Appendix Y



Physician-Administered Drug Prior Authorization Procedures, Coverage Policies and Drug Utilization Criteria For the Health First Colorado Medical Benefit

Physician-Administered Drugs (PADs) requiring a prior authorization (PA) for the Health First Colorado medical benefit are listed in this document. Prior authorization criteria is based on FDA product labeling, CMS approved compendia, clinical practice guidelines, and peer-reviewed medical literature.

Physician-Administered Drugs and Medical Billing

PADs include any medication or medication formulation that requires administration by a healthcare professional, including cases where FDA package labeling for a medication specifies that administration should be performed by or under the direct supervision of a healthcare professional. PADs administered in a provider's office or clinic should be billed through the Health First Colorado medical benefit using the standard buy-and-bill process following procedures in the PAD Billing Manual (found on the PAD Resources Page at https://www.colorado.gov/hcpf/physician-administered-drugs).

PAD criteria listed on Appendix Y applies specifically to medications billed through the Health First Colorado medical benefit.

• Only PADs administered by a healthcare professional in the member's home or in a long-term care facility should be billed through the Health First Colorado pharmacy benefit (see "Medical VS. Pharmacy Benefit Medication Coverage" section below).

Prior Authorization Procedures

• Prior authorization requests (PAR) may be submitted via the Acentra PAR portal at https://portal.kepro.com/. For PA assistance or questions, you may contact Acentra via the following methods:

Phone: (720) 689 - 6340 Fax: (833) 923 - 2359

Email: COproviderissue@kepro.com

- PA forms can be signed by anyone who has authority under Colorado law to prescribe the medication. Assistants of authorized persons cannot sign the PA form.
- Physicians or assistants who are acting as the agents of the physicians may request a PA by phone.
- Please note that initiating therapy with a requested drug product, including non-preferred drugs, prior to a PA request being reviewed and approved does not necessitate approval of the PA request. This includes initiating therapy by administration in the inpatient setting, by using office samples or by any other means.
- All PA requests are coded online into the PA system.

Trial and Failure

• Generally, failure is defined as lack of efficacy, allergy, intolerable side effects, contraindication to therapy or significant drug-drug interaction. For medications that use a varying definition of failure, the definition will be noted in the medication's specific criteria, below.

Medical VS. Pharmacy Benefit Medication Coverage

- For more information about pharmacy benefits versus medical benefits please see the Pharmaceutical Benefit Help Guide (found on the PAD resources page at https://hcpf.colorado.gov/physician-administered-drugs).
- Medications administered by a healthcare professional or self-administered in the member's home or long-term care facility should be billed through the Health First Colorado pharmacy benefit following the standards and procedures outlined in the Pharmacy Billing Manual (found on the Pharmacy Resources Page at https://hcpf.colorado.gov/pharmacy-resources).
- PADs are medications administered in a doctor's office, clinic, outpatient hospital or dialysis unit are only to be billed by those facilities through the Health First Colorado medical benefit using the standard buy-and-bill process and following procedures outlined in the PAD Billing Manual (located at https://www.colorado.gov/hcpf/physician-administered-drugs). PAD criteria listed on Appendix Y applies specifically to drug products when billed through the Health First Colorado medical benefit, when administered in the clinic or office setting.

HCPCS	Drug	Criteria	PAR Length
J0172	Aduhelm (aducanumab-avwa)	Aduhelm (aducanumab-avwa) may be approved if the member meets ALL the following criteria:	
		a. Member has documented diagnosis of mild cognitive impairment or mild dementia	See criteria
		stage of Alzheimer's disease, the population in which treatment was initiated in	
		clinical trials, as evidenced by ALL the following:	
		i. Positron Emission Tomography (PET) scan OR lumbar puncture	
		positive for amyloid beta plaque	
		ii. Clinical Dementia Rating global score (CDR-GS) of 0.5 or 1 (available	
		at https://otm.wustl.edu/cdr-terms-agreement/)	
		iii. Mini-Mental State Examination (MMSE) score of 24-30 OR Montreal	
		Cognitive Assessment (moCA) Test score of 19-25	
		AND	
		b. Member is ≥ 50 years of age AND	
		c. The prescriber attests that member has been counseled on the approval and safety	
		status of Aduhelm (aducanumab-avwa) being approved under accelerated approval	
		based on reduction in amyloid beta plaques AND	
		d. Prior to initiation of medication, the prescriber attests that the member meets ALL	
		the following:	
		i. Member has had a brain MRI within the prior one year to treatment	
		initiation, showing no signs or history of localized superficial siderosis,	
		≥ 10 brain microhemorrhages, and/or brain hemorrhage > 1 cm	
		ii. Attestation that MRI will be completed prior to the 7th (1st dose at 10	
		mg/kg) and 12th (6th dose at 10 mg/kg) infusion	

HCPCS	Drug	Criteria	PAR Length
	8	AND	
		e. Member does not have any of the following:	
		i. Any medical or neurological condition other than Alzheimer's Disease	
		that might be a contributing cause of the subject's cognitive	
		impairment including (but not limited to) stroke/vascular dementia,	
		tumor, dementia with Lewy bodies [DLB], frontotemporal dementia	
		[FTD] or normal pressure hydrocephalus	
		ii. Contraindications to PET, CT scan, or MRI	
		iii. History of or increased risk of amyloid related imaging abnormalities	
		ARIA-edema (ARIA-E) or ARIA-hemosiderin deposition (ARIA-H)	
		iv. History of unstable angina, myocardial infarction, chronic heart failure,	
		or clinically significant conduction abnormalities, stroke, transient	
		ischemic attack (TIA), or unexplained loss of consciousness within 1	
		year prior to initiation of medication	
		v. History of bleeding abnormalities or taking any form of	
		anticoagulation therapy	
		AND	
		f. Medication is prescribed by or in consultation with a neurologist	
		AND	
		g. The prescribed regimen meets FDA-approved labeled dosing:	
		 i. <u>Infusion 1 and 2</u>: 1 mg/kg over approximately 1 hour every 4 weeks ii. Infusion 3 and 4: 3 mg/kg over approximately 1 hour every 4 weeks 	
		ii. <u>Infusion 3 and 4</u> : 3 mg/kg over approximately 1 hour every 4 weeks iii. <u>Infusion 5 and 6</u> : 6 mg/kg over approximately 1 hour every 4 weeks	
		iv. Infusion 7 and beyond: 10 mg/kg over approximately 1 hour every 4 weeks	
		weeks	
		WEEKS	
		Initial approval period: 6 months	
		Second prior authorization: an additional 6 months of therapy may be approved with provider attestation that a follow-up MRI will be (or has been) completed prior to the 7th infusion	
		Subsequent approval: an additional 6 months of therapy may be approved with provider attestation that a follow-up MRI will be (or has been) completed prior to the 12th	
		infusion	

HCPCS	Drug	Criteria	PAR Length
J7171	Adzynma (apadamtase alfa)	Maximum dose: 10 mg/kg IV every 4 weeks The above coverage standards will continue to be reviewed and evaluated for any applicable changes due to the evolving nature of factors including disease course, available treatment options and available peer-reviewed medical literature and clinical evidence. If request is for use outside of stated coverage standards, support with peer reviewed medical literature and/or subsequent clinical rationale shall be provided and will be evaluated at the time of request. Continued approval for this indication may be contingent upon verification of clinical benefit in confirmatory trial(s). Adzynma (apadamtase alfa) may be approved if the following criteria are met: a. Member is ≥ 2 years of age AND b. Member has a diagnosis of congenital thrombotic thrombocytopenic purpura (cTTP) confirmed by genetic testing indicating severe deficiency of ADAMTS13 protease and/or based on clinical judgment, AND c. The requested medication is being prescribed by or in consultation with a	One year
J0179	Beovu (brolucizumab)	hematologist. Maximum dose: Prophylactic therapy: 40 IU/kg weekly On-demand therapy: 40 IU/kg/day Beovu (brolucizumab) may be approved if all the following criteria are met: a. Member is 18 years of age or older AND	See criteria
		b. Member has a diagnosis of one of the following: i. Neovascular (wet) age-related macular degeneration (nAMD) ii. Diabetic macular edema (DME) AND c. Member does not have any of the following: i. Ocular or periocular infection ii. Active intraocular inflammation iii. Hypersensitivity to the requested medication AND d. Member's best corrected visual acuity (BCVA) is measured at baseline and throughout treatment AND	

HCPCS	Drug	Criteria	PAR Length
		e. Requested medication will not be used with other ophthalmic vascular endothelial growth factor (VEGF) inhibitors AND f. Documentation of trial and failure* with bevacizumab containing product AND g. Documentation of the dosing regimen that is being requested	
		Initial approval: 6 months	
		Reauthorization may be approved if the following criteria are met: a. Documentation of improvement or stabilization of disease state and visual status AND b. Documentation of ongoing treatment regimen AND c. Prescriber attests that the member has had no significant adverse effects or drug toxicity, such as endophthalmitis and retinal detachments, increase in intraocular pressure, arterial thromboembolic events (ATE), retinal vasculitis and/or retinal vascular occlusion AND d. Prescriber attests requested medication will not be used with other ophthalmic VEGF inhibitors	
		Maximum dose: AMD: 6 mg every 4 weeks x 3 doses, then 6 mg every 8-12 weeks DME: 6 mg every 6 weeks x 5 doses, then 6 mg every 8-12 weeks	
		*Failure is defined as lack of efficacy with a three-month trial, contraindication to therapy, allergy, intolerable side effects, or significant drug-drug interaction	
	BONE RESORPTION INHIBITORS	Prolia (denosumab) may be approved for members meeting all the following criteria: a. Member has one of the following diagnoses:	One year
J0897	Prolia,	i. Postmenopausal osteoporosis with high fracture risk	
	Xgeva	ii. Osteoporosis	
	(denosumab)	iii. Bone loss in men receiving androgen deprivation therapy in prostate cancer	
		iv. Bone loss in women receiving adjuvant aromatase inhibitor therapy for	
		breast cancer	
		OR	
		b. Member is considered very high risk for fracture defined as any one of the	
		following: a fracture within the past 12 months, experience of fractures while receiving approved osteoporosis therapy (i.e.), a history of multiple fractures,	
		experience of a fracture while receiving medications that cause skeletal harm (e.g.	

HCPCS	Drug	Criteria	PAR Length
		long-term glucocorticoids), very low T-score (e.g. < -3.0), high risk for falls or a history of injurious falls, or very high fracture probability by FRAX® AND c. Member has serum calcium greater than 8.5mg/dL AND	7 67
		 d. Member is taking calcium 1000 mg daily and at least 400 IU vitamin D daily AND e. For members not considered very high risk of fracture, member has trial and failure of bisphosphonate for one year (Failure is defined as: lack of efficacy, allergy, intolerable side effects, or significant drug-drug interaction) AND 	
		f. Member meets ANY of the following criteria: i. has a history of an osteoporotic vertebral or hip fracture ii. has a pre-treatment T-score of < -2.5 iii. has a pre-treatment T-score of < -1 but > -2.5 AND either of the following: 1. Pre-treatment FRAX score of > 20% for any major fracture 2. Pre-treatment FRAX score of > 3% for hip fracture iv. Maximum dose of medication is 60mg every 6 months g. Member who is at very high risk of fracture and is currently stable on medication	
		 May continue to receive prior authorization approval to continue. Mgeva (denosumab) may be approved if member meets ONE of the following indications: a. Prevention of skeletal-related events in members with multiple myeloma or in members with bone metastasis from solid tumors b. Giant cell tumor of bone that is unresectable or where surgical resection is likely to result in severe morbidity c. Hypercalcemia of Malignancy, refractory to bisphosphonate therapy d. If member is currently receiving and stabilized on medication, they may continue to receive prior authorization approval to continue. 	
J0585, J0586, J0587, J0588	BOTULINUM TOXIN AGENTS Botox (onabotulinumtoxinA) Dysport (abobotulinumtoxinA) Myobloc (rimabotulinumtoxinB) Xeomin (incobotulinumtoxinA)	Botulinum toxin agents may be approved if the member meets the following criteria: Botox (onabotulinumtoxinA) may be approved if the member meets ALL the following criteria: a. If administered for Chronic Migraine, prophylaxis i. Member is 18 years of age or older AND	One year

HCPCS	Drug	Criteria	PAR Length
		ii. Member has a diagnosis of chronic migraine, which is defined as headaches	
		occurring 15 days or more monthly, where at least 8 of these days per	
		month for at least 3 months are migraine days with or without aura AND	
		iii. Member has trial and failure of topiramate AND	
		iv. Dosing interval no sooner than every 12 weeks	
		v. Reauthorization requests may be approved if member has shown a clinical	
		reduction in number of migraine days per month OR	
		b. If administered for one of the following indications, member must meet the	
		following age requirements and dosing must be no sooner than every 12 weeks	
		i. <u>Overactive Bladder</u>	
		1. Member is 18 years of age or older	
		ii. <u>Spasticity</u>	
		1. Member is 2 years of age or older	
		iii. <u>Cervical Dystonia</u>	
		1. Member is 16 years of age or older	
		iv. <u>Primary Axillary Hyperhidrosis</u> 1. Member is 18 years of age or older	
		71.1	
		v. <u>Blepharospasm and Strabismus</u> 1. Member is 12 years of age or older	
		1. Wichioel is 12 years of age of older	
		Dysport (abobotulinumtoxinA)may be approved if the member meets ALL the following criteria	
		for each indication:	
		a. If being administered for <u>cervical dystonia</u>	
		i. Member has a diagnosis of cervical dystonia AND	
		ii. Member is 18 years of age or older AND	
		iii. Dosing interval is no sooner than every 12 weeks AND	
		iv. Initial dose of 500 units followed by a maximum maintenance dose of 1000	
		units administered intramuscularly	
		OR	
		b. If being administered for spasticity	
		i. Member is 2 years of age or older AND	
		ii. Dosing interval is no sooner than every 12 weeks	
		iii. Maximum dose is 1500 units administered intramuscularly	
		Myobloc (rimabotulinumtoxinB) may be approved if the member meets ALL the following	
		criteria:	

HCPCS	Drug	Criteria	PAR Length
		a. Member is 18 years of age or older AND	
		b. If being administered for <u>cervical dystonia</u>	
		i. Member has a diagnosis of cervical dystonia AND	
		ii. Dosing interval is no sooner than every 12 weeks AND	
		iii. Maximum dose of 10,000 units	
		OR	
		c. If being administered for chronic sialorrhea	
		i. Member has a diagnosis of chronic sialorrhea AND	
		ii. Dosing interval is no sooner than every 12 weeks AND	
		iii. Maximum Initial dose is 3,000 units	
		Xeomin (incobotulinumtoxinA) may be approved if member meets ALL the following criteria for each indication:	
		a. If being administered for one of the following indications:	
		1. <u>Blepharospasm</u>	
		2. <u>Cervical dystonia</u>	
		ii. Member is at least 18 years of age AND	
		iii. Dosing frequency is no sooner than every 12 weeks AND	
		iv. If administered for blepharospasm, maximum dose 100 units per treatment	
		session	
		b. If being administered for the <u>chronic sialorrhea</u>	
		i. Member is 2 years of age or older AND	
		ii. Member weighs more than 12 kg AND	
		iii. Dosing frequency is no sooner than every 16 weeks AND	
		iv. Maximum dose of 100 units	
		c. If administered for the treatment of <u>upper limb spasticity</u>	
		i. Member is 2 years of age or older AND	
		ii. For members between 2 and 17 years of age, spasticity is not caused by	
		cerebral palsy AND	
		iii. Dosing frequency is no sooner than every 12 weeks AND	
		iv. Maximum dose of 200 units per single upper limb, or 400 units total	
		Not approved for Cosmetic Purposes	
J2786	Cinqair (reslizumab)	Cinqair (reslizumab) may be approved for members meeting all the following criteria:	One year
	_ ` ` ′	a. Member is 18 years of age or older AND	

HCPCS	Drug	Criteria	PAR Length
		b. Member has diagnosis of severe asthma with an eosinophilic phenotype AND c. Member has a blood eosinophil count of greater than or equal to 400 cells/mcL AND d. Medication is being used as a maintenance adjunctive therapy AND e. Member's symptoms remain uncontrolled despite adherence to concomitant treatment with a medium to high-dose inhaled corticosteroids and long acting beta2- agonist AND f. Member has uncontrolled disease characterized by the following: i. Asthmatic symptoms occurring throughout the day ii. Nighttime awakenings occurring 7 times per week iii. Use of Short Acting Beta-Agonist for symptom control several times per day iv. Lung Function, characterized by FEV1 is less than 60% v. Asthma exacerbations requiring oral systemic corticosteroids, occurring more frequently and intensely than mild or moderate asthma AND g. Baseline FEV1 and frequency of asthma exacerbations per month are provided AND h. Maximum dose of 3 mg/kg every 4 weeks i. Reauthorization may be approved if member meets one of the following: i. Improvement in lung function, measured in FEV1 OR ii. Reduction in the number of asthma exacerbations, defined as a decrease in use of oral or systemic corticosteroids and/or reduced asthma related hospitalizations and/or ER visits	
J1442 Q5125 Q5101 J2506 Q5108 Q5111 Q5122 Q5130 Q5127	COLONY STIMULATING FACTORS Neupogen (filgrastim) Releuko (filgrastim-ayow) Zarxio (filgrastim-sndz) Neulasta (pegfilrastim) Fulphila (pegfilgrastim-jmdb) Udenyca (pegfilgrastim-cbqv) Nyvepria (pegfilgrastim-apgf) Fylnetra (pegfilgrastim-pbbk) Stimufend (pegfilgrastim-fpgk)	Filgrastim (Neupogen brand/generic and filgrastim-containing products or biosimilars) may receive approval if the following criteria are met: a. The prescribed agent is one of the following preferred filgrastim products: Granix (tbo-filgrastim), Nivestym (filgrastim-aafi) OR b. If the prescribed agent is brand Neupogen or a filgrastim product formulation that is not a preferred filgrastim product, then the member has trialed and failed at least one favored filgrastim product. Failure is defined as lack of efficacy or intolerable side effects with the favored filgrastim product formulation.	One year

HCPCS	Drug	Criteria	PAR Length
		Pegfilgrastim (Neulasta brand/generic and pegfilgrastim-containing products or biosimilars) may receive approval if the following criteria are met: a. The prescribed agent is the following preferred pegfilgrastim product: Ziextenzo (pegfilgrastim-bmez) OR b. If the prescribed agent is brand Neulasta or a pegfilgrastim product formulation that is not the preferred pegfilgrastim product, then the member has trialed and failed the favored pegfilgrastim product. Failure is defined as lack of efficacy or intolerable side effects with the favored pegfilgrastim product formulation. NOTE: Granix (tbo-filgrastim), Nivestym (filgrastim-aafi), and Ziextenzo (pegfilgrastim-bmez) are preferred products and DO NOT require prior authorization.	
J1426 J1413 J1428 J1427 J1429	DUCHENNE MUSCULAR DYSTROPHY AGENTS Amondys 45 (casimersen) Elevidys (delandistrogene moxeparvovec-rokl) Exondys 51 (eteplirsen) Viltepso (viltolarsen) Vyondys 53 (golodirsen)	Amondys 45 (casimersen) may be approved when ALL the following criteria are met: a. Member has a diagnosis of Duchenne Muscular Dystrophy (DMD) AND b. Member must have genetic testing confirming mutation of the DMD gene that is amenable to exon 45 skipping AND c. Medication is prescribed by or in consultation with a provider who specializes in treatment of DMD (such as a neurologist, cardiologist, pulmonologist, or physical medicine and rehabilitation provider) AND d. Provider attests that serum cystatin C, urine dipstick, and urine protein-to-creatinine ratio (UPCR) and glomerular filtration rate (GFR) will be measured prior to initiation of and that the member will be monitored periodically for kidney toxicity during treatment AND e. The member must be on corticosteroids at baseline or prescriber provides clinical rationale for not using corticosteroids at baseline or prescriber provides clinical rationale for not using corticosteroids AND f. If the member is ambulatory, functional level determination of baseline assessment of ambulatory function is required OR if not ambulatory, member must have a baseline Brooke Upper Extremity Function Scale or Forced Vital Capacity (FVC) documented AND g. Provider and patient or caregiver are aware that continued US FDA approval of Amondys 45 (casimersen) for Duchenne muscular dystrophy (DMD) may be contingent upon verification and description of clinical benefit in a confirmatory trial.	Elevidys: one time approval Other DMD therapies: One Year
		Reauthorization: After one year of treatment with Amondys 45 (casimersen), the member may receive approval to continue therapy for one year if the following criteria are met:	

	DO MEDICAID FROGRAM	AFFENDICES	
HCPCS	Drug	Criteria	PAR Length
		 a. Member has shown no intolerable adverse effects related to Amondys 45 (casimersen) treatment at a dose of 30mg/kg IV once a week AND b. Member has normal renal function or stable renal function if known impairment AND c. Member demonstrates response to Amondys 45 (casimersen) treatment with clinical improvement in trajectory from baseline assessment in ambulatory function OR if not ambulatory, member demonstrates improvement from baseline on the Brooke Upper Extremity Function Scale or in Forced Vital Capacity (FVC). 	
		Maximum Dose: 30 mg/kg per week	
		Above coverage standards will continue to be reviewed and evaluated for any applicable changes due to the evolving nature of factors including disease course, available treatment options, and available peer-reviewed medical literature and clinical evidence.	
		Elevidys (delandistrogene moxeparvovec-rokl) may be approved if the following criteria are met: a. Member is aged 4 through 5 years AND b. Member has a diagnosis of Duchenne Muscular Dystrophy (DMD) with a confirmed mutation in the DMD gene AND c. Member is ambulatory and provider has performed and documented a functional level determination of baseline assessment of ambulatory function AND d. Provider attests that member does not have any of these conditions: i. elevated anti-AAVrh74 total binding antibody titers (≥1:400) based on ELISA testing ii. any deletion in exon 8 and/or exon 9 in the DMD gene iii. pre-existing liver impairment iv. recent or active infections e. Medication is prescribed by or in consultation with a provider who specializes in treatment of DMD (such as a neurologist, cardiologist, pulmonologist, or physical medicine and rehabilitation provider) AND f. Provider attests that baseline liver function (clinical exam, GGT, total bilirubin), platelet count, and troponin-I will be assessed prior to Elevidys infusion and also	
		monitored following the infusion according to product labeling AND g. The member must be on corticosteroids at baseline or prescriber provides clinical rationale for not using corticosteroids AND	

HCPCS	Drug	Criteria	PAR Length
		 h. Provider attests that member has received all age-appropriate vaccinations as recommended by current immunization guidelines at least 4 weeks prior to initiation of the corticosteroid regimen AND i. Provider and patient or caregiver are aware that continued US FDA approval of Elevidys (delandistrogene moxeparvovec) for Duchenne muscular dystrophy (DMD) may be contingent upon verification and description of clinical benefit in confirmatory trial(s). j. Above coverage standards will continue to be reviewed and evaluated for any applicable changes due to the evolving nature of factors including disease course, available treatment options, and available peer-reviewed medical literature and clinical evidence. 	
		Maximum dose: one kit containing 70 single-dose 10 mL vials Approval will be placed to allow for one treatment course	
		 Exondys 51 (eteplirsen) may be approved if the following criteria are met: a. Member must have genetic testing confirming mutation of the Duchenne Muscular Dystrophy (DMD) gene that is amenable to exon 51 skipping AND b. Medication is prescribed by or in consultation with a provider who specializes in treatment of DMD (such as a neurologist, cardiologist, pulmonologist, or physical medicine and rehabilitation provider) AND c. The member must be on corticosteroids at baseline or has a contraindication to corticosteroids AND d. If the member is ambulatory, functional level determination of baseline assessment of ambulatory function is required OR if not ambulatory, member must have a Brooke Upper Extremity Function Scale of five or less documented OR a Forced Vital Capacity (FVC) of 30% or more. Reauthorization may be approved if provider attests that treatment with Exondys 51 (eteplirsen) is necessary to help member improve or maintain functional capacity based on assessment of trajectory\ from baseline for ambulatory or upper extremity function or Forced Vital Capacity (FVC). 	
		Maximum Dose: 30 mg/kg per week (documentation of patient's current weight with the date the weight was obtained)	

HCPCS	Drug	Criteria	PAR Length
HCPCS	Drug	Exemption: Members currently stabilized on a Exondys 51 (eteplirsen) regimen administered in a physician's office or clinical setting that was initiated prior to 1/1/2023 may receive prior authorization approval for continuation of therapy without meeting the above criteria. Viltepso (viltolarsen) may be approved for members meeting the following criteria: a. Member must have genetic testing confirming mutation of the Duchenne muscular dystrophy (DMD) gene that is amenable to exon 53 skipping AND b. Medication is prescribed by or in consultation with a provider who specializes in treatment of DMD (such as a neurologist, cardiologist, pulmonologist, or physical medicine and rehabilitation provider) AND c. Serum cystatin C, urine dipstick, and urine protein-to-creatinine ratio should be measured before starting Viltepso (viltolarsen). Consider measurement of glomerular filtration rate prior to initiation of Viltepso (viltolarsen) AND d. Members with known renal function impairment should be closely monitored during treatment with Viltepso (viltolarsen), as renal toxicity has occurred with similar drugs AND e. If the member is ambulatory, functional level determination of baseline assessment of ambulatory function is required OR if not ambulatory, member must have a baseline Brooke Upper Extremity Function Scale score or Forced Vital Capacity (FVC) documented AND f. Provider and patient or caregiver are aware that continued US FDA approval of	PAR Length
		of ambulatory function is required OR if not ambulatory, member must have a baseline Brooke Upper Extremity Function Scale score or Forced Vital Capacity (FVC) documented AND	
		Reauthorization: After one year of treatment with Viltepso (viltolarsen), member may receive approval to continue therapy for one year if the following criteria are met: a. Member has shown no intolerable adverse effects related to Viltepso (viltolarsen) treatment at a dose of 80mg/kg IV once a week AND b. Member has normal renal function or stable renal function if known impairment AND c. Provider attests that treatment with Viltepso (viltolarsen) is necessary to help member improve or maintain functional capacity based on assessment of trajectory from baseline for ambulatory or upper extremity function or Forced Vital Capacity (FVC).	

HCPCS	Drug	Criteria	PAR Length
		Maximum dose: 80 mg/kg administered as an IV infusion once weekly (documentation of patient's current weight with the date the weight was obtained) Exemption: Members currently stabilized on a Viltepso (viltolarsen) regimen administered in a physician's office or clinical setting that was initiated prior to 1/1/2023 may receive prior	
		 authorization approval for continuation of therapy without meeting the above criteria. Vyondys 53 (golodirsen) may be approved if all the following criteria are met: a. Member must have genetic testing confirming mutation of the Duchenne Muscular Dystrophy (DMD) gene that is amenable to exon 53 skipping AND b. Medication is prescribed by or in consultation with a provider who specializes in treatment of DMD (such as a neurologist, cardiologist, pulmonologist, or physical medicine and rehabilitation provider) AND c. The member must be on corticosteroids at baseline or has a contraindication to corticosteroids AND d. If the member is ambulatory, functional level determination of baseline assessment of ambulatory function is required OR if not ambulatory, member must have a Brooke Upper Extremity Function Scale of five or less documented OR a Forced Vital Capacity of 30% or more. 	
		Reauthorization may be approved if provider attests that treatment with Vyondys 53 (golodirsen) is necessary to help member improve or maintain functional capacity based on assessment of trajectory from baseline for ambulatory or upper extremity function or Forced Vital Capacity (FVC). Maximum Dose: 30 mg/kg per week (documentation of patient's current weight with the date	
		Exemption: Members currently stabilized on a Vyondys 53 (golodirsen) regimen administered in a physician's office or clinical setting that was initiated prior to 1/1/2023 may receive prior authorization approval for continuation of therapy without meeting the above criteria.	
		*All above coverage standards for all above medications will continue to be reviewed and evaluated for any applicable changes due to the evolving nature of factors including disease	

HCPCS	Drug	Criteria	PAR Length
		course, available treatment options, and available peer-reviewed medical literature and clinical evidence.	
J2508	Elfabrio (pegunigalsidase alfa)	 Elfabrio (pegunigalsidase alfa) may be approved if the following criteria are met: a. Member is ≥ 18 years of age AND b. Member has a confirmed diagnosis of Fabry disease AND c. The medication is being prescribed by or in consultation with a neurologist or metabolic disease provider AND d. Member has an eGFR ≥ 30 mL/min AND e. Member has been counseled regarding use of highly effective contraceptive method(s) while receiving treatment Maximum dose: 1 mg/kg every two weeks, based on actual body weight 	One year
J3380	Entyvio (vedolizumab)	Entyvio (vedolizumab) may be approved for members meeting all the following criteria: a. Member is 18 years of age or older AND b. Member has a diagnosis of moderately-to-severely active ulcerative colitis or moderately-to-severely active Crohn's disease AND c. Member is not receiving medication in combination with Cimzia, Enbrel, Humira, infliximab, Simponi, or Tysabri AND d. For members with Crohn's disease i. Medication is initiated and titrated per FDA-labeled dosing for Crohn's Disease ii. Member has trial and failure‡ of one preferred adalimumab product e. For members with Ulcerative Colitis i. Medication is initiated and titrated per FDA-labeled dosing for Ulcerative Colitis ii. Member has trial and failure‡ of one preferred adalimumab product and XELJANZ IR †Failure is defined as lack of efficacy, allergy, intolerable side effects, contraindication to therapy, or significant drug-drug interaction. Maximum dose: 300mg IV infusion at 0, 2, and 6 weeks and then every 8 weeks	One year

HCPCS	Drug	Criteria	PAR Length
J1305	Evkeeza (evinacumab)	Evkeeza (evinacumab) may be approved if the following criteria are met: a. Member is ≥ 5 years of age AND b. Member has a diagnosis of homozygous familial hypercholesterolemia (HoFH) AND c. The requested drug is being prescribed by, or in consultation with a cardiologist, Certified Lipid Specialist (CLS) or an endocrinologist AND d. Member has failed to achieve desired LDL-C with three months of maximally tolerated therapy with one high-potency statin (atorvastatin or rosuvastatin) in combination with ezetimibe. Failure is defined as lack of efficacy (member with ASCVD and LDL-C >55 mg/dL or member with HoFH and LDL-C >100 mg/dL), allergy, intolerable side effects, contraindication, or significant drug-drug interaction. For members with past or current incidence of rhabdomyolysis, trial and failure of statin therapy is not required AND e. Member has trialed and failed therapy with a PCSK9 inhibitor (alirocumab or evolocumab). Failure is defined as lack of efficacy after a 3-month trial, allergy, intolerable side effects, contraindication, or significant drug-drug interaction AND f. Member is not pregnant and members of reproductive potential have been counseled regarding use of effective contraception during and for 5 months following treatment. Note: The safety and effectiveness of Evkeeza (evinacumab) have not been established in patients with other causes of hypercholesterolemia, including those with heterozygous familial hypercholesterolemia (HeFH). Initial approval: 1 year Reauthorization: Reauthorization may be approved for 1 year with provider attestation confirming efficacy in lowering LDL-C.	See criteria
J0178	Eylea (aflibercept)	Eylea (aflibercept) may be approved for members meeting all the following criteria: a. Member is 18 years of age or older AND b. Member has a definitive diagnosis of one of the following and dosing is appropriate for the specified diagnosis as follows: i. Neovascular (Wet) Age-Related Macular Degeneration	One year

HCPCS	Drug	Criteria	PAR Length
		1. Maximum dose of 2 mg (0.05 mL) administered every 4 weeks for the first 12 weeks, followed by 2 mg (0.05 mL) every 8 weeks thereafter ii. Diabetic macular edema 1. Maximum dose of 2 mg (0.05 mL) administered every 8 weeks iii. Macular edema following retinal vein occlusion 1. Maximum dose of 2 mg (0.05 mL) administered every 4 weeks iv. Diabetic retinopathy 1. Maximum dose of 2 mg (0.05 mL) administered every 8 weeks c. AND d. Medication is prescribed by or in consultation with an ophthalmologist AND e. Medication is not being used in combination with any other anti-vascular endothelial growth factor (VEGF) medication AND f. Member does not have any of the following: i. Ocular or periocular infection ii. Active intraocular inflammation iii. Hypersensitivity to requested medication Reauthorization may be approved if member met initial approval criteria at the time of initiation of therapy AND the provider attests that the member has shown clinical improvement defined as an improvement or stabilization in visual acuity	
J0517	Fasenra (benralizumab)	Fasenra (benralizumab) may be approved for members meeting all the following criteria: a. If being administered for the diagnosis of asthma: i. Member is ≥ 6 years of age AND ii. Member has diagnosis of severe asthma with eosinophilic phenotype based on a blood eosinophil level of ≥ 150/mcL AND iii. Member's severe asthma has been refractory to recommended evidence-based, guideline-supported pharmacologic therapies AND iv. The requested medication is being prescribed as add-on therapy to existing asthma regimen AND v. The requested medication will not be used concomitantly with other biologic products indicated for asthma b. If being administered for the diagnosis of eosinophilic granulomatosis with polyangiitis (EGPA), medication may be approved for patients ≥18 years old.	One year

HCPCS	Drug	Criteria	PAR Length
		Reauthorization may be approved if member meets one of the following:	
		a. Improvement in lung function, measured in FEV1 OR	
		b. Reduction in the number of asthma exacerbations, defined as a decrease in use of	
		oral or systemic corticosteroids and/or reduced asthma related hospitalizations	
		and/or ER visits	
		Maximum dose: 30mg subcutaneous injection every 4 weeks for 3 doses, then every 8 weeks	
		thereafter	
	IMMUNE GLOBULINS	May be approved for members meeting one of the approved conditions listed and for doses not	One year
J1459,	Privigen,	exceeding FDA-approved maximum (Table 1).	
J1552	Alyglo	a. Approved Conditions for Immune Globulin Use:	
J1554,	Bivigam,	i. Primary Humoral Immunodeficiency disorders including:	
J1556,	Gammaplex,	1. Common Variable Immunodeficiency (CVID)	
J1557,	Gammaked, Gamunex-C,	2. Severe Combined Immunodeficiency (SCID)	
J1561,	Gamunex,	3. X-Linked Agammaglobulinemia	
	Gammagard S/D,	4. X-Linked with Hyperimmunoglobulin M (IgM)	
	Octagam 5%, 10%,	Immunodeficiency	
J1566,	Gammagard Liquid,	5. Wiskott-Aldrich Syndrome	
J1568,	Flebogamma DIF,	6. Members < 13 years of age with pediatric Human	
J1569,	Panzyga	Immunodeficiency Virus (HIV) and CD-4 count > 200/mm3	
J1572,	Asceniv	ii. Neurological disorders including:	
J1576,		1. Guillain-Barré Syndrome	
J1599		2. Relapsing-Remitting Multiple Sclerosis	
		3. Chronic Inflammatory Demyelinating Polyneuropathy	
		4. Myasthenia Gravis	
		5. Polymyositis and Dermatomyositis	
		6. Multifocal Motor Neuropathy	
		iii. Kawasaki Syndrome	
		iv. Chronic Lymphocytic Leukemia (CLL)	
		v. Autoimmune Neutropenia (AN) with absolute neutrophil count < 800 mm	
		and history of recurrent bacterial infections	
		vi. Autoimmune Hemolytic Anemia (AHA)	
		vii. Liver or Intestinal Transplant	
		viii. Immune Thrombocytopenia Purpura (ITP) including:	

HCPCS	Drug	Criteria	PAR Length
		1. Requiring preoperative therapy for undergoing elective splenectomy with platelet count < 20,000 2. Members with active bleeding & platelet count <30,000 3. Pregnant members with platelet counts <10,000 in the third trimester 4. Pregnant members with platelet count 10,000 to 30,000 who are bleeding ix. Multisystem Inflammatory Syndrome in Children (MIS-C)	
		Table 1: FDA-Approved Maximum Immune Globulin Dosing	
		Alyglo 800 mg/kg every 3 weeks	
		Gammaked 2 g/kg	
		Gamunex-C 2 g/kg Octagam 2 g/kg	
		Octagam 2 g/kg Gammagard Liquid 2.4 g/kg/month	
		Gammaplex 5% - IV Infusion 2 g/kg	
		Privigen - IV Infusion 2 g/kg	
		Asceniv 800 mg/kg every 3 weeks	
		Panzyga 2 g/kg	
		Bivigam 800 mg/kg every 3 weeks	
		Flebogamma DIF 600 mg/kg every 3 weeks	
		Gammagard S/D 1 g/kg	
J0175	Kisunla (donanemab-azbt)	Kisunla (donanemab-azbt) may be approved if the member meets ALL the following criteria: a. For claims billed through the pharmacy benefit, prescriber verifies that the medication is being administered by a healthcare professional in the member's home or in a long-term care facility AND b. Member has documented diagnosis of mild cognitive impairment or mild dementia stage of Alzheimer's disease, the population in which treatment was initiated in clinical trials, as evidenced by ALL the following:	See criteria
		Positron Emission Tomography (PET) scan OR lumbar puncture positive for amyloid beta plaque	

HCPCS	Drug	Criteria	PAR Length
HCPCS	Drug	Criteria 2. Mini-Mental State Examination (MMSE) score of 20-28 OR Montreal Cognitive Assessment (MoCA) Test score of 19-25 3. Progressive change in memory function for at least 6 months AND c. Member is 60 years of age or older AND d. Prior to initiation of medication, the prescriber attests that the member meets ALL the following: 1. Member has had a baseline brain MRI within the prior one year to treatment initiation, showing no signs or history of microhemorrhages and/or superficial siderosis 2. Attestation that MRI will be completed prior to the 2nd, 3rd, 4th, and 7th infusions 3. Member is negative for apolipoprotein Ε ε4 (ApoE ε4) homozygotes AND e. Member does not have any of the following: i. Any medical or neurological condition other than Alzheimer's Disease that might be a contributing cause of the subject's cognitive impairment including (but not limited to) stroke/vascular dementia, tumor, dementia with Lewy bodies [DLB], frontotemporal dementia [FTD] or normal pressure hydrocephalus ii. Contraindications to PET, CT scan, or MRI iii. History of or increased risk of amyloid related imaging abnormalities ARIA-edema (ARIA-E) or ARIA-hemosiderin deposition (ARIA-H) iv. History of unstable angina, myocardial infarction, chronic heart failure, or clinically significant conduction abnormalities, stroke, transient ischemic attack (TIA), or unexplained loss of consciousness within 1 year prior to initiation of Kisunla (donanemab-azbt) v. History of bleeding abnormalities or taking any form of anticoagulation therapy f. Medication is prescribed by or in consultation with a neurologist	PAR Length
		Second prior authorization approvals: an additional 6 months of therapy may be approved with provider attestation that a follow-up MRI will be (or has been) completed prior to the 7th infusion	

HCPCS	Drug	Criteria	PAR Length
		Subsequent prior authorization approvals: may be approved if provider attests that the member has demonstrated a positive clinical response to treatment Maximum dose: 700 mg every 4 weeks for the first 3 doses, followed by 1,400 mg every 4 weeks	
J0174	Leqembi (lecanemab-rimb)	Leqembi (leanemab-irmb) may be approved if the member meets ALL the following criteria: a. Member has documented diagnosis of mild cognitive impairment or mild dementia stage of Alzheimer's disease as evidenced by ALL of the following: i. Positron Emission Tomography (PET) scan OR lumbar puncture positive for amyloid beta plaque ii. Clinical Dementia Rating global score (CDR-GS) of 0.5 or 1 (available at https://otm.wustl.edu/cdr-terms-agreement/) iii. Mini-Mental State Examination (MMSE) score of 24-30 OR Montreal Cognitive Assessment (moCA) Test score of 19-25 AND b. Member is ≥ 50 years of age AND c. The prescriber attests that member has been counseled on the approval and safety status of Leqembi (lecanemab-irmb) being approved under accelerated approval based on reduction in amyloid beta plaques AND d. Prior to initiation of Leqembi (lecanemab-irmb), the prescriber attests that the member meets ALL of the following: i. Member has had a brain MRI within the prior one year to treatment initiation, showing no signs or history of localized superficial siderosis, ≥ 10 brain microhemorrhages, and/or brain hemorrhage > 1 cm ii. Attestation that MRI will be completed prior to the 5th, 7th and 14th infusions AND e. Member does not have any of the following: i. Any medical or neurological condition other than Alzheimer's Disease that might be a contributing cause of the subject's cognitive impairment including (but not limited to) stroke/vascular dementia, tumor, dementia with Lewy bodies [DLB], frontotemporal dementia [FTD] or normal pressure hydrocephalus ii. Contraindications to PET, CT scan, or MRI	See criteria

HCPCS	Drug	Criteria	PAR Length
		iii. History of or increased risk of amyloid related imaging abnormalities ARIA-edema (ARIA-E) or ARIA-hemosiderin deposition (ARIA-H) iv. History of unstable angina, myocardial infarction, chronic heart failure, or clinically significant conduction abnormalities, stroke, transient ischemic attack (TIA), or unexplained loss of consciousness within 1 year prior to initiation of Leqembi (lecanemab-irmb) v. History of bleeding abnormalities or taking any form of anticoagulation therapy AND f. The medication is prescribed by or in consultation with a neurologist Initial approval period: 6 months Subsequent approval: an additional 6 months of Leqembi (lecanemab-irmb) therapy may be approved with provider attestation that a follow-up MRI will be (or has been) completed prior to the 14th infusion Maximum dose: 10 mg/kg IV every 2 weeks	
J2778	Lucentis (ranibizumab)	Lucentis (ranibizumab) may be approved if all the following criteria are met: a. Member is 18 years of age or older AND b. Member has a diagnosis of one of the following: i. Neovascular (wet) Age-related Macular Degeneration (nAMD) ii. Diabetic Macular Edema (DME) iii. Macular Edema following Retinal Vein Occlusion (RVO) iv. Diabetic Retinopathy (DR) v. Myopic Choroidal Neovascularization (mCNV) AND c. Member does not have any of the following: i. Ocular or periocular infection ii. Hypersensitivity to the requested medication AND d. Documented baseline visual status with notation of eye(s) being treated AND e. Requested medication will not be used with other ophthalmic vascular endothelial growth factor (VEGF) inhibitors AND f. Documentation of trial and failure* with bevacizumab containing product AND g. Documentation of the dosing regimen that is being requested	See criteria

HCPCS	Drug	Criteria	PAR Length
		Initial approval: 6 months	
		Reauthorization: may be approved for one year if the following criteria are met: a. Documentation of improvement or stabilization of disease state and visual status AND b. Documentation of ongoing treatment regimen AND c. Prescriber attests that the member has had no significant adverse effects or drug toxicity, such as endophthalmitis and retinal detachments, increase in intraocular pressure, arterial thromboembolic events (ATE), retinal vasculitis and/or retinal vascular occlusion AND d. Prescriber attests requested medication will not be used with other ophthalmic VEGF inhibitors AND e. For Myoptic Choroidal Neovascularization (mCNV) continued administration is necessary due to disease activity (such as visual symptoms or presence of intra-/sub-retinal fluid or active leakage) Maximum dose nAMD/RVO: 0.5 mg every 4 weeks DME/DR: 0.3 mg every 4 weeks DME/DR: 0.3 mg every 4 weeks mCNV: 0.5 mg every 4 weeks for up to 3 months *Failure is defined as lack of efficacy with a three-month trial, contraindication to therapy, allergy, intolerable side effects, or significant drug-drug interaction	
J0490 J0491	Lupus Agents Benlysta (belimumab) Saphnelo (anifrolumab)	 Benlysta (belimumab) may be approved if the following criteria are met: a. For claims billed through the pharmacy benefit, prescriber verifies that the medication is being administered by a healthcare professional in the member's home or in a long term care facility AND b. Member is age ≥ 5 years and has active, autoantibody-positive systemic lupus erythematosus (SLE) and receiving standard therapy OR has active lupus nephritis and is receiving standard therapy AND c. Member has incomplete response to standard therapy from at least two of the following therapeutic classes: antimalarials, immunosuppressants and glucocorticoids; AND d. Member maintains standard therapy while on medication AND 	One year

HCPCS	Drug	Criteria	PAR Length
		e. Member is not receiving other biologics or intravenous cyclophosphamide AND f. The product is NOT being prescribed for severe active lupus nephritis or severe active central nervous system lupus Maximum dose: 10 mg/kg at 2-week intervals for the first 3 doses and 4-week intervals thereafter	
		 Saphnelo (anifrolumab) may be approved if member meets the following criteria: a. Member is ≥ 18 years of age with active, autoantibody-positive, moderate to severe systemic lupus erythematosus (SLE) AND is currently receiving standard therapy AND b. The product is NOT being prescribed for severe active lupus nephritis or severe active central nervous system lupus AND c. Member has had incomplete response to standard therapy from at least two of the following therapeutic classes: antimalarials, immunosuppressants and glucocorticoids AND d. Member will maintain standard therapy for SLE while receiving requested medication therapy e. Prescriber acknowledges that there are limited human data available for the use of anifrolumab in pregnancy and data are insufficient to inform on drug-associated risks. A registry monitors pregnancy outcome in women exposed to anifrolumab during pregnancy. Maximum Dose: 300 mg IV every 4 weeks 	
J2329 J0202 J2350 J2323	Multiple Sclerosis Agents Briumvi (ublituximab) Lemtrada (alemtuzumab) Ocrevus (ocrelizumab) Tysabri (natalizumab)	 Briumvi (ublituximab) may be approved if the following criteria are met: a. Member is ≥ 18 years of age AND b. Member has a relapsing form of multiple sclerosis (MS) AND c. Member has experienced at least one relapse in the prior year or two relapses in the prior two years AND d. Member has had trial and failure of any two high efficacy disease modifying therapies (such as ofatumumab, fingolimod, rituximab, ocrelizumab, alemtuzumab). Failure is defined as allergy, intolerable side effects, significant drug-drug interaction, or lack of efficacy. Lack of efficacy is defined as one of the following: 	One Year

HCPCS Drug	Criteria	PAR Length
HCPCS Drug	i. On MRI, presence of any new spinal lesions, cerebellar or brainstem lesions, or change in brain atrophy OR ii. Signs and symptoms on clinical exam consistent with functional limitations that last one month or longer AND e. Member does not have active hepatitis B virus (HBV) infection AND f. Briumvi (ublituximab) is prescribed by or in consultation with a neurologist or a physician that specializes in the treatment of multiple sclerosis AND g. Member does not have low serum immunoglobulins, based on quantitative tests performed before initiating treatment, AND h. Prescriber attests that appropriate premedication (such as a corticosteroid and antihistamine) will be administered prior to each Briumvi (ublituximab) infusion AND i. For members of childbearing potential: i. Member is not pregnant and prescriber acknowledges that pregnancy testing is recommended for members of reproductive potential prior to each infusion AND ii. Member has been counseled regarding the use of highly effective contraceptive methods while receiving treatment with Briumvi and for at least 6 months after stopping Briumvi Quantity limit: Four 150 mg/6 mL single-dose vials for the first 2 weeks (initial dose), and three 150 mg/6 mL single-dose vials every 24 weeks thereafter Exemption: If member is currently receiving and stabilized on ublituximab, they may receive prior authorization approval to continue therapy. Lemtrada (alemtuzumab) may be approved if member meets the following criteria: a. Member is 18 years of age or older AND b. Member has a relapsing form of multiple sclerosis AND c. Member has experienced one relapse within the prior year or two relapses within the prior two years AND d. Member has trial and failure* of Tysabri (natalizumab), Ocrevus (ocrelizumab), or two preferred agents in the "Disease Modifying Therapies" PDL drug class that are	PAR Length

HCPCS	Drug	Criteria	PAR Length
		 e. Medication is administered by or in consultation with a neurologist or a physician that specializes in the treatment of multiple sclerosis AND f. For members with known psychiatric conditions, peer-to-peer consultation with member's behavioral health provider will be conducted prior to the member's receiving treatment with a high dose corticosteroid as part of the medication's premedication procedure AND g. Baseline skin exam and thyroid function assessment are completed and documented prior to initiation of treatment with the medication AND h. Prescriber is enrolled in the Lemtrada Risk Evaluation and Mitigation Strategy (REMS) program i. Exemption: If member is currently receiving and stabilized on Lemtrada (alemtuzumab), they may continue to receive prior authorization approval to continue. 	
		Ocrevus (ocrelizumab) may be approved for initial therapy if member meets the following criteria: a. Medication is administered by or in consultation with a neurologist or a physician that specializes in the treatment of multiple sclerosis AND b. If administered for Relapsing Forms of Multiple Sclerosis (MS) i. Member is 18 years of age or older AND ii. Member does not have active hepatitis B infection or hypogammaglobulinemia at baseline AND iii. Member has a relapsing form of multiple sclerosis AND iv. Member has experienced one relapse within the prior year or two relapses within the prior two years AND v. Request meets one of the following: 1. Member has had a trial and failure* of any high-efficacy diseasemodifying therapies OR trial and failure* of any preferred product in the PDL "Multiple Sclerosis Agents" drug class OR 2. Member with highly active relapsing MS (based on measures of relapsing activity and MRI markers of disease activity such as numbers of galolinium-enhanced lesions). OR c. If administered for Primary Progressive Multiple Sclerosis	
		i. Member is 18 years of age or older AND	

HCPCS	Drug	Criteria	PAR Length
		ii. Member is not concomitantly taking disease modifying therapies.	
		Maximum maintenance dose: 600 mg every 6 months	
		Exemption: If member is currently receiving and stabilized on Ocrevus, they may continue to	
		receive prior authorization approval to continue	
		Tysabri (natalizumab) may be approved for initial therapy if the following criteria are met:	
		a. Medication is not currently being used in combination with immunosuppressants	
		(azathioprine, 6-mercaptopurine, methotrexate) or TNF-alpha inhibitors	
		(adalimumab, certolizumab pegol, infliximab) AND	
		b. Member does not have anti-JC virus antibodies at baseline AND	
		c. <u>If administered for induction of remission of moderate to severe Crohn's disease</u>	
		i. The member is ≥ 18 years of age AND	
		ii. Prescriber and member are enrolled in the CD TOUCH® REMS program	
		AND	
		iii. Member has tried and failed aminosalicylates AND	
		iv. Member has tried and failed corticosteroids AND v. Member has tried and failed immunomodulators AND	
		vi. Member has tried and failed two TNF-alpha inhibitors (e.g. adalimumab,	
		certolizumab pegol, infliximab) AND	
		vii. Medication is administered by or in consultation with a gastroenterologist.	
		d. If administered for relapsing remitting multiple sclerosis (RRMS)	
		i. The member is ≥ 18 years of age AND	
		ii. Prescriber and member are enrolled in the MS TOUCH® REMS program	
		AND	
		iii. Medication is administered by or in consultation with a neurologist or a	
		physician that specializes in the treatment of multiple sclerosis	
		iv. Request meets one of the following:	
		1. Member has trial and failure* of any two high efficacy disease	
		modifying therapies (such as ofatumumab, fingolimod, rituximab, ocrelizumab, alemtuzumab)	
		OR	
		2. Member with highly active relapsing MS (based on measures of	
		relapsing activity and MRI markers of disease activity such as	
		numbers of galolinium-enhanced lesions) has had a trial and	

HCPCS	Drug	Criteria	PAR Length
		failure* of any high-efficacy disease-modifying therapy (such as ofatumumab, fingolimod, rituximab, alemtuzumab)	
		Exemption: If member is currently receiving and stabilized on Tysabri, they may continue to receive prior authorization approval to continue.	
		*Failure is defined as intolerable side effects, drug-drug interaction, contraindication, or lack of efficacy. Lack of efficacy is defined as one of the following:	
		1. One of the following on MRI: presence of any new spinal lesions, cerebellar or brainstem lesions, or change in brain atrophy OR	
		2. On clinical exam, signs and symptoms consistent with functional limitations that last one month or longer	
J2796	Nplate (romiplostim)	Nplate (romiplostim) may be approved if the member meets the following criteria: a. Member does not have thrombocytopenia due to myelodysplastic syndrome (MDS) or any cause of thrombocytopenia other than immune thrombocytopenia AND b. Medication is not being used in an attempt to normalize platelet counts AND c. If being administered for hematopoietic subsyndrome of acute radiation syndrome, member has been acutely exposed to myelosuppressive radiation levels greater than 2 gray (Gy) OR d. If being administered for immune thrombocytopenia (ITP) i. Member has had an insufficient response to corticosteroids, immunoglobulins, or splenectomy AND ii. Member has ITP whose degree of thrombocytopenia and clinical condition increases the risk for bleeding as indicated by a platelet count of ≤ 30,000/mm³ AND iii. Laboratory value for platelet count is current (e.g., drawn within the previous 28 days) AND iv. If being administered for Acute ITP 1. Member is at least 18 years of age or older OR If being administered for Chronic ITP 1. Member is at least 1 years of age or older AND 2. Member has had chronic ITP for at least 6 months	One year

HCPCS	Drug	Criteria	PAR Length
		Maximum dose: weekly dose of 10 mcg/kg $\frac{\text{Reauthorization}}{\text{Reauthorization}} \text{ may be approved for ITP if member met the initial indication-specific approval criteria above and member responded to treatment by achieving and maintaining a platelet count of \geq 50,000/\text{mm}^3, but <450,000/\text{mm}^3$	
J2182	Nucala (mepolizumab)	Nucala (mepolizumab) may be approved if member meets ALL the following criteria for the appropriate indication: a. Initial approval if administered for asthma:	One year

HCPCS	Drug	Criteria	PAR Length
HCPCS	Drug	ii. Member has been diagnosed with relapsing or refractory EGPA at least 6 months prior to request as demonstrated by ALL the following: 1. Member has a diagnosis of asthma AND 2. Member has a blood eosinophil count of greater than or equal to 1000 cells/mcL or a blood eosinophil level of 10% AND 3. Member has the presence of two of the following EGPA characteristics: Histopathological evidence of eosinophilic vasculitis, perivascular eosinophilic infiltration, or eosinophil-rich granulomatous inflammation Neuropathy Pulmonary infiltrates Sinonasal abnormality Cardiomyopathy Glomerulonephritis Alveolar hemorrhage Palpable purpura Antineutrophil cytoplasmic antibody (ANCA) positive 4. Member is on a stable dose of corticosteroids for at least 4 weeks prior to request AND 5. Dose of 300 mg once every 4 weeks iii. If administered for hypereosinophilic syndrome (HES): 1. Member is 12 years of age or older AND 2. Member has a diagnosis for HES for at least 6 months that is nonhematologic secondary HES AND 3. Member has a blood eosinophil count of greater than or equal to 1000 cells/mcL AND 4. Member has a history of two or more HES flares (defined as worsening clinical symptoms or blood eosinophil counts requiring an increase in therapy) AND 5. Member has been on stable dose of HES therapy for at least 4	PAR Length
		weeks, at time of request, including at least one of the following: Oral corticosteroids Immunosuppressive therapy	

HCPCS	Drug	Criteria	PAR Length
		☐ Cytotoxic therapy	
		AND	
		6. Dose of 300 mg once every 4 weeks	
J2267	Omvoh (mirikizumab-mrkz)	 Omvoh (mirikizumab-mrkz) may receive approval if the following criteria are met: a. The requested medication is being prescribed for treatment of moderately-to-severely active ulcerative colitis AND b. Member is ≥ 18 years of age AND c. Member has trial and failure‡ of one preferred adalimumab product AND XELJANZ IR AND ENTYVIO (vedolizumab) AND d. Prescriber acknowledges that administration of IV induction therapy prior to approval of OMVOH (mirikizumab-mrkz) pen for subcutaneous injection using the above criteria should be avoided and will not result in an automatic approval of requests for these formulations. 	One Year
J0129	Orencia (abatacept)	Orencia (abatacept) may be approved if meeting the following criteria: a. Member has a diagnosis of moderate to severe rheumatoid arthritis or polyarticular juvenile idiopathic arthritis (pJIA) AND has trialed and failed* all preferred agents in the "Targeted Immune Modulators" PDL drug class that are FDA-labeled for use for the prescribed indication (only one preferred adalimumab product trial required). OR b. Member is an adult with a diagnosis of psoriatic arthritis AND trial and failure of a preferred adalimumab product OR Enbrel and Xeljanz IR AND Taltz or Otezla OR c. The requested medication is being prescribed for the prophylaxis of acute graft versus host disease (aGVHD) in combination with a calcineurin inhibitor and methotrexate in patients undergoing hematopoietic stem cell transplantation (HSCT) from a matched or 1 allele-mismatched unrelated-donor.	One year
		 Exemption: Members currently stabilized on Orencia (abatacept) regimen that was initiated prior to 1/1/2023 may receive prior authorization approval for continuation of therapy without meeting the above criteria. *Failure is defined as lack of efficacy with a three-month trial, contraindication to therapy, allergy, intolerable side effects, or significant drug-drug interaction. Note that trial and failure of preferred TNF inhibitors will not be required when prescribed for pJIA in members with documented clinical features of lupus. 	

HCPCS	Drug			Criteria	PAR Length
J0224	Oxlumo (lumasiran)	Oxlumo (lumasiran) may be approved if all the following criteria are met: a. Member has a diagnosis of Primary hyperoxaluria type 1 (PH1) confirmed by either: i. Genetic testing that demonstrates a mutation of the alanine glyoxylate aminotransferase (AGXT) gene OR ii. Liver enzyme analysis demonstrating absent or significantly reduced AGXT b. Medication is being prescribed by, or in consultation with a nephrologist, neurologist, or other healthcare provider with expertise in treating PH1 c. Member has documented baseline urinary oxalate excretion or plasma oxalate concentrations Reauthorization: Member demonstrates response to medication as indicated by a positive clinical response from baseline urinary oxalate excretion or plasma oxalate concentration Maximum dose: weight-based dosing regimen as shown in the following table (documentation of patient's current weight with the date the weight was obtained)		One year	
		Body Weight	Loading Dose	Maintenance Dose	
		Less than 10 kg	6 mg/kg once monthly for three doses	3 mg/kg once monthly, beginning one month after the last loading dose	
		10 kg to less than 20 kg	6 mg/kg once monthly for three doses	6 mg/kg once every three months, beginning one month after the last loading dose	
		20 kg and above	3 mg/kg once monthly for three doses	3 mg/kg once every three months, beginning one month after the last loading dose	
		prior to 1/1/20		d on a Oxlumo (lumasiran) regimen that was initiated orization approval for continuation of therapy	
J1307	Piasky (crovalimab)	a. Mem b. Mem c. Mem	where is ≥ 13 years of age A where weighs at least 40 kg (88.2 pounds) AND oxysmal nocturnal hemoglobinuria (PNH) confirmed	See criteria

HCPCS	Drug	Criteria	PAR Length
		 d. Requested product is being prescribed by or in consultation with a hematologist, immunologist or nephrologist AND e. Member has a lactate dehydrogenase (LDH) level ≥ 2 times the upper limit of normal AND f. Member has had at least one PNH-related sign or symptom (such as hemoglobinuria, fatigue, dyspnea, abdominal pain, dysphagia) within the past 3 months AND g. Member has a hemoglobin level measured at baseline AND h. Member does not have any active infections caused by an encapsulated bacteria (such as <i>Streptococcus pneumoniae</i>, <i>Neisseria meningitidis</i>, and <i>Haemophilus influenzae</i> type b) AND i. Member has been vaccinated against <i>Neisseria meningitidis</i> (serogroups A, C, W, Y and B) within the 3 years prior to initiation of treatment with Piasky (crovalimab) OR will be vaccinated against <i>Neisseria meningitidis</i> within 7 days after starting treatment AND j. Member has been vaccinated against <i>Streptococcus pneumoniae</i> and <i>Haemophilus influenzae type b</i> (Hib) according to ACIP recommendations. If urgent Piasky (crovalimab) therapy is indicated in a patient who is not up to date with vaccines, or the vaccines were administered within the last 2 weeks, prescriber attests that the member will receive appropriate antibacterial drug prophylaxis and the vaccines will be administered as soon as possible AND k. Members of childbearing age have been counseled that fetal effects of Piasky (crovalimab) during pregnancy are unknown, and to avoid breastfeeding during treatment with Piasky (crovalimab) and for 9 months following the final dose AND l. Due to the risk of forming drug-target-drug complexes (DTDCs) and Type III hypersensitivity reactions, monitor patients switching from another C5 inhibitor to Piasky (crovalimab) or from Piasky (crovalimab) to another C5 inhibitor for at 30 	
		days as outlined in the full prescribing information. Maximum doses: 1,500 mg intravenous loading dose 340 mg subcutaneous loading doses (Days 2, 8, 15, 22) 1,020 mg subcutaneous maintenance doses (Day 29 and every 4 weeks thereafter) Quantity limits Initial IV loading dose: 5 single-dose 340 mg/2 mL vials Subcutaneous loading doses: 4 single-dose 340 mg/2 mL vials Subcutaneous maintenance doses: 3 single-dose 340 mg/2 mL vials every 28 days	

HCPCS	Drug	Criteria	PAR Length
		 Initial authorization: 6 months Reauthorization: Approval for 1 year may be given with prescriber attestation that member meets at least one of the following 6 months after initiation of treatment: a. Member has achieved BOTH of the following: i. Hemolysis control, defined as LDH ≤ 1.5× ULN during the first 6 months of treatment AND ii. Transfusion avoidance, defined as not receiving a transfusion of packed red blood cells during the first 6 months of treatment OR b. Member has been monitored for breakthrough hemolysis and meets BOTH of the following: i. Member has a documented initial reduction of LDH ≤ 1.5 × ULN while on treatment AND ii. Member has not experienced at least one new or worsening symptom or sign of intravascular hemolysis in the presence of elevated LDH ≥ 2 × ULN after the prior reduction of LDH ≤ 1.5 × ULN while on treatment OR c. Member has achieved hemoglobin stabilization, defined as avoidance of a ≥ 2 g/dL decrease in hemoglobin level from baseline in the absence of transfusion. 	
J0221 J0219	Pompe Disease Agents Lumizyme (alglucosidase alfa) Nexviazyme (avalglucosidase)	Lumizyme (alglucosidase alfa) may be approved if member meets the following criteria: a. Member has a definitive diagnosis of Pompe disease confirmed by one of the following: i. Deficiency of acid alpha-glucosidase (GAA) enzyme activity OR ii. Detection of biallelic pathogenic variants in the GAA by molecular genetic testing AND b. The Request meets one of the following based on indicated use: i. If being administered for infantile-onset Pompe disease 1. Member has documented baseline age appropriate assessments, including motor function tests, muscle weakness, respiratory function, cardiac involvement testing, percent predicted forced vital capacity (FVC), and 6-minute walk test (6MWT) OR	One year

HCPCS	Drug	Criteria	PAR Length
		ii. If being administered for <u>Late-onset Pompe disease</u>	
		1. Member has documented baseline age appropriate assessments,	
		including motor function tests, muscle weakness, respiratory	
		function, cardiac involvement testing, FVC and 6MWT	
		Reauthorization may be approved if member met initial approval criteria at the time of initiation of therapy AND meets the following: a. Member is being monitored for antibody formation and hypersensitivity AND b. Request meets the following based on indicated use: i. For infantile-onset disease: the member has shown clinical improvement defined as an improvement or stabilization in muscle weakness, motor function, respiratory function, cardiac involvement, percent predicted FVC, and/or 6MWT OR ii. For late-onset disease: the member has shown clinical improvement defined as an improvement or stabilization in percent predicted FVC and/or 6MWT	
		Maximum dose: 20 mg/kg administered every 2 weeks	
		Nexviazyme (avalglucosidase alfa-ngpt) may be approved if member meets the following criteria:	
		a. Member is 1 year of age or older AND	
		b. Member has a definitive diagnosis of Pompe disease confirmed by one of the following:	
		i. Deficiency of acid alpha-glucosidase (GAA) enzyme activity OR	
		ii. Detection of biallelic pathogenic variants in the GAA by molecular genetic testing AND	
		c. Member has a diagnosis of late-onset (non-infantile) Pompe disease AND	
		d. Medication is not being used in combination with other enzyme replacement therapies AND	
		e. Member has documented baseline age appropriate assessments, including motor	
		function tests, muscle weakness, respiratory function, cardiac involvement testing,	
		percent predicted FVC and 6MWT	
		f. Product is being prescribed by a provider specializing in the treatment of Pompe	
		disease AND	

HCPCS	Drug	Criteria	PAR Length
		g. Prescriber will consider administering antihistamines, antipyretics, and/or corticosteroids prior to Nexviazyme (avalglucosidase alpha) administration to reduce the risk of severe infusion-associated reactions.	
		Reauthorization may be approved if member met initial approval criteria at the time of initiation of therapy AND meets the following: a. Member has shown clinical improvement defined as an improvement or stabilization in percent predicted FVC and/or 6MWT AND b. Member is being monitored for antibody formation and hypersensitivity	
		Maximum weight dependent dosage: Members ≥30 kg, 20 mg/kg administered every 2 weeks Members ≤30 kg, 40 mg/kg administered every 2 weeks	
J1745	Remicade (infliximab)	Remicade (infliximab) may be approved with trial & failure of an infliximab biosimilar AND if meeting all the following criteria: a. Member has one of the following diagnoses: i. Crohn's disease and is 6 years or older ii. Ulcerative colitis and is 6 years or older iii. Rheumatoid arthritis and is 4 years or older iv. Psoriatic arthritis and is 18 years or older v. Ankylosing spondylitis and is 18 years or older vi. Juvenile idiopathic arthritis and is 4 years or older vii. Plaque psoriasis in adults viii. Hydradenitis suppurativa (HS) AND b. Member meets one of the following, based on prescribed indication: i. For continuation of infliximab therapy that was initiated in the hospital setting for treating severe ulcerative colitis, no additional medication trial is required OR ii. For treatment of moderate to severe hidradenitis suppurativa, no additional medication trial is required OR iii. For treatment of psoriatic arthritis member has trial and failure; of a preferred adalimumab product or Enbrel AND Xeljanz IR AND Taltz or Otezla. OR	One year

HCPCS	Drug	Criteria	PAR Length
		iv. For treatment of moderately-to-severely active Crohn's disease, member has trial and failure; of one preferred adalimumab product OR for treatment of moderately-to-severely active ulcerative colitis, member has trial and failure; of one preferred adalimumab product and XELJANZ IR. OR v. For all other prescribed indications, the member has trialed and failed†* all preferred agents in the Targeted Immune Modulators PDL drug class that are FDA labeled for use for the prescribed indication (with only one preferred adalimumab product trial required).	
		** Members \geq 50 years of age with an additional CV risk factor, will not need a trial and failure of Xeljanz IR.	
		*Renflexis does not require prior authorization on the medical benefit.	
J9312	Rituxan (rituximab)	Rituximab (Rituxan brand/generic and rituximab-containing products or biosimilars) may receive approval if the following criteria are met: a. The prescribed agent is one of the following preferred rituximab products: Ruxience (rituximab-pvvr), Riabni (rituximab-arrx), Truxima (rituximab-abbs) OR b. If the prescribed agent is brand Rituxan or a rituximab-containing product formulation that is not a preferred product, then the member has trialed and failed at least one favored rituximab product. Failure is defined as lack of efficacy or intolerable side effects with the favored rituximab product formulation. NOTE: Ruxience (rituximab-pvvr), Riabni (rituximab-arrx), Truxima (rituximab-abbs) are preferred products and DO NOT require prior authorization.	One year
J1412	Roctavian (valoctocogene roxaparvovec-rvox)	Roctavian (valoctocogene roxaparvovec-rvox) may be approved when ALL the following criteria are met: a. Member is 18 years of age or older AND b. Member has documented diagnosis of severe hemophilia A defined by both of the following: i. Factor VIII deficiency with factor VIII activity < 1 IU/dL AND	One time treatment

HCPCS	Drug	Criteria	PAR Length
	Diug	ii. Member has ≥ 10 bleeding events requiring factor replacement therapy per year AND c. Member has had a minimum of 150 exposure days per year to a factor VIII agent AND d. Member is currently using factor VIII prophylaxis therapy or emicizumab AND e. Member is adeno-associated virus serotype 5 negative as determined by an FDA approved test AND f. Member must have completed Bethesda assay results of < 0.6 Bethesda Units (BU) within the prior 12 months AND g. Prescribed by or in consultation with a hematologist AND h. Member has documented liver health assessments completed including:	TAK Edigii
J2802	Romiplostim	Romiplostim may be approved if the member meets the following criteria: e. Member does not have thrombocytopenia due to myelodysplastic syndrome (MDS) or any cause of thrombocytopenia other than immune thrombocytopenia AND f. 34Medication is not being used in an attempt to normalize platelet counts AND g. If being administered for hematopoietic subsyndrome of acute radiation syndrome , member has been acutely exposed to myelosuppressive radiation levels greater than 2 gray (Gy) OR h. If being administered for immune thrombocytopenia (ITP) i. Member has had an insufficient response to corticosteroids, immunoglobulins, or splenectomy AND	One year

	ii. Member has ITP whose degree of thrombocytopenia and clinical condition	
	increases the risk for bleeding as indicated by a platelet count of ≤ 30,000/mm³ AND iii. Laboratory value for platelet count is current (e.g., drawn within the previous 28 days) AND iv. If being administered for Acute ITP 1. Member is at least 18 years of age or older OR If being administered for Chronic ITP 3. Member is at least 1 years of age or older AND 4. Member has had chronic ITP for at least 6 months	
	Maximum dose: weekly dose of 10 mcg/kg Reauthorization may be approved for ITP if member met the initial indication-specific approval criteria above and member responded to treatment by achieving and maintaining a platelet count of ≥ 50,000/mm³, but <450,000/mm³	
stiggo (rozanolixizumab)	 Rystiggo (rozanolixizumab) may be approved if the following criteria are met: a. Member is ≥ 18 years of age AND b. Member has a diagnosis of generalized myasthenia gravis that falls within Myasthenia Gravis Foundation of America (MGFA) Class II to IVa disease, AND c. Member has a positive serologic test for anti-acetylcholine receptor (AChR) or anti-muscle-specific tyrosine kinase (MuSK) antibodies AND d. Requested product is being prescribed by or in consultation with a neurologist AND e. A baseline Quantitative Myasthenia Gravis (QMG) assessment has been documented, AND f. Patient has a MG-Activities of Daily Living (MG-ADL) total score of ≥3 (with at least 3 points from non-ocular symptoms), AND g. Patient has failed† treatment over at least 1 year with at least 2 immunosuppressive therapies (such as azathioprine, cyclosporine, tacrolimus, mycophenolate), or has failed at least 1 immunosuppressive therapy and required chronic therapeutic plasma exchange or intravenous immunoglobulin (IVIG) AND h. As a precaution, consider discontinuation or Rystiggo and use of alternative 	See criteria
	stiggo (rozanolixizumab)	iv. If being administered for Acute ITP 1. Member is at least 18 years of age or older OR If being administered for Chronic ITP 3. Member is at least 1 years of age or older AND 4. Member has had chronic ITP for at least 6 months Maximum dose: weekly dose of 10 mcg/kg Reauthorization may be approved for ITP if member met the initial indication-specific approval criteria above and member responded to treatment by achieving and maintaining a platelet count of ≥ 50,000/mm³, but <450,000/mm³ Rystiggo (rozanolixizumab) Rystiggo (rozanolixizumab) may be approved if the following criteria are met: a. Member is ≥ 18 years of age AND b. Member has a diagnosis of generalized myasthenia gravis that falls within Myasthenia Gravis Foundation of America (MGFA) Class II to IVa disease, AND c. Member has a positive serologic test for anti-acetylcholine receptor (AChR) or anti-muscle-specific tyrosine kinase (MuSK) antibodies AND d. Requested product is being prescribed by or in consultation with a neurologist AND e. A baseline Quantitative Myasthenia Gravis (QMG) assessment has been documented, AND f. Patient has a MG-Activities of Daily Living (MG-ADL) total score of ≥3 (with at least 3 points from non-ocular symptoms), AND g. Patient has failed† treatment over at least 1 year with at least 2 immunosuppressive therapies (such as azathioprine, cyclosporine, tacrolimus, mycophenolate), or has failed at least 1 immunosuppressive therapy and required chronic therapeutic plasma exchange or intravenous immunoglobulin (IVIG) AND

HCPCS	Drug	Criteria	PAR Length
		human Fc receptor (such as IVIG, other immunoglobulins, or other C5 complement inhibitors). † Failure is defined as lack of efficacy, allergy, intolerable side effects, or significant drug-drug interaction	
		Initial Approval: 6 months Reauthorization: Reauthorization for one year may be approved with prescriber attestation that member has experienced a positive clinical response to rozanolixizumab based on documented Quantitative Myasthenia Gravis (QMG) assessment AND/OR MG-Activities of Daily Living (MG-ADL) score Maximum dose: 840 mg (6 mL) by subcutaneous infusion every 6 weeks Quantity limit: three 280 mg/2 mL single-dose vials every 6 weeks Exemption: Members who are currently stabilized on the requested medication may receive approval to continue treatment on that medication	
J1602	Simponi (golimumab)	Simponi (golimumab) may receive approval if meeting the following: a. The request meets one of the following: i. Member has a diagnosis of moderate to severe rheumatoid arthritis, polyarticular juvenile idiopathic arthritis, or ankylosing spondylitis AND has trialed and failed‡ all preferred agents in the "Targeted Immune Modulators" PDL drug class that are FDA-labeled for use for the prescribed indication or have strong evidence supporting use for the prescribed indication from clinically recognized guideline compendia (only one preferred adalimumab product trial required). OR ii. Member is an adult with a diagnosis of psoriatic arthritis AND has trial and failure‡ of a preferred adalimumab product or Enbrel AND Xeljanz IR AND Taltz or Otezla. OR b. If the request is for use of the subcutaneous formulation for treating moderately to severely active ulcerative colitis, all the following criteria are met: i. Member is ≥ 18 years of age AND	One year

HCPCS	Drug	Criteria	PAR Length
		ii. Member has trial and failure‡ of one preferred adalimumab product and XELJANZ IR AND iii. Member has demonstrated corticosteroid dependence or has had an inadequate response to (or failed to tolerate) oral aminosalicylates, oral corticosteroids, azathioprine, or 6-mercaptopurine for inducing and maintaining clinical response, improving endoscopic appearance of the mucosa during induction, inducing clinical remission, or achieving and sustaining clinical remission in induction responders. Exemption: Members currently stabilized on a Simponi (golimumab) regimen that was initiated prior to 1/1/2023 may receive prior authorization approval for continuation of therapy without meeting the above criteria. ‡Failure is defined as lack of efficacy, allergy, intolerable side effects, contraindication to therapy, or significant drug-drug interaction. Note that trial and failure of Xeljanz IR will not be required when prescribed for ulcerative colitis for members ≥ 50 years of age that have an additional CV risk factor.	
J1300	Soliris (eculizumab)	Soliris (eculizumab) may be approved for members meeting all the following criteria: a. Member is diagnosed with either Paroxysmal Nocturnal Hemoglobinuria (PNH),	One year

HCPCS	Drug	Criteria	PAR Length
		c. Member demonstrate the presence of at least 2 different	
		glycosylphosphatidylinositol (GPI) protein deficiencies (e.g. CD55, CD59, etc.)	
		within at least 2 different cell lines (granulocytes, monocytes, erythrocytes) AND	
		d. Member has one of the following indications for therapy:	
		i. Presence of a thrombotic event	
		ii. Presence of organ damage secondary to chronic hemolysis	
		iii. Member is pregnant and potential benefit outweighs potential fetal risk	
		iv. Member is transfusion dependent	
		v. Member has high LDH activity (defined as ≥1.5 x ULN) with clinical	
		symptoms	
		AND	
		a. Member has documented baseline values for one or more of the following:	
		i. Serum lactate dehydrogenase (LDH)	
		ii. Hemoglobin level	
		iii. Packed RBC transfusion requirement	
		Atypical Hemolytic Uremic Syndrome	
		a. Member is 2 months or older AND	
		b. Thrombotic Thrombocytopenic Purpura (TTP) has been ruled out by evaluating	
		ADAMTS13 level (ADAMTS-13 activity level > 10%); AND	
		c. Shiga toxin E. coli related hemolytic uremic syndrome (STECHUS) has been ruled	
		out; AND	
		d. Other causes have been identified and are being treated appropriately such as	
		coexisting diseases or conditions (e.g. bone marrow transplantation, solid organ	
		transplantation, malignancy, autoimmune disorder, drug-induced, malignant	
		hypertension, HIV infection, etc.), Streptococcus pneumonia or Influenza A (H1N1)	
		infection, or cobalamin deficiency AND	
		e. Documented baseline values for one or more of the following:	
		i. Serum lactate dehydrogenase (LDH)	
		ii. Serum creatinine/eGFR	
		iii. Platelet count	
		iv. Plasma exchange/infusion requirement	
		Generalized Myasthenia Gravis	
		a. Member is 18 years or older AND	
		b. Member has Myasthenia Gravis Foundation of America (MGFA) Clinical	
		Classification of Class II to IV disease; AND	

HCPCS Drug Crit	teria PAR Length
c. Member has a positive serologic test f antibodies; AND d. Physician has assessed the baseline Qi AND e. Member has a MG-Activities of Daily f. Member has failed treatment over at let therapies (e.g. azathioprine, cyclospor 1 immunosuppressive therapy and req exchange (PE) or intravenous immune Neuromyelitis Optica Spectrum Disorder a. Member is 18 years or older AND b. Member has a past medical history of i. Optic neuritis ii. Acute myelitis iii. Acute myelitis iii. Area postrema syndrome; epi nausea and vomiting iv. Acute brainstem syndrome v. Symptomatic narcolepsy or a NMOSD-typical diencephali vi. Symptomatic cerebral syndre c. Member has a positive serologic test f (AQP4-IgG)/NMP-IgG antibodies; Al d. Diagnosis of multiple sclerosis or othe e. Member has an history of failure, contr therapy AND g. Member has at least one of the follow i. History of at least two relaps initiating medication ii. History of at least two relaps initiating medication ii. History of at least three relap relapse occurring within the 1 AND h. Member is not receiving medication ii i. Disease modifying therapies	tor anti-acetylcholine receptor (AchR) uantitative Myasthenia Gravis (QMG) score; y Living (MG-ADL) total score of ≥6; AND east 1 year with at least 2 immunosuppressive rine, mycophenolate, etc), or has failed at least quired chronic plasmapheresis or plasma oglobulin (IVIG) Tone of the following: isode of otherwise unexplained hiccups or acute diencephalic clinical syndrome with ic MRI lesions ome with NMOSD-typical brain lesions AND for anti-aquaporin-4 immunoglobulin G ND er diagnoses have been ruled out AND raindication, or intolerance to rituximab

HCPCS	Drug	Criteria	PAR Length
	J	ii. Anti-IL6 therapy Exemption: If a member is currently receiving and stabilized on Soliris, they may continue to receive prior authorization approval to continue if the member meets the appropriate diagnosis and age requirements Maximum dose: 900mg weekly for 4 weeks induction followed by 1200mg every 2 weeks maintenance dose	
J3357	Stelara (ustekinumab) subcutaneous injection	Stelara (ustekinumab) subcutaneous injection use may receive approval if meeting the following: a. If administered for Crohn's disease or Ulcerative Colitis i. The member has a diagnosis of moderate-to-severely active Crohn's disease or moderate-to-severely active ulcerative colitis AND ii. The member is ≥ 18 years of age AND iii. For treatment of moderately-to-severely active Crohn's disease, member has trial and failure‡ of one preferred adalimumab product and ENTYVIO (vedolizumab) OR For treatment of moderately-to-severely active ulcerative colitis, member has trial and failure‡ of one preferred adalimumab product and XELJANZ IR and ENTYVIO (vedolizumab) AND iv. Prescriber acknowledges that loading dose administration prior to approval of STELARA for maintenance therapy using the above criteria should be avoided and will not result in an automatic approval of STELARA for maintenance therapy AND v. Prior authorization approval may be given for an initial 16-week supply and authorization approval for continuation may be provided based on clinical response. b. If administered for psoriatic arthritis i. Member has trial and failure‡ of a preferred adalimumab product or ENBREL AND XELJANZ IR AND TALTZ or OTEZLA AND ii. Prior authorization approval may be given for an initial 16-week course and authorization approval for continuation may be provided based on clinical response c. If administered for plaque psoriasis	See criteria

HCPCS	Drug	Criteria	PAR Length
		 i. Member has trial and failure‡ of a preferred adalimumab product or ENBREL) AND two indicated second line agents (TALTZ, OTEZLA), AND ii. Prior authorization approval may be given for an initial 16-week course and authorization approval for continuation may be provided based on clinical response. 	
		*Members currently stabilized on a Stelara (ustekinumab) regimen that was initiated prior to 1/1/2023 may receive prior authorization approval for continuation of therapy without meeting the above criteria.	
		‡Failure is defined as lack of efficacy with a three-month trial, contraindication to therapy, allergy, intolerable side effects, or significant drug-drug interaction. Note that trial and failure of Xeljanz XR will not be required when prescribed for ulcerative colitis for members ≥ 50 years of age that have an additional CV risk factor.	
J3358	Stelara (ustekinumab) intravenous (IV) injection	 Stelara (ustekinumab) IV injection may be approved if meeting the following criteria: a. The member has a diagnosis of moderate-to-severely active Crohn's disease or moderate-to-severely active ulcerative colitis AND b. The member is ≥ 18 years of age AND c. For treatment of moderately-to-severely active Crohn's disease, member has trial and failure‡ of one preferred adalimumab product and ENTYVIO (vedolizumab) OR For treatment of moderately-to-severely active ulcerative colitis, member has trial and failure‡ of one preferred adalimumab product and XELJANZ IR and ENTYVIO (vedolizumab) AND d. If meeting criteria listed above, prior authorization approval will be placed based on the following: i. If maintenance subcutaneous therapy will be billed as a medical claim for administration in the doctor's office or other clinical setting, initial 16-week approval will be placed for initial IV dosage (one dose) and subcutaneous formulations (HCPCS J3357) and one-year prior authorization approval for continuation of subcutaneous maintenance therapy may be provided based on clinical response OR 	See criteria

Drug	Criteria	PAR Length
	ii. If maintenance subcutaneous therapy will be dispensed by a pharmacy for self-administration by the member or for administration in the member's home or LTCF, initial approval will be for initial intravenous dose only.	
	Maximum Dose: 520 mg initial IV dose for members weighing > 85 Kg (187 pounds)	
	Quantity Limit: For initial IV infusion, four 130 mg/26 mL single-dose vials	
	*Members currently stabilized on a Stelara (ustekinumab) regimen that was initiated prior to 1/1/2023 may receive prior authorization approval for continuation of therapy without meeting the above criteria.	
	‡Failure is defined as lack of efficacy, allergy, intolerable side effects, contraindication to therapy, or significant drug-drug interaction. Note that trial and failure of Xeljanz IR will not be required when prescribed for ulcerative colitis for members ≥ 50 years of age that have an additional CV risk factor.	
Tepezza (teprotumumab)	Tepezza (teprotumumab) may be approved if the member meets the following criteria: a. Member is 18 years of age or older AND b. Member has a documented diagnosis of Thyroid Eye Disease (TED) AND c. Member's prescriber must be in consultation with an ophthalmologist or endocrinologist AND d. Member does not require immediate surgical ophthalmological intervention AND e. Member does not currently require orbital (eye) surgery and is not planning corrective surgery/irradiation during therapy AND f. Member is euthyroid, mild hypothyroid, mild hyperthyroid (defined as free thyroxine (FT4) and free triiodothyronine (FT3) levels less than 50% above or below the normal limits) or seeking care for dysthyroid state from an endocrinologist or other provider experienced in the treatment of thyroid diseases AND g. Member does not have corneal decompensation unresponsive to medical management AND h. Member had an inadequate response, or there is a contraindication or intolerance, to	One year
		ii. If maintenance subcutaneous therapy will be dispensed by a pharmacy for self-administration by the member or for administration in the member's home or LTCF, initial approval will be for initial intravenous dose only. Maximum Dose: 520 mg initial IV dose for members weighing > 85 Kg (187 pounds) Quantity Limit: For initial IV infusion, four 130 mg/26 mL single-dose vials *Members currently stabilized on a Stelara (ustekinumab) regimen that was initiated prior to 1/1/2023 may receive prior authorization approval for continuation of therapy without meeting the above criteria. ‡Failure is defined as lack of efficacy, allergy, intolerable side effects, contraindication to therapy, or significant drug-drug interaction. Note that trial and failure of Xeljanz IR will not be required when prescribed for ulcerative colitis for members ≥ 50 years of age that have an additional CV risk factor. Tepezza (teprotumumab) may be approved if the member meets the following criteria: a. Member is 18 years of age or older AND b. Member has a documented diagnosis of Thyroid Eye Disease (TED) AND c. Member's prescriber must be in consultation with an ophthalmologist or endocrinologist AND d. Member does not require immediate surgical ophthalmological intervention AND e. Member does not require immediate surgical ophthalmological intervention AND d. Member does not require immediate surgical ophthalmological intervention AND e. Member does not currently require orbital (eye) surgery and is not planning corrective surgery/irradiation during therapy AND f. Member is euthyroid, mild hypothyroid, mild hypothyroid (defined as free thyroxine (FT3) levels less than 50% above or below the normal limits) or seeking care for dysthyroid state from an endocrinologist or other provider experienced in the treatment of thyroid diseases AND g. Member does not have comeal decompensation unresponsive to medical

HCPCS	Drug	Criteria	PAR Length
		dose of teprotumumab. If member becomes pregnant during treatment, Tepezza should be discontinued, AND j. If member is diabetic, member is being managed by an endocrinologist or other provider experienced in the treatment and stabilization of diabetes AND k. Authorization will be issued for one course of therapy of eight infusions Maximum Dose: Eight infusions per one year	
J2356	Tezspire (tezepelumab-ekko)	Tezspire (tezepelumab-ekko) may be approved if the following criteria are met: a. Member is 12 years of age or older AND b. Member has a diagnosis of severe asthma that is uncontrolled or inadequately controlled as demonstrated by i. 2 or more asthma exacerbations requiring use of oral or systemic corticosteroids and/or hospitalizations and/or ER visits in the year prior to medication initiation c. Medication is being administered as add-on therapy (not monotherapy) AND d. Member is taking a high dose inhaled corticosteroid and a long-acting beta agonist AND e. Medication will not be used in concomitantly with other biologics indicated for asthma AND f. Member has documented baseline FEV1 Reauthorization may be approved if member has shown clinical improvement as documented by one of the following a. Improvement in lung function, measured in FEV1 b. Reduction in the number of asthma exacerbations, defined as a decrease in use of oral or systemic corticosteroids and/or reduced asthma related hospitalizations and/or ER visits Maximum dose: 210 mg once every 4 weeks Exemption: Members currently stabilized on a Tezspire (tezepelumab-ekko) regimen that was initiated prior to 1/1/2023 may receive prior authorization approval for continuation of therapy without meeting the above criteria.	One year

HCPCS	Drug	Criteria	PAR Length
J1303	Ultomiris (ravulizumab-cwvz)	Ultomiris (ravulizumab-cwvz) may be approved if member meets the following criteria: a. Member is diagnosed with either Paroxysmal Nocturnal Hemoglobinuria (PNH), Atypical Hemolytic Uremic Syndrome (aHUS), Generalized Myasthenia Gravis (gMG), or Neuromyelitis Optica Spectrum Disorder (NMOSD) AND b. Member has been vaccinated for meningococcal disease according to current ACIP guidelines at least two weeks prior to medication initiation OR c. Member is receiving 2 weeks of antibacterial drug prophylaxis if meningococcal vaccination cannot be administered at least 2 weeks prior to starting requested medication AND d. Member does not have unresolved Neisseria meningitidis or any systemic infection e. Prescriber is enrolled in the Ultomiris Risk Evaluation and Mitigation Strategy (REMS) program AND f. Medication is administered by or in consultation with a hematologist for PNH and by or in consultation with a hematologist or nephrologist for aHUS and by or in consultation with a neurologist or pMG and by or in consultation with a neurologist or ophthalmologist for NMOSD AND g. Member meets criteria listed below for specific diagnosis: i. Paroxysmal nocturnal hemoglobinuria (PNH) 1. Member is one month of age or older if prescribing the IV formulation OR is ≥ 18 years of age if prescribing the subcutaneous formulation AND 2. Diagnosis of PNH must be accompanied by detection of PNH clones by flow cytometry diagnostic testing AND 3. Baseline values are documented for the following: □ Serum lactate dehydrogenase (LDH) □ Hemoglobin levels □ Packed RBC transfusion requirement	One year
		AND 4. Member has one of the following indications for therapy: □ Presence of a thrombotic event	
		Presence of a unombodic event Presence of organ dysfunction secondary to chronic hemolysis Member is transfusion dependent	

HCPCS	Drug	Criteria	PAR Length
		☐ Member has uncontrolled pain secondary to chronic hemolysis	
		ii. Atypical hemolytic uremic syndrome (aHUS)	
		1. Member is one month of age or older if prescribing the IV	
		formulation OR is ≥ 18 years of age if prescribing the	
		subcutaneous formulation AND	
		2. Member does not have Shiga toxin E. coli related HUS (STEC-	
		HUS) AND	
		3. Thrombotic Thrombocytopenic Purpura (TTP) has been ruled out	
		by evaluating ADAMTS13 level or a trial of plasma exchange did	
		not result in clinical improvement AND	
		4. Baseline values are documented for the following:	
		☐ Serum LDH	
		☐ Serum creatinine/eGFR	
		☐ Platelet count	
		☐ Dialysis requirement	
		iii. <u>Generalized myasthenia gravis</u>	
		1. Member is 18 years of age or older AND	
		2. Member has a positive serologic test for anti-acetylcholine	
		receptor (AchR) antibodies	
		3. Member has Myasthenia Gravis Foundation of America (MGFA)	
		Clinical Classification of Class II to IV disease; AND	
		4. Member has a MG-Activities of Daily Living (MG-ADL) total	
		score of ≥6; AND 5. Member has trial and failure of treatment over at least 1 year with	
		at least 2 immunosuppressive therapies (e.g., azathioprine,	
		cyclosporine, mycophenolate, etc.), or has failed at least 1	
		immunosuppressive therapy and required chronic plasmapheresis	
		or plasma exchange (PE) or intravenous immunoglobulin (IVIG)	
		iv. Neuromyelitis optica spectrum disorder (NMOSD):	
		1. Member is 18 years of age or older AND	
		2. Member has a positive test for anti-aquaporin-4 (AQP4)	
		antibodies AND	
		3. Exclusion of alternative diagnoses have been evaluated AND	
		4. Member has one of the following clinical characteristics:	

HCPCS	Drug	Criteria	PAR Length
		 □ Optic neuritis □ Acute myelitis □ Area postrema syndrome (episode of otherwise unexplained hiccups or nausea and vomiting) □ Acute brainstem syndrome □ Symptomatic narcolepsy or acute diencephalic clinical syndrome with NMOSD-typical diencephalic MRI lesions □ Symptomatic cerebral syndrome with NMOSD-typical brain lesions 	
		Maximum dose: 3.6 g every 8 weeks (IV infusion) 490 mg once weekly (subcutaneous administration)	
J2777	Vabysmo (faricimab)	Vabysmo (faricimab) may be approved if all the following criteria are met: a. Member is 18 years of age or older AND b. Member has a diagnosis of one of the following: i. Neovascular (wet) Age-Related Macular Degeneration (nAMD) ii. Diabetic Macular Edema (DME) iii. Macular Edema following Retinal Vein Occlusion (RVO) AND c. Member does not have any of the following: i. Ocular or periocular infection ii. Active intraocular inflammation iii. Hypersensitivity to the requested medication AND d. Member's best corrected visual acuity (BCVA) is measured at baseline and throughout treatment AND e. Requested medication will not be used with other ophthalmic vascular endothelial growth factor (VEGF) inhibitors AND f. Documentation of trial and failure* with bevacizumab containing product AND g. Documentation of the dosing regimen that is being requested	See criteria

HCPCS	Drug	Criteria	PAR Length
		Reauthorization: may be approved for one year if the following criteria are met: a. Documentation of improvement or stabilization of disease state and visual status AND b. Documentation of ongoing treatment regimen AND c. Prescriber attests that the member has had no significant adverse effects or drug toxicity, such as endophthalmitis and retinal detachments, increase in intraocular pressure, arterial thromboembolic events (ATE), retinal vasculitis and/or retinal vascular occlusion AND d. Prescriber attests requested medication will not be used with other ophthalmic VEGF inhibitors	
		Maximum dose: nAMD/DME: 6 mg every 4 weeks RVO: 6 mg every 4 weeks for 6 months *Failure is defined as lack of efficacy with a three-month trial, contraindication to therapy, allergy, intolerable side effects, or significant drug-drug interaction	
J3032	Vyepti (eptinezumab jjmr)	Vyepti (eptinezumab-jjmr) may be approved if member meets the following criteria: a. Member is 18 years of age or older AND b. Member has a diagnosis of episodic (fewer than 15 headache days monthly) or chronic migraine (headaches occurring 15 days or more monthly, where at least 8 of these days per month for at least 3 months are migraine days with or without aura) AND c. Member has tried and failed two oral preventive pharmacological agents listed as Level A per the most current American Headache Society/American Academy of Neurology guidelines (such as divalproex, topiramate, metoprolol, propranolol). Failure is defined as lack of efficacy, allergy, intolerable side effects, or significant drug-drug interaction AND d. The requested medication is not being used in combination with another CGRP medication AND e. Member has trial and failure of all preferred calcitonin gene-related peptide inhibitors (CGRPis) indicated for preventative therapy listed on the pharmacy benefit preferred drug list AND f. Initial dose is no more than 100 mg every 3 months	See criteria

HCPCS	Drug	Criteria	PAR Length
		 i. If 300 mg is requested, the member has tried and had an inadequate response (no less than 30% reduction in headache frequency in a 4-week period) to the 100 mg dosage. g. Initial authorization will be limited to 6 months. Continuation (12-month authorization) will require documentation of clinically relevant improvement with no less than 30% reduction in headache frequency in a 4-week period. 	
		Initial approval: 6 months	
		Reauthorization: One year	
		Maximum dose: 300 mg IV every 3 months	
J3401	Vyjuvek (beremagene geperpavec-svdt)	 Vyjuvek (beremagene geperpavec-svdt) may be approved if the following criteria are met: a. Member is ≥ 6 months of age, AND b. Member has a documented diagnosis of dystrophic epidermolysis bullosa AND c. Member must have undergone genetic testing confirming mutation(s) in the collagen type VII alpha 1 chain (COL7A1) gene AND d. The requested medication is being prescribed by or in consultation with a provider who has expertise in treating dystrophic epidermolysis bullosa AND e. Member has been counseled regarding use of highly effective contraceptive method(s) while receiving treatment Reauthorization: Prescribing provider attests that clinical condition is improving on Vyjevek therapy 	One year
J9332 J9334	Vyvgart (efgartigimod alfa) Vyvgart Hytrulo (efgartigimod alfa and hyaluronidase-qvfc)	Vyvgart (efgartigimod alfa) or Vyvgart Hytrulo (efgartigimod alfa/ hyaluronidase-qvfc) may be approved when ALL the following criteria are met:	One year
		 a. Member is ≥ 18 years of age AND b. The requested medication is being prescribed for treatment of generalized myasthenia gravis that is anti-acetylcholine receptor (AChR) antibody positive AND c. The member meets the criteria for Myasthenia Gravis Foundation of America (MGFA) clinical classification class II to IV AND 	

HCPCS	Drug	Criteria	PAR Length
		 d. The requested medication is being prescribed by or in consultation with a neurologist AND e. Provider will perform a myasthenia gravis functionality score (such as the MGADL or QMG) at baseline. 	
		Reauthorization: Additional one year approval may be granted with provider attestation that a follow-up myasthenia gravis functionality assessment indicates stable symptoms or clinical improvement.	
		Maximum Dose:	
		IV formulation: 1,200 mg weekly for 4 weeks	
		Subcutaneous formulation: 1,008 mg weekly for 4 weeks	
		Quantity Limit:	
		IV formulation: Twelve 400 mg/20 mL single-dose vials per 28 days	
		Subcutaneous formulation: Four 1,008 mg/5.6 mL single-dose vials per 28 days	
		Succession formation four 1,000 mg/510 m2 single dose viais per 20 days	
J2357	Xolair (omalizumab)	Xolair (omalizumab) may be approved if member meets ALL the following criteria for the	One year
		appropriate indication:	
		a. If administered for the treatment of <u>asthma</u> :	
		i. Member is 6 years of age or older AND	
		ii. Member has a diagnosis of moderate to severe asthma persistent asthma	
		whose symptoms are inadequately controlled with inhaled corticosteroids	
		with one of the following: 1. A pre-treatment IgE serum concentration greater than or equal to	
		30 IU per mL OR	
		2. A positive skin test or in vitro reactivity to a perennial inhaled	
		allergen AND	
		iii. Member's moderate to severe asthma has been refractory to recommended	
		evidence-based, guideline-supported pharmacologic therapies AND	
		iv. Medication is being prescribed as add-on therapy to existing asthma	
		regimen AND	
		v. Medication will not be used concomitantly with other biologics indicated	
		for asthma AND	
		vi. Maximum dose of 750mg every 4 weeks	

HCPCS	Drug	Criteria	PAR Length
		b. Reauthorization for <u>asthma</u> indication may be approved if member has shown	
		clinical improvement as documented by one of the following	
		i. Improvement in lung function, measured in FEV1 OR	
		ii. Reduction in the number of asthma exacerbations, defined as a decrease in	
		use of oral or systemic corticosteroids and/or reduced asthma related	
		hospitalizations and/or ER visits	
		c. If administered for the treatment of <u>chronic idiopathic urticaria</u> (CIU)	
		i. Member is 12 years of age or older AND	
		ii. Member is diagnosed with chronic idiopathic urticaria AND	
		iii. Member is symptomatic despite H1 antihistamine treatment AND	
		iv. Member has tried and failed at least three of the following:	
		1. Hydroxyzine or doxepin (must include)	
		2. High-dose second generation H1 antihistamine	
		3. H2 antihistamine	
		4. First-generation antihistamine	
		5. Leukotriene receptor antagonist	
		AND	
		v. Prescriber attests that the need for continued therapy will be periodically	
		reassessed (as the appropriate duration of therapy for CIU has currently not	
		been evaluated) AND	
		vi. Exemption: Member who is currently stable on Xolair for chronic idiopathic urticaria may continue to receive prior authorization approval to	
		continue.	
		d. If administered for the treatment of chronic rhinosinusitis with nasal polyps:	
		i. If the member has a concomitant diagnosis of asthma or chronic idiopathic	
		urticaria, then criteria listed above for the respective diagnoses are met	
		AND	
		ii. Member is 18 years of age or older AND	
		iii. Member has a pre-treatment IgE level greater than or equal to 30 IU per	
		mL AND	
		iv. Member has tried and failed at least two intranasal corticosteroids (see	
		Intranasal Rhinitis Agents PDL class). Failure is defined as lack of efficacy	
		with a 2-week trial, contraindication to therapy, allergy, intolerable side	
		effects, or significant drug-drug interaction	
		v. AND	
		vi. Member is <i>currently</i> adherent to intranasal corticosteroid therapy AND	

HCPCS	Drug	Criteria	PAR Length
J1748	Zymfentra (infliximab-dyyb)	viii. Member has a baseline bilateral endoscopic nasal polyps score indicating the need for treatment AND viii. Medication is being prescribed by or in consultation with a qualified subspecialist such as an allergist, ear/nose/throat specialist, immunologist, rheumatologist, or pulmonologist AND ix. Maximum dose for nasal polyps is 600 mg subcutaneously every 2 weeks e. Reauthorization for the chronic rhinosinusitis with nasal polyps indication may be approved if member has shown clinical improvement as indicated by the following: i. Initial approval criteria were met at the time of initiation of therapy AND ii. Provider attests that member has documented improvement in bilateral endoscopic nasal polyps score, AND iii. Provider attests that member is being periodically reassessed for need for continued therapy based on disease severity and/or level of symptom control Zymfentra (infliximab-dyyb) may be approved for members meeting all the following criteria: a. The requested medication is being prescribed for treating moderately-to-severely active Crohn's disease or moderately-to-severely active Ulcerative Colitis in alignment with indicated use outlined in FDA-approved product labeling AND b. The requested medication meets FDA-labeled indicated age for prescribed use AND c. For treatment of moderately-to-severely active Crohn's disease, member has trial and failure; of one preferred adalimumab product OR for treatment of moderately-to-severely active ulcerative colitis, member has trial and failure; of one preferred adalimumab product OR for treatment of moderately-to-severely active ulcerative colitis, member has trial and failure; of one preferred adalimumab product OR for treatment of moderately-to-severely active ulcerative colitis, member has trial and failure; of one preferred adalimumab product and XELJANZ IR.	One Year