

Prior Authorization Detail

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				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	-	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
ABOBOTULINUMTOXINA	1 - All FDA-approved	011 20001 0000	Exclusion enteria	Diagnosis.	Age Nestriction	Treseringer reserved	12 months	For reauthorization:	n o
(DYSPORT)	Indications.			Diagnosis.			12 1110111113	documentation from	Ĭ
	maleations.							prescriber indicating	
								stabilization or improvement	
								in condition.	
ACITRETIN (SORIATANE)	1 - All FDA-approved			Diagnosis. Must have a trial of			12 months	in condition.	0
ACTIVETIIN (SONIATANE)	Indications.			methotrexate or cyclosporine			12 1110111113		ľ
	malcations.			with inadequate response or					
				significant side effect/toxicity					
				or have a contraindication to					
				these therapies.					
				these therapies.					
ADALIMUMAB (HUMIRA)	3 - All Medically-accepted		Coverage is not provided for	Diagnosis. For rheumatoid	Member must be 2 years of	By or in consultation with a	12 months	For hidradenitis suppurativa	0
, ,	Indications.		use of once weekly doses of	arthritis (RA): history of trial	age or older.	rheumatologist,		(HS): moderate to severe	
			Humira in combination with	and failure, contraindication,		gastroenterologist,		disease with 3 active	
			methotrexate.	or intolerance to a 3 month		ophthalmologist, or		abscesses, inflammatory	
				trial with methotrexate or		dermatologist.		nodules, or lesions. For	
				another DMARD. For juvenile				uveitis: trial of a corticosteroid	1
				idiopathic arthritis (JIA) with				or immunomodulator with	
				polyarthritis: history of trial				inadequate response or side	
				and failure, contraindication,				effects/toxicities unless	
				or intolerance to a 3 month				contraindicated. For reauth:	
				trial with methotrexate,				must have documentation	
				leflunomide, or sulfasalazine.				from prescriber indicating	
				For JIA with oligoarthritis,				stabilization or improvement	
				enthesitis and/or sacroiliitis:				in condition.	
				history of trial and failure,					
				contraindication, or					
				intolerance to at least a 4					
				week trial of 2 different					
				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					

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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:	
				70mg/dL (w/ ASCVD), or				must have chart	
				55mg/dl if has extreme risk				documentation confirming	
				designation (see Other				history of at least one of the	
				Criteria). Must have failed to				following: myocardial	
				achieve goal LDL-C reduction				infarction or other acute	
				after a trial of a high intensity				coronary syndromes	
				statin (atorvastatin 40-80mg				(including ST-elevation	
				daily or rosuvastatin 20-40mg				myocardial infarction, non-ST	
				daily) OR 2 moderate-				elevation myocardial	
				intensity statins (atorvastatin				infarction, and unstable	
				or rosuvastatin) at the				angina), coronary or other	
				member's maximally tolerated				revascularization procedure,	
				dose OR documentation the				ischemic stroke or transient	
				member is determined to be				ischemic attack,	
				intolerant to statin therapy				atherosclerotic peripheral	
				with provider attestation of				arterial disease. For HeFH:	
				intolerance to statin therapy				must have chart	
				consisting of statin related				documentation of one of the	
				rhabdomyolysis or skeletal-				following: A score of greater	
				muscle related symptoms				than 8 using the Dutch Lipid	
ALOSETRON (LOTRONEX)	1 - All FDA-approved		Constipation. Concomitant	Diagnosis. Documentation of		By or in consultation with a	12 months		0
	Indications.		use of fluvoxamine. Male	chronic IBS symptoms	members 18 years of age and	Gastroenterologist		documentation from	
			gender. History of chronic or	diarrhea lasting at least 6	older.			prescriber indicating	
			severe constipation or	months. Gastrointestinal tract				stabilization or improvement	
			sequelae from constipation,	abnormalities have been ruled				in condition.	
			intestinal obstruction,	out. Must have trial of					
			stricture, toxic megacolon,	loperamide and dicyclomine					
			gastrointestinal perforation	used in the treatment of IBS-D					
			and/or adhesions, ischemic	with inadequate response or					
			colitis, impaired intestinal	significant side effects/toxicity					
			circulation, thrombophlebitis,	unless contraindicated					
			or hypercoagulable state,						
			Crohn's disease, ulcerative						
			colitis, diverticulitis, or severe						
ALPELISIB (VIJOICE)	1 - All FDA-approved		hepatic impairment.	Diagnosis of PIK3CA-Related	Coverage is provided for	By or in consultation with an	12 months	For reauthorization: must	0
ALFELISID (VIJUICE)	Indications.			Overgrowth Spectrum (PROS)		appropriate specialist	בב וווטוונווט	have documentation from	U
	mulcations.				older.	depending on the symptoms		prescriber indicating	
				Disease must be severe or life-		and part of the body that are		stabilization or improvement	
				threatening and require		affected.		in condition.	
				systemic treatment.		anecteu.		in condition.	
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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.	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.	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	
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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AMIKACIN INHALATION	1 - All FDA-approved			Diagnosis of Mycobacterium		By or in consultation with a	12 months	For reauth: must have	0
(ARIKAYCE)	Indications.			avium complex (MAC) lung		pulmonologist or infectious		attestation confirming	
				disease. Must be used as part		disease specialist		presence of a positive sputum	
				of a combination antibacterial				culture or that there have	
				drug regimen in patients who				been negative sputum	
				do not achieve negative				cultures for an insufficient	
				sputum cultures after a				period of time (e.g. less than	
				minimum of 6 consecutive				12 months).	
				months of a multidrug					
				background regimen therapy					
				containing at least 2 of the					
				following: a macrolide, a					
				rifamycin (rifampin or					
				rifabutin), and ethambutal.					
APREMILAST (OTEZLA)	1 - All FDA-approved			Diagnosis. For Psoriatic	Coverage is provided for	By or in consultation with a	12 months	For reauthorization: must	0
	Indications.			arthritis (PsA): for mild to	members 6 years of age or	dermatologist, rheumatologis	:	have documentation from	
				moderate axial or enthesitis,	older.			prescriber indicating	
				must have a history of trial				stabilization or improvement	
				and failure, contraindication,				in condition.	
				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: minimum BSA					
				involvement of at least 2% (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, skin lesions,					
				positive pathergy reaction,			<u> </u>		L

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ARIMOCLOMOL (MIPLYFFA)	1 - All FDA-approved				Member is 2 years of age and		12 months	Reauthorization:	0
	Indications.			diagnosis was confirmed by	older			Documentation the member	
				genetic testing demonstrating				is experiencing an	
				one of the following: 1. a				improvement or stabilization	
				mutation in both alleles of				in disease.	
				NPC1 or NPC2 OR 2. mutation					
				in one allele and either a positive filipin-staining or					
				elevated cholestance triol/oxysterols (greater than					
				2x ULN). Documentation the					
				member has at least one					
				neurological symptom of NPC					
				(e.g. decrease in motor skills,					
				ataxia, seizures, etc.). Must be					
				using in combination with					
				miglustat. Must not be used in					
				combination with Aqneursa.	1				
				combination with Aqueursa.					
				Diagnosis. Documentation the			12 months		0
SENSOR (ABILIFY MYCITE)	Indications.			member had at least a one-	members 18 years of age and				
				month trial of oral	older.				
				aripiprazole (Abilify) therapy.					
ARMODAFINIL (NUVIGIL)	1 - All FDA-approved			Diagnosis. Must have a history	′	By or in consultation with a	SWSD: 6 months. Narcolepsy,	For reauth: documentation of	0
	Indications.			of trial and failure,		sleep specialist, ENT (ear,	OSA: 12 months	improvement or stabilization.	
				contraindication, or		nose, and throat specialist),			
				intolerance to modafinil. For		neurologist, or pulmonologist			
				narcolepsy: Sleep Study (e.g.					
				Polysomnogram, Multiple					
				Sleep Latency Test)					
				confirming diagnosis. For					
				obstructive sleep apnea: Sleep					
				study (e.g. polysomnogram)					
				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					
	1			documentation of shift work				1	

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ATOGEPANT (QULIPTA)	1 - All FDA-approved Indications.			Diagnosis. For episodic migraine: Provider attestation the member has 4 to 14 headache days per month. For chronic migraine: Provider attestation 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.		Initial: 6 months Reauthorization: 12 months	For reauth: Provider attestation the member is having a reduced number of migraine/headache days per month or a decrease in migraine/headache severity. 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.	
	1 - All FDA-approved ndications.			Diagnosis of primary immunoglobulin A nephropathy (IgAN) that has been confirmed by biopsy. 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 both 1) an ACE inhibitor or ARB and 2) an SGLT-2 inhibitor (e.g. Farxiga).	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
	1 - All FDA-approved Indications.			Diagnosis of ANCA-associated vasculitis (GPA or MPA). Must be on concurrent therapy with glucocorticoids and immunosuppressants (e.g. cyclophosphamide, azathioprine, mycophenolate, rituximab).		By or in consultation with a rheumatologist, hematologist or oncologist.	12 Months	For reauthorization: documentation from prescriber indicating stabilization or improvement in condition.	0
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.	0
	3 - All Medically-accepted ndications.						NA		0

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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			3 months	For reauth: documentation of improvement or stabilization.	0
				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).					
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.	_	By or in consultation with a pulmonologist or infectious disease specialist	6 months		0
BELIMUMAB (BENLYSTA) (IV FORMULATION)	1 - All FDA-approved Indications.		Severe active central nervous system lupus. Combination therapy with other biologics or IV cyclophosphamide.			By or in consultation with a rheumatologist or hematologist	12 months	For reauth: documentation from the prescriber indicating stabilization or improvement in condition.	0

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BELIMUMAB (BENLYSTA) (SQ)	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 30 IU/ml to 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.	members 5 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
BELUMOSUDIL (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.		By or in consultation with an oncologist, hematologist, or transplant specialist	12 months	For reauth: documentation of improvement or stabilization.	0
BENRALIZUMAB (FASENRA)	1 - All FDA-approved Indications.			Diagnosis. For severe eosinophilic asthma: eosinophil blood count greater than or equal to 150cells/microliter. Documentation of inadequate response, intolerance, or contraindication to a highdose 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.	older.	By or in consultation with an allergist, immunologist, pulmonologist, or rheumatologist.	12 months	For reauth: documentation of improvement (e.g. reduced symptoms, reduced exacerbations, need for oral steroids).	0
BEREMAGENE 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.	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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BIRCH TRITERPENES	1 - All FDA-approved			Diagnosis of Dystrophic	Coverage is provided for	By or in consultation with a	6 months	Reauthorization: must have	0
(FILSUVEZ)	Indications.			Epidemolysis Bullosa (DEB) or	members 6 months of age or	dermatologist		documentation from	
				junctional epidermolysis	older.			prescriber indicating	
				bullosa (JEB) with an open				improvement in condition.	
				wound.					
BOSENTAN (TRACLEER)	1 - All FDA-approved		Pregnancy	Diagnosis. Pulmonary arterial		Prescribed by or in	Initial authorization: 3 months		0
	Indications.			hypertension (PAH) WHO		consultation with cardiologist	Reauthorization: 12 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	
				indicating a mean pulmonary				treatment	
				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.					
BUDESONIDE (EOHILIA)	1 - All FDA-approved		0	Diagnosis. For eosinophilic	Coverage is provided for	By or in consultation with an	3 months	Reauth: use beyond 3 months	0
	Indications.			esophagitis (EoE): must have	members 11 years of age or	allergist or gastroenterologist.		has not been studied.	
				at least 15 intraepithelial	older.				
				eosinophils per high-power					
				field (eos/hpf) following a					
				treatment course with a PPI.					
BUDESONIDE EXTENDED	1 - All FDA-approved			Diagnosis. Must have a trial	•	By or in consultation with a	8 weeks	For reauth: must have	0
RELEASE TABLETS (UCERIS)	Indications.			and failure, a	age or older.	rheumatologist or		documentation from	
				contraindication, or an		gastroenterologist.		prescriber indicating	
				intolerance to two (2) of the				stabilization or improvement	
				following therapy options:				in condition.	
				topical mesalamine, oral					
				aminosalicylate or					
				corticosteroids with					
				inadequate response or side					
				effects/toxicity unless					
				contraindicated.					

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BUROSUMAB-TWZA	1 - All FDA-approved		Use with oral phosphate or	Diagnosis. For X-linked		By or in consultation with a	12 months	Reauthorization:	0
(CRYSVITA)	Indications.		active vitamin D analogs	hypophosphatemia:		physician who is experienced		Documentation current	
				confirmation of the diagnosis		in the management of		(within the past 12 months)	
				by at least one of the		patients with metabolic bone		serum phosphorus level is not	
				following: A genetic test		disease.		above the upper limit of the	
				showing a PHEX gene				laboratory normal reference	
				mutation (phosphate				range and documentation the	
				regulating gene with				member has had a positive	
				homology to endopeptidase				clinical response or	
				on the X chromosome) or				stabilization in their disease.	
				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).					
BUT/APAP/CAF TAB	3 - All Medically-accepted			Diagnosis. This Prior	Coverage is provided for		12 months		0
	Indications.			Authorization requirement	members 12 years of age or				
				only applies to members	older.				
				when a non-FDA approved					
				diagnosis is submitted. FDA-					
				approved diagnosis codes					
				submitted will pay without					
				prior authorization					
				requirement.					
BUTAL/APAP TAB 50-325MG	3 - All Medically-accepted			Diagnosis. This Prior	Coverage is provided for		12 months		0
	Indications.			Authorization requirement	members 12 years of age or				
				only applies to members	older.				
				when a non-FDA approved					
				diagnosis is submitted. FDA-					
				approved diagnosis codes					
				submitted will pay without					
				prior authorization					
				requirement.					

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C1 ESTERASE INHIBITOR	1 - All FDA-approved			Diagnosis of HAE is confirmed		Prescribed by or in	Initial: 6 months	For reauth: must have	0
HAEGARDA)	Indications.			by laboratory values obtained	members 6 years of age or	consultation with an	Reauthorization: 12 months	documentation from	
,				on two separate instances	older.	allergist/immunologist,		prescriber indicating	
				(laboratory reports must		hematologist, dermatologist		improvement in condition.	
				contain reference ranges). For		The matologist, dermatologist		Improvement in condition.	
				Type I: Low C4 level and low					
				C1-INH antigenic level. For					
				Type II: Low C4 level and					
				normal or elevated C1-INH					
				antigenic level and low C1-INH					
				functional level. Must have					
				documentation of a previous					
				HAE attack in the absence of					
				hives or a medication known					
				to cause angioedema to					
				demonstrate member is					
				candidate for prophylactic					
				therapy. Member must not be					
				taking any medications that					
				may exacerbate HAE,					
				including angiotensin-					
				converting enzyme (ACE)					
				inhibitors, Tamoxifen, and					
				estrogen-containing					
				medications. Must be using as					
				prophylactic therapy for the					
				prevention of HAE attacks.					
				prevention of fixe attacks.					
									_
ANNABIDIOL (EPIDIOLEX)	1 - All FDA-approved			Diagnosis. Must have had an		'	12 months		0
	Indications.			inadequate response or	age or older	neurologist			
				intolerance to one generic					
				antiepileptic drug.					
ARGLUMIC ACID	1 - All FDA-approved			Diagnosis. This Prior			12 months		0
CARBAGLU)	Indications.			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.					
EFTAROLINE (TEFLARO)	1 - All FDA-approved			Diagnosis. For acute bacterial			14 days		0
	Indications.			skin and skin structure					
				infection (ABSSSI),					
				documentation of a history of					
				treatment failure with or					
				contraindication to					
				vancomycin.					
YSTEAMINE (CYSTAGON)	1 - All FDA-approved			Diagnosis. Must have		By or in consultation with a	Initial: 3 months	For reauth: must have	0
, ,	Indications.			documentation of CTNS gene		nephrologist or physician who		documentation from	
				mutation, elevated white		specializes in the treatment of		prescriber indicating	
									J
				blood cell cystine levels		inherited metabolic disorders		improvement in condition and	Ϊ
				greater than 2nmol per half-				a reduction in WBC cystine	
				cystine per mg of protein, or				levels since starting treatment	
				cystine corneal crystals by slit				with oral cysteamine	
				lamp examination.					
ALFAMPRIDINE (AMPYRA)	1 - All FDA-approved		History of seizure disorder,	Diagnosis of multiple sclerosis.		Neurologist	Initial: 3 months	For reauthorization: must	0
	Indications.		moderate to severe renal	Chart documentation of	members 18 years of age or		Reauthorization: 12 months	have documentation from	
			•	baseline motor disability or	older.			prescriber indicating	
			equal to 50 mL/min).	dysfunction.				stabilization or improvement	
	I		equal to 50 mil/min).	aystatiction.				in condition.	I

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
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.	0
DEFERASIROX (EXJADE)	1 - All FDA-approved Indications.		Glomerular Filtration Rate less than 40mL/min/1.73 m2. Concomitant advanced malignancy or high risk myelodysplastic syndrome. Platelet count less than 5000000000/L	Diagnosis. For chronic iron overload due to blood transfusions: pretreatment serum ferritin level is greater than 1000 mcg/L. For chronic iron overload due to nontransfusion-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		Prescribed by or in consultation with a hematologist	12 months	For reauth: documentation from prescriber indicating stabilization or improvement in condition.	0
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

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
DEUTETRABENAZINE	1 - All FDA-approved		Uncontrolled depression,	Diagnosis. For chorea: must	Coverage is provided for	By or in consultation with a	12 months	For reauthorization: must	0
(AUSTEDO)	Indications.		actively suicidal, hepatic	have confirmed Huntington's	members 18 years of age or	neurologist or psychiatrist		have documentation from	
			impairment, concurrent use		older.			prescriber indicating	
			with MAOI's, reserpine,	Disease Mutation analysis				stabilization or improvement	
			tetrabenazine, or valbenazine.	(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 (TD): must					
				have chart documentation of					
				involuntary athetoid or					
				choreiform movements and					
				has a history of treatment					
				with neuroleptic agent (i.e.					
				antipsychotic). Adjustments					
				to possible offending					
				medication such as dose					
				reduction or discontinuation					
DEUTIVACAFTOR/TEZACAFTO	'''			Diagnosis. Documentation of		By or in consultation with a	12 months	For reauthorization:	0
R/VANZACAFTOR (ALYFTREK)	Indications.			genetic test confirming the		cystic fibrosis specialist or		documentation indicating	
				member has at least one	older	pulmonologist		stabilization or improvement	
				F508del mutation or another				in condition.	
				responsive mutation in the					
				CFTR gene.					
DEXTROMETHORPHAN-	1 - All FDA-approved			Diagnosis. Pseudobulbar	Coverage is provided for	By or in consultation with	Initial: 3 months	For reauthorization:	0
QUINIDINE (NUEDEXTA)	Indications.			affect (PBA): documentation	members 18 years of age and	neurologist	Reauthorization: 12 months	Documentation indicating a	
				supporting the following:	older.			decrease in the number of	
				involuntary outbursts of				laughing and/or crying	
				laughing and/or crying that				episodes since starting the	
				are incongruent or				medication.	
				disproportionate to the					
				member's emotional state					
				AND other possible conditions					
				that could result in emotional					
				lability (e.g. depression,					
				bipolar disorder,					
				schizophrenia, epilepsy) have					
				been ruled out. Must have					
				underlying neurological					
				disorder such as amyotrophic					
				lateral sclerosis, multiple					
				sclerosis, Alzheimer's and					
				related diseases, Stroke,					
				Traumatic Brain Injury, or					
İ	1			Parkinsonian Syndrome.					1

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
	1 - All FDA-approved			Diagnosis. Documentation of	Coverage is provided for		12 months		0
OPION (AUVELITY)	Indications.			trial and failure of at least two	-				
				generic antidepressants	older.				
				alternatives such as an SSRI,					
				SNRI, bupropion, trazodone					
				or mirtazapine.					
DIAZOXIDE CHOLINE (VYKAT	1 - All FDA-approved			Diagnosis. Must have a	Member must be 4 years of	By or in consultation with an	Initial: 6 months	For reauth: must have	0
XR)	Indications.			diagnosis of Prader-Willi	age or older.	endocrinologist or geneticist	Reauthorization: 12 months	documentation from	
				syndrome (PWS) confirmed				prescriber indicating	
				by genetic testing and have				stabilization or improvement	
				symptoms associated with				in hyperphagia symptoms.	
				hyperphagia (i.e. persistent					
				sensation of hunger, food					
				preoccupations, an extreme					
				drive to consume food, food-					
				related behavior problems,					
				lack of normal satiety). Must					
				have baseline fasting plasma					
				glucose or hemoglobin A1c.					
DIHYDROERGOTAMINE NASAL			Members with hemiplegic or	Diagnosis. Documentation of	Coverage is provided for		12 months	For reauth: documentation	0
SPRAY (MIGRANAL)	Indications.		basilar migraine, ischemic	trial and failure of 1	members 18 years of age and			from prescriber indicating	
				medication from each of the	older.			stabilization or improvement	
			•	following classes: a NSAID and				in condition.	
			silent ischemia) or who have	a triptan unless					
			clinical symptoms or findings	contraindicated.					
			consistent with coronary						
			artery vasospasm (including						
			Prinzmetal's variant angina or						
			uncontrolled hypertension).						
DORNASE ALFA	1 - All FDA-approved			Diagnosis.		By or in consultation with a	12 months	For reauth: must have	0
(PULMOZYME)	Indications.			Diagnosis.		pulmonologist or cystic		documentation from	
	That cations:					fibrosis specialist		prescriber indicating	
						Thorosis specialist		stabilization or improvement	
								in condition.	
DRONABINOL	1 - All FDA-approved			Diagnosis. Nausea and			12 months	conditions	0
	Indications.			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.					
				amess contramulated.					
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				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
DROXIDOPA (NORTHERA)	1 - All FDA-approved			Diagnosis. Documentation of	Coverage is provided for		2 weeks	For reauth: rationale from the	0
	Indications.			a clinical diagnosis of	members 18 years of age and			provider for continuing	
				symptomatic neurogenic	older.			therapy beyond 2 weeks	
				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.					
DUPILUMAB (DUPIXENT)	1 - All FDA-approved			Diagnosis. For asthma: must	For atopic dermatitis: 6	By or in consultation with an	12 months	Reauth for asthma:	0
BOTTEON B (BOTTALINI)	Indications.			have either moderate to	months or older. For asthma:	allergist, dermatologist,	12 1110111113	documentation of	
	maications.			severe eosinophilic	6 years or older. For	immunologist, pulmonologist,		improvement (e.g. reduced	
				phenotype with an eosinophil	1	ear-nose/throat specialist, or		symptoms, reduced	
				count greater than or equal to		gastroenterologist.		exacerbations, need for oral	
				150 cells/microliter or oral	polyps: 12 years and older.	gasti denter diogist.		steroids). Reauth for all other	
				corticosteroid dependent	For all other indications: 18			indications: documentation	
				persistent asthma (chronic	years or older.			from prescriber indicating	
				oral corticosteroid use).	,			stabilization or improvement	
				Documentation of recent use				in condition.	
				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				1	
				intolerance to a topical				1	
				corticosteroid or topical				1	
				calcineurin inhibitor. For nasal					
				polyps: history of trial and					
				failure of Xhance (fluticasone					
				propionate). Must be used as					
	1			add-on maintenance therapy.	1				

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria		~	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
EDARAVONE (RADICAVA ORS)				Diagnosis of Amyotrophic		By or in consultation with a	12 months	Reauth: must provide	0
	Indications.			Lateral Sclerosis (ALS). Must	members 18 years of age and	neurologist		documentation of clinical	
				have normal respiratory	older			benefit based on the	
				function (defined as a forced				prescriber's assessment and	
				vital capacity (FVC) of at least				an ALSFRS-R score within the	
				80%), must be able to				past 12 months	
				perform activities of daily					
				living (ADLs) such as eating					
				and moving around					
				independently, must provide					
				a recent ALSFRS-R score.					
GARTIGIMOD	1 - All FDA-approved			Diagnosis. Memember must	Member must be 18 years of		12 months	For reauthorization:	0
FA/HYALURONIDASE-QVFC	Indications.			have generalized myasthenia	age or older.	neurologist.		Documentation from the	
YVGART HYTRULO)				gravis (gMG) who are anti-				provider that the member has	
				acetylcholine receptor (AChR)				experienced improvement in	
				antibody positive or chronic				signs and symptoms of	
				inflammatory demyelinating				generalized myasthenia gravis	
				polyneuropathy (CIDP). The				(for example, speech,	
				requested agent must not be				swallowing, mobility, or	
				used in combination with				respiratory function). The	
				another myasthenia gravis				member has also experienced	
				medication. The member has				a decrease in the number of	
				experienced therapeutic				exacerbations of generalized	
				failure, contradindication or				myasthenia gravis. For CIDP,	
				intolerance to generic				the member has experineced	
				pyridostigmine. For CIDP, the				improvement in their	
				member has experienced				functional ability or strength	
				progressive symptoms for at				from baseline.	
				least two (2) months. The				mom sasemie.	
				member has progressive or					
				relapsing motor sensory					
				dysfunction of more than one					
				limb or a peripheral nerve					
				nature, developing over at					
				least 2 months. The member					
				has hypo-or areflexia (usually					
				involves all four limbs). The					
				member has nerve					
				conduction studies strongly					
				supportive of demyelination					
				and meets one of the					
				following: motor distal latency	1				
				prolongation in at least 2					
				nerves, reduction of motor					
				conduction velocity in at least		<u> </u>	10 11		
EXACAFTOR/TEZACAFTOR/I				Diagnosis. Documentation of		By or in consultation with a	12 months	For reauthorization:	0
CAFTOR (TRIKAFTA)	Indications.			genetic test confirming the		cystic fibrosis specialist or		documentation from	
				member has at least one	older	pulmonologist		prescriber indicating	
				F508del mutation in the CFTR				stabilization or improvement	
				gene or a mutation in the				in condition.	
				CFTR gene that is responsive					
				based on in vitro data.					

Other Criteria 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
documentation of improvement in platelet count from baseline. For hepatitis C: documentation the member is still on antiviral	
improvement in platelet count from baseline. For hepatitis C: documentation the member is still on antiviral	
count from baseline. For hepatitis C: documentation the member is still on antiviral	
hepatitis C: documentation the member is still on antiviral	
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meet initial criteria.	
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m Ck ha 10 zio an Ck th Re	or reauth for CKD on dialysis: flust have a Hgb less than or qual to 11g/dl. For reauth for KD not on dialysis: must ave Hgb less than or equal to 0 g/dl. For reauth for dovudine treated members and pediatric members with KD: must have a Hgb less han or equal to 12 g/dl. eauth for all other dx must heet initial criteria.

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
ERENUMAB-AOOE (AIMOVIG)	1 - All FDA-approved			Diagnosis. For episodic	Coverage is provided for		Initial: 6 months	For reauth: Provider	0
	Indications.			migraine: Provider attestation	members 18 years of age and		Reauthorization: 12 months	attestation the member is	
				the member has 4 to 14	older			having a reduced number of	
				headache days per month. For	•			migraine/headache days per	
				chronic migraine: Provider				month or a decrease in	
				attestation the member has at				migraine/headache severity. A	· ·
				least 15 headache days per				migraine is defined as a	
				month for 3 or more months				headache that has at least	
				with at least 8 migraine days				two of the following	
				per month. For both: Must				characteristics: unilateral	
				have a trial and failure of one				location, pulsating/throbbing	
				beta-blocker and one				quality, moderate or severe	
				anticonvulsant unless				intensity (inhibits or prohibits	
				contraindicated or intolerant.				daily activities), is aggravated	
								by routine activity, nausea	
								and/or vomiting, photophobia	
								and phonophobia.	
ETANERCEPT (ENBREL)	3 - All Medically-accepted			Diagnosis. For rheumatoid	Member must be 2 years of	By or in consultation with a	12 months	For reauth: must have	0
	Indications.			arthritis (RA): history of trial	age or older.	rheumatologist or		documentation from	
				and failure, contraindication,		dermatologist.		prescriber indicating	
				or intolerance to a three-				stabilization or improvement	
				month trial with methotrexate				in condition.	
				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 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				1	

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria		Age Restriction		Coverage Duration	Other Criteria	Part B Prerequisite
ETRASIMOD (VELSIPITY)	1 - All FDA-approved Indications. 1 - All FDA-approved Indications.	Off-Label Uses	Exclusion Criteria	Information Diagnosis. For ulcerative colitis (UC): history of trial and failure, contraindication, or intolerance to 2 of the following therapy options: aminosalicylates, corticosteroids or immunomodulators with inadequate response or side effects/toxicity unless contraindicated. Documentation of a recent diagnosis of recurrent Clostridioides difficile infection (CDI) -AND- Will be used for prophylaxis and not treatment of recurrent CDI -AND- Attestation that antibiotic treatment for the most recent recurrent CDI is	Coverage is provided for	Prescriber Restriction By or in consultation with a gastroenterologist	Coverage Duration 12 months 1 month	For reauth: must have documentation from prescriber indicating stabilization or improvement in condition. For reauthorization, attestation of recurrent CDI episodes after administration of the initial fecal microbiota product -AND- Will be used for prophylaxis and not treatment of recurrent CDI - AND- Attestation that antibiotic treatment for the	0
				complete or will be completed.				most recent recurrent CDI is complete or will be completed.	
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
FENTANYL CITRATE (TRANSMUCOSAL)	1 - All FDA-approved Indications.		including headache/migraines and dental pain.	Diagnosis. Documentation the member has active cancer and is experiencing breakthrough pain despite being on around the clock opioid therapy. Must be opioid tolerant. Must currently be using a longacting opioid.		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 mcg 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 25 mg oral oxymorphone 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.	t
FILGRASTIM-SNDZ (ZARXIO)	3 - All Medically-accepted Indications.			Diagnosis.			6 months	For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	0
FLUTICASONE PROPIONATE (XHANCE)	1 - All FDA-approved Indications.			Diagnosis.	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

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
GALCANEZUMAB-GNLM	1 - All FDA-approved			Diagnosis. For episodic	Coverage is provided for		Initial: 6 months	For reauth: Provider	0
(EMGALITY)	Indications.				members 18 years of age and		Reauthorization: 12 months	attestation the member is	
,				the member has 4 to 14	older			having a reduced number of	
				headache days per month. Fo	-			migraine/headache days per	
				chronic migraine: Provider				month or a decrease in	
				attestation the member has a				migraine/headache severity. A	
				least 15 headache days per				migraine is defined as a	
				month for 3 or more months				headache that has at least	
				with at least 8 migraine days				two of the following	
				per month. For both: Must				characteristics: unilateral	
				have tried and failed one beta	-			location, pulsating/throbbing	
				blocker for at least 2 months				quality, moderate or severe	
				and one anticonvulsant for at				intensity (inhibits or prohibits	
				least 2 months unless				daily activities), is aggravated	
				contraindicated or intolerant.				by routine activity, nausea	
				For cluster headache:				and/or vomiting, photophobia	n l
				Provider attestation the				and phonophobia. A cluster	
				member has at least one				headache is defined as at leas	t
				cluster attack every other day				5 severe to very severe	
				and no more than 8 attacks a				unilateral headache attacks	
				day. Must have a trial and				lasting 15 to 180 minutes	
				failure of either verapamil for				untreated. Headaches occur	
				at least 2 weeks or a one-time				once every other day to 8	
				subocciptal steroid injection				times a day. The pain is	
				unless contraindicated or				associated with ipsilateral	
				intolerant.				conjunctival injection,	
								lacrimation, nasal congestion,	
								rhinorrhea, forehead and	
								facial sweating, miosis, ptosis	
								and/or eyelid edema, and/or	
								with restlessness or agitation.	
CANAVOLONE (ZTALBAV)	4. All FDA			Diama aria	Carrage as in a second and face	D in	42		
GANAXOLONE (ZTALMY)	1 - All FDA-approved			Diagnosis.	• .	By or in consultation with a	12 months		0
	Indications.					neurologist			
GLECAPREVIR-PIBRENTASVIR	1 All EDA annuariad		N. C. marks are suithly many disperses and	Critaria will be applied	older.	D	Criteria will be applied		
			Members with moderate or	Criteria will be applied	Coverage is provided for	By or in consultation with a			ľ
(MAVYRET)	Indications.		severe hepatic impairment	consistent with current	_	gastroenterologist,	consistent with current		
			(Child-Pugh C). Coadministration with	AASLD/IDSA guidance and/or		hepatologist, infectious	AASLD/IDSA guidance and/or		
				FDA approved labeling	AASLD/IDSA guidance and/or		FDA approved labeling		
GLP-1 RECEPTOR AGONISTS	1 - All FDA-approved		atazanavir and rifampin.	Diagnosis of Type 2 diabetes	FDA-approved labeling.	specialist.	12 months		0
GEF-1 NECEF TON AGONISTS	Indications.			or documented prior therapy			12 months		ľ
	marcations.			with a Type 2 diabetes					
				medication. Claims will					
				automatically pay on-line					
				without a requirement to					
				submit for prior authorization					
				when one of the following					
				criteria is met: 1. a Type 2					
				diabetes diagnosis code is					
				submitted at the point of sale					
				OR 2. a pharmacy claims					
				history of a Type 2 diabetes					
				medication within the past					
	!	<u> </u>		130 days.		l .	!	1	

C	La di cali ca la di cala ca	Off Labalitan	Fundamina Calibratia	Required Medical	A Doublishing	Durantila and Danatal at land	C	Out an Orithania	Don't D Don't and the
·	Indication Indicator	Off-Label Uses	Exclusion Criteria		Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
GLYCEROL PHENYLBUTYRATE (RAVICTI)	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.	U
GUSELKUMAB (TREMFYA)	1 - All FDA-approved Indications.			Diagnosis. 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).	Coverage is provided for members 18 years of age and older	By or in consultation with a rheumatologist, dermatologist, or gastroenterologist.	12 months	For reauth: must have documentation from prescriber indicating stabilization or improvement in condition.	0
ICATIBANT ACETATE	1 - All FDA-approved Indications.			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 functional level. There is a documented history of at least one symptom of a moderate to severe HAE attack (i.e. moderate to severe abdominal pain, facial swelling, airway swelling) in the absence of hives or a medication known to cause angioedema. Member must not be taking any medications that may exacerbate HAE, including angiotensinconverting enzyme (ACE) inhibitors, tamoxifen, or estrogen-containing medications.	members 18 years of age or older.	By or in consultation with an allergist, immunologist, hematologist, or dermatologist	12 months	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, paliperidone, risperidone, aripiprazole, or ziprasidone.	Coverage is provided for members 18 years of age or older.		12 months		0

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
INCOBOTULINUMTOXINA	1 - All FDA-approved			Diagnosis.			12 months	For reauthorization:	0
(XEOMIN)	Indications.							documentation from	
								prescriber indicating	
								stabilization or improvement	
								in condition.	
INFLIXIMAB-ABDA	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
(RENFLEXIS)	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			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,					1

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
FLIXIMAB-DYYB	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
IFLECTRA)	Indications.		moderate to severe heart	arthritis (RA): history of trial	Psoriasis: Coverage is	rheumatologist,		documentation from	
			failure.	and failure, contraindication,	provided for members 18	gastroenterologist, or		prescriber indicating	
				or intolerance to a three-	years of age or older. For CD,	dermatologist.		stabilization or improvement	
				month trial with methotrexate	UC: Coverage is provided for			in condition.	
				or another DMARD. For	members 6 years of age or				
				psoriatic arthritis (PsA) one of	older.				
				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,					
SULIN SUPPLIES	1 - All FDA-approved			Confirmation of insulin use			12 months		0
	Indications.			within the past 12 months					
				based on paid claims or					
				provider documentation.					

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction		Other Criteria	Part B Prerequisite
IPTACOPAN (FABHALTA)	1 - All FDA-approved		Initiation in patients with	Diagnosis. For paroxysmal	Coverage is provided for	By or in consultation with a	12 months	For reauth: documentation of	0
	Indications.		unresolved serious infection	nocturnal hemoglobinuria	members 18 years of age and			improvement.	
			caused by encapsulated	(PNH): confirmed diagnosis of	older	immunologist, nephrologist,			
			bacteria.	PNH by flow cytometry		or genetic specialist			
				testing. Flow Cytometry					
				pathology report must be					
				supplied and demonstrate at					
				least 2 different GPI protein					
				deficiencies within 2 different					
				cell lines from granulocytes,					
				monocytes, or erythrocytes.					
				Member is transfusion					
				dependent as defined by					
				having a transfusion within					
				the last 12 months and one of					
				the following: a hemoglobin is					
				less than or equal to 7 g per					
				dL or has symptoms of					
				anemia and the hemoglobin is					
				less than or equal to 10 g per					
				dL. Must have a Lactate					
				dehydrogenase (LDH) level at					
				least 1.5 times the upper limit					
				of the normal range.					
IVABRADINE (CORLANOR)	1 - All FDA-approved		Acute decompensated heart	Diagnosis. For Adult Chronic	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	members 18 years of age or	cardiologist		documentation from	
			than 90/50 mmHG, sick sinus	have left ventricular ejection	older. DCM: coverage is			prescriber indicating	
			syndrome, sinoatrial block, or		provided for members 6			stabilization or improvement	
			3rd degree AV block-unless a		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	carvedilol, metoprolol					
			(heart rate maintained	succinate) at the maximally					
			exclusively by the pacemaker)	, tolerated dose or has a					
			concomitant use of strong	contraindication to beta-					
			CYP3A4 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					
		1		105 beats per minute (BPM)					
				for 6-12 months of age,					
				greater than or equal to 95 for	1				
		1		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).					
			1		L	<u> </u>]		<u> </u>

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
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	members 1 month of age or	pulmonologist or cystic		documentation from	
				member has at least one	older.	fibrosis 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.					
L-GLUTAMINE (ENDARI)	1 - All FDA-approved			Diagnosis. Must be used to	Coverage is provided for	By or in consultation with a	12 months	For reauthorization:	0
	Indications.			reduce the acute	members 5 years of age and	physician who specializes in		Documentation there has	
				complications of sickle cell	older	SCD (e.g.a hematologist)		been a reduction in vaso-	
				disease (SCD) and the				occlusive painful events or an	
				member must have				improvement in condition.	
				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					
LANDESTIDE (CONTACTIVINE	4 411 50 4			hydroxyurea.		1	- 1 . 1		
LANREOTIDE (SOMATULINE	1 - All FDA-approved			Diagnosis. For acromegaly:	Coverage is provided for	By or in consultation with an	For oncology indications: 6	For reauth: documentation of	0
DEPOT)	Indications.			must have inadequate	-	endocrinologist or oncologist	months. All other indications:	improvement or stabilization.	
				response to surgery or	older.		12 months		
				radiotherapy 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.					
LEDISPASVIR-SOFOSBUVIR	1 - All FDA-approved			Criteria will be applied	Coverage is provided for	By or in consultation with a	Criteria will be applied		0
(HARVONI)	Indications.			consistent with current	members who are age-	gastroenterologist,	consistent with current		
				AASLD/IDSA guidance and/or	appropriate according to	hepatologist, infectious	AASLD/IDSA guidance and/or		
				FDA approved labeling	AASLD/IDSA guidance and/or	_	FDA-approved labeling.		
				T Break approved labeling	FDA-approved labeling.	specialist.	approved labeling.		
LENIOLISIB (JOENJA)	1 - All FDA-approved		<u> </u>	Diagnosis of activated	Coverage is provided for	By or in consultation with a	12 months		0
LEINOLISID (JOLINA)	Indications.			phosphoinositide 3-kinase	members 12 years of age or	hematologist, immunologist,	TE Mondis		ľ
	maications.			delta syndrome (APDS). Must	-	or geneticist.			
				have genetic testing	Joinet.	or genericist.			
				confirming the PI3K delta					
				mutation with a documented					
				variant in either PIK3CD or					
				PIK3R1. Documentation of					
				inadequate response to					
				immunoglobulins.					

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction		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.	200 days post-transplant	For reauth: no reauthorization after initial coverage period.	
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
LEVACETYLLEUCINE (AQNEURSA)	1 - All FDA-approved Indications.			Diagnosis. Documentation the diagnosis was confirmed by genetic testing demonstrating one of the following: 1. a mutation in both alleles of NPC1 or NPC2 OR 2. mutation in one allele and either a positive filipin-staining or elevated cholestance triol/oxysterols (greater than 2x ULN). Documentation the member has at least one neurological symptom of NPC (e.g. decrease in motor skills, ataxia, seizures, etc.). Must not be used in combination with Miplyffa.			12 months	Reauthorization: Documentation the member is experiencing an improvement or stabilization in disease.	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

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
LEVOMILNACIPRAN	1 - All FDA-approved			Diagnosis. Documentation of	Coverage is provided for		12 months		0
(FETZIMA)	Indications.				members 18 years of age and				
				generic antidepressants	older.				
				alternatives such as an SSRI,					
				SNRI, bupropion, trazodone					
LIDOCAINE PATCH	3 - All Medically-accepted			or mirtazapine Diagnosis. This Prior			12 months		0
LIDOCAINE PATCIT	Indications.			Authorization requirement			12 111011(115		
	maleucions.			only applies to members					
				when a non-FDA approved					
				diagnosis is submitted at the					
				point of sale. FDA-approved					
				diagnosis codes submitted wil	1				
				pay without prior					
				authorization requirement.					
LOTILANER (XDEMVY)	1 - All FDA-approved			Diagnosis of Demodex	•	Prescribed by or in	6 weeks		0
	Indications.			blepharitis confirmed by both	age and older	consultation with an			
				of the following: 1. Member		optometrist or			
				has at least mild erythema or itching of the upper eyelid		ophthalmologist			
				margin. 2. Mite presence (e.g.					
				collarettes) confirmed by slit					
				lamp examination of the					
				eyelashes.					
LUMACAFTOR/IVACAFTOR	1 - All FDA-approved			Diagnosis. Documentation of		By or in consultation with a	12 months	For reauthorization:	0
(ORKAMBI)	Indications.			a genetic test confirming that		pulmonologist or cystic		documentation from	
				the member is homozygous		fibrosis specialist		prescriber indicating	
				for the F508del mutation in				stabilization or improvement	
				the CFTR gene (has two copies				in condition.	
				of the F508del mutation in					
MACITENTAN (OPSUMIT)	1 - All FDA-approved		Drognongu	the CFTR gene). Diagnosis. Pulmonary arterial		Prescribed by or in	Initial: 3 months Reauth: 12	For reauth: documentation	0
IVIACITENTAN (OPSOIVIT)	Indications.		Pregnancy	hypertension (PAH) WHO		-		from prescriber that	l ^o
	malcations.			Group I confirmed by chart		or pulmonologist.	monuis	demonstrates member is	
				documentation of right-heart		or parmonologist.		tolerating and receiving	
				catheterization (RHC)				clinical benefit from	
				indicating a mean pulmonary				treatment	
				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.					
MANNITOL (BRONCHITOL)	1 - All FDA-approved			Diagnosis. Must have passed a			12 months	For reauth: documentation of	0
, , ,	Indications.			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.					
		1			1				

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
MARALIXIBAT (LIVMARLI)	1 - All FDA-approved Indications.		PFIC type 2 patients with specific ABCB11 variants resulting in non-functional or complete absence of bile salt export pump (BSEP) protein.	Diagnosis of pruritis caused by progressive familial intrahepatic cholestatis (PFIC) or Allagile syndrome (ALGS) which has been confirmed by genetic testing. Documentation of trial and failure of ursodiol and another medication for cholestatic pruritis (e.g.	Coverage is provided for members 3 months of age and older.	By or in consultation with a hepatologist or gastroenterologist.	12 months	For reauth: documentation of improvement in pruritis.	·
MARIBAVIR (LIVTENCITY)	1 - All FDA-approved Indications.			cholestyramine, rifampin). Diagnosis of post-transplant (solid organ or hematopoietic stem cell) cytomegaloviris (CMV) infection/disease that is refractory to treatment with ganciclovir, valganciclovir, cidofovir, or foscarnet. Must weight at least 35 kg. Must not be used concomitantly with ganciclovir or valganciclovir.		By or in consultation with a hematologist, oncologist, infectious disease physician, or transplant specialist.	3 months	For reauthorization: documentation from prescriber indicating stabilization or improvement in condition.	0
MAVORIXAFOR (XOLREMDI)	1 - All FDA-approved Indications.			Diagnosis. Confirmation of the diagnosis with a genetic test confirming pathogenic or likely pathogenic variants in the CXCR4 gene. Documentation of a baseline absolute neutrophil count (ANC) less than or equal to 400 cells/?L or absolute lymphocyte count (ALC) less than or equal to 650 cells/?L. Documentation of symptoms and complications associated with WHIM syndrome (e.g. warts, hypogammaglobulinemia, recurrent infections, and myelokathexis)	Members 12 years of age and older	By or in consultation with an immunologist, hematologist, or dermatologist	12 months	For reauthorization: Documentation of one of the following: 1. an improvement in ANC or ALC from baseline 2 A decrease in frequency or severity of infections since initiating therapy.	

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
MECASERMIN (INCRELEX)	1 - All FDA-approved		Coverage is not provided for	Diagnosis. Growth chart and	Coverage is provided for	By or in consultation with an	12 months	For reauth, must include a	0
	Indications.		members with active or	documentation that	members 2 years of age or	Endocrinologist		recent progress note from	
			suspected neoplasia, closed	epiphyses are open. For	older.			prescriber indicating growth	
			epiphyses.	growth hormone deletion:				and maturation as a result of	
				must have growth hormone				treatment and that epiphyses	
				(GH) gene deletion in gene				have not closed.	
				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 fo	r				
				age and sex, basal IGF-1 SD of					
				less than or equal to 3.0 based	d				
				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).					
METHYLNALTREXONE	1 - All FDA-approved		Known or suspected	Diagnosis. For opioid-induced	Coverage is provided for		12 months	For reauth: documentation	0
(RELISTOR)	Indications.		gastrointestinal obstruction	constipation and advanced	members 18 years of age and			from the prescriber indicating	
			and members at an increased	life-limiting illness: must have	older.			an improvement in condition	
			risk of recurrent obstruction.	documentation of previous				(both diagnoses) and must	
				trial of lactulose. For opioid-				continue to be on opioid	
				induced constipation with				therapy (non-cancer pain).	
				chronic non-cancer pain: mus	t				
				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.					
				lactarese.					
MIFEPRISTONE (KORLYM)	1 - All FDA-approved			Diagnosis. Must have failed	Coverage is provided for	By or in consultation with an	12 months		0
	Indications.			_	members 18 years of age and	-			
				for surgery. Female members					
				of reproductive potential:					
				must have baseline (within					
				previous month, must include	,				
				date of test) negative					
				pregnancy test prior to					
				starting mifepristone and	1				
				must be using nonhormonal					
				medically acceptable method					
				of contraception (unless					
				surgically sterilized) during					
				treatment and for 1 month					
	1			after mifepristone therapy.		1	I	ļ.	

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Group MIGLUSTAT (ZAVESCA)	Indication Indicator 1 - All FDA-approved	Off-Label Uses	Exclusion Criteria Miglustat is being used in	Information Diagnosis. Documentation the	Age Restriction Coverage is provided for	Prescriber Restriction By or in consultation with an	Coverage Duration 12 months	Other Criteria Reauthorization:	Part B Prerequisite
IVIIGEOSTAT (ZAVESCA)	Indications.		combination with another	member has at least one of		appropriate specialist (i.e.	12 months	Documentation from the	ľ
	malcations.		therapy for Gaucher's disease		older.	hematologist, geneticist,		prescriber indicating	
			incrupy for Gadefier's discuse	due to iron deficiency with a		radiologist, orthopedist,		improvement or stabilization	
				low hemoglobin for age and		endocrinologist,		in member's condition.	
				sex, 2) thrombocytopenia 3)		rheumatologist, hepatologist)			
				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).					
MITAPIVAT (PYRUKYND)	1 - All FDA-approved	+		Diagnosis of hemolytic anemia	Coverage is provided for	By or in consultation with a	12 months	For reauthorization:	0
	Indications.			with pyruvate kinase	members 18 years of age or	hematologist or a physician		documentation of	
				deficiency (PKD) confirmed by	older.	who specializes in the		improvement in condition.	
				genetic testing.		treatment of inherited			
						metabolic disorders.			
MODAFINIL (PROVIGIL)	1 - All FDA-approved			Diagnosis. For narcolepsy and		By or in consultation with a		For reauthorization: must	0
	Indications.			obstructive sleep apnea: Sleep		sleep specialist, ENT (ear,	OSA: 12 months	have documentation from	
				Study (e.g. Polysomnogram,		nose, and throat specialist),		prescriber indicating	
				Multiple Sleep Latency Test)		neurologist, or pulmonologist		stabilization or improvement	
				confirming diagnosis. For shift				in condition.	
				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).					
				and outing					

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
MULTIPLE SCLEROSIS	1 - All FDA-approved			Diagnosis. For multiple		By or in consultation with a	12 months	For reauthorization: must	0
THERAPIES	Indications.			sclerosis (MS), must have		neurologist or		have documentation from	
				relapsing Multiple Sclerosis		gastroenterologist		prescriber indicating	
				(including clinically isolated				stabilization or improvement	
				syndrome, relapsing-remitting				in condition.	
				disease, and active secondary					
				progressive disease) and					
				functional status must be					
				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. For ulcerative colitis					
				(UC): must have history of					
				trial and failure,					
				contraindication or					
				intolerance to an					
				immunomodulator (i.e.,					
				Azathioprine, 6-					
				Mercaptopurine,					
				Methotrexate).					
NETARSUDIL (RHOPRESSA)	1 - All FDA-approved			Diagnosis. Member must have			12 months	For reauthorization: must	0
	Indications.			a baseline intraocular	members 18 years of age and			have documentation from	
					older.			prescriber indicating	
				mmHg. Documentation of				stabilization or improvement	
				trial and failure,				in condition.	
				contraindication, or					
1				intolerance to timolol and					
				latanoprost.					

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
NINTEDANIB (OFEV)	1 - All FDA-approved	OII-Label Oses	Exclusion Citteria	Diagnosis. For a diagnosis of	Coverage provided for	By or in consultation with a	Initial: 6 months, Reauth: 12	For reauth: must have	n Prerequisite
MINIEDANIB (OFEV)	Indications.			Idiopathic Pulmonary Fibrosis	members age 18 years and	pulmonologist	months	documentation from	ľ
	malcations.			(IPF): Must have diagnosis	older.	pulliologist	montais	prescriber indicating that	
				confirmed by either high-	older.			member still is a candidate for	
				resolution computed				treatment.	
				tomography (HRCT) or				treatment.	
				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. Must have a					
				trial of pirfenidone (Esbriet).					
				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					
NITISINONE (ORFADIN)	1 - All FDA-approved			diagnosis of Chronic Fibrosing Diagnosis of hereditary			12 months	For reauth: Documentation	0
INTISINONE (OKI ADIN)	Indications.			tyrosinemia type 1 (HT-1)			12 months	from the prescriber indicating	ľ
	malcations.			confirmed by DNA testing or				improvement or stabilization	
				biochemical testing (ie. urine				in the member's condition	
				succinylacetone (SA) level).				in the member 5 condition	
NITROGLYCERIN 0.4%	1 - All FDA-approved		Severe anemia (defined as	Diagnosis. Must provide	Coverage is provided for		Initial: 2 months	For reauthorization:	0
OINTMENT (RECTIV)	Indications.		hemoglobin less than 8g/dL).	documentation that chronic	members 18 years of age or		Reauthorization: 12 months	documentation from	
			Increased intracranial	anal fissure symptoms have	older.			prescriber indicating	
			pressure. Concomitant use of	persisted for at least 6 weeks.				stabilization or improvement	
			a phosphodiesterase type 5					in condition.	
			(PDE5) inhibitor such as						
			sildenafil (Revatio, Viagra),						
			tadalafil (Adcirca, Cialis), or						
ODE: (1)((D (= / (2) ())))	4 All ED 6	1	vardenafil (Levitra, Staxyn).	D		B	42 1		
ODEVIXIBAT (BYLVAY)	1 - All FDA-approved		PFIC type 2 patients with	Diagnosis of pruritis caused by		By or in consultation with a	12 months	For reauth: documentation of	ľ
	Indications.		specific ABCB11 variants	progressive familial	members 3 months of age	hepatologist or		improvement in pruritis.	
			_	intrahepatic cholestatis (PFIC)	and older.	gastroenterologist.			
			complete absence of bile salt						
			export pump (BSEP) protein.	which has been confirmed by					
				genetic testing.					
				Documentation of trial and					
				failure of ursodiol and another medication for					
				cholestatic pruritis (e.g.					
				cholestyramine, rifampin).					
	1		1	ionorestyramine, mamping.	<u>I</u>	ı	<u>I</u>	<u> </u>	<u> </u>

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	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, paliperidone, risperidone, aripiprazole, or ziprasidone. If the member is 65 and older and not in hospice care and taking this medication at the same time as another anticholinergic medication, must provide documentation of the following: 1. Provider must acknowledge that the benefit or the combination of medication outweighs the potential risks, 2. The member has tried and failed monotherapy, 3. Clinical rationale for use of 2 or more anticholinergic medications.	members 18 years of age or older.		12 months		0
OLEZARSEN (TRYNGOLZA)	1 - All FDA-approved Indications.			Diagnosis. Confirmation of the diagnosis by at least one of the following: 1. a genetic test 2. a North American Familial Chylomicronemia Syndrome (NAFCS) score of greater than or equal to 60. 3. fasting triglycerides greater than 10 mmol/l or 880mg/dl and symptoms of the disease (e.g. acute pancreatitis, hepatosplenomegaly, abdominal pain, lipemia retinalis)	members 18 years of age and	By on in consultation with a lipidologist, geneticist cardiologist, or endocrinologist	12 months	For reauthorization: documentation indicating stabilization or improvement in condition.	0

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
OMALIZUMAB (XOLAIR)	1 - All FDA-approved			Diagnosis. For moderate to		By or in consultation with, for	12 months	For reauthorization:	0
	Indications.			severe allergic asthma: recent		Urticaria: allergist,		documentation from	
				total serum IgE level of		dermatologist, immunologist.		prescriber indicating	
				greater than 30 IU/ml and the		Asthma: pulmonologist or		stabilization or improvement	
				pre-treatment IgE levels do		allergist. Nasal Polyps:		in condition.	
				not exceed manufacturers		allergist, ear/nose/throat			
				dosing recommendations.		specialist, or immunologist.			
				Documentation of recent use		Allergy: allergist or			
				and failure to respond to		immunologist.			
				inhaled steroid in combo with					
				long acting beta agonist.					
				Documentation of a positive					
				skin or in vitro reactivity to					
				perennial aeroallergen. 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					
				spontaneou urticaria (CSU):					
				must have chart					
				documentation showing					
				history of urticaria w/					
				presence of hives, must have					
				trial of one 2nd generation H1					
OMAVELOXOLONE	1 - All FDA-approved			Diagnosis of Friedreich's	Coverage is provided for	By or in consultation with a	12 months		0
(SKYCLARYS)	Indications.			ataxia that has been	members 16 years of age or	neurologist.			
ľ ,				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%.					
OMNIPOD POD	1 - All FDA-approved			Must have documentation of			12 months		0
]
	Indications.			previous insulin use.					

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
ONABOTULINUMTOXINA	1 - All FDA-approved			Diagnosis. For migraine		By or in consultation with an	12 months	For reauth: documentation	0
(BOTOX)	Indications.			prophylaxis: must have		appropriate specialist (ie.		from prescriber indicating	
				adequate trial of two migraine		dermatologist, neurologist,		stabilization or improvement	
				prophylactic agents each from	n e	urologist).		in condition.	
				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.					
ONCOLOGY MEDICATIONS	3 - All Medically-accepted			Diagnosis. For Bosulif, Iclusig,		By or in consultation with an	6 months		0
	Indications.			and Tasigna for CML: must		oncologist, hematologist,			
				have had an inadequate		neurologist, transplant			
				response or intolerance to		specialist, allergist, or			
				imatinib or dasatinib.		immunologist.			
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 has tried and failed or had an				the taper will be reevaluated.	
				intolerance or				For all other ongoing therapy: documentation the member	
				contraindication to at least				has been treated with the	
				one non-benzodiazepine				requested agent within the	
				anticonvulsant. For sleep				past 90 days	
				disorder: documentation the				past 30 days	
				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-traumation					
				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					
				hospitalization within the past	:				
				3 months. 3. The requested					
				therapy is medically necessary					
				to prevent harm to the					
				member or others. For a					
<u> </u>	L			musculoskeletal disorder:	1	I	I .	1	1

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
PALIVIZUMAB (SYNAGIS)	1 - All FDA-approved		having received Beyfortus	DX. Must have documented	Less than 12 months or less		Minimum duration 1 month.		0
	Indications.		(nirsevimab-alip) for the	reason not being able to use	than 24 months of age at start	t	Maximum of 5 doses per RSV		
			current RSV season	Beyfortus (nirsevimab-alip). If	of RSV season depending on		season		
				under age 12 mo at start of	criteria.				
				RSV season w/ no other					
				medical dx: must have					
				gestational age (GA) less than					
				29 wks. If under age 24 mo at					
				start of RSV season during 1st					
				yr of life w/ Chronic Lung					
				Disease (CLD) of prematurity:					
				must have GA less than 32					
				wks 0 days & required greater					
				than 21% oxygen (O2) for at					
				least 1st 28 days of life. If					
				under age 24 mo at start of					
				RSV season during 2nd yr of					
				life w/ CLD of prematurity:					
				must have GA less than 32					
				wks 0 days & required greater					
				than 21% O2 for at least 1st					
				28 days of life & have					
				continued to require medical					
				support (chronic					
				corticosteroid therapy,					
				diuretic therapy,					
				supplemental O2) during 6 mo before start of 2nd RSV					
				season. If under age 12 mo. at					
				start of RSV season w/ heart					
				disease: must have					
				hemodynamically significant					
				Congenital Heart Disease					
				(CHD) (& be on drugs to					
PALOVAROTENE (SOHONOS)	1 - All FDA-approved			Diagnosis confirmed by	Members assigned female at	Prescribed by or in	12 months		0
, ,	Indications.			precesence of ACVR1	birth must be 8 years and	consultation with an			
				mutation.	older. Members assigned	orthopedist or			
					male at birth must be 10 years				
					and older.				
PAMIDRONATE (AREDIA)	1 - All FDA-approved			Diagnosis. For hypercalcemia	Coverage is provided for		12 months	For reauth: documentation	0
	Indications.			of malignancy: must be used	members 18 years of age or			from prescriber indicating	
				in conjunction with adequate	older.			stabilization or improvement	
				hydration in members with				in condition.	
				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					
	1		L	therapy .	l	L			<u> </u>

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria		Age Restriction	Prescriber Restriction	Coverage Duration		Part B Prerequisite
ASIREOTIDE (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.	members 18 years of age or	By or in consultation with an Endocrinologist	12 months	For reauth: documentation of improvement or stabilization.	
EGFILGRASTIM-BMEZ IEXTENZO)	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.		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

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
PIMAVANSERIN (NUPLAZID)	1 - All FDA-approved			Diagnosis. Must be using for	Coverage is provided for	By or in consultation with a	12 months		0
	Indications.			the treatment of	members 18 years of age or	neurologist or psychiatrist			
				hallucinations and delusions	older.				
				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					
PIRFENIDONE (ESBRIET)	1 - All FDA-approved			symptoms. Diagnosis. Must have	Coverage provided for	Pulmonologist	Initial: 6 months, Reauth: 12	For reauth: must have	0
EITIDOITE (EDDINET)	Indications.			diagnosis of idiopathic	members age 18 years and		months	documentation from	
	marcacions.			pulmonary fibrosis (IPF)	older.		months	prescriber indicating that	
				confirmed by either high-	oluci.			member still is a candidate for	
				resolution computed				treatment.	
				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%					

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
POLYPHARMACY - MULTIPLE ACH MEDICATIONS	1 - All FDA-approved Indications.			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 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.	
POSACONAZOLE (NOXAFIL)	1 - All FDA-approved Indications.		ergotamine, dihydroergotamine), HMG-	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
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

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
PREGABALIN (LYRICA)	1 - All FDA-approved			Diagnosis. For fibromyalgia:	For partial onset seizures,		12 months		0
	Indications.			must have trial and failure or	coverage is provided for				
				contraindication to	members 1 month of age and				
				gabapentin at a dose of at	older. For fibromyalgia, PHN,				
				least 1200mg/day or	DPN, and neuropathic pain				
				maximally tolerated dose in	associated with spinal cord				
				intolerant patients AND either					
				duloxetine or muscle relaxant	, ,				
				unless contraindicated. For	or older.				
				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.					
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		serum sickness or transfusion	
			infections, ocular herpes	sclerosis, member must have		neurologist or physician that		reaction due to serum protein	
			simplex, recent surgery,	tried and failed or have a		specializes in the treatment of		reaction, member must have	
			history of or the presence of a			multiple sclerosis, a		tried and failed 2	
			peptic ulcer, congestive heart			rheumatologist, allergist,		corticosteroids (e.g. IV	
			failure, hypertension, or	methylprednisolone, IV		dermatologist, immunologist,		methylprednisolone, IV	
				dexamethasone, or high dose		ophthalmologist,		dexamethasone, or high dose	
			from porcine sources, primary adrenocortical insufficiency or			pulmonologist, nephrologist		oral steroids) or has a	
			adrenocortical insufficiency of adrenocortical hyperfunction					contraindication to corticosteroid therapy. If the	
			are excluded.	medication for the treatment				member has a diagnosis of	
			are excluded.	of multiple sclerosis. For RA				atopic dermatitis, the membe	_
				(incl. Juvenile RA), psoriatic				is concurrently receiving	
				arthritis, ankylosing				maintenance therapy with	
				spondylitis, acute gouty				one (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	
				maintenance therapy with at				demonstrating neutropenia,	
				least one of the following: an				development of reactive	
				NSAID, DMARD (e.g.				plasmacytoid lymphocytes,	
				methotrexate, leflunomide,				and elevated erythrocyte	
				sulfasalazine) or biologic (e.g.				sedimentation rate or C-	
				adalimumab, etanercept,				reactive protein. For	
				infliximab, tofacitinib). For				ophthalmic diseases such as	
				collagen disease, member				severe acute and chronic	
				must have tried and failed or	1	l	<u> </u>	allergic and inflammatory	L

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
RESMETIROM (REZDIFFRA)	1 - All FDA-approved		Members with	Diagnosis. Medication must	Coverage is provided for	By or in consultation with a	12 months	For reauth: the member has	0
	Indications.		decompensated cirrhosis	be used in conjunction with	members 18 years of age and			received a clinical benefit	
				diet and exercise for the	older	gastroenterologist		demonstrated by either the	
				treatment of adults with				resolution of steatohepatitis	
				noncirrhotic nonalcoholic				and no worsening of liver	
				steatohepatitis (NASH) with				fibrosis or at least one stage	
				moderate to advanced liver				improvement in liver fibrosis	
				fibrosis (stage F2 to F3				and no worsening of	
				fibrosis) which has been				steatohepatitis.	
				confirmed by one of the					
				following (1, 2, or 3): 1) a liver					
				biopsy within the past 6					
				months with a NAFLD Activity					
				Score (NAS) of at least 4 and a					
				score of at least 1 in each NAS					
				component (steatosis,					
				ballooning degeneration, and					
				lobular inflammation) OR 2)					
				vibration-controlled transient					
				elastography (VCTE, e.g.					
				FibroScan) within the past 3					
				months with kPa greater than					
				or equal to 8.5 and controlled					
				attenuation parameter (CAP)					
				greater than or equal to 280					
				dB.m-1, OR 3) MRI with an					
				MRI-PDFF greater than or					
				equal to 8% liver fat.					
RIFAXIMIN (XIFAXAN)	1 - All FDA-approved			Diagnosis. For hepatic	Honotic oncombolomethy and	Honotic on conholomothy a by	Honotic oncombolomethy 12	For IBS-D: members who	0
RIFAXIIVIIN (XIFAXAN)	* *			-		Hepatic encephalopathy: by	Hepatic encephalopathy: 12		ľ
	Indications.			encephalopathy: must have		or in consultation with a	months, IBS-D: 2 weeks,	experience a recurrence of	
				trial and failure of lactulose.	older, Travelers diarrhea: 12	gastroenterologist,	Travelers diarrhea: 3 days	symptoms can be retreated	
				For diarrhea-predominant	years of age or older	hepatologist, or infectious		up to two times with the same	
				irritable bowel syndrome (IBS-		disease specialist, IBS-D:		dosage regimen. Reauth for	
				D): documentation of chronic		gastroenterologist		IBS-D: must have	
				IBS symptom diarrhea lasting				documentation from	
				at least 12 weeks and a trial				prescriber indicating	
				and failure of two				recurrence of IBS-D symptoms	5
				medications used in the				after a successful treatment	
				treatment of IBS-D (i.e.				with rifaximin.	
				loperamide, antispasmodics)					
				with inadequate responses or					
				significant side effect/toxicity					
				unless contraindicated. For					
				Traveler's diarrhea: must have					
				a trial and failure, intolerance,					
				or contraindication to one of					
				the following: a					
				fluoroquinolone (i.e.					
				ciprofloxacin, levofloxacin) or					
				azithromycin.					
				azidiromyciii.					
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Indication Indicator Off-Label Uses Exclusion Criteria Information Age Restriction Prescriber Restriction Coverage Duration Other Criteria Part Part	Part B Prerequisite
Associated Periodic Symdromes (CAPS), must have documented genetic mutation. Associated Periodic Symdromes (CAPS), must have documented genetic mutation in the Cold-Induced Auto-inflammatory Syndrome (ICAS1) also known as NLRP3 and a documented diagnosis of Familia Cold Autoinflammatory Syndrome (ECAS) or Muckle Wells Symdrome (MIVS). Member must have two or more of any of the CAPS-typical symptoms: urticaria-like rash, cold-triggered episodes, sensorineural hearing loss, musculoskeletal symptoms, chronic aseptic menigitis and skeletal abnormalities. Member must have documented baseline	
Syndromes (CAPS) , must have documented genetic mutation dollidren age 12 years and in the Cold-induced Auto- inflammatory Syndrome 1 (CIAS1) also known as NLRP3 and a documented diagnosis of Familial Cold Autoinflammatory Syndrome (FCAS) or Muckle Wells Syndrome (MWS). Member must have two or more of any of the CAPS-typical symptoms: uritcaria-like rash, cold-triggered episodes, sensorineural hearing loss, musculoskeletal symptoms, chronic asseptic meningitis and skeletal abnormalities. Member must have documented baseline	
documented genetic mutation in the Cold-Induced Auto- inflammatory Syndrome 1 (CIAS1) also known as NLRP3 and a documented diagnosis of Familial Cold Autoinflammatory Syndrome (FCAS) or Muckle Wells Syndrome (MWS), Member must have two or more of any of the CAPS-typical symptoms. uricaria-like rash, cold-triggered episodes, sensorineural hearing loss, musculoskeletal symptoms, chronic aseptic menigitis and skeletal abnormalities. Member must have documented baseline	
in the Cold-Induced Auto- inflammatory Syndrome 1 (CLAS1) also known as NLRP3 and a documented diagnosis of Familial Cold Autoinflammatory Syndrome (FCAS) or Muckle Wells Syndrome (MWS). Member must have two or more of any of the CAPS-typical symptoms: urticaria-like rash, cold-triggered episodes, sensorineural hearing loss, musculoskeletal symptoms, chronic aseptic meningitis and skeletal abnormalities. Member must have documented baseline	
inflammatory Syndrome 1 (CIASI) also known as NLRP3 and a documented diagnosis of Familial Cold Autoinflammatory Syndrome (FCAS) or Muckle Wells Syndrome (MWS). Member must have two or more of any of the CAPS-typical symptoms: urticaria-like rash, cold-triggered episodes, sensorineural hearing loss, musculoskeletal symptoms, chronic aseptic meningitis and skeletal abnormalities. Member must have documented baseline	
and a documented diagnosis of Familial Cold Autoinflammatory Syndrome (FCAS) or Muckle Wells Syndrome (MWS). Member must have two or more of any of the CAPS-typical symptoms: urticaria-like rash, cold-triggered episodes, sensorineural hearing loss, musculoskeletal symptoms, chronic aseptic meningitis and skeletal abnormalities. Member must have documented baseline	
and a documented diagnosis of Familial Cold Autoinflammatory Syndrome (FCAS) or Muckle Wells Syndrome (MWS). Member must have two or more of any of the CAPS-typical symptoms: urticaria-like rash, cold-triggered episodes, sensorineural hearing loss, musculoskeletal symptoms, chronic aseptic meningitis and skeletal abnormalities. Member must have documented baseline	l l
of Familial Cold Autoinflammatory Syndrome (FCAS) or Muckle Wells Syndrome (MWS). Member must have two or more of any of the CAPS-typical symptoms: urticaria-like rash, cold-triggered episodes, sensorineural hearing loss, musculoskeletal symptoms, chronic aseptic meningitis and skeletal abnormalities. Member must have documented baseline	
Autoinflammatory Syndrome (FCAS) or Muckle Wells Syndrome (MWS). Member must have two or more of any of the CAPS-typical symptoms: urticaria-like rash, cold-triggered episodes, sensorineural hearing loss, musculoskeletal symptoms, chronic aseptic meningitis and skeletal abnormalities. Member must have documented baseline	
(FCAS) or Muckle Wells Syndrome (MWS). Member must have two or more of any of the CAPS-typical symptoms: urticaria-like rash, cold-triggered episodes, sensorineural hearing loss, musculoskeletal symptoms, chronic aseptic meningitis and skeletal abnormalities. Member must have documented baseline	
Syndrome (MWS). Member must have two or more of any of the CAPS-typical symptoms: urticaria-like rash, cold-triggered episodes, sensorineural hearing loss, musculoskeletal symptoms, chronic aseptic meningitis and skeletal abnormalities. Member must have documented baseline	
must have two or more of any of the CAPS-typical symptoms: urticaria-like rash, cold-triggered episodes, sensorineural hearing loss, musculoskeletal symptoms, chronic aseptic meningitis and skeletal abnormalities. Member must have documented baseline	l
of the CAPS-typical symptoms: urticaria-like rash, cold-triggered episodes, sensorineural hearing loss, musculoskeletal symptoms, chronic aseptic meningitis and skeletal abnormalities. Member must have documented baseline	l
symptoms: urticaria-like rash, cold-triggered episodes, sensorineural hearing loss, musculoskeletal symptoms, chronic aseptic meningitis and skeletal abnormalities. Member must have documented baseline	
cold-triggered episodes, sensorineural hearing loss, musculoskeletal symptoms, chronic aseptic meningitis and skeletal abnormalities. Member must have documented baseline	
sensorineural hearing loss, musculoskeletal symptoms, chronic aseptic meningitis and skeletal abnormalities. Member must have documented baseline	
musculoskeletal symptoms, chronic aseptic meningitis and skeletal abnormalities. Member must have documented baseline	
chronic aseptic meningitis and skeletal abnormalities. Member must have documented baseline	
skeletal abnormalities. Member must have documented baseline	
Member must have documented baseline	
documented baseline	l
linflammatory markers	
including serum C-reactive	l
protein and serum amyloid A.	
For Deficiency of Interleukin-1	l
Receptor Antagonist (DIRA),	l
must have a confirmed	
diagnosis of DIRA as	
evidenced by a mutation in	l
the IL1RN gene. For recurrent	
pericarditis, must have a	l
history of trial and failure of at	l
Ieast 1 month,)
Indications.	l
the member has 4 to 14 older. and reauthorization: 12 having a reduced number of	
headache days per month.	
Must have a trial and failure month or a decrease in	l
of one beta-blocker and one migraine/headache severity. A	
anticonvulsant unless migraine is defined as a	
contraindicated or intolerant.	l
For acute treatment of two of the following	
migraine: Must have a history characteristics: unilateral	
of trial and failure,	
contraindication or quality, moderate or severe	
intolerance to at least one intensity (inhibits or prohibits	
triptan. daily activities), is aggravated	
by routine activity, nausea	
and/or vomiting, photophobia	
and phonophobia.	
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				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
RIOCIGUAT (ADEMPAS)	1 - All FDA-approved		Coverage will not be provided	Diagnosis. Pulmonary arterial		Prescribed by or in	Initial: 3 months, Reauth: 12	For reauth: documentation	0
	Indications.		for patients taking nitrates	hypertension (PAH) WHO		consultation with cardiologist	months	from prescriber that	
			(nitrates in any form) or a PDE	Group I confirmed by chart		or pulmonologist.		demonstrates member is	
			inhibitor (e.g. sildenafil).	documentation of right-heart				tolerating and receiving	
				catheterization (RHC)				clinical benefit from	
				indicating a mean pulmonary				treatment	
				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.					
RISANKIZUMAB-RZAA	1 - All FDA-approved			Diagnosis. For plaque		By or in consultation with a	12 months	For reauthorization: must	0
(SKYRIZI)	Indications.			psoriasis: minimum BSA	age or older.	rheumatologist, dermatologist		have 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):					
				history of trial and failure,					
				contraindication, or					
				intolerance to 2 of the					
				following therapy options:					
				aminosalicylates,					
				ammosancylates,	I .	I		1	l

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
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	autosomal recessive SMA.		consultation with neurologist,		that the patient is responding	
			concomitantly taking	Baseline assessment motor		or pediatric neurologist.		to the medication as	
			nusinersen.	milestone score from ONE of				demonstrated by clinically	
				the following assessments:				significant improvement or	
				Hammersmith Functional				maintenance of function from	
				Motor Scale Expanded				pretreatment baseline status	
				(HFMSE), Hammersmith				using the same exam as	
				Infant Neurologic Exam				performed at baseline	
				(HINE), Upper limb module				assessment (progression,	
				(ULM) score, Children?s				stabilization, or decreased	
				Hospital of Philadelphia Infant				decline in motor function).	
				Test of Neuromuscular					
				Disorders (CHOP INTEND), or					
				Six-minute walk test.					
ROFLUMILAST (DALIRESP)	1 - All FDA-approved		Moderate to sever liver	Diagnosis of GOLD Stage III or			12 months	For reauthorization must have	0
	Indications.			IV COPD associated with				documentation from	
				chronic bronchitis.				prescriber indicating	
				Documentation of COPD				improvement in condition.	
				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.	<u> </u>		1		

riber Restriction in consultation with a ologist. Coverage Duration 12 months	Other Criteria For reauthorization: Documentation from the provider that the member had	Part B Prerequisite 0
	Documentation from the	0
ologist.		
	provider that the member had	
	a positive clinical response	
	and tolerates therapy	
	supported by at least one of	
	the following: a 2 point	
	improvement in the member's	
	total MG-ADL score OR a 3 or	
	more point improvement in	
	QMG total score.	
in consultation with a 12 months		0
ologist.		
		the following: a 2 point improvement in the member's total MG-ADL score OR a 3 or more point improvement in QMG total score.

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	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.	0
SAPROTERIN DIHYDROCHLORIDE (KUVAN)				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) or rituximab.	members 18 years of age and 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	

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
SECUKINUMAB (COSENTYX)	1 - All FDA-approved			Diagnosis. For Psoriatic	Must be 2 years of age or	By or in consultation with a	12 months	For reauth: must have	0
	Indications.			arthritis (PsA): for mild to	older.	rheumatologist,		documentation from	
				moderate axial or enthesitis,		gastroenterologist, or		prescriber indicating	
				must have a history of trial		dermatologist.		stabilization or improvement	
				and failure, contraindication,				in condition.	
				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					
				ankylosing spondylitis (AS),					
				non-radiographic axial					
				spondyloarthritis (nr-axSpA),					
				and enthesitis-related arthritis	5				
				(ERA): history of trial and					
				failure, contraindication, or					
				intolerance to a four-week					
				trial each of at least 2 NSAIDs.					
				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	d				
				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,					
SELEXIPAG (UPTRAVI)	1 - All FDA-approved			Diagnosis. Pulmonary arterial		Prescribed by or in	Initial authorization: 3 months		0
	Indications.			hypertension (PAH) WHO		consultation with cardiologist	Reauthorization: 12 months	documentation from	
				Group I confirmed by chart		or pulmonologist.		prescriber that demonstrates	
				documentation of right-heart				member is tolerating and	
				catheterization (RHC)				receiving clinical benefit from	
				indicating a mean pulmonary				treatment	
				arterial pressure greater than					
				20 mmHg, pulmonary vascula	r				
				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.					
	1			recinculatiography.	1	1	ı	1	L

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
Group SILDENAFIL CITRATE REVATIO)	Indication Indicator 1 - All FDA-approved Indications.	Off-Label Uses				Prescriber Restriction Prescribed by or in consultation with a pulmonologist or cardiologist	Coverage Duration Initial: 3 months, Reauth: 12 months	Other Criteria For reauth: documentation from prescriber that demonstrates member is tolerating and receiving clinical benefit from treatment	Part B Prerequisite 0
SODIUM OXYBATE (XYREM)	1 - All FDA-approved Indications.			recommended, must have documentation of echocardiography. 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.	members 7 years of age or	By or in consultation with a neurologist or sleep specialist	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	members who are age- appropriate according to	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 ageappropriate according to	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

				Required Medical					
•	Indication Indicator	Off-Label Uses	Exclusion Criteria		Age Restriction	Prescriber Restriction	Coverage Duration		Part B Prerequisite
SOMATROPIN (GENOTROPIN)	· · ·		Coverage will not be provided			1 ·	6 months	For reauth for pediatric GHD,	0
	Indications.		for members with active	required for all diagnoses		endocrinologist or		Turner and Noonan	
			malignancy, active	except Adult Growth		neonatologist.		syndromes, SGA, Prader-Willi	
			proliferative or severe non-	Hormone Deficiency (GHD). Documentation that				syndrome, and ISS:	
			proliferative diabetic					Documentation the patient	
			retinopathy, pediatric member with closed	epiphyses are open for all pediatric indications. For				has open epiphyses. For reauth for adult GHD: current	
			epiphysis, members with	pediatric GHD: a height				IGF-1 level is normal for age	
			Prader-Willi who are severely	1 ·				and gender (does not apply to	
			obese or have severe	standard deviations below the				patients with structural	
			respiratory impairment.	mean for age and gender,				abnormality of the	
			respiratory impairment.	documentation of growth				hypothalamus/pituitary and 3	
				velocity, skeletal maturation,				or more pituitary hormone	
				2 provocative stimulation				deficiencies and childhood-	
				tests which demonstrate GHD				onset growth hormone	
				through peak growth				deficiency with congenital	
				hormone concentrations less				abnormality of the	
				than 10 ng/ml or IGF-1 or				hypothalamus/pituitary). For	
				IGFBP-3 levels or only one				reauth for Prader Willi:	
				stim test is needed in the				documentation growth	
				presence of a pituitary				hormone has resulted in an	
				abnormality. For Small for				increase in lean body mass or	
				Gestational Age (SGA), a				decrease in fat mass.	
				height greater than or equal					
				to 2 standard deviations					
				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					
SOTATERCEPT-CSRK	1 - All FDA-approved			Diagnosis. Pulmonary arterial		Prescribed by or in	Initial: 3 months, Reauth: 12		0
(WINREVAIR)	Indications.			hypertension (PAH) WHO		consultation with cardiologist or pulmonologist	months	from prescriber that	
				Group I confirmed by chart		or pulmonologist		demonstrates member is tolerating and receiving	
				documentation of right-heart				clinical benefit from	
				catheterization (RHC)					
				indicating a mean pulmonary arterial pressure greater than				treatment	
				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					
				standard of care therapy (e.g.					
				ERA or PDE-5 inhibitor)					

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Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
PARSENTAN (FILSPARI)	1 - All FDA-approved			Diagnosis of primary	Coverage is provided for	By or in consultation with a	Initial: 6 months. Reauth: 12	For reauth: must have a	0
	Indications.			immunoglobulin A	members 18 years of age or	nephrologist.	months	decrease from baseline in	
				nephropathy (IgAN) that has	older.			total urine protein or UPCR.	
				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					
ECOLINAND CD7O (CDEV//	GO) 1 - All FDA-approved			an ACE inhibitor or ARB. Diagnosis. For treatment of a	Coverage is provided for	By or in consultation with a	For a flare: one treatment	For reauth: documentation of	10
SOLIIVIAD-SBZO (SPEVIC	Indications.			generalized pustular psoriasis		dermatologist	course (up to 2 infusions over		•
	indications.			(GPP) flare, must have a	older and weighing at least 40	_	2 weeks). For maintenance:	flares while on treatment	
				moderate-to-severe flare			12 months	nares while on treatment	
					kg.		12 months		
				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).					
IRIPENTOL (DIACOMIT)	1 - All FDA-approved			Diagnosis. Must have had an	Member must be 6 months of	By or in consultation with a	12 months		0
2 22 (51/1001/111)	Indications.			inadequate response or	age or older	neurologist			-
				intolerance to two generic	-0-0.0.00				
				antiepileptic drugs (e.g.					
				valproate, topiramate,					
				clobazam). Must be using in					
				combination with clobazam.					

SUZETRIGINE (JOURNAVX)	1 - All FDA-approved	Off-Label Uses	Exclusion Criteria		Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
·									·
	Local transfer or a			Must have a diagnosis of	Coverage is provided for		14 Days	For reauthorization:	0
	Indications.			moderate-to-severe acute	members 18 years of age and			Documentation that the	
i i				pain. The prescriber attests	older			member is experiencing a new	
,				that the episode of acute pain				episode of moderate-to-	
				is anticipated to last less than				severe acute pain, separate	
				one month and the member				and distinct from the previous	
				has tried and failed within the				episode. The prescriber	
				previous 30 days or has a				attests that the episode of	
!				contraindication to either				acute pain is anticipated to	
!				TWO alternative pain				last less than one month and	
				medications for moderate				the member has tried and	
				pain (e.g. acetaminophen,				failed within the previous 30	
!				NSAIDs) or ONE alternative				days or has a contraindication	
				pain medication for severe				to either TWO alternative pain	
				pain (e.g. NSAID, opioid).				medications for moderate	
!								pain (e.g. acetaminophen,	
								NSAIDs) or ONE alternative	
								pain medication for severe	
								pain (e.g. NSAID, opioid).	
TADALAFIL (ADCIRCA)	1 - All FDA-approved		Coverage will not be provided	Diagnosis. Pulmonary arterial		Prescribed by or in	Initial: 3 months, Reauth: 12	For reauth: documentation	0
·	Indications.		for patients taking nitrates	hypertension (PAH) WHO		consultation with a	months	from prescriber that	
			(nitrates in any form) or a	Group I confirmed by chart		pulmonologist or cardiologist		demonstrates member is	
				documentation of right-heart				tolerating and receiving	
			(e.g. Adempas).	catheterization (RHC)				clinical benefit from	
			(e.g. Adempas).	indicating a mean pulmonary				treatment	
								l eatment	
				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.					
TADALAFIL (CIALIS)	1 - All FDA-approved			Diagnosis of benign prostatic			12 months		0
	Indications.			hyperplasia (BPH) and must					
				have a trial and failure of at					
				least two alternative					
				medications in the following					
				_					
				classes: alpha-1 adrenergic					
				blockers or 5-alpha reductase					
TACINAEL TECO: (************************************	4 411504			inhibitors.		B 1 10 10 10 10 10 10 10 10 10 10 10 10 1	12 11	5 B 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	
	1 - All FDA-approved			Diagnosis. Must submit chart	Coverage is provided for	By or in consultation with a	12 months	For Reauth: documentation	ľ
	Indications.			documentation describing	members 3 years of age or	neurologist or a physician		from prescriber indicating	
!				how diagnosis was confirmed	older.	who specializes in sleep		stabilization or improvement	
				(e.g. sleep-wake logs,		medicine		in condition.	
!				melatonin secretion					
				abnormalities, or progress					
				notes, etc.)					
TEDUGLUTIDE (GATTEX)	1 - All FDA-approved		Active intestinal obstruction	Diagnosis. For diagnosis of		By or in consultation with a	12 months	For reauthorization: must	0
	Indications.		or active gastrointestinal	short bowel syndrome,		gastroenterologists		have documentation from	
!			malignancy.	member must be receiving		3		prescriber indicating	
				parenteral support.				stabilization or improvement	
!				ραι επτεται δυρροιτ.					
TELOTRICTAT (VERNASI O)	1 All EDA amana :!		+	Diagnosis	Coverage is musual and form	Du on in consultation with	6 months	in condition.	0
	1 - All FDA-approved			Diagnosis.		·	6 months	For reauth: documentation of	U
	Indications.		1		members 18 years of age and	oncologist	1	improvement or stabilization.	

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria		Age Restriction	Prescriber Restriction	<u> </u>		Part B Prerequisite
TETREBENAZINE (XENAZINE)	1 - All FDA-approved Indications.	Off-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	Coverage is provided for members 18 years of age or older.	By or in consultation with a neurologist	12 months	Maximum dose approved is 100mg/day. For reauthorization: must have documentation from prescriber indicating stabilization or improvement in condition.	O Part B Prerequisite
TOFACITINIB (XELJANZ)	1 - All FDA-approved Indications.			Diagnosis. Must have history 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	dermatologist, rheumatologist or gastroenterologist.		Reauth: Documentation from the prescriber indicating stabilization or improvement in condition.	0
TOLVAPTAN (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 have an estimated glomerular filtration rate (eGFR) greater than or equal to 25 mL/min/1.73m^2 and at least one of the following: 1. Mayo classification 1C, 1D, or 1E 2. a historical rate of eGFR decline (greater than or equal to 3 ml/min /1.73 m^2 per year)	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
TRIENTINE 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

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
TROFINETIDE (DAYBUE)	1 - All FDA-approved		0	Diagnosis. Documentation of	Coverage is provided for	By or in consultation with a	12 months	0	0
	Indications.			a diagnosis of typical Rett	members 2 years of age or	pediatric neurologist or			
				syndrome according to the	older.	neurologist			
				Rett Syndrome Diagnostic					
				Criteria with a documented					
				disease-causing mutation in					
				the MECP2 gene.					
UBROGEPANT (UBRELVY)	1 - All FDA-approved			Diagnosis. Must have a history	Coverage is provided for		12 months	For reauth: documentation of	0
	Indications.			of trial and failure,	members 18 years of age and			improvement or stabilization.	
				contraindication, or	older.				
				intolerance to at least one					
				triptan.					
UPADACITINIB (RINVOQ)	1 - All FDA-approved			Diagnosis. For rheumatoid	For psoriatic arthritis and	By or in consultation with a	12 months	For reauthorization: must	0
	Indications.			arthritis (RA), psoriatic	polyarticular juvenile	rheumatologist,		have documentation from	
				arthritis (PsA), ankylosing	idiopathic arthritis: 2 years or	- ·		prescriber indicating	
				spondylitis (AS), non-	older, For atopic dermatitis:	gastroenterologist.		stabilization or improvement	
				radiographic axial	12 years or older. All other			in condition.	
				spondyloarthritis (nr-axSpA),	indications: 18 years and				
				ulcerative colitis (UC), and	older.				
				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).	1				1

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
USTEKINUMAB (STELARA) SQ	1 - All FDA-approved			Diagnosis. For Psoriatic	Must be 6 years of age or	By or in consultation with a	12 months	For reauth: must have	0
	Indications.			arthritis (PsA): one of the	older.	rheumatologist,		documentation from	
				following: 1) members with		gastroenterologist, or		prescriber indicating	
				axial or enthesitis must have a		dermatologist.		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					
				(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:					
V CO KIT	4. All EDA amanana	 		aminosalicylates,			42		
V-GO KIT	1 - All FDA-approved			Must have documentation of			12 months		0
	Indications.			previous insulin use.		<u> </u>	<u> </u>	<u> </u>	

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
VALBENAZINE (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	2				
				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: must have					
				chart documentation of					
				involuntary athetoid or					
				choreiform movements and					
				has a history of treatment					
				with neuroleptic agent (i.e.					
				antipsychotic). Adjustments					
				to possible offending					
				medication such as dose					
\(\(\text{FP}\)\(\text{C}\)\(\text{I}\)	4 411504			reduction or discontinuation			42		
VERICIGUAT (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					
			1	and a beta blocker.	<u> </u>	1	1		1

Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Required Medical Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
VIGABATRIN (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	members 1 month of age or older.	By or in consultation with a neurologist.	12 months		0
VILAZODONE (VIIBRYD)	1 - All FDA-approved Indications.			from trial above). 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
VORICONAZOLE 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
VORTIOXETINE (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.	micerious discuse specialist	12 months		0
VOSORITIDE (VOXZOGO)	1 - All FDA-approved Indications.			Diagnosis confirmed by documentation of one of the following: 1. genetic testing showing mutation in the FGFR3 gene or 2. radiographic assessment confirming achondroplasia (e.g. short, robust tubular bones, squared off iliac wings, flat horizontal acetabule, ect.). Documentation the member has open epiphyses.		Prescribed by or in consultation with an endocrinologist, geneticists, or other practitioner with expertise in the management of achondroplasia	12 Months	For reauth: documentation of both of the following: 1. improvement or stabilization. 2. The member's epiphyses remain open.	0
XANOMELINE/TROSPIUM (COBENFY)	1 - All FDA-approved Indications.			Diagnosis. Documentation of trial and failure of at least two of the following generic atypical antipsychotics: olanzapine, quetiapine, paliperidone, risperidone, aripiprazole, or ziprasidone.	•		12 months		0

				Required Medical					
Group	Indication Indicator	Off-Label Uses	Exclusion Criteria	Information	Age Restriction	Prescriber Restriction	Coverage Duration	Other Criteria	Part B Prerequisite
ZURANOLONE (ZURZUVAE)	1 - All FDA-approved			Diagnosis of postpartum	Coverage is provided for	Prescribed by or in	14 days		0
	Indications.			depression (PPD) with onset	members 18 years of age and	consultation with a			
				during pregnancy or within 4	older.	psychiatrist or OB/GYN			
				weeks postpartum.					
				Documentation of current					
				depressive symptoms					
				consistent with a diagnosis of					
				major depressive disorder					
				with peripartum onset.					
				Baseline assessment using a					
				validated depression rating					
				scale indicates at least					
				moderate severity depression					
				(e.g. PHQ-9 score of 10 or					
				higher, EPDS score of 14 or					
				higher).					