

An Independent Licensee of the Blue Cross Blue Shield Association **Prior Authorization Detail**Updated on 11/1/2024

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Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
ABOBOTULINUMTOXINA (DYSPORT)	1 - All FDA-approved Indications.			Diagnosis.			12 months	For reauthorization: documentation from prescriber indicating stabilization or improvement in condition.	0
ACITRETIN (SORIATANE)	1 - All FDA-approved Indications.			Diagnosis. Must have a trial of methotrexate or cyclosporine with inadequate response or significant side effect/toxicity or have a contraindication to these therapies.			12 months		0
ADALIMUMAB (HUMIRA)	3 - All Medically-accepted Indications.		Coverage is not provided for use of once weekly doses of Humira in combination with methotrexate.	Diagnosis. For rheumatoid arthritis (RA): history of trial and failure, contraindication, or intolerance to a 3 month trial with methotrexate or another DMARD. For juvenile idiopathic arthritis (JIA) with polyarthritis: history of trial and failure, contraindication, or intolerance to a 3 month trial with methotrexate, leflunomide, or sulfasalazine. For JIA with oligoarthritis, enthesitis and/or sacroillitis: history of trial and failure, contraindication, or intolerance to at least a 4 week trial of 2 different NSAIDS. For psoriatic arthritis (PsA) one of the following: 1.)members with axial or enthesitis must have a history of trial and failure, contraindication, or intolerance to a 4 week trial of 2 NSAIDS. 2.) the member has severe disease as defined by the prescriber. 3.) members with peripheral disease must have a history of a trial and failure, contraindication, or intolerance to a 12 week trial with methotrexate or another	Member must be 2 years of age or older.	By or in consultation with a rheumatologist, gastroenterologist, ophthalmologist, or dermatologist.	12 months	For hidradenitis suppurativa (HS): moderate to severe disease with 3 active abscesses, inflammatory nodules, or lesions with trial or an oral antibiotic with inadequate response or side effects/toxicities unless contraindicated. For uveitis: trial of a corticosteroid or immunomodulator with inadequate response or side effects/toxicities unless contraindicated. For reauth: must have documentation from prescriber indicating stabilization or improvement in condition.	

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ALIROCUMAB (PRALUENT)	1 - All FDA-approved			Diagnosis. Must have		By or in consultation with a	12 months	HoFH: must be confirmed by	0
	Indications.			confirmed diagnosis of		cardiologist, endocrinologist,		genetic testing with functional	
				heterozygous familial		or lipid specialist		mutation(s) in both LDL	
				hypercholesterolemia (see				receptor alleles or alleles	
				Other Criteria), homozygous				known to affect LDL receptor	
				familial hypercholesterolemia				functionality or have clinical	
				(HoFH, see Other criteria),				diagnosis defined as one of	
				clinical atherosclerotic				the following: untreated LDL	
				cardiovascular disease				greater than 500mg/dL or a	
				(ASCVD, see Other Criteria), or				treated LDL-C greater than	
				primary hyperlipidemia. Must				300mg/dL AND either	
				have baseline LDL-cholesterol				xanthoma before 10 years of	
				levels greater than or equal to				age or evidence of HeFH in	
				100 mg/dL (w/o ASCVD),				both parents. For ASCVD: must	
				70mg/dL (w/ ASCVD), or				have chart documentation	
				55mg/dl if has extreme risk				confirming history of at least	
				designation (see Other				one of the following:	
				Criteria). Must have failed to				myocardial infarction or other	
				achieve goal LDL-C reduction			1	acute coronary syndromes	
				after a trial of a high intensity			1	(including ST-elevation	
				statin (atorvastatin 40-80mg			1	myocardial infarction, non-ST	
				daily or rosuvastatin 20-40mg			1	elevation myocardial	
				daily) OR 2 moderate-intensity			1	infarction, and unstable	
				statins (atorvastatin or				angina), coronary or other	
				rosuvastatin) at the member's				revascularization procedure,	
				maximally tolerated dose OR				ischemic stroke or transient	
				documentation the member is				ischemic attack,	
				determined to be intolerant to				atherosclerotic peripheral	
				statin therapy with provider				arterial disease. For HeFH:	
				attestation of intolerance to				must have chart	
				statin therapy consisting of				documentation of one of the	
				statin related rhabdomyolysis				following: A score of greater	
				or skeletal-muscle related				than 8 using the Dutch Lipid	
ALOSETRON (LOTRONEX)	1 - All FDA-approved		Constipation. Concomitant use	Diagnosis. Documentation of	Coverage is provided for	By or in consultation with a	12 months	For reauth: must have	0
ALOSE TROM (LOTRONEX)	Indications.		of fluvoxamine. Male gender.	chronic IBS symptoms diarrhea		Gastroenterologist	12 11011(13	documentation from	0
	indications.		History of chronic or severe	lasting at least 6 months.	older.	dastroenterologist		prescriber indicating	
			constipation or sequelae from	Gastrointestinal tract	older.			stabilization or improvement	
			constipation, intestinal	abnormalities have been ruled				in condition.	
								in condition.	
			obstruction, stricture, toxic	out. Must have trial of					
			megacolon, gastrointestinal	loperamide and dicyclomine					
			perforation and/or adhesions,	used in the treatment of IBS-D					
			ischemic colitis, impaired	with inadequate response or					
			intestinal circulation,	significant side effects/toxicity					
			thrombophlebitis, or	unless contraindicated					
			hypercoagulable state, Crohn's						
			disease, ulcerative colitis,						
			diverticulitis, or severe hepatic						
			impairment.				1	1	
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							1	1	
VI DELICID (VIIO)CE)	1 All EDA and and a	+		Diagnosis of DIV2CA D-1-4	Causage to age the differ	December of the second section of the second section of the second section sec	12 manths	Fau annuth autor the control to	0
ALPELISIB (VIJOICE)	1 - All FDA-approved			Diagnosis of PIK3CA-Related	Coverage is provided for	By or in consultation with an	12 months	For reauthorization: must have	U
	Indications.			Overgrowth Spectrum (PROS)	members 2 years of age or	appropriate specialist	1	documentation from	
				confirmed by genetic testing.	older.	depending on the symptoms	1	prescriber indicating	
				Disease must be severe or life-		and part of the body that are	1	stabilization or improvement	
				threatening and require		affected.	1	in condition.	
				systemic treatment.					
				systemic treatment.					

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ALPHA-1 PROTEINASE INHIBITOR (PROLASTIN)	1 - All FDA-approved Indications.		Immunoglobulin A (IgA) deficient members with antibodies against IgA	Diagnosis. Member must have pre-treatment serum levels of alpha-1 antitrypsin (AAT) that are less than 11 micromoles per liter (80 milligrams per deciliter if measured by radial immunodiffusion or 57 milligrams per deciliter if measure by nephelometry) consistent with phenotypes PiZZ, PiZ (null) or Pi (null, null) of AAT. Member must have symptomatic emphysema confirmed with pulmonary function testing.	Coverage is provided for members 18 years of age and older.	By or in consultation with a pulmonologist	Initial: 6 months Reauthorization: 12 months	For reauth: documentation of improvement or stabilization of the signs and symptoms of emphysema associated with alpha-1 antitrypsin deficiency including slowed progression of emphysema as evidenced by annual spirometry testing or a decrease in frequency, duration or severity of pulmonary exacerbations	0
ALPHA-1 PROTEINASE INHIBITOR (ZEMAIRA)	1 - All FDA-approved Indications.		Immunoglobulin A (IgA) deficient members with antibodies against IgA	Diagnosis. Member must have pre-treatment serum levels of alpha-1 antitrypsin (AAT) that are less than 11 micromoles per liter (80 milligrams per deciliter if measured by radial immunodiffusion or 57 milligrams per deciliter if measure by nephelometry) consistent with phenotypes PiZZ, PiZ (null) or Pi (null, null) of AAT. Member must have symptomatic emphysema confirmed with pulmonary function testing.	Coverage is provided for members 18 years of age and older.	By or in consultation with a pulmonologist	Initial: 6 months, Reauthorization: 12 months	For reauth: documentation of improvement or stabilization of the signs and symptoms of emphysema associated with alpha-1 antitrypsin deficiency including slowed progression of emphysema as evidenced by annual spirometry testing or a decrease in frequency, duration or severity of pulmonary exacerbations	0
AMBRISENTAN (LETAIRIS)	1 - All FDA-approved Indications.		Pregnancy	Diagnosis. Pulmonary arterial hypertension (PAH) WHO Group I confirmed by chart documentation of right-heart catheterization (RHC) indicating a mean pulmonary arterial pressure greater than 20 mmHg, pulmonary vascular resistance greater than 2 wood units, and mean pulmonary capillary wedge pressure less than or equal to 15 mmHg. If provider indicates RHC is not recommended, must have documentation of an echocardiography.		Prescribed by or in consultation with cardiologist or pulmonologist.	Initial authorization: 3 months Reauthorization: 12 months	For reauth: documentation from prescriber that demonstrates member is tolerating and receiving clinical benefit from treatment	0

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AMOXAPINE (ASENDIN)	1 - All FDA-approved Indications.			Diagnosis. Requests for enrollees aged 65 or older	Prior authorization only applies to enrollees aged 65 or		12 months		0
	indications.			must document intolerance to					
				or clinical failure of 2 of the	age 65 are not subject to prior				
				following for depression: a	authorization.				
				SSRI (except paroxetine), SNRI,	authorization.				
				mirtazapine, or bupropion					
				init tazapine, or bupropion					
APREMILAST (OTEZLA)	1 - All FDA-approved			Diagnosis. For psoriatic	Plaque psoriasis: age 6 and	By or in consultation with a	12 months	For reauthorization: must have	0
	Indications.			arthritis (PsA), one of the	older, All other diagnoses: 18	dermatologist, rheumatologist		documentation from	
				following: 1) members with	years of age or older.			prescriber indicating	
				axial or enthesitis must have a				stabilization or improvement	
				history of trial and failure,				in condition.	
				contraindication, or					
				intolerance to a 4 week trial of					
				2 NSAIDs, 2) the member has					
				severe disease as defined by					
				the prescriber, 3) members					
				with peripheral disease must					
				have a history of a trial and					
				failure, contraindication, or intolerance to a 12 week trial					
				with methotrexate or another					
				DMARD. For plaque psoriasis:					
				minimum BSA involvement of					
				at least 3% (not required if on					
				palms, soles, head/neck,					
				genitalia), a history of trial and					
				failure of ONE of the following:					
				1) topical therapy (e.g.					
				corticosteroid, calcineurin					
				inhibitor, vitamin D analog), 2)					
				phototherapy, 3) systemic					
				treatment (e.g. methotrexate,					
				cyclosporine, oral retinoids).					
				For Behcet's disease: must					
				have recurrent oral ulceration					
				(at least 3 times within the					
				past year) plus 2 of the					
				following symptoms: recurrent					
				genital ulceration, eye lesions,					
ARIPIPRAZOLE INJECTION	1 - All FDA-approved			Diagnosis. Documentation of	Coverage is provided for		12 months		0
(ABILIFY MAINTENA)	Indications.			prior trial and failure of oral	members 18 years of age and				
				aripiprazole (Abilify) therapy.	older.				
									-
	1 - All FDA-approved			Diagnosis. Documentation the	Coverage is provided for		12 months		0
SENSOR (ABILIFY MYCITE)	Indications.			member had at least a one-	members 18 years of age and				
				month trial of oral aripiprazole	older.				
				(Abilify) therapy.					
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ARMODAFINIL (NUVIGIL) 1	1 - All FDA-approved	Off-Label Uses	Ens. asion criteria	Diagnosis. Must have a history	- Se vestiletion	By or in consultation with a		For reauth: documentation of	0
	Indications.			of trial and failure,		sleep specialist, ENT (ear,	OSA: 12 months	improvement or stabilization.	Ü
ı l'	aicacions.			contraindication, or		nose, and throat specialist),	05/11/12 1110111115	improvement or stabilization.	
ı				intolerance to modafinil. For		neurologist, or pulmonologist			
ı				narcolepsy: Sleep Study (e.g.		neurologist, or pulmonologist			
ı				Polysomnogram, Multiple					
ı				Sleep Latency Test) confirming					
ı				diagnosis. For obstructive					
ı				•					
				sleep apnea: Sleep study (e.g.					
				polysomnogram) confirming					
				diagnosis. For shift work sleep					
				disorder (SWSD): must meet					
ı				International Classification of					
				Sleep Disorders criteria for					
ı				SWSD (either primary					
ı				complaint of excessive					
				sleepiness or insomnia					
,		1		temporarily associated with					
,		1		work period that occurs during					
				habitual sleep phase OR					
		1		polysomnography and					
,		1		Multiple Sleep Latency Test					
ı				demonstrate loss of normal					
ı				sleep wake pattern, no other					
ı				medical or mental disorders					
ı				account for symptoms, and					
ı				symptoms do not meet criteria					
ı				for any other sleep disorder					
ı				producing insomnia or					
ı				excessive sleepiness such as					
ı				time zone change syndrome)					
ı				and must provide					
ı				documentation of shift work					
				cchadula chawing E or mara					
	1 - All FDA-approved			Diagnosis. Documentation of	Coverage provided for		12 months		0
, I'	Indications.			trial and failure of at least two	members age 10 years and				
ı					older.				
ı				atypical antipsychotics:					
ı				olanzapine, quetiapine,					
ı				clozapine, paliperidone,					
ı				risperidone, aripiprazole, or					
ı				ziprasidone.					
,		1							
		1							
	1 - All FDA-approved	1		Diagnosis. Requests for	Prior authorization only		12 months		0
(LOMOTIL)	Indications.	1		enrollees aged 65 or older	applies to enrollees aged 65 or				
,		1			older. All enrollees less than				
,		1		or clinical failure of	age 65 are not subject to prior				
		1		loperamide.	authorization.				
AVACODAN (TAVANCOC)	1 All FDA against d			Diamenia of ANCA	Coverage is not did at for	December of the control of the contr	12 Mantha	For required	0
	1 - All FDA-approved	1		Diagnosis of ANCA-associated		By or in consultation with a	12 Months	For reauthorization:	U
, l'	Indications.	1			members 18 years of age or	rheumatologist, hematologist		documentation from	
,		1		be on concurrent therapy with	older.	or oncologist.		prescriber indicating	
,		1		glucocorticoids and				stabilization or improvement	
,		1		immunosuppressants (e.g.				in condition.	
,		1		cyclophosphamide,					
,		1		azathioprine, mycophenolate,					
		1		rituximab).					
,		1							
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AVATROMBOPAG (DOPTELET)	1 - All FDA-approved Indications.			Diagnosis. For ITP, documentation of inadequate response to corticosteroids or immunoglobulins and documentation of a platelet count less than or equal to 30,000/microliter. For thrombocytopenia in adult patients with chronic liver disease who are scheduled to undergo a procedure, documentation of a platelet count less than 50,000/microliter.		By or in consultation with a hematologist, oncologist, hepatologist, or surgeon	Chronic ITP: 6 months. Thrombocytopenia in patients with chronic liver disease: 1 month	For reauth of chronic ITP: documentation of improvement in platelet count from baseline.	
BECAPLERMIN (REGRANEX)	1 - All FDA-approved Indications.		Neoplasm at application site. Treatment of pressure ulcers and venous stasis ulcers. Use on exposed joints, tendons, ligaments, and bone.	Diagnosis. Must have a lower extremity diabetic neuropathic ulcer that extends into the subcutaneous tissue or beyond and have an adequate blood supply. Must be used as adjunctive therapy to good ulcer care practices (i.e. debridement, infection control, pressure relief).			3 months	For reauth: documentation of improvement or stabilization.	0
BEDAQUILINE (SIRTURO)	1 - All FDA-approved Indications.			Diagnosis. Must have either inadequate response to a first-line tuberculosis (TB) regimen containing isoniazid and rifampin OR chart documentation of resistance to isoniazid and rifampin per susceptibility testing. Must weigh at least 15 kg. Must be used in combination with at least 3 other drugs indicated for the treatment of TB.	Member must be 5 years of age or older.	By or in consultation with a pulmonologist or infectious disease specialist	6 months		0

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BELIMUMAB (BENLYSTA) (IV FORMULATION)	1 - All FDA-approved Indications.		Severe active central nervous system lupus. Combination therapy with other biologics or IV cyclophosphamide.	Diagnosis of active, autoantibody-positive, systemic lupus erythematosus (SLE) or lupus nephritis. Must have ANA of at least 1:80 or anti-dsDNA of at least 3:80 or anti-dsDNA or support being autoantibody positive. Must be currently taking or has tried and failed or had an intolerance or contraindication to at least one standard therapy for systemic lupus erythematosus (e.g. corticosteroids, antimalarials, NSAIDS, or immunosuppressives) or lupus nephritis (e.g. corticosteroids, mycophenolate, cyclophosphamide, azathioprine). Diagnosis of active lupus nephritis. Documentation of a biopsyproved lupus nephritis Class III, IV or V.	Coverage is provided for members 5 years of age and older	By or in consultation with a rheumatologist or hematologist	12 months	For reauth: documentation from the prescriber indicating stabilization or improvement in condition.	0
BELIMUMAB (BENLYSTA) (SQ)	1 - All FDA-approved Indications.		Severe active central nervous system lupus. Combination therapy with other biologics or IV cyclophosphamide.	autoantibody-positive,	Coverage is provided for members 18 years of age and older.	By or in consultation with a rheumatologist, hematologist, or nephrologist	12 months	For reauth: documentation from the prescriber indicating stabilization or improvement in condition.	0

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ELUMOSUDIL (REZUROCK)	3 - All Medically-accepted Indications.			Diagnosis. For a diagnosis of chronic Graft versus host disease (GVHD), after a trial and failure of at least two prior lines of systemic therapy.	GVHD: age 12 years or older	By or in consultation with an oncologist, hematologist, or transplant specialist	12 months	For reauth: documentation of improvement or stabilization.	0
IENRALIZUMAB (FASENRA)	1 - All FDA-approved Indications.			Diagnosis. For severe eosinophilic asthma: eosinophilic asthma: eosinophil blood count greater than or equal to 150cells/microliter within the past 4 weeks. Documentation of inadequate response, intolerance, or contraindication to a high-dose ICS in combination with a LABA. Meets one of the following within the past year: one or more acute asthmarelated ED visit(s), one or more acute inpatient visits where asthma was the principal diagnosis, or two or more acute asthma exacerbations requiring oral systemic steroids.	Coverage is provided for members 6 years of age or older.	By or in consultation with an allergist, immunologist, or pulmonologist.	12 months	For reauth: documentation of improvement (e.g. reduced symptoms, reduced exacerbations, need for oral steroids).	0
EREMAGENE GEPERPAVEC VYJUVEK)	1 - All FDA-approved Indications.			Diagnosis of Dystrophic Epidemolysis Bullosa (DEB) with a mutation in the collagen type VII alpha 1 chain (COL7A1) gene confirmed by genetic testing. Must have a wound with no evidence or history of squamous-cell carcinoma or active infection.	Coverage is provided for members 6 months of age or older.	By or in consultation with a dermatologist	6 months	Reauthorization: must have documentation from prescriber indicating improvement in condition.	0
IIRCH TRITERPENES FILSUVEZ)	1 - All FDA-approved Indications.		0	Diagnosis of Dystrophic Epidemolysis Bullosa (DEB) or junctional epidermolysis bullosa (JEB) with an open wound.	Coverage is provided for members 6 months of age or older.	By or in consultation with a dermatologist.	6 months	Reauthorization: must have documentation from prescriber indicating improvement in condition.	0

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BOSENTAN (TRACLEER)	1 - All FDA-approved Indications.	On-Laure Oses	Pregnancy	Diagnosis. Pulmonary arterial hypertension (PAH) WHO Group I confirmed by chart documentation of right-heart catheterization (RHC) indicating a mean pulmonary arterial pressure greater than 20 mmHg, pulmonary vascular resistance greater than 2 wood units, and mean pulmonary capillary wedge pressure less than or equal to 15 mmHg. If provider indicates RHC is not recommended, must have documentation of an echocardiography.	Age nestriction	Prescribed by or in consultation with cardiologist or pulmonologist.	Reauthorization: 3 months Reauthorization: 12 months	For reauth: documentation from prescriber that demonstrates member is tolerating and receiving clinical benefit from treatment	0
BRIVARACETAM (BRIVIACT)	1 - All FDA-approved Indications.			Diagnosis. Must have had an inadequate response or intolerance to generic levetiracetam and at least one of the following generic anticonvulsant drugs: phenytoin, carbamazepine, oxcarbazepine, gabapentin, lamotrigine, valproate, or topiramate.	Coverage is provided for members 1 month of age and older	By or in consultation with a neurologist	12 months		0
BUDESONIDE (EOHILIA)	1 - All FDA-approved Indications.		0	Diagnosis. For eosinophilic esophagitis (EoE): must have at least 15 intraepithelial eosinophils per high-power field (eos/hpf) following a treatment course with a PPI.	Coverage is provided for members 11 years of age or older.	By or in consultation with an allergist or gastroenterologist.	3 months	Reauth: use beyond 3 months has not been studied.	0
BUDESONIDE EXTENDED RELEASE TABLETS (UCERIS)	1 - All FDA-approved Indications.			Diagnosis. Must have a trial and failure, a contraindication, or an intolerance to two (2) of the following therapy options: topical mesalamine, oral aminosalicylate or corticosteroids with inadequate response or side effects/toxicity unless contraindicated.	Member must be 18 years of age or older.	By or in consultation with a rheumatologist or gastroenterologist.	8 weeks	For reauth: must have documentation from prescriber indicating stabilization or improvement in condition.	0

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BUROSUMAB-TWZA (CRYSVITA)	1 - All FDA-approved Indications.		Use with oral phosphate or active vitamin D analogs	Diagnosis. For X-linked hypophosphatemia: confirmation of the diagnosis by at least one of the following: A genetic test showing a PHEX gene mutation (phosphate regulating gene with homology to endopeptidase on the X chromosome) or Serum fibroblast growth factor 23 (FGF23) level greater than 30 pg/mL. Documentation of a baseline fasting serum phosphorus concentration that is below the reference range for the members age (reference range must be provided). For FGF23-related hypophosphatemia in tumor-induced osteomalacia (TIO): documentation the member has a phosphaturic mesenchymal tumor that cannot be resected or localized. Documentation of a baseline fasting serum phosphorus concentration that is below the reference range for the members age (reference range must be provided).		By or in consultation with a physician who is experienced in the management of patients with metabolic bone disease.	12 months	Reauthorization: Documentation current (within the past 12 months) serum phosphorus level is not above the upper limit of the laboratory normal reference range and documentation the member has had a positive clinical response or stabilization in their disease.	0
BUT/APAP/CAF TAB	1 - All FDA-approved Indications.			Diagnosis. This Prior Authorization requirement only applies to members when a non-FDA approved diagnosis is submitted. FDA-approved diagnosis codes submitted will pay without prior authorization requirement.	Coverage is provided for members 12 years of age or older.		12 months		0
BUTAL/APAP TAB 50-325MG	1 - All FDA-approved Indications.			Diagnosis. This Prior Authorization requirement only applies to members when a non-FDA approved diagnosis is submitted. FDA-approved diagnosis codes submitted will pay without prior authorization requirement.			12 months		O

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C1 ESTERASE INHIBITOR ((HAEGARDA)	1 - All FDA-approved Indications.	Ori-Lauer Oses	LACIDIUM UNEFID	Diagnosis of HAE is confirmed	Coverage is provided for members 6 years of age or older.	Prescriber Mestriction Prescribed by or in consultation with an allergist/immunologist, hematologist, dermatologist	Initial: 6 months Reauthorization: 12 months	For reauth: must have documentation from prescriber indicating improvement in condition.	0
CANNABIDIOL (EPIDIOLEX)	1 - All FDA-approved Indications.			Diagnosis. Must have had an inadequate response or intolerance to one generic antiepileptic drug.	Member must be 1 year of age or older	By or in consultation with a neurologist	12 months		0
CARGLUMIC ACID (CARBAGLU)	1 - All FDA-approved Indications.			Diagnosis. This Prior Authorization requirement only applies to members when a non-FDA approved diagnosis is submitted at the point of sale. FDA-approved diagnosis codes submitted will pay without prior authorization requirement.			12 months		0
CEFTAROLINE (TEFLARO)	1 - All FDA-approved Indications.			Diagnosis. For acute bacterial skin and skin structure infection (ABSSSI), documentation of a history of treatment failure with or contraindication to vancomycin.			14 days		0

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CLOMIPRAMINE	1 - All FDA-approved Indications.			Diagnosis. Requests for enrollees aged 65 or older must document intolerance to or clinical failure of 2 of the following: fluoxetine, fluvoxamine, and sertraline.	Prior authorization only applies to enrollees aged 65 or older. All enrollees less than age 65 are not subject to prior authorization.		12 months		0
CYPROHEPTADINE	1 - All FDA-approved Indications.			Diagnosis. Requests for enrollees aged 65 or older must document contraindication to, intolerance to or clinical failure of cetirizine and levocetirizine for use as an antihistamine.	Prior authorization only applies to enrollees aged 65 or older. All enrollees less than age 65 are not subject to prior authorization.		12 months	For reauthorization: documentation from prescriber indicating stabilization or improvement in condition.	0
CYSTEAMINE (CYSTAGON)	1 - All FDA-approved Indications.			Diagnosis. Must have documentation of CTNS gene mutation, elevated white blood cell cystine levels greater than 2nmol per half-cystine per mg of protein, or cystine corneal crystals by slit lamp examination.		By or in consultation with a nephrologist or physician who specializes in the treatment of inherited metabolic disorders	Initial: 3 months Reauthorization: 12 months	For reauth: must have documentation from prescriber indicating improvement in condition and a reduction in WBC cystine levels since starting treatment with oral cysteamine	0
DALFAMPRIDINE (AMPYRA)	1 - All FDA-approved Indications.		History of seizure disorder, moderate to severe renal impairment (CrCl less than or equal to 50 mL/min).	Diagnosis of multiple sclerosis. Chart documentation of baseline motor disability or dysfunction.	Coverage is provided for members 18 years of age or older.	Neurologist	Initial: 3 months Reauthorization: 12 months	For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	0
DARBEPOETIN ALFA (ARANESP)	1 - All FDA-approved Indications.		Uncontrolled hypertension	Diagnosis. Must have Hgb level less than 10 g/dL.			6 months	For reauth for CKD on dialysis: must have a Hgb less than or equal to 11g/dl. For reauth for CKD not on dialysis: must have Hgb less than or equal to 10 g/dl. Reauth for pediatric members with CKD: must have a Hgb less than or equal to 12 g/dl. Reauth for all other dx must meet initial criteria.	
DEFERIPRONE (FERRIPROX)	1 - All FDA-approved Indications.			Diagnosis. Must have documentation of a trial and failure of Exjade (this requires a PA) unless contraindicated .		Prescribed by or in consultation with a hematologist	12 months	For reauth: documentation from prescriber indicating stabilization or improvement in condition.	0
DENOSUMAB (XGEVA)	3 - All Medically-accepted Indications.			Diagnosis.		Prescribed by or in consultation with a hematologist or oncologist	6 months		0

Group DESFERASIROX (EXJADE)	Indication Indicator 1 - All FDA-approved Indications.	Off-Label Uses	Exclusion Criteria Glomerular Filtration Rate less than 40mL/min/1.73 m2.	Required Medical Information Diagnosis. For chronic iron overload due to blood	Age Restriction	Prescriber Restriction Prescribed by or in consultation with a	Coverage Duration 12 months	Other Criteria For reauth: documentation from prescriber indicating	Part B Prerequisite 0
	maications.		Concomitant advanced malignancy or high risk	transfusions: pretreatment serum ferritin level is greater		hematologist		stabilization or improvement in condition.	
			myelodysplastic syndrome. Platelet count less than	than 1000 mcg/L. For chronic iron overload due to non-					
			50000000000/L	transfusion-dependent thalassemia (NTDT)					
				syndromes: pretreatment serum ferritin level is greater					
				than 300 mcg/L and a liver iron concentration of at least					
				5mg iron per gram dry weight.					
DESIPRAMINE (NORPRAMIN)	1 - All FDA-approved			Diagnosis. Requests for	Prior authorization only		12 months		0
	Indications.			enrollees aged 65 or older must document intolerance to					
				or clinical failure of 2 alternatives such as an SSRI	age 65 are not subject to prior authorization.				
				(except paroxetine), SNRI, bupropion, trazodone or mirtazapine for depression.					
				Thirt dzapine for depression.					
DEUTETRABENAZINE AUSTEDO)	1 - All FDA-approved Indications.		Uncontrolled depression, actively suicidal, hepatic	Diagnosis. For chorea: must have confirmed Huntington's	Coverage is provided for members 18 years of age or	By or in consultation with a neurologist or psychiatrist	12 months	For reauthorization: must have documentation from	0
			impairment, concurrent use with MAOI's, reserpine,	disease either by Huntington Disease Mutation analysis	older.			prescriber indicating stabilization or improvement	
			tetrabenazine, or valbenazine.	indicating expanded CAG				in condition.	
				repeat of greater than or equal to 36 in the Huntington					
				gene) or a positive family history of Huntington's					
				Disease with autosomal dominant inheritance pattern,					
				must have clinical signs of Huntington's Disease including chart documentation of a					
				clinical work-up showing one or more of the following signs:					
				motor (e.g. finger tapping, rigidity), oculomotor, bulbar					
				(e.g. dysarthria, dysphagia), affective (e.g. depression),					
				cognitive. Must have chart documentation of chorea. For					
				tardive dyskinesia (TD): must have moderate to severe TD					
				according to the DSM V criteria including involuntary					
				athetoid or choreiform movements and has a history					
				of treatment with neuroleptic agent (i.e. antipsychotic).					
				Adjustments to possible offending medication such as					
		1	<u> </u>	doco reduction or					

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
GFOUP DEXTROMETHORPHAN- QUINIDINE (NUEDEXTA)	1 - All FDA-approved Indications.	UTT-Label Uses	exclusion Criteria	Diagnosis. Pseudobulbar affect (PBA): documentation	-	By or in consultation with neurologist	Initial: 3 months Reauthorization: 12 months	For reauthorization: Documentation indicating a decrease in the number of laughing and/or crying episodes since starting the medication.	0
DEXTROMETHORPHAN/BUPR OPION (AUVELITY)	1 - All FDA-approved Indications.			Diagnosis. Documentation of trial and failure of at least two generic antidepressants alternatives such as an SSRI, SNRI, bupropion, trazodone or mirtazapine.	Coverage is provided for members 18 years of age or older.		12 months		0
DICLOFENAC TOPICAL GEL (SOLARAZE)	1 - All FDA-approved Indications.			Diagnosis.			90 days	For reauthorization: documentation from prescriber indicating stabilization or improvement in condition.	0
DICYCLOMINE (BENTYL)	1 - All FDA-approved Indications.				Prior authorization only applies to enrollees aged 65 or older. All enrollees less than age 65 are not subject to prior authorization.		12 months	For reauthorization: Prescriber must acknowledge that medication benefits outweigh potential risks in the member 65 years of age or older.	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information		Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
DIHYDROERGOTAMINE NASAL SPRAY (MIGRANAL)	1 - All FDA-approved Indications.		Members with hemiplegic or basilar migraine, ischemic heart disease (angina pectoris, history of MI, or documented silent ischemia) or who have clinical symptoms or findings consistent with coronary artery vasospasm (including Prinzmetal's variant angina or uncontrolled hypertension).	Diagnosis. Documentation of trial and failure of 1 medication from each of the following classes: a NSAID and a triptan unless contraindicated.	Coverage is provided for members 18 years of age and older.		12 months	For reauth: documentation from prescriber indicating stabilization or improvement in condition.	0
DIROXIMEL FUMARATE (VUMERITY)	1 - All FDA-approved Indications.		Coverage is not provided with coadministration with dimethyl fumarate or diroximel fumarate.	Diagnosis. Member must have relapsing Multiple Sclerosis (MS), including clinically isolated syndrome, relapsing-remitting disease, and active secondary progressive disease. Documentation that functional status is preserved and patient is either still able to walk at least a few steps or alternatively must have some functional arm/hand use consistent with performing activities of daily living.	Coverage is provided for members 18 years of age or older.	By or in consultation with a neurologist	12 months	For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	0
DORNASE ALFA (PULMOZYME)	1 - All FDA-approved Indications.			Diagnosis.		By or in consultation with a pulmonologist or cystic fibrosis specialist	12 months	For reauth: must have documentation from prescriber indicating stabilization or improvement in condition.	0
DRONABINOL	1 - All FDA-approved Indications.			Diagnosis. Nausea and vomiting associated with cancer chemotherapy: must have trial of two conventional antiemetic treatments (e.g., ondansetron, aprepitant, metoclopramide, dexamethasone, prochlorperazine) with inadequate response or significant side effects/toxicity unless contraindicated.			12 months		0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
DROXIDOPA (NORTHERA)	1 - All FDA-approved Indications.			Diagnosis. Documentation of a clinical diagnosis of symptomatic neurogenic orthostatic hypotension caused by one of the following: Primary autonomic failure (Parkinson's disease, multiple system atrophy, or pure autonomic failure), dopamine beta-hydroxylase deficiency or non-diabetic autonomic neuropathy. Must have a trial of midodrine with inadequate response or significant side effects/toxicity unless contraindicated.	Coverage is provided for members 18 years of age and older.		2 weeks	For reauth: rationale from the provider for continuing therapy beyond 2 weeks	
DUPILUMAB (DUPIXENT)	1 - All FDA-approved Indications.			Diagnosis. For asthma: must have either moderate to severe eosinophilic phenotype with an eosinophili count greater than or equal to 150 cells/microliter or oral corticosteroid dependent persistent asthma (chronic oral corticosteroid use). Documentation of recent use and failure to respond to inhaled steroid in combo with long acting beta agonist. Must have asthma symptoms that are inadequately controlled while on treatment (uncontrolled defined as having an asthma exacerbation requiring hospitalization in the past year, having 2 or more asthma exacerbations requiring oral systemic steroids, or inability to taper off daily corticosteroids). For atopic dermatitis: history of trial and failure, contraindication, or intolerance to a topical corticosteroid or topical calcineurin inhibitor. For nasal polyps: history or trial and failure of Xhance (fluticasone propionate). Must be used as	For atopic dermatitis: 6 months or older. For asthma: 6 years or older. For eosinophilic esophagitis: 1 year or older. For all other indications: 18 years or older.		12 months	Reauth for asthma: documentation of improvement (e.g. reduced symptoms, reduced exacerbations, need for oral steroids). Reauth for all other indications: documentation from prescriber indicating stabilization or improvement in condition.	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
EDARAVONE (RADICAVA ORS)	1 - All FDA-approved Indications.			Diagnosis of Amyotrophic Lateral Sclerosis (ALS). Must have normal respiratory function (defined as a forced vital capacity (FVC) of at least 80%), must be able to perform activities of daily living (ADLs) such as eating and moving around independently, must provide a recent ALSFRS-R score.	Coverage is provided for members 18 years of age and older	By or in consultation with a neurologist	12 months	Reauth: must provide documentation of clinical benefit based on the prescriber's assessment and an ALSFRS-R score within the past 12 months	0
ELEXACAFTOR/TEZACAFTOR/I VACAFTOR (TRIKAFTA)	1 - All FDA-approved Indications.			Diagnosis. Documentation of genetic test confirming the member has at least one F508del mutation in the CFTR gene or a mutation in the CFTR gene that is responsive based on in vitro data.	older	By or in consultation with a cystic fibrosis specialist or pulmonologist	12 months	For reauthorization: documentation from prescriber indicating stabilization or improvement in condition.	0
ELTROMBOPAG (PROMACTA)	1 - All FDA-approved Indications.			Diagnosis. For ITP, documentation of inadequate response to corticosteroids or immunoglobulins and documentation of a platelet count less than or equal to 30,000/microliter. For chronic hepatitis C, documentation that thrombocytopenia prevents the initiation of interferon-based therapy or limits the ability to maintain interferon-based therapy, and documentation of a platelet count less than 75,000/microliter. For severe aplastic anemia, documentation of a platelet count less than 30,000/microliter and one of the following: the member has had an insufficient response to immunosuppressive therapy or the members will be using the medication in combination with immunosuppressive therapy.		By or in consultation with a hematologist, oncologist, gastroenterologist, or hepatologist	6 months	For reauth: for all dx documentation of improvement in platelet count from baseline. For hepatitis C: documentation the member is still on antiviral therapy.	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
ENTECAVIR (BARACLUDE)	1 - All FDA-approved Indications.			Diagnosis. Member must have chronic hepatitis B virus (HBV) infection with evidence of active viral replication and either evidence of persistent elevations in serum aminotransferases (ALT or AST) or histologically active disease.	Member must be 2 years of age or older	By or in consultation with a gastroenterologist, hepatologist, infectious disease, HIV or transplant specialist.	12 months	For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	
EPOETIN ALFA-EPBX (RETACRIT)	3 - All Medically-accepted Indications.		Uncontrolled hypertension	Diagnosis. For Reduction of Allogeneic Red Blood Cell Transfusions in Members Undergoing Elective, Noncardiac, Nonvascular Surgery: must have hemoglobin (Hgb) greater than 10 and less than or equal to 13 g/dL, be at high risk for perioperative blood loss from surgery, and documentation that erythropoietin therapy will be used to decrease the need for transfusions associated with surgery in members unwilling or unable to undergo autologous blood donation prior to surgery. All other dx must have Hgb level less than 10 g/dL.			6 months	For reauth for CKD on dialysis: must have a Hgb less than or equal to 11g/dl. For reauth for CKD not on dialysis: must have Hgb less than or equal to 10 g/dl. For reauth for zidovudine treated members and pediatric members with CKD: must have a Hgb less than or equal to 12 g/dl. Reauth for all other dx must meet initial criteria.	0
ERENUMAB-AOOE (AIMOVIG)	1 - All FDA-approved Indications.			Diagnosis. For episodic migraine: Documentation the member has 4 to 14 headache days per month. For chronic migraine: Documentation the member has at least 15 headache days per month for 3 or more months with at least 8 migraine days per month. For both: Must have a trial and failure of one beta-blocker and one anticonvulsant unless contraindicated or intolerant.			Initial: 6 months Reauthorization: 12 months	A migraine is defined as a headache that has at least two of the following characteristics: unilateral location, pulsating/throbbing quality, moderate or severe intensity (inhibits or prohibits daily activities), is aggravated by routine activity, nausea and/or vomiting, photophobia and phonophobia. For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
TANERCEPT (ENBREL)	3 - All Medically-accepted Indications.	Un-table Uses	EALUSION LITTERIA	Diagnosis. For rheumatoid arthritis (RA): history of trial and failure, contraindication, or intolerance to a three-month trial with methotrexate or another DMARD. For juvenile idiopathic arthritis (JIA) with polyarthritis: history of trial and failure, contraindication, or intolerance to a 3 month trial with methotrexate, leflunomide, or sulfasalazine. For JIA with oligoarthritis, enthesitis and/or sacroilitis: history of trial and failure, contraindication, or intolerance to at least a 4 week trial of 2 different NSAIDS. For psoriatic arthritis (PsA) one of the following: 1) members with avial or enthesitis must have a history of trial and failure, contraindication, or intolerance to a 4 week trial of 2 NSAIDs. 2) the member has severe disease as defined by the prescriber. 3) members with peripheral disease must have a history of a trial and failure, contraindication, or intolerance to a 12 week trial and failure, contraindication, or intolerance to a 12 week trial and failure, contraindication, or intolerance to a 12 week trial and failure, contraindication, or intolerance to a 12 week trial and failure, contraindication, or intolerance to a 12 week trial.	Age kestriction Member must be 2 years of age or older.	By or in consultation with a rheumatologist or dermatologist.	12 months	For reauth: must have documentation from prescriber indicating stabilization or improvement in condition.	0
FELBAMATE (FELBATOL)	1 - All FDA-approved Indications.			Diagnosis. Must have had an inadequate response or intolerance to two of the following generic anticonvulsant drugs: levetiracetam, phenytoin, carbamazepine, oxcarbazepine, gabapentin, lamotrigine, valproate, or topiramate.	Coverage is provided for members 2 years of age or older.	By or in consultation with a neurologist.	12 months		0
FENFLURAMINE (FINTEPLA)	1 - All FDA-approved Indications.		Use of monoamine oxidase inhibitors within 14 days	Diagnosis. Must have had an inadequate response or intolerance to two generic antiepileptic drugs (e.g. valproate, lamotrigine, topiramate, clobazam).	Member must be 2 years of age or older	By or in consultation with a neurologist	12 months		0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
NTANYL CITRATE RANSMUCOSAL)	1 - All FDA-approved Indications.	On-Label Uses	Acute or postoperative pain including headache/migraines and dental pain.	Diagnosis. Documentation the	Age Restriction	By or in consultation with an oncologist, pain specialist, or hospice/palliative care specialist	12 months	Opioid tolerant is defined as being on around-the-clock medicine consisting of at least 60 mg of oral morphine per day, at least 25 mg of transdermal fentanyl per hour, at least 30 mg of oral oxycodone per day, at least 8 mg of oral hydromorphone per day, at least 56 mg oral hydromorphone per day, at least 60 mg oral hydrocodone per day, or an equianalgesic dose of another opioid daily for a week or longer. For reauthorization: Documentation the member still has active cancer and the member continues to have a medical need for the medication.	0
ERRIC CITRATE (AURYXIA)	1 - All FDA-approved Indications.		Members with Iron overload syndromes (e.g. hemochromatosis) are excluded from coverage. Use for iron deficiency anemia is excluded.	Diagnosis of hyperphosphatemia in adult patients with CKD on dialysis. Must have an intolerance, contraindication, or trial and failure of calcium acetate and sevelamer carbonate.	Coverage provided for members 18 years of age and older	By or in consultation with a hematologist or nephrologist	12 months	For reauth: documentation from prescriber indicating stabilization or improvement in condition.	0
ilgrastim-sndz (zarxio)	3 - All Medically-accepted Indications.			Diagnosis.			6 months	For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	0
LUTICASONE PROPIONATE XHANCE)	1 - All FDA-approved Indications.			Diagnosis of nasal polyps.	Coverage is provided for members 18 years of age or older.		12 months	For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	0

Group FREMANEZUMAB-VFRM (AJOVY)	1 - All FDA-approved Indications.	Off-Label Uses	Exclusion Criteria	Required Medical Information Diagnosis. For episodic migraine: Documentation the member has 4 to 14 headache days per month. For chronic migraine: Documentation the member has at least 15 headache days per month for 3 or more months with at least 8 migraine days per month. For both: Must have a trial and failure of one beta-blocker and one anticonvulsant unless contraindicated or intolerant.	Coverage is provided for members 18 years of age and older	Prescriber Restriction	Coverage Duration Initial: 6 months Reauthorization: 12 months	Other Criteria A migraine is defined as a headache that has at least two of the following characteristics: unilateral location, pulsating/throbbing quality, moderate or severe intensity (inhibits or prohibits daily activities), is aggravated by routine activity, nausea and/or vomiting, photophobia and phonophobia. For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	Part B Prerequisite 0
GANAXOLONE (ZTALMY)	1 - All FDA-approved Indications.			Diagnosis.	Coverage is provided for members 2 years of age or older.	By or in consultation with a neurologist	12 months		0
GIVOSIRAN (GIVLAARI)	1 - All FDA-approved Indications.			Diagnosis. This Prior Authorization requirement only applies to members when a non-FDA approved diagnosis is submitted at the point of sale. FDA-approved diagnosis codes submitted will pay without prior authorization requirement.	Coverage is provided for members 18 years of age or		12 months		0
GLECAPREVIR-PIBRENTASVIR (MAVYRET)	1 - All FDA-approved Indications.		Members with moderate or severe hepatic impairment (Child-Pugh C). Coadministration with atazanavir and rifampin.	Criteria will be applied consistent with current AASLD/IDSA guidance and/or FDA approved labeling	Coverage is provided for members who are age- appropriate according to AASLD/IDSA guidance and/or FDA-approved labeling.	By or in consultation with a gastroenterologist, hepatologist, infectious disease, HIV or transplant specialist.	Criteria will be applied consistent with current AASLD/IDSA guidance and/or FDA approved labeling		0
GLYCEROL PHENYLBUTYRATE (RAVICTI)	1 - All FDA-approved Indications.			Diagnosis. Documentation member has urea cycle disorders (UCDs). Must have a trial of sodium phenylbutyrate with inadequate response or significant side effects/toxicity unless contraindicated.		By or in consultation with a physician who specializes in the treatment of inherited metabolic disorders.	12 months	For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	0
GUANFACINE IR	1 - All FDA-approved Indications.			Diagnosis. Requests for enrollees aged 65 or older must document intolerance to or clinical failure of 2 alternatives such as nifedipine long-acting, amlodipine, felodipine.	Prior authorization only applies to enrollees aged 65 or older. All enrollees less than age 65 are not subject to prior authorization.		12 months	For reauthorization: documentation from prescriber indicating stabilization or improvement in condition.	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
GUSELKUMAB (TREMFYA)	1 - All FDA-approved Indications.			Diagnosis. For Psoriatic arthritis (PsA): for mild to moderate axial or enthesitis, must have a history of trial and failure, contraindication, or intolerance to a 4 week trial of 2 NSAIDs. For members with mild to moderate peripheral disease, must have a history of a trial and failure, contraindication, or intolerance to a 12 week trial with methotrexate or another DMARD. For plaque psoriasis (PsO): minmum BSA involvement of at least 3% (not required if on palms, soles, head/neck, genitalia), a history of trial and failure of ONE of the following: 1) topical therapy (e.g. corticosteroid, calcineurin inhibitor, vitamin D analog), 2) phototherapy, 3) systemic treatment (e.g. methotrexate, cyclosporine, oral retinoids).	Coverage is provided for members 18 years of age and older.	By or inconsultation with a rheumatologist or dermatologist.	12 months.	For reauth: must have documentation from prescriber indicating stabilization or improvement in condition.	
HIGH RISK MEDICATION	1 - All FDA-approved Indications.			Diagnosis. This Prior Authorization requirement only applies to members 65 years of age or older. Prescriber must acknowledge that medication benefits outweigh potential risks in the member 65 years of age or older.	Prior authorization only applies to enrollees aged 65 or older. All enrollees less than age 65 are not subject to prior authorization.		12 months	For reauthorization: Prescriber must acknowledge that medication benefits outweigh potential risks in the member 65 years of age or older.	0
HYDROXYZINE	1 - All FDA-approved Indications.			Diagnosis. Requests for enrollees aged 65 or older must document intolerance to or clinical failure of 2 alternatives such as buspirone or an SSRI (except paroxetine) for anxiety OR cetirizine, levocetirizine for pruritus.	Prior authorization only applies to enrollees aged 65 or older. All enrollees less than age 65 are not subject to prior authorization.		12 months	For reauthorization: documentation from prescriber indicating stabilization or improvement in condition.	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	-	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
ICATIBANT ACETATE	1 - All FDA-approved Indications.	On Liber 1965	and the state of t	Diagnosis of HAE is confirmed by laboratory values obtained on two separate instances (laboratory reports must contain reference ranges). For Type I HAE: Low C4 level and low C1-INH antigenic level. For Type II HAE: Low C4 level and Normal or elevated C1-INH antigenic level and low C1-INH contained the contained of the containe	Coverage is provided for members 18 years of age or older.	By or in consultation with an allergist, immunologist, hematologist, or dermatologist	12 months	Gree Cheria for reauthorization: documentation from prescriber indicating stabilization or improvement in condition.	0
ILOPERIDONE (FANAPT)	1 - All FDA-approved Indications.			Diagnosis. Documentation of trial and failure of at least two of the following generic, oral atypical antipsychotics: olanzapine, quetiapine, clozapine, paliperidone, risperidone, aripiprazole, or ziprasidone.	Coverage is provided for members 18 years of age or older.		12 months		0
IMIPRAMINE	1 - All FDA-approved Indications.			Diagnosis. Requests for enrollees aged 65 or older must document intolerance to or clinical failure of 2 alternatives such as an SSRI (except paroxetine), SNRI, bupropion, trazodone or mirtazapine for depression.	Prior authorization only applies to enrollees aged 65 or older. All enrollees less than age 65 are not subject to prior authorization.		12 months		0
INCOBOTULINUMTOXINA (XEOMIN)	1 - All FDA-approved Indications.			Diagnosis.			12 months	For reauthorization: documentation from prescriber indicating stabilization or improvement in condition.	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
INFLIXIMAB-ABDA (RENFLEXIS)	3 - All Medically-accepted		Doses greater than 5mg/kg in	Diagnosis. For rheumatoid	For RA, PsA, AS, Plaque	By or in consultation with a	12 months	For reauth: must have	0
	Indications.		moderate to severe heart	arthritis (RA): history of trial	Psoriasis: coverage is provided	rheumatologist,		documentation from	
			failure.	and failure, contraindication,	for members 18 years of age	gastroenterologist, or		prescriber indicating	
				or intolerance to a 3 month	or older. For CD, UC: coverage	dermatologist.		stabilization or improvement	
				trial with methotrexate or	is provided for members 6			in condition.	
				another DMARD. For psoriatic	years of age or older.				
				arthritis (PsA) one of the					
				following: 1.)members with					
				axial or enthesitis must have a					
				history of trial and failure,					
				contraindication, or					
				intolerance to a 4 week trial of					
				2 NSAIDs. 2.) the member has					
				severe disease as defined by					
				the prescriber. 3.) members					
				with peripheral disease must					
				have a history of a trial and					
				failure, contraindication, or					
				intolerance to a 12 week trial					
				with methotrexate or another					
				DMARD. For ankylosing					
				spondylitis (AS): history of trial					
				and failure, contraindication,					
				or intolerance to a four-week					
				trial each of at least 2 NSAIDs.					
				For plaque psoriasis: minimum					
				BSA involvement of at least 3%					
				(not required if on palms,					
				soles, head/neck, genitalia), a					
				history of trial and failure of					
				ONE of the following: 1)					
				topical therapy (e.g.					
				corticosteroid, calcineurin					
				inhibitor vitamin Danalog) 2)					

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
INFLIXIMAB-DYYB (INFLECTRA)			Doses greater than 5mg/kg in	Diagnosis. For rheumatoid	For RA, PsA, AS, Plaque	By or in consultation with a	12 months	For reauth: must have	0
	Indications.		moderate to severe heart	arthritis (RA): history of trial	Psoriasis: Coverage is provided			documentation from	
			failure.	and failure, contraindication,	for members 18 years of age	gastroenterologist, or		prescriber indicating	
				or intolerance to a three-	or older. For CD, UC: Coverage	dermatologist.		stabilization or improvement	
				month trial with methotrexate				in condition.	
				or another DMARD. For	years of age or older.				
				psoriatic arthritis (PsA) one of					
				the following: 1).members					
				with axial or enthesitis must					
				have a history of trial and					
				failure, contraindication, or					
				intolerance to a 4 week trial of					
				2 NSAIDs. 2.) the member has severe disease as defined by					
				the prescriber. 3.) members					
				with peripheral disease must					
				have a history of a trial and					
				failure, contraindication, or					
				intolerance to a 12 week trial					
		1		with methotrexate or another					
				DMARD. For ankylosing					
				spondylitis (AS): history of trial					
				and failure, contraindication,					
				or intolerance to a four-week					
				trial each of at least 2 NSAIDs.					
				For plaque psoriasis: minimum					
				BSA involvement of at least 3%					
				(not required if on palms,					
				soles, head/neck, genitalia), a					
				history of trial and failure of					
				ONE of the following: 1)					
				topical therapy (e.g.					
				corticosteroid, calcineurin					
IPTACOPAN (FABHALTA)	1 - All FDA-approved		Initiation in patients with	Diagnosis. For paroxysmal	Coverage is provided for	By or in consultaion with a	12 months	PNH Reauth: Documentation	0
	Indications.		unresolved serious infection	nocturnal hemoglobinuria	members 18 years of age and	hematologist, oncologist,		of LDH level (within 3 months)	
			caused by encapsulated	(PNH): confirmed diagnosis of	older.	immunologist, or genetic		that shows a reduction form	
			bacteria.	PNH by flow cytometry		specialist.		baseline and one of the	
				testing. Flow cytometry				following: 1) If baseline Hgb	
				pathology report must be				was 9 g/dL or higher, it has not	
				supplied and demonstrate at				dropped by more than 2g/dL	
				least 2 different GPI protein				from baseline. 2) If baseline	
				deficiencies within 2 different				Hgb was less than 10g/dL, it is	
				cell lines from granulocytes,				above 7g/dL.	
				monocytes, or erythrocytes.					
				Member is transfusion					
				dependent as defined by					
		1		having a transfusion within the					
		1		last 12 months and one of the follwoing: a hemoglobin is less					
				than or equal to 7 g per dL or					
				has symptoms of anemia and					
				the hemoglobin is less than or					
		1		equal to 10 g per dL. Must					
				have a lactate dehydrogenase					
				(LDH) level at least 1.5 times					
		1		the upper limit of the normal					
				range.					
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Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
IVABRADINE (CORLANOR)	1 - All FDA-approved		Acute decompensated heart	_	CHF: coverage is provided for	By or in consultation with a	12 months	For reauthorization:	0
	Indications.		failure, blood pressure less	Heart Failure (CHF): Must have		cardiologist		documentation from	
			than 90/50 mmHG, sick sinus		older. DCM: coverage is			prescriber indicating	
			syndrome, sinoatrial block, or		provided for members 6			stabilization or improvement	
			3rd degree AV block-unless a	equal to 35%, member is in	months of age or older.			in condition.	
			functioning demand	sinus rhythm and has a resting					
			pacemaker is present, resting	heart rate of greater than or					
			heart rate less than 60 bpm	equal to 70 beats per minute,					
			prior to treatment, severe	must currently be taking a					
			hepatic impairment,	beta-blocker (e.g., bisoprolol,					
			pacemaker dependence (heart						
			rate maintained exclusively by						
			the pacemaker), concomitant	tolerated dose or has a					
			use of strong CYP3A4	contraindication to beta-					
			inhibitors.	blocker use. For Pediatric					
				Dilated Cardiomyopathy					
				(DCM): Must have stable					
				symptomatic heart failure with					
				left ventricular ejection					
				fraction less than or equal to					
				45%, must be in sinus rhythm,					
				must have an elevated heart					
				rate (greater than or equal to					
				105 beats per minute (BPM)					
				for 6-12 months of age,					
				greater than or equal to 95 for					
				1-3 years of age, greater than					
				or equal to 75 for 3-5 years of					
				age, greater than or equal to					
				70 for 5-18 years of age).					
IVACAFTOR (KALYDECO)	1 - All FDA-approved			Diagnosis. Documentation of	Coverage is provided for	By or in consultation with a	12 months	For reauthorization:	0
, , , , , , , , , , , , , , , , , , , ,	Indications.			genetic test confirming the		pulmonologist or cystic fibrosis		documentation from	
					older.	specialist		prescriber indicating	
				mutation in the CFTR gene		-		stabilization or improvement	
				that is responsive to ivacaftor				in condition.	
				based on clinical and/or in					
				vitro assay data.					
				,					

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
oup EKIZUMAB (TALTZ)	1 - All FDA-approved Indications.	Off-Label Uses	Exclusion Criteria	Required Medical Information Diagnosis. For non- radiographic axial spondyloarthritis (nr-axSpA): history of trial and failure, contraindication, or intolerance to a four-week trial each of at least 2 NSAIDs. For ankylosing spondylitis (AS): history of trial and failure, contraindication, or intolerance to a 4 week trial each of at least 2 NSAIDs. For plaque psoriasis: minimum BSA involvement of at least 3% (not required if on palms, soles, head/neck, genitalia), a history of trial and failure of ONE of the following: 1) topical therapy (e.g. corticosteroid, calcineurin inhibitor, vitamin D analog), 2) phototherapy, 3) systemic treatment (e.g. methotrexate, cyclosporine, oral retinoids). For psoriatic arthritis (PsA) one of the following: 1) members with axial or enthesitis must have a history of trial and failure, contraindication, or intolerance to a 4 week trial of 2 NSAIDs. 2) the member has	For plaque psoriasis: member must be 6 years of age or older. All other diagnoses: member must be 18 years of age or older.	Prescriber Restriction By or in consultation with a rheumatologist or dermatologist.	12 months	For reauth: must have documentation from prescriber indicating stabilization or improvement in condition.	Part B Prerequisite 0
-GLUTAMINE (ENDARI)	1 - All FDA-approved			severe disease as defined by the processing. 2) members Diagnosis. Must be used to	Coverage is provided for	By or in consultation with a	12 months	For reauthorization:	0
	Indications.			reduce the acute complications of sickle cell disease (SCD) and the member must have experienced at least 2 painful episodes of sickle cell crises (SCC) in the previous 12 months. Member has had an adequate trial (3 months) of hydroxyurea unless the member has tried and failed or has a contraindication to hydroxyurea.	older	physician who specializes in SCD (e.g.a hematologist)		Documentation there has been a reduction in vaso-occlusive painful events or an improvement in condition.	

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Group LAMIVUDINE (EPIVIR-HBV)	Indication Indicator 1 - All FDA-approved Indications.	Off-Label Uses	Exclusion Criteria Coverage is not provided when coadministered with other medications that contain lamivudine or emtricitabine.	Required Medical Information Diagnosis. Member must have chronic hepatitis B virus (HBV) infection associated with evidence of hepatitis B viral replication and active liver inflammation. Member must have a previous trial and inadequate response or intolerance to or contraindication to an alternative antiviral agent with a higher genetic barrier resistance for Hepatitis B (such as entecavir or tenofovir).	Member must be 2 years of age or older	Prescriber Restriction By or in consultation with a gastroenterologist, hepatologist, infectious disease, HIV or transplant specialist.	Coverage Duration 12 months	Other Criteria For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	Part B Prerequisite 0
LANREOTIDE (SOMATULINE DEPOT)	1 - All FDA-approved Indications.			Diagnosis. For acromegaly: must have inadequate response to surgery or radiotherapy or documentation that these therapies are inappropriate, must have the following baseline labs: elevated serum 16F-1 level for gender/age range (including lab reference range) and elevated growth hormone level defined as GH at least 1ng/mL during oral glucose tolerance test.	Coverage is provided for members 18 years of age and older.	By or in consultation with an endocrinologist or oncologist	For oncology indications: 6 months. All other indications: 12 months	For reauth: documentation of improvement or stabilization.	0
LEDISPASVIR-SOFOSBUVIR (HARVONI)	1 - All FDA-approved Indications.			Criteria will be applied consistent with current AASLD/IDSA guidance and/or FDA approved labeling	Coverage is provided for members who are age- appropriate according to AASLD/IDSA guidance and/or FDA-approved labeling.	By or in consultation with a gastroenterologist, hepatologist, infectious disease, HIV or transplant specialist.	Criteria will be applied consistent with current AASLD/IDSA guidance and/or FDA-approved labeling.		0
LENIOLISIB (JOENJA)	1 - All FDA-approved Indications.		0	Diagnosis of activated phosphoinositide 3-kinase delta syndrome (APDS). Must have genetic testing confirming the PI3K delta mutation with a documented variant in either PIK3CD or PIK3RI. Documentation of inadequate response to immunoglobulins.	Coverage is provided for members 12 years of age or older.	By or in consultation with a hematologist, immunologist, or geneticist.	12 months.	0	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
LETERMOVIR (PREVYMIS)	1 - All FDA-approved Indications.		Use with pimozide or ergot alkaloids. Use with pitavastatin and simvastatin when co-administered with cyclosporine.	Diagnosis. Must have received either an allogeneic hematopoietic stem cell transplant (HSCT) and have tested CMV-seropositive (Recipient positive, R+) or received a kidney transplant and be a high risk donor (CMV seropositive D+/recipient CMV seronegative R-). Must be used for prophylaxis of CMV infection.		By or in consultation with a hematologist, infectious disease or transplant specialist.	HSCT: 4 mo, up to 100 days post-transplant. Kidney Transplant: 8 mo, up to 200 days post-transplant.	For reauth: no reauthorization after initial coverage period.	0
LEUPROLIDE ACETATE	1 - All FDA-approved Indications.			Diagnosis. For endometriosis: Documentation the member has tried and failed or has a contraindication to 2 conventional treatments such as oral contraceptives, non steroidal anti-inflammatory agents, progestins, or danazol. For CPP: Documentation that the age of onset of secondary sexual characteristics occurred at less than 8 years of age in a female child or less than 9 years of age in a male child.			Prostate cancer and endometriosis: 6 months. CPP or Fibroids: 3 months	For reauth: documentation indicating stabilization or improvement in condition. For endometriosis, a single retreatment course of not more than six months may be administered after the initial course of treatment if symptoms recur	0
LEUPROLIDE- NORETHINDRONE (LUPANETA)	1 - All FDA-approved Indications.		Current or history, known or suspected cancer of the breast or hormone sensitive. Hepatic disease or tumor. Breastfeeding or pregnancy (known or suspected). Thrombotic or thromboembolic disorder. Abnormal uterine bleeding.		Coverage is provided for members 18 years of age and older.		6 months	For reauth: documentation indicating stabilization or improvement in condition. A single retreatment course of not more than six months may be administered after the initial course of treatment if symptoms recur.	0
LEVETIRACETAM (SPRITAM)	1 - All FDA-approved Indications.			Diagnosis. Must have had an inadequate response or intolerance to generic levetiracetam and at least one of the following generic anticonvulsant drugs: phenytoin, carbamazepine, oxcarbazepine, gabapentin, lamotrigine, valproate, or topiramate.	Coverage is provided for members 4 years of age and older weighing more than 20kg.	By or in consultation with a neurologist.	12 months		0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
LEVOMILNACIPRAN (FETZIMA)				Diagnosis. Documentation of trial and failure of at least two generic antidepressants alternatives such as an SSRI, SNRI, bupropion, trazodone or mirtazapine	Coverage is provided for members 18 years of age and older.		12 months		0
LIDOCAINE PATCH	1 - All FDA-approved Indications.			Diagnosis. This Prior Authorization requirement only applies to members when a non-FDA approved diagnosis is submitted at the point of sale. FDA-approved diagnosis codes submitted will pay without prior authorization requirement.			12 months		0
LOTILANER (XDEMVY)	1 - All FDA-approved Indications.			Diagnosis of Demodex blepharitis confirmed by both of the following: 1. Member has at least mild erythema or itching of the upper eyelid margin. 2. Mite presence (e.g. collarettes) confirmed by slit lamp examination of the eyelashes.	Member must be 18 years of age and older	Prescribed by or in consultation with an optometrist or ophthalmologist	6 weeks		0
LUMACAFTOR/IVACAFTOR (ORKAMBI)	1 - All FDA-approved Indications.			Diagnosis. Documentation of a genetic test confirming that the member is homozygous for the F508del mutation in the CFTR gene (has two copies of the F508del mutation in the CFTR gene).		By or in consultation with a pulmonologist or cystic fibrosis specialist	12 months	For reauthorization: documentation from prescriber indicating stabilization or improvement in condition.	0
MACITENTAN (OPSUMIT)	1 - All FDA-approved Indications.		Pregnancy	Diagnosis. Pulmonary arterial hypertension (PAH) WHO Group I confirmed by chart documentation of right-heart catheterization (RHC) indicating a mean pulmonary arterial pressure greater than 20 mmHg, pulmonary vascular resistance greater than 2 wood units, and mean pulmonary capillary wedge pressure less than or equal to 15 mmHg. If provider indicates RHC is not recommended, must have documentation of echocardiography.		Prescribed by or in consultation with cardiologist or pulmonologist.	Initial: 3 months Reauth: 12 months	For reauth: documentation from prescriber that demonstrates member is tolerating and receiving clinical benefit from treatment	0

iroup	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
MANNITOL (BRONCHITOL)	1 - All FDA-approved			Diagnosis. Must have passed			12 months	For reauth: documentation of	0
	Indications.			a bronchitol tolerance test.				improvement	
				Must be used as add-on					
				maintenance treatment with					
				standard therapies (e.g.					
				bronchodilators, antibiotics,					
				anti-inflammatory therapy) to					
				improve pulmonary function.					
IARALIXIBAT (LIVMARLI)	1 - All FDA-approved		PFIC type 2 patients with	Diagnosis of pruritis caused by	Coverage is provided for	By or in consultation with a	12 months	For reauth: documentation of	0
TATALIXIDAT (LIVIVIANCI)	Indications.		specific ABSB11 variants	progressive familial	members 3 months of age and		12 months	improvement in pruritis.	o a
	indications.		resulting in non-functional or	intrahepatic cholestatis (PFIC)		gastroenterologist.		improvement in prunus.	
			complete absence of bile salt	or Allagile syndrome (ALGS)	older.	gasti deriter diogist.			
			export pump (BSEP) protein.	which has been confirmed by					
			export pump (B3EF) protein.						
				genetic testing. Documentation of trial and					
				failure of ursodiol and another					
				medication for cholestatic					
				pruritis (ie. cholestyramine,					
				colestipol, rifampin).					
MAVORIXAFOR (XOLREMDI)	1 - All FDA-approved			Diagnosis. Confirmation of the	Members 12 years of age and	By or in consultation with an	12 months	For reauthorization:	0
	Indications.			diagnosis with a genetic test	older	immunologist, hematologist,		Documentation of one of the	
				confirming pathogenic or likely		or dermatologist		following: 1. an improvement	
				pathogenic variants in the				in ANC or ALC from baseline 2.	
				CXCR4 gene. Documentation				A decrease in frequency or	
				of a baseline absolute				severity of infections since	
		1		neutrophil count (ANC) less	1			initiating therapy.	
				than or equal to 400 cells/?L	1				
				or absolute lymphocyte count	1				
				(ALC) less than or equal to 650					
				cells/?L. Documentation of	1				
				symptoms and complications	1				
		1		associated with WHIM	1				
				syndrome (e.g. warts,	1				
				hypogammaglobulinemia,					
				recurrent infections, and	1				
				myelokathexis)					
				,	1				
					1				
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Group MECASERMIN (INCRELEX)	Indication Indicator 1 - All FDA-approved Indications.	Off-Label Uses	Exclusion Criteria Coverage is not provided for members with active or suspected neoplasia, closed epiphyses.	Required Medical Information Diagnosis. Growth chart and documentation that epiphyses are open. For growth hormone deletion: must have growth hormone (GH) gene deletion in gene GH1 and developed neutralizing antibodies to GH therapy. For growth failure due to severe IGF-1 deficiency: must have dx of severe IGF-1 deficiency (defined as having all of the following: height below or equal to 3.0 standard deviation (SD) of the mean for age and sex, basal IGF-1 SD of less than or equal to 3.0 based on lab reference range, normal or elevated GH defined as stimulated serum GH level of greater than 10ng/mL or basal serum GH level greater than 5ng/mL).	Coverage is provided for members 2 years of age or older.	Prescriber Restriction By or in consultation with an Endocrinologist	12 months	Other Criteria For reauth, must include a recent progress note from prescriber indicating growth and maturation as a result of treatment and that epiphyses have not closed.	Part B Prerequisite 0
METHOTREXATE ORAL SOLUTION (XATMEP)	3 - All Medically-accepted Indications.			Diagnosis. Medical rationale why patient cannot take methotrexate tablet formulation.			6 months	For reauthorization: documentation from prescriber indicating stabilization or improvement in condition.	0
METHSCOPOLAMINE (PAMINE, PAMINE FORTE)	1 - All FDA-approved Indications.			Diagnosis. Requests for enrollees aged 65 or older must document intolerance to or clinical failure of 2 alternative antiulcer agents (such as omeprazole, sucralfate, famotidine).	Prior authorization only applies to enrollees aged 65 or older. All enrollees less than age 65 are not subject to prior authorization.		12 months	For reauthorization: documentation from prescriber indicating stabilization or improvement in condition.	0
METHYLNALTREXONE (RELISTOR)	1 - All FDA-approved Indications.		Known or suspected gastrointestinal obstruction and members at an increased risk of recurrent obstruction.	Diagnosis. For opioid-induced constipation and advanced life-limiting illness: must have documentation of previous trial of lactulose. For opioid-induced constipation with chronic non-cancer pain: must have documentation of current and ongoing opioid therapy and must have trials with inadequate responses or significant side effects/toxicity or have a contraindication to naloxegol (Movantik) and lactulose.	Coverage is provided for members 18 years of age and older.		12 months	For reauth: documentation from the prescriber indicating an improvement in condition (both diagnoses) and must continue to be on opioid therapy (non-cancer pain).	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
MIFEPRISTONE (KORLYM)	1 - All FDA-approved Indications.			Diagnosis. Must have failed surgery or not be a candidate vargery. Female members of reproductive potential: must have baseline (within previous month, must include date of test) negative pregnancy test prior to starting mifepristone and must be using nonhormonal medically acceptable method of contraception (unless surgically sterilized) during treatment and for 1 month after mifepristone therapy.		By or in consultation with an endocrinologist	12 months		0
MIGLUSTAT (ZAVESCA)	1 - All FDA-approved Indications.		Miglustat is being used in combination with another therapy for Gaucher's disease	Diagnosis. Documentation the member has at least one of the following: 1) anemia not due to iron deficiency with a low hemoglobin for age and sex, 2) thrombocytopenia 3) evidence of bone disease, 4) presence of hepatomegaly or splenomegaly. Enzyme replacement therapy must not be a therapeutic option for the member (i.e. due to allergy, hypersensitivity, or poor venous access).	older.	By or in consultation with an appropriate specialist (i.e. hematologist, geneticist, radiologist, orthopedist, endocrinologist, rheumatologist)	12 months	Reauthorization: Documentation from the prescriber indicating improvement or stabilization in member's condition.	o
MITAPIVAT (PYRUKYND)	1 - All FDA-approved Indications.			Diagnosis of hemolytic anemia with pyruvate kinase deficiency (PKD) confirmed by genetic testing.	Coverage is provided for members 18 years of age or older.	By or in consultation with a hematologist or a physician who specializes in the treatment of inherited metabolic disorders.	12 months	For reauthorization: documentation of improvement in condition.	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction		Coverage Duration	Other Criteria	Part B Prerequisite
MODAFINIL (PROVIGIL)	1 - All FDA-approved Indications.	Um-Label Uses		Diagnosis. For narcolepsy and obstructive sleep apnea: Sleep Study (e.g. Polysomnogram, Multiple Sleep Latency Test) confirming diagnosis. For shift work sleep disorder (SWSD): must meet International Classification of Sleep Disorders criteria for SWSD (either primary complaint of excessive sleepiness or insomnia temporarily associated with work period that occurs during habitual sleep phase OR polysomnography and Multiple Sleep Latency Test demonstrate loss of normal sleep wake pattern, no other medical or mental disorders account for symptoms, and symptoms do not meet criteria for any other sleep disorder producing insomnia or excessive sleepiness such as time zone change syndrome) and must provide documentation of shift work schedule showing 5 or more night shifts per month (defined as at least 4 hours of shift occurring between 10pm and 8am).		By or in consultation with a sleep specialist, ENT (ear, nose, and throat specialist), neurologist, or pulmonologist	SWSD: 6 months. Narcolepsy, OSA: 12 months	For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	0
MONOMETHYLFUMARATE (BAFIERTAM)	1 - All FDA-approved Indications.		Coverage is not provided with coadministration with dimethyl fumarate or diroximel fumarate.	relapsing Multiple Sclerosis (MS), including clinically isolated syndrome, relapsing-remitting disease, and active secondary progressive disease. Documentation that functional status is preserved and patient is either still able to walk at least a few steps or alternatively must have some functional arm/hand use consistent with performing activities of daily living.	Coverage is provided for members 18 years of age or older.	neurologist	12 months	For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	
MULTIPLE SCLEROSIS THERAPIES	1 - All FDA-approved Indications.			Diagnosis. Documentation that functional status is preserved and patient is either still able to walk at least a few steps or alternatively must have some functional arm/hand use consistent with performing activities of daily living.		By or in consultation with a neurologist or gastroenterologist	12 months	For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
ETARSUDIL (RHOPRESSA)	1 - All FDA-approved Indications.			Diagnosis. Member must have a baseline intraocular pressure of less than 30 mmHg. Documentation of trial and failure, contraindication, or intolerance to timolol and latanoprost.	Coverage is provided for members 18 years of age and older.		12 months	For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	0
IINTEDANIB (OFEV)	1 - All FDA-approved Indications.			Diagnosis. For a diagnosis of Idiopathic Pulmonary Fibrosis (IPF): Must have diagnosis confirmed by either high-resolution computed tomography (HRCT) or surgical lung biopsy and must have all other diagnoses ruled out (e.g., domestic and occupational environmental exposures, connective tissue disease, and drug toxicity). Must have a forced vital capacity (FVC) greater than or equal to 50% of predicted and a carbon monoxide diffusing capacity (DLCO) of at least 30% of predicted. For a diagnosis of Systemic Sclerosis-Associated Interstitial Lung Disease (SSc-ILD): Must have onset of disease (first non-Raynaud symptom) within the past 7 years and at least 10% fibrosis on a chest high-resolution computed tomography (HRCT) scan within the past 12 months. Must have a FVC greater than or equal to 40% of predicted and a DLCO of at least 30% of predicted. For a diagnosis of Chronic Fibrosing Interstitial	Coverage provided for members age 18 years and older.	By or in consultation with a pulmonologist	Initial: 6 months, Reauth: 12 months	For reauth: must have documentation from prescriber indicating that member still is a candidate for treatment.	0
ITISINONE (ORFADIN)	1 - All FDA-approved Indications.			Diagnosis of hereditary tyrosinemia type 1 (HT-1) confirmed by DNA testing or biochemical testing (ie. urine succinylacetone (SA) level).			12 months	For reauth: Documentation from the prescriber indicating improvement or stabilization in the member's condition	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
NITROGLYCERIN 0.4% SINTMENT (RECTIV)	1 - All FDA-approved Indications.		Acute circulatory failure or shock, allergy to corn or corn products, severe anemia (defined as hemoglobin less than 8g/dL). Increased intracranial pressure. Concomitant use of a phosphodiesterase type 5 (PDES) inhibitor such as sildenafil (Revatio, Viagra), tadalafil (Adcirca, Cialis), or vardenafil (Levitra, Staxyn) or riociguat use or other soluble guanylate cyclase inhibitors.	Diagnosis. Must provide documentation that chronic anal fissure symptoms have persisted for at least 6 weeks.	Coverage is provided for members 18 years of age or older.		Initial: 2 months Reauthorization: 12 months	For reauthorization: documentation from prescriber indicating stabilization or improvement in condition.	0
NORTRIPTYLINE (PAMELOR)	1 - All FDA-approved Indications.			Diagnosis. Requests for enrollees aged 65 or older must document intolerance to or clinical failure of 2 alternatives such as an SSRI (except paroxetine), SNRI, bupropion, trazodone or mirtazapine for depression.	Prior authorization only applies to enrollees aged 65 or older. All enrollees less than age 65 are not subject to prior authorization.		12 months		0
DDEVIXIBAT (BYLVAY)	1 - All FDA-approved Indications.			Diagnosis of pruritis caused by progressive familial intrahepatic cholestatis (PFIC) or Alagille syndrome (ALGS) which has been confirmed by genetic testing. Documentation of trial and failure of cholestyramine.	members 3 months of age and	By or in consultation with a hepatologist or gastroenterologist.	12 months	For reauth: documentation of improvement in pruritis.	0
OFATUMUMAB (KESIMPTA)	1 - All FDA-approved Indications.		Coverage is not provided for members with active HBV infection.	Diagnosis. Member must have relapsing forms of multiple sclerosis (MS), to include clinically isolated syndrome, relapsing-remitting disease, and active secondary progressive disease. Documentation that functional status is preserved and patient is either still able to walk at least a few steps or alternatively must have some functional arm/hand use consistent with performing activities of daily living.	Coverage is provided for members 18 years of age or older.	By or in consultation with a neurologist	12 months	For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	0
DLANZAPINE (ZYPREXA IELPREVV)	1 - All FDA-approved Indications.			Diagnosis. Documentation of prior oral olanzapine therapy.	Coverage is provided for members 18 years of age or older		12 months	For reauthorization: documentation from prescriber indicating stabilization or improvement in condition.	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
OLANZAPINE/SAMIDORPHAN (LYBALVI)	1 - All FDA-approved Indications.			Diagnosis. Documentation of trial and failure of at least two of the following generic, oral atypical antipsychotics: olanzapine, quetiapine, clozapine, paliperidone, risperidone, aripiprazole, or ziprasidone.	Coverage is provided for members 18 years of age or older.		12 months		0
OMALIZUMAB (XOLAIR)	1 - All FDA-approved Indications.		0	Diagnosis. For moderate to severe allergic asthma: recent total serum IgE level fo greater than 30 IU/ml and the pretreatment IgE levels do not exceed manufacturers dosing recommendations. Documentation of recent use and failure to respond to inhaled steroid in combo with long acting beta agonist. Documenation of a positive skin or in vitro reactivity to perennial aeroallergan. Must have asthma symptoms that are inadequately controlled while on treatment (uncontrolled defined as having an asthma exacerbation requiring hospitalization in the past year or having 2 or more asthma exacerbations requiring oral systemic steroids). Must follow recommended dosing guidelines based upon weight and IgE level. For chronic spontaneous urticaria (CSU): must have chart documentation showing history of urticaria with presence of hives, must have trial of one 2 nd generation H1	0	By or in consultation with, for Urticaria: allergist, dermatologist, immunologist. Asthma: pulmonologist or allergist. Nasal polyps: allergist, ear/nose/throat specialist, or immunologist. Allergy: allergist or immunologist.	12 months	For reauthroization: documentation from prescriber indicating stabilization or improvement in condition.	
OMAVELOXOLONE (SKYCLARYS)	1 - All FDA-approved Indications.			Diagnosis of Friedreich's ataxia that has been confirmed by genetic testing. Must have a modified Friedreich's Ataxia Rating Scale (mFARS) score between 20 and 80. Must have a left ventricular ejection fraction of at least 40%.	Coverage is provided for members 16 years of age or older.	By or in consultation with a neurologist.	12 months		0
OMNIPOD POD	1 - All FDA-approved Indications.			Must have documentation of previous insulin use.			12 months		0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
DIABOTULINUMTOXINA BOTOX)	1 - All FDA-approved Indications.			Diagnosis. For migraine prophylaxis: must have adequate trial of two migraine prophylactic agents each from a separate class (e.g., anticonvulsants, beta-blockers, tricyclic antidepressants) with inadequate response. For urinary incontinence or OAB with urge urinary incontinence, urgency, frequency: must have adequate trial (at least 4 weeks) at recommended dose of 2 anticholinergic meds (e.g., oxybutynin ER, oxybutynin, Toviaz) with inadequate response or intolerance unless contraindicated.			12 months	For reauth: documentation from prescriber indicating stabilization or improvement in condition.	0
DNCOLOGY MEDICATIONS	3 - All Medically-accepted Indications.			Diagnosis. For Bosulif, Iclusig, and Tasigna for CML: must have had an inadequate response or intolerance to imatinib or Sprycel. For Kisqali or Kisqali Femara copack: must have had an inadequate response or intolerance to Ibrance and Verzenio.		By or in consultation with an oncologist, hematologist, neurologist, rearsplant specialist, allergist, or immunologist.	6 months		0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information Age Restr	triction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
ORAL BENZODIAZEPINES	3 - All Medically-accepted			Prior authorization is only			12 months	Reauth: For ongoing opioid	0
	Indications.			required for requests greater				and benzodiazepine therapy:	
				than a 14 day supply in a 30				Documentation to taper the	
				day period and for members				benzodiazepine or opioid. If a	
				not in hospice care. Diagnosis.				taper is not appropriate at this	
				For seizure disorder:				time, documentation of when	
				documentation the member				the taper will be reevaluated.	
				has tried and failed or had an				For all other ongoing therapy:	
				intolerance or contraindication				documentation the member	
				to at least one non-				has been treated with the	
				benzodiazepine				requested agent within the	
				anticonvulsant. For sleep				past 90 days	
				disorder: documentation the					
				member has tried and failed or					
				had an intolerance to at least 2					
				non-benzodiazepine sleep					
				medications. For a psychiatric					
				disorder (e.g. generalized					
				anxiety disorder, panic					
				disorder, post-traumatic stress					
				disorder, etc.): documentation					
				of one of the following: 1. the					
				member tried and failed or					
				had an intolerance or					
				contraindication to at least 2					
				antidepressants. 2. The					
				request is related to a recent					
1				hospitalization within the past					
				3 months. 3. The requested					
İ				therapy is medically necessary					
İ				to prevent harm to the					
				member or others. For a					
İ				musculoskeletal disorder:					
				documentation the member				1	

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information		Coverage Duration	Other Criteria	Part B Prerequisite
PALOVAROTENE (SOHONOS)	1 - All FDA-approved Indications.		having received Beyfortus (nirsevimab-alip) for the current RSV season		Less than 12 months or less than 24 months of age at start of RSV season depending on criteria. Members assigned female at birth must be 8 years and older. Members assigned male at birth must be 10 years and older.	Minimum duration 1 month. Maximum of 5 doses per RSV season 12 months		0
PAMIDRONATE (AREDIA)	1 - All FDA-approved Indications.			Diagnosis. For hypercalcemia of malignancy: must be used in conjunction with adequate hydration in members with moderate or severe hypercalcemia associated with malignancy, with or without bone metastases. For Paget's disease: must have moderate to severe Paget's disease of bone. For osteolytic bone metastases of breast cancer and osteolytic lesions of multiple myeloma: must be used in conjunction with standard antineoplastic therapy.	Coverage is provided for members 18 years of age or older.	12 months	For reauth: documentation from prescriber indicating stabilization or improvement in condition.	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
PASIREOTIDE (SIGNIFOR)	1 - All FDA-approved Indications.			Diagnosis of Cushing's disease for whom pituitary surgery is not an option or has not been curative. Documentation of trial and failure with ketoconazole to reduce cortisol secretion.	Coverage is provided for members 18 years of age or older.	By or in consultation with an Endocrinologist	12 months	For reauth: documentation of improvement or stabilization.	· · · · · · · · · · · · · · · · · · ·
PEGFILGRASTIM-BMEZ (ZIEXTENZO)	3 - All Medically-accepted Indications.			Diagnosis.			6 months	For reauth: documentation from prescriber that demonstrates member is tolerating and receiving clinical benefit from treatment	0
PEGVISOMANT (SOMAVERT)	1 - All FDA-approved Indications.			Diagnosis of acromegaly. Must have inadequate response to surgery or radiation therapy or documentation that these therapies are inappropriate. Must have a trial and failure or inadequate response to one medical therapy (e.g. octreotide, octreotide LAR, lanreotide) or documentation that these therapies are inappropriate. Must have the following baseline labs: elevated serum IGF-1 level for gender/age range (including lab reference range) and elevated growth hormone level defined as GH at least 1ng/mL during oral glucose tolerance test.	members 18 years of age or older.	By or in consultation with an Endocrinologist	12 months	For reauth: documentation of improvement or stabilization.	0
PERAMPANEL (FYCOMPA)	1 - All FDA-approved Indications.			Diagnosis. Must have had an inadequate response or intolerance to two of the following generic anticonvulsant drugs: levetiracetam, phenytoin, carbamazepine, oxcarbazepine, gabapentin, lamotrigine, valproate, or topiramate.	Coverage is provided for members 4 years of age or older.	By or in consultation with a neurologist.	12 months		0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information		Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
PIMAVANSERIN (NUPLAZID)	1 - All FDA-approved Indications.			Diagnosis. Must be using for the treatment of hallucinations and delusions associated with Parkinson's disease psychosis. Must provide clinical rationale for diagnosis and exclusion of other diagnoses (e.g., dementia with Lewy bodies, visual processing deficits/loss of visual acuity, infectious causes). Must have tried to discontinue or reduce dose of any medication(s) that may cause or contribute to hallucinations and delusions (e.g., dopamine agonist, amantadine, monoamine oxidase B inhibitors, anticholinergics) or provide clinical rationale indicating why dose reduction or discontinuation of applicable medications would not be appropriate. Submission of a Mini-Mental State Examination (MMSE) score greater than or equal to 21 and documentation the member is able to self-report symptoms.	Coverage is provided for members 18 years of age or older.	By or in consultation with a neurologist or psychiatrist	12 months		
PIRFENIDONE (ESBRIET)	1 - All FDA-approved Indications.			Diagnosis. Must have diagnosis of idiopathic pulmonary fibrosis (IPF) confirmed by either high-resolution computed tomography (HRCT) or surgical lung biopsy. Must have all other diagnoses ruled out (e.g., domestic and occupational environmental exposures, connective tissue disease, and drug toxicity). Must have forced vital capacity (FVC) greater than or equal to 50% and a percent predicted diffusing capacity of the lungs for carbon monoxide (DLCO) greater than or equal to 30%		Pulmonologist	Initial: 6 months, Reauth: 12 months	For reauth: must have documentation from prescriber indicating that member still is a candidate for treatment.	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
POLYPHARMACY - MULTIPLE ACH MEDICATIONS	Indication in the control of the con		The second secon	This prior authorization requirement applies to members on 2 or more unique anticholinergic medications. Diagnosis. Provider must acknowledge that the benefit of the combination of the medications outweighs the potential risks. Documentation of both of the following: 1. the member has tried and failed monotherapy. 2. clinical rationale for use of 2 or more anticholinergic medications.	Prior authorization only applies to enrollees aged 65 or		12 months	Reauthorization: Documentation of one of the following: 1. attempt to taper of one of the medications OR 2. documentation of why tapering one of the medications is not appropriate at this time. Provider attestation the member continues to benefit from the combination of medications and this outweighs any potential risks.	0
POLYPHARMACY - MULTIPLE CNS MEDICATIONS	1 - All FDA-approved Indications.			This prior authorization requirement applies to members on 3 or more central nervous system (CNS) medications. Diagnosis. Provider must acknowledge that the benefit of the combination of the medications outweighs the potential risks. For a seizure diagnosis: no further criteria is required. For all other diagnoses: Documentation of both of the following: 1. the member has tried and failed monotherapy and dual therapy 2. clinical rationale for use of 3 or more CNS medications.	Prior authorization only applies to enrollees aged 65 or older not in hospice care.		12 months	Reauthorization: Documentation of one of the following: 1. attempt to taper of one of the medications OR 2. documentation of why tapering one of the medications is not appropriate at this time. Provider attestation the member continues to benefit from the combination of medications and this outweighs any potential risks.	0
POSACONAZOLE (NOXAFIL)	1 - All FDA-approved Indications.		Coadministration with sirolimus, ergot alkaloids (e.g., ergotamine, dihydroergotamine), HMG-CoA reductase inhibitors that are primarily metabolized through CYP3A4 (e.g., atorvastatin, lovastatin, simvastatin), or CYP3A4 substrates that prolong the QT interval (e.g., pimozide, quinidine), hypersensitivity to posaconazole, other azole antifungal agents, or any component of the formulation.	Diagnosis. For oropharyngeal candidiasis, must have at least a 2 week trial of fluconazole with an insufficient response, intolerable side effect, or have a contraindication.			12 months		0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
PRAMLINTIDE (SYMLIN)	1 - All FDA-approved Indications.			Diagnosis of Type 1 or Type 2 Diabetes Mellitus. Documentation the member uses mealtime insulin and has failed to achieve desired glycemic control despite optimal insulin therapy. Initial A1C greater than or equal to 6.5.			12 months	For reauth: if the patient has been receiving Symlin for at least 3 months, patient demonstrated a reduction in HbA1c since starting therapy with Symlin.	0
PREGABALIN (LYRICA)	1 - All FDA-approved Indications.			Diagnosis. For fibromyalgia: must have trial and failure or contraindication to gabapentin at a dose of at least 1200mg/day or maximally tolerated dose in intolerant patients AND either duloxetine or muscle relaxant unless contraindicated. For PHN: must have trial and failure, intolerance, or contraindication to gabapentin. For DPN: must have documented pharmacy claim history or prior therapy with a diabetic medication OR a medical/lab claim or physician chart note of diabetes diagnosis and must have trial and failure, intolerance, or contraindication to gabapentin.	For partial onset seizures, coverage is provided for members 1 month of age and older. For fibromyalgia, PHN, DPN, and neuropathic pain associated with spinal cord injury, coverage is provided for members 18 years of age or older.		12 months		0
PROTRIPTYLINE (VIVACTIL)	1 - All FDA-approved Indications.			Diagnosis. Requests for enrollees aged 65 or older must document intolerance to or clinical failure of 2 alternatives such as an SSRI (except paroxetine), SNRI, bupropion, trazodone or mirtazapine for depression.	Prior authorization only applies to enrollees aged 65 or older. All enrollees less than age 65 are not subject to prior authorization.		12 months		0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
PURIFIED CORTROPHIN GEL	1 - All FDA-approved		Members with scleroderma,	Diagnosis. For acute		Must be prescribed by or in	1 month	For allergic states such as	1
CORTICOTROPIN) INJECTION	Indications.		osteoporosis, systemic fungal	exacerbation of multiple		consultation with a neurologist		serum sickness or transfusion	
			infections, ocular herpes	sclerosis, member must have		or physician that specializes in		reaction due to serum protein	
			simplex, recent surgery,	tried and failed or have a		the treatment of multiple		reaction, member must have	
			history of or the presence of a	contraindication to 2		sclerosis, a rheumatologist,		tried and failed 2	
			peptic ulcer, congestive heart	corticosteroids (e.g. IV		allergist, dermatologist,		corticosteroids (e.g. IV	
			failure, hypertension, or	methylprednisolone, IV		immunologist,		methylprednisolone, IV	
			sensitivity to proteins derived	dexamethasone, or high dose		ophthalmologist,		dexamethasone, or high dose	
			from porcine sources, primary	oral steroids). Must have		pulmonologist, nephrologist		oral steroids) or has a	
			adrenocortical insufficiency or	documentation or claims				contraindication to	
			adrenocortical hyperfunction	verifying the member is on a				corticosteroid therapy. If the	
			are excluded.	medication for the treatment				member has a diagnosis of	
				of multiple sclerosis. For RA				atopic dermatitis, the member	
				(incl. Juvenile RA), psoriatic				is concurrently receiving	
				arthritis, ankylosing				maintenance therapy with one	
				spondylitis, acute gouty				(1) of the following, or is	
				arthritis: must be using as				contraindicated to all: topical	
				adjunctive therapy for short-				corticosteroid, topical	
				term administration (to tide				calcineurin inhibitor (e.g.,	
				over an acute episode or				tacrolimus, pimecrolimus),	
				exacerbation) and have a trial				topical PDE-4 inhibitor or	
				of 2 IV steroids w/ inadeq				Dupixent (dupilumab). For a	
				response or signif side				diagnosis of serum sickness,	
				effects/toxicity. The member				must provide laboratory	
				is concurrently receiving				documentation demonstrating	
				maintenance therapy with at				neutropenia, development of	
				least one of the following: an				reactive plasmacytoid	
				NSAID, DMARD (e.g.				lymphocytes, and elevated	
				methotrexate, leflunomide,				erythrocyte sedimentation	
				sulfasalazine) or biologic (e.g.				rate or C-reactive protein. For	
				adalimumab, etanercept,				ophthalmic diseases such as	
				infliximab, tofacitinib). For				severe acute and chronic	
				collagen disease, member				allergic and inflammatory	
			1	must have tried and failed or			1	processes involving the eve	1

Group Indi								
	dication Indicator	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
RIFAXIMIN (XIFAXAN) 1 - A	All FDA-approved dications. All FDA-approved dications.	Exclusion Criteria Members with decompensated cirrhosis.	Diagnosis. Medication must be used in conjunction with diet and exercise for the treatment of adults with noncirrhotic nonalcoholic steatohepatitis (NASH) with moderate to advanced liver fibrosis (consistent with stages F2 to F3 fibrosis). Must have a confirmed diagnosis of NASH by one of the following: a liver biopsy within the past 6 months confirming steatosis AND ALL of the following: NAFLD Activity Score (NAS) of at least 4, a score of at least 1 in each NAS component including steatosis, ballooning degeneration, lobular inflammation] OR the member has had ONE of the following imaging examps performed with 3 months prior to treatment including vibration-controlled transient elastography (VCTE, e.g. FibroScan) with kPa greater than or equal to 280 dB.m-1 OR computed tomography OR MRI. The member has an MRI-DECT secator than or causal to Diagnosis. For hepatic encephalopathy: must have	Coverage is provided for members 18 years of age and older. Hepatic encephalopathy and IBS-D: 18 years of age or older, Travelers diarrhea: 12 years of age or older	By or in consultation with a hepatologist or gastroenterologist. Hepatic encephalopathy: by or in consultation with a	Hepatic encephalopathy: 12 months, IBS-D: 2 weeks, Travelers diarrhea: 3 days	For IBS-D: members who experience a recurrence of symptoms can be retreated up to two times with the requested medical members who experience a recurrence of symptoms can be retreated up to two times with the same dosage regimen. Reauth for IBS-D: must have documentation from prescriber indicating recurrence of IBS-D symptoms after a successful treatment with rifaximin.	Part B Prerequisite 0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	<u> </u>	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
RILONACEPT (ARCALYST)	1 - All FDA-approved Indications.			documented genetic mutation in the Cold-Induced Auto- inflammatory Syndrome 1	children age 12 years and older. For DIRA: adults and pediatric members weighing 10kg or more.	By or in consultation with a hematologist, dermatologist, rheumatologist, neurologist, allergist, immunologist, cardiologist or a genetic specialist	12 months	For reauth: documentation from prescriber indicating stabilization or improvement in condition.	
RIOCIGUAT (ADEMPAS)	1 - All FDA-approved Indications.		Coverage will not be provided for patients taking nitrates (nitrates in any form) or a PDE inhibitor (e.g. sildenafil).	Diagnosis. Pulmonary arterial hypertension (PAH) WHO Group I confirmed by chart documentation of right-heart catheterization (RHC) indicating a mean pulmonary arterial pressure greater than 20 mmHg, pulmonary vascular resistance greater than 2 wood units, and mean pulmonary capillary wedge pressure less than or equal to 15 mmHg. If provider indicates RHC is not recommended, must have documentation of echocardiography.		Prescribed by or in consultation with cardiologist or pulmonologist.	Initial: 3 months, Reauth: 12 months	For reauth: documentation from prescriber that demonstrates member is tolerating and receiving clinical benefit from treatment	0

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Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information		Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
RISANKIZUMAB-RZAA	1 - All FDA-approved			Diagnosis. For plaque	Member must be 18 years of	By or in consultation with a	12 months	For reauthorization: must have	0
SKYRIZI)	Indications.			psoriasis: minimum BSA	age or older.	rheumatologist, dermatologist		documentation from	
				involvement of at least 3%		or gastroenterologist.		prescriber indicating	
				(not required if on palms,				stabilization or improvement	
				soles, head/neck, genitalia), a				in condition.	
				history of trial and failure of					
				ONE of the following: 1)					
				topical therapy (e.g.					
				corticosteroid, calcineurin					
				inhibitor, vitamin D analog), 2)					
				phototherapy, 3) systemic					
				treatment (e.g. methotrexate,					
				cyclosporine, oral retinoids).					
				For psoriatic arthritis (PsA),					
				one of the following: 1)					
				members with axial or					
				enthesitis must have a history					
				of trial and failure,					
				contraindication, or					
				intolerance to a 4 week trial of					
				2 NSAIDs, 2) the member has					
				severe disease as defined by					
				the prescriber, 3) members					
				with peripheral disease must					
				have a history of a trial and					
				failure, contraindication, or					
				intolerance to a 12 week trial					
				with methotrexate or another					
				DMARD. For Crohn's (CD) and					
				ulcerative colitis (UC): history					
				of trial and failure,					
				contraindication, or					
				intolerance to 2 of the					
RISDIPLAM (EVRYSDI)	1 - All FDA-approved		Coverage will be not be	Confirmed diagnosis fo 5q-		Prescribed by or in	12 months	For reauth: documentation	0
	Indications.		provided to members who are			consultation with neurologist,		that the patient is responding	
	marcacions.		concomitantly taking	Baseline assessment motor		or pediatric neurologist.		to the medication as	
			nusinersen.	milestone score from ONE of		or pediatric fledrologist.		demonstrated by clinically	
			nusinersen.	the following assessments:				significant improvement or	
				Hammersmith Functional				maintenance of function from	
				Motor Scale Expanded				pretreatment baseline status	
				(HFMSE), Hammersmith Infant				using the same exam as	
				Neurologic Exam (HINE),				performed at baseline	
				Upper limb module (ULM)				assessment (progression,	
				score, Children?s Hospital of				stabilization, or decreased	
				Philadelphia Infant Test of				decline in motor function).	
				Neuromuscular Disorders					
	1			(CHOP INTEND), or Six-minute	1				
	1			walk test.	1				
	1			1	1				
	1			1	1				
ISPERIDONE INJECTION	1 - All FDA-approved			Diagnosis. Documentation of	Coverage provided for		12 months		0
RISPERDAL CONSTA AND	Indications.			prior oral risperidone therapy.	members age 18 years and				
ERSERIS)	i i	1	ı		older.	1	1	I	

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Ago Bostriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
Group ROFLUMILAST (DALIRESP)	1 - All FDA-approved Indications.	Off-Label Uses	Moderate to sever liver impairment	Required Medical Information Diagnosis of GOLD Stage III or IV COPD associated with chronic bronchitis. Documentation of COPD exacerbation within the past year. Must have a trial and failure of an inhaled long- acting beta-agonist or inhaled long-acting anticholinergic. Must be used as add on therapy with a long-acting beta agonist or long-acting anti-muscarinic. Must have trial and failure of inhaled glucocorticosteroid or a contraindication to these agents.	Age Restriction	Prescriber Restriction	Coverage Duration 12 months	For reauthorization must have documentation from prescriber indicating improvement in condition.	•
ROTIGOTINE (NEUPRO)	1 - All FDA-approved Indications.			Diagnosis. For Parkinson's disease and primary restless legs syndrome: must have trial and failure, contraindication, or intolerance to both pramipexole and ropinirole.	Coverage is provided for members 18 years of age or older.		12 months	For reauthorization: documentation from prescriber indicating stabilization or improvement in condition.	0
RUFINAMIDE (BANZEL)	1 - All FDA-approved Indications.		Not covered for patients with Familial Short QT Syndrome	inadequate response or	Coverage is provided for members 1 year of age or older.	By or in consultation with a neurologist.	12 months		0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
RUXOLITINIB (JAKAFI)	1 - All FDA-approved Indications.			Diagnosis. Intermediate or high-risk myelofibrosis includes primary myelofibrosis, post-polycythemia vera myelofibrosis, and post-essential thrombocythemia myelofibrosis. For Polycythemia vera, must have trial and failure, intolerance, or contraindication of hydroxyurea. For acute Graft versus host disease (aGVHD), must have a trial and failure, intolerance, or contraindication to corticosteroids. For chronic Graft versus host disease (cGVHD), must have a trial and failure of at least two prior lines of systemic therapy.	GVHD: age 12 years or older All Others: age 18 years or older	By or in consultation with an oncologist, hematologist, or transplant specialist	6 months	For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	
SAPROTERIN DIHYDROCHLORIDE (KUVAN)	1 - All FDA-approved Indications.			Diagnosis. For treatment of Hyperphenylalaninemia. Clinically diagnosed with hyperphenylalaninemia due to tetrahydrobiopterin responsive phenylketonuria. Phe levels must be greater than 6 mg/dL (360 micromol/L).			Initial: 3 months, Reauth: 12 months	For reauthorization, must maintain Phe levels below member's baseline levels.	0
SATRALIZUMAB-MWGE (ENSPRYNG)	1 - All FDA-approved Indications.		Active hepatitis B infection, active or untreated latent tuberculosis	For Neuromyelitis Optica Spectrum Disorder (NMOSD): positive test for AQP4-IgG antibodies. At least 1 relapse in the last 12 months or 2 relapses in the last 24 months that required rescue therapy. Expanded Disability Status Scale (EDSS) score less than or equal to 6.5. Must have documentation of inadequate response, contraindication or intolerance to an immunosuppressant (e.g., mycophenolate mofetil, azathioprine, methotrexate) or rituximab.	older	By or in consultation with a neurologist or ophthalmologist	12 months	Part B before Part D Step Therapy. For reauth: documentation of stabilization or improvement in condition	1

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
SELEXIPAG (UPTRAVI)	1 - All FDA-approved Indications.			Diagnosis. Pulmonary arterial hypertension (PAH) WHO Group confirmed by chart documentation of right-heart catheterization (RHC) indicating a mean pulmonary arterial pressure greater than 20 mmHg, pulmonary vascular resistance greater than 2 wood units, and mean pulmonary capillary wedge pressure less than or equal to 15 mmHg. If provider indicates RHC is not recommended, must have documentation of an echocardiography.		Prescribed by or in consultation with cardiologist or pulmonologist.	Initial authorization: 3 months Reauthorization: 12 months	Reauthorization: documentation from prescriber that demonstrates member is tolerating and receiving clinical benefit from treatment	0
SILDENAFIL CITRATE (REVATIO)	1 - All FDA-approved Indications.		Coverage will not be provided for patients taking nitrates (nitrates in any form) or a guanylate cyclase stimulator (e.g. Adempas).	Diagnosis. Pulmonary arterial hypertension (PAH) WHO Group I confirmed by chart documentation of right-heart catheterization (RHC) indicating a mean pulmonary arterial pressure greater than 20 mmHg, pulmonary vascular resistance greater than 2 wood units, and mean pulmonary capillary wedge pressure less than or equal to 15 mmHg. If provider indicates RHC is not recommended, must have documentation of echocardiography.		Prescribed by or in consultation with a pulmonologist or cardiologist	Initial: 3 months, Reauth: 12 months	For reauth: documentation from prescriber that demonstrates member is tolerating and receiving clinical benefit from treatment	0
SIPONIMOD (MAYZENT)	1 - All FDA-approved Indications.		Coverage is not provided if the member has a CYP2C9*3/*3 genotype, presence of Mobitz type II second-degree, third-degree AV block, or sick sinus syndrome, unless patient has a functioning pacemaker or in the last 6 months, experienced myocardial infarction, unstable angina, stroke, TIA, decompensated heart failure requiring hospitalization, or Class III/IV heart failure.	Documentation that functional status is preserved and patient	Coverage is provided for members 18 years of age or older.	By or in consultation with a neurologist	12 months	For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
SODIUM OXYBATE (XYREM)	1 - All FDA-approved Indications.	Un-Lauer Uses	Exclusion Cheria	Diagnosis. For excessive daytime sleepiness associated with narcolepsy: a sleep study (e.g. polysomnogram, multiple sleep latency Test) confirming diagnosis. For cataplexy associated with narcolepsy: a sleep study confirming the diagnosis.	Coverage is provided for members 7 years of age or older	By or in consultation with a	Initial: 3 months, Reauthorization: 12 months	Reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	0
SODIUM PHENYLBUTYRATE	1 - All FDA-approved Indications.			Diagnosis.		By or in consultation with physician who specializes in the treatment of inherited metabolic disorders, a hematologist or a nephrologist.	12 months		0
SOFOSBUVIR-VELPATASVIR (EPCLUSA)	1 - All FDA-approved Indications.			Criteria will be applied consistent with current AASLD/IDSA guidance and/or FDA approved labeling	Coverage is provided for members who are age- appropriate according to AASLD/IDSA guidance and/or FDA-approved labeling.	By or in consultation with a gastroenterologist, hepatologist, infectious disease, HIV or transplant specialist.	Criteria will be applied consistent with current AASLD/IDSA guidance and/or FDA approved labeling		0
SOFOSBUVIR-VELPATASVIR- VOXILAPREVIR (VOSEVI)	1 - All FDA-approved Indications.		Coadministration with rifampin	Criteria will be applied consistent with current AASLD/IDSA guidance and/or FDA approved labeling	Coverage is provided for members who are age- appropriate according to AASLD/IDSA guidance and/or FDA-approved labeling.	By or in consultation with a gastroenterologist, hepatologist, infectious disease, HIV or transplant specialist.	Criteria will be applied consistent with current AASLD/IDSA guidance and/or FDA approved labeling		0

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Group SOMATROPIN (GENOTROPIN)	3 - All Medically-accepted	Off-Label Uses	Coverage will not be provided	Diagnosis. Growth chart	Age Restriction	By or in consultation with an	Coverage Duration 6 months	For reauth for pediatric GHD,	Part B Prerequisite
SOMATROPIN (GENOTROPIN)	Indications.		for members with active	required for all diagnoses		endocrinologist or	o months	Turner and Noonan	U
	marcacions.		malignancy, active	except Adult Growth Hormone		neonatologist.		syndromes, SGA, Prader-Willi	
			proliferative or severe non-	Deficiency (GHD).		l l l l l l l l l l l l l l l l l l l		syndrome, and ISS:	
			proliferative diabetic	Documentation that epiphyses				Documentation the patient	
			retinopathy, pediatric member					has open epiphyses. For	
			with closed epiphysis,	indications. For pediatric GHD:				reauth for adult GHD: current	
			members with Prader-Willi	a height greater than or equal				IGF-1 level is normal for age	
			who are severely obese or	to 2 standard deviations below				and gender (does not apply to	
			have severe respiratory	the mean for age and gender,				patients with structural	
			impairment.	documentation of growth				abnormality of the	
				velocity, skeletal maturation, 2				hypothalamus/pituitary and 3	
				provocative stimulation tests				or more pituitary hormone	
				which demonstrate GHD				deficiencies and childhood-	
				through peak growth hormone				onset growth hormone	
				concentrations less than 10				deficiency with congenital	
				ng/ml or IGF-1 or IGFBP-3				abnormality of the	
				levels or only one stim test is				hypothalamus/pituitary). For	
				needed in the presence of a				reauth for Prader Willi:	
				pituitary abnormality. For				documentation growth	
				Small for Gestational Age				hormone has resulted in an	
				(SGA), a height greater than or				increase in lean body mass or	
				equal to 2 standard deviations				decrease in fat mass.	
				below the mean for age and					
				gender, and EITHER a birth					
				weight less than 2500 g at a					
				gestational age greater than					
				37 weeks, OR weight or length					
				at birth greater than 2					
				standard deviations below the					
				mean for gestational age and					
				documentation that catch up					
				growth not achieved by age 2.					
SOTATERCEPT-CSRK	1 - All FDA-approved			Diagnosis. Pulmonary arterial		Prescribed by or in	Initial: 3 months, Reauth: 12	For reauth: documentation	0
(WINREVAIR)	Indications.			hypertension (PAH) WHO		consultation with cardiologist	months	from prescriber that	
(**************************************				Group I confirmed by chart		or pulmonologist		demonstrates member is	
				documentation of right-heart				tolerating and receiving	
				catheterization (RHC)				clinical benefit from treatment	
				indicating a mean pulmonary					
				arterial pressure greater than					
				20 mmHg, pulmonary vascular					
				resistance greater than 2					
				wood units, and mean					
				pulmonary capillary wedge					
				pressure less than or equal to					
				15 mmHg. If provider indicates					
				RHC is not recommended,					
				must have documentation of					
				echocardiography. Must be					
				used in combination with					
1				standard of care therapy (e.g.					
				ERA or PDE-5 inhibitor)					
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Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information		Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
SPARSENTAN (FILSPARI)	1 - All FDA-approved Indications.			Diagnosis of primary immunoglobulin A nephropathy (IgAN) that has been confirmed by biopsy. Must have an eGFR rate of at least 30 ml/min/1.73m^2. Must have a total urine protein of at least 1.0 g/day. Must be at risk of rapid disease progression defined as having a urine protein-to-creatinine ratio (UPCR) of at least 1.5 g/g. Must have tried and failed a stable and maximum tolerated dose of an ACE inhibitor or ARB.	Coverage is provided for members 18 years of age or older.	By or in consultation with a nephrologist.	Initial: 6 months. Reauth: 12 months	For reauth: must have a decrease from baseline in total urine protein or UPCR.	0
SPESOLIMAB-SBZO (SPEVIGO)	1 - All FDA-approved Indications.			Diagnosis. For treatment of a generalized pustular psoriasis (GPP) flare, must have a moderate-to-severe flare defined by ALL of the following: 1) GPPGA total score greater than or equal to 3 (moderate or severe), 2) presence of fresh pustules, 3) GPPGA postulation subscore of at least 2 (mild, moderate, or severe), and 4) at least 5% BSA covered with erythema and presence of pustules. For treatment of GPP when not experiencing a flare, must have a history of at least 2 moderate or severe GPP flares in the past and must have a history of flaring while on systemic treatment or upon reduction or discontinuation of systemic therapy for GPP (e.g. retinoids, methotrexate, cyclosporine).	Coverage is provided for members 12 years of age or older and weighing at least 40 kg.	By or in consultation with a dermatologist	For a flare: one treatment course (up to 2 infusions over 2 weeks). For maintenance: 12 months	For reauth: documentation of reduction in the frequency of flares while on treatment	0
STIRIPENTOL (DIACOMIT)	1 - All FDA-approved Indications.			Diagnosis. Must have had an inadequate response or intolerance to two generic antiepileptic drugs (e.g. valproate, topiramate, clobazam). Must be using in combination with clobazam.	Member must be 6 months of age or older	By or in consultation with a neurologist	12 months		0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
IDALAFIL (ADCIRCA)	1 - All FDA-approved Indications.		Coverage will not be provided for patients taking nitrates (nitrates in any form) or a guanylate cyclase stimulator (e.g. Adempas).	Diagnosis. Pulmonary arterial hypertension (PAH) WHO Group I confirmed by chart documentation of right-heart catheterization (RHC) indicating a mean pulmonary arterial pressure greater than 20 mmHg, pulmonary vascular resistance greater than 2 wood units, and mean pulmonary capillary wedge pressure less than or equal to 15 mmHg. If provider indicates RHC is not recommended, must have documentation of echocardiography.		Prescribed by or in consultation with a pulmonologist or cardiologist	Initial: 3 months, Reauth: 12 months	For reauth: documentation from prescriber that demonstrates member is tolerating and receiving clinical benefit from treatment	0
ASIMELTEON (HETLIOZ)	1 - All FDA-approved Indications.			Diagnosis. Must submit chart documentation describing how diagnosis was confirmed (e.g. sleep-wake logs, melatonin secretion abnormalities, or progress notes, etc.)	Coverage is provided for members 3 years of age or older.	By or in consultation with a neurologist or a physician who specializes in sleep medicine	12 months	For Reauth: documentation from prescriber indicating stabilization or improvement in condition.	0
EDUGLUTIDE (GATTEX)	1 - All FDA-approved Indications.		Active intestinal obstruction or active gastrointestinal malignancy.	Diagnosis. For diagnosis of short bowel syndrome, member must be receiving parenteral support.		By or in consultation with a gastroenterologists	12 months	For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	0
ELOTRISTAT (XERMELO)	1 - All FDA-approved Indications.			Diagnosis.	Coverage is provided for members 18 years of age and older.	By or in consultation with an oncologist	6 months	For reauth: documentation of improvement or stabilization.	0
ENOFOVIR ALAFENAMIDE VEMLIDY)	1 - All FDA-approved Indications.		0	Diagnosis. Member must have chronic hepatitis B virus (HBV) infection.	Coverage is provided for members 18 years of age or older.	By or in consultation with a gastroenterologist, hepatologist, infectious disease, HIV or transplant specialist.	12 months	For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
•	1 - All FDA-approved Indications.	Un-Label Uses	Uncontrolled depression, actively suicidal. Currently using a monoamine oxidase inhibitor or reserpine. Hepatic impairment. Concurrent use of deutetrabenazine or valbenazine.	Diagnosis. Must have confirmed Huntington's disease either by Huntington Disease Mutation analysis	Coverage is provided for members 18 years of age or older.	Prescriber Restriction By or in consultation with a neurologist	12 months	Other Criteria Maximum dose approved is 100mg/day. For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	O 0
TOFACITINIB (XELJANZ)	1 - All FDA-approved Indications.			of trial and failure, contraindication, or intolerance to a TNF blocker.	For Polyarticular course juvenile idiopathic arthritis: Coverage is provided for members 2 years of age and older. For all other diagnoses coverage is provided for members 18 years of age and older	By or in consultation with dermatologist, rheumatologist or gastroenterologist.	12 months	Reauth: Documentation from the prescriber indicating stabilization or improvement in condition.	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
ILVAPTAN (JYNARQUE)	1 - All FDA-approved Indications.		History of significant liver impairment or injury (not including uncomplicated polycystic liver disease), concomitant use of strong CYP3A inhibitors, uncorrected abnormal blood sodium concentrations, unable to sense or respond to thirst, hypovolemia, uncorrected urinary outflow obstruction, anuria	Diagnosis. Must meet one of the following criteria defining risk of rapidly progressing disease: (1) age 55 or younger and eGFR between 25 and 65 mL/min/1.73m^2, (2) age 56 to 65 and eGFR between 25 and 44 mL/min/1.73m^2 plus eGFR decline of greater than 2.0 mL/min/1.73m^2/year, (3) estimated CrCl greater than or equal to 60 mL/min, total kidney volume greater than or equal to 750 mL, and less than 51 years of age, (4) Mayo classification of 1C, 1D, or 1E disease.	Member must be 18 years of age or older	By or in consultation with a nephrologist	12 months	For reauthorization: documentation from prescriber indicating stabilization or improvement in condition.	0
ENTINE HCL (SYPRINE)	1 - All FDA-approved Indications.			Diagnosis. Must have a trial of penicillamine (Depen) with an inadequate response or significant side effects/toxicity or must have a contraindication to this therapy.		By or in consultation with a gastroenterologist, an ophthalmologist or a physician who specializes in the treatment of inherited metabolic disorders	12 months	For reauth: must have documentation from prescriber indicating improvement in condition.	0
IMIPRAMINE	1 - All FDA-approved Indications.			Diagnosis. Requests for enrollees aged 65 or older must document intolerance to or clinical failure of 2 alternatives such as an SSRI (except paroxetine), SNRI, bupropion, trazodone or mirtazapine for depression.	Prior authorization only applies to enrollees aged 65 or older. All enrollees less than age 65 are not subject to prior authorization.		12 months		0
OFINETIDE (DAYBUE)	1 - All FDA-approved Indications.		0	Diagnosis. Documentation of a diagnosis of typical Rett syndrome according to the Rett Syndrome Diagnostic Criteria with a documented disease-causing mutation in the MECP2 gene.	Coverage is provided for members 2 years of age or older.	By or in consultation with a pediatric neurologist or neurologist	12 months	0	0
BROGEPANT (UBRELVY)	1 - All FDA-approved Indications.			Diagnosis. Must have a history of trial and failure, contraindication, or intolerance to at least one triptan.	Coverage is provided for members 18 years of age and older.		12 months	For reauth: documentation of improvement or stabilization.	0

Curana	Indication Indicates	Off Label Harr	Fundamina Cultural	Demoised Madical Information	Ann Dostriction	Dunnaulh au Dant-d-d	Courses Durent's	Ohbar Criharia	Dank D. Duamano Initia
Group UPADACITINIB (RINVOQ)	Indication Indicator 1 - All FDA-approved Indications.	Off-Label Uses	Exclusion Criteria	Required Medical Information Diagnosis. For rheumatoid arthritis (RA), psoriatic arthritis (PsA), ankylosing spondylitis (AS), non- radiographic axial spondyloarthritis (nr-axSpA), polyarticular juvenile idiopathic arthritis (pJIA), ulcerative colitis (UC), and Crohn's disease: history of trial and failure, contraindication, or intolerance to a TNF blocker. For atopic dermatitis (AD): history of trial and failure, contraindication, or intolerance to 2 systemic products (immunosuppressant or biologic).	For PsA and pJIA, must be 2 years of age or older, For AD: must be 12 years of age or	Prescriber Restriction By or in consultation with a rheumatologist, dermatologist, or gastroenterologist.	Coverage Duration 12 months	Other Criteria For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	Part B Prerequisite 0
USTEKINUMAB (STELARA) SQ	1 - All FDA-approved Indications.			Diagnosis. For Psoriatic arthritis (PsA): for mild to moderate axial or enthesitis, must have a history of trial and failure, contraindication, or intolerance to a 4 week trial of 2 NSAIDs. For members with mild to moderate peripheral disease, must have a history of a trial and failure, contraindication, or intolerance to a 12 week trial with methotrexate or another DMARD. For plaque psoriasis (PsO): minimum BSA involvement of at least 3% (not required if on palms, soles, head/neck, genitalia), a history of trial and failure of ONE of the following: 1) topical therapy (e.g. corticosteroid, calcineurin inhibitor, vitamin D analog), 2 phototherapy, 3) systemic treatment (e.g. methotrexate, cyclosporine, oral retinoids). For Crohn's disease (CD): history of trial and failure, contraindication, or intolerance to 2 of the following therapy options: aminosalicylates, corticosteroids, or	Must be 6 years of age or older.	By or in consultation with a rheumatologist, gastroenterologist, or dermatologist.	12 months.	For reauth: must have documentation from the prescriber indicating stabilization or improvement in condition.	0
V-GO KIT	1 - All FDA-approved Indications.			Must have documentation of previous insulin use.			12 months		0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information		Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
/ALBENAZINE (INGREZZA)	1 - All FDA-approved			Diagnosis. For chorea: must	Coverage is provided for	By or in consultation with a	12 months	For reauthorization: must	0
	Indications.			have confirmed Huntington's	members 18 years of age or	neurologist or psychiatrist		have documentation from	
				disease either by Huntington	older			prescriber indicating	
				Disease Mutation analysis				stabilization or improvement	
				(with laboratory result				in condition.	
				indicating expanded CAG					
				repeat of greater than or					
				equal to 36 in the Huntington					
				gene) or a positive family					
				history of Huntington's					
				Disease with autosomal					
				dominant inheritance pattern,					
				must have clinical signs of					
				Huntington's Disease including					
				chart documentation of a					
				clinical work-up showing one					
				or more of the following signs:					
				motor (e.g. finger tapping,					
				rigidity), oculomotor, bulbar					
				(e.g. dysarthria, dysphagia),					
				affective (e.g. depression),					
				cognitive. Must have chart					
				documentation of chorea. For					
				Tardive Dyskinesia: member					
				has been diagnosed with					
				moderate to severe tardive					
				dyskinesia (TD) according to					
				the DSM V criteria including					
				involuntary athetoid or					
				*					
				choreiform movements and					
				has a history of treatment with					
				neuroleptic agent (i.e.					
				antipsychotic). Adjustments to					
/ERICIGUAT (VERQUVO)	1 - All FDA-approved			Diagnosis. Must have a left		Prescribed by or in	12 months	Reauthorization:	0
, ,,	Indications.			ventricular ejection fraction		consultation with cardiologist.		documentation from	
				(LVEF) less than or equal to				prescriber indicating	
				45%. Must have had a				stabilization or improvement	
				hospitalization for heart				in condition.	
				failure within the past 6					
				months or received outpatient					
				IV diuretics within the past 3					
				months. Documentation the					
				member is currently taking or					
				has had prior treatment with					
				an angiotensin-converting					
				enzyme inhibitor, angiotensin					
				II receptor blocker or Entresto					
				and a beta blocker.					
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Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
GABATRIN (SABRIL)	1 - All FDA-approved Indications.			Diagnosis. Must undergo vision testing prior to beginning treatment. For Refractory Complex Partial Seizures: must have inadequate response to at least two of the following anticonvulsant drugs: levetiracetam, phenytoin, carbamazepine, oxcarbazepine, gabapentin, lamotrigine, valproate, or topiramate. Must be using vigabatrin in combination with at least one other anticonvulsant medication (which can include medication from trial above).	Coverage is provided for members 1 month of age or older.	By or in consultation with a neurologist.	12 months		0
LAZODONE (VIIBRYD)	1 - All FDA-approved Indications.			Diagnosis. Documentation of trial and failure of at least two generic antidepressants alternatives such as an SSRI, SNRI, bupropion, trazodone or mirtazapine	Coverage is provided for members 18 years of age and older.		12 months		0
ORICONAZOLE INJECTION VFEND)	1 - All FDA-approved Indications.			Diagnosis.	2 years of age or older	Prescribed by or in consultation with an infectious disease specialist	12 months		0
ORTIOXETINE (TRINTELLIX)	1 - All FDA-approved Indications.			Diagnosis. Documentation of trial and failure of at least two generic antidepressants alternatives such as an SSRI, SNRI, bupropion, trazodone or mirtazapine	Coverage is provided for members 18 years of age and older.		12 months		0
OSORITIDE (VOXZOGO)	1 - All FDA-approved Indications.			Diagnosis. Documentation the member has open epiphyses.	Coverage is provided for members 5 years of age and older.		12 Months	For reauth: documentation of both of the following: 1. improvement or stabilization. 2. The member's epiphyses remain open.	0

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
VOXELOTOR (OXBRYTA)	1 - All FDA-approved Indications.				members 4 years of age and		Initial: 6 months Reauthorization: 12 months	For reauthorization: Documentation there has been a reduction in vaso- occlusive events or an improvement in condition.	0
ZURANOLONE (ZURZUVAE)	1 - All FDA-approved Indications.			depression (PPD) with onset	members 18 years of age and		14 days		0