

2025 PacificSource Health Plans Prior Authorization Criteria

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POLICY NAME: **ACTIMMUNE**

Affected Medications: ACTIMMUNE (interferon gamma 1b)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	Chronic Granulomatous Disease (CGD)
	 Severe, malignant osteopetrosis (SMO)
	NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A
	or higher
Required Medical	Patient's body surface area (BSA) must be documented along with the prescribed dose.
Information:	Pediatrics with BSA less than 0.5 m²: weight must be documented along with prescribed dose
	Chronic granulomatous disease
	 Diagnosis established by a molecular genetic test identifying a gene-related mutation associated with CGD
	Severe, malignant osteopetrosis
	 Diagnosis of severe infantile osteopetrosis established by ONE of the following: Radiographic imaging consistent with osteopetrosis
	OR
	 Molecular genetic test identifying a gene-related mutation associated with SMO
	Oncology indications
	 Documentation of performance status, disease staging, all prior therapies used, and anticipated treatment course
Appropriate	Chronic Granulomatous Disease
Treatment Regimen & Other	Patient is on a prophylactic regimen with an antibacterial agent and an antifungal agent
Criteria:	All indications
	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced
	Reauthorization: documentation of disease responsiveness to therapy
Exclusion Criteria:	Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
Age Restriction:	
Prescriber/Site of	CGD: prescribed by, or in consultation with, an immunologist
Care Restrictions:	SMO: prescribed by, or in consultation with, an endocrinologist
	Oncology indications: prescribed by, or in consultation with, an oncologist
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	CGD and SMO:
	Authorization: 12 months, unless otherwise specified



Oncology indications: Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: ADDYI & VYLEESI

Affected Medications: ADDYI (flibanserin), VYLEESI (bremelanotide injection)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Premenopausal women with acquired, generalized hypoactive sexual desire disorder (HSDD),* as characterized by low sexual desire that causes marked distress or interpersonal difficulty that is NOT due to any of the following: A coexisting medical or psychiatric condition Problems within a relationship
	 The effects of a medication or other drug substance *Also known as female sexual interest/arousal disorder
Paguirod Modical	
Required Medical Information:	 Documented mental health diagnosis of acquired, generalized HSDD meeting the Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition (DSM-5) criteria for female sexual interest/arousal disorder: Lack of, or significant reduction in, at least 3 of the following: interest in sexual activity sexual thoughts or fantasies initiation of sexual activity and responsiveness to a partner's initiation excitement or pleasure during all or almost all sexual activity interest or arousal in response to any sexual cues (e.g., written, verbal, visual) genital or non-genital sensations during sexual activity in all or almost all sexual encounters Symptoms have persisted for a minimum duration of 6 months Symptoms cause clinically significant distress Sexual dysfunction is not attributable to any of the following:
Appropriate	Reauthorization will require documentation of treatment success and confirmation that patient
Treatment	is still premenopausal
Regimen & Other Criteria:	
Exclusion Criteria:	Treatment of males or postmenopausal females
	Intended use is to enhance sexual performance
Age Restriction:	18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a mental health provider
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 2 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



ADENOSINE DEAMINASE (ADA) REPLACEMENT Affected Medications: REVCOVI (elapegademase-lvlr)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of adenosine deaminase severe combined immune deficiency (ADA-SCID) in pediatric and adult patients	
Required Medical Information:	 Diagnosis of ADA-SCID confirmed by genetic testing showing biallelic pathogenic variants in the ADA gene Laboratory findings show at least ONE of the following: Absent ADA levels in lysed erythrocytes A marked increase in deoxyadenosine triphosphate (dATP) levels in erythrocyte lysates A significant decrease in ATP concentration in red blood cells Absent or extremely low levels of N adenosylhomocysteine hydrolase in red blood cells Increase in 2'-deoxyadenosine in urine and plasma 	
Appropriate Treatment Regimen & Other Criteria:	Documentation showing that neither gene therapy nor a matched sibling or family donor for HCT (hematopoietic cell transplantation) is available, or that gene therapy or HCT was unsuccessful Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced authorization requires documentation of treatment success defined as disease stability d/or improvement as indicated by one or more of the following: Increase in plasma ADA activity Decrease in red blood cell dATP/dAXP level Improvement in immune function with diminished frequency/complications of infections	
Exclusion Criteria:	Other forms of autosomal recessive SCIDs All uses not listed under covered uses are considered experimental	
Age Restriction:		
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an immunologist or specialist experienced in the treatment of severe combined immune deficiency (SCID) All approvals are subject to utilization of the most cost-effective site of care 	
Coverage Duration:	Authorization: 12 months, unless otherwise specified	



POLICY NAME: **ADZYNMA**

Affected Medications: ADZYNMA (apadamtase alfa)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by				
	plan design				
	 Congenital thrombotic thrombocytopenic purpura (cTTP) 				
Required Medical	Diagnosis of severe cTTP confirmed by BOTH of the following:				
Information:	 Molecular genetic testing confirming presence of homozygous or compound 				
	heterozygous variants in the ADAMTS13 gene				
	 ADAMTS13 activity testing showing less than 10% of normal activity 				
	For on-demand treatment: documentation of current or past acute event with the				
	following:				
	 Reduction in platelet count by 50% or greater OR platelet count less than 				
	100,000/microliter				
	 Elevation in lactate dehydrogenase (LDH) level to more than 2x baseline or the 				
	upper limit of normal (ULN)				
	For prophylactic use:				
	 Must have history of at least one documented thrombotic thrombocytopenic 				
	purpura (TTP) event (past acute event or subacute event such as				
A	thrombocytopenia event or a microangiopathic hemolytic anemia event)				
Appropriate Treatment	Dosing: Description AC				
	Prophylactic: 40 IU/kg once every other week				
Regimen & Other Criteria:	 May be dosed weekly with documentation of appropriate prior dosing 				
Citteria.	regimen or clinical response o On-demand therapy: 40 IU/kg on day 1, 20 IU/kg on day 2, and 15 IU/kg on day				
	 On-demand therapy: 40 IU/kg on day 1, 20 IU/kg on day 2, and 15 IU/kg on day 3 and beyond until 2 days after the acute event is resolved 				
	3 and beyond until 2 days after the acute event is resolved				
	Reauthorization:				
	For prophylactic use: documentation of treatment success defined as an improvement in				
	the number or severity of TTP events, platelet counts, or clinical symptoms				
	For on-demand use: documentation of treatment success, defined as an increase in				
	platelet counts to at least 150,000/microliter, or counts returned to within 25% of baseline				
Exclusion Criteria:	Diagnosis of other TTP-like disorder, such as acquired or immune-mediated TTP				
Age Restriction:					
Prescriber/Site of	Prescribed by, or in consultation with, a hematologist, oncologist, intensive care				
Care Restrictions:	specialist, or specialist in rare genetic hematologic diseases				
	All approvals are subject to utilization of the most cost-effective site of care				
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified				
_	Reauthorization: 12 months, unless otherwise specified				



POLICY NAME: **AFAMELANOTIDE**

Affected Medications: SCENESSE (afamelanotide injection)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by				
	plan design				
	 Treatment of patients with erythropoietic protoporphyria (EPP) with phototoxic 				
	reactions				
Required Medical	Erythropoietic Protoporphyria (EPP)				
Information:	Documented diagnosis of EPP confirmed by biallelic loss-of-function mutation in the				
	ferrochelatase (FECH) gene				
	Documented increase in total erythrocyte protoporphyrin, with at least 85% metal-free protoporphyrin				
	Documented symptoms of phototoxic reactions, resulting in dysfunction and significant impact on activities of daily living				
Appropriate	Reauthorization:				
Treatment	Documentation of treatment success and clinically significant response to therapy (e.g.,				
Regimen & Other	decreased severity and number of phototoxic reactions, increased duration of sun				
Criteria:	exposure, increased quality of life, etc.)				
	AND				
	Continued implementation of sun and light protection measures during treatment to prevent phototoxic reactions				
Exclusion Criteria:	Cosmetic indications, such as vitiligo				
Age Restriction:	18 years of age and older				
Prescriber/Site of	Prescribed by, or in consultation with, a specialist at a recognized Porphyria Center				
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care				
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified				
	Reauthorization: 12 months, unless otherwise specified				



AFINITOR

Affected Medications: AFINITOR, AFINITOR DISPERZ (everolimus), EVEROLIMUS SOLUBLE TABLET

C) istant epilepsy, defined as lack of seizure control with 2 and meeting following criteria: ment failure with Epidiolex (cannabidiol solution) adjunct nitor Disperz (only form approved for TSC-seizures) is herapy for seizures
istant epilepsy, defined as lack of seizure control with 2 and meeting following criteria: ment failure with Epidiolex (cannabidiol solution) adjunct nitor Disperz (only form approved for TSC-seizures) is herapy for seizures
and meeting following criteria: ment failure with Epidiolex (cannabidiol solution) adjunct nitor Disperz (only form approved for TSC-seizures) is herapy for seizures
nitor Disperz (only form approved for TSC-seizures) is herapy for seizures
herapy for seizures
subependymal giant cell tumors (SGCTs) or TSC-cell astrocytoma (SEGA) in a patient who is not a good
tation of disease responsiveness to therapy
ess than or equal to 50% or ECOG performance score
ed by, or in consultation with, an oncologist or in consultation with, a neurologist or specialist in the
zation of the most cost-effective site of care
unless otherwise specified
less otherwise specified



POLICY NAME: **ALEMTUZUMAB**

Affected Medications: LEMTRADA (alemtuzumab)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of relapsing forms of multiple sclerosis (MS), including the following: Relapsing-remitting multiple sclerosis (RRMS) Active secondary progressive multiple sclerosis (SPMS) 		
Required Medical Information:	Diagnosis confirmed with magnetic resonance imaging (MRI) per revised McDonald diagnostic criteria for MS Clinical evidence alone will suffice; additional evidence desirable but must be consistent with MS		
Appropriate Treatment Regimen & Other Criteria:	Documentation of inadequate response to Tysabri (natalizumab) AND one additional medication indicated for MS Reauthorization requires provider attestation of treatment success Eligible for renewal 12 months after administration of last dose		
Exclusion Criteria:	Human immunodeficiency virus (HIV) infection Active infection Concurrent use of other disease-modifying medications indicated for the treatment of multiple sclerosis		
Age Restriction:			
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist or a multiple sclerosis specialist All approvals are subject to utilization of the most cost-effective site of care 		
Coverage Duration:	 Initial Authorization: 5 doses for 5 days, unless otherwise specified Reauthorization: 3 doses for 3 days, unless otherwise specified 		



ALGLUCOSIDASE ALFA

Affected Medications: LUMIZYME (alglucosidase alfa)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Pompe Disease		
Required Medical Information:	 Diagnosis of Pompe disease confirmed by an enzyme assay demonstrating a deficiency of acid α-glucosidase (GAA) enzyme activity or by DNA testing that identifies mutations in the GAA gene. Patient weight and planned treatment regimen 		
Appropriate Treatment Regimen & Other Criteria:	One or more clinical signs or symptoms of Pompe disease, including but not limited to: Readily observed evidence of glycogen storage (macroglossia, hepatomegaly, normal or increased muscle bulk) Involvement of respiratory muscles manifesting as respiratory distress (such as tachypnea) Profound diffuse hypotonia Proximal muscle weakness Reduced forced vital capacity (FVC) in upright or supine position Appropriate medical support is readily available when medication is administered in the event of anaphylaxis, severe allergic reaction, or acute cardiorespiratory failure Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced Reauthorization will require documentation of treatment success and a clinically significant response to therapy		
Exclusion Criteria:	Concurrent use of other enzyme replacement therapies such as Nexviazyme or Pombiliti and Opfolda		
Age Restriction:			
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a metabolic specialist, endocrinologist, biochemical geneticist, or physician experienced in the management of Pompe disease All approvals are subject to utilization of the most cost-effective site of care 		
Coverage Duration:	Authorization: 12 months, unless otherwise specified		



POLICY NAME: **ALOSETRON**

Affected Medications: ALOSETRON, LOTRONEX (alosetron)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Women with severe diarrhea-predominant irritable bowel syndrome (IBS) 			
Required Medical Information:	Female gender Chronic IBS syndrome lasting at least 6 months Diarrhea AND one or more of the following are present: o Frequent and severe abdominal pain/discomfort o Frequent bowel urgency or fecal incontinence o Disability or restriction of daily activities due to IBS Other anatomical or biochemical abnormalities of the gastrointestinal tract have been excluded as a cause of symptoms			
Appropriate Treatment Regimen & Other Criteria:	Documented inadequate response to all of the following:			
Exclusion Criteria:	 History of chronic or severe constipation or sequelae from constipation, intestinal obstruction, stricture, toxic megacolon, gastrointestinal perforation, and/or adhesions, ischemic colitis, impaired intestinal circulation, thrombophlebitis, or hypercoagulable state, Crohn's disease or ulcerative colitis, diverticulitis, or severe hepatic impairment Concomitant use of fluvoxamine 			
Age Restriction: Prescriber/Site of Care Restrictions:	 18 years of age and older Prescribed by, or in consultation with, a gastroenterologist All approvals are subject to utilization of the most cost-effective site of care 			
Coverage Duration:	Initial Authorization: 2 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified			



ALPHA-1 PROTEINASE INHIBITORS

Affected Medications: ARALAST NP, GLASSIA, PROLASTIN-C, ZEMAIRA

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded					
	by plan design.					
	o Chronic augmentation and maintenance therapy in adults with clinically evident					
	emphysema due to severe congenital alpha-1 antitrypsin (AAT) deficiency					
Required Medical	Documented diagnosis of severe congenital AAT deficiency, confirmed by BOTH of					
Information:	following (a and b):					
	a. Baseline AAT serum concentration of less than or equal to 11 mmol/L					
	(equivalent to 57 mg/dL or less via nephelometry, 80 mg/dL or less via radial					
	immunodiffusion)					
	b. One of the following high-risk phenotypic variants: PiZZ, PiSZ, Pi(null)(null), or other rare allelic mutation					
	Documentation of clinically evident emphysema or chronic pulmonary obstructive					
	disease (COPD), confirmed by ONE of the following (a or b):					
	a. Evidence of severe airflow obstruction, defined as forced expiratory volume in					
	one second (FEV1) of 30-65% predicted					
	b. Evidence of mild-moderate airflow obstruction, defined as an FEV1 between 66-					
	80% of predicted, but has demonstrated a rapid decline by at least 100 mL/year					
Appropriate Treatment	Documentation of non-smoker status					
Regimen & Other	Has not smoked for a minimum of 6 consecutive months leading up to therapy					
Criteria:	initiation and will continue to abstain from smoking during therapy					
	Design to CO may/log interconnected and a second					
	Dosing: 60 mg/kg intravenously once weekly Dose rounding to the pearest vial size within 10% of the prescribed dose will be					
	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced					
	Reauthorization will require documentation of treatment success and a clinically significant					
	response to therapy					
Exclusion Criteria:	Use in the management of lung disease in which severe AAT deficiency has not been					
	established					
	Patients with IgA deficiency or with the presence of IgA antibodies					
Ago Postriction	Prior liver transplant 19 years of are and older					
Age Restriction:	18 years of age and older					
Prescriber/Site of Care	Prescribed by, or in consultation with, a pulmonologist					
Restrictions:	All approvals are subject to utilization of the most cost-effective site of care					
Coverage Duration:	Authorization: 12 months, unless otherwise specified					
	1					



POLICY NAME: **AMIFAMPRIDINE**

Affected Medications: FIRDAPSE (amifampridine phosphate)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design			
Required Medical Information:	Documented diagnosis of LEMS confirmed by ONE of the following: Positive anti-P/Q-type voltage-gated calcium channel (VGCC) antibody test Repetitive nerve stimulation (RNS) abnormalities, such as an increase in compound muscle action potential (CMAP) amplitude at least 60 percent after maximum voluntary contraction (i.e., post-exercise stimulation) or at high frequency (50 Hz) Documentation of clinical signs and symptoms consistent with LEMS, as follows: proximal muscle weakness (without atrophy), with or without autonomic features and areflexia			
Appropriate Treatment Regimen & Other Criteria:	Documentation of inadequate clinical response or intolerance to ONE of the following (except in active small cell lung carcinoma [SCLC]-LEMS):			
Exclusion Criteria:	 Seizure disorder Active brain metastases Clinically significant long QTc interval on ECG in previous year OR history of additional risk factors for torsade de pointes 			
Age Restriction:	6 years of age or older			
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist or oncologist All approvals are subject to utilization of the most cost-effective site of care 			
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified 			



POLICY NAME: ANIFROLUMAB

Affected Medications: SAPHNELO (anifrolumab)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Systemic Lupus Erythematosus (SLE)			
Required Medical Information:	 Documentation of SLE with moderate to severe disease (significant but non-organ threatening disease including constitutional, cutaneous, musculoskeletal, or hematologi involvement) Autoantibody-positive SLE, defined as positive for antinuclear antibodies (ANA) and/or anti-double-stranded DNA (anti-dsDNA) antibody 			
Appropriate Treatment Regimen & Other Criteria:	 Failure with at least 12 weeks of combination therapy including hydroxychloroquine OR chloroquine with one of the following: Cyclosporine, azathioprine, methotrexate, or mycophenolate mofetil Documented failure with at least 12 weeks of subcutaneous Benlysta Reauthorization requires documentation of treatment success or a clinically significant improvement such as a decrease in flares or corticosteroid use 			
Exclusion Criteria:	 Use in combination with other biologic therapies Use in severe active central nervous system lupus 			
Age Restriction:	18 years of age and older			
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a rheumatologist or a specialist with experience in the treatment of systemic lupus erythematosus All approvals are subject to utilization of the most cost-effective site of care 			
Coverage Duration:	Authorization: 12 months, unless otherwise specified			



ANTIEMETICS

Affected Medications: AKYNZEO CAPSULES (netupitant-palonosetron), AKYNZEO INJECTION (fosnetupitant-palonosetron), VARUBI (rolapitant)

Covered Uses:	plan design Prevention courses of emetoger Prevention repeat co Prevention repeat co emetoger	on of delayed nausea and of emetogenic cancer chemic chemotherapy (arubi (rolapitant) on of acute and delayed nauses of highly emetoger akynzeo injection (fosneon of acute and delayed nauses of cancer chemotheric chemotherapy akynzeo capsules (neturally	I vomiting associated with the motherapy, including, but ausea and vomiting associated with the motherapetupitant-palonosetron) ausea and vomiting associated with the motherapy, including, but no	th initial and repeat out not limited to, high sociated with initial are.
Required Medical		ced Nausea and Vomiti		
Information:	Documentation of	f planned chemotherapy	regimen	
	 Akynzeo injection Documentation of a highly emetogenic chemotherapy regimen Akynzeo capsule Documentation of a highly OR moderately emetogenic chemotherapy regimen 			
		Highly Emetogen	ic Chemotherapy	
	Any regimen that contains an anthracycline and	Cyclophosphamide	Fam-trastuzumab deruxtecan-nxki	Sacituzumab govitecan-hziy
	cyclophosphamide			
		Dacarbazine	Ifosfamide	Streptozocin
	cyclophosphamide	Dacarbazine Doxorubicin	Ifosfamide Mechlorethamine	Streptozocin FOLFOX
	cyclophosphamide Carboplatin			· ·
	cyclophosphamide Carboplatin Carmustine Cisplatin	Doxorubicin	Mechlorethamine Melphalan	FOLFOX

Irinotecan

Oxaliplatin

Daunorubicin



	Moderately Emetogenic Chemotherapy				
	Aldesleukin	Cytarabine	Idarubicin	Mirvetuximab soravtansine-gynx	
	Amifostine	Dactinomycin	Irinotecan	Naxitamab-gqgk	
	Bendamustine	Daunorubicin	Irinotecan (liposomal)	Oxaliplatin	
	Busulfan	Dinutuximab	Lurbinectedin	Romidepsin	
	Clofarabine	Dual-drug liposomal encapsulation of cytarabine and daunorubicin	Methotrexate (250 mg/m² or greater)	Temozolomide	
Appropriate Treatment	Trabectedin	used Nauses and Vemitin	ag Prophylovic		
Regimen & Other Criteria:	 Chemotherapy induced Nausea and Vomiting Prophylaxis Varubi: Documented treatment failure with a 5-HT3 receptor antagonist (e.g., ondansetron, granisetron) in combination with dexamethasone while receiving the current chemotherapy regimen Akynzeo injection and capsule Documented treatment failure with both of the following while receiving the current chemotherapy regimen:				
	 Quantity Limit: Varubi: 1 dose per 14 days Akynzeo injection and capsule: 1 dose per 7 days Reauthorization requires documentation of treatment success and initial criteria to be met				
Exclusion Criteria:	Treatment of acute or breakthrough nausea and vomiting Used in anthracycline or cyclophosphamide-based chemotherapy (Akynzeo injection				
Age Restriction:	only) • 18 years of age and older				



Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an oncologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 6 months, unless otherwise specified



ANTIHEMOPHILIC FACTORS

Affected Medications: Advate, Adynovate, Afstyla, Alphanate, Alphanate/VWF Complex/Human, Alphanine SD, Alprolix, Altuviiio, Benefix, Corifact, Eloctate, Esperoct, Feiba NF, Helixate FS, Hemofil M, Humate P, Idelvion, Ixinity, Jivi, Koate DVI, Kogenate FS, Kovaltry, Monoclate-P, Mononine, Novoseven RT, NovoEight, Nuwiq, Obizur, Rebinyn, Recombinate, Riastap, Rixubis, Sevenfact, Tretten, Vonvendi, Wilate, Xyntha

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
Required Medical Information:	 Documentation of dose based on reasonable projections, current dose utilization, product labeling, diagnosis, baseline factor level, circulating factor activity (% of normal or units/dL), and rationale for use Current weight Documentation of Bethesda Titer level and number of bleeds in the past 3 months with severity and cause of bleed
	Documentation of one of the following diagnostic categories:
	Hemophilia A or Hemophilia B
	 Mild: factor levels greater than 5% and less than 30%
	 Moderate: factor levels of 1% to 5%
	 Severe: factor levels of less than 1%
	 Von Willebrand disease (VWD), which must be confirmed with plasma von Willebrand factor (VWF) antigen, plasma VWF activity, and factor VIII activity
	Documentation of one of the following indications:
	Acute treatment of moderate to severe bleeding in patients with:
	 Mild, moderate, or severe hemophilia A or B
	o Severe VWD
	 Mild to moderate VWD in clinical situations with increased risk of bleeding
	 Perioperative prophylaxis and/or treatment of acute, moderate to severe bleeding in patients with hemophilia A, hemophilia B, or VWD
	Routine prophylaxis in patients with severe hemophilia A, severe hemophilia B, or severe VWD
	 For Wilate and Vonvendi for routine prophylaxis: documentation of severe Type 3 VWD
Appropriate	Hemophilia A (factor VIII deficiency)
Treatment	Documentation indicates requested medication is to achieve or maintain but not to
Regimen & Other	exceed maximum functional capacity in performing daily activities
Criteria:	For mild disease: treatment failure or contraindication to Stimate (desmopressin)
	 Eloctate and Nuwiq require documented inadequate response, or documented intolerable adverse event, with all preferred products (Kogenate FS, Kovaltry, Novoeight, Jivi, Adynovate)
	Helixate FS requires documented treatment failure with Kogenate FS due to an intolerable adverse event and the prescriber has a compelling medical rationale for not expecting the same event to occur with Helixate FS



	 Altuviiio requires documentation of severe hemophilia or moderate hemophilia with a severe bleeding phenotype defined by frequent non-traumatic bleeds requiring prophylaxis
	 Hemophilia B (factor IX deficiency) For Benefix, Idelvion, and Rebinyn: documentation treatment failure or contraindication to Rixubis
	For Alprolix: documentation of contraindication to Rixubis for perioperative management
	von Willebrand disease (VWD) • For Vonvendi:
	 Documentation of treatment failure or contraindication to Humate P AND Alphanate for perioperative prophylaxis and/or treatment of acute, moderate to severe bleeding
	 Documentation of treatment failure or contraindication to Wilate for routine prophylaxis
	 All Indications Approval based on necessity and laboratory titer levels
	 Coverage for a non-preferred product requires documentation of one of the following: Documented intolerable adverse event to all preferred products, and the adverse event was not an expected adverse event attributed to the active ingredient Currently receiving treatment with a non-preferred product, excluding via samples or manufacturer's patient assistance programs
	<u>Reauthorization</u> : requires documentation of planned treatment dose, number of acute bleeds since last approval (with severity and cause of bleed), past treatment history, and titer inhibitor level to factor VIII and IX as appropriate
Exclusion Criteria:	 Acute thrombosis, embolism, or symptoms of disseminated intravascular coagulation Obizur for congenital hemophilia A or VWD
	 Tretten for congenital factor XIII B-subunit deficiency Jivi and Adynovate for VWD
	Idelvion for immune tolerance induction in patients with Hemophilia B
	Vonvendi for congenital hemophilia A or hemophilia B
	Afstyla and Nuwiq for VWD
Age Restriction:	Subject to review of FDA label for each product
	Jivi and Adynovate: 12 years of age and older
	 Vonvendi: 18 years of age and older Wilate for routine prophylaxis with von Willebrand disease: 6 years and older
Prescriber	Prescribed by, or in consultation with, a hematologist
Restrictions:	 Members who are on a State Based Drug List are required to utilize pharmacy benefits only
	All approvals are subject to utilization of the most cost-effective site of care



Coverage Duration:	•	Authorization: 12 months, unless otherwise specified
	•	Perioperative management: 1 month, unless otherwise specified



ANTITHYMOCYTE GLOBULINS

Affected Medications: ATGAM (antithymocyte globulin – equine), THYMOGLOBULIN (antithymocyte globulin – rabbit)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
0010.00 0000.	plan design
	Treatment of allograft rejection in renal transplant recipients (Atgam,
	Thymoglobulin)
	Treatment of moderate to severe aplastic anemia in patients unsuitable for bone
	marrow transplantation (Atgam)
	 Prophylaxis of acute rejection in renal transplant recipients (Thymoglobulin)
	National Comprehensive Cancer Network (NCCN) indications with evidence level of 2A
	or better
	Compendia-supported uses that will be covered (Thymoglobulin)
	 Prophylaxis and treatment of acute rejection in:
	 Heart transplant recipients
	 Liver transplant recipients
	 Lung transplant recipients
	 Pancreas transplant recipients
	 Intestinal transplant recipients
	 Prophylaxis of acute rejection in multivisceral transplant recipients
	 Prophylaxis of graft-versus-host disease in unrelated donor hematopoietic stem cell transplant recipients
Required	Oncology uses
Medical	Documentation of performance status, disease staging, all prior therapies used, and
Information:	anticipated treatment course
	All Indications
	Documentation of a complete treatment plan with planned dose, frequency and duration
	of therapy
	Current patient weight
	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced
	Prophylaxis of acute transplant rejection
	Patient must be considered high risk for acute rejection or delayed graft function based
	on one or more of either the following donor/recipient risk factors:
	Donor risk factors:
	o Donor cold ischemia for more than 24 hours
	 Donor age older than 50 years old Donor without a heartbeat
	D. W. ATM
	 Donor requiring high-dose inotropic support Recipient risk factors:

Panel-reactive antibody value exceeding 20% before transplant



	o Black race
	 One or more HLA antigen mismatches with the donor
Appropriate	Prophylaxis of acute transplant rejection
Treatment Regimen & Other Criteria:	 Documented treatment failure, intolerable adverse event, or contraindication to the use of basiliximab
	Treatment of allograft rejection in renal transplant recipients
	Requests for Atgam require documented treatment failure or rationale for avoidance of Thymoglobulin
Exclusion Criteria:	Oncology uses: Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
	Active acute or chronic infections which contraindicate additional immunosuppression
	Use in patients with aplastic anemia who are suitable candidates for bone marrow transplantation or in patients with aplastic anemia secondary to neoplastic disease, storage disease, myelofibrosis, Fanconi's syndrome, or in patients known to have been exposed to myelotoxic agents or radiation (Atgam)
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a specialist in oncology, hematology, nephrology or transplant medicine as appropriate for diagnosis All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 1 month, unless otherwise specified



POLICY NAME: ANTITHROMBIN III

Affected Medications: ANTITHROMBIN III (THROMBATE III)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design.
	 Indicated in patients with hereditary antithrombin deficiency (hATd) for:
	 Prevention of perioperative and peripartum thromboembolism
	 Prevention and treatment of thromboembolism
Required	All Indications
Medical	Documented diagnosis of hATd, confirmed by antithrombin (AT) activity levels below
Information:	70% on functional assay (not taken during acute illness, surgery, or thromboembolic
A	event that could give falsely low antithrombin levels)
Appropriate Treatment	Prevention of Perioperative Thromboembolism
Regimen &	Approved first-line for perioperative thromboprophylaxis in combination with heparin, with
Other Criteria:	or without intent to use as bridge to warfarin therapy
	Prevention of Peripartum Thromboembolism
	Documentation of ONE of the following:
	Personal or family history of thrombosis
	 Insufficient response to heparin AND intolerance to direct oral anticoagulants
	(DOACs)
	(50/103)
	Prevention of Thromboembolism
	 Documentation of inadequate clinical response, intolerance, or contraindication to BOTH
	of the following:
	o Warfarin
	At least one DOAC
	Treatment of Thromboembolism
	Approved first-line for treatment of thromboembolism as adjunct to anticoagulant
	therapy, unless coagulation is temporarily contraindicated
Exclusion	
Criteria:	
Age	
Restriction: Prescriber/Site of	Properihad by or in consultation with a homotologist geneticist or shotatricism
Care Restrictions:	Prescribed by, or in consultation with, a hematologist, geneticist, or obstetrician All approvals are subject to utilization of the most cost effective site of care.
	All approvals are subject to utilization of the most cost-effective site of care Position profits of particular appropriate through comboling transfer and the unless throughout the profits of
Coverage Duration:	Perioperative/peripartum prevention; thromboembolism treatment: 1 month, unless otherwise appairing.
Daration.	otherwise specified
	Thromboembolism prevention: 6 months, unless otherwise specified



ANTI-AMYLOID MONOCLONAL ANTIBODY

Affected Medications: LEQEMBI (lecanemab), KISUNLA (donanemab-azbt)

Covered Uses:	Leqembi (lecanemab) and Kisunla (donanemab-azbt) are not considered medically necessary due to insufficient evidence of therapeutic value.
Required Medical	
Information:	
Appropriate	
Treatment	
Regimen & Other	
Criteria:	
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of	
Care Restrictions:	
Coverage Duration:	



ANTI-TUBERCULOSIS AGENTS

Affected Medications: SIRTURO (bedaquiline), PRETOMANID

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise excluded by
	plan design.
	o Sirturo
	 Treatment of adult and pediatric patients with pulmonary tuberculosis
	(TB) due to <i>Mycobacterium tuberculosis</i> resistant to at least rifampin and isoniazid
	o Pretomanid
	 Treatment of adults with pulmonary TB resistant to isoniazid, rifamycins,
	a fluoroquinolone and a second line injectable antibacterial drug
	 Treatment of adults with pulmonary TB resistant to isoniazid and
	rifampin who are treatment-intolerant or nonresponsive to standard
	therapy
Required Medical	Sirturo
Information:	Documented diagnosis of multidrug resistant TB (MDR-TB), defined as resistance to at
	least isoniazid and rifampin
	Destauranid
	Pretomanid Documented diagnosis of one of the following:
	 Documented diagnosis of one of the following: Extensively drug resistant TB (XDR-TB)
	, ,
Appropriate	Treatment-intolerant or nonresponsive MDR-TB Sirturo
Treatment	 Documentation that this drug has been prescribed as part of a combination regimen with
Regimen & Other	other anti-tuberculosis agents
Criteria:	Documentation that this drug is being administered by directly observed therapy (DOT)
Omona.	Decamemation that the drag is being daministered by allocally essented therapy (Detr)
	Pretomanid
	Documentation that this drug has been prescribed as part of a combination regimen with
	Sirturo (bedaquiline) and linezolid
	Documentation that this drug is being administered by DOT
Exclusion Criteria:	Drug-sensitive (DS) pulmonary TB
	Latent infection due to Mycobacterium tuberculosis
	Extra-pulmonary infection due to Mycobacterium tuberculosis
	Infections caused by non-tuberculous mycobacteria
Age Restriction:	Sirturo: 5 years of age and older
	Pretomanid: 18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, an infectious disease specialist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Sirturo
1	Authorization: 24 weeks, unless otherwise specified



Authorization: 26 weeks, unless otherwise specified



POLICY NAME: **APOMORPHINE**

Affected Medications: KYNMOBI, APOKYN, APOMORPHINE SOLUTION

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Acute, intermittent treatment of hypomobility, "off" episodes in patients with advanced Parkinson's disease (PD)
Required Medical	Diagnosis of advanced PD
Information:	Documentation of acute, intermittent hypomobility, "off" episodes occurring for at least 2 hours per day while awake despite an optimized oral PD treatment regimen
Appropriate Treatment Regimen & Other Criteria:	Concurrent therapy with levodopa/carbidopa (at the maximum tolerated dose) and a second agent from one of the following alternate anti-Parkinson's drug classes:
Exclusion Criteria:	Use as monotherapy or first line agent
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: APROCITENTAN

Affected Medications: TRYVIO (aprocitentan)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
Covered Oses.	7 iii 7 ddd arid 27dg 7 idiriii iididaidi (1 27t) approved iididaidii 10t dirici iidd excidadd 2)
	plan design
	 Treatment of hypertension in combination with other antihypertensive drugs
Required Medical	Diagnosis of resistant hypertension
Information:	Blood pressure remains above target goal (as determined by treating provider) despite
	adherence to antihypertensive therapies
	Documentation of intent to use as an adjunct to current antihypertensive therapies
Appropriate	Documented treatment failure with concurrent use of at least four antihypertensive drugs
Treatment	(from different drug classes) at maximum tolerated doses, for a minimum of 12 weeks:
Regimen & Other	 Angiotensin-converting enzyme (ACE) inhibitor OR angiotensin II receptor
Criteria:	blocker (ARB)
	Calcium channel blocker (e.g. amlodipine, nifedipine, diltiazem, verapamil)
	Diuretic (e.g. hydrochlorothiazide, chlorthalidone)
	Beta-blocker (e.g. atenolol, carvedilol)
	 Mineralocorticoid receptor antagonist (e.g. spironolactone, eplerenone)
	Breed as a few and the second of the standard and the second of the seco
	Reauthorization requires documentation of treatment success and continued use of at least
Exclusion Criteria:	three background blood pressure therapies
exclusion Criteria.	Pregnancy
	 Concurrent use with an endothelin receptor antagonist (e.g. ambrisentan, bosentan, Opsumit, Filspari)
Age Restriction:	18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a cardiologist, nephrologist, or endocrinologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 3 months, unless otherwise specified
-	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **ARIKAYCE**

Affected Medications: ARIKAYCE (Amikacin inhalation suspension)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of <i>Mycobacterium avium</i> complex (MAC) lung disease as part of a combination antibacterial drug regimen in adults who have limited or no alternative treatment options, and who do not achieve negative sputum cultures after a minimum of 6 consecutive months of a multidrug background regimen therapy
Required Medical Information:	 Diagnosis of MAC lung disease confirmed by BOTH of the following: A MAC-positive sputum culture obtained within the last 3 months Evidence of underlying nodular bronchiectasis and/or fibrocavity disease on a chest radiograph or chest computed tomography The MAC isolate is susceptible to amikacin with a minimum inhibitory concentration (MIC) of less than or equal to 64 μg/mL Documentation of failure to obtain a negative sputum culture after a minimum of 6 consecutive months of a multidrug background regimen therapy for MAC lung disease such as clarithromycin (or azithromycin), rifampin and ethambutol
Appropriate Treatment Regimen & Other Criteria:	 Documentation of BOTH of the following: This drug has been prescribed as part of a combination antibacterial drug regimen This drug will be used with the Lamira® Nebulizer System Reauthorization requires documentation of negative sputum culture obtained within the last 30 days. The American Thoracic Society/Infectious Diseases Society of America (ATS/IDSA) guidelines state that patients should continue to be treated until they have negative cultures for 1 year. Treatment beyond the first reauthorization (after 18 months) will require documentation of a positive sputum culture to demonstrate the need for continued treatment. Patients that have had negative cultures for 1 year will not be approved for continued treatment.
Exclusion Criteria:	Diagnosis of non-refractory MAC lung disease
Age Restriction:	18 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an infectious disease specialist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **ASCIMINIB**

Affected Medications: SCEMBLIX (asciminib)

Covered Uses:	plan design	ninistration (FDA)-approved indications not otherwise excluded by rehensive Cancer Network) indications with evidence level of 2A
Required Medical Information:	 anticipated treatment of Documentation of Phila chronic myeloid leukem advanced phase CML- 	idelphia chromosome positive (Ph+) or BCR::ABL1- positive nia (CML) in chronic phase (May be appropriate in some cases of Check NCCN guidelines)
Appropriate		e or BCR::ABL1- positive chronic myeloid leukemia (CML) in
Treatment	chronic phase (CP) meeti	ng one of the following:
Regimen &	Law Biok Saara	
Other Criteria:	[TKI]) AND one or more	failure with imatinib (if used as initial tyrosine kinase inhibitor e additional tyrosine kinase inhibitor (TKI) bosutinib, dasatinib, or BL1 kinase domain mutation status for drug specific
	Intonio Patro a Districta	
	Intermediate or high-risk	
		failure with a second-generation tyrosine kinase inhibitor (TKI),
	specific contraindication	nilotinib. (Note BCR:ABL1 kinase domain mutation status for drugns)
	Drug	Contraindicated Mutations
	Asciminib	A337T, P465S, M244V, or F359V/I/C
	Bosutinib	T315I, V299L, G250E, or F317L
	Dasatinib	T315I/A, F317L/V/I/C, or V299L
	Nilotinib	T315I, Y253H, E255K/V, or F359V/C/I
	Ponatinib	None
	 OR Documented T315I positive AND Documented treatment Reauthorization requires of 	
Exclusion		e Status 50% or less or ECOG performance score 3 or greater
Criteria:		7T, P465S, M244V, or F359V/I/C BCR::ABL1 kinase domain
Age Restriction:		
Prescriber/Site of	Prescribed by, or in cor	nsultation with, an oncologist
Care Restrictions:	All approvals are subjective.	ct to utilization of the most cost-effective site of care



Coverage Duration:	•	Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified
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ATIDARSAGENE AUTOTEMCEL

Affected Medications: LENMELDY (atidarsagene autotemcel)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise excluded by
	plan design
	 Treatment of children with pre-symptomatic late-infantile (PSLI), pre- symptomatic early-juvenile (PSEJ), or early symptomatic early-juvenile (ESEJ) metachromatic leukodystrophy (MLD)
Required Medical	Diagnosis of metachromatic leukodystrophy (MLD) confirmed by the following:
Information:	 Arylsulfatase (ARSA) activity below the normal range in peripheral blood
	mononuclear cells or fibroblasts
	 Presence of two disease-causing mutations of either known or novel alleles
	 Presence of sulfatides in a 24-hour urine collection (to exclude MLD carriers and
	patients with ARSA pseudodeficiency)
	AND CALL OF CA
	 Diagnosis of the late-infantile subtype of MLD confirmed by two out of three of the following:
	 Age at onset of symptoms in the older sibling(s) less than or equal to 30 months
	 Two null (0) mutant ARSA alleles
	 Peripheral neuropathy as determined by electroneurographic study
	OR
	 Diagnosis of the early-juvenile subtype of MLD confirmed by two out of three of the following:
	 Age at onset of symptoms (in the patient or in the older sibling) between 30
	months and 6 years (has not celebrated their seventh birthday)
	 One null (0) and one residual (R) mutant ARSA allele(s)
	 Peripheral neuropathy as determined by electroneurographic study
Appropriate	
Treatment	
Regimen & Other	
Criteria:	Alleren St. Level and St. Control and Harris Heat of St. St. Heat on St. and St. and St.
Exclusion Criteria:	Allogeneic hematopoietic stem cell transplantation in the previous six months
	Previous gene therapy
	Documented HIV infection
	Documented history of a hereditary cancer
Age Restriction:	
Prescriber/Site of	Prescribed by or in consultation with a neurologist or hematologist/oncologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 2 months (for one time infusion), no reauthorization, unless otherwise specified



POLICY NAME: **AVACOPAN**

Affected Medications: TAVNEOS 10mg capsule

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design As an adjunctive treatment of adult patients with severe, active anti-neutrophil cytoplasmic autoantibody (ANCA)-associated vasculitis (AAV), including granulomatosis with polyangiitis (GPA) and microscopic polyangiitis (MPA), in	
	combination with standard therapy including glucocorticoids	
Required Medical Information:	 Diagnosis supported by at least one of the following: Tissue biopsy of kidney or other affected organs Positive ANCA, clinical presentation compatible with AAV, and low suspicion for secondary vasculitis Clinical presentation compatible with AAV, low suspicion for secondary vasculitis, and concern for rapidly progressive disease Documented severe, active disease (including major relapse), defined as: vasculitis with life- or organ-threatening manifestations (e.g., alveolar hemorrhage, glomerulonephritis, central nervous system vasculitis, subglottic stenosis, mononeuritis multiplex, cardiac involvement, mesenteric ischemia, limb/digit ischemia) Documentation of all prior therapies used and anticipated treatment course Baseline liver test panel: serum alanine aminotransferase, aspartate aminotransferase, alkaline phosphatase, and total bilirubin 	
	Current hepatitis B virus (HBV) status	
Appropriate	Will be used with a standard immunosuppressive regimen including glucocorticoids	
Treatment	Will be used during induction therapy only	
Regimen & Other Criteria:	 Will be used in any of the following populations/scenarios: In patients unable to use glucocorticoids at appropriate doses In patients with an estimated glomerular filtration rate less than 30 mL/min/1.73 m2 In patients who have experienced relapse following treatment with two or more different induction regimens, including both rituximab- and cyclophosphamide-containing regimens (unless contraindicated) During subsequent induction therapy in patients with refractory disease (failure to achieve remission with initial induction therapy regimen) Dosing: 30 mg (three 10 mg capsules) twice daily (once daily when used concomitantly 	
	with strong CYP3A4 inhibitors)	
Exclusion Criteria:	T	
Exclusion Ciliena.	 Treatment of eosinophilic-GPA (EGPA) Active, untreated and/or uncontrolled chronic liver disease (e.g., chronic active hepatitis B, untreated hepatitis C virus infection, uncontrolled autoimmune hepatitis) and cirrhosis Active, serious infections, including localized infections History of angioedema while receiving Tavneos, unless another cause has been established 	



	History of HBV reactivation while receiving Tavneos, unless medically necessary
Age Restriction:	18 years of age or older
Prescriber/Site of	Prescribed by, or in consultation with, a rheumatologist, nephrologist, or pulmonologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 6 months with no reauthorization, unless otherwise specified



AVALGLUCOSIDASE ALFA-NGPT

Affected Medications: NEXVIAZYME (avalglucosidase alfa-ngpt)

Covered Uses: Required Medical Information:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Late-Onset Pompe Disease Diagnosis of Pompe Disease confirmed by an enzyme assay demonstrating a deficiency of acid α-glucosidase (GAA) enzyme activity or by DNA testing that identifies mutations in the GAA gene. Patient weight and planned treatment regimen. 	
Appropriate Treatment Regimen & Other Criteria:	 One or more clinical signs or symptoms of Late-Onset Pompe Disease: Progressive proximal weakness in a limb-girdle distribution Delayed gross-motor development in childhood Involvement of respiratory muscles causing respiratory difficulty (such as reduced forced vital capacity [FVC] or sleep disordered breathing) Skeletal abnormalities (such as scoliosis or scapula alata) Low/absent reflexes Appropriate medical support is readily available when medication is administered in the event of anaphylaxis, severe allergic reaction, or acute cardiorespiratory failure. Patients weighing less than 30 kilograms will require documented treatment failure or intolerable adverse event to Lumizyme. Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced. Reauthorization will require documentation of treatment success and a clinically significant response to therapy. 	
Exclusion Criteria:	 Diagnosis of infantile-onset Pompe Disease Concurrent use of other enzyme replacement therapies such as Lumizyme or Pombiliti and Opfolda 	
Age Restriction:	1 year of age and older	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a metabolic specialist, endocrinologist, biochemical geneticist, or physician experienced in the management of Pompe disease All approvals are subject to utilization of the most cost-effective site of care 	
Coverage Duration:	Authorization: 12 months, unless otherwise specified	



POLICY NAME: **AVATROMBOPAG**

Affected Medications: DOPTELET (avatrombopag)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Thrombocytopenia in adult patients with chronic liver disease (CLD) who are
	scheduled to undergo a procedure
	 Thrombocytopenia in adult patients with chronic immune thrombocytopenia (ITP)
	who have had an insufficient response to a previous treatment
Required	Thrombocytopenia in patients with CLD undergoing a procedure
Medical	Documentation of planned procedure including date
Information:	Documentation of baseline platelet count of less than 50,000/microliter
	Thrombocytopenia in patients with chronic ITP
	Documentation of ONE of the following:
	Platelet count less than 20,000/microliter
	District and the soft and 00 000/via office AND and the office for
	 Platelet count less than 50,000/microliter AND increased risk for bleeding (such
	as peptic ulcer disease, use of antiplatelets or anticoagulants, history of bleeding
	at higher platelet count, need for surgery or invasive procedure)
Appropriate	Thrombocytopenia in patients with chronic (ITP):
Treatment	Documentation of inadequate response, defined as platelets did not increase to at least
Regimen &	50,000/microliter, to the following therapies:
Other Criteria:	 ONE of the following:
	 Inadequate response with at least 2 therapies for immune
	thrombocytopenia, including corticosteroids, rituximab, or
	immunoglobulin
	Splenectomy
	o Promacta
	Reauthorization (chronic ITP only)
	Response to treatment with platelet count of at least 50,000/microliter (not to exceed)
	400,000/microliter)
	OR
	The platelet counts have not increased to at least 50,000/microliter and the patient has
	NOT been on the maximum dose for at least 4 weeks
Exclusion	Use in combination with another thrombopoietin receptor agonist, spleen tyrosine kinase
Criteria:	inhibitor, or similar treatments (Promacta, Nplate, Tavalisse)
Age	
Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a hematologist or gastroenterology/liver specialist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage	Thrombocytopenia in patients with CLD undergoing a procedure:
Duration:	 1 month (for a one time 5-day regimen), unless otherwise specified
	Thrombocytopenia in patients with chronic ITP:
	 Initial Authorization: 4 months, unless otherwise specified
	 Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **AVONEX**

Affected Medications: AVONEX, AVONEX PEN

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design. Treatment of relapsing forms of Multiple Sclerosis (MS), including the following: Clinically isolated syndrome (CIS) Relapsing-remitting multiple sclerosis (RRMS) Active secondary progressive disease (SPMS)
Required Medical	Diagnosis confirmed with magnetic resonance imaging (MRI), per revised McDonald
Information:	diagnostic criteria for MS Clinical evidence alone will suffice; additional evidence desirable but must be consistent with MS
Appropriate Treatment Regimen & Other Criteria:	Reauthorization requires provider attestation of treatment success
Exclusion Criteria:	Concurrent use of other disease-modifying medications for treatment of MS
Age Restriction:	
Prescriber/Site of	All approvals are subject to utilization of the most cost-effective site of care
Care Restrictions:	Prescribed by, or in consultation with, a neurologist
Coverage Duration:	Approval: 12 months, unless otherwise specified



POLICY NAME: **AZTREONAM**

Affected Medications: CAYSTON (aztreonam)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Cystic fibrosis
Required Medical	Documentation of confirmed diagnosis of cystic fibrosis
Information:	Culture and sensitivity report confirming presence of Pseudomonas aeruginosa in the lungs
	Baseline FEV1 greater than 25% but less than 75% predicted
Appropriate	Documented failure, contraindication, or resistance to inhaled tobramycin.
Treatment	
Regimen & Other	Dosing: 28 days on and 28 days off
Criteria:	Reauthorization: requires documentation of improved respiratory symptoms and confirmed need for long-term use
Exclusion Criteria:	Baseline FEV1 less than 25% or greater than 75% predicted
Age Restriction:	Age 7 years of age and older
Prescriber/Site of	All approvals are subject to utilization of the most cost-effective site of care
Care Restrictions:	
Coverage Duration:	Initial approval: 6 months, unless otherwise specified
_	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **BELIMUMAB**

Affected Medications: BENLYSTA (belimumab)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Systemic Lupus Erythematosus (SLE) Lupus Nephritis (LN)
Required Medical Information:	 Documentation of current weight (intravenous requests only) Systemic Lupus Erythematosus: Documentation of active SLE with moderate classification (significant but non-organ threatening disease including constitutional, cutaneous, musculoskeletal, or hematologic involvement) Autoantibody-positive SLE, defined as positive for antinuclear antibodies (ANA) and/or anti-double-stranded DNA (anti-dsDNA) antibody Baseline measurement of ONE or more of the following:
	 Documentation of biopsy-proven active Class III, IV, and/or V disease Baseline measurement of one or more of the following: urine protein-creatinine ratio (uPCR), urine protein, estimated glomerular filtration rate (eGFR), or frequency of flares or corticosteroid use
Appropriate Treatment Regimen & Other Criteria:	All uses: ■ Use of intravenous formulation requires: □ Documented inability to use subcutaneous formulation OR □ Currently receiving treatment with the intravenous formulation, excluding via samples or manufacturer's patient assistance programs ■ Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced (intravenous requests only)
	Systemic Lupus Erythematosus: ■ Failure with at least 12 weeks of combination therapy including hydroxychloroquine OR chloroquine with one of the following: □ Cyclosporine, azathioprine, methotrexate, or mycophenolate mofetil
	 Reauthorization requires documentation of treatment success defined as ONE of the following: Clinically significant improvement in SRI-4, SLEDAI variant, or other validated scale for measurement of disease Decrease in frequency of flares or corticosteroid use
	Lupus Nephritis:



	 No dialysis in the past 12 months AND estimated glomerular filtration rate (eGFR) equal to or above 30 mL/min/1.73m² Failure of at least 12 weeks of mycophenolate mofetil AND cyclophosphamide
	 Reauthorization requires documentation of treatment success defined as ONE of the following: Improvement in eGFR Reduction in urinary protein-creatinine ratio or urine protein
	Decrease in flares or corticosteroid use
Exclusion Criteria:	 Use in combination with other biologic therapies for LN or SLE Use in severe active central nervous system lupus
Age Restriction:	5 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a nephrologist, rheumatologist, or specialist with experience in the treatment of systemic lupus erythematosus or lupus nephritis All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: **BELZUTIFAN**

Affected Medications: WELIREG (belzutifan)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or better
Required Medical Information:	 Von Hippel-Lindau (VHL) disease ● Diagnosis documented by the following: ○ Pathogenic VHL germline mutation diagnostic for VHL disease AND at least one of the following: ■ Presence of solid, locoregional tumor in kidney showing accelerated tumor growth (growth of 5 mm or more per year) ■ Presence of symptomatic and/or progressively enlarging central nervous system (CNS) hemangioblastomas not amenable to surgery ■ Presence of pancreatic solid lesion or pancreatic neuroendocrine tumor (pNET) with rapid tumor growth
	Treatment-refractory advanced or metastatic clear cell renal carcinoma ■ Advanced disease after use of the following treatments (per NCCN guidelines): □ A programmed death receptor-1 (PD-1) OR programmed death-ligand 1 (PD-L1) AND □ A vascular endothelial growth factor tyrosine kinase inhibitor (VEGF-TKI)
	Documentation of performance status, disease staging, all prior therapies used, and anticipated treatment course
Appropriate Treatment Regimen & Other Criteria:	Reauthorization: documentation of disease responsiveness to therapy
Exclusion Criteria:	 Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater Metastatic pNET disease Not to be used in combination with other oncologic agents for the treatment of VHL disease
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an oncologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **BENRALIZUMAB**

Affected Medications: FASENRA (benralizumab subcutaneous injection)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Add-on maintenance treatment of patients with severe asthma aged 6 years and older with an eosinophilic phenotype
	 Treatment of adult patients with eosinophilic granulomatosis with polyangiitis (EGPA)
Required Medical	Eosinophilic asthma
Information:	Diagnosis of severe asthma with an eosinophilic phenotype, defined by both of the following:
	 Baseline eosinophil count of at least 150 cells/μL OR dependent on daily oral corticosteroids
	AND
	 FEV1 less than 80% at baseline or FEV1/FVC reduced by at least 5% from normal
	EGPA
	Diagnosis of relapsing or refractory EGPA confirmed by all of the following:
	 Blood eosinophilia (at least 1,000 cells/mcL and/or greater than 10% of the total leukocyte count) at baseline
	Documented relapsing disease while on the highest tolerated oral corticosteroid or immunosuppressant dose
Appropriate	Eosinophilic asthma
Treatment	Documented use of high-dose inhaled corticosteroid (ICS) plus a long-acting beta
Regimen & Other Criteria:	agonist (LABA) for at least three months with continued symptoms AND
	Documentation of one of the following:
	 Documented history of 2 or more asthma exacerbations requiring oral or
	systemic corticosteroid treatment in the past 12 months while on combination
	inhaler treatment and at least 80% adherence
	 Documentation that chronic daily oral corticosteroids are required
	<u>EGPA</u>
	 Documented treatment failure or contraindication to at least two oral immunosuppressant drugs (azathioprine, methotrexate, mycophenolate) for at least 12 weeks each
	Reauthorization requires documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	Use in combination with another monoclonal antibody (e.g., Dupixent, Nucala, Xolair, Cinqair, Tezspire)
Age Restriction:	Eosinophilic asthma: 6 years of age and older
	EGPA: 18 years of age and older



Prescriber/Site of Care Restrictions:	 <u>Eosinophilic asthma</u>: prescribed by, or in consultation with, an allergist, immunologist, or pulmonologist <u>EGPA</u>: prescribed by, or in consultation with, a specialist in the treatment of EGPA (such as an immunologist or rheumatologist) All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



BEREMAGENE GEPERPAVEC-SVDT

Affected Medications: VYJUVEK (beremagene geperpavec-svdt)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not all as its analysis.
Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Dystrophic Epidermolysis Bullosa (DEB)
Required Medical	Diagnosis of recessive DEB confirmed by both of the following:
Information:	Skin biopsy of an induced blister with immunofluorescence mapping (IFM) and/or
	transmission electron microscopy (TEM)
	 Genetic test results documenting mutations in the COL7A1 gene
	Clinical signs and symptoms of DEB such as skin fragility, blistering, scarring, nail changes, and milia formation in the areas of healed blistering
Appropriate	Documentation of receiving standard of care preventative or treatment therapies for
Treatment	wound care, control of infection, nutritional support
Regimen & Other	Documented trial and failure of Filsuvez
Criteria:	
Ontona.	be in accordance with 1 27 classing and accorded the following.
	Maximum weekly volume of 2.5 mL (1.6 mL useable dose)
	Maximum of 12-week course per wound Maximum of 4 tyless per 32 days.
	Maximum of 4 tubes per 28 days
	Reauthorization will require documentation of treatment success defined as complete
	wound healing on a previous site and need for treatment on a new site
	would realing out a previous site and need for treatment out a new site
Exclusion Criteria:	Evidence or history of squamous cell carcinoma in the area that will undergo treatment
	Concurrent use with Filsuvez (birch triterpenes topical gel)
	Dominant DEB (DDEB)
Age Restriction:	6 months of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a dermatologist or a specialist experienced in the
Care Restrictions:	treatment of epidermolysis bullosa
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 3 months, unless otherwise specified
	Reauthorization: 3 months, unless otherwise specified
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BESREMI

Affected Medications: BESREMI (ropeginterferon alfa-2b)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Treatment of adults with polycythemia vera
	NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher
Required	Documentation of performance status, disease staging, all prior therapies used, and
Medical	anticipated treatment course
Information:	Evidence of increased red cell volume such as abnormal hemoglobin, hematocrit, or red
	cell mass AND one of the following:
	 Presence of JAK2 V617F or JAK2 exon 12 mutation
	 Subnormal serum erythropoietin level
Appropriate	Documentation of treatment failure, intolerance, or contraindication to hydroxyurea
Treatment	
Regimen &	Reauthorization requires documentation of disease responsiveness to therapy
Other Criteria:	<u> </u>
Exclusion	Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
Criteria:	
Age	18 years of age and older
Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist or hematologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage	Initial Authorization: 4 months, unless otherwise specified
Duration:	Reauthorization: 12 months, unless otherwise specified



BETAINE

Affected Medications: CYSTADANE (betaine), BETAINE

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Homocystinuria
Required Medical Information:	Diagnosis of homocystinuria associated with one of the following:
Appropriate Treatment Regimen & Other Criteria:	Documented trial and failure of ONE of the following forms of supplementation:
Exclusion Criteria:	Uncorrected vitamin B12 or folic acid levels
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a metabolic or genetic disease specialist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: **BETASERON**

Affected Medications: BETASERON (interferon beta-1b)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of relapsing forms of multiple sclerosis (MS), including the following: Clinically isolated syndrome (CIS) Relapsing-remitting multiple sclerosis (RRMS) Active secondary progressive disease (SPMS)
Required Medical Information:	Diagnosis confirmed with magnetic resonance imaging (MRI), per revised McDonald diagnostic criteria for MS Clinical evidence alone will suffice; additional evidence desirable but must be consistent with MS
Appropriate Treatment Regimen & Other Criteria:	Reauthorization: provider attestation of treatment success
Exclusion Criteria:	Concurrent use of other disease-modifying medications indicated for the treatment of multiple sclerosis
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist or multiple sclerosis specialist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Approval: 24 months, unless otherwise specified



BETIBEGLOGENE AUTOTEMCEL

Affected Medications: ZYNTEGLO (betibeglogene autotemcel)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design o Treatment of beta thalassemia in adult and pediatric patients who require regular red blood cell (RBC) transfusions
Required Medical Information:	 Documented diagnosis of transfusion dependent beta thalassemia (TDT), defined as: Requiring at least 100 mL/kg per year of packed red blood cells (pRBCs) or at least 8 transfusions per year of pRBCs in the 2 years preceding therapy Confirmed genetic testing based on the presence of biallelic mutations at the beta-globin gene (HBB gene) Clinically stable and eligible to undergo hematopoietic stem cell transplant (HSCT) Used as single agent therapy (not applicable to lymphodepleting or bridging therapy while awaiting manufacture) Females of reproductive potential must have negative pregnancy test prior to start of mobilization, reconfirmed prior to conditioning procedures, and again before administration of Zynteglo
Appropriate Treatment Regimen & Other Criteria:	Patients must weigh a minimum of 6 kilograms and be able to provide a minimum number of cells (5,000,000 CD34+ cells/kilogram)
Exclusion Criteria:	 Prior HSCT or other gene therapy Severe iron overload warranting exclusion from therapy, as determined by the treating physician Uncorrected bleeding disorder Cardiac T2* less than 10 milliseconds by magnetic resonance imaging (MRI) White blood cell count less than 3x109/L and/or platelet count less than 100x109/L that is unrelated to hypersplenism Positive for human immunodeficiency virus 1 & 2 (HIV-1/HIV-2), hepatitis B virus, or hepatitis C virus, advanced liver disease, or current or prior malignancy
Age Restriction:	4 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a hematologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 4 months (one-time infusion), unless otherwise specified



POLICY NAME: **BEVACIZUMAB**

Affected Medications: AVASTIN, MVASI, ZIRABEV, ALYMSYS, VEGZELMA

Covered Uses:	 NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher For the Treatment of Ophthalmic disorders: Neovascular (Wet) Age-Related Macular Degeneration (AMD) Macular Edema Following Retinal Vein Occlusion (RVO) Diabetic Macular Edema (DME) Diabetic Retinopathy (DR) in patients with Diabetes Mellitus
Required Medical	Documentation of disease staging, all prior therapies used, and anticipated treatment
Information:	course
Appropriate	Stage III or IV Epithelial Ovarian, Fallopian Tube, or Primary Peritoneal Cancer
Treatment	following initial surgical resection
Regimen & Other	Approval will be limited for up to 22 cycles of therapy
Criteria:	
G.1.66.1	All Indications
	Coverage for a non-preferred product (Avastin, Alymsys, Vegzelma) requires
	documentation of one of the following:
	 Use for an ophthalmic condition (Avastin only)
	 A documented intolerable adverse event to the preferred products, Mvasi and
	Zirabev, and the adverse event was not an expected adverse event attributed to the active ingredient
	Reauthorization requires documentation of disease responsiveness to therapy
Exclusion Criteria:	Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
Age Restriction:	
Prescriber/Site of	Oncologic indication: prescribed by, on in consultation with, an oncologist
Care Restrictions:	Ophthalmic indication: prescribed by, on in consultation with, an ophthalmologist
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 4 months, unless otherwise specified
-	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **BEZLOTOXUMAB**

Affected Medications: ZINPLAVA (bezlotoxumab)

Carrage I Hanne	AUG I ID ALITE (FDA)
Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	 Reduce recurrence of Clostridioides difficile infection (CDI) in patients who are
	receiving antibacterial drug treatment for CDI and are at a high risk for CDI
	recurrence
Required	Diagnosis of CDI confirmed by both of the following:
Medical	Presence of at least 3 unformed stools in 24 hours
Information:	 Positive stool test for toxigenic Clostridium difficile collected within 7 days prior to request
	Patient must be receiving concurrent CDI treatment when infusion is administered
Appropriate	Documentation of ONE of the following risk factors for CDI recurrence:
Treatment	o Age greater than 65
Regimen &	 One or more episodes of CDI in the past 6 months prior to the current episode
Other Criteria:	o Immunocompromised status
	 Clinically severe CDI (defined by Zar score greater than or equal to 2)
	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced
Exclusion	Previous treatment with Zinplava
Criteria:	'
Age	1 year of age and older
Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, an infectious disease specialist or
Care Restrictions:	gastroenterologist
	All approvals are subject to utilization of the most cost-effective site of care
Coverage	Authorization: 1 month (a single 10 mg/kg dose) with no reauthorization, unless
Duration:	otherwise specified



BIRCH TRITERPENES

Affected Medications: FILSUVEZ (birch triterpenes topical gel)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise excluded by
	plan design
	 Dystrophic Epidermolysis Bullosa (DEB)
	 Junctional Epidermolysis Bullosa (JEB)
Required Medical Information:	 Diagnosis of recessive DEB or JEB confirmed by skin biopsy of an induced blister with immunofluorescence mapping (IFM) and/or transmission electron microscopy (TEM) Genetic test results documenting mutations in one of the following genes: COL7A1, COL17A1, ITGB4, LAMA3, LAMB3, or LAMC2
	 Clinical signs and symptoms of EB such as skin fragility, blistering, scarring, nail changes, and milia formation in the areas of healed blistering Presence of open partial-thickness wounds that have been present for at least 21 days
Appropriate	Documentation of receiving standard of care preventative or treatment therapies for
Treatment	wound care, control of infection, nutritional support.
Regimen & Other	Dosing does not exceed the following:
Criteria:	Maximum of 1 mm layer to affected area(s)
	o Maximum of 28 tubes per 28 days
	Reauthorization requires documentation of treatment success defined as complete wound healing on a previous site and need for continued treatment on a new site
Exclusion Criteria:	Concurrent use with Vyjuvek (beremagene geperpavec-svdt)
	Dominant DEB (DDEB)
Age Restriction:	6 months of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a dermatologist or a specialist experienced in the
Care Restrictions:	treatment of epidermolysis bullosa
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 3 months, unless otherwise specified
	Reauthorization: 3 months, unless otherwise specified



BOTOX

Affected Medications: BOTOX (onabotulinum toxin A)

Covered Uses:	All Food and Drug Administration (FDA) approved or compandin supported indications
Covered Oses.	All Food and Drug Administration (FDA)-approved or compendia-supported indications act at a ruise evaluated by plan design.
	not otherwise excluded by plan design
	o Spasticity
	o Chronic migraine
	 Overactive bladder (OAB) with symptoms of urge urinary incontinence, urgency,
	and frequency
	 Neurogenic detrusor overactivity (NDO)
	o Focal dystonia
	 Cervical dystonia
	Blepharospasm
	 Laryngeal dystonia
	 Oromandibular dystonia
	 Severe brachial dystonia (writer's cramp)
	o Strabismus
	 Primary axillary hyperhidrosis
	o Achalasia
	o Anal fissure
Required Medical	Pertinent medical records and diagnostic testing
Information:	Complete description of the site(s) of injection
	Strength and dosage of botulinum toxin used
Appropriate Treatment	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \
Regimen & Other	indications not otherwise excluded by plan design that are not listed below, failure of
Criteria:	first-line recommended and conventional therapies is required
	 Approved first-line for: focal dystonia, hemifacial spasm, orofacial dyskinesia, upper/lower limb spasticity, or other conditions of focal spasticity wherein botulinum
	toxin is the preferred mode of therapy
	toxiir is the preferred mode of therapy
	Overactive bladder (OAB)/Neurogenic detrusor overactivity (NDO):
	Documentation of inadequate response or intolerance to at least two urinary
	incontinence anticholinergic agents (e.g., oxybutynin, solifenacin, tolterodine)
	Chuania minuaina
	 Chronic migraine Documentation of chronic migraine defined as headaches on at least 15 days per
	month, of which at least 8 days are with migraine
	 Documented failure with an adequate trial (at least 8 weeks) of a migraine preventative
	therapy, as follows:
	o Candesartan 16 mg daily
	 Antiepileptic (divalproex sodium 500 mg daily, valproic acid 500 mg daily,
	topiramate 50 mg daily)
	Beta-blocker (metoprolol 100 mg daily, propranolol 40 mg daily, timolol 20 mg
	daily, nadolol 80 mg daily)
	 Antidepressant (amitriptyline 25 mg daily, nortriptyline 25 mg daily, venlafaxine



	 75 mg daily, duloxetine 60 mg daily) Anti-calcitonin gene-related peptide (CGRP) monoclonal antibody or CGRP receptor antagonist (when used for prevention)
	Primary Axillary Hyperhidrosis
	 Thyroid-stimulating hormone (TSH) level AND inadequate response to two or more alternative therapies (topical aluminum chloride 20%, iontophoresis, oral glycopyrrolate, oral oxybutynin)
	Achalasia (Cardiospasm) - must meet 1 of the following
	Type I or II achalasia: Treatment failure with peroral endoscopic myotomy (POEM), laparoscopic Heller myotomy (LHM), and pneumatic dilation (PD)
	Type III achalasia: Treatment failure with tailored POEM and LHM
	 Not a candidate for POEM, surgical myotomy, or pneumatic dilation due to high risk of complications
	Anal fissure
	Documented failure or intolerance to an 8-week trial of each of the following: Rective ointment
	Topical diltiazem or topical nifedipine
	Number of treatments must not exceed the following:
	OAB/NDO: 4 treatments per 12 months
	Chronic migraine: initial treatment limited to two injections given 3 months apart,
	subsequent treatment approvals limited to 4 treatments per 12 months
	Primary axillary hyperhidrosis: 2 treatments per 12 months
	Anal fissure: 2 treatments per 12 months
	All other indications maximum of 4 treatments per 12 months unless otherwise specified
	Reauthorization:
	Chronic migraine continuation of treatment: Additional treatment requires that the
	member has achieved or maintained a 50% reduction in monthly headache frequency since starting therapy with Botox.
	All other indications: Documentation of treatment success and a clinically significant
	response to therapy
Exclusion Criteria:	Cosmetic procedures
	For intradetrusor injections: documented current/recent urinary tract infection or urinary
	retention
	Possible medication overuse headache: headaches occurring 15 or more days each
	month in a patient with pre-existing headache-causing condition possibly due to
	 Use of ergotamines, triptans, opioids, or combination analgesics at least 10 days per month for at least three months
	 Use of simple analgesics (acetaminophen, aspirin, or an NSAID) at least 15
	days per month for at least 3 months
	 Combined use of any of the previously mentioned products without overuse of
	any one agent if no causative pattern can be established
	Combined use with an anti-calcitonin gene-related peptide (CGRP) monoclonal antibody
	or an oral CGRP antagonist when used for migraine prevention
Age Restriction:	



Prescriber/Site of Care	Prescribed by, or in consultation with, a specialist for the following:
Restrictions:	 Blepharospasm, strabismus: ophthalmologist, optometrist, or neurologist
	 Chronic migraine: neurologist or headache specialist
	 OAB/NDO: urologist or neurologist
	 Anal fissure: gastroenterologist or colorectal surgeon
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Chronic migraine:
	Authorization: 12 months, unless otherwise specified
	OAB/NDO:
	Initial Authorization: 3 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified
	Anal Fissure:
	Authorization: 3 months (one treatment), unless otherwise specified
	All other indications:
	Authorization: 12 months, unless otherwise specified



POLICY NAME: **BUROSUMAB**

Affected Medications: CRYSVITA (burosumab-twza)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design X-linked hypophosphatemia (XLH) FGF23-related hypophosphatemia in tumor induced osteomalacia (TIO) associated with phosphaturic mesenchymal tumors
Required Medical Information:	All Indications ■ Documentation of diagnosis by: ○ A blood test demonstrating ALL of the following (in relation to laboratory reference ranges): ■ Low phosphate ■ Elevated FGF23 ■ Low 1,25-(OH)2D ■ Normal calcium or parathyroid hormone (PTH) ○ A urine test demonstrating decreased tubular reabsorption of phosphate corrected for glomerular filtration rate (TmP/GFR) ○ Evidence of skeletal abnormalities, confirmed by radiographic evaluation
	 Tumor-Induced Osteomalacia Documentation that tumor cannot be located or is unresectable Alternative renal phosphate-wasting disorders have been ruled out
Appropriate Treatment Regimen & Other Criteria:	 All Indications Documentation of treatment failure with at least 12 months of oral phosphate and calcitriol supplementation in combination, unless contraindicated or not tolerated Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced Reauthorization requires: Documentation of normalization of serum phosphate levels If established on therapy for 12 months or more, improvement in radiographic imaging of skeletal abnormalities
Exclusion Criteria:	
Age Restriction:	
Prescriber Restrictions:	 Prescribed by, or in consultation with, a nephrologist, endocrinologist, or a provider experienced in managing patients with metabolic bone disease All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: CALCIFEDIOL

Affected Medications: RAYALDEE (calcifediol extended-release)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of secondary hyperparathyroidism in adult patients with stage 3 or 4 chronic kidney disease (CKD) and serum total 25-hydroxyvitamin D levels less than 30 ng/mL
Required Medical Information:	 A confirmed diagnosis of secondary hyperparathyroidism with persistently elevated or progressively rising serum intact parathyroid hormone (iPTH) that is at least 2.3 times above the upper limit of normal for the assay used Documentation of all of the following prior to treatment initiation: Stage 3 or 4 CKD Serum total 25-hydroxyvitamin D level is less than 30 ng/mL Corrected serum calcium is below 9.8 mg/dL
Appropriate Treatment Regimen & Other Criteria:	Documentation of persistent vitamin D deficiency (level below 30 ng/mL), despite at least 12 weeks of adherent treatment with each of the following at an appropriate dose, unless contraindicated or not tolerated:
Exclusion Criteria:	A diagnosis of stage 1, 2, or 5 chronic kidney disease, or end-stage renal disease (ESRD) on dialysis
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a nephrologist or endocrinologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



CALCITONIN GENE-RELATED PEPTIDE (CGRP) INHIBITORS
Affected Medications: EMGALITY (galcanezumab), VYEPTI (eptinezumab)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Preventative treatment of migraine in adults Episodic cluster headaches (Emgality only)
Required Medical	Chronic migraine prevention:
Information:	Diagnosis of chronic migraine defined as headaches on at least 15 days per month of which at least 8 days are with migraine at baseline
	Fuir dis minorina massartian
	Episodic migraine prevention:
	Diagnosis of episodic migraine with at least 4 migraines per month at baseline
	Episodic cluster headaches (Emgality Only):
	History of episodic cluster headache with at least two cluster periods lasting from 7 days to 1 year (when untreated) separated by pain-free remission periods of at least one month
	Headaches are not due to medication overuse: headaches occurring 15 or more days each month in a patient with pre-existing headache-causing condition possibly due to: Use of ergotamines, triptans, opioids, or combination analgesics at least 10 days per month for at least three months Use of simple analgesics (acetaminophen, aspirin, or an NSAID) at least 15 days per month for at least 3 months Use of combination of any previously mentioned products without overuse of any one agent if no causative pattern can be established
Appropriate	Chronic or Episodic migraine:
Treatment	Documented treatment failure with an adequate trial (at least 8 weeks) of ONE oral
Regimen & Other	migraine preventive therapy as follows:
Criteria:	 Candesartan 16 mg daily
	 Propranolol 40 mg daily, metoprolol 100 mg daily, timolol 20 mg daily, nadolol 80 mg daily
	 Amitriptyline 25 mg daily, nortriptyline 25 mg daily, venlafaxine 75 mg daily, duloxetine 60 mg daily
	 Topiramate 50 mg daily, valproic acid 500 mg daily, divalproex sodium 500 mg daily
	 Requests for Vyepti: Documented treatment failure to an adequate 8-week trial of an oral preventive therapy AND a minimum 12-week trial with each of the following: Emgality Botox (chronic migraine only)
	Enjandia cluster handaches (Emgality Only):
	Episodic cluster headaches (Emgality Only):
	Documented treatment failure with an adequate trial of verapamil (dose of at least 480)



	Reauthorization requires documentation of treatment success defined as a 50% reduction in monthly headache frequency since starting therapy
Exclusion Criteria:	 Combined use with Botox Combined use with another anti-calcitonin gene-related peptide (CGRP) monoclonal antibody or CGRP receptor antagonist (acute or preventive)
Age Restriction:	18 years of age or older
Prescriber/Site of Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: CANNABIDIOL

Affected Medications: EPIDIOLEX (cannabidiol)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
Required Medical Information:	All Indications Patient weight Documentation that cannabidiol will be used as adjunctive therapy (LGS) Documentation of at least 8 drop seizures per month while on stable antiepileptic drug therapy (DS) Documentation of at least 4 convulsive seizures in the last month while on stable antiepileptic drug therapy (TSC) Documentation of at least 8 TSC-associated seizures (e.g., focal-onset seizures with and without impaired awareness, focal-onset to bilateral tonic-clonic seizures, subclinical seizures, generalized onset seizures) in the last month while on stable antiepileptic drug therapy
Appropriate Treatment Regimen & Other Criteria:	LGS ■ Documented treatment failure with at least three guideline directed therapies including: □ Valproate and □ Lamotrigine and □ Rufinamide, topiramate, felbamate, or clobazam DS ■ Documented treatment failure with at least four guideline directed therapies including: □ Valproate and □ Clobazam and □ Topiramate and □ Clonazepam, levetiracetam, or zonisamide TSC ■ Documented treatment failure with at least two antiepileptic drugs used as monotherapy
	AND Documented treatment failure with at least two antiephieptic drugs used as monotherapy AND Documented treatment failure with at least one adjunctive therapy Dosing:



	LGS or DS: Not to exceed 20 mg/kg per day
	TSC: Not to exceed 25 mg/kg per day
	Reauthorization requires documentation of treatment success and a reduction in seizure severity, frequency, and/or duration.
Exclusion Criteria:	Use as monotherapy for seizure control
Age Restriction:	1 year of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: CANTHARIDIN

Affected Medications: YCANTH

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	Molluscum contagiosum (MC)
Required Medical	Diagnosis of MC confirmed by one of the following:
Information:	 Presence of lesions that are consistent with MC (small, firm, pearly, with pitted centers, 2-5 millimeters in diameter, not associated with systemic symptoms such as fever) For lesions with unclear cause or otherwise not consistent with MC, confirmation of diagnosis using dermoscopy, microscopy, histological examination, or biopsy
	Documentation of persistent itching or pain AND one of the following:
	 Concomitant bacterial infection of the lesion
	 Concomitant atopic dermatitis
	 Significant concern for contagion (such as daycare setting) and prevention
	cannot be reasonably prevented through good hygiene and covering lesions with
	bandages or clothing
	 Continued presence of lesions after 12 months
Appropriate	Trial of at least two cycles of one of the following procedures for the removal of MC
Treatment	lesions:
Regimen & Other	 Cryotherapy
Criteria:	 Curettage
	 Laser therapy
	Adequate trial and failure of one additional treatment for MC that has evidence
	supporting use, such as:
	Topical podofilox for at least 1 month
	 Oral cimetidine for at least 2 months
	Dosing: Two applicators per treatment every 21 days, limit to 4 total treatments
Exclusion Criteria:	
Age Restriction:	2 years of age or older
Prescriber/Site of	Prescribed by, or in consultation with, a dermatologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 3 months, unless otherwise specified



CAPLACIZUMAB-YHDP

Affected Medications: CABLIVI (caplacizumab-yhdp)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of adult patients with acquired thrombotic thrombocytopenic purpura (aTTP), in combination with plasma exchange and immunosuppressive therapy
Required Medical Information:	 Diagnosis, or suspected diagnosis, of aTTP, meeting all of the following: Severe thrombocytopenia (platelet count less than 100 x 10⁹/L) Microangiopathic hemolytic anemia (MAHA) confirmed by red blood cell fragmentation (e.g., schistocytes) on peripheral blood smear Baseline ADAMTS13 activity level of less than 10% Documentation of <u>ONE</u> of the following: Failure of at least one initial treatment for aTTP, such as therapeutic plasma exchange (TPE), glucocorticoids, or rituximab Documentation of high-risk disease meeting <u>ONE</u> of the following:
Appropriate Treatment	Total treatment duration will be limited to 58 days beyond the last TPE treatment
Regimen & Other Criteria:	Reauthorization requires documented signs of ongoing disease (such as suppressed ADAMTS13 activity levels) and no more than 2 recurrences of aTTP while on Cablivi. Recurrence is defined as thrombocytopenia after initial recovery of platelet count (platelet count greater than or equal to 150,000) that requires re-initiation of daily plasma exchange.
Exclusion Criteria:	Use for other causes of thrombocytopenia, such as other TTP-like disorders (congenital or hereditary TTP)
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a hematologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 3 months, unless otherwise specified Reauthorization: 3 months (for new episode), unless otherwise specified



POLICY NAME: CAPSAICIN KIT

Affected Medications: QUTENZA (capsaicin kit)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Neuropathic pain associated with postherpetic neuralgia (PHN) Neuropathic pain associated with diabetic peripheral neuropathy (DPN) of the feet	
Required Medical Information:		
Appropriate Treatment Regimen & Other Criteria:	 Documented treatment failure with at least 12 weeks of ALL of the following: gabapentin pregabalin carbamazepine, oxcarbazepine, or valproic acid/divalproex sodium amitriptyline or nortriptyline topical lidocaine Dose limited to a single treatment (up to 4 patches) once every 90 days Reauthorization: requires documentation of treatment success and a clinically significant response to therapy as assessed by the prescribing provider 	
Exclusion Criteria:		
Age Restriction:		
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a pain management specialist All approvals are subject to utilization of the most cost-effective site of care 	
Coverage Duration:	 Initial Authorization: 3 months (single treatment), unless otherwise specified Reauthorization: 12 months (up to 4 treatments), unless otherwise specified 	



POLICY NAME: CARGLUMIC ACID

Affected Medications: CARBAGLU, CARGLUMIC ACID

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Acute hyperammonemia due to one of the following: N-Acetylglutamate Synthase (NAGS) deficiency Propionic Acidemia (PA) or Methylmalonic Acidemia (MMA) Chronic hyperammonemia due to N-Acetylglutamate Synthase (NAGS) deficiency
Required	Diagnosis is confirmed by enzymatic, biochemical, or genetic testing
Medical	Ammonia level above the upper limit of normal (ULN) reference range for the patient's
Information:	age
Appropriate Treatment	Current weight
Regimen &	Acute hyperammonemia
Other Criteria:	 Prescribed in combination with at least one other ammonia-lowering therapy (examples include: sodium phenylacetate and sodium benzoate, intravenous glucose, insulin, L- arginine, L-carnitine, protein restriction, dialysis)
	For disease due to PA or MMA: Prescribed treatment course does not exceed 7 days
	 Reauthorization for acute disease requires documentation of reoccurrence of acute hyperammonemia meeting initial criteria Chronic hyperammonemia due to N-Acetylglutamate Synthase (NAGS) deficiency Prescribed in combination with a protein-restricted diet Reauthorization for chronic disease requires: Documentation of treatment success and a clinically significant response to therapy as evidenced by reduction in ammonia levels Documentation of member's current weight and continuation of appropriate treatment course
Exclusion Criteria:	Hyperammonemia caused by other enzyme deficiencies in the urea cycle:
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a metabolic disease specialist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage	Acute Hyperammonemia due to PA or MMA:
Duration:	Authorization: 7 days, unless otherwise specified



Acute Hyperammonemia due to NAGs deficiency:

Authorization: 1 month, unless otherwise specified

Chronic Hyperammonemia:

- Initial Authorization: 3 months, unless otherwise specified
- Reauthorization: 12 months, unless otherwise specified



POLICY NAME: CERLIPONASE ALFA

Affected Medications: BRINEURA (cerliponase alfa)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design To slow the loss of ambulation in pediatric patients with neuronal ceroid lipofuscinosis type 2 (CLN2), also known as tripeptidyl peptidase-1 (TPP1) deficiency		
Required Medical Information:	 Diagnosis of CLN2 disease confirmed by BOTH of the following: Enzyme assay demonstrating deficient TPP1 activity Genetic testing that has detected two pathogenic variants/mutations in the TPP1/CLN2 gene (one on each parental allele of the TPP1/CLN2 gene) Documentation of mild to moderate functional impairment at baseline using the CLN2 Clinical Rating Scale, defined as ALL the following: Combined score of 3 to 6 in the motor and language domains Score of at least 1 in the motor domain Score of at least 1 in the language domain 		
Appropriate Treatment Regimen & Other Criteria:	Dosing is in accordance with FDA labeling Reauthorization:		
Exclusion Criteria:	Documentation of clinical responsiveness to therapy defined as disease stabilization OR a score of at least 1 in the motor domain of the CLN2 Clinical Rating Scale Any sign or symptom of acute or unresolved localized infection on or around the device insertion site (e.g., cellulitis or abscess); or suspected or confirmed CNS infection (e.g., cloudy CSF or positive CSF gram stain, or meningitis) Any acute intraventricular access device-related complication (e.g., leakage, extravasation of fluid, or device failure) Other forms of neuronal ceroid lipofuscinosis Patients with ventriculoperitoneal shunts		
Age Restriction:			
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist with expertise in the diagnosis of CLN2 All approvals are subject to utilization of the most cost-effective site of care 		
Coverage Duration:	Authorization: 6 months, unless otherwise specified		



CFTR MODULATORS

Affected Medications: ALYFTREK (vanzacaftor/tezacaftor/deutivacaftor), KALYDECO (ivacaftor), ORKAMBI (lumacaftor/ivacaftor), SYMDEKO (tezacaftor/ivacaftor), TRIKAFTA (elexacaftor/tezacaftor/ivacaftor)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by		
	plan design		
	 Cystic fibrosis (CF) in patients with mutation(s) in the F508del cystic fibrosis 		
	transmembrane conductance regulator (CFTR) gene or another responsive		
	mutation in the CFTR gene		
	 CF in patients who are homozygous for the F508del mutation in the CFTR gene (Orkambi) 		
Required Medical	Documentation of cystic fibrosis (CF) diagnosis confirmed by appropriate genetic or		
Information:	diagnostic testing (FDA-approved CF mutation test)		
	 Please provide the diagnostic testing report and/or Cystic Fibrosis Foundation 		
	Patient Registry Report		
	Documentation of mutation(s) in the CFTR gene for which the drug has been FDA-		
	approved to treat		
Appropriate	Reauthorization will require documentation of treatment success		
Treatment			
Regimen & Other			
Criteria:			
Exclusion Criteria:	Kalydeco: Homozygous F508del mutation		
	Concurrent use with another CFTR modulator		
Age Restriction:	Alyftrek: 6 years of age and older		
	Kalydeco: one month of age and older		
	Orkambi: 1 year of age and older		
	Symdeko: 6 years of age and older		
	Trikafta: 2 years of age and older		
Prescriber/Site of	Prescribed by, or in consultation with, a pulmonologist or provider who specializes in CF		
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care		
Coverage Duration:	Initial Authorization: 12 months, unless otherwise specified		
	Reauthorization: 24 months unless otherwise specified		
	<u>'</u>		



CHELATING AGENTS

Preferred drugs: deferasirox soluble tablet, deferasirox tablet Non-Preferred drugs: Ferriprox (deferiprone), deferiprone

1.	Is the request for continuation of therapy currently approved through insurance?	Yes – Go to renewal criteria	No – Go to #2	
2.	Is the request to treat a diagnosis according to one of the Food and Drug Administration (FDA)-approved indications?	Yes – Go to appropriate section below	No – Criteria not met	
Pr	Chronic Iron Overload Due to Blood Transfusions in Myelodysplastic Syndromes Preferred Drugs – deferasirox soluble tablet, deferasirox tablet Non -Preferred drugs: Ferriprox (deferiprone), deferiprone			
1.	Documentation of International Prognostic Scoring System (IPSS) low or intermediate-1 risk level?	Yes – Document and go to #2	No – Criteria not met	
2.	Documentation of a history of more than 20 red blood cell (RBC) transfusions OR that it is anticipated that more than 20 would be required?	Yes – Document and go to #3	No – Criteria not met	
3.	Documentation of serum ferritin levels greater than 2500 ng/ml?	Yes – Document and go to # 4	No – Criteria not met	
4.	Is the request for generic formulation of deferasirox (oral or soluble tablet)?	Yes – Go to #6	No- Go to #5	
5.	Is there documented failure to deferasirox and deferoxamine (Desferal)?	Yes – Document and go to #6	No – Criteria not met	
6.	Is the drug prescribed by, or in consultation with, a hematologist specialist?	Yes – Go to #7	No – Criteria not met	
7.	Is the requested dose within the Food and Drug Administration (FDA) approved label?	Yes – Approve up to 12 months	No – Criteria not met	
Ch	Chronic Iron Overload Due to Blood Transfusions in Thalassemia syndromes, Sickle Cell Disease, or other			

Chronic Iron Overload Due to Blood Transfusions in Thalassemia syndromes, Sickle Cell Disease, or other anemias

Preferred Drugs – deferasirox soluble tablet, deferasirox tablet **Non -Preferred drugs:** Ferriprox (deferiprone), deferiprone



1.	Documentation of pretreatment serum ferritin level within the last 60 days of at least 1000 mcg/L?	Yes – Document and go to #2	No – Criteria not met
2.	Is the request for generic formulation of deferasirox (oral or soluble tablet)?	Yes – Document and go to #4	No – Go to #3
3.	Is there documented failure to deferasirox and deferoxamine (Desferal)?	Yes – Document and go to #4	No – Criteria not met
4.	Documentation of platelet counts greater than 50,000 per microliter?	Yes – Go to #5	No – Criteria not met
5.	Is the drug prescribed by, or in consultation with, a hematologist specialist?	Yes – Document and go to #6	No – Criteria not met
6.	Is the requested dose within the Food and Drug Administration (FDA) approved label?	Yes – Approve up to 12 months	No – Criteria not met
	ronic Iron Overload in Non-Transfusion Dependent Thalaseferred Drugs –deferasirox soluble tablet, deferasirox tablet	ssemia Syndromes	
1.	Documentation of liver iron (Fe) concentration (LIC) levels consistently greater than or equal to 5 mg Fe per gram of dry weight	Yes – Document and go to #2	No – Criteria not met
2.	Documentation of serum ferritin levels consistently greater than 300 mcg/L prior to initiation of treatment	Yes – Document and go to #3	No – Criteria not met
3.	Is the requested dose within the Food and Drug Administration (FDA) approved label?	Yes – Approve up to 12 months	No – Criteria not met
Re	newal Criteria		
1.	Is there documentation of treatment success and a clinically significant response to therapy defined as a reduction from baseline liver iron concentration (LIC) or serum ferritin level (LIC and serum ferritin must still be above 3 mg Fe per gram of dry weight and 500 mcg/L, respectively)	Yes – Go to #2	No – Criteria not met
2.	Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations?	Yes – Approve up to 12 months	No – Criteria not met



Quantity Limitations

- Exjade (deferasirox soluble tablet) available in 125mg, 250mg, 500mg tablets
 - 20-40 mg/kg/day
- Jadenu (deferasirox tablet or granules) available in 90mg, 180mg, 360mg tablets
 - o 14-28 mg/kg/day
- Ferriprox (deferiprone) 100mg/ml oral solution, 500mg, 1000mg tablets
 - o 75-99 mg/kg/day
 - Can be used in adult and pediatric patients 8 years of age and older (tablets), or 3 years of age and older (solution)



POLICY NAME: CHOLBAM

Affected Medications: CHOLBAM (cholic acid)

All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of bile acid synthesis disorders due to single enzyme defects (SEDs) Adjunctive treatment of peroxisomal disorders, including Zellweger spectrum disorders and provided information: Pocumentation of all prior therapies, patient weight and anticipated treatment course Baseline liver function tests (AST, ALT, GGT, ALP, total bilirubin, INR) Bile acid synthesis disorder		
Baseline liver function tests (AST, ALT, GGT, ALP, total bilirubin, INR) Bile acid synthesis disorder Diagnosis confirmed by assessment of serum or urinary bile acid levels using mass spectrometry (Fast Atom Bombardment ionization - Mass Spectrometry (FAB-MS) analysis) Peroxisomal disorders including Zellweger spectrum disorders Diagnosis confirmed by clinical features, elevated very long-chain fatty acid (VLCFA) levels, peroxisomal biomarkers, genetic testing Prothrombin time (vitamin K), serum levels of vitamins A, D, and E Hepatic injury or at risk of liver injury (elevations in liver enzymes or atypical bile acids) OR If normal liver function tests, must show manifestations of liver disease, steatorrhea, or complications from decreased fat-soluble vitamin absorption Appropriate Treatment Regimen & Other Criteria: Pwill not be used for treatment of extrahepatic manifestations (such as neurologic symptoms) of bile acid synthesis disorders Reauthorization requires documentation of clinically significant improvement in liver function as determined by meeting TWO of the following criteria: Improvement in abnormal liver chemistries (AST, ALT, bilirubin) Reduction or stabilization of hepatic inflammation and fibrosis Reduced levels of the toxic C27-bile acid intermediates dihydroxycholestanoic acid (DHCA) and trihydroxycholestanoic acid (THCA) in plasma and urine Improvement in prothrombin time (as a result of improved vitamin K absorption) and serum levels of vitamins A, D, and E No evidence of cholestasis on liver biopsy Body weight increased or stabilized Treatment should be discontinued if liver function does not improve after 3 months of start of treatment Exclusion Criteria: Age	Covered Uses:	plan design Treatment of bile acid synthesis disorders due to single enzyme defects (SEDs) Adjunctive treatment of peroxisomal disorders, including Zellweger spectrum disorders, in patients who exhibit manifestations of liver disease, steatorrhea, or
Diagnosis confirmed by assessment of serum or urinary bile acid levels using mass spectrometry (Fast Atom Bombardment ionization - Mass Spectrometry (FAB-MS) analysis) Peroxisomal disorders including Zellweger spectrum disorders Diagnosis confirmed by clinical features, elevated very long-chain fatty acid (VLCFA) levels, peroxisomal biomarkers, genetic testing Prothrombin time (vitamin K), serum levels of vitamins A, D, and E Hepatic injury or at risk of liver injury (elevations in liver enzymes or atypical bile acids) OR If normal liver function tests, must show manifestations of liver disease, steatorrhea, or complications from decreased fat-soluble vitamin absorption Appropriate Treatment Regimen & Will not be used for treatment of extrahepatic manifestations (such as neurologic symptoms) of bile acid synthesis disorders Reauthorization requires documentation of clinically significant improvement in liver function as determined by meeting TWO of the following criteria: Improvement in abnormal liver chemistries (AST, ALT, bilirubin) Reduction or stabilization of hepatic inflammation and fibrosis Reduced levels of the toxic C27-bile acid intermediates dihydroxycholestanoic acid (DHCA) and trihydroxycholestanoic acid (THCA) in plasma and urine Improvement in prothrombin time (as a result of improved vitamin K absorption) and serum levels of vitamins A, D, and E No evidence of cholestasis on liver biopsy Body weight increased or stabilized Treatment should be discontinued if liver function does not improve after 3 months of start of treatment	Medical	· · · · · · · · · · · · · · · · · · ·
Diagnosis confirmed by assessment of serum or urinary bile acid levels using mass spectrometry (Fast Atom Bombardment ionization - Mass Spectrometry (FAB-MS) analysis) Peroxisomal disorders including Zellweger spectrum disorders Diagnosis confirmed by clinical features, elevated very long-chain fatty acid (VLCFA) levels, peroxisomal biomarkers, genetic testing Prothrombin time (vitamin K), serum levels of vitamins A, D, and E Hepatic injury or at risk of liver injury (elevations in liver enzymes or atypical bile acids) OR If normal liver function tests, must show manifestations of liver disease, steatorrhea, or complications from decreased fat-soluble vitamin absorption Appropriate Treatment Regimen & Will not be used for treatment of extrahepatic manifestations (such as neurologic symptoms) of bile acid synthesis disorders Reauthorization requires documentation of clinically significant improvement in liver function as determined by meeting TWO of the following criteria: Improvement in abnormal liver chemistries (AST, ALT, bilirubin) Reduction or stabilization of hepatic inflammation and fibrosis Reduced levels of the toxic C27-bile acid intermediates dihydroxycholestanoic acid (DHCA) and trihydroxycholestanoic acid (THCA) in plasma and urine Improvement in prothrombin time (as a result of improved vitamin K absorption) and serum levels of vitamins A, D, and E No evidence of cholestasis on liver biopsy Body weight increased or stabilized Treatment should be discontinued if liver function does not improve after 3 months of start of treatment		Rile acid synthesis disorder
Diagnosis confirmed by clinical features, elevated very long-chain fatty acid (VLCFA) levels, peroxisomal biomarkers, genetic testing Prothrombin time (vitamin K), serum levels of vitamins A, D, and E Hepatic injury or at risk of liver injury (elevations in liver enzymes or atypical bile acids) OR If normal liver function tests, must show manifestations of liver disease, steatorrhea, or complications from decreased fat-soluble vitamin absorption Appropriate Treatment Regimen & Other Criteria: Reauthorization requires documentation of clinically significant improvement in liver function as determined by meeting TWO of the following criteria: Improvement in abnormal liver chemistries (AST, ALT, bilirubin) Reduction or stabilization of hepatic inflammation and fibrosis Reduced levels of the toxic C27-bile acid intermediates dihydroxycholestanoic acid (DHCA) and trihydroxycholestanoic acid (THCA) in plasma and urine Improvement in prothrombin time (as a result of improved vitamin K absorption) and serum levels of vitamins A, D, and E No evidence of cholestasis on liver biopsy Body weight increased or stabilized Treatment should be discontinued if liver function does not improve after 3 months of start of treatment Exclusion Criteria: Age		 Diagnosis confirmed by assessment of serum or urinary bile acid levels using mass spectrometry (Fast Atom Bombardment ionization - Mass Spectrometry (FAB-MS)
Diagnosis confirmed by clinical features, elevated very long-chain fatty acid (VLCFA) levels, peroxisomal biomarkers, genetic testing Prothrombin time (vitamin K), serum levels of vitamins A, D, and E Hepatic injury or at risk of liver injury (elevations in liver enzymes or atypical bile acids) OR If normal liver function tests, must show manifestations of liver disease, steatorrhea, or complications from decreased fat-soluble vitamin absorption Appropriate Treatment Regimen & Other Criteria: Reauthorization requires documentation of clinically significant improvement in liver function as determined by meeting TWO of the following criteria: Improvement in abnormal liver chemistries (AST, ALT, bilirubin) Reduction or stabilization of hepatic inflammation and fibrosis Reduced levels of the toxic C27-bile acid intermediates dihydroxycholestanoic acid (DHCA) and trihydroxycholestanoic acid (THCA) in plasma and urine Improvement in prothrombin time (as a result of improved vitamin K absorption) and serum levels of vitamins A, D, and E No evidence of cholestasis on liver biopsy Body weight increased or stabilized Treatment should be discontinued if liver function does not improve after 3 months of start of treatment Exclusion Criteria: Age		Paravisamal disorders including Zellwager spectrum disorders
Hepatic injury or at risk of liver injury (elevations in liver enzymes or atypical bile acids) OR If normal liver function tests, must show manifestations of liver disease, steatorrhea, or complications from decreased fat-soluble vitamin absorption - Will not be used for treatment of extrahepatic manifestations (such as neurologic symptoms) of bile acid synthesis disorders - Will not be used for treatment of extrahepatic manifestations (such as neurologic symptoms) of bile acid synthesis disorders - Regimen & Other Criteria: - Reauthorization requires documentation of clinically significant improvement in liver function as determined by meeting TWO of the following criteria: - Improvement in abnormal liver chemistries (AST, ALT, bilirubin) - Reduction or stabilization of hepatic inflammation and fibrosis - Reduced levels of the toxic C27-bile acid intermediates dihydroxycholestanoic acid (DHCA) and trihydroxycholestanoic acid (THCA) in plasma and urine - Improvement in prothrombin time (as a result of improved vitamin K absorption) and serum levels of vitamins A, D, and E - No evidence of cholestasis on liver biopsy - Body weight increased or stabilized - Treatment should be discontinued if liver function does not improve after 3 months of start of treatment - Exclusion - Criteria: - Age		Diagnosis confirmed by clinical features, elevated very long-chain fatty acid (VLCFA)
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Other Criteria: Reauthorization as determined by meeting TWO of the following criteria: Improvement in abnormal liver chemistries (AST, ALT, bilirubin) Reduction or stabilization of hepatic inflammation and fibrosis Reduced levels of the toxic C27-bile acid intermediates dihydroxycholestanoic acid (DHCA) and trihydroxycholestanoic acid (THCA) in plasma and urine Improvement in prothrombin time (as a result of improved vitamin K absorption) and serum levels of vitamins A, D, and E No evidence of cholestasis on liver biopsy Body weight increased or stabilized Treatment should be discontinued if liver function does not improve after 3 months of start of treatment Exclusion Criteria: Age	Treatment	
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Treatment should be discontinued if liver function does not improve after 3 months of start of treatment Exclusion Criteria: Age		· ·
Start of treatment Exclusion Criteria: Age		Body weight increased or stabilized
Criteria: Age		·



Prescriber/Site of Care Restrictions:	Prescribed by, or in consultation with, a hepatologist, gastroenterologist, or metabolic specialist	
	All approvals are subject to utilization of the most cost-effective site of care	
Coverage	Initial Authorization: 3 months, unless otherwise specified	
Duration:	Reauthorization: 12 months, unless otherwise specified	



CHOLESTATIC LIVER DISEASE

Affected Medications: BYLVAY (odevixibat), LIVMARLI (maralixibat)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by Plan design		
	plan design		
	 Pruritus due to progressive familial intrahepatic cholestasis (PFIC) 		
	Cholestatic pruritus in patients with Alagille syndrome (ALGS)		
Required Medical	Documentation of experiencing moderate to severe pruritis associated with PFIC or		
Information:	ALGS		
	Documentation of serum bile acid concentration above the upper limit of normal (ULN) reference range for the reporting laboratory		
	PFIC		
	Documentation of confirmed molecular diagnosis of PFIC type 1 or type 2		
	 Documentation of absence of ABCB11 gene variant if PFIC type 2 		
	ALGS		
	Documentation of ALGS confirmed by:		
	 Genetic test detecting a JAG1 or NOTCH2 mutation OR 		
	 Liver biopsy and at least three clinical features: 		
	 Chronic cholestasis 		
	 Cardiac disease 		
	 Ocular or skeletal abnormalities 		
	 Characteristic facial features 		
	 Renal and vascular disease 		
Appropriate	Documentation of current weight and dosing in accordance with FDA labeling		
Treatment	Documented treatment failure with ALL of the following for at least 30 days:		
Regimen & Other	o Rifampin		
Criteria:	o Ursodiol		
	 Cholestyramine (or colesevelam if requesting for ALGS) 		
	Reauthorization requires documentation of treatment success and a clinically significant response to therapy		
Exclusion Criteria:	Prior hepatic decompensation events		
	Decompensated cirrhosis (such as ALT or total bilirubin greater than 10-times the ULN)		
	Concomitant liver disease (e.g., biliary atresia, liver cancer, non- PFIC related		
	cholestasis)		
	Prior liver transplant		
Age Restriction:	Age is in accordance with FDA labeling		
Prescriber/Site of	Prescribed by, or in consultation with, a hepatologist or a specialist with experience in		
Care Restrictions:	the treatment of PFIC or ALGS		
	All approvals are subject to utilization of the most cost-effective site of care		
Coverage Duration:	Initial Authorization: 4 months, unless otherwise specified		
Soverage Duration.	initial Additionzation. 4 months, unless otherwise specified		
	<u>l</u>		



Reauthorization: 12 months, unless otherwise specified



CIALIS

Affected Medications: CIALIS (2.5 mg, 5 mg), TADALAFIL (2.5 mg, 5 mg)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of symptomatic benign prostatic hyperplasia (BPH) Mental health diagnosis of erectile disorder (ED) meeting sexual dysfunction criteria	
Required	Benign Prostatic Hyperplasia	
Medical	Documented diagnosis of benign prostatic hyperplasia (BPH)	
Information:		
	Mental Health Diagnosis of Erectile Dysfunction	
	Documentation of a mental health diagnosis of erectile dysfunction meeting the	
	Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition (DSM-5) criteria:	
	 At least one of the three following symptoms must be experienced with 75% to 	
	100% of occasions of sexual activity:	
	 Marked difficulty in obtaining an erection during sexual activity 	
	 Marked difficulty in maintaining an erection until the completion of sexual 	
	activity	
	Marked decrease in erectile rigidity	
	 The above symptoms have persisted for a minimum duration of approximately 6 	
	months	
	 The above symptoms cause clinically significant distress in the individual 	
	 The sexual dysfunction is not attributable to any of the following: 	
	 A nonsexual medical or psychiatric condition 	
	 Severe relationship distress (e.g., partner violence) 	
	The effects of medication or other substance use	
	 Other clinically significant and relevant stressors 	
Appropriate	Benign Prostatic Hyperplasia	
Treatment	Documented treatment failure with at least two of the following: alfuzosin, doxazosin,	
Regimen &	silodosin, finasteride, tamsulosin, terazosin	
Other Criteria:		
	Reauthorization requires documentation of treatment success and a clinically significant	
	response to therapy	
	response to therapy	
	Limited to 1 tablet per day	
	·	
Exclusion	Freetile duefunction unrelated to a montal health diagnosis of covered duefunction	
	Erectile dysfunction unrelated to a mental health diagnosis of sexual dysfunction	
Criteria:	according to the DSM-5 diagnostic criteria	
Age		
Restriction:		
Prescriber/Site of	Mental health diagnosis of erectile dysfunction: prescribed by, or in consultation with, a	
Care Restrictions:	mental health provider	
	All approvals are subject to utilization of the most cost-effective site of care	
Coverage	Authorization: 12 months, unless otherwise specified	
Duration:		



POLICY NAME: **CLADRIBINE**

Affected Medications: MAVENCLAD (cladribine)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of relapsing forms of multiple sclerosis (MS), including the following: Clinically isolated syndrome (CIS) Relapsing-remitting multiple sclerosis (RRMS) Active secondary progressive disease (SPMS)
Required Medical Information:	Diagnosis confirmed with magnetic resonance imaging (MRI) per revised McDonald diagnostic criteria for MS Clinical evidence alone will suffice; additional evidence desirable but must be consistent with MS
Appropriate Treatment Regimen & Other Criteria:	Documented treatment failure with (or intolerance to) a minimum 12-week trial of at least two disease-modifying therapies for MS Reauthorization (one time only): provider attestation of treatment success Eligible to initiate second treatment cycle 43 weeks after last dose was administered
Exclusion Criteria:	 Current malignancy Human immunodeficiency virus (HIV) infection Active chronic infections (e.g., hepatitis, tuberculosis) Pregnancy Treatment beyond 2 years
Age Restriction:	
Prescriber/Site of Care Restrictions: Coverage Duration:	 Prescribed by, or in consultation with, a neurologist or MS specialist All approved are subject to utilization of the most cost-effective site of care Initial Authorization: 2 months, unless otherwise specified
Coverage Duration.	Reauthorization: 2 months, unless otherwise specified



POLICY NAME: COAGADEX

Affected Medications: COAGADEX (Factor X)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Indicated in children and adults with hereditary Factor X (FX) deficiency for: Routine prophylaxis to reduce frequency of bleeding episodes On-demand treatment and control of bleeding episodes Perioperative management of bleeding in mild, moderate, or severe disease 		
Required Medical	All Indications		
Information:	 Documented diagnosis of hereditary Factor X (FX) deficiency, confirmed by baseline plasma FX levels (FX:C) less than or equal to 10% Patient weight 		
	• Fallent Weight		
	Routine Prophylaxis		
	Documented baseline frequency of bleeding episodes		
	g opiocaco		
	Perioperative Management		
	Documentation of scheduled procedure with intent to use Coagadex for perioperative		
	management of bleeding episodes		
Appropriate	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced		
Treatment			
Regimen & Other	Reauthorization		
Criteria:	Prophylaxis: Reauthorization requires documentation of treatment plan and		
	responsiveness to therapy, defined as a reduction in spontaneous bleeds requiring treatment		
	Treatment: Reauthorization requires documentation of treatment plan, number of acute		
	bleeds since last approval, and number of doses on-hand (not to exceed 6 total doses)		
Exclusion Criteria:			
Age Restriction:			
Prescriber/Site of	Prescribed by, or in consultation with, a hematologist		
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care		
Coverage Duration:	Prophylaxis/On-demand:		
	 Initial Authorization: 3 months, unless otherwise specified 		
	Reauthorization: 12 months, unless otherwise specified		
	Perioperative:		
	Authorization: 1 month, unless otherwise specified		
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COMPOUNDED MEDICATION

Affected Medications: ALL COMPOUNDED MEDICATIONS

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design.
Required Medical Information:	All compounded ingredients must be submitted on the pharmacy claim
Appropriate Treatment Regimen & Other Criteria:	 Compounded medications will only be payable after <u>ALL</u> commercially available or formulary products have been exhausted In the case of payable claim, only compound ingredients that are covered on the applicable formulary will be reimbursed under this policy Compounds above a certain dollar threshold will be stopped by the claim adjudication system
Exclusion Criteria:	 Compounds for experimental or investigational uses will not be covered Compounds containing non-Food and Drug Administration (FDA) approved ingredients will not be covered Compounded medications will not be covered when an Food and Drug Administration (FDA) approved, commercially available medication is on the market for treatment of requested condition
Age Restriction: Prescriber/Site of Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 3 months, unless otherwise specified



CONTINUOUS GLUCOSE MONITORS

Preferred Products: Freestyle Libre 7, Freestyle Libre 2, Freestyle Libre 2 Plus, Freestyle Libre 3, Freestyle Libre 3 Plus, Dexcom G6, Dexcom G7

Non-Preferred Products: Medtronic Products (Enlite, Guardian, Minimed Guardian, Sof-sensor), Eversense Products

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
Required Medical Information:	 Documentation of diabetes mellitus diagnosis Currently on insulin treatment of at least 3 subcutaneous (SubQ) injections daily OR on an insulin pump Performing at least 4 blood glucose tests per day with a home blood glucose monitoring device
Appropriate Treatment Regimen & Other Criteria:	 Coverage for non-preferred continuous glucose monitoring devices and supplies (receiver, transmitter, sensor) must meet the following criteria: Current use of insulin pump that is only compatible with a non-preferred continuous glucose monitor
Exclusion Criteria:	 Type 2 diabetes not on intensive insulin therapy Use of continuous glucose monitor while on dialysis
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Must utilize pharmacy benefits only for coverage of all continuous glucose monitoring systems All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 2 years, unless otherwise specified



POLICY NAME: CORLANOR

Affected Medications: CORLANOR (ivabradine), IVABRADINE

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Stable symptometic charging boart failure with reduced direction fraction in adult.
	Stable, symptomatic chronic heart failure with reduced ejection fraction in adult
	patients (adjunctive therapy)
	Stable, symptomatic heart failure due to dilated cardiomyopathy (DCM) in
	pediatric patients 6 months and older
	Inappropriate sinus tachycardia
Required Medical	Chronic heart failure in adult patients
Information:	 Documentation of chronic heart failure with left ventricular ejection fraction (LVEF) 35% or less AND
	Resting heart rate of at least 70 beats per minute (bpm)
	Heart failure in pediatric patients
	Documentation of stable symptomatic disease due to DCM
	Currently in sinus rhythm with an elevated heart rate
	Inappropriate sinus tachycardia
	Documented resting heart of at least 100 beats per minute, with a mean heart rate of at
	least 90 beats per minute over 24 hours, that is not due to appropriate physiologic
	response or primary abnormality (such as hyperthyroidism or anemia)
	 Symptoms are present (such as palpitations, shortness of breath, dizziness, and/or decreased exercise capacity)
	Documented absence of identifiable causes of sinus tachycardia and exclusion of atrial
	tachycardia
Appropriate	Chronic heart failure in adult patients
Treatment	Documented treatment failure with a beta blocker (metoprolol succinate extended)
Regimen & Other	release, carvedilol, or carvedilol extended release) at the maximally tolerated dose for
Criteria:	heart failure treatment OR
	Documentation of contraindication to beta-blocker use
	Heart failure in pediatric patients
	Treatment failure with beta blocker or digoxin, or contraindication to beta blocker and
	digoxin use.
	All Indications
	Requests for brand Corlanor tablets will require documentation of an adverse event with
	generic ivabradine tablets (and the adverse event was not an expected adverse event
	attributed to the active ingredient)
	Requests for Corlanor oral solution will require at least ONE of the following:
	Request is for a pediatric patient
	 Request is for an adult patient who is unable to swallow tablets
	 Documentation of an adverse event with generic ivabradine tablets (and the
	adverse event was not an expected adverse event attributed to the active



	ingredient)
	 Reauthorization requires documentation of treatment success and a clinically significant response to therapy Development of atrial fibrillation while on therapy will exclude patient from reauthorization
Exclusion Criteria:	 Acute, decompensated heart failure Blood pressure less than 90/50 mm Hg Sick sinus syndrome, sinoatrial block, third-degree atrioventricular block (unless stable with functioning demand pacemaker) Severe hepatic impairment (Child-Pugh class C) Heart rate maintained exclusively by pacemaker
Age Restriction:	Heart failure due to DCM: 6 months to less than 18 years of age
Prescriber/Site of	Prescribed by, or in consultation with, a cardiologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



COVERAGE OF SELECT HIGH INTENSITY STATINS AT TIER 0 COPAY

Affected Medications: ATORVASTATIN (40 mg, 80 mg), ROSUVASTATIN (20 mg, 40 mg), SIMVASTATIN (80 mg)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Primary prevention of cardiovascular disease
Required Medical	Primary prevention of cardiovascular disease (must meet all of the following):
Information:	40 to 75 years of age
	Presence of at least one cardiovascular risk factor such as:
	o Dyslipidemia
	o Diabetes
	 Hypertension
	 Smoking
	Estimated 10-year risk of cardiovascular event of at least 10% or higher
Appropriate	
Treatment	
Regimen & Other	
Criteria:	
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of	All approvals are subject to utilization of the most cost-effective site of care
Care Restrictions:	
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: CRINECERFONT

Affected Medications: CRENESSITY (crinecerfont)

All Food and Drug Administration (FDA) approved indications not otherwise excluded by
plan design
Congenital adrenal hyperplasia (CAH)
Confirmed diagnosis of classic CAH due to 21-hydroxylase deficiency (21-OHD)
confirmed by one of the following
Elevated 17-hydroxyprogestone level
 Confirmed cytochrome CYP21A2 genotype
 Positive newborn screening with confirmatory second-tier testing (such as liquid
chromatography tandem mass spectrometry)
Cosyntropin stimulation test
Documentation of being used concurrently with a systemic glucocorticoid (such as budge participate produing an admiration of the system
hydrocortisone, prednisone, prednisolone, dexamethasone)
Body surface area (BSA)
Requests for oral solution must have documented inability to swallow tablets
Documentation of being on a supraphysiologic systemic glucocorticoid dose to control
disease (total glucocorticoid dose of at least 10 mg/m²/day in hydrocortisone dose
equivalents)
Dosing is in accordance with FDA labeling
Reauthorization requires documentation of treatment success defined by a reduction in serum androstenedione (A4) or reduction in glucocorticoid dose
Serum androstenedione (A4) or reduction in glucoconticold dose
4 years of age and older
4 years or age and order
Prescribed by, or in consultation with, an endocrinologist
All approvals are subject to utilization of the most cost-effective site of care
Initial Authorization: 6 months, unless otherwise specified
Reauthorization:12 months, unless otherwise specified



POLICY NAME: CRIZANLIZUMAB

Affected Medications: ADAKVEO (crizanlizumab)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design To reduce the frequency of vaso-occlusive crises (VOCs) in adults and pediatric patients aged 16 years and older with sickle cell disease
Required Medical Information:	 Diagnosis of sickle cell disease confirmed by genetic testing Two or more sickle cell-related crises in the past 12 months Therapeutic failure of 6-month trial on maximum tolerated dose of hydroxyurea or intolerable adverse event to hydroxyurea
Appropriate Treatment Regimen & Other Criteria:	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced Reauthorization requires documentation of treatment success defined by a decrease in the number of vaso-occlusive crises
Exclusion Criteria:	 Long-term red blood cell transfusion therapy Hemoglobin is less than 4.0 g/dL Chronic anticoagulation therapy (such as warfarin, heparin) other than aspirin History of stroke within the past 2 years Combined use with Endari (L-glutamine)
Age Restriction:	16 years of age and older
Prescriber Restrictions:	 Prescribed by, or in consultation with, a hematologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: CROVALIMAB

Affected Medications: PIASKY (crovalimab)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise excluded by
Covered Oses.	plan design
	5
Required Medical	
Information:	 Presence of at least 2 different glycosylphosphatidylinositol (GPI) protein deficiencies (e.g., CD55, CD59, etc.) within at least 2 different cell lines (e.g., granulocytes, monocytes, erythrocytes)
	Baseline lactate dehydrogenase (LDH) levels greater than or equal to 2 times the upper limit of normal range
	One of the following PNH-associated clinical findings: Processes of a through stip system
	 Presence of a thrombotic event Presence of organ damage secondary to chronic hemolysis
	 Presence of organ damage secondary to chronic hemolysis History of 4 or more blood transfusions required in the previous 12 months
	Body weight
Appropriate	Documented inadequate response, contraindication, or intolerance to ravulizumab-cwvz
Treatment	(Ultomiris)
Regimen & Other	Dosing is in accordance with FDA labeling and most recent body weight
Criteria:	
Ontona.	Reauthorization requires documentation of treatment success defined as a decrease in serum LDH, stabilized/improved hemoglobin, decreased transfusion requirement, and reduction in thromboembolic events compared to baseline
Exclusion Criteria:	 Concurrent use with other biologics for PNH (Soliris, Ultomiris, Empaveli, Fabhalta) Current meningitis infection or other unresolved serious infection caused by encapsulated bacteria
Age Restriction:	13 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a hematologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



CYSTARAN, CYSTADROPS

Affected Medications: CYSTARAN SOLUTION 0.44 % OPHTHALMIC (cysteamine hydrochloride solution), CYSTADROPS SOLUTION 0.37% OPHTHALMIC (cysteamine hydrochloride solution)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design. Ocular Cystinosis
Required Medical Information:	 Diagnosis of ocular cystinosis Documentation of slit-lamp examination showing corneal cystine crystal accumulation
Appropriate Treatment Regimen & Other Criteria:	Reauthorization requires documentation of treatment success defined as reduction in cystine crystals compared to baseline
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an ophthalmologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: CYSTEAMINE

Affected Medications: PROCYSBI (cysteamine bitartrate delayed release)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Nephropathic cystinosis
Required Medical Information:	Diagnosis of nephropathic cystinosis confirmed by ONE of the following:
Appropriate Treatment Regimen & Other Criteria: Exclusion	Documented treatment failure or intolerable adverse event with Cystagon
Criteria: Age Restriction:	
Prescriber/Site of Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: **DANICOPAN**

Affected Medications: VOYDEYA (danicopan)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise excluded by
	plan design
	 Treatment of extravascular hemolysis (EVH) in adults with paroxysmal nocturnal hemoglobinuria (PNH)
Required Medical	Patients must be administered a meningococcal vaccine at least two weeks prior to
Information:	initiation of the requested therapy and revaccinated according to current Advisory Committee on Immunization Practices (ACIP) guidelines
Appropriate	Must be used in combination with ravulizumab-cwvz (Ultomiris) or eculizumab (Soliris)
Treatment	[separate authorization required]
Regimen & Other	Documentation of clinically significant extravascular hemolysis (EVH) defined as
Criteria:	persistent anemia (Hgb less than or equal to 9.5 gram/deciliter) with absolute reticulocyte
	count greater than or equal to 120 x 109/liter despite use of Ultomiris or Soliris for at least
	6 months
	Reauthorization requires documentation of treatment success defined as a decrease in
	serum LDH, stabilized/improved hemoglobin, decreased transfusion requirement, and reduction in thromboembolic events compared to baseline
Exclusion Criteria:	Use without Ultomiris or Soliris
	Concurrent use with biologics for PNH other than Ultomiris and Soliris (such as
	pegcetacoplan or iptacopan)
	Current meningitis infection
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a hematologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Carrage Dringtham	
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **DASATINIB**

Affected Medications: DASATINIB

Covered Uses:	NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher
Required Medical Information:	 Documentation of performance status, all prior therapies used, and prescribed treatment regimen Documentation of Philadelphia chromosome or BCR::ABL1-positive mutation status
Appropriate Treatment Regimen & Other Criteria:	For patients with Chronic Myeloid Leukemia (CML) and low risk score, documented clinical failure with imatinib Reauthorization requires documentation of disease responsiveness to therapy (as applicable, BCR-ABL1 transcript levels, cytogenetic response)
Exclusion Criteria:	Karnofsky Performance Status less than or equal to 50% or ECOG performance score greater than or equal to 3
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **DEFIBROTIDE**

Affected Medications: DEFITELIO (defibrotide sodium)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of adult and pediatric patients with hepatic veno-occlusive disease (VOD), also known as sinusoidal obstruction syndrome (SOS), with renal or pulmonary dysfunction following hematopoietic stem-cell transplantation (HSCT)
Required	Diagnosis of, or high suspicion for, classical or late-onset hepatic VOD
Medical Information:	Weight prior to HSCT, dose, and frequency
Appropriate Treatment Regimen & Other Criteria:	Requested dose within the FDA-approved label
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 2 months with no reauthorization, unless otherwise specified



POLICY NAME: **DEFLAZACORT**

Affected Medications: DEFLAZACORT

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Duchenne muscular dystrophy (DMD) in patients 2 years of age and older
Required Medical Information:	 Laboratory confirmation of Duchenne muscular dystrophy (DMD) diagnosis by genetic testing and serum creatinine kinase at least 10 times the upper limit of normal prior to starting treatment Baseline motor function assessment from one of the following: 6-minute walk test North Star Ambulatory Assessment (NSAA) Motor Function Measure (MFM) Hammersmith Functional Motor Scale (HFMS)
Appropriate Treatment Regimen & Other Criteria:	Documented treatment failure with a 6-month trial of prednisone, or intolerable adverse event causing one of the following:
Exclusion Criteria:	
Age Restriction:	2 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



DELANDISTROGENE MOXEPARVOVEC-ROKL

Affected Medications: ELEVIDYS (delandistrogene moxeparvovec-rokl)

Covered Uses:	Comp Food and Drug Administration (FDA) approved indications not otherwise scaled a
Covered Uses.	 Some Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	 Treatment of ambulatory pediatric patients ages 4 and up with Duchenne muscular dystrophy (DMD)
Required Medical	Confirmed mutation of DMD gene between exons 18-58
Information:	Documentation of being ambulatory without needing an assistive device such as a
	wheelchair, walker, or cane
	North Star Ambulatory Assessment (NSAA) scale total score of 17 or more
	Receiving physical and/or occupational therapy
	 Baseline anti-AAVrh74 total binding antibody titer of less than 1:400 as measured by ELISA
	Current weight
Appropriate	Documentation of being on a stable dose of an oral corticosteroid such as prednisone for
Treatment	at least 12-weeks, and will continue prior to and following Elevidys infusion, according to
Regimen & Other	FDA approved labeling
Criteria:	Does not exceed FDA approved dosing based on weight and maximum of 70 vials
	Number of vials needed = patient body weight (kg) rounded to nearest number of vials
Exclusion Criteria:	Exon 8 and/or exon 9 deletion in DMD gene
	Concomitant therapy or within the past 6 months with DMD-directed antisense
	oligonucleotides such as golodirsen, casimersen, viltolarsen, eteplirsen
	Current active infection
	Previous Elevidys treatment in their lifetime
	Acute liver disease or impaired liver function
	Treatment in non-ambulatory patients – at this time, this indication is not considered medically necessary due to insufficient available evidence of therapeutic value
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 1 month (one-time dose, no reauthorization), unless otherwise specified



POLICY NAME: **DIFELIKEFALIN**

Affected Medications: KORSUVA (difelikefalin)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Chronic kidney disease-associated pruritus (CKD-aP) during hemodialysis (HD)
	Official kidney disease-associated pruntus (OND-air) during hemodialysis (110)
Required Medical Information:	 Documentation of chronic kidney disease confirmed by presence of kidney damage or decreased kidney function for three or more months Documentation of moderate to severe pruritus associated with HD Documentation of normal serum parathyroid hormone (PTH), phosphate, calcium, and magnesium levels Documentation of patient's current dry body weight
Appropriate	Documentation of inadequate relief with trial of all of the following therapies (minimum 1)
Treatment	month trial each):
Regimen & Other	A topical agent (such as an emollient or analgesic)
Criteria:	 An oral antihistamine (such as hydroxyzine or diphenhydramine) Gabapentin or pregabalin
	Reauthorization will require documentation of clinically significant improvement or stabilization in pruritus from baseline and continued hemodialysis use
Exclusion Criteria:	Peritoneal dialysis
	Severe hepatic impairment
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a nephrologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: **DINUTUXIMAB**

Affected Medications: UNITUXIN (dinutuximab)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher
Required Medical Information:	 Documentation of performance status, disease staging, all prior therapies used, and anticipated treatment course Documentation of high-risk neuroblastoma diagnosis as defined per the International Neuroblastoma Response Criteria (INRC): An unequivocal histologic diagnosis from tumor tissue by light microscopy [with or without immunohistochemistry, electron microscopy, or increased urine (or serum) catecholamines or their metabolites] OR Evidence of metastases to bone marrow on an aspirate or trephine biopsy with concomitant elevation of urinary or serum catecholamines or their metabolites Evidence of high-risk neuroblastoma, including: Stage 2/3/4/4S disease with amplified MYCN gene (any age) Stage 4 disease in patients greater than 18 months of age Documented history of previous treatment with at least a partial response to prior first-line multi-agent, multimodality therapy
Appropriate Treatment Regimen & Other Criteria:	Maximum duration: 5 cycles Reauthorization will require documentation of disease responsiveness to therapy
Exclusion Criteria:	Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
Age Restriction:	Under 18 years of age
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an oncologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 5 months, unless otherwise specified



DIROXIMEL FUMARATE

Affected Medications: VUMERITY (diroximel fumarate)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of relapsing forms of multiple sclerosis (MS), including the following: Clinically isolated syndrome (CIS) Relapsing-remitting multiple sclerosis (RRMS) Active secondary progressive disease (SPMS)
Required Medical Information:	Diagnosis confirmed with magnetic resonance imaging (MRI), per revised McDonald diagnostic criteria for MS Clinical evidence alone will suffice; additional evidence desirable but must be consistent with MS
Appropriate Treatment Regimen & Other Criteria:	Relapsing forms of MS Coverage of Vumerity (diroximel fumarate) requires documentation of one of the following: Documented disease progression or intolerable adverse event with one of the following: teriflunomide, dimethyl fumarate or fingolimod Currently receiving treatment with Vumerity (diroximel fumarate), excluding via samples or manufacturer's patient assistance program Reauthorization requires provider attestation of treatment success
Exclusion Criteria:	Concurrent use of other disease-modifying medications indicated for the treatment of multiple sclerosis
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist or a multiple sclerosis specialist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: **DOJOLVI**

Affected Medications: DOJOLVI (triheptanoin oral liquid)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design. A source of calories and fatty acids for the treatment of pediatric and adult patients with molecularly confirmed long-chain fatty acid oxidation disorders
Required Medical Information:	 Diagnosis of long chain fatty acid oxidation disorder (LC-FAOD) confirmed by molecular genetic testing or enzyme assay Documentation of total prescribed daily caloric intake Documentation of severe disease as evidenced by one of the following: Hypoglycemia after short periods of fasting Evidence of functional cardiomyopathy with poor ejection fraction requiring ongoing management Frequent severe major medical episodes requiring emergency room visits, acute care, or hospitalization (3 events within the past year, or 5 events within the past 2 years) Elevated creatinine kinase (chronic or episodic)
Appropriate Treatment Regimen & Other Criteria:	 Documentation of persistent symptoms despite dietary management and use of an over the counter (OTC) medium-chain triglyceride (MCT) product Dose not to exceed 35% of daily caloric intake Reauthorization will require documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	Concurrent use of another medium chain triglyceride product
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an endocrinologist or provider experienced in the management of metabolic disorders All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 3 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **DONISLECEL**

Affected Medications: LANTIDRA (donislecel solution)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
Required Medical Information:	 Diagnosis of type 1 diabetes for 5 or more years Documentation of inability to achieve target HbA1c despite adherence to intensive insulin management with all the following: Multiple daily injections of prandial and basal insulin or on an insulin pump Performing at least four blood glucose tests per day or using a continuous glucose monitor Documentation of 2 or more episodes of severe hypoglycemia (blood glucose level less than 50 mg/dL) in the past three years requiring assistance of another person with either an oral carbohydrate, intravenous glucose, or glucagon administration Documentation of hypoglycemia unawareness, defined by the absence of adequate autonomic symptoms during an episode of severe hypoglycemia
Appropriate Treatment Regimen & Other Criteria:	Reauthorization requires documentation of not achieving exogenous insulin independence within one year of infusion or within one year of losing independence from exogenous insulin (maximum of three infusions per lifetime)
Exclusion Criteria:	 Pregnancy Malignancy Active infection Previous kidney or pancreas transplant Prior portal vein thrombosis
Age Restriction:	18 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an endocrinologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 3 months (single treatment), unless specified otherwise



POLICY NAME: **DORNASE ALFA**

Affected Medications: PULMOZYME (dornase alfa)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
Required Medical Information:	 The diagnosis of Cystic Fibrosis (CF) has been confirmed by appropriate diagnostic or genetic testing Additional testing should include evaluation of overall clinical lung status and respiratory function (e.g., pulmonary function tests, lung imaging, etc.)
Appropriate Treatment Regimen & Other Criteria:	 Pulmozyme will be used in conjunction with standard therapies for cystic fibrosis Reauthorization will require documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	
Age Restriction:	1 month of age or older
Prescriber/Site of Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 24 months, unless otherwise specified



POLICY NAME: **DROXIDOPA**

Affected Medications: DROXIDOPA

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of orthostatic dizziness with symptomatic neurogenic orthostatic hypotension (nOH) caused by: Primary autonomic failure (Parkinson's disease [PD], multiple system atrophy [MSA], pure autonomic failure [PAF]) Dopamine beta-hydroxylase deficiency Non-diabetic autonomic neuropathy
Required Medical Information:	 Diagnosis of nOH caused by one of the following: Primary autonomic failure (such as PD, MSA, PAF) Dopamine beta-hydroxylase deficiency Non-diabetic autonomic neuropathy Documentation of severe symptomatic orthostatic hypotension, demonstrated by both of the following: Minimum 20 mmHg decrease in systolic blood pressure OR minimum 10 mmHg decrease in diastolic blood pressure within 3 minutes of standing Documentation of significant symptoms affecting activities of daily living
Appropriate Treatment Regimen & Other Criteria:	Documented treatment failure or intolerable adverse event with a minimum 30-day trial to both fludrocortisone and midodrine
	Reauthorization requires documentation of treatment success as determined by treating provider
Exclusion Criteria:	
Age Restriction:	18 years of age or older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist or cardiologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 1 month, unless otherwise specified Reauthorization: 3 months, unless otherwise specified



DUOPA

Affected Medications: DUOPA (carbidopa-levodopa enteral suspension)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	 Treatment of motor fluctuations in patients with advanced Parkinson's disease (PD)
Required Medical Information:	Documentation of all the following: Diagnosis of advanced PD Clear response to levodopa treatment with evidence of "On" periods Persistent motor fluctuations with "Off" time occurring 3 hours or more per day while awake despite an optimized PD treatment regimen Has undergone or has planned placement of a nasojejunal (NJ) tube for temporary administration of Duopa OR gastrostomy-jejunostomy (PEG-J) tube for long-term administration of Duopa
Appropriate Treatment Regimen & Other Criteria:	Documented treatment failure with both of the following:
Exclusion Criteria:	 response to therapy Atypical Parkinson's syndrome ("Parkinson's Plus" syndrome) or secondary Parkinson's Non-levodopa responsive PD Contraindication to percutaneous endoscopic gastro-jejunal (PEG-J) tube placement or long-term use of a PEG-J Concomitant use with nonselective MAO inhibitors or have recently (within 2 weeks) taken a nonselective MAO inhibitor
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: **DUPILUMAB**

Affected Medications: DUPIXENT (dupilumab)

Covered Uses:	All Food and Drug Administration (FDA)—approved indications not otherwise excluded by
	plan design
	 Moderate to severe eosinophilic phenotype or oral corticosteroid dependent asthma
	 Moderate to severe atopic dermatitis (AD)
	Chronic rhinosinusitis with nasal polyposis (CRSwNP)
	Eosinophilic esophagitis (EoE)
	Prurigo nodularis (PN)
	Chronic Obstructive Pulmonary Disease (COPD)
Required Medical	AD:
Information:	Documentation of severe inflammatory skin disease defined as functional impairment
	(inability to use hands or feet for activities of daily living or significant facial involvement
	preventing normal social interaction)
	Body surface area (BSA) involvement greater than or equal to 10% or hand, foot, or
	mucous membrane involvement
	Asthma:
	Documentation of BOTH of the following:
	 Baseline eosinophil count at least 150 cells/μL
	 Forced expiratory volume (FEV1) less than 80% at baseline or FEV1/FVC
	reduced by at least 5% from normal
	CRSwNP:
	Documentation of both of the following:
	 Diagnosis of chronic rhinosinusitis and has undergone prior bilateral total ethmoidectomy
	 Indicated for revision sinus endoscopic sinus surgery due to recurrent symptoms
	of nasal polyps (such as nasal obstruction/congestion, bilateral sinus obstruction)
	EoE:
	Diagnosis confirmed by endoscopic biopsy with greater than or equal to 15 eosinophils
	per high power field (HPF)
	Documentation of TWO or more dysphagia episodes per week despite current treatment
	PN:
	Documentation of all the following:
	o Diagnosis confirmed by skin biopsy
	Presence of at least 20 PN lesions for at least 3 months
	 Severe itching
	COPD



	Diagnosis of COPD with moderate to severe airflow limitation
	FEV1/FVC ratio less than 0.7 and FEV1 of 30-70% predicted
	Baseline eosinophil count of at least 300 cells/µL
	Symptoms of chronic productive cough for at least 3 months
Appropriate	Requested dosing according to the FDA label based on diagnosis
Treatment	
Regimen & Other	AD:
Criteria:	Documented treatment failure with at least 12 weeks of two of the following (1 in each
	category):
	 Tacrolimus ointment or pimecrolimus cream or Eucrisa
	 Phototherapy or cyclosporine or azathioprine or methotrexate or mycophenolate
	Asthma:
	Use of high-dose inhaled corticosteroid (ICS) plus a long-acting beta agonist (LABA) for
	at least three months with continued symptoms
	Documentation of one of the following:
	 Documented history of 2 or more asthma exacerbations requiring oral or
	systemic corticosteroid treatment in the past 12 months while on combination
	inhaler treatment with at least 80% adherence
	Documentation that chronic daily oral corticosteroids are required
	Bootimentation that official daily oral controls are required
	CRSwNP:
	Documented treatment failure with at least 1 intranasal corticosteroid (such as
	fluticasone) after ethmoidectomy
	Documented treatment failure with Sinuva implant
	Boodmonted treatment failule with omava implant
	EoE:
	Documented treatment failure with at least 12 weeks of BOTH of the following:
	High dose (twice daily dosing) proton pump inhibitor (e.g., omeprazole or
	esomeprazole)
	O allo a language and the second of the seco
	Swallowed corticosteroid therapy (such as fluticasone or budesonide)
	PN:
	Documented treatment failure with at least 2 weeks of a super high potency topical
	corticosteroid (such as clobetasol propionate 0.05%, halobetasol propionate 0.05%)
	Documentation of treatment failure with at least 12 weeks of one of the following:
	phototherapy, methotrexate, cyclosporine
	CORD
	COPD Description of inhelial triple the group consisting of a large acting revised in in-
	Documented use of inhaled triple therapy consisting of a long-acting muscarinic The population has a consist (LARA) and inhaled continuous actions have a consist (LARA).
	antagonist (LAMA), long-acting beta agonist (LABA), and inhaled corticosteroid (ICS) for
	at least 12 weeks with continued symptoms

Documentation of one of the following:

History of at least two moderate COPD exacerbations requiring treatment with a systemic corticosteroid and/or an antibiotic in the past year while adherent on



	triple therapy and at least 80% adherence
	 History of at least one severe COPD exacerbation requiring hospitalization in the
	past year while adherent on triple therapy and at least 80% adherence
	Reauthorization requires documentation of treatment success as determined by treating provider
Exclusion Criteria:	Concurrent use with another therapeutic immunomodulator agent utilized for the same indication
Age Restriction:	AD: 6 months of age and older
	Asthma: 6 years of age and older
	CRSwNP: 12 years of age and older
	EoE: 1 year of age and older
	PN: 18 years of age and older
	COPD: 18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a dermatologist, pulmonologist, otolaryngologist,
Care Restrictions:	gastroenterologist, allergist, or immunologist
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **ECULIZUMAB**

Affected Medications: SOLIRIS (eculizumab)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	 Paroxysmal nocturnal hemoglobinuria (PNH) to reduce hemolysis
	 Atypical hemolytic uremic syndrome (aHUS) to inhibit complement-mediated
	thrombotic microangiopathy
	 Generalized myasthenia gravis (gMG) in adults who are anti-acetylcholine
	receptor (AChR) antibody positive
	 Neuromyelitis optica spectrum disorder (NMOSD) in adult patients who are anti- aquaporin-4 (AQP4) antibody positive
Required Medical	<u>PNH</u>
Information:	Detection of PNH clones of at least 5% by flow cytometry diagnostic testing
	 Presence of at least 2 different glycosylphosphatidylinositol (GPI) protein
	deficiencies (e.g., CD55, CD59, etc.) within at least 2 different cell lines (e.g.,
	granulocytes, monocytes, erythrocytes)
	Baseline lactate dehydrogenase (LDH) levels greater than or equal to 1.5 times the upper
	limit of normal range
	One of the following PNH-associated clinical findings:
	 Presence of a thrombotic event
	 Presence of organ damage secondary to chronic hemolysis
	 History of 4 or more blood transfusions required in the previous 12 months
	aHUS
	Clinical presentation of microangiopathic hemolytic anemia, thrombocytopenia, and acute
	kidney injury
	Patient shows signs of thrombotic microangiopathy (TMA) (e.g., changes in mental)
	status, seizures, angina, dyspnea, thrombosis, increasing blood pressure, decreased
	platelet count, increased serum creatinine, increased LDH, etc.)
	ADAMTS13 activity level greater than or equal to 10%
	Shiga toxin E. coli related hemolytic uremic syndrome (ST-HUS) has been ruled out
	History of 4 or more blood transfusions required in the previous 12 months
	gMG
	 Diagnosis of gMG confirmed by ONE of the following:
	A history of abnormal neuromuscular transmission test
	 A positive edrophonium chloride test
	 Improvement in gMG signs or symptoms with an acetylcholinesterase inhibitor
	M
	Positive serologic test for AChR antibodies Decumentation of ONE of the following:
	Documentation of ONE of the following: MC Activities of Polly Living (MC API) total score of 6 or greater.
	MG-Activities of Daily Living (MG-ADL) total score of 6 or greater Quantitative Myesthopia Gravia (QMG) total score of 13 or greater.
	 Quantitative Myasthenia Gravis (QMG) total score of 12 or greater
	<u>NMOSD</u>



- Diagnosis of seropositive aquaporin-4 immunoglobulin G (AQP4-IgG) NMOSD confirmed by all the following:
 - o Documentation of AQP4-IgG-specific antibodies on cell-based assay
 - Exclusion of alternative diagnoses (such as multiple sclerosis)
 - At least ONE core clinical characteristic:
 - Acute optic neuritis
 - Acute myelitis
 - Acute area postrema syndrome (episode of otherwise unexplained hiccups or nausea/vomiting)
 - Acute brainstem syndrome
 - Symptomatic narcolepsy OR acute diencephalic clinical syndrome with NMOSD-typical diencephalic lesion on magnetic resonance imaging (MRI) [see table below]
 - Acute cerebral syndrome with NMOSD-typical brain lesion on MRI [see table below]

Clinical presentation	Possible MRI findings
Diencephalic syndrome	Periependymal lesionHypothalamic/thalamic lesion
Acute cerebral syndrome	 Extensive periependymal lesion Long, diffuse, heterogenous, or edematous corpus callosum lesion Long corticospinal tract lesion Large, confluent subcortical or deep white matter lesion

Appropriate Treatment Regimen & Other Criteria:

PNH

 Documented inadequate response, contraindication, or intolerance to ravulizumab-cwvz (Ultomiris)

aHUS

- Failure to respond to plasma therapy within 10 days
 - o Trial of plasma therapy not required if one of the following is present:
 - Life-threatening complications of HUS such as seizures, coma, or heart failure
 - Confirmed presence of a high-risk complement genetic variant (e.g., CFH or CFI)
- Documented inadequate response, contraindication, or intolerance to ravulizumab-cwvz (Ultomiris)

gMG

- Documentation of one of the following:
 - Treatment failure with an adequate trial (one year or more) of at least 2 immunosuppressive therapies (azathioprine, mycophenolate, tacrolimus, cyclosporine, methotrexate)
 - Has required three or more courses of rescue therapy (plasmapheresis/plasma exchange and/or intravenous immunoglobulin), while on at least one immunosuppressive therapy, over the last 12 months



	Documented inadequate response, contraindication, or intolerance to each of the following:
	 Reauthorization: gMG: documentation of treatment success defined as an improvement in MG-ADL and QMG scores from baseline NMOSD: documentation of treatment success defined as the stabilization or improvement in neurological symptoms as evidenced by a decrease in acute relapses, Expanded Disability Status Scale (EDSS) score, hospitalizations, or plasma exchange treatments PNH: documentation of treatment success defined as a decrease in serum LDH, stabilized/improved hemoglobin, decreased transfusion requirement, and reduction in thromboembolic events compared to baseline aHUS: documentation of treatment success defined as a decrease in serum LDH, stabilized/improved serum creatinine, increased platelet count, and decreased plasma exchange/infusion requirement compared to baseline
Exclusion Criteria:	 Concurrent use with other disease-modifying biologics for requested indication, unless otherwise indicated by the FDA for combination use with Soliris Current meningitis infection
Age Restriction:	 PNH, gMG and NMOSD: 18 years of age and older aHUS: 2 months of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a specialist PNH: hematologist aHUS: hematologist or nephrologist gMG: neurologist NMOSD: neurologist or neuro-ophthalmologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 3 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **EDARAVONE**

Affected Medications: RADICAVA (edaravone), RADICAVA ORS

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Amyotrophic lateral sclerosis (ALS)
Required Medical Information:	 Documentation of "definite" or "probable" ALS diagnosis based on revised El Escorial (Airlie House) or Awaji criteria Disease duration of 2 years or less Normal respiratory function defined as percent-predicted forced vital capacity values (% FVC) of at least 80% Patient currently retains most activities of daily living (ADLs), defined as at least 2 points on all 12 items of the ALS functional rating scale-revised (ALSFRS-R)
Appropriate Treatment Regimen & Other Criteria:	 Reauthorization requires both of the following: Documentation of treatment success, as determined by prescriber (e.g., retention of most ADLs) Patient is not dependent on invasive mechanical ventilation (e.g., intubation, tracheostomy)
Exclusion Criteria: Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist or provider with experience in treating ALS All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **EFLORNITHINE**

Affected Medications: IWILFIN (eflornithine)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
Required Medical Information:	 Maintenance therapy in patients with high-risk neuroblastoma who achieve at least a partial response to prior systemic agents and have completed maintenance immunotherapy with an anti-GD2 antibody NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher Documentation of performance status, disease staging, all prior therapies used, and anticipated treatment course Diagnosis of neuroblastoma as defined per the International Neuroblastoma Response Criteria (INRC): An unequivocal histologic diagnosis from tumor tissue by light microscopy [with or without immunohistochemistry, electron microscopy, or increased urine (or serum) catecholamines or their metabolites] OR Evidence of metastases to bone marrow on an aspirate or trephine biopsy with concomitant elevation of urinary or serum catecholamines or their metabolites Evidence of high-risk neuroblastoma, including: Stage 2/3/4/4S disease with amplified MYCN gene (any age) Stage 3 disease with MYCN gene NOT amplified in patients at least 18 months of age with International Neuroblastoma Pathology Classification (INPC) as unfavorable histology (UH) Stage 4 disease in patients greater than 12 months of age Staging studies documented by histology and/or appropriate imaging as follows:
	 Computed tomography (CT) or magnetic resonance imaging (MRI) scan of the primary site and nodal sites of metastatic disease Bone imaging (preferably with a metaiodobenzylguanidine [MIBG] scan and
	positron emission topography (PET) scan (if MIBG is negative)
Appropriate	Documentation of a partial response to prior systemic agents and completed
Treatment	maintenance immunotherapy with an anti-GD2 antibody (Dinutuximab, Naxitamab)
Regimen & Other	Reauthorization: documentation of disease responsiveness to therapy up to a total of 2
Criteria:	years of treatment
Exclusion Criteria:	Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial authorization: 4 months, unless otherwise specified Para the initial Continuous of the initial authorization (CO) was the formula of the initial authorization.
	 Reauthorization: One time reauthorization of 20 months to complete 2 years of treatment, unless otherwise specified



ELADOCAGENE EXUPARVOVEC-TNEQ

Affected Medications: KEBILIDI (eladocagene exuparvovec-tneq)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise excluded by
	plan design
	Treatment of aromatic L-amino acid decarboxylase (AADC) deficiency
Required Medical Information:	Diagnosis of AADC deficiency confirmed by genetic testing showing bilateral/biallelic mutations in the DDC gene Particle AADC agreement activities in places.
	Reduced AADC enzyme activity in plasma
	 Cerebrospinal fluid (CSF) shows all of the following: Reduced levels of 5-hydroxyindoleacetic acid (5-HIAA), homovanillic acid (HVA), and 3-methoxy-4-hydroxyphenylglycol (MHPG)
	 Elevated levels of 3-O-methyldopa (3-OMD), levodopa (L-Dopa), and 5- hydroxytryptophan (5-HTP)
	 Normal levels of pterins (neopterin and biopterin)
	Clinical symptoms of AADC deficiency such as movement disorders, hypotonia,
	autonomic dysfunction, and developmental delay
	Documented achieved skull maturity assessed by neuroimaging
Appropriate Treatment	Dosing is in accordance with FDA labeling
Regimen & Other	
Criteria:	
Exclusion Criteria:	Prior gene therapy administration
	Anti-AAV2 neutralizing antibody titer over 1,200 folds
Age Restriction:	1 to 17 years of age
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist or geneticist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 3 months (one-time infusion only), unless otherwise specified



POLICY NAME: **ELAGOLIX**

Affected Medications: ORILISSA (elagolix), ORIAHNN (elagolix/estradiol/norethindrone acetate)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Moderate to severe endometriosis-associated pain (Orilissa)
	 Heavy menstrual bleeding associated with uterine leiomyomas (Oriahnn)
Required Medical	Pain due to endometriosis
Information:	Documentation of both of the following:
	 Diagnosis of moderate to severe pain associated with endometriosis
	Attestation that patient is premenopausal
	Heavy menstrual bleeding due to uterine leiomyomas
	Documentation of both of the following:
	Diagnosis of heavy menstrual bleeding associated with uterine leiomyomas
	Attestation that patient is premenopausal
Appropriate	Pain due to endometriosis
Treatment	Documentation of a trial and inadequate relief (or contraindication) after at least 3
Regimen & Other	months of both of the following first-line therapies:
Criteria:	 Nonsteroidal anti-inflammatory drugs (NSAIDs)
	 Continuous (no placebo pills) hormonal contraceptives
	Board of a factor of the facto
	Reauthorization requires documentation of treatment success and a clinically significant response to therapy
	response to therapy
Exclusion Criteria:	History of osteoporosis
	Pregnancy
	Severe (Child-Pugh Class C) hepatic impairment (Orilissa)
	Mild, moderate, and severe (Child-Pugh Class A, B, and C) hepatic impairment
	(Oriahnn)
Age Restriction:	18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a specialist in obstetrics/gynecology or
Care Restrictions:	reproductive endocrinology
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization: 18 months (Orilissa 150 mg once daily* and Oriahnn only), unless
	otherwise specified



•	*Maximum treatment duration for Orilissa 150 mg once daily in patients with moderate hepatic impairment (Child-Pugh Class B) and Orilissa 200 mg twice daily is 6 months. Reauthorization not allowed
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ELIVALDOGENE AUTOTEMCEL

Affected Medications: SKYSONA (elivaldogene autotemcel)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Early, active cerebral adrenoleukodystrophy (CALD) in male patients
Required Medical Information:	 Confirmed diagnosis of CALD with all of the following: Confirmed ABCD1 gene mutation Elevated very-long-chain fatty acid (VLCFA) values for ALL of the following: Concentration of C26:0 Ratio of C24:0 to C22:0 Ratio of C26:0 to C22:0 Neurologic function score (NFS) less than or equal to 1 (asymptomatic or mildly symptomatic disease) Active central nervous system disease established by central radiographic review of brain magnetic resonance imaging (MRI) demonstrating both of the following:
Appropriate	Coverage of Skysona is provided if the patient does not have access to a hematopoietic
Treatment	stem cell transplant with a matched sibling donor
Regimen & Other Criteria:	Approved for one-time single infusion only
Exclusion Criteria:	 Female gender Previously received an allogeneic transplant or gene therapy
Age Restriction:	4 to 17 years of age
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist, endocrinologist, or
Care Restrictions:	hematologist/oncologist
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 4 months, unless otherwise specified (one infusion only)



ELTROMBOPAG DERIVATIVES

Affected Medications: PROMACTA (eltrombopag olamine), PROMACTA PACKET, ALVAIZ (eltrombopag choline)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Treatment of thrombocytopenia in patients with persistent or chronic immune
	thrombocytopenia (ITP)
	 Treatment of thrombocytopenia in patients with hepatitis C infection
	Treatment of severe aplastic anemia
Required Medical	Thrombocytopenia in patients with chronic ITP
Information:	Documentation of ONE of the following:
	 Platelet count less than 20,000/microliter
	Platelet count less than 30,000/microliter AND symptomatic bleeding Platelet count less than 50,000/microliter AND symptomatic bleeding
	Platelet count less than 50,000/microliter AND increased risk for bleeding (such
	as peptic ulcer disease, use of antiplatelets or anticoagulants, history of bleeding at higher platelet count, need for surgery or invasive procedure)
	at higher platelet count, fleed for surgery or invasive procedure/
	Thrombocytopenia in patients with chronic hepatitis C
	Documentation of plan to initiate interferon-based therapy
	Documentation of platelet count less than 75,000/microliter
	Severe aplastic anemia
	Diagnosis confirmed by bone marrow biopsy
	Documentation of at least two of the following:
	 Absolute reticulocyte count (ARC) less than 60,000/microliter
	 Platelet count less than 20,000/microliter
	Absolute neutrophil count (ANC) less than 500/microliter
Appropriate	Promacta packet formulation requires documented medical inability to use oral tablet
Treatment	formulation
Regimen & Other	
Criteria:	Thrombocytopenia in patients with persistent or chronic ITP
	 Documentation of inadequate response, defined as platelets did not increase to at least
	50,000/microliter, to the following therapies:
	o ONE of the following:
	 Inadequate response with at least 2 therapies for immune
	thrombocytopenia, including corticosteroids, rituximab, or
	immunoglobulin
	 Splenectomy
	Reauthorization:
	Response to treatment with platelet count of at least 50,000/microliter (not to exceed)
	400,000/microliter)
	OR
	The platelet counts have not increased to a platelet count of at least 50,000/microliter and
	the patient has NOT been on the maximum dose for at least 4 weeks



	Thrombocytopenia in patients with chronic hepatitis C
	Reauthorization:
	 Response to treatment with platelet count of at least 90,000/microliter (not to exceed 400,000/microliter) and eltrombopag is used in combination with antiviral therapy
	Severe aplastic anemia
	Documentation of refractory severe aplastic anemia as indicated by insufficient response to at least one prior immunosuppressive therapy OR
	For those less than 40 years of age without a rapidly available matched related donor (MRD) or 40 years of age and older: documentation that eltrombopag is being used as first line treatment in combination with standard immunosuppressive therapy (Atgam and cyclosporine)
	Reauthorization (refractory severe aplastic anemia only): Requires hematologic response to treatment defined as meeting ONE or more of the following criteria:
	Platelet count increases to 20,000/microliter above baseline, or stable platelet counts with transfusion independence for a minimum of 8 weeks
	Hemoglobin increases by greater than 1.5 g/dL or a reduction in greater than or equal to
	4 units red blood cell (RBC) transfusions for 8 consecutive weeks
	ANC increase of 100% or an ANC increase greater than 500/microliter
Exclusion Criteria:	Use in combination with another thrombopoietin receptor agonist, spleen tyrosine kinase inhibitor, or similar treatments (Doptelet, Nplate, Tavalisse)
Age Restriction:	Thrombocytopenia in patients with ITP
	1 year of age and older (Promacta)
	6 years of age and older (Alvaiz)
	Thrombocytopenia in patients with chronic hepatitis C and patients with severe
	aplastic anemia
	18 years of age and older (Promacta and Alvaiz)
	Severe Aplastic Anemia (initial therapy)
	2 years of age and older
	18 years of age and older (Alvaiz)
Prescriber/Site of	Prescribed by, or in consultation with, a hematologist or gastroenterology/liver specialist
Care Restrictions:	All approvals are subjects to utilization of the most cost-effective site of care
Coverage Duration:	Thrombocytopenia in patients with ITP
	Initial Authorization: 4 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified
	Thrombocytopenia in patients with chronic hepatitis C
	Initial Authorization: 2 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified
	Severe aplastic anemia



 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified
 Severe aplastic anemia in combination with cyclosporine and Atgam Authorization: 6 months, no reauthorization, unless otherwise specified



POLICY NAME: **EMAPALUMAB**

Affected Medications: GAMIFANT (emapalumab-lzsg)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise evaluded by
Covered Oses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	Treatment of primary hemophagocytic lymphohisticocytosis (HLH) in patients
	(newborn and older) intolerant to conventional HLH therapy or with refractory,
	recurrent, or progressive disease
Required	Diagnosis confirmed by presence of a genetic mutation known to cause primary HLH
Medical	(e.g., PRF1, UNC13D, STX11, STXBP2) OR documentation showing at least 5 of the
Information:	following are present:
	 Prolonged fever (lasting over 7 days)
	 Splenomegaly
	 Two of the following cytopenias in the peripheral blood:
	 Hemoglobin less than 9 g/dL
	 Platelet count less than 100,000/mcL
	 Neutrophils less than 100/mcL
	One of the following:
	 Hypertriglyceridemia defined as fasting triglycerides 3 mmol/L or higher
	(equivalent to 265 mg/dL or higher)
	 Hypofibrinogenemia defined as fibrinogen 1.5 g/L or lower
	 Hemophagocytosis in bone marrow, spleen, or lymph nodes (with no evidence of
	malignancy)
	 Low or absent natural killer cell activity (according to local laboratory reference)
	 Ferritin 500 mcg/L or higher
	 Soluble CD25 (i.e., soluble IL-2 receptor) 2,400 U/ml or higher
	Documentation confirming status as a hematopoietic stem cell transplant (HCST)
	candidate
Appropriate	Documentation of refractory, recurrent, or progressive disease (or intolerable adverse)
Treatment	event) on conventional HLH therapy (e.g., dexamethasone, etoposide, methotrexate,
Regimen &	hydrocortisone)
Other Criteria:	
	Must be used in combination with dexamethasone, unless currently established on and
	planning to continue one of the following: cyclosporine, glucocorticoids, and/or
	intrathecal methotrexate
	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced
	Reauthorization requires documentation of disease responsiveness to therapy AND patient
	has not yet received HSCT
Exclusion	
Criteria:	
Age	
Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a hematologist, oncologist, transplant specialist, or
Care Restrictions:	provider with experience in the management of HLH
	All approvals are subject to utilization of the most cost-effective site of care
Coverage	Initial Authorization: 2 months, unless otherwise specified
Duration:	Reauthorization: 4 months, unless otherwise specified
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POLICY NAME: **EMICIZUMAB**

Affected Medications: HEMLIBRA (emicizumab-kxwh)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design.
Required Medical	Documented diagnosis of hemophilia A with or without inhibitors
Information:	 Prescribed for routine prophylaxis to prevent or reduce the frequency of bleeding episodes
Appropriate	Baseline factor level less than 1% AND prophylaxis required OR
	Baseline factor level 1% to 3% AND a documented history of at least two episodes of
Regimen & Other	spontaneous bleeding into joints
Criteria:	 Prophylactic agents must be discontinued Factor VIII Inhibitors: after the first week of HEMLIBRA
	Bypassing Agents: one day before starting HEMLIBRA
	Loading Dose:
	3 mg/kg once every week for 4 weeks
	Maximum 1,380 mg per 28 day supply
	Maintenance dose:
	1.5 mg/kg once every week or
	3 mg/kg once every 2 weeks or
	6 mg/kg once every 4 weeks
	 Any increases in dose must be supported by an acceptable clinical rationale (i.e. weight gain, increase in breakthrough bleeding when patient is fully adherent to therapy, etc.)
	Product Availability
	 Single-dose vials for injection: 30 mg/mL, 60 mg/0.4 mL, 105 mg/0.7 mL, 150 mg/mL Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced
	Reauthorization requires documentation of treatment success defined as a reduction in spontaneous bleeds requiring treatment, as well as documentation of bleed history since last approval
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a hematologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 6 months, unless otherwise specified



EMSAM

Affected Medications: EMSAM (selegiline)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of major depressive disorder (MDD)
Required Medical Information:	Diagnosis of major depressive disorder (MDD)
Appropriate Treatment Regimen & Other Criteria:	 Documented treatment failure to an adequate trial (clinically sufficient doses for a minimum 6-week duration) to each of the following: A selective serotonin reuptake inhibitor (SSRI) A serotonin/norepinephrine reuptake inhibitor (SNRI) A tricyclic antidepressant or mirtazapine Bupropion Antidepressant augmentation therapy (e.g., second generation antipsychotic, thyroid hormone, lithium) OR Documentation of inability to take any oral preparations (including commercially available liquid antidepressants) Reauthorization will require documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	Pheochromocytoma
Age Restriction:	18 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a psychiatrist or behavioral health specialist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



ENDOTHELIN RECEPTOR ANTAGONISTS

Affected Medications: BOSENTAN, AMBRISENTAN, OPSUMIT (macitentan), OPSYNVI (macitentan and tadalafil)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Pulmonary artery hypertension (PAH) World Health Organization (WHO) Group 1
Required Medical Information:	 Documentation of Pulmonary Arterial Hypertension (PAH) World Health Organization (WHO) Group 1 confirmed by right heart catheterization meeting the following criteria: Mean pulmonary artery pressure of at least 20 mm Hg Pulmonary capillary wedge pressure less than or equal to 15 mm Hg Pulmonary vascular resistance of at least 2.0 Wood units New York Heart Association (NYHA)/WHO Functional Class II or higher symptoms Documentation of Acute Vasoreactivity Testing (positive result requires trial/failure to calcium channel blocker), unless there are contraindications: Low systemic blood pressure (systolic blood pressure less than 90) Low cardiac index Presence of severe symptoms (functional class IV)
Appropriate Treatment Regimen & Other Criteria:	 Documentation that the drug will be used in combination with a phosphodiesterase-5 (PDE-5) inhibitor Documentation of inadequate response or intolerance to oral calcium channel blocking agents if positive Acute Vasoreactivity Test For Opsumit (macitentan) and Opsynvi (macitentan and tadalafil) requests: documentation of inadequate response or intolerance to ambrisentan AND bosentan for 12 weeks is required Reauthorization requires documentation of treatment success defined as one or more of the following: Improvement in walking distance Improvement in pulmonary function Improvement or stability in WHO functional class
Exclusion Criteria:	improvement or classify in titre tanolicital class
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a cardiologist or pulmonologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: **ENFUVIRTIDE**

Affected Medications: FUZEON (enfuvirtide)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of human immunodeficiency virus type 1 (HIV-1) infection in combination with other antiretroviral agents in treatment-experienced patients with evidence of HIV-1 replication despite ongoing antiretroviral therapy
Required Medical Information:	 Documented weight greater than or equal to 11 kg Documentation of current (within past 30 days) HIV-1 RNA viral load of at least 200 copies/mL Documented treatment failure with minimum 12-weeks of antiretroviral therapy with at least one antiretroviral agent from three different classes (unless contraindicated or clinically significant adverse effects are experienced): Nucleoside reverse-transcriptase inhibitors (NRTIs) Non-nucleoside reverse-transcriptase inhibitors (NNRTIs) Integrase strand transfer inhibitors (INSTIs) Protease inhibitors (PIs)
Appropriate Treatment Regimen & Other Criteria:	 Prescribed in combination with an optimized background antiretroviral regimen Reauthorization requires documentation of all of the following: Treatment plan including continued use of optimized background antiretroviral regimen Documentation of treatment success as evidenced by one of the following:
Exclusion Criteria:	Initial therapy in patients who are antiretroviral naïve
Age Restriction:	6 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an infectious disease or HIV specialist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 24 months, unless otherwise specified



ENZYME REPLACEMENT THERAPY (ERT) FOR GAUCHER DISEASE TYPE 1
Affected Medications: CERDELGA (eliglustat), VPRIV (velaglucerase alfa), CEREZYME (imiglucerase), ELELYSO (taliglucerase alfa)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Vpriv: Gaucher disease type 1 (GD1) Elelyso: GD1 for ages 4 years and older Cerdelga: GD1 in adults who are CYP2D6 extensive metabolizers (EMs), intermediate metabolizers (IMs), or poor metabolizers (PMs) as detected by an FDA-cleared test Cerezyme: GD1 for ages 2 years and older that results in one or more of the following conditions: Anemia Thrombocytopenia Bone disease Hepatomegaly or splenomegaly
Required Medical	Diagnosis confirmed by enzyme assay showing deficiency of beta-glucocerebrosidase
Information:	glucosidase enzyme activity OR genetic testing indicating mutation of two alleles of the
	glucocerebrosidase genome
	 For Cerdelga, must also have documentation of cytochrome P450 2D6 (CYP2D6) genotype by an FDA-approved test indicating CYP2D6 EM, IM, or PM status
	Documentation of baseline tests such as hemoglobin level, platelet count, liver function
	tests, renal function tests
	Documentation of at least one clinically significant disease complication of GD1:
	 Anemia (low hemoglobin and hematocrit levels)
	 Thrombocytopenia (platelet count less than 120,000 mm³)
	 Bone disease (T-score less than -2.5 or bone pain)
	 Hepatomegaly or splenomegaly
	 For symptomatic children: symptoms of early presentation, such as
	malnutrition, growth retardation, impaired psychomotor development, and/or fatigue
Appropriate	<u>Cerdelga</u>
Treatment	Extensive or Intermediate Metabolizers of CVD2D6
Regimen & Other	 Extensive or Intermediate Metabolizers of CYP2D6 Quantity limit - 84 mg capsules #60 per 30 days
Criteria:	- Quantity little 04 mg capacitos #00 per 00 days
	Poor Metabolizers of CYP2D6
	Quantity limit - 84 mg capsules #30 per 30 days
	 Elelyso, Vpriv, and Cerezyme Dosing is in accordance with FDA labeling and patient's most recent weight Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced
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	Reauthorization will require documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	Concomitant use with another ERT for GD1 or with miglustat
	Cerdelga:
	CYP2D6 ultrarapid metabolizers
	Moderate or severe hepatic impairment
	Pre-existing cardiac disease (congestive heart failure, myocardial infarction, bradycardia, heart block, arrhythmias, and long QT syndrome)
Ana Destriction.	Presence of moderate to severe renal impairment or end stage renal disease
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a specialist in the management of Gaucher disease (hematologist, oncologist, hepatologist, geneticist or orthopedic specialist) All approvals are subjects to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



EPLONTERSEN, PATISIRAN, VUTRISIRAN

Affected Medications: WAINUA (eplontersen), ONPATTRO (patisiran), AMVUTTRA (vutrisiran)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Treatment of hereditary transthyretin amyloidosis with polyneuropathy (hATTR-
	PN) in adults
Required Medical	Documented diagnosis of hATTR confirmed by BOTH of the following:
Information:	 Amyloid deposition on biopsy
	Presence of pathogenic transthyretin (TTR) variant on genetic testing
	Presence of clinical manifestations of the disease, confirmed by presence of peripheral Presence of clinical manifestations of the disease, confirmed by presence of peripheral Presence of clinical manifestations of the disease, confirmed by presence of peripheral Presence of clinical manifestations of the disease, confirmed by presence of peripheral Presence of clinical manifestations of the disease, confirmed by presence of peripheral Presence of clinical manifestations of the disease, confirmed by presence of peripheral Presence of clinical manifestations of the disease, confirmed by presence of peripheral Presence of clinical manifestations of the disease, confirmed by presence of peripheral Presence of clinical manifestations of the disease, confirmed by presence of peripheral Presence of clinical manifestations of the disease, confirmed by presence of peripheral Presence of clinical manifestations of the disease of th
	neuropathy on nerve conduction studies OR 2 of the following: o Autonomic dysfunction (bladder/urinary tract infections, gastrointestinal
	disturbances, erectile dysfunction, orthostatic hypotension)
	Documented symptoms of sensorimotor polyneuropathy (e.g., paresthesia,
	balance issues, weakness/numbness in the hands/feet, or loss of sensation for
	pain, temperature, proprioception)
	Cardiomyopathy, ocular involvement, or renal involvement
	Documentation of ONE of the following: Output Description: Description: Output Description: Output Description: Output Description: Output Description: Description: Output Descr
	Baseline polyneuropathy disability (PND) score of less than or equal to IIIb
	 Baseline neuropathy impairment score (NIS) between 10 and 130
	 Baseline familial amyloid polyneuropathy (FAP) stage 1 or 2
Appropriate	Onpattro: Dose-rounding to the nearest vial size within 10% of the prescribed dose will
Treatment	be enforced
Regimen & Other	
Criteria:	Reauthorization:
	Documentation of a positive clinical response (e.g., stabilized or improved neurologic
	impairment, motor function, cardiac function, quality of life assessment, serum TTR
-	lovolo)
Lexclusion Criteria:	levels) • Prior or planned liver transplantation
Exclusion Criteria:	Prior or planned liver transplantation
Exclusion Criteria:	 Prior or planned liver transplantation New York Heart Association (NYHA) Functional Class III or IV
	Prior or planned liver transplantation
Age Restriction:	 Prior or planned liver transplantation New York Heart Association (NYHA) Functional Class III or IV
	 Prior or planned liver transplantation New York Heart Association (NYHA) Functional Class III or IV Combined use with TTR-lowering or stabilizing therapy 18 years of age and older
Age Restriction:	 Prior or planned liver transplantation New York Heart Association (NYHA) Functional Class III or IV Combined use with TTR-lowering or stabilizing therapy 18 years of age and older
Age Restriction: Prescriber/Site of	 Prior or planned liver transplantation New York Heart Association (NYHA) Functional Class III or IV Combined use with TTR-lowering or stabilizing therapy 18 years of age and older Prescribed by, or in consultation with, a neurologist or specialist experienced in the
Age Restriction: Prescriber/Site of	 Prior or planned liver transplantation New York Heart Association (NYHA) Functional Class III or IV Combined use with TTR-lowering or stabilizing therapy 18 years of age and older Prescribed by, or in consultation with, a neurologist or specialist experienced in the treatment of amyloidosis All approvals are subject to utilization of the most cost-effective site of care
Age Restriction: Prescriber/Site of	 Prior or planned liver transplantation New York Heart Association (NYHA) Functional Class III or IV Combined use with TTR-lowering or stabilizing therapy 18 years of age and older Prescribed by, or in consultation with, a neurologist or specialist experienced in the treatment of amyloidosis
Age Restriction: Prescriber/Site of Care Restrictions:	 Prior or planned liver transplantation New York Heart Association (NYHA) Functional Class III or IV Combined use with TTR-lowering or stabilizing therapy 18 years of age and older Prescribed by, or in consultation with, a neurologist or specialist experienced in the treatment of amyloidosis All approvals are subject to utilization of the most cost-effective site of care



POLICY NAME: **EPOPROSTENOL**

Affected Medications: EPOPROSTENOL, VELETRI (epoprostenol), FLOLAN (epoprostenol)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	 Pulmonary Arterial Hypertension (PAH) World Health Organization (WHO) Group
Required Medical	Pulmonary Arterial Hypertension (PAH) WHO Group 1
Information:	 Documentation of PAH confirmed by right-heart catheterization meeting the following criteria:
	Mean pulmonary artery pressure of at least 20 mm Hg
	 Pulmonary capillary wedge pressure less than or equal to 15 mm Hg
	 Pulmonary vascular resistance of at least 2.0 Wood units
	 New York Heart Association (NYHA)/World Health Organization (WHO) Functional Class III or higher symptoms
	 Documentation of Acute Vasoreactivity Testing (positive result requires trial/failure to calcium channel blockers) unless there are contraindications:
	 Low systemic blood pressure (systolic blood pressure less than 90)
	Low cardiac index OR
	 Presence of severe symptoms (functional class IV)
	Documentation of current patient weight
	· · · · · · · · · · · · · · · · · · ·
A	Documentation of a clear treatment plan
Appropriate	Documentation of inadequate response or intolerance to the following therapy classes is
Treatment	required:
Regimen & Other	o PDE5 inhibitors AND
Criteria:	 Endothelin receptor antagonists (exception WHO Functional Class IV)
	Reauthorization requires documentation of treatment success defined as one or more of the following:
	Improvement in exercise ability
	Improvement in pulmonary function
	Improvement or stability in WHO functional class
Exclusion Criteria:	Congestive heart failure due to severe left ventricular systolic dysfunction
	Long-term use in patients who develop pulmonary edema during dose initiation
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a cardiologist or pulmonologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months unless otherwise specified



ERECTILE DYSFUNCTION

Affected Medications: VIAGRA, SILDENAFIL (25 mg, 50 mg, 100 mg), CIALIS (10 mg and 20 mg), EDEX KIT, LEVITRA, MUSE PELLET, STAXYN, STENDRA, TADALAFIL (10 mg, 20 mg), VARDENAFIL, CAVERJECT

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Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Treatment for a mental health diagnosis of erectile dysfunction (ED), also known
	as erectile disorder, meeting sexual dysfunction criteria
Required	Documentation of a mental health diagnosis of erectile dysfunction meeting the
Medical	Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition (DSM-5) criteria:
Information:	 At least one of the three following symptoms must be experienced with 75% to 100% of occasions of sexual activity:
	 Marked difficulty in obtaining an erection during sexual activity
	 Marked difficulty in maintaining an erection until the completion of sexual activity
	 Marked decrease in erectile rigidity
	 The above symptoms have persisted for a minimum duration of approximately 6 months
	 The above symptoms cause clinically significant distress in the individual
	The sexual dysfunction is not attributable to any of the following:
	A nonsexual medical or psychiatric condition
	Severe relationship distress (e.g., partner violence)
	The effects of medication or other substance use
Annuanista	Other clinically significant and relevant stressors
Appropriate	Documentation of treatment failure with tadalafil 2.5 mg or 5 mg tablets
Treatment	
Regimen &	
Other Criteria:	
Exclusion	Erectile dysfunction unrelated to a mental health diagnosis of sexual dysfunction
Criteria:	according to the DSM-5 diagnostic criteria
Prescriber/Site of	Prescribed by, or in consultation with, a mental health provider
Care Restrictions	All approvals are subject to utilization of the most cost-effective site of care
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Age	
Restriction:	
Coverage	Authorization: 12 months, unless otherwise specified
Duration:	



POLICY NAME: ERGOT ALKALOIDS

Affected Medications: DIHYDROERGOTAMINE MESYLATE INJECTION, DIHYDROERGOTAMINE MESYLATE NASAL SOLUTION

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Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
Required Medical Information:	Documentation of moderate to severe migraines
Appropriate	• Documentation of treatment failure, intolerance, or contraindication to all of the following:
Treatment	 At least <u>two</u> prescription strength non-steroidal anti-inflammatory drugs (NSAIDs)
Regimen & Other	or combination analgesics (such as ibuprofen, naproxen,
Criteria:	acetaminophen/aspirin/caffeine)
	 At least <u>one</u> oral 5-hydroxytryptamine-1 (5-HT₁) receptor agonist (such as sumatriptan, naratriptan, rizatriptan, zolmitriptan)
	o At least <u>one</u> non-oral 5-HT₁ receptor agonist (such as sumatriptan, zolmitriptan)
	Reauthorization will require documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	Hemiplegic or basilar migraine
	Uncontrolled hypertension
	 Ischemic heart disease (e.g., angina pectoris, history of myocardial infarction, history of silent ischemia)
	Peripheral artery disease
	Pregnancy or breastfeeding
	Documented severe chronic liver disease
	Severe renal impairment
	Use in combination with 5HT1 receptor agonist such as sumatriptan
Age Restriction:	18 years of age and older
Prescriber/Site of	All approvals are subject to utilization of the most cost-effective site of care
Care Restrictions:	,,
Coverage Duration:	Authorization: 12 months, unless otherwise specified
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ERYTHROPOIESIS STIMULATING AGENTS (ESAs)

Affected Medications: ARANESP (darbepoetin alfa), EPOGEN (epoetin alfa), MIRCERA (methoxy polyethylene glycolepoetin beta), PROCRIT (epoetin alfa)

Required Medical Information:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Epogen & Aranesp & Procrit & Mircera Treatment of anemia due to chronic kidney disease (CKD), including patients on dialysis and not on dialysis to decrease the need for red blood cell (RBC) transfusion Epogen & Procrit & Aranesp Treatment of anemia in patients with non-myeloid malignancies where anemia is due to the effect of concomitant myelosuppressive chemotherapy, and upon initiation, there is a minimum of two additional months of planned chemotherapy Epogen & Procrit only To reduce the need for allogeneic RBC transfusions among patients with perioperative hemoglobin greater than 10 to 13 or less g/dL who are at high risk for perioperative blood loss from elective, noncardiac, nonvascular surgery Treatment of anemia due to zidovudine administered at ≤ 4200 mg/week in patients with HIV-infection with endogenous serum erythropoietin levels of ≤ 500 mUnits/mL Compendia-supported uses Symptomatic anemia in Myelodysplastic syndrome Allogenic bone marrow transplantation Anemia associated with Hepatitis C (HCV) treatment Anemia associated with chronic renal failure Anemia associated with chronic renal failure Anemia associated with chronic renal failure Anemia secondary to chemotherapy with a minimum of two additional months of planned chemotherapy Anemia secondary to zidovudine-treated Human Immunodeficiency Virus (HIV) patients Anemia in patients scheduled to undergo elective, non-cardiac, nonvascular surgery Symptomatic anemia in Myelodysplastic syndrome Allogenic bone marrow transplantation Anemia associated with Hepatitis C (HCV) treatment
	 Anemia associated with rheumatoid arthritis (RA)/ rheumatic disease
Appropriate Treatment Regimen & Other Criteria:	Coverage for the non-preferred drugs (Epogen, Procrit, Mircera) is provided when the following criteria is met:
Exclusion Criteria:	Use in combination with another erythropoiesis stimulating agent (ESA)
Age Restriction:	
Prescriber/Site of Care Restrictions:	Prescribed by, or in consultation with, a hematologist, oncologist, or nephrologist



Coverage Duration:	•	Authorization: 6 months, unless otherwise specified



POLICY NAME: **ETELCALCETIDE**

Affected Medications: PARSABIV (etelcalcetide)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Secondary hyperparathyroidism in adults with chronic kidney disease (CKD) on dialysis
Required Medical Information:	Documentation of both of the following:
Appropriate Treatment Regimen & Other Criteria:	Reauthorization will require documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	Diagnosis of parathyroid carcinoma, primary hyperparathyroidism or with chronic kidney disease who are not on hemodialysis
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an endocrinologist or nephrologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: **ETRANACOGENE**

Affected Medications: HEMGENIX (etranacogene dezaparvovec-drlb)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Hemophilia B (congenital factor IX deficiency)
Required Medical Information:	 Documentation of diagnosis of Hemophilia B Documentation of baseline circulating level of factor IX less than or equal to 2% as attested by the managing physician AND requiring prophylactic Factor IX treatment Documentation of negative Factor IX inhibitor titers (if test result is positive, re-test within 2 weeks with negative result) Baseline lab values (less than 2 times upper limit of normal): ALT AST Total bilirubin Alkaline phosphatase (ALP)
Appropriate Treatment Regimen & Other Criteria:	Documentation of plan to discontinue Factor IX prophylaxis therapy upon achieving circulating factor IX levels of 5%
Officia.	Dosing:
	2 x 10 ¹³ genome copies (gc) per kilogram of body weight
Exclusion Criteria:	Prior gene therapy administration
Age Restriction:	18 years of age and older
Prescriber/Site of	All approvals are subject to utilization of the most cost-effective site of care
Care Restrictions:	Prescribed by, or in consultation with, a hematologist or specialist with experience in the treatment of hemophilia
Coverage Duration:	Authorization: 2 months (one-time infusion only), unless otherwise specified



EVKEEZA and JUXTAPID

Affected Medications: EVKEEZA (evinacumab-dgnb), JUXTAPID (lomitapide)

Oncome al III	AUE I ID ALLICO (EDA)
Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design o Homozygous familial hypercholesterolemia (HoFH)
Required	Documentation of baseline untreated low-density lipoprotein cholesterol (LDL-C)
Medical	Diagnosis confirmed by ONE of the following:
Information:	Baseline LDL-C greater than 560 mg/dL
	Baseline LDL-C of 400 mg/dL and at least 1 parent with familial
	hypercholesterolemia
	Baseline LDL-C of 400 mg/dL with aortic valve disease or xanthomata in ages
	less than 20 years
	 Presence of two abnormal LDL-C-raising gene defects (excluding double-null LDL
	receptor [LDLR] mutations)
Appropriate	Documented intent to take alongside maximally tolerated doses of statin and/or ezetimibe,
Treatment	unless otherwise contraindicated
Regimen &	OR
Other Criteria:	History of statin intolerance requires documentation of ONE of the following:
	 Statin-associated rhabdomyolysis occurred with statin use and was confirmed by
	a creatinine kinase (CK) level at least 10 times the upper limit of normal
	 Statin-associated muscle symptoms (e.g., myopathy, myalgia) occurred with
	statin use and was confirmed by BOTH of the following:
	 A minimum of two different statin trials, with at least one being a
	hydrophilic statin (rosuvastatin, pravastatin)
	 A re-challenge of each statin (muscle symptoms stopped when each was
	discontinued and restarted upon re-initiation)
	 Documented treatment failure, defined as an inability to achieve LDL-C reduction of 50% or greater OR LDL-C less than 100 mg/dL despite at least six months of adherent therapy with all of the following, unless contraindicated or not tolerated: Maximally tolerated statin therapy Ezetimibe
	 PCSK9 monoclonal antibody, unless double-null or LDLR activity 15% or less
	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced
	Reauthorization requires documentation of treatment success and a clinically significant
	response to therapy defined by an LDL-C level at goal or decreased by at least 30% from
Parata de la constante de la c	baseline
Exclusion Criteria:	Combination therapy with Juxtapid and Evkeeza is considered experimental and is not a covered benefit
Age	Evkeeza: 5 years of age and older
Restriction:	Juxtapid: 18 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an endocrinologist, cardiologist, or lipid specialist All approvals are subject to utilization of the most cost-effective site of care



Coverage	Initial Authorization: 6 months, unless otherwise specified
Duration:	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **EVOLOCUMAB**

Affected Medications: REPATHA (evolocumab)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Secondary prevention in clinical atherosclerotic cardiovascular disease (ASCVD)
	 Primary hyperlipidemia (including heterozygous familial hypercholesterolemia
	[HeFH])
	Homozygous familial hypercholesterolemia (HoFH)
Required Medical	<u>All Indications</u>
Information:	Documentation of current complete lipid panel within last 3 months
	Documentation of baseline (untreated) low-density lipoprotein cholesterol (LDL-C)
	Clinical ASCVD
	Documentation of established ASCVD, confirmed by at least ONE of the following:
	 Acute coronary syndromes (ACS)
	 History of myocardial infarction (MI)
	 Stable or unstable angina
	 Coronary or other arterial revascularization
	 Stroke or transient ischemic attack
	 Peripheral artery disease (PAD) presumed to be of atherosclerotic origin
	Primary Hyperlipidemia (non-familial)
	Documentation of baseline (untreated) LDL-C of at least 190 mg/dL
	HeFH_
	Diagnosis confirmed by ONE of the following:
	 Minimum baseline LDL-C of 160 mg/dL in adolescents or 190 mg/dL in adults
	AND 1 first-degree relative affected
	Presence of one abnormal LDL-C-raising gene defect (e.g., LDL receptor)
	[LDLR], apolipoprotein B [apo B], proprotein convertase subtilisin kexin type 9
	[PCSK9] gain-of-function mutation, LDL receptor adaptor protein 1 [LDLRAP1])
	World Health Organization (WHO)/Dutch Lipid Network criteria score of at least 8
	points
	 Definite FH diagnosis per the Simon Broome criteria
	HoFH
	Diagnosis confirmed by ONE of the following:
	Baseline LDL-C greater than 560 mg/dL
	Baseline LDL-C of 400 mg/dL and at least 1 parent with familial hypercholesterolemia
	 Baseline LDL-C of 400 mg/dL with aortic valve disease or xanthomata in ages less than 20 years
	Presence of two abnormal LDL-C-raising gene defects (excluding double-null)
	LDLR mutations)



All Indications

Appropriate

Treatment Regimen & Other Criteria:	 Documented intent to take alongside maximally tolerated does of statin and/or ezetimibe, unless otherwise contraindicated OR History of statin intolerance requires documentation of ONE of the following:
	 History of statin intolerance requires documentation of ONE of the following: Statin-associated rhabdomyolysis occurred with statin use and was confirmed by a creatinine kinase (CK) level at least 10 times the upper limit of normal Statin-associated muscle symptoms (e.g., myopathy, myalgia) occurred with statin use and was confirmed by BOTH of the following:
	Major ASCVD Events High-Risk Conditions
	 ACS within the past 12 months History of MI (distinct from ACS event) Ischemic stroke Symptomatic PAD Age 65 years and older HeFH Prior coronary artery bypass or percutaneous intervention (outside of major ASCVD events) Diabetes Hypertension Chronic kidney disease Currently smoking History of congestive heart failure
	Primary Hyperlipidemia/HeFH/HoFH
	 Documented treatment failure, defined as an inability to achieve LDL-C reduction of 50% or greater OR LDL-C less than 100 mg/dL, with minimum 12 weeks of statin/ezetimibe combination therapy at maximally tolerated doses with consistent use
Exclusion Criteria:	Concurrent use with Leqvio
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a cardiologist, endocrinologist, or lipid specialist All approvals are subject to utilization of the most cost-effective site of care



Coverage Duration:	•	Authorization: 12 months, unless otherwise specified



EXAGAMGLOGENE AUTOTEMCEL

Affected Medications: CASGEVY (exagamglogene autotemcel)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	Treatment of sickle cell disease in adults and pediatric patients at least 12 years
	of age with recurrent vaso-occlusive crises.
	Treatment of transfusion-dependent beta-thalassemia in adults and pediatric petiants at least 12 years of age.
Deguired Medical	patients at least 12 years of age. SICKLE CELL DISEASE
Required Medical	SICKLE CELL DISEASE
Information:	Decumentation of cickle call disease confirmed by genetic tecting to about the presence
	 Documentation of sickle cell disease confirmed by genetic testing to show the presence of βS/βS, βS/β0 or βS/β+ genotype as follows:
	 Identification of significant quantities of HbS with or without an additional
	abnormal β-globin chain variant by hemoglobin assay
	OR
	 Identification of biallelic HBB pathogenic variants where at least one allele is the
	p.Glu6Val or p.Glu7Val pathogenic variant on molecular genetic testing
	AND
	 Patient does NOT have disease with more than two α-globin gene deletions
	Documentation of severe disease defined as 2 or more severe vaso-occlusive crises
	(VOCs) or vaso-occlusive events (VOEs) within the previous year (4 events over 2 years
	will also meet this requirement) VOC/VOEs defined as:
	 Acute pain event requiring a visit to a medical facility and administration of pain
	medications (opioids or IV NSAIDs) or RBC transfusions
	 Acute chest syndrome
	 Priapism lasting more than 2 hours and requiring visit to medical facility
	Splenic sequestration
	Clinically stable and eligible to undergo hematopoietic stem cell transplant (HSCT) but
	unable to find a human leukocyte antigen (HLA) matched, related donor
	Adequate bone marrow, lung, heart, and liver function to undergo myeloablative
	conditioning regimen
	TRANSFUSION REPENDENT RETAINING ACCEMIA
	TRANSFUSION DEPENDENT BETA THALASSEMIA
	Documented diagnosis of homozygous beta thalassemia or compound heterozygous
	beta thalassemia including β-thalassemia/hemoglobin E (HbE) (excludes alpha-
	thalassemia and hemoglobin S/ß-thalassemia variants) as outlined by the following:
	 Patient diagnosis is confirmed by HBB sequence gene analysis showing biallelic
	pathogenic variants
	OR
	 Patient has severe microcytic hypochromic anemia, anisopoikilocytosis with
	nucleated red blood cells on peripheral blood smear, and hemoglobin analysis
	that reveals decreased amounts or complete absence of hemoglobin A and
	increased amounts of hemoglobin F
	Documented transfusion-dependent disease defined as a history of transfusions of at least 100 ml //cg/year of period rad blood cells (pBBCs) are with 10 ar more transfusions.
	least 100 mL/kg/year of packed red blood cells (pRBCs) or with 10 or more transfusions
	of pRBCs per year in the 2 years preceding therapy



	Clinically stable and eligible to undergo hematopoietic stem cell transplant (HSCT) but unable to find a human leukocyte antigen (HLA) matched, related donor
Appropriate Treatment Regimen & Other Criteria:	 Must weigh a minimum of 6 kilograms and able to provide a minimum number of cells (3 x 10⁶ CD34+ cells/kg) Documentation that cardiac iron overload has been evaluated and there is no evidence of severe iron overload. (cardiac T2* less than 10 msec by magnetic resonance imaging [MRI] or left ventricular ejection fraction [LVEF] less than 45% by echocardiogram) No evidence of advanced liver disease [i.e., AST or ALT more than 3 times the upper limit of normal (ULN), or direct bilirubin value more than 2.5 times the ULN, or if a liver biopsy demonstrated bridging fibrosis or cirrhosis]
Exclusion Criteria:	Prior HSCT or other gene therapy
Age Restriction:	12 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a hematologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 6 months (one time infusion), unless otherwise specified



FABRY DISEASE AGENTS

Affected Medications: ELFABRIO (pegunigalsidase alfa), FABRAZYME (agalsidase beta), GALAFOLD (migalastat)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise evaluded by
Covered Oses.	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	Fabry disease
Required Medical	Diagnosis of Fabry disease confirmed by one of the following:
Information:	 Males: enzyme assay demonstrating undetectable (less than 3 percent) alpha-galactosidase A enzyme activity
	 Males: deficiency of alpha-galactosidase A enzyme activity (less than 35 percent) and genetic testing showing a mutation in the galactosidase alpha (GLA) gene
	 Females: genetic testing showing a mutation in the GLA gene
	For Galafold: Genetic testing confirming the presence of at least one amenable GLA variant
	Clinical signs and symptoms of Fabry disease, such as: Severe neuropathic pain
	 Dermatologic manifestations (telanglectasias and anglokeratomas) Corneal opacities
	 Kidney manifestations (proteinuria, polyuria, polydipsia)
	Cardiac involvement (left ventricular hypertrophy, myocardial fibrosis, heart)
	failure)
	 Cerebrovascular involvement (transient ischemic attacks, ischemic strokes) Other manifestations common in Fabry disease (sweating abnormalities, hearing loss, or intolerance to heat, cold, or exercise)
Appropriate	Dose-rounding to the nearest vial size within 10% of the prescribed dose will
Treatment	be enforced
Regimen & Other	
Criteria:	<u>Reauthorization</u> requires documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	Concurrent use with another agent on this policy (Galafold or enzyme replacement therapy for Fabry disease)
	For Galafold: Severe renal impairment (eGFR less than 30) or end-stage renal disease requiring dialysis
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a geneticist or a specialist experienced in the
Care Restrictions:	treatment of Fabry disease
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified
	- Roadmonzation. 12 months, diffess otherwise specified



FECAL MICROBIOTA

Affected Medications: REBYOTA (fecal microbiota, live-jslm), VOWST (fecal microbiota spores, live-brpk)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Prophylaxis of Clostridioides difficile (C.diff) infection recurrence following antibiotic treatment
Required Medical Information:	 Documentation confirming a current diagnosis of recurrent C.diff infection (CDI) with a history of at least 2 recurrent episodes (initial episode + a minimum of 2 recurrences) Recurrent CDI is defined as a resolution of CDI symptoms while on appropriate therapy, followed by a reappearance of symptoms within 8 weeks of discontinuing treatment Current episode of CDI must be controlled (less than 3 unformed or loose stools per day for 2 consecutive days) Administration will occur following completion of antibiotic course for CDI treatment Within 24 to 72 hours for Rebyota Within 2 to 4 days for Vowst Positive stool test for C.diff within the 30 days prior to request
Appropriate	Rebyota
Treatment	Previous treatment with at least TWO of the following in the setting of CDI recurrence:
Regimen & Other Criteria:	oral vancomycin, fidaxomicin (Dificid), or fecal microbiota transplant (FMT)
	<u>Vowst</u>
	 Previous treatment with at least TWO of the following in the setting of CDI recurrence: oral vancomycin, fidaxomicin (Dificid), or FMT
	Documented treatment failure with Rebyota
Exclusion Criteria:	Retreatment with Rebyota or Vowst
Age Restriction:	18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, an infectious disease specialist or
Care Restrictions:	gastroenterologist
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 1 month with no reauthorization, unless otherwise specified



POLICY NAME: **FENFLURAMINE**

Affected Medications: FINTEPLA (fenfluramine)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of seizures associated with Dravet syndrome (DS) Treatment of seizures associated with Lennox-Gastaut syndrome (LGS)
Required Medical Information:	 Documented diagnosis of Dravet syndrome (DS) or Lennox-Gastaut Syndrome (LGS) Current weight Documentation that therapy is being used as adjunct therapy for seizures
	 Dravet Syndrome Documentation of at least 6 convulsive seizures in the last 6 weeks while on stable antiepileptic drug therapy
	 Lennox-Gastaut Syndrome (LGS) Documentation of at least 8 drop seizures per month while on stable antiepileptic drug therapy
Appropriate Treatment Regimen & Other Criteria:	 Dravet Syndrome Documented treatment and inadequate control of seizures with Epidiolex AND at least four of the following therapies: ○ Valproate, clobazam, clonazepam, levetiracetam, zonisamide, or topiramate
	 Lennox-Gastaut Syndrome (LGS) Documented treatment and inadequate control of seizures with Epidiolex AND at least three guideline directed therapies: Valproate, lamotrigine, rufinamide, topiramate, felbamate, or clobazam
	Dosing: not to exceed 26 mg daily
	Reauthorization requires documentation of treatment success and a reduction in seizure severity, frequency, or duration
Exclusion	
Criteria: Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: FIDANACOGENE

Affected Medications: BEQVEZ (fidanacogene elaparvovec-dzkt)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise excluded by
	plan design
	 Hemophilia B (congenital factor IX deficiency)
Required Medical	Documentation of diagnosis of Hemophilia B
Information:	 Documentation of baseline circulating level of factor IX less than or equal to 2% of normal AND requiring prophylactic factor IX treatment for at least 6 months Documentation of negative factor IX inhibitor titers (less than 0.6 Bethesda units) Documentation of negative antibodies to AAVRh74var capsid per FDA approved
	diagnostic test
	 Baseline lab values (less than 2 times upper limit of normal): ALT AST
	All III I (ALD)
	Alkaline phosphatase (ALP) Bilirubin
Appropriate	Documentation of plan to discontinue factor IX prophylaxis therapy upon achieving
Treatment	circulating factor IX levels of 5%
Regimen & Other	onodiating factor fix fevers of 676
Criteria:	Dosing
	5 x 10 ¹¹ vector genomes per kilogram of body weight
Exclusion Criteria:	Prior gene therapy administration
	Unstable liver or biliary disease
	Active Hepatitis B or C infection
	HIV infection with CD4 cell count less than 200 mm³ or viral load greater than 20 copies/mL
Age Restriction:	18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation, with a hematologist or specialist with experience in
Care Restrictions:	treatment of hemophilia
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 2 months (one-time infusion)



POLICY NAME: FINERENONE

Affected Medications: KERENDIA (finerenone)

0	AUE I ID ALITY (CIPA)
Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Chronic kidney disease associated with type 2 diabetes to reduce the risk of:
	 Sustained estimated glomerular filtration rate (eGFR) decline
	End-stage kidney disease
	Cardiovascular death
	 Non-fatal myocardial infarction
	 Hospitalization for heart failure
Required Medical	Documentation of all the following:
Information:	 eGFR greater than or equal to 25 mL/min/1.73 m²
	 Urine albumin-to-creatinine ratio (UACR) greater than or equal to 30 mg/g
	 Serum potassium level less than or equal to 5.0 mEq/L
Appropriate	Currently receiving maximally tolerated dosage of an angiotensin converting enzyme
Treatment	(ACE) inhibitor or angiotensin receptor blocker (ARB), unless intolerant or
Regimen & Other	contraindicated
Criteria:	 Documented treatment failure or intolerable adverse event to at least 12 weeks of
Cilieria.	sodium-glucose cotransporter 2 (SGLT2) inhibitor therapy
	Reauthorization requires documentation of treatment success and a clinically significant
	response to therapy
Exclusion Criteria:	
Age Restriction:	18 years of age and older
Prescriber/Site of	Dropprihad by ar in consultation with a penhaginat and carinologist or conditions in
	Prescribed by, or in consultation with, a nephrologist, endocrinologist, or cardiologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified
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POLICY NAME: FLUCYTOSINE

Affected Medications: FLUCYTOSINE

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
Required Medical Information:	Susceptibility cultures matching flucytosine activity
Appropriate Treatment Regimen & Other Criteria:	Dosing: maximum 150 mg/kg/day
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an infectious disease specialist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 8 weeks, or lesser requested duration, unless otherwise specified



FLUOCINOLONE OCULAR IMPLANT

Affected Medications: ILUVIEN, RETISERT, YUTIQ

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Diabetic macular edema (DME) Chronic, non-infectious posterior uveitis
Required Medical Information:	 Iluvien Diagnosis of clinically significant diabetic macular edema Documentation of past treatment with corticosteroids without a clinically significant rise in intraocular pressure Retisert and Yutiq Diagnosis of chronic, non-infectious posterior uveitis confirmed by slit lamp and fundoscopic examination
Appropriate Treatment Regimen & Other Criteria:	Iluvien
	 Retisert and Yutiq Documentation of inadequate response or intolerance to all of the following: Minimum 12-week trial with oral systemic corticosteroid At least one corticosteroid-sparing immunosuppressive therapy (methotrexate, azathioprine, or mycophenolate mofetil) At least one calcineurin inhibitor (cyclosporine, tacrolimus) Retisert: Documentation of treatment failure with Yutiq
Exclusion Criteria:	Active or suspected ocular or periocular infections Concurrent use of intravitreal implants or injections (corticosteroid, anti-VEGF) Iluvien: Glaucoma (with cup to disc ratios greater than 0.8)
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, an ophthalmologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Iluvien: 36 months, unless otherwise specified Retisert: 30 months, unless otherwise specified Yutiq: 36 months, unless otherwise specified



Food and Drug Administration (FDA) APPROVED DRUG – Drug or Indication Not Yet Reviewed By Plan for Formulary Placement

Affected Medications: New Medications or Indications of Existing Drugs Not Yet Reviewed By Plan for Formulary Placement

Covered Uses:	Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
Required Medical	Documentation of disease state, level of control, and therapies failed
Information:	Documentation of failure with all available formulary products for treatment of disease state
	Documentation that delay in treatment will cause loss of life, limb, function or other extreme pain
Appropriate	Drug must be dosed according to package insert requirements
Treatment	
Regimen & Other	
Criteria:	
Exclusion Criteria:	Exclusion based on package insert requirements
Age Restriction:	Age based on package insert requirements
Prescriber/Site of	Prescriber restrictions based on package insert requirements
Care Restrictions:	All approvals are subject to utilization of the most cost effective site of care
Coverage Duration:	Case by case based on member need



POLICY NAME: FOSTAMATINIB

Affected Medications: TAVALISSE (fostamatinib)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Thrombocytopenia in adults with chronic immune thrombocytopenia (ITP) who have had an insufficient response to a previous treatment
Required Medical	Thrombocytopenia in patients with chronic ITP
Information:	Documentation of ONE of the following: Platelet count less than 20,000/microliter Platelet count less than 30,000/microliter AND symptomatic bleeding Platelet count less than 50,000/microliter AND increased risk for bleeding (such as peptic ulcer disease, use of antiplatelets or anticoagulants, history of bleeding at higher platelet count, need for surgery or invasive procedure)
Appropriate	Thrombocytopenia in patients with chronic ITP
Treatment	Documentation of inadequate response, defined as platelets did not increase to at least 50,000 (valence little), to the following at the applications: 6. **The control of the cont
Regimen & Other	50,000/microliter, to the following therapies: ONE of the following:
Criteria:	 Inadequate response with at least 2 therapies for immune thrombocytopenia, including corticosteroids, rituximab, or immunoglobulin Splenectomy Promacta
	Reauthorization:
	Response to treatment with platelet count of at least 50,000/microliter or above (not to exceed 400,000/microliter)
Exclusion Criteria:	Use in combination with a thrombopoietin receptor agonist, spleen tyrosine kinase inhibitor, or similar treatment for thrombocytopenia (such as Promacta, Doptelet, or Nplate)
Age Restriction:	
Prescriber	Prescribed by, or consultation with, a hematologist
Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



FYARRO

Affected Medications: FYARRO (nab-sirolimus)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design. NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A
	or better
Required Medical	Documentation of performance status, disease staging, all prior therapies used, and
Information:	anticipated treatment course
Appropriate	Perivascular Epithelioid Cell Tumor (PEComa)
Treatment	Presence of malignant locally advanced unresectable or metastatic disease confirmed by
Regimen & Other	pathology.
Criteria:	 History of intolerable adverse event with trial of each of the following agents: Sirolimus oral tablet
	Everolimus or temsirolimus
	Reauthorization: documentation of disease responsiveness to therapy
Exclusion Criteria:	Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
	History of disease progression with prior mechanistic target of rapamycin (mTOR) inhibitor treatment.
Age Restriction:	
Prescriber	Prescribed by, or in consultation with, an oncologist
Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial approval: 4 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



GABA-A RECEPTOR MODULATORS

Affected Medications: ZULRESSO (brexanolone), ZURZUVAE (zuranolone)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	Treatment of postpartum depression (PPD)
Required Medical	Documented major depressive episode with peripartum onset as defined by the
Information:	Diagnostic and Statistical Manual of Mental Health Disorders, Fifth Edition (DSM-5) criteria:
	 At least five of the following symptoms have been present during the same 2-
	week period and represent a change from previous functioning (must include either (1) depressed mood or (2) lack of interest or pleasure):
	(1). Depressed mood most of the day, nearly every day, as indicated by
	either subjective report or observation made by others (in adolescents, may present as irritable mood)
	(2). Markedly diminished interest or pleasure in all (or almost all) activities
	most of the day, nearly every day, as indicated by either subjective account or observation
	(3). Significant weight loss when not dieting, weight gain, or decrease or
	increase in appetite nearly every day (in adolescents, consider failure to make expected weight gain)
	(4). Insomnia or hypersomnia nearly every day
	(5). Psychomotor agitation or retardation nearly every day (observable by
	others, not merely subjective feelings of restlessness or being slowed down)
	(6). Fatigue or loss of energy nearly every day
	(7). Feelings of worthlessness, or excessive or inappropriate guilt nearly everyday
	(8). Diminished ability to think or concentrate, or indecisiveness, nearly everyday (subjective account or observed by others)
	(9). Recurrent thoughts of death (not just fear of dying), recurrent suicidal
	ideation without a specific plan, or a suicide attempt or a specific plan for committing suicide
	 Symptoms cause clinically significant distress or impairment in social,
	occupational, or other important areas of functioning
	 Episode is not attributable to the direct physiological effects of a substance or to another condition
	 Major depressive episode began no earlier than the third trimester and no later than the first 4 weeks following delivery
	Moderate to severe postpartum depression documented by one of the following rating
	scales:
	o Hamilton Rating Scale for Depression (HAM-D) score of greater than 17
	o Patient Health Questionnaire-9 (PHQ-9) score of greater than 10
	 Montgomery-Åsberg Depression Rating Scale (MADRS) greater than 20 points Edinburgh Postnatal Depression Scale (EPDS) score of greater than 13
Appropriate	
Treatment	Documented trial with an oral antidepressant for at least 8 weeks unless contraindicated or documentation shows that the severity of the depression would place the health of the
Regimen & Other	mother or infant at significant risk
Criteria:	For Zulresso requests: documented treatment failure with Zurzuvae
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Exclusion Criteria:	Greater than 6 months postpartum
Age Restriction:	15 years of age and older for Zulresso
	18 years of age and older for Zurzuvae
Prescriber/Site of	Prescribed by, or in consultation with, a psychiatrist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 1 month, one time approval per pregnancy, unless otherwise specified



POLICY NAME: GANAXOLONE

Affected Medications: ZTALMY (ganaxolone)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of seizures associated with cyclin-dependent kinase-like 5 (CDKL5) deficiency disorder (CDD) in patients 2 years of age and older
Required Medical Information:	 Documentation of CDKL5 mutation confirmed by genetic testing Documentation of inadequately controlled seizures despite current treatment
Appropriate Treatment Regimen & Other Criteria:	Documented treatment failure with at least two therapies for seizure management Reauthorization will require documentation of treatment success defined as a reduction in seizure frequency when compared to baseline
Exclusion Criteria:	West syndrome Seizures of a predominantly infantile spasm type
Age Restriction:	2 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: GIVINOSTAT

Affected Medications: DUVYZAT (givinostat)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise excluded by
	plan design
	 Duchenne muscular dystrophy (DMD) in patients 6 years of age and older
Required Medical	Genetically confirmed diagnosis of DMD
Information:	Documentation of being ambulatory without needing an assistive device such as a wheelchair, walker, or cane
	North Star Ambulatory Assessment (NSAA) scale total score of 17 or more
	Baseline motor function assessment from one of the following:
	o 4-stair climb (4SC) test
	o Time to Stand Test (TTSTAND)
	o 6-minute walk test (6MWT)
	North Star Ambulatory Assessment (NSAA)
	Motor Function Measure (MFM)
	Hammersmith Functional Motor Scale (HFMS)
	Current weight and planned treatment regimen
Appropriate	Documentation of being on a stable dose of an oral corticosteroid such as prednisone for
Treatment	at least 6 months, and will continue while on Duvyzat unless contraindicated
Regimen & Other	
Criteria:	<u>Reauthorization</u> requires a documented improvement from baseline or stabilization of motor function demonstrated by a motor function assessment tool
Exclusion Criteria:	Concomitant therapy or within the past 6 months with DMD-directed antisense
	oligonucleotides such as golodirsen, casimersen, viltolarsen, eteplirsen
	Platelet, white blood cell, or hemoglobin counts less than the lower limit of normal
	QTc is greater than 500 ms or the change from baseline is greater than 60 ms.
	History of additional risk factors for torsades de pointes (e.g. heart failure, hypokalemia,
	or family history of long QT syndrome)
Age Restriction:	6 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Care restrictions.	- 7 in approvate and subject to difficultion of the most obstraction of the most obstraction of the
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **GIVOSIRAN**

Affected Medications: GIVLAARI (givosiran)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of adults with acute hepatic porphyria (AHP)
Required Medical Information:	 Documentation of elevated urine porphobilinogen (PBG) levels based on specific lab test utilized Diagnosis confirmed based on Porphyria Genomic testing Documentation of baseline acute attack frequency Evaluation and elimination of exacerbating factors of porphyria attacks including certain medications, smoking, drinking, and infections
Appropriate Treatment Regimen & Other Criteria:	 Documentation of active disease defined as at least 2 documented porphyria attacks within the last six months, which can include hospitalization, urgent healthcare visits, or requiring intravenous Hemin administration Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced Reauthorization will require documentation of positive clinical response and a reduction in acute attack frequency from baseline
Exclusion Criteria:	 Active HIV, hepatitis C, or hepatitis B infection(s) History of pancreatitis Concomitant use with prophylactic hemin History of liver transplant
Age Restriction:	18 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, specialist in the treatment of acute hepatic porphyria All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



GLUCAGON-LIKE PEPTIDE (GLP-1) RECEPTOR AGONIST
Affected Medications: TRULICITY, OZEMPIC, RYBELSUS, MOUNJARO, LIRAGLUTIDE

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise excluded by
	plan design
	o Diabetes Mellitus, Type 2
Required Medical Information:	 Available information is reviewed, including previous fill history Diagnosis of Type 2 diabetes with a recent hemoglobin A1c greater than or equal to 7% despite current therapy Documented treatment failure with minimum of 12-week trial with metformin or metformin extended release 2000 mg daily (or if unable to tolerate 2000 mg daily, the maximum tolerated dose) defined as failure to achieve or maintain A1c less than 7% If intolerant to immediate release metformin, 12-week trial with metformin extended release must be trialed
Appropriate	
Treatment	Reauthorization requires documentation of disease responsiveness to therapy
Regimen & Other Criteria:	
Exclusion Criteria:	Use for weight loss or other excluded diagnosis
	 Dosing above Food and Drug Administration (FDA) approved label for treatment of diabetes
	Use in patients who have achieved remission of diabetes (defined as a return of HbA1c to less than 6.5% that occurs spontaneously or following an intervention and that persists for at least three months in the absence of usual glucose-lowering pharmacotherapy)
Age Restriction:	
Prescriber/Site of Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 24 months, unless otherwise specified



POLICY NAME: GONADOTROPIN

Affected Medications: CHORIONIC GONADOTROPIN, PREGNYL, NOVAREL

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Hypogonadotropic hypogonadism secondary to a pituitary deficiency in males Prepubertal cryptorchidism not caused by anatomic obstruction Perioperative use in male infants/toddlers with hypospadias and chordee OR total epispadias and bladder exstrophy
Required Medical Information:	Hypogonadotropic hypogonadism secondary to a pituitary deficiency in males: Documentation confirming the diagnosis
Appropriate	Reauthorization will require documentation of treatment success and a clinically significant
Treatment	response to therapy
Regimen & Other Criteria:	
Exclusion Criteria:	 Use for the diagnosis or treatment of infertility (if benefit exclusion) Obesity
	 Prevention of recurrent or habitual miscarriage Treatment or prevention of breast cancer
Age Restriction:	 Prepubertal cryptorchidism: generally, between 4 and 9 years of age Hypospadias or epispadias: infant or toddler
Prescriber/Site of	All approvals are subjects to utilization of the most cost-effective site of care
Care Restrictions:	
Coverage Duration:	Authorization: 12 months, unless otherwise specified



GOSERELIN ACETATE IMPLANT

Affected Medications: ZOLADEX (goserelin acetate implant)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Endometriosis Endometrial thinning NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A
	or better
Required Medical Information:	Endometriosis:
	Documentation of moderate to severe pain due to endometriosis
Appropriate Treatment	Endometriosis:
Regimen & Other Criteria:	 Documentation of a trial and inadequate relief (or contraindication) after at least 3 months of both of the following first-line therapies: Nonsteroidal anti-inflammatory drugs (NSAIDs) Continuous (no placebo pills) hormonal contraceptives
	Endometrial thinning:
	 Documentation of both of the following: Diagnosis of dysfunctional uterine bleeding Planning to use as an endometrial-thinning agent prior to endometrial ablation
	Reauthorization for oncologic uses require documentation of disease responsiveness to therapy
Exclusion Criteria:	 Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater For endometriosis, prior use of Zoladex for a 6-month period
Age Restriction:	18 years of age and older
Prescriber/Site of	For oncologic uses: Prescribed by, or in consultation with, an oncologist
Care Restrictions:	For gynecologic uses: Prescribed by, or in consultation with, a gynecologist
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Oncologic uses:
	Initial Authorization: 4 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified
	Endometriosis:
	Authorization: 6 months with no reauthorization, unless otherwise specified
	Endometrial thinning:
	Authorization: 4 months (up to 2 doses only), unless otherwise specified



GROWTH HORMONES

Affected Medications: GENOTROPIN, GENOTROPIN MINIQUICK, HUMATROPE, NORDITROPIN FLEXPRO, NUTROPIN AQ NUSPIN, OMNITROPE, SAIZEN, SKYTROFA, ZOMACTON, SOGROYA, NGENLA

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
Required Medical Information:	 All indications: Documentation of baseline height, height velocity, bone age (pediatrics), and patient weight
	 Growth hormone deficiency or Pituitary dwarfism For initial approval, documentation of the following is required: □ Diagnosis of growth hormone deficiency or pituitary dwarfism AND □ Low serum values for GH stimulation test, IGF-1, and IGFBP-3 with delayed bone age AND ■ Height standard deviation score (SDS) of -2.5 (0.6th percentile) OR ■ Height velocity impaired AND ■ Height SDS of -2 (2.3rd percentile) for bone age
	 Turner's syndrome For initial approval, documentation of the following is required: Diagnosis of Turner Syndrome done through genetic testing AND For patients less than 2 years of age:
	Noonan's syndrome ■ For initial approval, documentation of the following is required: □ Diagnosis of Noonan's syndrome done through genetic testing AND ■ Height standard deviation score (SDS) of -2.5 (0.6 th percentile) □ OR ■ Height velocity impaired AND ■ Height SDS of -2 (2.3rd percentile) for bone age Short stature homeobox-containing gene (SHOX) deficiency ■ For initial approval, documentation of the following is required: □ Diagnosis of SHOX deficiency done through genetic testing ■ Height standard deviation score (SDS) of -2.5 (0.6 th percentile) OR
	Height velocity impaired ANDHeight SDS of -2 (2.3rd percentile) for bone age



	 Chronic kidney disease stage 3 and greater OR kidney transplant For initial approval, documentation of the following is required: Diagnosis of chronic kidney disease stage 3 or higher (CrCl less than 60mL/min) Height velocity (SDS) less than -1.88 for bone age.
	Prader-Willi syndrome ■ For initial approval, documentation of the following is required: □ Diagnosis of Prader-Willi syndrome through genetic testing AND □ Height velocity impaired
	Short Stature born small for gestational age (SGA) with no catch-up growth by 2 years
	 to 4 years of age Birth weight and/or length of at least 2 standard deviations (-2 SD) from the mean for gestational age and sex Height standard deviation score (SDS) of -2.5 (0.6th percentile) Age at start of growth hormone therapy cannot be greater than 10 years Exclusion of other causes of short stature including growth-inhibiting medication, chronic disease, endocrine disorders
	Adult Growth Hormone Deficiency: ■ For initial approval, documentation of the following is required: □ Dose and frequency are appropriate AND □ Documented Growth Hormone Deficiency AND □ Documented IGF-1 outside reference range for patient's sex and age, AND the patient has failed one growth hormone stimulation test (insulin tolerance test-ITT or Glucagon stimulation test when ITT is contraindicated)
	 Reauthorization: Pediatric Indications: requires a documented growth rate increase of at least 2.5 cm over baseline per year AND evaluation of epiphyses (growth plates) documenting they remain open Adult Growth Hormone Deficiency: requires documented clinical improvement and IGF-I within normal reference range for age and sex
Appropriate Treatment Regimen & Other Criteria:	 Documented trial and failure of at least 12 weeks of Norditropin prior to any other daily growth hormone For Skytrofa and Sogroya: Documented trial and failure of at least 12 weeks of Norditropin and one additional daily growth hormone
Exclusion Criteria:	 Pregnancy Elderly adults with age-adjusted low IGF-1 levels and no history of pituitary or hypothalamic disease. Growth Hormone (GH) replacement to enhance athletic performance Diagnosis of: Idiopathic Short Stature (ISS), height standard deviation score (SDS) less than -2.25, and associated with growth rates unlikely to permit attainment of adult height in the normal range
Age	167



Restriction: Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an endocrinologist All approvals are subjects to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



HEPATITIS C DIRECT-ACTING ANTIVIRALS

Affected Medications: MAVYRET (glecaprevir & pibrentasvir), Vosevi (Sofosbuvir/Velpatasvir/Voxilaprevir), Sofosbuvir/Velpatasvir

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design.
	 AASLD (American Association for the Study of Liver Diseases)-supported use with class I or class IIa-Level A recommendation
Required Medical	 Documentation of chronic hepatitis C virus (HCV) by liver biopsy or by Food and Drug
Information:	Administration (FDA)-approved serum blood test
	Current HIV status
	Current Hepatitis B status
	Baseline HCV RNA level within last 3 months
	Documentation that patient is one of the following:
	 Treatment-naïve
	 Treatment experienced, including documentation of previous treatment regimen
	and outcome
	Current documentation of hepatic impairment severity with Child-Pugh Classification OR
	bilirubin, albumin, INR, ascites status, and encephalopathy status to calculate Child-Pugh
	score, within 12 weeks prior to anticipated start of therapy
	Expected survival from non-Hepatitis C-associated morbidity is greater than 12 months
Appropriate	Dose/duration or according to the most recently updated AASLD guideline
Treatment	recommendation (See table below)
Regimen & Other	
Criteria:	
Exclusion Criteria:	Mavyret is contraindicated in patients with moderate and severe hepatic impairment (Child-Pugh B and C)
	Vosevi is not recommended in patients with moderate or severe hepatic impairment
	(Child-Pugh class B or C)
	Concurrent use of Vosevi with rifampin is contraindicated
Age Restriction:	
Prescriber/Site of	All approvals are subject to utilization of the most cost-effective site of care
Care Restrictions:	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
Coverage Duration:	See Appropriate Treatment Regimen & Other Criteria

Recommended Treatment Regimens for Adults and Adolescents 12 years of age and older with Chronic Hepatitis C virus

Treatment History	Cirrhosis Status	Recommended Regimen
Treatment Naïve (Genotype 1-6)		
DAA-Treatment naïve, confirmed reinfection or prior treatment with		SOF/VEL x 12 weeks Mavyret x 8 weeks



Compensated Cirrhosis	SOF/VEL x 12 weeks
	Mavyret x 8 weeks
Decompensated Cirrhosis	SOF/VEL + RBV x 12 weeks SOF/VEL x 24 weeks (if ribavirin ineligible*)
e 1-6)	
t Non-cirrhotic or compensated cirrhosis	Vosevi x 12 weeks Mavyret x 16 weeks (except genotype 3)
Non-cirrhotic or compensated cirrhosis	Vosevi x 12 weeks
Non-cirrhotic or compensated cirrhosis	Mavyret + SOF + RBV x 16 weeks Vosevi x 12 weeks (plus RBV if compensated cirrhosis)
Non-cirrhotic or compensated cirrhosis	Mavyret + SOF + RBV x 16-24 weeks Vosevi + RBV x 24 weeks
	Decompensated Cirrhosis e 1-6) t Non-cirrhotic or compensated cirrhosis Non-cirrhotic or compensated cirrhosis Non-cirrhotic or compensated cirrhosis

Abbreviations: DAA = direct-acting antiviral; PEG = pegylated interferon; RBV = ribavirin; SOF/VEL = sofosbuvir/velpatasvir

*Ribavirin ineligible/intolerance may include: 1) neutrophils less than 750 mm3, 2) hemoglobin less than 10 g/dL, 3) platelets less than 50,000 cells/mm3, autoimmune hepatitis or other autoimmune condition, hypersensitivity or allergy to ribavirin

Recommended Treatment Regimens for children ages 3 to 12 years of age with Chronic Hepatitis C virus

Treatment History	Cirrhosis Status	Recommended Regimen
Treatment Naïve (Genotype 1-6	5)	
DAA-Treatment naïve, confirmed	Non-cirrhotic or compensated cirrhosis	SOF/VEL x 12 weeks
reinfection or prior treatment with PEG/RBV	·	Mavyret x 8 weeks
	Decompensated Cirrhosis	SOF/VEL + RBV x 12 weeks
Treatment Experienced		
treatment regimens in adults if Fl	DA approved for pediatric use. Recommend	
Abbreviations: DAA = direct-acting antiviral; PEG = pegylated interferon; RBV = ribavirin; SOF/VEL =		
sofosbuvir/velpatasvir		

Recommended dosage of SOF/VEL in pediatric patients 3 years of age and older

Body Weight	Dosing of SOF/VEL



Less than 17kg	One 150mg/37.5mg pellet packet once daily
17kg to less than 30kg	One 200mg/50mg pellet packet OR tablet once daily
IAT IPAST KUKO	Two 200mg/50mg pellet packets once daily OR one 400mg/100mg tablet once daily

Recommended dosage of Mavyret in pediatric patients 3 years of age and older

Body Weight	Dosing of Mavyret
Less than 20kg	Three 50mg/20mg pellet packets once daily
20kg to less than 30kg	Four 50mg/20mg pellet packets once daily
30kg to less than 45kg	Five 50mg/20mg pellet packets once daily
45kg and greater OR 12 years of age and older	Three 100mg/40mg tablets once daily



POLICY NAME: **HISTRELIN**

Affected Medications: SUPPRELIN LA (histrelin acetate)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Central precocious puberty (CPP) Gender dysphoria
Required Medical Information:	 Central Precocious Puberty: Documentation of CPP confirmed by basal luteinizing hormone (LH), follicle-stimulating hormone (FSH), and either estradiol or testosterone concentrations Gender Dysphoria: Documentation of all of the following:
Appropriate	All Indications:
Treatment	Approval requires documented treatment failure with leuprolide
Regimen & Other Criteria:	Reauthorization will require documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	
Age Restriction:	2 years of age and older
Prescriber/Site of Care Restrictions:	 Central Precocious Puberty: Prescribed by, or in consultation with, an endocrinologist Gender dysphoria: Diagnosis made and prescribed by, or in consultation with, a specialist in the treatment of gender dysphoria All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



HEREDITARY ANGIOEDEMA

Affected Medications: Berinert, Icatibant Acetate, Sajazir, Ruconest, Kalbitor, Cinryze, Haegarda, Takhzyro, Orladeyo

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Hereditary angioedema attacks, prophylaxis (Cinryze, Haegarda, Takhzyro, Orladeyo)
	 Hereditary angioedema attacks, acute treatment (Berinert, icatibant acetate,
	Sajazir, Kalbitor, Ruconest)
Required Medical	Diagnosis of hereditary angioedema (HAE) classified as one of the following:
Information:	Type I or II HAE confirmed by low C4 levels AND one of the following:
	 Low C1 inhibitor functional or antigenic level less than 50% of the lower limit of
	normal as defined by the laboratory performing test
	"Type III" HAE confirmed by normal C4, C1 inhibitor (functional and antigenic) with one
	of the following:
	 Genetic testing confirming presence of HAE causing mutation such as mutation
	of coagulation factor XII gene (F12 mutation), mutation in the angiopoietin-1
	gene, mutation in the plasminogen gene, mutation in the kininogen 1 gene,
	mutation in the myoferlin gene, mutation in the heparan sulfate 3-
	Osulfotransferase 6 gene
	 Family history of HAE AND documented recurring angioedema attacks that are
	refractory to high dose antihistamines (four times the usual dose)
	Documented full treatment plan and current body weight
	Documentation of number of attacks requiring treatment in the past year
Appropriate	Acute Treatment:
Treatment	Documented history of one of the following:
Regimen & Other	 Non-inflammatory subcutaneous angioedema (without hives) which is recurrent
Criteria:	and lasts greater than 12 hours
	 Abdominal pain without a clear organic cause lasting greater than 6 hours
	Coverage for non-preferred products (Berinert, Kalbitor, Ruconest) requires documentation of one of the following:
	Documented treatment failure to one of the preferred products: icatibant acetate or Sajazir
	Currently receiving treatment with a non-preferred product, excluding via samples or
	manufacturer's patient assistance programs
	For requests to treat more than 3 attacks per month:
	Documentation of current treatment with, or failure, intolerance, or clinical rationale for
	avoidance of, prophylactic therapies
	Authorization for acute treatment will provide a sufficient quantity to treat the average
	number of acute attacks per month plus 1 additional dose
	Prophylaxis Treatment:
L	l ————————————————————————————————————



	History of TWO or more severe attacks per month for the past 3 months (airway swelling,
	debilitating cutaneous or gastrointestinal episodes) despite short term treatment and at
	least one of the following:
	 Disabling symptoms for at least 5 days per month
	 History of at least one laryngeal attack caused by HAE
	Avoidance of possible triggers for HAE attacks such as
	 estrogen containing oral contraceptives/hormone replacement
	o angiotensin-converting-enzyme (ACE) inhibitors
	o dipeptidyl peptidase IV (DPP-4) inhibitors
	Neprilysin inhibitor
	Coverage for non-preferred products (Cinryze, Orladeyo) requires documentation of one of the following:
	Documented treatment failure to the preferred products Haegarda and Takhzyro
	Currently receiving treatment with a non-preferred product, excluding via samples or
	manufacturer's patient assistance programs
	Reauthorization requires documentation of number of acute HAE attacks treated in the past year AND documentation of treatment success defined as reduction of frequency and severity of HAE attack episodes requiring acute therapy by greater than or equal to 50% from baseline.
	Requested dose within the Food and Drug Administration (FDA)-approved label
	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced for all medical infusion drugs
Exclusion Criteria:	Concurrent use of multiple HAE prophylactic treatments (Orladeyo, Haegarda, Takhzyro, Cinryze)
	Concurrent use of multiple HAE acute treatments (Berinert, Kalbitor, Runconest, icatibant acetate, Sajazir)
Age Restriction:	Product specific per FDA labeled indication
Prescriber/Site of	Prescribed by, or in consultation with, an allergist, immunologist, or pulmonologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 3 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



HEREDITARY TYROSINEMIA (HT-1) AGENTS

Affected Medications: NITYR, ORFADIN, NITISINONE

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
Required Medical Information:	 Diagnosis of hereditary tyrosinemia type 1 confirmed by: Presence of succinylacetone (SA) in urine or blood Genetic testing showing a mutation in the gene encoding fumarylacetoacetate hydrolase (FAH) Current patient weight
Appropriate Treatment Regimen & Other Criteria:	 Use as an adjunct to dietary restriction of tyrosine and phenylalanine Orfadin requires: A documented intolerable adverse event to Nityr and the adverse event was not an expected adverse event attributed to the active ingredient
	 Reauthorization: documentation of treatment success confirmed by: Reduction in urine or plasma succinylacetone from baseline Documentation of dietary restriction of tyrosine and phenylalanine
Exclusion Criteria:	Use without dietary restriction of tyrosine and phenylalanine
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a specialist in the treatment of hereditary tyrosinemia or related disorders All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 3 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



Hormone Supplementation under 18 years of age

Affected Medications: Depo-Estradiol oil, Estradiol twice weekly patch, Estradiol weekly patch, Dotti patch, Estradiol tablets, Estradiol gel, Menest, Divigel transdermal, Elestrin gel, Estrogel, Evamist, Premarin tablets, Testosterone Cypionate solution, Testosterone enanthate, testosterone transdermal (gel, patch), Testred capsule, Methitest tablets, Alora Patches, Climara patches, Delestrogen oil, Estrace tablets, Estradiol valerate oil, Lyllana Patch, Menostar Patch, Minivelle Patch, Premarin solution, Vivelle-dot patches

Covered Uses:	Gender dysphoria
0010104 00001	Applies to patients under 18 years of age
	γ γγγιου το μετιουία απίσει το years or ago
Required Medical	Gender dysphoria
Information:	Documentation of all the following:
	 Current Tanner stage 2 or greater OR baseline and current estradiol and
	testosterone levels to confirm onset of puberty
	 Confirmed diagnosis of gender dysphoria that is persistent
	 The patient has the capacity to make a fully informed decision and to give
	consent for treatment
	 Any significant medical or mental health concerns are reasonably well controlled
	A comprehensive mental health evaluation has been completed by a licensed
	mental health professional (LMHP) and provided in accordance with the most
	current version of the World Professional Association for Transgender Health
	(WPATH) Standards of Care
	Note: For requests following pubertal suppression therapy, an updated or new
	comprehensive mental health evaluation must be provided prior to initiation of hormone
	supplementation
Appropriate	Reauthorization requires documentation of treatment success
Treatment	
Regimen & Other	
Criteria:	
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of	Gender Dysphoria: Diagnosis made and prescribed by, or in consultation with, a
Care Restrictions:	specialist in the treatment of gender dysphoria
ourc Nestrictions.	All approvals are subject to utilization of the most cost-effective site of care
	- 7 in approvate are subject to utilization of the most cost encourse site of care
Coverage Duration:	Authorization: 24 months, unless otherwise specified
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HYDROCORTISONE ORAL GRANULES

Affected Medications: ALKINDI SPRINKLE (hydrocortisone oral granules)

Covered Uses: Required Medical	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Glucocorticoid replacement therapy in pediatric patients with adrenocortical insufficiency Diagnosis of adrenal insufficiency confirmed with an adrenal stimulation test
Information:	 Current body surface area (or height and weight to calculate) Current height and weight velocity For adolescents, evaluation of epiphyses (growth plates) documenting they remain open Complete treatment plan including dose in mg/m²/day
Appropriate Treatment Regimen & Other Criteria:	Documented treatment failure with a 6-month trial of two or more of the following:
Exclusion Criteria:	 Use in adolescents who have achieved their adult height Use for stress dosing Use in acute treatment of adrenal crisis or acute adrenal insufficiency Long term use with strong CYP3A4 inducers, unless medically necessary
Age Restriction:	Less than 18 years of age
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a pediatric endocrinologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



HYFTOR

Affected Medications: HYFTOR (sirolimus gel)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design For the treatment of facial angiofibroma (FA) associated with tuberous sclerosis complex (TSC)
Required Medical Information:	 Documented diagnosis of FA associated with TSC which are: Rapidly changing in size and/or number Causing functional interference, pain or bleeding Inhibiting social interactions Current and baseline description of FA including lesion count, associated symptoms and
Appropriate Treatment Regimen & Other Criteria:	 complications, and overall severity Documented treatment failure with laser therapy and/or surgery (such as shave excision, cryotherapy, radiofrequency ablation, or dermabrasion), unless contraindicated Reauthorization requires documentation of a positive clinical response to therapy (decrease in size and/or redness of facial angiofibromas)
Exclusion Criteria: Age Restriction:	 Concurrent use of systemic mammalian target of rapamycin (mTOR) inhibitors Treatment of non-facial angiofibroma
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a dermatologist, oncologist, or neurologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 3 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



HYPOXIA-INDUCIBLE FACTOR PROLYL HYDROXYLASE (HIF PH) INHIBITORS

Affected Medications: JESDUVROQ (daprodustat), VAFSEO (vadadustat)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	 Anemia due to chronic kidney disease (CKD) in adults who have been receiving dialysis
Required Medical	Diagnosis of anemia due to CKD
Information:	Documentation of dialysis use for:
	 Jesduvroq: 4 or more months
	 Vafseo: 3 or more months
	 Documentation of pretreatment hemoglobin level greater than 8 g/dL and less than 12 g/dL
	Adequate iron stores as indicated by current (within the last three months) serum ferritin
	level greater than or equal to 100 mcg/L or serum transferrin saturation greater than or equal to 20%
Appropriate	Documentation of ONE of the following:
Treatment	 Documented hypo-responsiveness to an erythropoiesis stimulating agent (ESA),
Regimen & Other	defined as the need for ONE of the following:
Criteria:	 Greater than 300 IU/kg per week of epoetin alfa
	 Greater than 1.5 mcg/kg per week of darbepoetin
	 Intolerance to BOTH preferred ESA products epoetin alfa-epbx (Retacrit) and
	darbepoetin alfa (Aranesp)
	Reauthorization requires documentation of treatment success and hemoglobin of greater than 8 g/dL and less than 12 g/dL
Exclusion Criteria:	Use in combination with ESAs
	Current uncontrolled hypertension
	Active malignancy
	For Jesduvroq: Major adverse cardiac events (such as myocardial infarction, acute)
	coronary syndrome, stroke, transient ischemic attack, venous thromboembolism) within 3
Age Restriction:	months prior to starting treatment
_	
Prescriber/Site of	Prescribed by, or in consultation with, a specialist, such as a hematologist or
Care Restrictions:	nephrologist
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **IBREXAFUNGERP**

Affected Medications: BREXAFEMME (ibrexafungerp)

Covered Uses: Required Medical Information:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	 RVVC Documentation of three or more episodes of symptomatic vulvovaginal candidiasis infection within the past 12 months
Appropriate	VVC
Treatment	Documented treatment failure with both of the following for the current VVC episode:
Regimen & Other	Vaginally administered treatment (such as clotrimazole cream, miconazole)
Criteria:	cream, terconazole cream or suppository) A 7-day course of fluconazole taken orally every third day for a total of 3 doses (days 1, 4, and 7)
	 RVVC Documented disease recurrence following 10 to 14 days of induction therapy with a topical antifungal agent or oral fluconazole, followed by fluconazole 150 mg once per week for 6 months
	Reauthorization requires documentation of treatment success defined as a reduction in symptomatic vulvovaginal candidiasis episodes, and documentation supporting the need for additional treatment
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Authorization (VVC): 3 months, unless otherwise specified Authorization (RVVC): 6 months, unless otherwise specified



ILARIS

Affected Medications: ILARIS (canakinumab)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Tumor Necrosis Factor Receptor Associated Periodic Syndrome (TRAPS), Hyperimmunoglobulin D syndrome (HIDS)/Mevalonate Kinase Deficiency (MKD), Familial Mediterranean Fever (FMF), Adult-Onset Still's Disease (AOSD), Systemic Juvenile Idiopathic Arthritis (SJIA), Cryopyrin-Associated Periodic Syndromes (CAPS), Gout Flares
D : 114 !! !	T N 15 (B (A) (1B) (B (T) (T) (T)
Required Medical Information:	 Tumor Necrosis Factor Receptor Associated Periodic Syndrome (TRAPS) Confirmed diagnosis of TRAPS with frequent and/or severe recurrent disease (such as recurrent fevers, prominent myalgias, migratory rash, periorbital edema) AND documented genetic defect of TNFRSF1A gene Hyperimmunoglobulin D syndrome (HIDS)/ Mevalonate Kinase Deficiency (MKD) Confirmed diagnosis with one of the following: Elevated serum IgD with or without elevated IgA Genetic testing showing presence of heterozygous or homozygous mutation in the mevalonate kinase (MVK) gene Documentation of 3 or more febrile acute flares within a 6-month period Still's Disease Confirmed diagnosis of Still's Disease, including Adult-Onset Still's Disease (AOSD) and Systemic Juvenile Idiopathic Arthritis (SJIA) in patients 2 years of age and older Documented clinical signs and symptoms including fever, rash, arthritis, arthralgia, myalgia, pharyngitis, pulmonary disease, elevated liver enzymes, C-reactive protein (CRP), erythrocyte sedimentation rate (ESR), serum ferritin Cryopyrin-Associated Periodic Syndromes (CAPS) Confirmed diagnosis of CAPS in patients 4 years and older including Familial Cold Autoinflammatory Syndrome (FCAS) or Muckle-Wells Syndrome (MWS) with one of the following: Elevated inflammatory markers such as CRP and serum amyloid A with two of the following manifestations:
	abnormalities o Genetic testing showing presence of NALP3 mutations
	 Gout Flares Confirmed diagnosis of gout that is refractory to standard therapies Documentation of having 3 or more gout flares in the past 12 months
Appropriate Treatment Regimen & Other Criteria:	 TRAPS Documented clinical failure to episodic treatment with nonsteroidal anti-inflammatory drugs (NSAIDs), glucocorticoids (prednisone or prednisolone), and a minimum 12-week trial with Enbrel



	HIDS/MKD Documented treatment failure to episodic treatment with nonsteroidal anti-inflammatory drugs (NSAIDs), glucocorticoids, and anakinra EME
	 FMF Documented treatment failure with maximal tolerable dose of colchicine (3 mg daily in adults and 2 mg daily in children) Documentation of frequent and/or severe recurrence disease despite adequate treatment with at least 12 weeks of anakinra
	Still's Disease
	Documentation of frequent and/or severe recurrent disease despite adequate treatment with a minimum 12-week trial with each of the following:
	CAPS
	Documentation of treatment failure with a minimum 12-week trial with anakinra
	Gout Flares
	 Documented treatment failure with all of the following for the symptomatic treatment of gout flares:
	 Prescription strength NSAIDs (naproxen, indomethacin, diclofenac, meloxicam, or celecoxib) Colchicine Glucocorticoids (oral or intraarticular)
	Reauthorization requires documentation of treatment success
Exclusion Criteria:	 Treatment of neonatal onset multisystem inflammatory disorder (NOMID) or chronic infantile neurological cutaneous and articular syndrome (CINCA), rheumatoid arthritis, chronic obstructive pulmonary disease (COPD), type 2 diabetes mellitus Use in combination with tumor necrosis factor (TNF) blocking agents (e.g., Enbrel, Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Cimzia, Remicade, Simponi), Kineret, or Arcalyst
Age Restriction:	FMF, HIDS/MKD, juvenile idiopathic arthritis, TRAPS: 2 years of age and older
	CAPS: 4 years of age and older
	Gout Flares: 18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, an allergist, immunologist, or rheumatologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 6 months, unless otherwise specified



POLICY NAME: ILOPROST

Drug Name: VENTAVIS (iloprost)

	All Facility I Day A Indicator (FDA)	
Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design.	
	plan design	
	 Pulmonary Arterial Hypertension (PAH) World Health Organization (WHO) Group 1 	
	Pulmonary Arterial Hypertension (PAH) WHO Group 1	
Required	 Documentation of PAH confirmed by right-heart catheterization meeting the following 	
documentation:	criteria:	
	Mean pulmonary artery pressure of at least 20 mm Hg	
	 Pulmonary capillary wedge pressure less than or equal to 15 mm Hg 	
	 Pulmonary vascular resistance of at least 2.0 Wood units 	
	New York Heart Association (NYHA)/World Health Organization (WHO) Functional Class	
	Ill or higher symptoms	
	Documentation of Acute Vasoreactivity Testing (positive result requires trial/failure to	
	calcium channel blockers) unless there are contraindications:	
	 Low systemic blood pressure (systolic blood pressure less than 90) 	
	o Low cardiac index	
	OR	
	 Presence of severe symptoms (functional class IV) 	
Appropriate	Documentation of inadequate response or intolerance to the following therapy classes is	
Treatment	required:	
Regimen:	 PDE5 inhibitors AND 	
	 Endothelin receptor antagonists (exception WHO Functional Class IV) 	
	Reauthorization requires documentation of treatment success defined as one or more of the	
	following:	
	Improvement in walking distance	
	Improvement in exercise ability	
	Improvement in pulmonary function	
	Improvement or stability in WHO functional class	
Exclusion Criteria:		
Age Restriction:		
Prescriber/Site of	Prescribed by, or in consultation with, a cardiologist or a pulmonologist	
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care	
Coverage Duration:	Authorization: 12 months, unless otherwise specified	



IMMUNE GLOBULIN

Affected Medications: ASCENIV, BIVIGAM, FLEBOGAMMA, GAMMAGARD LIQUID/S-D, GAMMAPLEX, GAMUNEX-C, OCTAGAM, PANZYGA, PRIVIGEN, GAMMASTAN, ALYGLO

Covered Uses: All Food and Drug Administration (FDA)-approved and compendia-supported uses not otherwise excluded by plan design as follows: Primary immunodeficiency (PID)/Wiskott - Aldrich syndrome Idiopathic thrombocytopenia purpura (ITP) Chronic Inflammatory Demyelinating Polyneuropathy (CIDP) Guillain-Barre Syndrome (Acute inflammatory polyneuropathy) Multifocal Motor Neuropathy Pediatric HIV: Bacterial control or prevention Myasthenia Gravis 0 Dermatomyositis/Polymyositis Complications of transplanted solid organ (kidney, liver, lung, heart, pancreas) and bone marrow transplant Stiff-Person Syndrome Allogeneic Bone Marrow or Stem Cell Transplant Kawasaki's disease (Pediatric) Fetal alloimmune thrombocytopenia (FAIT) Hemolytic disease of the newborn Auto-immune Mucocutaneous Blistering Diseases Chronic lymphocytic leukemia with associated hypogammaglobulinemia (CLL) Toxic Shock Syndrome Pediatric Acute-Onset Neuropsychiatric Syndrome (PANS)/Pediatric Autoimmune Neuropsychiatric Disorder Associated with Streptococcal Infections (PANDAS) **Initial Approval** Primary immunodeficiency (PID)/Wiskott - Aldrich syndrome: Criteria: Includes but not limited to: X-linked agammaglobulinemia, common variable

Includes but not limited to: X-linked agammaglobulinemia, common variable immunodeficiency (CVID), transient hypogammaglobulinemia of infancy, IgG subclass deficiency with or without IgA deficiency, antibody deficiency with near normal immunoglobulin levels) and combined deficiencies (severe combined immunodeficiencies, ataxia-telangiectasia, x-linked lymphoproliferative syndrome)

- Documentation of one of the following:
 - o IgG level less than 200
 - Low IgG levels (below the laboratory reference range lower limit of normal) AND
 a history of multiple hard to treat infections as indicated by at least one of the
 following:
 - Four or more ear infections within 1 year
 - Two or more serious sinus infections within 1 year
 - Two or more months of antibiotics with little effect
 - Two or more pneumonias within 1 year
 - Recurrent or deep skin abscesses
 - Need for intravenous antibiotics to clear infections
 - Two or more deep-seated infections including septicemia

AND

- Documentation showing a deficiency in producing antibodies in response to vaccination including all the following:
 - Titers that were drawn before challenging with vaccination
 - Titers that were drawn between 4 and 8 weeks after vaccination



Idiopathic thrombocytopenia purpura (ITP):

For Acute disease state:

 Documented use to manage acute bleeding due to severe thrombocytopenia (platelet counts less than 30,000/microliter)

OR

• To increase platelet counts prior to invasive surgical procedures, such as splenectomy (platelet count less than 100,000/microliter)

OR

 Documented severe thrombocytopenia (platelet count less than 20,000/microliter) and is considered to be at risk for intracerebral hemorrhage

Chronic Immune Thrombocytopenia (CIT):

- Documentation of increased risk for bleeding as indicated by a platelet count less than 30,000/microliter
- History of failure, contraindication, or intolerance with corticosteroids
- Duration of illness more than 6 months

Chronic Inflammatory Demyelinating Polyneuropathy (CIDP):

- Documented baseline in strength/weakness using objective clinical measuring tool (INCAT, Medical Research Council (MRC) muscle strength, 6 MWT, Rankin, Modified Rankin)
- Documented disease course is progressive or relapsing and remitting for 2 months or longer
- Abnormal or absent deep tendon reflexes in upper or lower limbs
- Electrodiagnostic testing indicating demyelination with one of the following:
 - Motor distal latency prolongation in 2 nerves
 - Reduction of motor conduction velocity in 2 nerves
 - o Prolongation of F-wave latency in 2 nerves
 - o Absence of F-waves in at least 1 nerve
 - Partial motor conduction block of at least 1 motor nerve
 - Abnormal temporal dispersion in at least 2 nerves
 - Distal CMAP duration increase in at least 1 nerve
- Cerebrospinal fluid (CSF) analysis indicates all the following (if electrophysiologic findings are nondiagnostic):
 - CSF white cell count of less than 10 cells/mm3
 - CSF protein is elevated (greater than 45 mg/dL)
- Refractory to or intolerant of corticosteroids (prednisolone, prednisone) given in therapeutic doses over at least three months

Guillain-Barre Syndrome (Acute inflammatory polyneuropathy):

- Documentation that the disease is severe (aid required to walk)
- Onset of symptoms are recent (less than 1 month)

Multifocal Motor Neuropathy (MMN):



- Slowly progressive or stepwise progressive, focal, asymmetric limb weakness over at least one month
- Partial conduction block or abnormal temporal dispersion conduction must be present in at least 2 nerves
- Absence of upper motor neuron signs and bulbar involvement
- Baseline in strength/weakness has been documented using objective clinical measuring tool (e.g., Inflammatory Neuropathy Cause and Treatment (INCAT) Disability Score, Medical Research Council (MRC) muscle strength, 6 Minute walk test, Rankin, Modified Rankin

Pediatric HIV: Bacterial control or prevention:

- Approved for those 13 years of age and younger with HIV diagnosis
- Documented hypogammaglobulinemia (IgG less than 400 mg/dL) OR
- Functional antibody deficiency as demonstrated by either poor specific antibody titers or recurrent bacterial infections

Myasthenia Gravis:

- Documented myasthenic crisis (impending respiratory or bulbar compromise)
- Documented use for an exacerbation (difficulty swallowing, acute respiratory failure, functional disability leading to discontinuation of physical activity)
- Documented failure with conventional therapy alone (azathioprine, cyclosporine and/or cyclophosphamide)

Dermatomyositis/Polymyositis:

- Documented severe active disease state on physical exam
- Documentation of at least two of the following:
 - o Proximal muscle weakness in all upper and/or lower limbs
 - Elevated serum creatine kinase (CK) or aldolase level
 - Interstitial lung disease (ILD)
 - Skin findings such as Gottron papules, Gottron sign, heliotrope eruption, poikiloderma
 - Nailfold abnormalities
 - Hyperkeratosis and fissuring of palms and lateral fingers
- Documented failure with a trial of corticosteroids (such as prednisone)
- Documented failure with a trial of an immunosuppressant (methotrexate, azathioprine, cyclophosphamide)

<u>Complications of transplanted solid organ (kidney, liver, lung, heart, pancreas) and bone marrow transplant</u>:

Coverage is provided for one or more of the following:

- Suppression of panel reactive anti-HLA antibodies prior to transplantation
- Treatment of antibody mediated rejection of solid organ transplantation
- Prevention of cytomegalovirus (CMV) induced pneumonitis

Stiff-Person Syndrome:



- Documented anti-GAD antibodies
- Documented failure with at least 2 of the following treatments: benzodiazepines, baclofen, phenytoin, clonidine and/or tizanidine

Allogeneic Bone Marrow or Stem Cell Transplant:

- Approved in use for prevention of acute Graft- Versus- Host Disease (GVHD) or infection (such as cytomegalovirus)
- Documentation that the bone marrow transplant (BMT) was allogeneic
- Transplant was less than 100 days ago

Kawasaki's Disease (Pediatric):

- Diagnosis or suspected diagnosis of Kawasaki's disease
- 13 years of age and under

Fetal alloimmune thrombocytopenia (FAIT):

- Documentation of one or more of the following:
 - o Previous FAIT pregnancy
 - o Family history of the disease
 - Screening reveals platelet alloantibodies
- Authorization is valid until delivery date only

Hemolytic disease of the newborn:

Diagnosis or suspected diagnosis of hemolytic disease in newborn patient

Auto-immune Mucocutaneous Blistering Diseases:

- Diagnosis confirmed by biopsy of one of the following:
 - Pemphigus vulgaris
 - o Pemphigus foliaceus
 - Bullous Pemphigoid
 - o Mucous Membrane Pemphigoid (Cicatricial Pemphigoid)
 - o Epidermolysis bullosa aquisita
 - Pemphigus gestationis (Herpes gestationis)
 - Linear IgA dermatosis
- Documented severe disease that is extensive and debilitating
- Disease is progressive and refractory to a trial of conventional combination therapy with corticosteroids and immunosuppressive treatment (azathioprine, cyclophosphamide, mycophenolate mofetil)

Chronic lymphocytic leukemia (CLL) with associated hypogammaglobulinemia:

- Documentation of an IgG level less than 500 mg/dL
- Documented history of recurrent or chronic infections that have required intravenous antibiotics or hospitalization



Toxic Shock Syndrome:

Diagnosis or suspected diagnosis of toxic shock syndrome

<u>Pediatric Acute-Onset Neuropsychiatric Syndrome (PANS)/Pediatric Autoimmune</u> Neuropsychiatric Disorder Associated with Streptococcal Infections (PANDAS):

- A clinically appropriate trial of two or more less-intensive treatments was either not
 effective, not tolerated, or did not result in sustained improvement in symptoms, as
 measured by a lack of clinically meaningful improvement on a validated instrument
 directed at the patient's primary symptom complex. Treatments may be given
 concurrently or sequentially and may include:
 - Selective-serotonin reuptake inhibitor SSRI (e.g., fluoxetine, fluvoxamine, sertraline)
 - Behavioral therapy
 - o Nonsteroidal anti-inflammatory (NSAID) (e.g., naproxen, diclofenac, ibuprofen)
 - o Oral and IV corticosteroids (e.g., prednisone, methylprednisolone)
- Documentation of a consultation with a pediatric subspecialist (or adult subspecialist for adolescents) and the consulted subspecialist and the patient's primary care provider recommend the treatment

Renewal Criteria:

Primary immunodeficiency (PID)

 Renewal requires disease response as evidenced by a decrease in the frequency and/or severity of infections

Chronic Immune Thrombocytopenia (Chronic ITP or CIT)

 Renewal requires disease response as indicated by the achievement and maintenance of a platelet count of at least 50 as necessary to reduce the risk for bleeding

Chronic Inflammatory Demyelinating Polyneuropathy (CIDP)

 Renewal requires documentation of a documented clinical response to therapy based on an objective clinical measuring tool (e.g., INCAT, Medical Research Council (MRC) muscle strength, 6 Minute walk test, Rankin, Modified Rankin)

Multifocal Motor Neuropathy (MMN)

 Renewal requires documentation that there has been a demonstrated clinical response to therapy based on an objective clinical measuring tool (INCAT, Medical Research Council (MRC) muscle strength, 6 Minute walk test, Rankin, Modified Rankin)

Pediatric HIV: Bacterial control or prevention

13 years of age or less

Dermatomyositis/Polymyositis

 Renewal requires documentation that CPK (Creatine phosphokinase) levels are lower and documentation of clinically significant improvement above baseline per physical exam

Complications of transplanted solid organ (kidney, liver, lung, heart, pancreas) and bone marrow transplant

Renewal requires documentation of clinically significant disease response

Stiff Person Disease

 Renewal requires documentation of a clinically significant improvement over baseline per physical exam

Allogeneic Bone Marrow or Stem Cell Transplant



- Renewal requires documentation that the IgG is less than or equal to 400mg/dL; AND
- Therapy does not exceed one year past date of allogeneic bone marrow transplantation **Auto-immune mucocutaneous blistering diseases:**
- Renewal requires a documented clinically significant improvement over baseline per physical exam

Chronic lymphocytic leukemia (CLL) with associated hypogammaglobulinemia

 Renewal requires disease response as evidenced by a decrease in the frequency and/or severity of infections

Pediatric Acute-Onset Neuropsychiatric Syndrome (PANS)/Pediatric Autoimmune Neuropsychiatric Disorder Associated with Streptococcal Infections (PANDAS)

- Renewal requires all of the following:
 - o Documentation of a clinical reevaluation at three months after treatment initiation
 - Documentation of clinically meaningful improvement in the results of clinical testing with a validated instrument (which must be performed pretreatment and posttreatment)

Dosing and Coverage Duration:

- Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced
- Authorization durations are as stated below, unless otherwise specified

Indication	Dose	Approval Duration
PID	Up to 800 mg/kg every 3 to 4 weeks	Initial: up to 3 months Reauthorization: up to 12
		months
CIDP	2 g/kg divided over 2-5 days for one	Initial: up to 3 months
	dose then maintenance dosing of 1	Reauthorization: up to 12
	g/kg every 21 days	months
ITP	1 g/kg once daily for 1-2 days	Acute ITP:
		Approval: 1 month only
	May be repeated monthly for	Chronic ITP:
	chronic ITP	 Initial: up to 3 months
		Reauthorization: up to 12 months
FAIT	1 g/kg/week until delivery	Authorization is valid until
	gridgridgridgridgridgridgridgridgridgrid	delivery date only
Kawasaki's Disease	Up to 2 g/kg x 1 single dose	Approval: 1 month only
(pediatric		
patients)		
MMN	2 g/kg divided over 2-5 days in a	Initial approval: 1 month
	28-day cycle	Reauthorization: up to 12
	May be repeated monthly	months
CLL	400 mg/kg every 3 to 4 weeks	Approval: up to 6 months
Pediatric HIV	400 mg/kg every 28 days	Initial: up to 3 months
		Reauthorization: up to 12 months



	Guillain-Barre	400 mg/kg once daily for 5 days	Approval: maximum of 2 rounds of therapy within 6 weeks of onset; 2 months maximum
	Myasthenia Gravis	Up to 2 g/kg x 1 dose (acute attacks)	Approval: 1 month (one course of treatment)
	Auto- immune blistering diseases	Up to 2 g/kg divided over 5 days in a 28-day cycle	Approval: up to 6 months
	Dermatomyositis /Polymyositis	Up to 2 g/kg given over 2-5 days in a 28-day cycle	Initial: up to 3 months Reauthorization: up to 6 months
	Allogeneic Bone Marrow or Stem Cell Transplant	500 mg/kg/week x 90 days, then 500 mg/kg/month up to one-year post-transplant	Initial: up to 3 months Reauthorization: until up to one-year post-transplant
	Complications of transplanted solid organ: (kidney, liver, lung, heart, pancreas) transplant	2 g/kg divided over 5 days in a 28-day cycle	Initial: up to 3 months Reauthorization: up to 12 months
	Stiff Person Syndrome	2 g/kg divided over 5 days in a 28-day cycle	Initial: up to 3 months Reauthorization: up to 12 months
	Toxic shock syndrome	1 g/kg on day 1, followed by 500 mg/kg once daily on days 2 and 3	Approval: 1 month (one course of treatment)
	Hemolytic disease of the newborn	1 g/kg x 1 dose, may be repeated once if needed	Approval: 1 month (one course of treatment)
	PANS/PANDAS	Each dose: Up to 2 g/kg divided over 2-5 days	Initial: up to 3 months (3 monthly doses) Reauthorization: up to 3 months (3 monthly doses)
Care Restrictions:	rheumatologist, imn	by a specialist for the condition being nunologist, hematologist) bject to utilization of the most cost-eff	,



INCLISIRAN

Affected Medications: LEQVIO (inclisiran subcutaneous injection)

_	All Food and Drug Administration (FDA) approved indications not otherwise evaluded by			
Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design.			
	plan design			
	 Primary hyperlipidemia (including heterozygous familial hypercholesterolemia 			
	[HeFH])			
	 Secondary prevention in atherosclerotic cardiovascular disease (ASCVD) 			
Required Medical	All Indications			
Information:	Documentation of baseline (untreated) low-density lipoprotein cholesterol (LDL-C)			
	Primary Hyperlipidemia (non-familial)			
	Documentation of baseline (untreated) LDL-C of at least 190 mg/dL			
	<u>HeFH</u>			
	Diagnosis confirmed by ONE of the following:			
	 Minimum baseline LDL-C of 160 mg/dL in adolescents or 190 mg/dL in adults AND 1 first-degree relative affected 			
	 Presence of one abnormal LDL-C-raising gene defect (e.g., LDL receptor 			
	[LDLR], apolipoprotein B [apo B], proprotein convertase subtilisin kexin type 9			
	[PCSK9] loss-of-function mutation, or LDL receptor adaptor protein 1			
	[LDLRAP1])			
	 World Health Organization (WHO)/Dutch Lipid Network criteria score of at least 8 			
	points			
	Definite FH diagnosis per the Simon Broome criteria			
	Clinical ASCVD			
	Documentation of established ASCVD, confirmed by at least ONE of the following:			
	 Acute coronary syndromes (ACS) 			
	History of myocardial infarction (MI)			
	Stable or unstable angina			
	Coronary or other arterial revascularization			
	 Stroke or transient ischemic attack 			
	 Peripheral artery disease (PAD) presumed to be of atherosclerotic origin 			
Appropriate	All Indications			
Treatment	Documentation of intent to take alongside maximally tolerated doses of statin and/or			
Regimen & Other	ezetimibe, unless otherwise contraindicated			
Criteria:	OR			
	History of statin intolerance requires documentation of ONE of the following:			
	 Statin-associated rhabdomyolysis occurred with statin use and was confirmed by 			
	a creatinine kinase (CK) level at least 10 times the upper limit of normal			
	 Statin-associated muscle symptoms (e.g., myopathy, myalgia) occurred with 			
	statin use and was confirmed by BOTH of the following:			
	 A minimum of two different statin trials, with at least one being a 			



hydrophilic statin (rosuvastatin, pravastatin) A re-challenge of each statin (muscle symptoms stopped when each was discontinued and restarted upon re-initiation) Primary Hyperlipidemia/HeFH Documented treatment failure with minimum 12-week trial with ALL of the following, shown by an inability to achieve LDL-C reduction of 50% or greater OR LDL-C less than 100 ma/dL: 0 Maximally tolerated statin/ezetimibe therapy Repatha **Clinical ASCVD** Documented treatment failure with minimum 12 weeks of consistent maximally tolerated combination statin/ezetimibe therapy, as shown by **ONE** of the following: Current LDL-C of at least 70 mg/dL Current LDL-C of at least 55 mg/dL in patients at very high risk of future ASCVD events, based on history of multiple major ASCVD events OR 1 major ASCVD event + multiple high-risk conditions (see below) Documented treatment failure or intolerance to minimum 12-week trial of Repatha **Major ASCVD Events High-Risk Conditions** ACS within the past 12 months Age 65 years and older History of MI (distinct from HeFH ACS event) Prior coronary artery bypass or Ischemic stroke percutaneous intervention (outside of major ASCVD events) Symptomatic PAD **Diabetes** Hypertension Chronic kidney disease Current smoking History of congestive heart failure Reauthorization requires an updated lipid panel showing a clinically significant reduction in baseline LDL-C and continued adherence to therapy **Exclusion Criteria:** Concurrent use with PCSK9 monoclonal antibodies (e.g., Repatha, Praluent) Age Restriction: 18 years of age and older Prescriber/Site of Prescribed by, or in consultation with, a cardiologist, endocrinologist, or lipid specialist Care Restrictions: All approvals are subject to utilization of the most cost-effective site of care **Coverage Duration:** Authorization: 12 months, unless otherwise specified



INEBILIZUMAB-CDON

Affected Medications: UPLIZNA (inebilizumab-cdon)

Covered Uses:	plan design ○ Neuromyelitis o aquaporin-4 (AC	inistration (FDA)-approved indications not otherwise excluded by ptica spectrum disorder (NMOSD) in adults who are anti- QP4) antibody positive
Required Medical Information:	NMOSD	
	Clinical presentation	Possible MRI findings
	Diencephalic syndrome	Periependymal lesionHypothalamic/thalamic lesion
	Acute cerebral syndrome	 Extensive periependymal lesion Long, diffuse, heterogenous, or edematous corpus callosum lesion Long corticospinal tract lesion Large, confluent subcortical or deep white matter lesion
	History of at least 1 attac requiring rescue therapy	ck in the past year, or at least 2 attacks in the past 2 years,
Appropriate Treatment Regimen & Other Criteria:	 Documentation of inadequate response, contraindication, or intolerance to each of the following: Rituximab (preferred products: Riabni, Ruxience) Satralizumab-mwge (Enspryng) 	
		ocumentation of treatment success
Exclusion Criteria:	 Active Hepatitis B Virus Active or untreated later Concurrent use with other 	
Age Restriction:	18 years of age and older	



Prescriber/Site of Care Restrictions:	•	Prescribed by, or in consultation with, a neurologist or neuro-ophthalmologist. All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified	
	•	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: INHALED MANNITOL

Affected Medications: BRONCHITOL (mannitol)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	Add-on maintenance therapy to improve pulmonary function in cystic fibrosis
Required Medical	Documentation of cystic fibrosis (CF) diagnosis confirmed by appropriate genetic or
Information:	diagnostic testing
	 Additional testing should include evaluation of overall clinical lung status and respiratory function (e.g., pulmonary function tests, lung imaging, etc.)
Appropriate	Documented treatment failure with 6-month trial of twice daily inhaled hypertonic saline
Treatment	(at least 80% adherence), unless contraindicated or intolerable. Treatment failure
Regimen & Other	defined as one or more of the following:
Criteria:	 Increased pulmonary exacerbations from baseline
	 Decrease in FEV1
	Requests for Bronchitol 7-day and 4-week treatment packs for add-on maintenance
	therapy:
	 Documentation confirming successful completion of the Bronchitol Tolerance Test (BTT)
	 Prescribed in conjunction with a short-acting bronchodilator and standard therapies for CF
	Reauthorization requires documentation of a clinically significant response to therapy
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of	All approvals are subject to utilization of the most cost-effective site of care
Care Restrictions:	
Coverage Duration:	Authorization: 12 months, unless otherwise specified



INTRAVITREAL ANTI-VEGF THERAPY

Affected Medications: LUCENTIS (ranibizumab injection), EYLEA (aflibercept), EYLEA HD (aflibercept), BEOVU (brolucizumab), SUSVIMO (ranibizumab implant), VABYSMO (faricimab), PAVBLU (aflibercept-ayyh)

Covered Uses:	 All Food and Drug Administration (FDA)-approved, or compendia supported, indications not otherwise excluded by plan design Neovascular (Wet) Age-Related Macular Degeneration (AMD) Eylea, Eylea HD, Pavblu, Lucentis, Susvimo, Beovu, Vabysmo, Byooviz, Cimerli Macular Edema Following Retinal Vein Occlusion (RVO) Eylea, Pavblu, Lucentis, Byooviz, Cimerli, Vabysmo Diabetic Macular Edema (DME) Eylea, Eylea HD, Pavblu, Lucentis, Vabysmo, Beovu, Cimerli Diabetic Retinopathy (DR) in patients with Diabetes Mellitus Eylea, Eylea HD, Pavblu, Lucentis, Cimerli Myopic Choroidal Neovascularization (mCNV) Lucentis, Byooviz, Cimerli Retinopathy of Prematurity (ROP) Eylea, Lucentis, Byooviz, Cimerli
Required Medical	Anticipated treatment course with dose and frequency clearly stated in chart notes
Information:	
Appropriate	Eylea/Pavblu Dosing
Treatment	Coverage for the non-preferred products Eylea or Pavblu is provided when one of the following criteria is most:
Regimen & Other Criteria:	of the following criteria is met:
Other Officeria.	 Currently receiving treatment with Eylea or Pavblu, excluding when the product is obtained as samples or via manufacturer's patient assistance programs. A documented inadequate response or intolerable adverse event with all the preferred products (Avastin, AND Byooviz or Cimerli) Documentation of treatment-naïve retinopathy of prematurity (ROP) in a preterm infant 32 weeks or younger
	AMD – 2 mg (0.05 mL) every 4 weeks for the first 3 injections followed by 2 mg (0.05 mL) every 8 weeks
	 Continued every 4-week dosing requires documented clinical failure to every 8-week maintenance dosing
	 RVO - 2 mg (0.05 mL) every 4 weeks DME and DR - 2 mg (0.05 mL) every 4 weeks for the first 5 injections followed by 2
	mg (0.05 mL) every 8 weeks
	ROP – 0.4 mg (0.01 mL) as a single injection per affected eye(s); dose may be repeated up to 2 times with a minimum treatment interval between doses of at least 10 days (maximum of 3 doses total)
	Eylea HD Dosing
	Coverage for the non-preferred product Eylea HD is provided when one of the
	following criteria is met:
	 Currently receiving treatment with Eylea HD, excluding when the product is obtained as samples or via manufacturer's patient assistance programs.
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- A documented inadequate response or intolerable adverse event with all the preferred products (Avastin AND Byooviz or Cimerli)
- AMD and DME 8 mg (0.07 mL) every 4 weeks for the first 3 injections, followed by 8 mg (0.07 mL) every 8 to 16 weeks
 - Every 4-week dosing is limited to the first 3 injections only
- DR 8 mg (0.07 mL) every 4 weeks for the first 3 injections, followed by 8 mg (0.07 mL) every 8 weeks to 12 weeks
 - Every 4-week dosing is limited to the first 3 injections only

Lucentis Dosing

- Coverage for the non-preferred product Lucentis is provided when the following criteria is met:
 - A documented inadequate response or intolerable adverse event with all of the preferred products (Avastin, Byooviz, and Cimerli)
- AMD and RVO maximum 0.5 mg every 4 weeks
- DME and DR 0.3 mg every 4 weeks
- mCNV- 0.5 mg every 4 weeks for up to 3 months
- ROP 0.1 to 0.3 mg as a single injection in the affected eye(s); dose may be repeated up to 2 times with a minimum treatment interval between doses of 28 days (maximum of 3 doses total)

Beovu Dosing

- Coverage for the non-preferred product Beovu is provided when either of the following criteria is met:
 - Currently receiving treatment with Beovu, excluding when the product is obtained as samples or via manufacturer's patient assistance programs.
 - A documented inadequate response or intolerable adverse event with all the preferred products (Avastin, AND Byooviz or Cimerli)
- AMD 6 mg every month for the first three doses followed by 6 mg every 8 to 12 weeks
- DME 6 mg every six weeks for the first five doses followed by 6 mg every 8 to 12 weeks

Susvimo Dosing

- Coverage for the non-preferred product Susvimo is provided when the following criteria is met:
 - A documented inadequate response or intolerable adverse event with all of the preferred products (Avastin, Byooviz, and Cimerli)
- Must be established on ranibizumab (Lucentis, Byooviz, or Cimerli) injections with response to treatment for a minimum of 6 months at standard dosing (0.5 mg every 4 weeks)
- AMD 2 mg administered continuously via ocular implant with refills every 24 weeks.

Vabysmo Dosing

- Coverage for the non-preferred product Vabysmo is provided when either of the following criteria is met:
 - Currently receiving treatment with Vabysmo, excluding when the product is obtained as samples or via manufacturer's patient assistance programs.
 - A documented inadequate response or intolerable adverse event with all the



	preferred products (Avastin, AND Byooviz or Cimerli)
	AMD – 6 mg every 4 weeks for the first 4 injections followed by 6 mg every 8 to 16
	weeks
	 Some patients may require continued every 4-week injections following the
	initial doses
	DME
	by 6 mg every 8 weeks
	 Variable interval regimen: 6 mg once every 4 weeks for at least the first 4
	injections followed by 6 mg every 4 to 16 weeks (based on visual
	assessments)
	 Some patients may require continued every 4-week injections following the
	initial doses
	RVO - 6 mg (0.05 mL) every 4 weeks for up to 6 months
	Reauthorization requires documentation of vision stability defined as losing fewer than 15
	letters of visual acuity and/or improvements in visual acuity with evidence of decreased
	leakage and/or fibrosis (central retinal thickness)
Exclusion	Evidence of a current ocular or periocular infections
Criteria:	Active intraocular inflammation
Age	
Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, an ophthalmologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage	Macular Edema Following Retinal Vein Occlusion (RVO) for Vabysmo
Duration:	Authorization: 6 months with no reauthorization, unless otherwise specified
	Retinopathy of Prematurity (ROP)
	Authorization: 3 months with no reauthorization, unless otherwise specified
	Authorization: 5 months with no reauthorization, unless otherwise specified
	All other indications
	Initial Authorization: 6 months, unless otherwise specified
1	Reauthorization: 12 months, unless otherwise specified



INTRAVITREAL COMPLEMENT INHIBITORS

Affected Medications: SYFOVRE (pegcetacoplan), IZERVAY (avacincaptad pegol)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Treatment of geographic atrophy (GA) secondary to age-related macular
	degeneration (AMD)
Required Medical	Diagnosis of geographic atrophy (GA) secondary to age-related macular degeneration
Information:	(AMD) confirmed by all the following:
	 Fundus Autofluorescence (FAF) imaging showing:
	 Total GA area size between 2.5 and 17.5 mm²
	 If GA is multifocal, at least 1 focal lesion that is 1.25 mm² or greater
	Best-corrected visual acuity (BCVA) using Early Treatment Diabetic Retinopathy Study
	(ETDRS) charts
	 Must be 24 letters or greater (approximately 20/320 Snellen equivalent)
Appropriate	Dosing not to exceed:
Treatment	 Every 25-day dosing for Syfovre
Regimen & Other	 Every 30-day dosing with a maximum duration of 12 months for Izervay
Criteria:	
	Reauthorization
	Syfovre requires:
	 Documentation of treatment success as determined by treating provider
	o BCVA remains 24 letters or greater
	• Izervay:
	No reauthorization - maximum duration up to 12 months
Exclusion Criteria:	Presence of choroidal neovascularization in the eye(s) receiving treatment
Age Restriction:	60 years of age and older for Syfovre
	50 years of age and older for Izervay
Prescriber/Site of	Prescribed by, or in consultation with, an ophthalmologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



INTRON-A

Affected Medications: INTRON-A, INTRON-A WITH DILUENT (interferon alfa-2b)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher Hypereosinophilic Syndrome (HES) in patients that are consistently symptomatic or with evidence of end-organ damage.
Required Medical Information:	 For Hepatitis B and C: Documentation of intolerance to or clinical rationale for avoidance of PEGylated interferon. HES: documentation of steroid resistant disease OR disease responding only to high-dose steroids and the addition of a steroid-sparing agent would be beneficial. Non-lymphocytic variants of HES will also require documented failure with at least 12 weeks of hydroxyurea prior to interferon-alfa approval. Recent liver function tests, comprehensive metabolic panel, complete blood count with differential, TSH (within past 3 months) Documentation of performance status, disease staging, all prior therapies used, and anticipated treatment course Reauthorization: documentation of disease responsiveness to therapy
Appropriate Treatment Regimen & Other Criteria:	 Patients with preexisting cardiac abnormalities and/or advanced cancer: recent electrocardiogram Chest X ray for patients with pulmonary disorders Recent ophthalmologic exam at baseline for all patients Uncontrolled severe mental health illness should be addressed before use and monitored during treatment
Exclusion Criteria:	Autoimmune hepatitis Decompensated liver disease
Age Restriction:	 Hepatitis B: greater than or equal to 1 year of age Hepatitis C: greater than or equal to 3 years of age All other indications greater than or equal to 18 years of age
Prescriber/Site of Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



ISAVUCONAZONIUM SULFATE

Affected Medications: CRESEMBA (isavuconazonium sulfate)

Required Medical Information:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Invasive aspergillosis Invasive mucormycosis Diagnosis of invasive aspergillosis or invasive mucormycosis confirmed by one or more of the following: Sputum fungal staining and culture Biopsy showing aspergillosis or mucormycosis organisms Serum biomarkers such as galactomannan, beta-D-glucan assays, or polymerase chain reaction (PCR) testing
Appropriate	Aspergillosis
Treatment	Documented treatment failure or intolerable adverse event with at least a 6-week trial of
Regimen & Other	all of the following:
Criteria:	 Voriconazole
	o Posaconazole
	 Mucormycosis Documented treatment failure or intolerable adverse event with at least a 6-week trial of one of the following: ○ Amphotericin B (if request is for initial therapy) ○ Posaconazole (if request is for oral step-down therapy after initial therapy)
	Reauthorization will require documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	Familial short QT syndrome
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, an infectious disease specialist, transplant
Care Restrictions:	physician, or oncologist
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 3 months, unless otherwise specified Reauthorization: 3 months, unless otherwise specified



POLICY NAME: LAROTRECTINIB

Affected Medications: VITRAKVI (larotrectinib)

Covered Uses:	NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or better
Required Medical Information:	 Documentation of performance status, disease staging, all prior therapies used, and anticipated treatment course Documentation of positive neurotrophic tyrosine receptor kinase (NTRK) gene-fusion without a known acquired resistance mutation, as determined by an FDA approved test
Appropriate Treatment Regimen & Other Criteria:	Documentation of an intolerance to, or clinical rationale for avoidance of Rozlytrek (entrectinib) Reauthorization requires documentation of disease responsiveness to therapy
Exclusion Criteria:	Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **LAZERTINIB**

Affected Medications: LAZCLUZE (lazertinib)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or better
Required Medical Information:	Documentation of performance status, disease staging, all prior therapies used, and anticipated treatment course
	 Documentation of confirmed non-small cell lung cancer (NSCLC) that is metastatic or unresectable with epidermal growth factor receptor (EGFR) exon 19 deletions or exon 21 L858R substitution mutations
Appropriate	Documented intolerable adverse event to Tagrisso (osimertinib) with or without
Treatment	chemotherapy
Regimen & Other	
Criteria:	Reauthorization: documentation of disease responsiveness to therapy
Exclusion Criteria:	Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
Age Restriction:	18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 4 months, unless otherwise specified
_	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **LENACAPAVIR**

Affected Medications: SUNLENCA (lenacapavir)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of human immunodeficiency virus type 1 (HIV-1) infection, in combination with other antiretrovirals, in heavily treatment-experienced adults with multidrug resistant HIV-1 infection failing their current antiretroviral regimen due to resistance, intolerance, or safety considerations
Required Medical Information:	 Documentation of multidrug resistance within at least 3 of the 4 following antiretroviral classes (as defined by resistance to at least 2 agents within each of the 3 classes), unless contraindicated or clinically significant adverse effects are experienced: Nucleoside reverse-transcriptase inhibitors (NRTIs) Non-nucleoside reverse-transcriptase inhibitors (NNRTIs) Protease inhibitors (PIs) Integrase strand transfer inhibitors (INSTIs) Documentation of current (within the past 30 days) HIV-1 RNA viral load of at least 200 copies/mL
Appropriate Treatment Regimen & Other	Must be used in combination with an optimized background antiretroviral regimen that contains at least one agent demonstrating full viral susceptibility, as confirmed by resistance testing
Criteria:	 Reauthorization requires all of the following: Treatment plan includes continued use of optimized background antiretroviral regimen Documentation of treatment success, as evidenced by one of the following: Reduction in viral load from baseline or maintenance of undetectable viral load Absence of postbaseline emergence of lenacapavir resistance-associated mutations confirmed by resistance testing
Exclusion Criteria:	matations committed by resistance testing
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, an infectious disease or HIV specialist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Oral Tablet Initial Authorization: 1 month, unless otherwise specified
	Injection Initial Authorization: 6 months, unless otherwise specified
	Injection Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **LENIOLISIB**

Affected Medications: JOENJA (leniolisib)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Activated phosphoinositide 3-kinase delta syndrome (APDS)
Required Medical	Documentation of an APDS-associated PIK3CD/PIK3R1 mutation without concurrent
Information:	use of immunosuppressive medication
	Presence of at least one measurable nodal lesion on a CT or MRI scan
	Documentation of both of the following:
	 Nodal and/or extranodal lymphoproliferation
	 History of repeated oto-sino-pulmonary infections and/or organ dysfunction (e.g., lung, liver)
	Current weight (must be at least 45 kg)
Appropriate	Females of reproductive potential should have pregnancy ruled out and use effective
Treatment	contraception during therapy
Regimen & Other	
Criteria:	Reauthorization will require documentation of treatment success as shown by both of the
	following:
	Improvement in lymphoproliferation as measured by a change from baseline in
	 lymphadenopathy Normalization of immunophenotype as measured by the percentage of naïve B cells out
	of total B cells
Exclusion Criteria:	or total 2 dollo
Age Restriction:	12 to 75 years of age
Prescriber/Site of	Prescribed by, or in consultation with, an immunologist, hematologist/oncologist, or
Care Restrictions:	specialist with experience in the treatment of APDS
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 4 months, unless otherwise specified
-	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **LETERMOVIR**

Affected Medications: PREVYMIS (letermovir)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise excluded by plan design
	 Prophylaxis of cytomegalovirus (CMV) infection and disease in CMV-seropositive recipients [R+] of an allogeneic hematopoietic cell transplant for adults and pediatric patients 6 months of age and older and weighing at least 6 kg Prophylaxis of CMV disease in kidney transplant recipients at high risk for adult and pediatric patients 12 years of age and older and weighing at least 40 kg
Required Medical	Has received an allogeneic hematopoietic stem cell transplant (HSCT)
Information:	Is cytomegalovirus CMV-seropositive
	OR
	 Has received a kidney transplant and is at high risk (Donor CMV-seropositive/Recipient CMV-seronegative [D+/R-] of CMV infection
Appropriate	Documented trial and failure (or intolerable adverse event) with an adequate trial (at
Treatment	least 14 days) of at least one of the following: ganciclovir, valganciclovir, Foscarnet
Regimen & Other	(HSCT only)
Criteria:	
	HSCT Dosing : 480 mg (or 240 mg) once daily beginning between Day 0 and Day 28 post-transplantation and continued through Day 100 post-transplantation
	Kidney transplant Dosing : 480mg once daily beginning between Day 0 and Day 7 post kidney transplant for high-risk recipients (donor CMV-seropositive/recipient CMV-seronegative) and continue through day 200 post transplantation
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, an infectious disease provider or a specialist with
Care Restrictions:	experience in the prevention and treatment of CMV infection
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	HSCT Authorization: 4 months, unless otherwise specified
	 Kidney Transplant Authorization: 7 months, unless otherwise specified



LEUPROLIDE

Affected Medications: leuprolide acetate, LUPRON DEPOT, LUPRON DEPOT-PED, ELIGARD, LUPANETA (leuprolide-norethindrone), FENSOLVI, CAMCEVI

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
Required Medical	Endometriosis:
Information:	Documentation of moderate to severe pain due to endometriosis
	Uterine leiomyomata (fibroids):
	<u></u>
	Documentation of all of the following:
	 Preoperative anemia due to uterine leiomyomata (fibroids)
	 Planning to undergo leiomyomata-related surgery in the next 6 months or less Planning to use in combination with iron supplements
	Gender dysphoria:
	Documentation of all the following:
	 Current Tanner stage 2 or greater OR baseline and current estradiol and testosterone levels to confirm onset of puberty
	 Confirmed diagnosis of gender dysphoria that is persistent
	 The patient has the capacity to make a fully informed decision and to give consent for treatment
	 Any significant medical or mental health concerns are reasonably well controlled A comprehensive mental health evaluation has been completed by a licensed mental health professional (LMHP) and provided in accordance with the most current version of the World Professional Association for Transgender Health (WPATH) Standards of Care
	Central precocious puberty:
	Documentation of CPP confirmed by basal luteinizing hormone (LH), follicle-stimulating hormone (FSH), and either estradiol or testosterone concentrations
Appropriate Treatment	Endometriosis:
Regimen & Other Criteria:	 Documentation of a trial and inadequate relief (or contraindication) after at least 3 months of both of the following first-line therapies: Nonsteroidal anti-inflammatory drugs (NSAIDs) Continuous (no placebo pills) hormonal contraceptives



	Central precocious puberty:
	Approval of Fensolvi requires rationale for avoidance of Lupron and Supprelin LA
Exclusion Criteria:	Undiagnosed abnormal vaginal bleeding
	Management of uterine leiomyomata without intention of undergoing surgery.
	Pregnancy or breastfeeding
	Use for infertility (if benefit exclusion)
Age Restriction:	Endometriosis and preoperative uterine leiomyomata: 18 years of age and older
	Central precocious puberty (CPP): 11 years of age or younger (females), 12 years of age
	or younger (males)
Prescriber/Site of	Gender Dysphoria: Diagnosis made and prescribed by, or in consultation with, a
Care Restrictions:	specialist in the treatment of gender dysphoria
	All other indications: prescribed by, or in consultation with, an oncologist, endocrinologist,
	or gynecologist as appropriate for diagnosis
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Uterine leiomyomata: maximum of 6 months, unless otherwise specified
	Endometriosis: 6 months, unless otherwise specified
	All other diagnoses: 12 months, unless otherwise specified



LEVOKETOCONAZOLE

Affected Medications: RECORLEV (levoketoconazole)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Cushing syndrome
Required Medical	Diagnosis of Cushing's syndrome due to one of the following:
Information:	 Adrenocorticotropic hormone (ACTH)-secreting pituitary adenoma (Cushing's disease)
	 Ectopic ACTH secretion by a non-pituitary tumor
	 Cortisol secretion by an adrenal adenoma
	Mean 24-hour urine free cortisol (mUFC) greater than 1.5 times the upper limit of normal
	(ULN) for the assay (at least two measurements)
Appropriate	Documentation confirming surgery is not an option OR previous surgery has not been
Treatment	curative
Regimen & Other	Documentation of ONE of the following:
Criteria:	 Clinical failure to maximally tolerated dose of oral ketoconazole for at least 8 weeks
	 Intolerable adverse event to oral ketoconazole, and the adverse event was not
	an expected adverse event attributed to the active ingredient
	Reauthorization requires documentation of treatment success defined as mUFC normalization (i.e., less than or equal to the ULN)
Exclusion Criteria:	Adrenal or pituitary carcinoma
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, an endocrinologist, neurologist, or adrenal surgeon
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: LIFILEUCEL

Affected Medications: AMTAGVI (lifileucel)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	Diagnosis of unresectable or Stage IV metastatic melanoma
	NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or better
Required Medical	Documentation of performance status, disease staging, all prior therapies used, and
Information:	anticipated treatment course
	ECOG PS of 0 or 1
	Left ventricular ejection fraction (LVEF) greater than 45%
	Forced expiratory volume (FEV1) greater than 60%
	New York Heart Association (NYHA) classification not more than Class I
Appropriate	At least one resectable lesion (or aggregate of lesions resected) of 1.5 cm or more in
Treatment	diameter post-resection to generate tumor-infiltrating lymphocytes (TILs)
Regimen & Other	Disease progression after 1 or more prior systemic therapy including:
Criteria:	 a PD-1-blocking antibody; and
	 if BRAF V600 mutation–positive, a BRAF inhibitor or BRAF inhibitor plus a MEK inhibitor
Exclusion Criteria:	Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
	Melanoma of uveal or ocular origin
	Untreated or active brain metastasis
Age Restriction:	18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist.
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 6 months (one dose per patient's lifetime), unless otherwise specified



POLICY NAME: **LONAFARNIB**

Affected Medications: ZOKINVY (Ionafarnib)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design To reduce risk of mortality in Hutchinson-Gilford Progeria Syndrome For treatment of processing-deficient Progeroid Laminopathies
Required Medical Information:	A diagnosis of Hutchinson-Gilford Progeria Syndrome (HGPS) confirmed by mutational analysis (G608G mutation in the lamin A gene) OR
	 A diagnosis of processing-deficient Progeroid Laminopathies with one of the following: Heterozygous LMNA mutation with progerin-like protein accumulation Homozygous or compound heterozygous ZMPSTE24 mutations
Appropriate	Documented height and weight, or body surface area (BSA)
Treatment	Documentation of medication review and avoidance of drugs that significantly affect the
Regimen & Other	metabolism of Ionafarnib (e.g. strong or moderate CYP3A4 inhibitors/inducers)
Criteria:	Females of reproductive potential should have pregnancy ruled out and use effective contraception during treatment
	Labs:
	Absolute Phagocyte Count (sum of absolute neutrophil count, bands, and monocytes)
	greater than 1,000/microliters
	 Platelets greater than 75,000/microliters (transfusion independent) Hemoglobin greater than 9g/dl.
	Dosing:
	Available as oral capsules: 50 mg, 75 mg
	 Initial, 115 mg/m2/dose twice daily for 4 months, then increase to 150 mg/m2/dose twice daily
	 Do not exceed 115 mg/m2/dose twice daily when used in combination with a weak CYP3A4 inhibitor
	 Round all total daily doses to the nearest 25 mg increment
	Reauthorization requires documentation of treatment success and initial criteria to be met.
Exclusion Criteria:	Use for other progeroid syndromes or processing-proficient progeroid laminopathies
	Concomitant use with strong or moderate CYP3A4 inhibitors/inducers, midazolam,
	lovastatin, atorvastatin, or simvastatin
	Overt renal, hepatic, pulmonary disease or immune dysfunction
	BSA less than to 0.39 m2
Age Restriction:	Age 12 months or older with a BSA of greater than or equal to 0.39 m2
Prescriber/Site of	Prescribed by, or in consultation with, a provider with experience in treating progeria
Care Restrictions:	and/or progeroid laminopathies
Coverage Duration:	Initial Authorization: 4 months
	Reauthorization: 12 months
1	Troduction 12 months



POLICY NAME: **LOTILANER**

Affected Medications: XDEMVY

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Demodex blepharitis (DB)
Required Medical	Diagnosis of DB meeting both of the following criteria:
Information:	 Presence of erythema of the upper eyelid margin
	 Presence of mites upon examination of eyelashes by light microscopy OR
	presence of collarettes on slit lamp examination
	Documented trial and failure to oral ivermectin, 200 mcg/kg in a single dose and
	repeated at least once after 7 days
Appropriate	Reauthorization may be given at least 12 months after the first treatment and will require
Treatment	documentation of treatment success and returned presence of mites or collarettes requiring
Regimen & Other	retreatment
Criteria:	
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, an optometrist or ophthalmologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



LOVOTIBEGLOGENE AUTOTEMCEL

Affected Medications: LYFGENIA (lovotibeglogene autotemcel)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise excluded by
	plan design
	 Treatment of sickle cell disease in adults and pediatric patients at least 12 years
	of age with a history of recurrent vaso-occlusive crises
Required Medical	Documentation of sickle cell disease confirmed by genetic testing to show the presence
Information:	of $\beta S/\beta S$, $\beta S/\beta 0$ or $\beta S/\beta +$ genotype as follows:
	o Identification of significant quantities of HbS with or without an additional
	abnormal β-globin chain variant by hemoglobin assay
	OR
	o Identification of biallelic <i>HBB</i> pathogenic variants where at least one allele is the
	p.Glu6Val or p.Glu7Val pathogenic variant on molecular genetic testing AND
	 Patient does NOT have disease with more than two α-globin gene deletions
	Documentation of severe disease defined as 2 or more severe vaso-occlusive crises
	(VOCs) or vaso-occlusive events (VOEs) within the previous year (4 events over 2 years
	will also meet this requirement)
	VOC/VOEs defined as an event requiring a visit to a medical facility for
	evaluation AND necessitating subsequent interventions such as opioid pain
	management, non-steroidal anti-inflammatory drugs, red blood cell (RBC)
	transfusions, which results in a diagnosis of such being documented due to one
	(or more) of the following:
	 Acute pain event
	 Acute chest syndrome
	 Priapism lasting more than 2 hours
	 Acute splenic sequestration
	 Acute hepatic sequestration
	• For patients under 18 years of age, the patient does not have a known and suitable
	(10/10) human leukocyte antigen (HLA) matched related donor willing to participate in an
	allogeneic hematopoietic stem cell transplant (HSCT)
	Adequate bone marrow, lung, heart, and liver function to undergo myeloablative
	conditioning regimen
Annyanyiata	Confirmed HIV negative as confirmed by a negative HIV test prior to mobilization Abla to graphic the grid in
Appropriate	• Able to provide the minimum recommended dose of Lyfgenia- 3 x 10 ⁶ CD34+ cells/kg.
Treatment	
Regimen & Other	
Criteria:	
Exclusion Criteria:	Previous treatment with gene therapy for sickle cell disease
	Prior hematopoietic stem cell transplant (HSCT)
	History of hypersensitivity to dimethyl sulfoxide (DMSO) or dextran 40
Age Restriction:	12 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a hematologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
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Coverage Duration:	•	Authorization: 12 months (one-time infusion), unless otherwise specified



LUSPATERCEPT-AAMT

Affected Medications: REBLOZYL (luspatercept-aamt)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design o Treatment of anemia in adults with beta thalassemia who require regular red
	blood cell (RBC) transfusions
	Treatment of anemia in adults without previous erythropoiesis stimulating agent
	use (ESA-naïve) with very low- to intermediate-risk myelodysplastic syndromes
	(MDS) who may require regular RBC transfusions
	Treatment of anemia failing an ESA and requiring 2 or more RBC units over 8 weeks in adult national with your law to intermediate risk MDS with ring.
	weeks in adult patients with very low- to intermediate-risk MDS with ring sideroblasts (MDS-RS) or with myelodysplastic/myeloproliferative neoplasm with
	ring sideroblasts and thrombocytosis (MDS/MPN-RS-T)
Required Medical	Beta Thalassemia
Information:	Documented diagnosis of beta thalassemia OR hemoglobin E/beta thalassemia
	Documentation of transfusion dependence as evidenced by BOTH of the following in the
	previous 24 weeks:
	 Has required regular transfusions of at least 6 RBC units
	 No transfusion-free period greater than 35 days
	Pre-treatment or pre-transfusion hemoglobin (Hgb) level is less than or equal to 11 g/dL
	Myelodysplastic Syndromes
	Documented diagnosis of MDS, MDS-RS or MDS/MPN-RS-T with very low, low, or
	intermediate risk as classified by the International Prognostic Scoring System-Revised
	(IPSS-R)
	Documentation of requiring at least 2 RBC units over the previous 8 weeks
	Pre-treatment or pre-transfusion level is less than or equal to 11 g/dL
Appropriate	Myelodysplastic Syndromes
Treatment	For those with MDS-RS or MDS/MPN-RS-T, must have documentation of treatment
Regimen & Other	failure with an ESA (e.g., Retacrit, Procrit, Epogen, Mircera), unless intolerant or current
Criteria:	endogenous serum erythropoietin (sEPO) level is greater than 500 U/L
	Reauthorization
	Beta thalassemia: requires documentation of treatment success, defined as a reduction
	in RBC transfusion burden from baseline by at least 20%
	MDS: requires documentation of treatment success, defined as achieving transfusion
	independence and/or an improvement in Hgb level from baseline
Exclusion Criteria:	Diagnosis of non-transfusion-dependent beta thalassemia
	Use as immediate correction as a substitute for RBC transfusions
	Diagnosis of alpha thalassemia
Age Restriction:	 Known pregnancy 18 years of age and older
Age Nestriction.	18 years of age and older
L	



Prescriber/Site of Care Restrictions:	 Beta thalassemia: Prescribed by, or in consultation with, a hematologist MDS: Prescribed by, or in consultation with, a hematologist or oncologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 3 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: LUSUTROMBOPAG

Affected Medications: MULPLETA (lusutrombopag)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Thrombocytopenia in adult patients with chronic liver disease who are scheduled to undergo a procedure
Required Medical Information:	Documentation of ALL the following:
Appropriate Treatment Regimen & Other Criteria:	Approved for one time 7-day dosing regimen
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of Care Restrictions:	Prescribed by, or in consultation with, a hematologist or gastroenterology/liver specialist
Coverage Duration:	Authorization: 1 month (7 days of treatment), based on planned procedure date, unless otherwise specified



POLICY NAME: MARIBAVIR

Affected Medications: LIVTENCITY (maribavir)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	 Treatment of adults and pediatric patients (12 years of age and older and
	weighing at least 35 kg) with post-transplant cytomegalovirus (CMV)
	infection/disease that is refractory to treatment (with or without genotypic
	resistance) with ganciclovir, valganciclovir, cidofovir or foscarnet
Required Medical	Documentation of post-transplant CMV infection
Information:	Documentation of patient's current weight
Appropriate	Documented clinical failure (not due to drug intolerance) with an adequate trial (at least
Treatment	14 days) of at least one of the following: ganciclovir, valganciclovir, cidofovir or foscarnet
Regimen & Other	
Criteria:	Reauthorization:
	 Documented treatment success and a clinically significant response to therapy and continued need for treatment.
Exclusion Criteria:	CMV infection involving the central nervous system, including the retina.
Age Restriction:	12 years and older
Prescriber/Site of	Prescribed by an infectious disease provider or a specialist with experience in the
Care Restrictions:	treatment of CMV infection
Coverage Duration:	Authorization: 4 months, unless otherwise specified



POLICY NAME: MARSTACIMAB

Affected Medications: HYMPAVZI (marstacimab hncq)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	Hemophilia A (congenital factor VIII deficiency)
	 Hemophilia B (congenital factory IX deficiency)
Required Medical Information:	Diagnosis of congenital factor VIII deficiency (hemophilia A) or congenital factory IX deficiency (hemophilia B) without inhibitors
	Documentation of baseline factor level less than 1% AND prophylaxis required OR
	Baseline factor level 1% to 3% and a documented history of at least two episodes of spontaneous bleeding into joints
	 Prescribed for routine prophylaxis to prevent or reduce the frequency of bleeding episodes
Appropriate	Hemophilia A
Treatment	Documented treatment failure with Hemlibra (emicizumab-kxwh)
Regimen & Other	
Criteria:	Hemophilia B
	Documented treatment failure to factor IX prophylaxis for at least 6 months
	Dose escalation to 300 mg once weekly:
	Documentation of weighing at least 50 kg and experiencing at least 2 breakthrough
	bleeds while on 150 mg dose for at least 6 months
	Reauthorization requires documentation of treatment success defined as a reduction in spontaneous bleeds requiring treatment, and documentation of bleed history since last approval
Exclusion Criteria:	Concurrent use with bypassing agents
	Prior gene therapy administration
	Pregnancy
Age Restriction:	12 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a hematologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: MAVACAMTEN

Affected Medications: CAMZYOS (mavacamten)

All Food and Drug Administration (FDA)-approved indications not otherwise excluplan design. Hypertrophic cardiomyopathy with left ventricular outflow tract obstruction Required Medical Documented diagnosis of obstructive hypertrophic cardiomyopathy (OHCM)	-
 Hypertrophic cardiomyopathy with left ventricular outflow tract obstruction 	
Popular Modical - Decumented disapposis of shotrustive by partraphic cordinary enoting (OHCM)	
Required Medical • Documented diagnosis of obstructive hypertrophic cardiomyopathy (OHCM)	
Information: • New York Heart Association (NYHA) class II or III symptoms	
 Left ventricular ejection fraction (LVEF) of 55% or greater prior to starting therapy 	
 Valsalva left ventricular outflow tract (LVOT) peak gradient of 50 mmHg or greater or with provocation, prior to starting therapy 	at rest
Appropriate • Documentation of negative pregnancy test AND use of effective contraception in f	emales
Treatment of reproductive potential	
Regimen & Other • Documented treatment failure, intolerance, or contraindication, to ALL of the follow	ving:
Criteria: Non-vasodilating beta-blocker (e.g., atenolol, metoprolol, bisoprolol, propromoder (e.g., verapamil, diltiazem)	anolol)
Reauthorization will require documentation of symptomatic improvement and that LV remains above 50%	EF
Exclusion Criteria: • History of two measurements of LVEF less than 50% while on mavacamten 2.5 m tablets	g
Age Restriction:	
Prescriber/Site of Prescribed by, or in consultation with, a cardiologist or a specialist with experience	in the
Care Restrictions: treatment of obstructive hypertrophic cardiomyopathy	
All approvals are subject to utilization of the most cost-effective site of care	
Coverage Duration: • Initial Authorization: 6 months, unless otherwise specified	
Reauthorization: 12 months, unless otherwise specified	



POLICY NAME: **MAVORIXAFOR**

Affected Medications: XOLREMDI (mavorixafor)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Treatment of WHIM syndrome (warts, hypogammaglobulinemia, infections and myelokathexis) in patients 12 years of age and older to increase the number of circulating mature neutrophils and lymphocytes.
Required Medical Information:	 Diagnosis of WHIM syndrome confirmed by genotype variant of CXCR4 and ANC (absolute neutrophil count) of 400 cells/µL or less
	 Documentation of symptoms and complications associated with WHIM syndrome requiring medical treatment
Appropriate	Documentation of weight to assess appropriate dosing
Treatment	Documentation of baseline ALC (absolute lymphocyte count) and ANC (absolute)
Regimen & Other	neutrophil count) to assess clinical response to treatment
Criteria:	
	<u>Reauthorization</u> requires documentation of disease responsiveness to therapy with sustained improvement in ALC and ANC
Exclusion Criteria:	Concomitant use with drugs that are highly dependent on CYP2D6 for clearance.
Age Restriction:	12 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, an immunologist or hematologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
-	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **MECASERMIN**

Affected Medications: INCRELEX (mecasermin)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Severe primary insulin-like growth factor-1 (IGF-1) deficiency (Primary IGFD)
	 Patient with growth hormone (GH) gene deletion with neutralizing antibodies to GH
Required Medical	Prior to starting therapy, a height at least 3 standard deviations below the mean for
Information:	chronological age and sex, and an IGF-1 level at least 3 standard deviations below the mean for chronological age and sex.
	One stimulation test showing patient has a normal or elevated GH level
Appropriate	Initial: 0.04-0.08 mg/kg subcutaneously twice daily.
Treatment	Maintenance: Up to 0.12 mg/kg subcutaneously twice daily.
Regimen & Other	
Criteria:	• Reauthorization: requires a documented growth rate increase of at least 2.5 cm over baseline per year AND evaluation of epiphyses (growth plates) documenting they remain open.
Exclusion Criteria:	Epiphyseal closure, active or suspected neoplasia malignancy, or concurrent use with GH therapy.
	Patient has secondary causes of IGF1 deficiency (e.g., hypothyroidism, malignancy, chronic systemic disease, skeletal disorders, malnutrition, celiac disease)
Age Restriction:	For patients 2 to 18 years of age.
Prescriber/Site of	Prescribed by, or in consultation with, a pediatric endocrinologist
Care Restrictions:	All approvals are subjects to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



MEDICAL NECESSITY

Affected Medications: Abilify MyCitea, Abrilada, Absorica, Absorica LD, Acanya, Aciphex, Actemra SQ, Acthar Gel, Acuvail, Acyclovix, Aczone, Adalimumab-adbm, Adalimumab-fkjp, Adalimumab-ryvk, Adapalene pads, Adcirca, Adlarity, Adlyxin, Admelog, Advicor, Adzenys ER, Adzenys XR, Aerospan, Afrezza, Aimovig, AirDuo, AirDuo Digihaler, Airsupra, Ajovy, Aklief, Allopurinol 200 mg tablet, Allzital, Alprazolam Dispersible, Alprazolam Intensol, Altoprev, Alvesco, Ameluz, Amitiza, Amjevita, Amphetamine ER suspension, Ampyra, Amrix, Amturnide, Amzeeq, Ancobon, Androgel, Androxy, Apadaz, APAP-Caff-Dihydrocodeine, Apidra, Aplenzin, Arazlo, Aripiprazole Dispersible, Armonair Digihaler, Armonair Respiclick, Arymo ER, Asacol HD, Asmanex, Asmanex HFA, Aspruzyo, Astepro solution, Atorvaliq, Aubagio, Auvelity, Aveed, Azathioprine tablet (75 mg, 100 mg), Azelex, Azesco, Azmiro, Azstarys, Baclofen Oral Suspension, Basaglar, Basaglar Tempo pen, Baxdela, Beconase, Belbuca, Beser, Bevespi Aerophere, Bexagliflozin, BiDil, Biifenac, Bimzelx, Bismuth Subcitrate-Metronidazole-Tetracycline, Brenzavvy, Breztri, Bridion, Brisdelle, Briviact, Bryhali, Budesonide 9 mg ER tablet, Bunavail, Bupap, Buphenyl, Bupropion XL 450 mg, Butisol, Butrans patch, Bydureon, Bydureon BCise, Byetta, Bynfezia, Byvalson, Cabtreo, Calcipotriene-Betamethasone Dipropionate suspension, Cambia, Capex shampoo, Capital-Codeine, Carac, Carbinoxamine 6 mg tablet, Carisoprodol-ASA, Carisoprodol-ASA-Codeine, CaroSpir, Carticel implant, Cataflam, Cephalexin 750 mg capsule, Cephalexin tablet, Cequa, Chlorpheniramine-Codeine, Chlorzoxazone 250 mg tablet, Cibingo, Ciloxan, Cimzia, Ciprodex OTIC, Cipro HC Otic, Clemastine syrup, Clindamycin Phosphate-Benzoyl Peroxide gel 1.2-2.5 %, Clindavix, Clobetasol ophthalmic suspension, Clobetex, Clonidine ER 0.17 mg tablet, Codar AR, Colazal, Conjupri, Consensi, Conzip, Copaxone, Coreg CR, Cosopt PF, Cotempla XR ODT, Coxanto, Crexont, Crinone, Cuprimine, Cuyposa, Cyanocobalamin Nasal Spray, Cyclobenzaprine ER, Cyclosporine in Klarity, Cyltezo, Dapagliflozin, Dapagliflozin-Metformin ER, Dartisla ODT, Debacterol, Degludec, Delzicol, Demser, Depen, DermacinRx Lexitral cream pack, Dermalid, Desonate gel, Desonide gel, Desonide lotion, DesRx gel, Dexilant, Dhivy, Dichlorphenamide, Diclofenac 1.3 % patch, Diclofenac Potassium capsule, Diclofenac Potassium packet, Diclofenac Potassium 25 MG tablet, Diclofenac Sod soln 1.5 % & Capsaicin cream 0.025 % ther pack, Diclofex DC cream, Diclopak, Diclosaicin cream, Diclotral pack, Diclotrex, Diclovix DM pak, Diflorasone Diacetate, Dipentum, Dorvx MPC, Doxepin 5 % cream, Doxycyline Hyclate 50 mg tablet, Doxycycline Hyclate DR tablet (50 mg, 80 mg, 200 mg), Doxycycline Monohydrate DR 40 mg capsule, Duaklir Pressair, Duetact, Duexis, Dulera, Duobrii, Durlaza, Dutoprol, Duzallo, Dxevo, Dyanavel XR, Dymista, Dynabec, Ebglyss, Econasil, Edarbi, Edarbyclor, Egaten, Egrifta, Elepsia XR, Elidel, Elyxyb, Emend, Emflaza, Emflaza Suspension, Emrosi, Enalapril oral solution, Enstilar foam, Entadfi, Entyvio SQ, Eohilia, Epaned, Epanova, Epclusa, Eprontia, Equetro, Ergomar, Esbriet, Eskata, Evzio, Exjade, Exservan, Extavia, Extina foam 2 %, Fabior foam, Faslodex, Fenofibrate 120 mg, Fenortho, Firazyr, First-lansoprazole, Flector patch, Flegsuvy, Flolipid, Flowtuss, Fluopar kit, Fluorouracil 0.5 % cream, Flurandrenolide, Fluoxetine (PMDD) tablet, Forfivo XL, Fortamet, Fortesta gel, Fosamax Plus D, Fulyzag, Furoscix, Gabacaine pak, Gabapal, Giazo, Gilenya, Gimoti, Gleevec, Gloperba, Glumetza, Glycate, Glycopyrrolate 1.5 mg tablet, Gocovri, Gonitro, GPL pak, Halog, Halcinonide cream, Harvoni, Harvoni pak, Helidac, Hemady, Hemangeol, Hetlioz capsule, Hulio, Humalog, Humalog Junior KwikPen, Humatin, Humira, Humulin, Humulin, 70/30 KwikPen, Humulin N, Humulin R-100, Hycofenix, Hyrimoz (Sandoz), Ibrance, Ibsrela, Ibuprofen-Famotidine, Idacio, Igalmi, Iheezo, Ilumya, Imbruvica 70 mg capsule, Imbruvica 140 mg & 280 mg tablet, Imiquimod 3.75 %, Imkeldi, Impeklo, Impoyz, Imvexxy, Inbrija, Inderal LA, Indocin suppository, Indomethacin 20 mg capsule, Inflatherm kit, Inflatherm pak, Infugem, Ingrezza, Ingrezza Sprinkle, Innolet Insulin, Inpefa, Insulin Aspart, Insulin Aspart Protamine & Aspart 70/30, Insulin Degludec, Insulin Glargine, Insulin Glargine-yfgn, Insulin Lispro, Intrarosa, Invega ER, Invokamet, Invokamet XR, Invokana, Isordil Titradose, Isosorbide Dinitrate-Hydralazine, Isotretinoin 25 mg and 35 mg capsule, Iyuzeh, Jadenu, Jadenu sprinkle packet, Jentadueto, Jentadueto XR, Jublia, Jylamvo, Karbinal ER, Katerzia, Kazano, K-bicarb, Kenalog aerosol, Kenalog susp, Keragel, KeragelT, Kerydin, Kesimpta, Ketek, Ketorolac nasal spray, Keveyis, Kevzara, Kineret, Klisyri, Kombiglyze XR, Konvomep, Korlym, Kyzatrex, Lampit, Latuda, Lescol XL, Letairis, Levamlodipine, Levorphanol Tartrate, Lexette, Lexuss, Lialda, Libervant, Licart, Lido GB 300 kit, Lidostream, Lidotin Pak, Lifems, Likmez, Lipritin Pak, Liptruzet, Lithostat, LMR Plus Lidocaine, Lodoco, Lofena, Lonhala Magnair, Loreev XR, Lucemyra, Luzu, Lybalvi, Lyrica, Lyrica CR tablet, Lyumjev, Lyumjev Kwikpen, Lyvispah, Meclofen, Meloxicam capsule, Mentax cream 1 %, Mesalamine DR 800 mg tablet, Metaclopramide disintegrating tablet, Metaxall, Metaxall CP, Metformin ER (OSM), Metformin solution, Methadone Intensol, MethylTESTOSTERone capsule, Metyrosine, Miebo, Mifepristone, Migraine pack, Minocycline ER, Minolira, Mitigare, Monocycline ER, MorphaBond, MorphaBond ER, Motegrity, Motofen, Motpoly XR, Mycapssa, Myfembree, Myhibbin, Myrbetriq, Mytesi, Nalocet, Namenda XR, Namzaric, Naprelan, Naproxen-Esomeprazole, Nascobal, Natesto



gel, Neo-Synalar cream, Nesina, Nexiclon XR, Nexletol, Nexlizet, Nitisinone, Nocdurna, Noctiva, Nolix, Nopioid TC kit, Norgesic Forte, Noritate, Norligva, Noroxin, Northera, Nourianz, Novolin 70/30 Relion, Novolin N Relion, Novolin R Relion, Noxafil, NuDiclo Solupak, Nurtec, Nuvakaan kit, Nuvakaan II kit, Nuvigil, Nuzyra, Ofloxacin tablet, Ohtuvayre, Olpruva, Olumiant, Olysio, Omeprazole-Sodium Bicarb, Omnaris, Omvoh SQ, Ondansetron 24 mg tablet, Onexton, Onfi, Onglyza, Onmel, Onyda XR, Onzetra Xsail, Opipza, Oracea, Oralair, Orencia SQ, Ormalvi, Orphenadrine-Aspirin-Caffeine tablet, Orphengesic Forte, Ortikos, Oseni, Otrexup, Oxaprozin capsule, Oxaydo, Oxycodone-Acetaminophen (2.5 mg-300 mg, 5 mg-300 mg, 7.5 mg-300 mg, 10 mg-300 mg), Ozobax, Pamelor, Panlor, Panretin gel, Paromomycin, Pazeo, Pedizolpak, Penicillamine tablet, Pennsaid solution, Pentican pak, Percocet, Pertzye, Pheburane, Picato, Pioglitazone-Glimepiride, Pirfenidone 534 mg tablet, Plaquenil, Pradaxa, Praluent, Prevacid SoluTab, Prevpac, Prialt, Prilo Patch, Prilopentin, Primlev, Primsol, Pristiq, ProAir Digihaler, Prolate, Prudoxin, Purified Cortrophin gel, Purixan, Qbrelis, Qbrexza, Qdolo, Qelbree, Qmiiz, QNASL, Qtern, Qudexy XR, QuilliChew ER, Quillivant XR, Quinixil, Quinosone, Qulipta, Qwo, Ranexa, Rasuvo, Rayos, Recarbrio, Reditrex, Relexxii, Relion Insulins, Relprevy, Reltone, Retin-A Micro pump gel (0.06 %, 0.08 %), Revatio, Rezvoglar, Rhofade, Ribasphere, Ridaura, Riomet, Riomet ER, Rocklatan, Ryaltris, Ryvent, Ryzodeg 70/30, Sabril, Samsca, Saphris, Sarafem, Savaysa, Saxagliptin-Metformin ER, Seconal, Seebri Neohaler, Seglentis, Segluromet, Semglee, Sensipar, Sernivo, Seysara, Siklos, Silenor, Sila III pak, Siliq subcutaneous injection, Simlandi, Simponi, Simvastatin suspension, Skelaxin, Skelid, Soaanz, Sofdra, Soliqua, Solodyn, Solosec, Soolantra, Sorilux, Sotyktu, Sovaldi, Sovaldi pak, Spevigo Subcutaneous, Spironolactone suspension, Sporanox solution, Spritam, Sprix, Sprycel, Steglatro, Steglujan, Stegeyma, Striant, Striant buccal, Suboxone, Sumatriptan-Naproxen, Sure Result DSS premium pack, Symbyax, Sympazan, Symproic, Synalar, Syndros, Syprine, Taclonex suspension, Talicia, Taltz, Tanzeum, Targadox, Tascenso ODT, Tasoprol, Tavaborole, Tazarotene foam, Tazarotene cream 0.05%, Tazorac Cream, Tazorac Gel, Tecfidera, Technivie, Thalitone, Thiola, Thiola EC, Thyquidity, Ticlopidine, Tiglutik, Tiopronin, Tivorbex, Tolak, Tolsura, Topiramate ER, Tosymra, Tovet kit, Tracleer, Tradjenta, Tramadol oral solution, Tretinoin Microsphere Gel 0.08 %, Treximet, Tri-Luma, Trixylitral kit, Trokendi XR, Trudhesa, Trulance, Tudorza Pressair, Twyneo, Tyrvaya, Tyzeka, Tyzine, Ubrelvy, Ultravate, Ultresa, Uptravi, Ursodiol capsule (200 mg, 400 mg), Utibron Neohaler, Uzedy, Valsartan oral solution, Vanatol LQ, Vanos, Varophen, Vasotec, Vecamyl, Vectical, Velsipity, Veltassa, Venlafaxine Besylate ER, Veozah, Veramyst, Veregen, Verkazia, Versacloz, Vesicare LS, Vevye, Vexasyn, Vexasyn gel, Vfend oral suspension, V-Go, Viberzi, Vibramycin, Victoza, Victrelis, Viekira, Vigafyde, Viibryd, Viibryd Starter Pack, Vimovo, Viokace, Vivlodex, Vogelxo, Voquezna dual pak, Voriconazole oral suspension, Vtol LQ solution, Vyzulta, Wakix, Wegovy, Wezlana, Winlevi, Wynorza, Xaciato, Xadago, Xartemis XR, Xatmep, Xcopri, Xelitral pack, Xeloda, Xelstrym, Xenazine, Xenleta, Xerese, Xermelo, Xhance, Ximino, Xtampza ER, Xultophy, Xyosted, Yosprala, Yuflyma, Yupelri, Yusimry, Zanaflex capsule, Zavzpret, Zcort, Zebutal, Zecuity, Zelnorm, Zembrace, Zenevix, Zepatier, Zetonna, Zileuton ER, Zinbryta, Zipsor, Zituvimet, Zituvimet XR, Zituvio, Zolpak, Zolpidem capsule, Zolpimist, Zonalon, Zonisade, Zorvolex, ZTLido, Z-Tuss, Zyclara, Zymfentra, Zypitamag, Zytiga

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
Required Medical Information:	Documented intolerance or treatment failure with the formulary alternatives for the submitted diagnosis
Appropriate Treatment Regimen & Other Criteria:	Food and Drug Administration (FDA)-approved compendia supported dosing
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of Care Restrictions:	All approvals are subjects to utilization of the most cost-effective site of care



Coverage Duration:	•	Dependent on expected duration of therapy and necessity of documentation of response to therapy



POLICY NAME: **MEPOLIZUMAB**

Affected Medications: NUCALA (mepolizumab)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Add-on maintenance treatment of patients with severe asthma aged 6 years and older with an eosinophilic phenotype
	 Treatment of adult patients with eosinophilic granulomatosis with polyangiitis
	(EGPA)
	 Treatment of patients aged 12 years and older with hypereosinophilic syndrome (HES)
	 Add-on maintenance treatment of chronic rhinosinusitis with nasal polyps (CRSwNP) in adult patients 18 years of age and older with inadequate response to nasal corticosteroids (NCS)
Required Medical	Eosinophilic asthma
Information:	Diagnosis of severe asthma with an eosinophilic phenotype, defined by both of the
	following:
	 Baseline eosinophil count of at least 150 cells/µL OR dependent on daily oral
	corticosteroids
	AND
	 FEV1 less than 80% at baseline or FEV1/FVC reduced by at least 5% from normal
	<u>EGPA</u>
	Diagnosis of relapsing or refractory EGPA confirmed by all of the following:
	 Chronic rhinosinusitis
	o Asthma
	 Blood eosinophilia (at least 1,500 cells/mcL and/or 10% eosinophils on differential) at baseline
	 Diagnosis must be confirmed by a second clinical opinion
	Documented relapsing disease while on the highest tolerated oral corticosteroid dose
	HES
	Diagnosis of HES with all of the following: Plead equipophil count greater than or equal to 1 000 cells/mall.
	 Blood eosinophil count greater than or equal to 1,000 cells/mcL Disease duration greater than 6 months
	At least Offices within the enact 40 months
	 At least 2 flares within the past 12 months Lab work showing Fip1-like1-platelet-derived growth factor receptor alpha
	(FIP1L1-PDGFRα) mutation negative disease
	Non-hematologic secondary HES (e.g., drug hypersensitivity, parasitic helminth)
	infection, HIV infection, non-hematologic malignancy) has been ruled out
	Documentation that disease is currently controlled on the highest tolerated glucocorticoid
	dose (defined as an improvement in clinical symptoms and a decrease in eosinophil count by at least 50% from baseline)



	CRSwNP
	Documentation of both of the following:
	 Diagnosis of chronic rhinosinusitis and has undergone prior bilateral total
	ethmoidectomy
	 Indicated for revision sinus endoscopic sinus surgery due to recurrent symptoms of nasal polyps (such as nasal obstruction/congestion, bilateral sinus
	obstruction)
Appropriate	Eosinophilic asthma
Treatment	Documented use of high-dose inhaled corticosteroid (ICS) plus a long-acting beta
Regimen & Other	agonist (LABA) for at least three months with continued symptoms
Criteria:	Documentation of one of the following:
	 Documented history of 2 or more asthma exacerbations requiring oral or systemic corticosteroid treatment in the past 12 months while on combination inhaler treatment with at least 80% adherence
	 Documentation that chronic daily oral corticosteroids are required
	<u>EGPA</u>
	Documented treatment failure or contraindication to at least two oral immunosuppressant
	drugs (azathioprine, methotrexate, mycophenolate) for at least 12 weeks each
	HEC
	HES - Decumented treatment failure or contraindication to at least 12 weeks of hydroxyuras
	Documented treatment failure or contraindication to at least 12 weeks of hydroxyurea (not required if notions has a hymphonytic verient of HES II, HES))
	(not required if patient has a lymphocytic variant of HES [L-HES])
	Documented treatment failure with interferon alfa
	CRSwNP
	Documented treatment failure with at least 1 intranasal corticosteroid (such as
	fluticasone) after ethmoidectomy
	Documented treatment failure with Sinuva implant
	Bocumented treatment failure with Sindva implant
	Reauthorization: documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	Use in combination with another monoclonal antibody (e.g., Dupixent, Fasenra, Xolair, Cinqair, Tezspire)
Age Restriction:	Eosinophilic asthma: 6 years of age and older
	EGPA: 18 years of age and older
	HES: 12 years of age and older
	CRSwNP: 18 years of age and older
Prescriber/Site of	Eosinophilic asthma: prescribed by, or in consultation with, an allergist, immunologist,
Care Restrictions:	or pulmonologist
	EGPA: prescribed by, or in consultation with, a specialist in the treatment of EGPA (such
	as an immunologist or rheumatologist)
	HES: prescribed by, or in consultation with, a specialist in the treatment of HES (such as
	an immunologist or hematologist)
	CRSwNP: prescribed by, or in consultation with, an otolaryngologist



	•	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	•	Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



METHYLNALTREXONE

Affected Medications: RELISTOR (methylnaltrexone bromide)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Opioid-induced constipation in adult patients with advanced illness or pain caused by active cancer who require opioid dosage escalation for palliative care Opioid-induced constipation in adult patients with chronic non-cancer pain, including patients with chronic pain related to prior cancer or its treatment who do not require frequent (e.g., weekly) opioid dosage escalation
Required Medical Information:	Documentation of treatment of opioid-induced constipation (OIC) in an adult with:
Appropriate Treatment Regimen & Other Criteria:	OIC in adults with chronic non-cancer pain Documented treatment failure or contraindication to a trial of all of the following: Lubiprostone Linzess Movantik
	Reauthorization will require documentation of treatment success, a clinically significant response to therapy, and documentation of continued opioid use
Exclusion Criteria:	 Known or suspected mechanical gastrointestinal obstruction or increased risk for recurrent obstruction
Age Restriction:	
Prescriber/Site of Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **METRELEPTIN**

Affected Medications: MYALEPT (metreleptin)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Congenital or acquired generalized lipodystrophy as a result of leptin deficiency
Required Medical Information:	 Current weight Baseline serum leptin levels, hemoglobin A1c (HbA1c), fasting glucose, fasting triglycerides, fasting serum insulin Prior Myalept use will require testing for anti-metrepeptin antibodies Documented leptin deficiency confirmed by laboratory testing (serum leptin of less than 12 ng/mL) Documentation of congenital or acquired generalized lipodystrophy with least ONE of the following: Concurrent hypertriglyceridemia Concurrent diabetes
Appropriate Treatment Regimen & Other Criteria:	 Generalized lipodystrophy with concurrent hypertriglyceridemia Triglycerides of 500 mg/dL or higher despite optimized therapy with at least two triglyceride-lowering agents from different classes (e.g., fibrates, statins) at maximum tolerated doses for at least 12 weeks each Generalized lipodystrophy with concurrent diabetes Persistent hyperglycemia (HbA1c 7 percent or greater) despite dietary intervention and optimized insulin therapy at maximally tolerated doses for at least 12 weeks
	Reauthorization will require documentation of treatment success and a clinically significant response to therapy documented by increased metabolic control defined by improvement in HbA1c, fasting glucose, and fasting triglyceride levels
Exclusion Criteria:	 Partial lipodystrophy General obesity not associated with leptin deficiency HIV-related lipodystrophy Metabolic disease, including diabetes mellitus and hypertriglyceridemia, without concurrent documentation of generalized lipodystrophy
Age Restriction:	de la company de
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an endocrinologist All approvals are subjects to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



MIACALCIN

Affected Medications: MIACALCIN injection (calcitonin-salmon)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	o Paget's disease of bone
	o Hypercalcemia
Required	<u>Hypercalcemia</u>
Medical	Documented calcium level greater than or equal to 14 mg/dL (3.5 mmol/L)
Information:	
	Paget's disease of bone
	Documented baseline radiographic findings of osteolytic bone lesions
	Abnormal liver function test (LFT), including alkaline phosphatase
	Documented lack of malignancy within the past 3 months
Appropriate	<u>Hypercalcemia</u>
Treatment	Documentation that additional methods for lowering calcium (such as intravenous fluids)
Regimen &	did not result in adequate efficacy OR
Other Criteria:	Clinical judgement necessitated immediate administration without waiting for other
	methods to show efficacy
	Paget's disease of bone
	 Documented trial and failure (or intolerable adverse event) with an adequate trial of both
	of the following:
	Zoledronic acid (at least one dose)
	o Oral bisphosphonate (e.g., alendronate, risedronate) for at least 8 weeks
	OR
	Documentation that the patient has severe renal impairment (e.g., creatinine clearance)
	less than 35 mL/min)
	AND
	Documentation of all of the following:
	 Normal vitamin D and calcium levels and/or supplementation
	 Symptoms that necessitate treatment with medication (e.g., bone pain, bone
	deformity)
	Poputherization Paget's disease of honor
	 Reauthorization - Paget's disease of bone: Documentation of treatment success and a clinically significant response to therapy
	Documentation of treatment success and a clinically significant response to therapy (such as stable or lowered alkaline phosphatase level, resolution of bone pain or other
	symptoms)
Exclusion	Related to Paget's disease of bone
Criteria:	History of a skeletal malignancy or bone metastases
- Cinonai	 Concurrent use of zoledronic acid or oral bisphosphonates
	Asymptomatic Paget's Disease of the bone
	Treatment or prevention of osteoporosis
Age	18 years of age or older - for Paget's disease of bone only
Restriction:	
Prescriber/Site of	All approvals are subject to utilization of the most cost-effective site of care
Care Restrictions:	



Coverage Duration:	•	Authorization: 12 months, unless otherwise specified



POLICY NAME: MIGLUSTAT

Affected Medications: MIGLUSTAT

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of adult patients with mild to moderate type 1 Gaucher disease Compendia-supported uses that will be covered: Niemann-Pick disease type C (NPC)
Required Medical Information:	Gaucher Disease Diagnosis of Gaucher disease confirmed by ONE of the following:
	 NPC Diagnosis of NPC confirmed by genetic testing showing biallelic pathogenic variants in either the NPC1 gene or NPC2 gene Documentation of at least one neurological symptom of Niemann-Pick disease type C, such as: Loss of motor function Problems with swallowing or speech Cognitive impairment Documentation of being ambulatory without needing an assistive device such as a wheelchair, walker, or cane Documentation of baseline signs and symptoms of NPC
Appropriate Treatment Regimen & Other Criteria:	Gaucher Disease: Reauthorization will require documentation of treatment success and a clinically significant response to therapy NPC:
	 Reauthorization requires: Documentation of treatment success defined as stability or improvement of Niemann-Pick disease type C signs and symptoms Documentation that patient is still ambulatory
Exclusion Criteria:	Female of childbearing potential who is pregnant or planning a pregnancy
Age Restriction:	



Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, one of the following: A specialist in the management of Gaucher disease (hematologist, oncologist, hepatologist, geneticist or orthopedic specialist) A specialist in the management of NPC (such as a geneticist, endocrinologist, metabolic disorder subspecialist, or neurologist) All approvals are subject to utilization of the most cost-effective site of care
Coverage	Initial Authorization: 4 months, unless otherwise specified
Duration:	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **MILTEFOSINE**

Affected Medications: IMPAVIDO (miltefosine)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of the following in adults and pediatric patients 12 years of age and older weighing greater than or equal to 30 kg (66 lbs):
Required Medical	All Indications
Information:	Current weight
	 Visceral Leishmaniasis Documentation of diagnosis confirmed by smear or culture in tissue (usually bone marrow or spleen) Cutaneous and Mucosal Leishmaniasis Documentation of diagnosis confirmed by histology, culture, or molecular analysis via
	polymerase chain reaction (PCR)
Appropriate	Dosing:
Treatment	30 to 44 kg: 50 mg twice daily for 28 days
Regimen & Other Criteria:	45 kg or greater: 50 mg three times daily for 28 days
Exclusion Criteria:	 Pregnancy Sjögren-Larsson syndrome Weight less than 30 kg (66 lbs) Treatment of leishmaniasis outside of the visceral, cutaneous, or mucosal settings Treatment of other <i>Leishmania</i> species
Age Restriction:	12 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, an infectious disease specialist
Care Restrictions:	All approvals are subjects to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 1 month, unless otherwise specified
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POLICY NAME: MITAPIVAT

Affected Medications: PYRUKYND (mitapivat tablet)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	Hemolytic anemia due to pyruvate kinase deficiency (PKD)
Required Medical Information:	 Documented diagnosis of pyruvate kinase deficiency (PKD), confirmed by BOTH of the following: Presence of at least 2 variant alleles in the pyruvate kinase liver and red blood cell (PLKR) gene At least one variant allele is a missense mutation Documentation of ONE of the following: Regularly receiving red blood cell (RBC) transfusions, defined as 6 or more transfusions in the previous 12 months Baseline hemoglobin (Hb) level of less than or equal to 10 g/dL with a history of no more than 4 transfusions in the previous 12 months Documentation of baseline transfusion count, including dates and number of units transfused
Appropriate Treatment Regimen & Other Criteria:	 Reauthorization requires documentation of treatment success and a clinically significant response to therapy, defined as: For patients receiving regular transfusions at baseline: must document greater than or equal to a 33% reduction in RBC units transfused compared to baseline For patients not receiving regular transfusions at baseline: must document greater than or equal to a 1.5 g/dL increase in Hb from baseline sustained at 2 or more scheduled visits AND no transfusions were needed
Exclusion Criteria:	 Splenectomy scheduled during treatment or have undergone within the 12-month period prior to starting treatment Previous bone marrow or stem cell transplant
Age Restriction:	Must be 18 years or older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a hematologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



MOMETASONE SINUS IMPLANT

Affected Medications: SINUVA (mometasone sinus implant)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of chronic rhinosinusitis with nasal polyps in patients who have had ethmoid sinus surgery
Required Medical Information:	 Documentation of a diagnosis of chronic rhinosinusitis and has undergone prior bilateral total ethmoidectomy Indication for revision endoscopic sinus surgery due to recurrent symptoms of nasal polyps (such as nasal obstruction/congestion, bilateral sinus obstruction)
Appropriate Treatment Regimen & Other Criteria:	 Documented treatment failure to an adequate trial (minimum of 3 months each) with two nasal corticosteroid sprays Documented treatment failure of a minimum 14-day trial with an oral corticosteroid Reauthorization: documented presence of ethmoid sinus polyps, grade 1 or higher, at least 90 days after previous treatment with Sinuva
Exclusion Criteria:	 Known history of resistant or poor response to oral steroids Acute bacterial or invasive fungal sinusitis Immune deficiency (including cystic fibrosis)
Age Restriction:	18 years of age or older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an otolaryngologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 1 month, unless otherwise specified Reauthorization: 1 month, unless otherwise specified



MONOMETHYL FUMARATE

Affected Medications: BAFIERTAM (monomethyl fumarate)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design. Treatment of relapsing forms of multiple sclerosis (MS), including the following: Clinically isolated syndrome (CIS) Relapsing-remitting multiple sclerosis (RRMS) Active secondary progressive disease (SPMS)
Required Medical Information:	Diagnosis confirmed with magnetic resonance imaging (MRI), per revised McDonald diagnostic criteria for MS Clinical evidence alone will suffice; additional evidence desirable but must be consistent with MS
Appropriate Treatment Regimen & Other Criteria:	Relapsing forms of MS Coverage of Bafiertam (monomethyl fumarate) requires documentation of one of the following: Documented disease progression or intolerable adverse event with one of the following: teriflunomide, dimethyl fumarate or fingolimod Currently receiving treatment with Bafiertam (monomethyl fumarate), excluding via samples or manufacturer's patient assistance program Reauthorization requires provider attestation of treatment success
Exclusion Criteria:	Concurrent use of other disease-modifying medications indicated for the treatment of multiple sclerosis
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist or a multiple sclerosis specialist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified.



POLICY NAME: **MOTIXAFORTIDE**

Affected Medications: APHEXDA (motixafortide)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan
	 In combination with filgrastim (granulocyte colony-stimulating factor [G-CSF]) to
	mobilize hematopoietic stem cells (HSCs) to the peripheral blood circulation to
	facilitate their collection for subsequent autologous stem cell transplantation
	(ASCT) in patients with multiple myeloma (MM)
	NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A
	or better (autologous HSCT must be NCCN recommended)
Required Medical	Documentation of performance status, disease staging, all prior therapies used, and
Information:	anticipated treatment course
	Documentation of diagnosis of multiple myeloma in first or second remission
	Eligible for Autologous stem cell transplantation (ASCT)
	At least 7 days from most recent high dose induction therapy
	No single agent chemotherapy or maintenance therapy within 7 days
	Eastern Cooperative Oncology Group (ECOG) performance status (PS) of 0 or 1
Appropriate	Inadequate stem cell collection amount despite previous trial with ALL the following:
Treatment	 Single agent granulocyte colony stimulating factor (G-CSF)
Regimen & Other	 G-CSF in combination with plerixafor
Criteria:	No reauthorization
Exclusion Criteria:	Karnofsky Performance Status 50% or less or Eastern Cooperative Oncology Group
	(ECOG) performance status (PS) of 2 or greater
Age Restriction:	18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist
Care Restrictions:	
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 2 months unless otherwise specified



MUCOPOLYSACCHARIDOSIS (MPS) AGENTS

Affected Medications: VIMIZIM (elosulfase alfa), NAGLAZYME (galsulfase), MEPSEVII (vestronidase alfa-vjbk), ALDURAZYME (laronidase), ELAPRASE (idursulfase)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Vimizim: Mucopolysaccharidosis type IVA (MPS IVA; Morquio A syndrome)
	 Naglazyme: Mucopolysaccharidosis type VI (MPS VI, Maroteaux-Lamy
	syndrome)
	 Mepsevii: Mucopolysaccharidosis VII (MPS VII; Sly Syndrome)
	o Aldurazyme:
	 Hurler Mucopolysaccharidosis type I (MPS I H)
	 Herler-Scheie Mucopolysaccharidosis type I (MPS I H/S)
	 Scheie form of Mucopolysaccharidosis (MPS I S) with moderate to
	severe symptoms
	Elaprase: Mucopolysaccharidosis type II (MPS II; Hunters syndrome)
Required Medical	Diagnosis of specific MPS type confirmed by enzyme assay showing deficient activity of
Information:	the relevant enzyme OR detection of pathogenic mutations in the relevant gene by
	molecular genetic testing, as follows:
	 For Vimizim: N-acetylgalactosamine 6-sulfatase (GALNS) enzyme or GALNS
	gene For Noglozymo: N. gostylgologtosomino 4 gulfatago (ASP) anzymo ar
	 For Naglazyme: N-acetylgalactosamine 4-sulfatase (ASB) enzyme or Arylsulfatase B (ARSB) gene
	For Mepsevii: beta-glucuronidase (GUSB) enzyme or GUSB gene
	 For Aldurazyme: alpha-L-iduronidase (IDUA) enzyme or IDUA gene
	 For Elaprase: iduronate 2-sulfatase (I2S or IDS) enzyme or IDS gene
	Documented clinical signs and symptoms of MPS, such as soft tissue abnormality,
	skeletal abnormality, joint abnormality, respiratory disease, gait abnormality, motor
	issues, or cardiac abnormality
	Baseline value for one or more of the following:
	 Function test such as the Bruininks-Oseretsky Test of Motor Proficiency (BOT-
	2), 6-minute walk test (6MWT), three-minute stair climb test (3-MSCT), or
	pulmonary function tests (PFTs)
	 Liver and/or spleen volume
	Urinary glycosaminoglycan (GAGs) level
Appropriate	Dose does not exceed the recommended dosing according to the FDA label
Treatment	Dose-rounding to the nearest vial size within 10% of the prescribed dose will
Regimen & Other	be enforced
Criteria:	Deput having tion requires desumentation of two transferred and fined as ONE
	Reauthorization requires documentation of treatment success defined as ONE or more of the following:
	 Stability or improvement in function tests such as BOT-2, 6MWT, 3-MSCT, or PFTs
	Reduction in liver and/or spleen volume
	 Reduction in urinary GAG level Other clinically significant improvement in MPS signs and symptoms
L	Other clinically significant improvement in MP3 signs and symptoms



Exclusion Criteria:	 Treatment of central nervous system manifestation of the disorder Severe, irreversible cognitive impairment
Age Restriction:	 Vimizim and Naglazyme: 5 years of age and older Elaprase: 16 months of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a specialist in the treatment of inherited metabolic disorders, such as a geneticist or endocrinologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



MUSCULAR DYSTROPHY RNA THERAPY

Affected Medications: AMONDYS 45 (casimersen), EXONDYS 51 (eteplirsen), VYONDYS 53 (golodirsen), VILTEPSO (viltolarsen)

Covered Uses:	Casimersen (Amondys 45), eteplirsen (Exondys 51), golodirsen (Vyondys 53), and viltolarsen (Viltepso) are not considered medically necessary due to insufficient evidence of therapeutic value.
Required Medical	
Information:	
Appropriate	
Treatment	
Regimen & Other	
Criteria:	
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of	
Care Restrictions:	
Coverage Duration:	



MYELOID GROWTH FACTORS

Affected Medications: UDENYCA (pegfilgrastim-cbqv), FULPHILA (pegfilgrastim-jmdb), NEULASTA (pegfilgrastim), ZIEXTENZO (pegfilgrastim-bmez), NYVEPRIA (pegfilgrastim-apgf), NEUPOGEN (filgrastim), ZARXIO (filgrastim-sndz), GRANIX (tbo-filgrastim), LEUKINE (sargramostim), NIVESTYM (filgrastim-aafi), RELEUKO (filgrastim-ayow), FYLNETRA (pegfilrastim-pbbk), ROLVEDON (eflapegrastim-xnst), STIMUFEND (pegfilgrastim-fpgk), NYPOZI (filgrastim-txid)

Covered Uses:

 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design

Neupogen, Nivestym, Nypozi, Releuko and Zarxio

Patients with Cancer Receiving Myelosuppressive Chemotherapy

Indicated to decrease the incidence of infection, as manifested by febrile neutropenia, in
patients with non-myeloid malignancies receiving myelosuppressive anti-cancer drugs
associated with a significant incidence of severe neutropenia with fever.

<u>Patients With Acute Myeloid Leukemia Receiving Induction or Consolidation</u> Chemotherapy

• Indicated for reducing the time to neutrophil recovery and the duration of fever, following induction or consolidation chemotherapy treatment of adults with acute myeloid leukemia.

Patients with Cancer Receiving Bone Marrow Transplant

 Indicated to reduce the duration of neutropenia and neutropenia-related clinical sequelae, (e.g., febrile neutropenia) in patients with non-myeloid malignancies undergoing myeloablative chemotherapy followed by marrow transplantation.

<u>Patients Undergoing Autologous Peripheral Blood Progenitor Cell Collection and Therapy (Neupogen, Nivestym, Nypozi, Zarxio)</u>

 Indicated for the mobilization of autologous hematopoietic progenitor cells into the peripheral blood for collection by leukapheresis.

Patients With Severe Chronic Neutropenia

• Indicated for chronic administration to reduce the incidence and duration of sequelae of neutropenia (e.g., fever, infections, oropharyngeal ulcers) in symptomatic patients with congenital neutropenia, cyclic neutropenia, or idiopathic neutropenia.

<u>Patients Acutely Exposed to Myelosuppressive Doses of Radiation (Hematopoietic</u> Syndrome of Acute Radiation Syndrome) (Neupogen)

 Indicated to increase survival in patients acutely exposed to myelosuppressive doses of radiation.

Leukine

<u>Use Following Induction Chemotherapy in Acute</u> <u>Myelogenous Leukemia</u>

 Indicated for use following induction chemotherapy in older adult patients with acute myelogenous leukemia to shorten time to neutrophil recovery and to reduce the incidence of severe and life-threatening infections and infections resulting in death.



<u>Use in Mobilization and Following Transplantation of Autologous Peripheral Blood</u> Progenitor Cells

 Indicated for the mobilization of hematopoietic progenitor cells into peripheral blood for collection by leukapheresis.

Use in Myeloid Reconstitution After Autologous Bone Marrow Transplantation

 Indicated for acceleration of myeloid recovery in patients with non-Hodgkin's lymphoma (NHL), acute lymphoblastic leukemia (ALL) and Hodgkin's disease undergoing autologous bone marrow transplantation (BMT).

Use in Myeloid Reconstitution After Allogeneic Bone Marrow Transplantation

 Indicated for acceleration of myeloid recovery in patients undergoing allogeneic BMT from human leukocyte antigen (HLA)-matched related donors.

Use in Bone Marrow Transplantation Failure or Engraftment Delay

• Indicated in patients who have undergone allogeneic or autologous BMT in whom engraftment is delayed or has failed.

Neulasta, Fulphila, Udenyca, Ziextenzo, Nyvepria, Fylnetra, Stimufend, and Rolvedon

Patients with Cancer Receiving Myelosuppressive Chemotherapy

Indicated to decrease the incidence of infection, as manifested by febrile neutropenia, in
patients with non-myeloid malignancies receiving myelosuppressive anti-cancer drugs
associated with a significant incidence of severe neutropenia with fever.

<u>Patients with Hematopoietic Subsyndrome of Acute Radiation Syndrome (Neulasta, Udenyca, Ziextenzo)</u>

 Indicated to increase survival in patients acutely exposed to myelosuppressive doses of radiation

Granix

 Indicated to reduce the duration of severe neutropenia in patients with non-myeloid malignancies receiving myelosuppressive anti-cancer drugs associated with a clinically significant incidence of febrile neutropenia.

Compendia supported uses that will be covered (if applicable) Neupogen/Granix/Zarxio/Nypozi/Nivestym/Leukine:

- Treatment of chemotherapy-induced febrile neutropenia in patients with non-myeloid malignancies
- Treatment of anemia in patients with myelodysplastic syndromes (MDS)
- Treatment of neutropenia in patients with MDS
- Following chemotherapy for acute lymphocytic leukemia (ALL)
- Stem cell transplantation-related indications
- Agranulocytosis
- Aplastic anemia
- Neutropenia related to human immunodeficiency virus (HIV)
- Neutropenia related to renal transplantation

Required Medical Information:

 Complete blood counts with differential and platelet counts will be monitored at baseline and regularly throughout therapy



•	Documentation of therapy intention (curative, palliative) for prophylaxis of febrile
	neutropenia

- Documentation of patient specific risk factors for febrile neutropenia
- Documentation of febrile neutropenia risk associated with the chemotherapy regimen.
- Documentation of planned treatment course
- Documentation of current patient weight

Appropriate Treatment Regimen & Other Criteria:

Filgrastim products: Neupogen, Nivestym, Releuko, Zarxio, Granix, Nypozi

When requested via the MEDICAL benefit:

Coverage for the non-preferred products, Neupogen, Nypozi, Releuko and Granix, is provided when the member meets the following criteria:

Documented treatment failure or intolerable adverse event to Zarxio and Nivestym

When requested through the specialty PHARMACY benefit:

Coverage for the non-preferred products, Neupogen, Nypozi, Zarxio, Releuko and Granix, is provided when the member meets the following criteria:

Documented treatment failure or intolerable adverse event to Nivestym

Sargramostim product: Leukine

Coverage for the non-preferred product, Leukine, is provided when the member meets one of the following criteria:

- Leukine will be used for myeloid reconstitution after autologous or allogenic bone marrow transplant or bone marrow transplant engraftment delay or failure
- A documented treatment failure or intolerable adverse event to preferred products listed above

<u>Pegfilgrastim products: Neulasta, Fulphila, Udenyca, Ziextenzo, Nyvepria, Fylnetra, Stimufend, Rolvedon</u>

Coverage for the non-preferred products, Neulasta, Fylnetra, Rolvedon, Stimufend, Ziextenzo and Nyvepria is provided when the member meets the following criteria:

Documented treatment failure or intolerable adverse event to Fulphila and Udenyca

Eflapegrastim product: Rolvedon

Coverage for the non-preferred product, Rolvedon, is provided when the member meets the following criteria:

 Documented treatment failure or intolerable adverse event to the preferred pegfilgrastim products Fulphila and Udenyca

For prophylaxis of febrile neutropenia (FN) or other dose-limiting neutropenic events for patients receiving myelosuppressive anticancer drugs:

Meets **ONE** of the following:

- Curative Therapy:
 - High risk (greater than 20% risk) for febrile neutropenia based on chemotherapy regimen OR
 - Intermediate risk (10-20% risk) for febrile neutropenia based on chemotherapy regimen with documentation of significant patient risk factors for serious medical consequences OR



	 Has experienced a dose-limiting neutropenic event on a previous cycle of current chemotherapy to be continued 		
	Palliative Therapy:		
	Myeloid growth factors will not be approved upfront for prophylaxis of febrile neutropenia in the palliative setting. Per the NCCN (National Comprehensive Cancer Network), chemotherapy regimens with a 20% or greater risk of neutropenic events should not be used. If however, a dose limiting neutropenic event occurs on a previous cycle of chemotherapy, and the effectiveness of chemotherapy will be reduced with dose reduction, growth factor will be approved for secondary prophylaxis on a case by case basis.		
	For Treatment of Severe Chronic Neutropenia		
	Must meet ALL the following:		
	Congenital neutropenia, cyclic neutropenia, OR idiopathic neutropenia		
	 Current documentation of absolute neutrophil count (ANC) less than 500 cells/microliter Neutropenia symptoms (fever, infections, oropharyngeal ulcers) 		
Exclusion	14Cuttoponia symptoms (tevet, infections, otopharyngeal dicers)		
Criteria:			
Age			
Restriction:			
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist or hematologist		
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care		
Coverage Duration:	Authorization: 6 months, unless otherwise specified		
Duration:			



POLICY NAME: **NAFARELIN**

Affected Medications: SYNAREL (nafarelin)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Central Precocious Puberty (CPP) Endometriosis 		
Required Medical Information:	Central Precocious Puberty: Documentation of CPP confirmed by basal luteinizing hormone (LH), follicle-stimulating hormone (FSH), and either estradiol or testosterone concentrations Endometriosis: Documentation of moderate to severe pain due to endometriosis		
Appropriate	Endometriosis:		
Treatment			
Regimen & Other Criteria:	 Documentation of a trial and inadequate relief (or contraindication) after at least 3 months of both of the following first-line therapies: Nonsteroidal anti-inflammatory drugs (NSAIDs) Continuous (no placebo pills) hormonal contraceptives Maximum treatment duration 6 months total 		
Exclusion Criteria:	Use for infertility (if benefit exclusion) Undiagnosed abnormal vaginal bleeding		
Age Restriction:	 Endometriosis: 18 years of age and older Central precocious puberty (CPP): 11 years of age or younger (females), 12 years of age or younger (males) 		
Prescriber/Site of	Prescribed by, or in consultation with, an endocrinologist or gynecologist		
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care		
Coverage Duration:	 Endometriosis: 6 months (no reauthorization), unless otherwise specified CPP: 12 months, unless otherwise specified 		



POLICY NAME: NALOXEGOL

Affected Medications: MOVANTIK (naloxegol)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design. Opioid-induced constipation
Required Medical Information:	Documentation supporting a diagnosis of opioid-induced constipation in a patient with chronic, non-cancer pain that has been taking opioids for at least 4 weeks
Appropriate Treatment Regimen & Other Criteria:	Documented treatment failure or intolerable adverse event to polyethylene glycol 3350 (PEG 3350) and one other laxative (such as lactulose) Reauthorization will require documentation of treatment success and a clinically significant response to therapy, AND documented continued use of opioid pain medication
Exclusion Criteria:	Known or suspected mechanical gastrointestinal obstruction.
Age Restriction:	
Prescriber/Site of Care Restrictions:	All approvals are subjects to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: **NATALIZUMAB**

Affected Medications: TYSABRI (natalizumab)

Covered Uses: Required Medical Information:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of relapsing forms of multiple sclerosis (MS), including the following: Clinically isolated syndrome (CIS) Relapsing-remitting multiple sclerosis (RRMS) Active secondary progressive multiple sclerosis (SPMS) Crohn's disease (CD) Screening for anti-JC virus (JCV) antibodies prior to initiating Tysabri therapy
	Diagnosis confirmed with magnetic resonance imaging (MRI), per revised McDonald diagnostic criteria for MS Clinical evidence alone will suffice; additional evidence desirable but must be consistent with MS
	Crohn's disease
Appropriate	Moderate to severely active disease despite current treatment Relapsing Forms of MS
Treatment	Documentation of treatment failure (or documented intolerable adverse event) to:
Regimen & Other	Rituximab (preferred biosimilar products: Riabni and Ruxience) OR
Criteria:	 Ocrevus (ocrelizumab) if previously established on treatment OR
	 Documentation of pregnancy and severe disease
	 Crohn's disease Documented treatment failure or intolerable adverse event with at least 12 weeks of TWO oral treatments: corticosteroids, azathioprine, 6-mercaptopurine, sulfasalazine, balsalazide, or methotrexate AND Documented clinical failure with at least 12 weeks of infliximab (preferred biosimilar products: Inflectra and Renflexis)
	Reauthorization:
	 Anti-JCV antibody <u>negative</u>: documentation of positive clinical response to therapy Anti-JCV antibody <u>positive</u>: documentation of positive clinical response to therapy and periodic MRI to monitor for progressive multifocal leukoencephalopathy (PML)
Exclusion Criteria:	Current or prior history of PML
	MS: concurrent use of other disease-modifying medications indicated for the treatment of
	 multiple sclerosis CD: concurrent use of other targeted immune modulators for the treatment of Crohn's disease
Age Restriction:	
Prescriber/Site of	MS: prescribed by, or in consultation with, a neurologist or a MS specialist
Care Restrictions:	CD: prescribed by, or in consultation with, a gastroenterologist



	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Relapsing Forms of MS:
	Authorization: 12 months, unless otherwise specified
	Crohn's Disease:
	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **NAXITAMAB**

Affected Medications: DANYELZA (naxitamab)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of relapsed or refractory high-risk neuroblastoma in the bone or bone marrow (in combination with granulocyte-macrophage colony-stimulating factor [GM-CSF]) in patients who have demonstrated a partial response, minor response, or stable disease to prior therapy NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher
Required Medical	Documentation of performance status, disease staging, all prior therapies used, and
Information:	anticipated treatment course.
	Diagnosis of neuroblastoma as defined per the International Neuroblastoma Response Criteria (INRC):
	 An unequivocal histologic diagnosis from tumor tissue by light microscopy [with or without immunohistochemistry, electron microscopy, or increased urine (or serum) catecholamines or their metabolites] OR
	 Evidence of metastases to bone marrow on an aspirate or trephine biopsy with concomitant elevation of urinary or serum catecholamines or their metabolites
	 Evidence of high-risk neuroblastoma, including: Stage 2/3/4/4S disease with amplified MYCN gene (any age)
	 Stage 4 disease in patients greater than 18 months of age
	Disease is evaluable in the bone and/or bone marrow, as documented by histology and/or appropriate imaging [e.g., metaiodobenzylguanidine (MIBG) scan and positron emission topography (PET) scan if MIBG is negative]
	Documented history of previous treatment with at least one systemic therapy to treat disease outside of the bone or bone marrow
	Documentation of clinical rationale for avoiding use of dinutuximab plus chemotherapy (if under 18 years of age)
Appropriate Treatment	Must be used in combination with granulocyte-macrophage colony-stimulating factor (GM-CSF).
Regimen & Other	Poputhorization will require documentation of disease responsiveness to therepy
Criteria:	Reauthorization will require documentation of disease responsiveness to therapy
Exclusion Criteria:	 Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater Patients with progressive disease
Age Restriction:	1 year of age or older
Prescriber/Site of Care Restrictions:	 Must be prescribed by, or in consultation with, a hematologist/oncologist with expertise in neuroblastoma All approvals are subject to utilization of the most cost-effective site of care



Coverage Duration:	•	Initial Authorization: 4 months, unless otherwise specified
	•	Reauthorization: 12 months, unless otherwise specified



NEMOLIZUMAB-ILTO

Affected Medications: NEMLUVIO (nemolizumab-ilto)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise excluded by			
	plan design			
	 Prurigo nodularis (PN) 			
	Atopic dermatitis (AD)			
Required Medical	<u>PN</u>			
Information:	Documentation of all the following:			
	 Diagnosis confirmed by skin biopsy 			
	 Presence of at least 20 PN lesions for at least 3 months 			
	 Severe itching 			
	<u>AD</u>			
	Documentation of severe inflammatory skin disease defined as functional impairment			
	(inability to use hands or feet for activities of daily living or significant facial involvement			
	preventing normal social interaction) AND			
	Body Surface Area (BSA) of at least 10% OR Hand foot or much mambrane involvement.			
Appropriate	Hand, foot or mucous membrane involvement PN			
Treatment				
	Documented treatment failure with at least 2 weeks of a super high potency topical article state of the state of transition state 0.05% ballot state of the state of			
Regimen & Other Criteria:	corticosteroid (such as clobetasol propionate 0.05%, halobetasol propionate 0.05%)			
Criteria:	Documentation of treatment failure with at least 12 weeks of one of the following:			
	phototherapy, methotrexate, cyclosporine			
	Documented treatment failure with at least 12 weeks of Dupixent (dupilumab)			
	<u>AD</u>			
	Documentation of treatment failure with at least 6 weeks of one of the following: tacrolimus ointment, pimecrolimus cream, Eucrisa			
	Documentation of treatment failure with at least 12 weeks of one of the following:			
	phototherapy, methotrexate, cyclosporine			
	Documented treatment failure with at least 12 weeks of Dupixent (dupilumab)			
Exclusion Criteria:	Concurrent use with another therapeutic immunomodulator agent			
Age Restriction:	PN: 18 years of age and older			
	AD: 12 years of age and older			
Prescriber/Site of	 Prescribed by, or in consultation with, a dermatologist, allergist, or immunologist 			
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care			
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified			
	Reauthorization: 12 months, unless otherwise specified			
	<u>'</u>			



NEONATAL FC RECEPTOR ANTAGONISTS

Affected Medications: VYVGART (efgartigimod alfa), VYVGART HYTRULO (efgartigimod alfa and hyaluronidase), RYSTIGGO (rozanolixizumab)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by			
	plan design			
	Vyvgart			
	 Generalized myasthenia gravis (gMG) in adult patients who are anti- 			
	acetylcholine receptor (AChR) antibody positive			
	Rystiggo			
	 Generalized myasthenia gravis (gMG) in adult patients who are AChR or anti- muscle-specific tyrosine kinase (MuSK) antibody positive 			
	Vyvgart Hytrulo			
	Generalized myasthenia gravis (gMG) in adult patients who are anti-			
	acetylcholine receptor (AChR) antibody positive			
	Chronic Inflammatory Demyelinating Polyneuropathy (CIDP)			
Required Medical	Myasthenia Gravis			
Information:	 Diagnosis of generalized Myasthenia Gravis (gMG) confirmed by one of the following: A history of abnormal neuromuscular transmission test A positive edrophonium chloride test 			
	 Improvement in gMG signs or symptoms with an acetylcholinesterase inhibitor Myasthenia Gravis Foundation of America (MGFA) Clinical Classification Class II to IV Positive serologic test for AChR or MuSK antibodies (for Rystiggo) 			
	 Documentation of ONE of the following: MG-Activities of Daily Living (MG-ADL) total score of 6 or greater Quantitative Myasthenia Gravis (QMG) total score of 12 or greater 			
	CIDP (Vyvgart Hytrulo only)			
	Documented baseline in strength/weakness using an objective clinical measuring tool (INCAT, Medical Research Council (MRC) muscle strength, 6 Minute Walk Test, Rankin, Modified Rankin)			
	Documented disease course is progressive or relapsing and remitting for 2 months or			
	longer			
	Abnormal or absent deep tendon reflexes in upper or lower limbs			
	Electrodiagnostic evidence of demyelination indicated by one of the following:			
	o Motor distal latency prolongation in 2 nerves			
	 Reduction of motor conduction velocity in 2 nerves Prolongation of F-wave latency in 2 nerves 			
	 Prolongation of F-wave latency in 2 nerves Absence of F-waves in at least 1 nerve 			
	Partial motor conduction block of at least 1 motor nerve			
	 Abnormal temporal dispersion in at least 2 nerves 			
	 Distal CMAP duration increase in at least 1 nerve 			
	Cerebrospinal fluid (CSF) analysis indicates all of the following (if electrophysiologic			
	findings are non-diagnostic):			
	CSF white cell count of less than 10 cells/mm³ SSF protein is also at all (greater than or agual to 45 mg/dl.)			
Appropriate	CSF protein is elevated (greater than or equal to 45mg/dL) Currently on a stable dose of at least one gMC thorapy (acetylchelinestorase inhibitor).			
Appropriate	 Currently on a stable dose of at least one gMG therapy (acetylcholinesterase inhibitor, corticosteroid, or non-steroidal immunosuppressive therapy (NSIST)) that will be 			
Treatment	controsteroid, or non-steroidal infinitionosuppressive therapy (NSIST)) that will be			



Regimen & Other	continued during initial treatment with Vyvgart, Vyvgart Hytrulo, or Rystiggo
Criteria:	Documentation of ONE of the following:
	 Coverage for Rystiggo is provided when one of the following is met: Currently receiving treatment with Rystiggo, excluding when the product is obtained as samples or via manufacturer's patient assistance programs Documented treatment failure or intolerable adverse event with Vyvgart for AChR antibody positive gMG Documented treatment failure with rituximab for MuSK antibody positive gMG (preferred products: Riabni, Ruxience)
	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced Reauthorization:
	Documentation of treatment success and clinically significant response to therapy defined as:
	 A minimum 2-point reduction in MG-ADL score from baseline or improvement in QMG total score Absent or reduced need for rescue therapy compared to baseline
	Documentation that the patient requires continuous treatment, after an initial beneficial response, due to new or worsening disease activity
	Note : a minimum of 50 days for Vyvgart/Vyvgart Hytrulo or 63 days for Rystiggo must have elapsed from the start of the previous treatment cycle
	CIDP (Vyvgart Hytrulo only) Documented trial and failure of at least 3 months of intravenous or subcutaneous immune globulin
	 Reauthorization: Documentation of a clinical response to therapy based on an objective clinical measuring tool (e.g., INCAT, Medical Research Council (MRC) muscle strength, 6-Minute walk test, Rankin, Modified Rankin)
Exclusion Criteria:	 Immunoglobulin G (IgG) levels less than 600 mg/dL at baseline Concurrent use with other disease-modifying biologics for the treatment of gMG
Age Restriction:	18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care



Coverage Duration: • Initial Authorization: 4 months, unless otherwise specified		Initial Authorization: 4 months, unless otherwise specified
	•	Reauthorization: 12 months, unless otherwise specified



NIEMANN-PICK DISEASE TYPE C (NPC) AGENTS
Affected Medications: MIPLYFFA (arimoclomol citrate), AQNEURSA (levacetylleucine)

All Food and Drug Administration (FDA)-approved indications not otherwise.				
	plan design			
	Niemann-Pick disease type C (NPC)			
Required Medical	Diagnosis of NPC confirmed by genetic testing showing biallelic pathogenic variants in			
Information:	either the NPC1 gene or NPC2 gene			
	Documentation of at least one neurological symptom of Niemann-Pick disease type C,			
	such as:			
	 Loss of motor function 			
	 Problems with swallowing or speech 			
	 Cognitive impairment 			
	Documentation of being ambulatory without needing an assistive device such as a			
	wheelchair, walker, or cane			
	Documentation of baseline signs and symptoms of NPC			
Appropriate	For Miplyffa:			
Treatment	Documentation that patient has been receiving miglustat with a stable dose for at least			
Regimen & Other	the past 6 consecutive months			
Criteria:	Documentation that Miplyffa will be taken in combination with miglustat			
	Reauthorization requires:			
 Documentation of treatment success defined as stability or improvement of 				
	Pick disease type C signs and symptoms			
	Documentation that patient is still ambulatory			
	For Miplyffa: that the drug continues to be used in combination with miglustat			
Exclusion Criteria:	Use of Miplyffa and Aqneursa in combination			
Age Restriction:	Miplyffa: 2 years of age and older			
	Aqneursa: Adults and pediatric patients weighing 15 kilograms or greater			
Prescriber/Site of	Prescribed by, or in consultation with, a specialist in the management of NPC (such as a			
Care Restrictions:	geneticist, endocrinologist, metabolic disorder subspecialist, or neurologist)			
	All approvals are subject to utilization of the most cost-effective site of care			
Coverage Duration:	Authorization: 12 months, unless otherwise specified			



POLICY NAME: **NILOTINIB**

Affected Medications: TASIGNA (nilotinib)

Covered Uses:	NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher	
Required Medical Information:	 Documentation of performance status, all prior therapies used, and prescribed treatment regimen Documentation Philadelphia chromosome or BCR::ABL1-positive mutation status 	
Appropriate Treatment	For patients with Chronic Myeloid Leukemia (CML) and low-risk score, documented clinical failure with imatinib	
Regimen & Other Criteria:	<u>Reauthorization</u> requires documentation of disease responsiveness to therapy (as applicable, BCR-ABL1 transcript levels, cytogenetic response)	
Exclusion Criteria:	Karnofsky Performance Status 50% or less, ECOG performance score 3 or greater	
Age Restriction:		
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an oncologist All approvals are subject to utilization of the most cost-effective site of care 	
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified 	



POLICY NAME: NIROGACESTAT

Affected Medications: OGSIVEO (nirogacestat)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design			
	 Progressive desmoid tumor(s) requiring systemic therapy 			
	 NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher 			
Required Medical Information:	 Documentation of performance status, disease staging, all prior therapies used, and anticipated treatment course 			
	 Diagnosis of biopsy proven desmoid tumor/aggressive fibromatosis (DT/AF) with documentation of tumor progression (tumor growth causing chronic pain, disfigurement, internal bleeding, and/or impaired range of motion) 			
Appropriate	Documentation of clinical failure with sorafenib			
Treatment				
Regimen & Other	Reauthorization: documentation of disease responsiveness to therapy			
Criteria:				
Exclusion Criteria:	Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater			
Age Restriction:	18 years of age and older			
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist			
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care			
Coverage Duration:	Initial Authorization: 4 months, unless otherwise specified			
	Reauthorization: 12 months, unless otherwise specified			



NON-Preferred HYALURONIC ACID DERIVATIVES

Affected Medications: DUROLANE (hyaluronic acid), EUFLEXXA (1% sodium hyaluronate), GEL-ONE (cross-linked hyaluronate), GELSYN-3 (sodium hyaluronate 0.84%), GENVISC 850 (sodium hyaluronate), HYALGAN (sodium hyaluronate), HYMOVIS (high molecular weight viscoelastic hyaluronan), MONOVISC (high molecular weight hyaluronan), SUPARTZ (sodium hyaluronate), SYNOJOYNT (sodium hyaluronate), TRIVISC (sodium hyaluronate), VISCO-3 (sodium hyaluronate)

1.	Is this the first time a Hyaluronic Acid (HA) derivative product is being used in this member for this indication?	Yes – Go to #2	No – Document date of last use and go to Renewal criteria		
2.	Is the request for a Food and Drug Administration (FDA)-approved indication: Treatment of osteoarthritis pain of the knee?	Yes – Go to #3	No – Criteria not met		
3.	Is there documented failure to respond to conservative non- pharmacologic therapy (such as ice, physical therapy) and simple analgesics (such as acetaminophen)?	Yes – Document and go to #4	No – Criteria not met		
4.	Has there been a documented intolerable adverse event to Synvisc, Synvisc-One, and Orthovisc with date and description of reactions?	Yes – Go to #6	No – Go to #5		
5.	Is the member currently undergoing treatment and coverage is required to complete the current course of treatment?	Yes – Document and go to #6	No – Criteria not met		
6.	Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations?	Yes – Document and approve up to 6 months	No – Criteria not met		
Re	Renewal for hyaluronic acid (HA) after previous administration of HA product				
1.	Is there documentation of treatment success that lasted at least 6 months from date of previous HA administration AND documented intolerable adverse event to Synvisc, Synvisc-One, and Orthovisc with date and description of reactions?	Yes – Go to #2	No – Criteria not met		
2.	Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations?	Yes – Approve up to 6 months	No – Criteria not met		



Quantity Limitations

Durolane: 1 injection per courseEuflexxa: 3 injections per course

Euflexxa: 3 injections per courseGel-One: 1 injection per course

Gelsyn-3: 3 injections per course

GenVisc 850: 3 to 5 injections per course

Hyalgan: 5 injections per course

Hymovis: 2 injections per course

Monovisc: 1 injection per course

Supartz: 3 to 5 injections per courseSynojoynt: 3 injections per course

Triluron: 3 injections per course

Trivisc: 3 injections per courseVisco-3: 3 injections per course



NON-PREFERRED MEDICAL DRUG CODES

Affected Medications: BORTEZOMIB, PEMETREXED

Covered Uses: Required Medical Information:	plan design	ations: National Comprehe	oved indications not otherwise exnsive Cancer Network (NCCN) in	_
Appropriate Treatment Regimen & Other Criteria: • Approval of a non-preferred medical drug listed below requires documentation intolerable adverse event to all the preferred alternatives, and the adverse event attributed to the active ingredient				
	Drug	Non-Preferred code (Manufacturer)	Preferred Alternatives	
	Bortezomib (Boruzu, Velcade)	J9046 (Dr. Reddy's, Boruzu)	J9041, J9048, J9049	
	Pemetrexed (Pemfexy, Alimta, Pemrydi RTU, Axtle)	J9304 (Apotex), J9292 (Axtle)	J9294, J9296, J9297, J9305, J9314, J9324	
	Reauthorization: docu	onsiveness to therapy	_	
Exclusion Criteria:				
Age Restriction:				
Prescriber/Site of Care Restrictions:	All approvals are su	ubject to utilization of the m	nost cost-effective site of care	
Coverage Duration:	Authorization: 12 months, unless otherwise specified			



POLICY NAME: **NUEDEXTA**

Affected Medications: NUEDEXTA (dextromethorphan hydrobromide/quinidine sulfate)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design. Treatment of pseudobulbar affect (PBA)
Required Medical Information:	Documentation of at least ONE underlying neurological condition associated with PBA such as: amyotrophic lateral sclerosis (ALS) extrapyramidal and cerebellar disorders (Parkinson's disease, multiple system atrophy, progressive supranuclear palsy) multiple sclerosis (MS) traumatic brain injury Alzheimer's disease and other dementias stroke. Baseline Center for Neurologic Study-Lability Scale (CNS-LS) score of 13 or greater Documentation of treatment failure to a 30-day trial of each of the following: serotonin reuptake inhibitor (SSRI) tricyclic antidepressant (TCA)
Appropriate Treatment Regimen & Other Criteria: Exclusion Criteria:	Reauthorization requires documentation of treatment success defined as decreased frequency of pseudobulbar affect (PBA) episodes.
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Approval: 12 months, unless otherwise specified



NULIBRY

Affected Medications: NULIBRY (fosdenopterin)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design To reduce the risk of mortality in patients with molybdenum cofactor deficiency (MoCD) Type A	
Required Medical Information:	 Documentation of presumptive or genetically confirmed molybdenum cofactor deficiency (MoCD) Type A diagnosis Presumptive diagnosis of Molybdenum cofactor deficiency (MoCD) Type A Documentation of family history meeting ONE of the following: Affected sibling(s) with confirmed MoCD Type A; or a history of deceased sibling(s) with classic MoCD presentation One or both parents are known to carry a copy of the mutated gene [Molybdenum Cofactor Synthesis 1 (MOCS1)] Child has consanguineous parents with a family history of MoCD Onset of clinical and/or laboratory signs and symptoms consistent with MoCD Type A, such as: Clinical presentation: intractable seizures, exaggerated startle response, high-pitched cry, axial hypotonia, limb hypertonia, feeding difficulties Biochemical findings: elevated urinary sulfite and/or S-sulfocysteine (SSC), elevated xanthine in urine or blood, or low/absent uric acid in the urine or blood Genetic testing to confirm diagnosis of MoCD Type A is scheduled or in progress 	
	 Confirmed diagnosis of MoCD Type A: Diagnosis of MoCD Type A confirmed by genetic testing showing the presence of mutation in molybdenum cofactor synthesis gene 1 (MOSC1) 	
Appropriate Treatment Regimen & Other Criteria:	Reauthorization: Documentation of clinically significant response to therapy as determined by prescribing provider Documentation of genetically confirmed MoCD Type A (MOCS1 mutation) if initially approved for presumptive diagnosis	
Exclusion Criteria:	 Molybdenum cofactor deficiency (MoCD) Type B (MOCS2 mutation) MoCD Type C (gephyrin or GPHN mutation) 	
Age Restriction: Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neonatologist, pediatrician, pediatric neurologist, neonatal neurologist, or geneticist All approvals are subject to utilization of the most cost-effective site of care 	
Coverage Duration:	Presumptive diagnosis:	



Authorization: 1 month, unless otherwise specified. Must have confirmed diagnosis for
continued approval.
Confirmed diagnosis:
Authorization: 12 months, unless otherwise specified



POLICY NAME: **NUPLAZID**

Affected Medications: NUPLAZID (pimavanserin tartrate)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of hallucinations and delusions associated with Parkinson's disease (PD) psychosis
Required Medical Information:	 Diagnosis of Parkinson's disease (PD) Presence of psychotic symptoms: hallucinations and/or delusions described as severe and frequent that started after the PD diagnosis
Appropriate Treatment Regimen & Other	Documentation of treatment failure or contraindication to a 30-day trial of quetiapine Reauthorization requires documentation of treatment success and a clinically significant response to therapy
Criteria: Exclusion Criteria:	response to therapy
Age Restriction: Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: **NUSINERSEN**

Affected Medications: SPINRAZA (nusinersen)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Spinal muscular atrophy (SMA)
Required Medical	Diagnosis of SMA type 1, 2, or 3 confirmed by genetic testing of chromosome 5q13.2
Information:	demonstrating ONE of the following:
	 Homozygous gene deletion of SMN1 (survival motor neuron 1)
	 Homozygous gene mutation of SMN1
	 Compound heterozygous gene mutation of SMN1
	Documentation of 2 or more copies of the SMN2 (survival motor neuron 2) gene
	Documentation of previous treatment history
	Documentation of one of the following baseline motor assessments appropriate for
	patient age and motor function:
	 Hammersmith Infant Neurological Examination (HINE-2)
	 Hammersmith Functional Motor Scale (HFSME)
	 Children's Hospital of Philadelphia Infant Test of Neuromuscular Disorders
	(CHOP-INTEND)
	 Upper Limb Module (ULM) test
	o 6-Minute Walk Test (6MWT)
	Documentation of ventilator use status
	 Patient is NOT ventilator-dependent (defined as using a ventilator at least 16
	hours per day on at least 21 of the last 30 days)
	 This does not apply to patients who require non-invasive ventilator assistance
	Planned treatment regimen
Appropriate	Documented treatment failure with or intolerable adverse event on Evrysdi
Treatment	
Regimen & Other	Reauthorization requires documentation of improvement in baseline motor assessment
Criteria:	score, clinically meaningful stabilization, or delayed progression of SMA-associated signs and
	symptoms
Exclusion Criteria:	SMA type 4
	Advanced SMA at baseline (complete paralysis of limbs, permanent ventilation support)
	Prior treatment with SMA gene therapy (i.e., onasemnogene abeparvovec-xioi)
	Will not use in combination with other agents for SMA (e.g., onasemnogene
	abeparvovec-xioi, risdiplam, etc.)
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist or provider who is experienced in
Care Restrictions:	treatment of spinal muscular atrophy
Jaie Nestrictions.	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 8 months, unless otherwise specified
Jordiago Daration.	Reauthorization: 12 months, unless otherwise specified
	1 Teauthonzation. 12 months, unless otherwise specified



POLICY NAME: OCRELIZUMAB

Affected Medications: OCREVUS (ocrelizumab), OCREVUS ZUNOVO (ocrelizumab hyaluronidase)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design. Primary progressive multiple sclerosis (PPMS) Treatment of relapsing forms of multiple sclerosis (MS), including the following: Clinically isolated syndrome (CIS) Relapsing-remitting multiple sclerosis (RRMS) Active secondary progressive disease (SPMS)
Required Medical	All Indications:
Information:	Diagnosis confirmed with magnetic resonance imaging (MRI) per revised McDonald diagnostic criteria for MS Clinical evidence alone will suffice; additional evidence desirable but must be consistent with MS
	 Primary Progressive MS: Documentation of at least one year of disease progression and baseline Expanded Disability Status Scale (EDSS) of 3.0 to 6.5
Appropriate	Relapsing forms of MS:
Treatment Regimen & Other Criteria:	Coverage of Ocrevus (ocrelizumab) or Ocrevus Zunovo (ocrelizumab hyaluronidase) requires documentation of one of the following: Documented disease progression or intolerable adverse event with rituximab
	 (biosimilar products, Riabni and Ruxience, preferred) Currently receiving treatment with Ocrevus (ocrelizumab) or Ocrevus Zunovo (ocrelizumab hyaluronidase), excluding via samples or manufacturer's patient assistance program
	Reauthorization requires documentation of treatment success
Exclusion Criteria:	Active hepatitis B infection
	Concurrent use of other disease-modifying medications indicated for the treatment of MS
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist or MS specialist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



OFEV

Affected Medications: OFEV (nintedanib esylate)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded
	by plan design
	Idiopathic pulmonary fibrosis (IPF)
	Chronic fibrosing interstitial lung disease (ILD) with a progressive phenotype
	Systemic sclerosis-associated interstitial lung disease (SSc-ILD)
Required Medical	Idiopathic Pulmonary Fibrosis (IPF)
Information:	Documented diagnosis of idiopathic pulmonary fibrosis (IPF) confirmed by ONE of the
	following:
	 Usual interstitial pneumonia (UIP) pattern demonstrated on high-resolution
	computed tomography (HRCT)
	 UIP pattern demonstrated on surgical lung biopsy
	 Probable UIP pattern demonstrated on BOTH HRCT and surgical lung biopsy
	Documentation confirming known causes of interstitial lung disease have been ruled out
	(e.g., rheumatic disease, environmental exposure, drug toxicity)
	Documentation of BOTH of the following:
	The state of the s
	 Baseline forced vital capacity (FVC) greater than or equal to 50 percent
	predicted
	Baseline diffusing capacity for carbon monoxide (DLCO) greater than or equal
	to 30 percent predicted
	Systemic Sclerosis-Associated Interstitial Lung Disease (SSc-ILD)
	Documented diagnosis of SSc-ILD
	Documentation of greater than or equal to 10% fibrosis on a chest high resolution
	computed tomography (HRCT) scan conducted within the previous 12 months
	Documentation of baseline FVC greater than or equal to 40% of predicted
	Documentation of predicted DLCO 30-89% of predicted
	Chronic Fibrosing Interstitial Lung Disease (ILD) with a Progressive Phenotype
	Documented diagnosis of chronic fibrosing ILD with a progressive phenotype (aka)
	progressive pulmonary fibrosis), confirmed by at least two of the following:
	 Worsening respiratory symptoms
	 Physiological evidence of disease progression (defined as DLCO reduced by
	10% or greater OR FVC reduced by 5% or greater)
	Radiological evidence of disease progression (eg, increased traction
	bronchiectasis, new ground-glass opacity or fine reticulation, new/increased
	 honeycombing) Documentation of relevant fibrosis (greater than 10% fibrotic features) on chest HRCT
	Documentation of relevant fibrosis (greater than 10% fibrotic features) on chest HRC1 scan
	Baseline FVC greater than or equal to 45% of predicted
	Baseline PVC greater than or equal to 45% or predicted Baseline DLCO 30% to less than 80% of predicted
Appropriate	IPF:
Treatment	Documented treatment failure, contraindication, or intolerance to pirfenidone
	boournemed treatment failure, contraindication, or intolerance to pitteriluone
Regimen & Other	SSc-ILD:
Criteria:	



	Documented treatment failure with mycophenolate (MMF)
	Reauthorization requires documentation of treatment success
Exclusion Criteria:	Documentation of airway obstruction (such as pre-bronchodilator FEV/FVC less than 0.7)
	Combined use with pirfenidone (Esbriet)
Age Restriction:	18 years of age or older
Prescriber/Site of	Prescribed by, or in consultation with, a pulmonologist or rheumatologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: OLEZARSEN

Affected Medications: TRYNGOLZA (olezarsen sodium)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise excluded by
00 TC1 Cu 03C3.	
	plan design
	Reduce triglycerides as an adjunct to diet in adults with familial
	chylomicronemia syndrome (FCS)
Required Medical	Diagnosis of FCS (type 1 hyperlipoproteinemia) confirmed by genetic testing showing a
Information:	pathogenic gene mutation in LPL, APOC2, APOA5, GPIHBP1 or LMF1 genes
	Fasting triglyceride level of at least 880 mg/dL
	Will be used as an adjunct to diet
Appropriate	Documentation of following a low-fat diet with less than 20 grams of fat per day
Treatment	
Regimen & Other	Reauthorization requires documentation of treatment success defined as a decrease in
Criteria:	triglycerides since starting therapy
Exclusion Criteria:	History of acute coronary syndrome
Age Restriction:	18 years of age or older
Prescriber/Site of	Prescribed by, or in consultation with, a cardiologist or endocrinologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: OLIPUDASE ALFA

Affected Medications: XENPOZYME (olipudase alfa-rpcp)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded
	by plan design
	 Treatment of non-central nervous system manifestations of acid
	sphingomyelinase deficiency (ASMD) in adult and pediatric patients
Required Medical Information:	 Documentation of acid sphingomyelinase deficiency as evidenced by one of the following: Enzyme assay showing diminished (less than 10% of controls) or absent acid
	sphingomyelinase (ASM) activity
	 Gene sequencing showing biallelic pathogenic sphingomyelin phosphodiesterase-1 (SMPD1) mutation
	 Documentation of clinical presentation outside the central nervous system (e.g., hepatosplenomegaly, interstitial lung disease, liver fibrosis, growth restriction of childhood)
	Documentation of current body mass index (BMI), weight, and height
	For adults 18 years of age and older, documentation of both of the following:
	 Diffusion capacity of lungs (DLCO) is less than or equal to 70% of the predicted normal value
	 Spleen volume greater than or equal to 6 multiples of normal (MN) measured by magnetic resonance imaging (MRI)
	For pediatrics 18 years of age and younger, documentation of both of the following:
	 Spleen volume greater than or equal to 5 MN measured by MRI
	Height Z-score -1 or lower
Appropriate	Dosing: Dosed every two weeks based on FDA label
Treatment	Body mass index (BMI) less than or equal 30, the dosage is based on actual body
Regimen & Other	weight (kg)
Criteria:	BMI of greater than 30 is dosed based on adjusted body weight
	Adjusted body weight = (height in m²) x 30
	Availability: 20 mg single-dose vials
	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be
	enforced
	Reauthorization requires documentation of improvement in patient specific disease presentation such as:
	Improvement in PFT or DLCO
	Improvement in spleen and/or liver volume or function
	Improvement/stability in platelet counts
	Improvement in linear growth progression (pediatric)
Exclusion Criteria:	Exclusive central nervous system manifestations
	<u> </u>
Age Restriction:	



Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a metabolic specialist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: OMALIZUMAB

Affected Medications: XOLAIR (omalizumab)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise evaluded by
Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	Treatment of moderate to severe allergic asthma in adults and pediatric patients
	6 years of age and older
	 Add-on maintenance treatment of chronic rhinosinusitis with nasal polyps
	(CRSwNP) in adult patients
	 Treatment of symptomatic chronic spontaneous urticaria (CSU) in patients 12
	years of age and older
	 Reduction of allergic reactions (Type I), including anaphylaxis, that may occur
	with accidental exposure to one or more foods in adults and pediatric patients
B : 134 !! !	aged 1 year and older with an IgE-mediated food allergy
Required Medical	Allergic Asthma
Information:	Documentation of moderate to severe allergic asthma defined by all of the following:
	A positive skin test or in vitro reactivity to a perennial aeroallergen (e.g., house
	dust mite, animal dander [dog, cat], cockroach, feathers, mold spores)
	A serum total IgE level at baseline of: At least 20 III/rel, and least then 700 III/rel, in national 40 years of a re-
	 At least 30 IU/mL and less than 700 IU/mL in patients 12 years of age
	and older OR
	 At least 30 IU/mL and less than 1,300 IU/mL in patients 6 to 11 years of
	age ○ FEV1 less than 80% at baseline or FEV1/FVC reduced by at least 5% from
	o FEV1 less than 80% at baseline or FEV1/FVC reduced by at least 5% from normal
	CRSwNP
	Documentation of both of the following:
	 Diagnosis of chronic rhinosinusitis and has undergone prior bilateral total ethmoidectomy
	 Indicated for revision sinus endoscopic sinus surgery due to recurrent symptoms
	of nasal polyps (such as nasal obstruction/congestion, bilateral sinus obstruction)
	<u>csu</u>
	Documentation of active CSU where the underlying cause is not considered to be any
	other allergic condition or other form of urticaria
	Documentation of presence of recurrent urticaria, angioedema, or both, for a period of
	six weeks or longer
	Documented avoidance of triggers (such as nonsteroidal anti-inflammatory drugs
	[NSAIDs])
	Documented baseline score from an objective clinical evaluation tool, such as:
	 Urticaria Activity Score (UAS7) (Score of 28 or higher)
	 Urticaria Control Test (UCT)) (Score under 12)
	 Dermatology Life Quality Index (DLQI) (Score of 21 or higher)



	 Chronic Urticaria Quality of Life Questionnaire (CU-QoL) (Score of 75 or higher)
	IgE-Mediated Food Allergy
	Serum total IgE level between 30 and 1850 IU/mL
	Body weight between 10 and 150 kg
	Diagnosis of IgE-mediated food anaphylactic allergy to three or more foods with
	documented positive skin prick test and positive serum IgE
	Documentation of past IgE-mediated food anaphylactic reactions requiring use of
	epinephrine despite avoidance of food allergen and modifications to diet
	Documentation that avoidance of food allergen alone is not feasible based on the
	number of allergens, malnutrition due to nutritional restrictions, and impaired quality of
	life causing food allergy-related anxiety
Appropriate	Allergic Asthma
Treatment	Documented use of high-dose inhaled corticosteroid (ICS) plus a long-acting beta
Regimen & Other	agonist (LABA) for at least three months with continued symptoms
Criteria:	Documentation of one of the following:
	A documented history of 2 or more asthma exacerbations requiring oral or
	systemic corticosteroid treatment in the past 12 months while on combination
	inhaled treatment with at least 80% adherence.
	Documentation that chronic daily oral corticosteroids are required
	Boodinemation that officine daily of all controlled are required
	CRSwNP
	Documented treatment failure with at least 1 intranasal corticosteroid (such as
	fluticasone) after ethmoidectomy
	Documented treatment failure with Sinuva implant
	<u>CSU</u>
	Documented treatment failure with up to 4-fold standard dosing (must be scheduled) of
	one of the following second generation H1-antihistamine products for at least one month:
	cetirizine, fexofenadine, loratadine, desloratadine, or levocetirizine
	Documented treatment failure with scheduled dosing of ALL of the following for at least
	one month each:
	 Add-on therapy with a leukotriene antagonist (montelukast or zafirlukast)
	 Add-on therapy with a H2-antagonist (famotidine or cimetidine)
	 Add-on therapy with cyclosporine A
	Lot Ma Pata I Fa a I Allagon
	IgE-Mediated Food Allergy
	Trial and failure of oral immunotherapy (OIT)
	Reauthorization: documentation of treatment success and a clinically significant response
	to therapy
Exclusion Criteria:	 Use in combination with another monoclonal antibody (e.g., Fasenra, Nucala, Tezspire,
	Dupixent, Cinqair)
Age Restriction:	Allergic Asthma: 6 years of age and older
	CRSwNP: 18 years of age and older
	CSU: 12 years of age and older



Prescriber/Site of Care Restrictions:	 Allergic Asthma: prescribed by, or in consultation with, an allergist, immunologist, or pulmonologist CRSwNP: prescribed by, or in consultation with, an otolaryngologist CSU/IgE-Mediated Food Allergy: prescribed by, or in consultation with, an allergist or immunologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: OMAVELOXOLONE

Affected Medications: Skyclarys (omaveloxolone)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of Friedreich's ataxia in adults and adolescents aged 16 years and older
Required Medical	Genetically confirmed diagnosis of Friedreich's Ataxia
Information:	Documentation of baseline modified Friedreich's Ataxia Rating Scale (mFARS) score under 81
	Documentation that the patient is still ambulatory or retains enough activity to assist in activities of daily living
Appropriate	Reauthorization will require documentation of treatment success, such as a reduction in the
Treatment	rate of decline, as determined by prescriber
Regimen & Other	
Criteria:	
Exclusion Criteria:	
Age Restriction:	16 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: OMIDUBICEL

Affected Medications: OMISIRGE (Omidubicel)

Covered Uses:	 NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or better All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
Required Medical	Documentation of performance status, disease staging, all prior therapies used, and
Information:	anticipated treatment course
	Documented diagnosis of a hematologic malignancy Olivinally at the good a limit to forward little department of the property of the second state of the second s
	 Clinically stable and eligible for umbilical cord blood transplantation (UCBT) following myeloablative conditioning
Appropriate	Must NOT have a matched related donor (MRD), matched unrelated donor (MUD),
Treatment	mismatched unrelated donor (MMUD), or haploidentical donor readily available.
Regimen & Other	Documentation that NONE of the following are present:
Criteria:	 Other active malignancy
	 Active or uncontrolled infection
	 Active central nervous system (CNS) disease
	Reauthorization: None - Omisirge will be used as a one-time treatment
Exclusion Criteria:	 Karnofsky Performance Status (KPS) of 50% or less or Eastern Cooperative Oncology Group (ECOG) score of 3 or greater
	HLA (human leukocyte antigen)-matched donor able to donate
	Prior allo-HSCT (hematopoietic stem cell transplantation)
	Pregnancy or lactation
Age Restriction:	12 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 2 months for 1 time administration, unless otherwise specified
	L



ONASEMNOGENE ABEPARVOVEC XIOI

Affected Medications: ZOLGENSMA (onasemnogene abeparvovec xioi)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Spinal muscular atrophy (SMA)
Required Medical Information:	 Diagnosis of SMA type 1 confirmed by genetic testing of chromosome 5q13.2 demonstrating ONE of the following: Homozygous gene deletion of SMN1 (survival motor neuron 1) Homozygous gene mutation of SMN1 Compound heterozygous gene mutation of SMN1 Documentation of 2 or fewer copies of the SMN2 (survival motor neuron 2) gene Documentation of previous treatment history Documentation of ventilator use status: Patient is NOT ventilator-dependent (defined as using a ventilator at least 16 hours per day on at least 21 of the last 30 days) This does not apply to patients who require non-invasive ventilator assistance Documentation of anti-adeno-associated virus (AAV) serotype 9 antibody titer less than or equal 1:50 Patient weight and planned treatment regimen
Appropriate Treatment Regimen & Other Criteria:	
Exclusion Criteria:	 Prior treatment with SMA gene therapy (i.e., onasemnogene abeparvovec-xioi) Will not use in combination with other agents for SMA (e.g., nusinersen, risdiplam, etc.) Advanced SMA at baseline (complete paralysis of limbs, permanent ventilation support)
Age Restriction:	Children less than 2 years of age
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a pediatric neurologist or provider who is experienced in treatment of spinal muscular atrophy All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Approved for one dose only per lifetime, unless otherwise specified



ONCOLOGY AGENTS

Affected Medications: ABECMA, ABRAXANE, AUCATZYL, ADCETRIS, ADSTILADRIN, AKEEGA, ALECENSA, ALIQOPA, ALKERAN, ALUNBRIG 180MG ORAL TABLET, ANKTIVA, ARZERRA, ASPARLAS, AUGTYRO, AYVAKIT, AZEDRA, BALVERSA, BAVENCIO, BELEODAQ, BELRAPZO, BENDAMUSTINE, BENDEKA, BESPONSA, BIZENGRI, BLENREP, BLINCYTO, BOSULIF, BRAFTOVI, BREYANZI, BRUKINSA, CABOMETYX, CALQUENCE, CAPRELSA, CARVYKTI, CLOFARABINE, CLOLAR, COLUMVI, COMETRIQ, COPIKTRA, COSELA, COTELLIC, CYRAMZA, DACOGEN, DANZITEN, DARZALEX, DARZALEX FASPRO, DAURISMO, DOXIL, DOXORUBICIN LIPOSOMAL, ELAHERE, ELREXFIO, EMPLICITI, ENHERTU, EPKINLY, ERBITUX, ERIVEDGE, ERLEADA, ERLOTINIB, ERWINAZE, EVOMELA, FOTIVDA, FRUZAQLA, GAZYVA, GAVRETO, GEFITINIB, GILOTRIF, HEPZATO, HYCAMTIN, IBRUTINIB, ICLUSIG, IDHIFA, IMBRUVICA, IMDELLTRA, IMFINZI, IMJUDO, IMLYGIC IRESSA, INLYTA, INQOVI, INREBIC, IOBENGUANE I-131, ISTODAX, ITOVEBI, IXEMPRA, JAKAFI, JAYPIRCA, JELMYTO, JEMPERLI, JEVTANA, KADCYLA, KEYTRUDA, KIMMTRAK, KISQALI, KISQALI FEMARA, KRAZATI, KYMRIAH, KYPROLIS, LAPATINIB, LARTRUVO, LENALIDOMIDE, LENVIMA, LIBTAYO, LONSURF, LOQTORZI, LORBRENA, LUMAKRAS, LUMOXITI, LUNSUMIO, LUTATHERA, LYNPARZA, LYTGOBI, MARGENZA, MARQIBO, MATULANE, MEKINIST, MEKTOVI, MELPHALAN, MONJUVI, MYLOTARG, NAB-PACLITAXEL, NEXAVAR, NERLYNX, NILANDRON, NINLARO, NIVOLUMAB, NUBEQA, ODOMZO, OJEMDA, OJJAARA, ONCASPAR, ONIVYDE, ONUREG, OPDIVO, OPDIVO QVANTIG, OPDUALAG, ORSERDU, PADCEV, PAZOPANIB, PEMAZYRE, PEPAXTO, PERJETA, PHOTOFRIN, PIQRAY, PLUVICTO, POLIVY, POMALYST, POTELIGEO, PROLEUKIN, PROVENGE, QINLOCK, RETEVMO, REVLIMID, REVUFORJ, REZLIDHIA, REZUROCK, ROMIDEPSIN, ROZLYTREK, RUBRACA, RYBREVANT, RYDAPT, RYLAZE, RYTELO, SARCLISA, SORAFENIB, STIVARGA, SUNITINIB, SUTENT, SYNRIBO, TABRECTA, TAFINLAR, TAGRISSO, TALVEY, TALZENNA, TARCEVA, TAZVERIK, TECARTUS, TECELRA, TECENTRIQ, TECENTRIQ HYBREZA, TECVAYLI, TEMODAR, TEMOZOLOMIDE, TEPADINA, TEPMETKO, TEVIMBRA, TIBSOVO, TIVDAK, TORISEL, TREANDA, TRODELVY, TRUQAP, TURALIO, TYKERB, VANFLYTA, VECTIBIX, VENCLEXTA, VERZENIO, VIDAZA, VIVIMUSTA, VIZIMPRO, VONJO, VORANIGO, VOTRIENT, VYXEOS, XALKORI, XOFIGO, XOSPATA, XPOVIO, XTANDI, YERVOY, YESCARTA, YONDELIS, ZALTRAP, ZEJULA TABLETS, ZELBORAF, ZEPZELCA, ZOLINZA, ZYDELIG, ZYKADIA, ZYNLONTA, ZYNYZ

Covered Uses:	NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher
Required Medical Information:	Documentation of performance status, disease staging, all prior therapies used, and anticipated treatment course
Appropriate Treatment Regimen & Other Criteria:	Reauthorization: documentation of disease responsiveness to therapy
Exclusion Criteria:	Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



OPIOID QUANTITY ABOVE 90 MORPHINE MILLIGRAM EQUIVALENTS (MME)

Affected Medications: ALL OPIOIDS

Covered Uses:	All Food and Drug Administration (FD) plan design	A)-approved indications not otherwise excluded by
Required Medical Information:	following: Recent surgery Acute injury Chronic use of opioids with a Morphine 90 MME requires: A comprehensive individual tremanagement agreement betw Continued assessment and do	apers have been attempted or documentation of a
Appropriate	Calculating morphine milligram equival	
Treatment	[autorial	Forton
Regimen & Other Criteria:	Opioid Methadone	Factor 4.7
Orneria.	Codeine	0.15
	Fentanyl transdermal (mcg/hr)	2.4
	Hydrocodone	1
	Hydromorphone	5
	Morphine	1
	Oxycodone (Roxicodone, Oxycontin)	1.5
	Oxymorphone	3
	Tramadol	0.2
	Buprenorphine patch	**
	Tapentadol	0.4
	Oxycodone myristate	1.67
	 One milligram of parenteral buprenorp and One patch delivers the dispensed micro Example: 5 mcg/hr buprenorphine patch X 24 hr 	rphine patches is based on the assumption that: hine is equivalent to 75 milligrams of oral morphine rograms (mcg) per hour over a 24-hour day. s = 120 mcg/day buprenorphine = 0.12 mg/day phine=75 mg morphine) = 9 mg/day oral MME.



	In other words, the conversion factor not accounting for days of use would be 9/5 or 1.8.
	• Since the buprenorphine patch remains in place for 7 days, we have multiplied the conversion factor by 7 (1.8 X 7 = 12.6). In this example, MME/day for four 5 mcg/hr buprenorphine patches dispensed for use over 28 days would work out as follows:
	Example: 5 mcg/hr buprenorphine patch X (4 patches/28 days) X 12.6 = 9 MME/day.
	Please note that because this allowance has been made based on the typical dosage of one buprenorphine patch per 7 days. You should first change all days supply in your prescription data to follow this standard, i.e., days supply for buprenorphine patches= # of patches x 7
Exclusion Criteria:	Pain related to current active cancer
	Chronic pain related to sickle cell disease
	Pain related to hospice care
	Surgery or documented acute injury – 1 month approval
Age Restriction:	
Prescriber/Site of	All approvals are subject to utilization of the most cost-effective site of care
Care Restrictions:	
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: OPZELURA

Affected Medications: OPZELURA CREAM (1.5%)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Atopic dermatitis
Required Medical Information:	 Documentation of affected body surface area (BSA) and areas of involvement Documentation of severe atopic dermatitis, resulting in functional impairment as defined by one of the following: Inability to use hands or feet for activities of daily living Significant facial involvement preventing normal social interaction Documentation of one or more of the following: BSA of at least 10% Hand, foot, or mucous membrane involvement
Appropriate Treatment Regimen & Other Criteria:	 Documented treatment failure with a minimum 6-week trial with two of the following: tacrolimus ointment, pimecrolimus cream, Eucrisa Documented treatment failure with a minimum 12-week trial of two of the following: phototherapy, cyclosporine, azathioprine, methotrexate, mycophenolate Documented treatment failure with a minimum 12-week trial with each of the following: Dupixent, Adbry Reauthorization: No reauthorization permitted.
Exclusion Criteria:	 Combined use with a biologic or Janus kinase (JAK) inhibitor Previous 8-week treatment course Cosmetic indications, such as vitiligo
Age Restriction:	12 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a dermatologist, allergist, or immunologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 8 weeks (no reauthorization), unless otherwise specified.



ORAL-INTRANASAL FENTANYL

Affected Medications: ABSTRAL, ACTIQ, FENTORA, FENTANYL CITRATE BUCCAL TABLET, LAZANDA, SUBSYS, FENTANYL CITRATE LOZENGE ON A HANDLE

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Management of breakthrough pain in cancer patients who are already receiving and who are tolerant to around-the-clock opioid therapy for their underlying persistent cancer pain
Required Medical Information:	 Documentation of ALL of the following: This drug is being prescribed for breakthrough cancer-related pain The patient is currently receiving, and will continue to receive, around-the-clock opioid therapy for underlying persistent cancer pain The patient is opioid tolerant, defined as taking one of the following for one week or longer:
Appropriate Treatment Regimen & Other Criteria:	 An equianalgesic dose of another opioid Documentation of ONE of the following: The patient is unable to swallow, or has dysphagia, esophagitis, mucositis, or uncontrollable nausea/vomiting The patient has documented intolerance or allergies to two other short-acting narcotics (such as oxycodone, morphine sulfate, hydromorphone, etc.) PDL only: Actiq requests will require documentation of clinical trial and failure with fentanyl citrate lozenge on a handle Reauthorization requires documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an oncologist or specialist in the treatment of cancer-related pain All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: ORENITRAM

Affected Medications: ORENITRAM (Treprostinil oral)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	 Pulmonary Arterial Hypertension (PAH) World Health Organization (WHO) Group 1
Required	Pulmonary Arterial Hypertension (PAH) WHO Group 1
Medical Information:	Documentation of PAH confirmed by right-heart catheterization meeting the following criteria:
	 Pulmonary capillary wedge pressure less than or equal to 15 mm Hg Pulmonary vascular resistance of at least 2.0 Wood units
	Etiology of PAH: idiopathic, heritable, or associated with connective tissue disease
	PAH secondary to one of the following conditions:
	 Connective tissue disease
	 Human immunodeficiency virus (HIV) infection
	o Cirrhosis
	 Anorexigens
	 Congenital left to right shunts
	 Schistosomiasis
	 Drugs and toxins
	 Portal hypertension
	New York Heart Association (NYHA)/World Health Organization (WHO) Functional Class
	II or higher symptoms
	Documentation of Acute Vasoreactivity Testing (positive result requires trial/failure to
	calcium channel blockers) unless there are contraindications:
	 Low systemic blood pressure (systolic blood pressure less than 90)
	 Low cardiac index
	OR
	 Presence of severe symptoms (functional class IV)
Appropriate	Documentation of failure with Remodulin
Treatment Regimen &	The pulmonary hypertension has progressed despite maximal medical and/or surgical treatment of the identified condition
Other Criteria:	 Documentation that treprostinil is used as a single route of administration (Remodulin, Tyvaso, Orenitram should not be used in combination)
	 Not recommended for PAH secondary to pulmonary venous hypertension (e.g., left sided atrial or ventricular disease, left sided valvular heart disease, etc) or disorders of the respiratory system (e.g., chronic obstructive pulmonary disease, interstitial lung disease, obstructive sleep apnea or other sleep disordered breathing, alveolar hypoventilation disorders, etc.)
	Reauthorization requires documentation of treatment success defined as one or more of the following:
	Improvement in walking distance
	Improvement in exercise ability
	Improvement in pulmonary function
	Improvement or stability in WHO functional class
	,



Exclusion Criteria:	Severe hepatic impairment (Child Pugh Class C)
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a cardiologist or pulmonologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified.



POLICY NAME: ORGOVYX

Affected Medications: ORGOVYX (relugolix)

Covered Uses:	NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher
Required Medical Information:	
Appropriate Treatment Regimen & Other Criteria:	Prostate Cancer Documented treatment failure or intolerable adverse event with leuprolide or degarelix Reauthorization: documentation of disease responsiveness to therapy
Exclusion Criteria:	Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an oncologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: OSILODROSTAT

Affected Medications: ISTURISA (osilodrostat)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by	
	plan design	
	Cushing's disease	
Required Medical	Documented diagnosis of Cushing's disease	
Information:	Documentation of at least TWO of the following:	
	 Mean (at least two measurements) 24-hour urine free cortisol (mUFC) greater 	
	than 1.5 times the upper limit of normal (ULN) for the assay	
	 Bedtime salivary cortisol (at least two measurements) greater than 145 ng/dL 	
	 Overnight dexamethasone suppression test (DST) with a serum cortisol greater than 1.8 mcg/dL 	
Appropriate	Documentation confirming pituitary surgery is not an option OR previous surgery has not	
Treatment	been curative	
Regimen & Other		
Criteria:	Reauthorization Reauthorization requires documentation of treatment success defined as mUFC normalization (i.e., less than or equal to the ULN)	
Exclusion Criteria:		
Age Restriction:	18 years of age and older	
Prescriber/Site of	Prescribed by, or in consultation with, an endocrinologist, neurologist, or adrenal surgeon	
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care	
Coverage Duration:	Authorization: 12 months, unless otherwise specified	



POLICY NAME: OTESECONAZOLE

Affected Medications: VIVJOA (oteseconazole)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design To reduce the incidence of recurrent vulvovaginal candidiasis (RVVC) in females with a history of RVVC who are not of reproductive potential, alone or in combination with fluconazole
Required Medical Information:	 Diagnosis of RVVC defined as three or more episodes of symptomatic vulvovaginal candidiasis infection within the past 12 months Documented presence of signs/symptoms of current acute vulvovaginal candidiasis with a positive potassium hydroxide (KOH) test Documentation confirming that the patient is permanently infertile (e.g., due to tubal ligation, hysterectomy, salpingo-oophorectomy) or postmenopausal
Appropriate Treatment Regimen & Other Criteria:	 Documented disease recurrence following 10 to 14 days of induction therapy with a topical antifungal agent or oral fluconazole, followed by fluconazole 150 mg once per week for 6 months Not to exceed one treatment course per year Reauthorization requires documentation of treatment success defined as a reduction in symptomatic vulvovaginal candidiasis episodes, and documentation supporting the need for
Exclusion Criteria:	Women of reproductive potential or who are pregnant or breastfeeding
Age Restriction:	18 years of age and older
Prescriber/Site of Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 3 months, unless otherwise specified



POLICY NAME: **OXERVATE**

Affected Medications: OXERVATE (cenegermin-bkbj)

	T		
Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by		
	plan design		
	 Treatment of neurotrophic keratitis 		
Required	 Documentation of decreased corneal sensitivity (≤ 4 cm using the Cochet-Bonnet [CB] 		
Medical	aesthesiometer) within the area of the recurrent/persistent epithelial defect (PED) or		
Information:	corneal ulcer AND outside of the area of the defect in at least one corneal quadrant		
	Documentation of one of the following:		
	 Stage 2 neurotrophic keratitis, confirmed by presence of recurrent or persistent 		
	corneal epithelial defect		
	 Stage 3 neurotrophic keratitis, confirmed by presence of corneal ulceration (with 		
	or without stromal melting and perforation)		
Appropriate	Documentation of treatment failure (e.g., persistent epithelial defects or corneal		
Treatment	ulceration) with preservative-free artificial tears/ointments and TWO of the following:		
Regimen &	 Therapeutic contact lenses (TCLs) (e.g., corneal or scleral contact lenses, soft 		
Other Criteria:	bandage contact lenses)		
	,		
	 Amniotic membrane transplantation 		
	 Tarsorrhaphy 		
	Conjunctival flap surgery		
	Dose may not exceed more than 1 vial per eye per day		
	<u>Reauthorization</u> requires documentation of treatment response, as shown by a reduction in corneal staining with fluorescein		
Exclusion	Active or suspected ocular or periocular infections		
Criteria:	, tours or edoposion or periodian initiation		
Age			
Restriction:			
Prescriber/Site of	Prescribed by, or in consultation with, an ophthalmologist		
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care		
Coverage	Initial Authorization: 8 weeks, unless otherwise specified		
Duration:	Reauthorization: 8 weeks, unless otherwise specified		
	Lifetime Limit: 16 weeks (per affected eye)		
	Lifetime Limit. To weeks (per anected eye)		



OXYBATES

Affected Medications: LUMRYZ (sodium oxybate extended release), XYREM (sodium oxybate), XYWAV (oxybate salts), SODIUM OXYBATE

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Narcolepsy with cataplexy Narcolepsy with excessive daytime sleepiness (EDS) Idiopathic Hypersomnia (IH) (Xywav only)
Required Medical Information:	 Diagnosis confirmed by polysomnography and multiple sleep latency test Other causes of sleepiness have been ruled out or treated (including but not limited to obstructive sleep apnea, insufficient sleep syndrome, shift work, the effects of substances or medications, or other sleep disorders)
	Narcolepsy with cataplexy: Documentation of cataplexy episodes defined as more than one episode of sudden loss of muscle tone with retained consciousness
	 Narcolepsy with EDS or IH: Current evaluation of symptoms and Epworth Sleepiness Scale (ESS) score of more than 10 despite treatment
Appropriate	Narcolepsy with cataplexy:
Treatment	Documented treatment failure with TWO of the following for at least 1 month each:
Regimen & Other Criteria:	 Venlafaxine Fluoxetine Duloxetine
	 Tricyclic antidepressant (such as clomipramine, protriptyline)
	Narcolepsy or IH, with EDS: Documented treatment failure to all of the following (1 in each category required) for at least 1 month each: Modafinil or armodafinil Methylphenidate, or dextroamphetamine, or lisdexamfetamine Sunosi (Narcolepsy with EDS only)
	 Reauthorization: Narcolepsy with cataplexy: requires clinically significant reduction in cataplexy episodes Narcolepsy or IH, with EDS: requires clinically significant improvement in activities of daily living and in Epworth Sleepiness Scale (ESS) score
Exclusion Criteria:	Concurrent use of alcohol, sedative/hypnotic drugs, or other central nervous system depressants. Use for other untreated causes of sleepings.
Age Restriction:	 Use for other untreated causes of sleepiness 7 years of age and older for cataplexy or EDS due to narcolepsy
. 130 11001110110111	18 years of age and older for EDS due to Harcolepsy 18 years of age and older for EDS due to IH



Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a sleep specialist or neurologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 4 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **OZANIMOD**

Affected Medications: ZEPOSIA (ozanimod)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design: Treatment of relapsing forms of multiple sclerosis (MS), including the following: Clinically isolated syndrome (CIS) Relapsing-remitting multiple sclerosis (RRMS) Active secondary progressive disease (SPMS) Ulcerative Colitis
Required	Multiple Sclerosis
Medical	Diagnosis confirmed with magnetic resonance imaging (MRI), per revised McDonald
Information:	diagnostic criteria for MS Clinical evidence alone will suffice; additional evidence desirable but must be consistent with MS
	Illegrative Colitie
	Ulcerative Colitis Diagnosis supported by endoscopy/colonoscopy/sigmoidoscopy or biopsy with moderate to severely active disease despite current treatment
Appropriate	Relapsing forms of MS
Treatment	Coverage of Zeposia (ozanimod) requires documentation of one of the following:
Regimen &	 Documented disease progression or intolerable adverse event with one of the
Other Criteria:	following: teriflunomide, dimethyl fumarate or fingolimod
	Currently receiving treatment with Zeposia (ozanimod), excluding via samples or
	manufacturer's patient assistance program
	 Ulcerative Colitis Documented failure with at least two oral treatments for a minimum of 12 weeks each: corticosteroids, sulfasalazine, azathioprine, mesalamine, balsalazide, cyclosporine, 6-mercaptopurine AND
	 Documented treatment failure with or intolerable adverse event with all preferred pharmacy drugs (Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Xeljanz, Stelara, Rinvoq)
	Reauthorization requires provider attestation of treatment success
Exclusion	MS: concurrent use of other disease-modifying medications indicated for the treatment of
Criteria:	multiple sclerosis
	UC: concurrent use with a JAK inhibitor or biologic medication for the treatment of ulcerative colitis
Age Restriction:	
Prescriber/Site of	MS: prescribed by, or in consultation with, a neurologist or a multiple sclerosis specialist
Care Restrictions:	UC: prescribed by, or in consultation with, a gastroenterologist
	1 • Oo. prosonibed by, or in consultation with, a gastrochicrologist



Coverage Duration:	Initial Authorization:	
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POLICY NAME: PALFORZIA

Affected Medications: PALFORZIA (peanut allergen powder)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
Covered Oses.	· , , , , , , , , , , , , , , , , , , ,
	plan design
	 Mitigation of allergic reactions, including anaphylaxis, that may occur with accidental exposure to peanut
Required Medical	Documented treatment plan, including dose and frequency
Information:	Diagnosis of peanut allergy confirmed by one of the following:
	A positive skin prick test (SPT) response to peanut with a wheal diameter at
	least 3 mm larger than the control
	 Serum peanut-specific IgE level greater than or equal to 0.35 kUA/L
	Documented history of an allergic reaction to peanut with all of the following:
	 Signs and symptoms of a significant systemic allergic reaction to peanut (e.g.,
	hives, swelling, wheezing, hypotension, gastrointestinal symptoms)
	The reaction occurred within a short period of time following a known ingestion
	of peanut or peanut-containing food
	The reaction was severe enough to warrant a prescription for an epinephrine
	injection
	Documentation indicating a significant impact on quality of life due to peanut allergies
Appropriate	Dosing:
Treatment	Requests for initial dose escalation: must be between 1 and 17 years of age
Regimen & Other	Requests for up-dosing and maintenance phase: 1 year of age and older
Criteria:	
	Reauthorization requires documentation of completion of the appropriate initial dose
	escalation and up-dosing phases prior to moving on to the maintenance phase AND
	documentation of treatment success and a clinically significant response to therapy, defined
	by one or more of the following:Improvement in quality of life
	Reduction in severe allergic reactions Reduction in eninephrine uses
	Reduction in epinephrine use Reduction in physician office visite. ER visite or been talligations due to people ellers.
Exclusion Criteria:	 Reduction in physician office visits, ER visits, or hospitalizations due to peanut allergy Use for the emergency treatment of allergic reactions, including anaphylaxis
Exclusion officia.	
	History of eosinophilic esophagitis (EoE) and other eosinophilic gastrointestinal disease History of conditionable disease, including uncentralled or including uncentralled.
	History of cardiovascular disease, including uncontrolled or inadequately controlled bypartonian
	hypertension
	History of a mast cell disorder, including mastocytosis, urticarial pigmentosa, and hereditary or idiopathic angioedema
Age Restriction:	1 year of age and older (see Appropriate Treatment Regimen & Other Criteria for specific
	age-related dosing requirements)
Prescriber/Site of	Prescribed by, or in consultation with, an allergist or immunologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care



Coverage Duration:	•	Initial Authorization: 6 months, unless otherwise specified
	•	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: PALIVIZUMAB

Affected Medications: SYNAGIS (palivizumab)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by Plan design.
	plan design.
Required	Documentation of one of the following conditions:
Medical	Congenital heart disease (CHD):
Information:	a. With cardiac transplantation, cardiac bypass, or extra-corporeal membrane
	oxygenation
	 That is hemodynamically significant (e.g., acyanotic heart disease, congestive heart failure, or moderate to severe pulmonary hypertension)
	2. Chronic lung disease (CLD) of prematurity:
	 a. In the first year of life, born less than 32 weeks gestation and requiring greater that 21% oxygen for at least the first 28 days of life
	b. In the second year of life necessitating continued medical support within the 6
	month period prior to RSV season (e.g. corticosteroids, diuretics, supplemental oxygen)
	3. Cystic Fibrosis and:
	a. Clinical evidence of CLD and/or nutritional compromise
	b. Severe lung disease (e.g., previous hospitalization for pulmonary exacerbation
	in the first year of life or abnormalities on chest radiography or computed
	tomography that persist when stable)
	c. A weight for length less than the 10 th percentile
	4. Congenital airway abnormality or neuromuscular condition (not cystic fibrosis) that
	impairs the ability to clear airway secretions
	5. Premature infants without above conditions
Appropriate	Prevention of serious lower respiratory tract disease caused by respiratory syncytial
Treatment	virus (RSV)
Regimen &	The first dose of Synagis should be administered prior to commencement of the RSV
Other Criteria:	season
	Remaining doses should be administered monthly throughout the RSV season
	(Exception: dose administration should occur immediately post cardiopulmonary bypass
	surgery, even if dose is administered earlier than a month from previous dose, then
	dosing schedule should resume monthly)
	No more than 5 monthly doses During the RSV season, November 1 through March 31
	Discontinue prophylaxis therapy if hospitalized for RSV
Exclusion	For use in the treatment of RSV disease
Criteria:	Received Beyfortus during the current RSV season
Age	Refer to numbered conditions above in "Required Medical Information":
Restriction:	1a. Less than 2 years of age 1b. Less than 1 year of age 1c. The Less than 1 years of age 1c. The Less th
	1b. Less than 1 year of age 2c. Less than 1 year of age: Costational Age less than 22 weeks.
	2a. Less than 1 year of age; Gestational Age less than 32 weeks 3b. Less than 3 years of age; Costational Age less than 32 weeks
	2b. Less than 2 years of age; Gestational Age less than 32 weeks 3a Less than 1 year of age.
	3a. Less than 1 year of age 3b. Less than 2 years of age
	 3b. Less than 2 years of age 3c. Less than 2 years of age
	3c. Less than 2 years of age4. Less than 1 year of age
	● 4. Less man i year or age



	5. Less than 1 year of age; Gestational Age less than 29 weeks	
Prescriber/Site of	All approvals are subject to utilization of the most cost-effective site of care	
Care Restrictions:		
Coverage	Authorization:	
Duration:	• 5 months (November 1 through March 31) [5 monthly doses], unless otherwise specified	
	1 month for off-season when RSV activity greater than or equal to 10% for the region	
	according to the CDC [1 monthly dose], unless otherwise specified	



POLICY NAME: PALOVAROTENE

Affected Medications: SOHONOS (palovarotene)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
Oovered Oses.	plan design
	To reduce the volume of new heterotopic ossification in patients with
	fibrodysplasia ossificans progressiva (FOP)
Required Medical	Documentation of genetic testing confirming a diagnosis of FOP with an activin receptor
Information:	type 1 (ACVR1) R206H mutation
	Radiographic testing has confirmed the presence of both of the following:
	 Heterotopic ossification (HO)
	 Joint malformations (such as hallux valgus deformity, malformed first metatarsal,
	absent or fused interphalangeal joint)
	Documentation of at least two flare-ups in the past 12 months requiring prescription
	strength non-steroidal anti-inflammatory drugs (NSAIDs) and oral glucocorticoids (e.g.,
	prednisone)
Appropriate	Reauthorization requires documentation of treatment success defined as a decrease in HO
Treatment	volume or number of flare-ups compared to baseline
Regimen & Other	
Criteria:	
Exclusion Criteria:	Patients weighing less than 10 kg
	Pregnancy
Age Restriction:	Females: 8 years of age and older
	Males: 10 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a specialist in rare connective tissue diseases
Care Restrictions:	(e.g., endocrinologist, geneticist, orthopedist, rheumatologist)
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified
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POLICY NAME: **PALYNZIQ**

Affected Medications: PALYNZIQ (pegvaliase-pqpz)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Reduce phenylalanine (Phe) blood concentrations in adults with phenylketonuria (PKU) who have uncontrolled blood Phe greater than 600 micromol/L on existing management
Required Medical Information:	 Documentation of a diagnosis of PKU Documentation of treatment failure with dual therapy of sapropterin and a Phe restricted diet as shown by a blood Phe level greater than 600 micromol/L (10 mg/dL) despite treatment
Appropriate Treatment Regimen & Other Criteria:	 Documentation that Palynziq will not be used in combination with sapropterin Reauthorization requires documentation of one of the following: Reduction in baseline Phe levels by 20 percent Increase in dietary Phe tolerance Improvement in clinical symptoms
Exclusion Criteria:	
Age Restriction:	18 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a specialist in metabolic disorders or an endocrinologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 3 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



PARATHYROID HORMONE

Affected Medications: YORVIPATH (palopegteriparatide)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise evaluated by		
OUVEIEU USES.	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design.		
	plan design		
	Treatment of hypoparathyroidism		
Required	Documentation of the following lab values while on standard of care calcium and active		
Medical	vitamin D treatment:		
Information:	 25-hydroxyvitamin D levels between 20-80 ng/mL 		
	 Total serum calcium (albumin-corrected) greater than 7.8 mg/dL 		
Appropriate	Documented failure with at least 12 weeks of a consistent supplementation regimen as		
Treatment	follows:		
Regimen &	 Calcium 1000-2000 mg (elemental) daily 		
Other Criteria:	 Vitamin D metabolite (calcitriol) OR vitamin D analog 		
	Reauthorization will require documentation of treatment success defined as total serum		
	calcium (albumin-corrected) within the lower half of the normal range (approximately 8-9		
	mg/dL)		
Exclusion			
Criteria:			
Age	18 years of age and older		
Restriction:			
Prescriber/Site of	Prescribed by, or in consultation with, an endocrinologist or nephrologist		
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care		
Coverage	Initial Authorization: 6 months, unless otherwise specified		
Duration:	Reauthorization: 12 months, unless otherwise specified		
	1 Modulion 2 months, unless otherwise specified		



PARATHYROID HORMONE ANALOGS

Affected Medications: TERIPARATIDE, TYMLOS (abaloparatide), FORTEO (teriparatide)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of osteoporosis in men and postmenopausal women at high risk for fracture (teriparatide, Tymlos, and Forteo) Treatment of glucocorticoid-induced osteoporosis in men and women at high risk for fracture (teriparatide and Forteo only)
Required Medical Information:	 Diagnosis of osteoporosis as defined by at least one of the following: T-score –2.5 or lower (current or past) at the lumbar spine, femoral neck, total hip, or 1/3 radius site T-score between –1.0 and –2.5 at the lumbar spine, femoral neck, total hip, or 1/3 radius site AND increased risk of fracture as defined by at least one of the following Fracture Risk Assessment Tool (FRAX) scores:
Appropriate Treatment Regimen & Other Criteria:	Documentation of one of the following: Treatment failure (new fracture or worsening T-score despite adherence to an adequate trial of therapy), contraindication, or intolerance to BOTH of the following: Oral or intravenous bisphosphonate (such as, alendronate, risedronate, zoledronic acid or ibandronate) Prolia (denosumab) High risk of fracture defined as T-score -3.5 or lower, OR T-score -2.5 or lower with a history of fragility fractures
	For Forteo requests: Documented treatment failure with Tymlos and teriparatide Total duration of therapy with parathyroid hormone analogs should not exceed 2 years in a lifetime • Forteo or teriparatide may be reauthorized for up to one additional year beyond two years of parathyroid hormone analog use (maximum of 3 total years) if meeting the following criteria: • Documentation of treatment success with parathyroid hormone use, defined as reduced frequency of fragility fractures or stable T-score while on Forteo or teriparatide • Documentation that after 24 months of parathyroid hormone analog use, the patient remains at or has returned to having a high risk for fracture as evidenced
Exclusion	 by new fragility fracture or decline in T-score Paget's Disease



Criteria:	 Open epiphyses (such as, pediatric or young adult patient) Bone metastases or skeletal malignancies Hereditary disorders predisposing to osteosarcoma Prior external beam or implant radiation therapy involving the skeleton Concurrent use of bisphosphonates, other parathyroid hormone analogs, or RANK ligand inhibitors Preexisting hypercalcemia Pregnancy
Age Restriction:	
Prescriber/Site of Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 24 months (no reauthorization), unless otherwise specified



POLICY NAME: **PEDMARK**

Affected Medications: PEDMARK (sodium thiosulfate)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design To reduce the risk of ototoxicity associated with cisplatin in pediatric patients 1 month of age and older with localized, non-metastatic solid tumors.
Required Medical Information:	Documentation of a treatment plan that is a cisplatin-based regimen treating a localized, non-metastatic solid tumor
Appropriate Treatment Regimen & Other Criteria:	
Exclusion Criteria:	Metastatic disease
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an oncologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 6 months or duration of cisplatin regimen, unless otherwise specified



POLICY NAME: **PEGASYS**

Affected Medications: PEGASYS

Covered Uses:		d and Drug Administration (FDA)-a erwise excluded by plan design	oproved and compendia-supported in	ndications
Required Medical Information:	Documentation of anticipated treatment course, to include full antiviral regimen, and duration of therapy			
	 Chronic Hepatitis C (CHC): Documentation chronic hepatitis C virus (HCV) genotype by liver biopsy or by Food and Drug Administration (FDA)-approved serum test Baseline HCV RNA level 			
	 Chronic Hepatitis B (CHB): Documentation of HBeAg-positive or HBeAg-negative chronic hepatitis B virus (HBV) infection Baseline HBV DNA level Current (within 12 weeks) alanine transaminase (ALT) level 			
	 Chronic Hepatitis C and B: Current documentation of hepatic impairment severity with Child-Pugh Classification OR bilirubin, albumin, INR, ascites status, and encephalopathy status to calculate Child-Pugh score within 12 weeks prior to anticipated start of therapy Documentation if HIV/HCV/HBV coinfection 			
	Chronic Hepatitis C: Approve if used in combination with Food and Drug Administration (FDA)- and/or AASLD/IDSA- recommended regimen and if not otherwise excluded from PacificSource policies of other medications in the regimen			
Appropriate Treatment Regimen & Other Criteria:	Approve AASLD	e if used in combination with Food /IDSA- recommended regimen and	if not otherwise excluded from Pacif	
Treatment Regimen & Other	Approve AASLD policies Chronic He	e if used in combination with Food /IDSA- recommended regimen and of other medications in the regime	if not otherwise excluded from Pacif	
Treatment Regimen & Other	Approve AASLD policies Chronic He	e if used in combination with Food /IDSA- recommended regimen and of other medications in the regime	if not otherwise excluded from Pacif	
Treatment Regimen & Other	 Approve AASLD, policies Chronic He Docume 	e if used in combination with Food /IDSA- recommended regimen and of other medications in the regime epatitis B: entation of ONE of the following some HBV DNA	if not otherwise excluded from Pacifn	
Treatment Regimen & Other	 Approve AASLD, policies Chronic He Docume HBeAg 	e if used in combination with Food /IDSA- recommended regimen and of other medications in the regime epatitis B: entation of ONE of the following some HBV DNA	if not otherwise excluded from Pacifn	
Treatment Regimen & Other	 Approve AASLD, policies Chronic He Docume HBeAg Without c 	e if used in combination with Food /IDSA- recommended regimen and of other medications in the regime epatitis B: entation of ONE of the following so HBV DNA irrhosis	if not otherwise excluded from Pacifin enarios: ALT Greater than 2 times the upper	
Treatment Regimen & Other	 Approve AASLD, policies Chronic He Docume HBeAg Without c 	e if used in combination with Food /IDSA- recommended regimen and of other medications in the regime epatitis B: entation of ONE of the following sc HBV DNA irrhosis Greater than 20,000 copies/mL	if not otherwise excluded from Pacifin enarios: ALT Greater than 2 times the upper limit of normal (ULN)	
Treatment Regimen & Other	 Approve AASLD, policies Chronic He Docume HBeAg Without c Positive Negative 	e if used in combination with Food /IDSA- recommended regimen and of other medications in the regime epatitis B: entation of ONE of the following sci HBV DNA irrhosis Greater than 20,000 copies/mL Greater than 2,000 copies/mL	enarios: ALT Greater than 2 times the upper limit of normal (ULN) Greater than 2 times the ULN 1-2 times the ULN and moderate/severe liver	
Treatment Regimen & Other Criteria:	 Approve AASLD, policies Chronic He Docume HBeAg Without c Positive Negative 	e if used in combination with Food /IDSA- recommended regimen and of other medications in the regime epatitis B: entation of ONE of the following screen HBV DNA irrhosis Greater than 20,000 copies/mL Greater than 2,000 copies/mL Greater than 2,000 copies/mL	enarios: ALT Greater than 2 times the upper limit of normal (ULN) Greater than 2 times the ULN 1-2 times the ULN and moderate/severe liver	
Treatment Regimen & Other	Approve AASLD policies Chronic He Docume HBeAg Without c Positive Negative Negative With com Either Autoimn	e if used in combination with Food /IDSA- recommended regimen and of other medications in the regime epatitis B: entation of ONE of the following screen HBV DNA irrhosis Greater than 20,000 copies/mL Greater than 2,000 copies/mL Greater than 2,000 copies/mL pensated cirrhosis Greater than 2,000 copies/mL ent of patients with CHC who have mune hepatitis	Greater than 2 times the upper limit of normal (ULN) Greater than 2 times the ULN 1-2 times the ULN and moderate/severe liver inflammation/fibrosis Any ALT had solid organ transplantation	
Treatment Regimen & Other Criteria:	Approve AASLD, policies Chronic He Docume HBeAg Without c Positive Negative With com Either Autoimr Hepatic	e if used in combination with Food /IDSA- recommended regimen and of other medications in the regime epatitis B: entation of ONE of the following sci HBV DNA irrhosis Greater than 20,000 copies/mL Greater than 2,000 copies/mL Greater than 2,000 copies/mL pensated cirrhosis Greater than 2,000 copies/mL ent of patients with CHC who have	Greater than 2 times the upper limit of normal (ULN) Greater than 2 times the ULN 1-2 times the ULN and moderate/severe liver inflammation/fibrosis Any ALT had solid organ transplantation	



Prescriber/Site of	Prescribed by, or in consultation with, a gastroenterologist, hepatologist, or infectious
Care Restrictions:	disease specialist
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	CHC: 12 weeks, unless otherwise specified (depends on regimen and diagnosis)
	CHB: 12 months, unless otherwise specified



POLICY NAME: **PEGLOTICASE**

Affected Medications: KRYSTEXXA (pegloticase)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Chronic gout in adults refractory to conventional therapy
Required Medical Information:	Baseline serum uric acid (SUA) level greater than 8 mg/dL Documentation of ONE of the following:
Appropriate Treatment Regimen & Other Criteria:	 Documented contraindication, intolerance or clinical failure (defined as inability to reduce SUA level to less than 6 mg/dL) following a 12-week trial at maximum tolerated dose to BOTH: Xanthine oxidase inhibitor (allopurinol or febuxostat) Combination of a xanthine oxidase inhibitor AND a uricosuric agent (such as probenecid). If xanthine oxidase inhibitor is contraindicated, trial with uricosuric agent required. Documentation Krystexxa will be used in combination with oral methotrexate 15 mg weekly unless contraindicated Reauthorization will require ALL of the following: Documentation of SUA less than 6 mg/dL prior to next scheduled Krystexxa dose Documentation of response to treatment such as reduced size of tophi or number of flares or affected joints Rationale to continue treatment after resolution of tophi or reduction in symptoms
Exclusion Criteria:	Concurrent use with oral urate-lowering therapies
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a nephrologist or rheumatologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 6 months, unless otherwise specified



POLICY NAME: **PEMIVIBART**

Affected Medications: PEMGARDA (pemivibart)

Covered Uses:	All Food and Drug Administration (FDA) or compendia supported indications not
Covered Oses.	otherwise excluded by plan design
	,, ,
	 Pre-exposure prophylaxis (PrEP) of coronavirus disease 2019 (COVID-19) in moderate-to-severe immune compromised individuals 12 years of age and older
	weighing at least 40 kg
Required Medical	Documentation of moderate-to-severe immune compromise due to a medical condition
Information:	or receipt of immunosuppressive medications or treatments, and are unlikely to mount an
	adequate response to COVID-19 vaccination, meeting one of the following:
	Active treatment for solid tumor and hematologic malignancies Hematologic malignancies appropriate with poor responses to COVID 10.
	 Hematologic malignancies associated with poor responses to COVID-19 vaccines regardless of current treatment status (e.g., chronic lymphocytic
	leukemia, non-Hodgkin lymphoma, multiple myeloma, acute leukemia)
	Receipt of solid-organ transplant or an islet transplant and taking
	immunosuppressive therapy
	 Receipt of chimeric antigen receptor (CAR)-T-cell or hematopoietic stem cell
	transplant (within 2 years of transplantation or taking immunosuppressive therapy)
	Moderate or severe primary immunodeficiency (e.g., common variable
	immunodeficiency disease, severe combined immunodeficiency, DiGeorge
	syndrome, Wiskott-Aldrich syndrome)
	 Advanced or untreated human immunodeficiency viruses (HIV) infection (people
	with HIV and CD4 cell counts less than 200/mm³, history of an AIDS-defining
	illness without immune reconstitution, or clinical manifestations of symptomatic HIV)
	 Active treatment with high-dose corticosteroids (at least 20 mg prednisone or
	equivalent per day when administered for 2 or more weeks), alkylating agents,
	antimetabolites, transplant-related immunosuppressive drugs, cancer
	chemotherapeutic agents classified as severely immunosuppressive, and biologic agents that are immunosuppressive or immunomodulatory (such as B-
	cell depleting agents)
	Documentation of prophylactic use
	Baseline SARS-CoV-2 titers that show undetectable antibodies
	Weight of 40 kg or more
Appropriate	Dosing is in accordance with FDA labeling and does not exceed 4500 mg once every 3
Treatment	months
Regimen & Other	
Criteria:	Reauthorization requires documentation of continued immune compromise and low SARS-CoV-2 titers
Exclusion Criteria:	Positive SARS-CoV-2 antigen test or PCR test within the last 3 months
	Received COVID-19 vaccine within the last 3 months
Age Restriction:	12 years of age and older
Prescriber/Site of	All approvals are subject to utilization of the most cost-effective site of care
Care Restrictions:	7 an approvate and easign to anneation of the most open ended to one of our



Coverage Duration:	•	Authorization: 3 months, unless otherwise specified



POLICY NAME: **PENICILLAMINE**

Affected Medications: PENICILLAMINE CAPSULE

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Cystinuria Wilson's Disease Rheumatoid arthritis Copper measurement in urine	
Required Medical Information:	Documented treatment plan including routine urinalysis, WBCs, hemoglobin, platelet count, liver function tests, renal function tests due to risk of fatalities due to aplastic anemia, agranulocytosis, thrombocytopenia, myasthenia gravis, and Goodpasture's Syndrome	
	 Wilson's Disease Diagnosis confirmed by ONE of the following: Genetic testing results confirming biallelic pathogenic ATP7B mutations (in either symptomatic or asymptomatic individuals) Liver biopsy findings consistent with Wilson's disease Presence of Kayser-Fleischer (KF) rings AND serum ceruloplasmin level less than 20 mg/dL AND 24-hour urinary copper excretion greater than 40 mcg Presence of Kayser-Fleischer (KF) rings AND 24-hour urinary copper excretion greater than 100 mcg Absence of KF rings with serum ceruloplasmin level less than 10 mg/dL AND 24-hour urinary copper excretion greater than 100 mcg Rheumatoid arthritis Documentation of severe, active disease defined by one of the following: 	
	 The Disease Activity Score derivative for 28 joints (DAS-28) greater than 3.2 The Simplified Disease Activity Index (SDAI) greater than 11 The Clinical Disease Activity Index (CDAI) greater than 10 Weighted Routine Assessment of Patient Index Data 3 (RAPID3) of at least 2.3 	
Appropriate Treatment Regimen & Other Criteria:	Rheumatoid arthritis Has failed to respond to an adequate trial of conventional therapies (such as methotrexate, sulfasalazine, hydroxychloroquine, leflunomide, Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Enbrel, Xeljanz, Rinvoq, and Inflectra) Reauthorization requires documentation of disease responsiveness to therapy	
Exclusion Criteria:	 For Wilson's disease, must have normalization of free serum copper (non-ceruloplasmin bound copper) to less than 15 mcg/dL and 24-hour urinary copper in the range of 200 to 500 mcg Use of penicillamine during pregnancy (except for treatment of Wilson's disease or exceptions) 	
Age Restriction:	cystinuria)	



Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a specialist familiar with the toxicity and dosage considerations (such as a hepatologist, gastroenterologist, or liver transplant physician for Wilson's Disease) All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months unless otherwise specified Reauthorization: 12 months, unless otherwise specified



PHENOXYBENZAMINE

Affected Medications: PHENOXYBENZAMINE, DIBENZYLINE (phenoxybenzamine)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of sweating and hypertension associated with pheochromocytoma
Required Medical Information:	 Documented diagnosis of pheochromocytoma that requires treatment to control episodes of hypertension and sweating This drug will be used for one of the following: Preoperative preparation for a scheduled surgical resection Chronic treatment of pheochromocytoma that is not amenable to surgery
Appropriate Treatment Regimen & Other Criteria:	Documentation of treatment failure, intolerance, or contraindication to a selective alpha-1 adrenergic receptor blocker (e.g., doxazosin, terazosin, prazosin) Reauthorization will require documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an endocrinologist or a specialist with experience in the management of pheochromocytoma All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Preoperative preparation: 1 month, unless otherwise specified Chronic treatment: 12 months, unless otherwise specified



PHESGO

Affected Medications: PHESGO (pertuzumab-trastuzumab-hyaluronidase-zzxf)

Covered Uses:	NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or better
Required Medical Information:	 Documentation of performance status, disease staging, all prior therapies used, and prescribed dosing regimen Documentation of HER2 positivity based on: 3+ score on immunohistochemistry (IHC) testing OR Positive gene amplification by fluorescence in situ hybridization (FISH) test
Appropriate Treatment Regimen & Other Criteria:	Documentation of an intolerable adverse event to all of the preferred products (Perjeta in combination with Kanjinti, Perjeta in combination with Ogivri) and the adverse event was not an expected adverse event attributed to the active ingredients Reauthorization requires documentation of disease responsiveness to therapy
Exclusion Criteria:	Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an oncologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



suspension)

PHOSPHODIESTERASE-5 (PDE-5) ENZYME INHIBITORS FOR PULMONARY ARTERIAL HYPERTENSION
Affected Medications: ALYQ (tadalafil 20 mg tablet), TADALAFIL (PAH) 20 MG TABLET, TADLIQ (tadalafil 20 mg/5 ml suspension), SILDENAFIL 20 MG TABLET, SILDENAFIL 10 MG/ML SUSPENSION, LIQREV (sildenafil 10 mg/mL

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Pulmonary Arterial Hypertension (PAH) World Health Organization (WHO) Group 1
Required Medical Information:	 Diagnosis of World Health Organization (WHO) Group 1 PAH confirmed by right heart catheterization meeting the following criteria: Mean pulmonary artery pressure of at least 20 mm Hg Pulmonary capillary wedge pressure less than or equal to 15 mm Hg Pulmonary vascular resistance of at least 2.0 Wood units New York Heart Association (NYHA)/WHO Functional Class II or higher symptoms Documentation of Acute Vasoreactivity Testing (positive result requires trial/failure to calcium channel blocker) unless there are contraindications: Low systemic blood pressure (systolic blood pressure less than 90) Low cardiac index Presence of severe symptoms (functional class IV)
Appropriate	For all brand requests: Documented inadequate response or intolerance to sildenafil items 20 mg tablets and tadelafil 20 mg tablets.
Treatment Regimen & Other	 citrate 20 mg tablets and tadalafil 20 mg tablets Requests for oral suspension must have documented inability to swallow tablets
Criteria:	Reauthorization requires documentation of treatment success defined as one or more of the following: Improvement in walking distance Improvement in exercise ability Improvement in pulmonary function Improvement or stability in WHO functional class
Exclusion Criteria:	 Concomitant nitrate therapy on a regular or intermittent basis Concomitant use of a guanylate cyclase stimulator (such as riociguat or vericiguat) Use for erectile dysfunction
Age Restriction:	CCC 10. C. COMIC MYSTATIONS.
Prescriber/Site of	Prescribed by, or in consultation with, a cardiologist or pulmonologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: **PIRFENIDONE**

Affected Medications: PIRFENIDONE (267 and 801 mg)

Cayarad Hasar	All Food and Drum Administration (FDA) approved indications not all the little		
Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by		
	plan design		
	o Idiopathic Pulmonary Fibrosis (IPF)		
Required	Documented diagnosis of idiopathic pulmonary fibrosis (IPF) confirmed by ONE of the		
Medical	following:		
Information:	 Usual interstitial pneumonia (UIP) pattern demonstrated on high-resolution 		
	computed tomography (HRCT)		
	 UIP pattern demonstrated on surgical lung biopsy 		
	 Probable UIP pattern demonstrated on BOTH HRCT and surgical lung biopsy 		
	(e.g., rheumatic disease, environmental exposure, drug toxicity)		
	Documentation of BOTH of the following:		
	 Baseline forced vital capacity (FVC) greater than or equal to 50 percent predicted 		
	 Baseline diffusing capacity for carbon monoxide (DLCO) greater than or equal to 		
	30 percent predicted		
Appropriate	Reauthorization requires documentation of treatment success		
Treatment			
Regimen &			
Other Criteria:			
Exclusion	Combined use with nintedanib (Ofev)		
Criteria:			
Age	18 years of age or older		
Restriction:			
Prescriber/Site of	Prescribed by, or in consultation with, a pulmonologist		
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care		
Coverage	Initial Authorization: 6 months, unless otherwise specified		
Duration:	Reauthorization: 12 months, unless otherwise specified		



POLICY NAME: **PLEGRIDY**

Affected Medications: PLEGRIDY (peglyated interferon beta-1a)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of relapsing forms of multiple sclerosis (MS), including the following: Clinically isolated syndrome (CIS) Relapsing-remitting multiple sclerosis (RRMS) Active secondary progressive disease (SPMS)
Required Medical Information:	Diagnosis confirmed with magnetic resonance imaging (MRI), per revised McDonald diagnostic criteria for MS Clinical evidence alone will suffice; additional evidence desirable but must be consistent with MS
Appropriate Treatment Regimen & Other Criteria:	Reauthorization: provider attestation of treatment success
Exclusion Criteria:	Concurrent use of other disease-modifying medications indicated for the treatment of multiple sclerosis
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist or multiple sclerosis specialist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 24 months, unless otherwise specified



POMBILITI and OPFOLDA

Affected Medications: POMBILITI (cipaglucosidase alfa-atga intravenous injection), OPFOLDA (miglustat oral capsule)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Late-onset Pompe disease for patients weighing 40 kg or more and who are not improving on their current enzyme replacement therapy (ERT)
Required Medical Information:	 Diagnosis of late-onset Pompe disease confirmed by one of the following: Enzyme assay demonstrating a deficiency of acid alpha-glucosidase (GAA) enzyme activity DNA testing that identifies mutations in the GAA gene One or more clinical signs or symptoms of late-onset Pompe disease: Progressive proximal weakness in a limb-girdle distribution Delayed gross-motor development in childhood Involvement of respiratory muscles causing respiratory difficulty (such as reduced forced vital capacity [FVC] or sleep disordered breathing) Skeletal abnormalities (such as scoliosis or scapula alata) Low/absent reflexes Documentation that patient has a 6-minute walk test (6MWT) of 75 meters or more Documentation of a sitting percent predicted forced vital capacity (FVC) of 30% or more Patient weight
Appropriate Treatment Regimen & Other Criteria:	 Documentation of planned treatment regimen for both Pombiliti and Opfolda which are within FDA-labeling Documentation that patient is no longer improving after at least one year of current enzyme replacement therapy (ERT) with Lumizyme (alglucosidase alfa) or Nexviazyme (avalglucosidase alfa-ngpt) Reauthorization will require documentation of treatment success and a clinically significant response to therapy as evidenced by an improvement, stabilization, or slowing of progression in percent-predicted FVC and/or 6MWT
Exclusion Criteria:	 Pregnancy or, if female of reproductive potential, not using effective contraception during treatment Use of invasive or noninvasive ventilation support for more than 6 hours a day while awake Diagnosis of infantile-onset Pompe disease Concurrent treatment with Lumizyme or Nexviazyme Pombiliti or Opfolda as monotherapy Use of Opfolda for Gaucher disease
Age Restriction:	18 years of age or older



Prescriber/Site of Care Restrictions:	•	Prescribed by, or in consultation with, a metabolic specialist, endocrinologist, biochemical geneticist, or provider experienced in the management of Pompe disease All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	•	Authorization: 12 months, unless otherwise specified



POLICY NAME: **PONVORY**

Affected Medications: Ponvory (ponesimod)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of relapsing forms of multiple sclerosis (MS), including the following: Clinically isolated syndrome (CIS) Relapsing-remitting multiple sclerosis (RRMS) Active secondary progressive disease (SPMS)
Required Medical Information:	Diagnosis confirmed with magnetic resonance imaging (MRI), per revised McDonald diagnostic criteria for MS Clinical evidence alone will suffice; additional evidence desirable but must be consistent with MS
Appropriate Treatment Regimen & Other Criteria:	Documented treatment failure with TWO of the following (minimum 12-week trial each): fingolimod, teriflunomide, Mayzent Reauthorization: provider attestation of treatment success
Exclusion Criteria:	Concurrent use of other disease-modifying medications indicated for the treatment of multiple sclerosis
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist or a multiple sclerosis specialist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: POSACONAZOLE

Affected Medications: NOXAFIL (posaconazole), POSACONAZOLE

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
Required Medical Information:	 Susceptibility cultures matching posaconazole activity Current body weight (for pediatric patients)
Appropriate Treatment Regimen & Other Criteria:	 Treatment of invasive aspergillosis Documentation of resistance (or intolerable adverse event) to voriconazole Prophylaxis of invasive Aspergillus and Candida infections
other orienta.	 Documentation of severely immunocompromised state, such as hematopoietic stem cell transplant (HSCT) recipients with graft versus-host disease (GVHD) or those with hematologic malignancies with prolonged neutropenia from chemotherapy Documentation of resistance (or intolerable adverse event) to one other compendia-supported systemic agent (e.g. fluconazole, itraconazole, voriconazole) Treatment of oropharyngeal candidiasis (OPC): Documented failure (or intolerable adverse event) to 10 days or more of treatment with all of the following:
Exclusion Criteria:	0 Itraconazore
Age Restriction:	 Posaconazole delayed release tablets – 2 years of age and older, who weigh greater than 40 kg Noxafil oral suspension – 13 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an infectious disease specialist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 6 months, unless otherwise specified



POLICY NAME: **POZELIMAB**

Affected Medications: VEOPOZ (pozelimab-bbfg)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of CD55-deficient protein-losing enteropathy (PLE) or CHAPLE disease
Required Medical Information:	 Diagnosis of CD-55-deficient PLE confirmed by biallelic CD55 loss-of-function mutation using molecular genetic testing Documentation of hypoalbuminemia (serum albumin of 3.2 g/dL or less) Clinical signs and features of active PLE including abdominal pain, diarrhea, peripheral edema, or facial edema Documentation of at least two albumin transfusions or hospitalizations in the past year
Appropriate Treatment Regimen & Other Criteria:	Dosing is in accordance with FDA labeling and does not exceed the following: Loading Dose: 30 mg/kg by intravenous infusion for 1 dose Maintenance Dose: Starting on day 8; 10 mg/kg as a subcutaneous injection once weekly May be increased to 12 mg/kg starting week 4 Maximum maintenance dosage of 800 mg once weekly Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced Reauthorization requires documentation of positive clinical response with all the following: Improvement or stabilization of clinical symptoms Improvement or normalization of serum albumin concentrations Reduction in albumin transfusion requirements and/or hospitalizations
Exclusion Criteria:	 Receiving concurrent therapy with Soliris (eculizumab) Unresolved Neisseria meningitidis, Streptococcus pneumoniae, or Haemophilus influenzae type b (Hib) infection
Age Restriction:	1 year of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a hematologist, gastroenterologist, or provider that specializes in rare genetic hematologic diseases All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **PRAMLINTIDE**

Affected Medications: SYMLINPEN (pramlintide)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Type 1 diabetes mellitus Type 2 diabetes mellitus
Required Medical Information:	 Documentation of inadequate glycemic control (HbA1c greater than 7 percent) on optimized insulin therapy AND Patient will take SymlinPen in addition to mealtime insulin therapy
Appropriate	Reauthorization will require documentation of treatment success and a clinically significant
Treatment	response to therapy
Regimen & Other Criteria:	
Exclusion Criteria:	HbA1c level greater than 9 percent
	Weight loss treatment
Age Restriction:	
Prescriber/Site of Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



PROLIA

Affected Medications: PROLIA (denosumab)

Covered Uses:	•	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design. Osteoporosis/bone loss
Appropriate Treatment Regimen & Other Criteria:	•	Dosage is 60 mg once every 6 months
Coverage Duration:	•	Initiation Authorization: 24 months, unless otherwise specified Reauthorization: 24 months, unless otherwise specified



PROSTAGLANDIN IMPLANTS

Affected Medications: Durysta (bimatoprost intracameral implant), iDose TR (travoprost intracameral implant)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	 Reduction of intraocular pressure (IOP) in patients with open angle glaucoma (OAG) or ocular hypertension (OHT)
Required Medical	Diagnosis of OAG or OHT with a baseline IOP of at least 22 mmHg
Information:	Documentation of clinical justification for inability to manage routine topical therapy (e.g., due to progression of glaucoma, aging, comorbidities, and administration difficulties that cannot be addressed through instruction and technique)
Appropriate	Documented treatment failure or intolerable adverse event with at least two IOP-lowering
Treatment	agents with different mechanisms of action, (used concurrently), one of which must
Regimen & Other	include a prostaglandin analog such as latanoprost, bimatoprost, tafluprost, travoprost
Criteria:	For iDose TR requests:
Evaluaian Critaria	Documented treatment failure to the preferred product Durysta
Exclusion Criteria:	Repeat implantation with the same prostaglandin implant
	Diagnosis of corneal endothelial cell dystrophy (e.g., Fuchs' Dystrophy)
	 Prior corneal or endothelial cell transplantation (e.g., Descemet's Stripping Automated Endothelial Keratoplasty [DSAEK])
	Active or suspected ocular or periocular infections
	Absent or ruptured posterior lens capsule (Durysta)
Age Restriction:	18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, an ophthalmologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 1 month (one implant per impacted eye), unless otherwise specified



PROXIMAL COMPLEMENT INHIBITOR

Affected Medications: EMPAVELI (pegcetacoplan), FABHALTA (iptacopan)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of adults with paroxysmal nocturnal hemoglobinuria (PNH) Reduce proteinuria in adults with primary immunoglobulin A nephropathy (IgAN) at risk of rapid disease progression, generally a urine protein-to-creatinine ratio (UPCR) ≥1.5 g/g (Fabhalta) Patients must be administered a meningococcal vaccine at least two weeks prior to
Medical Information:	initiation of the requested therapy and revaccinated according to current Advisory Committee on Immunization Practices (ACIP) guidelines
	 PNH Detection of PNH clones of at least 5% by flow cytometry diagnostic testing Presence of at least 2 different glycosylphosphatidylinositol (GPI) protein deficiencies (e.g., CD55, CD59, etc.) within at least 2 different cell lines (e.g., granulocytes, monocytes, erythrocytes) Baseline lactate dehydrogenase (LDH) levels greater than or equal to 1.5 times the upper limit of normal range. One of the following PNH-associated clinical findings: Presence of a thrombotic event Presence of organ damage secondary to chronic hemolysis History of 4 or more blood transfusions required in the previous 12 months IgAN (Fabhalta) Diagnosis of IgAN confirmed with biopsy Documentation of one of the following (with labs current within 30 days of request):
Appropriate Treatment Regimen & Other Criteria:	 UPCR greater than 1.5 g/g PNH For Empaveli: documented inadequate response, contraindication, or intolerance to ravulizumab (Ultomiris) For Fabhalta: documented inadequate response, contraindication, or intolerance to another complement inhibitor such as ravulizumab (Ultomiris) or Empaveli Reauthorization requires documentation of treatment success defined as a decrease in serum LDH, stabilized/improved hemoglobin, decreased transfusion requirement, and reduction in thromboembolic events compared to baseline IgAN (Fabhalta) Documented treatment failure (defined as proteinuria equal to or greater than 1 g/day OR UPCR greater than 1.5 g/g) with a minimum of 12 weeks of all of the following:



	o Filspari (sparsentan)
	<u>Reauthorization</u> requires documentation of treatment success defined as reduction in UPCR or proteinuria from baseline
Exclusion Criteria:	 Concurrent use with other biologics for PNH (Soliris, Ultomiris, Empaveli, or Fabhalta) except when cross tapering according to FDA approved dosing Current meningitis infection or other unresolved serious infection caused by encapsulated
Age Restriction:	bacteria 18 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a hematologist or a nephrologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 3 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



PRIMARY BILIARY CHOLANGITIS AGENTS

Affected Medications: OCALIVA (obeticholic acid), IQIRVO (elafibranor), LIVDELZI (seladelpar)

Covered Uses: Required Medical Information:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Primary biliary cholangitis (PBC) Liver function tests (including alkaline phosphatase and bilirubin) Child-Pugh score 	
Appropriate Treatment Regimen & Other Criteria:	 Documentation that after at least 12 months of adherent therapy with ursodiol or clinical inability to tolerate ursodiol, the patient has ONE of the following: Alkaline phosphatase level (ALP) at least 1.67 times the upper limit of normal (ULN) of the reference lab Total bilirubin above the ULN of the reference lab 	
	Reauthorization will require documentation of treatment success defined as a significant reduction in alkaline phosphatase (ALP) and/or bilirubin levels	
Exclusion Criteria:	 Complete biliary obstruction Decompensated cirrhosis (e.g., Child-Pugh Class B or C) or a prior decompensation event For Ocaliva: Compensated cirrhosis with evidence of portal hypertension (e.g., ascites, gastroesophageal varices, persistent thrombocytopenia) Use in combination with another drug on this policy (Ocaliva, Iqirvo, Livdelzi) 	
Age Restriction:		
Prescriber/Site of	Prescribed by, or in consultation with, a hepatologist	
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care	
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified 	



POLICY NAME: **PYRIMETHAMINE**

Affected Medications: Daraprim, pyrimethamine

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Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by		
	plan design		
	 Toxoplasmosis 		
Required	Documentation of recent <i>Toxoplasma</i> infection		
Medical	Documentation of one of the following:		
Information:	 Severe symptoms (pneumonitis, myocarditis, etc) or prolonged symptoms 		
	greater than 4 weeks with significant impact on quality of life		
	o Immunocompromised status		
Appropriate			
Treatment	Dosing Regimen (adult):		
Regimen &			
Other Criteria:	 Day 1: Pyrimethamine 100 mg, sulfadiazine 2-4 gm divided four times daily, 		
	leucovorin 5-25 mg		
	 Day 2: Pyrimethamine 25-50 mg, sulfadiazine 2-4 gm divided four times daily, 		
	leucovorin 5-25 mg		
	 Day 3 and beyond: Pyrimethamine 25-50 mg, sulfadiazine 500 mg-1 gm divided 		
	four times daily, leucovorin 5-25 mg		
Exclusion	Treatment regimen does not contain leucovorin and a sulfonamide (or alternative if		
Criteria:	allergic to sulfa)		
Age			
Restriction:			
Prescriber/Site of	All approvals are subject to utilization of the most cost-effective site of care		
Care Restrictions:	,		
Coverage	Authorization: Up to 6 weeks, with no reauthorization unless otherwise specified		
Duration:			



RAVICTI

Affected Medications: RAVICTI (glycerol phenylbutyrate)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Chronic management of patients with urea cycle disorders (UCDs) who cannot be managed by dietary protein restriction and/or amino acid supplementation alone
Required Medical Information:	Diagnosis confirmed by enzymatic, biochemical, or genetic testing
Appropriate Treatment Regimen & Other Criteria:	 Documented treatment failure with dietary protein restriction and/or amino acid supplementation alone Documented treatment failure (or intolerable adverse event) to sodium phenylbutyrate or documented comorbid condition with high risk of sodium-induced fluid retention such as heart failure, renal impairment, or edema Must be used in combination with dietary protein restriction Reauthorization will require BOTH of the following: Documentation of treatment success defined as ammonia levels maintained within normal limits That this drug continues to be used in combination with dietary protein restriction
Exclusion Criteria:	 Known hypersensitivity to phenylbutyrate Use for treatment of acute hyperammonemia or N-acetylglutamate synthase (NAGS) deficiency
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a specialist experienced in the treatment of metabolic diseases All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 3 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



RAVULIZUMAB-CWVZ

Affected Medications: ULTOMIRIS (ravulizumab-cwvz)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise evaluded by
Covered Uses.	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	Paroxysmal nocturnal hemoglobinuria (PNH) to reduce hemolysis
	 Atypical hemolytic uremic syndrome (aHUS) to inhibit complement-mediated
	thrombotic microangiopathy
	 Generalized myasthenia gravis (gMG) in adult patients who are anti-acetylcholine
	receptor (AChR) antibody positive
	 Neuromyelitis optica spectrum disorder (NMOSD) who are anti-aquaporin-4
Required Medical	(AQP4) antibody positive for adult patients PNH
Information:	Detection of PNH clones of at least 5% by flow cytometry diagnostic testing
information:	Presence of at least 2 different glycosylphosphatidylinositol (GPI) protein
	deficiencies (e.g., CD55, CD59, etc.) within at least 2 different cell lines (e.g.,
	granulocytes, monocytes, erythrocytes)
	Baseline lactate dehydrogenase (LDH) levels greater than or equal to 1.5 times the upper
	limit of normal range.
	One of the following PNH-associated clinical findings:
	Presence of a thrombotic event
	Presence of organ damage secondary to chronic hemolysis
	 History of 4 or more blood transfusions required in the previous 12 months
	aHUS_
	Clinical presentation of microangiopathic hemolytic anemia, thrombocytopenia, and acute
	kidney injury
	Patient shows signs of thrombotic microangiopathy (TMA) (e.g., changes in mental
	status, seizures, angina, dyspnea, thrombosis, increasing blood pressure, decreased
	platelet count, increased serum creatinine, increased LDH, etc.)
	ADAMTS13 activity level greater than or equal to 10%
	Shiga toxin E. coli related hemolytic uremic syndrome (ST-HUS) has been ruled out
	History of 4 or more blood transfusions required in the previous 12 months
	<u>gMG</u>
	Diagnosis of gMG confirmed by ONE of the following:
	A history of abnormal neuromuscular transmission test
	 A positive edrophonium chloride test
	 Improvement in gMG signs or symptoms with an acetylcholinesterase inhibitor
	Myasthenia Gravis Foundation of America (MGFA) Clinical Classification Class II to IV
	Positive serologic test for AChR antibodies
	Documentation of ONE of the following:
	MG-Activities of Daily Living (MG-ADL) total score of 6 or greater Overtitative Mysetheria Greyin (OMC) total score of 13 or greater
	 Quantitative Myasthenia Gravis (QMG) total score of 12 or greater
	NMOSD
	 Diagnosis of NMOSD with aquaporin-4 immunoglobulin G (AQP4- IgG) antibody positive
	disease confirmed by all of the following:
	Documentation of positive test for AQP4-IgG antibodies via cell-based assay
	1 5 Decementation of positive test for Agri 4 190 antibodies via cell based assay



0	Exclusion of	alternative	diagnoses	(such as	s multiple	sclerosis)
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- At least ONE core clinical characteristic:
 - Acute optic neuritis
 - Acute myelitis
 - Area postrema syndrome (episode of otherwise unexplained hiccups or nausea/vomiting)
 - Acute brainstem syndrome
 - Symptomatic narcolepsy OR acute diencephalic clinical syndrome with NMSOD-typical diencephalic MRI lesions
 - Symptomatic cerebral syndrome with NMOSD-typical lesion on magnetic resonance imaging (MRI) [see table below]
 - Acute cerebral syndrome with NMOSD-typical brain lesion on MRI [see table below]

Clinical presentation	Possible MRI findings
Diencephalic syndrome	Periependymal lesion
	Hypothalamic/thalamic lesion
Acute cerebral syndrome	Extensive periependymal lesion
	 Long, diffuse, heterogenous, or edematous corpus callosum lesion
	Long corticospinal tract lesion
	Large, confluent subcortical or deep white matter lesion

Appropriate Treatment Regimen & Other Criteria:

aHUS

- Failure to respond to plasma therapy within 10 days
 - Trial of plasma therapy not required if one of the following is present:
 - Life-threatening complications of HUS such as seizures, coma, or heart failure
 - Confirmed presence of a high-risk complement genetic variant (e.g., CFH or CFI)

<u>gM</u>G

- Documentation of one of the following:
 - Treatment failure with an adequate trial (one year or more) of at least 2 immunosuppressive therapies (azathioprine, mycophenolate, tacrolimus, cyclosporine, methotrexate)
 - Has required three or more courses of rescue therapy (plasmapheresis/plasma exchange and/or intravenous immunoglobulin), while on at least one immunosuppressive therapy, over the last 12 months
- Documented inadequate response, contraindication, or intolerance to efgartigimod-alfa (Vyvgart)

NMOSD

 Documented inadequate response, contraindication, or intolerance to ALL of the following:



Rituximab (preferred products: Riabni, Ruxience)Satralizumab-mwge (Enspryng)	
o Inebilizumab-cdon (Uplizna)	
Reauthorization requires:	
 gMG: documentation of treatment success defined as an improvement in MG- QMG scores from baseline 	·ADL and
 PNH: documentation of treatment success defined as a decrease in serum LD stabilized/improved hemoglobin, decreased transfusion requirement, and redu thromboembolic events compared to baseline 	,
 aHUS: documentation of treatment success defined as a decrease in serum L stabilized/improved serum creatinine, increased platelet count, and decreased exchange/infusion requirement compared to baseline 	,
 NMOSD: documentation of treatment success defined as the stabilization or 	
improvement in neurological symptoms as evidenced by a decrease in acute of Expanded Disability Status Scale (EDSS) score, hospitalizations, or plasma extreatments	
Exclusion Criteria: • Current meningitis infection	
 Concurrent use with other disease-modifying biologics for requested indication otherwise specified 	ո, unless
Age Restriction: • PNH, aHUS: 1 month of age and older	
gMG: 18 years of age and older	
Prescriber/Site of • Prescribed by, or in consultation with, a specialist	
Care Restrictions: o PNH: hematologist	
 aHUS: hematologist or nephrologist 	
o gMG: neurologist	
NMOSD: neurologist or neuro-ophthalmologist	
All approvals are subject to utilization of the most cost-effective site of care	
Coverage Duration: • Initial Authorization: 3 months, unless otherwise specified	
Reauthorization: 12 months, unless otherwise specified	



REBIF

Affected Medications: REBIF, REBIF TITRATION PACK

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of relapsing forms of multiple sclerosis (MS), including the following: Clinically isolated syndrome (CIS) Relapsing-remitting multiple sclerosis (RRMS) Active secondary progressive disease (SPMS)
Required Medical Information:	Diagnosis confirmed with magnetic resonance imaging (MRI), per revised McDonald diagnostic criteria for MS Clinical evidence alone will suffice; additional evidence desirable but must be consistent with MS
Appropriate Treatment Regimen & Other Criteria:	Reauthorization: provider attestation of treatment success
Exclusion Criteria:	Concurrent use of other disease-modifying medications for the treatment of MS
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified.



POLICY NAME: **RELYVRIO**

Affected Medications: RELYVRIO (sodium phenylbutyrate-taurursodiol)

	T		
Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by		
	plan design		
	 Amyotrophic lateral sclerosis (ALS) 		
Required Medical	Definite or probable Amyotrophic lateral sclerosis (ALS) based on El Escorial revised		
Information:	(Airlie House) criteria		
	Symptom onset within 18 months		
	Slow vital capacity (SVC) of at least 60 percent		
	Patient currently retains most activities of daily living defined as at least 2 points on all 12		
	items of the ALS functional rating scale-revised (ALSFRS-R)		
Appropriate	Documentation of one of the following:		
Treatment	 Member is stable on riluzole 		
Regimen & Other Criteria:	Prescriber has indicated clinical inappropriateness of riluzole		
	Reauthorization: Documentation of treatment success as determined by prescriber		
	including retaining most activities of daily living		
Exclusion Criteria:	Presence of a tracheostomy		
	Use of permanent assisted ventilation		
Age Restriction:	18 years of age and older		
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist		
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care		
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified		
	Reauthorization: 12 months, unless otherwise specified		



POLICY NAME: **REMODULIN**

Affected Medications: REMODULIN INJECTION (treprostinil)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by			
	plan design			
	 Pulmonary Arterial Hypertension (PAH) World Health Organization (WHO) Group 			
	7			
	 Pulmonary Arterial Hypertension in patients requiring transition from 			
D'	epoprostenol (PAII) M/IIO Committee			
Required	Pulmonary Arterial Hypertension (PAH) WHO Group 1			
Medical Information:	Documentation of PAH confirmed by right-heart catheterization meeting the following			
information:	criteria:			
	Mean pulmonary artery pressure of at least 20 mm Hg			
	Pulmonary capillary wedge pressure less than or equal to 15 mm Hg Pulmonary capillary register as a feet least 2.0 West during			
	o Pulmonary vascular resistance of at least 2.0 Wood units			
	Etiology of PAH: idiopathic PAH, hereditary PAH, OR			
	PAH secondary to one of the following conditions:			
	Connective tissue disease			
	Human immunodeficiency virus (HIV) infection			
	o Cirrhosis			
	Anorexigens Congenite Left to right abunta			
	 Congenital left to right shunts Schistosomiasis 			
	Drugs and toxins Dortel hypertension			
	Portal hypertension New York Heart Association (NYHA) (Morald Health Organization (MHQ) Functional Class			
	New York Heart Association (NYHA)/World Health Organization (WHO) Functional Class If or higher symptoms			
	If or higher symptoms			
	 Documentation of Acute Vasoreactivity Testing (positive result requires trial/failure to calcium channel blockers) unless there are contraindications: 			
	Low systemic blood pressure (systolic blood pressure less than 90)			
	Low systemic blood pressure (systolic blood pressure less than 90) Low cardiac index			
	OR			
	Presence of severe symptoms (functional class IV)			
Appropriate	The pulmonary hypertension has progressed despite maximal medical and/or surgical			
Treatment	treatment of the identified condition			
Regimen &	Documentation that treprostinil is used as a single route of administration (Remodulin,			
Other Criteria:	Tyvaso, Orenitram should not be used in combination)			
	Treatment with oral calcium channel blocking agents has been tried and failed, or has			
	been considered and ruled out			
	Treatment with combination of endothelin receptor antagonist (ERA) and			
	phosphodiesterase 5 inhibitor (PDE5I) has been tried and failed for WHO functional class			
	II and III			
	Reauthorization requires documentation of treatment success defined as one or more of the following:			
	Improvement in walking distance Improvement in exercise shility			
	Improvement in exercise ability			
	Improvement in pulmonary function			



	Improvement or stability in WHO functional class
Exclusion Criteria:	PAH secondary to pulmonary venous hypertension (e.g., left sided atrial or ventricular disease, left sided valvular heart disease, etc) or disorders of the respiratory system (e.g., chronic obstructive pulmonary disease, interstitial lung disease, obstructive sleep apnea or other sleep disordered breathing, alveolar hypoventilation disorders, etc.)
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a cardiologist or pulmonologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **RESLIZUMAB**

Affected Medications: CINQAIR (reslizumab)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by	
	plan design	
	 Add-on maintenance treatment of adult patients with severe asthma with an 	
	eosinophilic phenotype	
Required Medical	Diagnosis of severe asthma with an eosinophilic phenotype, defined by both of the	
Information:	following:	
	 Baseline eosinophil count of at least 400 cells/μL 	
	 FEV1 less than 80% at baseline or FEV1/FVC reduced by at least 5% from normal 	
Appropriate	Documented use of high-dose inhaled corticosteroid (ICS) plus a long-acting beta	
Treatment	agonist (LABA) for at least three months with continued symptoms	
Regimen & Other	Documentation of one of the following:	
Criteria:	 Documented history of 2 or more asthma exacerbations requiring oral or 	
	systemic corticosteroid treatment in the past 12 months while on combination	
	inhaler treatment and at least 80% adherence	
	 Documentation that chronic daily oral corticosteroids are required 	
	Documented treatment failure or intolerable adverse event with all of the preferred	
	products (Dupixent, Fasenra, Nucala, and Xolair)	
	Availability: 100 mg/10 mL vials	
	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced	
	Reauthorization: documentation of treatment success and a clinically significant response to therapy	
Exclusion Criteria:	Use in combination with another monoclonal antibody (e.g., Dupixent, Nucala, Xolair, Fasenra, Tezspire)	
Age Restriction:	18 years of age and older	
Prescriber/Site of	Prescribed by, or in consultation with, an allergist, immunologist, or pulmonologist	
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care	
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified	
	Reauthorization: 12 months, unless otherwise specified	



POLICY NAME: **RESMETIROM**

Affected Medications: REZDIFFRA (resmetirom)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
0010104 00001	plan design
	Treatment of adults with noncirrhotic nonalcoholic steatohepatitis (NASH) with
	moderate to advanced liver fibrosis (consistent with stages F2 to F3 fibrosis), in
	conjunction with diet and exercise
Required Medical	Diagnosis of NASH or metabolic dysfunction–associated steatohepatitis (MASH) with
Information:	moderate to advanced (F2 to F3) liver fibrosis confirmed by ONE of the following:
	 Conclusive result from a well-validated non-invasive test such as:
	 Fibroscan-AST (FAST) score
	 MAST (score from MRI–proton density fat fraction, Magnetic resonance
	elastography [MRE], and serum AST)
	 MEFIB (Fibrosis-4 Index ≥1.6 and MRE ≥3.3 kPa)
	 Liver biopsy (also required if non-invasive testing is inconclusive or other causes
	for liver disease have not been ruled out)
	Other causes for liver steatosis have been ruled out (such as alcohol-associated liver)
	disease, chronic hepatitis C, Wilson disease, drug-induced liver disease)
	Baseline lab values for AST and ALT
Appropriate	Documentation of abstinence from alcohol consumption
Treatment	Documentation of comprehensive comorbidity management being undertaken, including
Regimen & Other	all of the following:
Criteria:	 Use of diet and exercise for weight management
	 Medications to manage associated comorbid conditions, such as thyroid disease
	(must not have active disease), diabetes, dyslipidemia, hypertension, or
	cardiovascular conditions
	Reauthorization requires documentation of disease responsiveness to therapy based on
	improvements or stability in laboratory results, such as ALT and AST, or fibrosis as evaluated
	by a non-invasive test
Exclusion Criteria:	History of excessive alcohol use or alcohol-associated liver disease
	Current excessive alcohol use
	Continued use of medications associated with liver steatosis
	Stage 4 liver disease or cirrhosis
	Use for other liver disease
	Active or untreated thyroid disease
Age Restriction:	
Prescriber/Site of	 Prescribed by, or in consultation with, a hepatologist or gastroenterologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: **RETHYMIC**

Affected Medications: RETHYMIC (allogeneic processed thymus tissue-agdc)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Immune reconstitution in pediatric patients with congenital athymia	
Required Medical Information:	Documentation of congenital athymia associated with one of the following:	
Appropriate Treatment Regimen & Other Criteria:	 Congenital athymia confirmed by flow cytometry that demonstrates: Fewer than 50 naïve T cells/mm3 in the peripheral blood OR Less than 5% of total T cells being naïve T cells 	
Exclusion Criteria:	 Treatment of patients with severe combined immunodeficiency (SCID) Prior thymus transplant 	
Age Restriction:		
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a pediatric immunologist or prescriber experienced in the treatment of congenital athymia All approvals are subject to utilization of the most cost-effective site of care 	
Coverage Duration:	Authorization: 1 month (1 treatment only), unless otherwise specified	



POLICY NAME: RILONACEPT

Affected Medications: ARCALYST (rilonacept)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of Cryopyrin-Associated Periodic Syndromes (CAPS), including 		
	 Treatment of Cryopyrin-Associated Periodic Syndromes (CAPS), including Familial Cold Autoinflammatory Syndrome (FCAS), and Muckle-Wells Syndrome 		
	(MWS) in adults and pediatric patients 12 years and older		
	The maintenance of remission of Deficiency of Interleukin-1 Receptor Antagonist (DIRA) in adults and podiatric patients weighing at least 10 kg.		
	(DIRA) in adults and pediatric patients weighing at least 10 kg		
	Treatment of recurrent pericarditis (RP) and reduction in risk of recurrence in		
Required	adults and pediatric patients 12 years and older Documentation confirming one of the following:		
Medical	Diagnosis of Cryopyrin-Associated Periodic Syndromes (CAPS), including Familial Cold		
Information:	Autoinflammatory Syndrome (FCAS), and Muckle-Wells Syndrome (MWS)		
	Diagnosis of Deficiency of Interleukin-1 Receptor Antagonist (DIRA)		
	Must include genetic testing results which confirm the presence of homozygous		
	mutations in the interleukin-1 receptor antagonist (IL1RN) gene		
	Disease must currently be in remission		
	Diagnosis of Recurrent Pericarditis with an inflammatory phenotype shown by one of the		
	following:		
	 Fever, elevated C-Reactive protein (CRP), elevated white blood cell count, 		
	elevated erythrocyte sedimentation rate (ESR), pericardial late gadolinium		
	enhancement (LGE) on cardiac magnetic resonance (CMR), or pericardial		
A	contrast enhancement on computed tomography (CT) scan		
Appropriate Treatment	All Indications: Documented treatment failure or intolerable adverse event with trial of Kineret (anakinra)		
Regimen &	Documented treatment failure or intolerable adverse event with trial of Kineret (anakinra)		
Other Criteria:	Recurrent Pericarditis:		
	 Documented treatment failure or intolerable adverse event to triple therapy with all of the 		
	following:		
	Colchicine		
	Non-steroidal anti-inflammatory (NSAID) or aspirin		
	Glucocorticoid		
	Reauthorization:		
	All indications: documentation of treatment success and a clinically significant response		
	to therapy		
	Recurrent pericarditis: documentation that the patient is unable to remain		
	asymptomatic with normal CRP levels upon trial of an appropriate tapering regimen		
Exclusion	Active or chronic infection		
Criteria:	Concurrent therapy with anakinra, tumor necrosis factor (TNF) inhibitors, or other		
	1		
	biologics		



Age Restriction:	CAPS or Recurrent Pericarditis: 12 years of age and older
Prescriber/Site of Care Restrictions:	Prescribed by, or in consultation with, a rheumatologist, immunologist, cardiologist, or dermatologist
	All approvals are subject to utilization of the most cost-effective site of care
Coverage	Initial Authorization: 3 months, unless otherwise specified
Duration:	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: RIOCIGUAT

Affected Medications: ADEMPAS (riociguat)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Pulmonary Arterial Hypertension (PAH) World Health Organization (WHO) Group 1 Chronic-Thromboembolic Pulmonary Hypertension (WHO Group 4)
Required	Chronic Thromboembolic Pulmonary Hypertension (CTEPH)
Medical	Documentation of CTEPH (WHO Group 4) meeting the following criteria:
Information:	 Evidence of thromboembolic occlusion of proximal or distal pulmonary vasculature on CT/MRI or V/Q scan
	Mean pulmonary arterial pressure greater than 20 mm Hg
	PAWP less than 15 mm Hg
	Elevated pulmonary vascular resistance over 2 Wood units
	Pulmonary Arterial Hypertension (PAH)
	Documentation of PAH confirmed by right-heart catheterization meeting the following criteria:
	 Mean pulmonary artery pressure of at least 20 mm Hg
	 Pulmonary capillary wedge pressure less than or equal to 15 mm Hg
	 Pulmonary vascular resistance of at least 2.0 Wood units
	Etiology of PAH (idiopathic, heritable, or associated with connective tissue disease)
	New York Heart Association (NYHA)/World Health Organization (WHO) Functional Class
	II or higher symptoms
	Documentation of Acute Vasoreactivity Testing (positive result requires trial/failure to
	calcium channel blocker) unless there are contraindications:
	 Low systemic blood pressure (systolic blood pressure less than 90) Low cardiac index OR
	 Presence of severe symptoms (functional class IV)
Appropriate	СТЕРН
Treatment Regimen &	 Documentation of failure of or inability to receive pulmonary endarterectomy surgery Current therapy with anticoagulants
Other Criteria:	
	PAH District of the state of th
	 Documented failure to the following therapy classes: Phosphodiesterase type 5 (PDE5) inhibitors AND endothelin receptor antagonists
	Reauthorization requires documentation of treatment success defined as one or more of the
	following:
	Improvement in walking distance Improvement in exercise ability
	Improvement in exercise ability
	Improvement in pulmonary function
Evaluaier	Improvement or stability in WHO functional class
Exclusion Critoria:	Concomitant use with nitrates or nitric oxide donors (such as amyl nitrite)
Criteria:	



	Concomitant use with specific PDE-5 inhibitors (such as sildenafil, tadalafil, or vardenafil) or non-specific PDE inhibitors (such as dipyridamole or theophylline)
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a cardiologist or a pulmonologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: RISDIPLAM

Affected Medications: EVRYSDI (risdiplam)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Spinal muscular atrophy (SMA)
Required Medical Information:	Diagnosis of SMA type 1, 2, or 3 confirmed by genetic testing of chromosome 5q13.2 demonstrating ONE of the following: Homozygous gene deletion of SMN1 (survival motor neuron 1) Homozygous gene mutation of SMN1 Compound heterozygous gene mutation of SMN1 Documentation of 4 or fewer copies of the SMN2 (survival motor neuron 2) gene Documentation of one of the following baseline motor assessments appropriate for patient age and motor function: Hammersmith Infant Neurological Examination (HINE-2) Hammersmith Functional Motor Scale (HFSME) Children's Hospital of Philadelphia Infant Test of Neuromuscular Disorders (CHOP-INTEND) Upper Limb Module (ULM) test 6-Minute Walk Test (6MWT) Documentation of previous treatment history Documentation of ventilator use status: Patient is NOT ventilator-dependent (defined as using a ventilator at least 16 hours per day on at least 21 of the last 30 days) This does not apply to patients who require non-invasive ventilator assistance
Appropriate Treatment Regimen & Other Criteria:	Reauthorization requires documentation of improvement in baseline motor assessment score, clinically meaningful stabilization, or delayed progression of SMA-associated signs and symptoms
Exclusion Criteria:	 SMA type 4 Advanced SMA at baseline (complete paralysis of limbs, permanent ventilation support) Prior treatment with SMA gene therapy (i.e., onasemnogene abeparvovec-xioi) Will not use in combination with other agents for SMA (e.g., onasemnogene abeparvovec-xioi, nusinersen, etc.)
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist or provider who is experienced in treatment of spinal muscular atrophy All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



RITUXIMAB

Affected Medications: RITUXAN (rituximab), RITUXAN HYCELA (rituximab and hyaluronidase human), TRUXIMA (rituximab-abbs), RUXIENCE (rituximab-pvvr), RIABNI (rituximab-arrx)

Covered Uses:	All Food and Drug Administration (FDA)-approved and compendia supported indications
	not otherwise excluded by plan design
	Rheumatoid arthritis (RA)
	Microscopic Polyangiitis (MPA)
	 Granulomatosis with Polyangiitis (GPA)
	 Eosinophilic granulomatosis with polyangiitis (EGPA)
	 Relapsing forms of multiple sclerosis (MS)
	 Clinically isolated syndrome (CIS)
	 Relapsing-remitting multiple sclerosis (RRMS)
	 Active secondary progressive disease (SPMS)
	 Neuromyelitis Optica Spectrum Disorder (NMOSD)
	 Pemphigus Vulgaris (PV) and other autoimmune blistering skin diseases
	 Thrombocytopenia in patients with immune thrombocytopenia (ITP)
	NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A
	or higher
Required Medical	 Documentation of disease staging, all prior therapies used, and anticipated treatment course
Information:	DA.
	RA Desumentation of moderate to severe disease despite surrent treatment
	 Documentation of moderate to severe disease despite current treatment Documented current level of disease activity with one of the following (or equivalent
	objective scale):
	 Disease Activity Score derivative for 28 joints (DAS-28) greater than 3.2
	 Simplified Disease Activity Index (SDAI) greater than 11
	 Clinical Disease Activity Index (CDAI) greater than 10
	 Weighted RAPID3 of at least 2.3
	MPA or GPA
	Documentation of active GPA or MPA
	EGPA FORA
	Non-severe disease: documentation of active EGPA
	OR
	Severe disease: documentation of organ or life-threatening manifestations as defined by the American College of Pharmetalagu/Acapulitia Foundation (ACRA/F)
	the American College of Rheumatology/Vasculitis Foundation (ACR/VF)
	Relapsing Forms of MS
	Diagnosis confirmed with magnetic resonance imaging (MRI) per revised McDonald
	diagnostic criteria for multiple sclerosis (MS)
	 Clinical evidence alone will suffice; additional evidence desirable but must be consistent with MS
	NMOSD



- Diagnosis of seropositive aquaporin-4 immunoglobulin G (AQP4-IgG) NMOSD confirmed by all of the following:
 - o Documentation of AQP4-IgG-specific antibodies on cell-based assay
 - Exclusion of alternative diagnoses (such as multiple sclerosis)
 - o At least **one** core clinical characteristic:
 - Acute optic neuritis
 - Acute myelitis
 - Acute area postrema syndrome (episode of otherwise unexplained hiccups or nausea/vomiting)
 - Acute brainstem syndrome
 - Symptomatic narcolepsy OR acute diencephalic clinical syndrome with NMOSD-typical diencephalic lesion on magnetic resonance imaging (MRI) [see table below]
 - Acute cerebral syndrome with NMOSD-typical brain lesion on MRI [see table below]

Clinical presentation	Possible MRI findings	
Diencephalic syndrome	Periependymal lesionHypothalamic/thalamic lesion	
Acute cerebral syndrome	 Extensive periependymal lesion Long, diffuse, heterogenous, or edematous corpus callosum lesion Long corticospinal tract lesion Large, confluent subcortical or deep white matter lesion 	

<u>PV and other autoimmune blistering skin diseases (such as but not limited to pemphigus foliaceus, bullous pemphigoid, cicatricial pemphigoid, epidermolysis bullosa acquisita, and paraneoplastic pemphigus)</u>

- Diagnosis confirmed by biopsy
- Documented severe or refractory disease with failure to conventional topical and oral systemic therapies

Thrombocytopenia in patients with ITP

- Platelet count less than 20,000/mcL AND
- One of the following:
 - Documented steroid dependence to maintain platelets/prevent bleeding for at least 3 months
 - Lack of clinically meaningful response to corticosteroids (defined as inability to increase platelets to at least 50,000/mcL)

Appropriate Treatment Regimen & Other Criteria:

All Uses

- Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced
 - Coverage of Truxima, Rituxan, or Rituxan Hycela requires documentation of one of the following:



 A documented intolerable adverse event to the preferred products, Riabni and Ruxience, and the adverse event was not an expected adverse event attributed to the active ingredient

Oncology Uses

 Documentation of ECOG performance status of 1 or 2 OR Karnofsky performance score greater than 50%

<u>RA</u>

- Initial Course: Documented failure with two of the preferred pharmacy drugs (Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Enbrel, Xeljanz, Rinvoq)
 - o Dose is approved for up to 2 doses of 1,000 mg given 2 weeks apart
- Repeat Course: Approve if 16 weeks or more after the first dose of the previous rituximab regimen and the patient has responded (e.g., less joint pain, morning stiffness, or fatigue, or improved mobility, or decreased soft tissue swelling in joints or tendon sheaths) as determined by the prescribing physician.

Relapsing Forms of MS

- Initial: May include one-time induction dose (e.g., 1,000 mg once every 2 weeks for 2 doses)
- Maintenance: Approvable up to 2,000 mg annually. Higher doses will require documentation to support

NMOSD

- Initial: May include one-time induction dose (e.g., 1,000 mg once every 2 weeks for 2 doses)
- Maintenance: Approvable up to 2,000 mg annually. Higher doses will require documentation to support (e.g., detection of CD19+ lymphocytes)

MPA and GPA

- Initial: May include one-time induction dose (e.g., 1,000 mg once every 2 weeks for 2 doses or 375 mg/m² once weekly for 4 doses), to be used in combination with a systemic glucocorticoid
- Maintenance: Approvable for up to 1,000 mg annually. Higher doses will require documentation to support (e.g., positive ANCA titers, detection of CD19+ lymphocytes)

EGPA

- Non-severe disease:
 - Documented treatment failure with a corticosteroid
 - Documented treatment failure to an adequate trial (at least 12 weeks) with an oral immunosuppressive therapy: azathioprine, methotrexate, mycophenolate, leflunomide
- Severe disease:
 - Documentation that rituximab will be administered in combination with a systemic glucocorticoid



	PV and other autoimmune blistering skin diseases		
	 Documentation that rituximab will be administered in combination with a systemic glucocorticoid (if appropriate) 		
	Documented treatment failure with 12 weeks of a corticosteroid AND		
	 Documented treatment failure with 12 weeks of an immunosuppressant at an adequate dose (e.g., azathioprine, mycophenolate, methotrexate, etc.) or other appropriate corticosteroid-sparing therapy 		
	All other indications		
	A Food and Drug Administration (FDA)-approved or compendia supported dose, frequency, and duration of therapy		
	Documented treatment failure of first-line recommended and conventional therapies		
	Reauthorization: documentation of disease responsiveness to therapy		
Exclusion Criteria:	MS: Concurrent anti-CD20-directed therapy or other disease-modifying medications indicated for the treatment of MS		
	Other non-oncology indications: Concurrent use with targeted immune modulators		
Age Restriction:			
Prescriber/Site of	For RA, MPA, GPA, EGPA: Prescribed by, or in consultation with, a rheumatologist		
Care Restrictions:	For CLL, NHL: Prescribed by, or in consultation with, an oncologist		
	For MS, NMOSD: Prescribed by, or in consultation with, a neurologist or MS specialist		
	For PV: Prescribed by, or in consultation with, a dermatologist		
	All approvals are subjects to utilization of the most cost-effective site of care		
Coverage	Initial Authorization:		
Duration:	 PV, MPA, GPA, EGPA – 3 months, unless otherwise specified 		
	Oncology – 4 months, unless otherwise specified PA MS NMOSP – 6 months, unless otherwise specified.		
	o RA, MS, NMOSD – 6 months, unless otherwise specified		
	Reauthorization: 12 months, unless otherwise specified		



RNA INTERFERENCE DRUGS FOR PRIMARY HYPEROXALURIA 1

Affected Medications: OXLUMO (lumasiran), RIVFLOZA (nedosiran)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Primary hyperoxaluria type 1 (PH1)
Required Medical Information:	 A diagnosis of primary hyperoxaluria type 1 (PH1) confirmed by genetic testing confirming presence of AGXT gene mutation Metabolic testing demonstrating elevated urinary oxalate excretion Presence of clinical manifestations diagnostic of PH1 such as: Metabolic testing demonstrating elevated urinary glycolate excretion Normal levels of L-glyceric acid (elevation indicates PH type 2) Normal levels of hydroxy-oxo-glutarate (elevation indicates PH type 3) For Rivfloza: eGFR of 30 or more
Appropriate Treatment Regimen & Other Criteria:	 For Rivfloza: Trial and failure or contraindication with Oxlumo <u>Reauthorization</u> requires documentation of the following criteria related to treatment success: Reduction from baseline in urine or plasma oxalate levels Improvement, stabilization, or slowed worsening of one or more clinical manifestation of PH1 (i.e., nephrocalcinosis, renal stone events, renal impairment, systemic oxalosis)
Exclusion Criteria:	 Diagnosis of primary hyperoxaluria type 2 or type 3 Secondary hyperoxaluria Concurrent use of another RNA interference drug for PH1
Age Restriction Prescriber/Site of Care Restrictions:	 For Rivfloza: age in accordance with FDA labeling Prescribed by, or in consultation with, a nephrologist, urologist, geneticist, or specialist in the treatment of PH1 All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: ROMIPLOSTIM

Affected Medications: NPLATE (romiplostim)

All Food and Drug Administration (FDA)-approved indications not otherwise excluded plan design Adult patients with immune thrombocytopenia (ITP) who have had an insuffice response to corticosteroids, immunoglobulins, or splenectomy Pediatric patients 1 year of age and older with ITP or at least 6 months who have had an insufficient response to corticosteroids, immunoglobulins, or splenectomy Adult and pediatric patients (including term neonates) with acute exposure to myelosuppressive radiation doses Thrombocytopenia in patients with ITP Documentation of ONE of the following: Platelet count less than 20,000/microliter AND symptomatic bleeding Platelet count less than 50,000/microliter AND increased risk for bleeding (su as peptic ulcer disease, use of antiplatelets or anticoagulants, history of bleed at higher platelet count, need for surgery or invasive procedure) Hematopoletic syndrome of acute radiation syndrome Suspected or confirmed exposure to radiation levels greater than 2 gray (Gy)	overed Uses:	All Food and Drug Administration (FDA) approved indications not otherwise evaluded by
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Hematopoietic syndrome of acute radiation syndrome		
Approved for one-time single subcutaneous injection of 10 mcg/kg		
• Treatment of thrombocytopenia due to myelodysplastic syndrome (MDS)		Treatment of thrombocytopenia due to myelodysplastic syndrome (MDS)
Criteria: • Use in combination with another thrombopoietin receptor agonist, spleen tyrosine kind	riteria:	Use in combination with another thrombopoietin receptor agonist, spleen tyrosine kinase
inhibitor, or similar treatments (Promacta, Nplate, Tavalisse)		
Age		,
Restriction:	.ge	



Prescriber/Site of	Prescribed by, or in consultation with, a hematologist	
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care	
Coverage	Thrombocytopenia in patients with ITP	
Duration:	Initial Approval: 4 months, unless otherwise specified	
	Reauthorization: 12 months, unless otherwise specified	
	Hematopoietic syndrome of acute radiation syndrome	
	Authorization: 1 month, unless otherwise specified	



POLICY NAME: ROMOSOZUMAB

Affected Medications: EVENITY (romosozumab-aqqg)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of osteoporosis in postmenopausal women at high risk for fracture, defined as one of the following:
Required Medical Information:	Diagnosis of osteoporosis as defined by at least one of the following: T-score less than or equal to -2.5 (current or past) at the lumbar spine, femoral neck, total hip, or 1/3 radius site T-score between -1.0 and -2.5 at the lumbar spine, femoral neck, total hip, or 1/3 radius site AND increased risk of fracture as defined by at least one of the following Fracture Risk Assessment Tool (FRAX) scores: FRAX 10-year probability of major osteoporotic fracture is 20% or greater FRAX 10-year probability of hip fracture is 3% or greater History of non-traumatic fractures in the absence of other metabolic bone disorders
Appropriate Treatment Regimen & Other Criteria:	Treatment failure, contraindication, or intolerance to all of the following:
Exclusion Criteria:	 Heart attack or stroke event within the preceding year Concurrent use of bisphosphonates, parathyroid hormone analogs, or RANK ligand inhibitors Hypocalcemia that is uncorrected prior to initiating Evenity
Age Restriction: Prescriber/Site of Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months (no reauthorization), unless otherwise specified



POLICY NAME: **RUFINAMIDE**

Affected Medications: BANZEL (rufinamide), RUFINAMIDE SUSPENSION, RUFINAMIDE TABLET

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Lennox-Gastaut Syndrome(LGS)
Required Medical	All Indications
Information:	Patient weight
	Documentation that rufinamide will be used as adjunctive therapy
	Lennox-Gastaut Syndrome (LGS)
	Documentation of at least 8 drop seizures per month while on stable antiepileptic drug thorapy
	 therapy Documented treatment and inadequate seizure control with at least three guideline directed therapies including:
	 Valproate and
	 Lamotrigine and
	 Topiramate, felbamate, or clobazam
Appropriate	Dosing: not to exceed 3200 mg daily
Treatment	
Regimen & Other	
Criteria:	<u>Reauthorization</u> requires documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	Familial Short QT syndrome
	Use as monotherapy for seizure control
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: RYPLAZIM

Affected Medications: RYPLAZIM

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by		
	plan design		
	 Plasminogen Deficiency Type 1 		
Required Medical Information:	 Diagnosis of symptomatic congenital plasminogen deficiency (C-PLGD) type 1, as evidenced by documentation of all of the following: Clinical signs and symptoms of the disease (such as ligneous conjunctivitis, gingivitis, tonsillitis, abnormal wound healing) Presence of (ligneous) pseudomembranous lesions with documentation of size location, and total number of lesions Baseline plasminogen activity level less than or equal to 45% of laboratory 		
Appropriate	standard Dosing		
Treatment	Dosing may not exceed 6.6 mg/kg every 2 days.		
Regimen & Other	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be		
Criteria:	enforced.		
	Reauthorization requires documentation of disease responsiveness to therapy, defined as the following:		
	Trough plasminogen activity level (taken 72 hours after dose) increased by 10% or		
	greater above baseline		
	Improvement (reduction) in lesion number/size from baseline		
Exclusion Criteria:	Prior treatment failure with Ryplazim		
	Treatment of idiopathic pulmonary fibrosis		
Age Restriction:			
Prescriber/Site of	Prescribed by, or in consultation with, a hematologist		
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care		
Coverage Duration:	Initial Authorization: 4 months, unless otherwise specified		
	Reauthorization: 12 months, unless otherwise specified		



POLICY NAME: SACROSIDASE

Affected Medications: SUCRAID (sacrosidase)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by		
	plan design		
	o Oral replacement therapy for congenital sucrase-isomaltase deficiency (CSID)		
Required Medical Information:	 Documentation of confirmed congenital sucrose-isomaltase deficiency, diagnosed by one of the following: Small bowel biopsy Sucrose breath test Genetic test Documentation of current symptoms (e.g., diarrhea, abdominal pain or cramping, bloating, gas, loose stools, nausea, vomiting) Reauthorization: requires documentation of treatment success and a clinically significant response to therapy (fewer stools, lower number of symptoms) 		
Appropriate	- sapanas to masapy (taken etcolo), letter hamber of cymptomor		
Treatment			
Regimen & Other Criteria:			
Exclusion Criteria:			
Age Restriction:	5 months of age or older		
Prescriber/Site of	Prescribed by, or in consultation with, a gastroenterologist or genetic specialist		
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care		
Coverage Duration:	Initial Authorization: 3 months, unless otherwise specified		
	Reauthorization: 12 months, unless otherwise specified		



POLICY NAME: **SAPROPTERIN**

Affected Medications: KUVAN (sapropterin), SAPROPTERIN

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise evaluded by
Covered Oses.	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by Plan design.
	plan design
	Reduce phenylalanine (Phe) levels in those that are one month of age and older with the pullbate and older.
	with phenylketonuria (PKU)
Required Medical	Documentation of a diagnosis of PKU
Information:	Baseline (pre-treatment) blood Phe level greater than or equal to 360 micromol/L (6)
	mg/dL)
	Documentation of failure to Phe restricted diet as monotherapy
Appropriate	Documentation of continuation on a Phe restricted diet
Treatment	
Regimen & Other	Reauthorization requires documentation of one of the following:
Criteria:	 Reduction in baseline Phe levels by 30 percent or levels maintained between 120 - 360
	micromol/L (2 - 6 mg/dL)
	Increase in dietary Phe tolerance
	Improvement in clinical symptoms
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a specialist in metabolic disorders or an
Care Restrictions:	endocrinologist
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 2 months, unless otherwise specified
-	Reauthorization: 12 months, unless otherwise specified



SATRALIZUMAB-MWGE

Affected Medications: ENSPRYNG (satralizumab-mwge)

Covered Uses: Required Medical Information:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Neuromyelitis optica spectrum disorder (NMOSD) in adults who are antiaquaporin-4 (AQP4) antibody positive NMOSD Diagnosis of seropositive aquaporin-4 immunoglobulin G (AQP4-IgG) NMOSD confirmed by all the following:
	 Acute optic neuritis Acute myelitis Acute area postrema syndrome (episode of otherwise unexplained hiccups or nausea/vomiting) Acute brainstem syndrome Symptomatic narcolepsy OR acute diencephalic clinical syndrome with NMOSD-typical diencephalic lesion on magnetic resonance imaging (MRI) [see table below] Acute cerebral syndrome with NMOSD-typical brain lesion on MRI [see table below]
	Clinical presentation Possible MRI findings
	Diencephalic syndrome Periependymal lesion Hypothalamic/thalamic lesion
	Acute cerebral syndrome Extensive periependymal lesion Long, diffuse, heterogenous, or edematous corpus callosum lesion Long corticospinal tract lesion Large, confluent subcortical or deep white matter lesion
	History of at least 1 attack in the past year, or at least 2 attacks in the past 2 years, requiring rescue therapy
Appropriate Treatment Regimen &	Documented inadequate response, contraindication, or intolerance to rituximab (preferred agents Riabni and Ruxience)
Other Criteria:	Reauthorization requires documentation of treatment success
Exclusion Criteria:	 Active Hepatitis B Virus (HBV) infection Active or untreated latent tuberculosis Concurrent with other disease-modifying biologics for requested indication
Age Restriction:	18 years of age and older



Prescriber/Site of Care Restrictions:		Prescribed by, or in consultation with, a neurologist or neuro-ophthalmologist All approvals are subject to utilization of the most cost-effective site of care
Coverage	•	Initial Authorization: 6 months, unless otherwise specified
Duration:	•	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **SEBELIPASE ALFA**

Affected Medications: KANUMA (sebelipase alfa)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	Treatment of Lysosomal Acid Lipase (LAL) deficiency
Required Medical Information:	 Diagnosis of LAL deficiency or Rapidly Progressive LAL deficiency within the first 6 months of life confirmed by one of the following: Absence or deficiency in lysosomal acid lipase activity
	 Mutation in the lipase A, lysosomal acid type (LIPA) gene
	Documentation of patient weight
	Documentation of prescribed treatment regimen (dose and frequency)
	Baseline fasting lipid panel including LDL-c prior to initiating therapy (not required for Rapidly Progressive LAL deficiency)
Appropriate	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced
Treatment	
Regimen & Other	Reauthorization:
Criteria:	Rapidly Progressive LAL deficiency: documentation of improvement in weight-for-age Z-score
	LAL deficiency: documentation of improvement in LDL-c
Exclusion Criteria:	
Age Restriction:	1 month of age or older
Prescriber/Site of	Prescribed by, or in consultation with, an endocrinologist or metabolic specialist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 3 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



SELF-ADMINISTERED DRUGS (SAD)

Affected Medications: Please refer to package insert for directions on self-administration.

Covered Uses:	
Required Medical Information:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
Appropriate Treatment Regimen & Other	Pharmaceuticals covered under your pharmacy benefit are in place of, not in addition to, those same covered supplies under the medical plan. Please refer to your benefit book for more information.
Criteria: Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of Care Restrictions:	
Coverage Duration:	



POLICY NAME: **SELUMETINIB**

Affected Medications: KOSELUGO (selumetinib)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Neurofibromatosis type 1 with symptomatic, inoperable plexiform neurofibromas in pediatric patients 2 years of age and older NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A
	or better
Required Medical Information:	Documented body surface area (BSA) and requested dose
	 Neurofibromatosis type 1 (NF1) with inoperable plexiform neurofibromas Documentation of diagnosis of symptomatic and/or progressive, inoperable NF1, defined as one or more plexiform neurofibromas that cannot be completely removed without risk for substantial morbidity due to encasement of, or close proximity to, vital structures, invasiveness, or high vascularity Documentation of 2 or more of the following clinical diagnostic criteria as evaluated by a multidisciplinary specialist care team (A child of a parent with NF1 can be diagnosed if one or more of these criteria are met): Six or more café-au-lait macules over 5 mm in greatest diameter in prepubertal individuals and over 15 mm in greatest diameter in post pubertal individuals Freckling in the axillary or inguinal region Two or more neurofibromas of any type or one plexiform neurofibroma Optic pathway glioma
	 Two or more iris Lisch nodules identified by slit lamp examination or two or more choroidal abnormalities A distinctive osseous lesion such as sphenoid dysplasia, anterolateral bowing of the tibia, or pseudarthrosis of a long bone A heterozygous pathogenic NF1 variant with a variant allele fraction of 50% in apparently normal tissue such as white blood cells NCCN Indications Documentation of performance status, disease staging, all prior therapies used, and anticipated treatment course
Appropriate Treatment Regimen & Other Criteria:	Reauthorization: documentation of disease responsiveness to therapy For NF1: defined as a decrease in tumor volume from baseline and improvement in symptoms, such as pain
Exclusion Criteria:	NCCN Indications ■ Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
Age Restriction:	Neurofibromatosis type 1 (NF1) with inoperable plexiform neurofibromas • 2 to 18 years of age



Prescriber/Site of Care Restrictions:	Neurofibromatosis type 1 (NF1) with inoperable plexiform neurofibromas Prescribed by, or in consultation with, a pediatric oncologist or specialist with experience in the treatment of neurofibromatosis
	 NCCN Indications Prescribed by, or in consultation with, an oncologist
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **SEROSTIM**

Affected Medications: SEROSTIM (somatropin)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	o HIV (human immunodeficiency virus)-associated wasting, cachexia
Required Medical	Documentation of current body mass index (BMI), actual body weight, and ideal body
Information:	weight (IBW)
	Serostim is used in combination with antiretroviral therapy to which the patient has
	documented compliance
	Alternative causes of wasting (e.g., inadequate nutrition intake, malabsorption,
	opportunistic infections, hypogonadism) have been ruled out or treated appropriately
	Prior to somatropin, patient had a suboptimal response to at least 1 other therapy for
	wasting or cachexia (e.g., megestrol, dronabinol, cyproheptadine, or testosterone
	therapy if hypogonadal) unless contraindicated or not tolerated
	Diagnosis of HIV-association wasting syndrome or cachexia confirmed by one of the
	following:
	 Unintentional weight loss greater than or equal to 10% of body weight over prior
	12 months
	 Unintentional weight loss greater than or equal to 5% of body weight over prior 6
	months
	○ BMI less than 20 kg/m²
	 Weight is less than 90% of IBW
Appropriate	Reauthorization:
Treatment	Documentation of treatment success and clinically significant response to therapy (e.g.,
Regimen & Other Criteria:	improved or stabilized BMI, increased physical endurance compared to baseline, etc.)
Criteria:	
	Documentation of continued compliance to antiretroviral regimen
Exclusion Criteria:	Acute critical illness due to complications following open heart or abdominal surgery,
	multiple accidental trauma or acute respiratory failure
	Active malignancy
	Acute respiratory failure
	Active proliferative or severe non-proliferative diabetic retinopathy
Age Restriction:	
Dreceriber/Cite of	Described by an in approximation with an infrared at Process and Pro-
Prescriber/Site of	Prescribed by, or in consultation with, an infectious disease specialist All approvals are subject to utilization of the most cost effective site of core.
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an infectious disease specialist All approvals are subject to utilization of the most cost-effective site of care
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care



POLICY NAME: **SIGNIFOR**

Affected Medications: SIGNIFOR (pasireotide)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Cushing's disease
Required Medical Information:	Documented diagnosis of Cushing's disease Documentation of at least TWO of the following:
Appropriate Treatment Regimen & Other Criteria:	 Documented treatment failure or intolerable adverse event to ketoconazole and cabergoline Documentation confirming pituitary surgery is not an option OR previous surgery has not been curative Reauthorization requires documentation of treatment success defined as mUFC normalization (i.e., less than or equal to the ULN)
Exclusion Criteria:	Severe hepatic impairment (Child Pugh C)
Age Restriction:	18 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an endocrinologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: SIGNIFOR LAR

Affected Medications: SIGNIFOR LAR (pasireotide)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	o Acromegaly
	Cushing's disease
Required	Acromegaly
Medical	Documentation confirming clinical manifestations of disease
Information:	Diagnosis of acromegaly confirmed by ONE of the following:
	 Elevated pre-treatment serum insulin-like growth factor-1 (IGF-1) level for
	age/gender
	 Serum growth hormone (GH) level of 1 microgram/mL or greater after an oral
	glucose tolerance test (OGTT)
	Cushing's Disease
	Documented diagnosis of Cushing's disease
	Documentation of at least TWO of the following:
	 Mean 24-hour urine free cortisol (mUFC) greater than 1.5 times the upper limit of
	normal (ULN) for the assay (at least two measurements)
	 Bedtime salivary cortisol greater than 145 ng/dL (at least two measurements)
	 Overnight dexamethasone suppression test (DST) with a serum cortisol greater
	than 1.8 mcg/dL
Appropriate	Acromegaly
Treatment	Documented treatment failure or intolerance to ONE of the following: lanreotide
Regimen & Other Criteria:	(Somatuline Depot), Sandostatin LAR, or pegvisomant (Somavert)
Other Criteria.	Documentation confirming ONE of the following:
	 Inadequate response to surgery or radiotherapy
	 Not a candidate for surgical management or radiotherapy (e.g., medically
	unstable, high risk for complications under anesthesia, major systemic
	complications of acromegaly, severe hypertension, uncontrolled diabetes, etc.)
	Dosing: Not to exceed 60 mg every 4 weeks (after 3 months of 40 mg)
	Reauthorization requires documentation of treatment success shown by
	Reauthorization requires documentation of treatment success shown by decreased/normalized IGF-1 or GH levels
	decreased/normalized IGF-1 or GH levels
	decreased/normalized IGF-1 or GH levels Cushing's Disease
	decreased/normalized IGF-1 or GH levels Cushing's Disease Documentation confirming pituitary surgery is not an option OR previous surgery has not
	decreased/normalized IGF-1 or GH levels Cushing's Disease Documentation confirming pituitary surgery is not an option OR previous surgery has not been curative
	Cushing's Disease Documentation confirming pituitary surgery is not an option OR previous surgery has not been curative Documented treatment failure or intolerance to ketoconazole and cabergoline
	decreased/normalized IGF-1 or GH levels Cushing's Disease Documentation confirming pituitary surgery is not an option OR previous surgery has not been curative



	Reauthorization requires documentation of treatment success defined as mUFC normalization (i.e., less than or equal to the ULN)
Exclusion Criteria:	Severe hepatic impairment (Child Pugh C)
Age Restriction:	18 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an endocrinologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **SILTUXIMAB**

Affected Medications: SYLVANT (siltuximab)

Required Medical Information:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of patients with multicentric Castleman's disease (MCD) who are human immunodeficiency virus (HIV) negative and human herpesvirus-8 (HHV-8) negative NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher Documentation of performance status, disease staging, all prior therapies used, and anticipated treatment course The diagnosis was confirmed by biopsy of lymph gland Documented negative tests for HIV and HHV-8
Appropriate Treatment Regimen & Other Criteria:	 Patient weight Dosing MCD: 11 mg/kg intravenous (IV) infusion once every 3 weeks until treatment failure Cytokine release syndrome (CRS): 11 mg/kg IV one time only Availability: 100 mg and 400 mg vials Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced
Exclusion Criteria:	Reauthorization requires documentation of disease responsiveness to therapy
Age Restriction:	18 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an oncologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 MCD: Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified CRS: 1 month (1 dose only), unless otherwise specified



POLICY NAME: **SIPONIMOD**

Affected Medications: MAYZENT (siponimod)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of relapsing forms of multiple sclerosis (MS), including the following: Clinically isolated syndrome (CIS)
	 Relapsing-remitting multiple sclerosis (RRMS) Active secondary progressive multiple sclerosis (SPMS)
Required Medical Information:	Diagnosis confirmed with magnetic resonance imaging (MRI), per revised McDonald diagnostic criteria for MS Clinical evidence alone will suffice; additional evidence desirable but must be consistent with MS
Appropriate Treatment Regimen & Other Criteria:	Coverage of Mayzent (siponimod) requires documentation of one of the following: Documented disease progression or intolerable adverse event with one of the following: teriflunomide, dimethyl fumarate or fingolimod Currently receiving treatment with Mayzent (siponimod), excluding via samples or manufacturer's patient assistance program
	Reauthorization requires provider attestation of treatment success
Exclusion Criteria:	 Presence of CYP2C9*3/*3 genotype Concurrent use of other disease-modifying medications indicated for the treatment of MS
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist or a MS specialist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



SODIUM PHENYLBUTYRATE

Affected Medications: SODIUM PHENYLBUTYRATE

Covered Uses:	
Covered Oses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Adjunctive therapy in the chronic management of patients with urea cycle disorders (UCDs) involving deficiencies of carbamylphosphate synthetase (CPS), ornithine transcarbamylase (OTC), or argininosuccinic acid synthetase (AS) Neonatal-onset deficiency (complete enzymatic deficiency, presenting within the first 28 days of life) Late-onset disease (partial enzymatic deficiency, presenting after the first month of life) with history of hyperammonemic encephalopathy
Required Medical Information:	Diagnosis confirmed by blood, enzymatic, biochemical, or genetic testing
Appropriate Treatment Regimen & Other Criteria:	 Oral tablets require documented inability to use sodium phenylbutyrate powder Documented treatment failure with dietary protein restriction and/or amino acid supplementation alone Must be used in combination with dietary protein restriction Reauthorization will require BOTH of the following: Documentation of treatment success defined as ammonia levels maintained within normal limits That this drug continues to be used in combination with dietary protein restriction
Exclusion Criteria:	Use for management of acute hyperammonemia
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a specialist experienced in the treatment of metabolic diseases All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: **SOLRIAMFETOL**

Affected Medications: SUNOSI (solriamfetol)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Excessive daytime sleepiness associated with narcolepsy
	Excessive daytime sleepiness associated with obstructive sleep apnea
Required Medical	<u>Narcolepsy</u>
Information:	Diagnosis confirmed by polysomnography and multiple sleep latency test
	Symptoms of excessive daytime sleepiness consistent with narcolepsy have been
	present for at least 3 months
	An Epworth Sleepiness Scale score of more than 10 despite treatment
	Obstructive Sleep Apnea (OSA)
	Diagnosis confirmed by sleep study
	An Epworth Sleepiness Scale score of more than 10 despite drug treatment and current
	use of continuous positive airway pressure (CPAP) for at least 3 months
	Documentation that CPAP use will be continued during treatment with solriamfetol
	All indications:
	Documentation that other causes of sleepiness have been treated or ruled out (including)
	but not limited to insufficient sleep syndrome, shift work, the effects of substances or
	medications, or other sleep disorders)
Appropriate	Documented trial and failure or contraindication to modafinil OR armodafinil
Treatment	For narcolepsy only, documented trial and failure or contraindication to ONE of the
Regimen & Other	following: methylphenidate, dextroamphetamine, lisdexamfetamine, amphetamine-
Criteria:	dextroamphetamine
	Reauthorization requires clinically significant improvement in activities of daily living and in
	Epworth Sleepiness Scale score
Exclusion Criteria:	Use for other untreated causes of sleepiness
	Concurrent use of sedative/hypnotic drugs or other central nervous system depressants
Age Restriction:	18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a sleep specialist or neurologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 4 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



SOMATOSTATIN ANALOGS

Affected Medications: OCTREOTIDE, SANDOSTATIN LAR, LANREOTIDE, SOMATULINE DEPOT (lanreotide)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by Plan design.
	Octreotide, Sandostatin LAR: Acromegaly Symptomatic treatment of metastatic carcinoid tumors (carcinoid syndrome) Symptomatic treatment of vasoactive intestinal peptide tumors (VIPomas) Lanreotide, Somatuline Depot: Acromegaly Carcinoid syndrome (to reduce the frequency of short-acting somatostatin analog rescue therapy) Unresectable, well- or moderately-differentiated, locally advanced or metastatic gastroenteropancreatic neuroendocrine tumors (GEP-NETs) NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher
Required Medical Information:	Acromegaly Documentation confirming clinical manifestations of disease Diagnosis of acromegaly confirmed by ONE of the following: Elevated pre-treatment serum insulin-like growth factor-1 (IGF-1) level for age/gender Serum growth hormone (GH) level of 1 microgram/mL or greater after an oral glucose tolerance test (OGTT)
	All other indications Documentation of performance status, disease staging, all prior therapies used, and anticipated treatment course
Appropriate Treatment Regimen & Other Criteria:	Acromegaly Documentation confirming ONE of the following: Inadequate response to surgery or radiotherapy Not a candidate for surgical management or radiotherapy (e.g., medically unstable, high risk for complications under anesthesia, major systemic complications of acromegaly, severe hypertension, uncontrolled diabetes, etc.)
	 Sandostatin LAR Coverage for the non-preferred product Sandostatin LAR is provided when ONE of the following criteria is met: Currently receiving treatment with Sandostatin LAR, excluding when the product is obtained as samples or via manufacturer's patient assistance programs Documented inadequate response or intolerable adverse event with one of the following: Lanreotide, Somatuline Depot, OR Somavert (Note: Somavert indicated for acromegaly only)



	Lanreotide, Somatuline Depot
	GEP-NETs must use 120 mg injection
	Reauthorization: • Acromegaly: requires documentation of treatment success shown by decreased/normalized IGF-1 or GH levels
	All other indications: requires documentation of disease responsiveness to therapy
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist, endocrinologist, or
Care Restrictions:	gastroenterologist
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **SOMAVERT**

Affected Medications: SOMAVERT (pegvisomant)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design	
Required Medical Information:	Documentation confirming clinical manifestations of disease Diagnosis of acromegaly confirmed by ONE of the following:	
Appropriate Treatment Regimen & Other Criteria:	Documented treatment failure or intolerance to octreotide or lanreotide (Somatuline Depot) Documentation confirming one of the following:	
Exclusion Criteria:		
Age Restriction:		
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an endocrinologist All approvals are subject to utilization of the most cost-effective site of care 	
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified 	



SOTATERCEPT-CSRK

Affected Medications: WINREVAIR (sotatercept-csrk)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise excluded by plan design	
	 Pulmonary Arterial Hypertension (PAH) World Health Organization (WHO) Group 1 	
Required Medical Information:	 Documentation of PAH confirmed by right-heart catheterization meeting the following criteria: Mean pulmonary artery pressure of at least 20 mm Hg 	
	 Pulmonary capillary wedge pressure less than or equal to 15 mm Hg Pulmonary vascular resistance of at least 5 Wood units 	
	Etiology of PAH: idiopathic PAH, hereditary PAH OR	
	 PAH secondary to one of the following conditions: Connective tissue disease 	
	 Simple, congenital systemic to pulmonary shunts at least 1 year following repair Drugs and toxins 	
	New York Heart Association (NYHA)/World Health Organization (WHO) Functional Class II or III symptoms	
	Documentation of Acute Vasoreactivity Testing (positive result requires trial/failure to calcium channel blockers) unless there are contraindications:	
	 Low systemic blood pressure (systolic blood pressure less than 90) Low cardiac index (cardiac index less than 2 L/min/m²) OR 	
	 Presence of severe symptoms (functional class IV) Baseline 6-minute walk test (6MWD) 	
Appropriate	Documentation that drug will be used as an add-on treatment with all the following (one	
Treatment Regimen & Other Criteria:	from each category) at therapeutic doses for at least 90 days: o Phosphodiesterase-5 (PDE-5) inhibitor: sildenafil, tadalafil o Endothelin Receptor Antagonist: ambrisentan, bosentan, Opsumit o Prostacyclin: treprostinil, epoprostenol, Ventavis	
	Documentation of inadequate response or intolerance to oral calcium channel blocking agents (nifedipine, diltiazem) if positive Acute Vasoreactivity Test	
	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced	
	Reauthorization requires documentation of treatment success defined as one or more of the following:	
	Improvement in walking distance (6MWD)	
Exclusion Criteria:	 Improvement or stability in WHO functional class Human immunodeficiency virus (HIV)-associated PAH 	
	PAH associated with portal hypertension	
	Schistosomiasis-associated PAH	
	Pulmonary veno occlusive disease	
	Platelet count less than 50,000/mm³ (50 x 109/L)	
	Hemoglobin (Hgb) at screening above gender-specific upper limit of normal (ULN)	



Age Restriction:	18 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a cardiologist or pulmonologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: SPARSENTAN

Affected Medications: FILSPARI (sparsentan)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Reduce proteinuria in adults with primary immunoglobulin A nephropathy (IgAN) at risk of rapid disease progression
Required Medical Information:	 Diagnosis of primary immunoglobulin A nephropathy (IgAN) confirmed with biopsy Documentation of ONE of the following (with labs current within 30 days of request): Proteinuria defined as equal to or greater than 1 g/day Urine protein-to-creatinine ratio (UPCR) greater than 1.5 g/g
Appropriate Treatment Regimen & Other Criteria:	Documented treatment failure (defined as proteinuria equal to or greater than 1 g/day OR UPCR greater than 1.5 g/g) with a minimum of 12 weeks of each of the following: Maximum tolerated dose of an angiotensin-converting enzyme (ACE) inhibitor or angiotensin receptor blocker (ARB) High dose glucocorticoid therapy such as oral prednisone or methylprednisolone (or an adverse effect to two or more glucocorticoid therapies that is not associated with the corticosteroid class)
Exclusion Criteria:	 Hepatic impairment (Child-Pugh class A-C) Estimated glomerular filtration rate (eGFR) that is less than 30 mL/min/1.73 m²
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a nephrologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **SPESOLIMAB**

Affected Medications: SPEVIGO INTRAVENOUS (IV) SOLUTION

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Generalized pustular psoriasis flares (GPP, also called von Zumbusch psoriasis) 		
Required Medical Information:	 Diagnosis of generalized pustular psoriasis as confirmed by the following: The presence of widespread sterile pustules arising on erythematous skin Pustulation is not restricted to psoriatic plaques Signs and symptoms of an acute GPP flare of moderate-to-severe intensity as follows: A Generalized Pustular Psoriasis Physician Global Assessment (GPPGA) total score of greater than or equal to 3 A GPPGA pustulation category subscore of greater than or equal to 2 Greater than or equal to 5% body surface area (BSA) covered with erythema and the presence of pustules 		
Appropriate Treatment Regimen & Other Criteria:	Documented treatment failure of acute disease flare (or documented intolerable adverse event) with: A one-week trial of cyclosporine AND Infliximab (preferred biosimilars Inflectra, Renflexis) Treatment for each flare is limited to two 900 mg infusions of Spevigo separated by 1 week		
Exclusion Criteria:	 Previous use of Spevigo Erythrodermic plaque psoriasis without pustules or with pustules restricted to psoriatic plaques Synovitis-acne-pustulosis-hyperostosis-osteitis syndrome Drug-induced acute generalized exanthematous pustulosis 		
Age Restriction:	18 years of age and older		
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a dermatologist All approvals are subject to utilization of the most cost-effective site of care 		
Coverage Duration:	Authorization: 1 month with no reauthorization, unless otherwise specified		



POLICY NAME: **SPRAVATO**

Affected Medications: SPRAVATO (esketamine nasal spray)

Covered Uses: Required Medical Information:	plan design	junction with an oral antidep nt-resistant depression (TRI ive symptoms in adults with te suicidal ideation or behaves sistant Depression (TRD) risk for abuse or misuse	major depressive disorder (: MDD)
	Diagnosis of Major Depressibehavior: Assessment of patient's of Montgomery-Asberg Depressions	sive Disorder (MDD) with a risk for abuse or misuse pression Rating Scale (MAD		28,
Appropriate			<u> </u>	
Treatment Regimen & Other Criteria:	 Treatment-Resistant Depression: Documented treatment failure (defined by less than 50% improvement in depression symptom severity using a standard rating scale such as a PHQ-9) to an adequate trial least 6 weeks each), or intolerance, of at least three antidepressants from at least two different classes, during the current depressive episode Failure to respond to augmentation therapy such as:		e trial (at st two htly htly ral	
		T		
	Induction Phase	Weeks 1 to 4	Day 1 starting dose: 56 mg	
		Administer twice per week	Subsequent doses: 56 mg or 84 mg	
	Maintenance Phase	Weeks 5 to 8		
		Administer once weekly	56 mg or 84 mg]



		Weeks 9 and after		
		Administer every 2 weeks or once weekly*	56 mg or 84 mg	
	*Dosing frequency should be individualized to the least frequent dosing to maintain remission/response			
	depression compared to symptoms Spravato continues to be Major depressive disorder (Documentation of current documentation of why pa Spravato will be used in compared to symptoms.	ent success defined as at le baseline using a standard raused in combination with a used in combination with a (MDD) with acute suicidal inpatient psychiatric hospit tient is not currently at inpaticombination with an oral antikly for 4 weeks maximum (1)	ideation or behavior: calization OR adequate tient level of care tidepressant	
Exclusion Criteria:	 Concomitant psychotic di Bipolar or related disorde History of substance use 	ers		
	Use as an anesthetic agePregnancy	ent		
	peripheral arterial vesselsHistory of intracerebral he	s) or arteriovenous malform		ii, and
Age Restriction:	18 years of age and olde	r		
Prescriber/Site of Care Restrictions:		ultation with, a psychiatrist value to utilization of the most cost		
Coverage Duration:	#24 nasal spray devicesTRD: 2 months (Induction	in 28 days of treatment only n phase – maximum of 23 n maintenance phase), unles		,



POLICY NAME: **STIRIPENTOL**

Affected Medications: DIACOMIT (stiripentol)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of seizures associated with Dravet syndrome (DS) 	
Required Medical Information:	 Current Weight Documentation that therapy is being used as adjunct to clobazam for seizures Documentation of at least 4 generalized clonic or tonic-clonic seizures in the last month while on stable antiepileptic drug therapy 	
Appropriate Treatment Regimen & Other Criteria:	Documented treatment and inadequate control of seizures with at least four guideline directed therapies including:	
Exclusion Criteria:		
Age Restriction:	6 months of age and older	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a neurologist All approvals are subject to utilization of the most cost-effective site of care 	
Coverage Duration:	Authorization: 12 months, unless otherwise specified	



POLICY NAME: **STRENSIQ**

Affected Medications: STRENSIQ (asfotase alfa)

	7		
Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by		
	plan design		
	 Perinatal/infantile or Juvenile onset hypophosphatasia (HPP) 		
Required Medical	Diagnosis of Perinatal/Infantile or Juvenile onset hypophosphatasia (HPP) with ALL of		
Information:	the following:		
	Age of onset less than 18 years		
	One of the following:		
	 Clinical manifestations consistent with hypophosphatasia at onset prior to age 18, such as: vitamin B6 dependent seizures, respiratory insufficiency, failure to thrive, non-traumatic fracture, dental abnormalities, low score on 6-minute walk 		
	test, low bone density score		
	 Skeletal abnormalities confirmed with radiographic imaging (such as flared and frayed metaphyses, widened growth plate, bowed arms or legs, rachitic chest deformity, craniosynostosis) 		
	Genetic test confirming mutation of tissue-non-specific alkaline phosphatase (TNSALP) gene		
	Low level of serum alkaline phosphatase (ALP) evidenced by lab result below reference		
	range for patient's age and genderElevated levels of one of the following:		
	 Elevated levels of one of the following: Urine or serum concentration of phosphoethanolamine (PEA) 		
	 Serum concentration of phosphoethanolamine (PEA) Serum concentration of pyridoxal 5'-phosphate (PLP) in the absence of vitamin 		
	supplements within one week prior to the test		
	 Urinary inorganic pyrophosphate (PPi) 		
	Simaly morganic pyrophicophiate (i. 1.)		
Appropriate	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced.		
Treatment Regimen & Other Criteria:	 Please note: the 80 mg/0.8 mL vial is for patients weighing greater than 40 kilograms only 		
	Poputhorization requires decumentation of:		
	 Reauthorization requires documentation of: Laboratory results confirming a decrease in urine concentration of urine or serum 		
	phosphoethanolamine (PEA), serum concentration of pyridoxal 5'-phosphate (PLP), or		
	urinary inorganic pyrophosphate (PPi)		
	Improvement or stabilization in the clinical signs and symptoms of hypophosphatasia,		
	such as:		
	 Radiographic evidence of improvement in skeletal deformities or growth 		
	 Improvement in 6-minute walk test 		
	 Improved bone density 		
	o Reduction in fractures		
	 Respiratory function/breathing 		
	 Improvement in developmental milestones 		



Exclusion Criteria:	Other types of osteomalacia or hypophosphatasia, including adult onset hypophosphatasia
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an endocrinologist or specialist experienced in the treatment of metabolic bone disorders All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



SUBCUTANEOUS IMMUNE GLOBULIN

Affected Medications: CUTAQUIG, CUVITRU, GAMUNEX-C, HIZENTRA, HYQVIA, XEMBIFY

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	Primary immunodeficiency (PID)/Wiskott-Aldrich syndrome
	Such as: x-linked agammaglobulinemia, common variable
	immunodeficiency (CVID), transient hypogammaglobulinemia of infancy,
	immunoglobulin G (IgG) subclass deficiency with or without
	immunoglobulin A (IgA) deficiency, antibody deficiency with near normal
	immunoglobulin levels) and combined deficiencies (severe combined
	immunodeficiencies, ataxia-telangiectasia, x-linked lymphoproliferative
	syndrome) [list not all inclusive]
	 Chronic Inflammatory Demyelinating Polyneuropathy (CIDP)
	o Thomathinatory Demyelinating Folyhedropathy (Olbi)
Required Medical	Monthly intravenous immune globulin (IVIG) dose for those transitioning
Information:	Patient weight
	Primary Immunodeficiency (PID)
	Type of immunodeficiency
	Documentation of one of the following:
	 Recent IgG level less than 200
	 Low IgG levels (below the laboratory reference range lower limit of normal) AND
	a history of multiple hard to treat infections as indicated by at least one of the
	following:
	 Four or more ear infections within 1 year
	 Two or more serious sinus infections within 1 year
	Two or more months of antibiotics with little effect
	Two or more pneumonias within 1 year
	Recurrent or deep skin abscesses
	Need for intravenous antibiotics to clear infections
	Two or more deep-seated infections including septicemia
	Documentation showing a deficiency in producing antibodies in response to vaccination including all of the following:
	including all of the following:
	 Titers that were drawn before challenging with vaccination
	 Titers that were drawn between 4 and 8 weeks after vaccination
	Chronic Inflammatory Demyelinating Polyneuropathy (CIDP)
	Documented baseline in strength/weakness has been documented using an objective
	clinical measuring tool (INCAT, Medical Research Council (MRC) muscle strength, 6
	Minute Walk Test, Rankin, Modified Rankin)
	Documented disease course is progressive or relapsing and remitting for 2 months or
	longer
	Abnormal or absent deep tendon reflexes in upper or lower limbs
	1 - Abriornial of absent deep tendori reflexes in upper of lower littles



	Electrodiagnostic evidence of demyelination indicated by one of the following:
	 Motor distal latency prolongation in 2 nerves
	 Reduction of motor conduction velocity in 2 nerves
	 Prolongation of F-wave latency in 2 nerves
	 Absence of F-waves in at least 1 nerve
	 Partial motor conduction block of at least 1 motor nerve
	 Abnormal temporal dispersion in at least 2 nerves
	 Distal CMAP duration increase in at least 1 nerve
	Cerebrospinal fluid (CSF) analysis indicates all of the following (if electrophysiologic
	findings are non-diagnostic):
	 CSF white cell count of less than 10 cells/mm³
	 CSF protein is elevated (greater than or equal to 45mg/dL)
Appropriate	Meets all criteria for IVIG approval
Treatment	Exceptions may be given for patients without prior intravenous (IV) or subcutaneous
Regimen & Other	(SC) immune globulin use
Criteria:	DID
	 PID Documentation of at least 3 months of IVIG therapy
	CIDP
	HyQvia, Hizentra and Gamunex-c only
	Refractory to or intolerant of corticosteroids (prednisolone, prednisone) given in
	therapeutic doses over at least three months
	Reauthorization:
	PID: Documented disease response defined as a decrease in the frequency or severity
	of infections • CIDP:
	 Documentation of a beneficial clinical response to maintenance therapy, without
	relapses, based on an objective clinical measuring tool
	OR
	 Re-initiating maintenance therapy after experiencing a relapse while on Hizentra;
	AND documented improvement and stability on IVIG treatment AND was NOT
Fredrice Onitonia	receiving maximum dosing of Hizentra prior to relapse
Exclusion Criteria:	IgA deficiency with antibodies to IgA
	History of hypersensitivity to immune globulin or product components Hypersystinamic type Lev II.
Age Restriction:	 Hyperprolinemia type I or II PID: 2 years of age and older
Age Restriction.	CIDP: 18 years of age and older
	OIDI : 10 years or age and older
Prescriber/Site of	PID: prescribed by, or in consultation with, an immunologist
Care Restrictions:	CIDP: prescribed by, or in consultation with, a neurologist or rheumatologist with CIDP
	expertise
Coverage Duration:	Initial Authorization:
	CIDP: 3 months, unless otherwise specified
	PID: 12 months, unless otherwise specified



Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **SUTIMLIMAB**

Affected Medications: ENJAYMO (sutimlimab-jome)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of hemolysis in adults with cold agglutinin disease (CAD)	
Required Medical Information:	 Cold Agglutinin Disease (CAD) Documentation of current weight Diagnosis of CAD as confirmed by all of the following: Chronic hemolysis as confirmed by hemoglobin level of 10 g/dL or less AND elevated indirect bilirubin level Positive monospecific direct antiglobulin test (DAT) or Coombs test for C3d A positive DAT or Coombs test for IgG of 1+ or less Cold agglutinin titer of greater than or equal to 64 at 4°C 	
Appropriate Treatment Regimen & Other Criteria:	Cold Agglutinin Disease (CAD) ■ Dosing: □ 39 kg to less than 75 kg: 6,500 mg/dose □ 75 kg or greater: 7,500 mg/dose □ Administered weekly for the first two weeks, then every two weeks thereafter Reauthorization: documentation of disease responsiveness to therapy (e.g., increased hemoglobin, normalized markers of hemolysis [bilirubin, lactate dehydrogenase, reticulocyte count], reduced blood transfusion requirements)	
Exclusion Criteria:	Disease secondary to infection, rheumatologic disease, systemic lupus erythematosus, or overt hematologic malignancy Concomitant use of rituximab with or without cytotoxic agents	
Age Restriction:	18 years of age or older	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a hematologist All approvals are subject to utilization of the most cost-effective site of care 	
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified 	



TTR STABILIZERS

Affected Medications: VYNDAQEL (tafamidis meglumine 20 mg), VYNDAMAX (tafamidis 61 mg), ATTRUBY (acoramidis hydrochloride)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by		
	plan design		
	Treatment of wild type or hereditary transthyretin amyloid cardiomyopathy		
	(ATTR-CM) to reduce cardiovascular mortality and cardiovascular-related		
	hospitalizations in adults		
	Troophanization in addition		
Required Medical Information:	Diagnosis of ATTR-CM supported by ONE of the following (a, b, or c): a. Cardiac tissue biopsy confirms presence of ATTR amyloid deposits by		
	immunohistochemistry (IHC) or mass spectrometry		
	b. Documentation of BOTH of the following (i and ii):		
	i. Noncardiac tissue biopsy confirms presence of ATTR amyloid deposits		
	by IHC or mass spectrometry		
	ii. Imaging consistent with cardiac amyloidosis (echocardiogram [ECG],		
	cardiac magnetic resonance [CMR], or positron emission tomography		
	[PET])		
	c. Documentation of ALL the following (i, ii, and iii):		
	i. Grade 2 to 3 uptake on cardiac scintigraphy (utilizing Tc-PYP, Tc-DPD,		
	or Tc-HMDP radiotracers)		
	ii. Normal serum kappa/lambda free light chain (sFLC) ratio, serum protein		
	immunofixation, AND urine protein immunofixation		
	iii. Imaging consistent with cardiac amyloidosis (ECG, CMR, or PET)		
	Documentation of New York Heart Association (NYHA) Functional Class I to III		
Appropriate	Coverage for Vyndaqel or Vyndamax is provided when the following is met:		
Treatment	Documented treatment failure with Attruby (acoramidis)		
Regimen & Other			
Criteria:	Reauthorization requires documentation of disease responsiveness (improvement in		
	symptoms, quality of life, or 6-Minute Walk Test; slowing or stabilization of disease		
	progression; reduced cardiovascular-related hospitalizations, etc.)		
Exclusion Criteria:	NYHA Functional Class IV heart failure		
	Presence of light-chain (primary) amyloidosis		
	Prior liver or heart transplant		
	Implanted cardiac mechanical assist device		
	Combined use with another TTR stabilizer or TTR silencer (such as eplontersen,		
	patisiran, vultrisiran)		
	1 ,		
Age Restriction:	18 years of age and older		
Prescriber/Site of	Prescribed by, or in consultation with, a cardiologist or specialist experienced in the		
Care Restrictions:	treatment of amyloidosis		
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All approvals are subject to utilization of the most cost-effective site of care	
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
Reauthorization: 12 months, unless otherwise specified	



TAGRAXOFUSP-ERZS

Affected Medications: ELZONRIS (tagraxofusp-erzs)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of blastic plasmacytoid dendritic cell neoplasm (BPDCN) in adults and in pediatric patients at least 2 years of age NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or better
Required Medical Information:	Diagnosis of BPDCN is confirmed by ALL of the following: A biopsy showing the morphology of plasmacytoid dendritic blast cells At least 3 of the following plasmacytoid dendritic cell (pDC) markers are expressed by immunohistochemistry (IHC) or flow cytometry: CD123 CD4 CD56 TCF4 TCL1 CD303 CD304 The following pDC markers are negative: CD3, CD14, CD19, CD34, lysozyme, myeloperoxidase Diagnosis is made by a board-certified hematopathologist or dermatopathologist Documentation of performance status, disease staging, all prior therapies used, and anticipated treatment course
Appropriate Treatment Regimen & Other Criteria:	Reauthorization requires documentation of disease responsiveness to therapy
Exclusion Criteria:	Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater Pregnancy
Age Restriction:	2 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a prescriber experienced in the treatment of BPDCN All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



TARGETED IMMUNE MODULATORS

PA Policy Applicable to:

Preferred Drugs: Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Enbrel, Cosentyx, Otezla, Tremfya, Stelara, Xeljanz, Skyrizi, Rinvoq

Preferred Medical Drugs: Inflectra, Renflexis, Skyrizi Intravenous, Stelara, Simponi Aria Intravenous, Tofidence Intravenous, Tyenne Intravenous, Tremfya Intravenous

Non-preferred Medical Drugs: Remicade, Entyvio, Orencia Intravenous, Actemra Intravenous, Avsola, Infliximab (J1745), Cosentyx Intravenous, Steqeyma, Wezlana

1.	Is the request for continuation of currently approved therapy?	Yes – Go to renewal criteria	No – Go to #2
2.	Is the request for combined treatment with multiple targeted immune modulators (E.g., Hadlima plus Otezla)	Yes – Criteria not met, experimental	No – Go to #3
3.	Is the request to treat a diagnosis according to one of the Food and Drug Administration (FDA)-approved or compendia supported indications?	Yes – Go to appropriate section below	No – Criteria not met
Rheumatoid Arthritis (RA) Preferred Pharmacy Drugs – Hadlima, Hyrimoz (Cordavis), A Preferred Medical Drugs –Inflectra, Renflexis, Simponi Aria, Non-Preferred Medical Drugs – Remicade, Actemra IV, Oren		Tofidence IV, Tyenne	e IV
1.	Is there documented current disease activity with one of the following (or equivalent objective scale)? Disease Activity Score derivative for 28 joints (DAS-28) greater than 3.2 Clinical Disease Activity Index (CDAI) greater than 10 Weighted Routine Assessment of Patient Index Data 3 (RAPID3) of at least 2.3	Yes – Document and go to #2	No – Criteria not met
2.	Is there documented treatment failure with minimum 12- week trial with methotrexate? If contraindicated or unable to tolerate, is there evidence of 12-week treatment failure with sulfasalazine, hydroxychloroquine, or leflunomide?	Yes – Go to #3	No – Criteria not met
3.	Is the request for a non-preferred medical drug?	Yes – Go to #4	No – Go to #5
4.	Is there documented treatment failure with each of the following: One of the preferred pharmacy drugs: Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Enbrel, Xeljanz, Rinvoq AND One of the preferred medical drugs: Inflectra,	Yes – Document and Go to #5	No – Criteria not met



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	Renflexis, Simponi Aria, Tofidence IV, Tyenne IV				
5.	Is the drug prescribed by, or in consultation with, a rheumatology specialist?	Yes – Go to #6	No – Criteria not met		
6.	Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations?	Yes – Approve up to 6 months	No – Criteria not met		
Pre Ste	Plaque Psoriasis (PP) Preferred Pharmacy Drugs –Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Enbrel, Cosentyx, Otezla, Stelara, Skyrizi, Tremfya Preferred Medical Drugs – Inflectra, Renflexis, Stelara Non-Preferred Medical Drugs – Remicade, Infliximab (J1745), Avsola, Steqeyma, Wezlana				
	Is there documentation that the skin disease meets one of the following: At least 10% body surface area involvement despite current treatment Hand, foot, or mucous membrane involvement	Yes – Document and go to #2	No – Criteria not met		
2.	Is there documented treatment failure with 12 weeks of at least two systemic therapies: methotrexate, cyclosporine, Acitretin, phototherapy (UVB, PUVA)?	Yes – Document and go to #3	No – Criteria not met		
3.	Is the request for a non-preferred medical drug?	Yes – Go to #4	No – Go to #5		
4.	Is there documented treatment failure with each of the following: One of the preferred pharmacy drugs: Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Enbrel, Cosentyx, Otezla, Stelara, Skyrizi, Tremfya AND One of the preferred medical drugs: Inflectra, Renflexis	Yes – Go to #5	No – Criteria not met		
5.	Is the drug prescribed by, or in consultation with, a dermatology specialist?	Yes – Go to #6	No – Criteria not met		
6.	Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations?	Yes – Approve up to 6 months	No – Criteria not met		



Psoriatic Arthritis (PsA)

Preferred Pharmacy Drugs – Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Enbrel, Otezla, Cosentyx, Xeljanz, Stelara, Tremfya, Rinvoq, Skyrizi Preferred Medical Drugs – Inflectra, Renflexis, Stelara, Simponi Aria Non-Preferred Medical Drugs – Remicade, Orencia IV, Infliximab (J1745), Avsola, Cosentyx Intravenous, Steqeyma, Wezlana

 Is there documentation of Classification for Psoriatic Arthritis (CASPAR) criteria score 3 or greater based on chart notes: Skin psoriasis: present – two points, OR previously present by history – one point, OR a family history of psoriasis, if the patient is not affected – one point Nail lesions (onycholysis, pitting): one point Dactylitis (present or past, documented by a rheumatologist): one point Negative rheumatoid factor (RF): one point Juxta-articular bone formation on radiographs (distinct from osteophytes): one point 	Yes – Document and go to #2	No – Criteria not met
2. Is there documented failure with at least 12 weeks of treatment with methotrexate, or if unable to tolerate methotrexate or contraindications apply, another disease modifying antirheumatic drug (sulfasalazine, cyclosporine, leflunomide)?	Yes – Document and go to #3	No – Criteria not met
3. Is the request for a non-preferred medical drug?	Yes – Go to #4	No – Go to #5
4. Is there documented treatment failure with each of the following: One of the preferred pharmacy drugs: Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Enbrel, Otezla, Cosentyx, Xeljanz, Stelara, Tremfya, Rinvoq, Skyrizi AND One of the preferred medical drugs: Inflectra, Renflexis, Simponi Aria	Yes – Go to #5	No – Criteria not met
Is the drug prescribed by, or in consultation with, a rheumatology specialist?	Yes – Go to #6	No – Criteria not met
Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource	Yes – Approve up to 6 months	No – Criteria not met



quantity limitations?			
Ankylosing Spondylitis (AS) & Non-radiographic Axial Spondyloarthritis (nr-axSpA) & Psoriatic Arthritis with Axial Involvement Preferred Pharmacy Drugs – Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Enbrel, Cosentyx, Xeljanz, Rinvoq Preferred Medical Drugs –Inflectra, Renflexis, Simponi Aria Non-preferred Medical Drugs –Remicade, Infliximab (J1745), Avsola, Cosentyx Intravenous			
1. Is there a diagnosis of axial spondyloarthritis (SpA) confirmed by sacroiliitis on imaging AND at least 1 Spondyloarthritis (SpA) feature: Inflammatory back pain (4 of 5 features met): Onset of back discomfort before the age of 40 years Insidious onset Improvement with exercise No improvement with rest Pain at night (with improvement upon arising) Arthritis Enthesitis Uveitis Dactylitis (inflammation of entire digit) Psoriasis Crohn's disease/ulcerative colitis Good response to NSAIDs Family history of SpA Elevated CRP OR HLA-B27 genetic test positive AND at least 2 SpA features	Yes – Go to #2	No – Criteria not met	
Is there documentation of active disease defined by Bath ankylosing spondylitis disease activity index (BASDAI) at least 4 or equivalent objective scale?	Yes – Document and go to #3	No – Criteria not met	
3. Is there documented failure with two daily prescription strength nonsteroidal anti-inflammatory drugs (ibuprofen, naproxen, diclofenac, meloxicam, etc.) with minimum 1 month trial each? OR For isolated sacroiliitis, enthesitis, peripheral arthritis: documented treatment failure with locally administered parenteral glucocorticoid?	Yes – Document and go to #4	No – Criteria not met	



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4.	Is the request for a non-preferred medical drug?	Yes – Go to #5	No – Go to #6
5.	Is there documented treatment failure with each of the following: One of the preferred pharmacy drugs: Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Enbrel, Cosentyx, Xeljanz, Rinvoq AND One of the preferred medical drugs: Inflectra, Renflexis, Simponi Aria	Yes – Go to #6	No – Criteria not met
6.	Is the drug prescribed by, or in consultation with, a rheumatology specialist?	Yes – Go to #7	No – Criteria not met
7.	Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations?	Yes – Approve up to 6 months	No – Criteria not met
Pre Pre	Crohn's Disease (CD) Preferred Pharmacy Drugs – Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Stelara, Skyrizi, Rinvoq Preferred Medical Drugs – Inflectra, Renflexis, Skyrizi Intravenous, Stelara Non-preferred Medical Drugs –Remicade, Entyvio, Infliximab (J1745), Avsola, Steqeyma, Wezlana		
1.	Is there a diagnosis supported by endoscopy/colonoscopy/sigmoidoscopy or biopsy with moderate to severely active disease despite current treatment?	Yes – Go to #2	No – Criteria not met
2.	Is there documented failure with at least two oral treatments for a minimum of 12 weeks: corticosteroids, azathioprine, 6-mercaptopurine, methotrexate, sulfasalazine, balsalazide? OR Documentation of previous surgical intervention for Crohn's disease?	Yes – Document and go to #4	No –Go to #3
3.	Is there documented severe, high-risk disease on colonoscopy defined by one of the following: Fistulizing disease Stricture Presence of abscess/phlegmon Deep ulcerations Large burden of disease including ileal, ileocolonic, or proximal gastrointestinal involvement	Yes – Document and go to #4	No – Criteria not met
4.	Is the request for a non-preferred medical drug?	Yes – Go to #5	No – Go to #6



5.	Is there documented treatment failure with each of the following: One of the preferred pharmacy drugs: Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Stelara, Skyrizi, Rinvoq AND One of the preferred medical drugs: Inflectra, Renflexis	Yes – Go to #6	No – Criteria not met
6.	Is the drug prescribed by, or in consultation with, a gastroenterology specialist?	Yes – Go to #7	No – Criteria not met
7.	Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations?	Yes – Approve up to 6 months	No – Criteria not met
Pre Sky Pre No	cerative Colitis (UC) eferred Pharmacy Drugs – Hadlima, Hyrimoz (Cordavis), A yrizi, Tremfya eferred Medical Drugs –Inflectra, Renflexis, Stelara, Skyriz n-Preferred Medical Drugs –Remicade, Entyvio, Omvoh, In	zi Intravenous, Tremf	ya Intravenous
	zlana		
1.	Is there a diagnosis supported by endoscopy/colonoscopy/sigmoidoscopy or biopsy with moderate to severely active disease despite current treatment?	Yes – Go to #2	No – Criteria not met
2.	Is there a diagnosis supported by endoscopy/colonoscopy/sigmoidoscopy or biopsy with moderate to severely active disease despite current	Yes – Go to #2 Yes – Document and got to #4	No – Criteria not met No – Go to #3
	Is there a diagnosis supported by endoscopy/colonoscopy/sigmoidoscopy or biopsy with moderate to severely active disease despite current treatment? Is there severely active disease despite current treatment, defined by one of the following: O Greater than or equal to 6 bloody, loose stools per day with severe cramps and evidence of systemic toxicity (fever, tachycardia, anemia, and/or elevated CRP/ESR)	Yes – Document	



following: o One of the Hyrimoz (Skyrizi, R AND	ted treatment failure with each of the e preferred pharmacy drugs: Hadlima, Cordavis), Adalimumab-adaz, Stelara, invoq, Xeljanz, Tremfya	Yes – Go to #6	No – Criteria not met
6. Is the drug prescr gastroenterology	ibed by, or in consultation with, a specialist?	Yes – Go to #7	No – Criteria not met
•	lose within the Food and Drug DA)-approved label and PacificSource s?	Yes – Approve up to 6 months	No – Criteria not met
Preferred Medical Di	Arthritis (JIA) Drugs – Hadlima, Hyrimoz (Cordavis), A rug – Simponi Aria, Tofidence IV, Tyenno ral Drugs – Orencia IV, Actemra IV		brel, Xeljanz, Rinvoq
	ted current level of disease activity with ssessment (MD global score) or active	Yes – Document and go to #2	No – Criteria not met
 Glucocorticoio AND 	ted failure with each of the following: d joint injections or oral corticosteroids week trial with methotrexate or	Yes – Go to #3	No – Criteria not met
3. Is the request for	a non-preferred medical drug?	Yes – Go to #4	No – Go to #5
following: One of the Hyrimoz (Xeljanz, F AND One of the	ted treatment failure with each of the e preferred pharmacy drugs: Hadlima, Cordavis), Adalimumab-adaz, Enbrel, Rinvoq e preferred medical drugs: Simponi Aria, e IV, Tyenne IV	Yes – Go to #5	No – Criteria not met
5. Is the drug prescr rheumatologist? s	ibed by, or in consultation with, a pecialist?	Yes – Go to #6	No – Criteria not met



Administration (FDA)-approved label and PacificSource quantity limitations? Uveitis – Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz 1. Is there a confirmed diagnosis of noninfectious uveitis? Yes – Go to #2 No – Criteria not me 2. Is the diagnosis being treated intermediate or panuveitis? Yes – Go to #5 No – Go to #3 3. Is the diagnosis being treated posterior uveitis? Yes – Go to #6 No – Go to #4 4. Is the diagnosis being treated anterior uveitis? Yes – Criteria not met 5. Is there documented treatment failure with the following: o One immunosuppressive agent: methotrexate, azathioprine, mycophenolate AND One systemic calcineurin inhibitor (cyclosporine, tacrolimus) 6. Is there documented treatment failure with Yutiq AND Retisert? 7. Is the drug prescribed by, or in consultation with, an ophthalmology specialist? 8. Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations? Hidradenitis Suppurativa (HS) Preferred Pharmacy Drugs – Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Cosentyx Preferred Medical Drugs –Remicade, Infliximab (J1745), Avsola						
1. Is there a confirmed diagnosis of noninfectious uveitis? 2. Is the diagnosis being treated intermediate or panuveitis? 3. Is the diagnosis being treated posterior uveitis? 4. Is the diagnosis being treated anterior uveitis? 5. Is there documented treatment failure with the following: One immunosuppressive agent: methotrexate, azathioprine, mycophenolate AND One systemic calcineurin inhibitor (cyclosporine, tacrolimus) 6. Is there documented treatment failure with Yutiq AND Retisert? 7. Is the drug prescribed by, or in consultation with, an ophthalmology specialist? 8. Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations? Hidradenitis Suppurativa (HS) Preferred Pharmacy Drugs — Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Cosentyx Preferred Medical Drugs —Remicade, Infliximab (J1745), Avsola 1. Is there a diagnosis of moderate to severe Hidradenitis Yes — Document No — Criteria not me	6.	Administration (FDA)-approved label and PacificSource		No – Criteria not met		
2. Is the diagnosis being treated intermediate or panuveitis? 3. Is the diagnosis being treated posterior uveitis? 4. Is the diagnosis being treated anterior uveitis? 5. Is there documented treatment failure with the following: One immunosuppressive agent: methotrexate, azathioprine, mycophenolate AND One systemic calcineurin inhibitor (cyclosporine, tacrolimus) 6. Is there documented treatment failure with Yutiq AND Retisert? 7. Is the drug prescribed by, or in consultation with, an ophthalmology specialist? 8. Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations? Hidradenitis Suppurativa (HS) Preferred Pharmacy Drugs – Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Cosentyx Preferred Medical Drugs –Remicade, Infliximab (J1745), Avsola 1. Is there a diagnosis of moderate to severe Hidradenitis Yes – Document No – Criteria not me	Uv	eitis – Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz				
3. Is the diagnosis being treated posterior uveitis? 4. Is the diagnosis being treated anterior uveitis? 5. Is there documented treatment failure with the following: 6. One immunosuppressive agent: methotrexate, azathioprine, mycophenolate AND 7. One systemic calcineurin inhibitor (cyclosporine, tacrolimus) 6. Is there documented treatment failure with Yutiq AND Retisert? 7. Is the drug prescribed by, or in consultation with, an ophthalmology specialist? 8. Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations? Find a provided in the proof of the preferred Pharmacy Drugs — Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Cosentyx Preferred Medical Drugs —Remicade, Infliximab (J1745), Avsola 1. Is there a diagnosis of moderate to severe Hidradenitis Yes — Go to #7 No — Criteria not me to 6 months No — Criteria not me to 6 months No — Criteria not me to 6 months	1.	Is there a confirmed diagnosis of noninfectious uveitis?	Yes – Go to #2 No – Criteria not met			
4. Is the diagnosis being treated anterior uveitis? 5. Is there documented treatment failure with the following: One immunosuppressive agent: methotrexate, azathioprine, mycophenolate AND One systemic calcineurin inhibitor (cyclosporine, tacrolimus) 6. Is there documented treatment failure with Yutiq AND Retisert? 7. Is the drug prescribed by, or in consultation with, an ophthalmology specialist? 8. Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations? Hidradenitis Suppurativa (HS) Preferred Pharmacy Drugs – Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Cosentyx Preferred Medical Drugs –Inflectra, Renflexis Non-Preferred Medical Drugs –Remicade, Infliximab (J1745), Avsola	2.	Is the diagnosis being treated intermediate or panuveitis?	Yes – Go to #5	No – Go to #3		
met	3.	Is the diagnosis being treated posterior uveitis?	Yes – Go to #6	No – Go to #4		
 One immunosuppressive agent: methotrexate, azathioprine, mycophenolate AND One systemic calcineurin inhibitor (cyclosporine, tacrolimus) Is there documented treatment failure with Yutiq AND Retisert? Is the drug prescribed by, or in consultation with, an ophthalmology specialist? Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations? Hidradenitis Suppurativa (HS) Preferred Pharmacy Drugs – Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Cosentyx Preferred Medical Drugs –Inflectra, Renflexis Non-Preferred Medical Drugs –Remicade, Infliximab (J1745), Avsola Is there a diagnosis of moderate to severe Hidradenitis Yes – Go to #8 No – Criteria not method No – Criteria not method No – Criteria not method 	4.	Is the diagnosis being treated anterior uveitis?				
7. Is the drug prescribed by, or in consultation with, an ophthalmology specialist? 8. Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations? Hidradenitis Suppurativa (HS) Preferred Pharmacy Drugs – Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Cosentyx Preferred Medical Drugs –Inflectra, Renflexis Non-Preferred Medical Drugs –Remicade, Infliximab (J1745), Avsola 1. Is there a diagnosis of moderate to severe Hidradenitis Yes – Go to #8 No – Criteria not me	5.	 One immunosuppressive agent: methotrexate, azathioprine, mycophenolate AND One systemic calcineurin inhibitor (cyclosporine, 	Yes – Go to #7	No – Criteria not met		
8. Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations? Hidradenitis Suppurativa (HS) Preferred Pharmacy Drugs – Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Cosentyx Preferred Medical Drugs –Inflectra, Renflexis Non-Preferred Medical Drugs –Remicade, Infliximab (J1745), Avsola 1. Is there a diagnosis of moderate to severe Hidradenitis Yes – Document No – Criteria not me	6.		Yes – Go to #7	No – Criteria not met		
Administration (FDA)-approved label and PacificSource quantity limitations? Hidradenitis Suppurativa (HS) Preferred Pharmacy Drugs – Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Cosentyx Preferred Medical Drugs –Inflectra, Renflexis Non-Preferred Medical Drugs –Remicade, Infliximab (J1745), Avsola 1. Is there a diagnosis of moderate to severe Hidradenitis Yes – Document No – Criteria not me	7.		Yes – Go to #8	No – Criteria not met		
Preferred Pharmacy Drugs – Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Cosentyx Preferred Medical Drugs –Inflectra, Renflexis Non-Preferred Medical Drugs –Remicade, Infliximab (J1745), Avsola 1. Is there a diagnosis of moderate to severe Hidradenitis Yes – Document No – Criteria not me	8.	Administration (FDA)-approved label and PacificSource		No – Criteria not met		
	Pre Pre	eferred Pharmacy Drugs – Hadlima, Hyrimoz (Cordavis), A eferred Medical Drugs –Inflectra, Renflexis		sentyx		
	1.			No – Criteria not met		



	Documentation of baseline count of abscess and inflammatory nodules?		
2.	Is there documented treatment failure with each of the following: Output Minimum 90-day trial with oral antibiotics: tetracycline/doxycycline/minocycline OR clindamycin with rifampin Minimum 8-week oral retinoid trial: isotretinoin OR acitretin	Yes – Document and go to #3	No – Criteria not met
3.	Is the request for a non-preferred medical drug?	Yes – Go to #4	No- Go to #5
4.	Is there documented treatment failure with each of the following: One of the preferred pharmacy drugs: Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz, Cosentyx AND One of the preferred medical drugs: Inflectra, Renflexis	Yes – Go to #5	No – Criteria not met
5.	Is the drug prescribed by, or in consultation with, a dermatology specialist?	Yes – Go to #6	No – Criteria not met
6.	Is the age of the member and requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations?	Yes – Approve up to 6 months	No – Criteria not met
Gia	ant Cell Arteritis (GCA) & Cytokine Release Syndrome (CR	S) – Actemra, Tofide	nce IV, Tyenne IV
1.	Is there a confirmed diagnosis of Cytokine Release Syndrome (CRS)?	Yes – Go to #4	No – Go to #2
2.	Is there a confirmed diagnosis of Giant Cell Arteritis (GCA) based on temporal artery biopsy or color doppler ultrasound OR	Yes – Go to #3	No – Criteria not met



	Large vessel GCA diagnosis by advanced imaging of the vascular tree with computed tomography (CT), magnetic resonance imaging (MRI), magnetic resonance angiography (MRA), positron emission tomography (PET) or PET with CT?		
3.	Is there documentation of disease refractory to treatment with glucocorticoids?	Yes – Go to #4	No – Criteria not met
4.	Is the drug prescribed by, or in consultation with, a rheumatology specialist?	Yes – Go to #5	No – Criteria not met
5.	Is the age of the member and requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations?	Yes – Approve up to 6 months (Maximum 4 doses for CRS)	No – Criteria not met
Ora	al Ulcers Associated with Behcet's Disease – Otezla		
1.	Is there a diagnosis of Behcet's with documentation of recurrent oral aphthae at least 3 times in a year AND two of the following: Output Recurrent genital aphthae Output Eye lesions Output Skin lesions Output Positive pathergy test defined by a papule 2 mm or greater	Yes – Go to #2	No – Criteria not met
2.	Is there documented treatment failure to a minimum 12- week trial to one of the following: colchicine, prednisone, azathioprine	Yes – Go to #3	No – Criteria not met
3.	Is the drug prescribed by, or in consultation with, a specialist with experience in treating Behcet's?	Yes – Go to #4	No – Criteria not met
4.	Is the age of the member and requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations?	Yes – Approve up to 6 months	No – Criteria not met
Ac	ute Graft Versus Host Disease (GVHD) Prophylaxis – Orer	ncia Intravenous	



1.	Is there documentation of a planned hematopoietic stem cell transplant (HSCT) including procedure date, patient weight, and planned dose?	Yes – Document and go to #2	No – Criteria not met
2.	Is there documentation that the drug will be used in combination with a systemic calcineurin inhibitor (tacrolimus, cyclosporine) AND methotrexate?	Yes – Document and go to #3	No – Criteria not met
3.	Is there documentation of a prior allogeneic hematopoietic stem cell transplant (HSCT), human immunodeficiency virus (HIV) infection, or any uncontrolled active infection (viral, bacterial, fungal, or protozoal)?	Yes – Criteria not met	No – Go to #4
4.	Is the drug prescribed by, or in consultation with, a hematologist or oncologist?	Yes – Approve up to 1 month (4 days of treatment maximum) with no reauthorization, unless otherwise specified	No – Criteria not met
Ato	opic Dermatitis (AD) - Rinvoq		
1.	Is there documentation of severe inflammatory skin disease defined as functional impairment (inability to use hands or feet for activities of daily living, or significant facial involvement preventing normal social interaction)?	Yes – Document and go to #2	No – Criteria not met
	Is there documentation of severe inflammatory skin disease defined as functional impairment (inability to use hands or feet for activities of daily living, or significant facial		No – Criteria not met No – Criteria not met
1.	Is there documentation of severe inflammatory skin disease defined as functional impairment (inability to use hands or feet for activities of daily living, or significant facial involvement preventing normal social interaction)? Is there a documented body surface area (BSA) affected of at least 10% OR hand, foot or mucous membrane	and go to #2 Yes – Document	



5.	Is the drug prescribed by, or in consultation with, a specialist in the treatment of atopic dermatitis (such as a dermatologist)?	Yes – Approve up to 6 months	No – Criteria not met
	thesitis-Related Arthritis (ERA) Preferred Drugs - Cosenty venile Psoriatic Arthritis (JPsA) Preferred Drugs – Cosenty		
1.	Is there diagnosis of ERA confirmed by presence of the following: • Arthritis persisting at least 6 weeks AND enthesitis present OR • Arthritis or enthesitis with two of the following features: • Sacroiliac tenderness or inflammatory lumbosacral pain • Positive HLA-B27 • Onset of arthritis in males greater than 6 years of age • Acute symptomatic anterior uveitis • First-degree relative with ERA, sacroiliitis associated with inflammatory bowel disease, reactive arthritis, or acute anterior uveitis	Yes – Document and go to #2	No – Go to #2
2.	Is there diagnosis of JPsA confirmed by presence of: • Arthritis and psoriasis OR • Arthritis and at least 2 of the following: • Dactylitis • Nail pitting or onycholysis • Psoriasis in a first-degree relative	Yes – Document and go to #3	No – Criteria not met
3.	Is there documented treatment failure with a nonsteroidal anti-inflammatory drug (ibuprofen, naproxen, celecoxib, meloxicam, etc.) with a minimum trial of 1 month?	Yes – Document and go to #4	No – Criteria not met
4.	Is there documented treatment failure with at least one of the following disease-modifying antirheumatic drugs (DMARDs) with a minimum trial of 12 weeks: methotrexate, sulfasalazine, leflunomide.	Yes – Document and go to #5	No – Criteria not met



5.	Is the drug prescribed by, or in consultation with, a rheumatologist?	Yes – Document and go to #6	No – Criteria not met
6.	Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations?	Yes – Approve up to 6 months	No – Criteria not met
Pre	neralized Pustular Psoriasis (GPP) Flare ferred Medical Drugs – Inflectra, Renflexis n-Preferred Medical Drugs – Remicade, Avsola, Infliximab	(J1745)	
1.	Is there documentation of a diagnosis of generalized pustular psoriasis (GPP) confirmed by the following:a. The presence of widespread sterile pustules arising on erythematous skinb. Pustulation is not restricted to psoriatic plaques	Yes – Document and go to #2	No – Criteria not met
2.	 Are there signs and symptoms of an acute GPP flare of moderate-to-severe intensity as follows: a. A Generalized Pustular Psoriasis Physician Global Assessment (GPPGA) total score of greater than or equal to 3 b. A GPPGA pustulation category subscore of greater than or equal to 2 c. Greater than or equal to 5% body surface are (BSA) covered with erythema and the presence of pustules 	Yes – Document and go to #3	No – Criteria not met
3.	Is there documented 1-week treatment failure with cyclosporine?	Yes – Document and go to #4	No – Criteria not met
4.	Is the request for Remicade, Avsola, or Infliximab (J1745)?	Yes – Go to #5	No – Go to #6
5.	Is there documented failure with one of the preferred medical drugs (Inflectra, Renflexis)?	Yes – Go to #6	No – Criteria not met
6.	Is the drug prescribed by, or in consultation with, a dermatology specialist?	Yes – Go to #7	No – Criteria not met



7.	Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations?	Yes – Approve up to 6 months	No – Criteria not met
Re	newal Criteria		
1.	Is there documentation of treatment success and a clinically significant response to therapy as assessed by the prescribing provider, with clinical documentation to support?	Yes – Go to #2	No – Criteria not met
2.	Is the request for combined treatment with multiple targeted immune modulators? (E.g., Hadlima plus Otezla)	Yes – Criteria not met	No – Go to #3
3.	Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations?	Yes – Approve up to 12 months	No – Criteria not met

Quantity Limitations

• Hadlima, Hyrimoz (Cordavis), Adalimumab-adaz

- Induction
 - PP/Uveitis: 80 mg as a single dose, followed by 40 mg every other week beginning 1 week after initial dose (160 mg total in first 28 days)
 - CD/UC/HS: 160 mg on day 1, followed by 80 mg on day 15, then maintenance dosing beginning on day 29
- o Maintenance
 - RA/PP/PsA/CD/UC/AS/nr-axSpA/Uveitis/JIA: 40 mg every other week
 - HS: 40 mg every week OR 80 mg every other week
- Dose escalation (40 mg every week OR 80 mg every other week)
 - RA/PP/CD/UC: Approval will require documentation of lost or inadequate response after a minimum of 16 weeks with standard maintenance dosing

• Enbrel

- Induction
 - PP: 50 mg twice weekly for 3 months (8 injections per 28 days for 3 months)
- Maintenance (All indications):
 - PP/JPsA: 50 mg once weekly (4 injections per 28 days)
 - RA/PP/PsA/AS/nr-axSpA/JIA:
 - 25 mg twice weekly (8 injections per 28 days)
 - 50 mg once weekly (4 injections per 28 days)

Cosentyx



Induction

- Adult Plaque Psoriasis: 4 two-packs (300 mg) in first 28 days
- Pediatric Plaque Psoriasis/Pediatric Psoriatic Arthritis/Pediatric Enthesitis-Related Arthritis:
 - Less than 50 kg: four 75 mg doses in the first 28 days
 - Greater than or equal to 50 kg: four 150 mg doses in the first 28 days
 - Hidradenitis Suppurativa: 4 two-packs (300 mg) in first 28 days

Maintenance

- Adult Plaque Psoriasis: 1 two-pack (300 mg) per 28 days
- Pediatric Plaque Psoriasis/Pediatric Psoriatic Arthritis/Pediatric Enthesitis-Related Arthritis:
 - Less than 50 kg: 75 mg per 28 days
 - Greater than or equal to 50 kg: 150 mg per 28 days
- Psoriatic arthritis without plaque psoriasis/AS/Nr-axSpA: 1 injection (150 mg) per 28 days
 - If a patient continues to have active disease, a dosage of 300 mg may be considered
- Hidradenitis Suppurativa: 1 two-pack (300 mg) per 28 days

Otezla

- Induction (All indications): Titration pack
- Maintenance (All indications): 60 tablets per 30 days

Stelara

- o Induction
 - Plaque Psoriasis: One 45 mg injection (0.5 mL) in first 28 days for those weighing 60 to 100 kg, one 90 mg injection (1 mL) in first 28 days for those weighing over 100 kg
 - For those under 60kg, the dose is 0.75 mg/kg
 - Psoriatic Arthritis: One 45 mg injection (0.5 mL) in the first 28 days
 - For coexistent moderate to severe PP and weight greater than 100kg: one 90 mg injection (1 mL) in first 28 days
 - Crohn's Disease and Ulcerative Colitis: A single intravenous infusion per below
 - 55 kg or less: 260 mg
 - 55 kg to 85 kg: 390 mg
 - More than 85 kg: 520 mg

Maintenance

- Plaque Psoriasis: One 45 mg injection (0.5 mL) per 84 days for those weighing 100 kg or less; one 90 mg injection (1 mL) per 84 days for those weighing over 100 kg
- Psoriatic Arthritis: 45 mg (0.5 mL) per 84 days
 - For coexistent moderate-to-severe plaque psoriasis weighing more than 100 kg:
 90 mg (1 ml) per 84 days
- Crohn's Disease and Ulcerative Colitis: 90 mg (1 mL) per 56 days starting 8 weeks after the initial IV dose

Tremfya

- o PP/PsA:
 - Induction: 100 mg (one injection) in first 28 days



- Maintenance: 100 mg (one injection) per 56 days
- o Ulcerative Colitis:
 - Induction: 200 mg intravenous at week 0, week 4, and week 8
 - Maintenance: 100 mg subcutaneously every 8 weeks, beginning week 16
 - For consideration of every 4 week dosing, must meet all of the following:
 - Documented clinical failure to Tremfya 100 mg every 8 week dosing for at least 3 months

• Skyrizi

- o PP/PsA:
 - Induction: 150 mg in the first 28 daysMaintenance: 150 mg per 84 days
- o Crohn's Disease:
 - Induction: 600 mg intravenous at week 0, week 4, and week 8
 - Maintenance: 360 mg subcutaneously every 8 weeks, beginning week 12
- Ulcerative Colitis:
 - Induction: 1200 mg intravenous at week 0, week 4, and week 8
 - Maintenance: 360 mg subcutaneously every 8 weeks, beginning week 12

Rinvoq

- RA/PsA/AS/nr-axSpA: 15 mg once daily (30 tablets per 30 days)
- AD: 15 mg once daily, may increase to 30 mg once daily if inadequate response (30 tablets per 30 days)
- UC: 45 mg once daily for 8 weeks then 15 mg once daily. May increase to 30 mg once daily if inadequate response (30 tablets per 30 days).
 - **45mg limited to 56 tablets (first 8 weeks of treatment)
- CD: 45 mg once daily for 12 weeks, then 15 mg once daily. May increase to 30 mg once daily for patients with refractory, severe or extensive disease.
 - **45mg limited to 84 tablets (first 12 weeks of treatment)
- Polyarticular JIA/Pediatric Psoriatic Arthritis: 10 kg to <20 kg: 3 mg (3 mL solution) twice daily; 20 kg to <30 kg: 4 mg (4 mL solution) twice daily; 30 kg and greater: 6 mg (6 mL solution) twice daily or 15 mg tablet once daily

Xeljanz

- RA/PsA/AS: 60 tablets per 30 days (5 mg IR) OR 30 tablets per 30 days (11 mg XR)
- UC: 60 tablets per 30 days (5 mg or 10 mg IR tablets) OR 30 tablets per 30 days (11 mg or 22 mg XR)
- JIA: 10 kg to less than 20 kg: 3.2 mg (3.2 mL oral solution) twice daily; 20 kg to less than 40 kg: 4 mg (4 mL oral solution) twice daily; 40 kg or greater: 5 mg (one 5 mg tablet or 5 mL oral solution) twice daily
 - Oral solution available as 240 mL bottle
- Infliximab (Remicade, Inflectra, Renflexis, Avsola, Infliximab (J1745))*
 - Availability: 100 mg single-dose vials
 - Crohn's/UC/HS: 5 mg/kg at 0, 2 and 6 weeks followed by 5 mg/kg every 8 weeks thereafter. For those who respond and lose response, consideration may be given to treatment with 10 mg/kg



- Psoriatic Arthritis/Plaque Psoriasis/Generalized Pustular Psoriasis: 5 mg/kg at 0, 2 and 6 weeks followed by 5 mg/kg every 8 weeks thereafter
- RA: 3 mg/kg at 0, 2 and 6 weeks followed by 3 mg/kg every 8 weeks thereafter. For those with an
 incomplete response, consideration may be given for dosing up to 10 mg/kg or as often as every 4
 weeks
- AS: 5 mg/kg at 0, 2 and 6 weeks followed by 5 mg/kg every 6 weeks thereafter

Simponi Aria Intravenous*

- o Availability: 50 mg single-dose vials
- o RA/PsA/AS: 2 mg/kg at weeks 0 and 4, then every 8 weeks thereafter
- Pediatric PsA and JIA: 80 mg/m2 at weeks 0 and 4, then every 8 weeks thereafter

Orencia Intravenous*

- o Availability: 250 mg single-use vials
- RA/PsA: <60 kg: 500 mg, 60-100 kg: 750 mg, >100 kg: 1,000 mg at 0, 2, and 4 weeks followed by every 4 weeks thereafter
- JIA: 6 years and older and <75 kg: 10 mg/kg; 75-100 kg: 750 mg; >100 kg: 1,000 mg (maximum dose) at 0, 2 and 4 weeks followed by every 4 weeks thereafter
- Acute GVHD Prophylaxis:
 - 2 to less than 6 years: 15 mg/kg on day -1 (day before transplantation) followed by 12 mg/kg on days 5, 14, and 28 post-transplant
 - 6 years and older: 10 mg/kg on day -1 (day before transplantation) followed by 10 mg/kg on days 5, 14, and 28 post-transplant (maximum: 1,000 mg/dose)

Entyvio*

- Availability: 300 mg single-use vials
- Crohn's/UC: 300 mg at 0, 2 and 6 weeks followed by every 8 weeks thereafter
- For Consideration of every 4 week dosing must meet all of the following:
 - Documented clinical failure to Entyvio at standard dosing for at least 6 months
 - Clinical failure defined as failure to achieve a clinical response (greater than or equal to 70 point improvement in CDAI score for Crohn's)
 - Documented failure to minimum of 12 weeks on two alternative Tumor necrosis factor alpha (TNF) inhibitors

Actemra Intravenous, Tofidence Intravenous, Tyenne Intravenous*

- Availability: 400 mg, 200 mg & 80 mg single-dose vials
- RA: 4 mg/kg once every 4 weeks; may be increased to 8 mg/kg once every 4 weeks based on clinical response (maximum dose: 800 mg)
- o GCA: 6mg/kg every 4 weeks
- CRS: For patients less than 30kg, recommended dose is 12mg/kg; patients 30kg or greater recommended dose is 8mg/kg up to maximum of 800mg (Maximum 4 doses)
- Polyarticular JIA: <30 kg: 10 mg/kg every 4 weeks; 30 kg or greater: 8 mg/kg every 4 weeks
- Systemic JIA: <30 kg: 12 mg/kg every 2 weeks; 30 kg or greater: 8 mg/kg every 2 weeks

^{*}Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced for all medical infusion drugs



Drug Name	Ankylosin g Spondyliti s	Crohn's Disease	Juvenile Idiopathic Arthritis	Plaque Psoriasis	Psoriati c Arthritis	Rheumatoi d Arthritis	Ulcerativ e Colitis	Other
Abatacept (Orencia SQ & Orencia IV)			≥2 yo		≥2 yo	≥18 yo		Acute GVHD prophylaxis: IV: ≥2 yo
Adalimumab (Hadlima, Hyrimoz (Cordavis), Adalimumabadaz)	≥18 yo	≥6 yo ≥18 yo (biosimilar s)	≥2 yo ≥4 yo (biosimilar s)	≥18 yo	≥18 yo	≥18 yo	≥5 yo	Uveitis (noninfectiou s) ≥2 yo HS ≥12 yo
Anakinra (Kineret)						≥18 yo		NOMID
Apremilast (Otezla)				≥6 yo	≥18 yo			Behçet's Disease
Baricitinib (Olumiant)						≥18 yo		
Brodalumab (Siliq)				≥18 yo				
Canakinumab (Ilaris) [See standalone policy]			≥2 yo					FCAS ≥4 yo MWS ≥4 yo TRAPS ≥2 yo HIDS ≥2 yo MKD ≥2 yo FMF ≥2 yo
Certolizumab (Cimzia)	≥18 yo	≥18 yo		≥18 yo	≥18 yo	≥18 yo		Nr-axSpA ≥18 yo
Etanercept (Enbrel)	≥18 yo		≥2 yo	≥4 yo (Enbrel) ≥18 yo (biosimilar s)	≥18 yo	≥18 yo		JPsA ≥2 yo
Golimumab (Simponi & Simponi Aria)	≥18 yo		≥2 yo (Simponi Aria)		≥18 yo (Simponi) ≥2 yo (Simponi Aria)	≥18 yo	≥18 yo (Simponi)	



Guselkumab (Tremfya)				≥18 yo	≥18 yo		≥18 yo	
Infliximab (J1745), Remicade, Inflectra, Renflexis, Avsola	≥18 yo	≥6 yo		≥18 yo	≥18 yo	≥18 yo	≥6 yo	GPP≥18 yo
lxekizumab (Taltz)	≥18 yo			≥6 yo	≥18 yo			Nr-axSpA ≥18 yo
Rituximab (Rituxan) [See standalone policy]						≥18 yo		CLL ≥18 yo NHL ≥18 yo; ≥6 yo (Rituxan) GPA ≥18 yo; ≥2 yo (Rituxan) Pemphigus Vulgaris ≥18 yo RRMS ≥18
Risankizuma b-rzaa <mark>(Skyrizi)</mark>		≥18 yo		≥18 yo	≥18 yo		≥18 yo	
Sarilumab (Kevzara)						≥18 yo		
Secukinumab (Cosentyx)	≥18 yo			≥6 yo	≥2 yo			Nr-axSpA ≥18 yo ERA ≥ 4 yo JPsA ≥ 2 yo HS ≥18 yo
Tildrakizuma b-asmn (Ilumya)				≥18 yo				
Tocilizumab (Actemra SQ & Actemra IV, Tofidence IV, Tyenne IV SQ)			≥2 yo			≥18 yo		CRS >2 yo GCA >18 yo



Tofacitinib (Xeljanz)	≥18 yo		≥2 yo		≥18 yo	≥18 yo	≥18 yo	
Upadacitinib (Rinvoq)	≥18 yo	≥18 yo			≥18 yo	≥18 yo	≥18 yo	AD ≥12 yo Nr-axSpA ≥18 yo
Ustekinumab (Stelara)		≥18 yo		≥6 yo	≥18 yo		≥18 yo	
Vedolizumab (Entyvio)		≥18 yo					≥18 yo	

Yellow: Preferred Pharmacy Drugs Green: Medical Infusion Drugs

Abbreviations: AD = Atopic Dermatitis; CLL = Chronic Lymphocytic Leukemia; CRS = Cytokine Release Syndrome; ERA= Enthesitis-Related Arthritis; FCAS = Familial Cold Autoinflammatory Syndrome; FMF = Familial Mediterranean Fever; GCA = Giant Cell Arteritis; GPA = Granulomatosis with Polyangiitis (Wegener's Granulomatosis); HIDS: Hyperimmunoglobulin D Syndrome; HS = Hidradenitis Suppurativa; JPsA= Juvenile Psoriatic Arthritis; MKD = Mevalonate Kinase Deficiency; MPA = Microscopic Polyangitis; MWS = Muckle-Wells Syndrome; NHL = Non-Hodgkin's Lymphoma; NOMID = Neonatal Onset Multi-Systemic Inflammatory Disease; Nr-axSpA = nonradiographic Axial Spondyloarthritis; Still's dx = Adult-onset Still's disease; TRAPS = Tumor Necrosis Factor Receptor Associated Periodic Syndrome; RRMS = Relapsing-Remitting Multiple Sclerosis; yo = years



POLICY NAME: **TARPEYO**

Affected Medications: TARPEYO (Budesonide Delayed Release Capsule 4 mg)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	 Reduce the loss of kidney function in adults with primary immunoglobulin A nephropathy (IgAN) who are at risk for disease progression
Required Medical Information:	 Diagnosis of primary immunoglobulin A nephropathy (IgAN) confirmed with biopsy Documentation of risk of rapid disease progression with a urine protein-to-creatinine ratio (UPCR) equal to or greater than 1.5 g/g (labs current within 30 days of request) OR Proteinuria defined as equal to or greater than 1 g/day (labs current within 30 days of request)
Appropriate Treatment Regimen & Other Criteria:	 Documentation of treatment failure with a minimum of 12 weeks of an angiotensin-converting enzyme (ACE) inhibitor or angiotensin receptor blocker (ARB) AND Documentation of treatment failure with a minimum of 12 weeks of glucocorticoid therapy such as oral prednisone or methylprednisolone (treatment failure defined as proteinuria equal to or greater than 1 g/day or an adverse effect to two glucocorticoid therapies that is not associated with the corticosteroid class) AND Documentation of treatment failure with a minimum of 12 weeks of Filspari (treatment failure defined as proteinuria equal to or greater than 1 g/day or an adverse effect to Filspari) No reauthorization – Recommended duration of therapy is 9 months followed by a 2-week dose taper prior to discontinuation
Exclusion Criteria:	Other glomerulopathies or nephrotic syndrome
Age Restriction:	18 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a nephrologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 10 months, unless otherwise specified



POLICY NAME: TASIMELTEON

Affected Medications: HETLIOZ LQ SUSPENSION, TASIMELTEON

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Treatment of Non-24-Hour Sleep-Wake Disorder (Non-24)
	 Treatment of nighttime sleep disturbances in Smith-Magenis Syndrome (SMS)
Required Medical	Non-24
Information:	Documentation of being totally blind with no light perception
	 Diagnosis of Non-24 hour sleep wake disorder meeting ALL of the following: Documented history of insomnia, excessive daytime sleepiness, or both, that alternates with asymptomatic periods
	 Symptoms have been present for at least three months Drift in rest-activity patterns demonstrated by at least 4 weeks of data from daily sleep logs and actigraphy
	 Documentation that other sleep disorders were treated or ruled out using a sleep study
	Smith-Magenis Syndrome (SMS)
	 Diagnosis of Smith-Magenis Syndrome (SMS) confirmed by both of the following: Genetic test showing mutation or deletion of the retinoic acid-induced 1 (RAI1) gene
	Documentation of significant nighttime sleep disturbances
Appropriate	<u>Non-24</u>
Treatment	Documentation of treatment failure with at least 12 weeks of melatonin
Regimen & Other	Smith-Magenis Syndrome (SMS)
Criteria:	Documented trial and failure with treatment regimen that includes both melatonin taken at bedtime AND acebutolol taken during daytime for at least 12 weeks
	Reauthorization requires documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	Sleep disorders other than Non-24 and SMS such as insomnia, shift work disorder, jet lag
	disorder, irregular sleep-wake rhythm disorder, delayed sleepwake phase disorder,
	advanced sleep-wake rhythm disorder
	Sleep disturbances caused by taking sedative or stimulant central nervous system-active
	drugs
	Sleep disturbances caused by other conditions
Age Restriction:	Non-24: 18 years of age and older
	SMS:
	Capsules: 16 years of age and older
	Suspension: 3 to 15 years of age



Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with a neurologist or sleep specialist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **TEDIZOLID**

Affected Medications: SIVEXTRO injection, SIVEXTRO tablets

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Acute bacterial skin and skin structure infections (ABSSSI) caused by susceptible isolates of the following Gram-positive microorganisms: Staphylococcus aureus (including methicillin-resistant [MRSA] and methicillin-susceptible [MSSA] isolates) Streptococcus pyogenes Streptococcus agalactiae Streptococcus anginosus Group (including Streptococcus anginosus, Streptococcus intermedius, and Streptococcus constellatus) Enterococcus faecalis
Required	Documentation of confirmed or suspected diagnosis
Medical	Documentation of treatment history and current treatment regimen
Information:	Documentation of culture and sensitivity data
	Documentation of culture and sensitivity data Documentation of planned treatment duration
	Bocumentation of planned treatment duration
Appropriate Treatment	Dosing: 200 mg once daily for 6 days
Regimen &	Requests for the intravenous formulation will require both of the following:
Other Criteria:	 Documentation of treatment failure, contraindication, or intolerable adverse event with intravenous linezolid AND
	 Documentation of treatment failure, contraindication, or intolerable adverse event with at least 2 of the following drugs/drug classes: Vancomycin
	 Avoidance of vancomycin due to nephrotoxicity will require documentation of multiple (at least 2 consecutive) increased serum creatinine concentrations (increase of 0.5 mg/dL [44 mcmol/L] or at least 50 percent increase from baseline, whichever is greater), without an alternative explanation Daptomycin
	Cephalosporin (cefazolin)
	Requests for the oral tablet formulation will require both of the following: • Documentation of treatment failure, contraindication, or intolerable adverse event with oral linezolid AND
	 Documentation of treatment failure, contraindication, or intolerable adverse event with at least 2 of the following drugs/drug classes: Trimethoprim-sulfamethoxazole
	 Trimetrioprim-sullametrioxazole Tetracycline (doxycycline, minocycline) Clindamycin
Exclusion Criteria:	
Age Restriction:	12 years of age and older



Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a infectious disease specialist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 1 month, unless otherwise specified



POLICY NAME: **TEDUGLUTIDE**

Affected Medications: GATTEX (teduglutide)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of Short Bowel Syndrome (SBS)
Required Medical Information:	 Documentation of confirmed SBS diagnosis Dependence on parenteral nutrition (PN) and/or intravenous (IV) fluids at least 12 consecutive months continuously Receiving three or more days per week of PN support such as fluids, electrolytes, and/or nutrients
Appropriate Treatment Regimen & Other Criteria:	 Documentation of inability to be weaned from PN despite use of the following conventional measures: Dietary manipulations, oral rehydration solutions Antidiarrheal/motility agents: loperamide or diphenoxylate Antisecretory agents: H2 receptor antagonists or proton pump inhibitors Developed significant complications or severe impairment in quality of life related to parenteral nutrition use (such as loss of vascular access sites, recurrent catheter-related bloodstream infections, and liver disease) Dose does not exceed 0.05 mg/kg daily Reauthorization requires documentation of clinically significant benefit defined by parenteral support reduction of 1 day or greater a week
Exclusion Criteria: Age Restriction:	 Weight of less than 10 kg Onset or worsening of gallbladder/biliary disease Onset or worsening of pancreatic disease Presence of any gastrointestinal malignancy Presence of intestinal or stomal obstruction 1 year of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a gastroenterologist or SBS specialist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 6 months, unless otherwise specified



POLICY NAME: **TENAPANOR**

Affected Medications: XPHOZAH (tenapanor)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of hyperphosphatemia associated with chronic kidney disease (CKD)
Required Medical Information:	 Diagnosis of hyperphosphatemia associated with CKD and currently on dialysis treatment
	 Documentation of progressively or persistently elevated serum phosphate that is greater than 5.5 mg/dL over the past 6 months despite adherence to phosphate binders and dietary restrictions Documentation that Xphozah (tenapanor) will be used as add-on therapy to phosphate binder therapy unless contraindicated or clinically significant adverse effects were experienced
Appropriate	Documented treatment failure with at least an 8-week trial, at maximally indicated doses,
Treatment	of at least two of the following:
Regimen & Other	o calcium acetate
Criteria:	o lanthanum carbonate
	o sevelamer
	 Velphoro (sucroferric oxyhydroxide)
	Auryxia (ferric citrate)
	Reauthorization requires documentation of treatment success defined as reduction in serum phosphorus from pretreatment level and maintenance of serum phosphorus level at 5.5 mg/dL or lower
Exclusion Criteria:	Known or suspected mechanical gastrointestinal obstruction
Age Restriction:	18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a nephrologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
_	Reauthorization: 12 months, unless otherwise specified



TENOFOVIR ALAFENAMIDE

Affected Medications: VEMLIDY (tenofovir alafenamide)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	 For the treatment of chronic hepatitis B virus (HBV) infection in adults and
	pediatric patients 6 years of age and older with compensated liver disease
Required Medical	Documentation confirming diagnosis of chronic hepatitis B infection
Information:	 Documentation of compensated liver disease (Child-Pugh A) within 12 weeks prior to anticipated start of therapy
Appropriate	Documentation of one or more of the following:
Treatment	o Inadequate virologic response or intolerable adverse event to tenofovir disoproxil
Regimen & Other	fumarate
Criteria:	 CrCl less than or equal to 80 mL/min within 12 weeks prior to anticipated start date OR high risk for acute renal injury (i.e., nephrotoxic medications)
	 Diagnosis of osteoporosis, osteopenia, or high risk for developing osteoporosis with supporting documentation (i.e., chronic use of steroids or other drugs that worsen bone density, poor nutrition, early menopause)
	Reauthorization: documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	Decompensated hepatic impairment (Child-Pugh B or C)
Age Restriction:	6 years of age and older
Prescriber	Prescribed by, or in consultation with, a hepatologist, gastroenterologist, or infectious
Restrictions:	disease specialist
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: **TEPLIZUMAB-MZWV**

Affected Medications: TZIELD (teplizumab-mzwv)

All Food and Drug Administration (FDA)-approved indication plan design Type 1 diabetes mellitus, to delay the onset of Stage and pediatric patients with Stage 2 type 1 diabetes Required Medical Information: Diagnosis of Stage 2 type 1 diabetes, confirmed by both of Positive for two or more of the following pancreatic the past 6 months: Glutamic acid decarboxylase 65 (GAD) aut Insulin autoantibody (IAA) Insulin autoantibody (IAA) Islet cell autoantibody (ICA) Dysglycemia on oral glucose tolerance testing (OG as shown by one of the following: Fasting blood glucose between 110 mg/dL All Food and Drug Administration (FDA)-approved indication plan design Documentation that the patient has a first-degree or second diabetes and one of the following:	e 3 type 1 diabetes in adults, the following:
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Documentation that the patient has a first-degree or second	man or oqual to 200 mg/ac
	-degree relative with type 1
diabetes and one of the following.	degree relative with type 1
 If first-degree relative (brother, sister, parent, offspr 	ng) natient must be between
8 and 45 years of age	ng), patient must be between
	le grandchild cousin) nationt
o If second-degree relative (niece, nephew, aunt, und must be between 8 and 20 years of age	e, grandemid, codsin), patient
, ,	CA) or beight and weight to
Documentation of the patient's current body surface area (E	SA) or neight and weight to
calculate BSA	
Treatment plan, including planned dose and frequency	
Appropriate Treatment Approved for one-time 14-day infusion only, based on the f	ollowing dosing schedule:
Regimen & Other Criteria: Treatment Day Dose	
Day 1 65 mcg/m ²	
Day 2 125 mcg/m ²	
Day 3 250 mcg/m ²	
Day 4 500 mcg/m ²	
Days 5- 14 1,030 mcg/m ²	
 Availability: 2 mg/2 mL (1 mg/mL) single-dose vials 	
Dose-rounding to the nearest vial size within 10% of the pre-	
Exclusion Criteria: • Prior treatment with Tzield	scribed dose will be enforced
Diagnosis of Stage 3 type 1 diabetes (clinical type 1 diabetes)	scribed dose will be enforced



	Diagnosis of Type 2 diabetes
	Current active serious infection or chronic infection
	Pregnant or lactating
Age Restriction:	8 to 45 years of age
	See Required Medical Information for age requirements based on first-degree or second- degree relative
Prescriber/Site of Care	Prescribed by, or in consultation with, an endocrinologist
Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 3 months, unless otherwise specified (one 14-day infusion only)



TEPROTUMUMAB-TRBW

Affected Medications: TEPEZZA (teprotumumab-trbw)

	plan design
	Thyroid Eye Disease (TED) regardless of TED activity or duration
Required Medical Information:	 Documentation that baseline disease is under control prior to starting therapy, as defined by one of the following: Patient is euthyroid (thyroid function tests are within normal limits) Patient has recent and mild hypo- or hyperthyroidism (thyroid function tests show free thyroxine (T4) and free triiodothyronine (T3) levels less than 50% above or below normal limits) and will undergo treatment to maintain euthyroid state TED has an appreciable impact on daily life, defined as: Proptosis greater than or equal to 3 mm increase from baseline (prior to diagnosis of TED) and/or proptosis greater than or equal to 3 mm above normal for race and gender OR Current moderate-to-severe active TED with a Clinical Activity Score (CAS)
	greater than or equal to 4 (on the 7-item scale) for the most severely affected eye and symptoms such as: lid retraction greater than or equal to 3 mm, moderate or severe soft tissue involvement, diplopia, and/or proptosis greater than or equal to 3 mm above normal for race and gender
Appropriate Treatment	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced
Regimen & Other Criteria:	 Evidence of stable, well-controlled disease if comorbid inflammatory bowel disease (IBD) or diabetes Documented failure to intravenous or oral steroid at appropriate dose over 12 weeks
	., ,
Exclusion Criteria:	Use of more than one course of Tepezza treatment
	Prior orbital irradiation, orbital decompression, or strabismus surgery
	Decreasing visual acuity, new defect in visual field, color vision defect from optic nerve
	involvement within the previous 6 months
	Corneal decompensation that is unresponsive to medical management
Age Restriction:	18 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an ophthalmologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 7 months, maximum approval (total of 8 doses) with no reauthorization, unless otherwise specified



TESTOSTERONE

Affected Medications: TESTOPEL (testosterone pellets), JATENZO (testosterone undecanoate capsules), TLANDO (testosterone undecanoate capsules)

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Required Medical Information:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Testosterone replacement therapy in adult males for conditions associated with a deficiency or absence of endogenous testosterone: primary hypogonadism or hypogonadotropic hypogonadism Gender Dysphoria All Indications If 65 years of age and older, must provide documentation of a yearly evaluation that includes All of the following:
miormation.	includes ALL of the following: The need for continued hormone replacement therapy Education on the risks of hormone replacement therapy (heart attack, stroke) Discussion about the limited efficacy and safety for hormone replacement therapy in patients experiencing an age-related decrease in testosterone levels
	 Hypogonadism in Adults Confirmed low testosterone level (total testosterone less than 300 ng/dl or morning free or bioavailable testosterone less than 5 ng/dL) or absence of endogenous testosterone
	 Gender Dysphoria Documented diagnosis of gender dysphoria If under 18 years of age, documentation of all of the following: Current Tanner stage 2 or greater OR baseline and current estradiol and testosterone levels to confirm onset of puberty Confirmed diagnosis of gender dysphoria that is persistent The patient has the capacity to make a fully informed decision and to give consent for treatment Any significant medical or mental health concerns are reasonably well controlled A comprehensive mental health evaluation has been completed by a licensed mental health professional (LMHP) and provided in accordance with the most current version of the World Professional Association for Transgender Health (WPATH) Standards of Care Note: For requests following pubertal suppression therapy, an updated or new comprehensive mental health evaluation must be provided prior to initiation of hormone supplementation
Appropriate Treatment Regimen & Other Criteria:	All Indications Requests for oral testosterone (e.g., Jatenzo, Tlando) require documented treatment failure with testosterone injections AND generic transdermal testosterone Requests for Testopel require all of the following:



	 Documented treatment plan, including dosage in milligrams or number of pellets to be administered and frequency Maximum dosage: 450 mg per treatment Maximum of 4 treatments in 12 months
	 Reauthorization: Hypogonadism in Adults: Documentation of a recent testosterone level within normal limits Gender Dysphoria: Documentation of treatment success
Exclusion Criteria:	Treatment of sexual dysfunction Treatment of symptoms of menopause
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Gender Dysphoria: Diagnosis made and prescribed by, or in consultation with, a specialist in the treatment of gender dysphoria All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Testopel: • Authorization: 12 months (maximum of 4 treatments), unless otherwise specified All other formulations: • Authorization: 24 months, unless otherwise specified



TEZEPELUMAB-EKKO

Affected Medications: TEZSPIRE (tezepelumab-ekko)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
OOVERED OSCS.	plan design
	 Add-on maintenance treatment of patients aged 12 years and older with severe asthma
Required Medical	Diagnosis of severe asthma defined by the following:
Information:	
illorillation.	 For adults: FEV1 less than 80% at baseline or FEV1/FVC reduced by at least 5% from normal
	 For adolescents aged 12 to 17: FEV1 less than 90% at baseline or FEV1/FVC reduced by at least 5% from normal
Appropriate	Documented use of high-dose inhaled corticosteroid (ICS) plus a long-acting beta
Treatment	agonist (LABA) for at least three months with continued symptoms
Regimen & Other	A documented history of 2 or more asthma exacerbations requiring oral or systemic
Criteria:	corticosteroid treatment in the past 12 months while on combination inhaled treatment
	with at least 80% adherence
	<u>Reauthorization</u> : documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	Use in combination with another monoclonal antibody (e.g., Fasenra, Nucala, Xolair, Dupixent, Cinqair)
Age Restriction:	12 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, an allergist, immunologist, or pulmonologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **THALIDOMIDE**

Affected Medications: THALOMID (thalidomide)

All Food and Drug Administration (FDA)-approved or compendia-supported indications not otherwise excluded by plan design Multiple Myeloma (MM) Erythema Nodosum (ENL) Systemic light chain amyloidosis AIDS-related aphthous stomatitis Waldenström macroglobulinemia Graft-versus-host disease, chronic (refractory) NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher Required Medical Information: Appropriate Treatment Regimen & Other Criteria: MCCN (National Comprehensive Cancer Network) regimen with evidence level of 2A or higher NCCN (National Comprehensive Cancer Network) regimen with evidence level of 2A or higher NCCN (National Comprehensive Cancer Network) regimen with evidence level of 2A or higher Systemic light chain amyloidosis NCCN (National Comprehensive Cancer Network) regimen with evidence level of 2A or higher Waldenström Macroglobulinemia NCCN (National Comprehensive Cancer Network) regimen with evidence level of 2A or higher AIDS-related or Severe recurrent aphthous stomatitis Documented trial and failure with BOTH topical and systemic corticosteroids Erythema Nodosum Leprosum (ENL) Acute treatment of the cutaneous manifestations of moderate to severe ENL (Type 2 reaction) Maintenance therapy for prevention and suppression of the cutaneous manifestations of ENL recurrence Reauthorization: Documentation of disease responsiveness to therapy Exclusion Criteria: Pregnancy Karnofsky Performance Status less than or equal to 50% or ECOG performance score greater than or equal to 3 Age Restriction: Prescriber/Site of Care Restrictions: Initial Authorization: 4 months, unless otherwise specified		
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Appropriate Treatment Treatment Treatment Treatment Treatment Nultiple Myeloma Nultiple Myel	Required Medical	Documentation of performance status, disease staging, all prior therapies used, and
Appropriate Treatment Regimen & Other Criteria: Systemic light chain amyloidosis	-	· · · · · · · · · · · · · · · · · · ·
NCCN (National Comprehensive Cancer Network) regimen with evidence level of 2A or higher Systemic light chain amyloidosis NCCN (National Comprehensive Cancer Network) regimen with evidence level of 2A or higher Waldenström Macroglobulinemia NCCN (National Comprehensive Cancer Network) regimen with evidence level of 2A or higher AIDS-related or Severe recurrent aphthous stomatitis Documented trial and failure with BOTH topical and systemic corticosteroids Erythema Nodosum Leprosum (ENL) Acute treatment of the cutaneous manifestations of moderate to severe ENL (Type 2 reaction) Maintenance therapy for prevention and suppression of the cutaneous manifestations of ENL recurrence Reauthorization: Documentation of disease responsiveness to therapy Exclusion Criteria: Pregnancy Karnofsky Performance Status less than or equal to 50% or ECOG performance score greater than or equal to 3 Age Restriction: Prescriber/Site of Care Restrictions: All approvals are subject to utilization of the most cost-effective site of care	Appropriate	
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reaction) • Maintenance therapy for prevention and suppression of the cutaneous manifestations of ENL recurrence Reauthorization: Documentation of disease responsiveness to therapy • Pregnancy • Karnofsky Performance Status less than or equal to 50% or ECOG performance score greater than or equal to 3 Age Restriction: Prescriber/Site of Care Restrictions: • 12 years of age and older • Prescribed by, or in consultation with, an oncologist or infectious disease specialist • All approvals are subject to utilization of the most cost-effective site of care		Erythema Nodosum Leprosum (ENL)
 Maintenance therapy for prevention and suppression of the cutaneous manifestations of ENL recurrence Reauthorization: Documentation of disease responsiveness to therapy Exclusion Criteria: Pregnancy Karnofsky Performance Status less than or equal to 50% or ECOG performance score greater than or equal to 3 Age Restriction: Prescriber/Site of Care Restrictions: All approvals are subject to utilization of the most cost-effective site of care 		Acute treatment of the cutaneous manifestations of moderate to severe ENL (Type 2)
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 Prescriber/Site of Care Restrictions: Prescribed by, or in consultation with, an oncologist or infectious disease specialist All approvals are subject to utilization of the most cost-effective site of care 		
 Prescriber/Site of Care Restrictions: Prescribed by, or in consultation with, an oncologist or infectious disease specialist All approvals are subject to utilization of the most cost-effective site of care 	Age Restriction:	12 years of age and older
• All approvals are subject to utilization of the most cost-effective site of care		
Coverage Duration: • Initial Authorization: 4 months, unless otherwise specified		
	Coverage Duration:	Initial Authorization: 4 months, unless otherwise specified



•	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: TIRZEPATIDE

Affected Medications: ZEPBOUND (tirzepatide)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise excluded by plan design Treat moderate to severe obstructive sleep apnea (OSA) in combination with a reduced-calorie diet and increased physical activity in adults with obesity
Required Medical Information:	 Diagnosis of moderate to severe obstructive sleep apnea (OSA) with Apnea-Hypopnea Index (AHI) of at least 15 on polysomnography Body mass index (BMI) of 30 or greater Documentation of being used in combination with caloric restriction (diet), increased physical activity, and behavioral modification
Appropriate Treatment Regimen & Other Criteria:	Reauthorization requires documentation of treatment success defined by an improvement in AHI score and OSA symptoms (such as less daytime sleepiness, fewer sleep arousals, fewer pauses in breathing)
Exclusion Criteria:	 Diagnosis of type 1 or type 2 diabetes with or without OSA Diagnosis of central or mixed sleep apnea Diagnosis of obesity hypoventilation syndrome or daytime hypercapnia History of ketoacidosis Personal or family history of medullary thyroid carcinoma (MTC) or Multiple Endocrine Neoplasia syndrome type 2 (MEN 2)
Age Restriction:	18 years of age and older
Prescriber/Site of Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



TOBRAMYCIN INHALATION

Affected Medications: BETHKIS (tobramycin), KITABIS PAK (tobramycin), TOBI (tobramycin), TOBI PODHALER (tobramycin), TOBRAMYCIN NEBULIZED SOLUTION

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Management of Cystic Fibrosis (CF) patients with Pseudomonas aeruginosa
Required Medical Information:	 Diagnosis of Cystic Fibrosis (phenotyping not required). Culture and sensitivity report confirming presence of pseudomonas aeruginosa in the lungs Baseline forced expiratory volume in 1 second (FEV1) Tobi Podhaler: FEV1 equal to or between 25% and 80% Bethkis: FEV1 equal to or between 40% and 80% Kitabis Pak: FEV1 equal to or between 25% and 75%
Appropriate Treatment Regimen & Other Criteria:	 For Tobi Podhaler, Kitabis Pak, Bethkis, and Tobi: Documentation of failure with nebulized tobramycin or clinical rationale for avoidance Use is limited to a 28 days on and 28 days off regimen Reauthorization requires documentation of improved respiratory symptoms and need for long-term use
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a pulmonologist, or provider who specializes in CF All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



POLICY NAME: **TOFERSEN**

Affected Medications: QALSODY (tofersen)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	o Amyotrophic lateral sclerosis (ALS) associated with a mutation in the superoxide
	dismutase 1 (SOD1) gene (SOD1-ALS)
Required Medical Information:	Documentation of "definite" or "probable" ALS diagnosis based on revised El Escorial (Airlie House) or Awaji criteria
	Documentation of a confirmed SOD1 genetic mutation
	• Forced vital capacity (FVC) greater than or equal to 50% as adjusted for age, sex, and height (from a sitting position)
	Baseline plasma neurofilament light chain (NfL) value
	 Patient currently retains most activities of daily living defined as at least 2 points on all 12 items of the ALS functional rating scale-revised (ALSFRS-R)
Appropriate	Reauthorization requires documentation of treatment success and a clinically significant
Treatment	response to therapy, defined as both of the following:
Regimen & Other	Reduction in plasma NfL from baseline
Criteria:	The patient's baseline functional status has been maintained at or above baseline level
Oritoria.	or not declined more than expected given the natural disease progression
	 Patient is not dependent on invasive mechanical ventilation (e.g., intubation, tracheostomy)
Exclusion Criteria:	
Age Restriction:	18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist, neuromuscular specialist, or
Care Restrictions:	specialist with experience in the treatment of ALS
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
-	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **TOLVAPTAN**

Affected Medications: JYNARQUE, TOLVAPTAN (15 mg, 30 mg)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Tolvaptan: treatment of clinically significant hypervolemic and euvolemic hyponatremia (serum sodium less than 125 mEq/L OR less marked hyponatremia that is symptomatic and has resisted correction with fluid restriction), including patients with heart failure and Syndrome of Inappropriate Antidiuretic Hormone (SIADH) Jynarque: to slow kidney function decline in adults at risk of rapidly progressing autosomal dominant polycystic kidney disease (ADPKD)
Required Medical Information:	Hyponatremia Serum sodium less than 125 mEq/L at baseline OR Serum sodium less than 135 mEq/L at baseline and symptomatic (nausea, vomiting, headache, lethargy, confusion)
	 ADPKD Diagnosis of typical ADPKD confirmed by family history, imaging, and if applicable, genetic testing Estimated glomerular filtration rate (eGFR) greater than or equal to 25 mL/min/1.73m² High risk for rapid progression determined by Mayo imaging class 1C, 1D, or 1E
Appropriate Treatment Regimen & Other Criteria:	Hyponatremia Treatment is initiated or re-initiated in a hospital setting prior to discharge ADPKD Documentation of intensive blood pressure control with an angiotensin-converting enzyme (ACE) inhibitor or angiotensin receptor blocker (ARB), unless contraindicated Reauthorization (for ADPKD) requires documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	 Patients requiring intervention to raise serum sodium urgently to prevent or treat serious neurological symptoms Patients who are unable to sense or respond to thirst Hypovolemic hyponatremia Anuria Uncorrected urinary outflow obstruction
Age Restriction:	18 years of age and older



Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a nephrologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 1 month (no reauthorization), unless otherwise specified
	<u>ADPKD</u>
	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



TOPICAL AGENTS FOR CUTANEOUS T-CELL LYMPHOMA (including Mycosis fungoides and Sézary syndrome)
Affected Medications: VALCHLOR (mechlorethamine topical gel), TARGRETIN (bexarotene gel)

Covered Uses: Required Medical	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher Documentation of performance status, disease staging, all prior therapies used, and
Information:	anticipated treatment course
	Documentation of cutaneous T-cell lymphoma (CTCL), stage and type confirmed by biopsy.
	Extent of skin involvement (limited/localized or generalized)
Appropriate	Limited/localized skin involvement (topical bexarotene and mechlorethamine)
Treatment	Documented clinical failure to ALL of the following:
Regimen & Other	 Topical corticosteroids (high or super-high potency) such as clobetasol,
Criteria:	betamethasone, fluocinonide, halobetasol
	Topical imiquimod
	o Phototherapy
	Generalized skin involvement (topical mechlorethamine only)
	Documentation of failure or contraindication to at least 1 skin-directed therapy
	Reauthorization: documentation of disease responsiveness to therapy
Exclusion Criteria:	Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
	Pregnancy
Age Restriction:	18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 4 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



TOPICAL DERMATITIS AND PSORIATIC AGENTS

Affected Medications: VTAMA (tapinarof 1% cream), ZORYVE (roflumilast 0.3% cream), ZORYVE (roflumilast 0.3% foam), ZORYVE (roflumilast 0.15% cream)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Plaque psoriasis (Vtama and Zoryve 0.3% cream) Seborrheic dermatitis (Zoryve 0.3% foam) Atopic dermatitis (Vtama and Zoryve 0.15% cream)
Required Medical Information:	All Indications Documentation of affected body surface area (BSA) and areas of involvement
	 Plaque Psoriasis Documentation of chronic plaque psoriasis that meets <u>ONE</u> of the following: At least 10% BSA involvement despite current treatment Hand, foot, face, or mucous membrane involvement
	 Seborrheic Dermatitis Diagnosis of moderate to severe seborrheic dermatitis with presence of lesions that are characteristic of the condition (such as erythematous plaques and yellowish scales distributed on areas with sebaceous glands) Documentation of persistent itching, scaling, and erythema despite current therapy
	 Atopic Dermatitis Documentation of atopic dermatitis that meets <u>ONE</u> of the following: At least 10% BSA involvement despite current treatment Hand, foot, face, or mucous membrane involvement
Appropriate Treatment Regimen &	All Indications Documented treatment failure with a high or super-high potency topical corticosteroid
Other Criteria:	 Plaque Psoriasis Documented treatment failure with each of the following for a minimum of 4-weeks: Topical vitamin D analog (e.g., calcipotriene, calcitriol) Tazarotene <u>Vtama</u>: Requires additional treatment failure with 8 weeks of Zoryve 0.3% cream
	Reauthorization requires documentation of disease responsiveness to therapy, defined as a decrease in affected BSA from baseline
	Seborrheic Dermatitis Documented failure with ALL the following: Minimum 6-week trial of one topical calcineurin inhibitor (e.g., tacrolimus, pimecrolimus) Topical antifungal (such as ketoconazole, ciclopirox, or selenium sulfide)



	Reauthorization requires documentation of disease responsiveness to therapy, defined as a reduction in itching, scaling, erythema, and number of affected areas compared to baseline
	 Atopic Dermatitis Documented treatment failure with a minimum 6-week trial of one of the following: tacrolimus ointment or pimecrolimus cream Vtama: Requires additional treatment failure with 4 weeks of Zoryve 0.15% cream or Eucrisa
	Reauthorization requires documentation of disease responsiveness, defined as a decrease in affected BSA from baseline
Exclusion Criteria:	
Age Restriction:	Vtama: 18 years of age and older (plaque psoriasis) 2 years of age and older (atopic dermatitis) 7 years of age and older (atopic dermatitis)
	Zoryve cream: 6 years of age and olderZoryve foam: 9 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a dermatologist, allergist, or immunologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 6 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: TRALOKINUMAB

Affected Medications: ADBRY (tralokinumab)

1.	Is the request for continuation of therapy currently approved through insurance?	Yes – Go to renewal criteria	No – Go to #2	
2.	Is the request to treat a diagnosis according to one of the Food and Drug Administration (FDA)-approved indications? - Treatment of moderate to severe atopic dermatitis in adults	Yes – Go to appropriate section below	No – Criteria not met	
Мо	derate to Severe Atopic Dermatitis			
1.	Is there documentation of severe inflammatory skin disease defined as functional impairment (inability to use hands or feet for activities of daily living, or significant facial involvement preventing normal social interaction)?	Yes – Document and go to #2	No – Criteria not met	
2.	Is there a documented body surface area (BSA) effected of at least 10% OR hand, foot or mucous membrane involvement?	Yes – Document and go to #3	No – Criteria not met	
3.	Is there documented failure with at least 6 weeks of treatment with one of the following: tacrolimus ointment, pimecrolimus cream, Eucrisa?	Yes – Document and go to #4	No – Criteria not met	
4.	Is there documented treatment failure with two of the following for at least 12 weeks: Phototherapy, cyclosporine, azathioprine, methotrexate, mycophenolate?	Yes – Document and go to #5	No – Criteria not met	
5.	Is the drug prescribed by, or in consultation with, a specialist in the treatment of atopic dermatitis (Such as a dermatologist)?	Yes – Approve up to 6 months	No – Criteria not met	
Re	Renewal Criteria			
1.	Is there documentation of treatment success and a clinically significant response to therapy as assessed by the prescribing provider?	Yes – Go to #2	No – Criteria not met	
2.	Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations?	Yes – Approve up to 12 months	No – Criteria not met	



Quantity Limitations

- Adbry
 - Availability: 150 mg/mL prefilled syringes, 300 mg/2 mL autoinjectors
 - o Dosing:
 - Adults 18 years and older: 600 mg as single dose, then 300 mg every 2 weeks.
 - If less than 100 kg and clear/almost clear is achieved, dosing may be reduced to 300 mg every 4 weeks
 - Pediatric patients 12 to 17 years old: 300 mg as a single dose, then 150 mg every 2 weeks.



TRASTUZUMAB

Affected Medications: HERCEPTIN IV (trastuzumab), HERCEPTIN HYLECTA SQ (trastuzumab and hyaluronidase), KANJINTI (trastuzumab-anns), OGIVRI (trastuzumab-dkst), TRAZIMERA (trastuzumab-qyyp), HERZUMA (trastuzumab-pkrb), ONTRUZANT (trastuzumab-dttb), HERCESSI (Trastuzumab-strf)

	-
Covered Uses:	NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher
Required Medical Information:	 Documentation of performance status, disease staging, all prior therapies used, and prescribed dosing regimen Documentation of HER2 positivity based on: 3+ score on immunohistochemistry (IHC) testing OR Positive gene amplification by fluorescence in situ hybridization (FISH) test
Appropriate	Maximum duration for adjuvant breast cancer therapy is 12 months
Treatment	
Regimen & Other	All Indications
Criteria:	Coverage for a non-preferred product (Trazimera, Herzuma, Ontruzant, Herceptin, Herceptin, Hylosta) requires desumptation of the following:
	Hercessi or Herceptin Hylecta) requires documentation of the following:
	 A documented intolerable adverse event to the preferred products Kanjinti and Ogivri and the adverse event was not an expected adverse event attributed to the
	active ingredient
	double miground
	Reauthorization requires documentation of disease responsiveness to therapy
Exclusion Criteria:	Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	For new starts to adjuvant breast cancer therapy – approve for 12 months with no
-	reauthorization
	For all other clinical scenarios:
	 Initial Authorization: 4 months, unless otherwise specified
	 Reauthorization: 12 months, unless otherwise specified



TRIENTINE

Affected Medications: TRIENTINE HYDROCHLORIDE, CUVRIOR (trientine tetrahydrochloride)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
Covered Oses.	plan design.
	o Wilson's disease
	o Wilson's disease
Required Medical	Diagnosis of Wilson's disease confirmed by ONE of the following:
Information:	 Genetic testing results confirming biallelic pathogenic ATP7B mutations (in either
	symptomatic or asymptomatic individuals)
	 Liver biopsy findings consistent with Wilson's disease
	 Presence of Kayser-Fleischer (KF) rings AND serum ceruloplasmin level less
	than 20 mg/dL AND 24-hour urinary copper excretion greater than 40 mcg
	 Presence of Kayser-Fleischer (KF) rings AND 24-hour urinary copper excretion
	greater than 100 mcg
	 Absence of KF rings with serum ceruloplasmin level less than 10 mg/dL AND 24-
	hour urinary copper excretion greater than 100 mcg
Appropriate	For Cuvrior, must meet BOTH of the following:
Treatment	 Documented treatment failure with a minimum 6-month trial of penicillamine that
Regimen & Other	was not due to tolerability
Criteria:	 Documented intolerable adverse event to a maximally tolerated dosage of
	generic trientine hydrochloride and the adverse event was not an expected
	adverse event attributed to the active ingredient
	Reauthorization: Documentation of treatment success and a clinically significant response to
	therapy as shown by normalization of free serum copper (non-ceruloplasmin bound copper)
	to less than 15 mcg/dL and 24-hour urinary copper in the range of 200 to 500 mcg
	to less than 13 mog/de and 24-notifically copper in the range of 200 to 300 mog
Exclusion Criteria:	For trientine hydrochloride:
	Treatment of rheumatoid arthritis
	Treatment of cystinuria
	Treatment of biliary cirrhosis
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a hepatologist, gastroenterologist, or liver
Care Restrictions:	transplant provider
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization:12 months, unless otherwise specified
	- ReductionZacion. 12 months, unicos outorwise specifica



POLICY NAME: **TRIPTORELIN**

Affected Medications: TRELSTAR, TRIPTODUR (triptorelin)

Required Medical Information:	 NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Prostate Cancer (Trelstar) Central Precocious Puberty (Triptodur) Gender Dysphoria Central Precocious Puberty (CPP) Documentation of CPP confirmed by one of the following labs: Elevated basal luteinizing hormone (LH) level greater than 0.2 - 0.3 mlU/L Elevated leuprolide-stimulated LH level greater than 3.3 - 5 lU/L (dependent on type of assay used) Bone age greater than 2 standard deviations (SD) beyond chronological age Gender Dysphoria
	Documentation of all the following: Current Tanner stage 2 or greater OR baseline and current estradiol and testosterone levels to confirm onset of puberty Confirmed diagnosis of gender dysphoria that is persistent The patient has the capacity to make a fully informed decision and to give consent for treatment Any significant medical or mental health concerns are reasonably well controlled A comprehensive mental health evaluation has been completed by a licensed mental health professional (LMHP) and provided in accordance with the most current version of the World Professional Association for Transgender Health (WPATH) Standards of Care
Appropriate Treatment Regimen & Other Criteria:	For all Triptodur requests: Documentation of treatment failure with leuprolide Reauthorization will require documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria: Age Restriction:	 Use as neoadjuvant androgen deprivation therapy (ADT) for radical prostatectomy CPP: 2 years of age through 11 years for females, 2 years of age through 12 years for
	males
Prescriber/Site of Care Restrictions:	 Oncology: prescribed by, or in consultation with, an oncologist CPP: prescribed by, or in consultation with, a pediatric endocrinologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Oncology Initial Authorization: 4 months, unless otherwise specified CPP Approval/Oncology Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **TROFINETIDE**

Affected Medications: DAYBUE (trofinetide)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of Rett syndrome (RTT)
Required Medical Information:	 Documented diagnosis of typical RTT (per the revised diagnostic criteria for Rett Syndrome) AND a period of regression followed by recovery or stabilization Documented presence of mutation in the MECP2 gene Documentation of all the following: Partial or complete loss of acquired purposeful hand skills Partial or complete loss of acquired spoken language Gait abnormalities: Impaired (dyspraxic) or absence of ability Stereotypic hand movements such as hand wringing/squeezing, clapping/tapping, mouthing, and washing/rubbing automatisms Current weight (within past 30 days) Must weigh minimum of 9 kilograms
Appropriate	Reauthorization requires documentation of treatment success determined by treating
Treatment	provider
Regimen & Other Criteria:	
Exclusion Criteria:	Brain injury secondary to trauma or severe infection
	Grossly abnormal psychomotor development in first 6 months of life
Age Restriction:	2 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist or provider experienced in the
Care Restrictions:	management of Rett syndrome
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: TROGARZO

Affected Medications: TROGARZO (ibalizumab-uiyk/IV infusion)

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Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of human immunodeficiency virus type 1 (HIV-1) infection, in combination with other antiretrovirals, in heavily treatment-experienced adults with multidrug resistant HIV-1 infection failing their current antiretroviral regimen
Required Medical	Documentation of all prior therapies used
Information:	Documentation of active antiretroviral therapy for at least 6 months
	Documented resistance to at least one antiretroviral agent from three different classes:
	N. J. C.
	, , ,
	Non-nucleoside reverse-transcriptase inhibitors (NNRTIs) Integrand strengther inhibitors (INCTIs)
	Integrase strand transfer inhibitors (INSTIs)
	o Protease inhibitors (PIs)
	Documentation of current (within the past 30 days) HIV-1 RNA viral load of at least 200
	copies/mL
Appropriate Treatment	Prescribed in combination with an optimized background antiretroviral regimen
Regimen & Other	
Criteria:	Reauthorization requires all of the following:
	Treatment plan includes continued use of optimized background antiretroviral regimen
	Documentation of treatment success as evidenced by one of the following:
	 Reduction in viral load from baseline or maintenance of undetectable viral load
	 Absence of postbaseline emergence of ibalizumab resistance-associated
	mutations confirmed by resistance testing
Exclusion Criteria:	
Age Restriction:	18 years of age and older
Prescriber/Site of Care	Prescribed by, or in consultation with, an infectious disease or HIV specialist
Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 3 months, unless otherwise specified
	Reauthorization 12 months, unless otherwise specified
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POLICY NAME: **TUCATINIB**

Affected Medications: TUKYSA (tucatinib)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A
	or better
Required Medical Information:	Documentation of performance status, disease staging, all prior therapies used, and anticipated treatment course
	Documentation of RAS wild-type, human epidermal growth factor receptor-2 (HER2) positive, unresectable or metastatic colorectal cancer that has progressed following treatment with fluoropyrimidine, oxaliplatin, and irinotecan-based chemotherapy OR
	Advanced, unresectable or metastatic, HER2-positive breast cancer with prior treatment of 1 or more anti-HER2-based regimens in the metastatic setting
Appropriate	Colorectal cancer
Treatment	Documented intolerable adverse event to Lapatinib
Regimen & Other Criteria:	Reauthorization: documentation of disease responsiveness to therapy
Exclusion Criteria:	 Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater Colorectal cancer ONLY: previous treatment with a HER2 inhibitor
Age Restriction:	18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 4 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



TYVASO

Affected Medications: TYVASO (treprostinil inhalation)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Pulmonary Arterial Hypertension (PAH) World Health Organization (WHO) Group
	7 Pulmonom: Antonial I han antonoion (DAII) Mondal Hoolth Organization (M/HO) Croun
	 Pulmonary Arterial Hypertension (PAH) World Health Organization (WHO) Group
Required	Pulmonary Arterial Hypertension (PAH) WHO Group 1
Medical	Documentation of PAH confirmed by right-heart catheterization meeting the following
Information:	criteria:
	 Mean pulmonary artery pressure of at least 20 mm Hg
	 Pulmonary capillary wedge pressure less than or equal to 15 mm Hg
	 Pulmonary vascular resistance of at least 2.0 Wood units
	Etiology of PAH: idiopathic PAH, hereditary PAH, OR
	PAH secondary to one of the following conditions:
	Connective tissue disease
	 Human immunodeficiency virus (HIV) infection
	o Drugs
	Congenital left to right shunts
	Schistosomiasis Portal hypothesian
	 Portal hypertension New York Heart Association (NYHA)/World Health Organization (WHO) Functional Class
	III or higher symptoms
	Documentation of Acute Vasoreactivity Testing (positive result requires trial/failure to
	calcium channel blockers) unless there are contraindications:
	 Low systemic blood pressure (systolic blood pressure less than 90)
	o Low cardiac index
	OR
	 Presence of severe symptoms (functional class IV)
	Pulmonary Hypertension Associated with Interstitial Lung Disease WHO Group 3 Documentation of diagnosis of idiopathic pulmonary fibrosis confirmed by presence of usual interstitial pneumonia (UIP) on high resolution computed tomography (HRCT), and/or surgical lung biopsy OR
	Pulmonary fibrosis and emphysema OR
	Connective tissue disorder
Appropriate	The pulmonary hypertension has progressed despite maximal medical and/or surgical
Treatment	treatment of the identified condition
Regimen &	Documentation that treprostinil is used as a single route of administration (Remodulin,
Other Criteria:	Tyvaso, Orenitram should not be used in combination)
	WHO Group 1 only:



Exclusion	Treatment with oral calcium channel blocking agents has been tried and failed, or has been considered and ruled out Treatment with combination of endothelin receptor antagonist (ERA) and phosphodiesterase 5 inhibitor (PDE5I) has been tried and failed for WHO functional class II and III Ambrisentan and tadalafil Bosentan and riociguat Macitentan and sildenafil Reauthorization requires documentation of treatment success defined as one or more of the following: Improvement in walking distance Improvement in exercise ability Improvement in pulmonary function Improvement or stability in WHO functional class PAH secondary to pulmonary venous hypertension such as left sided atrial or ventricular
Criteria:	disease, left sided valvular heart disease, or disorders of the respiratory system such as chronic obstructive pulmonary disease, obstructive sleep apnea or other sleep disordered breathing, alveolar hypoventilation disorders, etc.
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a cardiologist or pulmonologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage	Initial Authorization: 6 months unless otherwise specified
Duration:	Reauthorization: 12 months unless otherwise specified



POLICY NAME: UBLITUXIMAB-XIIY

Affected Medications: BRIUMVI (ublituximab-xiiy)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Treatment of relapsing forms of multiple sclerosis (MS), including the following:
Required Medical	Relapsing Forms of MS
Information:	 Diagnosis confirmed with magnetic resonance imaging (MRI), per revised McDonald diagnostic criteria for MS
	 Clinical evidence alone will suffice; additional evidence desirable, but must be consistent with MS
Appropriate	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced
Treatment	
Regimen & Other	Documentation of one of the following:
Criteria:	 Documented disease progression or intolerance to rituximab (preferred products: Riabni and Ruxience)
	 Currently receiving treatment with Briumvi, excluding via samples or manufacturer's patient assistance program
	Reauthorization requires documentation of treatment success
Exclusion Criteria:	Active hepatitis B infection
	Concurrent use of disease-modifying medications indicated for the treatment of multiple sclerosis
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist or a multiple sclerosis specialist
Care Restrictions:	All approved are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



UPNEEQ

Affected Medications: UPNEEQ (oxymetazoline opthalmic solution)

Upneeq (oxymetazoline opthalmic solution) is not considered medically necessary insufficient evidence of therapeutic value. Required Medical Information:	due to
·	
Information:	
mornation.	
Appropriate	
Treatment	
Regimen & Other	
Criteria:	
Exclusion Criteria:	
Age Restriction:	
Prescriber/Site of	
Care Restrictions:	
Coverage Duration:	



VAGINAL PROGESTERONE

Affected Medications: FIRST-PROGESTERONE VGS 100 MG, FIRST-PROGESTERONE VGS 200 MG

Covered Uses:	Prevention of preterm birth in pregnancy		
Required Medical Information:	Documentation of a current pregnancy with one or more risk factor(s) for preterm birth, including but not limited to: Ethnicity (e.g., African American, American Indian/Alaska Native) Lifestyle factors (e.g., smoking, drinking alcohol, using illegal drugs) Being underweight or obese before pregnancy Prior preterm delivery Having multiple gestations (e.g., twins, triplets) Short time period between pregnancies (less than 6 months between a birth and the beginning of the next pregnancy) Documentation of a short cervix (defined as cervical length less than or equal to 25 mm) confirmed by ultrasound Current week of gestation and estimated delivery date		
Appropriate Treatment Regimen & Other Criteria:	May continue until completion of 36 weeks gestation		
Exclusion Criteria:	Treatment of infertility		
Age Restriction:			
Prescriber/Site of	Prescribed by, or in consultation with, a gynecologist or obstetrician		
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care		
Coverage Duration:	Authorization: up to 6 months, unless otherwise specified		



VALOCTOCOGENE ROXAPARVOVEC-RVOX

Affected Medications: ROCTAVIAN (valoctocogene roxaparvovec-rvox) - Available on Medical Benefit only

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Hemophilia A (Factor VIII deficiency) 		
Required Medical Information:	 Documentation of diagnosis of Hemophilia A Documentation of current testing with negative results for active factor VIII inhibitors on 2 consecutive occasions (at least one week apart within the past 12 months) and is not receiving a bypassing agent (e.g., Feiba) Documentation of baseline circulating level of factor with Factor VIII activity level equal to or less than 1 IU/dL or 1% endogenous factor VIII Evidence of any bleeding disorder NOT related to hemophilia A has been ruled out No detectable antibodies to AAV5 as determined by an FDA-approved/CLIA-compliant test Has received stable dosing of prophylactic Factor VIII replacement therapy on a regular basis for at least 1 year Baseline lab values (must be less than 2 times upper limit of normal): ALT AST Total bilirubin Alkaline phosphatase (ALP) ALP 		
Appropriate Treatment Regimen & Other Criteria:	 Dosing 6 x 10¹³ vector genomes/kg (which is 3 mL/kg) as a single one-time dose 		
Exclusion Criteria:	 History of or current presence of Factor VIII inhibitors Prior gene therapy administration Active Hepatitis B or C infection or other active acute or uncontrolled chronic infection Cirrhosis Female gender at birth Allergy to mannitol 		
Age Restriction:	18 years of age and older		
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation, with a hematologist or specialist with experience in treatment of hemophilia All approvals are subject to utilization of the most cost-effective site of care 		
Coverage Duration:	Authorization: 2 months (one time infusion), unless otherwise specified		



POLICY NAME: **VAMOROLONE**

Affected Medications: AGAMREE (vamorolone)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by				
	plan design				
	 Duchenne muscular dystrophy (DMD) in patients 2 years of age and older 				
Required Medical	Confirmation of Duchenne muscular dystrophy (DMD) diagnosis by genetic testing or				
Information:	biopsy showing lack of muscle dystrophin				
	Documentation of being ambulatory without needing an assistive device such as a				
	wheelchair, walker, or cane				
	Baseline motor function assessment from one of the following:				
	 Time to Stand Test (TTSTAND) 				
	o 6-minute walk test				
	North Star Ambulatory Assessment (NSAA)				
	Motor Function Measure (MFM)				
	Hammersmith Functional Motor Scale (HFMS)				
	Patient weight and planned treatment regimen				
Appropriate	Documented treatment failure with a 6-month trial of prednisone, or intolerable adverse				
Treatment	event causing one of the following:				
Regimen & Other	 Clinically significant weight gain defined as greater than or equal to 10% of body 				
Criteria:	weight gain over a 6-month period				
	 Psychiatric/behavioral issues (e.g., abnormal behavior, aggression, irritability) 				
	that persists beyond the first six weeks of prednisone treatment				
	Reauthorization requires a documented improvement from baseline or stabilization of motor				
	function demonstrated by a motor function assessment tool				
Exclusion Criteria:					
Age Restriction:	2 years of age and older				
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist				
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care				
Coverage Duration:	Initial Authorization: 6 months, unless otherwise specified				
	Reauthorization: 12 months, unless otherwise specified				



VARIZIG

Affected Medications: VARIZIG (varicella zoster immune globulin (human) IM injection)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design For post exposure prophylaxis of varicella in high-risk individuals 		
Required Medical Information:	 Documentation of immunocompromised patient, defined as: Newborns of mothers with signs and symptoms of varicella shortly before or after delivery (five days before to two days after delivery) Hospitalized premature infants born at least 28 weeks of gestation who are exposed during their hospitalization and whose mothers do not have evidence of immunity Hospitalized premature infants less than 28 weeks of gestation or who weigh 1000 grams or less at birth and were exposed to varicella during hospitalization, regardless of mother's immunity status to varicella Immunocompromised children and adults who lack evidence of immunity to varicella Pregnant women who lack evidence of immunity to varicella Lack evidence of immunity to varicella is defined as: those who are seronegative for varicella zoster antibodies OR those with unknown history of varicella 		
Appropriate Treatment Regimen & Other Criteria:	If repeat dose is necessary due to re-exposure, use more than 3 weeks after initial administration		
Exclusion Criteria:	Coagulation disorders		
Age Restriction:			
Prescriber/Site of Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care		
Coverage Duration:	Authorization: 6 months, unless otherwise specified		



VELMANASE ALFA-TYCV

Affected Medications: LAMZEDE (velmanase alfa-tycv)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design The treatment of non-central nervous system manifestations of alphamannosidosis			
Required Medical Information:	 Diagnosis of alpha-mannosidosis (AM) confirmed by enzyme assay demonstrating alpha-mannosidase activity less than 10% of normal activity Documentation of symptoms consistent with AM such as hearing impairment, difficulty walking, skeletal abnormalities, or intellectual disabilities 			
Appropriate Treatment Regimen & Other Criteria:	Reauthorization will require documentation of treatment success such as improvement in motor function, forced vital capacity (FVC), or reduction in frequency of infections			
Exclusion Criteria:	AM with only central nervous system manifestations and no other symptoms			
Age Restriction:				
Prescriber/Site of Care Restrictions:	 All approvals are subject to utilization of the most cost-effective site of care Prescribed by, or in consultation with, a specialist familiar with the treatment of lysosomal storage disorders 			
Coverage Duration:	Authorization: 12 months, unless otherwise specified			



POLICY NAME: **VERTEPORFIN**

Affected Medications: VISUDYNE (verteporfin)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by			
	plan design			
	Treatment of predominantly classic subfoveal choroidal neovascularization (CNN) due to see at the fall principle.			
	(CNV) due to one of the following:			
	Age-related macular degeneration (AMD)Pathologic myopia			
	Presumed ocular histoplasmosis			
Required Medical	Documented diagnosis of subfoveal CNV due to one of the following:			
Information:	Neovascular AMD			
	5 , ,			
	 Presumed ocular histoplasmosis Documentation of current body surface area (BSA) 			
Appropriate	Neovascular AMD and Pathologic Myopia			
Treatment	Documentation of one of the following:			
Regimen & Other	Currently receiving treatment with Visudyne, excluding via samples or			
Criteria:	manufacturer's patient assistance program			
Ontena.	Documented treatment failure or intolerance following a minimum 3-month trial			
	with both of the following: Avastin and ranibizumab (preferred products: Byooviz,			
	with both of the following: Avastin and ranibizumab (preferred products: Byooviz, Cimerli)			
	Oillielli)			
	Dosing			
	• 6 mg/m² BSA			
	Every 3 month dosing is permitted with evidence of choroidal neovascular			
	leakage (see reauthorization criteria)			
	Dose-rounding to the nearest vial size within 10% of the prescribed dose will be enforced.			
	Reauthorization requires documentation of the following:			
	Positive response to therapy (e.g., improved or stable visual acuity, reduced central			
	macular thickness)			
	Evidence of recurrent or persistent leakage on fluorescein angiogram or optical			
	coherence tomography (OCT), performed at least 3 months after the last treatment			
Exclusion Criteria:	Concurrent therapy with vascular endothelial growth factor (VEGF) inhibitors			
	Treatment of non-neovascular (dry) AMD			
Age Restriction:				
Prescriber/Site of	Prescribed by, or in consultation with, an ophthalmologist			
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care			
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Coverage Duration:				
	Reauthorization: 12 months, unless otherwise specified			



POLICY NAME: VIGABATRIN

Affected Medications: VIGABATRIN, VIGADRONE (vigabatrin)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Refractory complex partial seizures (focal seizures with impaired awareness) 			
	 Infantile spasms 			
Required	Infantile Spasms			
Medical Information:	Used as monotherapy for pediatric patients (1 month to 2 years of age)			
	Refractory Complex Partial Seizures (focal seizures with impaired awareness)			
	Used as adjunctive therapy only			
Appropriate	Refractory complex partial seizures (focal seizures with impaired awareness)			
Treatment	Documentation of treatment failure with at least 2 alternative therapies: carbamazepine,			
Regimen & Other Criteria:	phenytoin, levetiracetam, topiramate, oxcarbazepine, or lamotrigine			
	<u>Reauthorization</u> will require documentation of treatment success and a reduction in seiz severity, frequency, and/or duration			
Exclusion Criteria:	Use as a first line agent for complex partial seizures (focal seizures with impaired awareness)			
Age	Infantile Spasms: 1 month to 2 years of age			
Restriction:	 Refractory complex partial seizures (focal seizures with impaired awareness): greater than 2 years of age 			
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist			
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care			
Coverage	Infantile Spasms			
Duration:	Initial Authorization: 6 months, unless otherwise specified			
	Reauthorization: 12 months (or up to 2 years of age), unless otherwise specified			
	Refractory Complex Partial Seizures (focal seizures with impaired awareness)			
	Authorization: 12 months, unless otherwise specified			



POLICY NAME: **VIJOICE**

Affected Medications: VIJOICE (alpelisib)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by			
0010.00 0000.	plan design			
	 Treatment of severe manifestations of PIK3CA-related overgrowth spectrum 			
	(PROS) in patients who require systemic therapy			
Required	Documented diagnosis of PROS, to include any of the following:			
Medical	CLAPOS syndrome			
Information:	CLOVES syndrome			
	 Diffuse capillary malformation with overgrowth (DCMO) 			
	Dysplastic megalencephaly (DMEG)			
	Facial infiltrating lipomatosis (FIL)			
	 Fibroadipose hyperplasia (FAH)/fibroadipose overgrowth (FAO)/hemihyperplasia 			
	multiple lipomatosis (HHML) syndrome			
	Fibroadipose vascular anomaly (FAVA)			
	Hemimegalencephaly (HMEG)			
	Klippel-Trenaunay syndrome (KTS)			
	Lipomatosis of nerve (LON)			
	Megalencephaly-capillary malformation (MCAP) syndrome			
	Muscular hemihyperplasia (HH)			
	Documentation of PIK3CA gene mutation			
	Documentation of clinical manifestations that were assessed by the treating provider as			
	severe or life-threatening and necessitating systemic treatment			
	Documentation that clinical manifestations are a direct result of a lesion that is both of the			
	following:			
	 Inoperable, as defined by the treating provider 			
	Causing functional impairment			
	Documentation of one or more target lesion(s) identified on imaging within 6 months prior			
	to request, including location(s) and volume of lesion(s)			
Appropriate	Treatment failure (or intolerable adverse event) with sirolimus for at least 6 months at a			
Treatment	dose of at least 2 mg daily in patients with lymphatic, venous, or combined manifestations			
Regimen & Other Criteria:	of disease			
	Reauthorization will require documentation of both of the following:			
	 Radiological response, defined as greater than or equal to a 20% reduction from 			
	baseline in the sum of measurable target lesion volume, confirmed by at least			
	one subsequent imaging assessment			
	 Absence of greater than or equal to a 20% increase from baseline in any target 			
	lesion, progression of non-target lesions, or appearance of a new lesion			
Exclusion Criteria:	Treatment of PIK3CA-mutated conditions other than PROS			
Age Restriction:	2 years of age and older			



Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, a specialist with experience in the treatment of PROS All approvals are subject to utilization of the most cost-effective site of care 	
Coverage	Initial Authorization: 6 months, unless otherwise specified	
Duration:	Reauthorization: 12 months, unless otherwise specified	



POLICY NAME: VISTOGARD

Affected Medications: VISTOGARD (uridine triacetate)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design			
	 For the emergency treatment of adult and pediatric patients: Following a fluorouracil or capecitabine overdose regardless of the presence of symptoms, OR Who exhibit early-onset, severe, or life-threatening toxicity affecting the cardiac or central nervous system, and/or early-onset, unusually severe adverse reactions (e.g., gastrointestinal toxicity and/or neutropenia) within 96 hours following the end of fluorouracil or capecitabine administration 			
Required Medical	Documentation of fluorouracil or capecitabine administration			
Information:	Documentation of overdose OR early-onset, severe adverse reaction, or life-threatening toxicity			
Appropriate	Dosing is in accordance with FDA labeling			
Treatment				
Regimen & Other				
Criteria:				
Exclusion Criteria:	Non-emergent treatment of adverse events associated with fluorouracil or capecitabine Use more than 96 hours following the end of fluorouracil or capecitabine administration			
Age Restriction:				
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist			
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care			
Coverage Duration:	Authorization: 7 days, unless otherwise specified			



POLICY NAME: VMAT2 INHIBITORS

Affected Medications: TETRABENAZINE, AUSTEDO (deutetrabenazine), AUSTEDO XR (deutetrabenazine)

Covered Uses:	 All Food and Drug Administration (FDA)-approved and compendia supported indications not otherwise excluded by plan design Chorea associated with Huntington's disease Tardive dyskinesia 				
Required Medical	Chorea related to Huntington's Disease				
Information:	Diagnosis of Huntington's Disease with Chorea requiring treatment				
	Tardive Dyskinesia				
	Diagnosis of moderate to severe tardive dyskinesia including all of the following:				
	 A history of at least one month of ongoing or previous dopamine receptor- 				
	blocking agent exposure				
	 Presence of dyskinetic or dystonic involuntary movements that developed either 				
	while exposed to a dopamine receptor-blocking agent, or within 4 weeks of				
	discontinuation from an oral agent (8 weeks from a depot formulation)				
	Other causes of abnormal movements have been excluded				
	Baseline evaluation of the condition using one of the following: Absolute Alexander Management Cooks (AIMC)				
	Abnormal Involuntary Movement Scale (AIMS) Superstant Region (FSRS)				
	Extrapyramidal Symptom Rating Scale (ESRS)				
Appropriate	Tardive Dyskinesia				
Treatment	Persistent dyskinesia despite dose reduction or discontinuation of the offending agent				
Regimen & Other	OR				
Criteria:	Documented clinical inability to reduce dose or discontinue the offending agent				
	Reauthorization requires documentation of treatment success and a clinically significant				
	response to therapy				
	Tardive Dyskinesia: must include an improvement in AIMS or ESRS score from baseline				
Exclusion Criteria:	Use for Huntington's comorbid with untreated or inadequately treated depression or				
	actively suicidal				
	Concomitant use with another VMAT2 inhibitor or reserpine				
	Hepatic impairment				
Age Restriction:	18 years of age and older				
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist or psychiatrist				
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care				
	,				
Coverage Duration:	Initial Authorization: 3 months, unless otherwise specified				
	Reauthorization: 12 months, unless otherwise specified				
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POLICY NAME: VOCLOSPORIN

Affected Medications: LUPKYNIS CAPSULE 7.9 MG ORAL

Is the request for continuation of therapy currently approved through insurance?	Yes – Go to renewal criteria	No – Go to #2	
Is the request to treat a diagnosis according to the Food and Drug Administration (FDA)-approved indication? a. For use in combination with a background immunosuppressive therapy regimen for the treatment of adult patients with active lupus nephritis	Yes – Go to appropriate section below	No – Criteria not met	
Lupus Nephritis (LN)			
Is there documented International Society of Nephrology/Renal Pathology Society (ISN/RPS) biopsy- proven active class III, IV and/or V disease?	Yes – Document and go to #2	No – Criteria not met	
Are there documented current baseline values (within the last 3 months) for all of the following? a. Estimated glomerular filtration rate (eGFR) b. Urine protein to creatinine ratio (uPCR) c. Blood pressure	Yes – Document and go to #3	No – Criteria not met	
Is there documented treatment failure with at least 12 weeks of standard therapy with both mycophenolate mofetil (MMF) AND cyclophosphamide?	Yes – Document and go to #4	No – Criteria not met	
Is there documented treatment failure with at least 12 weeks of subcutaneous Benlysta?	Yes – Document and go to #5	No – Criteria not met	
Will Lupkynis be used in combination with MMF and corticosteroids or other background immunosuppressive therapy, other than cyclophosphamide?	Yes – Document and go to #6	No – Criteria not met	
Is the drug prescribed by, or in consultation with, a rheumatologist, immunologist, nephrologist or kidney specialist?	Yes – Go to #7	No – Criteria not met	
7. Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations?	Yes – Approve up to 12 months	No – Criteria not met	
Renewal Criteria			



1.	Is there documentation of treatment success defined as an increase in eGFR, decrease in uPCR, or decrease in flares and corticosteroid use?	Yes – Go to #2	No – Criteria not met
2.	Is the requested dose within the Food and Drug Administration (FDA)-approved label and PacificSource quantity limitations?	Yes – Approve up to 12 months	No – Criteria not met

Quantity Limitations

• Lupkynis

- Starting dose: 23.7 mg twice daily (BID)
- Starting dose must be reduced in the below situations as follows:
 - eGFR 45 mL/min/1.73 m² or less at initiation: 15.8 mg BID
 - Mild-to-moderate hepatic impairment (Child-Pugh A or B): 15.8 mg BID
 - Concomitant use with moderate CYP3A4 inhibitors: 15.8 mg in morning and 7.9 mg in afternoon.



VORETIGENE NEPARVOVEC

Affected Medications: LUXTURNA (Voretigene neparvovec-rzyl intraocular suspension for subretinal injection)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design. Inherited Retinal Dystrophies (IRD) caused by mutations in the retinal pigment epithelium-specific protein 65kDa (RPE65) gene
Required Medical Information:	 Diagnosis of a confirmed biallelic RPE65 mutation-associated retinal dystrophy (e.g., Leber's congenital amaurosis [LCA], Retinitis pigmentosa [RP], Early Onset Severe Retinal Dystrophy [EOSRD], etc.) Genetic testing documenting biallelic mutations of the RPE65 gene Visual acuity of at least 20/800 OR have remaining light perception in the eye(s) receiving treatment Visual acuity of less than 20/60 OR a visual field of less than 20 degrees Presence of neural retina and a retinal thickness greater than 100 microns within the posterior pole as assessed by optical coherence tomography with AND have sufficient viable retinal cells as assessed by the treating physician
Appropriate Treatment Regimen & Other Criteria:	
Exclusion Criteria:	 Patient has been previously enrolled in clinical trials of gene therapy for retinal dystrophy RPE65 mutations or has been previously treated with gene therapy for retinal dystrophy in the eye(s) receiving treatment Patient has other pre-existing eye conditions or complicating systemic diseases that would eventually lead to irreversible vision loss and prevent the patient from receiving full benefit from treatment (e.g., severe diabetic retinopathy)
Age Restriction:	12 months of age and older
Prescriber/Site of Care Restrictions:	Ophthalmologist or retinal surgeon with experience providing sub-retinal injections
Coverage Duration:	Authorization: 1 month - 1 injection per eye per lifetime, unless otherwise specified



POLICY NAME: **VOSORITIDE**

Affected Medications: VOXZOGO (vosoritide)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 To increase linear growth in pediatric patients with achondroplasia with open
	epiphyses
Required Medical	Diagnosis of achondroplasia confirmed by molecular genetic testing showing a mutation
Information:	in the fibroblast growth factor receptor type 3 (FGFR3) gene
	Baseline height, growth velocity, and patient weight
Appropriate	Documentation of all the following:
Treatment	 Evaluation of epiphyses (growth plates) documenting they are open
Regimen & Other	 Growth velocity greater than or equal to 1.5 cm/yr
Criteria:	
	Reauthorization:
	Evaluation of epiphyses (growth plates) documenting they remain open
	Growth velocity greater than or equal to 1.5 cm/yr
Exclusion Criteria:	Hypochondroplasia
	Other short stature condition other than achondroplasia
	Evidence of growth plate closure
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a pediatric orthopedist, endocrinologist, or a
Care Restrictions:	provider with experience in treating skeletal dysplasia
	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 12 months, unless otherwise specified
_	Reauthorization: 12 months, unless otherwise specified



VYALEV

Affected Medications: VYALEV (carbidopa-levodopa infusion)

Covered Uses:	All Food and Drug Administration (FDA) approved indications not otherwise excluded by plan design Treatment of motor fluctuations in adults with advanced Parkinson's disease
	(PD)
Required Medical	Diagnosis of advanced PD
Information:	Clear response to levodopa treatment with evidence of "On" periods
	Persistent motor fluctuations with "Off" time occurring 2.5 hours or more per day while
	awake despite an optimized PD treatment regimen
Appropriate	Documented treatment failure with both of the following:
Treatment	 Oral carbidopa/levodopa extended release
Regimen & Other	 Two additional agents from different anti-PD drug classes:
Criteria:	 Monoamine oxidase-B (MAO-B) inhibitors (ex: selegiline, rasagiline) Dopamine agonists (ex: amantadine, pramipexole, ropinirole) Catechol-O-methyltransferase (COMT) inhibitors (ex: entacapone) Dosing is in accordance to FDA labeling and does not exceed 3,525 mg of foslevodopa component per day
	Reauthorization requires documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	Non-levodopa responsive PD
	Concomitant or recent (within 2 weeks) use of nonselective MAO inhibitors
	Concomitant use with carbidopa/levodopa extended release products
Age Restriction:	18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



XEOMIN, DYSPORT, MYOBLOC, and DAXXIFY

Affected Medications: XEOMIN (incobotulinum toxin A), DYSPORT (abobotulinumtoxinA), MYOBLOC (rimabotulinumtoxinB), JEUVEAU (prbotulinumtoxinA-xvfs), DAXXIFY (daxibotulinumtoxinA-lanm)

Covered Uses:	All Food and Drug Administration (FDA)-approved and compendia-supported indications	
	not otherwise excluded by plan design	
	o Dysport	
	 Focal dystonia (cervical dystonia, blepharospasm, laryngeal spasm, 	
	oromandibular dystonia, severe writer's cramp)	
	 Hemifacial spasm 	
	 Upper/lower limb spasticity 	
	○ Xeomin	
	 Cervical dystonia 	
	 Blepharospasm 	
	 Upper limb spasticity 	
	Chronic sialorrhea	
	Myobloc, Daxxify	
	Cervical dystonia	
Required Medical	Pertinent medical records and diagnostic testing	
Information:	Complete description of the site(s) of injection	
iniormation.	Strength and dosage of botulinum toxin used	
Appropriate	Dysport Dysport	
Treatment	Approved first-line for focal dystonia, hemifacial spasm, drug-induced orofacial	
Regimen & Other	dyskinesia, upper or lower limb spasticity	
Criteria:	, and a parameters, appearance, and a parameter, and a pa	
	<u>Xeomin</u>	
	Approved first-line for cervical dystonia, blepharospasm, upper limb spasticity, chronic	
	sialorrhea	
	Myobloc	
	Cervical dystonia: Documentation of treatment failure with Botox, Dysport, and Xeomin	
	Axillary hyperhidrosis: Documentation of treatment failure with Botox	
	Chronic sialorrhea: Documentation of treatment failure with glycopyrrolate oral tablets	
	<u>Daxxify</u>	
	Cervical dystonia: Documentation of treatment failure with Botox, Dysport, and Xeomin	
	Quantity limitations	
	Maximum of 4 treatments per 12 months	
	Reauthorization requires documentation of treatment success and a clinically significant	
	response to therapy	
Exclusion Criteria:	Cosmetic procedures (including glabellar lines, horizontal forehead lines, lateral canthal	
	lines)	
	Migraine headache use (Botox is preferred product)	



Age Restriction:	Myobloc, Daxxify: 18 years of age and older
Prescriber/Site of Care Restrictions:	 Blepharospasm: Prescribed by, or in consultation with, a neurologist, ophthalmologist, or optometrist Other indications: Prescribed by, or in consultation with, a neurologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



XGEVA

Affected Medications: XGEVA (denosumab)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design. One of these diagnoses: Giant cell tumor Bone metastases from solid tumors Hypercalcemia of malignancy Multiple myeloma NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher 	
Required Medical Information:	 7. Giant cell tumor Unresectable disease or surgical resection would likely result in severe morbidity. Bone metastases from solid tumors Hypercalcemia of malignancy Refractory to bisphosphonate therapy or contraindication Multiple myeloma Requires failure of zoledronic acid or pamidronate OR creatinine clearance less than 30 mL/min 	
Appropriate	Reauthorization requires documentation of treatment success and a clinically significant	
Treatment	response to therapy	
Regimen & Other Criteria:		
Exclusion Criteria:		
Age Restriction:	 Giant cell tumor: Adults and adolescents at least 12 years of age and skeletally mature weighing at least 45 kg All other indications: 18 years of age and older 	
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist	
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care	
Coverage Duration:	Authorization: 12 months, unless otherwise specified	



XIAFLEX

Affected Medications: XIAFLEX (collagenase clostridium histolyticum)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by	
	plan design	
	 Dupuytren's contracture with a palpable cord 	
	o Peyronie's disease	
Required Medical	Peyronie's disease:	
Information:	 Documented diagnosis of Peyronie's disease with a palpable plaque Curvature deformity is at least 30 degrees at the start of therapy Documentation of stable disease defined as symptoms that have remained unchanged for at least 3 months 	
Appropriate	Dupuytren's:	
Treatment Regimen & Other Criteria:	Authorization will be limited per joint as follows: One injection per month for a maximum of three injections per cord	
	Reauthorization will require documentation of treatment success and a clinically significant response to therapy	
	Peyronie's disease:	
	One treatment cycle consists of two Xiaflex injection procedures	
	Reauthorization for additional treatment cycles may be given if the curvature deformity is more than 15 degrees after the first, second or third treatment cycle, or if the prescribing healthcare provider determines that further treatment is clinically indicated	
	Maximum of 4 treatment cycles per plaque, administered at 6-week intervals	
Exclusion Criteria:	Peyronie's plaques that involve the penile urethra	
Age Restriction:		
Prescriber/Site of	Peyronie's: prescribed by, or in consultation with, a urologist	
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care	
Coverage Duration:	 Dupuytren's: 12 weeks, unless otherwise specified Peyronie's: 6 weeks, unless otherwise specified 	



XIFAXAN

Affected Medications: XIFAXAN (rifaximin)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	o Prevention of hepatic encephalopathy (HE)
	 Treatment of Travelers' Diarrhea caused by noninvasive strains of Escherichia coli (E. coli)
	 Treatment of Irritable Bowel Syndrome with Diarrhea (IBS-D)
	Compendia-supported uses that will be covered (if applicable)
	Treatment of HE
	Treatment of recurrent Clostridium difficile (C. diff)-associated diarrhea
	Treatment of Small Intestinal Bacterial Overgrowth (SIBO)
Required Medical	Documentation of complete & current treatment course required
Information:	Documentation of Complete & Current treatment course required Documentation of E-coli bacterial cultures for travelers' diarrhea
	Previous antibiotic history and documented allergies/hypersensitivity
Appropriate	Recurrent C. diff
Treatment	Documentation confirming a current diagnosis of recurrent C. diff infection (CDI) with
Regimen & Other	ALL of the following:
Criteria:	 CDI symptoms resolved on prior appropriate therapy and have reappeared
	within 8 weeks of completing prior therapy
	 Presence of at least 3 unformed stools in 24 hours
	Positive stool test for toxigenic Clostridium difficile
	Documented treatment failure with oral vancomycin
	Documented treatment failure with oral varicomyem
	<u>HE</u>
	Documented treatment failure with at least 1 month of lactulose therapy defined as
	continued altered mental status or elevated ammonium levels despite adequate upward
	titration
	Travelers' Diarrhea
	Documentation of ALL of the following:
	 Travelers' diarrhea is caused by noninvasive strains of E. coli
	 Systemic signs of infection (fever or blood in stool) are not present
	Member is returning from an area of high fluoroquinolone resistance On the standard description of the standard fluorogen in the standard fluo
	Documented treatment failure with a fluoroquinolone (e.g., ciprofloxacin, levofloxacin) and paithtenning.
	and azithromycin
	SIBO
	Documented diagnosis confirmed by a carbohydrate breath test
	Documented treatment failure with trial of at least one of the following antibiotics:
	amoxicillin/clavulanic acid, ciprofloxacin, metronidazole
	IBS-D
	Documentation confirming a Rome IV diagnosis with recurrent abdominal pain, on
	average, at least one day per week in the last 3 months, associated with two or more of
	the following:



	 Related to defecation Associated with a change in stool frequency Associated with a change in stool form (appearance) Symptom onset at least six months prior to diagnosis Documented treatment failure with ALL of the following: Loperamide Dicyclomine or hyoscyamine Tricyclic antidepressant (e.g., amitriptyline, nortriptyline) Retreatment criteria for IBS-D: Patient must have responded to the initial treatment for at least 4 weeks with either greater than or equal to 30% improvement from baseline in the weekly average abdominal pain score OR at least a 50% reduction in number of days in a week with a daily stool consistency of Bristol Stool Scale type 6 or 7 compared with baseline (6: fluffy pieces with ragged edges, a mushy stool; 7: watery stool, no solid pieces; entirely liquid). Retreatment can be approved when recurrence of symptoms (abdominal pain or mushy/watery stool consistency) occur for 3 weeks of a rolling 4-week period. Retreatment can be approved twice per lifetime.
	Reauthorization will require documentation of treatment success and a clinically significant response to therapy
Exclusion Criteria:	Recurrent C. diff Xifaxan exceeding 400 mg three times per day for 20 days HE Xifaxan exceeding the recommended dose of 550 mg twice daily or 400 mg 3 times daily
	for the treatment or prevention of hepatic encephalopathy Travelers' Diarrhea Xifaxan exceeding 200 mg three times per day for total of 3 days Diarrhea complicated by fever or bloody stool, or caused by bacteria other than noninvasive strains of E. coli
	SIBO • Xifaxan exceeding 550 mg three times per day for 14 days
	 IBS-D Mild cases of irritable bowel syndrome or diagnosis of irritable bowel syndrome with constipation Xifaxan exceeding 550 mg three times per day for 14 days
Age Restriction:	12 years of age and older
Prescriber/Site of Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Recurrent C. diff Authorization: 20 days, unless otherwise specified HE Authorization: 12 months, unless otherwise specified Travelers' Diarrhea Authorization: 7 days, unless otherwise specified
	SIBO AGT



 Authorization: 14 days, unless otherwise specified (one treatment per lifetime)
<u>IBS-D</u>
Authorization: 14 days, unless otherwise specified (maximum of 3 treatment courses per
lifetime)



POLICY NAME: **XURIDEN**

Affected Medications: XURIDEN (uridine triacetate)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design Hereditary orotic aciduria
Required Medical Information:	Diagnosis of hereditary orotic aciduria confirmed by ONE of the following: Molecular genetic testing confirming biallelic pathogenic mutation in the UMPS gene Urinary orotic acid level above the normal reference range Clinical manifestations consistent with disease such as: Megaloblastic anemia Leukopenia Developmental delays Failure to thrive
Appropriate	Reauthorization requires documentation of treatment success based on ONE of the
Treatment	following:
Regimen &	Improvement of hematologic abnormalities such as megaloblastic anemia and
Other Criteria:	leukopenia
	Reduction of urinary orotic acid levels
Exclusion Criteria:	
Age	
Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, a metabolic specialist or geneticist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Authorization: 12 months, unless otherwise specified



YONSA

Affected Medications: YONSA (abiraterone)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or higher
Required Medical Information:	Documentation of performance status, disease staging, all prior therapies used, and anticipated treatment course
Appropriate Treatment	Documented inadequate response or intolerable adverse event with the preferred product abiraterone acetate
Regimen & Other Criteria:	Reauthorization requires documentation of disease responsiveness to therapy
Exclusion Criteria:	 Child-Pugh Class C Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
Age Restriction:	18 years of age and older
Prescriber/Site of Care Restrictions:	 Prescribed by, or in consultation with, an oncologist All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	 Initial Authorization: 4 months, unless otherwise specified Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **ZANIDATAMAB**

Affected Medications: ZIIHERA (zanidatamab)

Covered Uses:	 All Food and Drug Administration (FDA)-approved indications not otherwise excluded by plan design
	NCCN (National Comprehensive Cancer Network) indications with evidence level of 2A or better
Required Medical Information:	Documentation of performance status, disease staging, all prior therapies used, and anticipated treatment course
	Documentation that Ziihera will be administered as monotherapy.
	Documentation of previously treated unresectable or metastatic human epidermal growth
	factor receptor 2 (HER2)-positive biliary tract cancer (BTC) that has progressed following at least 1 prior systemic therapy
	Documentation of HER2 positivity with a score of 3+ on immunohistochemistry (IHC) testing
Appropriate	Documented treatment failure or intolerable adverse event with Enhertu (fam-
Treatment	trastuzumab deruxtecan)
Regimen & Other Criteria:	Reauthorization requires documentation of disease responsiveness to therapy
Exclusion Criteria:	Karnofsky Performance Status 50% or less or ECOG performance score 3 or greater
Age Restriction:	
Prescriber/Site of	Prescribed by, or in consultation with, an oncologist.
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 4 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified



POLICY NAME: **ZILUCOPLAN**

Affected Medications: ZILBRYSQ (zilucoplan)

Covered Uses:	All Food and Drug Administration (FDA)-approved indications not otherwise excluded by
	plan design
	 Generalized myasthenia gravis (gMG) in adult patients who are anti- acetylcholine receptor (AChR) antibody positive
Required Medical	Diagnosis of generalized myasthenia gravis (gMG) confirmed by one of the following:
Information:	 A history of abnormal neuromuscular transmission test
	 A positive edrophonium chloride test
	 Improvement in gMG signs or symptoms with an acetylcholinesterase inhibitor
	Myasthenia Gravis Foundation of America (MGFA) Clinical Classification Class II to IV
	Positive serologic test for AChR antibodies
	 MG-Activities of Daily Living (MG-ADL) total score of 6 or greater OR Quantitative Myasthenia Gravis (QMG) total score of 12 or greater
Appropriate	Currently on a stable dose of at least one gMG therapy (acetylcholinesterase inhibitor,
Treatment	corticosteroid, or non-steroidal immunosuppressive therapy (NSIST)) that will be
Regimen & Other	continued during initial treatment with Zilbrysq
Criteria:	Documentation of one of the following: Transfer of the following:
	 Treatment failure with an adequate trial (one year or more) of at least two immunosuppressive therapies (azathioprine, mycophenolate, tacrolimus, cyclosporine, methotrexate)
	 Has required three or more courses of rescue therapy (plasmapheresis/plasma exchange and/or intravenous immunoglobulin), while on at least one immunosuppressive therapy, over the last 12 months
	Reauthorization:
	 Documentation of treatment success and clinically significant response to therapy defined as:
	 A minimum 2-point reduction in MG-ADL score from baseline AND Absent or reduced need for rescue therapy compared to baseline
	That the patient requires continuous treatment, after an initial beneficial response, due to
	new or worsening disease activity
Exclusion Criteria:	Current or recent systemic infection within 2 weeks
	Concurrent use with other biologics (rituximab, eculizumab, IVIG, etc)
Age Restriction:	18 years of age and older
Prescriber/Site of	Prescribed by, or in consultation with, a neurologist
Care Restrictions:	All approvals are subject to utilization of the most cost-effective site of care
Coverage Duration:	Initial Authorization: 4 months, unless otherwise specified
	Reauthorization: 12 months, unless otherwise specified