Appendix P



#### Colorado Medical Assistance Program Prior Authorization Procedures, Coverage Policies and Drug Utilization Criteria Health First Colorado Pharmacy Benefit For Physicians and Pharmacists

Drug products requiring a prior authorization for the Health First Colorado pharmacy benefit are listed in this document. Prior authorization criteria are based on FDA product labeling, CMS approved compendia, clinical practice guidelines, and peer-reviewed medical literature.

#### **Prior Authorization Procedures:**

• Prior authorizations may be called or faxed to the helpdesk at:

Phone:1-800-424-5725Fax:1-888-424-5881

- Products qualify for a 3-day emergency supply in an emergency situation. In this case, call the helpdesk for an override.
- Prior authorization (PA) forms are available by visiting <u>https://www.colorado.gov/hcpf/pharmacy-resources</u> .
- 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.
- Pharmacists from long-term-care pharmacies and infusion pharmacy must obtain a signature from someone who is authorized to prescribe drugs before they submit PA forms.
- Pharmacists from long-term-care pharmacies and infusion pharmacies can request a PA by phone if specified in the criteria.
- 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.

#### **Early Refill Limitations:**

• Non-controlled prescriptions may be refilled after 75% of previous fill is used. Controlled substance prescriptions (DEA Schedule 2 through 5) may be refilled after 85% of the previous fill is used. Synagis may be refilled after 92.5% of the previous fill is used.

#### **Medical Supply Products and Medications:**

- All supplies, including insulin needles, food supplements and diabetic supplies are not covered under the pharmacy benefit, but are covered as medical supply items through the Durable Medical Equipment (DME) benefit.
- If a medical benefit requires a PA, the PA request can be submitted through the provider application available at <a href="http://www.coloradopar.com/">http://www.coloradopar.com/</a>
- DME questions should be directed to Gainwell Technologies (Formerly DXC Technology) 1-844-235-2387. Only policy questions regarding Durable Medical Equipment should be directed to the state at 303-866-3406.

#### **Physician Administered Drugs and Medical Billing:**

• Physician administered drugs (PADs) include any medication or medication formulation that is administered intravenously or 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). PAD criteria listed on Appendix P apply specifically to drug products when billed through the Health First Colorado pharmacy 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 "Physician Administered Drugs" section below). PADs administered by a healthcare professional in the office, clinic, dialysis unit, or outpatient hospital settings should be billed through the Health First Colorado medical benefit using the standard buy-and-bill process and following procedures outlined in the PAD Billing Manual (found on the PAD Resources Page at <a href="https://www.colorado.gov/hcpf/physician-administered-drugs">https://www.colorado.gov/hcpf/physician-administered-drugs</a>).

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	ROGRAM APPENDICES Criteria	DAD
Drug	Criteria	PAR Length
Drug classes that have been migrated to the Preferred Drug List (PDL)	Anticoagulants (oral), Antidepressants, Antiemetics, Antiherpetics, Antihistamines with decongestants, Antihypertensives, Antiplatelets, Atypical Antipsychotics (oral), Bisphosphonates (oral), Constipation (opioid-induced), Diabetes Management Classes, Erythropoiesis Stimulating Agents, Fibromyalgia Agents,	
https://www.colorado.gov/hc pf/pharmacy-resources	Filgrastim/Pegfilgrastim/Sargromastim/Filgrastim-SNDZ, Fluoroquinolones, Growth Hormones, Hepatitis C Virus Treatments, Insulin, Intranasal Corticosteroids, Leukotrienes, Multiple Sclerosis Agents, Neurocognitive Disorder Agents, Ophthalmic Allergy Products, Otezla (apremilast), Overactive Bladder Agents, Pancreatic Enzymes, Proton Pump Inhibitors, Pulmonary Arterial Hypertension	
	Therapies, Respiratory Inhalants, Sedative Hypnotics, Skeletal Muscle Relaxants, Stimulants and other ADHD Agents, Targeted Immune Modulators (self- administered), Testosterone Products, Topical Immunomodulators, Triptans	
ACETAMINOPHEN CONTAINING PRODUCT	A prior authorization is required for dosages of acetaminophen exceeding 4000mg/day.	N/A
MAXIMUM DOSING ADAKVEO (crizanlizumab-tmca)	Doses over 4000mg/day are not qualified for emergency 3 day supply approval <b>ADAKVEO</b> (crizanlizumab-tmca) may be approved for members meeting the following criteria: • Medication is being administered in the member's home or in a	One year
	<ul> <li>Medication is being administered in the memoer's none of in a long-term care facility by a healthcare professional AND</li> <li>Medication is being used to reduce the frequency of vasoocclusive crises (VOCs) in adults and pediatric patients aged 16 years and older with sickle cell disease.</li> </ul>	
	Maximum dose: Adakveo 5mg/kg every 2 weeks (IV Infusion)	
ALBUMIN	Albumin products may be approved if meeting the following criteria: Medication is given in the member's home or in a long-term care facility AND Administration is for one of the following FDA-approved indications: Hypoproteinemia Burns Shock due to: Burns Trauma Surgery Infection Erythrocyte resuspension Acute nephrosis Renal dialysis Hyperbilirubinemia Erythroblastosis fetalis	One year
ALDURAZYME (laronidase)	<ul> <li>ALDURAZYME (laronidase) may be approved for members meeting the following criteria:</li> <li>Aldurazyme (laronidase) is being administered in a long-term care facility or in a member's home by a healthcare professional AND</li> <li>Member is 6 months of age or older AND</li> <li>Member does not have acute febrile or respiratory illness AND</li> <li>Member does not have progressive/irreversible severe cognitive impairment AND</li> <li>Member has a diagnosis of Mucopolysaccharidosis, Type 1 confirmed by one of the following:</li> </ul>	One year

COLORADO MEDICAID	PROGRAM	Appendices	
	0	Detection of pathogenic mutations in the IDUA gene by molecular genetic testing OR Detection of deficient activity of the α-L-iduronidase lysosomal	
		enzyme	
	AND		
	• Membe	er has a diagnosis of one of the following subtypes:	
	0	Diagnosis of Hurler (severe) or Hurler-Scheie (attenuated) forms of disease OR	
	0	Diagnosis of Scheie (attenuated) form of disease with moderate to severe symptoms	
	AND		
		yme (laronidase) is being prescribed by or in consultation with a er who specializes in inherited metabolic disorders AND	
		er has a documented baseline value for urinary glycosaminoglycan	
		er has a documented baseline value for one of the following based on	
	0 o	Members $\geq$ 6 years of age: percent predicted forced vital capacity (FVC) and/or 6- minute walk test OR	
	0	Members 6 months to 6 years of age: cardiac status, upper airway obstruction during sleep, growth velocity, mental development, FVC, and/or 6-minute walk test	
	Doouthorization	Critorio	
	<u>Reauthorization</u>		
	following:	nember may receive approval to continue therapy if meeting the	
		cumented reduction in uGAG levels AND	
		monstrated stability or improvement in one of the following based on	
	age:		
	0	Members $\geq$ 6 years of age: stability or improvement in percent predicted FVC and/or 6-minute walk test OR	
	0	Members 6 months to less than 6 years of age: stability or	
		improvement in cardiac status, upper airway obstruction during	
		sleep, growth velocity, mental development, FVC and/or 6-minute walk test	
	Max dose: 0.58	mg/kg as a 3 to 4-hour infusion weekly.	
ALINIA (nitazoxanide)	ALINIA (nitazo	exanide) may be approved if meeting the following criteria:	
		being prescribed for diarrhea caused by Giardia lamblia or	
		dium parvum AND	
	••••	1 year of age or older AND	
		iarrhea due to C. parvum in members with Human Immunodeficiency	
	U	) infection, the member is receiving antiretroviral therapy AND	
		meets the following FDA-labeled dosing:	
	Age (years)	Dosage of Nitazoxanide Duration	
	1-3	5 mL (100mg) oral suspension every 12 hours with food	
	4-11	10 mL (200mg) oral suspension every 12 hours with food 3 days	
	>11	500mg orally every 12 hours with food	

#### APPENDICES

	Note: The tablet product formulation is currently not reported as an active drug in the Medicaid Drug Rebate Program (MDRP) and will not be covered until such a time that there is change made to rebate status for this product.	
ALLERGY EXTRACT PRODUCTS (Oral)	<ul> <li>GRASTEK (timothy grass pollen allergen extract):</li> <li>Must be between 5 and 65 years old.</li> <li>Must not be pregnant or nursing.</li> <li>Must be prescribed by an allergist.</li> <li>Must have a documented diagnosis to ONLY timothy grass pollen allergen extract or the Pooideae family (meadow fescue, orchard, perennial rye, Kentucky blue, and red top grasses) confirmed by positive skin test or IgE antibodies.</li> <li>Must have tried and falled allergy shots for reasons other than needle phobia. Failure is defined as: lack of efficacy, allergy, intolerable side effects, or significant drug-drug interaction.</li> <li>Must be willing to administer epinephrine in case of severe allergic reaction.</li> <li>Must be started 12 weeks prior to the season if giving only seasonally.</li> <li>Must be taken daily for up to 3 consecutive years.</li> <li>Must NOT have:</li> <li>Severe, unstable or uncontrolled asthma</li> <li>Had an allergic reaction in the past that included trouble breathing, dizziness or fainting, rapid or weak hearbeat</li> <li>Ever had difficulty with breathing due to swelling of the throat or upper airway after using any sublingual immunotherapy before</li> <li>Been diagnosed with eosinophilic esophagitis</li> <li>Allergic to any of the inactive ingredients contained in Grastek which include gelatin, mannitol, and sodium hydroxide</li> <li>A medical condition that may reduce the ability to survive a serious allergic reaction including but not limited to: markedly compromised lung function, unstable angina, recent myocardial infarction, significant arrhythmia, and uncontrolled hypertension.</li> <li>Taking medications that can potentiate or inhibit the effect of epinephrine including but not limited to: beta-adrenergic blockers, alpha-adrenergic blockers, ergot alkaloids, tricyclic antidepressants, levothyroxine, monoamine oxidase inhibitors, certain antihistamines, cardiac glycosides, and diuretics.</li> <li>Be taken with other immunotherapy (oral or injectable)</li> <li>ORALAIR (sweet v</li></ul>	One year
	Must NOT have:	

COLORADO MEDICAID PROG	RAM APPENDICES	
• • • • • • • • • • • • • • • • •	Severe, unstable or uncontrolled asthma Had an allergic reaction in the past that included trouble breathing, dizziness or ainting, rapid or weak heartbeat Ever had difficulty with breathing due to swelling of the throat or upper airway after using any sublingual immunotherapy before Been diagnosed with eosinophilic esophagitis Allergic to any of the inactive ingredients contained in Oralair which include nannitol, microcrystalline cellulose, croscarmellose sodium, colloidal anhydrous silica, magnesium stearate, and lactose monohydrate. A medical condition that may reduce the ability to survive a serious allergic reaction including but not limited to: markedly compromised lung function, instable angina, recent myocardial infarction, significant arrhythmia, and uncontrolled hypertension. Faking medications that can potentiate or inhibit the effect of epinephrine ncluding but not limited to: beta-adrenergic blockers, alpha-adrenergic blockers, ergot alkaloids, tricyclic antidepressants, levothyroxine, monoamine oxidase nhibitors, certain antihistamines, cardiac glycosides, and diuretics. Be taken with other immunotherapy (oral or injectable) <b>EWITEK</b> (short ragweed pollen allergen extract): the between 18 and 65 years old. be started 12 weeks prior to the season and only prescribed seasonally. not be pregnant or nursing. be prescribed by an allergist. have a documented diagnosis to ONLY short ragweed pollen allergen extract or ambrosia family (giant, false, and western ragweed) confirmed by positive skin or IgE antibodies. have tried and failed allergy shots for reasons other than needle phobia. Failure fined as: lack of efficacy, allergy, intolerable side effects, or significant drug- interaction. be willing to administer epinephrine in case of a severe allergic reaction. take first dose in physician's office.	
	NOT have: Severe, unstable or uncontrolled asthma Had an allergic reaction in the past that included trouble breathing, dizziness or fainting, rapid or weak heartbeat Ever had difficulty with breathing due to swelling of the throat or upper airway after using any sublingual immunotherapy before Been diagnosed with eosinophilic esophagitis Allergic to any of the inactive ingredients contained in Ragwitek which include gelatin, mannitol, and sodium hydroxide A medical condition that may reduce the ability to survive a serious allergic reaction including but not limited to: markedly compromised lung function, unstable angina, recent myocardial infarction, significant arrhythmia, and uncontrolled hypertension. Faking medications that can potentiate or inhibit the effect of epinephrine ncluding but not limited to: beta-adrenergic blockers, alpha-adrenergic blockers, ergot alkaloids, tricyclic antidepressants, levothyroxine, monoamine oxidase nhibitors, certain antihistamines, cardiac glycosides, and diuretics. Be taken with other immunotherapy (oral or injectable)	
ALPHA-1 PROTEINASE FDA INHIBITORS facilit	*	Lifetime

COLORADO MEDICAID P	ROGRAM APPENDICES	
	<ul> <li>Aralast: Chronic augmentation therapy in members having congenital deficiency of Alpha –1 Proteinase Inhibitor with clinically evident emphysema</li> <li>Prolastin: Emphysema associated with Alpha-1 Antitrypsin Deficiency</li> <li>Zemaira: Chronic augmentation and maintenance therapy in members with Alpha-1 Proteinase Inhibitor deficiency with clinically evident emphysema</li> </ul>	
ANOREXIANTS	Weight loss medications are not a covered benefit. Adipex P (phentermine) Belviq (lorcaserin) Contrave (naltrexone/bupropion) Lomaira (phentermine) Phentermine Qsymia (phentermine/topiramate ER) Saxenda (liraglutide) Xenical (Orlistat)	Weight loss drugs are not a covered benefit.
ANTI-ANEMIA MEDICATIONS	<ul> <li>Oral prescription iron products may be approved for members with a diagnosis of iron deficient anemia (applies to products available by prescription only)</li> <li>Injectable anti-anemia agents (such as Infed®, Ferrlecit®, Venofer®, Dexferrum®) may be approved for members meeting the following criteria: <ul> <li>Member has a diagnosis of iron deficient anemia AND</li> <li>Oral preparations are ineffective or cannot be used AND</li> <li>Medication is being administered in a long-term care facility or in the member's home by a home healthcare provider</li> </ul> </li> <li>Note: For coverage criteria for OTC ferrous sulfate and ferrous gluconate, refer to "OTC Products" section.</li> </ul>	Lifetime
ATYPICAL ANTIPSYCHOTIC INJECTABLES Abilify Maintena, Aristada, Geodon injection, Invega Sustenna, Invega Trinza, Perseris ER, Risperdal Consta, Zyprexa Relprevv	A prior authorization may be approved for when the medication is administered in a long-term care facility or in a member's home by a healthcare professional. Oral atypical antipsychotic criteria can be found on the preferred drug list.	One year
AVEED (testosterone undecanoate)	<ul> <li>Aveed<sup>®</sup> (testosterone undecanoate) prior authorization may be approved for members who are receiving the injection in their home or in a long-term care facility and have met all of the following criteria:</li> <li>Male patient ≥ 18 years of age AND</li> <li>Has a documented diagnosis of hypogonadotropic or primary hypogonadism (Patients with other diagnoses will require a manual review) AND</li> <li>Has two documented low serum testosterone levels below the lower limit of normal range for testing laboratory prior to initiation of therapy AND</li> <li>Does not have a diagnosis of breast or prostate cancer AND</li> <li>Does not have a palpable prostate nodule or prostate-specific antigen (PSA) &gt; 4ng/mL AND</li> <li>Has normal liver function tests prior to initiation of therapy AND</li> <li>Has trail and failure of two preferred agents from PDL class "Androgenic Agents," one trial must be testosterone cypionate injection.</li> </ul>	One year
BACTROBAN (mupirocin) Cream and Nasal Ointment	<b>Bactroban Cream</b> (mupirocin calcium cream) must be prescribed for the treatment of secondarily infected traumatic skin lesions (up to 10 cm in length or 100 cm <sup>2</sup> in	Cream: One year

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	total area), impetigo, infected eczema or folliculitis caused by susceptible strains of Staphylococcus aureus and Streptococcus pyogenes. <b>Bactroban Nasal Ointment</b> (mupirocin calcium) must be prescribed for the eradication of nasal colonization with methicillin-resistant Staphylococcus aureus in adult patients and health care workers as part of a comprehensive infection control program to reduce the risk of infection among patients at high risk of methicillin- resistant S. aureus infection during institutional outbreaks of infections with this pathogen.	Nasal Ointment Lifetime
BARBITURATES Coverage for Medicare dual- eligible members	Dual-eligible Medicare-Medicaid Beneficiaries: Beginning on January 1, 2013 Colorado Medicaid will no longer cover barbiturates for Medicare-Medicaid enrollees (dual-eligible members). For Medicaid primary members, barbiturates will be approved for use in epilepsy, cancer, chronic mental health disorder, sedation, treatment of insomnia, tension headache, muscle contraction headache and treatment of raised intracranial pressure. All other uses will require manual review	(3 month for neonatal narcotic abstinence syndrome
BENLYSTA (belimumab)	<ul> <li>Benlysta (belimumab) prior authorization may be approved only when documentation has been received indicating that the drug is being administered in the member's home or long-term care facility. The member must also meet the following criteria:</li> <li>Member is age ≥ 5 years with active, autoantibody-positive systemic lupus erythematosus (SLE) and receiving standard therapy OR member is an adult with active lupus nephritis who are receiving standard therapy AND</li> <li>Member has incomplete response to standard therapy from at least two of the following therapeutic classes: antimalarials, immunosuppressants and glucocorticoids; AND</li> <li>Member maintains standard therapy while on BENLYSTA (belimumab).</li> </ul>	One year
<b>BENZODIAZEPINES</b> Dual-eligible Medicare- Medicaid Beneficiaries	Dual-eligible Medicare-Medicaid Beneficiaries: Benzodiazepines will no longer be a Medicaid benefit for Medicare-Medicaid enrollees (dual-eligible members). The claims are no longer excluded from Medicare part D coverage and therefore must be billed to Medicare part D. Colorado Medicaid will no longer cover these medications for these members beginning on January 1, 2013.	One year
BONE RESORPTION SUPPRESSION AND RELATED AGENTS (Injectable Formulations) Boniva, Aredia, Miacalcin, Zemplar, Hectorol, Zometa, Reclast, Pamidronate, Prolia, Ganite	<ul> <li>A prior authorization will only be approved as a pharmacy benefit when the medication is administered in a long-term care facility or in a member's home.</li> <li>Prolia<sup>®</sup> (denosumab) will be approved if the member Meets the following criteria: <ul> <li>Member is in a long term care facility or home health (this medication is required to be administered by a healthcare professional) AND</li> <li>Member has one of the following diagnoses: <ul> <li>Postmenopausal osteoporosis with high fracture risk</li> <li>Osteoporosis</li> <li>Bone loss in men receiving androgen deprivation therapy in prostate cancer</li> <li>Bone loss in women receiving adjuvant aromatase inhibitor therapy for breast cancer</li> <li>AND</li> </ul> </li> <li>Member has serum calcium greater than 8.5mg/dL AND</li> <li>Member is taking calcium 1000 mg daily and at least 400 IU vitamin D daily AND</li> <li>Has trial and failure of preferred bisphosphonate for one year AND (Failure is defined as: lack of efficacy, allergy, intolerable side effects, or significant drug-drug interaction)</li> <li>Member meets ANY of the following criteria: <ul> <li>has a history of an osteoporotic vertebral or hip fracture</li> </ul> </li> </ul></li></ul>	One year

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	$\circ$ has a pre-treatment T-score of < -2.5	
	$\circ$ has a pre-treatment T-score of < -1 but > -2.5 AND either of the following:	
	• Pre-treatment FRAX score of > 20% for any major fracture	
	• Pre-treatment FRAX score of > 3% for hip fracture	
	Maximum dose of Prolia is 60mg every 6 months	
BLOOD PRODUCTS	FDA approved indications if given in the member's home or in a long-term care facility:	Lifetime
	• Plasma protein fraction; shock due to burns, trauma, surgery; hypoproteinemia; adult respiratory distress syndrome; cardiopulmonary bypass; liver failure; renal dialysis; or hemophilia.	
<b>BOTULINUM TOXIN</b> Botox, Dysport, Myobloc, Xeomin	<ul> <li>Botulinium toxin agents may receive approval if meeting the following criteria:</li> <li>Medication is being administered in a long-term care facility or the member's home by a healthcare professional AND</li> </ul>	One year
	• Member has a diagnosis of cervical or facial dystonia	
	Not approved for Cosmetic Purposes	
BOWEL PREPERATION AGENTS	For the following Bowel Preparation Agents, members will require a prior authorization for quantities exceeding 2 units in 30 days.	30 days
	<ul><li>Colyte</li><li>Gavilyte-C</li></ul>	
	<ul><li>Gavilyte-C</li><li>Gavilyte-H</li></ul>	
	Gavilyte-N	
	Gialax	
	• Golytely <sup>®</sup>	
	Moviprep	
	• Peg-Prep	
	• Suprep	
	• Sutab	
	• Trilyte	
BRAND FAVORED	See "Brand Favored Product List" on the Pharmacy Resources webpage at	
MEDICATIONS	https://www.colorado.gov/pacific/hcpf/pharmacy-resources .	
<b>BUPRENORPHINE-</b>	Bunavail <sup>®</sup> (buprenorphine/naloxone) buccal film will be approved for members who	One year
CONTAINING	meet all of the following criteria:	
PRODUCTS	• Approval will be granted if the prescriber meets the qualification criteria under	
(used for opioid use disorder/opioid dependency*)	the Drug Addiction Treatment Act (DATA) of 2000 and has been issued a unique DEA identification number by the DEA, indicating that he or she is qualified	
	under the DATA to prescribe Subutex® or Suboxone® AND	
	• The member has a diagnosis of opioid dependence AND	
	• The member is 16 years of age or older AND	
	<ul> <li>No claims data show concomitant use of opiates in the preceding 30 days unless</li> </ul>	
	the physician attests the member is no longer using opioids AND	
	• The member must have tried and failed, intolerant to, or has contraindication to generic buprenorphine/naloxone SL tablets or Suboxone® films.	
	<b>Buprenorphine/Naloxone</b> sublingual film will be approved if the all of following criteria are met:	
	• Effective 10/01/19: Brand Suboxone <sup>®</sup> sublingual film is covered as a favored	
	product, and for members meeting all of the following criteria (or members with	
	current prior authorization approval on file), claims for brand Suboxone®	
	sublingual film will pay with submission of DAW code 0, 1, or 9. Prior authorization for generic buprenorphine/naloxone sublingual film will require	

	prescriber verification that there is clinical necessity for use of the generic	
	product in addition to meeting all of the following:	
	• The prescriber is authorized to prescribe Suboxone AND	
	• The member has an opioid dependency AND	
	• The member is not currently receiving an opioid or opioid combination	
	product unless the physician attests the member is no longer using	
	opioids AND	
	• Will not be approved for the treatment of pain AND	
	• Opioid claims will not be allowed for members with a claim for	
	Suboxone in the preceding 30 days AND	
	• Will not be approved for more than 24mg of buprenorphine/day	
	Buprenorphine/Naloxone sublingual tablet will be approved if all of the following	
	criteria are met:	
	• The prescriber is authorized to prescribe buprenorphine/naloxone AND	
	• The member has an opioid dependency AND	
	• The member is not currently receiving an opioid or opioid combination product	
	unless the physician attests the member is no longer using opioids AND	
	• Will not be approved for the treatment of pain AND	
	• Opioid claims will not be allowed for members with a claim for Suboxone in the	
	preceding 30 days AND	
	<ul> <li>Will not be approved for more than 24mg of buprenorphine/day</li> </ul>	
	Sublocade <sup>®</sup> (buprenorphine extended-release) injection will be approved for	
	members who meet all of the following criteria:	
	• Sublocade is being administered in a long-term care facility or in a member's	
	home by a home healthcare provider (all other claims must be submitted through	
	the medical benefit) AND	
	• Sublocade is being dispensed directly to the home healthcare professional	
	(medication should not be dispensed directly to the member) AND	
	• Provider attests to member's enrollment in a complete treatment program	
	including counseling and psychosocial support AND	
	<ul> <li>Member must have documented diagnosis of moderate to severe opioid use</li> </ul>	
	disorder AND	
	• Member must have initiated therapy with a transmucosal buprenorphine-	
	containing product, and had dose adjustment for a minimum of 7 days AND	
	• Maximum dose is 300 mg injection every month	
	Suboxone <sup>®</sup> sublingual film (brand name) will be approved if all of the following	
	criteria are met:	
	• The prescriber is authorized to prescribe Suboxone AND	
	• The member has an opioid dependency AND	
	<ul> <li>The member is not currently receiving an opioid or opioid combination product</li> </ul>	
	unless the physician attests the member is no longer using opioids AND	
	<ul> <li>Will not be approved for the treatment of pain AND</li> </ul>	
	<ul> <li>Opioid claims will not be allowed for members with a claim for Suboxone in the</li> </ul>	
	preceding 30 days AND	
	• Will not be approved for more than 24mg of buprenorphine/day	
	Subutex <sup>®</sup> (buprenorphine) sublingual tablet will be approved if all of the following	
	criteria are met:	
	The prescriber is authorized to prescribe Subutex AND	
	<ul> <li>The member has an opioid dependency AND</li> <li>The member is program or the member is allerois to Nelsyons AND</li> </ul>	
	The member is pregnant or the member is allergic to Naloxone AND     Subuty will not be approved for the treatment of poin AND	
	Subutex will not be approved for the treatment of pain AND	
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	• Subutex will not be approved for more than 24mg/day	
	<b>Zubsolv</b> <sup>®</sup> (buprenorphine/naloxone) sublingual tablet will be approved if all of the following criteria are met:	
	• Approval will be granted if the prescriber meets the qualification criteria under the Drug Addiction Treatment Act (DATA) of 2000 and has been issued a unique DEA identification number by the DEA, indicating that he or she is qualified under the DATA to prescribe Subutex or Suboxone AND	
	<ul> <li>The member has a diagnosis of opioid dependence AND</li> </ul>	
	<ul> <li>The member is 16 years of age or older AND</li> </ul>	
	<ul> <li>No claims data show concomitant use of opiates in the preceding 30 days unless the physician attests the member is no longer using opioids AND</li> <li>The member must have tried and failed, intolerant to, or has a contraindication to generic buprenorphine/naloxone SL tablets or Suboxone films.</li> </ul>	
	*Buprenorphine products indicated for treating pain are located on the preferred drug list (PDL)	
BYNFEZIA (octreotide acetate)	<b>BYNFEZIA</b> (octreotide acetate) may be approved if all of the following criteria are met:	One year
	<ul> <li>Member is an adult (≥ 18 years of age) with a confirmed diagnosis of acromegaly OR severe diarrhea and flushing episodes associated with metastatic carcinoid tumors OR vasoactive intestinal peptide tumors (VIPomas) AND</li> <li>Bynfezia (octreotide acetate) is prescribed by, or in consultation with, an endocrinologist or oncologist AND</li> <li>Member has trialed and failed octreotide acetate injection solution (vial). Failure is defined as lack of efficacy, contraindication to therapy, allergy, intolerable side effects, or significant drug-drug interaction AND</li> <li>Provider confirms that member has had a baseline thyroid function test drawn prior to the initiation of Bynfezia (octreotide) and plans to monitor periodically during treatment AND</li> <li>For treatment indication acromegaly, the following criteria are met:         <ul> <li>The member has trialed and failed bromocriptine mesylate at maximally tolerated doses. Failure is defined as lack of efficacy, contraindication to therapy, allergy, intolerable side effects, or significant drug-drug interaction AND</li> </ul> </li> </ul>	
	<ul> <li><u>Maximum Dose</u>:</li> <li>Acromegaly: 1500 mcg/day (doses &gt; 300 mcg/day may not result in additional benefit)</li> <li>Carcinoid Tumors: 750 mcg/day</li> <li>VIPomas: 750 mcg/day (doses &gt; 450 mcg/day are generally not required)</li> </ul>	
CERDELGA (eliglustat)	<ul> <li>Cerdelga® (eliglustat) may be approved if all of the following criteria are met:</li> <li>Member has a diagnosis of Gaucher disease type 1 AND</li> <li>Documentation has been provided to the Department that the member is a CYP2D6 extensive, intermediate, or poor metabolizer as detected by an FDA cleared test AND</li> <li>Members who are CYP2D6 intermediate or poor metabolizers are not taking a strong CYP3A inhibitor (e.g., indinavir, nelfinavir, ritonavir, saquinavir,</li> </ul>	One year

COLORADO MEDICAID		1
	<ul> <li>suboxone, erythromycin, clarithromycin, telithromycin, posaconazole, itraconazole, ketoconazole, nefazodone) AND</li> <li>Members who are CYP2D6 extensive or intermediate metabolizers are not receiving strong or moderate CYP2D6 inhibitors (e.g, sertraline, duloxetine, quinidine, paroxetine, fluoxetine, buproprion, terbinafine) AND a strong or moderate CYP3A inhibitor (e.g, indinavir, nelfinavir, ritonavir, saquinavir, suboxone, erythromycin, clarithromycin, telithromycin, posaconazole, itraconazole, ketoconazole, fluconazole, nefazodone, verapamil, diltiazem)</li> <li>Quantity Limits: Max 60 tablets/30 days</li> </ul>	
CHLOROQUINE	Effective 03/24/20: Prior authorization may be approved for FDA-labeled indication, dose, age, and role in therapy as outlined in product package labeling.	Chronic conditions: One year
		Acute conditions: Duration of acute use
CLIENT OVERUTILIZATION PROGRAM (COUP)	Effective 9/14/19, pharmacy claims for members enrolled in Health First Colorado's COUP (Client Overutilization Program) program may deny for these members when filling prescriptions at a pharmacy that is not their designated COUP lock-in pharmacy or filling a medication prescribed by a provider that is not their designated COUP lock-in prescriber.	
	Health First Colorado Reginal Accountable Entity (RAE) organizations work with members enrolled in COUP to assist with coordinating care and improving services provided to these members. <u>Members and providers should contact the member's RAE organization for questions regarding the COUP program</u> . <sup>*</sup> Contact information for Health First Colorado RAE regions can be found at <u>https://www.colorado.gov/pacific/hcpf/accphase2</u> .	
	Additional information regarding the COUP program and enrollment criteria can be accessed at <u>https://www.colorado.gov/pacific/hcpf/client-overutilization-program</u> .	
	*For questions regarding pharmacy claims denials <u>that are unable to be addressed</u> during normal RAE organizational business hours (M-F 8:00 AM – 4:00 PM Mountain Standard Time), members and providers may contact the Magellan Helpdesk at 1-800-424-5725.	
CONTRACEPTIVE TWELVE-MONTH SUPPLY	Prescription Contraceptive Products (oral and topical):         Initial fills may be dispensed for up to a three-month supply to establish tolerance         (lack of adverse events). If the prescribed medication is tolerated for at least three         months of therapy, subsequent fills of that medication will be eligible to be filled for         up to a twelve-month supply.	One year
	Effective 01/20/2020, brand Nuvaring is covered as favored product and claims for brand will pay with submission of DAW code 0, 1, or 9. Generic equivalent etonorgetstral/ethinyl estradiol vaginal ring products require prior authorization and may be approved based on prescriber verification that there is clinical necessity of use of the generic product.	
COUGH AND COLD (Prescription Products)	Depot and IUD formulations are billed through the medical benefit.           Effective 03/19/20*, select prescription cough and cold products are covered for members of all ages without prior authorization. Eligible products include: <ul></ul>	One year

#### COLORADO MEDICAID PROGRAM APPENDICES Coverage of all other prescription cough and cold medications (not identified above) will be subject to meeting the following criteria: For members < 21 years of age, no prior authorization is required OR For members $\geq 21$ years of age, prior authorization may be approved with diagnosis of a chronic condition (such as COPD or asthma). For members with dual Medicare eligibility, pharmacy claims for prescription cough and cold medications prescribed for chronic conditions should be billed to Medicare. Prescription cough and cold medications prescribed for dual Medicare eligible members for acute conditions are covered through the Health First Colorado pharmacy benefit with completion of prior authorization verifying use for acute illness. Note: For OTC cough and cold product coverage, see "OTC Products" section. \*Until such time changes are implemented in the claims system, pharmacies may call the Magellan helpdesk at 1-800-424-5725 for prior authorization overrides for eligible products. **DALIRESP** (roflumilast) DALIRESP (roflumilast) tablets may be approved for members that meet the One year following criteria: Member has a diagnosis for severe COPD with history of COPD exacerbations (2 or more per year) and chronic bronchitis AND Member must be greater than 18 years of age AND • Member must have failed a trial of two of the following: long-acting beta2 agonist, preferred anticholinergic/anticholinergic combination, or preferred inhaled anticholinergic/anticholinergic combinations due to lack of efficacy, allergy, intolerable side effects or significant drug-drug interaction AND Member must not have moderate to severe liver disease (Child Pugh B or C). Note: this medication is not a bronchodilator and cannot be used for acute bronchospasms DARAPRIM **DARAPRIM** (pyrimethamine) may be approved if all the following criteria are met: 8 weeks (pyrimethamine) Member is being treated for toxoplasmic encephalitis or congenital toxoplasmosis or receiving prophylaxis for congenital toxoplasmosis AND Daraprim is prescribed in conjunction with an infectious disease specialist AND Member does not have megaloblastic anemia due to folate deficiency AND For prophylaxis, member has experienced intolerance to prior treatment with trimethoprim-sulfamethoxazole (TMP-SMX) meeting one of the following: Member has been re-challenged with trimethoprim-sulfamethoxazole (TMP-SMX) using a desensitization protocol and is still unable to tolerate Member has evidence of life threatening-reaction to trimethoprim-0 sulfamethoxazole (TMP-SMX) in the past (e.g. toxic epidermal necrolysis (TEN), Stevens-Johnson syndrome) OR Member is being treated for acute malaria due to susceptible strains of plasmodia • AND Member has tried and had an inadequate response or intolerant to two other malaria treatment regimens (such as but not limited to atovaquone/proguanil, Coartem, chloroquine, hydroxychloroquine, chloroquine plus Primaquine, quinine plus clindamycin, quinidine plus doxycycline) AND Daraprim is prescribed in conjunction with an infectious disease specialist with travel/tropical medicine expertise AND Member does not have megaloblastic anemia due to folate deficiency

	Note The Control of the Control of the American American Structure and the Structure	
	Note: The Center for Disease Control does not recommend Daraprim for the	
DEGLODING	prevention or the treatment of malaria	
DESI DRUGS	DESI drugs (Drugs designated by the Food and Drug Administration as Less Than	
	Effective Drug Efficacy Study Implementation medications) are not a covered	
DIFICID (fidementicity)	benefit.	1
DIFICID (fidoxomicin)	<b>DIFICID</b> (fidoxomicin) may be approved if all the following criteria are met:	1 month
	• Member is age $\geq 6$ months <b>AND</b>	
	Member has a documented diagnosis (including any applicable labs and/or tests)     for Clostridium difficile-associated diarrhea <b>AND</b>	
	• Prescribed by or in conjunction with a gastroenterologist or an infectious disease specialist <b>AND</b>	
	*	
	• Member has failed at least a 10 day treatment course of oral vancomycin.	
	Failure is defined as lack of efficacy, allergy, intolerable side effects, or significant drug-drug interaction.	
	significant drug-drug interaction.	
	Maximum quantity:	
	20 tablets per 30 days	
	136 mL per 10 days	
	150 mL per 10 days	
DIHYDROERGOTAMINE	MIGRANAL and dihydroergotamine product formulations will be approved if	One year
PRODUCTS	member meets ALL of the following criteria:	One year
inobeens	Member is not currently taking a potent CYP 3A4 inhibitor (for	
	example, protease inhibitor, macrolide antibiotic) AND	
	• Member does not have uncontrolled hypertension or ischemic heart	
	disease AND	
	• Product is being prescribed for cluster headache (vial only) or acute	
	migraine treatment (vial and nasal spray) AND	
	• Intranasal dihydroergotamine generic and Migranal <sup>®</sup> will be approved	
	with adequate trial and/or failure of dihydroergotamine vial (Failure is	
	defined as: lack of efficacy with 10 day trial, allergy, intolerable side	
	effects or significant drug-drug interactions)	
	AND	
	• If dihydroergotamine product is being prescribed for acute migraine	
	treatment, member has adequate trial and/or failure of 2 triptan agents	
	(for example sumatriptan, naratriptan)and 1 NSAID medication. Failure	
	is defined as lack of efficacy with 10 day trial, allergy, intolerable side	
	effects or significant drug-drug interactions.	
	OR	
	• If dihydroergotamine product is being prescribed for cluster headaches,	
	member has adequate trial and/or failure of 2 triptan agents. Failure is	
	defined as: lack of efficacy with 10 day trial, allergy, intolerable side	
	effects or significant drug-drug interactions.	
	Crondfotharing	
	Grandfathering: Members currently utilizing Migranel or a dihydrograptamine formulation (based on	
	Members currently utilizing Migranal or a dihydroergotamine formulation (based on recent claims history) may receive one year approval to continue therapy with that	
	medication.	
	Maximum Dosing:	
	Dihydroergotamine nasal spray and Migranal: 16mg per 28 days	
	Dihydroergotamine vial: 24mg per 28 days	
	_ 2mj aroongoumine that 2 mg por 20 aujo	1

DOPTELET (avatrombopag)	<ul> <li>DOPTELET (avatrombopag) prior authorization may be approved for members meeting the following criteria:</li> <li>Member is 18 years of age or older AND</li> <li>Member has a confirmed diagnosis of thrombocytopenia with chronic liver disease who is scheduled to undergo an elective procedure AND</li> <li>Member has trial and failure of Mulpleta (lusutrombopag). Failure is defined as a lack of efficacy, allergy, intolerable side effects, or significant drug-drug interactions.</li> <li>Quantity Limit: 5 day supply per procedure OR</li> <li>Member has a documented diagnosis of chronic immune thrombocytopenia AND</li> <li>Member has trial and failure of Promacta (eltrombopag). Failure is defined as a lack of efficacy, allergy, intolerable side effects, or significant drug-drug interactions.</li> <li>Quantity Limit: 40mg daily</li> </ul>	One year
DOXEPIN TOPICAL PRODUCTS	<ul> <li>Prudoxin<sup>®</sup> and generic doxepin 5% cream may be approved if the member meets the following criteria:         <ul> <li>Member is 18 years of age or older AND</li> <li>Member has a diagnosis of moderate pruritis with atopic dermatitis or lichen simplex chronicus AND</li> <li>Member has trial and failure<sup>‡</sup> of one prescription-strength topical corticosteroid AND one topical immunomodulator product (see PDL for preferred products)</li> </ul> </li> <li>Zonalon<sup>®</sup> may be approved if member has trial and failed<sup>‡</sup> either doxepin 5% cream or Prudoxin<sup>®</sup> and meets all of the following criteria.         <ul> <li>Member has a diagnosis of moderate pruritis with atopic dermatitis or lichen simplex chronicus AND</li> <li>Member has a diagnosis of moderate pruritis with atopic dermatitis or lichen simplex chronicus AND</li> <li>Member has a diagnosis of moderate pruritis with atopic dermatitis or lichen simplex chronicus AND</li> <li>Member has trial and failure<sup>‡</sup> of one prescription-strength topical corticosteroid AND one topical immunomodulator product (see PDL for preferred products)</li> </ul> </li> <li>Quantity Limit for Topical Doxepin Products:         <ul> <li>8 days-supply per 30 day period</li> <li>‡Failure is defined as: lack of efficacy of a three-month trial, allergy, intolerable side effects or significant drug-drug interaction</li> </ul> </li> </ul>	One year
DUPIXENT (dupilumab)	effects or significant drug-drug interaction         Dupixent® (dupilumab) may be approved for members meeting the following criteria:         *Atopic Dermatitis:         • Member is 6 years of age or older AND         • Member has a diagnosis of moderate to severe chronic atopic dermatitis AND         • Member has baseline Investigator Global Assessment (IGA) score for atopic dermatitis severity of at least 3 (Scored 0-4, 4 being most severe) OR moderate erythema and moderate papulation/infiltration AND	Initial: See criteria Continued: One year

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<ul> <li>Member has been educated by provider regarding the elimination of exacerbating factors including aeroallergens, food allergens, and contact allergens AND</li> <li>Member has been educated by provider regarding the appropriate use of emollients and moisturizers for promotion of skin hydration AND</li> <li>Member has trialed and failed<sup>‡</sup> the following agents:         <ul> <li>Two medium potency to very-high potency topical corticosteroids [such as mometasone furoate, betamethasone dipropionate, or fluocinonide (see PDL for list of preferred products)] AND</li> <li>Two topical calcineurin inhibitors (see PDL for list of preferred products) AND</li> <li>Must be prescribed by or in conjunction with a dermatologist, allergist/immunologist, or rheumatologist AND</li> <li>Initial authorization will be for 18 weeks. Continuation will be authorized for 12 months with prescriber attestation to 16-week IGA score showing improvement by at least 2 points OR clinically significant improvement with Dupixent® regimen.</li> </ul> </li> <li>Member has a diagnosis of moderate to severe asthma (on medium to high dose inhaled corticosteroid and a long-acting beta agonist) with eosinophilic phenotype OR <u>oral</u> corticosteroid dependent asthma AND</li> <li>Member has had at least one asthma exacerbation in the past year requiring systemic corticosteroids or emergency department visit or hospitalization OR dependence on daily <u>oral</u> corticosteroid Herapy PLUS regular use of high dose inhaled corticosteroid plus an additional controller medication AND</li> <li>Metication is being prescribed as add-on therapy to existing regimen AND</li> </ul>	
*Asthma:	
• Member has a diagnosis of moderate to severe asthma (on medium to high dose inhaled corticosteroid and a long-acting beta agonist) with eosinophilic phenotype OR <u>oral</u> corticosteroid dependent	
• Member has had at least one asthma exacerbation in the past year requiring systemic corticosteroids or emergency department visit or hospitalization OR dependence on daily <u>oral</u> corticosteroid therapy PLUS regular use of high dose inhaled corticosteroid PLUS an	
• Medication is being prescribed by or in conjunction with a rheumatologist, allergist, or pulmonologist AND	
<ul> <li>For indication of moderate to severe asthma with eosinophilic phenotype:         <ul> <li>baseline lung function (FEV1) is provided and baseline eosinophils are greater than 300 cells/mcL AND</li> </ul> </li> </ul>	
<ul> <li>Initial authorization will be for 12 weeks. Continued authorization will require prescriber attestation of improvement in FEV<sub>1</sub> of 25% from baseline and will be for 12 months</li> </ul>	
<ul> <li>For indication of oral corticosteroid dependent asthma:         <ul> <li>Dosing of the oral corticosteroid is provided AND</li> <li>Initial authorization will be 24 weeks. Continued authorization will require prescriber attestation of a reduction of oral corticosteroid by at least 50% and will be for 12 months</li> </ul> </li> </ul>	
<ul> <li><u>*Chronic Rhinosinusitis with Nasal Polyposis:</u></li> <li>If member has a diagnosis of asthma or atopic dermatitis, they must</li> </ul>	
meet listed criteria for that indication	

	ROGRAM APPENDICES	
	<ul> <li>Member is 18 years of age or older AND</li> <li>Medication is being prescribed as an add-on maintenance treatment in adult patients with inadequately controlled chronic rhinosinusitis with nasal polyposis (CRSwNP) AND</li> <li>Member has a baseline bilateral endoscopic nasal polyps score (NPS; scale 0-8) AND nasal congestion/obstruction score (NC; scale 0-3) averaged over 28-day period AND</li> <li>Member has trialed and failed‡ therapy with three intranasal corticosteroids (see PDL Class) AND</li> <li>Medication is being prescribed by or in conjunction with a rheumatologist, allergist, ear/nose/throat specialist or pulmonologist AND</li> <li>Dose of Dupixent (dupilumab) 300mg every 2 weeks is used AND</li> <li>Initial authorization will be for 24 weeks, for additional approval member must meet the following criteria:         <ul> <li>NC and NPS scores are provided and show a 20% reduction in symptoms AND</li> <li>Member continues to use primary therapies such as intranasal corticosteroids</li> </ul> </li> <li>Quantity Limit: 2 syringes every 28 days after initial 14 days of therapy (first dose is twice the regular scheduled dose)</li> <li>*For members that have a diagnosis of asthma and/or atopic dermatitis in addition to another indicated diagnosis.</li> </ul>	
EGRIFTA (tesamorelin acetate)	<ul> <li>‡Failure is defined as a lack of efficacy with one month trial, allergy, intolerable side effects, contraindication to, or significant drug-drug interactions.</li> <li>Egrifta® or Egrifta SV® will be approved if all the following criteria is met:</li> <li>Must be prescribed in consultation with a physician who specializes in HIV/AIDS AND</li> <li>Member is 18 years of age or older AND</li> <li>Member has a diagnosis of HIV-related lipodystrophy with excess abdominal fat meeting the following criteria: <ul> <li>Male member must have a waist circumference of at least 95cm (37.4in) and a waist to hip ratio of at least 0.94 OR</li> <li>Female member must have a waist circumference of at least 94cm (37in) and a waist to hip ratio of at least 0.88 AND</li> <li>Baseline waist circumference and waist to hip ratio must be provided</li> </ul> </li> <li>Member is currently receiving highly active antiretroviral therapy including protease inhibitors, nucleoside reverse transcriptase inhibitor, or non-nucleoside reverse transcriptase inhibitors AND</li> <li>Member does not have a diagnosis of hypophysectomy, hypopituitarism, pituitary surgery, head irradiation or head trauma AND</li> <li>Member does not have any active malignancy or history of malignancy AND</li> <li>For women of childbearing potential, member must have a negative pregnancy test within one month of therapy initiation</li> </ul>	6 months
ELESTRIN GEL (estradiol)	A prior authorization will only be approved if a member has tried and failed on generic oral estradiol therapy and diagnosed with moderate-to-severe vasomotor symptoms (hot flashes) associated with menopause. (Failure is defined as: lack of efficacy, allergy, intolerable side effects or significant drug-drug interactions)	One year

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EMFLAZA (deflazacort)		pproved if all the	following criteria are met	APPENDICES	One year
EMIFLAZA (uenazacort)	<ul> <li>Member is at lear</li> </ul>			•	One year
			ne muscular dystrophy an	d a documented	
	mutation in the d				
			(per claims history or prov		
			therapy, adequate trial dur		
			ck of efficacy, allergy, into t drug-drug interactions) A		
			or in consultation with a pl		
			ichenne muscular dystroph		
	neuromuscular d				
			at least 10 times the upper	limit of normal at	
	some stage in the		ding tubanoulogic and han	atitic D minus	
	Absence of activ	e miection metu	ding tuberculosis and hepa	autis D virus	
	Maximum dose	of 0.9mg/kg dail	y for tablets and suspensio	on, may be rounded up	
	to nearest ml				
EMVERM (mebendazole)	Table 1. Emuran I	DA Annuoved D	osing and Duration in Adult	ta and Children	See Table
ENT VERMI (medendazore)	Table 1; Eniverni r	DA Approved Do	osing and Duration in Adult	is and Chhuren	See Table
	Diagnosis	Dose	Duration	Quantity Limits	
	Ancylostoma	100 mg	3 consecutive days,	6 tablets/member	
	duodenale or	twice daily	may be repeated in 3		
	Necator		weeks in needed.		
	americanus (hookworm)				
	Ascariasis	100 mg	3 consecutive days,	6 tablets/member	
	(roundworm)	twice daily	may be repeated in 3 weeks if needed.		
	Enterobiasis	100 mg	May give second dose	2 tablets/member	
	(pinworm)	once	in three weeks if		
	Trichuriasis	100 mg	needed.	6 tablets/member	
	(whipworm)	100 mg twice daily	3 consecutive days, may be repeated in 3	o tablets/member	
	(winpworm)	twice dury	weeks in needed.		
	<b>Emverm</b> ® will be a	nproved for mem	bers that meet the followi	ng criteria:	
	<ul> <li>Member is 2 year</li> </ul>			115 ci ittila.	
	•		f the following: Ancylosto	oma duodenale or	
			, Ascariasis (roundworm),	Enterobiasis	
	(pinworm), or Tr	-			
			ndazole for FDA approved ined as lack of efficacy, al		
			nteractions) AND	lergy, intolerable side	
			n, Emverm is being prescr	ribed by an infectious	
	disease specialis	t AND		-	
		-	e pregnancy test AND	11 2	
	• Emverm® Is bei (Table 1)	ng prescribed in	accordance to FDA dosing	g and duration	
	Quantity limits: Base	ed on indication (	(Table 1)		

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ENSPRYNG	ENSPRYNG (satralizumab-mwge) may be approved if meeting the following	Initial:
(satralizumab-mwge)	<ul> <li>criteria:</li> <li>Member is an adult (≥ 18 years of age) AND</li> <li>Member has a documented diagnosis of neuromyelitis optica spectrum disorder (NMOSD) that includes a positive serologic test for anti-aquaporin-4 (AQP4) antibodies AND</li> <li>Member has a past medical history of <u>at least one</u> of the following: <ul> <li>Optic neuritis</li> <li>Acute myelitis</li> <li>Acute myelitis</li> <li>Acute brainstem syndrome; episode of otherwise unexplained hiccups or nausea and vomiting</li> <li>Acute brainstem syndrome</li> <li>Symptomatic narcolepsy or acute diencephalic clinical syndrome with NMOSD-typical diencephalic MRI lesions</li> <li>Symptomatic cerebral syndrome with NMOSD-typical brain lesions AND</li> </ul> </li> <li>Member does not have any active infections, including localized infections AND</li> <li>Member does not have active Hepatitis B infection, as confirmed by negative surface antigen [HBsAg] and anti-HBV tests AND</li> <li>Member does not have active or untreated latent tuberculosis AND</li> <li>Provider confirms that member has a baseline Liver Function Panel drawn prior to initiation of ENGSPYNG treatment and member does not has an AST or ALT level greater than 1.5 times the upper limit of normal AND</li> <li>Provider confirms that neutrophil counts will be checked 4 to 8 weeks after initiation of ENSPRYNG therapy, and thereafter at regular clinically determined intervals to monitor for decreased neutrophil counts AND</li> <li>Provider has screened for immunizations the member is due to receive according to immunization guidelines AND</li> <li>Any live or live-attenuated vaccines will be administered at least 4 weeks prior to initiation of ENSPRYNG AND</li> <li>Any non-live vaccines will be administered at least 2 weeks prior to initiation of ENSPRYNG (whenever possible) AND</li> </ul>	6 months Continued: One year
	<ul> <li>Reauthorization Criteria: After receiving initial six month approval, EYNSPRYNG (satralizumab-mwge) may be approved for one year if the following criteria:</li> <li>Member has shown no adverse effects to ENGSPYNG treatment at a maintenance dose of 120 mg subcutaneously every 4 weeks AND</li> </ul>	
	<ul> <li>Member does not have any active infections (including localized infections) AND</li> <li>Member does not have an AST or ALT level greater than 1.5 times the upper limit of normal AND</li> <li>Provider confirms that neutrophil counts are currently within normal limits and will continue to be monitored at clinically determined intervals during ENSPRYNG therapy.</li> </ul>	
	Maximum dose: 120 mg subcutaneously every 2 weeks for three doses, followed by 120 mg subcutaneously every 4 weeks maintenance dose.	
ENTRESTO (sacubitril/valsartan)	<ul> <li>ENTRESTO (sacubitril/valsartan) may be approved for members if the following criteria has been met:</li> <li>Member has a diagnosis of heart failure with reduced ejection fraction and NYHA Class II to IV AND</li> </ul>	One year

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ERECTILE DYSFUNCTION OR SEXUAL DYSFUNCTION PRODUCTS	<ul> <li>Member is NOT currently on ACE-inhibitor or Angiotensin Receptor Blocking agent AND</li> <li>Member does not have history of angioedema related to previous ACE inhibitor or ARB therapy</li> <li>Medications prescribed for use for erectile dysfunction or other sexual dysfunction diagnoses are not covered.</li> <li>Yohimbine prior authorization may be approved for use as a mydriatic agent or a</li> </ul>	Not covered Do not
Caverject, Cialis, Edex, Imvexxy, Levitra, Muse, Viagra, Addyi, Osphena, Premarin Cream, Sildenafil, Tadalafil (generic Cialis), Staxyn, Stendra, Xiaflex, Yohimbine	<ul> <li>vasodilator (not related to erectile dysfunction). Prior authorizations for use of yohimbine for erectile dysfunction will not be approved.</li> <li>Sildenafil prior authorization may be approved for off-label use for Raynaud's disease.</li> </ul>	qualify for emergency 3 day supply
ERGOMAR (ergotamine tartrate)	<ul> <li>Ergomar<sup>®</sup> (ergotamine tartrate) sublingual tablet may be approved for members meeting the following criteria:</li> <li>Ergomar (ergotamine tartrate) is being prescribed to prevent or treat vascular headache (migraine, migraine variants or so-called "histaminic cephalalgia") AND</li> <li>Member has a negative pregnancy test within 30 days of receipt of Ergomar AND</li> <li>Member is not taking a potent CYP 3A4 inhibitor (ritonavir, nelfinavir, indinavir, erythromycin, clarithromycin and troleandomycin) AND</li> <li>Member has adequate trial and/or failure of 2 triptan agents (see PDL class) AND</li> <li>Member has adequate trial and/or failure of 2 NSAIDs (see PDL class) AND</li> <li>Member has adequate trial and/or failure of dihydroergotamine vial. Failure is defined as lack of efficacy with 10 day trial, allergy, intolerable side effects or significant drug-drug interactions.</li> <li>Maximum quantity: 20 tablets per 28 days (40mg per 28 days)</li> </ul>	One year
ESBRIET (pirenidone)	<ul> <li>Esbriet® may be approved if the following criteria are met:</li> <li>Member has been diagnosed with idiopathic pulmonary fibrosis AND</li> <li>Is being prescribed by or in conjunction with a pulmonologist AND</li> <li>Member is 18 years or older AND</li> <li>Member has baseline ALT, AST, and bilirubin prior to starting therapy AND</li> <li>Member does not have severe (Child Pugh C) hepatic impairment, severe renal impairment (Crcl&lt;30 ml/min), or end stage renal disease requiring dialysis AND</li> <li>Female members of reproductive potential must have been counseled regarding risk to the fetus AND</li> <li>Member is not receiving a strong CYP1A2 inducer (e.g, carbamazepine, phenytoin, rifampin)</li> </ul>	One year
EUCRISA (crisaborole)	<ul> <li>Eucrisa® may be approved if the following criteria are met:</li> <li>Member is at least 3 months of age and older AND</li> <li>Member has a diagnosis of mild to moderate atopic dermatitis AND</li> </ul>	One year

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	<ul> <li>Member has a history of failure, contraindication, or intolerance to at least two medium- to high-potency topical corticosteroid for a minimum of 2 weeks, or is not a candidate for topical corticosteroids AND</li> <li>Member must have trialed and/or failed pimecrolimus and tacrolimus. Failure is defined as a lack of efficacy, allergy, intolerable side effects, contraindication to, or significant drug-drug interactions. AND</li> <li>Must be prescribed by or in conjunction with a dermatologist or allergist/immunologist.</li> </ul>	
EVRYSDI (risdiplam)	EVRYSDI (risdiplam) may be approved if the following criteria are met:	15 months
EVRYSDI (risdiplam)	<ul> <li>Member is between 2 months of age and 25 years old AND</li> <li>Member has documented diagnosis of 5q-autosomal recessive spinal muscular atrophy (SMA) by genetic testing and SMN1 mutation (two or more SMN2 gene copies must be specified) AND</li> <li>Treating and prescribing provider(s) is a neurologist or pediatrician experienced in treatment of SMA AND</li> <li>The prescriber attests that the member will be assessed by <u>at least one</u> of the following exam scales at baseline and during subsequent office visits:         <ul> <li>Hammersmith Infant Neurological Examination Module 2 (HINE2)</li> <li>Children's Hospital of Philadelphia Infant Test of Neuromuscular Disorders (CHOP-INTEND)</li> <li>Hammersmith Functional Motor Scale Expanded (HFMSE)</li> <li>Bayley Scales of Infant and Toddler Development, Third Edition (BSID-III)</li> <li>Motor Function Measure (MFM-32)</li> <li>Revised Upper Limb Module (RULM)</li> </ul> </li> <li>AND</li> <li>Prior to the start of EVRYSDI treatment, the provider attests that the member meets all of the following:         <ul> <li>Female members of childbearing potential have a documented negative pregnancy test within 2 weeks of initiating EVRYSDI therapy AND</li> <li>Female members of childbearing potential have been instructed to use effective contraception during treatment AND</li> <li>Male members have been advised prior to initiation of therapy that their fertility may be compromised while being treated with EVRYSDI AND</li> <li>Baseline liver function panel has been drawn and does not indicate hepatic impairment (EVRYSDI is extensively metabolized by the liver) AND</li> <li>Drug-drug interactions including (but not limited to) MATE substrates such as metformin, cimetidine, and acyclovir, have been screened for, addressed if needed, and will be continually monitored</li> </ul> </li> <l< th=""><th>15 months</th></l<></ul>	15 months
	• The member is not receiving concomitant treatment with SPINRAZA (nusinersen) <b>OR</b> the member was treated with SPINRAZA previously and had to discontinue use due to lack of efficacy, allergy, intolerable side effects, or a contraindication to receiving intrathecal injections <b>AND</b>	

COLORADO MEDICAID F	ORADO MEDICAID PROGRAM A		
	<ul> <li>The member's weight is provided an dosing:</li> </ul>	nd meets recommended daily	
	Age and Body Weight	Recommended Daily Dosage	
	2 months to less than 2 years of age	0.2 mg/kg	
	2 years and older, weighing less than 20 kg	0.25 mg/kg	
	2 years and older, weighing 20 kg or more	5 mg	
	<ul> <li>Reauthorization criteria: After 15 months, memory if the following criteria are met:</li> <li>The member has shown no adverse events to</li> <li>The member has demonstrated response to traclinical improvement or no decline document the same exam scale(s) used prior to initiating number 4 of initial authorization criteria). Im symptoms must be compared to the baseline are be measured against the degenerative effects</li> <li>The prescriber provides the following inform <ul> <li>A brief explanation, including the provider other than the one who init completes any follow-up exam(s) A</li> <li>A brief explanation must be submitt scale used for initial authorization is</li> <li>The member does not have hepatic i</li> <li>Member weight is provided and mediated and mediate</li></ul></li></ul>	EVRYSDI treatment <b>AND</b> eatment by showing significant ted using quantitative scores using g EVRYSDI treatment (please see provement of SMA-related assessment and motor function must of SMA <b>AND</b> ation: rovider name, must be submitted if a ially performed the motor exam <b>ND</b> ed if an exam scale other than the used for reassessment <b>AND</b> mpairment <b>AND</b>	
	Age and Body Weight	Recommended Daily Dosage	
	2 months to less than 2 years of age	0.2 mg/kg	
	2 years and older, weighing less than 20 kg	0.25 mg/kg	
	2 years and older, weighing 20 kg or more	5 mg	
	Maximum dose: 5mg/day Above coverage standards will continue to be rev applicable changes due to the evolving nature of f available treatment options, and available peer-re- clinical evidence.	actors including disease course,	
EXJADE (deferasirox)	Please see "Jadenu and Exjade"		
EXONDYS 51 (eteplirsen)	<ul> <li>Exondys 51<sup>®</sup> may be approved if the following cr.</li> <li>Medication is being administered in the r care facility by a healthcare professional</li> <li>Member has a diagnosis of Duchenne M</li> <li>Member must have genetic testing confine that is amenable to exon 51 skipping AN</li> <li>Medication is prescribed by or in consult provider who specializes in treatment of cardiologist or pulmonary specialist) AN</li> <li>The member must be on corticosteroids a to corticosteroids AND</li> </ul>	member's home or in a long-term AND uscular Dystrophy (DMD) AND rming mutation of the DMD gene D tation with a neurologist or a DMD (i.e. pediatric neurologist, D	One year

Effective 04/01/2021 Revised 02/26/2021

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<ul> <li>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.</li> <li>Maximum Dose: 30 mg/kg per week</li> </ul>	0.000
<ul> <li>following criteria:</li> <li>Fasenra<sup>®</sup> is being administered by a healthcare professional in the member's home or in a long-term care facility (all other claims are billed through the Health First Colorado medical benefit) AND</li> <li>Member is 12 years of age or older AND</li> <li>Member has diagnosis of severe asthma with eosinophilic phenotype AND</li> <li>Member has eosinophil count of at least 300 cells/µl AND</li> <li>Fasenra is being prescribed as add-on therapy (not monotherapy) AND</li> <li>Member is taking a high dose inhaled corticosteroids and a long-acting beta agonist AND</li> <li>Member has had at least 2 asthma exacerbations requiring systemic corticosteroid therapy in the past 12 months</li> </ul>	One year
<ul> <li>Ferriprox® may be approved if the following criteria are met:</li> <li>Must be prescribed in conjunction with a hematologist or oncologist AND</li> <li>Member's weight must be provided AND</li> <li>Member has a diagnosis of transfusion-related iron overload due to thalassemia syndrome or sickle cell disease AND</li> <li>Member has an absolute neutrophil count &gt; 1.5 x 109 AND</li> <li>Member has failed or has had an inadequate response to Desferal (deferoxamine) AND Exjade (deferasirox) as defined by serum ferritin &gt;2,500mcg/L before treatment with Ferriprox OR member has been intolerant to or experienced clinically significant adverse effects to Desferal (deferoxamine) or Exjade (deferasirox) such as evidence of cardiac iron overload or iron-induced cardiac dysfunction.</li> </ul>	One year
<ul> <li>Firdapse<sup>®</sup> (amifampridine) may be approved for members meeting the following criteria:</li> <li>Member is an adult ≥ 18 years of age AND</li> <li>Member has a diagnosis of Lambert-Eaton myasthenic syndrome (LEMS)</li> <li>Max Dose: 80mg daily</li> </ul>	One year
<ul> <li><u>Prescription fluoride products:</u> <ul> <li>Prescription fluoride products will be approved for members less than 21 years of age without a prior authorization.</li> <li>For members 21 years of age or older approval will be granted if using well water or living in an under-fluoridated area designated by the CDC*.</li> <li>Approval for members not meeting these criteria will require a letter of necessity and will be individually reviewed.</li> </ul> </li> </ul>	One year
	<ul> <li>If the member is ambulatory, functional level determination of baseline assessment of ambulatory function is required QR if not ambulatory, member must have a Brooke Upper Extremity Function Scale of five or less documented QR a Forced Vital Capacity of 30% or more.</li> <li>Maximum Dose: 30 mg/kg per week</li> <li>Fasenra® prior authorization may be approved for member's meeting all of the following criteria:</li> <li>Fasenra® is being administered by a healthcare professional in the member's home or in a long-term care facility (all other claims are billed through the Health First Colorado medical benefit) AND</li> <li>Member is 12 years of age or older AND</li> <li>Member has diagnosis of severe asthma with eosinophilic phenotype AND</li> <li>Member has cosinophil count of at least 300 cells/µl AND</li> <li>Fasenra is being prescribed as add-on therapy (not monotherapy) AND</li> <li>Member is taking a high dose inhaled corticosteroids and a long-acting beta agonist AND</li> <li>Member has had at least 2 asthma exacerbations requiring systemic corticosteroid therapy in the past 12 months</li> <li>Maximum dose: 30mg subcutaneous injection every 4 weeks for 3 doses, then every 8 weeks thereafter</li> <li>Ferriprox® may be approved if the following criteria are met:</li> <li>Must be prescribed in conjunction with a hematologist or oncologist AND</li> <li>Member has a diagnosis of transfusion-related iron overload due to thalassemia syndrome or sickle cell disease AND</li> <li>Member has failed or has had an inadequate response to Desferal (deferoxamine) AND Exjade (deferasirox) such derives to Desferal (deferoxamine) AND Exjade (deferasirox) such arevise to Desferal (deferoxamine) AND Exjade (deferasirox) as defined by serum ferritin &gt;&gt;2,50mcg/L. before treatment with Ferriprox OR member has been intolerant to or experienced clinically significant adverse effects to Desferal (deferoxamine) with Seriprox OR member has been intolerant to or expe</li></ul>

ROGRAM APPENDICES	
<ul> <li>The following OTC fluoride products are eligible for prior authorization approval for all members using well water or living in an under-fluoridated area designated by the CDC*: fluoride chewable tablets, ludent fluoride chewable tablets, sodium fluoride 0.5mg/mL drops</li> <li>Approval for members not meeting these criteria will require a letter of necessity and will be individually reviewed.</li> <li>*Information and reports regarding water fluoridation can be found on the CDC website at: <a href="https://nccd.cdc.gov/DOH_MWF/Default/CountyList.aspx?state=Coloradateid=8&amp;st_ateabbr=CO&amp;reportLevel=2">https://nccd.cdc.gov/DOH_MWF/Default/CountyList.aspx?state=Coloradateid=8&amp;st_ateabbr=CO&amp;reportLevel=2</a>.</li> </ul>	
<ul> <li>If administered in the physician's office or delivered to physician's office, physician must bill as a medical claim on the 1500 claim form (no PA required).</li> <li>If administered in the member's home or in a long-term care facility, a prior authorization is required and must meet the criteria below for approval.</li> <li>Based on clinical trial data, ENF should be used as part of an <i>optimized</i> background regimen for treatment-experienced members:</li> <li>For treatment-experienced members with evidence of HIV-1 replication, treatment should include at least one antiretroviral agent with demonstrated HIV-1 susceptibility on the basis of genotypic/phenotypic <i>resistance</i> assays, and <i>two</i> "active" antiretroviral agents.</li> <li>Members must have limited treatment options among currently commercially available agents.</li> <li>Members must be 18 years of age or older with advanced HIV-1 infection, and not responding to approved antiretroviral therapy.</li> <li>Members must be 18 years of age or older with advanced HIV-1 infection, and not responding to approved antiretroviral therapy.</li> <li>Members must be active a CD4 lymphocyte count less than 100 cells/mm3 and a viral load greater than 10,000 copies/ml (measurement within the last 90 days).</li> <li>Past adherence must be demonstrated based on:</li> <li>Attendance at scheduled appointments, and/or</li> <li>Prior antiretroviral regimen adherence, and/or</li> <li>Utilization data from pharmacy showing member's use of medications as prescribed</li> <li>Ability to reconstitute and self-administer ENF therapy.</li> <li>At 24 weeks, members must experience at least ≥ 1 log<sub>10</sub> decrease in HIV RNA or have HIV RNA below quantifiable limits to continue treatment with ENF.</li> <li>Members are not eligible if antiretroviral treatment-naive and/or infected with HIV-2.</li> <li>Pre-approval is necessary</li> <li>Practitioner must either be Board Certified in Infectious Disease, or be an HIV experienced practitioner. Verification must be produced with the</li></ul>	Six months
the emergence of new data.	
<ul> <li>GALAFOLD (migalastat hydrochloride) prior authorization may be approved for members meeting the following criteria:</li> <li>Member is ≥ 12 years of age AND</li> <li>The medication is being prescribed by or in consultation with a neurologist AND</li> </ul>	One year
	<ul> <li>The following OTC fluoride products are eligible for prior authorization approval for all members using well water or living in an under-fluoridated area designated by the CDC*: fluoride chewable tablets, ludent fluoride chewable tablets, sodium fluoride 0.5mg/mL drops</li> <li>Approval for members not meeting these criteria will require a letter of necessity and will be individually reviewed.</li> <li>*Information and reports regarding water fluoridation can be found on the CDC website at: https://nccd.cdc.gov/DOH_MWF/Default/CountyList.aspx?state=Coloradateid=8&amp;st ateabbr=CO&amp;reportLevel=2.</li> <li>If administered in the physician's office or delivered to physician's office, physician must bill as a medical claim on the 1500 claim form (no PA required). If administered in the member's home or in a long-term care facility, a prior authorization is required and must meet the criteria below for approval.</li> <li>Based on clinical trial data, ENF should be used as part of an <i>optimized</i> background regimen for treatment-experienced members:         <ul> <li>For treatment-experienced members:</li> <li>For treatment-experienced members:</li> <li>Members must be ls of genotypic/phenotypic <i>resistance</i> assays, and <i>two</i> "active" anticroviral agents.</li> <li>Members must be 8 years of age or older with advanced HIV-1 infection, and not responding to approved antiretroviral therapy.</li> <li>Members must be 8 lyears of age or older with advanced HIV-1 infection, and not responding to approved antiretroviral therapy.</li> </ul> </li> <li>Members must be 18 years of age or older with advanced HIV-1 infection, and not responding to approved antiretroviral therapy.</li> <li>Members must be 8 demonstrated based on:         <ul> <li>Attendance at scheduled appointments, and/or</li> <li>Prior antiretroviral regime adherence, and/or</li> <li>With NA below quantifiable limits to cont</li></ul></li></ul>

COLORADO MEDICAID P		
	<ul> <li>Member has a confirmed diagnosis of Fabry's disease with an amenable galactose alpha gene (GLA) variant per in vitro assay data. (Amenable GLA variants are those determined by a clinical genetics professional as pathologic or likely pathologic) AND</li> <li>Member does not have severe renal impairment or end-stage renal disease requiring dialysis.</li> </ul>	
	Maximum dose: 123 mg once every other day	
GAMASTAN (immune globulin)	Prior authorization may be approved for FDA-labeled indication, dose, age, and role in therapy as outlined in package labeling.	One year
GATTEX (teduglutide)	<ul> <li>GATTEX (teduglitide) may be approved if all of the following criteria are met:</li> <li>Member is one year of age or older AND</li> <li>Member has documented short bowel syndrome AND</li> <li>Member is dependent on parenteral nutrition for twelve consecutive months AND</li> <li>The prescribing physician is a gastroenterologist AND</li> <li>Medical necessity documentation has been received and approved by Colorado Medicaid clinical staff (please fax to 303-866-3590 attn: Clinical Pharmacy Staff)</li> </ul>	Two months initially; may be approved by State for up to one year
GENERIC MANDATE	The initial prior authorization will be limited to a two month supply. Brand Name Medications and Generic Mandate:	
	<ul> <li>Brand name drug products that have a therapeutically equivalent generic drug product (as determined by the FDA) will require prior authorization for brand product coverage and will be covered without a prior authorization if meeting one of the following exceptions: <ul> <li>The brand name drug is prescribed for the treatment of (and the prescriber has indicated dispense as written on the brand name prescription):</li> <li>Biologically based mental illness defined in 10-16-104 (5.5) C.R.S.</li> <li>Cancer</li> <li>Epilepsy</li> <li>HIV/AIDS</li> </ul> </li> <li>The Department has determined that the brand name product is lower cost than the therapeutically equivalent generic</li> <li>Prior authorization for use of a brand name drug product that has a therapeutically equivalent generic (and does not meet exceptions above) may also be approved if: <ul> <li>The prescriber is of the opinion that a transition to the generic equivalent of the brand name drug would be unacceptably disruptive to the patient's stabilized drug regimen</li> <li>The patient is started on the generic equivalent drug but is unable to continue treatment on the generic drug as determined by the prescriber</li> </ul> </li> </ul>	
GLYCATE (glycopyrollate)	<ul> <li>Glycate<sup>®</sup> (glycopyrollate) may be approved for members meeting the following criteria:</li> <li>Member is 18 years of age or older AND</li> <li>Member has a diagnosis of peptic ulcer disease AND</li> <li>Member does not have any of the following conditions: <ul> <li>Glaucoma</li> <li>Obstructive uropathy (such as bladder neck obstruction due to prostatic hypertrophy)</li> <li>Obstructive disease of the gastrointestinal tract (such as achalasia, pyloroduodenal stenosis, etc.)</li> <li>Paralytic ileus</li> </ul> </li> </ul>	One year

<ul> <li>Intestinal atony of the clicrity or debilitated patient         <ul> <li>Unstable cardiovascular status in acute hemorrhage</li> <li>Severe ulcerative colitis</li> <li>Toxic megacolon complicating ulcerative colitis</li> <li>Myasthenia gravis</li> <li>AND</li> </ul> </li> <li>Member has tried and failed at least two proton pump inhibitors (failure is defined as lack of efficacy with 4 week trial, allergy, intolerable side effects, or significant drog-drug interaction) AND</li> <li>Glycate (glycopyrollate) is being prescribed by or in consultation by a gastroenterologist</li> <li>HETLIOZ (tasimeteon)</li> <li>Hetlioz (tasimeleon) may be approved for members meeting the following criteria:</li> <li>Have a documented diagnosis of non-24-hour sleep wake disorder (non-24 or N24) by a sleep specialist AND</li> <li>Member is completely blind</li> </ul> <li>HIGH COST CLAIMS</li> <li>Hetlioz (tasimeleon) may be approved for members meeting the following partDavia sleep specialist AND</li> <li>Member is completely blind</li> <li>Pharmacy claust meet the following per FDA product package labeling:</li> <li>Diagnosis for labeled indication AND</li> <li>Bused on prescribed indication, prescription meets the following per label:             <ul> <li>Dosing</li> <li>Strongth</li> <li>Dosing form</li> <li>Quantity</li> <li>Days Supply</li> </ul> </li> <li>Homozygous Familial</li> <li>Hardge@(ontipatio)</li> <li>Member is a storign section, Stappistion administered drugs (see "Physician Administered Drugs" section, Ministered Drugs" section, Ministered Drugs (see "Physician Administered Drugs" section, Ministered Drugs (see "Physician Administered Drugs" section).</li> <li>Member is 18 years of age or older;</li> <ul> <li>Member has fale dubrap with high dose statin thera</li></ul>	COLORADO MEDICAID P	PROGRAM APPENDICES	
<ul> <li>Have a documented diagnosis of non-24-hour sleep wake disorder (non-24 or N24) by a sleep specialist AND</li> <li>Member is completely bilid</li> <li>HIGH COST CLAIMS</li> <li>Pharmacy claims exceeding \$19,999.00 may be approved following pharmacist review if the product meets current criteria (on the PDL/Appendix P when listed) OR if not listed, must meet the following per FDA product package labeling:         <ul> <li>Diagnosis for labeled indication AND</li> <li>Based on prescribed indication, prescription meets the following per label:                 <ul> <li>Doing</li> <li>Strength</li> <li>Doage form</li> <li>Quantity</li> <li>Day Supply</li></ul></li></ul></li></ul>		<ul> <li>Unstable cardiovascular status in acute hemorrhage</li> <li>Severe ulcerative colitis</li> <li>Toxic megacolon complicating ulcerative colitis</li> <li>Myasthenia gravis</li> </ul> AND Member has tried and failed at least two proton pump inhibitors (failure is defined as lack of efficacy with 4 week trial, allergy, intolerable side effects, or significant drug-drug interaction) AND Glycate (glycopyrollate) is being used as adjunctive therapy AND Glycate (glycopyrollate) is being prescribed by or in consultation by a	
review if the product meets current criteria (on the PDL/Appendix P when listed) OR         if not listed, must meet the following per FDA product package labeling:         • Diagnosis for labeled indication AND         • Based on prescribed indication, prescription meets the following per label:         • Dosing         • Strength         • Days Supply         AND         • If product is an IV formulation or product labeling indicates that the medication should be administered by a healthcare professional, must meet approval criteria for physician administered drugs (see "Physician Administered Drugs" section).         Homozygous Familial Hypercholesterolemia (HoFH)         Hypercholesterolemia (HoFH)         (HoFH)         Kielde (Interpreted)         Member has failed therapy with high dose statin therapy (e.g. atorvastatin 40mg or higher, Crestor 20mg or higher)         • The prescribing physician is enrolled in the Juxtapid REMS program.         Kynamro@ (mipomersen) may be approved for members meeting all of the following criteria:         • Confirmed diagnosis of homozygous familial hypercholesterolemia (HoFH) as determined by either a or b         • The prescribing physician is enrolled in the Juxtapid B (HoFH) as determined by either a or b         • Laboratory tests confirming diagnosis of HoFH: LDLR DNA Sequence Analysis for large gene rearrangement testingonly if the Sequence Analysis for Barge gene rearrangement testingonly if the Sequence Analysis in negative OR APOB and dPCSK9 testing if both of th	HETLIOZ (tasimelteon)	• Have a documented diagnosis of non-24-hour sleep wake disorder (non-24 or N24) by a sleep specialist AND	One year
Hypercholesterolemia (HoFH)       Prior authorization may be approved if all of the following criteria are met:         • Member is 18 years of age or older;       • Member has documented diagnosis of homozygous familial hypercholesterolemia (HoFH);         • Member has failed therapy with high dose statin therapy (e.g. atorvastatin 40mg or higher, Crestor 20mg or higher)       • The prescribing physician is enrolled in the Juxtapid REMS program.         Kynamro® (mipomersen) may be approved for members meeting all of the following criteria:       • Confirmed diagnosis of homozygous familial hypercholesterolemia (HoFH) as determined by either a or b         a.       Laboratory tests confirming diagnosis of HoFH: LDLR DNA Sequence Analysis OR LDLR Deletion/Duplication Analysis for large gene rearrangement testing only if the Sequence Analysis is negative OR APOB and dPCSK9 testing if both of the above tests are negative but a	HIGH COST CLAIMS	<ul> <li>review if the product meets current criteria (on the PDL/Appendix P when listed) OR</li> <li>if not listed, must meet the following per FDA product package labeling: <ul> <li>Diagnosis for labeled indication AND</li> </ul> </li> <li>Based on prescribed indication, prescription meets the following per label: <ul> <li>Dosing</li> <li>Strength</li> <li>Dosage form</li> <li>Quantity</li> <li>Days Supply</li> </ul> </li> <li>AND</li> <li>If product is an IV formulation or product labeling indicates that the medication should be administered by a healthcare professional, must meet approval criteria for physician administered drugs (see "Physician")</li> </ul>	
b. Documentation is received confirming a clinical or laboratory diagnosis of	Hypercholesterolemia	<ul> <li>Prior authorization may be approved if all of the following criteria are met:</li> <li>Member is 18 years of age or older;</li> <li>Member has documented diagnosis of homozygous familial hypercholesterolemia (HoFH);</li> <li>Member has failed therapy with high dose statin therapy (e.g. atorvastatin 40mg or higher, Crestor 20mg or higher)</li> <li>The prescribing physician is enrolled in the Juxtapid REMS program.</li> <li>Kynamro® (mipomersen) may be approved for members meeting all of the following criteria:</li> <li>Confirmed diagnosis of homozygous familial hypercholesterolemia (HoFH) as determined by either a or b <ul> <li>a. Laboratory tests confirming diagnosis of HoFH:</li> <li>LDLR DNA Sequence Analysis OR</li> <li>LDLR Deletion/Duplication Analysis for large gene rearrangement testingonly if the Sequence Analysis is negative OR</li> <li>APOB and dPCSK9 testing if both of the above tests are negative but a strong clinical picture exists.</li> </ul> </li> </ul>	One year

COLORADO MEDICAID	PROGRAM APPENDICES	
	<ul> <li>Has a history of therapeutic failure, contraindication, or intolerance to high dose statin therapy or cholesterol absorption inhibitor (ezetimibe or bile acid resin) AND</li> <li>Is being prescribed by a physician specializing in metabolic lipid disorders AND</li> <li>The prescriber is enrolled in the REMS program AND</li> <li>Is not being used as monotherapy AND</li> <li>Has baseline liver function (AST, ALT, ALK, and total bilirubin) AND</li> <li>Does not have moderate or severe hepatic impairment or active liver disease.</li> </ul>	
HORIZANT (gabapentil enacarbil)	<ul> <li>Horizant® may be approved for members who have a diagnosis of <u>Restless Leg</u> <u>Syndrome</u> and who meet the following criteria:</li> <li>Member has failed a one month trial of Mirapex® (pramipexole) and Requip® (ropinorole) AND</li> <li>Member has had a positive therapeutic response to generic gabapentin but incomplete response due to duration of action.</li> <li><u>Max quantity</u>: 30 tablets/30 days</li> <li>Horizant® will be approved for members who have a diagnosis of <u>Post Herpetic</u> <u>Neuralgia</u> and who meet the following criteria:</li> <li>Member has failed a one month trial of tricyclic antidepressant, pregabalin and gabapentin</li> <li><u>Max quantity</u>: 60 tablets / 30 days</li> </ul>	One year
HORMONE THERAPY	<ul> <li>Depo Provera (medroxyprogesterone)/ Lunelle (estradiol cipionate/ medroxyprogesterone)</li> <li>FDA approved indication if given in a long-term care facility or in the members home:</li> <li>Females: Contraception, uterine bleeding, amenorrhea, endometrial cancer</li> <li>Males: Sexual aggression / Pedophilia – Only Depo-Provera will be approved</li> <li>Not approved for administration in the physician's office – these must be billed through medical.</li> <li>Implanon (etonogestrel)</li> <li>See PHYSICIAN ADMINISTERED DRUGS. Not a covered pharmacy benefit when implanted in the clinic or hospital outpatient center.</li> <li>Nexplanon (etonogestrel)</li> <li>See PHYSICIAN ADMINISTERED DRUGS. Not a covered pharmacy benefit when implanted in the clinic or hospital outpatient center.</li> </ul>	One year
HP ACTHAR (corticotropin)	<ul> <li>HP Acthar® may be approved for members that meet the following criteria:         <ul> <li>Member has a diagnosis of Infantile Spasms (West Syndrome) and meets <u>all</u> the criteria below:                 <ul> <li>Member is &lt; 2 years of age</li> <li>Member has electroencephalogram documenting diagnosis</li></ul></li></ul></li></ul>	4 week supply

COLORADO MEDICAID	PROGRAM	APPENDICES	
	Member does not have one         Scleroderma, ostec         herpes simplex, rec         failure, uncontrolle         porcine origin. AN	of the following concomitant diagnoses: opporosis, systemic fungal infections, ocular, cent surgery, history of peptic ulcer disease, heart ed hypertension, or sensitivity to proteins of <b>D</b> I based on the following FDA recommended	
	Diagnosis	Dose	
	Infantile Spasms under Age of 2 years	75 units/m <sup>2</sup> IM twice daily for two weeks; After two weeks, dose should be tapered according to the following schedule: 30 U/m <sup>2</sup> IM in the morning for 3 days; 15 units/m <sup>2</sup> IM in the morning for 3 days; 10 units/m <sup>2</sup> IM in the morning for 3 days; and 10 units/m <sup>2</sup> IM every other morning for 6 days (3 doses).	
	Acute Exacerbation of Multiple Sclerosis	80-120 units IM or SQ daily for 2-3 weeks	
	Quantity Limits: 4 week supply		
HUNTINGTON'S CHOREA / TARDIVE DYSKINESIA AGENTS	<ul> <li>been met:</li> <li>Member is 18 years and older way Tardive Dyskinesia AND <ul> <li>For chorea secondary to Hu and/or failed tetrabenazine, defined as a lack of efficacy contraindication to, or signifies</li> <li>For tardive dyskinesia a bas the 12 week AIMS does not authorization will no longer</li> </ul> </li> <li>Member does not have untreated suicide attempt AND</li> <li>Member has been informed of the Member does not have severe here</li> <li>Maximum dose 48mg/day, 120 the Xenazine® (tetrabenazine) may be met:</li> <li>Member is 18 years and older way AND</li> <li>Member does not have a history</li> </ul>	I depression, suicidal thoughts, or a history of ne risks of depression and suicidality AND epatic impairment ablets per month approved if all the following criteria have been ith chorea secondary to Huntington's Disease of suicide or untreated depression AND ne risks of depression and suicidality AND	One year unless AIMS follow-up required

#### COLORADO MEDICAID PROGRAM APPENDICES Ingrezza® (valbenazine) may be approved if all the following criteria have been met: Member is 18 years or older AND • Member has been diagnosed with tardive dyskinesia clinically AND • Has a baseline Abnormal Involuntary Movement Scale (AIMS) AND • If there is no improvement at 6 weeks of therapy per AIMS, the medication will • be discontinued Quantity limit of 60 capsules per 30 days HYDROXYCHLOROQUINE Effective 03/24/20: Prior authorization may be approved for FDA-labeled indication, Chronic conditions: dose, age, and role in therapy as outlined in product package labeling. One year Acute conditions: Duration of acute use **ILUMYA** Ilumya<sup>®</sup> prior authorization may be approved for members meeting all of the Initial: (tildrakizumab-asmn) 12 weeks following criteria: Medication is being administered in the member's home or in a long-term care Continued: facility by a healthcare professional AND One year Member is 18 years of age or older and has diagnosis of moderate to severe plaque psoriasis who are candidates for systemic therapy or phototherapy AND Member does not have guttate, erythrodermic, or pustular psoriasis AND Provider attests to: Baseline Provider Global Assessment (PGA) score for plaque • psoriasis severity of at least 3 (Scored 0-4, 4 being most severe) OR Baseline Psoriasis Area and Severity Index (PASI) score of 12 or greater AND Medication is being prescribed by or in conjunction with a rheumatologist, allergist, or dermatologist AND Member has tried and failed<sup>‡</sup> ALL preferred agents in the "Targeted Immune Modulators" PDL drug class that are FDA-labeled for use for the same prescribed indication AND Initial authorization will be for 12 weeks Continued authorization for 12 months will require prescriber attestation to PGA score reduction of 2 or more points OR PASI score reduction of 75% OR prescriber attestation to clinically meaningful improvement with Ilumya® regimen. Claims for medications administered in a clinic or medical office are billed through the Health First Colorado medical benefit. **JADENU and EXJADE** Jadenu (deferasirox) or Exjade (deferasirox) may be approved for members that One year (deferasirox) meet the following criteria: Must be prescribed in conjunction with a hematologist or oncologist AND ٠ Member's weight must be provided AND • Member has a diagnosis for chronic iron overload due to blood transfusion AND Member is 2 years of age or older AND • Member has consistently high serum ferritin levels > 1000 mcg/L (demonstrated by at least 2 values in the prior three months OR

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	<ul> <li>Member has a diagnosis for chronic iron overload due to non-transfusion dependent thalassemia syndromes AND</li> <li>Member is 10 years of age or older AND</li> <li>Member has liver iron levels &gt; 5 mg iron per gram of dry weight and serum ferritin levels &gt; 300 mcg/L document in the prior three months</li> <li>Members must also meet the following additional criteria for all Jadenu and Exjade approvals:         <ul> <li>Member does not have advanced malignancies and/or high-risk myelodysplastic syndromes AND</li> <li>Member has a creatinine clearance &gt; 40 ml/min AND</li> <li>Member has a platelet count &gt; 50 x 10<sup>9</sup>/L</li> </ul> </li> <li>Maximum Dosing:         <ul> <li>Maximum dose of Jadenu (deferasirox): 28mg/kg/day</li> <li>Maximum dose of Exjade (deferasirox): 40mg/kg/day</li> </ul> </li> </ul>	
JYNARQUE (tolvaptan)	<ul> <li>JYNARQUE (tolvaptan) may be approved if the following criteria are met:</li> <li>Member is an adult (≥ 18 years of age) and has been diagnosed with slow kidney function decline and at risk of rapidly progressing autosomal dominant polycystic kidney disease (ADPKD) AND</li> <li>Medication is being prescribed by a nephrologist AND</li> <li>Member does not have a history or sign/symptoms of significant liver impairment or injury (uncomplicated polycystic liver disease is not a contraindication for therapy) AND</li> <li>Member is not taking a strong Cytochrome 3A inhibitor (such as erythromycin, clarithromycin, telithromycin, itraconazole, ketoconazole, posaconazole, fluconazole, voriconazole, lopinavir/ritonavir, indinavir/ritonavir, ritonavir, conivaptan, delavirdine and milk thistle) AND</li> <li>Member is not taking a OATP1B1/B3 or OAT3 substrate (such as a statin, bosentan, glyburide, nateglinide, repaglinide, methotrexate or furosemide) AND</li> <li>Member is not taking a BCRP substrate (such as rosuvastatin, prazosin, pantoprazole, rosuvastatin, teriflunomide, or chlorthiazide) AND</li> <li>Member is not using desmopressin (dDAVP) AND</li> <li>If member is taking a moderate Cytochrome 3A inhibitor (such as erythromycin, fluconazole, or verapamil) JYNARQUE (tolvaptan) will be prescribed at a reduced dose AND</li> <li>Member has normal blood sodium concentrations, is able to sense or respond to thirst, and has a normal blood volume AND</li> <li>Member does not have urinary outflow obstruction or anuria</li> </ul>	One year
KALYDECO (ivacaftor)	<ul> <li>120mg per day</li> <li>Kalydeco (ivacaftor) may be approved if all of the following criteria are met: <ul> <li>Member has been diagnosed with cystic fibrosis AND</li> <li>Member is an adult or pediatric patient 4 months of age or older AND</li> <li>Documentation has been provided to indicate one of the following gene mutation: in the CFTR gene: G551D, G1244E, G1349D, G178R, G551S, S1251N, S1255P, S549N, R117H, S549R or another FDA approved gene mutation.* AND</li> </ul> </li> </ul>	One year

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KUVAN (sapropterin dihydrochloride)	<ul> <li>Documentation has been provided that baseline ALT and AST have been accessed and are within 2x normal limits (AST and ALT should be examined every 3 months for the first year and annually after that).</li> <li>* If the member's genotype is unknown, an FDA-cleared CF mutation test should be used to detect the presence of a CFTR mutation followed by verification with bidirectional sequencing when recommended by the mutation test instructions for use.</li> <li>Kalydeco® will only be approved at doses no more than 150 mg twice daily. Prior Authorizations need to be obtained yearly.</li> <li>Kalydeco® will not be approved for members who are concurrently receiving rifampin, rifabutin, phenobarbital, carbamazepine, phenytoin, or St. John's Wort.</li> <li>Kuvan@ may be approved if all the following criteria are met:</li> <li>Member is &gt; 1 month old AND</li> <li>Member has been diagnosed with hyperphenylalaninemia due to tetrahydrobiopterin responsive phenylketonuria AND</li> <li>Prescriber is a metabolic specialist AND</li> <li>Phenylalanine levels must be greater than 10 mg/dL for members 18 years and older AND</li> <li>Must be in conjunction with dietary restriction of phenylalanine</li> <li>Initial approval will be for 1 month. Authorization may be extended if: <ul> <li>Members on the 10mg/kg/day dose whose blood phenylalanine levels have not decreased from baseline after 1 month of treatment should increase to 20mg/kg/day. These members will be approved for another 1 month trial at the higher dose.</li> <li>Members on the 20mg/kg/day dose whose blood phenylalanine levels have not decreased from baseline after 1 month are considered non-responders, and treatment will be discontinued.</li> </ul> </li> </ul>	Initial approval one month
LAMPIT (nifurtimox)	intervals.         LAMPIT (nifurtimox) may be approved if the following criteria are met:	One year
	<ul> <li>Lampit (nifurtimox) is prescribed by or in conjunction with an infectious disease specialist, cardiologist or gastroenterologist AND</li> <li>The member's age falls between term newborn and &lt; 18 years of age AND</li> <li>The member's weight is provided and is at least 2.5 kg (5.5 pounds) AND</li> <li>The member has a diagnosis, documented and confirmed by blood smear, of Chagas disease (American Trypanosomiasis) caused by <i>Trypanosoma cruzi</i> AND</li> <li>For pediatric members 2 to 12 years of age, the member has trialed and failed treatment with benznidazole. Failure is defined as lack of efficacy, contraindication to therapy, allergy, intolerable side effects, or significant drug-drug interaction AND</li> <li>For female members of childbearing potential, a documented negative pregnancy test is obtained within 2 weeks of initiating therapy AND</li> <li>The member has received counseling (when appropriate) to not consume alcohol during treatment with Lampit (nifurtimox) AND</li> <li>The prescription meets the following recommended daily dosing:</li> </ul>	

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	Lampit (nifurtimov) D	osing in Pediatric Patients	
	Body weight group	Total daily dose	
	40 kg or greater	8 to 10 mg/kg	
	Less than 40 kg	10 to 20 mg/kg	
	Maximum Dosing:		
	300mg three times a day (900mg/day) for	r 60 days	
LIPIDS/AMINO ACIDS/PLASMA	<ul> <li>to be billed through the medical benefit. (may only receive approval if the medicat home by a home health agency/provider of (see "Physician Administered Drugs" see</li> <li>Prior authorization may be approved for I</li> <li>Eligard® (leuprolide): Palliative tree</li> <li>Fensolvi® (leuprolide acetate): Cente</li> <li>Lupaneta Pack® (leuprolide and note)</li> <li>Lupron® (leuprolide): Prostate cance (fibroids), precocious puberty. Luprobased on the following criteria: <ul> <li>The member has a diagnosis of a health professional with experie available, the mental health profand adolescent developmental p</li> <li>The member should have at leass testing for gender identity prior</li> <li>The prescribing provider has tra gonadotropin releasing hormone</li> <li>Lupron may not be started until puberty (confirmed by levels of no earlier than Tanner stages 2-3 tripling testicular size to 4-8 cc)</li> <li>Duration of treatment: Lupron w age for gender dysphoria.</li> </ul> </li> <li>Synarel® (nafarelin): Endometriosis</li> <li>Trelstar® (triptorelin): Palliative tree</li> <li>Triptodur® (triptorelin): Palliative tree</li> </ul>	FDA-labeled indications only. eatment of advanced prostate cancer ral precocious puberty orethindrone): Endometriosis eer, endometriosis, uterine leiomyomata on may be approved for gender dysphoria gender dysphoria which is made by a ment nce in treating gender dysphoria. Where essional should ideally have training in chi sychology AND t 6 months of counseling and psychometric to initiation of Lupron AND ining in puberty suppression using a agonist AND girls and boys exhibit physical changes of estradiol and testosterone, respectively) and 8 (bilateral breast budding or doubling to will be covered to a maximum of 16 years of the member's home or in a long-term care	al ld c f
PROTEINS	medical expense.	,	
LUCEMYRA (lofexidine)	<ul> <li>Lucemyra (lofexidine) may receive prior meeting all of the following criteria:</li> <li>Member is 18 years of age or old</li> <li>Lucemyra® is prescribed for mi facilitate abrupt opioid discontin</li> </ul>	der <b>AND</b> tigation of opioid withdrawal symptoms to	14 days

	ROGRAINI APPENDICES	
	<ul> <li>Member is not pregnant or nursing AND</li> <li>Member is not experiencing withdrawal symptoms from substances other than opioids AND</li> <li>Member is not currently taking monoamine oxidase inhibitors or allergic to imidazole drugs AND</li> <li>Member does not have an abnormal cardiovascular exam prior to treatment:         <ul> <li>Clinically significant abnormal ECG (e.g., second or third degree heart block, uncontrolled arrhythmia, or QTc interval &gt; 450 msec for males, and &gt; 470 msec for females)</li> <li>Heart rate less than 45 bpm or symptomatic bradycardia</li> <li>Systolic blood pressure &lt; 90 mm Hg or symptomatic hypotension (diastolic blood pressure &lt; 60 mm Hg)</li> <li>Blood pressure &gt; 160/100 mm Hg</li> <li>Prior history of myocardial infarction AND</li> </ul> </li> <li>Member has two-day trial and failed clonidine IR for opioid withdrawal symptoms. Failure is defined as lack of efficacy, allergy, intolerable side effects, or significant drug-drug interaction.</li> </ul>	
	Approval for Lucemyra (lofexidine) will be 14 days	<u> </u>
LUMIZYME (alglucosidase alfa)	<ul> <li>Lumizyme (alglucosidase alfa) may be approved for members meeting all of the following criteria:</li> <li>Medication is being administered in the member's home or in a long-term care facility by a healthcare professional AND</li> <li>Member has diagnosis of Pompe disease (acid α-glucosidase [GAA] deficiency).</li> </ul>	One year
	Maximum dose: Lumizyme 20mg/kg every 2 weeks (IV Infusion)	
MAKENA (hydroxyprogesterone caproate)	<ul> <li>Makena® may be approved for members that meet the following criteria:</li> <li>The drug is being administered in the home or in long-term care setting</li> <li>Member has a Singleton pregnancy and a history of singleton spontaneous preterm birth</li> <li>Therapy is being initiated between 16 weeks gestation and 20 weeks 6 days gestation and continued through 36 weeks 6 days gestation or delivery (whichever occurs first)</li> <li>Dose is administered by a healthcare professional.</li> </ul> Maximum Dosing: Makena vial: 250mg IM once weekly Makena autoinjector: 275mg SubQ once weekly	See criteria
MALARIA PROPHYLAXIS EXCEEDING THIRTY DAYS	<ul> <li>Prior authorization is required for claims exceeding a 30-day supply for medications used for malaria prophylaxis (e.g. atovaquone/proguanil, chloroquine, doxycycline, mefloquine, primaquine, tafenoquine) and may be approved for members meeting the following: <ul> <li>Prescriber verification that the member is traveling to a malaria endemic area for a period of time that requires duration of therapy exceeding thirty days.</li> <li>Prescriber verification of member's duration of stay in the malaria endemic area and the total days needed for the malaria prophylaxis medication regimen.</li> </ul> </li> <li>Note: The Centers for Disease Control and Prevention recommendations for malaria prophylaxis therapy based on country of travel are available at www.cdc.gov</li> </ul>	See criteria

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MIFEPRISTONE and MISOPROSTOL	<b>Mifeprex</b> <sup>®</sup> (mifepristone) is excluded from coverage under the pharmacy benefit.	One year
	<b>Korlym</b> <sup>®</sup> (mifepristone) – Prior authorization may be approved for members meeting the following:	
	<ul> <li>Mifepristone is not being prescribed for use related to termination of pregnancy AND</li> </ul>	
	• Mifepristone is being prescribed for use for hyperglycemia secondary to hypercortisolism in adult patients with Cushing's Syndrome who have type 2 diabetes or glucose intolerance and have failed or are not candidates for surgery.	
	<b>Cytotec</b> <sup>®</sup> (misoprostol) – ( <i>Effective 07/18/19</i> ) Prior authorization may be approved for members meeting the following:	
	<ul> <li>Misoprostol is not being prescribed for use related to termination of pregnancy AND</li> </ul>	
	• Misoprostol is being prescribed for use as prophylaxis for reducing risk of NSAID-induced gastric ulcers in patients at high risk of complications from gastric ulceration OR is being prescribed for use for off-label indications supported by clinical compendia and peer-reviewed medical literature.	
	<i>Note: See PDL for coverage information for misoprostol/NSAID combination products.</i>	
MIGERGOT (ergotamine/caffeine)	<ul> <li>Migergot® (ergotamine/caffeine) suppository may be approved for members meeting the following criteria:</li> <li>Migergot (ergotamine/caffeine) suppository is being prescribed to prevent or treat vascular headache (migraine, migraine variants or so-called "histaminic cephalalgia") AND</li> <li>Member has a negative pregnancy test within 30 days of receipt of Ergomar AND</li> <li>Member is not taking a potent CYP 3A4 inhibitor (ritonavir, nelfinavir, indinavir, erythromycin, clarithromycin and troleandomycin) AND</li> <li>Member has adequate trial and/or failure of 2 triptan agents (see PDL class) AND</li> <li>Member has adequate trial and/or failure of dihydroergotamine vial. Failure is defined as lack of efficacy with 10 day trial, allergy, intolerable side effects or significant drug-drug interactions.</li> <li>Maximum quantity: 20 suppositories per 28 days</li> </ul>	One year
MOXATAG (amoxicillin)	A prior authorization will only be approved if a member has an allergic/intolerance to	One year
MULPLETA	inactive ingredients in immediate release amoxicillin. <b>Mulpleta</b> <sup>®</sup> (lusutrombopag) prior authorization may be approved for members	One year
(lusutrombopag)	meeting the following criteria:	che your
-	• Member is 18 years of age or older AND	
	• Member has a confirmed diagnosis of thrombocytopenia with chronic liver	
	disease who is scheduled to undergo an elective procedure AND	
	• Member has trialed and failed both dexamethasone and methylprednisolone (Failure is defined as a lack of efficacy, allergy, intolerable side effects, or significant drug-drug interactions) AND	

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	<ul> <li>Mulpleta is being prescribed by or in consultation with a hematologist, hepatologist, or gastroenterologist AND</li> <li>Member has a baseline platelet count no more than 2 days before procedure. AND</li> <li>Mulpleta (lusutrombopag) will not be administered with a thrombopoietic agent or spleen tyrosine kinase inhibitor (such as Promacta (eltrombopag), Nplate (romiplostim), or Tavalisse (fotamatinib)</li> </ul>	
	Quantity limit: 7 day supply per procedure	
MYALEPT (metreleptin)	<ul> <li>Myalept<sup>®</sup> may be approved if all of the following criteria are met:</li> <li>Prescriber is an endocrinologist who is enrolled in the Myalept REMS program AND</li> <li>Member has a diagnosis of congenital or acquired generalized lipodystrophy AND</li> <li>Member does not have HIV-related lipodystrophy AND</li> <li>Member has a diagnosis of leptin deficiency AND</li> <li>Member has been diagnosed with poorly controlled diabetes (HgA1c &gt; 7) and/or hypertriglyceridemia (&gt; 500 mg/dl) AND</li> <li>Member has tried and failed two standard therapies for diabetes and/or hypertriglyceridemia</li> </ul>	Six Months
NAGLAZYME (galsulfase)	<ul> <li>Naglazyme<sup>®</sup> (galsulfase) may be approved for members meeting the following criteria:</li> <li>Naglazyme (galsulfase) is being administered in a long-term care facility or in a member's home by a healthcare professional AND</li> <li>Member is 5 years of age or older AND</li> <li>Member has a confirmed diagnosis of Mucopolysaccharidosis, Type VI confirmed by the following: <ul> <li>Detection of pathogenic mutations in the ARSB gene by molecular genetic testing OR</li> <li>Arylsulfatase B (ASB) enzyme activity of &lt;10% of the lower limit of normal in cultured fibroblasts or isolated leukocytes AND</li> <li>Member has an elevated urinary glycosaminoglycan (uGAG) level above the upper limit of normal as defined by the reference laboratory AND</li> <li>Member has a documented baseline 12-minute walk test (12-MWT), 3-minute stair climb test, and/or pulmonary function tests (such as FEV1) AND</li> <li>Member has a documented baseline value for uGAG AND</li> <li>Naglazyme (galsulfase) is being prescribed by or in consultation with a provider who specializes in inherited metabolic disorders</li> </ul> </li> <li>Reauthorization Criteria: <ul> <li>After one year, member may receive approval to continue therapy if meeting the following:         <ul> <li>Has documented reduction in uGAG levels AND</li> </ul> </li> </ul></li></ul>	One year

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	Max dose: 1 mg/kg as a 4-hour infusion weekly	
NALOXONE and NALTREXONE	Narcan <sup>®</sup> (naloxone) intranasal <u>does not</u> require prior authorization.	
	<b>Revia</b> <sup>®</sup> (naltrexone) tablet <u>does not</u> require prior authorization.	
	Naloxone vial/prefilled syringe:	
	• <u>does not</u> require prior authorization.	
	• The atomizer device for use with naloxone can be obtained by the pharmacy billing as a DME claim code A4210. The unit limit is 1 atomizer per vial/syringe dispensed up to a total of 15 per year. A prior authorization is not required.	
	<b>Vivitrol</b> <sup>®</sup> (naltrexone ER) injection:	
	• Prior authorization for claims submitted under the pharmacy benefit may be approved when Vivitrol is administered by a healthcare professional in the member's home or in a long-term care facility. All other Vivitrol claims must be billed through the medical benefit.	
	• Effective 01/01/2019, pharmacies that have entered into a collaborative practice agreement with one or more physicians for administration of Vivitrol may receive reimbursement for enrolled pharmacists to administer Vivitrol with appropriate claim submission through the Health First Colorado medical benefit (claims for pharmacist administration of Vivitrol are not covered under the pharmacy benefit). Additional information regarding pharmacist enrollment and medical claims billing can be found at <a href="https://www.colorado.gov/hcpf/otc-">https://www.colorado.gov/hcpf/otc-</a>	
	immunizations .	
	<b>Evzio</b> <sup>®</sup> (naloxone) autoinjector – Product is not Medicaid rebate eligible per current status in Medicaid Drug Rebate Program (MDRP); product excluded	
	*For buprenorphine/naloxone products, see "Buprenorphine-containing Products" section	
NAYZILAM (midazolam)	<b>Nayzilam<sup>®</sup></b> (midazolam) may be approved for members meeting the following criteria:	One Year
	• Member is 12 years of age or older AND	
	• Nayzilam is being prescribed for the acute treatment of intermittent,	
	stereotypic episodes of frequent seizure activity (i.e., seizure clusters,	
	acute repetitive seizures) that are distinct from a patient's usual seizure	
	<ul> <li>pattern and medical records are provided supporting this diagnosis AND</li> <li>Member is stable on regimen of antiepileptic medications AND</li> </ul>	
	<ul> <li>Medication is being prescribed by or in conjunction with the same</li> </ul>	
	provider/provider team who manages the member's anti-epileptic	
	regimen AND	
	• Member is educated on appropriate identification of seizure cluster and Nayzilam (midazolam) administration not exceeding 2 doses per seizure cluster.	
	Maximum dose: 4 nasal spray units per year unless used / damaged / lost	
	Members are limited to one prior authorization approval on file for Valtoco (diazepam) and Nayzilam (midazolam).	

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	Grandfathering: If member is currently receiving Nayzilam (midazolam) intranasal, they may receive prior authorization approval to continue.	
NEWLY APPROVED PRODUCTS AND CHANGE IN PRODUCT PRIOR AUTHORIZATION STATUS	Newly marketed or approved products that fall within a PDL drug class will be subject to non-preferred prior authorization criteria for the drug class and will be included as part of the next regularly scheduled P&T Committee and DUR Board reviews for that class. Newly marketed or approved products that fall within a drug category on appendix P (such as "Blood Products" or "Atypical Antipsychotic Injectables") will be subject to prior authorization criteria listed for medications in that drug category on Appendix P.	
	For change in prior authorization status for a product that is not included in a PDL drug class or on Appendix P, notice will be given regarding DUR Board review of prior authorization criteria for the product as part of the posted DUR Board meeting agenda located at <u>https://www.colorado.gov/pacific/hcpf/drug-utilization-review-board</u> and posted at least 30 days prior to the DUR Board meeting during which the product is scheduled to be reviewed. Until such time that DUR Board review is conducted, products may receive prior authorization approval based on FDA-labeled indication, dose, age, and role in therapy as outlined in product package labeling. IV formulations or products where labeled use indicates that the medication should be administered by a healthcare professional will also be subject to meeting criteria for	
NORTHERA (droxidopa)	<ul> <li>physician administered drugs (see "Physician Administered Drugs" section).</li> <li>Northera® (droxidopa) will be approved if all the following is met:</li> <li>Member has a diagnosis of symptomatic neurogenic orthostatic hypotension</li> </ul>	3 months
	<ul> <li>(NOH) as defined by one of the following when an upright position is assumed or when using a head-up tilt table testing at an angle of at least 60 degrees.</li> <li>At least a 20 mmHg fall is systolic pressure</li> <li>At least a 10 mmHg fall in diastolic pressure</li> <li>At least a 10 mmHg fall in diastolic pressure</li> <li>At least a 10 mmHg fall in diastolic pressure</li> <li>At least a 10 mmHg fall in diastolic pressure</li> <li>At least a 10 mmHg fall in diastolic pressure</li> <li>Primary autonomic failure (e.g., Parkinson's disease, multiple system atrophy, and pure autonomic failure</li> <li>Dopamine beta-hydroxylase deficiency</li> <li>Non-diabetic autonomic neuropathy</li> <li>AND</li> <li>Member does not have orthostatic hypotension due to other causes (e.g., heart failure, fluid restriction, malignanacy) AND</li> <li>Members has tried at least three of the following non-pharmacological interventions:</li> <li>Discontinuation of drugs which can cause orthostatic hypotension [e.g., diuretics, antihypertensive medications (primarily sympathetic blockers), anti-anginal drugs (nitrates, excluding SL symptom treatment formulations), alpha-adrenergic antagonists, and antidepressants]</li> <li>Raising the head of the bed 10 to 20 degrees</li> <li>Compression stockings</li> <li>Increased salt and water intake, if appropriate</li> <li>Avoiding precipitating factors (e.g., overexertion in hot weather, arising too quickly from supine to sitting or standing)</li> <li>AND</li> <li>Northera (droxidopa) is being prescribed by either a cardiologist, neurologist, or nephrologist AND</li> <li>Member has failed a 30 day trail, has a contraindication, or intolerance to both Florinef (fludrocortisone) and ProAmatine (midodrine).</li> </ul>	

#### COLORADO MEDICAID PROGRAM APPENDICES A prior authorization will only be approved as a pharmacy benefit when the NUCALA (mepolizumab) One year medication is administered in a long-term care facility. Medications administered in a physician's office must be billed as a medical expense. Because this medication has a FDA-labeled boxed warning requiring the administration under the supervision of a physician, a prior authorization will not be approved if administered in a member's home. Nuedexta (dextromethorphan/quinidine) may be approved for members who meet the Initial **NUEDEXTA** Approval: (dextromethorphan following criteria: 3 months /quinidine) Nuedexta is being prescribed for diagnosis of pseudobulbar affect caused by an underlying neurologic condition (such as MS, ALS, or other underlying Continuation neurologic condition) AND Approval: One year Member has a Center for Neurologic Study-Lability Scale (CNS-LS) score of 13 . or higher AND Member has at least 10 episodes of inappropriate laughing or crying per day . before therapy AND Member has a baseline electrocardiogram (ECG) with no significant abnormalities and no history of QT prolongation syndrome AND Nuedexta is prescribed by a neurologist or in conjunction with a neurologist AND Member has trailed and failed one tricyclic antidepressant and one selective serotonin reuptake inhibitor within the past year (failure is defined as lack of efficacy, allergy, intolerable side effects, contraindication to therapy, or significant drug-drug interactions) Initial approval will be given for 3 months and continued approval for one year may be given if member has 50% reduction in daily episodes at 3 months of therapy Nuedexta® Max Dose: 2 capsules (dextromethorphan 20mg/quinidine 10mg) per day given every 12 hours Renewal: members currently stabilized on this medication may continue to receive it with a documented diagnosis of pseudobulbar affect and evidence of efficacy (documentation of decrease in pseudobulbar episodes by 50% from baseline) **OCREVUS** (ocrelizumab) One year Ocrevus® (ocrelizumab) may be approved if the following criteria are met: Ocrevus is being administered in a LTCF or in the member's home AND If prescribed for Relapsing Forms of Multiple Sclerosis (MS) Member is 18 years of age or older AND 0 Member has a relapsing form of multiple sclerosis AND 0 Member has experienced one relapse within the prior year or two 0 relapses within the prior two years AND Member has trial and failure of three of the following agents: 0 Avonex (interferon beta-1a), Rebif (interferon beta 1-a), Betaseron/Extavia (interferon beta-1b), Plegridy (peginterferon beta1a), Copaxone/Glatopa (glatiramer acetate), Aubagio (teriflunomide tablets), Gilenya (fingolimod capsules), Tecfidera (dimethyl fumarate delayed-release capsules), Tysabri (Natalizumab) or Lemtrada (alemtuzumab). Failure will be defined as intolerable side effects, drug-drug interaction, or lack of efficacy. Lack of efficacy will be defined as one of the following: One of the following on MRI: presence of any new spinal lesions, 0 cerebellar or brain stem lesions, or change in brain atrophy On clinical exam, signs and symptoms consistent with functional 0 limitations that last one month or longer AND Ocrevus is prescribed by a neurologist or is prescribed in conjunction with a

neurologist AND

	PROGRAMI APPENDICES	
	<ul> <li><u>If prescribed for Primary Progressive Multiple Sclerosis</u> <ul> <li>Member is 18 years of age or older AND</li> <li>Member is not concomitantly taking: Avonex (interferon beta-1a), Rebif (interferon beta 1-a), Betaseron/Extavia (interferon beta-1b), Plegridy (peginterferon beta1a), Copaxone/Glatopa (glatiramer acetate), Aubagio (teriflunomide tablets), Gilenya (fingolimod capsules), Tecfidera (dimethyl fumarate delayed-release capsules), Tysabri (Natalizumab) or Lemtrada (alemtuzumab) AND</li> </ul> </li> <li>Member does not have active hepatitis B infection AND</li> <li>Ocrevus is prescribed by a neurologist or is prescribed in conjunction with a neurologist</li> <li>Maximum maintenance dose: 600mg every 6 months</li> </ul>	
OFEV (nintedanib)	<ul> <li>Ofev (nintedanib) may be approved if all of the following criteria are met:</li> <li>Member has been diagnosed with idiopathic pulmonary fibrosis, chronic fibrosing interstitial lung disease with a progressive phenotype, or systemic sclerosis-associated interstitial lung disease (SSC-ILD) AND</li> <li>Is being prescribed by or in conjunction with a pulmonologist AND</li> <li>Member is 18 years or older AND</li> <li>Member has baseline ALT, AST, and bilirubin prior to starting therapy AND</li> <li>Member does not have moderate (Child Pugh B) or severe (Child Pugh C) hepatic impairment AND</li> <li>Female members of reproductive potential must have been counseled regarding risk to the fetus and to avoid becoming pregnant while receiving treatment with Ofev and to use adequate contraception during treatment and at least 3 months after the last dose of Ofev AND</li> <li>Member is not taking a P-gp or CYP3A4 inducer (e.g, rifampin, carbamazepine, phenytoin, St. John's Wort)</li> </ul>	One year
ORILISSA (elagolix)	<ul> <li>Orilissa® (elagolix) may be approved for members meeting the following criteria:</li> <li>Member is a premenopausal woman 18-49 years of age AND</li> <li>Orilissa® is not being prescribed for dyspareunia or any other sexual function related indication AND</li> <li>Member has a definitive diagnosis of endometriosis as noted by surgical histology of lesions AND</li> <li>Member has failed a 6-month trial of contraceptive agents (progestins, combined contraceptives, medroxyprogesterone acetate, levonorgestrel IUD). Failure is defined as lack of efficacy, allergy, intolerable side effects, significant drug-drug interaction, or contraindication to therapy AND</li> <li>Member has failed a 1 month trial of NSAIDs. Failure is defined as lack of efficacy, allergy, intolerable side effects, significant drug-drug interaction, or contraindication to therapy AND</li> <li>Member has failed a 3 month trial with a GnRH agonist (such as leuprolide). Failure is defined as lack of efficacy, allergy, intolerable side effects, significant drug-drug interaction, or contraindication to therapy AND</li> <li>Member has failed a 3 month trial with a GnRH agonist (such as leuprolide). Failure is defined as lack of efficacy, allergy, intolerable side effects, significant drug-drug interaction, or contraindication to therapy AND</li> <li>Member is not pregnant, breast feeding, planning a pregnancy within the next 24 months, or less than 6 months post-partum, post-abortion, or post-pregnancy AND</li> </ul>	One year 6 months for moderate hepatic impairment (Child Pugh Class B)

#### APPENDICES

<ul> <li>criteria has been met:</li> <li>Member must have diagnosis of cystic fibrosis with genetic testing performed to confirm that member is homozygous for the F508del mutation in the CFTR gene AND</li> <li>Member is 6 years of age or older AND</li> <li>Member has 5 times upper limit of normal (ULN) AST/ALT or &lt; 3 times ULN AST/ALT if concurrently has &gt; 2 times ULN bilirubin at time of initiation AND</li> <li>Member has serum transaminase and bilirubin measured before initiation and every 3 months during the first year of treatment</li> <li>ORIAHNN (elagolix, estradiol, norethindrone acetate) prior authorization may be approved for members meeting the following criteria:         <ul> <li>Member has a confirmed diagnosis of heavy menstrual bleeding associated with uterine leiomyomas (fibroids) AND</li> <li>Member has tried and failed treatment with an estrogen-progestin contraceptive (oral tablets, vaginal ring, transdermal patch) OR a progestin-releasing intrauterine device (UD). Failure is defined as lack of efficacy, allergy, intolerable side effects, significant drug-drug interaction, or contraindication to therapy AND</li> <li>The medication is prescribed by or in consultation with an obstetrician/gynecologist AND</li> <li>Women over 35 years of age who smoke OR</li> <li>Women with a past or current history of the following:                 <ul> <li>DVT, PE, or cerebrovascular disease, (such as cerebrovascular disease, coronary artery disease, peripheral vascular disease, or orangt afformation) or or contraindiction in the age of the heart (such as subacute bacterial endocarditis with valvular disease, or artial fibrillation) OR</li></ul></li></ul></li></ul>		PROGRAM APPENDICES	
<ul> <li>criteria has been met:</li> <li>Member must have diagnosis of cystic fibrosis with genetic testing performed to confirm that member is homozygous for the F508del mutation in the CFTR gene AND</li> <li>Member is 6 years of age or older AND</li> <li>Member has 5 times upper limit of normal (ULN) AST/ALT or &lt; 3 times ULN AST/ALT if concurrently has &gt; 2 times ULN bilirubin at time of initiation AND</li> <li>Member has serum transaminase and bilirubin measured before initiation and every 3 months during the first year of treatment</li> <li>ORIAHNN (elagolix, estradiol, norethindrone acetate) prior authorization may be approved for members meeting the following criteria:         <ul> <li>Member has a confirmed diagnosis of heavy menstrual bleeding associated with uterine leiomyomas (fibroids) AND</li> <li>Member has tried and failed treatment with an estrogen-progestin contraceptive (oral tablets, vaginal ring, transdermal patch) OR a progestin-releasing intrauterine device (UD). Failure is defined as lack of efficacy, allergy, intolerable side effects, significant drug-drug interaction, or contraindication to therapy AND</li> <li>The medication is prescribed by or in consultation with an obstetrician/gynecologist AND</li> <li>Women over 35 years of age who smoke OR</li> <li>Women with a past or current history of the following:                 <ul> <li>DVT, PE, or cerebrovascular disease, (such as cerebrovascular disease, coronary artery disease, peripheral vascular disease, or orangt afformation) or or contraindiction in the age of the heart (such as subacute bacterial endocarditis with valvular disease, or artial fibrillation) OR</li></ul></li></ul></li></ul>		<ul> <li>should be used during therapy and for at least 1 week following discontinuation AND</li> <li>Member does not have osteoporosis or severe hepatic impairment (Child-Pugh Class C) AND</li> <li>Member is not concomitantly taking a OATP 1B1 inhibitor (such as gemfibrozil, cyclosporine, ritonavir, rifampin).</li> <li>Orilissa® Maximum Dose: 150mg tablet daily, or 200mg tablet twice daily</li> <li>Orilissa® limited to a maximum treatment duration of 6 months for members with</li> </ul>	
<b>ORIAHNN</b> (elagolix, estradiol, norethindrone acetate) prior authorization may be approved for members meeting the following criteria: <ul> <li>Member is a woman 18 years of age or older <b>AND</b></li> <li>Member has a confirmed diagnosis of heavy menstrual bleeding associated with uterine leiomyomas (fibroids) <b>AND</b></li> <li>Member has tried and failed treatment with an estrogen-progestin contraceptive (oral tablets, vaginal ring, transdermal patch) OR a progestin-releasing intrauterine device (IUD). Failure is defined as lack of efficacy, allergy, intolerable side effects, significant drug-drug interaction, or contraindication to therapy <b>AND</b></li> <li>The medication is prescribed by or in consultation with an obstetrician/gynecologist <b>AND</b></li> <li>Member does not have a high risk of arterial, venous thrombotic, or thromboembolic disorder, including:                 <ul> <li>Women over 35 years of age who smoke <b>OR</b></li> <li>Women with a past or current history of the following:</li> <li>DVT, PE, or cerebrovascular disease (such as cerebrovascular disease) <b>OR</b></li></ul></li></ul>	ORKAMBI lumacaftor/ivacaftor)	<ul> <li>criteria has been met:</li> <li>Member must have diagnosis of cystic fibrosis with genetic testing performed to confirm that member is homozygous for the F508del mutation in the CFTR gene AND</li> <li>Member is 6 years of age or older AND</li> <li>Member is being treated by a pulmonologist AND</li> <li>Member has &lt; 5 times upper limit of normal (ULN) AST/ALT or &lt; 3 times ULN AST/ALT if concurrently has &gt; 2 times ULN bilirubin at time of initiation AND</li> <li>Member has serum transaminase and bilirubin measured before initiation and</li> </ul>	One year
• Member is not pregnant <b>AND</b>	ORIAHNN (elagolix, estradiol, norethindrone acetate)	<ul> <li>every 3 months during the first year of treatment</li> <li>ORIAHNN (elagolix, estradiol, norethindrone acetate) prior authorization may be approved for members meeting the following criteria:         <ul> <li>Member is a woman 18 years of age or older AND</li> <li>Member has a confirmed diagnosis of heavy menstrual bleeding associated with uterine leiomyomas (fibroids) AND</li> <li>Member has tried and failed treatment with an estrogen-progestin contraceptive (oral tablets, vaginal ring, transdermal patch) OR a progestin-releasing intrauterine device (IUD). Failure is defined as lack of efficacy, allergy, intolerable side effects, significant drug-drug interaction, or contraindication to therapy AND</li> <li>The medication is prescribed by or in consultation with an obstetrician/gynecologist AND</li> <li>Momen over 35 years of age who smoke OR                 <ul> <li>Women over 35 years of age who smoke OR</li> <li>Women with a past or current history of the following:</li></ul></li></ul></li></ul>	One year
• 1		• 1	<ul> <li>Uncontrolled hypertension OR</li> <li>Headaches with focal neurological symptoms OR migraine headaches with aura if over age 35</li> </ul>

COLORADO MEDICAID P	ROGRAM APPENDICES	
	<ul> <li>Member does not have current or history of breast cancer or other hormonally-sensitive malignancies AND</li> <li>Member does not have known liver impairment or disease AND</li> <li>Member is not concomitantly taking not an OATP 1B1 inhibitor (such as gemfibrozil, ritonavir, rifampin, cyclosporine) AND</li> <li>Member has been counseled that that Oriahnn does not prevent pregnancy AND</li> <li>Member has been instructed that only non-hormonal contraceptives should be used during Oriahnn therapy and for at least 1 week following discontinuation AND</li> <li>Prescriber acknowledges that assessment of bone mineral density (BMD) by dual-energy X-ray absorptiometry (DXA) is recommended at baseline and periodically thereafter, and discontinuation of Oriahnn should be considered if the risk associated with bone loss exceeds the potential benefit of treatment.</li> </ul>	
	Reauthorization: Members with current one-year prior authorization approval on file may receive additional one-year prior authorization approval to continue therapy. Total duration for prior authorization approvals is limited to 2 years (or two one-year approvals).	
OTC PRODUCTS*	Maximum dose: 2 capsules daily (AM and PM daily doses supplied in blister pack)         The following OTC products do not require a prior authorization for coverage: <ul> <li>Aspirin</li> <li>Oral emergency contraceptive products</li> <li>Polyethylene glycol powder laxatives</li> <li>Docusate (oral) <i>Effective 03/01/19</i></li> <li>Bisocodyl (oral and suppository) <i>Effective 03/01/19</i></li> <li>Children's liquid and chewable acetaminophen for ages 2-11 years</li> <li>Children's liquid and chewable ibuprofen for ages 6 months – 11 years</li> <li>Children's dextromethorphan suspension for ages 4-11 years</li> <li>Nicotine replacement therapies (OTC patch, gum, and lozenge)</li> </ul> <li>The following OTC products may be covered with a prior authorization:</li> <li>L-methylfolate may be approved for members with depression who are currently taking an antidepressant and are partial or non-responders</li> <li>Nicomide may be approved for the treatment of acne</li> <li>Crapherry tablets may be approved for urinary tract infections</li>	One year
	<ul> <li>Cranberry tablets may be approved for urinary tract infections</li> <li>Cough and Cold Products may be approved for members with a diagnosis of a chronic respiratory condition for which these medications may be prescribed or based on medical necessity supported by clinical practice recommendations</li> <li>Guaifenesin 600mg LA may be approved for members having an abnormal amount of sputum</li> <li>Bisacodyl enema may be approved following adequate trial and/or failure with a bisocodyl oral formulation and bisocodyl suppository (Failure is defined as lack of efficacy with 10 day trial, allergy, intolerable side effects, or significant drug-drug interactions). <i>Effective 03/01/19</i></li> <li>Docusate enema may be approved following adequate trial and with a docusate oral formulation (Failure is defined as lack of efficacy with 10 day trial, allergy, intolerable side effects, or significant drug-drug interactions). <i>Effective 03/01/19</i></li> </ul>	

COLORADO MEDICAID P	PROGRAM APPENDICES	
	<ul> <li>Ferrous sulfate and ferrous gluconate may be approved with diagnosis iron deficient anemia OR iron deficiency verified by low serum ferritin. <i>Effective 03/01/19</i></li> <li>Members with erythema bullosum (EB) may be approved to receive OTC medications (any Medicaid rebate-eligible OTC medications)</li> <li>Other OTC product coverage information: <ul> <li>Diabetic needles and supplies are covered under the DME benefit</li> <li>Broncho saline: <i>See Sodium Chloride section</i></li> <li>Fluoride supplements: <i>See Fluoride Products section</i></li> <li>OTC Proton Pump Inhibitors: <i>See PDL</i></li> <li>OTC Combination Antihistamine/Decongestant Products: <i>See PDL</i></li> <li>Long Term Care Facilities (LTCFs): Various OTC drugs and supplies for LTCF residents shall be furnished by the facility, within the per diem rate, at no charge to the resident pursuant to 10 CCR 2505-10 Skilled Nursing Facility: 8.440 NURSING FACILITY BENEFITS. These OTC drugs and supplies, known as products on a "floor stock list", are not covered or eligible for prior authorization under the pharmacy benefit for LTCF members.</li> </ul> </li> </ul>	
	* Coverage criteria outlined in this section apply to prescriptions written by non-pharmacist prescribers. For coverage relating to pharmacist prescribers please see "Pharmacist	
	Prescribers. For coverage relating to pharmacist prescribers please see "Pharmacist Prescriptions" section.	
OXANDRIN (oxandrolone)	<ul> <li>Oxandrin<sup>®</sup> (oxandrolone) may be approved if meeting all of the following criteria:</li> <li>Medication is being prescribed for one of the following indications: <ul> <li>As adjunctive therapy to promote weight gain after weight loss following extensive surgery, chronic infections, severe trauma, and without definite pathophysiologic reasons to fail to gain or maintain normal weight</li> <li>To offset the protein catabolism associated with prolonged administration of corticosteroids</li> <li>For the relief of bone pain frequently accompanying osteoporosis AND</li> </ul> </li> <li>Member does not have any of the following medical conditions: <ul> <li>Hypercalcemia</li> <li>Known or suspected carcinoma of the prostate or the male breast</li> <li>Carcinoma of the breast in females with hypercalcemia</li> <li>Nephrosis, the nephrotic phase of nephritis AND</li> </ul> </li> <li>If member is female, has had a negative pregnancy test within the past month AND</li> <li>Medication is being prescribed by or in consultation with an endocrinologist.</li> </ul>	One Year
OXBRYTA (voxelotor)	<ul> <li>OXBRYTA (voxelotor) prior authorization may be approved for members meeting the following criteria:</li> <li>Member is ≥ 12 years of age AND</li> <li>Member has a confirmed diagnosis of sickle cell disease AND</li> <li>Member has a hemoglobin ≥ 5.5 g/dL AND</li> <li>OXBRYTA is prescribed by or in consultation with hematologist/oncologist</li> </ul>	Initial: 6 months Continued: One year
	<ul> <li>or sickle cell disease specialist AND</li> <li>Prior to initiation of therapy, member had at least two episodes of sickle cell</li> </ul>	
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	ROOMAIN AFFEINDICES	
OXERVATE (cenegermin-bkbj)	<ul> <li>related pain crises in the past 12 months AND</li> <li>Member has trialed and failed a six-month trial of hydroxyurea (intolerance or contraindication) or is continuing concomitant hydroxyurea (herapy following a six-month trial. Failure is defined a lack of efficacy, allergy, intolerable side effects, significant drug-drug interaction, or contraindication to therapy AND</li> <li>Member is not receiving chronic transfusion therapy OR</li> <li>Member has severe renal disease (GFR &lt;30 mL/min)</li> <li>Initial approval: 6 months</li> <li>Reauthorization: Member may receive reauthorization approval for 1 year if meeting the following:</li> <li>Member has a reduction in vasoocclusive events and/or increased hemoglobin response rate defined as a hemoglobin increase of more than 1 g/dL.</li> <li>Maximum dose: 1,500 mg per day (2,500 mg per day may be approved for members taking concomitant strong or moderate CYP3A4 inducers (such as carbamazepine, oxarbazepine, phenytoin, phenobarbital, rifaximin, rifampin or dexamethasone-containing products).</li> <li>OXERVATE (conegermin-bkbi) prior authorization may be approved for members meeting the following criteria:</li> <li>Member is 2 years of age or older AND</li> <li>Member 's PED and/or corneal ulcer have been present for at least two weeks AND</li> <li>Member has trialed and failed one of the following conventional non-surgical treatments; preservative-free lubricant eye drops or ointment, therapeutic soft contact lenses, or topical autologous serun application. Failure is defined as also tift (24 cm using the Cachet-Bonnet esthsiometer) within the area of the PED or ulcer and outside the area of defect in at least one corneal sensitivity (≤4 cm using the Cachet-Bonnet esthsiometer) within the area of the PED or ulcer and outside the area of defect in at least one corneal sensitivity (≤4 cm using the Cachet-Bonnet esthsiometer) within the area of the PED or ulcer and outside the area of defect in at least one cor</li></ul>	8 weeks
	<ul> <li>Any ocular surgery in the affected eye within the past 90 days that has not been determined to be the cause of NK</li> <li>Corneal perforation, ulceration involving the posterior third of the corneal stroma, or corneal melting</li> </ul>	
	<ul> <li>has not been determined to be the cause of NK</li> <li>Corneal perforation, ulceration involving the posterior third of the</li> </ul>	
OXSORALEN	<ul> <li>has not been determined to be the cause of NK</li> <li>Corneal perforation, ulceration involving the posterior third of the corneal stroma, or corneal melting</li> </ul>	One year

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	ROGRAM APPENDICES	
PALFORZIA (arachis hypogaea allergen powder-dnfp)	<ul> <li>PALFORZIA (arachis hypogaea allergen powder-dnfp) prior authorization may be approved for members meeting the following criteria:</li> <li>Member is 4 -17 years of age at initiation of therapy AND</li> <li>Member has a documented diagnosis of peoput allergy within the past 2</li> </ul>	One year
	<ul> <li>Member has a documented diagnosis of peanut allergy within the past 2 years (ICD-10 Z91.010) AND</li> <li>Diagnosis of peanut allergy is made by or in consultation with an allergist or immunologist AND</li> <li>Palforzia will be used in conjunction with a peanut-avoidant diet AND</li> <li>Member does not have a past or current history of any of the following:         <ul> <li>Severe, unstable or uncontrolled asthma</li> <li>Eosinophilic esophagitis or other eosinophilic gastrointestinal disease</li> <li>Mast cell disorder including mastocytosis, urticarial pigmentosa, and hereditary or idiopathic angioedema</li> <li>Severe or life-threatening anaphylaxis within the previous 60 days</li> </ul> </li> <li>AND</li> <li>Member has injectable epinephrine available for immediate use at all times and counseling regarding proper use has been provided AND</li> <li>Prescriber acknowledges member preparedness to adhere to complex up-dosing schedule and frequent visits to the administering healthcare facility AND</li> <li>Prescriber acknowledges that Palforzia doses administered by a</li> </ul>	
	<ul> <li>healthcare provider in the doctor's office or clinic are to be billed through the Health First Colorado medical benefit through the standard buy-and-bill process.</li> <li>Reauthorization: Member may receive reauthorization approval for 1 year if meeting</li> </ul>	
	<ul> <li>the following:</li> <li>Palforzia continues to be used in conjunction with a peanut-avoidant diet AND</li> <li>Member continues to tolerate the prescribed daily doses of Palforzia AND</li> <li>Member continues to have injectable epinephrine available for immediate use at all times AND</li> <li>Member has not experienced recurrent asthma exacerbations AND</li> <li>Member does not have eosinophilic esophagitis or other eosinophilic gastrointestinal disease AND</li> </ul>	
	<ul> <li>Member does not have a mast cell disorder including mastocytosis, urticarial pigmentosa, and/or hereditary/idiopathic angioedema AND</li> <li>Member has not experienced any treatment-restricting adverse effects (such as repeated systemic allergic reaction and/or severe anaphylaxis)</li> </ul>	
PALYNZIQ (pegvaliase-pqpz)	Maximum dose (maintenance): 300 mg daily         PALYNZIQ (pegvaliase-pgpz) prior authorization may be approved for members meeting the following criteria: <ul> <li>Member is at 18 years of age or older AND</li> </ul>	One year
	<ul> <li>Member has a diagnosis of phenylketonuria (PKU) AND</li> <li>Member has a blood phenylalanine concentration &gt; 600 mcmol/L AND</li> <li>Member is not receiving Palynziq in combination with Kuvan (sapropterin dihydrochloride) AND</li> <li>Member is actively on a phenylalanine-restricted diet AND</li> <li>Member will have a phenylalanine blood level measured at baseline prior to initiation and every four weeks until a maintenance dose is established AND</li> </ul>	

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	ROGRAM APPENDICES	
	<ul> <li>Prescriber acknowledges that first dose is being administered under the supervision of a healthcare provider equipped to manage anaphylaxis <b>AND</b></li> <li>Prescriber acknowledges that any doses administered in the doctor's office or clinic are to be billed to the Health First Colorado medical benefit through the standard buy-and-bill process.</li> <li>Reauthorization: Member may receive reauthorization approval for 1 year if meeting the following:         <ul> <li>Member is showing signs of continuing improvement, as evidenced by one of the following:                 <ul> <li>Blood phenylalanine level decrease of at least 20% from pretreatment baseline <b>OR</b></li> <li>Reduction of blood phenylalanine below 600 mcmol/L at current dose or maximum dose after 16 weeks of treatment.</li></ul></li></ul></li></ul>	
PCSK9 INHIBITORS Praluent, Repatha	<ul> <li>Maximum dose: 40 mg per day</li> <li>PCSK9 inhibitors may be approved for members that meet the following criteria:</li> <li>Medication is prescribed for one of the following diagnoses:         <ul> <li>Praluent® (alirocumab): heterozygous familial hypercholesterolemia or clinical atherosclerotic cardiovascular disease</li> <li>Repatha® (evolocumab): heterozygous familial hypercholesterolemia or homozygous familial hypercholesterolemia or clinical atherosclerotic cardiovascular disease</li> </ul> </li> </ul>	Initial Approval: 3 months Continuation Approval: One year
	Conditions Which Define Clinical Atherosclerotic Cardiovascular Disease         • Acute Coronary Syndrome         • History of Myocardial Infarction         • Stable or Unstable Angina         • Coronary or other Arterial Revascularization         • Stroke         • Transient Ischemic Attach         • Peripheral Arterial Disease of Atherosclerotic Origin	
	<ul> <li>PCSK9 inhibitor therapy is prescribed by, or in consultation with, one of the following providers:         <ul> <li>Cardiologist</li> <li>Certified Lipid Specialist</li> <li>Endocrinologist AND</li> </ul> </li> <li>Member is concurrently adherent (&gt;80% of the past 180 days) on maximally tolerated dose (see table below) of statin therapy (must include atorvastatin and rosuvastatin). If intolerant to a statin due to side effects, member must have a one month documented trial with at least two other statins. For members with a past or current incidence of rhabdomyolysis, one month failure is not required AND</li> <li>Member must be concurrently treated (in addition to maximally tolerated statin) with ezetimibe AND have a treated LDL ≥ 70 mg/dl for a clinical history of ASCVD or LDL ≥ 100 mg/dl if familial hypercholesterolemia AND</li> <li>PA will be granted for 3 months initially. Additional one year approval for continuation will be granted with provider attestation of safety and efficacy with initial medication therapy</li> </ul>	
	Atorvastatin 80mg Fluvastatin 80 mg	

COLORADO MEDICAID P	PROGRAM APPENDICES	
	Lovastatin 80 mg Pravastatin 80 mg	
	Rosuvastatin 40 mg	
	Simvastatin 40 mg (80 mg not used in practice)	
PHARMACIST PRESCRIPTIONS	The following OTC products will be covered with a written prescription by a pharmacist:	
	<ul> <li>Oral emergency contraceptive products</li> <li>Nicotine replacement therapy products including:</li> </ul>	
	• Nicotine gum (up to 200 units/fill)	
	<ul> <li>Nicotine patch (up to 30 patches/30days)</li> <li>Nicotine large (up to 288 up is (fill))</li> </ul>	
	<ul> <li>Nicotine lozenge (up to 288 units/fill)</li> <li>Children's dextromethorphan suspension for members age 4-11 years (up to</li> </ul>	
	150 ml per 30 days)	
	• Children's liquid and chewable acetaminophen for members age 2-11 years	
	<ul> <li>(up to 240 ml per 30 days)</li> <li>Children's liquid and chewable ibuprofen for members age 6 months – 11</li> </ul>	
	years (up to 240 mL per 30 days)	
PHYSICIAN ADMINISTERED DRUGS	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 <u>https://www.colorado.gov/hcpf/physician-</u> <u>administered-drugs</u> ).	
	<ul> <li>Physician administered drugs include any medication or medication formulation that is administered intravenously or 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) and may only be billed through the pharmacy benefit when given in a long-term care facility or when administered in the member's home by a healthcare professional or home health service. Prior authorization for physician administered drugs requires documentation of the following (in addition to meeting any other prior authorization criteria if listed):</li> <li>For drugs administered in the member's home by a home health agency or healthcare professional (home health administered): <ol> <li>Name of home health agency or healthcare professional</li> <li>Phone number</li> <li>Date and authorization number for home health agencies)</li> </ol> </li> <li>For drugs administered in a long-term care facility: <ol> <li>Name of long-term care facility</li> <li>Phone number of long-term care facility</li> </ol> </li> </ul>	
PRETOMANID	<ul> <li>PRETOMANID prior authorization may be approved for members meeting the following criteria:         <ul> <li>Member is an adult (≥ 18 years of age) AND</li> <li>Member has a confirmed diagnosis of multidrug resistant tuberculosis AND</li> <li>Pretomanid is prescribed by or in conjunction with an infectious disease specialist AND</li> <li>Pretomanid is prescribed in combination with bedaquiline and linezolid by</li> </ul> </li> </ul>	One year
	<ul> <li>Prescriber acknowledges member readiness and anticipated compliance with undergoing directly observed therapy (DOT) AND</li> </ul>	

COLORADO MEDICAID P	ROGRAM APPENDICES	
PREVYMIS (letermovir)	<ul> <li>Prescriber acknowledges that Pretomanid doses administered by a healthcare provider in a hospital, doctor's office, or clinic are to be billed through the Health First Colorado medical benefit through the standard buy-and-bill process.</li> <li>Maximum dose: 200 mg orally once daily</li> <li>Prevymis® (letermovir) may be approved for members that meet the following criteria:         <ul> <li>Member is a CMV-seropositive transplant recipient and meets ALL of the following: AND             <ul> <li>Member is 18 years or older.</li></ul></li></ul></li></ul>	100 days
	<ul> <li>Member has received an allogeneic hematopoietic stem cell transplant.</li> <li>Member does not have severe hepatic impairment (Child-Pugh Class C).</li> <li>Member is not receiving pitavastatin or simvastatin co-administered with cyclosporine.</li> <li>Member is not receiving pimozide or ergot alkaloids.</li> <li>Prevymis® is being prescribed by or in consultation with an oncologist, hematologist, infectious disease specialist, or transplant specialist. AND</li> <li>Provider agrees to monitor for CMV reactivation. AND</li> <li>Prevymis® dose does not exceed 480 mg orally or dose does not exceed 240mg if co-administered with cyclosporine. AND</li> <li>If request is for IV injectable Prevymis®, must provide medical justification why the patient cannot use oral therapy. AND</li> <li>If request is for IV injectable Prevymis®, must be administered in a long-term care facility or in a member's home by a home healthcare provider</li> </ul>	
	Length of Approval: Prevymis® will only be approved for 100 days Renewal: Authorization may be reviewed every 100 days to confirm that current medical necessity criteria are met and that the medication is effective (e.g. no evidence of CMV viremia).	
PROCYSBI (cysteamine)	Approval will be granted if the member is 2 years of age or older <b>AND</b> Has a diagnosis of nephropathic cystinosis <b>AND</b> documentation is provided to the Department that treatment with cysteamine IR (Cystagon®) was ineffective, not tolerated, or is contraindicated.	One year
PROMACTA (eltrombopag)	<ul> <li>Promacta@ (eltrombopag) prior authorization may be approved for members meeting criteria for the following diagnoses:</li> <li><u>Chronic immune idiopathic thrombocytopenia purpura:</u></li> <li>Confirmed diagnosis of chronic (&gt; 3 months) immune idiopathic thrombocytopenia purpura AND</li> <li>Must be prescribed by a hematologist AND</li> <li>Member is at risk (documented) of spontaneous bleed as demonstrated by the following labs: AND <ul> <li>Platelet count less than 20,000/mm3 or</li> <li>Platelet count less than 30,000/mm3 accompanied by signs and symptoms of bleeding</li> </ul> </li> <li>In the past 6 months, member has tried and failed (failure is defined as lack of efficacy, allergy, intolerable side effects or significant drug-drug interactions) systemic corticosteroids (e.g. prednisone 1 to 2 mg/kg for 2 to 4 weeks, or pulse dexamethasone 40 mg daily for 4 days), immunoglobulin replacement, or splenectomy.</li> </ul>	One year*
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	Thrombocytopenia associated with hepatitis C:	
	• Member must have confirmed diagnosis of chronic hepatitis C associated	
	thrombocytopenia AND	
	• Must be prescribed by a gastroenterologist, infectious disease specialist,	
	transplant specialist or hematologist AND	
	• Member has clinically documented thrombocytopenia defined as platelets <	
	60,000 microL AND	
	• Patients' degree of thrombocytopenia prevents the initiation of interferon-based	
	therapy or limits the ability to maintain interferon-based therapy	
	Severe aplastic anemia:	
	• Member must have confirmed diagnosis of severe aplastic anemia AND	
	• Must be prescribed by a hematologist AND	
	• Member must have had a documented insufficient response to	
	immunosuppressive therapy [antithymocyte globulin (ATG)] alone or in	
	combination with cyclosporine and/or a corticosteroid	
	*All initial prior authorization approvals will be granted for 12 months. Further	
	approvals for a maximum of 6 months require lab results and documentation for	
	efficacy.	
PROMETHAZINE	A Prior authorization is required for all routes of administration for members under	One year
	the age of two. Children under the age of two should not use Promethazine.	
	Promethazine is contraindicated in such patients because of the potential for fatal	
	respiratory depression.	
	Not qualified for emergency 3 day supply PA	
PROPECIA (finasteride)	Not covered for hair loss	One year
	Not qualified for emergency 3 day supply PA	
PULMOZYME (dornase	Pulmozyme® (dornase alfa) may be approved for members that meet the following	
alfa)	criteria:	
	<ul> <li>Member has a diagnosis of cystic fibrosis AND</li> <li>Member is five years of age or older</li> </ul>	
	<ul> <li>Member is five years of age or older</li> <li>For children &lt; 5 years of age, Pulmozyme will be approved if the member</li> </ul>	
	has severe lung disease as documented by bronchoscopy or CT scan	
	Pulmozyme twice daily will only be approved if patient has tried and foiled on adoptive trial of once daily design for one month	
	failed an adequate trial of once daily dosing for one month	
	All prior authorization renewals are reviewed on an annual basis to determine the	
	Medical Necessity for continuation of therapy. Authorization may be extended at 1-	
	year intervals based upon documentation from the prescriber that the member	
	continues to benefit from Pulmozyme therapy.	
	Quantity Limits: 30 ampules (2.5 mg/2.5 ml) per month	
QBREXZA	<b>QBREXZA</b> (glycopyrronium) prior authorization may be approved for members	Initial:
(glycopyrronium)	meeting the following criteria:	3 months
~~ · · /	• Member is 9 years of age or older <b>AND</b>	
	· · ·	
		Continued: One year

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	RUGRAIVI APPENDICES	
	<ul> <li>Member has a diagnosis of primary hyperhidrosis occurring more than once weekly and symptoms cease at night AND</li> <li>Member has a documented Hyperhidrosis Disease Severity Scale (HDSS) score of 3 or 4 AND</li> <li>There is documentation that the axillary hyperhidrosis is severe, intractable and disabling in nature as documented by at least one of the following:         <ul> <li>Significant disruption of professional and/or social life as a result of excessive sweating OR</li> <li>The condition is causing persistent or chronic cutaneous conditions (such as skin maceration, dermatitis, fungal infections, secondary microbial infections)</li> </ul> </li> <li>AND         <ul> <li>Prescriber has considered a trial of OTC topical antiperspirants (such as 20% aluminum chloride hexahydrate, 15% aluminum chloride hexahydrate, or 6.25% aluminum chloride hexahydrate)</li> </ul> </li> <li>Initial approval: 3 months</li> <li>Reauthorization: Member may receive reauthorization approval for 1 year if meeting the following:             <ul> <li>Member has documented improvement of at least two points in Hyperhidrosis Disease Severity Scale (HDSS) score following initiation</li> </ul> </li> </ul>	
	(or ongoing use) of Qbrexza regimen.	
	Manimum daare 1 slath aan daa	
RADICAVA (edaravone)	Maximum dose: 1 cloth per day <b>Radicava®</b> (edaravone) may be approved for members that meet the following	6 months
	<ul> <li>criteria:</li> <li>RADICAVA is being administered in a long-term care facility or in a member's home by a home healthcare provider AND</li> <li>Member has a "definite" or "probable" diagnosis of amyotrophic lateral sclerosis (ALS) based on medical history and diagnostic testing which may include imaging and nerve conduction conditions studies AND</li> <li>Member meets ALL of the following: <ul> <li>Member has a diagnosis of ALS for 2 or less years (for new starts only).</li> <li>Diagnosis has been established by or with the assistance of a neurologist with expertise in ALS using El Escorial or Airlie House diagnostic criteria (ALSFRS-R).</li> <li>Member has normal respiratory function as defined as having a percent-predicated forced vital capacity of greater than or equal to 80%.</li> <li>The ALSFRS-R score is greater than or equal to 2 for all items in the criteria.</li> <li>Member does not have severe renal impairment (CrCl&lt; 30 ml/min) or end stage renal disease</li> <li>Member does not have moderate or severe hepatic impairment (Child-Pugh Class C) AND</li> </ul> </li> </ul>	
	Length of Approval: 6 months. Quantity Limits: For patients initiating therapy, approval will include 28 bags per 28 days (initial dose) for the first month and 20 bags per 28 days for the remainder of the 6 months.	

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COLORADO MEDICAID H	PROGRAM APPENDICES	
	Renewal: Authorization may be reviewed every six months to confirm that current medical necessity criteria are met and that the medication is effective per improvement in ALSFRS-R score.	
RANITIDINE Capsule/Solution	Prescription ranitidine capsule and liquid formulations require prior authorization.	One year
	<u>Ranitidine capsule</u> : Require the prescribing provider to certify that capsules are medically necessary and that the member cannot use the tablets.	
	<u>Ranitidine liquid</u> : A prior authorization will be approved for members with a feeding tube or who have difficulty swallowing. A prior authorization is not required for children under 12 years of age.	
RAVICTI (glycerol phenylbutyrate)	<b>Ravicti</b> (glycerol phenylbutyrate) will only be approved for members meeting the following criteria:	One year
	<ul> <li>Member must have a documented diagnosis of urea cycle disorder (UCD)</li> <li>Member must be on a dietary protein restriction (verified by supporting documentation)</li> </ul>	
	• Member must have tried and failed Buphenyl as evidenced by uncontrolled hyperammonia over the past 365 days	
	• Medication must be prescribed by a physician experienced in the management of UCD (e.g., geneticist)	
REBATE DISPUTE DRUGS	Medical necessity.	One year
	Not qualified for emergency 3 day supply PA	
REVCOVI (elapegademase-lvlr)	<b>REVCOVI</b> (elepegademase-lvlr) may be approved for members meeting the following criteria:	One year
	f adenosine deaminase severe combined immune deficiency (ADA-SCID).	
	Maximum dose: Revcovi 0.4mg/kg per week (based on ideal body weight, IM administration)	
RUZURGI (amifampridine)	<b>Ruzurgi</b> (amifampridine) may be approved for members meeting the following criteria:	One year
	• Member is 6 to less than 17 years of age AND	
	• Member has a diagnosis of Lambert-Eaton myasthenic syndrome (LEMS)	
	Maximum dose: 100mg daily	
SANDOSTATIN (octreotide)	Approved for acromegaly; carcinoid tumors; and vasoactive intestinal peptide tumors.	Lifetime
SILENOR (doxepin tablet)	<b>Silenor</b> (doxepin) <u>tablets</u> may be approved if a member meets ONE of the following criteria:	One year
	• Contraindication to preferred oral sedative hypnotics (see preferred drug list "Sedative Hypnotic" class for list of preferred products) OR	
	<ul> <li>Prescriber attests to the medical necessity for use of doxepin dose &lt; 10 mg OR</li> <li>Member age is greater than 65 years</li> </ul>	
SIVEXTRO (tedizolid)	Sivextro may be approved for members $\geq 12$ years of age if all of the following criteria are met:	Six months
	• Member has diagnosis of acute bacterial skin and skin structure	
	• Member has diagnosis of acute bacterial skin and skin structure infection (ABSSSI) caused by one of the following Gram-positive	
	• Member has diagnosis of acute bacterial skin and skin structure	

APPENDICES

	AFFEIDICES	
	(including Streptococcus anginosus, Streptococcus intermedius, and	
	Streptococcus constellatus), and Enterococcus faecalis. AND	
	• Member has adequate trial and/or failure of linezolid 600mg twice daily	
	for 10 days. Failure is defined as: lack of efficacy with 10 day trial,	
	allergy, intolerable side effects or significant drug-drug interactions	
	anergy, intolerable side effects of significant drug-drug interactions	
	Maximum dosing: 200mg daily for 6 days total duration	
SODIUM CHLORIDE	Broncho Saline is not covered under the pharmacy benefit.	N/A
(Inhalation)	broneno sume <u>is not</u> covered under the pharmacy benefit.	1 1/2 1
(Innuturion)	Sodium chloride (inhalation use) must be billed through medical.	
SOLARAZE 3% GEL	A prior authorization will only be approved if the member has a diagnosis of Actinic	One year
(diclofenac sodium)	Keratoses (AK).	One year
SOLIRIS (eculizumab)	Soliris (ecluizumab) may be approved for members meeting all of the following	One year
Solinis (ccuizuillas)	criteria:	One year
	<ul> <li>Medication is being administered in the member's home or in a long-term care</li> </ul>	
	-	
	facility by a healthcare professional AND	
	• Member is diagnosed with either Paroxysmal Nocturnal Hemoglobinuria (PNH),	
	Atypical Hemolytic Uremic Syndrome (aHUS), Generalized Mysthenia Gravis	
	(gMG), or Neuromyleitis Optica Spectrum Disorder (NMOSD) AND	
	Member does not have a systemic infection AND	
	• Member must be administered a meningococcal vaccine at least two weeks prior	
	to initiation of Soliris therapy and revaccinated according to current medical	
	guidelines for vaccine use AND	
	• Prescriber is enrolled in the Soliris (eculizumab) Risk Evaluation and Mitigation	
	Strategy (REMS) program AND	
	<ul> <li>Medication is prescribed by or in conjunction with a hematologist for PNH and</li> </ul>	
	by or in conjunction with a hematologist or nephrologist for aHUS and by or in	
	conjunction with a neurologist for gMG or NMOSD AND	
	Member meets criteria listed below based on specific diagnosis:	
	Paroxysmal Nocturnal Hemoglobinuria	
	Member is 18 years of age or older AND	
	<ul> <li>Diagnosis of PHN must be accompanied by detection of PNH</li> </ul>	
	clones by flow cytometry diagnostic testing AND	
	• 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	
	• Member has one of the following indications for therapy:	
	• Presence of a thrombotic event	
	• Presence of organ damage secondary to chronic hemolysis	
	• Patient is pregnant and potential benefit outweighs	
	potential fetal risk	
	• Patient is transfusion dependent	
	• Patient has high LDH activity (defined as $\geq 1.5 \text{ x ULN}$ )	
	with clinical symptoms	
	AND	
	<ul> <li>Member has documented baseline values for one or more of the</li> </ul>	
	following:	
	•	
	• Serum lactate dehydrogenase (LDH)	
L	<ul> <li>Hemoglobin level</li> </ul>	

• Packed RBC transfusion requirement	
Atypical Hemolytic Uremic Syndrome	
<ul> <li>Member is 2 months or older AND</li> <li>Thromhotic Thromhogytononic Dumung (TTD) has been ruled out</li> </ul>	
<ul> <li>Thrombotic Thrombocytopenic Purpura (TTP) has been ruled out by evaluating ADAMTS13 level (ADAMTS-13 activity level &gt;</li> </ul>	
10%); AND	
<ul> <li>Shiga toxin E. coli related hemolytic uremic syndrome (STEC-</li> </ul>	
HUS) has been ruled out; AND	
• Other causes have been ruled out 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	
• Documented baseline values for one or more of the following:	
• Serum lactate dehydrogenase (LDH)	
• Serum creatinine/eGFR	
• Platelet count	
<ul> <li>Plasma exchange/infusion requirement</li> </ul>	
Generalized Myasthenia Gravis	
Member is 18 years or older AND	
• Patient has Myasthenia Gravis Foundation of America (MGFA)	
Clinical Classification of Class II to IV disease; AND	
• Patient has a positive serologic test for anti-acetylcholine receptor	
(AchR) antibodies; AND	
Physician has assessed the baseline Quantitative Myasthenia Gravis	
(QMG) score; AND	
• Patient has a MG-Activities of Daily Living (MG-ADL) total score	
of $\geq 6$ ; AND	
• Patient has failed 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)	
Neuromyelitis Optica Spectrum Disorder	
• Member is 18 years or older AND	
• Member has a past medical history of one of the following:	
<ul><li>Optic neuritis</li><li>Acute myelitis</li></ul>	
<ul> <li>Acute myelitis</li> <li>Area postrema syndrome; episode of otherwise</li> </ul>	
unexplained hiccups or nausea and vomiting	
<ul> <li>Acute brainstem syndrome</li> </ul>	
<ul> <li>Symptomatic narcolepsy or acute diencephalic clinical</li> </ul>	
syndrome with NMOSD-typical diencephalic MRI lesions	
• Symptomatic cerebral syndrome with NMOSD-typical	
brain lesions	
AND	
• Member has a positive serologic test for anti-aquaporin-4	
immunoglobulin G (AQP4-IgG)/NMP-IgG antibodies; AND	

COLORADO MEDICAID F	PROGRAM APPENDICES	
SOLOSEC (secnidazole)	<ul> <li>ADPENDICES         <ul> <li>Diagnosis of multiple sclerosis or other diagnoses have been ruled out AND</li> <li>Member has not failed a previous course of Soliris (eculizumab) therapy AND</li> <li>Member has a history of failure, contraindication, or intolerance to rituximab therapy AND</li> <li>Member has at least one of the following:                 <ul> <li>History of at least two relapses during the previous 12 months prior to initiating Soliris (eculizumab)</li></ul></li></ul></li></ul>	One year
	interaction, or contraindication to therapy) Solosec® Maximum Quantity: 1 packet of 2 grams per 30 days	
STRENSIQ (asfotase alfa)	<ul> <li>Strensiq® (asfotase alfa) may be approved if all of the following criteria are met:</li> <li>Member has a diagnosis of either perinatal/infantile- OR juvenile-onset hypophosphatasia (HPP) based on all of the following <ul> <li>a. Member was ≤ 18 years of age at onset</li> <li>b. Member has/had clinical manifestations consistent with hypophosphatasia at the age of onset prior to age 18 (e.g. vitamin B6-dependent seizures, skeletal abnormalities: such as rachitic chest deformity leading to respiratory problems or bowed arms/legs, "failure to thrive").</li> <li>c. Member has/had radiographic imaging to support the diagnosis of hypophosphatasia at the age of onset prior to age 18 (e.g. infantile rickets, alveolar bone loss, craniosynostosis)</li> <li>d. Member has one of the following: elevated urine concentration of phosphoethanolamine (PEA), elevated serum concentration of pyridoxal 5'-phosphate (PLP) in the absence of vitamin supplements within one week prior to the test, or elevated urinary inorganic pyrophosphate (PPi) AND</li> </ul> </li> </ul>	Six months
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#### COLORADO MEDICAID PROGRAM **APPENDICES** Molecular genetic test has been completed confirming mutations in the ALPL e. gene that encodes the tissue nonspecific isoenzyme of ALP (TNSALP) within 30 days of initiation. If genetic test is negative, approval will not be granted past 30 days. Prescriber is a specialist in the area of the members disease (such as an f. endocrinologist) Symdeko® (tezacaftor/ivacaftor and ivacaftor) may be approved for members that **SYMDEKO** One year (tezacaftor/ivacaftor and meet the following criteria: ivacaftor) The member has a diagnosis of cystic fibrosis AND The member is 6 years of age or older AND • The member has one of the following mutations: Homozygous for the F508del mutation in the CFTR gene 2 OR 0 Heterozygous for the F508del mutation in the CFTR gene and one 0 of the following mutations: E56K, P67L, R74W, D110E, D110H, R117C, E193K, L206W, R347H, R352Q, A455E, D1270N, D579G, 711+3A-G, E831X, S945L, S977F, F1052V, K1060T, A1067T, R1070W, F1074L, D1152H, 3272-26A-G, 2789+5G-A, 3849-10kbC-T, or another FDA approved gene mutation AND Member has ALT, AST, and bilirubin at baseline and tested every 3 months for the first year AND Member has a baseline ophthalmological examination and periodic followup exams for cataracts AND Must be prescribed by or in consultation with a pulmonologist or gastroenterologist AND Member is not receiving dual therapy with another cystic fibrosis transmembrane conductance regulator (CFTR) potentiator AND Member has had 2 negative respiratory cultures for any of the following organisms: Burkholeria cenocepacia, Burkholderia dolosa, or Mycobacterium abscessus in the past 12 months. SYNAGIS (palivizumab) Pharmacy prior authorization requests for Synagis must be submitted by fax Maximum using the Synagis prior authorization form found at of 5 doses https://www.colorado.gov/hcpf/provider-forms. Medical prior authorization per season requests must be submitted at http://coloradopar.com/. Synagis season will begin November 16, 2020 and end April 16, 2021. Prior authorization may be requested beginning November 2, 2020. Synagis given in a doctor's office, hospital or dialysis unit is to be billed directly by those facilities as a medical benefit. Synagis may only be a pharmacy benefit if the medication is administered in the member's home or long-term care facility. **Key Points** 1. No more than 5 doses per season. 5 doses provide more than 6 months of protective serum concentration. 2. Synagis is not recommended for controlling outbreaks of health care-associated disease. Synagis is not recommend for prevention of health care-associated RSV disease. 3. 4. Infants born later in the season may require less than 5 doses to complete therapy to the end of the season. 5. Monthly prophylaxis should be discontinued in any child who experiences a breakthrough RSV hospitalization. Effective 04/01/2021 Revised 02/26/2021 Page A-53

COLORADO MEDICAID P	RO	GRAM APPENDICES	
	6.	Synagis is not recommended to prevent wheezing, nosocomial disease, or treatment of RSV	
	7.	Synagis is not routinely recommended for patients with a diagnosis of Down	
	1.		
	0	syndrome unless they also have a qualifying indication listed below.	
	8.	In the <b>first year of life</b> Synagis is recommended:	
		a. For infants born before 29w 0d gestation.	
		b. For infants born before 32w 0d AND with chronic lung disease (CLD) of	
		prematurity <b>AND</b> requirements of >21% oxygen for at least 28 days after birth.	
		c. For infants with hemodynamically significant heart disease ( <u>a</u> cyanotic heart disease who are reaciving mediantion to control concessive heart foilure)	
		disease who are receiving medication to control congestive heart failure	
		(CHF) and will require cardiac surgical procedures or infants with moderate	
		to severe pulmonary hypertension) <b>AND</b> born within 12 months of onset of the RSV season.	
		d. Infants who undergo cardiac transplantation during the RSV season.	
		e. For infants with cyanotic heart defects <b>AND</b> in consultation with a pediatric	
		cardiologist <b>AND</b> requirements of >21% oxygen for at least 28 days after	
		birth AND continue to require medical intervention (supplemental oxygen,	
	1	chronic corticosteroid, or diuretic therapy)	
		f. If an infant has neuromuscular disease or pulmonary abnormality <b>AND</b> is	
		unable to clear secretions from the upper airways	
		g. An infant who will be profoundly immunocompromised during the RSV	
		season (solid organ or hematopoietic stem cell transplantation, receiving	
		chemotherapy)	
		h. An infant with cystic fibrosis with clinical evidence of CLD AND/OR	
		nutritional compromise	
	9.	In the second year of life Synagis is recommended for:	
		a. Children born before 32w 0d AND with CLD of prematurity AND	
		requirements of >21% oxygen for at least 28 days after birth AND continue	
		to require medical intervention (supplemental oxygen, chronic corticosteroid,	
		or diuretic therapy)	
		b. A child who will be profoundly immunocompromised during the RSV	
		season (solid organ or hematopoietic stem cell transplantation, receiving	
		chemotherapy)	
		c. Children with manifestations of severe lung disease (previous hospitalization	
		for pulmonary exacerbation in the first year of life or abnormalities of chest	
		radiography or chest computed tomography that persist when stable) <b>OR</b>	
		weight for length less than the 10 <sup>th</sup> percentile.	
		d. Children who undergo cardiac transplantation during the RSV season.	
SYPRINE (trientine)	Sw	prine® (trientine) may be approved if all of the following criteria are met:	One year
STI KINE (trientine)	Sy		One year
		Must be prescribed in conjunction with a gastroenterologist, hepatologist, or liver transplant specialist. <b>AND</b>	
		1 1	
	•	Member has a diagnosis of Wilson's Disease meeting at least one of the	
		following criteria:	
	1	• Hepatic parenchymal copper content of $\geq 250 \mu g/g$ dry weight	
		• Presence of Kayser-Fleischer Ring in cornea	
		• Serum ceruloplasmin level <50mg/L	
		<ul> <li>Basal 24-hour urinary excretion of copper &gt;100μg (1.6 μmoles)</li> </ul>	
		• Genetic testing results indicating mutation in ATP7B gene	
		AND	
	•	Member has failed a three-month trial or is intolerant to penicillamine. Failure is	
		defined as a lack of efficacy, allergy, intolerable side effects, contraindication to,	
		or significant drug-drug interactions <b>AND</b>	
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COLORADO MEDICAID	PROGRAM APPENDICES	
	• Member has failed a three-month trial or is intolerant to generic trientine. Failure is defined as a lack of efficacy, allergy, intolerable side effects, contraindication to, or significant drug-drug interactions.	
TAMIFLU (oseltamivir) capsules	Effective 10/15/2019: Claims for brand Tamiflu® capsules require prior authorization approval (see section "Brand Name Medications and Generic Mandate" for brand product coverage details). Generic equivalent oseltamivir formulations do not require prior authorization.	
TAVALISSE (fostamatinib)	<ul> <li>Tavalisse<sup>®</sup> (fostamatinib) prior authorization may be approved for members meeting the following criteria:</li> <li>Member is 18 years of age or older AND</li> <li>Member has a documented diagnosis of chronic immune thrombocytopenia AND</li> <li>Member has trialed and failed at least ONE of the following therapies (Failure is defined as a lack of efficacy, allergy, intolerable side effects, or significant drug-drug interactions): <ul> <li>Promacta (eltrombopag) or other thrombopoietin receptor agonist</li> <li>Corticosteroids</li> <li>Immunoglobulin</li> <li>Splenectomy</li> </ul> </li> <li>AND</li> <li>Baseline platelet count prior to initiation is less than 30x10<sup>9</sup>/L or 30x10<sup>9</sup>/L to 50x10<sup>9</sup>/L with symptomatic bleeding AND</li> <li>Prescriber attests to monitoring liver function tests and CBC monthly until a stable dose is achieved AND</li> <li>Tavalisse (fostamatinib) is not being used as dual therapy with a thrombopoietin receptor agonist AND</li> <li>Initial prior authorization approval will be for 3 months. Continuation may be approved with verification of documented platelet response (platelet count ≥50x109/L)</li> </ul>	Initial Approval: 3 months Continuation Approval: One year
	Quantity Limit: 60 tablets per 30 days	
TARGETED IMMUNE MODULATORS (IV and physician-administered products)	<ul> <li>Actemra (tocilizumab) IV injection may be approved if meeting the following criteria:</li> <li>Actemra is being prescribed for an FDA-labeled indication (per product package labeling) AND</li> <li>Member has trialed and failed ALL preferred agents in the "Targeted Immune Modulators" PDL drug class that are FDA-labeled for use for the same prescribed indication (failure is defined as lack of efficacy of a three-month trial, contraindication to therapy, allergy, intolerable side effects or significant drug-drug interaction) AND</li> <li>Actemra IV injection is being administered by a healthcare professional in the member's home or in a long-term care facility</li> </ul>	One year (for Stelara, see criteria)
	<ul> <li>Entyvio (vedolizumab) may be approved for members who are receiving infusion in their home or in a long-term care facility and who meet the following criteria:</li> <li>Medication is being used in an adult member with ulcerative colitis or Crohn's disease AND</li> </ul>	

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ORADO MEDICAID PI	ROGRAM APPE	NDICES
ORADO MEDICAID P	<ul> <li>For diagnosis of Crohn's disease, have trialed and failed<sup>‡</sup> Humira and OR for a diagnosis of ulcerative colitis, have trialed and failed<sup>‡</sup> Humi Simponi AND</li> <li>Member has had an inadequate response with, intolerance to, or demon dependence on corticosteroids AND</li> <li>Member is not receiving Entyvio in combination with Humira, Simpon Tysabri AND</li> <li>Medication is initiated and titrated per FDA-labeled dosing for Crohn' and Ulcerative Colitis up to a maximum of 300mg IV infusion every 8</li> <li>Inflectra (infliximab dyyb) may be approved with trial &amp; failure<sup>‡</sup> of Renfl (infliximab abda) AND if meeting all of the following criteria:</li> <li>Medication is being administered in the member's home or in a long-t facility AND</li> <li>Member has one of the following diagnoses:         <ul> <li>Crohn's disease and is 6 years or older</li> <li>Ulcerative colitis and is 6 years or older</li> <li>Rheumatoid arthritis in adults</li> <li>Ankylosing spondylitis in adults</li> <li>Juvenile idiopathic arthritis</li> <li>Plaque psoriasis in adults</li> <li>Plaque psoriasis in adults</li> <li>Orencia (abatacept) – may be approved for members who are receiving th in their home or in long-term care and who meet one of the following:</li> </ul> </li> <li>Member has a diagnosis of moderate to severe rheumatoid arthritis or polyarticular juvenile idiopathic arthritis AND has trialed and failed<sup>‡</sup> preferred agents in the "Targeted Immune Modulators" PDL drug class that are FDA-labeled for use for the sam prescribed indication</li> <li>Member has a diagnosis of moderate to severe rheumatoid arthritis or polyarticular juvenile idiopathic arthritis AND has trialed and failed<sup>‡</sup> preferred agents in the "Targeted Immune Modulators" PDL drug class abda) AND is frieded for use for the prescribed indication OR</li> <li>Member has a diagnosis of psoriatic arthritis AND ha</li></ul>	I Cimzia ra and nstrated a ni, or 's Disease B weeks exis erm care e infusion all s that are aled and (infliximab
	<ul> <li>Member has one of the following diagnoses:</li> <li>Crohn's disease and is 6 years or older</li> </ul>	e

	<ul> <li>Medication is being administered in the member's home or long-term care facility AND</li> <li>Member has one of the following diagnoses:         <ul> <li>Crohn's disease and is 6 years or older</li> <li>Ulcerative colitis and is 4 years or older</li> <li>Rheumatoid arthritis and is 4 years or older</li> <li>Psoriatic arthritis in adults</li> <li>Ankylosing spondylitis in adults</li> <li>Juvenile idiopathic arthritis</li> <li>Plaque psoriasis in adults</li> <li>Ankylosing spondylitis in adults</li> <li>Juvenile idiopathic arthritis</li> <li>Plaque psoriasis in adults</li> <li>AND</li> </ul> </li> <li>Member has tried and failed<sup>‡</sup> all preferred agents in the "Targeted Immune Modulators" PDL drug class that are FDA-labeled for use for the prescribed indication.</li> <li>Rituxan (rituximab) IV and subcutaneous - will be approved for administration in a long-term care facility or in a member's home by a home healthcare provider AND for members who meet one of the following:         <ul> <li>Have diagnosis of moderate to severe rheumatoid arthritis AND have tried and failed both Enbrel and Humira OR</li> <li>Have diagnosis of chronic lymphocytic leukemia OR</li> <li>Have diagnosis of Non-Hodgkins Lymphoma</li> </ul> </li> <li>Stelara is being prescribed for an FDA-labeled indication (per product package labeling) AND         <ul> <li>Member has trialed and failed<sup>‡</sup> ALL preferred agents in the "Targeted Immune Modulators" PDL drug class that are FDA-labeled for use for the same prescribed indication (ND alabeled for use for the same prescribed indication (per product package labeling) AND</li> <li>Member has tried and failed<sup>‡</sup> ALL preferred agents in the "Targeted Immune Modulators" PDL drug class that are FDA-labeled for use for the same prescribed indication for one year m</li></ul></li></ul>	
	<ul> <li>Member has tried and rando<sup>*</sup> an preferred agents in the "Fargeted infinute Modulators" PDL drug class that are FDA-labeled for use for the prescribed indication.</li> </ul>	
	<sup>‡</sup> Failure is defined as: lack of efficacy, allergy, intolerable side effects, or significant drug-drug interaction. Note that trial and failure of Xeljanz IR will not be required when prescribed for ulcerative colitis for members $\geq 50$ years of age that have an additional CV risk factor.	
THIOLA EC (tiopronin DR)	<ul><li>Thiola EC<sup>®</sup> (tiopronin DR) may be approved for members meeting the following criteria:</li><li>Member is an adult or pediatric weighing 20kg or more AND</li><li>Member has severe homozygous cystinuria AND</li></ul>	One year
	· · · · · · · · · · · · · · · · · · ·	

COLORADO MEDICAID F	PROGRAM APPENDICES	
	<ul> <li>Member has increased fluid intake and diet modifications have been implemented for the prevention of cysteine stone formation AND</li> <li>Member has trial and failure of urinary alkalization agent (such as potassium citrate or potassium bicarbonate) AND</li> <li>Member has trial and failure of Thiola IR (tiopronin). Failure is defined as lack of efficacy with 14 day trial, allergy, intolerable side effects or significant drug-drug interactions.</li> <li>Maximum dose: Thiola EC 1500mg per day</li> </ul>	
THROMBOLYTIC ENZYMES	Approved for <b>IV Catheter Clearance or Occluded AV Cannula</b> if given in member's home or long term care facility.	One year
TOBACCO CESSATION	Effective 11/01/18 prior authorization will not be required for tobacco cessation medications including nicotine gum, nicotine patch, nicotine lozenge, nicotine inhaler (Nicotrol <sup>®</sup> ), varenicline (Chantix <sup>®</sup> ), and bupropion SR (Zyban <sup>®</sup> ). Smoking and tobacco cessation resources are available at no charge to members or providers through the Colorado QuitLine found at coquitline.org or by calling 1-800-QUIT-NOW.	
TRIKAFTA (elexacaftor, tezacaftor, ivacaftor)	<ul> <li>Trikafta® may be approved for members meeting the following criteria:</li> <li>Member is 12 years of age or older AND</li> <li>Member has at least one F508del mutation in the cystic fibrosis transmembrane conductance regulator (CTFR) gene or a mutation in the CFTR gene that is responsive based on in vitro data AND</li> <li>Member continues to receive standard of care CF therapies (such as bronchodilators, inhaled antibiotics, dornase alfa, and hypertonic saline) AND</li> <li>Member must have liver function tests checked within 3 months without abnormal results (ALT, AST, ALP, or GGT ≥ 3 × ULN, or total bilirubin ≥2 × ULN) AND</li> <li>Baseline Forced Expiratory Volume (FEV1) must be collected</li> </ul>	One year
TPN PRODUCTS	Approval will be given if included as part of TPN therapy administered in the member's home or in a long-term care facility by a home healthcare provider. If given in the hospital or physician's office, the claim must be billed as a medical expense.	Lifetime
TYBOST (cobicistat)	<ul> <li>Tybost® may be approved for members meeting the following criteria:</li> <li>Member has a diagnosis of HIV-1 AND</li> <li>Member is currently being treated with atazanavir or darunavir only AND</li> <li>Member is not taking cobicistat-containing drugs, or ritonavir-containing drugs AND</li> <li>Member has failed treatment with ritonavir (failure defined as intolerable side effect, allergy, or lack of efficacy).</li> </ul>	One year
TYSABRI (natalizumab)	<ul> <li>TYSABRI (natalizumab) will be approved for initial therapy if the following criteria are met:</li> <li>Tysabri is being administered in a long-term care facility or in home-health setting AND</li> <li>Medication is not currently being used in combination with immunosuppresants (azathioprine, 6-mercaptopurine, methotrexate) or TNF-alpha inhibitors (adalimumab, certolizumab pegol, infliximab) AND</li> </ul>	One year

### APPENDICES

	<ul> <li>If prescribed for induction of remission of moderate to severe Crohn's disease</li> <li>The patient is ≥ 18 years of age AND</li> <li>Member has tried and failed Aminosalicylates AND</li> <li>Member has tried and failed Corticosteroids AND</li> <li>Member has tried and failed immunomodulators AND</li> <li>Member has tried and failed two TNF-alpha inhibitors (e.g. adalimumab, certolizumab pegol, infliximab) AND</li> <li>Tysabri is prescribed by or in consultation with a gastroenterologist.</li> </ul>	
ULTOMIRIS (ravulizumab)	If prescribed for relapsing remitting multiple sclerosis (RRMS)         • The patient is ≥ 18 years of age; AND         • Member has trial and failure of three of the following agents: Avonex (interferon beta-1a), Rebif (interferon beta 1-a), Betaseron/Extavia (interferon beta-1b), Plegridy (peginterferon beta 1a), Copaxone/Glatopa (glatiramer acetate), Aubagio (teriflunomide tablets), Gilenya (fingolimod capsules), Tecfidera (dimethyl fumarate delayed-release capsules), Ocrevus (ocrelizumab) or Lemtrada (alemtuzumab). Failure will be defined as intolerable side effects, drug-drug interaction, or lack of efficacy indicated by one of the following: • One of the following on MRI: presence of any new spinal lesions, cerebellar or brain stem lesions, or change in brain atrophy • On clinical exam, signs and symptoms consistent with functional limitations that last one month or longer AND         • Tysabri is prescribed by or in consultation with a neurologist or a physician that specializes in the treatment of multiple sclerosis         Ultomiris <sup>®</sup> (ravulizumab) may be approved for members meeting the following criteria: Medication is being administered in the member's home or in a long-term care facility by a healthcare professional AND	One year
	<ul> <li>Member has a diagnosis of either paroxysmal nocturnal hemoglobinuria (PNH) OR atypical hemolytic uremic syndrome (aHUS).</li> <li>Maximum dose: Ultomiris 3.6g every 8 weeks (IV infusion)</li> </ul>	
VACCINES	<ul> <li>Pharmacies that have entered into a collaborative practice agreement with one or more physicians may receive reimbursement (with claim submission through the Health First Colorado <u>medical</u> benefit) for enrolled pharmacists to administer the following vaccines (claims for pharmacist administration of vaccines are not covered under the pharmacy benefit): <ul> <li>Covid-19</li> <li>Influenza</li> <li>Pneumococcal</li> <li>Shingles</li> <li>Tdap</li> <li>Td</li> </ul> </li> <li>Additional information regarding pharmacist enrollment and vaccine medical claims billing can be found at <a href="https://www.colorado.gov/hcpf/otc-immunizations">https://www.colorado.gov/hcpf/otc-immunizations</a>.</li> <li>Vivotif oral typhoid vaccine may be approved under the pharmacy benefit for outpatient administration.</li> </ul>	

### APPENDICES

COLORADO MEDICAID P	RUGRAIN	APPENDICES	
	administered in a long-term care facility. I	-	
	Not qualified for emergency 5 day suppry	IA	
VALCYTE (valganciclovir hydrochloride)	Effective 10/15/19: Brand Valcyte <sup>®</sup> soluti product (see section "Brand Name Medica product coverage details).		One year
	Valcyte® will be approved for members w Cytomegalovirus (CMV) retinitis AND ac Syndrome (AIDS) per dosing guidelines b OR	quired immunodeficiency	
	For members that require prophylactic treatment for CMV post kidney, heart or kidney-pancreas transplant per dosing guidelines below OR		
	For members $\leq 16$ years of age that are at and need prophylactic treatment post hear per dosing guidelines below	•	
	Adu	lt Dosage	
	Treatment of CMV retinitis	Induction: 900 mg (two 250 mg tablets) twice a day for 21 days Maintenance: 900 mg once a day	
	Prevention of CMV disease in heart or kidney-pancreas patients	900 mg once a day within 10 days of transplantation 100 days post- transplantation	
	Prevention of CMV disease in kidney transplant patients	900 mg once a day within 10 days of transplantation until 200 days post- transplantation	
		tric Dosage	
	Prevention of CMV disease in kidney transplant patients 4 month to 16 years of age	Dose once daily within 10 days of transplantation until 200 days post- transplantation	
	Prevention of CMV disease in heart transplant patients 1 month to 16 years of age	Dose once a day within 10 days of transplantation until 100 days post- transplantation	
VALTOCO (diazepam)	Valtoco <sup>®</sup> (diazepam) may be approved for • Member is 6 years of age or	r members meeting the following criteria:	One year
	stereotypic episodes of frequ acute repetitive seizures) tha	t are distinct from a patient's usual seizure are provided supporting this diagnosis AND	
	<ul><li>Member is stable on regimer</li><li>Medication is being prescrib</li></ul>	n of antiepileptic medications AND ed by or in conjunction with the same manages the member's anti-epileptic	
	<ul><li>regimen AND</li><li>Member is educated on appr</li></ul>	opriate identification of seizure cluster and tration and not to exceed 2 doses per seizure	

COLORADO MEDICAID P		
	Maximum dose: 4 nasal spray units per year unless used / damaged / lost	
	Members are limited to one prior authorization approval on file for Valtoco (diazepam) and Nayzilam (midazolam).	
	Grandfathering: If member is currently receiving Valtoco (diazepam) intranasal, they may receive prior authorization approval to continue.	
VELTASSA (patiromer)	<ul> <li>Veltassa® prior authorization will be approved for members that meet the following criteria:</li> <li>Documented diagnosis of hyperkalemia (serum potassium &gt; 5 mEq/L) AND</li> <li>Veltassa is not being used for emergent hyperkalemia AND</li> <li>Member does not have severe gastrointestinal motility dysfunction AND</li> <li>Member does not have hypomagnesemia (serum magnesium &lt; 1.4 mg/dL)</li> </ul>	One year
VERIPRED (prednisolone)	A prior authorization will only be approved if a member has tried and failed on a generic prednisolone product (Failure is defined as: lack of efficacy, allergy, intolerable side effects or significant drug-drug interactions.)	One year
VERSED (midazolam)	Effective 09/25/2019 prior authorization is no longer required for generic midazolam	
Injection	vial/syringe formulations.	0
VIMIZIM (elosulfase alfa)	<ul> <li>VIMIZIM (elosulfase alfa) prior authorization may be approved for members meeting the following criteria:</li> <li>Member is ≥ 5 years of age AND</li> <li>Member has a confirmed diagnosis of mucopolysaccharidosis (MPS)</li> </ul>	One year
	<ul> <li>Type IV A (Morquio A syndrome) AND</li> <li>Medication is being administered by a healthcare provider in the member's home or in a long-term care facility (and meets approval criteria listed in "Physician Administered Drug" section of Appendix P) AND</li> </ul>	
	<ul> <li>Vimizim is prescribed by or in consultation with an endocrinologist         AND     </li> <li>Prescriber acknowledges that Vimizim will be administered under close</li> </ul>	
	medical observation due to risk of life-threatening anaphylactic reactions.	
VITAMINS* (prescription vitamins)	*Coverage criteria outlined in this section apply to vitamin products available as prescription drugs. For over-the-counter product coverage, please see "OTC Products" section.	One year
	<ul> <li>The following prescription vitamin products will be covered without prior authorization:</li> <li>Vitamin D</li> <li>Vitamin K</li> </ul>	
	<ul> <li>**General prescription vitamin criteria: Prescription vitamin products will be approved for:</li> <li>ESRD, CRF, renal insufficiency, diabetic neuropathy or renal transplant OR</li> <li>Members under the age of 21 with a disease state or clinical diagnosis associated with prohibited nutritional absorption processes as a secondary effect OR</li> <li>Members with Erythema Bullosum</li> </ul>	
	<ul> <li>Hydroxocobalamin injection will be approved for:</li> <li>Members meeting any general prescription vitamin criteria** OR</li> <li>Methylmalonic acidemia (MMA)</li> </ul>	
	<ul> <li>Cyanocobalamin will be approved for:</li> <li>Members meeting any general prescription vitamin criteria** OR</li> </ul>	

COECITA ADO MIEDIO AID I		1
	Vitamin B12 deficiency	
	<ul> <li>Folic acid prescription products will be approved for: <ul> <li>Members meeting any general prescription vitamin criteria** OR</li> <li>Folic acid 1mg will be approved for female members without a prior authorization OR</li> <li>Members currently taking methotrexate or pemetrexed OR</li> <li>Documented folic acid deficiency by the treating clinician (megaloblastic and macrocytic anemia are the most common. Some drugs or other conditions may cause deficiency as well) OR</li> <li>Homocysteinemia OR</li> <li>Sickle cell disease OR</li> <li>Female members prescribed folic acid for the prevention of a neural tube defect during pregnancy or for the prevention of miscarriage</li> </ul> </li> </ul>	
	<ul> <li>Cyanocobalamin/folic acid/pyridoxine prescription products will be approved for:</li> <li>Members meeting any general prescription vitamin criteria** ORMembers meeting any general prescription vitamin criteria* OR</li> <li>Members with Homocysteinemia or Homocystinuria OR</li> <li>Members on dialysis OR</li> <li>Members with (or at risk for) cardiovascular disease</li> </ul>	
	For prescription iron-containing products see "Anti-anemia Medications"	
	Metanx will be approved for members with non-healing diabetic wounds	
VUSION OINTMENT (miconazole/zinc oxide/white petrolatum)	A prior authorization will only be approved if a member has failed on an OTC antifungal <b>and</b> a generic prescription antifungal. (Failure is defined as: lack of efficacy, allergy, intolerable side effects or significant drug-drug interactions)	One year
VYNDAMAX (tafamidis)	<ul> <li>Vyndamax<sup>®</sup> (tafamidis) may be approved for members meeting the following criteria:</li> <li>Member is an adult ≥ 18 years of age AND</li> <li>Member has a diagnosis of cardiomyopathy of wild type or hereditary transthyretin-mediated amyloid cardiomyopathy (ATTR-CM) AND</li> <li>Member has a documented history of heart failure with NYHA functional class I-III</li> <li>Maximum dose: Vyndamax (tafamidis) 61mg daily</li> </ul>	One year
VYNDAQEL (tafamidis meglumine)	<ul> <li>Vyndaqel<sup>®</sup> (tafamidis meglumine) may be approved for members meeting the following criteria:</li> <li>Member is an adult ≥ 18 years of age AND</li> <li>Member has a diagnosis of cardiomyopathy of wild type or hereditary transthyretin-mediated amyloid cardiomyopathy (ATTR-CM) AND</li> <li>Member has a documented history of heart failure with NYHA functional class I-III</li> </ul>	One year
	Maximum dose: Vyndaqel (tafamidis meglumine) 80mg daily	
VYONDYS 53 (golodirsen)	<b>Vyondys 53</b> <sup>®</sup> may be approved if all the following criteria are met:	One year

	APPENDICES	
	<ul> <li>Medication is being administered in the member's home or in a long-term care facility by a healthcare professional AND</li> <li>Member has a diagnosis of Duchenne Muscular Dystrophy (DMD) AND</li> <li>Member must have genetic testing confirming mutation of the DMD gene that is amenable to exon 53 skipping AND</li> <li>Medication is prescribed by or in consultation with a neurologist or a provider who specializes in treatment of DMD (i.e. pediatric neurologist, cardiologist or pulmonary specialist) AND</li> <li>The member must be on corticosteroids at baseline or has a contraindication to corticosteroids AND</li> <li>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.</li> </ul>	
	Maximum Dose: 30 mg/kg per week	
XERMELO (telotristat ethyl)	<ul> <li>XERMELO (telotristat ethyl) prior authorization may be approved for members meeting the following criteria:</li> <li>Member is at 18 years of age or older AND</li> <li>Member has a diagnosis of carcinoid syndrome diarrhea AND</li> <li>Member has trialed and failed three months of somatostatin analog therapy (such as octreotide). Failure is defined as lack of efficacy, allergy, intolerable side effects, or significant drug-drug interaction AND</li> <li>Xermelo is being used in combination with somatostatin analog therapy</li> </ul>	One year
	Maximum dose: 750 mg per day	
XIFAXAN (rifaximin)	<ul> <li>Xifaxan® prior authorization will be approved for members meeting the following criteria:</li> <li>For members prescribed Xifaxan for prophylaxis of hepatic encephalopathy (HE) in adults:         <ul> <li>Member must be concomitantly taking lactulose or other non-</li> </ul> </li> </ul>	See Criteria
	<ul> <li>absorbable disaccharide AND         <ul> <li>Member must not have undergone transjugular intrahepatic portosystemic shunt (TIPS) procedure within the last 3 months AND</li> <li>Xifaxan is being prescribed for secondary prophylaxis of HE (member has experienced previous episode of HE) AND</li> <li>Maximum dosing regimen is 550mg twice daily</li> <li>Members meeting criteria will receive approval for one year</li> </ul> </li> <li>For members prescribed Xifaxan for irritable bowel syndrome with diarrhea (IBS-D):         <ul> <li>Maximum dosing regimen is 550mg three times daily for 14 days AND</li> <li>Approval is limited to two 14-day treatment courses per 14 week time period</li> </ul> </li> </ul>	
	<ul> <li>For members prescribed Xifaxan for traveler's diarrhea:         <ul> <li>Member must be ≥ 12 years of age AND</li> <li>Maximum dosing regimen is 200mg three times daily for 3 days</li> <li>Members meeting criteria will receive approval for one year</li> </ul> </li> </ul>	
XOLAIR (omalizumab)	A prior authorization will only be approved as a pharmacy benefit when the medication is administered in a long-term care facility. Medications administered in a	One year

### APPENDICES

	Because this medication has an FDA Boxed Warning requiring administration under	
	the supervision of a physician, a PA will not be approved if administered in a	
	member's home.	
XYREM (sodium oxybate)	<b>Xyrem</b> (sodium oxybate) may be approved for <u>adults and children 7 to 17 years of</u>	Initial
	age if all the following criteria are met:	Approval:
	Member has a diagnosis of cataplexy or excessive daytime sleepiness	30 days
	with narcolepsy (confirmed by one of the following):	Continuation
	<ul> <li>Cataplexy episodes occurring three or more times per month</li> </ul>	Approval:
		One year
	OR UNITED STATES OF	
	• Hypocretin deficiency OR	
	<ul> <li>Nocturnal sleep polysomnography showing rapid eye</li> </ul>	
	movement (REM) sleep latency less than or equal to 15	
	minutes, or a Multiple Sleep Latency Test (MSLT) showing a	
	mean sleep latency less than or equal to 8 minutes and two or	
	more sleep-onset REM periods	
	AND	
	• Baseline excessive daytime sleepiness is measured using the Epworth	
	Sleepiness Scale or cataplexy episode count AND	
	<ul> <li>Member has adequately trialed and failed therapy with 3 stimulants for</li> </ul>	
	narcolepsy (examples include methylphenidate and amphetamine salts)	
	Failure is defined as: lack of efficacy with 2 week trial, allergy,	
	intolerable side effects, or significant drug-drug interactions. AND	
	• Member must not have recent (within 1 year) history of substance abuse	
	AND	
	• Member is not taking opioids, benzodiazepines, sedative hypnotics	
	(such as zolpidem, zaleplon, eszopiclone, chloral hydrate, etc.) or	
	consuming alcohol concomitantly with Xyrem (sodium oxybate)	
	AND	
	<ul> <li>Prescriber is enrolled in corresponding REMS program AND</li> </ul>	
	and failure of therapy with 3 sedative hypnotic medications (examples	
	include zolpidem and eszopiclone). Failure is defined as: lack of	
	efficacy with 2 week trial, allergy, intolerable side effects or significant	
	drug-drug interactions.	
	Initial and Continuation Prior Authorization Approval:	
	Initial prior authorization approval will be for 30 days. For continuation approval for	
	one year, the following information must be provided:	
	• Verification of Epworth Sleepiness Scale score reduction on follow-up OR	
	• Verification of cataplexy episode count reduction on follow-up	
	Maximum Desing:	
	Maximum Dosing: 9 grams/day	
	7 grams/uay	
XYWAV	<b>XYWAV</b> (calcium, magnesium, potassium, sodium oxybates) may be approved if the	Initial
(calcium, magnesium,	following criteria are met:	Approval:
potassium, sodium	• Member is $\geq$ 7 years of age AND	30 days
oxybates)	<ul> <li>Member has a diagnosis of excessive daytime sleepiness with narcolepsy</li> </ul>	Continuation
	(confirmed by one of the following):	Approval:
1	• Hypocretin deficiency OR	One year

ROGRAM APPENDICES	
<ul> <li>Nocturnal sleep polysomnography showing rapid eye movement (REM) sleep latency less than or equal to 15 minutes, or a Multiple Sleep Latency Test (MSLT) showing a mean sleep latency less than or equal to 8 minutes and two or more sleep-onset REM periods AND</li> <li>Baseline excessive daytime sleepiness is measured using the Epworth Sleepiness Scale or cataplexy episode count AND</li> <li>Member has adequately trialed and failed therapy with 3 stimulants for narcolepsy (examples include methylphenidate and amphetamine salts) Failure is defined as: lack of efficacy with 2 week trial, contraindication to therapy, allergy, intolerable side effects, or significant drug-drug interactions AND</li> <li>Member must not have recent (within 1 year) history of substance abuse AND</li> <li>Member is not taking opioids, benzodiazepines, sedative hypnotics (such as zolpidem, zaleplon, eszopiclone, chloral hydrate, etc.) or consuming alcohol while receiving Xywav (calcium, magnesium, potassium, sodium oxybates) therapy AND</li> <li>Prescriber is enrolled in corresponding REMS program AND</li> <li>If member is an adult (≥ 18 years of age), they have had an adequate trial and failure of therapy with 3 sedative hypnotic medications (examples include zolpidem and eszopiclone). Failure is defined as: lack of efficacy with 2 week trial, contraindication to therapy, intolerable side effects or significant drug-drug interactions.</li> </ul>	
<b>Yosprala</b> <sup>®</sup> will be approved for members who meet the following criteria:	One year
cerebrovascular events AND	
years of age or has documented history of gastric ulcers) AND	
• Member has failed treatment with three preferred proton pump inhibitors in the last 6 months (Failure is defined as: lack of efficacy of a seven-day trial, allergy, intolerable side effects, or significant drug-drug interaction.)	
	<ul> <li>Nocturnal sleep polysomnography showing rapid eye movement (REM) sleep latency less than or equal to 15 minutes, or a Multiple Sleep Latency Test (MSLT) showing a mean sleep latency less than or equal to 8 minutes and two or more sleep-onset REM periods AND</li> <li>Baseline excessive daytime sleepiness is measured using the Epworth Sleepiness Scale or cataplexy episode count AND</li> <li>Member has adequately trialed and failed therapy with 3 stimulants for narcolepsy (examples include methylphenidate and amphetamine salts) Failure is defined as: lack of efficacy with 2 week trial, contraindication to therapy, allergy, intolerable side effects, or significant drug-drug interactions AND</li> <li>Member must not have recent (within 1 year) history of substance abuse AND</li> <li>Member is not taking opioids, benzodiazepines, sedative hypnotics (such as zolpidem, zaleplon, eszopiclone, chloral hydrate, etc.) or consuming alcohol while receiving Xywav (calcium, magnesium, potassium, sodium oxybates) therapy AND</li> <li>Prescriber is enrolled in corresponding REMS program AND</li> <li>If member is an adult (≥ 18 years of age), they have had an adequate trial and failure of therapy with 3 sedative hypnotic medications (examples include zolpidem and eszopiclone). Failure is defined as: lack of efficacy with 2 week trial, contraindication to therapy, allergy, intolerable side effects or significant drug-drug interactions.</li> <li>Initial and Continuation Prior Authorization Approval:</li> <li>Nerification of Epworth Sleepiness Scale score reduction on follow-up OR</li> <li>Verification of Epworth Sleepiness Scale score reduction on follow-up OR</li> <li>Verification of approved for members who meet the following criteria:</li> <li>Member requires aspirin for secondary prevention of cardiovascular or cerebrovascular events AND</li> <li>Member requires appring for secondary prevention of acadiovascular or cerebrovascular events AND</li> <li>Member has f</li></ul>