.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Other Criteria	Adjunctive tx and individual as a high risk for infection-associated complications. For dose dense therapy (treatment given more frequently, such as every two weeks instead of every three weeks) for adjuvant treatment of breast cancer. For acute myeloid leukemia and using shortly after completion of induction or repeat induction chemo of AML. For myelodysplastic syndromes (MDS) with severe neutropenia (absolute neutrophil count (ANC) less than or equal to 500mm3 or experiencing recurrent/resistant infection. For mobilization of hematopoietic progenitor cells into peripheral blood for collection by leukapheresis and autologous transplantation. For acceleration of myeloid reconstitution after autologous or allogenic bone marrow transplantation or peripheral blood progenitor cell transplantation. For delayed neutrophil recovery/graft failure after autologous or allogenic bone marrow transplantation. Used to increase survival in individual exposed to myelosuppressive doses of radiation such as Hematopoietic Syndrome of Acute Radiation Syndrome. For malignant melanoma.
Indications	All Medically-accepted Indications.
Off Label Uses	

Levoleucovorin

Products Affected

- FUSILEV
- KHAPZORY
- levoleucovorin calcium
- levoleucovorin calcium pf

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Libtayo

Products Affected

• LIBTAYO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Lidocaine Topical

Products Affected

- lidocaine external ointment
- lidocaine hcl external cream
- lidocaine hcl external solution
- *lidocaine hcl mouth/throat*

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual is using for local analgesia OR Individual is using for relief of pain and itching due to minor cuts, minor scrapes, minor skin irritations, minor burns, and insect bites.
Indications	All Medically-accepted Indications.
Off Label Uses	

• lidocaine pak

Lidoderm Patch

Products Affected

- lidocaine external patch
- LIDODERM
- ZTLIDO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Lonsurf

Products Affected

• LONSURF

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Lorbrena

Products Affected

 LORBRENA ORAL TABLET 100 MG, 25 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Lotronex

Products Affected

- alosetron hcl
- LOTRONEX

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual is a Female with Severe diarrhea-predominant Irritable Bowel Syndrome (IBS) defined as including diarrhea and 1or more of the following: Frequent and severe abdominal pain/discomfort, Frequent bowel urgency or fecal incontinence, Disability or restriction of daily activities due to IBS. Member is female AND Member has chronic symptoms of IBS that have persisted for 6 months or longer AND does not have an anatomic or biochemical abnormality of the gastrointestinal tract.
Age Restrictions	18 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a documented trial of, an inadequate response or intolerance TWO (2) of the following medications: (a) Loperimide (b) antispasmodics (for example, dicyclomine), or (c) tricyclic antidepressants (AGA 2014).
Indications	All Medically-accepted Indications.
Off Label Uses	

Lucentis

Products Affected

• LUCENTIS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Lumizyme

Products Affected

• LUMIZYME

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For infantile-onset Pompe disease, dx is confirmed with acid alpha- glucosidase deficiency (GAA) activity in skin fibroblasts of less than 1% of the normal mean or by GAA gene sequencing AND presence of symptoms (for example respiratory and/or skeletal muscle weakness) of infantile-onset Pompe disease AND evidence of hypertrophic cardiomyopathy. For non-infantile onset (late-onset) Pompe disease, dx is confirmed by GAA enzyme assay which shows reduced enzyme activity less than 40% of the lab specific normal mean value AND confirmed by a second GAA enzyme activity assay in a separate sample (from purified lymphocytes, fibroblasts or muscle) or by GAA gene sequencing AND forced vital capacity (FVC) 30 -79% of predicted value while in the sitting position AND ability to walk 40 meters on a 6-minute walk test (assistive devices permitted) AND muscle weakness in the lower extremities.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Lumoxiti

Products Affected

• LUMOXITI

PA Criteria	Criteria Details
Exclusion Criteria	Individuals with severe renal impairment (CrCl less than 29 mL/min).
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Lupaneta

Products Affected

 LUPANETA PACK COMBINATION KIT 11.25 & 5 MG, 3.75 & 5 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	6 months
Other Criteria	For initial or retreatment of endometriosis. Individual has utilized Lupaneta for a combined total duration of less than 12 months (initial and retreatment).
Indications	All Medically-accepted Indications.
Off Label Uses	

Lupron Depot

Products Affected

- FENSOLVI (6 MONTH)
- LUPRON DEPOT (1-MONTH)
- LUPRON DEPOT (3-MONTH)
- LUPRON DEPOT (4-MONTH)
- LUPRON DEPOT (6-MONTH)
- LUPRON DEPOT-PED (1-MONTH)
- LUPRON DEPOT-PED (3-MONTH)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For Prostate cancer: Clinically localized dz with intermediate (T2b to T2c cancer, Gleason score of 7/Gleason grade group 2-3, or prostate specific antigen (PSA) value of 10-20 ng/mL) or higher risk of recurrence as neoadjuvant therapy with radiation therapy or cryosurgery OR Other advanced, recurrent, or metastatic disease. For Gynecology Uses: Initial treatment/retreatment of endometriosis (not to continue beyond 6 months) OR Dysfunctional uterine bleeding OR Preoperative tx as adjunct to surgical tx of uterine fibroids (leiomyoma uteri). May be used to reduce size of fibroids to allow for a vaginal procedure, prior to surgical tx (myomectomy or hysterectomy) in patients with documented anemia (Letheby et al. 2001). To induce amenorrhea in women (such as but not limited to menstruating women diagnosed with severe thrombocytopenia or aplastic anemia). For Endocrine Uses: Precocious puberty, defined as beginning of secondary sexual characteristics before age 8 in girls and before age 9 in boys. Ovarian Cancer (including fallopian tube cancer and primary peritoneal cancer): Hormonal therapy for clinical relapse in individuals with stage II-IV granulosa cell tumors or Hormonal therapy for treatment of epithelial ovarian cancer, fallopian tube cancer, primary peritoneal cancer as a single agent for persistent dz or recurrence.
Age Restrictions	
Prescriber Restrictions	

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Coverage Duration	1 year, except for Endometriosis:6months, Uterine Fibroids:3months
Other Criteria	For Gender Dysphoria in Adolescents (Hembree 2009)m 2017): Fulfills the DSM V criteria for gender dysphoria (American Psychiatric Assoc 2013) AND has experienced puberty to at least Tanner stage 2 (Hembree 2009, 2017) AND has (early) pubertal changes that have resulted in an increase of their gender dysphoria (Hembree 2009, 2017) AND does not suffer from a psychiatric comorbidity that interferes with the diagnostic work-up or treatment (Hembree 2009, 2017) AND has psychological and social support during treatment (Hembree 2009, 2017) AND demonstrates knowledge and understanding of the expected outcomes of GnRH analog treatment(Hembree 2009, 2017).
Indications	All Medically-accepted Indications.
Off Label Uses	

Lupron Kit IR

Products Affected

• *leuprolide acetate*

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For Prostate cancer: Clinically localized disease with intermediate (T2b to T2c cancer, Gleason score of 7/Gleason grade group 2-3, or prostate specific antigen (PSA) value of 10-20 ng/mL) or higher risk of recurrence as neoadjuvant therapy with radiation therapy or cryosurgery OR Other advanced, recurrent, or metastatic disease.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Lynparza

Products Affected

• LYNPARZA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Test results from an FDA-approved test confirms (written or verbal) the BRCA mutation for individuals with ovarian cancer or individuals with metastatic human epidermal growth factor receptor 2 (HER2) negative breast cancer.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Lyrica CR

Products Affected

• LYRICA CR

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has had a prior trial of immediate-release form of Lyrica (pregabalin) AND Documentation (verbal or written) has been provided which defines the following: (a.) The inadequate response to Lyrica (pregablin) AND (b.) The medical reason extended release Lyrica CR is clinically necessary, and the same medical reason and clinical reason benefits are not expected with Lyrica.
Indications	All Medically-accepted Indications.
Off Label Uses	

Makena

Products Affected

- hydroxyprogesterone caproate
- MAKENA INTRAMUSCULAR
- MAKENA SUBCUTANEOUS

PA Criteria	Criteria Details
Exclusion Criteria	Women with multiple gestations.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	6 months
Other Criteria	Therapy initiated between 16 weeks, 0 days and 20 weeks, 6 days of gestation and Singleton pregnancy and absence of preterm labor within the current pregnancy and individual is between 16 and 36 weeks of gestation with a singleton pregnancy. Prior history of a preterm singleton delivery before 37 weeks gestation due to either spontaneous preterm labor or premature rupture of membranes.
Indications	All Medically-accepted Indications.
Off Label Uses	

Mavenclad

Products Affected

- MAVENCLAD (10 TABS)
- MAVENCLAD (4 TABS)
- MAVENCLAD (5 TABS)
- MAVENCLAD (6 TABS)

- MAVENCLAD (7 TABS)
- MAVENCLAD (8 TABS)
- MAVENCLAD (9 TABS)

PA Criteria	Criteria Details
Exclusion Criteria	Concurrent use with other MS disease modifying agents (such as Aubagio, Gilenya, Mayzent, Tecfidera, Tysabri, Lemtrada, Ocrevus, Copaxone/Glatopa, Extavia, Rebif, Avonex, Plegridy, or Betaseron). Individual with clinically isolated syndrome (CIS) OR human immunodeficiency virus (HIV) infection. Individual with an active acute or chronic infection at the initiation of therapy OR moderate to severe renal impairment (creatinine clearance less than 60 mL/min) OR moderate to severe hepatic impairment (Child-Pugh class B or C) OR has completed two treatment courses (two years) of Mavenclad therapy.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual has a diagnosis of relapsing multiple sclerosis (RMS), including relapsing-remitting disease and active secondary progressive disease AND has had a trial and inadequate response or intolerance to at least one alternative drug indicated for the treatment of multiple sclerosis.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Mavyret

Products Affected

• MAVYRET

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Documentation is provided for a diagnosis of chronic hepatitis C (CHC) infection, which includes a genotype and a positive HCV RNA result (AASLD/IDSA 2017, CDC 2013) AND Individual has received baseline evaluation for liver fibrosis to guide appropriate therapy AND Individual does not have a short life expectancy (less than 12 months owing to non-liver related comorbid conditions) that cannot be remediated by treating HCV, by transplantation or other directed therapy (AASLD/IDSA 2017).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Criteria will be applied consistent with current AASLD/IDSA guidance.
Other Criteria	Criteria will be applied consistent with current AASLD/IDSA guidance.
Indications	All Medically-accepted Indications.
Off Label Uses	

Mayzent

Products Affected

• MAYZENT ORAL TABLET 0.25 MG, 2 MG

PA Criteria	Criteria Details
Exclusion Criteria	Concurrent use with other MS disease modifying agents (such as Aubagio, Gilenya, Mavenclad, Tecfidera, Tysabri, Lemtrada, Ocrevus, Copaxone/Glatopa, Extavia, Rebif, Avonex, Plegridy, or Betaseron). Individual who has been tested for CYP2C9 genotype and is homozygous for CYP2C9*3 (ie, CYP2C9*3/*3 genotype). Individual has had a recent (within the past 6 months) occurrence of one of the following: Myocardial infarction or Unstable angina or Stroke or Transient ischemic attack (TIA) or Decompensated heart failure requiring hospitalization or Class III/IV heart failure.Individual has history or presence of Mobitz Type II second- or third-degree atrioventricular (AV) block or sick sinus syndrome, unless individual has a functioning pacemaker. Individual has an active acute or chronic infection at the initiation of therapy.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Megace Suspension HRM

Products Affected

megestrol acetate oral suspension 40
 mg/ml, 400 mg/10ml

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual is using for the treatment of anorexia, cachexia, or unexplained weight loss in individuals with HIV/AIDS. Prescriber must acknowledge this is a high risk medication (HRM) [as identified by American Geriatric Society] for individuals greater than 65 and medication benefits outweigh potential risk for this individual
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Megace Tabs HRM

Products Affected

• megestrol acetate oral tablet

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Prescriber must acknowledge this is a high risk medication (HRM) [as identified by American Geriatric Society] for individuals greater than 65 and medication benefits outweigh potential risk for this individual.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has advanced, inoperable, recurrent breast cancer and using for palliative management. Individual has endometrial/uterine cancer and is suing for palliative management.
Indications	All Medically-accepted Indications.
Off Label Uses	

Mekinist

Products Affected

• MEKINIST ORAL TABLET 0.5 MG, 2 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	BRAF V600E or V600K mutation results must be confirmed (written or verbal attestation).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 Year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Mektovi

Products Affected

• MEKTOVI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	BRAF V600E or V600K mutation results must be confirmed (written or verbal attestation is acceptable).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Mepron

Products Affected

- atovaquone
- MEPRON

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Intolerance to trimethoprim-sulfamethoxazole (TMP-SMX)
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Mepsevii

Products Affected

• MEPSEVII

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Initial 6 months. Continuation 6 months.
Other Criteria	For initial use in individuals with a diagnosis of Mucopolysaccharidosis type VII (Sly syndrome) and confirmed (written or verbal) diagnosis of leukocyte or fibroblast glucuronidase enzyme assay or genetic testing and individual has elevated urine glycosaminoglycans excretion at a minimum of 3- fold over the mean normal for age at screening. For continued use, there is confirmation (written or verbal attestation) of clinically significant improvement or stabilization in clinical signs and symptoms of disease compared to the predicted natural history trajectory of disease.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Methylphenidate

Products Affected

- ADHANSIA XR
- APTENSIO XR
- CONCERTA ORAL TABLET EXTENDED RELEASE 18 MG, 27 MG, 36 MG, 54 MG
- COTEMPLA XR-ODT
- JORNAY PM ٠
- metadate er
- METHYLIN ORAL SOLUTION 10 MG/5ML, 5 MG/5ML
- methylphenidate hcl er (cd)
- methylphenidate hcl er (la) oral capsule extended release 24 hour 10 mg, 20 mg, • QUILLIVANT XR *30 mg, 40 mg, 60 mg*
- *methylphenidate hcl er (xr)*
- methylphenidate hcl er oral tablet extended release 10 mg, 18 mg, 20 mg, 27 mg, 36 mg, 54 mg, 72 mg

- methylphenidate hcl er oral tablet ٠ extended release 24 hour 18 mg, 27 mg, 36 mg, 54 mg
- methylphenidate hcl oral solution 10 *mq/5ml, 5 mq/5ml*
- methylphenidate hcl oral tablet
- methylphenidate hcl oral tablet chewable ٠ 10 mg, 2.5 mg, 5 mg
- QUILLICHEW ER ORAL TABLET CHEWABLE EXTENDED RELEASE 20 MG, 30 MG, 40 MG
- RELEXXII
- RITALIN
- RITALIN LA ORAL CAPSULE EXTENDED RELEASE 24 HOUR 10 MG, 20 MG, 30 MG, 40 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual is using for Attention Deficit Hyperactivity Disorder/Attention Deficit Disorder (ADHD/ADD) or Narcolepsy.
Age Restrictions	6 years of age and older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Off Label Uses	

Mircera

Products Affected

• MIRCERA

PA Criteria	Criteria Details
Exclusion Criteria	Continued use when the hemoglobin level exceeds 11.0 g/dL (except when the dose of methoxy polyethylene glycol-epoetin beta is adjusted to achieve and maintain target hemoglobin not to exceed 11.0 g/dL). Uncontrolled hypertension. Use beyond 12 weeks in the absence of response in individuals with chronic kidney disease. As treatment in the presence of a sudden loss of response with severe anemia and a low reticulocyte count.
Required Medical Information	Hemoglobin (Hgb) levels are less than 10 g/dL, prior to initiation of therapy AND (prior to initiation) the individuals iron status, including transferrin saturation or serum ferritin or bone marrow, (baseline) evaluation reveals: transferrin saturation at least 20% or ferritin at least 80 ng/mL or evidence of bone marrow demonstrates adequate iron stores And individual is using for one of the following: anemia associated with chronic kidney disease (CKD) for Individual on dialysis to achieve and maintain hemoglobin levels within the range of 10.0 to 11.0 g/dL OR anemia associated with CKD for Individual not on dialysis to achieve and maintain hemoglobin levels of 10.0 g/dL.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	12 weeks.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Modafinil

Products Affected

- modafinil oral tablet 100 mg, 200 mg
- PROVIGIL ORAL TABLET 100 MG, 200 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For Narcolepsy type 1: confirmed by the presence of daily periods of irrepressible need to sleep or daytime lapses into sleep occurring for at least 3 months and at least one of the following: 1. Clear cataplexy (defined as more than one episode of generally brief) usually bilaterally symmetrical, sudden loss of muscle tone with retained consciousness) AND 2.Multiple Sleep Latency Test (MSLT) showing one of the following: a. Mean sleep latency of less than 8 minutes with evidence of two sleep-onset rapid eye movement periods (SOREMPs) (ICSD-3, 2014)OR b. At least one SOREMP on MSLT and a SOREMP (less than 15 minutes) on the preceding overnight polysomnography (PSG) OR 3.Cerebrospinal fluid hypocretin-1 deficiency (less than 110 pg/mL or less than one-third of the normative values with the same standardized assay). For Narcolepsy type 2: confirmed by 1.Multiple sleep latency of less than 8 minutes with and evidence of two sleep-onset rapid eye movement periods (SOREMPs) ICSD-3, 2014) OR b. At least one SOREMP on MSLT and a SOREMP (less than 15 minutes) on the preceding overnight polysomnography (PSG) AND 2.The absence of cataplexy AND 3.Exclusion of alternative causes of excessive daytime sleepiness by history, physical exam and Polysomnography.
Age Restrictions	
Prescriber Restrictions	

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Coverage Duration	1 year
Other Criteria	For obstructive sleep apnea/hypopnea syndrome objectively confirmed by polysomnography (PSG) or home testing with portable monitor showing one of the following (AASM 2009): (1).Greater than 15 obstructive events (defined as apneas, hypopneas plus respiratory event related arousal) per hour of sleep OR (2)Greater than 5 obstructive events per hour of sleep and individual reports any of the following: a.Unintentional sleep episodes during wakefulness or b. Daytime sleepiness or c. Unrefreshing sleep or d. Fatigue or e. Insomnia or f.Waking up breath holding, gasping or choking or g.Bed partner describing loud snoring, breathing interruptions or both or h. Presence of comorbid conditions including hypertension, mood disorder, cognitive dysfunction, coronary artery disease, stroke, congestive heart failure, atrial fibrillation or type 2 diabetes mellitus. And has an Epworth Sleepiness Scale score greater than or equal to 10. For Shift-Work Sleep Disorder (SWSD) confirmed by all of the following: (1)No other medical disorder or mental disorder accounts for the symptoms AND (2)Symptoms do not meet criteria for any other sleep disorder (i.e. jet lag) AND (3)Symptoms have occurred for at least 3 months, AND (4)Individual has one of the following: a.Individual has excessive sleepiness or insomnia associated with a work period that occurs during the usual sleep phase OR b.Polysomnography demonstrates loss of a normal sleep- wake pattern (such as, disturbed chronobiological rhythmicity).
Indications	All Medically-accepted Indications.
Off Label Uses	

Mozobil

Products Affected

• MOZOBIL

PA Criteria	Criteria Details
Exclusion Criteria	Using as a mobilizing agent for an allogeneic stem cell donor (NCCN, ASBMT 2014), mobilizer of leukemic cells or as a component of a conditioning regimen prior to an allogeneic hematopoietic stem cell transplant.
Required Medical Information	Using in combination with granulocyte colony stimulating factor (G-CSF) (such as Neupogen, Nivestym, Zarxio or Granix) and after stem cell mobilization and collection, a subsequent autologous hematopoietic stem cell transplant is anticipated.
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	3 months
Other Criteria	Individual may use a maximum of up to four consecutive doses of plerixafor (Mozobil) injections per cycle for up to 2 cycles.
Indications	All Medically-accepted Indications.
Off Label Uses	

Mulpleta

Products Affected

• MULPLETA

PA Criteria	Criteria Details
Exclusion Criteria	Used to normalize platelet counts in those with chronic liver disease
Required Medical Information	Individual has a platelet count of less than 50 X 109/L
Age Restrictions	18 years of age
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Myalept

Products Affected

• MYALEPT

PA Criteria	Criteria Details
Exclusion Criteria	Individual is using for the treatment of complications of partial lipodystrophy. Individual is using for the treatment of liver disease, including nonalcoholic steatohepatitis (NASH). Individual is using for the treatment of HIV-related lipodystrophy. Individual is using for treatment in patients with general obesity or metabolic disease, including diabetes mellitus and hypertriglyceridemia, without concurrent evidence of congenital or acquired generalized lipodystrophy
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Mycapssa

Products Affected

• MYCAPSSA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For continuation therapy, individual meets initial criteria AND IGF-1 levels remain less than 1.3 X the upper limit of normal (ULN) and a serum growth hormone level less than 2.5ng/mL (Melmed 2015).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a diagnosis of acromegaly AND has responded to and tolerated treatment with octreotide or lanreotide (defined as currently receiving a stable dose of either for at least the previous 3 months (Label, Melmed 2015).
Indications	All Medically-accepted Indications.
Off Label Uses	

Mylotarg

Products Affected

• MYLOTARG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	Relapsed or refractory CD33-positive AML: 2 years and older. For newly diagnosed CD33-positive AML: 1 month and older.
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Naglazyme

Products Affected

• NAGLAZYME

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Mucopolysaccharidosis VI is confirmed: (a) with an increase in dermatan sulfate in the urine and (b) Decrease in the activity of N- acetylgalactosamine-4-sulfatase (arylsulfatase B) enzyme as measured in fibroblasts or leukocytes combined with normal enzyme activity level of another sulfatase OR (c) N- acetylgalactosamine-4-sulfatase (arylsulfatase B) gene mutation confirmed (written or verbal attestation).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Namenda Line

Products Affected

- memantine hcl er
- memantine hcl oral solution
- memantine hcl oral tablet 10 mg, 28 x 5 NAMENDA XR mg & 21 x 10 mg, 5 mg
- NAMENDA ORAL TABLET 10 MG, 5 MG
- NAMENDA TITRATION PAK

 - NAMENDA XR TITRATION PACK

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	Individuals that are 50 years of age or older are NOT subject to the prior authorization requirements. Prior Authorization applies to individuals that are 49 years of age or younger.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a diagnosis of moderate to severe dementia of the Alzheimer's type.
Indications	All Medically-accepted Indications.
Off Label Uses	

Natpara

Products Affected

• NATPARA

PA Criteria	Criteria Details
Exclusion Criteria	Serum corrected total calcium levels maintained within therapeutic range on calcium supplements and active vitamin D forms alone OR serum corrected total calcium level of less than or equal to 7.5 mg/dL at initiation of therapy. Individual is using to treat hypoparathyroidism caused by a gene mutation in the calcium- sensing receptor OR using to treat acute (duration of less than 6 months, Bilezikian et al. 2011) postoperative hypoparathyroidism OR Individual is at increased risk for osteosarcoma (such as but not limited to, concomitant Paget's disease of bone, open epiphyses, prior history of skeletal external beam or implant radiation therapy).
Required Medical Information	
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Diagnosis of chronic (duration of greater than or equal to 18 months, mannstadt et, al 2013) hypoparathyroidism.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Nerlynx

Products Affected

• NERLYNX

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Nexavar

Products Affected

• NEXAVAR

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Confirmed (verbal or written attestation) results of FLT3-ITD mutation with acute myeloid leukemia, relapsed/refractory disease.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Ninlaro

Products Affected

• NINLARO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	For multiple myeloma, individual received at least two prior therapies, including immunomodulatory agent and a proteasome inhibitor AND demonstrated disease progression on or within 60 days of completion therapy AND Ninlaro is given as part of a treatment regimen containing dexamethasone and pomalidomide.
Indications	All Medically-accepted Indications.
Off Label Uses	

Nityr

Products Affected

• NITYR

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual's plasma tyrosine level is maintained below 500 micromol/L to reduce risk of ocular symptoms, developmental delay, or hyperkeratotic plaques.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Northera

Products Affected

 NORTHERA ORAL CAPSULE 100 MG, 200 MG, 300 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	Individual is 18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has had a trial (resulting in inadequate response, therapeutic failure or intolerance) of at least one prior pharmacologic therapy (which may include midodrine or fludrocortisone) for treatment of symptoms of NOH.
Indications	All Medically-accepted Indications.
Off Label Uses	

Nourianz

Products Affected

• NOURIANZ ORAL TABLET 20 MG, 40 MG

PA Criteria	Criteria Details
Exclusion Criteria	Individuals with severe hepatic impairment (Child Pugh Class C)
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Noxafil

Products Affected

- NOXAFIL ORAL
- posaconazole

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

NP CSF SA Agents

Products Affected

- NEUPOGEN
- NIVESTYM

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Febrile neutropenic individuals who are at risk for infection- associated complications or have any of the following: Expected prolonged (greater than 10 day) and profound (less than 0.1 x 10to the power of 9/L) neutropenia, Age greater than 65 years, Pneumonia or other clinically documented infection, Hypotension and multi organ dysfunction (sepsis syndrome), Invasive fungal infection, prior episode of FN, or Hospitalized at the time of the development of fever. Primary prophylaxis of FN in patients who have a risk of FN of 20% or greater based on chemotherapy regimens. OR when the risk of developing FN is greater than or equal to 10% and less than 20% based on chemotherapy in patients who have other patient specific risk factors for FN including any of the following: Individual age greater than 65 years, Poor performance status (ECOG status 3-4), prior chemotherapy or radiation therapy, Bone marrow involvement by tumor producing cytopenias, persistent neutropenia (ANC less than 1500mm3), poor renal function (GFR less than 60mL/min), liver dysfunction (liver function tests at least 2x upper limit of normal) or recent surgery and/or presence of open wounds.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Other Criteria	Individual has had a trial and inadequate response to intolerance to Zarxio (Filgrastim-sndz). Secondary Prophylaxis for patients who experienced a neutropenic complication from a prior cycle of chemotherapy (for which primary prophylaxis was not received), in which a reduced dose may compromise disease-free or overall survival or treatment outcome. Using for adjunctive tx for FN and has been prophylactic therapy with GCSF agent or has not received prophylactic therapy with a GCSF and who are at high risk for infection-associated complications. Use in acute lymphocytic leukemia (ALL) after completion of the first few days of initial induction chemotherapy or first post-remission course of chemotherapy. Use in individuals with acute myeloid leukemia (AML) shortly after the completion of induction or repeat induction chemotherapy or after the completion of consolidation chemotherapy for AML. For tx of moderate to severe aplastic anemia. Tx of severe neutropenia in individuals with hairy cell leukemia. For myelodysplastic syndromes (MDS) with severe neutropenia (absolute neutrophil count (ANC) less than or equal to 500 mm3 or experiencing recurrent infection. For dose dense therapy (treatment given more frequently, such as every two weeks instead of every three weeks) for adjuvant treatment of breast cancer. For chronic administration to reduce the incidence and duration of sequelae of neutropenia (e.g., fever, infections, oropharyngeal ulcers) in symptomatic individuals with congenital neutropenia, cyclic neutropenia, or idiopathic neutropenia. For tx of (non-chemotherapy) drug-induced neutropenia. For tx of low neutrophil counts in individuals with glycogen storage disease type 1b. For tx of neutropenia associated with human immunodeficiency virus (HIV) infection and antiretroviral therapy. In individuals receiving radiation therapy in the absence of chemotherapy if prolonged delays secondary to neutropenia are expected. After accidental or intentional total body radiation of myleosuppressive doses (greater than

PA Criteria	Criteria Details
	Syndrome of Acute Radiation Syndrome). After hematopoietic progenitor stem cell transplant (HPCT/HSCT) to promote myeloid reconstitution or when engraftment is delayed or has failed. To mobilize progenitor cells into peripheral blood for collection by leukapheresis, as an adjunct to peripheral blood/hematopoietic stem cell transplantation (PBSCT/PHSCT). Use as an alternate or adjunct to donor leukocyte infusions (DLI) in individuals with leukemic relapse after an allogeneic hematopoietic stem cell transplant.
Indications	All Medically-accepted Indications.
Off Label Uses	

NP Human Growth Hormone

Products Affected

- GENOTROPIN
- GENOTROPIN MINIQUICK
- HUMATROPE
- NUTROPIN AQ NUSPIN 10
- NUTROPIN AQ NUSPIN 20
- NUTROPIN AQ NUSPIN 5

- SAIZEN
- SAIZENPREP
- SEROSTIM
- ZOMACTON
- ZORBTIVE

PA Criteria	Criteria Details
Exclusion Criteria	Use of GH tx when applicable criteria have not been met may not be approved for, but not limited to, the following: Renal transplant, Anabolic tx, except for AIDS, provided to counteract acute or chronic catabolic illness (e.g. surgery, trauma, cancer,chronic hemodialysis) producing catabolic (protein wasting) changes in both adult and ped. Anabolic tx to enhance body mass or strength for professional, recreational or social reasons, Constitutional delay of growth and development. Cystic fibrosis, GH Tx in combo with GnRH agonist (Lupron-leuprolide) as a Tx of precocious puberty. Hypophosphatemic rickets, osteogenesis imperfecta, osteoporosis, Short Stature associated with GH insensitivity (Laron Syndrome). Therapy in older adults with normally occurring decrease in GH, who are not congenitally GH deficient and who have no evidence of organic pituitary dz (this is referred to as age-related GH deficiency). Tx of CHF, burn patients, fibromyalgia, glucocorticoid- induced growth failure. Tx of HIV lipodystrophy (fat redistribution syndrome), also referred to as altered body habitus (e.g. buffalo hump), associated with antiviral therapy in HIV-infected patients. Tx of intrauterine growth restriction (IUGR)or Russell-Silver Syndrome that does not result in SGA. Tx of obesity. Other etiologies of SS where GH has not been shown to be associated with an increase in final height, including but not limited to achondrodoplasia and other skeletal dysplasias. CONT therapy: evaluated annually AND growth rate above 2.5cm/yr (not for child w/prior documented hypopituitarism)(Grimberg 2016) AND Child over 12: an X-ray report with evidence that epiphyses have closed or SMR of 3 or more. GH tx for reconstruction is terminated when BA =16 yr (M), or = 14 yr (F) is reached OR Epiphyseal fusion has occurred OR Mid-parenteral height is achieved.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Required Medical Information	For initial idiopathic GHD requests, has signs/sym sx of GHD, GV 2 SD below age-appropriate mean or ht 2.25 SD below age- appropriate mean and A subnormal (SubNL) response (less than10ng/ml)to 2GH stim tests OR Neonates with hypoglycemia and clinical/hormone evidence of hypopit and low GH (less than 10ng/ml)OR 20ther pit hormone deficiencies and low IGF-1 OR mbr has cranial irradiation with low IGF1 and normal thyroid tests. Child born SGA(birth wt or length 2 or more SD below the mean for gest age),Child fails to manifest catch up growth by age 2yr(ht 2 or more SD below the mean for age,gender)AND Other causes for SS have been ruled out.Transitioning adolescent: individual completed linear growth (less than 2cm/yr) AND either GH tx has been stopped for at least a month, and GHD has been reconfirmed:idiopathic isolated GHD(SubNL response to 2 GH stim tests, OR SubNL response (GH conc of less than10 ng/mL)to 1 provocative test and low IGF-I/IGFBP-3)OR multiple pit hormone deficiency,(SubNL response to 1 provocative GH test and/or low IGF-I/IGFBP-3)or for mbr with cranial irradiation, low IGF with normal thyroid or any of the following, known genetic mutation associated with def GH production or secretion or Hypothalamic-pit tumor or structural defect or 3 other pit hormone deficiencies.Adult GHD must be confirmed/reconfirmed:SubNL response in adults to 2 GH stim tests (GH conc of less than or equal to 5ng/ml when using insulin induced hypoglycemia OR GH conc of less than or equal to 4.1ng/ml when using arginine)OR SubNL response to 1 stim test for adults with hypothalamic or pit dz and 1 pit hormone deficits OR 3 other pit hormone deficiencies.Initial request for Reconstructive GH tx in child w/ mean ht is at least 2.25 but less than 2.55D below the mean for age, gender and GV is less than 10th percentile over 1yr or mean ht at least 2.55D below the mean for age, gender for conditions known responsive to GH.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For Non Preferred Growth hormone agents, individual has had trial of TWO preferred GH agents (Norditropin AND Omnitrope) or preferred GH agent is not FDA-approved and does not have an accepted off-label use per CMS recognized compendia for the prescribed indication and the requested non-preferred agent is. GH may be approved for Adolescents with childhood onset GHD who have completed linear growth. GH tx in other populations approved when: individual has AIDS wasting (defined as greater than 10% of baseline wt loss that is not explained by concurrent illness other than HIV) AND is being treated with antiviral therapy AND will continue tx until definition not met OR individual dx with short bowel syndrome AND is receiving specialized nutritional support.
Indications	All Medically-accepted Indications.
Off Label Uses	

NP Interferon for MS

Products Affected

EXTAVIAREBIF

• REBIF TITRATION PACK

- REBIF REBIDOSE
- REBIF REBIDOSE TITRATION PACK

PA Criteria	Criteria Details
Exclusion Criteria	Individuals with primary progressive MS. Individuals with secondary progressive MS without relapsing disease. Concurrent use with other MS disease modifying agents (such as Aubagio, Gilenya, Tecfideria, Tysabri, Lemtrada, Ocrevus, Copaxone/Glatopa, Extavia, Rebif, Avonex, Plegridy or Betaseron).
Required Medical Information	Individual has experienced first clinical episode and has MRI features consistent with multiple sclerosis OR individual has MS with relapsing disease (RMS) OR individual with secondary progressive MS (SPMS) with a history of superimposed relapses.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has been on Extavia or Rebif in the past 180 days OR member has tried therapy with ONE of the following agents: Avonex (interferon beta-1a) OR Plegridy (interferon beta-1a) OR Betaseron (interferon beta-1b) OR Tecfidera (dimethyl fumarate) OR Copaxone/Glatopa (glatiramer).
Indications	All Medically-accepted Indications.
Off Label Uses	

NP IVIG

Products Affected

- ASCENIV
- BIVIGAM
- CARIMUNE NF
- CUTAQUIG
- CUVITRU
- FLEBOGAMMA DIF
- GAMMAGARD
- GAMMAGARD S/D LESS IGA

- GAMMAKED
- GAMMAPLEX
- HIZENTRA
- HYQVIA
- PANZYGA
- PRIVIGEN
- XEMBIFY

PA Criteria	Criteria Details
Exclusion Criteria	

Y0114_20_114077_I_012

Criteria Details
For initial CIDP trial (up to 12wks): has muscle weakness/sensory dysfx caused by neuropathy in more than 1 limb, nerve conduction studies or diagnostic criteria confirm demyelinating neuropathy and other polyneuropathies was ruled out. For CIDP continued use, there is improvement in neurological sx on exam AND continued need shown by attempts annually to titrate dose or interval of therapy result is worsening of sx. For initial trial (up to 12wks) to tx Multifocal Motor Neuropathy (MMN) when clinical presentation combined w/ electrodiagnositic test, labs or diagnostics confirm/suggest MMN. CONT MMN use: mbr had strength improvement and fx after initial tx and CONT need is shown by attempts on yearly basis to titrate dose or interval of therapy result in worsening of sx. Desensitization prior to Solid Organ transplantation for suppression on panel reactive anti-HLA AB in mbr w/PRA level to HLA (AAAAI 2016). Transplant recipient having AB-mediated rejection w/donor specific AB (KDIGO 2009). Tx of ITP with either active bleeding or platelet count less than 30,000mcL (ASH 2011). Tx of fetal alloimmune TCP with AB to paternal platelet antigen found in maternal serum AND one of the following: a previously affected pregnancy or family hx of maternofetal alloimmune TCP or Fetal blood samples shows TCP. For isoimmune hemolytic dz of newborn, tx of severe hyperbilirubinemia. Tx of Lambert-Eaton myasthenic (AAAAI 2016) syndrome: has muscle weakness and dx confirmed by either eletrodiagnostic findings using nerve conduction tests, RNS, exercise testing or SFEMG OR presence of AB directed against VGCC. Stiff person syndrome when mbr had inadeq response/intolerance/contraind to other tx such as benzo or baclofen (AAAAI 2016). Kawasaki Syndrome tx no more than 5dys (AHA 2004).
1 year

PA Criteria	Criteria Details
Other Criteria	NP IG may be approved if trial/inadeq response/intol to 1 PF IG (Gammunex, Gammunex-C, Octagam) OR PF Ig is not FDA/Off- label approved or due to concomitant clinical condition(s) such as but not limited to: Renal insuff/impairmt, Non-O blood type, Severe IgA def, DM/pre-DM, CVD, Hyper-prolinemia, Hypernatremia, hi- risk of thrombosis (such as but not limited to hyperviscosity syn OR hypercoagulable cond), documented hypersensitivity manifested by severe systemic/allergic or anaphylactic rxn to any ingred not also present in requested NP agent OR if SCIG-only dose forms are designated as NP, may be approved for difficult vein access that precludes use of any IVIG or hx of serious systemic rxn to IVIG expected to be avoided by using SCIG or hx of inconsistent serum levels of IgG with IVIG. Tx of PI when hx of recurrent sinopulmonary infection req abx tx AND lack of/inadeq response to immunization AND no evidence of renal and GI as causes of HGG AND initial pre-tx total serum IgG is BLL of age adj lab ref range or more than 2SD below adj mean. Tx of other PI when no evidence of renal/GI causes of HGG AND initial pre-tx total serum IgG is below lower limit of age adj lab ref range or more than 2 SD below age adj mean AND hx of recurrent sinopulmonary infect requiring abx therapy AND lack of/inadeq response to immunization. For autoimmune neutropenia, active infect is excluded as the cause. For ELS: has muscle weakness, characteristic electromyography and presence of AB directed against voltage-gated Ca-channels. Autoimmune muccoutaneous blistering dz or dermatomyosistiis(DM)/polymyositis when mbr had inadeq response/intolerance/contraind to other tx such as corticosteroids or immunosuppressive agents (AAAAI 2016). Myasthenia Gravis (AAAAI 2016): clinical presentation is characteristic AND dx confirmed by AB against the acetylcholine receptor or muscle specific tyrosine kinase OR electrodiagnostic
	finding RNS or SFEMG AND using for worsening sx or short term therapy as immunosuppressive tx or MAINT therapy when mbr had inadeq response/intolerance/contraind to other tx such as steroids or immunosuppressants.
Indications	All Medically-accepted Indications.

PA Criteria	Criteria Details
Off Label Uses	

NP LA Opioid

Products Affected

- CONZIP
- DOLOPHINE
- hydromorphone hcl er
- KADIAN
- methadone hcl oral tablet
- morphine sulfate er beads
- morphine sulfate er oral capsule extended release 24 hour
- morphine sulfate er oral tablet extended tramadol hcl er (biphasic)

release 100 mg, 15 mg, 200 mg, 30 mg, 60 mg

- MS CONTIN ORAL TABLET EXTENDED RELEASE 100 MG, 15 MG, 200 MG, 30 MG, 60 MG
- NUCYNTA ER
- oxymorphone hcl er
- tramadol hcl er

PA Criteria	Criteria Details
Exclusion Criteria	Individual is requesting or using as an as-needed analgesic OR Individual has one of the following conditions: (a) Significant respiratory depression OR (b) Acute or severe bronchial asthma or hypercarbia OR (c) Known or suspected paralytic ileus.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Initial 3 months. Maintenance 6 months. Cancer Pain/Terminal Dx or Palliative Care 1 Year.

PA Criteria	Criteria Details
Other Criteria	For initial use, Individual has one of the following: (a) Diagnosis of cancer related pain and/or is actively undergoing cancer therapy (provide cancer diagnosis) OR (b) Diagnosis of terminal illness and is receiving palliative/end-of-life care (provide terminal diagnosis) OR Individual has pain severe enough to require daily, around-the-clock, long term opioid treatment (provide diagnosis) AND Individual has one of the following: (a) An inadequate response to ONE alternative treatment options, such as but not limited to non-opioid analgesics and immediate-release opioids OR (b) Alternative treatment options would otherwise be inadequate to provide sufficient management of pain OR (c) Individual has contraindications to non-opioid analgesics (such as NSAID use in individuals with active ulcer condition/gastrointestinal bleeding, renal failure) AND for initial therapy, individual is not opioid analyse as noted by the following: (a) Individual is currently using a short-acting opioid analgesic, including use of opioid analgesia as an inpatient for post-surgical pain OR (b) Individual is transitioning from one long-acting opioid analgesic to another long-acting opioid analgesic to another long-acting opioid therapy has provided meaningful improvement in pain and/or function compared to baseline AND Prescriber has consulted individual regarding risks of opioid therapy AND clear treatment goals have been defined and outline as part of overall plan AND has one of the following: Diagnosis of cancer related pain and/or is actively undergoing cancer therapy (provide cancer diagnosis) OR diagnosis of terminal illness and is receiving palliative/end-of-life care (provide terminal diagnosis) OR has pain severe enough to require daily, around-the-clock, long term opioid treatment (provide diagnosis).
Indications	All Medically-accepted Indications.
Off Label Uses	

NP LA Opioid Abuse Deterrent

Products Affected

- ARYMO ER
- BELBUCA
- buprenorphine transdermal patch weekly hydrocodone bitartrate er 10 mcg/hr, 15 mcg/hr, 20 mcg/hr, 5 mcg/hr, 7.5 mcg/hr
- BUTRANS TRANSDERMAL PATCH WEEKLY 10 MCG/HR, 15 MCG/HR, 20
- MCG/HR, 5 MCG/HR, 7.5 MCG/HR
- EMBEDA
- HYSINGLA ER
- MORPHABOND ER
- XTAMPZA ER
- ZOHYDRO ER

PA Criteria	Criteria Details
Exclusion Criteria	Individual is requesting or using as an as-needed analgesic OR Individual has one of the following conditions: (a) Significant respiratory depression OR (b) Acute or severe bronchial asthma or hypercarbia OR (c) Known or suspected paralytic ileus.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Initial 3 months. Maintenance 6 months. Cancer Pain/Terminal Dx or Palliative Care 1 Year.

PA Criteria	Criteria Details
Other Criteria	For initial use, Individual has one of the following: (a) Diagnosis of cancer related pain and/or is actively undergoing cancer therapy (provide cancer diagnosis) OR (b) Diagnosis of terminal illness and is receiving palliative/end-of-life care (provide terminal diagnosis) OR Individual has pain severe enough to require daily, around-the- clock, long term opioid treatment (provide diagnosis) AND Individual has one of the following: (a) An inadequate response to ONE alternative treatment options, such as but not limited to non-opioid analgesics and immediate-release opioids OR (b) Alternative treatment options would otherwise be inadequate to provide sufficient management of pain OR (c) Individual has contraindications to non-opioid analgesics (such as NSAID use in individuals with active ulcer condition/gastrointestinal bleeding, renal failure) AND for initial therapy, individual is not opioid anaive as noted by the following: (a) Individual is currently using a short-acting opioid analgesic, including use of opioid analgesia as an inpatient for post-surgical pain OR (b) Individual is transitioning from one long-acting opioid analgesic to another long-acting opioid analgesic. AND If an abuse deterrent formulation is needed [such as but not limited to Embeda ER, Hysingla ER, Targiniq ER, Troxyca ER, Xtampza ER and Zohydro ER], and individual has a history of substance abuse disorder OR If is there is concern for abuse or dependence with pure opioid agents. For continued use, Attestation (verbal or written) that long-acting opioid therapy has provided meaningful improvement in pain and/or function compared to baseline AND Prescriber has consulted individual regarding risks of opioid therapy AND clear treatment goals have been defined and outline as part of overall plan AND has one of the following: Diagnosis of cancer related pain and/or is actively undergoing cancer therapy (provide cancer diagnosis) OR diagnosis of terminal illness and is receiving palliative/end-of-life care
	(provide terminal diagnosis) OR has pain severe enough to require daily, around-the-clock, long term opioid treatment (provide diagnosis).
Indications	All Medically-accepted Indications.
Off Label Uses	
Y0114 20 114077	7 I 012 December 2020

NP SGLT2

Products Affected

- INVOKAMET
- INVOKAMET XR
- INVOKANA ORAL TABLET 100 MG, 300 MG
- SEGLUROMET
- STEGLATRO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual has had a trial and inadequate response or intolerance to metformin OR has a contraindication to metformin therapy [such as but not limited to, renal insufficiency (eGFR is less than 45 mL/minute/1.73m2)]. AND has had a trial and inadequate response or intolerance to Jardiance (empagliflozin), Synjardy (empagliflozin/metformin), or Synjardy XR (empagliflozin/metformin extended-release).
Indications	All Medically-accepted Indications.
Off Label Uses	

NP SGLT2 DPP4 Combo

Products Affected

- GLYXAMBI
- QTERN
- STEGLUJAN
- TRIJARDY XR ORAL TABLET EXTENDED

RELEASE 24 HOUR 10-5-1000 MG, 12.5-2.5-1000 MG, 25-5-1000 MG, 5-2.5-1000 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual has had a trial and inadequate response or intolerance to ONE preferred DPP4 AND ONE preferred SGLT2 AND Individual has had an adequate response (achieved glucose control) with trial of the DPP-4 inhibitor and SGLT2 inhibitor at the same time AND Documentation (Verbal or Written) has been provided for why the combination agent is clinically necessary and not for convenience.
Indications	All Medically-accepted Indications.
Off Label Uses	

NP Statin

Products Affected

ALTOPREV

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has had a trial of generic statin at any dose and provider attests the member has experienced failure, contraindication, or intolerance to a generic statin. Or Individual is currently on a product that interacts with generic statin.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

NP Topical Androgens

Products Affected

- ANDRODERM
- ANDROGEL TRANSDERMAL GEL 25 MG/2.5GM (1%), 50 MG/5GM (1%)
- FORTESTA
- TESTIM
- testosterone transdermal gel 10 mg/act

PA Criteria **Criteria Details** Exclusion Criteria Required Medical Information 18 years of age or older. For transgender use, individual is 16 Age Restrictions vears of age or older. Prescriber Restrictions Coverage 1 year Duration **Other Criteria** Individual has had a trial of androgel 1.62% AND Individual has a dx of one: (1) primary hypogonadism (congenital or acquired) (for example, bilateral torsion, cryptorchidism, chemotherapy, Klinefleter Syndrome, orchitis, orchiectomy, toxic damage from alcohol or heavy metals, vanishing testis syndrome, idiopathic primary hypogonadism, age-related hypogonadism [also referred to as late-onset hypogonadism]) OR (2) Hypogonadotropic hypogonadism (also called secondary hypogonadism) (congenital or acquired) [for example, idiopathic gonadotropin or luteinizing hormone-relapsing hormone (LHRH) deficiency, pituitaryhypothalamic injury] OR (3) Individual is transgender AND is using for a diagnosis of gender Dysphoria or gender Identity Disorder AND Goal of treatment is female-to-male gender reassignment. Indications All Medically-accepted Indications.

Y0114_20_114077_I_012

December 2020

(2%), 12.5 mg/act (1%), 25 mg/2.5gm (1%), 50 mg/5gm (1%)

- testosterone transdermal solution
- VOGELXO
- VOGELXO PUMP

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

NP TZD

Products Affected

• AVANDIA ORAL TABLET 2 MG, 4 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has had a trial and inadequate response or intolerance to metformin OR Individual has a contraindication to metformin therapy [such as but not limited to, renal insufficiency (eGFR is less than 45mL/minute/1.73m2)] AND Individual has had a trial with ONE of the following: dipeptidyl peptidase-4 (DPP-4), glucagon-like peptide-1 (GLP-1), or a sodium-glucose co-transporter-2 (SGLT2) inhibitor.
Indications	All Medically-accepted Indications.
Off Label Uses	

Nplate

Products Affected

• NPLATE

PA Criteria	Criteria Details
Exclusion Criteria	Using Nplate to normalize platelet counts. Using for low platelet count caused by any condition other than chronic ITP.
Required Medical Information	For initial therapy, individual's degree of thrombocytopenia (platelet count less than 30,000/mm3) and clinical condition increase the risk for bleeding AND individual demonstrated an insufficient response to corticosteroids, immunoglobulins (for example, IVIg or anti-D), or splenectomy. For maintenance therapy, individual demonstrated response to therapy as evidenced by increased platelet counts, and the goal of ongoing treatment is to maintain an adequate platelet count (50,000-100,000/mm3) to decrease the risk of bleeding.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	For ITP: Initial 6 months, renewal 1 year. For all other diagnoses: 1 year
Other Criteria	For treatment of myelodysplastic syndrome (MDS) in individual with severe or refractory thrombocytopenia following disease progression or no response to hypomethylating agents or immunosuppressive therapy.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Nubeqa

Products Affected

• NUBEQA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Nucala

Products Affected

NUCALA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For severe eosinophilic asthma: In the absence of other potential causes of eosinophilia, including hypereosinophilic syndromes, neoplastic disease and known or suspected parasitic infections, the individual has a blood eosinophil count that is either greater than or equal to 150 cells/microliter at ignition of therapy OR greater than or equal 300 cells/microliter in the prior 12 months. Evidence of asthma is demonstrated by the following (NAEPP 2008): The individual has a pretreatment forced expiratory volume in one second (FEV1) less than 80% predicted AND FEV1 reversibility of at least 12% and 200mL after albuterol (salbutamol) administration.
Age Restrictions	For eosinophilic asthma: 6 years old or older. For eosinophilic granulomatosis with polyangitis (EGPA): 18 years old or older.
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For severe eosinophilic asthma, individual has had a 3 month trial/inadequate response to combination controller therapy (high dose inhaled corticosteroids plus LA beta2-agonist, leukotriene modifiers, theophylline or oral corticosteroids) (ERS/ATS 2013) AND has experienced 2 or more asthma exacerbations in the prior 12 months requiring use of a systemic corticosteroid or temporary increase in the individual's usual maintenance of oral corticosteroids (ERS/ATS 2013). For Continuation Therapy after 12 months in individuals with severe eosinophilic asthma: Treatment has resulted in clinical improvement as confirmed by either i) Decreased utilization of rescue medications OR ii) A decreased frequency of exacerbation (defined as worsening of asthma that requires increase in inhaled corticosteroid dose or treatment with systemic corticosteroid) OR iii) An increase in percent predicted FEV1 from pretreatment baseline OR iv) A reduction in reported asthma-related symptoms, such as, to wheezing, shortness of breath, coughing, fatigue, sleep disturbance or asthmatic symptoms upon awakening. For individuals with relapsing or refractory eosinophilic granulomatosis with polyangiitis for 6 months or greater and 1) a history or presence of asthma and 2) blood eosinophil level of greater than or equal to 10% of leucocytes or an absolute eosinophilic syndromes, neoplastic disease and known or suspected parasitic infection) and 2) the presence of 2 or more features of eosinophilic granulomatosis with polyangiitis (such as, biopsy showing histopathological evidence of eosinophili-rich granulomatosis inflammation, neuropathy, mono or poly(motor deficit or nerve conduction abnormality), pulmonary infiltrates, non-fixed sino-nasal abnormality, cardiomyopathy, glomerulonephritis, alveolar hemorrhage, palpable purpura or antineutrophil cytoplasmic antibody positive status. For

PA Criteria	Criteria Details
	Continuation Therapy after 12 months in individuals with eosinophilic granulomatosis with polyangiitis when treatment has resulted in clinical improvement as confirmed by the achievement of remission at some point during tx, defines as the following: Birmingham Vasculitis Activity Score, version 3, of zero on scale from 0 to 63 and receipt of prednisolone or prednisone at dose of 4mg or less per day.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Nuedexta

Products Affected

• NUEDEXTA

PA Criteria	Criteria Details
Exclusion Criteria	Concomitant use with any of the following: (i.) Agents containing quinidine, quinine, or mefloquine OR (ii.) Agents that both prolong the QT interval and are metabolized by CYP2D6 (for example, thioridazine, pimozide) OR Concomitant monoamine oxidase inhibitor (MAOI) use or use in the preceding 14 days OR Individual has any of the following cardiovascular conditions: (i.)Prolonged QT interval, congenital long QT syndrome, or history suggestive of torsades de pointes OR (ii.) Heart failure OR (iii.) Complete atrioventricular (AV) block without an implanted pacemaker or at high-risk of a complete AV block.
Required Medical Information	
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual has a diagnosis of pseudobulbar affect (PBA) AND has a concomitant diagnosis with an unrelated neurologic disease or injury [amyotrophic lateral sclerosis (AAN 2014, Pioro et al. 2010), multiple sclerosis (AAN 2016, Pioro et al, 2010), stroke (2016 AHA/ASA)].
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Nulojix

Products Affected

• NULOJIX

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Nuplazid

Products Affected

• NUPLAZID

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Initial:3 months, Maintenance: 1 Year
Other Criteria	Initial therapy: Individual has a diagnosis of Parkinson's disease AND Symptoms of psychosis developed after the PD diagnosis AND Symptoms of psychosis include at least one of the following: (1) Visual hallucinations, (2) Auditory hallucination OR (3) Delusions AND Symptoms have been present for at least one month AND Individual has experienced symptoms at least once weekly. Psychiatric symptoms cannot be attributed to disorders such as schizophrenia, schizoaffective disorder, delusional disorder, or mood disorder with psychotic features, or a general medical condition including delirium. For continued therapy, the individual has had a reduction in symptoms of psychosis compared to baseline.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Nurtec

Products Affected

• NURTEC

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has had a trial of and inadequate response or intolerance to two oral triptans (AHS 2019) OR has one of the following cardiovascular or non-coronary vascular contraindications to use of triptans: (a) Ischemic coronary artery disease (CAD) including angina pectoris, history of myocardial infarction, documented silent ischemia, coronary artery vasospasm (including Prinzmetal's angina) or (b) History of stroke or transient ischemic attack (TIA) or (c) Peripheral vascular disease or (d) Ischemic bowel disease or (e) Uncontrolled hypertension.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Nuvigil

Products Affected

- armodafinil oral tablet 150 mg, 200 mg, 250 mg, 50 mg
- NUVIGIL ORAL TABLET 150 MG, 200 MG, 250 MG, 50 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For Narcolepsy type 1: confirmed by the presence of daily periods of irrepressible need to sleep or daytime lapses into sleep occurring for at least 3 months and at least one of the following: 1. Clear cataplexy (defined as more than one episode of generally brief) usually bilaterally symmetrical, sudden loss of muscle tone with retained consciousness) AND 2.Multiple Sleep Latency Test (MSLT) showing one of the following: a. Mean sleep latency of less than 8 minutes with evidence of two sleep-onset rapid eye movement periods (SOREMPs) (ICSD-3, 2014)OR b. At least one SOREMP on MSLT and a SOREMP (less than 15 minutes) on the preceding overnight polysomnography (PSG) OR 3.Cerebrospinal fluid hypocretin-1 deficiency (less than 110 pg/mL or less than one-third of the normative values with the same standardized assay). For Narcolepsy type 2: confirmed by 1.Multiple sleep latency of less than 8 minutes with and evidence of two sleep-onset rapid eye movement periods (SOREMPs) ICSD-3, 2014) OR b. At least one SOREMP on MSLT and a SOREMP (less than 15 minutes) on the preceding overnight polysomnography (PSG) AND 2.The absence of cataplexy AND 3.Exclusion of alternative causes of excessive daytime sleepiness by history, physical exam and Polysomnography.
Age Restrictions	
Prescriber Restrictions	

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Coverage Duration	1 year
Other Criteria	For obstructive sleep apnea/hypopnea syndrome objectively confirmed by polysomnography (PSG) or home testing with portable monitor showing one of the following (AASM 2009): (1).Greater than 15 obstructive events (defined as apneas, hypopneas plus respiratory event related arousal) per hour of sleep OR (2)Greater than 5 obstructive events per hour of sleep and individual reports any of the following: a.Unintentional sleep episodes during wakefulness or b. Daytime sleepiness or c. Unrefreshing sleep or d. Fatigue or e. Insomnia or f.Waking up breath holding, gasping or choking or g.Bed partner describing loud snoring, breathing interruptions or both or h. Presence of comorbid conditions including hypertension, mood disorder, cognitive dysfunction, coronary artery disease, stroke, congestive heart failure, atrial fibrillation or type 2 diabetes mellitus. And has an Epworth Sleepiness Scale score greater than or equal to 10. For Shift-Work Sleep Disorder (SWSD) confirmed by all of the following: (1) No other medical disorder or mental disorder accounts for the symptoms AND (2)Symptoms do not meet criteria for any other sleep disorder (i.e. jet lag) AND (3) Symptoms have occurred for at least 3 months, AND (4) Individual has one of the following: a.Individual has excessive sleepiness or insomnia associated with a work period that occurs during the usual sleep phase OR b.Polysomnography demonstrates loss of a normal sleep- wake pattern (such as, disturbed chronobiological rhythmicity).
Indications	All Medically-accepted Indications.
Off Label Uses	

Ocaliva

Products Affected

OCALIVA

PA Criteria	Criteria Details
Exclusion Criteria	Individual has a diagnosis of nonalcoholic steatohepatitis (NASH), nonalcoholic fatty liver disease (NAFLD), primary sclerosing cholangitis (PSC), or biliary atresia. Individual has complete biliary obstruction.
Required Medical Information	Individual has a diagnosis of primary biliary cholangitis (PBC) as confirmed by TWO of the following (Lindor, 2009): (a) Elevated alkaline phosphatase. (b) Positive antimitochondrial antibodies (AMA) titer. (c) Liver biopsy with findings consistent with PBC.
Age Restrictions	Individual is 18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For Initial request, Individual has had a one year trial of ursodiol (Urso 250, Urso Forte) with an inadequate response as demonstrated by one of the following (FDA Ad Com, Lindor, 2009): (a) Alkaline phosphatase greater than or equal to 1.67 times the upper limit of normal OR (b) Total bilirubin greater than the upper limit of normal but less than two times the upper limit of normal) AND Individual will be utilizing Ocaliva (obeticholic acid) in combination with ursodiol (Urso 250, Urso Forte) OR has an intolerance to ursodiol (Urso 250, Urso Forte). For continuing treatment with Ocaliva (obeticholic acid), individual has previously met the initiation criteria above and: (a) Individual has achieved an adequate response of alkaline phosphatase or total bilirubin AND (b) Documentation has been provided.
Indications	All Medically-accepted Indications.
Y0114_20_114077	_I_012 December 2020

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

Ocrevus

Products Affected

OCREVUS

PA Criteria	Criteria Details
Exclusion Criteria	Diagnosis of hepatitis B virus infection or hepatitis C virus infection or another active infection at initiation of therapy OR History of life- threatening infusion reaction of ocrelizumab OR treating secondary progressive multiple sclerosis (MS), systemic lupus erythematosus or rheumatoid arthritis OR concurrent use with other MS disease modifying agents (such as Aubagio, Gilenya, Tecifdera, Tysabri, Lemtrada, Copaxone/Glatopa, Extavia, Rebif, Avonex, Plegridy or Betaseron).
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For diagnosis of primary progressive multiple sclerosis (PPMS) and individual able to ambulate more than 5 meters (not considered wheelchair bound). For diagnosis of relapsing multiple sclerosis (RMS) and individual able to ambulate without aid or rest for at least 1000 meters AND individual has not experienced a least 2 relapses within the previous 2 years or one relapse within the previous year.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Octreotide Line

Products Affected

- BYNFEZIA PEN
- octreotide acetate
- SANDOSTATIN
- SANDOSTATIN LAR DEPOT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.

PA Criteria	Criteria Details
Other Criteria	Individual has (A) diagnosis of carcinoid tumors for any of the following: Metastatic carcinoid tumors OR Carcinoid syndrome to suppress or inhibit severe diarrhea and flushing episodes associated with the disease OR Prophylactic administration prior to biopsy in an individual with a suspected functioning carcinoid tumor OR Prophylactic administration prior to induction of anesthesia in an individual with a functional carcinoid tumor OR Prophylactic administration prior to induction of anesthesia in an individual with a functional carcinoid tumor OR Prophylactic administration prior to induction of anesthesia in an individual with a functional carcinoid tumor OR Prophylactic administration to a surgical procedure in an individual with a functional carcinoid tumor. OR (B) Diagnosis of Bleeding gastroesophageal varices when GE varices are associated with liver disease AND octreotide acetate is used in combination with endoscopic therapy or alone if endoscopic therapy is not available. OR (C) Chemotherapy or radiation-induced diarrhea that is unresponsive to conventional antidiarrheal medications (for example, diphenoxylate and atropine or loperamide) OR (D) Malignant bowel obstruction to manage GI symptoms (such as nausea, vomiting or pain). OR (E) central nervous system meningiomas that are surgically inaccessible, recurrent, or progressive and is not a candidate for further radiation therapy OR (F) thymic carcinoma or thymoma with or without prednisone OR (G) Using for rapid relief of symptoms or for breakthrough symptoms in individuals taking long-acting octreotide acetate when any of the criteria are met for the above uses OR (H) Neuroendocrine Tumors: (i) Management of unresectable locoregional disease or distant metastasis or (ii) As treatment of the profuse watery diarrhea associated with VIPomas or (iii) Treatment of underlying hypergastrinemic Zollinger-Ellison syndrome or (iv) Prophylactic treatment prior to surgery for gastrinoma.
Indications	All Medically-accepted Indications.
Off Label Uses	

Odactra

Products Affected

• ODACTRA

PA Criteria	Criteria Details
Exclusion Criteria	Severe, unstable or uncontrolled asthma. History of any severe systemic allergic reaction. History of eosinophilic esophagitis. Or, individual is receiving concomitant therapy with other allergen immunotherapy product.
Required Medical Information	For house dust mite-induced allergic rhinitis, individual has a documented positive skin test OR positive in vitro testing for pollen-specific IgE antibodies to Dermatophagoides farinae or Dermatophagoides pteronyssinus house dust mites. Individual has had a trial of, and inadequate symptom control or intolerance to (1) nasal steroid and (1) non-sedating antihistamine AND individual has a confirmed (verbal or written attestation) prescription for an auto-injectable epinephrine product.
Age Restrictions	Individual is between the ages of 18 years and 65 years old.
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Odomzo

Products Affected

• ODOMZO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 Year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Ofev

Products Affected

• OFEV

PA Criteria	Criteria Details
Exclusion Criteria	Using in combination with Esbriet (pirfenidone). Individuals with severe hepatic impairment (child pugh class C) or end-stage liver disease.
Required Medical Information	Diagnosis of idiopathic pulmonary fibrosis (IPF) is confirmed by: Exclusion of other known causes of interstitial lung disease (ILD) such as domestic and occupational environmental exposures, connective tissue disease, and drug toxicity AND High resolution computed tomography (HRCT) with or without surgical lung biopsy. Individual has documented pulmonary function tests within prior 60 days with a Forced Vital Capacity (% FVC) greater than or equal to 50%.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Olumiant

Products Affected

• OLUMIANT

PA Criteria	Criteria Details
Exclusion Criteria	Using in combination with other JAK inhibitors (such as Xeljanz), biologic drugs (such as but not limited to, TNF agents, anti-CD20 monoclonal antibodies, IL-1R antagonists, selective co-stimulation modulators) or potent immunosuppressants (such as azathioprine and cyclosporine). At initiation of therapy, absolute neutrophil count (ANC) less than 1000 cells/mm3, lymphocyte count less than 500 cells/mm3, or hemoglobin less than 8 g/dL. Tuberculosis or other active serious infections or a history of recurrent infection. Individual has not had a tuberculin skin test (TST) or a Centers for Disease Control (CDC-) and Prevention -recommended equivalent to evaluate for latent tuberculosis prior to initiating Olumiant. Individual has severe hepatic impairment (Child Pugh class C) OR has a diagnosis of moderate [30-59 mL/min/1.73 m2 (KDIGO 2012)] or severe [less than 30 mL/min/1.73 m2 (KDIGO 2012)] renal impairment.
Required Medical Information	
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year.

PA Criteria	Criteria Details
Other Criteria	For moderate to severe RA, individual has had an inadequate reponse to/intolerant of/contraindication to conventional therapy [nonbiologics DMARDS 9such as methotrexate, sulfasalazine, leflunomide or hydroxychloroquine)] AND a trial and inadequate response/intolerance to Enbrel(etanercept) OR Humira(adalimumab) OR agents tried and failed (Humira(adalimumab) and /Enbrel(etanercept)) are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Olumiant (baricitinib) or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis.
Indications	All Medically-accepted Indications.
Off Label Uses	

Oncaspar

Products Affected

• ONCASPAR

PA Criteria	Criteria Details
Exclusion Criteria	History of serious thrombosis with prior L-asparaginase therapy. History of pancreatitis with prior L-asparaginase therapy. History of serious hemorrhagic events with prior L-asparaginase therapy.
Required Medical Information	Individual is using Oncaspar as a component of a multi-agent chemotherapeutic regimen AND is using for Acute lymphoblastic lymphoma or acute lymphocytic leukemia (ALL) or Extranodal natural killer T-cell lymphoma, nasal type (ENKL).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Onfi

Products Affected

- clobazam oral suspension
- clobazam oral tablet 10 mg, 20 mg
- ONFI ORAL SUSPENSION
- ONFI ORAL TABLET 10 MG, 20 MG
- SYMPAZAN ORAL FILM 10 MG, 20 MG, 5 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Diagnosis (all ages) and if over 65 years of age or older, Prescriber must acknowledge this is a high risk medication (HRM) [as identified by American Geriatric Society] and medication benefits outweigh potential risk for this individual.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Onpattro

Products Affected

• ONPATTRO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has TTR (transthyretin) mutation confirmed by genotyping
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Opdivo

Products Affected

OPDIVO

PA Criteria	Criteria Details
Exclusion Criteria	Receiving therapy for an autoimmune disease or chronic condition requiring treatment with an immunosuppressant. Previous treatment with another programmed death receptor-1 (PD-1) blocking antibody agent.
Required Medical Information	Current ECOG performance status 0-2. For renal cell carcinoma, histologic confirmation with clear-cell component.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For unresectable or metastatic melanoma: used as a single agent or in combination with Yervoy, as first-line therapy for untreated melanoma OR used as a single agent or in combination with Yervoy, as second-line or subsequent therapy for documented disease progression while receiving or since completing most recent therapy if PD-1 (programed death receptor -1) agent not previously used. For resected advanced melanoma for up to 12 months of adjuvant therapy when individual has resected state IIIB, IIIC or stage IV disease AND nivolumab is used as a single agent. For malignant pleural mesothelioma, used as subsequent therapy OR individual is ineligible for platinum-based therapy, defined as having one or more of the following risk factors for platinum-based chemotherapy toxicity: ECOG performance status equal to 2, Glomerular filtration rate less than 60ml/min, hearing loss (measured at audiometry) of 25 dB at two contiguous frequencies, or Grade 2 or greater peripheral neuropathy. For first- line tx of stage IV or recurrent NSCLC when: agent is used in combination with ipilmumab AND cytologically confirmed stage IV or recurrent NSCLC AND high tumor mutation burden (greater than or equal to 10 mutations per megabase) AND no sensitizing epidermal growth factor receptor mutations or anaplastic lymphoma kinase translation in nonsquamous carcinoma AND individual has not received prior systemic therapy as primary therapy for advanced or metastatic NSCLC, prior adjuvant or neoadjuvant chemotherapy is permitted as long as the administration of the prior regimen occurred at least 6 months prior. For renal cell carcinoma, agent used as single agent OR used in combination with ipilmumab if no checkpoint blockade (PD-1, PD-L1 or CTLA-4) antibody treatment has been previously administered.
Indications	All Medically-accepted Indications.
Off Label Uses	

Opsumit

Products Affected

• OPSUMIT

PA Criteria	Criteria Details
Exclusion Criteria	Individual is initiating therapy and has a diagnosis of clinically significant/severe anemia or in combination with other endothelin receptor antagonist (ERA) agents, such as but not limited to Letairis (ambrisentan) or Tracleer (bosentan).
Required Medical Information	Individual has catheterization-proven diagnosis of pulmonary arterial hypertension (PAH) (WHO Group 1) AND Individual has WHO Functional Class II-IV symptoms.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Oralair

Products Affected

- ORALAIR
- ORALAIR CHILDRENS STARTER PACK

PA Criteria	Criteria Details
Exclusion Criteria	Severe, unstable or uncontrolled asthma. History of any severe systemic allergic reaction. Or, individual is receiving concomitant therapy with other allergen immunotherapy product. History of eosinophillic esophagitis.
Required Medical Information	For grass pollen induced allergic rhinitis, individual has a confirmed (verbal or written attestation) positive skin test OR positive in vitro testing for pollen-specific IgE antibodies for at least one of the following grass pollens: Sweet vernal, orchard, perennial rye, Timothy, or Kentucky blue grass pollen. Individual has had a trial of, and inadequate symptom control or intolerance to one (1) nasal steroid and (1) non-sedating antihistamine AND individual has a confirmed (verbal or written attestation) prescription for an auto- injectable epinephrine product.
Age Restrictions	Individual is between the ages of 10 years and 65 years old.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Treatment is initiated at least 16 weeks before the expected onset of grass pollen season and is continued throughout the season.
Indications	All Medically-accepted Indications.
Off Label Uses	

Orencia

Products Affected

• ORENCIA CLICKJECT

MG/0.4ML, 87.5 MG/0.7ML

- ORENCIA INTRAVENOUSORENCIA SUBCUTANEOUS SOLUTION
- PREFILLED SYRINGE 125 MG/ML, 50

PA Criteria	Criteria Details
Exclusion Criteria	Using in combination with TNF antagonists or other biologic RA therapy, such as anakinra. Tuberculosis, other active serious infections or a history of recurrent infections. Individual has not had a tuberculin skin test (TST) or Centers for Disease Control (CDC)-recommended equivalent to evaluate for latent tuberculosis.
Required Medical Information	
Age Restrictions	For RA, Patient is 18 years of age or older. For JIA, Patient is 2 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For RA, Individual has had an inadequate response to ONE conventional therapy [non-biologic DMARD (such as methotrexate, sulfasalazine, leflunomide, or hydroxychloroquine)] or a tumor necrosis factor (TNF) antagonist AND has had a trial and inadequate response or intolerance to Humira (adalimumab) OR Enbrel (etanercept). For PsA, individual has had an inadequate response to ONE conventional therapy [nonbiologic DMARDs (such as methotrexate, sulfasalazine, or leflunomide)] or a tumor necrosis factor antagonist AND has had a trial and inadequate response or intolerance to Humira (adalimumab) OR Enbrel (etanercept). For JIA, Individual has had an inadequate response or intolerance to Humira (adalimumab) OR Enbrel (etanercept). For JIA, Individual has had an inadequate response to, is intolerant of, or has a contraindication to ONE conventional Therapy [non-biologic DMARD such as methotrexate)] or a TNF antagonist AND has had a trial and inadequate response or intolerance to Humira (adalimumab) OR Enbrel (etanercept). For any of the above indications, if the TNF agent (Enbrel(etanercept)/Humira(adalimumab)) tried and failed are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Orencia or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis OR mbr has either concomitant clinical condition: (a) Demyelinating disease or (b) Heart failure with documented left ventricular dysfunction, Orencia may be allowed without trial of preferred TNF agents (Enbrel(etanercept)/Humira(adalimumab)).
Indications	All Medically-accepted Indications.
Off Label Uses	

Orenitram

Products Affected

• ORENITRAM

PA Criteria	Criteria Details
Exclusion Criteria	Moderate (child-Pugh Class B) or severe hepatic impairment (Child Pugh Class C). Using in combination with other treprostinil dosage forms (SQ, IV, and inhalation) unless transitioning from one dose form to another. Using in combination with other prostacyclin analogs [such as but not limited to epoprostenol (Flolan, Veletri, Ventavis (iloprost)] or prostacyclin receptor agonists [such as but not limited to Uptravi (selexipag)].
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Orfadin

Products Affected

- nitisinone
- ORFADIN

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual's plasma tyrosine level is maintained below 500 micromol/L to reduce risk of ocular symptoms, developmental delay, or hyperkeratotic plaques.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Orilissa

Products Affected

ORILISSA ORAL TABLET 150 MG, 200
 MG

PA Criteria	Criteria Details
Exclusion Criteria	Individual has any of the following: (a) osteoporosis (b) Individual has severe hepatic impairment [Child-Pugh class C] (c) Individual is requesting in concurrent therapy with hormonal contraceptives (d) Individual is requesting in concurrent therapy with contraindicated agents, such as but not limited to, cyclosporine or gemfibrozil.
Required Medical Information	
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	Initial approval is 6 months, Renewal is 6 months. Requests to continue therapy beyond 24 months (2

PA Criteria	Criteria Details
Other Criteria	For initial requests, Individual is using for moderate or severe endometriosis-associated pain AND has had a trial and inadequate response or intolerance to two of the following medications or has a contraindication to all of the following medications (ACOG 2010): (a) Nonsteroidal antiinflammatory drugs (NSAIDs) OR (b) Combined oral contraceptives (OCs) OR (c) Oral or depot medroxyprogesterone (Provera, Depo-Provera) OR (d) Oral norethindrone. AND one of the following: (a) is naive to Orilissa (elagolix) OR (b) is using low dose (150 mg once daily), has mild (Child-Pugh class A) or no hepatic impairment, and has utilized Orilissa (elagolix) for a combined total duration of less than 24 months in their lifetime OR (c) is using high dose (200 mg twice daily) or has moderate hepatic impairment (Child-Pugh class B), and has utilized Orilissa (elagolix) for a combined total duration of less than 6 months in their lifetime. For continuation requests, Individual is using low dose (150 mg once daily) and does not have moderate hepatic impairment (Child-Pugh class B) AND has experienced a clinically significant improvement in endometriosis- associated pain.
Indications	All Medically-accepted Indications.
Off Label Uses	

Orkambi

Products Affected

- ORKAMBI ORAL PACKET
- ORKAMBI ORAL TABLET

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Mutation testing confirms (verbal or written) the individual has two copies of the F508del mutation in the cystic fibrosis transmembrane conductance regulator (CFTR) gene.
Age Restrictions	Individual is 2 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Products Affected

- OTEZLA ORAL TABLET
- OTEZLA ORAL TABLET THERAPY PACK

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual is unable to take biologic agent due to product warning or contraindication for any of the following: Serious infection or sepsis, Chronic or recurrent infection, Tuberculosis infection, OR Malignancy. For plaque psoriasis (Ps) involves greater than five percent (5%) body surface area (BSA) or involves less than or equal to five percent (5%) BSA involving sensitive areas or areas that significantly impact daily function (such as palms, soles of feet, head/neck, or genitalia).
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For Psoriatic Arthritis (PsA), Individual has had an inadequate response to, is intolerant of, or has a contraindication to conventional therapy [nonbiologic DMARDS (such as methotrexate, sulfasalazine, leflunomide)] AND individual has had a trial and an inadequate response or is intolerant to: Humira (adalimumab) OR Enbrel (etanercept). For plaque psoriasis (Ps), Individual has had an inadequate response to, is intolerant of, or has a contraindication to phototherapy or other systemic therapy (such as acitretin, cyclosporine or methotrexate) AND individual has had a trial and an inadequate response or is intolerant to: Humira (adalimumab) OR Enbrel (etanercept). For any of the above indications, if the TNF agent (Enbrel(etanercept)/Humira(adalimumab)) tried and failed are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Otezla or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis.
Indications	All Medically-accepted Indications.
Off Label Uses	

Oxandrin

Products Affected

• oxandrolone oral tablet 10 mg, 2.5 mg

PA Criteria	Criteria Details
Exclusion Criteria	Carcinoma of the prostate or breast in male individuals OR Carcinoma of the breast in females with hypercalcemia. Using to enhance athletic performance or physique. Individual has a diagnosis of nephrosis (nephrotic phase of nephritis). Individual has a diagnosis of hypercalcemia.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	6 months
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

OxyContin

Products Affected

- oxycodone hcl er
- OXYCONTIN

PA Criteria	Criteria Details
Exclusion Criteria	Individual is requesting or using as an as-needed analgesic OR Individual has one of the following conditions: (a) Significant respiratory depression OR (b) Acute or severe bronchial asthma or hypercarbia OR (c) Known or suspected paralytic ileus.
Required Medical Information	
Age Restrictions	Individual is 11 years of age or older
Prescriber Restrictions	
Coverage Duration	Initial 3 months. Maintenance 6 months. Cancer Pain/Terminal Dx or Palliative Care 1 Year.

PA Criteria	Criteria Details
Other Criteria	For initial use, Individual has one of the following: (a) Diagnosis of cancer related pain and/or is actively undergoing cancer therapy (provide cancer diagnosis) OR (b) Diagnosis of terminal illness and is receiving palliative/end-of-life care (provide terminal diagnosis) OR Individual has pain severe enough to require daily, around-the-clock, long term opioid treatment (provide diagnosis) AND Individual has one of the following: (a) An inadequate response to ONE alternative treatment options, such as but not limited to non-opioid analgesics and immediate-release opioids OR (b) Alternative treatment options would otherwise be inadequate to provide sufficient management of pain OR (c) Individual has contraindications to non-opioid analgesics (such as NSAID use in individuals with active ulcer condition/gastrointestinal bleeding, renal failure). Individual is not opioid analgesia as an inpatient for post-surgical pain OR (b) Individual is transitioning from one long-acting opioid analgesic to another long-acting opioid analgesic OR (c) already receiving and tolerating a minimum daily opioid dose of at least 20mg oxycodone orally or its equivalent. For continued use, Attestation (verbal or written) that long-acting opioid therapy has provided meaningful improvement in pain and/or function compared to baseline AND Prescriber has consulted with individual regarding risks of opioid therapy AND Clear treatment goals have been defined and outlined as part of overall plan.
Indications	All Medically-accepted Indications.
Off Label Uses	

Ozurdex Implant

Products Affected

OZURDEX

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Padcev

Products Affected

• PADCEV

PA Criteria	Criteria Details
Exclusion Criteria	Individuals have moderate or severe hepatic impairment (Child- Pugh B or C) OR the criteria not met.
Required Medical Information	Individual has diagnosis of locally advanced or metastatic urothelial cancer AND using as subsequent therapy after progression with anti-PD-1 and anti-PD-L1 agent AND individual has current ECOG performance status of 0 - 2.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

PAH - B

Products Affected

- epoprostenol sodium
- FLOLAN
- VELETRI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Diagnostic Criteria for PAH: right heart catheterization which shows a mean pulmonary artery pressure (mPAP) greater than 25mmhg, a pulmonary capillary wedge pressure (PCWP), left atrial pressure or left ventricular end diastolic pressure (LVEDP) less than or equal to 15mmhg AND a pulmonary vascular resistance (PVR) greater than 3 wood units. Criteria for Vasodilator Response: a favorable response is defined as a fall in mPAP of at least 10mmhg to an absolute mPAP of less than 40mmhg without a decrease in cardiac output, when challenged with inhaled nitric oxide, intravenous epoprostenol or intravenous adenosine.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For continuous intravenous infusion of Epoprostenol, individual must meet all of the following: meets diagnostic criteria for PAH AND demonstrates an unfavorable acute response to vasodilators AND individual has New York Heart Association (NYHA) functional class III, or IV symptoms AND World health Organization (WHO) Group I PAH (idiopathic PAH, PAH associated with connective tissue disorders, PAH associated with congenital heart defects, and all Group 1 subtypes).
Indications	All Medically-accepted Indications.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

Palynziq

Products Affected

• PALYNZIQ

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has a confirmed prescription for an auto-injectable epinephrine agent.
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For initial requests, individual has a diagnosis of phenylketonuria (PKU) and has uncontrolled blood phenylalanine (PHE) concentrations (greater than 600 micromol/L) on existing management, including but not limited to the following: (a) Dietary phenylalanine and/or protein restriction (b) Kuvan (sapropterin dihydrochloride) agents. For continued use, Individual is showing signs of continuing improvement, as evidenced by blood PHE levels decrease of at least 20% from pretreatment baseline or a reduction below 600micromol/L after 16 weeks of treatment at a maximum dose of 40mg/day.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Pegfilgrastim Agents

Products Affected

• FULPHILA

• ZIEXTENZO

- NEULASTA
- NEULASTA ONPRO
- UDENYCA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individuals who are at high risk for infection-associated complications by any of the following: Expected prolonged (greater than 10 day) and profound (less than 0.1 x 10 to the power of 9/L) neutropenia, Age greater than 65 years, Pneumonia or other clinically documented infections, Hypotension and multi organ dysfunction (sepsis syndrome), Invasive fungal infection, Prior episode of febrile neutropenia, Hospitalized at the time of the development of fever. Primary prophylaxis of FN in patients who have a risk of FN of 20% or greater based on chemotherapy regimens. OR when the risk of developing FN is greater than or equal to 10% and less than 20% based on chemotherapy in patients who have risk factors for FN including any of the following: Individual age greater than 65 years, Poor performance status (ECOG status 3-4), Prior chemotherapy or radiation therapy, Bone marrow involvement by tumor producing cytopenias, persistent neutropenia (ANC less than 1500mm3), poor renal function (GFR less than 60mL/min), liver dysfunction, recent surgery and or presence of open wounds.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Other Criteria	Individual with nonmyeloid malignancy is using for Secondary Prophylaxis for patients who experienced a neutropenic complication from a prior cycle of chemotherapy (for which primary prophylaxis was not received), in which a reduced dose may compromise disease-free or overall survival or treatment outcome. Adjunctive use in neutropenic patients who are at high risk for infection-associated complications. In an individual with acute lymphocytic leukemia (ALL) after completion of the first few days of initial induction chemotherapy or first post-remission course of chemotherapy. For use after accidental or intentional total body radiation of myelosuppressive doses (greater than 2 Grays [Gy]) (such as Hematopoietic Syndrome of Acute Radiation Syndrome). For dose dense therapy (treatment given more frequently, such as every two weeks instead of every three weeks) for adjuvant treatment of breast cancer. After a hematopoietic progenitor stem cell transplant (HPCT/HSCT) to promote myeloid reconstitution OR when engraftment is delayed or has failed.
Indications	All Medically-accepted Indications.
Off Label Uses	

Pemazyre

Products Affected

• PEMAZYRE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual is using as monotherapy AND has confirmed disease progression (written or verbal) after one or more prior lines of systemic therapy AND confirmation (written or verbal) of fibroblast growth factor receptor 2 (FGFR2) fusion or non-fusion rearrangement as detected by an FDA-approved test.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Perjeta

Products Affected

• PERJETA

PA Criteria	Criteria Details
Exclusion Criteria	If administered after Herceptin (trastuzumab) is discontinued or as part of a regimen without Herceptin (trastuzumab). Concomitant use with other targeted biologic agents (including but not limited to erlotinib, cetuximab, panitumumab, bevacizumab, ziv-aflibercept and lapatinib).
Required Medical Information	Tumor(s) have been evaluated with an assay validated to predict HER2 positive (HER2+) protein overexpression. Individuals are considered HER2 positive (HER2+) as documented by one of the following, immunohistochemistry (IHC) 3+ or In Situ hybridization (ISH) positive by any of the following: Single probe average HER2 copy number greater than or equal to 6.0 signals/cell OR Dual- probe HER2/CEP 17 ratio greater than or equal to 2.0 OR Dual- probe HER2/CEP 17 ratio less than 2.0 with an average HER2 copy number greater than or equal to 6.0 signals/cell.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For metastatic breast cancer use Perjeta will be used in combination with trastuzumab AND either docetaxel or paclitaxel. (Note If docetaxel or paclitaxel treatment is discontinued (for example, related to toxicity), treatment with Perjeta and trastuzumab may continue.) AND combination chemotherapy with Perjeta (pertuzumab) will be used as single line anti-HER2 chemotherapy for metastatic disease until progression OR individual has early stage, locally advanced or inflammatory breast cancer and will undergo neoadjuvant therapy (prior to surgery) or adjuvant systemic therapy AND primary tumor is larger than 2cm or individual is lymph node positive (for neoadjuvant therapy: clinically evident by palpation or imaging) AND used in combination with trastuzumab and with one of the following: docetaxel with or without carboplatin or paclitaxel AND pertuzumab is used for a maximum of 18 cycles (12 month course).
Indications	All Medically-accepted Indications.
Off Label Uses	

Phesgo

Products Affected

• PHESGO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has HER2-positive breast cancer confirmed (verbal or written) by EITHER immunohistochemistry (IHC) of 3+ OR positive In situ hybridization (ISH).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Piqray

Products Affected

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

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has confirmed (written or verbal attestation is acceptable) PIK3CA mutation using an FDA-approved test (such as the therascreen PIK3CA RGQ PCR Kit).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Pomalyst

Products Affected

 POMALYST ORAL CAPSULE 1 MG, 2 MG, 3 MG, 4 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Praluent

Products Affected

• PRALUENT

PA Criteria	Criteria Details
Exclusion Criteria	Concurrent use with Juxtapid or Kynamro.
Required Medical Information	For (A) Heterozygous Familial Hypercholesterolemia (HeFH) confirmed by: 1.Presence of a mutation in LDLR, apolipoprotein B (ApoB), PCSK9 or ARH adaptor protein gene (LDLRAP1) gene OR 2. World Health Organization (WHO)/Dutch Lipid Network Criteria with score of gtr than 8 points OR (B) History of Clinical Atherosclerotic Cardiovascular Disease (ASCVD), including one of the following: 1.Acute coronary syndromes 2.Coronary Artery Disease (CAD) 3. History of myocardial infarction (MI) 3.Stable or unstable angina 4.Coronary or other arterial revascularization 5.Stroke 6.Transient ischemic attack (TIA) 7.Peripheral arterial disease (PAD). OR (C) Primary hyperlipidemia alone or in combination with other lipid lowering agents. OR (D) using prophylactically for Established CVD.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Initial 3 month. Continuation 1 yr.

PA Criteria	Criteria Details
Other Criteria	For initial request, individual meets one of the following: (A) Individual is a on high intensity statin therapy, or statin therapy at the maximum tolerated dose, where high intensity statin is defined as atorvastatin 40 mg or higher OR rosuvastatin 20 mg or higher OR (B) Individual is statin intolerant OR (C) Individual has a condition that is a contraindication for statin therapy including active liver disease, unexplained persistent elevation of serum hepatic, or pregnancy. Individual also has had an adequate trial and titration of a Repatha (evolocumab) and has achieved suboptimal lipid lowering response despite at least 90 days of Repatha (evolocumab) therapy. For continuation, Individual continues to receive concomitant maximally tolerated statin (unless contraindication or individual is statin intolerant) AND there is confirmation (verbal or written attestation) of LDL reduction.
Indications	All Medically-accepted Indications.
Off Label Uses	

Probuphine Implant

Products Affected

• PROBUPHINE IMPLANT KIT

PA Criteria	Criteria Details
Exclusion Criteria	New entrants to treatment. Treatment for longer than 12 months with Probuphine. Retreatment with Probuphine after a prior 12- month treatment period. For individuals who have not achieved and sustained prolonged clinical stability while being maintained on buprenorphine 8 mg per day or less of a sublingual tablet or its transmucosal buprenorphine product equivalent.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	6 months
Other Criteria	Individual has been diagnosed with opioid dependence (opioid use disorder) and individual has been treated with a stable transmucosal buprenorphine dose (of 8 mg per day or less of a sublingual tablet or its transmucosal buprenorphine product equivalent) for 3 months or longer without any need for supplemental dosing or adjustments and individual is currently on a maintenance dose of 8 mg per day or less of a sublingual tablet or its transmucosal buprenorphine product equivalent to achieve sustained prolonged clinical stability on transmucosal buprenorphine and Probuphine is used as part of a substance use disorder treatment program to include counseling and psychosocial support.
Indications	All Medically-accepted Indications.
	Z_I_012 December 2020

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

Prolia

Products Affected

PROLIA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Osteoporosis is defined as a BMD T-Score of less than or equal to - 2.5 OR a clinical diagnosis based on a history of a low trauma fracture (fragility fracture) at high risk for fracture. Risk factors for osteoporotic fracture is defined as: Hypogonadism or premature ovarian failure, Low body mass, Smoking, Rheumatoid arthritis, Alcohol intake of 3 or more drinks/day, Vitamin D deficiency, Low calcium intake, Hyperkyphosis, Parental hip fracture, Multiple falls, Medication: Anticoagulants, anticonvulsants, aromatase inhibitors, cancer chemotherapeutic drugs, gonadotropin-releasing hormone agonists, (glucocorticoid daily dosage equivalent to 5 mg or greater of prednisone for at least 3 months). Glucocorticoid-induced osteoporosis defined as a T score -2.5 or less and is initiating or continuing systemic glucocorticoids in a daily dosage equivalent to 7.5mg or greater of prednisone and expected or remain on glucocorticoids for a least 6 months.
Age Restrictions	For Osteoporosis 18 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For osteoporosis/ glucocorticoid-induced osteoporosis treatment, individual has had at least ONE osteoporotic (minimal trauma) fracture OR has two or more risk factors for osteoporotic fracture OR Individual has failed or is intolerant to or has a medical contraindication to other available osteoporosis therapies (such as, bisphosphonates). For male receiving androgen deprivation therapy for non- metastatic prostate cancer, individual has had at least ONE osteoporotic (minimal trauma) fracture OR has one or more risk factors for osteoporotic fracture. Individual is a postmenopausal (natural or induced) female receiving adjuvant aromatase inhibitor therapy for the treatment of breast cancer.
Indications	All Medically-accepted Indications.
Off Label Uses	

Promacta

Products Affected

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

PA Criteria	Criteria Details
Exclusion Criteria	Using Promacta to normalize platelet counts. Use in individuals with ITP whose degree of thrombocytopenia and clinical condition (for example, platelet count greater than 30 x 109/L or active bleeding) do not increase the risk of bleeding. Use in individuals with chronic hepatitis C whose degree of thrombocytopenia does not prevent the initiation of peginterferon therapy or limits the ability to maintain an optimal peginterferon-based therapy. Used in individuals with chronic hepatitis C who are no longer on a peginterferon and ribavirin based regimen. Used concomitantly with other thrombopoietin receptor agonists such as romiplostim (Nplate). Used in individuals taking in combination with directacting antiviral agents used without concomitant use of a peginterferon agent for treatment of thrombocytopenia associated with chronic hepatitis C infection.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For initial therapy: 1) Diagnosis of chronic immune (idiopathic) thrombocytopenia (ITP) AND individual has platelet count of less than 30 x 109/L or active bleeding (ASH, 2011. Hicks et al., 2014) AND has had a prior trial and insufficient response to one of the following: a) corticosteroids or b) immunoglobulins [for example, intravenous, anti-D] or c) splenectomy. OR, 2) Diagnosis of severe aplastic anemia AND individual has a platelet count of less than or equal to $30 \times 109/L$ (Olnes et al., 2012.Desmond et al., 2014) AND individual has had a prior trial and insufficient response to an immunosuppressive therapy [such as, anti-thymocte globulin (ATG)]. OR 3) dx of severe aplastic anemia AND is being used in combination with standard immunosuppressive therapy for first-line treatment. For maintenance therapy, individual has demonstrated a response to therapy as evidenced by increased platelet counts AND to maintain an adequate platelet count (50 - 200 x 109/L) to decrease the risk of bleeding.
Indications	All Medically-accepted Indications.
Off Label Uses	

Protopic

Products Affected

- PROTOPIC
- tacrolimus external

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual had a trial of and inadequate response or intolerance to one topical prescription strength corticosteroid.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Purixan

Products Affected

• PURIXAN

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Qinlock

Products Affected

• QINLOCK

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

quinine

Products Affected

- QUALAQUIN
- quinine sulfate

PA Criteria	Criteria Details
Exclusion Criteria	Treatment or prevention for nocturnal recumbancy leg muscle cramps or related conditions such as but not limited to: Leg cramps, muscle cramps, myoclonus, Periodic Movements of Sleep, Periodic Limb Movements of Sleep (PLMS), Restless Leg Syndrome (RLS), severe hepatic impairment (Child-Pugh C), known prolongation of the QT interval, initial treatment of severe or complicated P. falciparum malaria, prevention of malaria, individuals with glucose-6-phosphate dehydrogenase (G6PD) deficiency, individuals with myasthenia gravis, or individuals with optic neuritis .
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has been diagnosed with uncomplicated malaria caused by one of the following: Plasmodium falciparum known or suspected to be resistant to chloroquine (CDC 2013) OR chloroquine-resistant Plasmodium vivax (CDC 2013) OR an unidentified plasmodial species known or suspected to be resistant to chloroquine (CDC 2013).
Indications	All Medically-accepted Indications.
Off Label Uses	
Y0114 20 114077	I 012 December 2020

Y0114_20_114077_I_012

Ragwitek

Products Affected

• RAGWITEK

PA Criteria	Criteria Details
Exclusion Criteria	Severe, unstable or uncontrolled asthma. History of any severe systemic allergic reaction. History of eosinophilic esophagitis. Or, individual is receiving concomitant therapy with other allergen immunotherapy product.
Required Medical Information	For short ragweed pollen induced allergic rhinitis, individual has a documented positive skin test OR positive in vitro testing for pollen-specific IgE antibodies for short ragweed pollen. Individual has had a trial of, and inadequate symptom control or intolerance to (1) nasal steroid and (1) non-sedating antihistamine AND individual has a confirmed (verbal or written attestation) prescription for an auto-injectable epinephrine product.
Age Restrictions	Individual is between the ages of 18 years and 65 years old.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Treatment is initiated at least 12 weeks before the expected onset of ragweed pollen season and is continued throughout the season.
Indications	All Medically-accepted Indications.
Off Label Uses	

Ravicti

Products Affected

• RAVICTI

PA Criteria	Criteria Details
Exclusion Criteria	Using for the management of acute hyperammonemia. Using to treat N-acetylglutamate synthase deficiency (NAGS).
Required Medical Information	
Age Restrictions	2 months of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has had a trial and inadequate response or intolerance to sodium phenylbutyrate (Buphenyl) OR Individual has any of the following: (a) Congestive heart failure or (b) Severe renal insufficiency or (c) A clinical state where there is sodium retention with edema.
Indications	All Medically-accepted Indications.
Off Label Uses	

Rayos

Products Affected

RAYOS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Confirmation (Verbal or written) has been provided for why the delayed-release agent is clinically necessary and not for convenience.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual has had a prior trial and inadequate response or intolerance to one generic immediate-release oral prednisone agent AND has had a prior consecutive trial and inadequate response to an additional generic oral corticosteroid agents (such as but not limited to, prednisolone, methylprednisolone, hydrocortisone).
Indications	All Medically-accepted Indications.
Off Label Uses	

Reblozyl

Products Affected

REBLOZYL

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For dx beta thalassemia/hg E beta thalassemia, Individual required regular red blood cell transfusions at initiation, defined as both of the following: individual received six to twenty RBC units in the last 24 weeks and had no transfusion-free period greater than 35 days in the last 24 weeks AND individual has a baseline hemoglobin level less than or equal to 11 g/dL. For MDS or MDS/MPN RS T, Individual has required regular red blood cell transfusions of two (2) or more RBC units over eight (8) weeks in the last 16 weeks AND has a baseline hemoglobin (Hgb) level less than or equal to 11 g/dL. AND has one of the following (A or B) (A) has a diagnosis very low to intermediate risk myelodysplastic syndromes with ring sideroblasts (MDS-RS) greater than or equal to 15% (or ring sideroblasts 5% to 14% with an SF3B1 mutation) AND meets one of the following criteria: (1) Serum erythropoietin (EPO) level of greater than 500 mU/mL OR (2) Serum EPO level of less than or equal to 500 mU/mL following no response to combination treatment with erythropoiesis-stimulating agent (ESA) and granulocyte-colony stimulating factor (G-CSF). OR (B) Individual has a diagnosis of myelodysplastic/myeloproliferative neoplasm with ring sideroblasts and thrombocytosis (MDS/MPN-RS-T) with all of the following: (1) Ring sideroblasts greater than or equal to 15% (WHO 2017) AND (2) Thrombocytosis (defined as platelets greater than or equal to 450 x109/L) (WHO 2017) AND (3) Insufficient response to ESAs.
Indications	All Medically-accepted Indications.
Off Label Uses	

Reclast

Products Affected

- RECLAST
- zoledronic acid intravenous solution 5 mg/100ml

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Regranex

Products Affected

• REGRANEX

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	3 months
Other Criteria	Individual is using as adjunctive therapy with good ulcer care practices including, but not limited to sharp debridement of the ulcer
Indications	All Medically-accepted Indications.
Off Label Uses	

RELISTOR

Products Affected

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

PA Criteria	Criteria Details
Exclusion Criteria	Individual has a known or suspected mechanical gastrointestinal obstruction.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual must have a previous trial of or insufficient response to polyethylene glycol (generic MiraLax) (AGA 2013) AND a trial and inadequate response or intolerance to a preferred agent (Movantik/Amitiza) OR the preferred agent (Movantik/Amitiza) is not acceptable due to concomitant clinical situations, warnings or contraindications, such as but not limited to the following: (1) individual is taking a diphenylheptane opioid (e.g., methadone), where effectiveness has not been established in the treatment of OIC (Amitiza) OR (2) individual has disruption to the blood-brain barrier and may be at increased risk for opioid withdrawal and/or reduced analgesia (Movantik) OR (3) individual is taking strong CYP3A4 inhibitors and may be at increased risk for opioid withdrawal and/or reduced analgesia (Movantik).
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Remicade

Products Affected

- AVSOLA
- INFLECTRA
- REMICADE
- RENFLEXIS

PA Criteria	Criteria Details
Exclusion Criteria	Tuberculosis, other active serious infections or a history of recurrent infection. Individuals who have not had a tuberculin skin test or CDC-recommended equivalent to evaluate for latent tuberculosis prior to initiating infliximab. Using in combination with other TNF antagonists, JAK inhibitors, or other biologic drugs (such as, abatacept, anakinra, tocilizumab, or vedolizumab).
Required Medical Information	For chronic moderate to severe plaque psoriasis: Greater than 5% of body surface area with plaque psoriasis OR Less than or equal to 5% body surface area with plaque psoriasis involving sensitive areas or areas that would significantly impact daily function (such as palms, soles of feet, head/neck, or genitalia).
Age Restrictions	For Crohn's Disease or Ulcerative colitis, 6 yr of age or older. For JIA, 2 yr of age or older. For all other indications 18 yr of age.
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For RA, individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE Conventional Therapy [nonbiological DMARDs (such as methotrexate, sulfasalazine, leflunomide, or hydroxychloroquine)] (ACR 2015). For Crohn's Disease, individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional therapy (such as 5-ASA products, sulfasalazine, systemic corticosteroids, or immunosuppressive drugs). For moderately to severely active Ulcerative Colitis, individual has failed to respond to, is intolerant of, or has a medical contraindication to ONE conventional therapy (such as 5-ASA products, sulfasalazine, systemic corticosteroids, or immunosuppressive drugs). For Ankylosing Spondylitis individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional therapy (such as 5-ASA products, sulfasalazine, systemic corticosteroids, or immunosuppressive drugs). For Ankylosing Spondylitis individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional therapy [such as NSAIDs, or nonbiologic DMARDs(such as sulfasalazine)] (ACR 2015)]. For Psoriatic Arthritis, individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional therapy [nonbiological DMARDs (such as methotrexate, sulfasalazine, or leflunomide)]. For chronic plaque psoriasis, individual has had an inadequate response to, is intolerant of, or has a medical contraindication to phototherapy or ONE other systemic therapy (such as methotrexate, acetretin, or cyclosporine). For Refractory Wegener's Granulomatosis, individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional Therapy [nonbiologic DMARD (such as methotrexate)]. For chronic, recurrent, treatment- refractory or vision-threatening, non-infectious uveitis, individual has had an inadequate response to, is intolerant
	AND has had an inadequate response to, is intolerant of, or has a contraindication to nonbiologic DMARDs (such as methotrexate or azathioprine).
Indications	All Medically-accepted Indications.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

Remodulin

Products Affected

- REMODULIN
- treprostinil

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Diagnostic Criteria for PAH: right heart catheterization which shows a mean pulmonary artery pressure (mPAP) greater than or equal to 25mmhg, a pulmonary capillary wedge pressure (PCWP), left atrial pressure or left ventricular end diastolic pressure (LVEDP) less than or equal to 15mmhg AND a pulmonary vascular resistance (PVR) greater than 3 wood units. Criteria for Vasodilator Response: a favorable response is defined as a fall in mPAP of at least 10mmhg to an absolute mPAP of less than 40mmhg without a decrease in cardiac output, when challenged with inhaled nitric oxide, intravenous epoprostenol or intravenous adenosine.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For Remodulin patient must meet all of the following: meets diagnostic criteria for PAH AND demonstrates an unfavorable acute response to one vasodilator AND Individual has New York Heart Association (NYHA) functional class II, III, or IV symptoms: AND World health Organization (WHO) Group I PAH (idiopathic PAH, PAH associated with connective tissue disorders, PAH associated with congenital heart defects, and all Group 1 subtypes).
Indications	All Medically-accepted Indications.
Off Label Uses	
 Y0114_20_114077	_I_012 December 2020

Repatha

Products Affected

- REPATHA
- REPATHA PUSHTRONEX SYSTEM
- REPATHA SURECLICK

PA Criteria	Criteria Details
Exclusion Criteria	Concurrent use with Juxtapid or Kynamro.
Required Medical Information	For (A). Homozygous Familial Hypercholesterolemia (HoFH) confirmed by: 1.Presence of two (2) mutant alleles at the LDLR, apoB, PCSK9 or ARH adaptor gene locus OR 2.untreated LDL-C concentration greater than 500 mg/dL (13 mmol/L) or treated LDL- C greater than or equal to 300 mg/dL (7.76 mmol/L) AND one of the following: i.Cutaneous or tendonous xanthoma before age of 10 years OR ii.Untreated LDL-C levels consistent with heterozygous familial hypercholesterolemia in both parents (greater than190 mg/dL) OR (B). Heterozygous Familial Hypercholesterolemia (HeFH) with diagnosis confirmed by: 1.Presence of a mutation in LDLR, apolipoprotein B (ApoB), PCSK9 or ARH adaptor protein gene (LDLRAP1) gene OR 2. World Health Organization (WHO)/Dutch Lipid Network Criteria with score of gtr than 8 points OR (C). History of Clinical Atherosclerotic Cardiovascular Disease (ASCVD), including one of the following: 1.Acute coronary syndromes 2.Coronary Artery Disease (CAD) 3. History of myocardial infarction (MI) 3.Stable or unstable angina 4.Coronary or other arterial revascularization 5.Stroke 6.Transient ischemic attack (TIA) 7.Peripheral arterial disease (PAD). OR (D) Primary hyperlipidemia alone or in combination with other lipid lowering agents. OR (E) using prophylactically for Established CVD.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Initial 3 month. Continuation 1 yr.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Other Criteria	For initial HoFH request, individual meets the following: (A) Individual is a on high intensity statin therapy, or statin therapy at the maximum tolerated dose, where high intensity statin is defined as atorvastatin 40 mg or higher OR rosuvastatin 20 mg or higher OR (B) Individual is statin intolerant OR (C) Statin associated rhabdomyolysis after a trial of a statin OR (D) Individual has a contraindication for statin therapy including active liver disease, unexplained persistent elevation of serum hepatic transaminases, or pregnancy AND Individual is on ezetimibe (applies to individuals on statin therapy only). For initial HeFH or ASCVD requests, individual meets the following: (A) Individual is a on high intensity statin therapy, or statin therapy at the maximum tolerated dose, where high intensity statin is defined as atorvastatin 40 mg or higher OR rosuvastatin 20 mg or higher OR (B) Individual is statin intolerant OR (C) Statin associated rhabdomyolysis after a trial of a statin OR (D) Individual has a contraindication for statin therapy including active liver disease, unexplained persistent elevation of serum hepatic transaminases, or pregnancy. For continuation (HeFH, HoFH, ASCVD), mbr continues to receive concomitant maximally tolerated statin (unless contraindication or individual is statin intolerant) AND there is confirmation (verbal or written attestation) of LDL reduction. For continuation (established CVD or Primary Hyperlipidemia), confirmation (verbal or written attestation) of LDL reduction.
Indications	All Medically-accepted Indications.
Off Label Uses	

Retevmo

Products Affected

• RETEVMO ORAL CAPSULE 40 MG, 80 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Written or verbal attestation is provided for confirmation of mutations where applicable based on use/diagnosis.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Retisert Implant

Products Affected

• RETISERT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Revatio

Products Affected

- REVATIO ORAL SUSPENSION RECONSTITUTED
- REVATIO ORAL TABLET
 - sildenafil citrate oral suspension

PA Criteria **Criteria Details** Exclusion Use in combination with phosphodiesterase-5 (PDE5) inhibitors Criteria [such as but not limited to, Viagra (sildenafil)]. Use in combination with organic nitrates, such as but not limited to, isosoribide mono/dinitrate or nitroglycerin. Use in combination with guanylate cyclase stimulators [such as but not limited to, Adempas (riociguat)]. Use in individuals requesting for the treatment of erectile dysfunction. Use in individuals with severe hepatic impairment (Child-Pugh Class C). Use in individual has a diagnosis of pulmonary veno-occlusive disease (PVOD). Use in individual has a known hereditary degenerative retinal disorder (such as but not limited to, retinitis pigmentosa). Required Catheterization-proven diagnosis of Pulmonary Arterial Medical Hypertension (PAH) World Health Organization (WHO Group I) and Information WHO Functional Class II-IV symptoms. Age Restrictions Prescriber Restrictions Coverage 1 year Duration **Other Criteria** Indications All Medically-accepted Indications. **Off Label Uses**

reconstituted

sildenafil citrate oral tablet 20 mg

Revatio IV

Products Affected

- REVATIO INTRAVENOUS
- sildenafil citrate intravenous

PA Criteria	Criteria Details
Exclusion Criteria	Use in combination with phosphodiesterase-5 (PDE5) inhibitors [such as but not limited to, Viagra (sildenafil)]. Use in combination with organic nitrates, such as but not limited to, isosoribide mono/dinitrate or nitroglycerin. Use in combination with guanylate cyclase stimulators [such as but not limited to, Adempas (riociguat)]. Use in individuals requesting for the treatment of erectile dysfunction. Use in individuals with severe hepatic impairment (Child-Pugh Class C). Use in individual has a diagnosis of pulmonary veno-occlusive disease (PVOD). Use in individual has a known hereditary degenerative retinal disorder (such as but not limited to, retinitis pigmentosa).
Required Medical Information	Catheterization-proven diagnosis of Pulmonary Arterial Hypertension (PAH) World Health Organization (WHO Group I) and WHO Functional Class II-IV symptoms.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For sildenafil INJ, individual is temporarily unable to take oral dose forms and requires continued therapy. For Individual with a diagnosis of persistent pulmonary hypertension of the newborn (AHA/ATS 2015) AND Individual was started and stabilized on Revatio (sildenafil) in the hospital and requires continued outpatient therapy.
Indications	All Medically-accepted Indications.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

Revlimid

Products Affected

• REVLIMID ORAL CAPSULE 10 MG, 15 MG, 2.5 MG, 20 MG, 25 MG, 5 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Reyvow

Products Affected

• REYVOW ORAL TABLET 100 MG, 50 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has had a trial/inadequate response or intolerance to 2 oral triptans (AHS 2019) OR Individual has one of the following CV or non-coronary vascular contraindications to use of triptans: Ischemic coronary artery disease (CAD) including angina pectoris, history of MI, documented silent ischemia, coronary artery vasospasm (including Prinzmetal's angina), history of stroke or TIA, PVD, ischemic bowel disease, or uncontrolled hypertension.
Indications	All Medically-accepted Indications.
Off Label Uses	

Rinvoq

Products Affected

RINVOQ

PA Criteria	Criteria Details
Exclusion Criteria	Using in combination with other JAK inhibitors (such as Xeljanz), biologic drugs (such as but not limited to, TNF antagonists, anti- CD20 monoclonal antibodies, IL-1R antagonists, selective co- stimulation modulators) or potent immunosuppressants (such as azathioprine and cyclosporine). At initiation of therapy, absolute neutrophil count (ANC) less than 1000 cells/mm3, lymphocyte count less than 500 cells/mm3, or hemoglobin less than 8 g/dL. Tuberculosis or other active serious infections or a history of recurrent infection. Individual has not had a tuberculin skin test (TST) or a Centers for Disease Control (CDC-) and Prevention - recommended equivalent to evaluate for latent tuberculosis prior to initiating upadacitinib. Individual has severe hepatic impairment (Child Pugh class C). Individual has end stage renal disease [less than 15 mL/min/1.73 m2 (KDIGO 2012)].
Required Medical Information	
Age Restrictions	Individual is 18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year.

PA Criteria	Criteria Details
Other Criteria	of, or has a contraindication to conventional therapy [nonbiologic DMARDS (such as methotrexate, sulfasalazine, leflunomide, or hydroxychloroquine)] AND has had an inadequate response to Enbrel (etanercept) OR Humira (adalimumab) OR agents tried and failed (Humira (adalimumab)/Enbrel (etanercept)) are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with upadacitinib or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis.
Indications	All Medically-accepted Indications.
Off Label Uses	

Rozlytrek

Products Affected

ROZLYTREK ORAL CAPSULE 100 MG, 200 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	For metastatic non-small cell lung cancer (NSCLC), 18 years of age or older. For a diagnosis of a solid tumor, 12 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For solid tumors, the individual has a neurotrophic receptor tyrosine kinase (NTRK) gene fusion without a known acquired resistance mutation with confirmed genetic test results.
Indications	All Medically-accepted Indications.
Off Label Uses	

Rubraca

Products Affected

 RUBRACA ORAL TABLET 200 MG, 250 MG, 300 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Written or verbal attestation is provided for confirmation of mutations where applicable based on use/diagnosis.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Ruconest

Products Affected

RUCONEST

PA Criteria	Criteria Details
Exclusion Criteria	Individual is using for prophylaxis or in individuals with laryngeal attacks.
Required Medical Information	Hereditary Angioedema (HAE) is confirmed by a C4 level below the lower limit of normal as defined by the laboratory performing the test AND ONE of the following (a or b): a) C1 inhibitor (C1-INH) antigenic level below the lower limit of normal as defined by the laboratory performing the test OR b) C1-INH functional level below the lower limit of normal as defined by the laboratory performing the test AND Individual has a history of moderate or severe attacks (for example, airway swelling, severe abdominal pain, facial swelling, nausea and vomiting, painful facial distortion).
Age Restrictions	13 years and older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Rydapt

Products Affected

• RYDAPT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Sabril

Products Affected

- SABRIL
- vigabatrin
- vigadrone

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	For infantile spasm 1 month to 2yr old. For seizure 10 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Samsca

Products Affected

- SAMSCA ORAL TABLET 15 MG, 30 MG
- tolvaptan

PA Criteria	Criteria Details
Exclusion Criteria	Individual has an acute, urgent need to raise serum sodium OR is unable to sense/appropriately respond to thirst OR is anuric. Diagnosis of hypovolemic hyponatremia. Individual has underlying liver disease, including cirrhosis OR using to treat autosomal dominant polycystic kidney disease OR is currently receiving a strong CYP3A inhibitor (such as clarithromycin, ketoconazole, itraconazole, ritonavir. indinavir, nelfinavir, saquinavir, nefazodone and telithromycin).
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	30 Days
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

SANCUSO

Products Affected

• SANCUSO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has had a trial of and inadequate response or intolerance to EITHER generic ondansetron or oral granisetron OR individual is unable to take oral medications due to the following: (A)The presence of head and neck cancer OR (B)Mucositis due to recent radiation to the head and neck area.
Indications	All Medically-accepted Indications.
Off Label Uses	

Sarclisa

Products Affected

• SARCLISA

PA Criteria	Criteria Details
Exclusion Criteria	Individual has a diagnosis of multiple myeloma AND has not received treatment with isatuximab or another anti-CD38 agent such as daratumumab) AND has relapsed or refractory disease following treatment with at least two prior lines of therapy including lenalidomide and a proteasome inhibitor (for example, bortezomib, carfilzomib, or ixazomib).
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Signifor IR

Products Affected

• SIGNIFOR

PA Criteria	Criteria Details
Exclusion Criteria	Individual has a diagnosis of severe hepatic impairment (Child- Pugh C)
Required Medical Information	
Age Restrictions	Individual is 18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Signifor LAR

Products Affected

• SIGNIFOR LAR

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Siklos

Products Affected

• SIKLOS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual has a diagnosis of sickle cell anemia with recurrent moderate to severe painful crises AND is unable to swallow the oral tablet dose form due to a clinical condition such as but not limited to the following: (a) Dysphagia or (b) Individual's age.
Indications	All Medically-accepted Indications.
Off Label Uses	

Siliq

Products Affected

• SILIQ

PA Criteria	Criteria Details
Exclusion Criteria	Use of brodalumab in combination with phototherapy OR Use of brodalumab in combination with IL-17 inhibitors or other biologic drugs OR Tuberculosis, other active serious infections, or a history of recurrent infections OR Individual has not had a tuberculin skin test (TST) or a Centers for Disease Control and Prevention (CDC)- recommended equivalent test to evaluate for latent tuberculosis prior to initiating brodalumab OR Individual has Crohn's disease.
Required Medical Information	Diagnosis of moderate to severe plaque psoriasis with either of the following: Patient has greater than 5% of body surface area with plaque psoriasis OR Less than or equal to 5% of body surface area with plaque psoriasis involving sensitive areas or areas that would significantly impact daily function (such as palms, soles of feet, head/neck, or genitalia).
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year.

PA Criteria	Criteria Details
Other Criteria	For a dx of chronic plaque psoriasis, individual has had an inadequate response to, is intolerant of, or has a medical contraindication to phototherapy or any ONE systemic therapy (for example acitretin, methotrexate, cyclosporine) AND individual has had a trial of and an inadequate response or is intolerant to: Humira(adalimumab) or Enbrel (etanercept) OR if the TNF agent (Enbrel(etanercept)/Humira(adalimumab)) tried and failed are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Siliq or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis OR individual has either concomitant clinical condition: (a) Demyelinating disease or (b) Heart failure with documented left ventricular dysfunction, Siliq may be allowed without trial of preferred TNF agents (Enbrel(etanercept)/ Humira(adalimumab)).
Indications	All Medically-accepted Indications.
Off Label Uses	

SIMPONI

Products Affected

 SIMPONI SUBCUTANEOUS SOLUTION AUTO-INJECTOR 100 MG/ML, 50 MG/0.5ML PREFILLED SYRINGE 100 MG/ML, 50 MG/0.5ML

SIMPONI SUBCUTANEOUS SOLUTION

PA Criteria	Criteria Details
Exclusion Criteria	Using golimumab in combination with other TNF antagonists, JAK inhibitors, or other biologic drugs (such as, abatacept, anakinra, or vedolizumab). Tuberculosis, other active serious infections, or a history of recurrent infections. Individuals who have not had a TST or a CDC-recommended equivalent to evaluate for latent tuberculosis prior to initiating Simponi (golimumab).
Required Medical Information	
Age Restrictions	Individual is 18 years or older
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For RA, individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE Conventional Therapy [non-biologic DMARD (such as methotrexate, sulfasalazine, leflunomide, or hydroxychloroquine)] (ACR 2015) AND individual has had a trial of and inadequate response or intolerance to: Humira (adalimumab) OR Enbrel (etanercept). For Psoriatic Arthritis, individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional therapy [nonbiologic DMARDs (such as methotrexate, sulfasalazine, or leflunomide)] AND individual has had a trial of and an inadequate response or intolerance to Humira(adalimumab) OR Enbrel(etanercept). For Ankylosing Spondylitis, had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional therapy (e.g. NSAIDs or nonbiologic DMARDs) AND individual has had a trial of and an inadequate response to: Humira(adalimumab) OR Enbrel(etanercept). For UC, individual has had a trial of and an inadequate response or intolerance to: Humira(adalimumab) OR Enbrel(etanercept). For UC, individual has had a trial of and an inadequate response or intolerante to: Humira(adalimumab) OR Enbrel(etanercept). For UC, individual has had a ninadequate response, is intolerant of, or has a medical contraindication to ONE conventional therapy (such as 6-mercaptopurine, azathioprine, oral aminosalicylates, or oral corticosteroids) AND individual has had a trial of and an inadequate response or is intolerant to Humira(adalimumab). For any of the above indications, if the TNF agent (Enbrel(etanercept)/Humira(adalimumab)) tried and failed are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Simponi or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis. Simponi may be allowed without trial of p
Indications	All Medically-accepted Indications.
Off Label Uses	

Simponi ARIA

Products Affected

• SIMPONI ARIA

PA Criteria	Criteria Details
Exclusion Criteria	Use in combination with other TNF antagonists, JAK inhibitors, or other biologic drugs (such as, abatacept, anakinra, or vedolizumab). Tuberculosis, other active serious infections, or a history of recurrent infections. Individuals who have not had a TST or a CDC-recommended equivalent to evaluate for latent tuberculosis prior to initiating Simponi (golimumab).
Required Medical Information	
Age Restrictions	Individual is 18 years or older
Prescriber Restrictions	
Coverage Duration	1 year.

PA Criteria	Criteria Details
Other Criteria	For RA, individual had an inadequate response to, is intolerant of, or has a medical contraindication to ONE Conventional Therapy [non-biologic DMARD (such as methotrexate, sulfasalazine, leflunomide, or hydroxychloroquine)] (ACR 2015) AND individual has had a trial of and an inadequate response or intolerance to: Humira (adalimumab) OR Enbrel(etanercept) OR the TNF agent (Enbrel(etanercept)/Humira(adalimumab)) tried and failed are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Simponi Aria or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis, Simponi Aria may be allowed without trial of preferred TNF agents (Enbrel(etanercept)/Humira(adalimumab)). For PsA, individual has had an inadequate response to, is intolerant of or has a medical contraindication to ONE conventional therapy [nonbiologic DMARDs (such as methotrexate, sulfasalazine, or leflunomide)] AND has had a trial of and an inadequate response or intolerance to: Humira (adalimumab) OR Enbrel(etanercept) OR the TNF agent (Enbrel(etanercept)/Humira(adalimumab)) tried and failed are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Simponi Aria or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis. For Ankylosing Spondylitis, individual has had an inadequate response to, is intolerance to Humira (adalimumab) OR Enbrel(etanercept) OR thes not also associated with Simponi Aria or (b) Individual has had an inadequate response to, is intolerance to Humira (adalimumab) OR Enbrel(etanercept) PMD individual has had an inadequate response to, is intolerance to response to respise. For Ankylosing Spondylitis, individual has had an ina
	limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Simponi Aria or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis.
Indications	All Medically-accepted Indications.

PA Criteria	Criteria Details
Off Label Uses	

Sirturo

Products Affected

• SIRTURO

PA Criteria	Criteria Details
Exclusion Criteria	Latent infection due to Mycobacterium tuberculosis OR Drug- sensitive tuberculosis OR Extra-pulmonary tuberculosis OR Infections caused by non-tuberculosis mycobacteria.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a diagnosis of pulmonary multi-drug resistant tuberculosis AND is unable to use an effective regimen for treatment AND the individual is using Sirturo (bedaquiline) with at least 3 drugs to which the multi-drug resistant tuberculosis isolate is susceptible in vitro OR with at least 4 drugs to which the multi- drug resistant tuberculosis isolate is likely to be susceptible if in vitro testing results are unavailable.
Indications	All Medically-accepted Indications.
Off Label Uses	

Sivextro

Products Affected

- SIVEXTRO INTRAVENOUS
- SIVEXTRO ORAL

PA Criteria	Criteria Details
Exclusion Criteria	Treatment of gram-negative infections.
Required Medical Information	Individual has been diagnosed with acute bacterial skin and skin structure infection (ABSSSI) defined as one of the following (FDA, 2013): Cellulitis/erysipelas OR Wound infection OR Major cutaneous abscess. AND Individual has at least 1 regional or 1 systemic sign of infection as defined by: Lymphadenopathy OR temperature greater than or equal to 38 degrees Celsius OR White blood cell count greater than or equal to 10,000 per microliter OR White blood cell count less than 4000 per microliter OR Greater than 10% of immature neutrophils.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	30 days
Other Criteria	Individual has had a trial and inadequate response or intolerance to of or has contraindications to an alternative antibiotic that the organism is susceptible to (depending on manifestation, severity of infection and culture or local sensitivity patterns, examples of alternative antibiotics may include, but are not limited to: TMP/SMX, doxycycline, vancomycin, daptomycin, televancin, clindamycin) (IDSA 2014) OR Individual started treatment with intravenous antibiotic(s) in the hospital and requires continued outpatient therapy for an organism susceptible to Sivextro (tedizolid).

PA Criteria	Criteria Details
Indications	All Medically-accepted Indications.
Off Label Uses	

Skyrizi

Products Affected

• SKYRIZI (150 MG DOSE)

PA Criteria	Criteria Details
Exclusion Criteria	Use in combination with other biologic drugs or phototherapy OR Tuberculosis, other active serious infections, or a history of recurrent infections OR Individual has not had a tuberculin skin test (TST) or a Centers for Disease Control (CDC-) and Prevention - recommended equivalent to evaluate for latent tuberculosis prior to initiating risankizumab-rzaa.
Required Medical Information	Dx of chronic moderate to severe (that is, extensive or disabling) plaque Ps with either of the following (AAD 2011): 1. Plaque Ps involving greater than five percent (5%) body surface area (BSA) OR 2. Plaque Ps involving less than or equal to five percent (5%) BSA involving sensitive areas or areas that significantly impact daily function (such as palms, soles of feet, head/neck, or genitalia).
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year.

PA Criteria	Criteria Details
Other Criteria	For dx of chronic moderate to severe plaque Ps, individual has had an inadequate response to, is intolerant of, or has a contraindication to phototherapy or other systemic therapy (such as acitretin, cyclosporine, or methotrexate) AND Individual has had a trial and inadequate response or intolerance to Enbrel (etanercept) OR Humira (adalimumab) OR if the agents tried and failed (Humira (adalimumab)/Enbrel (etanercept)) are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Skyrizi (risankizumab-rzaa) or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis OR individual has either concomitant clinical condition: (a) Demyelinating disease or (b) Heart failure with documented left ventricular dysfunction.
Indications	All Medically-accepted Indications.
Off Label Uses	

Solaraze

Products Affected

• diclofenac sodium transdermal gel 3 %

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Dx of Actinic Keratosis
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Soliris

Products Affected

• SOLIRIS

PA Criteria	Criteria Details
Exclusion Criteria	

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Required Medical Information	For initial tx of Paroxysmal nocturnal hemoglobinuria (PNH) as documented by flow cytometry, including the presence: (1.) Paroxysmal nocturnal hemoglobinuria type III red cell clone or a measurable granulocyte or monocyte clone OR (2.) Glycosylphosphatidylinositol-anchored proteins (GPI-AP)-deficient polymorphonuclear cells (PMNs) AND Individual has been immunized with a meningococcal vaccine at least 2 weeks prior to administration of the first dose of Soliris (eculizumab) (unless the clinical record documents that the risks of delaying Soliris (eculizumab) outweigh the risk of meningococcal infection) AND There is NO evidence of an active meningococcal infection AND Individual has (a)Hemoglobin that is less than or equal to 7 g/dl, or the individual has symptoms of anemia and the hemoglobin is less than or equal to 9 g/dl or (b)Lactate dehydrogenase is greater than 1.5 times the upper limit of normal. OR (c) Documented history of a major adverse vascular event from thromboembolism. For the initial tx of atypical hemolytic uremic syndrome (aHUS) when the following criteria are met: (A) dx of aHUS is supported by the absence of Shiga toxin-producing E. coli infection AND (B) Thrombotic thrombocytopenic purpura has been ruled out (for example, normal ADAMTS 13 activity and no evidence of an ADAMTS 13 inhibitor), or if thrombotic thrombocytopenic purpura cannot be ruled out by laboratory and clinical evaluation, a trial of plasma exchange did not result in clinical improvement AND (C) Individual has been immunized with a meningococcal vaccine at least 2 weeks prior to administration of the first dose of Soliris (eculizumab) (unless the clinical record documents that the risks of delaying Soliris (eculizumab) outweigh the risk of meningococcal infection.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	For PNH 1yr. For aHUS Initial 3 mon. For MG Initial 7 mon. Continuation is 1 year for all dx.

PA Criteria	Criteria Details
Other Criteria	For initial tx of generalized myasthenia gravis (MG) when the following criteria are met: (A) Individual has Myasthenia Gravis Foundation of America Clinical Classification Class II to IV disease AND (B) has a documented positive serologic test for binding anti- acetylcholine receptor antibodies (AChR-ab) AND (C) has had an inadequate response to, is intolerant of, or has a medical contraindication to two or more immunosuppressive drug agents (such as, azathioprine, cyclosporine, or methotrexate) as monotherapy or in combination therapy for greater than or equal to 12 months OR (D) has had an inadequate response to, is intolerant of, or has a medical contraindication to one or more immunosuppressive drug agents as monotherapy or in combination therapy and requires chronic plasma exchange or plasmapheresis or intravenous immunoglobulin therapy AND (E)has been immunized with a meningococcal vaccine at least 2 weeks prior to administration of the first dose of eculizumab (unless the clinical record documents that the risk of delaying eculizumab outweigh the risk of meningococcal infection. For Continuation following initial tx of aHUS may be approved when the following is met: There is clinical improvement after the initial trial (for example, increased platelet count or laboratory evidence of reduced hemolysis) until an individual becomes a candidate for physician directed cessation as evidenced by the following: (a) Complete clinical remission has been achieved (that is, resolution of thrombocytopenia and mechanical hemolysis, and normalization or new baseline plateau of renal function) and improvement of precipitating illness is clinically apparent AND (b) Duration of clinical remission has been stable for 2 months. Resumption of Soliris (eculizumab) in aHUS may be approved if the following criteria are met (Fakhouri 2017): (A) Individual experienced a relapse after discontinuation of therapy as defined by: (1) Reduction in platelet count to less than 150,000/mm3 or greater

PA Criteria	Criteria Details
	than 25% from baseline OR (2) Mechanical hemolysis (having 2 or more features of hemoglobin less than 10 g/dL, lactate dehydrogenase greater than 2 times upper limit of normal, undetectable haptoglobin, or presence of schistocytes on smear) OR (3) Acute kidney injury with serum creatinine increase greater than 15% from baseline levels. For Continuation for the tx of gMG may be approved when Individual has experienced a clinical response (that is, a reduction in signs or symptoms that impact daily function).
Indications	All Medically-accepted Indications.
Off Label Uses	

Somatuline Depot

Products Affected

• SOMATULINE DEPOT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Somavert

Products Affected

• SOMAVERT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Dx of acromegaly AND member has had an inadequate response to surgery and/or radiation OR Surgery and/or radiation therapy are not an option.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Sovaldi

Products Affected

• SOVALDI

PA Criteria	Criteria Details
Exclusion Criteria	Individual has severe renal impairment (CrCl less than 30 mL/min), end stage renal disease, or requires dialysis (AASLD/IDSA 2014). Individual is using in combination with Daklinza (daclatasvir) and a known NS5A polymorphism is present.
Required Medical Information	Documentation is provided for a diagnosis of chronic hepatitis C (CHC) infection, which includes genotype and a positive HCV RNA result (AASLD/IDSA 2017, CDC 2013) AND Individual has received baseline evaluation for liver fibrosis to guide appropriate therapy AND Individual does not have a short life expectancy (less than 12 months owing to non-liver related comorbid conditions) that cannot be remediated by treating HCV, by transplantation or other directed therapy (AASLD/IDSA 2016).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Criteria will be applied consistent with current AASLD/IDSA guidance.

PA Criteria	Criteria Details
Other Criteria	Criteria will be applied consistent with current AASLD/IDSA guidance. For GT 1, Individual has had a prior trial and inadequate response to Harvoni(sofosbuvir/ledipasvir) OR individual is currently on and completing a course of therapy with the requested regimen OR Individual has a documented hypersensitivity, as manifested by a severe allergic reaction, to any ingredient in Harvoni (ledipasvir/sofosbuvir) which is not also in Sovaldi OR Individual is concurrently using an agent that cannot be substituted with another agent or temporarily discontinued and is contraindicated or not recommended for concomitant use with Harvoni(sofosbuvir/ledipasvir). For GT 4, individual has had a prior trial and inadequate response to Harvoni(sofosbuvir/ledipasvir) or Epclusa(sofosbuvir/velpatasvir). OR individual is currently on and completing a course of therapy with the requested regimen OR Individual has a documented hypersensitivity, as manifested by a severe allergic reaction, to any ingredient in Harvoni(sofosbuvir/ledipasvir) or Epclusa (sofosbuvir/velpatasvir) which is not also in Sovaldi OR Individual is concurrently using an agent that cannot be substituted with another agent or temporarily discontinued and is contraindicated or not recommended for concomitant use with Harvoni (sofosbuvir/ledipasvir) or Epclusa(sofosbuvir/velpatasvir).
Indications	All Medically-accepted Indications.
Off Label Uses	

Spinraza

Products Affected

• SPINRAZA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For initial requests, Individual has documentation (written or verbal attestation is acceptable) of a confirmatory diagnosis by either: (1) Spinal Muscular Atrophy (SMA) diagnostic test results confirming 0 copies of SMN1 OR (2) Molecular genetic testing of 5q SMA for any of the following: (a) homozygous gene deletion or (b) homozygous conversion mutation or (c) compound heterozygote. AND Individual has documentation (written or verbal attestation is acceptable) of either: (1) Genetic testing confirming no more than 2 copies of SMN2 (Finkel 2017) OR (2) Onset of SMA-associated signs and symptoms before 21 months of age (Mercuri 2018). For individuals using Spinraza following treatment with Zolgensma (onasemnogene abeparvovec-xioi), mbr meets the above criteria AND (1) Individual does not require use of invasive ventilatory support (tracheotomy with positive pressure) or use of non-invasive ventilator support (BiPAP) for more than 16 hours per day as a result of advanced SMA disease AND (2) Individual has not achieved the expected benefit from gene therapy, as shown by the following: (a) within 3 months of gene therapy, individual has not achieved and sustained a CHOP INTEND score of more than 40 points.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	6 Months.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Other Criteria	For continuation requests, meets initial criteria AND Individual has documentation (written or verbal) of clinically significant improvement in spinal muscular atrophy-associated signs and symptoms (i.e., progression, stabilization, or decreased decline in motor function) compared to the predicted natural history trajectory of disease.
Indications	All Medically-accepted Indications.
Off Label Uses	

Spravato

Products Affected

- SPRAVATO (56 MG DOSE)
- SPRAVATO (84 MG DOSE)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	Initial 3 months, continuation 1 year.
Other Criteria	For initial use, individual has been diagnosed with moderate to severe major depressive disorder AND had an inadequate response to the maximum tolerated dose of two antidepressant therapies during the current major depressive episode (MDE) as defined by less than 50% reduction in symptom severity using a standard rating scale that reliably measures depressive symptoms AND will use Spravato in addition to antidepressant therapy. For continuation, individual has had at least a 50% reduction in symptoms of depression compared to baseline using a standard rating scale that reliably measures depressive symptoms AND will use Spravato in addition to antidepressant therapy.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Spritam

Products Affected

 SPRITAM ORAL TABLET DISINTEGRATING SOLUBLE 1000 MG, 250 MG, 500 MG, 750 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has a diagnosis of partial onset seizures OR primary generalized tonic-clonic seizures AND weighs more than 20 kg.
Age Restrictions	Partial onset seizures: 4 years old. Juvenile myoclonic epilepsy: 12 years old. Primary generalized tonic-clonic seizures: 6 years old.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Sprycel

Products Affected

• SPRYCEL

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Written or verbal attestation is provided for confirmation of mutations where applicable based on use/diagnosis.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Stelara

Products Affected

- STELARA INTRAVENOUS
- STELARA SUBCUTANEOUS

PA Criteria	Criteria Details
Exclusion Criteria	Known to have reversible posterior leukoencephalopathy syndrome (RPLS) while on tx with Stelara. Individual has not had a tuberculin skin test (TST) or a Centers for Disease Control (CDC) and Prevention-recommended equivalent test to evaluate for latent tuberculosis prior to initiating Stelara (ustekinumab). Individual has tuberculosis, other active serious infections or a history of recurrent infections. Using ustekinumab in combination with phototherapy. In combination with JAK inhibitors or other biologic drugs (such as TNF antagonists).
Required Medical Information	Diagnosis of moderate to severe plaque psoriasis with either of the following: Patient has greater than 5% of body surface area with plaque psoriasis OR Less than or equal to 5% of body surface area with plaque psoriasis involving sensitive areas or areas that would significantly impact daily function (such as palms, soles of feet, head/neck, or genitalia).
Age Restrictions	Individual is 18 years of age or older. For Plaque Psoriasis, age 6 and older.
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For chronic plaque psoriasis, individual has had an inadequate response to, is intolerant of, or has a medical contraindication to phototherapy or any ONE systemic therapy (for example acitretin, methotrexate, cyclosporine) AND individual has had a trial of and an inadequate response or is intolerant to Humira (adalimumab) or Enbrel (etanercept). For psoriatic arthritis, individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional therapy ([nonbiologic DMARDS] such as methotrexate, sulfasalazine, or leflunomide) AND individual has had a trial of and an inadequate response or is intolerant to Humira (adalimumab) or Enbrel (etanercept). For Crohns disease, individual has had an inadequate response or is intolerant to Humira (adalimumab) or Enbrel (etanercept). For Crohns disease, individual has had an inadequate response to, has lost response to, is intolerant of, or has a medical contraindication to conventional therapy (such as 5-Aminosalicylic acid products, systemic corticosteroids, or immunosuppressants) or a tumor necrosis factor (TNF) antagonist AND individual has had a trial and inadequate response or intolerance to Humira (adalimumab). For any of the above indications, if the TNF agent (Enbrel(etanercept)/Humira(adalimumab)) tried and failed are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Stelara or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis OR individual has either concomitant clinical condition: (a) Demyelinating disease or (b) Heart failure with documented left ventricular dysfunction, Stelara may be allowed without trial of preferred TNF agents (Enbrel(etanercept)/Humira(adalimumab)). For Crohns, if the TNF agent tried and failed are not acceptable due to additional concomitant clinical conditions including: (c) Malig
Indications	All Medically-accepted Indications.
Off Label Uses	

Stivarga

Products Affected

• STIVARGA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Strensiq

Products Affected

• STRENSIQ

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Total serum alkaline phosphatase is below the lower limit of normal for the individual's age and gender at diagnosis and Plasma pyridoxal 5'-phosphate levels are greater than the upper limit of normal at the time of diagnosis.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Initial 3 mon. Continuation 1 year
Other Criteria	For initial treatment of perinatal/infantile or juvenile onset hypophosphatasia, individual has onset of symptoms occurred prior to 6 months of age and has one or more of the following: (a) Radiographic evidence of poor bone mineralization such as flared and frayed metaphyses, severe/ generalized osteopenia,or widened growth plates or (b) Genetic test results that confirm infantile HPP or (c) one or more of the following: (1) History or presence of nontraumatic postnatal fracture healing or (2) History of elevated serum calcium or (3) Functional craniosynostosis with decreased head circumference growth or (4) Nephrocalcinosis or (5) Rachitic chest deformity or (6) Respiratory compromise or (7) Vitamin B6- responsive seizures or (8) Failure to thrive. For Continuation of Therapy: The individual has demonstrated clinical improvement in symptoms following asfotase alfa therapy.
Indications	All Medically-accepted Indications.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Off Label Uses	

Subsys

Products Affected

• SUBSYS

PA Criteria	Criteria Details
Exclusion Criteria	Using for the treatment of acute or postoperative pain. Using for treatment of migraine headache pain. Using for non-cancer related breakthrough pain.
Required Medical Information	
Age Restrictions	Individual is 18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a diagnosis of active cancer (such as, metastatic or locally invasive cancer for which the individual is currently seeking treatment) with breakthrough cancer pain (provide diagnosis) AND has tried and failed fentanyl citrate lozenge (generic Actiq) AND Individual is already receiving opioid therapy and is TOLERANT to opioid therapy as defined as receiving around the clock medicine consisting of one of the following: At least 60mg morphine per day, OR At least 25mcg/hr transdermal fentanyl/hour, OR At least 30mg of oxycodone daily, OR At least 8mg of oral hydromorphone daily, OR At least 25mg of oral oxymorphone daily, OR An equianalgesic dose of another opioid for a week or longer. Individual will also continue around the clock opioids when taking Subsys (fentanyl) for cancer related breakthrough pain.
Indications	All Medically-accepted Indications.
Off Label Uses	
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Y0114_20_114077_I_012

Supprelin LA

Products Affected

• SUPPRELIN LA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has precocious puberty defined as: Beginning of secondary sexual characteristics before age 8 in girls and age 9 in boys.
Indications	All Medically-accepted Indications.
Off Label Uses	

Sutent

Products Affected

 SUTENT ORAL CAPSULE 12.5 MG, 25 MG, 37.5 MG, 50 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Sylatron

Products Affected

• SYLATRON

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Member is being treated for melanoma with microscopic or gross nodal involvement AND Treatment is initiated within 84 days after definitive surgical resection.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Sylvant

Products Affected

• SYLVANT

PA Criteria	Criteria Details
Exclusion Criteria	Individual does not have a concurrent clinically significant infection (for example, Hepatitis B or C) AND No concurrent lymphoma.
Required Medical Information	Individual is negative for human immunodeficiency virus (HIV) and human herpesvirus-8 (HHV-8).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Symdeko

Products Affected

• SYMDEKO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	6 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual has a diagnosis of cystic fibrosis (CF) AND has a confirmed mutation-positive result in the cystic fibrosis transmembrane conductance regulator (CFTR) gene and the mutation type is provided and responsive to Symdeko.
Indications	All Medically-accepted Indications.
Off Label Uses	

Symlin

Products Affected

- SYMLINPEN 120
- SYMLINPEN 60

PA Criteria	Criteria Details
Exclusion Criteria	May not be approved if individual has any of the following: receiving drugs that stimulate gastric motility (i.e. metoclopramide), diagnosis of severe gastroparesis, hypoglycemia unawareness or recent hypoglycemia requiring assistance within past 6 months
Required Medical Information	Type 1 or type 2 diabetes AND taking mealtime insulin therapy AND failed to achieve glucose control AND HBA1C is less than or equal to 9.
Age Restrictions	18 or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Synagis

Products Affected

• SYNAGIS

PA Criteria	Criteria Details
Exclusion Criteria	Administration of more than 5 doses of palivizumab in one RSV season. Children who reach 24 months of age prior to the commencement of the RSV season. Treatment in children or infants with known RSV disease. Continued RSV immunoprophylaxis for children who experience breakthrough RSV hospitalization. Primary asthma prevention or to reduce subsequent episodes of wheezing. Children with surgically corrected congenital heart disease or hemodynamically insignificant heart disease (including secundum atrial septal defect, small ventricular septal defect, uncomplicated pulmonic stenosis, uncomplicated aortic stenosis, mild coarctation of the aorta, and patent ductus arteriosus) who do not otherwise meet approval criteria. Children with Down syndrome who do not otherwise meet approval criteria.

PA Criteria	Criteria Details
Required Medical Information	Individual is using when the following are met: A) Maximum of Five (5) doses of palivizumab for infants during the first year of life: Born before 29 weeks 0 days gestation (up to and including 28 weeks, 6 days) and younger than 12 months of age at the start of the RSV season OR Chronic lung disease (CLD) of prematurity (defined as birth at less than 32 weeks, 0 days gestation and a requirement for greater than 21% oxygen for at least 28 days after birth) OR Hemodynamically significant congenital heart disease (CHD) (for example, but not limited to, infants with acyanotic heart disease who are receiving medication to control congestive heart failure and will require cardiac surgical procedures and infants with moderate to severe pulmonary hypertension OR infants with anatomic pulmonary abnormalities (i.e., tracheal ring) or a neuromuscular condition that impairs the ability to clear secretions from the upper airway because of ineffective cough OR Cystic fibrosis with clinical evidence of chronic lung disease or nutritional compromise (weight for length less than tenth percentile). B) Maximum of five (5) doses of palivizumab for children younger than 24 months of age with any of the following: Profoundly immunocompromised, such as severe combined immunodeficiency, advanced acquired immunodeficiency syndrome undergoing organ or hematopoietic stem cell transplant, or an absolute lymphocyte count of less than 100 cell/mm3 OR undergoing cardiac transplantation.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	5 months

PA Criteria	Criteria Details
Other Criteria	C) An additional dose of palivizumab may be allowed for children younger than 24 months of age who have approval for a course of treatment and who undergo cardiopulmonary bypass for surgical procedures. D) A maximum of 5 doses of palivizumab prophylaxis may be approved for children in the second year of life with any of the following: (i) for preterm infants born at less than 32 weeks, 0 days gestation who required at least 28 days of oxygen after birth and who continue to require medical intervention within 6 months of the start of the second RSV season (including, supplemental oxygen, chronic systemic corticosteroid therapy, or diuretics) or (ii) Cystic fibrosis with severe lung disease (history of hospitalization, abnormal chest x-ray or CT scan) or weight for length less than tenth percentile.
Indications	All Medically-accepted Indications.
Off Label Uses	

Synarel Nasal Solution

Products Affected

• SYNAREL

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Precocious puberty, defined as the beginning of secondary sexual maturation characteristics before age 8 in girls and before age 9 in boys.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Endometriosis: 6 months, all other indications: 1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Synribo

Products Affected

• SYNRIBO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Tabrecta

Products Affected

• TABRECTA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has confirmation (written or verbal) of mesenchymal- epithelial transition (MET) exon 14 skipping positive tumors as detected by an FDA-approved test AND individual has not received treatment with another MET exon 14 skipping-targeted agent, such as crizotinib.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual is using Tabrecta (capmatinib) as monotherapy
Indications	All Medically-accepted Indications.
Off Label Uses	

Tafamidis Agents

- VYNDAMAX
- VYNDAQEL

PA Criteria	Criteria Details
Exclusion Criteria	Individual has a history of liver or heart transplantation.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual has a diagnosis of wild type or hereditary transthyretin amyloid cardiomyopathy confirmed by biopsy and DNA mutation analysis (Bozkurt, 2016, Maurer, 2018) AND is using for the treatment of New York Heart Association class I, II or III heart failure symptoms (Maurer, 2018).
Indications	All Medically-accepted Indications.
Off Label Uses	

Tafinlar

Products Affected

• TAFINLAR

PA Criteria	Criteria Details
Exclusion Criteria	Tafinlar may not be approved for the treatment of individuals with wild type BRAF melanoma.
Required Medical Information	BRAF V600E or V600K mutation results must be confirmed (verbal or written).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 Year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Tagrisso

Products Affected

• TAGRISSO ORAL TABLET 40 MG, 80 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has either: (a) EGFR (epidermal growth factor receptor) T790M mutation is confirmed (verbal or written) OR (b) EGFR exon 19 deletions or exon 21 L858R mutations is confirmed (verbal or written)
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Takhzyro

Products Affected

• TAKHZYRO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	12 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	1.Using for prophylaxis against acute attacks of hereditary angioedema (HAE) for either: of the following: a. Short term prophylaxis prior to surgery, dental procedures or intubation OR b. Long term prophylaxis and individual has failed, or is intolerant to, or has contraindication (such as pregnant, or breastfeeding) to 17 alpha-alkylated androgens (e.g., danazol) or antifibrinolytic agents (e.g., aminocaproic acid) AND 2.Diagnosis is confirmed by C4 level below the lower limit of normal as defined by lab test and any of the following: a. C1 inhibitor antigenic level below the lower limit or normal as defined by lab test OR b. C1 inhibitor functional level below the lower limit of normal as defined by lab test OR c. Presence of a known HAE causing C1-INH mutation AND 3. Individual has a history of moderate or severe attacks (such as airway swelling, severe abdominal pain, facial swelling, nausea and vomiting, painful facial distortion).
Indications	All Medically-accepted Indications.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

Taltz

Products Affected

• TALTZ

PA Criteria	Criteria Details
Exclusion Criteria	Use of Taltz (ixekizumab) in combination with phototherapy. Use of Taltz (ixekizumab) in combination with IL-17 inhibitors or other biologic drugs. Individual with Tuberculosis, other active serious infections, or a history of recurrent infections. Individual has not had a tuberculin skin test (TST) or a Centers for Disease Control (CDC)-recommended equivalent test to evaluate for latent tuberculosis prior to initiating Taltz (ixekizumab).
Required Medical Information	Diagnosis of moderate to severe plaque psoriasis with either of the following: Individual has greater than 5% of body surface area with plaque psoriasis OR Less than or equal to 5% of body surface area with plaque psoriasis involving sensitive areas or areas that would significantly impact daily function (such as palms, soles of feet, head/neck, or genitalia).
Age Restrictions	Individual is 6 years of age or older for plaque psoriasis, 18 years of age or older for other indications
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For a dx of moderate to severe psoriatic arthritis, the individual has had an inadequate response to, is intolerant of, or has a medical contraindication to conventional drug therapy [nonbiologic DMARDS (such as methotrexate, sulfasalazine, or leflunomide)] or TNF Antagonists (AAD 2011) AND individual has tried and failed: Humira(adalimumab) OR Enbrel(etanercept). For a dx of moderate to severe plaque psoriasis, had an inadequate response, is intolerant of, or has a medical contraindication to phototherapy or any ONE systemic therapy (for example acitretin, methotrexate, cyclosporine) AND individual has tried and failed: Humira(adalimumab)OR Enbrel(etanercept). For either of the above indications, if the TNF agent (Humira/Enbrel) tried and failed are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Taltz (ixekizumab) or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis OR The individual has either concomitant clinical condition: (a) Demyelinating disease or (b) Heart failure with documented left ventricular dysfunction.
Indications	All Medically-accepted Indications.
Off Label Uses	

Talzenna

Products Affected

 TALZENNA ORAL CAPSULE 0.25 MG, 1 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Written or verbal attestation provided to confirm deleterious or suspected deleterious germline BRCA-mutation (gBRCAm) and human epidermal growth factor receptor 2-negative (HER2) breast cancer.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Tarceva

- erlotinib hcl oral tablet 100 mg, 150 mg, 25 mg
- TARCEVA ORAL TABLET 100 MG, 150 MG, 25 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For NSCLC tumors that have EGFR exon 19 deletions or exon 21 (L858R) substitution mutations, a copy of the test results must be provided
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Targretin

Products Affected

- bexarotene
- TARGRETIN EXTERNAL
- TARGRETIN ORAL

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Tasigna

Products Affected

• TASIGNA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Tasmar

Products Affected

- TASMAR
- tolcapone

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Tavalisse

Products Affected

• TAVALISSE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has a platelet count of less than 50 X 109/L
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Tazorac

- tazarotene
- TAZORAC

DA Cuitouia	Criteria Detaile
PA Criteria	Criteria Details
Exclusion Criteria	May not be approved for cosmetic purposes such as, but not limited to the following: Photoaging, Wrinkles, Hyperpigmentation, Sun damage, or Melasma.
Required Medical Information	For psoriasis, individual has up to 20% of body surface area involvement.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For psoriasis, individual has had a prior trial and inadequate response to either of the following (AAD 2009): Any two (2) topical corticosteroids or any one (1) topical corticosteroids plus calcipotriene. For psoriasis use greater than 1 year, efficacy must be documented for continued use. Documentation may include chart notes, consultation notes.
Indications	All Medically-accepted Indications.
Off Label Uses	

Tazverik

Products Affected

• TAZVERIK

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For Epithelioid Sarcoma, individual has a histologically confirmed (written or verbal) diagnosis and has a current ECOG performance status of 0-2. For follicular lymphoma, tumor is positive for EZH2 mutation as detected by an FDA approved test.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Tecentriq

Products Affected

 TECENTRIQ INTRAVENOUS SOLUTION 1200 MG/20ML, 840 MG/14ML

PA Criteria	Criteria Details
Exclusion Criteria	Individual has received treatment with another PD-1 agent or PD- L1 (for example, nivolumab or pembrolizumab) and Individual is receiving therapy for an autoimmune disease or chronic condition requiring treatment with a systemic immunosuppressant.
Required Medical Information	Written or verbal attestation is provided for confirmation of mutations where applicable based on use/diagnosis.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For locally advanced or metastatic urothelial carcinoma, has a current Eastern Cooperative Oncology Group (ECOG) performance status of 0-2. For metastatic non-small cell lung cancer (NSCLC), extensive-stage small cell lung cancer has a current Eastern Cooperative Oncology Group (ECOG) performance status of 0-2.
Indications	All Medically-accepted Indications.
Off Label Uses	

Tecfidera

- dimethyl fumarate
- *dimethyl fumarate starter pack*
- TECFIDERA

PA Criteria	Criteria Details
Exclusion Criteria	Using in combination with other immunomodulatory products (such as Aubagio, Gilenya, Tysabri, Copaxone/Glatopa, Extavia, Rebif, Avonex, Plegridy, Lemtrada, Ocrevus or Betaseron).
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Tegsedi

Products Affected

• TEGSEDI

PA Criteria	Criteria Details
Exclusion Criteria	Individual has a history of acute glomerulonephritis caused by Tegsedi (inotersen)
Required Medical Information	Individual has a baseline platelet count greater than or equal to 100×10 9/L AND urinary protein to creatinine ratio (UPCR) less than 1000 mg/g AND Individual has a TTR mutation confirmed by genotyping.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has diagnosis of hereditary transthyretin (hATTR) amyloidosis or familial amyloid polyneuropathy (FAP) AND associated mild to moderate polyneuropathy.
Indications	All Medically-accepted Indications.
Off Label Uses	

Testosterone Inj

- AVEED
- DEPO-TESTOSTERONE
- TESTONE CIK
- testosterone cypionate

- testosterone enanthate
- testosterone propionate
- XYOSTED

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For initial use of replacement therapy, Individual has a dx of (1) primary hypogonadism (congenital or acquired) such as but not limited to: Cryptorchidism OR Bilateral torsion OR Vanishing testis syndrome OR orchitis OR Orchiectomy OR Klinefelter Syndrome OR Chemotherapy OR Toxic damage from alcohol or heavy metals OR idiopathic primary hypogonadism. Or (2) Hypogonadotropic hypogonadism (congenital or acquired), such as but not limited to: Idiopathic luteinizing hormone-releasing hormone (LHRH deficiency) OR Pituitary-hypothalamic injury AND Individual presents with symptoms associated with hypogonadism, such as but not limited to, at least one of the following: (a) Reduced sexual desire (libido) and activity or (b) Decreased spontaneous erections or (c) Breast discomfort/gynecomastia or (d) Loss of body (axillary and pubic) hair, reduced need for shaving or (e) Very small (especially less than 5 mL) or shrinking testes or (f) Inability to father children or low/zero sperm count or (g) Height loss, low trauma fracture, low bone mineral density or (h)Hot flushes, sweats or (i) Other less specific signs and symptoms including decreased energy, depressed mood/dysthymia, irritability, sleep disturbance, poor concentration/memory, diminished physical or work performance. For Continuation of Testosterone Inj agents for replacement therapy, (a) Individual met all diagnostic criteria for initial therapy and (b) Individual has had serum testosterone level measured in the previous 180 days and (c) Individual has obtained clinical benefits as noted by symptom improvement. For treatment of delayed puberty when ALL the criteria below are met: Individual is using secondarily for advanced inoperable metastatic (skeletal) breast cancer OR Premenopausa female who has benefited from oophorectomy and is considered to have a hormone responsive tumor. For treatment of HIV-infected male adults with low
	testosterone and HIV-associated weight loss and wasting. For transgender individuals who meet ALL the following criteria: Individual has a diagnosis of gender dysphoria or gender identity disorder and goal of treatment is female-to-male gender reassignment.
Indications	All Medically-accepted Indications.
Y0114_20_114077	I_012 December 2020

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

Thalomid

Products Affected

 THALOMID ORAL CAPSULE 100 MG, 150 MG, 200 MG, 50 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Thiola

Products Affected

- THIOLA
- THIOLA EC

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	Individual is 9 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a diagnosis of severe homozygous cystinuria AND Individual has urinary cystine greater than 500 mg/day.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Tibsovo

Products Affected

TIBSOVO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Documented susceptible isocitrate dehydrogenase-1 (IDH1) (written or verbal attestation is acceptable)
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Topical Acne Antibiotic

- clindamycin-tretinoin
- VELTIN
- ZIANA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For dx of Acne, Individual has had a prior trial and inadequate response to the following: (1) One preferred generic topical tretinoin agent AND (2) One preferred generic erythromycin/benzoyl peroxide combination agent OR (3) One preferred generic clindamycin/benzoyl peroxide combination agent.
Indications	All Medically-accepted Indications.
Off Label Uses	

Topical Androgens

Products Affected

- ANDROGEL PUMP
- ANDROGEL TRANSDERMAL GEL 20.25 MG/1.25GM (1.62%), 40.5 MG/2.5GM (1.62%)
- testosterone transdermal gel 1.62 %, 20.25 mg/1.25gm (1.62%), 20.25 mg/act (1.62%), 40.5 mg/2.5gm (1.62%)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	18 years of age or older. For transgender use, individual is 16 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a dx of (1) primary hypogonadism (congenital or acquired) [for example, Cryptorchidism OR Bilateral torsion OR orchitis OR Vanishing testis syndrome OR Orchiectomy OR Klinefelter Syndrome OR Chemotherapy OR Toxic damage from alcohol or heavy metals OR idiopathic primary hypogonadism]. Or (2) Hypogonadotropic hypogonadism (congenital or acquired) [for example, Idiopathic luteinizing hormone-releasing hormone (LHRH deficiency), OR Pituitary-hypothalamic injury.] Individual is transgender AND is using for a diagnosis of gender Dysphoria or gender Identity Disorder AND Goal of treatment is female-to-male gender reassignment.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Topical Onychomycosis

- JUBLIA
- KERYDIN
- tavaborole

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has a confirmed fungal infection (i.e. physical exam).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has had a trial of and inadequate response or intolerance to oral itraconazole and terbinafine. Or has a, contraindication, drug interaction or concomitant clinical condition (such as but not limited history of liver disease or concerns over hepatotoxicity, history of CHF) which make use of oral itraconazole and terbinafine unacceptable OR Individual has used requested medication within the previous 6 months.
Indications	All Medically-accepted Indications.
Off Label Uses	

Topical Tretinoin Agents

- ALTRENO
- ATRALIN
- avita
- REFISSA
- RETIN-A
- RETIN-A MICRO

- RETIN-A MICRO PUMP
- tretinoin (emollient)
- tretinoin external
- tretinoin microsphere
- tretinoin microsphere pump

PA Criteria	Criteria Details
Exclusion Criteria	Topical tretinoin agents may not be approved for cosmetic purposes such as, but not limited to the following: Photoaging, Wrinkles, Hyperpigmentation, Sun damage, Melasma.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Torisel

Products Affected

- temsirolimus
- TORISEL

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For advanced renal cell carcinoma, individual is using for either of the following (A or B): (A) As first-line therapy as a single agent (monotherapy) for (either i or ii): (i) Relapsed metastatic disease or (ii) Surgically unresectable stage IV renal carcinoma in individuals with a poor prognosis as manifested by having at least 3 of the following (1 through 6): 1. Lactate dehydrogenase greater than 1.5 times the upper limit of normal or 2. Hemoglobin less than the lower limit of normal or 3. Corrected calcium level greater than 10mg/dL (2.5mmol/liter) or 4. Interval of less than a year from original diagnosis to the start of systemic therapy or 5. Karnofsky performance status less than or equal to 70 or ECOG performance score of 2 - 4 or 6. Greater than or equal to 2 sites of metastases. OR (B) For subsequent (second-line) therapy as a single agent (monotherapy) for relapsed metastatic or for surgically unresectable stage IV disease.
Indications	All Medically-accepted Indications.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

Tracleer

- bosentan
- TRACLEER ORAL TABLET
- TRACLEER ORAL TABLET SOLUBLE

PA Criteria	Criteria Details
Exclusion Criteria	Individual is concomitantly taking cyclosporine A or glyburide. Individual has a diagnosis of moderate (Child-Pugh Class B) or severe (Child-Pugh Class C) hepatic impairment or in the treatment of congestive heart failure with left ventricular dysfunction. Or Individual is initiating therapy and has elevated [greater than 3 times the upper limit of normal (ULN)] baseline aminotransferase levels OR In combination with other endothelin receptor antagonist (ERA) agents, such as but not limited to Letairis (ambrisentan) or Opsumit (macitentan).
Required Medical Information	Catheterization-proven diagnosis of Pulmonary Arterial Hypertension (WHO Group I), and WHO Functional Class II-IV symptoms.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Transmucosal Fentanyl Citrate

Products Affected

• fentanyl citrate buccal

PA Criteria	Criteria Details
Exclusion Criteria	Using for the treatment of acute or postoperative pain. Using for treatment of migraine headache pain Using for non-cancer related breakthrough pain.
Required Medical Information	
Age Restrictions	Individual is 16 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has a diagnosis of active cancer with breakthrough cancer (such as, metastatic or locally invasive cancer for which the individual is currently seeking treatment) with breakthrough cancer pain AND Individual is already receiving opioid therapy and is TOLERANT to opioid therapy as defined as receiving around the clock medicine consisting of one of the following: At least 60mg morphine per day, OR At least 25mcg/hr transdermal fentanyl/hour, OR At least 30mg of oxycodone daily, OR At least 8mg of oral hydromorphone daily, OR At least 25mg of oral oxymorphone daily, OR An equianalgesic dose of another opioid for a week or longer. Individual will also continue around the clock opioids when taking transmucosal fentanyl Citrate for cancer related breakthrough pain.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Trelstar Line

Products Affected

 TRELSTAR MIXJECT INTRAMUSCULAR SUSPENSION RECONSTITUTED 11.25 MG, 22.5 MG, 3.75 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For Prostate cancer: Clinically localized disease with intermediate (T2b to T2c cancer, Gleason score of 7/Gleason grade group 2-3, or prostate specific antigen (PSA) value of 10-20 ng/mL) OR higher risk of recurrence as neoadjuvant therapy with radiation therapy or cryosurgery OR Following radical prostatectomy as adjuvant therapy when lymph node metastases are present OR Locally advanced disease OR Other advanced, recurrent, or metastatic disease.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Tremfya

Products Affected

TREMFYA

PA Criteria	Criteria Details
Exclusion Criteria	Using in combination with other biologic drugs or phototherapy. Tuberculosis, other active serious infections, or a history of recurrent infections. Individual has not had a tuberculin skin test or a Centers for Disease Control and Prevention-recommended equivalent test to evaluate for latent tuberculosis prior to initiating Tremfya.
Required Medical Information	Diagnosis of moderate to severe plaque psoriasis with either of the following: Patient has greater than 5% of body surface area with plaque psoriasis OR Less than or equal to 5% of body surface area with plaque psoriasis involving sensitive areas or areas that would significantly impact daily function (such as palms, soles of feet, head/neck, or genitalia).
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For a dx of chronic plaque psoriasis, individual has failed to respond to, is intolerant of, or has a medical contraindication to phototherapy or any ONE systemic therapy (for example acitretin, methotrexate, cyclosporine) AND individual has had a trial of and had an inadequate response or is intolerant to either: Humira (adalimumab) or Enbrel (etanercept). For any of the above indications, if the TNF agent (Enbrel(etanercept)/Humira(adalimumab)) tried and failed are not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Tremfya or (b) Individuals age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis OR individual has either concomitant clinical condition: (a) Demyelinating disease or (b) Heart failure with documented left ventricular dysfunction, Tremfya may be allowed without trial of preferred TNF agents (Enbrel(etanercept)/Humira(adalimumab)).
Indications	All Medically-accepted Indications.
Off Label Uses	

Trikafta

Products Affected

• TRIKAFTA

PA Criteria	Criteria Details
Exclusion Criteria	Concurrent use with Kalydeco (ivacaftor), Orkambi (ivacaftor/lumacaftor) or Symdeko (tezacaftor/ivacaftor) OR Individual with severe hepatic impairment (Child-Pugh Class C).
Required Medical Information	
Age Restrictions	Individual is 12 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	For dx of CF, mutation testing confirms (verbal or written attestation) the individual has two copies of the F508del mutation in the cystic fibrosis transmembrane conductance regulator (CFTR) gene OR testing confirms (verbal or written attestation) the individual has one copy of the F508del mutation in the cystic fibrosis transmembrane conductance regulator (CFTR) gene and one other mutation that is unresponsive to Ivacaftor or Tezacaftor/Ivacaftor.
Indications	All Medically-accepted Indications.
Off Label Uses	

Triptodur

Products Affected

• TRIPTODUR

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Precocious puberty, defined as beginning of secondary sexual characteristics before age 8 in girls and before age 9 in boys.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For Gender Dysphoria use (Hembree 2009, 2017), individual fulfills the DSM V criteria for gender dysphoria (American Psychiatric Association 2013) AND has experienced puberty to at least Tanner stage 2 (Hembree 2009, 2017) AND has (early) pubertal changes that have resulted in an increase of their gender dysphoria (Hembree 2009, 2017) AND Individual does not suffer from a psychiatric comorbidity that interferes with the diagnostic work-up or treatment (Hembree 2009, 2017) AND has psychological and social support during treatment (Hembree 2009, 2017) AND demonstrates knowledge and understanding of the expected outcomes of GnRH analog treatment (Hembree 2009, 2017).
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Trodelvy

Products Affected

TRODELVY

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has recurrent or metastatic, histologically confirmed triple-negative Breast Cancer (lack of estrogen- and progesterone- receptor expression and no overexpression of HER2) AND Individual has confirmation of disease progression (written or verbal) after two prior therapies.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Trogarzo

Products Affected

• TROGARZO

PA Criteria	Criteria Details
Exclusion Criteria	Individual who has received immunomodulating therapy within 12 weeks of initiating treatment with Trogarzo (for example, interferon, systemic steroids or systemic chemotherapy) (NCT00784147) OR Individual is being treated for an acute infection secondary to HIV infection (NCT00784147).
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual is using to treat human immunodeficiency virus (HIV) infection AND has a viral load of greater than 1000 copies/mL AND has a history of at least 6 months of antiretroviral treatment AND is receiving a failing antiretroviral regimen or has failed and is off therapy AND has confirmed resistance to at least one antiretroviral agent from three different classes as measured by resistance testing (FDA Summary, 2018) AND Individual is using in combination with other antiretroviral agents and has confirmed full viral sensitivity/susceptibility to at least one antiretroviral agent (other than Trogarzo) as determined by resistance testing.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Tukysa

Products Affected

• TUKYSA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	HER2-positive breast cancer confirmed (verbal or written) by one of the following: Immunohistochemistry (IHC) is 3+ or In situ hypridization (ISH) positive.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Turalio

Products Affected

TURALIO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Tykerb

- lapatinib ditosylate
- TYKERB

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Cancer has been confirmed HER2 positive
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Tymlos

Products Affected

• TYMLOS

PA Criteria	Criteria Details
Exclusion Criteria	Individual has utilized abaloparatide and a parathyroid hormone analog (for example, teriparatide [Forteo]) for a combined total lifetime duration of 2 years or longer. Individual is using Tymlos in combination with any of the following: (1) Prolia (denosumab) OR (2) Bisphosphonate OR (3) Evista (raloxifene) OR (4) Miacalcin/Fortical (calcitonin nasal spray) OR (5) Reclast (zoledronic acid) OR (6) Forteo (teriparatide).
Required Medical Information	Individual is a postmenopausal female with one of the following: (A) dx of osteoporosis (defined as a bone mineral density [BMD] T- score in the spine, femoral neck, total hip or distal 1/3 of the radius of less than or equal to -2.5 as compared to a young-adult reference population) OR (B) dx of osteoporosis based on history of an osteoporotic low trauma fracture (fragility fracture) and considered at high risk for additional fracture AND Individual has had one of the following: (a) trial of an oral bisphosphonate OR (b) individual is intolerant to or has a contraindication to oral bisphosphonate therapy as defined by one of the following (1 through 5): (1) Hypersensitivity to TWO oral bisphosphonates (one of which must be generic alendronate) OR (2) Inability to stand or sit upright for at least 30 minutes OR (3) A pre-existing gastrointestinal disorder (for example, Barrett's esophagus, hypersecretory disorders, delayed esophageal emptying, etc.) OR (4) Uncorrected hypocalcemia OR (5) Severe renal insufficiency as defined by creatinine clearance less than 35 mL/min for alendronate agents or creatinine clearance less than 30 mL/min for risedronate and ibandronate.
Age Restrictions	
Prescriber Restrictions	
Y0114_20_114077	Z_I_012 December 2020

PA Criteria	Criteria Details
Coverage Duration	2 years. Requests to continue therapy beyond 24 months (2 years) should not be approved.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Tysabri

Products Affected

• TYSABRI

PA Criteria	Criteria Details
Exclusion Criteria	Using for Types of MS other than relapsing forms. Currently responsive to and tolerating another treatment for MS or CD. Current or prior history of progressive multifocal leukoencephalopathy (PML). Medical condition which significantly compromises the immune system including HIV infection or AIDS, leukemia, lymphoma or organ transplantation. Concurrent use with chronic antineoplastics or immunosuppressants (for example, azathioprine) or TNF inhibitors. Concurrent use with other MS disease modifying agents (such as Aubagio, Gilenya, Tecfidera, Lemtrada, Ocrevus, Copaxone/Glatopa, Extavia, Rebif, Avonex, Plegridy, or Betaseron). Positive test results for anti- John Cunningham virus (JCV) antibodies.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual is using as monotherapy for relapsing forms of multiple sclerosis (MS) who have had an inadequate response to, or are unable to tolerate, alternative treatments for MS. For diagnosis of Crohns disease, individual is enrolled in and met all conditions of the CD or MS Touch Prescribing Program.
Indications	All Medically-accepted Indications.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Off Label Uses	

Y0114_20_114077_I_012

Tyvaso

Products Affected

- TYVASO
- TYVASO REFILL
- TYVASO STARTER

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Diagnostic Criteria for PAH: right heart catheterization which shows a mean pulmonary artery pressure (mPAP) greater than or equal to 25mmhg, a pulmonary capillary wedge pressure (PCWP), left atrial pressure or left ventricular end diastolic pressure (LVEDP) less than or equal to 15mmhg AND a pulmonary vascular resistance (PVR) greater than 3 wood units. Criteria for Vasodilator Response: a favorable response is defined as a fall in mPAP of at least 10mmhg to an absolute mPAP of less than 40mmhg without a decrease in cardiac output, when challenged with inhaled nitric oxide, intravenous epoprostenol or intravenous adenosine.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For Tyvaso, patient must meet all of the following: meets diagnostic criteria for PAH AND demonstrates an unfavorable acute response to vasodilators AND Individual has New York Heart Association (NYHA) functional class III, or IV symptoms: AND World health Organization (WHO) Group I PAH (idiopathic PAH, PAH associated with connective tissue disorders, PAH associated with congenital heart defects, and all Group 1 subtypes).
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Ubrelvy

Products Affected

• UBRELVY

PA Criteria	Criteria Details
Exclusion Criteria	Using a strong CYP3A4 inhibitor (such as ketoconazole, itraconazole, clarithromycin).
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has had a trial/inadequate response or intolerance to 2 oral triptans (AHS 2019) OR Individual has one of the following CV or non-coronary vascular contraindications to use of triptans: Ischemic coronary artery disease (CAD) including angina pectoris, history of MI, documented silent ischemia, coronary artery vasospasm (including Prinzmetal's angina), history of stroke or TIA, PVD, ischemic bowel disease, or uncontrolled hypertension.
Indications	All Medically-accepted Indications.
Off Label Uses	

Uceris

Products Affected

- budesonide er
- UCERIS ORAL

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	18 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Ultomiris

Products Affected

• ULTOMIRIS

PA Criteria	Criteria Details
Exclusion Criteria	Individual has evidence of active meningococcal infection. Individual has not been immunized with meningococcal vaccine at least 2 weeks prior to administration of the first dose of Ultomiris unless clinical record documents the risks of delaying Ultomiris outweigh the risk of meningococcal infection.
Required Medical Information	Individual has Paroxysmal Nocturnal hemoglobinuria (PNH) as documented by flow cytometry (Written or Verbal attestation is acceptable)in the presence of 1) PNH type III red cell clone or a measurable granulocyte or monocyte clone OR 2) Glycosylphosphatidylinositol-anchored proteins (GPI-AP)-deficient polymorphonuclear cells(PMNs).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has Lactate dehydrogenase greater than 1.5 times upper limit of normal AND has one or more PNH related symptoms (such as but not limited to anemia or history of major adverse vascular event from thromboembolism)
Indications	All Medically-accepted Indications.
Off Label Uses	

Uptravi

- UPTRAVI ORAL TABLET
- UPTRAVI ORAL TABLET THERAPY PACK

PA Criteria	Criteria Details
Exclusion Criteria	Individual has a diagnosis of severe hepatic impairment (Child- Pugh Class C). In combination with prostacyclin analogs [such as but not limited to treprostinil (Orenitram, Remodulin, Tyvaso), Epoprostenol (Flolan, Veletri), Ventavis (iliprost)]
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has catheterization-proven diagnosis of pulmonary arterial hypertension (PAH) [World Health Organization (WHO) Group 1) AND individual has WHO functional class II-IV symptoms.
Indications	All Medically-accepted Indications.
Off Label Uses	

Valchlor

Products Affected

• VALCHLOR

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Vancocin

- FIRVANQ
- VANCOCIN
- VANCOCIN HCL ORAL CAPSULE 125 MG, vancomycin hcl oral solution 250 MG
- vancomycin hcl oral capsule 125 mg, 250 ٠ тg
 - reconstituted

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual is being treated for enterocolitis caused by Staphylococcal aureus including methicillin-resistant strains. Individual is being treated for clostridium difficile.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	14 days
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Vectibix

Products Affected

• VECTIBIX

PA Criteria	Criteria Details
Exclusion Criteria	Individual has received prior treatment with cetuximab (Erbitux) [Note: a course of cetuximab discontinued because of an adverse reaction is not considered prior treatment] OR Vectibix is used in combination with other anti-VEGF agents (e.g., bevacizumab) OR Vectibix is being used for more than one line (course) of therapy.
Required Medical Information	Extended RAS gene mutation testing with an FDA approved test and results confirm (written or verbal) the tumor is RAS wild-type.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	6 months
Other Criteria	Used as a single agent or as part of combination therapy for stage IV colon, rectal, colorectal, small bowel or anal adenocarcinoma.
Indications	All Medically-accepted Indications.
Off Label Uses	

Velcade

Products Affected

- BORTEZOMIB
- VELCADE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Vemlidy

Products Affected

• VEMLIDY

PA Criteria	Criteria Details
Exclusion Criteria	Individual has end stage renal disease (estimated creatinine clearance below 15 mL/min). Individual has decompensated (Child-Pugh B or C) hepatic impairment
Required Medical Information	
Age Restrictions	Individual is 18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Venclexta

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

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Ventavis

Products Affected

• VENTAVIS

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Diagnostic Criteria for PAH: right heart catheterization which shows a mean pulmonary artery pressure (mPAP) greater than or equal to 25mmhg, a pulmonary capillary wedge pressure (PCWP), left atrial pressure or left ventricular end diastolic pressure (LVEDP) less than or equal to 15mmhg AND a pulmonary vascular resistance (PVR) greater than 3 wood units. Criteria for Vasodilator Response: a favorable response is defined as a fall in mPAP of at least 10mmhg to an absolute mPAP of less than 40mmhg without a decrease in cardiac output, when challenged with inhaled nitric oxide, intravenous epoprostenol or intravenous adenosine.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For Ventavis, patient must meet all of the following: meets diagnostic criteria for PAH AND demonstrates an unfavorable acute response to vasodilators AND Individual has New York Heart Association (NYHA) functional class III, or IV symptoms: AND World health Organization (WHO) Group I PAH (idiopathic PAH, PAH associated with connective tissue disorders, PAH associated with congenital heart defects, and all Group 1 subtypes).
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Verzenio

Products Affected

• VERZENIO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Vfend

- VFEND
- voriconazole oral

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual is currently transitioning from inpatient treatment (hospital/medical facility) with IV antifungal (voriconazole) to an outpatient (home) setting.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Vibativ

Products Affected

• VIBATIV

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	30 day supply/One time only
Other Criteria	Individual has started therapy in an inpatient setting and requires continued outpatient therapy for an organism susceptible to VIBATIV (telavancin). For hospital-acquired or ventilator-associated bacterial pneumonia (HABP or VABP), Individual has had a trial and an inadequate response or intolerance to or has a contraindication to at least one alternative antibiotic (such as but not limited to, intravenous vancomycin) (IDSA 2011, ATS/IDSA 2005). For complicated skin and skin structure infections (cSSSI), Individual has had an inadequate response to at least one alternative therapy (such as but not limited to, intravenous vancomycin) (IDSA 2014, 2011).
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

VIBERZI

Products Affected

• VIBERZI

PA Criteria	Criteria Details
Exclusion Criteria	Individual has a history of severe constipation or sequelae from constipation OR Biliary duct obstruction or sphincter of Oddi dysfunction OR History of pancreatitis or structural disease of the pancreas OR Excessive alcohol intake (more than 3 alcoholic beverages per day) OR Severe hepatic impairment (Child-Pugh Class C) OR Concomitant use with Lotronex (alosetron) OR history of cholecystectomy or absence of a gallbladder.
Required Medical Information	
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 Year
Other Criteria	Individual is using for the treatment of irritable bowel syndrome with diarrhea (IBS-D) AND has had a trial and inadequate response or intolerance to two of the following medications or has a contraindication to all of the following medications: 1. Loperamide OR 2. Antispasmodics (such as dicyclomine) OR 3. Tricyclic antidepressants (AGA 2014).
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Vidaza

- azacitidine
- VIDAZA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Viekira

Products Affected

• VIEKIRA PAK

PA Criteria	Criteria Details
Exclusion Criteria	Individual is requesting in concurrent therapy with alfuzosin, carbamazepine, phenytoin, phenobarbital, gemfibrozil, rifampin, ergot derivatives, ethinyl estradiol-containing agents, St. Johns Wort, lovastatin, simvastatin, pimozide, efavirenz, Revatio (sildenafil), triazolam, and oral midazolam. Individual is using in combination with another NS3/4A protease inhibitor [such as but not limited to Olysio (simeprevir). Individual is using in combination with another non-nucleoside NS5B polymerase inhibitor. Individual is using in combination with another NS5A inhibitor (such as but not limited to, Harvoni [ledipasvir/sofosbuvir]. Individual is requesting the regimen for re-treatment and either failed to achieve a SVR (defined as a lower limit HCV RNA of 25 IU/mL) or relapsed after achieving a SVR during a prior successfully completed treatment regimen consisting of a serine NS3/4A protease inhibitor [such as but not limited to, Olysio (simeprevir), or paritaprevir] (AASLD/IDSA 2014). Individual is requesting the regimen for re-treatment and either failed to achieve a SVR (defined as a lower limit HCV RNA of 25 IU/mL) or relapsed after achieving a SVR during a prior successfully completed treatment regimen for re-treatment and either failed to achieve a SVR (defined as a lower limit HCV RNA of 25 IU/mL) or relapsed after achieving a SVR during a prior successfully completed treatment regimen consisting of ombitasvir. Individual is requesting the regimen for re-treatment and either failed to achieve a SVR (defined as a lower limit HCV RNA of 25 IU/mL) or relapsed after achieving a SVR during a prior successfully completed treatment regimen consisting of a non-nucleoside NS5B polymerase inhibitor, such as dasabuvir or a regimen containing a nucleotide NS5B polymerase inhibitor, such as Sovaldi or Harvoni (AASLD/IDSA 2014).

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Required Medical Information	Documentation is provided for a diagnosis of chronic hepatitis C (CHC) infection, which includes genotype and positive HCV RNA result (AASLD/IDSA 2017, CDC 2013) AND Individual has received baseline evaluation for liver fibrosis to guide appropriate therapy AND Individual does not have a short life expectancy (less than 12 months owing to non-liver related comorbid conditions) that cannot be remediated by treating HCV, by transplantation or other directed therapy (AASLD/IDSA 2016).
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	Criteria will be applied consistent with current AASLD/IDSA guidance.
Other Criteria	Criteria will be applied consistent with current AASLD/IDSA guidance. Individual has had a prior trial and inadequate response to Harvoni (sofosbuvir/ledipasvir) OR individual is currently on and completing a course of therapy with the requested regimen OR Individual has a documented hypersensitivity, as manifested by a severe allergic reaction, to any ingredient in Harvoni (ledipasvir/sofosbuvir) which is not also in Viekira OR Individual is concurrently using an agent that cannot be substituted with another agent or temporarily discontinued and is contraindicated or not recommended for concomitant use with Harvoni (sofosbuvir/ledipasvir).
Indications	All Medically-accepted Indications.
Off Label Uses	

Vimizim

Products Affected

• VIMIZIM

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Confirmed diagnosis of Morquio A syndrome (Hendriksz 2015, Wood 2013) by documented reduced fibroblast or leukocyte N- acetylgalactosamine-6-sulfatase (GALNS) enzyme activity combined with normal enzyme activity level or another sulfatase or by genetic testing and Documented clinical signs and symptoms of Morquio A syndrome (for example, knee deformity, corneal opacity or pectus carinatum) (Hendriksz 2015, Wood 2013).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Vimovo

Products Affected

- naproxen-esomeprazole
- VIMOVO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual has had a trial and inadequate response or intolerance to one (1) oral generic prescription Non-Steroidal Anti-Inflammatory Drugs (NSAID) AND has had a trial and inadequate response or intolerance to one (1) of the following (Lanza 2009): (a) preferred proton pump inhibitor (PPI) OR (b) Generic misoprostol AND Individual has had an adequate response (pain relief and appropriate gastro protection) with a trial of naproxen and a proton pump inhibitor (such as esomeprazole) used at the same time AND Documentation has been provided for why the combination agent is clinically necessary and not for convenience.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Virazole

Products Affected

- ribavirin inhalation
- VIRAZOLE

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual is hospitalized and will receive treatment in an inpatient setting.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Vitrakvi

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

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Written or verbal attestation to confirm genetic test results show the tumor has a neurotrophic receptor tyrosine kinase (NTRK) gene fusion without a known acquired resistance mutation.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Vizimpro

Products Affected

 VIZIMPRO ORAL TABLET 15 MG, 30 MG, 45 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	genetic mutations test result is confirmed by written or verbal attestation.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Vosevi

Products Affected

• VOSEVI

PA Criteria	Criteria Details
Exclusion Criteria	Individual has severe renal impairment (eGFR less than 30 mL/min/1.73m2), end stage renal disease, or requires dialysis OR Individual has moderate or severe hepatic impairment (Child-Pugh B or C).
Required Medical Information	Documentation is provided for a diagnosis of chronic hepatitis C (CHC) infection, which includes a genotype and positive HCV RNA result (AASLD/IDSA 2017, CDC 2013) AND Individual has received baseline evaluation for liver fibrosis to guide appropriate therapy AND Individual does not have a short life expectancy (less than 12 months owing to non-liver related comorbid conditions) that cannot be remediated by treating HCV, by transplantation or other directed therapy (AASLD/IDSA 2016).
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	Criteria will be applied consistent with current AASLD/IDSA guidance.

PA Criteria	Criteria Details
Other Criteria	Criteria will be applied consistent with current AASLD/IDSA guidance. For Genotype 1, 1a Individual has had a trial of and inadequate response to Harvoni (sofosbuvir/ledipasvir) OR Individual is currently on and completing a course of therapy with Vosevi OR has one of the following: (1) Documented hypersensitivity, as manifested by a severe allergic reaction to any ingredient in Harvoni(sofosbuvir/ledipasvir) which is not also in Vosevi OR (2) concurrently using an agent that cannot be substituted with another agent or temporarily discontinued and is contraindicated or not recommended for concomitant use with the preferred regimen or regimens OR (3) Individual failed to achieve a sustained viral response (SVR) or relapsed after achieving a SVR during a prior successfully completed Hepatitis C regimen containing an NS5A inhibitor. For Genotype 4 Individual has had a trial of and inadequate response to Harvoni(sofosbuvir/ledipasvir) or Epclusa(sofosbuvir/velpatasvir) OR Individual is currently on and completing a course of therapy with Vosevi OR has one of the following: (1) Documented hypersensitivity, as manifested by a severe allergic reaction to any ingredient in Harvoni(sofosbuvir/ledipasvir) or Epclusa(sofosbuvir/velpatasvir) which is not also in Vosevi OR (2) concurrently using an agent that cannot be substituted with another agent or temporarily discontinued and is contraindicated or not recommended for concomitant use with the preferred regimen or regimens OR (3) Individual failed to achieve a sustained viral response (SVR) or relapsed after achieving a SVR during a prior successfully completed Hepatitis C regimen containing an NS5A inhibitor.
Indications	All Medically-accepted Indications.
Off Label Uses	

VOTRIENT

Products Affected

• VOTRIENT

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Vumerity

- VUMERITY
- VUMERITY (STARTER)

PA Criteria	Criteria Details
Exclusion Criteria	Concurrent use with other MS disease modifying agents (including Aubagio, Avonex, Betaseron, Copaxone/Glatiramer/Glatopa, Extavia, Gilenya, Lemtrada, Mavenclad, Mayzent, Ocrevus, Plegridy, Rebif, Tecfidera and Tysabri). Individual with moderate or severe renal impairment (creatinine clearance less than 60 mL/min). Individual is using to treat non-active secondary progressive multiple sclerosis.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Vyepti

Products Affected

• VYEPTI

PA Criteria	Criteria Details
Exclusion Criteria	Individual is using concomitantly with botulinum toxin for migraine prophylaxis.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Initial 3 month, Continuation 1 year.

PA Criteria	Criteria Details
Other Criteria	For Initial requests: Individual has dx of one of the following: (a)Episodic migraine defined as at least 4 and fewer than 15 migraine days per month and fewer than 15 headache days per month on average during the previous 3 month period OR (b) Chronic migraine defined as a headache occurring on 15 or more days per month for more than 3 months, which, on at least 8 days per month, has features of a migraine headache (ICHD-3). AND Individual has had a trial of and inadequate response or intolerance to two agents for migraine prophylaxis (at least one agent in any two of the following classes) or has a contraindication to all of the following medications (AAN/AHA 2012/2015, Level A and B evidence, ICSI 2013, high quality evidence): (a) The following antidepressants: amitriptyline, venlafaxine or (b)One of the following beta blockers: Metoprolol, propranolol, timolol (oral), nadolol, atenolol, nebivolol or (c) The following antiepileptic agents: valproate sodium, divalproex sodium, topiramate, gabapentin or (e) Botox (for chronic migraine). For Renewal requests: Individual has a reduction in the overall number of migraine days or reduction in number of severe migraine days per month AND Individual has obtained clinical benefit deemed significant by individual or prescriber.
Indications	All Medically-accepted Indications.
Off Label Uses	

Vyleesi

Products Affected

• VYLEESI

PA Criteria	Criteria Details
Exclusion Criteria	Treatment of HSDD in postmenopausal women. Treatment of HSDD in men. Enhancement of sexual performance. Individual has uncontrolled hypertension (systolic blood pressure greater than or equal to 140 mmHg, or diastolic blood pressure greater than or equal to 90 mmHg) (Clayton 2017 and ACC/AHA 2017). Individual has known cardiovascular disease (including, but not limited to, coronary heart disease, cerebrovascular disease, peripheral vascular disease, heart failure, and cardiomyopathies) (WHO 2019). Individuals who will be using Vyleesi (bremelanotide) in combination with Addyi (flibanserin).
Required Medical Information	
Age Restrictions	Individual is 18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year.

PA Criteria	Criteria Details
Other Criteria	For diagnosis of acquired, generalized hypoactive sexual desire disorder (HSDD) or acquired Female Sexual Interest Arousal Disorder (FSIAD) for at least 24 weeks characterized low sexual desire that causes marked distress or interpersonal difficulty. Individual is female AND is premenopausal AND it is confirmed that the diagnosis of HSDD or FSIAD is not caused by any of the following: (A) A co-existing psychiatric condition OR (B) A co- existing medical condition that could contribute to sexual dysfunction OR (C) Problems within a relationship OR (D) Other co- existing psychological conditions, such as loss of income or bereavement (NCT02333071, APA 2013). (E) Effects of a medication or other drug substance.
Indications	All Medically-accepted Indications.
Off Label Uses	

Vyondys 53

Products Affected

• VYONDYS 53

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For initial therapy, individual has a confirmed diagnosis of Duchenne muscular dystrophy (DMD) AND a genetic mutation that is amendable to exon 53 skipping.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Continuation of therapy may be approved when initial therapy criteria are met and individual remains ambulatory with or without needing an assistive device, such as a cane or walker.
Indications	All Medically-accepted Indications.
Off Label Uses	

Vytorin

- ezetimibe-simvastatin
- VYTORIN

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has had a trial of TWO generic statin (at any dose) and did not achieve LDL cholesterol goal OR Individual is currently on an agent that interacts with both preferred generics.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Vyvanse

Products Affected

• VYVANSE

PA Criteria	Criteria Details
Exclusion Criteria	Individual is using for weight loss.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual has diagnosis of binge-eating disorder OR has a diagnosis of attention deficit hyperactivity disorder (ADHD) AND Individual has had a trial of and insufficient response or intolerance to one of the following: (1) Methylphenidate extended-release or (2) Extended-release amphetamine/dextroamphetamine salt combination OR Individual has been diagnosed with coexisting ADHD and substance use disorder.
Indications	All Medically-accepted Indications.
Off Label Uses	

Wakix

Products Affected

• WAKIX

PA Criteria	Criteria Details
Exclusion Criteria	Individual has severe hepatic impairment. Individual has a risk factor for prolonged QT interval. Is using with another drug that increases the QT interval.
Required Medical Information	For Narcolepsy type 1 confirmed by the following (ICSD-3): (1) Daily periods of irrepressible need to sleep or daytime lapses into sleep occurring for at least 3 months and at least one of the following: (a) Clear cataplexy (defined as more than one episode of generally brief [less than 2 min]) usually bilaterally symmetrical, sudden loss of muscle tone with retained consciousness) AND (b) Multiple Sleep Latency Test (MSLT) with one of the following: (i) Mean sleep latency of less than 8 minutes with evidence of two sleep-onset rapid eye movement periods (SOREMPs) OR (ii) At least one SOREMP on MSLT and a SOREMP (less than 15 minutes) on the preceding overnight polysomnography (PSG) OR (c) Cerebrospinal fluid hypocretin-1 deficiency (less than 100 pg/mL or less than one-third of the normative values with the same standardized assay). Narcolepsy type 2 confirmed by the following (ICSD-3): (1) Daily periods of irrepressible need to sleep or daytime lapses into sleep occurring for at least 3 months AND (2) Multiple sleep latency test (MSLT) with one of the following: (a) MSLT of less than 8 minutes and evidence of two sleep-onset rapid eye movement periods (SOREMPs) (ICSD-3, 2014) OR (b) At least one SOREMP on MSLT and a SOREMP (less than 15 minutes) on the preceding overnight polysomnography (PSG) AND (3) The absence of cataplexy AND (4) Exclusion of alternative causes of excessive daytime sleepiness by history, physical exam, and PSG.
Age Restrictions	Individual is 18 years of age or older
Prescriber Restrictions	
Y0114 20 114077	I 012 December 2020

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Xalkori

Products Affected

• XALKORI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Documentation is provided that tumor is anaplastic lymphoma kinase (ALK)-positive or c-ros oncogene 1 (ROS1) positive.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Xeljanz PDP

- XELJANZ
- XELJANZ XR

PA Criteria	Criteria Details
Exclusion Criteria	Using in combination with, other JAK inhibitors (such as Olumiant), biologic disease-modifying antirheumatic drug (DMARDs) (such as but not limited to, TNF agents, anti-CD20 monoclonal antibodies, IL-1R antagonists, selective co-stimulation modulators) or potent immunosuppressants such as azathioprine and cyclosporine. At initiation of therapy, absolute neutrophil count (ANC) less than 1000 cells/mm3, lymphocyte count less than 500 cells/mm3, or hemoglobin less than 9 g/dL. Tuberculosis or other active serious infections or a history of recurrent infections. Individual has not had a tuberculin skin test (TST) or a Centers for Disease Control (CDC)-recommended equivalent to evaluate for latent tuberculosis prior to initiating Xeljanz. Individual has severe hepatic impairment (Child Pugh class C).
Required Medical Information	
Age Restrictions	18 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year.

PA Criteria	Criteria Details
Other Criteria	For RA and PsA, Individual had an inadequate response to, is intolerant of, or has a contraindication ONE conventional therapy [nonbiologic DMARDS (such as methotrexate, sulfasalazine, leflunomide, or hydroxychloroquine)] AND individual has had a trial of and an inadequate response or is intolerant to: Humira(adalimumab) OR the TNF agent (Humira (adalimumab)) tried and failed is not acceptable due to concomitant clinical conditions, such as but not limited to any of the following: (a) Known hypersensitivity or any active or inactive component which is not also associated with Xeljanz or (b) Individual's age or (c) Pregnant or planning or becoming pregnant or (d) Serious infections or concurrent sepsis. For UC, Individual has had an inadequate response to, is intolerant of, or has a medical contraindication to ONE conventional therapy (such as 6- mercaptopurine, azathioprine, oral aminosalicylates, or oral corticosteroids) AND has had a trial of and an inadequate response or is intolerant to Humira (adalimumab) OR Humira (adalimumab) is not acceptable due to concomitant clinical conditions, such as but not limited to any of the following listed above (a-d)
Indications	All Medically-accepted Indications.
Off Label Uses	

XENAZINE

- tetrabenazine oral tablet 12.5 mg, 25 mg
- XENAZINE ORAL TABLET 12.5 MG, 25
 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Xermelo

Products Affected

XERMELO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	Initial: 3 months. Continuation: 1 year
Other Criteria	For initial therapy: Individual is using in combination with somtostatin analog (SSA) therapy (such as but not limited to, lanreotide(Somatuline Depot), cotreotide (Sandostatin)) AND individual has had an inadequate response on a stable dose of SSA monotherapy for at least 3 months. For continuation therapy: Individual has previously met the initiation criteria AND if clinically significant improvements are confirmed after 12 weeks pf treatment with Xermelo (telotristat ethyl) when added to SSA therapy
Indications	All Medically-accepted Indications.
Off Label Uses	

Xgeva

Products Affected

• XGEVA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Skeletally mature adolescent is defined by at least one mature long bone [for example, closed epiphyseal growth plate of the humerus]. Hypercalcemia of malignancy is defined as an albumin corrected serum calcium level greater than 12.5 mg/dL (3.1 mmol/L).
Indications	All Medically-accepted Indications.
Off Label Uses	

Xiaflex

Products Affected

• XIAFLEX

PA Criteria	Criteria Details
Exclusion Criteria	Repeat injection of a previously treated cord within one year of a prior course for treating Dupuytren's contracture.
Required Medical Information	For Peyronie disease, stable disease as define by symptoms (that is, penile curvature and pain) for at least 6 months and Penile curvature greater than or equal to 30 and less than or equal to 90 degrees and Intact erectile function with or without use of medications and Palpable penile plaque. For Dupuytren's contracture, there is documented impairment to the individual's functional activities which measures either: 20 degrees or more at the metacarpophalangeal (MP) joint or 20 degrees or more at the proximal interphalangeal (PIP) joint.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Xifaxan - HE

Products Affected

• XIFAXAN ORAL TABLET 550 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	18 years of age or older.
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Xifaxan 200mg

- AEMCOLO
- XIFAXAN ORAL TABLET 200 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	For Xifaxan 200mg, travelers diarrhea (TD) caused by noninvasive strains of Escherichia coli AND Individual has already been started on Xifaxan and needs to complete treatment OR Individual has had a trial and inadequate response or intolerance to one of the following medications or has contraindications to all of the following medications (CDC, 2018): (1)Generic Fluoroquinolone OR(2)Azithromycin.
Indications	All Medically-accepted Indications.
Off Label Uses	

Xolair

- XOLAIR SUBCUTANEOUS SOLUTION
 PREFILLED SYRINGE
- XOLAIR SUBCUTANEOUS SOLUTION RECONSTITUTED

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has Moderate to Severe Persistent Asthma AND has a positive skin test or in vitro reactivity to a perennial aeroallergen, AND individual has an FEV1 less than 80% predicted AND IgE level is equal to or greater than 30 IU/ml. Severe asthma as defined by the National Heart, Lung, and Blood Institute: Severe Persistent Asthma: symptoms throughout the day, extremely limited normal activity. Nocturnal symptoms are frequent, FEV1 or PEF is less than or equal to 60% predicted. Moderate Persistent Asthma as defined by the National Heart, Lung, and Blood Institute: Daily symptoms, daily use of inhaled short-acting beta2-agonist, somewhat limited activity, Nocturnal symptoms occur greater than 1 time per week, FEV1 or PEF is greater than 60% and less than 80% predicted, FEV1 FVC is reduced 5 percent or exacerbations requiring oral systemic corticosteroids use for more than or equal to 2 times per year.
Age Restrictions	Patient is 12 years of age or older for urticaria and 6 years of age or older for moderate to severe persistent asthma
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For moderate to severe persistent asthma, individual has had a minimum of 3 month trial and inadequate response or intolerance to ONE combination controller therapy (medium to high doses of inhaled corticosteroids plus long acting beta-2 agonists or Leukotriene modifiers)(GINA 2018). Continued treatment beyond 12 months is allowed when treatment has resulted in clinical improvement as documented by one or more of the following: Decreased utilization of rescue medications OR Decreased frequency of exacerbations (defined as worsening of asthma that requires increase in inhaled corticosteroid dose or treatment with systemic corticosteroids) OR Increase in percent predicted FEV1 from pretreatment baseline OR Reduction in reported asthmarelated symptoms, such as, but not limited to, wheezing, shortness of breath, coughing, fatigue, sleep disturbance, or asthmatic symptoms upon awakening. For chronic idiopathic urticaria, individual has had trial and inadequate response or intolerance to ONE potent antihistamine (AAAAI/ACAAI 2014).
Indications	All Medically-accepted Indications.
Off Label Uses	

Xospata

Products Affected

• XOSPATA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Individual has confirmed FMS-like tyrosine kinase 3 (FLT3) mutation (written or verbal attestation is acceptable).
Age Restrictions	18 years of age and older.
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Xpovio

- XPOVIO (100 MG ONCE WEEKLY)
- XPOVIO (60 MG ONCE WEEKLY)
- XPOVIO (80 MG ONCE WEEKLY)
- XPOVIO (80 MG TWICE WEEKLY)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Xpovio

- XPOVIO (40 MG ONCE WEEKLY)
- XPOVIO (40 MG TWICE WEEKLY)
- XPOVIO (60 MG TWICE WEEKLY)

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Xtandi

Products Affected

• XTANDI

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual is concomitantly receiving a gonadotropin-releasing hormone (GnRH) analog or has had a bilateral orchiectomy.
Indications	All Medically-accepted Indications.
Off Label Uses	

Xuriden

Products Affected

• XURIDEN

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Xyrem

- XYREM
- XYWAV

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	For initial tx of Narcolepsy type 1 (narcolepsy with cataplexy) confirmed by the presence of daily periods of irrepressible need to sleep or daytime lapses into sleep occurring for at least 3 months and at least one of the following: (1) Clear cataplexy (defined as more than one episode of generally brief [less than 2 min]) usually bilaterally symmetrical, sudden loss of muscle tone with retained consciousness) AND (2) Multiple Sleep Latency Test (MSLT) showing one of the following: (a) MSLT of less than 8 minutes with evidence of two sleep-onset rapid eye movement periods (SOREMPs) (ICSD-3, 2014) or (b) At least one SOREMP on MSLT and a SOREMP (less than 15 minutes) on the preceding overnight polysomnography (PSG) OR (3) Cerebrospinal fluid hypocretin-1 deficiency (less than 110 pg/mL or less than one-third of the normative values with the same standardized assay).
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	Initial Request 3 months, Renewal is 6 months.

PA Criteria	Criteria Details
Other Criteria	For initial tx, of Narcolepsy type 2 (narcolepsy without cataplexy) confirmed by the following: (1) Daily periods of irrepressible need to sleep or daytime lapses into sleep occurring for at least 3 months AND (2) MSLT showing one of the following: (a) MSLT of less than 8 minutes with evidence of two SOREMPs (ICSD-3, 2014) or (b) At least one SOREMP on MSLT and a SOREMP (less than 15 minutes) on the preceding overnight PSG AND (3) absence of cataplexy AND (4) Exclusion of alternative causes of excessive daytime sleepiness by history, physical exam and PSG. AND (5) Mbr has had a previous trial of and inadequate response or intolerance to TWO of the following medications: (A) One of the following wakefulness promoting medications: (i) Modafinil or (ii) Nuvigil (armodafinil) AND (B) One of the following stimulants: (i) Methylphenidate (ii) Dextroamphetamine or (iii) Amphetamine/dextroamphetamine salt immediate-release OR (6) Trials of wakefulness promoting agents and stimulant agents are not acceptable due to concomitant clinical situations including but not limited to the following: (1) Cardiovascular disease or (2) Drug interactions. For Renewal of Narcolepsy type I or II, Xyrem (sodium oxybate) use has resulted in a reduction in frequency of cataplexy attacks OR use has resulted in a reduction in excessive daytime sleepiness (EDS) as measured by improvement in Epworth Sleepiness Test (MWT). For continuation, use has resulted in a reduction in frequency of cataplexy attacks compared to baseline OR use has resulted in a reduction in excessive daytime sleepiness (EDS) as measured by improvement in Epworth Sleepiness Cale (ESS) measurements or Maintenance of Wakefulness Test (MWT) compared to baseline
Indications	All Medically-accepted Indications.
Off Label Uses	

Yervoy

Products Affected

• YERVOY

PA Criteria	Criteria Details
Exclusion Criteria	Individual has autoimmune disease which requires treatment with immunosuppressant drugs.
Required Medical Information	For small cell lung cancer, unresectable or metastatic melanoma (cutaneous or uveal), colorectal cancer, renal cell carcinoma, or first line treatment of stage IV/recurrent NSCLC, individual has an Eastern Cooperative Oncology Group (ECOG) performance status of 0-2.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year

PA Criteria	Criteria Details
Other Criteria	For the treatment of unresectable or metastatic melanoma (cutaneous and uveal): Used in combination with nivolumab (Opdivo) as: (a) First-line therapy or (b) Second-line or subsequent therapy for disease progression if nivolumab was not previously used or Ipilimumab is used as a single agent for one of the following: (a) First line therapy as a single course of 4 treatments or (b) Second-line or subsequent lines of therapy as a single course of 4 treatments or (c) Retreatment, consisting of a 4-dose limit, for an individual who had no significant systemic toxicity during prior ipilimumab therapy, and whose disease progressed after being stable for greater than 6 months following completion of a prior course of ipilimumab, and for whom no intervening therapy has been administered. OR used for the adjuvant treatment of cutaneous melanoma in individuals with pathologic involvement of regional lymph nodes of more than 1 millimeter who have undergone complete resection, including lymphadenetomy. For the treatment of small cell lung cancer (SCLC): Yervoy is used in combination with nivolumab (Opdivo) as subsequent therapy for one of the following complete or partial response or stable disease with initial treatment, OR 2) no response with initial treatment, OR 3) primary progressive disease. For colorectal cancer AND meets one of the following criteria: (a) Primary tx used in combination with nivolumab for unresectable metachronous metastases (defective mismatch repair/high microsatellite instability [dMMR/MSIH] only) and previous adjuvant FOLFOX (fluorouracil, leucovorin, and oxaliplatin) or CapeOX (capecitabine and oxaliplatin) within the past 12 months or (b) Ipilimumab is used in combination with nivolumab as subsequent therapy for unresectable advanced or metastatic colorectal cancer with defective mismatch repair (dMMR) or high microsatellite instability (MSIH) mutations that has progressed following treatment with fluoropyrimidine and oxaliplatin or irinotecan. For Renal cell

PA Criteria	Criteria Details
	carcinoma, when: (a) used in combination with nivolumab, as first- line therapy for previously untreated RCC or (b) used in subsequent therapy with nivolumab if no checkpoint blockade (PD- 1, PD-L1, or CTLA-4) antibody treatment has been previously administered and (c)Histologic confirmation of RCC with clear-cell component. For stage IV/recurrent NSCLC when: used in combination with nivolumab and Cytologically confirmed stage IV or recurrent NSCLC and High tumor mutation burden (greater than or equal to 10 mutations per megabase) and No sensitizing epidermal growth factor receptor (EGFR) mutations or anaplastic lymphoma kinase (ALK) translocations in nonsquamous carcinoma and Has not received prior systemic therapy as primary therapy for advanced or metastatic NSCLC, prior adjuvant or neoadjuvant chemotherapy is permitted as long as the last administration of the prior regimen occurred at least 6 months prior.
Indications	All Medically-accepted Indications.
Off Label Uses	

Yonsa

Products Affected

• YONSA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual is concomitantly receiving a gonadotropin-releasing hormone (GnRH) analog or has had a bilateral orchiectomy.
Indications	All Medically-accepted Indications.
Off Label Uses	

Zaltrap

Products Affected

• ZALTRAP

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Diagnosis of metastatic anal adenocarcinoma or metastatic appendicle adenocarcinoma or metastatic small bowel adenocarcinoma or metastatic colorectal cancer AND used in combination with an irinotecan based regimen AND individual is resistant to or has disease progression following treatment with an oxaliplatin containing regimen AND Zaltrap will be used in a single line of therapy.
Indications	All Medically-accepted Indications.
Off Label Uses	

Zarxio

Products Affected

• ZARXIO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Febrile neutropenic individuals who are at risk for infection- associated complications or have any of the following: Expected prolonged (greater than 10 day) and profound (less than 0.1 x 10to the power of 9/L) neutropenia, Age greater than 65 years, Pneumonia or other clinically documented infection, Hypotension and multi organ dysfunction (sepsis syndrome), Invasive fungal infection, prior episode of FN, or Hospitalized at the time of the development of fever. Primary prophylaxis of FN in patients who have a risk of FN of 20% or greater based on chemotherapy regimens. OR when the risk of developing FN is greater than or equal to 10% and less than 20% based on chemotherapy in patients who have other patient specific risk factors for FN including any of the following: Individual age greater than 65 years, Poor performance status (ECOG status 3-4), prior chemotherapy or radiation therapy, Bone marrow involvement by tumor producing cytopenias, persistent neutropenia (ANC less than 1500mm3), poor renal function (GFR less than 60mL/min), liver dysfunction (liver function tests at least 2x upper limit of normal) or recent surgery and/or presence of open wounds.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Other Criteria	Secondary Prophylaxis for patients who experienced a neutropenic complication from a prior cycle of chemotherapy (for which primary prophylaxis was not received), in which a reduced dose may compromise disease-free or overall survival or treatment outcome. Using for adjunctive tx for FN and has been prophylactic therapy with GCSF agent or has not received prophylactic therapy with GCSF and who are at high risk for infection-associated complications. Use in acute lymphocytic leukemia (ALL) after completion of the first few days of initial induction chemotherapy or first post-remission course of chemotherapy. Use in individuals with acute myeloid leukemia (AML) shortly after the completion of induction chemotherapy, or after the completion of consolidation chemotherapy, or after the completion of consolidation chemotherapy for AML. For tx of moderate to severe aplastic anemia. Tx of severe neutropenia in individuals with hairy cell leukemia. For myelodysplastic syndromes (MDS) with severe neutropenia (absolute neutrophil count (ANC) less than or equal to 500 mm3 or experiencing recurrent infection. For dose dense therapy (treatment given more frequently, such as every two weeks instead of every three weeks) for adjuvant treatment of breast cancer. For chronic administration to reduce the incidence and duration of sequelae of neutropenia (e.g., fever, infections, oropharyngeal ulcers) in symptomatic individuals with congenital neutropenia, cyclic neutropenia, or idiopathic neutropenia. For tx of low neutrophil counts in individuals with glycogen storage disease type 1b. For tx of neutropenia associated with human immunodeficiency virus (HIV) infection and antiretroviral therapy. In individuals receiving radiation therapy in the absence of chemotherapy if prolonged delays secondary to neutropenia are expected. After accidental or intentional total body radiation of myleosuppressive doses (greater than 2 Grays [Gy] (such as Hematopoietic Syndrome of Acute Radiation Syndrome). After hematopoietic progenitor stem cell

PA Criteria	Criteria Details
	to promote myeloid reconstitution or when engraftment is delayed or has failed. To mobilize progenitor cells into peripheral blood for collection by leukapheresis, as an adjunct to peripheral blood/hematopoietic stem cell transplantation (PBSCT/PHSCT). Use as an alternate or adjunct to donor leukocyte infusions (DLI) in individuals with leukemic relapse after an allogeneic hematopoietic stem cell transplant.
Indications	All Medically-accepted Indications.
Off Label Uses	

Zavesca

Products Affected

- miglustat
- ZAVESCA

PA Criteria	Criteria Details
Exclusion Criteria	May not be approved for use in conjunction with Cerdelga (eliglustat) or enzyme replacement therapy (ERT) agents (Cerezyme, Elelyso or Vpriv). Severe Type 1 Gaucher disease (hemoglobin less than 9 g/dL, platelet count less than 50,000 mm3 or those at risk developing new bone complications) (Weinreb et al. 2005). Individual has severe renal impairment (less than 30 mL/min/1.73 m2). Individual has mild, moderate or severe hepatic impairment or cirrhosis.
Required Medical Information	Presence of type 1 Gaucher disease is confirmed by either of the following (Weinreb et al. 2004, Wang et al. 2011): Glucocerebrosidase activity in the white blood cells or skin fibroblasts less than or equal to 30% of normal activity, OR genotype testing indicating mutation of two alleles of the glucocerebrosidase genome. And patient has clinically significant manifestations of gauchers disease including any of the following: skeletal disease (demonstrated by radiologic evidence of ANY of the following (Weineb et al. 2004):: avascular necrosis, erlenmeyer flask deformity, lytic disease, marrow infiltration, osteopenia, osteosclerosis, pathological fracture, joint deterioration or replacement) OR patient presents with at least 2 of the following (Weinreb et al. 2004, Mistry et al. 2015): clinically significant hepatomegaly as confirmed by medical imaging (such as but limited to, volumetric MRI), clinically significant splenomegaly as confirmed by medical imaging (such as but limited to, volumetric MRI), spleess than or equal to 11.5 grams per deciliter below lower limit for normal for age and sex, platelet counts less than or equal to 120,000 mm3.
Age Restrictions	Individual is 18 years of age or older

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Enzyme replacement therapy with Cerezyme, ELELYSO or VPRIV is not a therapeutic option for reasons such as but limited to any of the following (Label, Weinreb et al. 2005): (a) Medically unmanageable hypersensitivity or (b) Development of therapy- limiting inhibitory antibodies or (c) Poor peripheral or central venous access.
Indications	All Medically-accepted Indications.
Off Label Uses	

Zejula

Products Affected

• ZEJULA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	In the last 8 weeks, the individual has had a complete or partial response to a platinum-based chemotherapy.
Indications	All Medically-accepted Indications.
Off Label Uses	

Zelboraf

Products Affected

• ZELBORAF

PA Criteria	Criteria Details
Exclusion Criteria	Individuals with wild-type BRAF melanoma.
Required Medical Information	Individual has BRAF mutation and a copy of the BRAF test results must be provided.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Zepatier

Products Affected

• ZEPATIER

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Documentation is provided for a diagnosis of chronic hepatitis C (CHC) infection, which includes a genotype and positive HCV RNA result (AASLD/IDSA 2017, CDC 2013) AND Individual has received baseline evaluation for liver fibrosis to guide appropriate therapy AND Individual does not have a short life expectancy (less than 12 months owing to non-liver related comorbid conditions) that cannot be remediated by treating HCV, by transplantation or other directed therapy (AASLD/IDSA 2017).
Age Restrictions	18 years of age or older.
Prescriber Restrictions	
Coverage Duration	Criteria will be applied consistent with current AASLD/IDSA guidance.

PA Criteria	Criteria Details
Other Criteria	Criteria will be applied consistent with current AASLD/IDSA guidance. For GT 1, Individual has had a prior trial and inadequate response to Harvoni (sofosbuvir/ledipasvir) OR individual is currently on and completing a course of therapy with the requested regimen OR Individual has a documented hypersensitivity, as manifested by a severe allergic reaction, to any ingredient in Harvoni (ledipasvir/sofosbuvir) which is not also in Zepatier OR Individual is concurrently using an agent that cannot be substituted with another agent or temporarily discontinued and is contraindicated or not recommended for concomitant use with Harvoni (sofosbuvir/ledipasvir) OR Individual has concomitant severe or end-stage CKD or requires dialysis. For GT 4, individual has had a prior trial and inadequate response to Harvoni (sofosbuvir/ledipasvir) or Epclusa(sofosbuvir/velpatasvir) OR individual is currently on and completing a course of therapy with the requested regimen OR Individual has a documented hypersensitivity, as manifested by a severe allergic reaction, to any ingredient in Harvoni(sofosbuvir/ledipasvir) or Epclusa(sofosbuvir/velpatasvir) which is not also in Zepatier OR Individual is concurrently using an agent that cannot be substituted with another agent or temporarily discontinued and is contraindicated or not recommended for concomitant use with Harvoni(sofosbuvir/ledipasvir) or Epclusa(sofosbuvir/velpatasvir) or Epclusa(sofosbuvir/velpatasvir) which is not also in Zepatier OR Individual is concurrently using an agent that cannot be substituted with another agent or temporarily discontinued and is contraindicated or not recommended for concomitant use with Harvoni(sofosbuvir/ledipasvir) or Epclusa(sofosbuvir/velpatasvir) OR Individual has concomitant severe or end-stage CKD or requires dialysis.
Indications	All Medically-accepted Indications.
Off Label Uses	

Zeposia

Products Affected

- ZEPOSIA
- ZEPOSIA 7-DAY STARTER PACK
- ZEPOSIA STARTER KIT

PA Criteria	Criteria Details
Exclusion Criteria	Concurrent use with other MS disease modifying agents (including Aubagio, Avonex, Betaseron, Copaxone/Glatiramer/Glatopa, Extavia, Gilenya, Lemtrada, Mavenclad, Mayzent, Ocrevus, Plegridy, Rebif, Tecfidera, Tysabri and Vumerity). Individual has had a recent (within the past 6 months) occurrence of one of the following: (a) Myocardial infarction OR (b) Unstable angina OR (c) Stroke OR (d) Transient ischemic attack (TIA) OR (e) Decompensated heart failure requiring hospitalization OR (f) Class III/IV heart failure. Individual has history or presence of Mobitz Type II second- or third-degree atrioventricular (AV) block, sick sinus syndrome or sino-atrial block, unless individual has a functioning pacemaker. Individual has severe untreated sleep apnea. Concurrent use with a monoamine oxidase (MAO) inhibitor (including but not limited to selegiline, phenelzine and linezolid). Individual is using to treat non-active secondary progressive multiple sclerosis.
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	
Indications	All Medically-accepted Indications.

Y0114_20_114077_I_012

PA Criteria	Criteria Details
Off Label Uses	

Zinplava

Products Affected

• ZINPLAVA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual has confirmed Clostridium difficile infection when the following are met: (a) Passage of three or more loose stools within 24 hours or less AND (b) Positive stool test for toxigenic Clostridium difficile from a stool sample collected not more than 7 days prior to scheduled infusion AND (c) currently receiving antibacterial therapy for Clostridium difficile infection AND (d) Individual is at high risk of Clostridium difficile infection recurrence meeting any one of the following: (1) Individual 65 years of age or older, or (2) history of Clostridium difficile infection in the past 6 months or (3) Immunocompromised state or (4) Severe Clostridium difficile infection at presentation or (4 5) Clostridium difficile ribotype 027.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Zolinza

Products Affected

• ZOLINZA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Zometa

Products Affected

- zoledronic acid intravenous concentrate
- zoledronic acid intravenous solution 4 mg/100ml
- ZOMETA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Bone metastases documented on imaging or bone pain associated with imaging-documented metastases from breast, prostate, lung, kidney, thyroid, or other solid tumors.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	For Breast cancer, prevention of bone loss secondary to adjuvant hormone therapy OR for early stage, premenopausal breast cancer, prevention of bone loss secondary to ovarian dysfunction induced by adjuvant chemotherapy therapy OR Hypercalcemia of malignancy, treatment or Multiple myeloma OR Prevention of osteoporosis during androgen deprivation therapy in prostate cancer.
Indications	All Medically-accepted Indications.
Off Label Uses	

Zulresso

Products Affected

• ZULRESSO

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	18 years of age or older
Prescriber Restrictions	
Coverage Duration	1 year.
Other Criteria	Individual is 6 months postpartum or less AND has a diagnosis of moderate to severe postpartum depression consistent with qualifying score using a standardized screening tool for depression (such as, but not limited to, Hamilton Rating Scale for Depression [HAM-D], Patient Health Questionnaire [PHQ-9], Beck Depression Inventory [BDI], Montgomery-Asberg Depression Rating Scale [MADRS], Edinburgh Postnatal Depression Scale [EPDS]).
Indications	All Medically-accepted Indications.
Off Label Uses	

Zydelig

Products Affected

• ZYDELIG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Zykadia

Products Affected

• ZYKADIA

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Zytiga

Products Affected

- abiraterone acetate
- ZYTIGA ORAL TABLET 250 MG, 500 MG

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	1 year
Other Criteria	Individual is concomitantly receiving a gonadotropin-releasing hormone (GnRH) analog or has had a bilateral orchiectomy.
Indications	All Medically-accepted Indications.
Off Label Uses	

Y0114_20_114077_I_012

Zyvox

Products Affected

- Inezolid oral suspension reconstituted
 YVOX ORAL TABLET
- linezolid oral tablet
- ZYVOX ORAL SUSPENSION RECONSTITUTED

PA Criteria	Criteria Details
Exclusion Criteria	
Required Medical Information	Confirmed vancomycin-resistant enterococcus faecium (VRE) infection. Confirmed methicillin-resistant S. aureus (MRSA) infection AND individual has had a trial and inadequate response or intolerance to or has contraindications to an alternative antibiotic that the organism is susceptible to (depending on manifestation, severity of infection and culture or local sensitivity patterns, examples of alternative antibiotics may include, but are not limited to: vancomycin, TMP-SMX, clindamycin, doxycycline, tetracycline (Based on 2011 IDSA MRSA guideline recommendations)). Individual started treatment with Zyvox in the hospital and requires continued outpatient therapy. Isolates of MRSA have a vancomycin minimum inhibitory concentration (MIC) of greater than 2.
Age Restrictions	
Prescriber Restrictions	
Coverage Duration	30 days
Other Criteria	
Indications	All Medically-accepted Indications.
Off Label Uses	

Index

abiraterone acetate	595
ABRAXANE	1
ABSTRAL	3
ACTEMRA	
ACTEMRA ACTPEN	4
ACTHAR	
ACTIMMUNE	
ACTIQ	
ACTIVELLA	
ADCETRIS	
ADCIRCA	
ADDERALL ORAL TABLET 10 MG, 12.5	
MG, 15 MG, 20 MG, 30 MG, 5 MG, 7.5	
MG	
ADDERALL XR	
ADDYI	
adefovir dipivoxil	
ADEMPAS	14
ADHANSIA XR	
ADZENYS ER	
ADZENYS XR-ODT	
AEMCOLO	
AFINITOR	
AFINITOR DISPERZ	16
AFREZZA INHALATION POWDER 12	
UNIT, 4 & 8 & 12 UNIT, 4 UNIT, 8	,
UNIT, 90 X 4 UNIT & 90X8 UNIT, 90 X	
8 UNIT & 90X12 UNIT	
ALDURAZYME	
ALECENSA	
ALIMTA	
ALIQOPA	
ALLZITAL	
alogliptin benzoate oral tablet 12.5 mg	
25 mg, 6.25 mg	106
alogliptin-metformin hcl	
alogliptin-pioglitazone oral tablet 12.5-	-
15 mg, 12.5-30 mg, 12.5-45 mg, 25-	100
15 mg, 25-30 mg, 25-45 mg	
ALORA	
alosetron hcl	
ALOXI	. 24

ALTOPREV	325
ALTRENO	.499
ALUNBRIG ORAL TABLET 180 MG, 30	
MG, 90 MG	26
ALUNBRIG ORAL TABLET THERAPY	-
PACK	26
ALYQ	. 11
AMABELZ	
AMBIEN	
AMBIEN CR	
ambrisentan	
amitriptyline hcl	
amoxapine	
amphetamine er	
amphetamine sulfate oral tablet 10	
mg, 5 mg	143
amphetamine-dextroamphet er	
amphetamine-dextroamphetamine ora	
tablet 10 mg, 12.5 mg, 15 mg, 20 mg,	
30 mg, 5 mg, 7.5 mg	
AMPYRA	
AMRIX	
AMYTAL SODIUM	193
ANADROL-50	31
ANAFRANIL	.192
ANDRODERM	326
ANDROGEL PUMP	497
ANDROGEL TRANSDERMAL GEL 20.25	
MG/1.25GM (1.62%), 40.5 MG/2.5GM	
(1.62%)	497
ANDROGEL TRANSDERMAL GEL 25	
MG/2.5GM (1%), 50 MG/5GM (1%)	326
ANGELIQ	193
APOKYN	
APTENSIO XR	288
ARALAST NP	25
ARANESP (ALBUMIN FREE)	33
ARCALYST	35
armodafinil oral tablet 150 mg, 200	
mg, 250 mg, 50 mg	
ARMOUR THYROID	193
ARYMO ER	
ARZERRA	36

ASCENIV. 315 ASCOMP-CODEINE 193 atovaquone 286 ATRALIN 499 AUBAGIO 37 AURYXIA. 38 AUSTEDO 39 AVANDIA ORAL TABLET 2 MG, 4 MG. 328 AVASTIN 41 AVEED 490 avita. 499 AVONEX 218 AVONEX PEN 218 AVONEX PREFILLED 218 AVONEX PREFILLED 218 AVSOLA 402 AYVAKIT 42 azacitidine 534 BAFIERTAM 43 BALVERSA ORAL TABLET 3 MG, 4 MG, 5 S MG 44 BANZEL ORAL SUSPENSION 45 BARACLUDE 46 BAVENCIO 47
BELEODAQ
<i>benztropine mesylate oral</i> 193 BEOVU52
BERINERT53
BETASERON
BIJUVA
BIVIGAM
BLINCYTO54
BONJESTA
BORTEZOMIB
BOSULIF ORAL TABLET 100 MG, 400
MG, 500 MG
BOTOX
BRAFTOVI ORAL CAPSULE 50 MG, 75
MG58

BRIVIACT INTRAVENOUS	
BRIVIACT ORAL SOLUTION	. 59
BRIVIACT ORAL TABLET 10 MG, 100	
MG, 25 MG, 50 MG, 75 MG	. 59
BRUKINSA	60
budesonide er	520
BUPAP	193
BUPHENYL	
buprenorphine transdermal patch	
weekly 10 mcg/hr, 15 mcg/hr, 20	
mcg/hr, 5 mcg/hr, 7.5 mcg/hr	321
butalbital-acetaminophen	193
butalbital-apap-caff-cod	193
butalbital-apap-caffeine	193
butalbital-asa-caff-codeine	
butalbital-asa-caffeine	193
butalbital-aspirin-caffeine	193
BUTISOL SODIUM	193
BUTRANS TRANSDERMAL PATCH	
WEEKLY 10 MCG/HR, 15 MCG/HR, 20	
MCG/HR, 5 MCG/HR, 7.5 MCG/HR	321
BYNFEZIA PEN	343
CABOMETYX	62
CALQUENCE	
CAPLYTA	
CAPRELSA ORAL TABLET 100 MG, 300	
MG	
CARBAGLU	
carbinoxamine maleate	193
CARIMUNE NF	315
carisoprodol	
carisoprodol-aspirin	
carisoprodol-aspirin-codeine	
CAYSTON	
CELEBREX	. 68
celecoxib	68
CEQUA	69
CERDELGA	70
CEREZYME	
CHANTIX CONTINUING MONTH PAK	
CHANTIX ORAL TABLET 0.5 MG, 1 MG	
CHANTIX STARTING MONTH PAK	
CHENODAL	
chlordiazepoxide-amitriptyline	192

chlordiazepoxide-clidinium
<i>mg</i> 193
CHLORZOXAZONE ORAL TABLET 375
MG, 750 MG
CHOLBAM
<i>chorionic gonadotropin</i> 75 CIALIS ORAL TABLET 2.5 MG, 5 MG76
CIMZIA
CIMZIA PREFILLED
CIMZIA STARTER KIT77
CINQAIR
CINRYZE82
clemastine fumarate
CLIMARA
CLIMARA PRO193 clindamycin-tretinoin496
clobazam oral suspension
clobazam oral tablet 10 mg, 20 mg351
clomipramine hcl
COMBIPATCH193
COMETRIQ (100 MG DAILY DOSE)83
COMETRIQ (140 MG DAILY DOSE)83
COMETRIQ (60 MG DAILY DOSE)83
CONCERTA ORAL TABLET EXTENDED
RELEASE 18 MG, 27 MG, 36 MG, 54 MG288
CONZIP
COPAXONE SUBCUTANEOUS
SOLUTION PREFILLED SYRINGE 20
MG/ML, 40 MG/ML 84
COPIKTRA85
CORLANOR ORAL SOLUTION
CORLANOR ORAL TABLET
COSENTYX
COSENTYX SENSOREADY (300 MG)88
COSENTYX SENSOREADY PEN
COTELLIC
COTEMPLA XR-ODT
CRESEMBA
CRINONE
CUTAQUIG
CUVITRU

cyclobenzaprine hcl	
cyclobenzaprine hcl er	193
cyproheptadine hcl	193
CYRAMZA	. 93
D.H.E. 45	
DAKLINZA	97
dalfampridine er	30
DALIRESP	
DARZALEX FASPRO	100
DARZALEX INTRAVENOUS SOLUTION	
100 MG/5ML	100
darzalex intravenous solution 400	
mg/20ml	.100
DAURISMO ORAL TABLET 100 MG, 25	
MG	101
deferasirox148,	228
deferasirox granules	228
deferiprone	
DEMEROL	
DEPO-TESTOSTERONE	490
desipramine hcl	
DESOXYN	
dexchlorpheniramine maleate	.193
DICLEGIS	193
diclofenac epolamine	163
diclofenac sodium transdermal gel 3 %	6
-	
DIFICID	
digitek oral tablet 250 mcg	.193
digox oral tablet 250 mcg	193
digoxin injection	193
digoxin oral tablet 250 mcg	193
dihydroergotamine mesylate injection.	
dimethyl fumarate	
dimethyl fumarate starter pack	
diphenhydramine hcl oral	
dipyridamole oral	
disopyramide phosphate	
DIVIGEL	
DOLOPHINE	319
DOPTELET	
DOTTI	
doxepin hcl oral capsule	
doxepin hcl oral concentrate	192

doxepin hcl oral tablet
DOXIL
doxorubicin hcl liposomal105
doxylamine-pyridoxine193
DUAVEE
DUEXIS108
DUOBRII
DUOPA
DUPIXENT SUBCUTANEOUS
SOLUTION PEN-INJECTOR
DUPIXENT SUBCUTANEOUS
SOLUTION PREFILLED SYRINGE 200
MG/1.14ML, 300 MG/2ML 111
DURAGESIC-100
DURAGESIC-12113
DURAGESIC-25113
DURAGESIC-50113
DURAGESIC-75113
DYANAVEL XR27
DYSPORT
EDLUAR193
EGRIFTA115
EGRIFTA SV115
ELAPRASE
ELELYSO
ELESTRIN
ELESTRIN
ELIGARD SUBCUTANEOUS KIT 22.5
MG, 30 MG, 45 MG, 7.5 MG118
ELITEK119
ELZONRIS120
EMBEDA
EMFLAZA121
EMPLICITI
EMSAM124
ENBREL MINI
ENBREL SUBCUTANEOUS SOLUTION
PREFILLED SYRINGE 25 MG/0.5ML, 50
MG/ML
ENBREL SUBCUTANEOUS SOLUTION
RECONSTITUTED
ENBREL SURECLICK
ENHERTU127
entecavir

ENTRESTO	.128
ENTYVIO	. 129
EPCLUSA	. 131
EPIDIOLEX	. 132
EPOGEN	. 133
epoprostenol sodium	. 371
ERAXIS	
ERBITUX	. 137
ergoloid mesylates	. 193
ERIVEDGE	. 138
ERLEADA	. 139
erlotinib hcl oral tablet 100 mg, 150	
mg, 25 mg	.480
ERWINAZE	
ESBRIET ORAL CAPSULE	. 141
ESBRIET ORAL TABLET 267 MG, 801	
MG	.141
ESGIC	. 193
ESTRACE ORAL	. 193
estradiol oral	.193
estradiol transdermal patch twice	
weekly	.193
estradiol transdermal patch weekly	
estradiol-norethindrone acet	
ESTROGEL	.193
ETHYOL	.142
EVAMIST	. 193
EVEKEO ODT ORAL TABLET	
DISPERSIBLE 10 MG, 15 MG, 20 MG,	5
	.143
EVEKEO ORAL TABLET 10 MG, 5 MG.	. 143
EVENITY	
everolimus oral tablet 2.5 mg, 5 mg,	
7.5 mg	16
EVRYŠDI	
EXJADE	. 148
EXONDYS 51	
EXTAVIA	
EYLEA	. 150
ezetimibe-simvastatin	
FABRAZYME	
FARYDAK ORAL CAPSULE 10 MG, 15	
MG, 20 MG	. 152
FASENRA	

FASENRA PEN 153 FASLODEX 155 FEMHRT LOW DOSE 193 FENSOLVI (6 MONTH) 273 fentanyl 113 fentanyl citrate buccal 156, 503 FENTORA 156 FERRIPROX 157 FERRIPROX TWICE-A-DAY 157 FETZIMA ORAL CAPSULE EXTENDED 157
RELEASE 24 HOUR 120 MG, 20 MG, 40
MG, 80 MG
FEXMID
FINTEPLA159
FIORICET193
FIORICET/CODEINE
FIORINAL 193
FIORINAL/CODEINE #3193 FIRAZYR160
FIRAZIR
FIRMAGON (240 MG DOSE)
FIRMAGON SUBCUTANEOUS
SOLUTION RECONSTITUTED 120 MG,
80 MG162
FIRVANQ524
FLEBOGAMMA DIF
FLECTOR
FLOLAN
FORTESTA 326
FULPHILA
fulvestrant155
FUSILEV
<i>fyavolv</i>
GALAFOLD
GAMASTAN
GAMIFANT
GAMMAGARD
GAMMAGARD S/D LESS IGA
GAMMAKED
GAMMAPLEX
GAMUNEX-C

GATTEX171
GAVRETO174
GAZYVA175
GENOTROPIN
GENOTROPIN MINIQUICK
GILENYA176
GILOTRIF178
GIVLAARI179
GLASSIA25
glatiramer acetate subcutaneous
solution prefilled syringe 20 mg/ml, 40
<i>mg/ml</i> 84
glatopa subcutaneous solution prefilled
<i>syringe 20 mg/ml, 40 mg/ml</i> 84
GLEEVEC ORAL TABLET 100 MG, 400
MG180
GLEOSTINE
glyburide micronized oral tablet 1.5
<i>mg, 3 mg, 6 mg</i> 193
glyburide oral tablet 1.25 mg, 2.5 mg,
5 mg
glyburide-metformin oral tablet 1.25-
250 mg, 2.5-500 mg, 5-500 mg193
GLYNASE ORAL TABLET 1.5 MG, 3 MG,
6 MG193
GLYXAMBI
GRANIX
GRASTEK184
guanfacine hcl 193
guanfacine hcl er219
HAEGARDA185
HALAVEN
HARVONI187
HEPSERA188
HETLIOZ 189
HIZENTRA
HORIZANT ORAL TABLET EXTENDED
RELEASE 300 MG, 600 MG190
HUMATROPE311
HUMIRA PEDIATRIC CROHNS START
SUBCUTANEOUS PREFILLED SYRINGE
KIT 40 MG/0.8ML, 40 MG/0.8ML (6
PACK), 80 MG/0.8ML, 80 MG/0.8ML &
40MG/0.4ML

HUMIRA PEN	199
HUMIRA PEN-CD/UC/HS STARTER	
SUBCUTANEOUS PEN-INJECTOR KIT	
40 MG/0.8ML, 80 MG/0.8ML	199
HUMIRA PEN-PS/UV/ADOL HS START	
SUBCUTANEOUS PEN-INJECTOR KIT	
40 MG/0.8ML, 80 MG/0.8ML &	
40MG/0.4ML	199
HUMIRA SUBCUTANEOUS PREFILLED	
SYRINGE KIT 10 MG/0.1ML, 10	
MG/0.2ML, 20 MG/0.2ML, 20	
MG/0.4ML, 40 MG/0.4ML, 40	
MG/0.8ML	199
HUMULIN R U-500 (CONCENTRATED)	202
HUMULIN R U-500 KWIKPEN	202
hydrocodone bitartrate er	321
hydromorphone hcl er	319
hydroxyprogesterone caproate	278
hydroxyzine hcl	
hydroxyzine pamoate	193
HYQVIA	
HYSINGLA ER	
IBRANCE	203
icatibant acetate	
ICLUSIG ORAL TABLET 15 MG, 45 MG	204
IDHIFA ORAL TABLET 100 MG, 50 MG	205
ILARIS	206
ILUMYA	208
imatinib mesylate oral tablet 100 mg,	
400 mg	180
IMBRUVICA ORAL CAPSULE 140 MG,	
70 MG	210
IMBRUVICA ORAL TABLET 140 MG,	
280 MG, 420 MG, 560 MG	
IMFINZI	211
imipramine hcl	
imipramine pamoate	
IMLYGIC	
INCRELEX	
INDOCIN ORAL	
indomethacin	
indomethacin er	
INFLECTRA	402

INGREZZA ORAL CAPSULE 40 MG, 80	
	214
MG INGREZZA ORAL CAPSULE THERAPY	214
	214
PACK	
INLYTA ORAL TABLET 1 MG, 5 MG	
INQOVI	
INREBIC	
INTERMEZZO	
INTUNIV	
INVOKAMET	323
INVOKAMET XR	
INVOKANA ORAL TABLET 100 MG, 300	
MG	323
ISTODAX (OVERFILL)	
ISTURISA ORAL TABLET 1 MG, 10 MG,	,
5 MG	221
itraconazole oral capsule	222
IXEMPRA KIT	
JADENU	
JADENU SPRINKLE	
JAKAFI ORAL TABLET 10 MG, 15 MG,	
20 MG, 25 MG, 5 MG.	229
JETREA	
JEVTANA	
jinteli	
JORNAY PM	
JUBLIA	
JUXTAPID ORAL CAPSULE 10 MG, 20	150
MG, 30 MG, 40 MG, 5 MG, 60 MG	222
JYNARQUE ORAL TABLET	
JYNARQUE ORAL TABLET THERAPY	ررے
PACK	722
KADCYLA	
KADIAN	
KALYDECO ORAL PACKET	
KALYDECO ORAL TABLET	
KANUMA	
KARBINAL ER	
KAZANO	
KERYDIN	
ketorolac tromethamine injection	
ketorolac tromethamine intramuscular	
ketorolac tromethamine oral	193

KEVEYIS KEVZARA KEYTRUDA KHAPZORY KINERET KISQALI (200 MG DOSE) KISQALI (400 MG DOSE) KISQALI (600 MG DOSE) KISQALI FEMARA (400 MG DOSE) KISQALI FEMARA (600 MG DOSE) KISQALI FEMARA (200 MG DOSE)	. 239 241 262 244 246 246 246 246 246 246
KOMBIGLYZE XR ORAL TABLET	
EXTENDED RELEASE 24 HOUR 2.5- 1000 MG, 5-1000 MG, 5-500 MG	106
KORLYM	
KOSELUGO	
KRYSTEXXA	
KUVAN	
KYNAMRO	
KYNMOBI	
KYPROLIS	
LANOXIN INJECTION	
LANOXIN ORAL TABLET 187.5 MCG,	
250 MCG	193
250 MCG lapatinib ditosylate	
lapatinib ditosylate	513
	513 254
lapatinib ditosylate	. 513 254 .256
lapatinib ditosylate LARTRUVO LAZANDA ledipasvir-sofosbuvir LEMTRADA	513 254 .256 .187 .257
lapatinib ditosylate LARTRUVO LAZANDA ledipasvir-sofosbuvir LEMTRADA LENVIMA (10 MG DAILY DOSE)	513 254 .256 .187 .257 .258
lapatinib ditosylate LARTRUVO LAZANDA ledipasvir-sofosbuvir LEMTRADA	513 254 .256 .187 .257 .258
lapatinib ditosylate LARTRUVO LAZANDA ledipasvir-sofosbuvir LEMTRADA LENVIMA (10 MG DAILY DOSE)	513 254 .256 .187 .257 .258 .258
lapatinib ditosylate LARTRUVO LAZANDA ledipasvir-sofosbuvir LEMTRADA LENVIMA (10 MG DAILY DOSE) LENVIMA (12 MG DAILY DOSE)	513 254 .256 .187 .257 .258 .258 .258
lapatinib ditosylate LARTRUVO LAZANDA ledipasvir-sofosbuvir LEMTRADA LENVIMA (10 MG DAILY DOSE) LENVIMA (12 MG DAILY DOSE) LENVIMA (14 MG DAILY DOSE) LENVIMA (18 MG DAILY DOSE) LENVIMA (20 MG DAILY DOSE)	. 513 . 254 . 256 . 187 . 257 . 258 . 258 . 258 . 258 . 258
lapatinib ditosylate LARTRUVO LAZANDA ledipasvir-sofosbuvir LEMTRADA LENVIMA (10 MG DAILY DOSE) LENVIMA (12 MG DAILY DOSE) LENVIMA (14 MG DAILY DOSE) LENVIMA (18 MG DAILY DOSE) LENVIMA (20 MG DAILY DOSE) LENVIMA (24 MG DAILY DOSE)	513 254 .256 .187 .258 .258 .258 .258 .258 .258 .258
lapatinib ditosylate LARTRUVO LAZANDA ledipasvir-sofosbuvir LEMTRADA LENVIMA (10 MG DAILY DOSE) LENVIMA (12 MG DAILY DOSE) LENVIMA (14 MG DAILY DOSE) LENVIMA (18 MG DAILY DOSE) LENVIMA (20 MG DAILY DOSE) LENVIMA (24 MG DAILY DOSE)	.513 .254 .256 .187 .257 .258 .258 .258 .258 .258 .258 .258 .258
lapatinib ditosylate LARTRUVO LAZANDA ledipasvir-sofosbuvir LEMTRADA LENVIMA (10 MG DAILY DOSE) LENVIMA (12 MG DAILY DOSE) LENVIMA (14 MG DAILY DOSE) LENVIMA (18 MG DAILY DOSE) LENVIMA (20 MG DAILY DOSE) LENVIMA (24 MG DAILY DOSE) LENVIMA (4 MG DAILY DOSE) LENVIMA (8 MG DAILY DOSE)	513 254 .256 .187 .258 .258 .258 .258 .258 .258 .258 .258
lapatinib ditosylate LARTRUVO LAZANDA ledipasvir-sofosbuvir LEMTRADA LENVIMA (10 MG DAILY DOSE) LENVIMA (12 MG DAILY DOSE) LENVIMA (14 MG DAILY DOSE) LENVIMA (18 MG DAILY DOSE) LENVIMA (20 MG DAILY DOSE) LENVIMA (24 MG DAILY DOSE) LENVIMA (4 MG DAILY DOSE) LENVIMA (8 MG DAILY DOSE) LENVIMA (8 MG DAILY DOSE)	513 254 .256 .187 .258 .258 .258 .258 .258 .258 .258 .258
lapatinib ditosylate LARTRUVO LAZANDA ledipasvir-sofosbuvir LEMTRADA LENVIMA (10 MG DAILY DOSE) LENVIMA (12 MG DAILY DOSE) LENVIMA (12 MG DAILY DOSE) LENVIMA (14 MG DAILY DOSE) LENVIMA (18 MG DAILY DOSE) LENVIMA (20 MG DAILY DOSE) LENVIMA (24 MG DAILY DOSE) LENVIMA (4 MG DAILY DOSE) LENVIMA (8 MG DAILY DOSE) LETAIRIS LEUKINE	.513 .254 .256 .187 .257 .258 .258 .258 .258 .258 .258 .258 .258
lapatinib ditosylate LARTRUVO LAZANDA ledipasvir-sofosbuvir LEMTRADA LENVIMA (10 MG DAILY DOSE) LENVIMA (12 MG DAILY DOSE) LENVIMA (14 MG DAILY DOSE) LENVIMA (18 MG DAILY DOSE) LENVIMA (20 MG DAILY DOSE) LENVIMA (24 MG DAILY DOSE) LENVIMA (24 MG DAILY DOSE) LENVIMA (4 MG DAILY DOSE) LENVIMA (8 MG DAILY DOSE) LETAIRIS LEUKINE LEUKINE	513 254 .256 .187 .258 .258 .258 .258 .258 .258 .258 .258
lapatinib ditosylate LARTRUVO LAZANDA ledipasvir-sofosbuvir LEMTRADA LENVIMA (10 MG DAILY DOSE) LENVIMA (12 MG DAILY DOSE) LENVIMA (12 MG DAILY DOSE) LENVIMA (14 MG DAILY DOSE) LENVIMA (18 MG DAILY DOSE) LENVIMA (20 MG DAILY DOSE) LENVIMA (24 MG DAILY DOSE) LENVIMA (4 MG DAILY DOSE) LENVIMA (8 MG DAILY DOSE) LEUKINE LEUKINE leuprolide acetate levoleucovorin calcium	.513 .254 .256 .187 .258 .258 .258 .258 .258 .258 .258 .258
lapatinib ditosylate LARTRUVO LAZANDA ledipasvir-sofosbuvir LEMTRADA LENVIMA (10 MG DAILY DOSE) LENVIMA (12 MG DAILY DOSE) LENVIMA (14 MG DAILY DOSE) LENVIMA (18 MG DAILY DOSE) LENVIMA (20 MG DAILY DOSE) LENVIMA (24 MG DAILY DOSE) LENVIMA (24 MG DAILY DOSE) LENVIMA (4 MG DAILY DOSE) LENVIMA (8 MG DAILY DOSE) LENVIMA (8 MG DAILY DOSE) LENVIMA (8 MG DAILY DOSE) LENVIMA (8 MG DAILY DOSE) LEUKINE LEUKINE leuprolide acetate levoleucovorin calcium pf	513 254 .256 .187 .258 .258 .258 .258 .258 .258 .258 .258
lapatinib ditosylate LARTRUVO LAZANDA ledipasvir-sofosbuvir LEMTRADA LENVIMA (10 MG DAILY DOSE) LENVIMA (12 MG DAILY DOSE) LENVIMA (12 MG DAILY DOSE) LENVIMA (14 MG DAILY DOSE) LENVIMA (18 MG DAILY DOSE) LENVIMA (20 MG DAILY DOSE) LENVIMA (24 MG DAILY DOSE) LENVIMA (4 MG DAILY DOSE) LENVIMA (8 MG DAILY DOSE) LEUKINE LEUKINE leuprolide acetate levoleucovorin calcium	513 254 256 187 257 258 258 258 258 258 258 258 258 258 258

LIBTAYO	263
LICART	
lidocaine external ointment	264
lidocaine external patch	265
lidocaine hcl external cream	264
lidocaine hcl external solution	
lidocaine hcl mouth/throat	264
lidocaine pak	
LIDODERM	
linezolid oral suspension reconstituted	596
linezolid oral tablet	
LONSURF	266
LOPREEZA	193
LORBRENA ORAL TABLET 100 MG, 25	
MG	267
LORZONE	193
LOTRONEX	268
LUCENTIS	269
LUMIZYME	270
LUMOXITI	271
LUPANETA PACK COMBINATION KIT	
11.25 & 5 MG, 3.75 & 5 MG	272
LUPRON DEPOT (1-MONTH)	273
LUPRON DEPOT (3-MONTH)	273
LUPRON DEPOT (4-MONTH)	273
LUPRON DEPOT (6-MONTH)	273
LUPRON DEPOT-PED (1-MONTH)	273
LUPRON DEPOT-PED (3-MONTH)	
LYNPARZA	276
LYRICA CR	
MAKENA INTRAMUSCULAR	
MAKENA SUBCUTANEOUS	278
MAVENCLAD (10 TABS)	
MAVENCLAD (4 TABS)	279
MAVENCLAD (5 TABS)	
MAVENCLAD (6 TABS)	279
MAVENCLAD (7 TABS)	
MAVENCLAD (8 TABS)	
MAVENCLAD (9 TABS)	
MAVYRET	
MAYZENT ORAL TABLET 0.25 MG, 2	
MG	
MEGACE ES	193

megestrol acetate oral suspension 40	
mg/ml, 400 mg/10ml	282
megestrol acetate oral suspension 625	
mg/5ml	
megestrol acetate oral tablet	
MEKINIST ORAL TABLET 0.5 MG, 2 MG	
MEKTOVI	
memantine hcl er	
memantine hcl oral solution	
memantine hcl oral tablet 10 mg, 28 x	
5 mg & 21 x 10 mg, 5 mg	
MENEST	
MENOSTAR	
meperidine hcl injection	
meperidine hcl oral solution	
meperidine hcl oral tablet	
meprobamate	
MEPRON	
MEPSEVII	
metadate er	
METAXALL	
metaxalone	
methadone hcl oral tablet	
methamphetamine hcl	
methocarbamol	193
methyldopa	
methyldopa-hydrochlorothiazide	
METHYLIN ORAL SOLUTION 10	
MG/5ML, 5 MG/5ML	288
methylphenidate hcl er (cd)	288
methylphenidate hcl er (la) oral	
capsule extended release 24 hour 10	
mg, 20 mg, 30 mg, 40 mg, 60 mg	288
methylphenidate hcl er (xr)	
methylphenidate hcl er oral tablet	
extended release 10 mg, 18 mg, 20	
mg, 27 mg, 36 mg, 54 mg, 72 mg	288
methylphenidate hcl er oral tablet	
extended release 24 hour 18 mg, 27	
mg, 36 mg, 54 mg	288
methylphenidate hcl oral solution 10	
mg/5ml, 5 mg/5ml	
methylphenidate hcl oral tablet	288

methylphenidate hcl oral tablet	
chewable 10 mg, 2.5 mg, 5 mg	288
miglustat	
MIMVEY	
MIMVEY LO	193
MINIVELLE	
MIRCERA	
modafinil oral tablet 100 mg, 200 mg.	
MORPHABOND ER	
morphine sulfate er beads	
morphine sulfate er oral capsule	515
extended release 24 hour	319
morphine sulfate er oral tablet	515
extended release 100 mg, 15 mg, 200	
mg, 30 mg, 60 mg	319
MOTOFEN	
MOZOBIL	
MS CONTIN ORAL TABLET EXTENDED	255
RELEASE 100 MG, 15 MG, 200 MG, 30	
MG, 60 MG	310
MULPLETA	
MVASI	
MYALEPT	
MYCAPSSA	
MYDAYIS	
MYLOTARG	
MYOBLOC	
NAGLAZYME	
NAMENDA ORAL TABLET 10 MG, 5 MG	
NAMENDA TITRATION PAK	200
NAMENDA XR	
NAMENDA XR TITRATION PACK	
naproxen-esomeprazole	
NATPARA	
NEMBUTAL	
NERLYNX	
NESINA ORAL TABLET 12.5 MG, 25	201
MG, 6.25 MG	106
NEULASTA	
NEULASTA ONPRO	
NEUPOGEN	
NEOPOGEN	
nifedipine	193

NINLARO	. 303
nitisinone	.360
NITYR	
NIVESTYM	. 308
NORDITROPIN FLEXPRO	.196
norethindrone-eth estradiol	.193
norgesic forte	
NORPACE	193
NORPACE CR	. 193
NORPRAMIN	
NORTHERA ORAL CAPSULE 100 MG,	
200 MG, 300 MG	. 305
nortriptyline hcl	
NOURIANZ ORAL TABLET 20 MG, 40	
MG	.306
NOVAREL	75
NOXAFIL ORAL	307
np thyroid	
NPLATE	
NUBEQA	. 330
NUCALA	
NUCYNTA ER	319
NUEDEXTA	
NULOJIX	
NUPLAZID	336
NURTEC	
NUTROPIN AQ NUSPIN 10	. 311
NUTROPIN AQ NUSPIN 20	. 311
NUTROPIN AQ NUSPIN 5	
NUVIGIL ORAL TABLET 150 MG, 200	
MG, 250 MG, 50 MG	338
OCALIVA	
OCREVUS	342
OCTAGAM	.223
octreotide acetate	343
ODACTRA	
ODOMZO	. 346
OFEV	.347
OLUMIANT	
OMNITROPE	
ONCASPAR	
ONFI ORAL SUSPENSION	.351
ONFI ORAL TABLET 10 MG, 20 MG	
·	

ONGLYZA ORAL TABLET 2.5 MG, 5 MG	
ONPATTRO OPDIVO OPSUMIT	352 353 355
ORALAIR	
ORALAIR CHILDRENS STARTER PACK.	
ORENCIA CLICKJECT	
ORENCIA INTRAVENOUS	357
ORENCIA SUBCUTANEOUS SOLUTION	
PREFILLED SYRINGE 125 MG/ML, 50	
MG/0.4ML, 87.5 MG/0.7ML	357
ORENITRAM	359
ORFADIN	
ORILISSA ORAL TABLET 150 MG, 200	
MG	361
ORKAMBI ORAL PACKET	363
ORKAMBI ORAL TABLET	363
orphenadrine citrate	193
orphenadrine citrate er	193
orphenadrine-asa-caffeine	193
orphenadrine-aspirin-caffeine	193
ORPHENGESIC FORTE	193
OSENI ORAL TABLET 12.5-15 MG,	
12.5-30 MG, 12.5-45 MG, 25-15 MG,	
25-30 MG, 25-45 MG	106
OTEZLA ORAL TABLET	364
OTEZLA ORAL TABLET THERAPY PACK	(
	364
oxandrolone oral tablet 10 mg, 2.5 mg	366
oxycodone hcl er	367
OXYCONTIN	
oxymorphone hcl er	319
OZURDEX	369
PADCEV	370
palonosetron hcl	. 24
PALYNZIQ	373
PAMELOR	192
PANZYGA	315
PEMAZYRE	376
pentazocine-naloxone hcl	193
pentobarbital sodium	192
PERJETA	377
perphenazine-amitriptyline	192

PHENADOZ1PHENERGAN1phenobarbital oral elixir1phenobarbital oral solution1phenobarbital oral tablet 100 mg, 15	193 192
mg, 16.2 mg, 30 mg, 32.4 mg, 60 mg, 64.8 mg, 97.2 mg	192 379
pimecrolimus	380 380 380
PLEGRIDY STARTER PACK	218 381
posaconazole	882 193 .75
PREMARIN ORAL	193 193
PROBUPHINE IMPLANT KIT	884 193 133
PROLASTIN-C PROLIA	886
PROMACTA ORAL TABLET 12.5 MG, 25 MG, 50 MG, 75 MG	193
promethazine-phenylephrine	193 193 193
PROTOPIC	192

PURIXAN	391
QINLOCK	. 392
QTERN	324
QUALAQUIN	393
QUILLICHEW ER ORAL TABLET	
CHEWABLE EXTENDED RELEASE 20	
MG, 30 MG, 40 MG	
QUILLIVANT XR	288
quinine sulfate	393
RAGWITEK	394
RAVICTI	395
RAYOS	396
REBIF	
REBIF REBIDOSE	314
REBIF REBIDOSE TITRATION PACK	.314
REBIF TITRATION PACK	
REBLOZYL	397
RECLAST	399
REFISSA	. 499
REGRANEX	400
RELEXXII	
RELISTOR ORAL	
RELISTOR SUBCUTANEOUS SOLUTIO	N
12 MG/0.6ML, 8 MG/0.4ML	
REMICADE	.402
REMODULIN	
RENFLEXIS	402
REPATHA	
KEPAI NA	.406
REPATHA PUSHTRONEX SYSTEM	406
REPATHA PUSHTRONEX SYSTEM REPATHA SURECLICK	406
REPATHA PUSHTRONEX SYSTEM REPATHA SURECLICK RETACRIT	406
REPATHA PUSHTRONEX SYSTEM REPATHA SURECLICK RETACRIT RETEVMO ORAL CAPSULE 40 MG, 80	406 .406 .133
REPATHA PUSHTRONEX SYSTEM REPATHA SURECLICK RETACRIT RETEVMO ORAL CAPSULE 40 MG, 80 MG	406 .406 .133 .408
REPATHA PUSHTRONEX SYSTEM REPATHA SURECLICK RETACRIT RETEVMO ORAL CAPSULE 40 MG, 80 MG RETIN-A	406 .406 .133 .408 .499
REPATHA PUSHTRONEX SYSTEM REPATHA SURECLICK RETACRIT RETEVMO ORAL CAPSULE 40 MG, 80 MG RETIN-A RETIN-A MICRO	406 .406 .133 .408 .499 .499
REPATHA PUSHTRONEX SYSTEM REPATHA SURECLICK RETACRIT RETEVMO ORAL CAPSULE 40 MG, 80 MG RETIN-A RETIN-A MICRO RETIN-A MICRO PUMP	406 .406 .133 .408 .499 .499 .499
REPATHA PUSHTRONEX SYSTEM REPATHA SURECLICK RETACRIT RETEVMO ORAL CAPSULE 40 MG, 80 MG RETIN-A RETIN-A MICRO RETIN-A MICRO PUMP RETISERT	406 .406 .133 .408 .499 .499 .499 .499
REPATHA PUSHTRONEX SYSTEM REPATHA SURECLICK RETACRIT RETEVMO ORAL CAPSULE 40 MG, 80 MG RETIN-A RETIN-A MICRO RETIN-A MICRO PUMP RETISERT REVATIO INTRAVENOUS	406 .406 .133 .408 .499 .499 .499 .499
REPATHA PUSHTRONEX SYSTEM REPATHA SURECLICK RETACRIT RETEVMO ORAL CAPSULE 40 MG, 80 MG RETIN-A RETIN-A MICRO RETIN-A MICRO PUMP RETISERT REVATIO INTRAVENOUS REVATIO ORAL SUSPENSION	406 .406 .133 .408 .499 .499 .499 .499 .409 .411
REPATHA PUSHTRONEX SYSTEM REPATHA SURECLICK RETACRIT RETEVMO ORAL CAPSULE 40 MG, 80 MG RETIN-A RETIN-A MICRO RETIN-A MICRO PUMP RETISERT REVATIO INTRAVENOUS REVATIO ORAL SUSPENSION RECONSTITUTED	406 .406 .133 .408 .499 .499 .499 .499 .409 411 .410
REPATHA PUSHTRONEX SYSTEM REPATHA SURECLICK RETACRIT RETEVMO ORAL CAPSULE 40 MG, 80 MG RETIN-A RETIN-A MICRO RETIN-A MICRO PUMP RETISERT REVATIO INTRAVENOUS REVATIO INTRAVENOUS REVATIO ORAL SUSPENSION RECONSTITUTED REVATIO ORAL TABLET	406 .406 .133 .408 .499 .499 .499 .499 .409 411 .410
REPATHA PUSHTRONEX SYSTEM REPATHA SURECLICK RETACRIT RETEVMO ORAL CAPSULE 40 MG, 80 MG RETIN-A RETIN-A MICRO RETIN-A MICRO PUMP RETISERT REVATIO INTRAVENOUS REVATIO ORAL SUSPENSION RECONSTITUTED REVATIO ORAL TABLET REVATIO ORAL TABLET REVLIMID ORAL CAPSULE 10 MG, 15	406 .406 .133 .408 .499 .499 .499 .499 .409 411 .410 .410
REPATHA PUSHTRONEX SYSTEM REPATHA SURECLICK RETACRIT RETEVMO ORAL CAPSULE 40 MG, 80 MG RETIN-A RETIN-A MICRO RETIN-A MICRO PUMP RETISERT REVATIO INTRAVENOUS REVATIO INTRAVENOUS REVATIO ORAL SUSPENSION RECONSTITUTED REVATIO ORAL TABLET	406 .406 .133 .408 .499 .499 .499 .499 .409 411 .410 .410

REYVOW ORAL TABLET 100 MG, 50	
MG43	14
ribavirin inhalation53	39
RINVOQ	
RITALIN	
RITALIN LA ORAL CAPSULE	
EXTENDED RELEASE 24 HOUR 10 MG,	
20 MG, 30 MG, 40 MG	28
ROBAXIN	
ROBAXIN-750	
romidepsin	
ROZLYTREK ORAL CAPSULE 100 MG,	10
200 MG	17
RUBRACA ORAL TABLET 200 MG, 250	L/
,	10
MG, 300 MG42 RUCONEST	
RUZURGI	
RYCLORA	
RYDAPT42	
RYVENT	
SABRIL	
SAIZEN	
SAIZENPREP	
SAMSCA ORAL TABLET 15 MG, 30 MG 42	
SANCUSO	
SANDOSTATIN	
SANDOSTATIN LAR DEPOT	
sapropterin dihydrochloride25	
SARCLISA	24
SECONAL19) 3
SEGLUROMET	23
SEROSTIM	1
SIGNIFOR42	25
SIGNIFOR LAR	26
SIKLOS42	27
sildenafil citrate intravenous41	1
sildenafil citrate oral suspension	
reconstituted4	10
sildenafil citrate oral tablet 20 mg4	
SILENOR	
SILIQ	
SIMPONI ARIA	

SIMPONI SUBCUTANEOUS SOLUTION	
AUTO-INJECTOR 100 MG/ML, 50	
MG/0.5ML	430
SIMPONI SUBCUTANEOUS SOLUTION	
PREFILLED SYRINGE 100 MG/ML, 50	
MG/0.5ML	430
SIRTURO	435
SIVEXTRO INTRAVENOUS	436
SIVEXTRO ORAL	436
SKELAXIN	193
SKYRIZI (150 MG DOSE)	438
sodium phenylbutyrate	
sofosbuvir-velpatasvir	
SOLIRIS	
SOMA	
SOMATULINE DEPOT	
SOMAVERT	
SOVALDI	
SPINRAZA	
SPORANOX ORAL CAPSULE	
SPORANOX PULSEPAK	
SPRAVATO (56 MG DOSE)	
SPRAVATO (84 MG DOSE)	451
SPRITAM ORAL TABLET	
DISINTEGRATING SOLUBLE 1000 MG,	
250 MG, 500 MG, 750 MG	
SPRYCEL	453
STEGLATRO	
STEGLUJAN	
STELARA INTRAVENOUS	
STELARA SUBCUTANEOUS	
STIVARGA	
STRENSIQ	
SUBSYS	
SUPPRELIN LA.	460
SUTENT ORAL CAPSULE 12.5 MG, 25	
MG, 37.5 MG, 50 MG	
SYLATRON	
SYLVANT	
SYMDEKO	
SYMLINPEN 120	
SYMLINPEN 60	
SYMPAZAN ORAL FILM 10 MG, 20 MG,	
5 MG	351

SYNAGIS 460	6
SYNAREL	
SYNRIBO470	0
TABRECTA	1
-	
tacrolimus external	U
tadalafil (pah)1	1
tadalafil oral tablet 2.5 mg, 5 mg	
TAFINLAR	3
TAGRISSO ORAL TABLET 40 MG, 80	
MG474	Δ
	т Е
TAKHZYRO	5
TALTZ	7
TALZENNA ORAL CAPSULE 0.25 MG, 1	
•	~
MG479	9
TARCEVA ORAL TABLET 100 MG, 150	
MG, 25 MG 480	n
TARGRETIN EXTERNAL48	
TARGRETIN ORAL	1
TASIGNA	
TASMAR	
tavaborole	8
TAVALISSE	4
tazarotene	
TAZORAC	5
TAZVERIK480	6
TECENTRIQ INTRAVENOUS SOLUTION	-
	_
1200 MG/20ML, 840 MG/14ML 48	
TECFIDERA488	
	8
1F(-SFI)I 489	
TEGSEDI	9
temsirolimus	9 0
	9 0
temsirolimus	9 0 3
temsirolimus	9 0 3 4
temsirolimus	9 0 3 4 6
temsirolimus	9 0 3 4 6
temsirolimus500TENCON193teriparatide (recombinant)164TESTIM320TESTONE CIK490	9 0 3 4 6 0
temsirolimus500TENCON193teriparatide (recombinant)164TESTIM320TESTONE CIK490testosterone cypionate490	9 0 3 4 6 0
temsirolimus500TENCON193teriparatide (recombinant)164TESTIM320TESTONE CIK490testosterone cypionate490testosterone enanthate490	9 0 3 4 6 0 0
temsirolimus500TENCON193teriparatide (recombinant)164TESTIM320TESTONE CIK490testosterone cypionate490testosterone enanthate490testosterone propionate490	9 0 3 4 6 0 0
temsirolimus500TENCON193teriparatide (recombinant)164TESTIM320TESTONE CIK490testosterone cypionate490testosterone enanthate490testosterone propionate490	9 0 3 4 6 0 0
temsirolimus500TENCON193teriparatide (recombinant)164TESTIM320TESTONE CIK490testosterone cypionate490testosterone enanthate490testosterone propionate490testosterone propionate490testosterone transdermal gel 1.62 %,	9 0 3 4 6 0 0
temsirolimus500TENCON193teriparatide (recombinant)164TESTIM320TESTONE CIK490testosterone cypionate490testosterone enanthate490testosterone propionate490testosterone transdermal gel 1.62 %,20.25 mg/1.25gm (1.62%), 20.25	9 0 3 4 6 0 0
temsirolimus500TENCON193teriparatide (recombinant)164TESTIM320TESTONE CIK490testosterone cypionate490testosterone enanthate490testosterone propionate490testosterone transdermal gel 1.62 %,20.25 mg/1.25gm (1.62%), 20.25mg/act (1.62%), 40.5 mg/2.5gm	9 0 3 4 6 0 0 0
temsirolimus500TENCON193teriparatide (recombinant)164TESTIM320TESTONE CIK490testosterone cypionate490testosterone enanthate490testosterone propionate490testosterone transdermal gel 1.62 %,20.25 mg/1.25gm (1.62%), 20.25mg/act (1.62%), 40.5 mg/2.5gm	9 0 3 4 6 0 0 0
temsirolimus 500 TENCON 191 teriparatide (recombinant) 164 TESTIM 320 TESTONE CIK 490 testosterone cypionate 490 testosterone enanthate 490 testosterone transdermal gel 1.62 %, 20.25 mg/1.25gm (1.62%), 20.25 mg/act (1.62%), 40.5 mg/2.5gm 491	9 0 3 4 6 0 0 0
temsirolimus 500 TENCON 191 teriparatide (recombinant) 164 TESTIM 320 TESTONE CIK 490 testosterone cypionate 490 testosterone enanthate 490 testosterone propionate 490 testosterone transdermal gel 1.62 %, 20.25 mg/1.25gm (1.62%), 20.25 mg/act (1.62%), 40.5 mg/2.5gm 491 testosterone transdermal gel 10 491	9 0 3 4 6 0 0 0
temsirolimus 500 TENCON 191 teriparatide (recombinant) 164 TESTIM 320 TESTONE CIK 490 testosterone cypionate 490 testosterone enanthate 490 testosterone propionate 490 testosterone transdermal gel 1.62 %, 20.25 mg/1.25gm (1.62%), 20.25 mg/act (1.62%), 40.5 mg/2.5gm 491 testosterone transdermal gel 10 491 mg/act (2%), 12.5 mg/act (1%), 25 491	9 0 3 4 6 0 0 0 7
temsirolimus 500 TENCON 191 teriparatide (recombinant) 164 TESTIM 320 TESTONE CIK 490 testosterone cypionate 490 testosterone enanthate 490 testosterone propionate 490 testosterone transdermal gel 1.62 %, 20.25 mg/1.25gm (1.62%), 20.25 mg/act (1.62%), 40.5 mg/2.5gm 491 testosterone transdermal gel 10 491	9 0 3 4 6 0 0 0 7

testosterone transdermal solution	326
tetrabenazine oral tablet 12.5 mg, 25	
<i>mg</i>	558
THALOMID ORAL CAPSULE 100 MG,	
150 MG, 200 MG, 50 MG	<u>4</u> 03
THIOLA	
THIOLA	
	494
thyroid oral tablet 120 mg, 15 mg, 30	
<i>mg, 60 mg, 90 mg</i>	193
THYROID ORAL TABLET 65 MG	193
TIBSOVO	495
TOFRANIL	192
tolcapone	
tolsura	
tolvaptan	
TORISEL	
TRACLEER ORAL TABLET	
TRACLEER ORAL TABLET SOLUBLE	
tramadol hcl er	
tramadol hcl er (biphasic)	319
TRELSTAR MIXJECT INTRAMUSCULAR	
SUSPENSION RECONSTITUTED 11.25	
MG, 22.5 MG, 3.75 MG	504
TRÉMFYA	
treprostinil	
tretinoin (emollient)	
tretinoin external	400
tretinoin microsphere	
tretinoin microsphere pump	
trihexyphenidyl hcl	193
TRIJARDY XR ORAL TABLET	
EXTENDED RELEASE 24 HOUR 10-5-	
1000 MG, 12.5-2.5-1000 MG, 25-5-	
1000 MG, 5-2.5-1000 MG	324
TRIKAFTA	507
TRIPTODUR	508
TRODELVY	509
TROGARZO	
TUKYSA	
TURALIO	
TYKERB	
	212
TYMLOS	E 1 4
TYSABRI TYVASO	516

TYVASO REFILL	
TYVASO STARTER	
UBRELVY	. 519
UCERIS ORAL	. 520
UDENYCA	374
ULTOMIRIS	
UPTRAVI ORAL TABLET	
UPTRAVI ORAL TABLET THERAPY	. JZZ
PACK	522
VALCHLOR	
VALCHLOR.	
VANADON VANATOL LQ	
VANATOL LQ	
VANATOL S	
	.324
VANCOCIN HCL ORAL CAPSULE 125	F24
MG, 250 MG	.524
vancomycin hcl oral capsule 125 mg,	FQ 4
250 mg	524
vancomycin hcl oral solution	
reconstituted	
VECTIBIX	
VELCADE	
VELETRI	
VELTIN	
VEMLIDY	
VENCLEXTA ORAL TABLET 10 MG, 10	
MG, 50 MG	
VENCLEXTA STARTING PACK	. 528
VENTAVIS	.529
VERZENIO	. 530
VFEND	.531
VIBATIV	. 532
VIBERZI	533
VIDAZA	. 534
VIEKIRA PAK	
vigabatrin	
vigadrone	
VIMIZIM	
VIMOVO	
VIRAZOLE	
VISTARIL	
VITRAKVI ORAL CAPSULE 100 MG, 25	
MG	
VITRAKVI ORAL SOLUTION	

VIZIMPRO ORAL TABLET 15 MG, 30 MG, 45 MG. 541 VOGELXO 326 VOGELXO PUMP. 326 voriconazole oral. 531 VOSEVI. 542 VOTRIENT. 544 VPRIV. 172 VTOL LQ. 193 VUMERITY 545 VUMERITY (STARTER) 546 VYLEESI 548 VYNDAMAX. 472 VYNDAQEL 472 VYONDYS 53 550 VYTORIN. 551 VYVANSE 552 WAKIX. 553 XALKORI 556 XELJANZ XR 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 56 XEOMIN. 56 XEOMIN 56 XERMELO 559
MG, 45 MG. 541 VOGELXO 326 VOGELXO PUMP. 326 voriconazole oral. 531 VOSEVI. 542 VOTRIENT 544 VPRIV. 172 VTOL LQ. 193 VUMERITY 545 VUMERITY (STARTER) 545 VYLEESI 546 VYNDAQEL 472 VYNDAQEL 472 VYONDYS 53 550 VYTORIN 551 VYVANSE 552 WAKIX 553 XALKORI 556 XELJANZ XR 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 568 XEOMIN 568
VOGELXO 326 VOGELXO PUMP 326 voriconazole oral 531 VOSEVI 542 VOTRIENT 544 VPRIV 172 VTOL LQ 193 VUMERITY 545 VUMERITY (STARTER) 545 VYLEESI 548 VYNDAMAX 472 VYONDYS 53 550 VYTORIN 551 VYVANSE 552 WAKIX 553 XALKORI 556 XELJANZ 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 568 XEOMIN 568
voriconazole oral. 531 VOSEVI. 542 VOTRIENT. 544 VPRIV. 172 VTOL LQ. 193 VUMERITY 545 VUMERITY (STARTER). 545 VYEPTI. 546 VYLEESI. 548 VYNDAMAX. 472 VYNDAQEL. 472 VYONDYS 53. 550 VYTORIN. 551 VYANSE. 552 WAKIX. 553 XALKORI. 555 XELJANZ XR. 556 XEMBIFY. 315 XENAZINE ORAL TABLET 12.5 MG, 25 56 XEOMIN. 56
voriconazole oral. 531 VOSEVI. 542 VOTRIENT. 544 VPRIV. 172 VTOL LQ. 193 VUMERITY 545 VUMERITY (STARTER). 545 VYEPTI. 546 VYLEESI. 548 VYNDAMAX. 472 VYNDAQEL. 472 VYONDYS 53. 550 VYTORIN. 551 VYANSE. 552 WAKIX. 553 XALKORI. 555 XELJANZ XR. 556 XEMBIFY. 315 XENAZINE ORAL TABLET 12.5 MG, 25 56 XEOMIN. 56
VOTRIENT 544 VPRIV 172 VTOL LQ 193 VUMERITY 545 VUMERITY (STARTER) 546 VYEPTI 546 VYLEESI 548 VYNDAMAX 472 VYONDYS 53 550 VYTORIN 551 VYVANSE 552 WAKIX 553 XALKORI 555 XELJANZ 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 56 XEOMIN 56
VPRIV. 172 VTOL LQ. 193 VUMERITY 545 VUMERITY (STARTER) 546 VYEPTI. 546 VYLEESI 548 VYNDAMAX 472 VYNDAQEL 472 VYONDYS 53 550 VYTORIN 551 VYVANSE 552 WAKIX 553 XALKORI 555 XELJANZ XR 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 56 XEOMIN 56
VPRIV. 172 VTOL LQ. 193 VUMERITY 545 VUMERITY (STARTER) 546 VYEPTI. 546 VYLEESI 548 VYNDAMAX 472 VYNDAQEL 472 VYONDYS 53 550 VYTORIN 551 VYVANSE 552 WAKIX 553 XALKORI 555 XELJANZ XR 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 56 XEOMIN 56
VUMERITY 545 VUMERITY (STARTER) 545 VYEPTI 546 VYLEESI 548 VYNDAMAX 472 VYNDAQEL 472 VYONDYS 53 550 VYTORIN 551 VYVANSE 552 WAKIX 553 XALKORI 555 XELJANZ 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 558 XEOMIN 568 XEOMIN 568
VUMERITY 545 VUMERITY (STARTER) 545 VYEPTI 546 VYLEESI 548 VYNDAMAX 472 VYNDAQEL 472 VYONDYS 53 550 VYTORIN 551 VYVANSE 552 WAKIX 553 XALKORI 555 XELJANZ 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 558 XEOMIN 568 XEOMIN 568
VYEPTI
VYEPTI
VYNDAMAX 472 VYNDAQEL 472 VYONDYS 53 550 VYTORIN 551 VYVANSE 552 WAKIX 553 XALKORI 555 XELJANZ 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 558 XEOMIN 568
VYNDAQEL 472 VYONDYS 53 550 VYTORIN 551 VYVANSE 552 WAKIX 553 XALKORI 555 XELJANZ 556 XELJANZ XR 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 558 XEOMIN 568
VYONDYS 53 550 VYTORIN 551 VYVANSE 552 WAKIX 553 XALKORI 555 XELJANZ 556 XELJANZ XR 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 558 XEOMIN 568
VYONDYS 53 550 VYTORIN 551 VYVANSE 552 WAKIX 553 XALKORI 555 XELJANZ 556 XELJANZ XR 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 558 XEOMIN 568
VYVANSE 552 WAKIX 553 XALKORI 555 XELJANZ 556 XELJANZ XR 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 558 XEOMIN 568
WAKIX 553 XALKORI 555 XELJANZ 556 XELJANZ XR 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 558 MG 558 XEOMIN 56
XALKORI 555 XELJANZ 556 XELJANZ XR 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 MG MG 558 XEOMIN 56
XELJANZ 556 XELJANZ XR 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 MG MG 558 XEOMIN 56
XELJANZ 556 XELJANZ XR 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 MG MG 558 XEOMIN 56
XELJANZ XR 556 XEMBIFY 315 XENAZINE ORAL TABLET 12.5 MG, 25 MG MG 558 XEOMIN 56
XENAZINE ORAL TABLET 12.5 MG, 25 MG558 XEOMIN
MG
XEOMIN
XGEVA560
XIAFLEX
XIFAXAN ORAL TABLET 200 MG 563
XIFAXAN ORAL TABLET 550 MG 562
XOLAIR SUBCUTANEOUS SOLUTION
PREFILLED SYRINGE
XOLAIR SUBCUTANEOUS SOLUTION
RECONSTITUTED564
XOSPATA
XPOVIO (100 MG ONCE WEEKLY)567
XPOVIO (40 MG ONCE WEEKLY) 568
XPOVIO (40 MG TWICE WEEKLY)568
XPOVIO (OU MG OINCE WEEKLY)
XPOVIO (60 MG ONCE WEEKLY) 567 XPOVIO (60 MG TWICE WEEKLY) 568
XPOVIO (60 MG ONCE WEEKLY) 567 XPOVIO (60 MG TWICE WEEKLY)

XTAMPZA ER
XTANDI
XURIDEN
XYOSTED
XYREM
XYWAV
YERVOY
YONSA
ZALTRAP
ZARXIO
ZAVESCA
ZEBUTAL
ZEJULA
ZELBORAF
ZEMAIRA
ZEPATIER
ZEPOSIA
ZEPOSIA 7-DAY STARTER PACK
ZEPOSIA STARTER KIT
ZIANA
ZIEXTENZO
ZINPLAVA
ZIRABEV
ZOHYDRO ER
zoledronic acid intravenous
concentrate
zoledronic acid intravenous solution 4
<i>mg/100ml</i>
zoledronic acid intravenous solution 5
<i>mg/100ml</i>
ZOLINZA
zolpidem tartrate193
<i>zolpidem tartrate er</i> 193
ZOLPIMIST
ZOMACTON
ZOMETA
ZORBTIVE
ZTLIDO
ZULRESSO
ZYDELIG
ZYKADIA
ZYTIGA ORAL TABLET 250 MG, 500
MG

ZYVOX ORAL SUSPENSION	
RECONSTITUTED	. 596
ZYVOX ORAL TABLET	. 596