

Pharmacy Prior Authorization Non-Formulary and Prior Authorization Guidelines

Scroll down to see PA Criteria by drug class, or Ctrl+F to search document by drug name

Non-preferred	Medication
Guideline	

Following criteria guidelines will be applied to all Non-preferred drugs. In addition, some drugs classes will have additional criteria that will apply. Please see drug specific guidelines.

Is there any reason the member cannot be changed to a preferred drug within the same class?

Acceptable reasons include:

- Allergy to preferred drug.
- Contraindication to or drug-to-drug interaction with preferred drug.
- History of unacceptable/toxic side effects preferred drug.
- Member's condition is clinically stable; changing to a preferred drug might cause deterioration of the member's condition.
- The requested drug may be approved if both of the following are true:
 - There has been a therapeutic failure of at least **two** preferred drugs **within the same class as appropriate for diagnosis unless otherwise noted in the clinical criteria**. A therapeutic failure of only one preferred drug is required when there is only one preferred drug within a therapeutic class.
 - The requested drug's corresponding generic (if a generic is available and covered by the State) has been attempted and failed or is contraindicated.

Initial Approval:

 Minimum of 3 months, depending on the diagnosis, to determine adherence, efficacy and patient safety monitoring

Renewal:

• Minimum of 6 months; up to 1 year

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Medications requiring Prior	Requests for Medications requiring Prior Authorization (PA) will be reviewed based on	As documented in the individual guideline
Authorization	the PA Guidelines/Criteria for that medication. Scroll down to view the PA Guidelines	
	for specific medications. Medications that do not have a specific PA guideline will	
	follow the Non-Preferred Medication Guideline. Additional information may be	
	required on a case-by-case basis to allow for adequate review.	
Medications requiring Step	Medications that require Step Therapy (ST) require trial and failure of formulary agents	Initial Approval:
Therapy	prior to their authorization. If the prerequisite medications have been filled within the	One year
	specified time frame, the prescription will automatically process at the pharmacy. Prior	
	Authorization will be required for prescriptions that do not process automatically at the	
	pharmacy.	
Quantity Level Limits	Requests that exceed established Quantity Level Limits will require prior authorization	Initial Approval:
	Drugs subject to additional utilization management requirements (for example, non-	One year
	formulary, clinical prior authorization, and step therapy) must meet clinical criteria and	B
	medical necessity for approval, in addition to any established Quantity Level Limit	Renewal Approval:
	Approval of Quantity Level Limit exceptions are considered after medication specific	One year
	prior authorization guideline and medical necessity review	
	Authorization Criteria for Quantity Limit Exceptions:	
	Quantities that Exceed Food and Drug Administration (FDA) Maximum Dose:	
	 Member is tolerating medication with no side effect, but had inadequate 	
	response at lower dose, and the inadequate response is not due to medication	
	non-adherence	

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	 Request meets one of the following: Dose is included in drug compendia or evidence-based clinical practice guidelines for same indication Published randomized, double blind, controlled trial, demonstrating safety and efficacy of requested dose is submitted with request Quantities that do not Exceed Food and Drug Administration (FDA) Maximum Dose (Dose Optimization): Request meets one of the following:	
Abilify Mycite	 Clinical Criteria for Abilify Mycite: Member is 18 years of age or older Member has tolerability to oral aripiprazole with suboptimal effects (as assessed by prescriber) that may be due to adherence problems Member has a smart phone compatible with the device 	Initial Approval: 3 months Renewals: 3 months Requires:

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- Member has given consent to a healthcare provider and caregiver (if applicable) to monitor the portal
- There will be documented intervention by prescriber if nonadherence is detected

In addition, clinical criteria for non-preferred agents:

- Must meet general non-preferred guideline
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs

- Member must:
- Continue to meet initial criteria
- Have prescriber attestation that member benefited from therapy
- Have prescriber attestation that there is a continued need for device (e.g., continued suboptimal effects and/or compliance)
- Have a healthcare provider and caregiver (if applicable) agree to continue to monitor device
- Not have worsened target symptoms
- Not have had any treatment-limited adverse effects (e.g., hypersensitivity, suicidality, neuroleptic malignant syndrome, tardive dyskinesia, metabolic changes, pathological gambling and other compulsive behaviors, orthostatic hypotension, falls, seizures, cognitive and motor impairment, dysphagia, disruption in body temperature regulation, and leukopenia, neutropenia, and agranulocytosis)
- Have a healthcare provider state reason why the member cannot use long-acting injectable atypical antipsychotic if there is continued nonadherence.

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Acne Agents, Topical	Clinical criteria for Dermatologic Acne agents:	Initial approval:
	 For members over the age of 18 years: Products are intended for acne only. Prior authorization for a cosmetic 	1 year
	indication cannot be approved	Renewal:
	In addition, clinical criteria for non-preferred agents:	1 year
	Must meet general non-preferred guideline:	1 year
	Had failure to respond to a therapeutic trial of at least two preferred	Requires:
	drugs.	Member is responding to treatment
Adbry	Clinical criteria for Adbry:	Initial Approval:
	Atopic Dermatitis	1 year
	 Member must have an FDA approved diagnosis: Atopic dermatitis that is moderate to severe Member is 12 years of age or older Prior documented trial and failure of 30-day trial of (or contraindication) of: One (1) topical corticosteroid of medium to high potency (for example, 	Renewals: 1 year Requires: Response to therapy
	mometasone, fluocinolone) and One (1) topical calcineurin inhibitor (tacrolimus or pimecrolimus) 	Quantity Level Limit: 4 syringes/14 days (initial dose); 4 syringes/28 days
Aemcolo	Clinical Criteria for Aemcolo:	Approval:
		3 days

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	 Diagnosis of travelers' diarrhea with moderate diarrhea that is distressing or interferes with planned activities Documentation of a history of failure, contraindication, or intolerance to one or more of the following: Azithromycin (generic Zithromax), Ciprofloxacin (generic Cipro), Levofloxacin (generic Levaquin), Ofloxacin (generic Floxin) 	Quantity Limit: 6 tablets per 3 days (dosing is 2 x 194mg [or 388mg] tablets twice daily for 3 days)
Analgesics Opioids – Long/Short- Acting All schedule II and III opiate	All opioids will be subject to a greater than or equal to 90 cumulative morphine milligram equivalent (MME) per day edit. This may require additional medical necessity. Prescribers shall order naloxone for any member with risk factors of substance use disorder, or daily morphine equivalent exceeding 90 mg per Virginia Board of Medicine	 Approvals: 3 months for chronic pain (includes HIV/AIDS, Chronic back pain, Arthritis, Fibromyalgia, Diabetic neuropathy,
narcotics	(BOM) regulations.	Postherpetic Neuralgia)
except Fentanyl Transmucosal Products, methadone	The General Authorization criteria is not required for members with pain associated with cancer, or in remission from cancer with a tapering plan safely weaning off opioids, palliative care (treatment of symptoms associated with life limiting-illnesses)	6 months for cancer pain, sickle cell disease, palliative care, hospice, and end- of-life care l
Tramadol	sickle cell disease, hospice care, or in a long-term care setting. Additional Prior Authorization criteria will still be required for non-preferred long-acting opioids and non-preferred short-acting opioids	Opioid Quantity Limits
Pentazocine	Horr-preferred short-acting opiolas	
	General Authorization Criteria for ALL opioids:	
	 Prescriber agrees to ALL of the following: Prescriber has checked the Virginia Prescription Monitoring Program (PMP); PMP website: (https://virginia.pmpaware.net/login) 	
	 Documents the morphine milligram equivalent (MME)/day and: For those with MME greater than or equal to 90 prescriber attests that he/she will be managing the member's opioid therapy long term, has 	

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reviewed the Virginia Board of Medicine (BOM) Regulations for Opioid Prescribing, has prescribed naloxone, and acknowledges the warnings associated with high dose opioid therapy including fatal overdose, and that therapy is medically necessary for this member

- Prescriber must agree to the following for history of benzodiazepine filled within the past 30 days:
 - Counseled member on the Food and Drug Administration (FDA) black box warning on the dangers of prescribing opioids and benzodiazepines including fatal overdose
 - Documented that treatment is medically necessary and has recorded a tapering plan to achieve the lowest possible effective dose of both opioids and benzodiazepines per the Virginia Board of Medicine Opioid Prescribing Regulations
- Naloxone been prescribed for members with risk factors of overdose. Risk factors
 include substance use disorder, doses in excess of 50 MME/day, antihistamines,
 antipsychotics, benzodiazepines, gabapentin, pregabalin, tricyclic
 antidepressants or the "Z" drugs (zopiclone, zolpidem, or zaleplon)
- For female members ages 18 45 years old, the prescriber has discussed the risk of neonatal abstinence syndrome and provided counseling on contraceptive options
- The prescriber has used at least one non-opioid therapy prior to consideration of an opioid (for example, oral NSAIDs, gabapentin, baclofen, capsaicin gel, duloxetine, lidocaine 5% patch, tricyclic antidepressants [nortriptyline], physical therapy, or cognitive behavioral therapy)

Additional Prior Authorization Criteria:

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Long-Acting Opioids

Documentation to support member meets the following:

- Diagnosis of one of the following:
 - Pain associated with cancer
 - Member is in remission from cancer with a plan to taper
 - Member is in palliative care, hospice, or a long-term care facility
 - Sickle cell disease

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- Diagnosis of chronic pain (related to fibromyalgia, diabetic neuropathy, arthritis, postherpetic neuralgia, HIV/AIDS, etc.) and
- For non-preferred long-acting opioids
 - Documentation to support an adequate trial and failure of TWO preferred formulary alternatives or contraindication to all of the agents (must include drug name, length of trial, and reason for discontinuation)

Short-Acting Opioids

Initial prescriptions for short-acting opiate containing medications will be allowed, up to a 7-day supply, without prior authorization. The member will be allowed one additional 7-day supply within 60 days of the original prescription fill date. Any additional prescriptions within 60 days from the fill date of the original prescription will require prior authorization.

Documentation to support member meets all of the following:

- Diagnosis of one of the following:
 - o Pain associated with cancer.

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	 Member is in remission from cancer with a plan to taper Member is in palliative care, hospice or a long-term care facility Sickle cell disease Or Diagnosis of chronic pain (related to fibromyalgia, diabetic neuropathy, arthritis, postherpetic neuralgia, HIV/AIDS, etc.) and For non-preferred short-acting opioids: Documentation to support an adequate trial and failure of TWO preferred short acting opioids or contraindication to all of the formulary short acting opioids (must include drug name, length of trial, and reason for discontinuation) 	
Anti-allergens and Palforzia	Clinical Criteria for Anti-allergens:	Anti-allergens –
3	Grastek:	Initial Approval:
Grastek	 Member has a diagnosis of grass pollen-induced allergic rhinitis with or 	1 year
Odactra	without conjunctivitis	
Oralair	Odactra:	Renewals:
Ragwitek	 Member has a diagnosis of house dust mite (HDM)-induced allergic rhinitis with or without conjunctivitis 	1 year
	Oralair:	Requires:
	 Member has a diagnosis of grass pollen-induced allergic rhinitis with or without conjunctivitis 	Member responding to therapy
	Ragwitek:	
	 Member has a diagnosis of short ragweed pollen-induced allergic rhinitis with or without conjunctivitis 	

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•	Documentation member has had a treatment failure with (or contraindication) to
	antihistamines (e.g., diphenhydramine, loratadine, etc.) and Montelukast/Singulair

Documentation of clinical reason why the member cannot use allergy shots

Clinical Criteria for Palforzia:

- Medication is being requested by or in consultation with an allergy or immunology specialist
- Member is between 4 and 17 years of age
- Member has a clinical history of allergy to peanuts or peanut-containing foods
- Physician verifies that they have reviewed the member's history and that the member is a candidate for Palforzia treatment following the REM requirements
- Palforzia will be initiated at a REMS-certified healthcare facility and the initial dose escalation phase and the first dose of each of the 11 up-dosing phases will be given at a REMS-certified healthcare facility

Palforzia -

Initial Approval:

6 months

Renewal:

12 months

Requires:

- Member meets initial criteria
- Member continues to tolerate the prescribed daily doses of Palforzia
- Member has not experienced recurrent asthma exacerbations
- Member has not experienced any treatment-restricting adverse effects (for example, repeated systemic allergic reaction and/or severe anaphylaxis)

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		Note: Members 18 years of age or older who met the initial approval criteria may continue
		maintenance treatment upon renewal
Anticonvulsants	Clinical criteria for Epidiolex:	Initial Approval:
	Member is 1 year of age or older	• 1 year
Preferred:	Member has a diagnosis of Epilepsy and recurrent seizures including Lennox-	
clobazam tab/susp	Gastaut syndrome (LGS), Dravet syndrome (DS), or tuberous sclerosis complex	Renewal:
clonazepam tab		• 1 year
Diastat rectal	Clinical Criteria for Fintepla®:	
Diastat AcuDial Rectal	Member is two years of age or older	Requires:
diazepam rectal & Device rectal	Member has a diagnosis of Dravet syndrome or Lennox-Gastaut syndrome	Ztalmy only: Documentation of positive clinical response as demonstrated by low
Epidiolex		disease activity and/or improvements in
	Clinical Criteria for Ztalmy:	the condition's signs and symptoms (e.g.,
Non-preferred:	Member is 2 years of age or older	reduced seizure activity, frequency,
clonazepam ODT	Prescribed by or in consultation with a neurologist, geneticist, or physician who	and/or duration)
clorazepate	specialized in treatment of epileptic disorders	 All others: Member is responding to
Fintepla	Documented diagnosis of cyclin-dependent kinase-like 5 deficiency disorder	treatment
Klonopin Tab	Documentation that seizures have been inadequately controlled by a trial of at least	
Onfi susp /tab	2 antiepileptic drugs (e.g., clobazam, valproate, lamotrigine, levetiracetam,	
Sympazan	topiramate, felbamate, vigabatrin) or member has labeled contraindications to	
Tranxene	other antiepileptic drugs	
Valtoco Nasal		
	Clinical Criteria for Valtoco:	
**Please refer to PDL for full	Not required to meet general non-preferred guideline if:	
list of preferred vs. non-	 Member has an FDA approved diagnosis; AND 	
preferred medications as	o Member is 6 – 11 years of age	

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the one here is not all-		
inclusive	Clinical Criteria for Non-Preferred Agents:	
	Must meet general non-preferred guideline	
	 Had failure to respond to a therapeutic trial of at least two preferred drugs 	
Antiemetic/Antivertigo	Clinical criteria for Non-Preferred Antiemetic Agents:	Approval duration for 5HT3 Receptor
Agents	Member has a diagnosis of one of the following: 1) severe, chemotherapy induced nausea and vomiting 2) nausea or vomiting related to radiation therapy, moderate	Blockers:
	to highly emetogenic chemotherapy, or post-operative nausea and vomiting 3)	Initial Approval:
	hyperemesis (i.e., pregnancy-related nausea/vomiting)	3 months, unless otherwise noted
	 Member has tried and failed therapeutic doses of, or has adverse effects or 	
	contraindications to, 2 different conventional antiemetics (e.g., promethazine,	Renewal:
	prochlorperazine, meclizine, metoclopramide, dexamethasone, etc.) Note: clinical evidence must be provided stating that the preferred agent(s) will not	3 months, unless otherwise noted
	provide adequate benefit, what pharmaceutical agents were attempted, and what outcomes were achieved;	Approval duration for Cannabinoids:
	OR	Initial approval:
	 Member has a diagnosis of diabetic gastroparesis Provider must specify why oral metoclopramide cannot be used; 	6 months
	OR	Renewal:
	 Diagnosis of AIDS-relating wasting (Dronabinol/Marinol/Syndros only) Member has tried and failed megestrol acetate oral suspension or has a 	6 months
	contraindication, intolerance, or drug-drug interaction o Marinol and Syndros require trial and failure of dronabinol	NK-1 Receptor Antagonists:
	OR	Initial Approval:
	For ondansetron 16mg ODT:	Length of chemotherapy regimen or a
	 Member must have tried and failed or been intolerant to ondansetron 8mg ODT OR 	maximum of 6 months

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	Must meet general non-preferred guideline: Had failure to respond to a therapeutic trial of at least two preferred drugs **Promethazine products require that member is 2 years of age or older	Renewal: Length of chemotherapy regimen or a maximum of 6 months Others: 1 year, unless otherwise noted Requires: Member is responding to treatment
Antimigraine	Clinical Criteria for Antimigraine Agents: • Preventive treatment of migraine (Aimovig, Ajovy, Ajovy autoinjector, Emgality pen/	Initial Approval: 6 months
Preferred: Aimovig Ajovy Ajovy autoinjector Emgality pen and syringe (120 mg) Nurtec ODT Qulipta Ubrelvy Non-preferred: Emgality syringe (100 mg) Reyvow Vyepti	 syringe (120 mg), Nurtec ODT, Emgality syringe (100 mg), Qulipta): Prescriber has assessed baseline disease severity utilizing an objective measure/tool (e.g., International Classification of Headache Disorders (ICHD-III); Headache Impact Test [HIT-6]; monthly headache day [MHD]; Migraine Disability Assessment [MIDAS]; Migraine Physical Function Impact Diary [MPFID])AND Member greater than or equal to 18 years of age; AND Member has greater than or equal to four migraine days per month for at least three months: AND Member has tried and failed a greater than or equal to 1 month trial of any 2 of the following oral generic medications: Antidepressants (for example, amitriptyline, venlafaxine) Beta blockers (for example, propranolol, metoprolol, timolol, atenolol) Anti-epileptics (for example, valproate, topiramate) 	Renewals: 12 months Requires: Vyepti: Member continues to meet initial criteria Member has absence of unacceptable toxicity from the drug Member experienced a clinical response as evidenced by: Reduction in mean monthly headache days (MHD) of at least moderate severity of greater than

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Zavzpret

- Angiotensin converting enzyme inhibitors/angiotensin II receptor blockers (for example, lisinopril, candesartan)
- o Non-preferred medications require trial and failure of 2 preferred agents
- Treatment of acute migraine (Nurtec ODT, Ubrelvy, Reyvow, Zavzpret):
 - Member has a diagnosis of migraine with or without aura; AND
 - Member greater than or equal to 18 years of age; AND
 - Member has tried and failed (or has contraindications to) two preferred triptan medication
 - o Non-preferred medications require trial and failure of 2 preferred agents
- Treatment of episodic cluster headaches (Emgality syringe (100 mg)):
 - o Member greater than or equal to 18 years of age; AND
 - Member experienced at least 2 cluster periods lasting from 7 days to 365 days, separated by pain-free periods lasting at least three months
 - Medication will not be used in combination with another CGRP antagonist or inhibitor used for the preventive treatment of migraines
 - Member tried and failed (or has contraindications to) at least one standard prophylactic (preventive) pharmacologic therapy for cluster headache
- Vyepti only:
 - o Member is 18 years of age or older
 - Prescriber has assessed baseline disease severity utilizing an objective measure/tool (e.g., International Classification of Headache Disorders (ICHD-III); Headache Impact Test [HIT-6]; monthly headache day [MHD]; Migraine Disability Assessment [MIDAS]; Migraine Physical Function Impact Diary [MPFID])
 - Member has been utilizing prophylactic intervention modalities (e.g., pharmacotherapy, behavioral therapy, physical therapy, etc.)

- or equal to 50% relative to the pretreatment baseline (diary documentation or medical professional attestation); OR
- A clinically meaningful improvement in ANY of the following validated migrainespecific member-reported outcome measures: 1) Reduction of greater than or equal to 5 points when baseline score is 11-20 OR Reduction of ≥30%when baseline score is >20 in the MIDAS (Migraine Disability Assessment) scores; OR 2) Reduction of greater than or equal to 5 points in the MPFID (Migraine Physical Function Impact Diary) score; OR 3) Reduction of greater than or equal to 5 points in the HIT-6 (Headache Impact Test) score

Others:

 Member demonstrated significant decrease in the number, frequency, and/or intensity of headaches

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- Member has a diagnosis of chronic migraines defined as 15 or more headache (tension-type-like and/or migraine-like) days per month for greater than 3 months
 - Member has had at least five attacks with features consistent with migraine (with and/or without aura); AND
 - On at least 8 days per month for greater than 3 months:
 - Headaches have characteristics and symptoms consistent with migraine; OR
 - Member suspected migraines are relieved by a triptan or ergot derivative medication; AND
 - Member has failed at least an 8-week trial of any two oral medications for the prevention of migraines (e.g antidepressants, beta blockers, antiepileptics) prior to initiation of eptinezumab; AND
 - Member had an inadequate response (or unable to tolerate) a minimum trial of at least two preferred self-injectable CGRP options

OR

- Member has a diagnosis of frequent episodic migraines defined as at least 5 headache attacks lasting 4-72 hours (when untreated or unsuccessfully treated)
 - Headaches have characteristics and symptoms consistent with migraine without aura; AND
 - Medication overuse headache has been ruled out by trial and failure of titrating off acute migraine treatments in the past
- Will not be used in combination with prophylactic calcitonin gene-related peptide (CGRP) inhibitors? (e.g., erenumab, galcanezumab, fremanezumab, atogepant, rimegepant, etc.)

Quantity Level Limits:

Prevention:

- Aimovig: 70 mg/mL autoinjector = 1 mL per month or 140 mg/mL autoinjector = 1 mL per month
- Ajovy: 1 Injection: 225 mg/1.5 mL singledose prefilled autoinjector month.
- Emgality: 120 mg/mL pen and syringe = 1 mL per month 100 mg/1 mL syringe = 1 mL per 30 days
- Nurtec® ODT: 16 tabs per 30 days
- Qulipta[™] 34 for 34-day supply

Acute treatment:

- Nurtec® ODT: 16 tabs per 30 days
- Reyvow®: 8 tabs per 30 days
- Ubrelvy®: 50mg or 100mg can have up to 16 tabs per strength per 30 days

Episodic cluster headache:

 Emgality Episodic cluster headache recommended dosage: 300 mg (administered as three consecutive injections of 100 mg each) at the onset of the cluster period, and then monthly until the end of the cluster period

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Antipsychotics In Children	Clinical criteria for antipsychotics in children less than 18 years of age:	Initial Approval:
Less Than 18 Years	 Prior authorization is required for all agents when prescribed for patients who are under 18 years of age (typical and atypical antipsychotic agents): Antipsychotic is being prescribed by, or in consultation with a Psychiatrist, Neurologist, or a Developmental/Behavioral Pediatrician. Documentation of a developmentally-appropriate, comprehensive psychiatric assessment with diagnoses, impairments, treatment target and treatment plans has been done. Patient had inadequate clinical response to a psychosocial treatment and psychosocial treatment with parental involvement will continue for the duration of medication therapy. Parent or guardian informed consent has been obtained for this medication. A family assessment has been done and includes parental psychopathology and treatment needs and evaluation for family functioning and parent-child relationship. In addition clinical criteria for non-preferred agents: 	1 year Renewal: 1 year Requires: • Member is responding to treatment
	 Must meet general non-preferred guideline Had failure to respond to a therapeutic trial of at least one preferred drug. 	
Attention Deficit Hyperactivity Disorder (ADHD) (non- stimulants/stimulants) medications	Preferred stimulants/Attention Deficit Hyperactivity Disorder (ADHD) medications for individuals age 4-17 years do not require prior authorization. Member must meet the minimum FDA-approved age. Non-preferred agents must meet age edit and non-preferred clinical criteria for approval. Age Edits and clinical criteria for Attention Deficit Hyperactivity Disorder (ADHD)	Approvals: • 1 year
	mediations:	

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Stimulants for children less than 4 years of age (does not apply to non-stimulant ADHD medications (such as atomoxetine, clonidine ER, Kapvay®, guanfacine ER, Intuniv®, Qelbree®, etc.)):

• The medication is being prescribed by a pediatric psychiatrist, pediatric neurologist, developmental/behavioral pediatrician, or in consultation with one of these specialists

Stimulants/ADHD medications for adults age 18 and older (does not apply to nonstimulant ADHD medications (such as atomoxetine, clonidine ER, Kapvay®, guanfacine ER, Intuniv®, Qelbree®, etc.)):

- Primary care provider has used the *Diagnostic and Statistical Manual of Mental Disorders*, 5th Edition and determined that criteria have been met (including documentation of impairment in more than 1 major setting) to make the diagnosis of Attention Deficit Hyperactivity Disorder (ADHD)
- The practitioner has regularly evaluated the member for stimulant or other substance use disorder, and, if present, initiated specific treatment, consulted with an appropriate health care provider, or referred the member for evaluation for treatment if indicated

Clinical Criteria for Vyvanse chewable tab:

• Member must have tried and failed methylphenidate solution

In addition, clinical criteria for non-preferred agents:

- Must meet general non-preferred guideline
 - Had failure to respond to a therapeutic trial of at least two preferred drugs (note: outcome of failed agents and pertinent information to support the use of the requested stimulant/ADHD medication must be provided)

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Buprenorphine Products	Authorization Criteria for INITIAL Treatment:	Initial approval:
	Note: oral buprenorphine products do not require PA if: 1) It is for a preferred product Suboxone® SL film or buprenorphine/naloxone tablets; 2) The member must be 16 years	• 3 months
	of age or older 3) The prescribed dose must be less than or equal to 24 mg/day	Renewal: • 6 months
	 Requests for plain buprenorphine monotherapy (without naloxone): will be approved if the member has a pregnancy confirmed by a positive laboratory test and the expected date of delivery (EDD) is provided (Buprenorphine mono-product will only be covered for pregnant women for a maximum of 10 months) Member is at least 16 years of age and diagnosed with Opioid Use Disorder using Diagnostic and Statistical Manual of Mental Disorders (DSM) 5: 	 10 months maximum duration for plain buprenorphine for pregnancy (10 months total, including initial authorization) Requires: Response to therapy
	 https://pcssnow.org/resource/opioid-use-disorder-opioid-addiction/ Non-preferred agents: documentation as to why the member cannot be prescribed a preferred agent. Include details and a completed FDA MedWatch Form (https://www.accessdata.fda.gov/scripts/medwatch/index.cfm) is required to be 	 Quantity Limits: buprenorphine/naloxone SL film 2 mg/0.5 mg; 3/day buprenorphine/naloxone SL film 4 mg/1
	 attached for adverse reactions to combination products The buprenorphine dose does not exceed 24 mg/day. Doses greater than 24 mg/day will not be approved 	 mg; 1/day buprenorphine/naloxone SL film 8 mg/2 mg; 3/day Zubsolv® SL tab 0.7 mg/0.18 mg; 2/day
		 Zubsolv® SL tab 1.4 mg/0.36 mg; 2/day Zubsolv® SL tab 2.9 mg/0.71 mg; 2/day Zubsolv® SL tab 5.7 mg/1.4 mg; 2/day Zubsolv® SL tab 8.6 mg/2.1 mg; 2/day Zubsolv® SL tab 11.4 mg/2.9 mg; 2/day
Cholestatic Pruritus Agents	Clinical Criteria for Cholestatic Pruritus Agents:	Approval: 12 months

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Bylvay Livmarli		Renewal: Member is responding to therapy
Cialis for Benign Prostatic Hypertrophy (BPH)	 Clinical criteria for Cialis 2.5mg and 5mg: Patient must try and fail (or have contraindications) to both Alpha Blockers (e.g. alfuzosin, tamsulosin) and Androgen Inhibitors (e.g. finasteride) for BPH and The prescriber must attest that the patient is not on the state list of sex offenders and The patient must have had a consult or been evaluated by a Urologist. 	Initial Approval: •1 year Renewal: •1 year Requires: •Patient is responding to treatment
Colony Stimulating Factors	 Clinical Criteria for Non-preferred Colony Stimulating Factors: Member has an FDA-approved indication and one of the following criteria is met 	Initial Approval: 6 months
Preferred: Fulphila Neupogen Disp Syrin Neupogen Vial	 The member's age is within FDA labeling for the requested indicalon for the requested agent The prescriber has provided information in support of using the requested agent for the member's age for the requested indication 	Renewal: 12 months
Non-preferred: Fylnetra Granix Syringe Granix Vial Leukine Neulasta Kit Neulasta Syringe Nivestym Syringe Nivestym Vial	Compendia allowed: DrugDex 1, 2a or 2b level of evidence, NCCN 1, 2a or 2b recommended use.	 Requires: Member continues to meet the initial criteria Member has an absence of unacceptable toxicity from the drug Member is being appropriately monitored for a beneficial response to therapy

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Nyvepria Releuko Syringe Releuko Vial Rolvedon Syringe Stimufend Syringe Udenyca Udenyca Autoinjector Udenyca Onbody Zarxio Ziextenzo Syringe	Clinical Edit for Cough and Cold Amonto	
Cough and Cold Products	 Clinical Edit for Cough and Cold Agents Patient is 6 years of age and older; AND Had failure to respond to a therapeutic trial of at least one preferred drug. Note: Children under the age of 6 years are not eligible for cough and cold products. 	Approval duration: •1 time (date of service)
Cytokine and CAM Antagonists and Related Agents Preferred: Enbrel Humira Infliximab (generic Remicade)	 Enbrel, Humira, and infliximab (generic Remicade) are preferred agents without PA. Non-preferred agents must meet drug specific criteria and general non-preferred criteria for approval. Clinical criteria for Actemra (tocilizumab) and Tyenne (tocilizumab-aazg): Diagnosis of moderately to severely active rheumatoid arthritis in adults, active polyarticular juvenile idiopathic arthritis (PJIA) in members 2 years of age or older, or active systemic juvenile idiopathic arthritis (SIJA) in member 2 years of age or older Trial and failure with methotrexate, requested medication will be used in conjunction with methotrexate, OR member has a contraindication to 	 Initial Approval: Initial: 3 months for Crohn's or Ulcerative Colitis; 1 year for all other indications Renewal: 1 year dependent upon medical records supporting response to therapy and review of Rx history Renewal for Kevzara and Siliq also require member is not receiving the medication in combination with any of the following: Biologic DMARD [for example, Humira (adalimumab), Cimzia (certolizumab), Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)] Janus kinase inhibitor [for example,

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Current Effective Date: 7/1/2025



- methotrexate (for example, alcohol abuse, cirrhosis, chronic liver disease, or other contraindication)
- Member has tried and failed another DMARD (other than methotrexate), such as azathioprine, d-penicillamine, cyclophosphamide, cyclosporine, gold salts, hydroxychloroquine, leflunomide, sulfasalazine, or tacrolimus
- o Had failure to respond to a therapeutic trial of at least two preferred drugs; OR
- Diagnosis of Cytokine Release Syndrome (Actemra only)
 - Member is 2 years of age or older with chimeric antigen receptor (CAR) T cellinduced severe or life-threatening cytokine release syndrome
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs; OR
- Diagnosis of Giant Cell Arteritis (GCA) in adults or Systemic Sclerosis-Associated Interstitial Lung Disease (SSc-ILD) to slow the rate of decline in pulmonary function (Actemra only)

Clinical criteria for Arcalyst (rilonacept):

- Diagnosis of Cryopyrin-Associated Periodic Syndromes (CAPS), including Familial Cold Auto-inflammatory Syndrome (FCAS) and Muckle-Wells Syndrome (MWS) in those 12 years of age or older
 - For those 18 and older:
 - Loading dose will be 320 mg, delivered as two 160 mg (2 mL) injections
 - Maintenance dose will be a 160 mg (2 mL) injection once weekly
 - o For those 12 to 17 years of age:
 - Loading dose will be 4.4 mg/kg, up to a maximum of 320 mg, delivered as 1 or 2 injections (up to 2 mL/injection)
 - Maintenance dose will be 2.2 mg/kg, up to a maximum of a 160 mg (2 mL) injection once weekly
 - \circ Had failure to respond to a therapeutic trial of at least two preferred drugs; OR

Xeljanz (tofacitinib)]

- Phosphodiesterase 4 (PDE4) inhibitor [for example Otezla (apremilast)]
- Renewal for Sotyktu: Documentation (e.g., progress note) of response to therapy compared to baseline, such as redness, thickness, scaliness, amount of surface area involvement, and/or PASI score.

Rasuvo/Otrexup:

Initial:

RA: 6 months

Psoriasis: 6 months

Quantity Limit = 4 auto-injectors per month

For renewal:

- Compliant and appropriate monitoring occurs
- Member must be followed by a physician for monitoring of renal and hepatic function and complete blood counts with differential and platelet count (Rasuvo only)

RA: 1 year

Psoriasis: 6 months

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- Maintenance of remission of deficiency of interleukin-1 receptor antagonist (DIRA) in adults and pediatric members weighing greater than or equal to 10 kg
 - Dosing will be 4.4mg/kg up to a maximum of 320 mg delivered as 1 or 2 subcutaneous injections once weekly
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Treatment of recurrent pericarditis (RP) and reduction in risk of recurrence in adults and children 12 years and older

Clinical criteria for Asvola (infliximab-axxq):

- Diagnosis of Crohn's disease, pediatric Cohn's disease, ulcerative colitis (reducing signs and symptoms, inducing, and maintaining clinical response), pediatric ulcerative colitis, rheumatoid arthritis in combination with methotrexate, ankylosing spondylitis, psoriatic arthritis, plague psoriasis
 - Had failure to respond to a therapeutic trial of at least two preferred drugs

Clinical criteria for Bimzelx (bimekizumab):

- Diagnosis of plaque psoriasis in adults
 - Must have a previous failure on a topical psoriasis agent and be a candidate for phototherapy or systemic therapy
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of active psoriatic arthritis, active ankylosing spondylitis, nonradiographic axial spondyloarthritis (nr-axSpA) with objective signs of inflammation, or moderate to severe hidradenitis suppurativa (HS)
 - Member is 18 years of age or older
 - Had failure to respond to a therapeutic trial of at least two preferred drugs

Clinical criteria for Cibingo (abrocitinib):

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- Diagnosis of refractory, moderate-to-severe atopic dermatitis in members 12 years of age or older
 - Prior documented trial and failure (or contraindication) of 1 topical corticosteroid of medium to high potency (for example, mometasone, fluocinolone) and 1 topical calcineurin inhibitor (tacrolimus or pimecrolimus)
 - Inadequate response to a 3-month minimum trial of at least 1 immunosuppressive systemic agent (for example, cyclosporine, azathioprine, methotrexate, mycophenolate mofetil, etc.)
 - Inadequate response (or is not a candidate) to a 3-month minimum trial of phototherapy (for example, psoralens with UVA light [PUVA], UVB, etc) provided member has reasonable access to photo treatment
 - Prescriber attestation that Cibinqo will not be used in combination with other
 JAK inhibitors, biologic immunomodulators, or with other immunosuppressants
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs

Clinical criteria for Cimzia (certolizumab):

- Diagnosis of moderately to severely active Crohn's Disease (reducing signs and symptoms, and maintaining clinical response) in adult members
 - Trial and failure of a compliant regimen of oral corticosteroids (moderate to severe CD) unless contraindicated or intravenous corticosteroids (severe and fulminant CD or failure to respond to oral corticosteroids)
 - Trial and failure of a compliant regimen of azathioprine or mercaptopurine for three consecutive months
 - Trial and failure of a compliant regimen of parenteral methotrexate for three consecutive months
 - $_{\odot}$ $\,$ Had failure to respond to a the rapeutic trial of at least two preferred drugs
- Diagnosis Moderately to severely active RA

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0	Trial and failure of, contraindication, or adverse reaction to methotrexate and at		
	least one other DMARD (sulfasalzine, hydroxychloroguine, minocycline)		

- o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of psoriatic arthritis
 - Trial and failure of methotrexate, requested medication will be used in conjunction with methotrexate, or member has a contraindication to methotrexate (for example, alcohol abuse, cirrhosis, chronic liver disease, or other contraindication)
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of ankylosing spondylitis
 - Trial and failure of an adequate trial of at least two NSAIDs or use of NSAIDs is contraindicated in the member
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of Active Non-radiographic Axial Spondyloarthritis (nr-axSpA)
 - o Member has objective signs of inflammation
 - Inadequate response, intolerance, or contraindication to at least two nonsteroidal anti-inflammatory drugs (NSAIDs)
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Treatment of adults with moderate-to-severe plaque psoriasis who are candidates for systemic therapy or phototherapy
- Treatment of active polyarticular juvenile idiopathic arthritis (pJIA) for patients greater than or equal to 2 years of age
 - Trial and failure of Methotrexate; OR requested medication will be used in conjunction with Methotrexate; OR member has a contraindication to Methotrexate
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs

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Clinical criteria for Cosentyx (secukinumab):

- Diagnosis of active enthesitis-related arthritis (ERA) in members 4 years of age and older
 - Trial and failure or failed an adequate trial of at least two NSAIDs; OR use of NSAIDs is contraindicated in member
- Diagnosis of Moderate to severe Plaque Psoriasis in adults and children 6 years of age and older who are candidates for systemic therapy or phototherapy
 - o Must have a previous failure on a topical psoriasis agent
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of active psoriatic arthritis in adults, active ankylosing spondylitis in adults, or adults with moderate to severe hidradenitis suppurativa (HS)
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of active non-radiographic axial spondyloarthritis (nr-axSpA) with objective signs of inflammation in adults

Clinical criteria for Enspryng (satralizumab-mwge):

- Diagnosis of Neuromyelitis optica spectrum disorder (NMOSD) in adult members who are anti-aquaporin-4 (AQP4) antibody positive (NMOSD)
 - Will be given as three 120 mg loading doses, administered at weeks 0, 2, and 4, with subsequent maintenance doses of 120 mg given every 4 weeks
 - o Member has a confirmed diagnosis based on the following:
 - Member was found to be seropositive for aquaporin-4 (AQP4) IgG antibodies: AND
 - Member has greater than or equal to 1 core clinical characteristic (for example, optic neuritis, acute myelitis, area postrema syndrome, acute brainstem syndrome, symptomatic narcolepsy or acute diencephalic clinical

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syndrome with NMOSD-typical diencephalic MRI lesions, symptomatic cerebral syndrome with NMOSD-typical brain lesions); AND

 Alternative diagnoses have been excluded (for example, multiple sclerosis, sarcoidosis, cancer, chronic infection);

Clinical criteria for Entyvio (vedolizumab):

- Diagnosis of moderately to severely active Crohn's disease or moderately to severely active UC in adults
 - Trial and failure of a compliant regimen of oral corticosteroids (moderate to severe Crohn's disease) unless contraindicated or intravenous corticosteroids (severe and fulminant Crohn's disease or failure to respond to oral corticosteroids)
 - Trial and failure of a compliant regimen of azathioprine or mercaptopurine for three consecutive months
 - Trial and failure of a compliant regimen of parenteral methotrexate for three consecutive months
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs

Clinical criteria for Eticovo (etanercept-ykro):

- Diagnosis of rheumatoid arthritis, juvenile idiopathic arthritis, psoriatic arthritis, or ankylosing spondylitis
 - o Member is 18 years of age or older
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of juvenile psoriatic arthritis
 - o Member is 2 years of age or older
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of plaque psoriasis (PsO)

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- o Member is 4 years of age or older
- o Had failure to respond to a therapeutic trial of at least two preferred drugs

Clinical criteria for Humira (adalimumab) biosimilars:

- Diagnosis of rheumatoid arthritis, juvenile idiopathic arthritis, psoriatic arthritis, ankylosing spondylitis, Crohn's disease, ulcerative colitis, plaque psoriasis, hidradenitis suppurativa, or uveitis
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs

Clinical criteria for Ilaris (canakinumabl):

- Diagnoses of the following require confirmation of the diagnosis and no trial of preferred agents:
 - Periodic Fever Syndromes
 - Cryopyrin-Associated Periodic Syndromes (CAPS), in adults and children 4 years of age and older including Familial Cold Autoinflammatory Syndrome (FCAS)
 - Muckle-Wells Syndrome (MWS)
 - Tumor Necrosis Factor Receptor Associated Periodic Syndrome (TRAPS) in adult and pediatric members
 - Hyperimmunoglobulin D Syndrome (HIDS)/Mevalonate Kinase Deficiency (MKD) in adult and pediatric members
 - o Familial Mediterranean Fever (FMF) in adult and pediatric members
- Diagnosis of Active Still's disease, including Adult-Onset Still's Disease (AOSD) or Active Systemic Juvenile Idiopathic Arthritis (SJIA) in members aged 2 years and older
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 12/1/2018, 2/4/19, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/24/2019, 11/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/20/2023, 3/24/2023, 3/30/2023, 4/6/2023, 4/15/2023, 4/20/2023, 5/15/2023, 7/1/2023, 7/6/2023, 7/20/2023, 8/10/2023, 8/17/2023, 8/17/2023, 8/17/2023, 10/1/2024, 1/1/2024, 1/1/2025, 4/1/2025



 Diagnosis of gout flares in adults in whom NSAIDs and colchicine are contraindicated, are not tolerated, or do not provide an adequate response, and in whom repeated courses of corticosteroids are not appropriate

Clinical criteria for Ilumya (tildrakizumab-asmn):

- Diagnosis of Moderate-to severe plaque psoriasis (PSO)
 - Have moderate to severe plaque psoriasis for at least 6 months and are candidates for systemic therapy or phototherapy with at least 1 of the following:
 - Involvement of at least 10% of body surface area (BSA)
 - Psoriasis Area and Severity Index (PASI) score of 10 or greater
 - Incapacitation due to plaque location (e.g., head and neck, palms, soles or genitalia)
 - Has not responded adequately (or is not a candidate) to a 3-month minimum trial of topical agents (for example, anthralin, coal tar preparations, corticosteroids, emollients, immunosuppressives, keratolytics, retinoic acid derivatives, and/or Vitamin D analogues)
 - Has not responded adequately (or is not a candidate) to a 3-month minimum trial of at least 1 systemic agent (for example Immunosuppressives, retinoic acid derivatives, and/or methotrexate)
 - Has not responded adequately (or is not a candidate) to a 3 month minimum trial of phototherapy (for example Psoralens with UVA light (PUVA) OR UVB with coal tar or dithranol)
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs

Clinical criteria for Jylamvo (methotrexate):

- Medication is used for one of the following diagnoses:
 - Treatment of adults and pediatric members with acute lymphoblastic leukemia (ALL) as part of a combination chemotherapy maintenance regimen

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- o Treatment of adults with mycosis fungoides
- o Treatment of adults with relapsed or refractory non-Hodgkin lymphoma as part of a metronomic combination regimen
- o Treatment of adults with rheumatoid arthritis
- Treatment of pediatric members with polyarticular juvenile idiopathic arthritis (pJIA)
- Treatment of adults with severe psoriasis
- Member meets one of the following:
 - Dosing will not allow the use of preferred methotrexate tablets
 - Member is unable to swallow methotrexate tablets

Clinical criteria for Kevzara (sarilumab):

- Diagnosis of moderately to severely active rheumatoid arthritis (RA) or polyarticular juvenile idiopathic arthritis (pJIA)
 - o RA: Member is 18 years of age or older
 - o pJIA: Member weighs 63kg or more
 - o Prescribed by or in consultation with a rheumatologist
 - History of failure, contraindication, or intolerance to one non-biologic disease modifying anti-rheumatic drug (DMARD) [for example, Rheumatrex /Trexall (methotrexate), Arava (leflunomide), Azulfidine (sulfasalazine)]
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of polymyalgia rheumatica (PMR)
 - o Member is 18 years of age or older
 - History of failure, contraindication, or intolerance to corticosteroids or member cannot tolerate a steroid taper
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 12/1/2019, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/24/2019, 1/1/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2024, 1/1/2024, 1/1/2024, 1/1/2025, 4/1/2025



Clinical criteria for Kineret (anakinra):

- Diagnosis Moderately to severely active RA to reduce the signs and symptoms and slow the progression of structural damage in members 18 years of age and older
 - o Trial and failure of, contraindication, or adverse reaction to methotrexate and at least one other DMARD (sulfasalzine, hydroxychloroquine, minocycline)
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of Deficiency of Interleukin-1 Receptor Antagonist (DIRA)
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of Cryopyrin-Associated Periodic Syndromes (CAPS), specifically Neonatal-Onset Multisystem Inflammatory Disease
 - Approvable with confirmation of this diagnosis and no trial of preferred agents required

Clinical criteria for Olumiant (baricitnib):

- Diagnosis of moderately to severely active rheumatoid arthritis (RA) in adults
 - Prescriber acknowledgement that use in combination with other JAK inhibitors, biologic disease-modifying antirheumatic drugs (DMARDs), or with potent immunosuppressants, such as azathioprine and cyclosporine, is not recommended
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs

Clinical criteria for Omvoh (mirikizumab-mrkz):

- Diagnosis of moderate to severe ulcerative colitis or moderate to severe Crohn's disease
 - Member is 18 years of age or older
 - O Had failure to respond to a therapeutic trial of at least two preferred drugs

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Clinical criteria for Orencia (abatacept):

- Moderately to severely active RA in adults
 - o Trial and failure of, contraindication, or adverse reaction to methotrexate and at least one other DMARD (sulfasalzine, hydroxychloroquine, minocycline)
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Active psoriatic arthritis (PsA) in members 2 years of age or older
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Moderate to severely active polyarticular juvenile Idiopathic Arthritis (JIA) in members 2 years and older
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Medication will be used for prophylaxis of acute graft versus host disease (aGVHD), in combination with a calcineurin inhibitor and methotrexate, in adults and pediatric members 2 years of age and older undergoing hematopoietic stem cell transplantation (HSCT) from a matched or 1 allele-mismatched unrelated donor

Clinical criteria for Otezla (apremilast):

- Diagnosis of active psoriatic arthritis in adults
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of moderate to severe plaque psoriasis in members 6 years of age or older weighing 20kg or more
 - Must have a previous failure on a topical psoriasis agent and be a candidate for phototherapy or systemic therapy
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Adult members with plaque psoriasis who are candidates for phototherapy or systemic therapy

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- Must have a previous failure on a topical psoriasis agent and be a candidate for phototherapy or systemic therapy
- o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Oral ulcers associated with Behcet's Disease in adults
 - No trial of preferred agents is required

Clinical criteria for Otrexup (methotrexate):

- Management of severe, active rheumatoid arthritis (RA)
 - o 18 Years of age or older, AND
 - Had an inadequate response, intolerance, or contraindication to NSAIDs, and oral methotrexate; AND
 - Must have allergy or contraindication to benzoyl alcohol or other preservative contained in generic injectable methotrexate
- Polyarticular juvenile idiopathic arthritis (pJIA), who are intolerant of or had an inadequate response to first-line therapy
 - o Has had therapeutic failure to two preferred NSAIDS agents; AND
 - Must have allergy or contraindication to benzoyl alcohol or other preservative contained in generic injectable methotrexate
- Symptomatic control of severe, recalcitrant, disabling psoriasis in adults who are not adequately responsive to other forms of therapy
 - A therapeutic trial and failure on topical therapies such as topical emollients and/or topical corticosteroids, topical retinoids, topical vitamin D analogs, and topical tacrolimus AND pimecrolimus; AND
 - Must have allergy or contraindication to benzoyl alcohol or other preservative contained in generic injectable methotrexate

Clinical criteria for Rasuvo (methotrexate):

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- Management of severe, active rheumatoid arthritis (RA)
 - o Has had therapeutic failure to two preferred DMARD agents
 - Must have allergy or contraindication to benzoyl alcohol or other preservative contained in generic injectable methotrexate
- Polyarticular juvenile idiopathic arthritis (pJIA), in members who are intolerant of or had an inadequate response to first-line therapy
 - Has had therapeutic failure to two preferred NSAID agents
 - Must have allergy or contraindication to benzoyl alcohol or other preservative contained in generic injectable methotrexate
- Symptomatic control of severe, recalcitrant, disabling psoriasis in adults who are not adequately responsive to other forms of therapy
 - A therapeutic trial and failure on topical therapies such as topical emollients and/or topical corticosteroids, topical retinoids, topical vitamin D analogs, and topical tacrolimus and pimecrolimus
 - Must have allergy or contraindication to benzoyl alcohol or other preservative contained in generic injectable methotrexate

Clinical criteria for Remicade and biosimilars (Avsola, Inflectra, Renflexis, Zymfentra):

- Diagnosis of Crohn's disease, pediatric Crohn's disease, ulcerative colitis, pediatric ulcerative colitis, Rheumatoid Arthritis in combination with methotrexate, Ankylosing Spondylitis, Psoriatic Arthritis, Plaque Psoriasis
 - Member is 18 years of age or older for all diagnoses except Crohn's disease and ulcerative colitis, for which member must be 6 years of age or older
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs

Clinical criteria for Rinvoq & Rinvoq LQ (upadacitinib):

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- Moderately to severely active rheumatoid arthritis in adults who have had an inadequate response or intolerance to one or more TNF blockers
- Members 2 years of age or older with active psoriatic arthritis who have had an inadequate response or intolerance to one or more TNF blockers
- Members 2 years of age and older with active polyarticular juvenile idiopathic arthritis (pJIA) who have had an inadequate response or intolerance to one or more TNF blockers
- Adults and pediatric members 12 years of age and older with refractory, moderate to severe atopic dermatitis whose disease is not adequately controlled with other systemic drug products, including biologics, or when use of those therapies is inadvisable
- Adults with active ankylosing spondylitis who have had an inadequate response or intolerance to one or more TNF blockers, or
- Adults with moderately to severely active ulcerative colitis who have had an inadequate response or intolerance to one or more TNF blockers
- Adults with moderately to severely active Crohn's Disease who have had an inadequate response or intolerance to one or more TNF blockers
- Diagnosis of active non-radiographic axial spondylarthritis in adults with objective signs of inflammation who have had an inadequate response or intolerance to TNF blocker therapy
- Diagnosis of giant cell arteritis in adults AND
- Prescriber acknowledgement that use in combination with other JAK inhibitors, biologic disease-modifying antirheumatic drugs (DMARDs), or with potent immunosuppressants, such as azathioprine and cyclosporine, is not recommended
- Had failure to respond to a therapeutic trial of at least two preferred drugs

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Clinical criteria for Siliq (brodalumab):

- Diagnosis of Psoriatic Arthritis (PsA) in adults who are candidates for systemic therapy or phototherapy and have failed to respond or have lost response to other systemic therapies
 - Dosing will be 210 mg of SQ (1 prefilled syringe) at Weeks 0, 1, and 2 followed by 210 mg every 2 weeks
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of moderate to severe plaque psoriasis in adults
 - Greater than or equal to 5% body surface area involvement, palmoplantar, facial, or genital involvement, or severe scalp psoriasis
 - History of failure, contraindication, or intolerance to both of the following conventional therapies:
 - Topical therapy with one of the following:
 - Corticosteroids (for example, betamethasone, clobetasol, desonide)
 - Vitamin D analogs (for example, calcitriol, calcipotriene)
 - Tazarotene
 - Calcineurin inhibitors (for example, tacrolimus, pimecrolimus)
 - Anthralin
 - Coal tar
 - Systemic therapy of at least 3 months duration with methotrexate
 - History of failure, contraindication, or intolerance to both of the following preferred biologic products (document drug, date, and duration of trial):
 - Humira (adalimumab)
 - Enbrel (etanercept)
 - Member is not receiving Siliq in combination with any of the following:

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•	Biologic DMARD [for example, Humira (adalimumab), Cimzia (certolizumal		
	Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)]		

- Janus kinase inhibitor [for example, Xeljanz (tofacitinib)]
- Phosphodiesterase 4 (PDE4) inhibitor [for example Otezla (apremilast)]
- Had failure to respond to a therapeutic trial of at least two preferred drugs

OR

- Member is currently on Siliq therapy
- o Member is not receiving Siliq in combination with any of the following:
 - Biologic DMARD [for example, Humira (adalimumab), Cimzia (certolizumab),
 Simponi (golimumab), Cosentyx (secukinumab), Orencia (abatacept)]
 - Janus kinase inhibitor [for example, Xeljanz (tofacitinib)]
 - Phosphodiesterase 4 (PDE4) inhibitor [for example Otezla (apremilast)]
- Had failure to respond to a therapeutic trial of at least two preferred drugs

Clinical criteria for Simponi (golimumab) and Simponi Aria (golimumab):

- Diagnosis of Moderately to severely active Rheumatoid Arthritis (RA) in adults
 - Trial and failure of, contraindication, or adverse reaction to methotrexate alone and at least one other DMARD (sulfasalazine, hydroxychloroquine, minocycline).
 - o Must be in combination with methotrexate
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of Active Psoriatic Arthritis (PsA) in adults, alone or in combination with methotrexate, or Active Ankylosing Spondylitis in adults
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of Moderately to severely active Ulcerative Colitis
 - Trial and failure of a compliant regimen of oral or rectal aminosalicylates (for example, sulfasalazine or mesalamine) for two consecutive months

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 12/1/2019, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/24/2019, 1/1/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2024, 1/1/2024, 1/1/2024, 1/1/2025, 4/1/2025



0	Trial and failure of a compliant regimen of oral corticosteroids (for moderate to
	severe CD) unless contraindicated, or intravenous corticosteroids (for severe
	and fulminant CD or failure to respond to oral corticosteroids)

- Trial and failure of a compliant regimen of azathioprine or mercaptopurine for three consecutive months
- O Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of moderately or severely active juvenile RA or juvenile idiopathic arthritis (Simponi Aria only)
 - Member has at least five swollen joints
 - Member has three or more joints with limitation of motion and pain, tenderness, or both
 - Member has had an inadequate response to one DMARD
 - Member is 2 years of age or older

Clinical criteria for Skyrizi (risankizumab-rzaa):

- Diagnosis of Moderate-to-severe plaque psoriasis (PSO) in adults
 - Diagnosis of moderate to severe plaque psoriasis for greater than or equal to 6 months with 1 or more of the following:
 - Affected body surface area (BSA) of 10% or more
 - Psoriasis Area and Severity Index (PASI) score 10 or more
 - Incapacitation due to plaque location (for example, head and neck, palms, soles or genitalia)
 - Member did not respond adequately (or is not a candidate) to a 3-month minimum trial of topical agents (for example, anthralin, coal tar preparations, corticosteroids, emollients, immunosuppressives, keratolytics, retinoic acid derivatives, and/or Vitamin D analogues)

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 12/1/2019, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 11/2020, 1/1/2020, 1/1/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/15/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/1/2023, 2/10/2023, 3/24/2023, 3/30/2023, 4/6/2023, 4/15/2023, 5/15/2023, 5/25/2023, 7/1/2023, 7/6/2023, 7/20/2023, 8/10/2023, 8/17/2023, 8/11/2024, 10/1/2024, 10/1/2024, 10/1/2024, 1/1/2025, 4/1/2025



0	Member did not respond adequately (or is not a candidate) to a 3-month
	minimum trial of at least 1 systemic agent (for example Immunosuppressives,
	retinoic acid derivatives, and/or methotrexate)

 Member did not respond adequately (or is not a candidate) to a 3-month minimum trial of phototherapy (for example, psoralens with UVA light (PUVA) or UVB with coal tar or dithranol)

OR

- Diagnosis of moderate to severe psoriatic arthritis
 - o Member is 18 years of age or older
 - Member did not respond adequately (or is not a candidate) to a 3-month minimum trial of greater than or equal to 1 systemic agent (e.g. Immunosuppressives, and/or methotrexate)

OR

- Diagnosis of moderate to severe Crohn's disease
 - o Member is 18 years of age or older
 - Trial and failure of a compliant regimen of oral corticosteroids unless contraindicated or intravenous corticosteroids

OR

- Diagnosis of moderate to severe ulcerative colitis
 - o Member is 18 years of age or older
 - Trial and failure to ONE conventional agent (i.e., 6-mercaptopurine, azathioprine, balsalazide, corticosteroids, cyclosporine, mesalamine, sulfasalazine) used in the treatment of UC after at least a 3-month duration of therapy

AND

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 12/1/2019, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/24/2019, 11/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/20/202



- Member is not receiving risankizumab-rzaa in combination with another biologic agent for the requested diagnosis or non-biologic immunomodulator (for example apremilast, tofacitinib, baricitinib)
- Had failure to respond to a therapeutic trial of at least two preferred drugs

Clinical criteria for Sotyktu (deucravacitinib):

- Diagnosis of moderate to severe plaque psoriasis; AND
- Prescribed by or in consultation with, a dermatologist, rheumatologist, or other specialist in the treatment of psoriasis; AND
- Symptoms persistent for greater than or equal to 6 months with at least 1 of the following:
 - o Involvement of at least 3% of body surface area (BSA); OR
 - o Psoriasis Area and Severity Index (PASI) score of 10 or greater; OR
 - Incapacitation due to plaque location (i.e., head and neck, palms, soles, or genitalia); AND
- Trial and failure (at least 3 months) of ≥ 1 conventional therapy:
 - o Disease-modifying anti-rheumatic drug (DMARD), such as methotrexate
 - o Immunosuppressant (e.g., cyclosporine)
 - o Oral retinoid (e.g., acitretin); AND
- Not used in combination with any other biologic agent; AND
- Trial and failure (at least 3 months) unless contraindication or intolerance to, ≥ 1
 preferred cytokine or CAM antagonist indicated for the treatment of this condition;
 AND
- Member must meet the minimum age recommended by the package insert for this FDA-approved indication

Clinical criteria for Spevigo (spesolimab):

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 12/1/2019, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/24/2019, 1/1/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2024, 1/1/2024, 1/1/2024, 1/1/2025, 4/1/2025



- Member is 12 years of age or older
- Member weighs 40kg or more
- Prescribed by or in consultation with dermatologist, rheumatologist, or other specialist in the treatment of psoriasis; AND
- Member has a known documented history of pustular psoriasis (GPP) flares (either relapsing [greater than 1 episode] or persistent [greater than 3 months])
- Had failure to respond to a therapeutic trial of at least two preferred drugs
- For members *with* flares related to GPP:
 - Member is presenting with primary, sterile, macroscopically visible pustules on non-acral skin (excluding cases where pustulation is restricted to psoriatic plaques); AND
 - o Member has at least one of the following documented:
 - IL36RN, CARD14, or AP1S3 gene mutation; or
 - Skin biopsy confirming presence of Kogoj's spongiform pustules; or
 - Systemic symptoms or laboratory abnormalities commonly associated with GPP flare (e.g., fever, asthenia, myalgia, elevated C-reactive protein [CRP], leukocytosis, neutrophilia [above ULN]); or
 - GPP flare of moderate-to-severe intensity (e.g., at least 5% body surface area is covered with erythema and the presence of pustules; Generalized Pustular Psoriasis Physician Global Assessment [GPPPGA] total score of greater or equal to 3).

Clinical criteria for Stelara (ustekinumab) and Biosimilars:

 Diagnosis of moderate to severe plaque psoriasis for adolescents (6 years of age and older) and adults who are candidates for phototherapy or systemic therapy, active psoriatic arthritis in adults, moderately to severely active Crohn's disease in adults, moderately to severely active ulcerative colitis in adults, or psoriatic arthritis in pediatric members 6 years of age or older

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 12/1/2019, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 11/2020, 1/1/2020, 1/1/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/15/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/1/2023, 2/10/2023, 3/24/2023, 3/30/2023, 4/6/2023, 4/15/2023, 5/15/2023, 5/25/2023, 7/1/2023, 7/6/2023, 7/20/2023, 8/10/2023, 8/17/2023, 8/11/2024, 10/1/2024, 10/1/2024, 10/1/2024, 1/1/2025, 4/1/2025



O Had failure to respond to a therapeutic trial of at least two preferred drugs

Clinical criteria for Taltz (ixekizumab):

- Diagnosis of moderate-to-severe plaque psoriasis in adolescents (6 years of age or older) and adults who are candidates for systemic therapy or phototherapy
 - Member has tried and failed at least 2 topical treatments, such as corticosteroids, calcipotriene, coal tar, tazarotene, or anthralin
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of active psoriatic arthritis in adults, active ankylosing spondylitis in adults, or active non-radiographic axial spondyloarthritis (nr-axSpA) with objective signs of inflammation in adults
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs

Clinical criteria for Tremfya (guselkumab):

- Diagnosis of moderate-to-severe plaque psoriasis in adults who are candidates for systemic therapy or phototherapy
 - Diagnosis has been present for greater than or equal to 6 months with 1 or more of the following:
 - Affected body surface area (BSA) of 10% or more
 - Psoriasis Area and Severity Index (PASI) score 10 or more
 - Incapacitation due to plaque location (for example, head and neck, palms, soles or genitalia)
 - Member did not respond adequately (or is not a candidate) to a 3-month minimum trial of topical agents (for example, anthralin, coal tar preparations, corticosteroids, emollients, immunosuppressives, keratolytics, retinoic acid derivatives, and/or Vitamin D analogues)

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 2/4/19, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/2/2019, 11/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/24/2023, 3/20/2023, 3/24/2023, 3/20/2023, 4/15/2023, 4/15/2023, 5/15/2023, 5/25/2023, 7/1/2023, 7/6/2023, 7/20/2023, 8/10/2023, 8/17/2023, 8/11/2024, 10/1/2024, 10/1/2024, 1/1/2025, 4/1/2025



- Member did not respond adequately (or is not a candidate) to a 3-month minimum trial of at least 1 systemic agent (for example Immunosuppressives, retinoic acid derivatives, and/or methotrexate)
- Member did not respond adequately (or is not a candidate) to a 3 month minimum trial of phototherapy (for example, psoralens with UVA light (PUVA) or UVB with coal tar or dithranol)
- Member is not receiving guselkumab in combination with another biologic agent for psoriasis or non-biologic immunomodulator (for example, apremilast, tofacitinib, baricitinib)
- o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of active psoriatic arthritis in adults
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs
- Diagnosis of moderately to severely active ulcerative colitis (UC) in adults
 - Trial and failure to ONE conventional agent (i.e., 6-mercaptopurine, azathioprine, balsalazide, corticosteroids, cyclosporine, mesalamine, sulfasalazine) used in the treatment of UC after at least a 3-month duration of therapy
 - Member is not receiving guselkumbab in combination with another biologic agent for ulcerative colitis or non-biologic immunomodulator (e.g., upadacitinib)
- Diagnosis of moderate to severe Crohn's disease
 - o Member is 18 years of age or older
 - Trial and failure of a compliant regimen of oral corticosteroids (unless contraindicated) or intravenous corticosteroids
 - o Member is not receiving guselkumab in combination with another biologic agent for Crohn's disease or non-biologic immunomodulator (e.g., upadacitinib)
 - o Had failure to respond to a therapeutic trial of at least two preferred drugs

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 2/4/19, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/2/2019, 12/2/2019, 11/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/24/2023, 3/20/2023, 3/24/2023, 3/30/2023, 4/6/2023, 4/15/2023, 5/15/2023, 5/25/2023, 7/1/2023, 7/6/2023, 7/20/2023, 8/10/2023, 8/17/2023, 8/17/2023, 9/14/2023, 10/1/2024, 1/1/2025, 4/1/2025



Clinical criteria for Trexall (methorexate):

• Had failure to respond to a therapeutic trial of at least two preferred drugs

Clinical criteria for Uplizna (inebilizumab-cdon):

- Diagnosis neuromyelitis optica spectrum disorder (NMOSD) in an adult member confirmed by blood serum test for anti-aquaporin- 4 antibody positive (AQP4-IgG)
 - Prescriber attests that member has been screened for hepatitis B virus (HBV) and tuberculosis (TB) prior to initiating treatment and member does not have an active infection
 - Prescriber attestation that member is not concomitantly receiving therapy with other immunosuppressant type drugs
 - Prescriber attestation that member will not be using in combination with complement-inhibitor (for example, eculizumab, ravulizumab) or anti-CD20directed antibody (for example, rituximab) therapies
 - Documentation history of: a) one or more relapses that required rescue therapy within the previous 12 months OR b) 2 or more relapses that required rescue therapy in 2 years prior to screening
 - Documentation that member has a baseline Expanded Disability Status Scale (EDSS) score less than or equal to 8
 - o Documentation of baseline relapse rate and visual acuity
 - Had failure to respond to a therapeutic trial of at least two preferred drugs

Clinical criteria for Xatmep (methorexate):

- Member is 12 years of age or older
- Dosing will not allow the use of preferred methotrexate tablets or member is unable to swallow methotrexate tablets

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 12/1/2019, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 11/2020, 1/1/2020, 1/1/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/15/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/1/2023, 2/10/2023, 3/24/2023, 3/30/2023, 4/6/2023, 4/15/2023, 5/15/2023, 5/25/2023, 7/1/2023, 7/6/2023, 7/20/2023, 8/10/2023, 8/17/2023, 8/11/2024, 10/1/2024, 10/1/2024, 10/1/2024, 1/1/2025, 4/1/2025



	 Clinical criteria for Xeljanz (tofacitinib) & Xeljanz XR (tofacitinib): Diagnosis of Moderate to severe active Rheumatoid Arthritis in adult members, psoriatic arthritis in adults (in combination with nonbiologic DMARDs), or Polyarticular Course Juvenile Idiopathic Arthritis (pcJIA) in members 2 years of age or older Trial and failure of, contraindication, or adverse reaction to methotrexate and at least one other DMARD (sulfasalazine, hydroxychloroquine, minocycline) Trial and failure or inadequate response or intolerant to TNF blockers Had failure to respond to a therapeutic trial of at least two preferred drugs Diagnosis of moderately to severely active ulcerative colitis in adults or active ankylosing spondylitis in adults Trial and failure or inadequate response or intolerant to TNF blockers Had failure to respond to a therapeutic trial of at least two preferred drugs 	
Dalfampridine ER	 Clinical Criteria for Dalfampridine ER: Diagnosis of multiple sclerosis with a gait disorder or difficulty walking Member does not have a history of seizures Member does not have moderate to severe renal impairment (Creatinine Clearance less than 50 mL/min) Baseline timed 25-foot walk test and date are submitted 	Initial Approval: 1 year Renewals: 1 year Requires: • Current timed 25-foot walk test and date are submitted
Daliresp (roflumilast)	Roflumilast is the preferred agent while Daliresp is non-preferred. Non-preferred agents must meet non-preferred clinical criteria for approval.	Initial Approval: 1 year

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 2/4/19, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/24/2019, 1/1/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/20/2023, 3/24/2023, 3/30/2023, 4/6/2023, 4/15/2023, 5/15/2023, 5/25/2023, 7/1/2023, 7/6/2023, 7/20/2023, 8/10/2023, 8/17/2023, 8/31/2023, 9/14/2023, 10/1/2024, 1/1/2024, 1/1/2025, 4/1/2025



	 Clinical criteria for Daliresp (roflumilast): If the member has a diagnosis of severe Chronic Obstructive Pulmonary Disease (COPD) associated with chronic bronchitis and a history of exacerbations Trial/failure on at least one first-line or second-line agent (inhaled anticholinergics, long-acting beta agonists or inhaled corticosteroids) Adjunctive therapy (Daliresp® must be used in conjunction with first-line or second-line agent) In addition, clinical criteria for non-preferred agents: Must meet general non-preferred guideline Had failure to respond to a therapeutic trial of at least two preferred drugs 	Renewals: 1 year Requires: Response to therapy
Diclegis & Bonjesta Preferred: Diclegis Non-preferred: Bonjesta doxylamine succinate/vit B6 (pyridoxine)	 Clinical criteria for Diclegis & Bonjesta: Member is pregnant and greater than or equal to 18 years of age Expected delivery date must be provided In addition, clinical criteria for non-preferred agents: Must meet general non-preferred guideline Had failure to respond to a therapeutic trial of at least two preferred drugs 	Approval: Duration of the pregnancy
Duchenne Muscular Dystrophy	Clinical Criteria for Agamree: Age requirements: Agamree: Member is 2 years of age or older	Initial approval: 12 months

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 2/4/19, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/24/2019, 1/1/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/20/2023, 3/24/2023, 3/30/2023, 4/6/2023, 4/15/2023, 5/15/2023, 5/25/2023, 7/1/2023, 7/6/2023, 7/20/2023, 8/10/2023, 8/17/2023, 8/31/2023, 9/14/2023, 10/1/2024, 1/1/2024, 1/1/2025, 4/1/2025



Preferred:	Duvyzat: Member is 6 years of age or older	Renewal:
Emflaza	Member has a diagnosis of Duchenne Muscular Dystrophy (DMD)	12 months
	Member has tried and failed or is intolerant to prednisone or prednisolone	
Non-preferred:	Member has tried and failed or is intolerant to Emflaza	Requires:
Agamree		 Antisense Oligonucleotides:
Amondys-45	Clinical Criteria for Antisense Oligonucleotides:	 Member continues to meet initial
Deflazacort	Amondys 45:	criteria
Duvyzat	 A confirmed mutation of the DMD gene that is amendable to exon 45 skipping 	 There is an absence of unacceptable
Exondys-51	• Exondys 51:	toxicity to the drug
Viltepso	 A confirmed mutation of the DMD gene that is amendable to exon 51 skipping 	 Member is being appropriately
Vyondys 53	Vyondys 53 or Viltepso:	monitored for a beneficial response to
	 A confirmed mutation of the DMD gene that is amendable to exon 53 skipping 	therapy
	Member has been on a stable dose of corticosteroids unless there is a	All others:
	contraindication or intolerance	 Member is responding to therapy
	The requested agent will be used as the only exon skipping therapy for the member's DMD	
	Clinical Criteria for Emflaza:	
	Brand Emflaza is preferred; use of the generic requires rationale for inability to use the	
i	brand	
	Member is 2 years of age or older	
	Member has a diagnosis for treatment of Duchenne muscular dystrophy (DMD)	
	Member has tried and failed or is intolerant to prednisone or prednisolone	
Dupixent	Clinical criteria for Dupixent:	Initial Approval:
	Asthma	1 year
	 Member is 6 years of age or older 	

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 2/4/19, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/24/2019, 1/1/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/20/2023, 3/24/2023, 3/30/2023, 4/6/2023, 4/15/2023, 5/15/2023, 5/25/2023, 7/1/2023, 7/6/2023, 7/20/2023, 8/10/2023, 8/17/2023, 8/31/2023, 9/14/2023, 10/1/2024, 1/1/2024, 1/1/2025, 4/1/2025



- o Diagnosis of Moderate to severe Asthma with
 - Eosinophilic phenotype with eosinophil count ≥ 150 cells/mcL; OR
 - Oral corticosteroid dependent asthma with at least 1 month of daily oral corticosteroid use within the last 3 months

• Eosinophilic Esophagitis (EoE);

- o Member is greater than or equal to 1 years old; AND
- Member weighs greater than or equal to 15 kg; AND
- o Prescribed by or consultation with an allergist or gastroenterologist; AND
- Member did not respond clinically to treatment with a topical glucocorticosteroid or proton pump inhibitor

• Atopic Dermatitis

- Member must have an FDA approved diagnosis: Atopic dermatitis that is moderate to severe
- o Member is 6 months of age or older
- o Prior documented trial & failure of 30-day trial (or contraindication) of:
 - One (1) topical corticosteroid of medium to high potency (e.g., mometasone, fluocinolone) OR
 - One (1) topical calcineurin inhibitors (tacrolimus or pimecrolimus)

• Chronic Rhinosinusitis with Nasal Polyposis

- o Member is 18 years of age or older
- Member has inadequate response after 3 consistent months use of preferred PDL intranasal steroids or oral corticosteroids; AND
- o Member is concurrently treated with intranasal corticosteroids

Renewals:

1 year

Requires:

- Member has experienced therapeutic benefit from the requested medication
- Member is free from toxicity

Quantity Level Limit:

Atopic Dermatitis – 2 syringes/pens for 14 days (initial dose), then 2 syringes/pens every 28 days

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	 Prurigo Nodularis Age 18 years of age or older Diagnosis of Prurigo Nodularis (PN) Inadequately Controlled Chronic Obstructive Pulmonary Disease (COPD) and an Eosinophilic Phenotype Member is 18 years of age or older Medication is prescribed by or in consultation with a pulmonologist Member has a diagnosis of COPD that is inadequately controlled and a minimum blood eosinophil count of 300 cells/mcL at screening, measured within the past 12 months Member is receiving maximal inhaled therapy consisting of a long-acting muscarinic antagonist (LAMA), long-acting beta agonist (LABA), and inhaled corticosteroid (ICS) (or therapy of LAMA plus LABA if ICS is contraindicated) Member has a history of at least 2 moderate (requiring treatment with systemic corticosteroids and/or antibiotics) or 1 severe exacerbation(s) (resulting in hospitalization or observation for over 24 hours in an emergency department or urgent care facility) in the previous year, with 1 exacerbation occurring while the patient was on maximal inhaled therapy 	
Ebglyss	 Clinical Criteria for Eblgyss: Member must have an FDA approved diagnosis: Atopic dermatitis that is moderate to severe Member is 12 years of age or older Prior documented trial & failure of 8 weeks of each: One (1) topical corticosteroid of medium to high potency (for example, mometasone, fluocinolone) and One (1) topical calcineurin inhibitors (tacrolimus or pimecrolimus) AND 	Initial Approval: 1 year Renewals: 1 year Requires: Response to therapy

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	A trial and failure of Dupixent®	
Enstilar Foam	Clinical Criteria for Enstilar Foam:	Initial Approval:
	Diagnosis of plaque psoriasis; AND	4 weeks
	Minimum age of 18 years; AND	
	In addition, clinical criteria for non-preferred agents:	Renewal:
	Must meet general non-preferred guideline	4 weeks
	 Had failure to respond to a therapeutic trial of at least two preferred drugs 	
Entresto Sprinkle	Clinical Criteria for Entresto Sprinkle Capsule:	Initial Approval:
	Member is intolerant to solid oral dosage forms	• 1 year
		Renewal:
		• 1 year
		1 year
		Requires:
		Member is responding to treatment
GI Motility agents	Clinical Criteria for GI Motility Agents:	Initial Approval:
ar would agone	Amitiza:	6 months
Preferred:	Must have one of the following diagnoses:	o monare
Linzess	Chronic idiopathic constipation (CIC)	Renewal Approval:
Lubiprostone	 Constipation Predominant Irritable Bowel Syndrome (IBS-C) 	6 months
Movantik	 Opioid induced constipation in chronic non-cancer pain (OIC) 	
	Member has tried and failed both polyethylene glycol AND lactulose	Requires:
Non-preferred:	 Treatment failure of at least ONE product from TWO of the following classes: 	Member is responding to treatment

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Alosetron	 Osmotic Laxatives (i.e., lactulose, polyethylene glycol (PEG), sorbitol)
Amitiza	■ Bulk Forming Laxatives (i.e., psyllium, fiber)
Lotronex	Stimulant Laxatives (i.e., bisacodyl, senna)
Ibsrela	
Motegrity	• Linzess:
Prucalopride	Must have one of the following diagnoses:
Relistor	 Chronic idiopathic constipation (CIC)
Symproic	 Functional constipation (FC) in pediatric patients 6 to 17 years of age and
Trulance	other causes of constipation have been ruled out (only 72mcg capsule can
Viberzi	be approved for this diagnosis)
	 Constipation predominant irritable bowel syndrome (IBS-C)
	 Treatment failure of at least ONE product from TWO of the following classes:
	 Osmotic Laxatives (i.e., lactulose, polyethylene glycol (PEG), sorbitol)
	 Bulk Forming Laxatives (i.e., psyllium, fiber)
	Stimulant Laxatives (i.e., bisacodyl, senna)
	Movantik, Relistor, & Symprioc:
	o Diagnosis of opioid induced constipation in chronic non-cancer pain
	Member has tried and failed both polyethylene glycol AND lactulose
	member has and and sear perjoury, one gry services
	Alosetron, Lotronex, & Viberzi
	 Diagnosis of severe diarrhea predominant irritable bowel syndrome (IBS-D)
	 Member has tried and failed at least three agents from the following classes
	(one from each class):
	 Bulk-forming laxatives (i.e., psyllium, fiber)
	 Antispasmodic agents (i.e., dicyclomine, hyoscyamine)
	 Antidiarrheal agents (i.e., loperamide, diphenoxaylate/atropine)

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- Motegrity, prucalopride:
 - o Diagnosis of chronic idiopathic constipation (CIC)
 - o Member has had treatment failure with both of the following:
 - Two or more preferred traditional laxative therapies (e.g., polyethylene glycol, lactulose)
 - One or more preferred newer products indicated for CIC (e.g., linaclotide, lubiprostone)
- Trulance:
 - Must have one of the following diagnoses:
 - Chronic idiopathic constipation (CIC)
 - Constipation predominant irritable bowel syndrome (IBS-C)
 - Treatment failure of at least ONE product from TWO of the following classes:
 - Osmotic Laxatives (i.e., lactulose, polyethylene glycol (PEG), sorbitol)
 - Bulk Forming Laxatives (i.e., psyllium, fiber)
 - Stimulant Laxatives (i.e., bisacodyl, senna)
- Ibsrela:
 - Diagnosis of constipation predominant irritable bowel syndrome (IBS-C
 - o Treatment failure of at least ONE product from TWO of the following classes:
 - Osmotic Laxatives (i.e., lactulose, polyethylene glycol (PEG), sorbitol)
 - Bulk Forming Laxatives (i.e., psyllium, fiber)
 - Stimulant Laxatives (i.e., bisacodyl, senna)

In addition, clinical criteria for non-preferred agents (excluding Motegrity):

Must meet general non-preferred guideline

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	 Had failure to respond to a therapeutic trial of at least two preferred drugs (if up to two preferred drugs are indicated) 	
GLP-1 RAs for	Clinical criteria for GLP-1 RAs for CV RR:	Initial approval:
Cardiovascular Risk Reduction	 Member is 45 years of age or older Medication is prescribed by a cardiologist or vascular specialist Member has a clinical history of one of the following: 	• 6 months
Wegovy SQ	 Myocardial infarction (MI) defined as cardiac biomarkers, an electrocardiogram or cardiac imaging Stroke defined as neurological dysfunction as a result of a hemorrhage or infarction Peripheral artery disease as defined by intermittent claudication with ankle-brachial index less than 0.85 at rest, or peripheral arterial revascularization procedure, or amputation due to atherosclerotic disease The member has not had a MI, stroke, transient ischemic attack or hospitalization for unstable angina in the last 60 days The member has a BMI ≥ 27 kg/m² The provider attests that the member received individualized healthy lifestyle counseling The member does not have a previous diagnosis of diabetes The member does not have pancreatitis, acute suicidal behavior/ideation, personal or family history of medullary thyroid cancer or multiple endocrine neoplasia 2 syndrome 	 Requires: The member continues to meet initial criteria The member is being treated with a maintenance dosage of the requested drug
GLP-1 RAs for Obstructive	Clinical criteria for GLP-1 RAs for OSA:	Initial approval:
Sleep Apnea	Member is 18 years of age or older	• 6 months

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Dysphoria	services, will be covered as specified below.	
•	normone therapy, genuer-armining normone therapy and associated taboratory	Official
GnRH Analogs for Gender	Medical (hormonal) therapy for gender dysphoria, including puberty suppressing hormone therapy, gender-affirming hormone therapy and associated laboratory	Initial Approval: 6 months
GnRH Analogs for Gender	 The member has a body mass index (BMI) of greater than or equal to 30kg/m² The member does not have craniofacial abnormalities that may affect breathing The member does not have diagnosis of central or mixed sleep apnea or Cheyne-Stokes respiration The member is not using any other GLP-1 product The member does not have pancreatitis, acute suicidal behavior/ideation, gastroparesis or using prokinetic drugs (i.e metoclopramide), personal or family history of medullary thyroid cancer or multiple endocrine neoplasia 2 syndrome Documentation requirements: Submission of polysomnography conducted within the last 12 months Submission of weight loss treatment plan within the past 6 months Medical (hormonal) therapy for gender dysphoria, including puberty suppressing 	Initial Approval:
Zepbound	 Must be prescribed by an otolaryngologist (ENT), neurologist, pulmonologist or sleep apnea specialist for the member to receive authorization The requesting provider is managing the member's obstructive sleep apnea The member has a diagnosis of moderate to severe obstructive sleep apnea (OSA) defined by an apnea-hypopnea index greater than or equal to 15 events/hour confirmed by polysomnography The member is currently on or has tried, failed or is unable to tolerate continuous positive airway pressure therapy (CPAP) (an adequate trial is defined as CPAP use for greater than or equal to 4 hours per night on greater than or equal to 70% of nights for two or more months) Note: documentation required if unable to tolerate CPAP therapy The member has a body mass index (RMI) of greater than or equal to 30kg/m² 	Renewal: • 12 months Requires: • The member continues to meet initial criteria

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Eligard	Puberty-suppressing and gender-affirming hormonal therapy for gender dysphoria	12 months
Supprelin LA	 is considered medically necessary when ALL of the following criteria are met: The member has been assessed and diagnosed with gender dysphoria according to DSM-V criteria, by one of the following provider types; and A licensed mental health provider; or If the member is over the age of 18, a gender dysphoria-informed hormone prescriber, as defined previously Medication is recommended and prescribed by, or in consultation with, an endocrinologist or other medical provider experienced in gender dysphoria hormone therapy; and Coexisting behavioral health and medical comorbidities or social problems that may interfere with diagnostic procedures or treatment are being appropriately treated and are not causing symptoms of gender dysphoria; and Member has experienced puberty development to at least Tanner stage 2 (stage 2 through 4) or has lab values for Luteinizing Hormone (LH), Follicle Stimulating Hormone (FSH), and the endogenous sex hormones consistent with at least Tanner stage 2; and The member has capacity to make informed treatment decisions and has assented to treatment after discussion of the potential benefits and risks. The process should include parental or legal guardian consent for unemancipated members under the age of 18. 	 Requires: Lab results to support response to treatment (for example, follicle-stimulating hormone (FSH), luteinizing hormone (LH), weight, height, tanner stage, bone age)
Growth Hormone	Preferred agents are Genotropin, Norditropin FlexPro, and Nutropin AQ NuSpin. Non-preferred agents must meet GH and non-preferred clinical criteria for approval.	Approval duration for PEDIATRIC Members (18 years of age and under):
Preferred:	profession agents made mode arranament profession ad announcement to approval.	Initial:
Genotropin Cartridge	Clinical Criteria for PEDIATRIC Members (18 years of age and under):	1 year
Genotropin Miniquick	Prescriber is an endocrinologist, nephrologist, other appropriate specially, or one	
Norditropin FlexPro	has been consulted on this case	Renewal:

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Non-preferred:

Humatrope cartridge

Humatrope vial

Ngenla

Nutropin AQ NuSpin

Omnitrope cartridge

Omnitrope vial

Saizen cartridge

Saizen vial

Serostim vial

Skytrofa

Zomacton vial

The member has open epiphysis and one of the following diagnoses:

- o Turner Syndrome
- o Prader-Willi Syndrome
- o Pediatric chronic kidney disease or renal insufficiency
- Small for gestational age (SGA)
- Idiopathic short stature
- Growth hormone deficiency
- Noonan Syndrome
- SHOX deficiency
- Familial short stature
- Documentation of the member's pretreatment age and height
- Pretreatment height is greater than or equal 2 SD (standard deviations) below average for the population mean height for age and gender
- Documentation showing one of the following:
- Pretreatment height velocity greater than or equal to 1 SD below the mean for age and gender
- At least 2 heights measured by an endocrinologist at least 6 months apart (data for at least 1 year) or at least 4 heights measured by a primary care physician at least 6 months apart (data for at least 2 years)
- For pediatric growth hormone deficiency:
 - o Member meets one of the following:
 - Documentation member had a growth hormone response of less than 10ng/mL (or otherwise abnormal as determined by the lab) of at least 2 GH stimulation tests
 - Documentation member had growth hormone response of less than 15 ng/mL on at least 1 GH stimulation test and a defined Central Nervous

1 year

Requires:

- Documentation showing growth velocity is least 2cm/year) while on growth hormone therapy
- Growth plates are open
- Documentation of member's current age and height

<u>Approval duration for adults (greater than 18 years of age):</u>

Initial:

1 year

Renewal:

1 year

Requires:

Member is responding to treatment

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System pathology, history of cranial irradiation, or genetic condition	
associated GH deficiency	

- Documentation member has both IGF-1 and IGFBP-3 levels below normal for age and gender
- Diagnosis of neonatal hypoglycemia with documentation of growth hormone level
- Member has at least 2 or more documented pituitary hormone deficiencies other than GH
- For pediatric chronic kidney disease or renal insufficiency:
 - Creatinine clearance of 75 mL/min/1.73m² or less, dialysis dependency, or serum creatinine greater than 3.0 g/dL

Clinical Criteria for ADULTS (Greater than 18 years of age):

- Prescriber is an endocrinologist
- Member does not have a defect in GH synthesis or irreversible hypothalamic/pituitary structural lesions or ablation
- Member meets one of the following:
- o GH deficiency diagnosed during childhood
- 3 or more pituitary hormone deficiencies and there is documentation the pretreatment IGF-1 level is below the laboratory's range of normal
- Member was retested after an at least 1-month break in GH therapy and GH peak level is provided
 - Insulin: less than or equal to 5 ng/ml
 - Glucagon: less than or equal to 3 ng/ml
 - Arginine: less than or equal to 0.4 ng/ml
 - Clonidine or Levadopa: not ideal agents for determining GH deficiency

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	 Diagnosis of growth hormone deficiency confirmed by growth hormone stimulation tests and rule-out of other hormonal deficiency, as follows: growth hormone response of fewer than five nanograms per mL to at least two provocative stimuli of growth hormone release: insulin, levodopa, L-Arginine, clonidine or glucagon when measured by polyclonal antibody (RIA) or fewer than 2.5 nanograms per mL when measured by monoclonal antibody (IRMA); AND Cause of growth hormone deficiency is Adult Onset Growth Hormone Deficiency (AO-GHD), alone or with multiple hormone deficiencies, such as hypopituitarism, as a result of hypothalamic or pituitary disease, radiation therapy, surgery or trauma; OR Other hormonal deficiencies (thyroid, cortisol or sex steroids) have been ruled out or stimulation testing would not produce a clinical response such as in a diagnosis of panhypopituitarism. 	
Hemangeol	Clinical criteria for Hemangeol:	Initial Approval:
	 Diagnosis treatment of proliferating infantile hemangioma requiring systemic therapy; AND 	•1 year
	Patient's age must be between 5weeks and 5 months.	Renewal:
		•1 year
		Requires: •Patient is responding to treatment
Hepatitis C Agents	Clinical Criteria for Direct-Acting Antivirals (DAAs) (EXCEPT Mavyret and	Approval duration:
	sofosbuvir/velpatasvir (generic Epclusa))	
Preferred: Mavyret,	Member is 12 years of age for ledipasvir/sofosbuvir (Harvoni) and 18 years of age or	<u>Initial</u> : 8 weeks (for all diagnoses)
Mavyret Pellet pack, and	older for all other agents	
sofosbuvir/velpatasvir	Prescriber must be a gastroenterologist, hepatologist, infectious disease specialist	Renewal Criteria
(generic Epclusa)	or transplant specialist or in consultation with one of the above	

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Epclusa® Harvoni® Ledipasvir/Sofosbuvir (generic Harvoni®) Olysio™ Pegasys® Proclick/syringe/kit/vial Sovaldi® Technivie™ Viekira Pak™ Viekira XR™ Vosevi™ Zepatier®	 Members must be evaluated for decompensated cirrhosis (which is defined as a Child-Pugh score greater than 6 [class B or C]) Members must be evaluated for severe renal impairment (eGFR <30 mL/min/1.73m2) or end stage renal disease (ESRD) requiring hemodialysis ***Note: Only non-preferred Hepatitis C Drugs require the submission of a prior authorization 	Member is compliant with drug therapy regimen (per pharmacy paid claims history)
Hereditary Angioedema	Preferred agents are Berinert, Cinryze, Icatibant, Kalbitor, and Sajazir. Non-preferred	Approval duration:
Agents (HAE)	agents must meet criteria for HAE agents and non-preferred agents for approval.	1 year
Preferred Berinert Cinryze Icatibant Kalbitor Sajazir Non-preferred Firazyr	 Clinical Criteria for Blood Modifiers: Must be prescribed by, or in consultation with, a specialist in allergy, immunology, hematology, pulmonology, or medical genetics Diagnosis has been confirmed by C1 inhibitor (C1-INh) deficiency or dysfunction (type I or II HAE) as documented by one of the following: C1-INh antigenic level below the lower limit of normal; OR C1-INh functional level below the lower limit of normal 	 Quantity Limits Cinryze – 20 vials per 34 days Haegarda – 2,000 IU SDV kit = 16 kits per 28 days; 3,000 IU SDV kit = 8 kits per 28 days Orladeyo – 34 capsules per 34 days Takhzyro – 2 vials per 28 days

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Haegarda	 For treatment of acute HAE attacks (Berinert, Firazyr, Icatibant, Kalbitor, Ruconest 	t,
Orladeyo	Sajazir): requested medication will be used as mono therapy to treat acute HAE	
Ruconest	attacks	
Takhzyro	For prophylaxis (Cinryze, Haegarda, Orladeyo, Takhzyro):	
	 Medication will be used solely for prophylaxis of HAE attacks 	
Hetlioz	Clinical Criteria for Hetlioz	Length of Authorizations: 6 months
	For the treatment of Non-24-Hour Sleep-Wake Disorder (Non-24), AND	
	Member must be age 18 years of age or older.	For Renewal: must document therapeutic
	Quantity limit = 1 capsule per day.	benefit and confirm compliance
	Clinical Criteria for Hetlioz LQ oral suspension	
	For the treatment Nighttime sleep disturbances in SMS in pediatric patients AN	D
	Mamber must be 2 years to 15 years of age	
	 Member must be 3 years to 15 years of age 	
	• Wember must be 3 years to 13 years of age	
Immunomodulators	Clinical Criteria for Verkazia:	Initial approval:
Immunomodulators	Clinical Criteria for Verkazia:	Initial approval: 1 year
Immunomodulators Preferred Restasis	Clinical Criteria for Verkazia: Patient is greater than or equal to 4 years of age	
<u>Preferred</u>	 Clinical Criteria for Verkazia: Patient is greater than or equal to 4 years of age Diagnosis of moderate to severe vernal keratoconjunctivitis Trial and failure, contraindication, or intolerance to one of the following: 	1 year
<u>Preferred</u>	 Clinical Criteria for Verkazia: Patient is greater than or equal to 4 years of age Diagnosis of moderate to severe vernal keratoconjunctivitis Trial and failure, contraindication, or intolerance to one of the following: 	1 year Renewals:
Preferred Restasis	 Clinical Criteria for Verkazia: Patient is greater than or equal to 4 years of age Diagnosis of moderate to severe vernal keratoconjunctivitis Trial and failure, contraindication, or intolerance to one of the following: Topical ophthalmic "dual-action" mast cell stabilizer and antihistamine (e.g., 	1 year Renewals:
Preferred Restasis	 Clinical Criteria for Verkazia: Patient is greater than or equal to 4 years of age Diagnosis of moderate to severe vernal keratoconjunctivitis Trial and failure, contraindication, or intolerance to one of the following: Topical ophthalmic "dual-action" mast cell stabilizer and antihistamine (e.g., olopatadine, azelastine) 	1 year Renewals: 1 year
Preferred Restasis Xiidra	 Clinical Criteria for Verkazia: Patient is greater than or equal to 4 years of age Diagnosis of moderate to severe vernal keratoconjunctivitis Trial and failure, contraindication, or intolerance to one of the following: Topical ophthalmic "dual-action" mast cell stabilizer and antihistamine (e.g., olopatadine, azelastine) Topical ophthalmic mast cell stabilizers (e.g., cromolyn) 	1 year Renewals: 1 year Requires:
Preferred Restasis Xiidra Non-preferred	 Clinical Criteria for Verkazia: Patient is greater than or equal to 4 years of age Diagnosis of moderate to severe vernal keratoconjunctivitis Trial and failure, contraindication, or intolerance to one of the following: Topical ophthalmic "dual-action" mast cell stabilizer and antihistamine (e.g., olopatadine, azelastine) Topical ophthalmic mast cell stabilizers (e.g., cromolyn) Prescribed by ophthalmologist or optometrist in consultation with an 	1 year Renewals: 1 year Requires:
Preferred Restasis Xiidra Non-preferred Cequa	 Clinical Criteria for Verkazia: Patient is greater than or equal to 4 years of age Diagnosis of moderate to severe vernal keratoconjunctivitis Trial and failure, contraindication, or intolerance to one of the following: Topical ophthalmic "dual-action" mast cell stabilizer and antihistamine (e.g., olopatadine, azelastine) Topical ophthalmic mast cell stabilizers (e.g., cromolyn) Prescribed by ophthalmologist or optometrist in consultation with an 	1 year Renewals: 1 year Requires: • Member is responding to treatment

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Restasis Multidose Tyrvaya Nasal Spray Verkazia	o Had failure to respond to a therapeutic trial of at least two preferred drugs	
Immunomodulators for	Clinical Criteria for Asthma Immunomodulators:	Initial approval:
Asthma	Age restrictions:	6 months
	o Cinqair: Member is 18 years of age or older	
Preferred:	o Tezspire: Member is 12 years of age or older	Renewals:
Fasenra pen	o Fasenra, Nucala, Xolair: Member is 6 years of age or older	1 year
Fasenra syringe	Member has a diagnosis of severe asthma	
Xolair Autoinjector	o Components of severity for classifying asthma as severe may include any of the	Requires:
Xolair syringe	following (not all-inclusive):	Asthma
Xolair vial	 Asthma that remains uncontrolled despite optimized treatment with high- dose ICS-LABA 	 Member has been assessed for toxicity Member has improvement in asthma
Non-preferred:	 Asthma that requires high-dose ICS-LABA to prevent it from being 	symptoms or asthma exacerbations as
Cinqair	uncontrolled	evidenced by decrease in one or more of
Nucala auto-injector	 Symptoms throughout the day 	the following:
Nucala syringe	 Nighttime awakenings, often 7 times/week 	 Use of systemic corticosteroids
Nucala vial	 SABA use for symptom control occurs several times per day 	 Hospitalizations
Tezspire pen	 Extremely limited normal activities 	o ER visits
Tezspire syringe	Lung function (percent predicted FEV1) < 60%	Unscheduled visits to healthcare
	 Exacerbations requiring oral systemic corticosteroids are generally more 	provider
	frequent and intense relative to moderate asthma	o Improvement from baseline in forced
	Member has an eosinophilic phenotype defined as blood eosinophils greater than	expiratory volume in 1 second (FEV1)
	or equal to 150 cells/μL (does not apply to Tezspire or Xolair)	5004
	Xolair only:	EGPA

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- o Member has a positive skin test or in vitro reactivity to a perennial aero-allergen
- o Member weighs between 20 kg (44 lbs.) and 150 kg (330 lbs.)
- Member has serum total IgE level, measured before the start of treatment, of either:
 - Greater than or equal to 30 IU/mL and less than or equal to 700 IU/mL in members age greater than or equal to 12 years; OR
 - Greater than or equal to 30 IU/mL and less than or equal to 1300 IU/mL in members age 6 to less than 12 years;
- Coadministration with another monoclonal antibody will be avoided (e.g., omalizumab, mepolizumab, reslizumab, benralizumab, dupilumab, tezepelumabekko)
- Will be used for add-on maintenance treatment in members regularly receiving
 both (unless otherwise contraindicated) of the following:
 - Medium- to high-dose inhaled corticosteroids; AND
 - An additional controller medication (e.g., long-acting beta agonist, leukotriene modifiers)
- Member has had two or more exacerbations in the previous year requiring oral or injectable corticosteroid treatment (in addition to the regular maintenance therapy defined above) or one exacerbation resulting in a hospitalization
- Member has at least one of the following for assessment of clinical status:
 - Use of systemic corticosteroids
 - Use of inhaled corticosteroids
 - Number of hospitalizations, ER visits, or unscheduled visits to healthcare provider due to condition
 - o Forced expiratory volume in 1 second (FEV₁)

- Member has been assessed for toxicity
- Member has disease response as indicated by improvement in signs and symptoms compared to baseline as evidenced in one or more of the following:
 - Member is in remission [defined as a Birmingham Vasculitis Activity Score (BVAS) score=0 and a prednisone/prednisolone daily dose of less than or equal to 7.5 mg]
 - Decrease in maintenance dose of systemic corticosteroids
 - Improvement in BVAS score compared to baseline
 - Improvement in asthma symptoms or asthma exacerbations
 - Improvement in duration of remission or decrease in the rate of relapses

HES

- Member has been assessed for toxicity
- Member has disease response as indicated by a decrease in HES flares from baseline

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- For non-preferred agents:
 - Member tried and failed an adequate trial of the 2 different preferred products;
 OR
 - Tezspire only:
 - Member lacks an eosinophilic phenotype with blood eosinophils greater than or equal to 150 cells/µL; AND
 - Member lacks a serum IgE level less than 30 IU/mL; OR
 - Member has another predicted intolerance the preferred agents (documentation must be included)

Clinical Criteria for Eosinophilic Granulomatosis with Polyangiitis (EGPA) [Nucala only]:

- Member is 18 years of age or older
- Member has a confirmed diagnosis of EGPA (aka Churg-Strauss Syndrome)
- Member has blood eosinophils greater than or equal to 1000 cells/ μL or greater than or equal to 10% eosinophils on white blood cell differential count
- Member has been on stable doses of concomitant oral corticosteroid therapy for at least 4 weeks (i.e.,prednisone or prednisolone at a dose of 7.5 mg/day)
- Physician has assessed baseline disease severity utilizing an objective measure/tool (e.g., BirminghamVasculitis Activity Score [BVAS], history of asthma symptoms and/or exacerbations, duration of remission,rate of relapses)
- Member tried and failed an adequate trial of the preferred product Fasenra

Clinical Criteria for Hypereosinophilic Syndrome (HES) [Nucala only]:

Member is 12 years of age or older

 Note: An HES flare is defined as worsening of clinical signs and symptoms of HES or increasing eosinophils (on at least 2 occasions), resulting in the need to increase oral corticosteroids or increase/add cytotoxic or immunosuppressive HES therapy

CRSwNP

- Member has been assessed for toxicity
- Member has disease response as indicated by improvement in signs and symptoms compared to baseline in one or more of the following: nasal/obstruction symptoms, improvement of sinus opacifications as assessed by CT-scans and/or an improvement on a disease activity scoring tool [e.g., nasal polyposis score (NPS), nasal congestion (NC) symptom severity score, sinonasal outcome test-22 (SNOT-22), etc.]
- Member had improvement in at least one of the following response criteria:
 - Reduction in nasal polyp size

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- Member has been diagnosed with HES (without an identifiable non-hematologic secondary cause (e.g., drug hypersensitivity, parasitic helminth infection, HIV infection, non-hematologic malignancy) or FIP1L1- PDGFRα kinase-positive HES) for at least 6 months prior to starting treatment
- Member has had a history of 2 or more HES flares within the previous 12 months (e.g., documented HES-related worsening of clinical symptoms or blood eosinophil counts requiring an escalation in therapy)
- Will be used in combination with stable doses of at least one other HES therapy,
 (e.g., oral corticosteroids, immunosuppressive agents, cytotoxic therapy) unless the member cannot tolerate other therapy

Clinical Criteria for Chronic Rhinosinusitis with Nasal Polyps (CRSwNP) [Nucala and Xolair only]:

- Member is 18 years of age or older
- Member failed at least 8 weeks of intranasal corticosteroid therapy
- Will be used in combination with intranasal corticosteroids unless member is unable to tolerate or use is contraindicated
- Xolair only:
 - Member has at least 3 of the following indicators for biologic treatment (note: members with a history of sino-nasal surgery are only required to have at least 3 of the indicators):
 - Patient has evidence of type 2 inflammation (e.g., tissue eosinophils greater than or equal to 10/hpf, blood eosinophils greater than or equal to 150 cells/μL, or total IgE greater than or equal to 100 IU/mL)

- Reduction in need for systemic corticosteroids
- o Improvement in quality of life
- Improvement in sense of smell
- Reduction of impact of comorbidities

CIU/CSU

- Member has been assessed for toxicity
- Member has a clinical improvement as documented an objective clinical evaluation tool? (e.g., UAS7, AAS, DLQI, AE-QoL, UCT, AECT, CU-Q2oL, etc.)

Food Allergy

- Member has been assessed for toxicity
 - Member is experiencing a clinical response and improvement as attested by the prescriber

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 12/1/2019, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/24/2019, 1/1/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2023, 3/20/2024, 1/1/2024, 1/1/2024, 1/1/2025, 4/1/2025



- Patient has required greater than or equal to 2 courses of systemic corticosteroids per year or greater than 3 months of low dose corticosteroids, unless contraindicated
- Disease significantly impairs the patient's quality of life
- Patient has experienced significant loss of smell
- Patient has a comorbid diagnosis of asthma
- o Member does not have any of the following:
 - Antrochoanal polyps
 - Nasal septal deviation that would occlude at least one nostril
 - Disease with lack of signs of type 2 inflammation
 - Cystic fibrosis
 - Mucoceles
- Other causes of nasal congestion/obstruction have been ruled out (e.g., acute sinusitis, nasal infection or upper respiratory infection, rhinitis medicamentosa, tumors, infections, granulomatosis)
- o Physician assessed baseline disease severity utilizing an objective measure/tool
- Nucala only:
 - Member has bilateral symptomatic sino-nasal polyposis with symptoms lasting at least 8 weeks
 - o Member tried and failed an adequate trial of the preferred product Xolair

<u>Clinical Criteria for Chronic Idiopathic Urticartia/Chronic Spontaneous Urticaria</u> (CIU/CSU) [Xolair only]:

- Member is 12 years of age or older
- Underlying cause of the patient's condition is NOT considered to be any other allergic condition(s) or other form(s) of urticaria

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 2/4/19, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/2/2019, 11/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/24/2023, 3/30/2023, 4/6/2023, 4/15/2023, 5/15/2023, 5/25/2023, 7/1/2023, 7/6/2023, 7/20/2023, 8/10/2023, 8/17/2023, 8/11/2024, 10/1/2024, 10/1/2024, 10/1/2024, 1/1/2025, 4/1/2025



- Member is avoiding triggers (e.g., NSAIDs, etc.)
- Documented baseline score from an objective clinical evaluation tool, such as: urticaria activity score (UAS7), angioedema activity score (AAS), Dermatology Life Quality Index (DLQI), Angioedema Quality of Life (AE-QoL), urticaria control test (UCT), angioedema control test (AECT), or Chronic Urticaria Quality of Life Questionnaire (CU-Q2oL)
- Member had an inadequate response to a one or more-month trial on previous therapy with scheduled dosing of a second-generation H1-antihistamine product
- Has the member had an inadequate response to a one or more-month trial on previous therapy with scheduled dosing of at least one of the following:
 - Up-dosing/dose advancement (up to 4-fold) of a second generation H1antihistamine
 - Add-on therapy with a leukotriene antagonist (e.g., montelukast, zafirlukast, etc.)
 - o Add-on therapy with another H1-antihistamine
 - o Add-on therapy with a H2-antagonist (e.g. ranitidine, famotidine, etc.)

Clinical Criteria for IgE-Mediated Food Allergy (Xolair only):

- Member is 1 year of age or older
- Prescribing physician is an allergist or immunologist or an allergist or immunologist has been consulted
- Member has a diagnosed food allergy as confirmed by:
 - o A positive skin prick test under a drop of allergen extract; OR
 - o A positive IgE screening (greater than or equal to kUA/L) to identified foods
- Member will continue to practice allergen avoidance

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 2/4/19, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/2/2019, 11/2020, 1/1/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/20/2023, 3/24/2023, 3/30/2023, 4/6/2023, 4/15/2023, 5/15/2023, 5/25/2023, 7/1/2023, 7/6/2023, 7/20/2023, 8/10/2023, 8/17/2023, 8/31/2023, 9/14/2023, 10/1/2024, 10/1/2024, 1/1/2025, 4/1/2025



Immunomodulators for Atopic Dermatitis

Clinical Criteria for Atopic Dermatitis Agents:

- Member must have an FDA approved diagnosis: Atopic dermatitis
- Elidel®: mild to moderate for ages greater than 2 years
- Protopic® 0.03%: moderate to severe for ages greater than 2 years
- Protopic® 0.1%: moderate to severe for ages greater than 16 years
- Zoryve cream, 0.15%: mild to moderate for ages greater than or equal to 6 years

Elidel,® pimecrolimus (AG), or tacrolimus:

 Prior documented trial & failure of 8 weeks of one (1) topical corticosteroid of medium to high potency (for example, mometasone, fluocinolone)

Eucrisa™:

- Eucrisa™: mild to moderate for ages equal to or greater than 3 months
- Member must have an FDA approved diagnosis: Atopic dermatitis
- Prior documented trial & failure of 30-day trial (or contraindication) of:
 - One (1) topical corticosteroid of medium to high potency (for example, mometasone, fluocinolone) OR
 - o One (1) topical calcineurin inhibitors (tacrolimus or pimecrolimus)

Opzelura:

- Opzelura[™]: for ages equal to or greater than 12 years
- Member must have an FDA approved diagnosis: Atopic dermatitis that is mild to moderate
- Prior documented trial & failure of 8 weeks of each:
 - One (1) topical corticosteroid of medium to high potency (for example, mometasone, fluocinolone) and
 - \circ One (1) topical calcineurin inhibitors (tacrolimus or pimecrolimus) AND

Initial Approval:

• 1 year

Renewal:

• 1 year

Requires:

• Member is responding to treatment

Quantity Limits

- Elidel 30gm per 30 days
- Eucrisa 100gm per 30 days
- Protopic 30 gm per 30 days
- Opzelura 240 gm (4 x 60gm) per 30 days

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 12/1/2018, 2/4/19, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 11/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 3/1/2021, 6/28/2021, 7/1/2021, 8/1/2021, 10/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/20/2023, 3/24/2023, 3/30/2023, 4/6/2023, 4/15/2023, 5/15/2023, 5/25/2023, 7/1/2023, 7/20/2023, 8/10/2023, 8/17/2023, 8/31/2023, 9/14/2023, 10/1/2024, 1/1/2024, 1/1/2024, 1/1/2025, 4/1/2025

Current Effective Date: 7/1/2025



Incretin Mimetics Preferred: Byetta Trulicity Victoza Non-preferred: Liraglutide Mounjaro Ozempic	 A trial and failure of Dupixent * Note: Opzelura cream can not be approved for the indication of nonsegmental vitiligo in adult and pediatric members greater than or equal to 12 years old Zoryve foam 0.3%: Member is 9 years of age or older and has a diagnosis of seborrheic dermatitis Clinical Criteria for Non-Preferred Incretin Mimetics: Member has a diagnosis of type 2 diabetes mellitus with documentation of the member's A1c value (must be greater than or equal to 6.5 for first starts) from within the last 12 months Non-preferred medications:	Approval: 12 months Renewal: Member is responding to therapy
Mounjaro Ozempic Rybelsus Tab Soliqua 100/33 Tanzeum Xultophy	**Preferred incretin mimetics require a diagnosis of type 2 diabetes	
Inhaled Antibiotics	Age requirements for Inhaled antibiotics:	Initial Approval:
Preferred Agents: Bethkis 300 mg/4 mL	Bethkis, Kitabis Pak, Tobi and Tobi Podhaler: Minimum age for use is 6 years for all tobramycin inhalation nebulizer solution	•1 year Renewal: •1 year

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 2/4/19, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/24/2019, 1/1/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/20/2023, 3/24/2023, 3/30/2023, 4/6/2023, 4/15/2023, 5/15/2023, 5/25/2023, 7/1/2023, 7/6/2023, 7/20/2023, 8/10/2023, 8/11/2023, 8/31/2023, 9/14/2023, 10/1/2024, 1/1/2024, 1/1/2025, 4/1/2025



Kitabis Pak 300 mg/5mL Tobi Podhaler tobramycin inhalation neb soln (generic Tobi® inh)

Cayston:

• Minimum age for use is 7 years

Clinical criteria for Bethkis, Kitabis pak:

• Member must have minimum age of 6 years

Clinical criteria for Tobi Podhaler:

- Member must have minimum age of 6 years AND
- Requires a clinical reason as to why one of the preferred tobramycin inhalation nebulizer solutions cannot be used

Clinical criteria for Arikayce

- Member is greater than or equal to 18 years of age; AND
- Diagnosis of Mycobacterium avium complex (MAC) lung disease as determined by the following:
 - chest radiography or high-resolution computed tomography (HRCT) scan;
 AND
 - o at least 2 positive sputum cultures; AND
 - other conditions such as tuberculosis and lung malignancy have been ruled out; AND
- Member has failed a multi-drug regimen with a macrolide (clarithromycin or azithromycin), rifampin, and ethambutol. (Failure is defined as continual positive sputum cultures for MAC while adhering to a multi-drug treatment regimen for a minimum duration of 6 months); AND
- Member has documented failure or intolerance to aerosolized administration of amikacin solution for injection, including pretreatment with a bronchodilator; AND

Requires:

Member is responding to treatment

Quantity Limitations:

Arikayce = 590 mg/8.4 mL (28 vials)/28 days (Each carton contains a 28-day supply of medication (28 vials))

Bethkis = 224mL (56 amps)/28 days

Cayston = 84ml/28 days Kitabis Pak = 280mL (56 amps)/28 days Tobi Podhaler = 224 capsule/28 day Tobi inhalation neb, generic tobramycin solution = 280mL (56 amps)/28 days

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 12/1/2019, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/24/2019, 1/1/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/20/2023, 3/24/2023, 3/30/2023, 4/6/2023, 4/15/2023, 5/15/2023, 5/25/2023, 7/1/2023, 7/6/2023, 7/20/2023, 8/10/2023, 8/11/2023, 8/11/2024, 1/1/2024, 1/1/2024, 1/1/2025, 4/1/2025



	 Arikayce will be prescribed in conjunction with a multi-drug antimycobacterial regimen Clinical criteria for Non-preferred Inhaled antibiotics: Minimum age for use is 6 years for all tobramycin inhalation nebulizer solution and 7 years for Cayston; AND Had failure to respond to a therapeutic trial of at least two preferred agents 	
Teriparatide & Tymlos – Injectable Osteoporosis	 Clinical Criteria for Teriparatide & Tymlos Member is 18 years of age or older Member has a confirmed diagnosis of osteoporosis Member has experienced a therapeutic failure or inadequate response to at least two bisphosphonates or member is unable to receive or has a contraindication to a bisphosphonate (Note: If unable to receive or these is a contraindication documentation as to why must be provided) Member will be taking calcium and vitamin D supplementation if dietary intake is inadequate One of the following: Member has a documented Hip DXA (femoral neck or total hip) or lumbar spine T-score -2.5 (standard deviations) or below and Bone Mineral Density (BMD) of -3 or worse Male members requiring increased bone mass with primary or hypogonadal osteoporosis must be at high risk of fracture (teriparatide only; Tymlos is not approved for this diagnosis) 	Approvals: 1 year Renewals require that member continues to meet the initial authorization criteria

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 2/4/19, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/24/2019, 1/1/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/20/2023, 3/24/2023, 3/30/2023, 4/6/2023, 4/15/2023, 5/15/2023, 5/25/2023, 7/1/2023, 7/6/2023, 7/20/2023, 8/10/2023, 8/17/2023, 8/31/2023, 9/14/2023, 10/1/2024, 1/1/2024, 1/1/2025, 4/1/2025



	 For postmenopausal women with a history of non-traumatic fractures two or more of the following risk factors: Family history of non-traumatic fracture(s) DXA BMD T-score ≤-2.5 at any site More than 2 alcohol beverages per day Glucocorticoid use (≥ 6 months of use at 7.5 dose of prednisolone equivalent) History of non-traumatic fracture(s) Rheumatoid Arthritis Current smoker Member is not at increased risk of osteosarcoma (for example, Paget's disease of bone, bone metastases or skeletal malignancies, etc.) Member has not received therapy with parathyroid hormone analogs (for example, teriparatide) in excess of 24 months in total 	
Journavx	 Clinical Criteria for Journavx: Member is 18 years of age or older Prescriber attest that the member has moderate to severe acute pain Member has tried and failed two of the following non-opioid therapies covered without prior authorization in the past 30 days (Must specify which agents): Diclofenac sodium gel Acetaminophen NSAIDs (oral) Lidocaine patch Other For members of childbearing potential and between 18 and 45 years old: prescriber has advised members using hormonal contraceptives containing progestins other than levonorgestrel and norethindrone to use an additional nonhormonal contraceptive or to use alternative contraceptives during Journavx treatment and for 	Approval: 14 days; may only be approved every 30 days

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 2/4/19, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/24/2019, 1/1/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/20/2023, 3/24/2023, 3/30/2023, 4/6/2023, 4/15/2023, 5/15/2023, 5/25/2023, 7/1/2023, 7/6/2023, 7/20/2023, 8/10/2023, 8/17/2023, 8/31/2023, 9/14/2023, 10/1/2024, 1/1/2024, 1/1/2025, 4/1/2025



Juxtapid	 28 days after discontinuation of Journavx Prescriber attests that the member is not pregnant, planning to become pregnant, or breastfeeding Clinical Criteria for Juxtapid: Member has a diagnosis of homozygous familial hypercholesterolemia (HoFH) Member is 18 years of age or older Provider is certified with the applicable REMS program Member has had a treatment failure, maximum dosing with, or contraindication to statins, ezetimibe, niacin, fibric acid derivatives, omega-3 agents, and bile acid sequestrants 	Approval: 1 year
Preferred: PEG 3350/electrolytes powder pack (AG) lactulose Solution Non-preferred: PEG 3350/electrolytes powder pack Kristalose Packet **See PDL for full list – this one is not all-inclusive	Clinical Criteria for Kristalose: Must have tried, failed, or be intolerant to generic lactulose	Initial Approval: 1 year Renewal: 1 year Requires: Member is responding to treatment

Previous Effective Date: 08/1/17 (02/1/17 PDL criteria), 12/1/17, 2/1/18, 3/1/18, 6/1/18, 7/1/18, 8/1/18, 10/1/2018, 12/1/2018, 2/4/19, 3/1/19, 4/1/19, 5/1/2019, 6/3/2019, 7/1/2019, 8/23/2019, 9/9/2019, 10/1/2019, 12/2/2019, 12/24/2019, 1/1/2020, 1/20/2020, 2/17/2020, 2/28/2020, 3/30/2020, 4/1/2020, 6/8/2020, 7/1/2020, 8/18/2020, 9/1/2020, 10/5/2020, 1/1/2021, 3/1/2021, 4/30/2021, 6/28/2021, 7/1/2021, 8/1/2021, 9/13/2021, 10/13/2021, 10/19/2021, 1/1/2022, 1/9/2022, 2/1/2022, 4/1/2022, 5/2/2022, 6/7/2022, 7/1/2022, 8/1/2022, 9/2/2022, 12/16/2022, 1/1/2023, 2/10/2023, 2/10/2023, 3/20/2023, 3/24/2023, 3/30/2023, 4/6/2023, 4/15/2023, 5/15/2023, 5/25/2023, 7/1/2023, 7/6/2023, 7/20/2023, 8/10/2023, 8/17/2023, 8/31/2023, 9/14/2023, 10/1/2024, 1/1/2024, 1/1/2025, 4/1/2025



Lucemyra	Clinical Criteria for Lucemyra:	Approval:
	Member is 18 years of age or older AND	3 days (to allow receipt of prescription)
	 Medication used for the mitigation of opioid withdrawal symptoms to facilitate 	
	abrupt opioid discontinuation	Quantity Level Limits:
	NOTE: PDL criteria does not apply	224 tablets per 180 days
Methadone	All opioids will be subject to a >/= 90 cumulative morphine milligram equivalent (MME)	Initial Approval:
	per day edit. This may require additional medical necessity. Prescribers should consider	6 months for chronic pain
	offering a prescription for naloxone and provide overdose prevention education; plus	Up to 1 years of age for infants discharged
	consider consultation with a pain specialist for MME/day exceeding 90. For 51 – 90	on methadone for neonatal abstinence
	MME/day prescriber should consider offering a prescription for naloxone and overdose	syndrome
	prevention education.	
		Renewals:
	The General Authorization criteria is not required for members with intractable pain	6 months for chronic pain
	associated with active cancer, palliative care (treatment of symptoms associated with	
	life limiting illnesses), or hospice care.	
		Requires:
	General Authorization Criteria:	Prescriber has reviewed and documented
	Prescriber agrees to ALL of the following:	information required from PMP
	 Prescribed by or in consultation with one of the following specialists: oncologist, 	UDS results (see criteria for specific
	sickle cell specialist, chronic pain specialist, or palliative care	requirements)
	 Prescriber has checked the Virginia Prescription Monitoring Program (PMP) on 	
	the date of the request to determine whether the member is receiving opioid	
	dosages or dangerous combinations (such as opioids and benzodiazepines) that	
	put them at high risk for fatal overdose	
	■ PMP website:	
	https://www.pmp.dhp.virginia.gov/VAPMPWebCenter/login.aspx	

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 Documents the MME/day and date of last opioid and benzodiazepine 	
filled	
■ For MME:	
 If 51 to 90 MME/day prescriber should consider offering a 	
prescription for naloxone and overdose prevention education	
o If greater than 90 MME/day prescriber should consider	
offering a prescription for naloxone and provide overdose	
prevention education; plus consider consultation with a pain	
specialist	
Note: Naloxone injection 0.4 mg/mL and 1 mg/mL vials and	
syringes and Narcan Nasal Spray (4 mg of naloxone	
hydrochloride/0.1 mL spray) are available without a	
service/prior authorization. Evzio requires a service	
authorization	
 Prescriber must agree to having counseled the member of the risks 	
associated with combined use of benzodiazepines and opioids if they	
will be given concomitantly	
 Prescriber attests that a treatment plan with goals that addresses benefits and 	
harm has been established with the member and the following bullets are	
included:	
 Established expected outcome and improvement in both pain relief 	
and function or just pain relief as well as limitations (for example,	
function may improve yet pain persist OR pain may never be totally	
eliminated)	
 Established goals for monitoring progress toward member-centered 	
functional goals (for example, walking the dog or walking around the	

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 There is a signed agreement with the member. A sample Physician/Patient Agreement may be found at: www.drugabuse.gov/sites/default/files/files/samplepatientagreementforms.pd f A presumptive urine drug screen (UDS) must be done at least annually. The UDS must check for the prescribed drug plus a minimum of 10 substances including heroin, prescription opioids, cocaine, marijuana, benzodiazepines, amphetamines, and metabolites. A copy of the most recent UDS must be submitted with the fax form. Member does not have a history of, or received treatment for, drug dependency or drug abuse Documentation to support an adequate 2-week trial and failure of ALL preferred formulary alternatives (for example, Oxymorphone ER, buprenorphine patch, fentanyl patch, and morphine sulfate ER) or contraindication to all of the agents (if contraindication to all agents must submit MEDWATCH form)
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	 Documentation showing whether or not the member is on any of the following concomitant therapies: single entity immediate release or extend release opioids, benzodiazepines, barbiturates, carisoprodol, meprobamate Note: methadone will only be approved in children discharged from the hospital (under 1 year of age; does not require prior authorization when a diagnosis of neonatal abstinence syndrome is submitted) and for those requiring around the clock analgesia i.e. chronic pain. Methadone is not covered under the pharmacy benefit for the treatment of opioid addiction. 	
Preferred Austedo tab Austedo XR tab Austedo XR titration pack Ingrezza cap Ingrezza Initiation Pack Ingrezza Sprinkle Tetrabenazine tab	Clinical Criteria for Movement Disorders: Diagnoses of Tardive Dyskinesia or Huntington's disease Prescribed by or in consult with a neurologist or psychiatrist	Initial approval: 1 year Renewals: 1 year Requires: • Member is responding to treatment Quantity limit:
Non-preferred Xenazine tab		 4 tabs/day Austedo 1 tab/day Austedo XR 42 tablets/365 days Austedo XR titration pack 1 cap/day Ingrezza cap/sprinkle 4 tabs/day Xenazine

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Multiple sclerosis (MS) Agents

Clinical criteria for preferred products and Kesimpta:

Preferred products may process through Auto-PA. For requests that don't pay use the criteria below.

- Member is greater than or equal to the age defined in the package insert; and
- Member has had a baseline magnetic resonance imaging (MRI) before initiating the first treatment course (within 3 months prior to start of therapy); and
- Member has been diagnosed with a relapsing form of multiple sclerosis (for example, relapsing remitting disease (RRMS), or active secondary progressive disease (SPMS), OR clinically isolated syndrome (CIS)) as indicated in the package insert; and
- For Kesimpta:
 - Member has tried and failed an injectable preferred product or dimethyl fumarate (generic Tecfidera)

Clinical criteria for non-preferred products without specific criteria listed below:

- Member is greater than or equal to the age defined in the package insert; and
- Member has had a baseline magnetic resonance imaging (MRI) before initiating the first treatment course (within 3 months prior to start of therapy); and
- Member has been diagnosed with a relapsing form of multiple sclerosis (for example, relapsing remitting disease (RRMS), or active secondary progressive disease (SPMS), OR clinically isolated syndrome (CIS)) as indicated in the package insert; and
- Member has tried and failed at least two preferred agents

Clinical criteria for Mavenclad:

• Member is greater than or equal to 18 years of age; and

Approval duration:

Initial:

• Briumvi: 6 months

Others: 1 year

Renewal: 1 year

- Briumvi & Ocrevus (including Zunovo):
 - Member continues to meet the relevant criteria identified in the initial criteria
 - Member has an absence of unacceptable toxicity from the drug
 - Member is being continuously monitored for response to therapy that indicates a beneficial response
- Others: Member is responding to treatment

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- Member has had a baseline magnetic resonance imaging (MRI) before initiating the first treatment course (within 3 months prior to start of therapy); and
- Member has been diagnosed with a relapsing form of multiple sclerosis (for example, relapsing remitting disease (RRMS) or active secondary progressive disease (SPMS)) as indicated in the package insert; and
- Lymphocyte count is greater than or equal to 800 cells per microliter prior to start of therapy; and
- Member does not have human immunodeficiency virus (HIV) infection; and
- Member has been tested for antibodies to the varicella zoster virus (VZV) or received immunization for VZV four weeks prior to beginning therapy; and
- Member has been screened for tuberculosis according to local guidelines; and
- Member has been evaluated and screened for the presence of hepatitis B and hepatitis C virus (HBV/HCV) prior to initiating treatment; and
- Member has tried and failed at least two preferred agents; and
- · Mavenclad will be used as single-agent therapy; and
- Prescriber attestation that women of childbearing age are not pregnant and that members of reproductive potential must use effective contraception during treatment with therapy and for at least six months after the last dose

Clinical criteria for Mayzent, Ponvory, and Zeposia:

- Member is greater than or equal to 18 years of age; and
- Member has had a baseline magnetic resonance imaging (MRI) before initiating the first treatment course (within 3 months prior to start of therapy); and
- Member has been diagnosed with a relapsing form of multiple sclerosis (for example, relapsing remitting disease (RRMS), or active secondary progressive disease (SPMS), OR clinically isolated syndrome (CIS)) as indicated in the package insert; and

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•	Member has obtained a baseline electrocardiogram	(ECG): and
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- Member has had a baseline ophthalmic evaluation of the fundus, including the macula, before starting treatment; and
- Member has been tested for antibodies to the varicella zoster virus (VZV) or received immunization for VZV four weeks prior to beginning therapy; and
- Member has been screened for tuberculosis according to local guidelines; and
- Member has been evaluated and screened for the presence of hepatitis B and hepatitis C virus (HBV/HCV) prior to initiating treatment; and
- Member has tried and failed at least two preferred agents; and
- Mayzent, Ponvory, Zeposia will be used as single-agent therapy; and
- Prescriber attestation that women of childbearing age are not pregnant and that members of reproductive potential must use effective contraception during treatment with therapy; and
- Prescriber attestation that member does **not** have any of the following:
 - o Recent myocardial infarction
 - o Unstable angina
 - Stroke
 - Transient ischemic attack
 - Decompensated heart failure with hospitalization
 - Class III/IV heart failure within the previous 6 months
 - o Prolonged QTc interval at baseline (> 500 msec)
 - CYP2C9*3/*3 genotype (Mayzent only)
 - History of Mobitz Type II second or third-degree atrioventricular block or sick sinus syndrome (unless treated with a functioning pacemaker)
- Additional criteria for Mayzent:
 - The member has been tested for CYP2C9 variant status to determine genotyping; and

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0	 Confirmation member will not be using Mayzent in combination with any of 		
	following:		

- Moderate or strong CYP3A4 inducers (e.g., modafinil, efavirenz) in members with a CYP2C9*1/*3 and CYP2C9*2/*3 genotypes; OR
- Drug regimens that contain CYP2C9/CY3A4 dual inhibitors (e.g., fluconazole); OR
- Moderate CYP2C9 inhibitor plus a moderate-to-strong CYP3A4 inhibitor; OR
- Other antineoplastic, immunosuppressive or immunomodulating drugs.

• Additional criteria for Zeposia:

- Confirmation that Zeposia will **not** be used in combination with the following:
 - Will not be initiating therapy after previous treatment with alemtuzumab; OR
 - Monoamine oxidase inhibitor (MAOI) (e.g., selegiline, phenelzine, linezolid);
 OR
 - Drugs known to prolong the QT-interval (e.g., fluoroquinolone or macrolide antibiotics, venlafaxine, fluoxetine, quetiapine, ziprasidone, sumatriptan, zolmitriptan); OR
 - Strong cytochrome p450 2C8 (CYP2C8) inhibitors (e.g., gemfibrozil) or inducers (e.g., rifampin); OR
 - BCRP inhibitors (e.g., cyclosporine, eltrombopag); OR
 - Adrenergic or serotonergic drugs which can increase norepinephrine or serotonin (e.g., opioids, selective serotonin reuptake inhibitors [SSRIs], selective norepinephrine reuptake inhibitors [SNRIs], tricyclics, tyramine); OR
 - Foods with large amounts of tyramine (e.g., > 150 mg), such as aged cheeses, cured meats, craft/unfiltered beers, beans); OR

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- Other antineoplastic, immunosuppressive or immunomodulating drugs (Note: if there is a history of prior use of these drugs, consider possible unintended additive immunosuppressive effects); AND
- Member will not receive live vaccines during and at least 4 weeks prior to and 12 weeks after treatment; AND
- Member does not have an active infection, including clinically important localized infections

Clinical criteria for Briumvi and Ocrevus:

- Member is at least 18 years of age
- Member has been screened for the presence of Hepatitis B virus (HBV) prior to initiating treatment AND does not have active disease (i.e., positive HBsAg and anti-HBV tests)
- Member's baseline serum immunoglobulin has been assessed
- Member will not receive live or live attenuated vaccines while on therapy or withing
 4 weeks prior to the initiation of treatment
- Member is free of an active infection.
- Medication will be used as a single therapy
- Member has not received a dose of ocrelizumab or ublituximab within the past 5 months

• Additional criteria for Briumvi:

- Member has a confirmed diagnosis of multiple sclerosis (MS) as documented by laboratory report (i.e., MRI)
- Member has a diagnosis of a relapsing form of MS [i.e., relapsing-remitting MS (RRMS), active secondary progressive disease (SPMS), or clinically isolated syndrome (CIS)]

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	 Additional criteria for Ocrevus: Member has a confirmed diagnosis of multiple sclerosis (MS) as documented by laboratory report (i.e., MRI); AND Member has a diagnosis of a relapsing form of MS [i.e., relapsing-remitting MS (RRMS), active secondary progressive disease (SPMS), or clinically isolated syndrome (CIS)]; OR Member has a diagnosis of primary progressive MS (PPMS), is less than 65 years of age, and has an expanded disability status scale (EDSS) score of less than or equal to 6.5 	
Preferred: Armodafinil Modafinil Sunosi Non-Preferred: Nuvigil Provigil Wakix	Stimulants are also preferred medications and include, but are not limited to: Adderall XR, amphetamine salts combo (generic for Adderall IR), and all methylphenidate IR generics. Clinical Criteria for Narcolepsy Medications: Member is 18 years of age or older Approvable diagnoses include: Narcolepsy: Documentation/confirmation of diagnosis via sleep study Excessive daytime sleepiness (EDS) in adult members with narcolepsy: Documentation/confirmation of diagnosis via sleep study Obstructive Sleep Apnea: Documentation/confirmation of diagnosis via sleep study Sudden onset of weak or paralyzed muscles (cataplexy) Shift Work Sleep disorder: Documentation showing current shift schedule Symptoms do not occur during the course of another sleep disorder or mental disorder and are not due to the direct physiological effects of a medication or a general medical condition	Initial approval: 1 year Renewals: 1 year Requires: Member continues to meet initial criteria Member reports a reduction in excessive daytime sleepiness from pre-treatment baseline Member has not experienced any treatment-related adverse effects

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- Sunosi only:
 - o Member has tried and failed either modafinil or armodafinil
- Wakix only:
 - Member has an International Classification of Sleep Disorders (ICSD-3) or Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition (DSM-5) diagnosis of narcolepsy
 - Member has a baseline daytime sleepiness as measured by a validated scale (e.g., Epworth Sleepiness Scale, Stanford Sleepiness Scale, Karolinska Sleepiness Scale, Cleveland Adolescent Sleepiness Questionnaire, or a Visual Analog Scale)
 - A mean sleep latency of less than or equal to 8 minutes AND greater than or equal to 2 sleep onset REM periods (SOREMPs) are found on a mean sleep latency test (MSLT) performed according to standard techniques (A SOREMP [within 15 minutes of sleep onset] on the preceding nocturnal polysomnogram may replace one of the SOREMPs on the MSLT)
 - Either cerebrospinal fluid (CSF) hypocretin-1 concentration has not been measured OR CSF hypocretin-1 concentration measured by immunoreactivity is either greater than 110 pg/mL OR greater than 1/3 of mean values obtained in normal subjects with the same standardized assay
 - The hypersomnolence and/or MSLT findings are not better explained by other causes such as insufficient sleep, obstructive sleep apnea, delayed sleep phase disorder, or the effect of medication or substances or their withdrawal
 - Member has daily periods of irrepressible need to sleep or daytime lapses into sleep occurring for greater than or equal to 3 months
 - Member must not be receiving treatment with sedative hypnotic agents (e.g., zolpidem, eszopiclone, zaleplon, benzodiazepines, barbiturates)
 - Member will not use histamine-1 (H1) receptor antagonists (e.g., pheniramine maleate, diphenhydramine, promethazine, imipramine, clomipramine, mirtazapine) concomitantly

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	 Member does not have a history of prolonged QTc interval (e.g., QTc interval greater than 450 milliseconds) Therapy will not be used in members with severe hepatic impairment (Child-Pugh C) Member does not have end stage renal disease (ESRD) (e.g., eGFR less than 15 mL/minute/1.73 m2) Brand Nuvigil and Provigil only: Member tried and failed the preferred generics for the requested products 	
	In addition, clinical criteria for non-preferred agents:	
	Must meet general non-preferred guideline	
	 Had failure to respond to a therapeutic trial of at least two preferred drugs 	
	NOTE: Sunosi is indicated only for narcolepsy and obstructive sleep apnea (OSA). Wakix is approved only for excessive daytime sleepiness or sudden onset of weak or paralyzed muscles (cataplexy) in patients with narcolepsy. Provigil (modafinil) and Nuvigil (Armodafinil) are indicated for narcolepsy-related excessive daytime sleepiness, OSA, and shift work sleep disorder.	
Nemluvio	Clinical Criteria for Nemluvio:	Initial Approval:
	Atopic Dermatitis:	1 year
	Member must have an FDA approved diagnosis: Atopic dermatitis that is	
	moderate to severe	Renewals:
	Member is 12 years of age or older	1 year
	o Prior documented trial & failure of 8 weeks of each:	Requires:
	 One (1) topical corticosteroid of medium to high potency (for example, mometasone, fluocinolone) and 	Response to therapy
	 One (1) topical calcineurin inhibitors (tacrolimus or pimecrolimus) AND 	

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	 A trial and failure of Dupixent® Prurigo nodularis: Member is 18 years of age or older Member has a diagnosis of prurigo nodularis Member had an 8-week trial and failure of Dupixent 	
Non-preferred Antibiotics –	Clinical Criteria for Cephalosporins, Macrolides, Ketolides, and Quinolones:	Approval duration:
Cephalosporins	 Infection caused by an organism resistant to preferred drugs, OR 	
Macrolides, Ketolides, and	 A therapeutic failure to no less than a three-day trial of <u>one preferred drug</u> 	Date of service only; no refills.
Quinolones	 within the same class; OR The member is completing a course of therapy with a non-preferred drug which was initiated in the hospital. 	
Non-preferred Steroids	Clinical Criteria for non-preferred steroids:	Approval duration:
Sernivo	 Must meet general non-preferred guideline Had failure to respond to a therapeutic trial of no less than a one-month trial of at least at least two preferred drugs within the same class. Clinical Criteria for Sernivo: Minimum age restriction of 18 years of age; AND 	Sernivo: • 4 weeks (Treatment beyond 4 weeks is not recommended.)
	 Indicated for the treatment of mild to moderate plaque psoriasis; AND A therapeutic failure of at least TWO preferred drugs within the same class. 	Others: Initial/renewal duration: 1 year Renewal requires: Patient is responding to treatment

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Nuplazid	 Clinical Criteria for Nuplazid: Member is 18 years or older Indicated for the treatment of hallucinations and delusions associated with Parkinson's disease psychosis. 	Initial Approval: 1 year Renewal: 1 year Requires: Patient is responding to treatment Quantity Limit = 2 per day
Oral Antifungals	Clinical criteria for non-preferred oral antifungal agents:	Initial Approval:
	Documentation member has tried and failed two preferred oral antifungals	Duration of the prescription (up to 12 months)
Preferred:	OR	
fluconazole tab/susp	Documentation member has contraindications or intolerances to preferred agents	Renewal:
griseofulvin ^{susp}	or member has a diagnosis for which none of the preferred oral antifungals are	1 year
nystatin tab/susp	indicated or widely medically-accepted such as, but not limited to:	
terbinafine	o aspergillosis	Requires:
	o blastomycosis	Patient is responding to treatment
Non-Preferred:	o coccidioidomycosis	
Ancobon	o cryptococcosis	
Clotrimazole (mucous mem)	o febrile neutropenia	
Cresemba	o fungal infection caused by S. apiospermum or Fusarium species, including F.	
Diflucan tab/susp	solani	
flucytosine	o histoplasmosis	
Gris-Peg	o mucormycosis	
griseofulvin	Documentation of the member's diagnosis and planned duration of treatment must	
tab/ultramicrosize	be submitted	
itraconazole		

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itraconazole solution (generic for Sporanox® soln) ketoconazole Lamisil tab/granules Noxafil Onmel Sporanox cap/soln Talsura Vfend tab/susp voriconazole tab & powder for susp		
Otezla	 Psoriatic Arthritis Member must meet all the following criteria: Diagnosis of moderate to severe Psoriatic Arthritis Member is 18 years of age or older Prescribed by or in consultation with a Rheumatologist Member has active Psoriatic Arthritis despite a three-month trial with one of the following: Methotrexate (leflunomide or sulfasalazine if methotrexate is contraindicated) Anti-tumor necrosis factor antagonists such as Humira or Enbrel. Otezla will not be used in combination with a targeted synthetic Disease-Modifying Anti-Rheumatic Drug (for example Xeljanz), or a biologic Disease-Modifying Anti-Rheumatic Drug (for example Actemra, Kineret, Orencia, Rituxin), or a Tumor Necrosis Factor antagonist (for example Cimzia, Enbrel, Humira, Remicade, or Simponi) 	Initial Approval: 4 months Renewal: 12 months Requires: Member is responding to treatment Quantity Level Limit (QLL): 60 tablets per 30 days after initial 5-day titration

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	(NOTE: Anti-Tumor Necrosis Factors (TNFs) require prior authorization)	
	Plaque Psoriasis	
	 Member must meet all the following criteria: Diagnosis of moderate to severe Plaque Psoriasis Member is 18 years of age or older Prescribed by or in consultation with a dermatologist Documentation to support an adequate 3-month trial and failure or intolerance to methotrexate or cyclosporine or there is a true contraindication to both. Attestation to one of the following: More than 10% of body surface area affected Less than 10% body surface area affected, but involves sensitive areas (for example: hands, feet, face or genitals) that interferes with daily activities Psoriasis Area and Severity Index score of more than 10 Trial and failure of 2 month of phototherapy (PUVA (psoralen ultraviolet type A), UVB (ultraviolet type B)) Otezla will not be used in combination with a targeted synthetic Disease-Modifying Anti-Rheumatic Drug (for example Xeljanz), or a biologic Disease-Modifying Anti-Rheumatic Drug (for example Actemra, Kineret, Orencia, Rituxin), or a Tumor Necrosis Factor antagonist (for example Cimzia, Enbrel, Humira, Remicade, or Simponi) 	
Pancreatic Enzymes	Clinical criteria for preferred pancreatic enzymes:	Initial Approval:
Preferred:	 Diagnosis of pancreatic insufficiency due to cystic fibrosis or chronic pancreatitis or pancreatectomy. 	1 year
Creon	 If member has a feeding tube then two different pancreatic enzymes can be 	Renewal:
Viokace	approved for use together.	1 year
Zenpep		

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Non-Preferred: Pancreaze pancrelipase Pertzye Ultresa	 In addition, clinical criteria for non-preferred agents: Must meet general non-preferred guideline Had failure to respond to a therapeutic trial of at least two preferred drugs; OR Member has a diagnosis of Cystic Fibrosis If member has a feeding tube then two different pancreatic enzymes can be approved for use together 	Requires: Member is responding to treatment
Phosphodiesterase 5 Inhibitors (PDE-5) & Combinations Preferred: Alyq (tadalafil) sildenafil tab sildenafil suspension tadalafil	 Clinical criteria for all preferred and non-preferred PDE-5s: Diagnosis of pulmonary hypertension in members greater than 18 years of age is required (greater than or equal to 1 years for oral Revatio only) The prescriber must be a pulmonary specialist or cardiologist Must have rationale for not taking the sildenafil tab to receive injectable Revatio Clinical Criteria for Non-Preferred Agents: Must meet general non-preferred guideline Had failure to respond to a therapeutic trial of at least two preferred drugs 	Initial Approval: 1 year Renewal: 1 year Requires: Member is responding to treatment
Non-preferred: Adcirca Liqrev Revatio tablet Revatio injection Revatio suspension Tadliq suspension Opsynvi		

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Proprotein Convertase Subtilisin/Kexin Type 9 Inhibitors (PCSK9 Inhibitors) & Legvio

Leqvio Praluent Repatha

Clinical Criteria for PCSK9 Inhibitors & Leqvio:

- Member's pre-treatment LDL-C level (that is, prior to starting PCSK9 therapy) is provided (Note: Please specify value)
- Medication is used for one of the following diagnoses:
 - To reduce the risk of myocardial infarction, stroke, and coronary revascularization in adults with established cardiovascular disease
 - As an adjunct to diet, alone or in combination with other lipid-lowering therapies (for example, statins, ezetimibe), for treatment of adults with primary hyperlipidemia (including heterozygous familial hypercholesterolemia [HeFH]) to reduce low-density lipoprotein cholesterol (LDL-C)
 - As an adjunct to diet and other LDL-lowering therapies (for example, statins, ezetimibe, LDL apheresis) in patients with homozygous familial hypercholesterolemia (HoFH) who require additional lowering of LDL-C
 - o The member has had prior treatment history with highest available dose or maximally-tolerated dose of high intensity statin (atorvastatin or rosuvastatin) and ezetimibe for at least three continuous months with failure to reach target LDL-C and is in one of the three groups identified by NLA (that is, extremely high risk ASCVD members with LDL-C ≥ 70 mg/dL, very high risk atherosclerotic cardiovascular disease [ASCVD] members with LDL-C ≥ 100 mg/dL, and high risk members with LDL-C ≥ 130 mg/dL)
- Repatha:
 - Member is 10 years of age or older for diagnoses of heterozygous familial hypercholesterolemia (HeFH) or homozygous familial hypercholesterolemia (HoFH)

Initial Approval:

3 months

Renewal Approval:

6 months

Requires:

- Member continues to meet initial diagnosis criteria
- Member achieved at least a 30%
 reduction in LDL-C since the beginning of
 treatment with Praluent, Repatha, or
 Leqvio (Note: please attach clinical notes
 and laboratory results that support
 reduction in LDL-C after initiation of
 therapy)
- Member continues to benefit from treatment as measured by either continued decrease in LDL-C levels or maintenance of optimum LDL-C levels (Note: please attach clinical notes and laboratory results that support continued benefit of Praluent, Repatha, or Leqvio therapy)
- If member is unable to use a maximum dose of atorvastatin or rosuvastatin due to

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 Member is 18 years of age or older when medication is used to reduce the risk of myocardial infarction, stroke, and coronary revascularization in established cardiovascular disease

Praluent:

- Member is 8 years of age or older for diagnoses of heterozygous familial hypercholesterolemia (HeFH)
- Member is 18 years of age of older for a diagnosis of homozygous familial hypercholesterolemia (HoFH) or when medication is used to reduce the risk of myocardial infarction, stroke, and coronary revascularization in established cardiovascular disease
- Leqvio: member is 18 years of age or older

For treatment of Heterozygous Familial Hypercholesterolemia:

- Member has a definite diagnosis of heterozygous familial hypercholesterolemia (HeFH) as defined by the Dutch Lipid Clinic Network criteria (total score greater than 8) (Note: please provide a copy of the lab report with LDL-C level at time of diagnosis and other documentation supporting clinical/family history and/or physical findings (For example, chart notes, medical records)); OR
- Member has a definite diagnosis of HeFH as defined by Simon Broome diagnostic criteria

For treatment of Homozygous Familial Hypercholesterolemia:

Member is diagnosed with homozygous familial hypercholesterolemia (HoFH)

muscle symptoms, documentation of a causal relationship must be established between statin use and muscle symptoms. Documentation must demonstrate that the member experienced pain, tenderness, stiffness, cramping, weakness, and/or fatigue, and all of the following (Note: documentation showing details must be provided):

- Muscle symptoms resolved after discontinuation of statin
- Muscle symptoms occurred when rechallenged at a lower dose of the same statin
- Muscle symptoms occurred after switching to an alternative statin
- Documentation ruling out non-statin causes of muscle symptoms (for example, hypothyroidism, reduced renal function, reduced hepatic function, rheumatologic disorders [for example, polymyalgia rheumatica], steroid myopathy, vitamin D deficiency, or primary muscle disease)

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- Genetic testing has confirmed the presence of two mutant alleles at the LDLR, APOB, PCSK9, or LDLRAP1 gene locus (Note: Please attach a copy of genetic testing result)
- Diagnosis of HoFH has been confirmed by any of the following (Note: Please specify and provide a copy of the laboratory report with LDL-C level at time of diagnosis and other documentation supporting the presence of xanthoma or family history of HoFH (for example, chart notes, medical records)):
 - Untreated LDL-C > 500 mg/dL and cutaneous or tendon xanthoma before age 10 years
 - Untreated LDL-C > 500 mg/dL and untreated elevated LDL-C levels consistent with heterozygous familial hypercholesterolemia in both parents
 - Treated LDL-C ≥ 300 mg/dL and cutaneous or tendon xanthoma before age 10 years
 - Treated LDL-C ≥ 300 mg/dL and untreated elevated LDL-C levels consistent with heterozygous familial hypercholesterolemia in both parents

For treatment of established cardiovascular disease:

- Member has a history of clinical ASCVD or a cardiovascular event listed below (Note: Please specify which):
 - o Acute coronary syndromes
 - o Stable or unstable angina
 - Stroke of presumed atherosclerotic origin
 - Coronary or other arterial revascularization procedure (for example, percutaneous transluminal coronary angioplasty [PTCA], coronary artery bypass graft [CABG])

 The member has been diagnosed with statin-induced rhabdomyolysis

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	 Peripheral arterial disease of presumed atherosclerotic origin Findings from a computerized tomography (CT) angiogram or catheterization consistent with clinical ASCVD Myocardial infarction Transient ischemic attack (TIA) Member's pre-treatment LDL-C level must be provided 	
Sickle Cell Disease Drugs	Clinical Criteria for Sickle Cell Disease Drugs:	Initial approval:
	Medication is prescribed by or in consultation with an oncologist, hematologist or	6 months
Preferred: Droxia	sickle cell specialist	
Endari	Member has a diagnosis of Sickle Cell Disease presenting as one of following: HbSS,	Renewal:
Siklos** (PA not required	HbSC, HbSβ ⁰ -thalassemia, or HbSβ ⁺ -thalassemia	1 year
unless member is greater	Dose is proper for the member's age or other conditions affecting the dose,	
than 18 years of age)	according to the product package insert approved by the FDA	Requires:
	Adakveo:	Member continues to meet initial
Non-preferred:	o Member had an insufficient response to a minimum 3-month trial of	approval criteria
Adakveo	hydroxyurea (unless contraindicated or intolerant)	Member had disease response
Glutamine	 Member has experienced TWO or more vaso-occlusive crises (VOC) in the 	improvement with treatment
	previous year despite adherence to hydroxyurea therapy	Adakveo:
	Siklos:	 Member's response compared to pre-
	o Member is 18 years of age or older (if member is 2 to 17 years of age medication	treatment baseline is evidenced by a
	does not require PA)	decrease in the frequency of vaso-
	o Brand Siklos is medically necessary	occlusive crises (VOC) necessitating
	Glutamine:	treatment, reduction in number or
	 Member had an insufficient response to a minimum 3-month trial of brand 	duration of hospitalizations, and/or
	name Endari (unless contraindicated or intolerant)	reduction in severity of VOC
Topical Antifungals	Clinical criteria for Topical Antifungals:	Approval:

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	Luliconazole: member is 18 years of age or older	1 year
Non-preferred:	Ciclopirox: member is 12 years of age or older	
Ciclopirox 8% kit	Jublia: member is 6 years of age or older	
Jublia	Onychomycosis: ciclopirox 8%, Jublia	
Luliconazole	 Must have failure of an adequate trial of ONE oral alternative – terbinafine (6 weeks for fingernail infections; 1 week for toenail infections); fluconazole (6 months); itraconazole (60 days for fingernail infections; 90 days for toenail infections) 	
	Tinea pedis, cruris, or corporis: luliconazole	
	 Must have failure of an adequate trial of TWO preferred topical antifungal medications OR allergy or contraindication to oral terbinafine, fluconazole, or itraconazole 	
Tranexamic Acid Tablets ⁱⁱ	Member is 12 years of age or older	Initial Approval:
	Treatment is for cyclic heavy menstrual bleeding	90 days
	 Prescriber attestation that member has no fibroids, or fibroids are less than 3 cm in size 	Renewal Approval:
	There was inadequate response, intolerable side effect, or contraindication to one	6 months
	oral Non-Steroidal Anti-inflammatory Drug (NSAID)	Requires:
	 Member had inadequate response, intolerable side effect, or contraindication to one of the following: 	Reduction in menstrual blood loss
	 Oral hormonal cycle control combinations Oral progesterone 	 Quantity Level Limit: Menstrual bleeding: 20 to blets now 20 down
	Progesterone-containing intrauterine device (IUD)Medroxyprogesterone depot	30 tablets per 30 days • Hemophilia:
	 Member does not have history of thrombosis or thromboembolism (including retinal vein or artery occlusion) 	84 tablets per 30 days

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	Approved for treatment and prevention of acute bleeding episodes, such as dental surgery, in members with hemophilia.	
Tysabri	Clinical criteria for Tysabri: Member is 18 years of age or older Prescriber and member enrolled in and meet the conditions of the TOUCH (applicable to Tysabri) or REMS (applicable to Tyruko) programs Member has a documented negative JCV antibody ELISA test within the past 6 months Requested product will not be used in combination with antineoplastic, immunosuppressant, or immunomodulating agents Member is immunocompetent Multiple sclerosis: Medication will be used as a single therapy Member has a confirmed diagnosis of multiple sclerosis (MS) as documented by laboratory report (i.e., MRI) Member has a diagnosis of a relapsing form of MS [i.e., relapsing-remitting MS (RRMS), active secondary progressive disease (SPMS), or clinically isolated syndrome (CIS) Crohn's disease: Member has moderate to severe active disease Physician has assessed baseline disease severity utilizing an objective measure/tool Member has a documented trial and failure on ONE oral immunosuppressive therapy for at least 3 months, unless use is contraindicated, such as corticosteroids, methotrexate, azathioprine, and/or 6-mercaptopurine	Initial approval: 6 months Renewals: 1 year Requires: • Multiple sclerosis: • Member continues to meet the relevant criteria identified in the initial criteria • Member has an absence of unacceptable toxicity from the drug • Member is being continuously monitored for response to therapy and it indicates a beneficial response • Crohn's disease: • Disease has responded as indicated by improvement in signs and symptoms compared to baseline such as endoscopic activity, number of liquid stools, presence and severity of abdominal pain, presence of
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	 Member has a trial of two of the preferred Cytokine and CAM antagonist agents for Crohn's Disease (see Cytokine and CAM Antagonists on the PDL) Will be used as single agent therapy [Not used concurrently with another biologic drug or immunosuppressant (e.g., 6-mercaptopurine, azathioprine, cyclosporine, methotrexate, etc.) used for Crohn's Disease 	abdominal mass, body weight compared to IBW, hematocrit, presence of extra intestinal complications, tapering or discontinuation of corticosteroid therapy, use of anti-diarrheal drugs, and/or an improvement on a disease activity scoring tool Initial renewal only (6-month approval): Member has been tapered off of oral corticosteroids within 6 months of starting Tysabri Subsequent renewals (12-month approval): Member does not require additional steroid use that exceeds 3 months in a calendar year to control their Crohn's disease
Vivjoa	Clinical Criteria for Vivjoa	Approvals:
	 Member is 10 years of age or older Documentation member has diagnosis of recurrent vulvovaginal candidiasis with ≥3 	Date of service (1 day)
	episodes of vulvovaginal candidiasis (VVC) in a 12-month period	Quantity Level Limits:
	Member is a biological female who is postmenopausal or has another reason for permanent infertility (e.g., tubal ligation, hysterectomy, salpingo-oophorectomy)	18 tablets per treatment course

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	Member has tried and failed or has a contraindication or intolerance to	
	maintenance antifungal therapy with oral fluconazole	
Vraylar	 Clinical Criteria for Vraylar: Member is 18 years of age or older Member has one of the following diagnoses: Bipolar disorder, major depressive disorder, or schizophrenia For diagnosis of major depressive disorder member must be adherent to complementary therapy with a selective serotonin reuptake inhibitor (SSRI) or serotonin/norepinephrine reuptake inhibitor (SNRI) (80% of fills in the last 90 days) 	Approval: 1 year
Weight Management	Clinical criteria for weight loss agents:	Initial approval:
Medications Preferred:	 Phentermine (min age 17 years), phendimetrazine tablet (min age 18 years), phendimetrazine ER capsule (min age 17 years), and orlistat (min age 12 years): Body mass index (BMI) ≥ 30 kg/m²; OR 	 Benzphetamine, diethylpropion, phendimetrazine, phentermine: 3 months GLP-1 receptor agonists: 6 months
Orlistat Phendimetrazine IR Phendimetrazine ER	 Member has a BMI of ≥ 27 kg/m² with at least one weight-related comorbidity (i.e. coronary heart disease, dyslipidemia, hypertension, sleep apnea, type 2 diabetes) Benzphetamine (min age 17 years) and diethylpropion (min age 16 years): 	Orlistat: 6 months
Phentermine	o Body mass index (BMI) ≥ 30 kg/m²	Renewal requests: Varies (drug specific):
Benzphetamine Diethylpropion IR Diethylpropion ER Non-preferred:	 Imcivree (min age 6 years): Body mass index (BMI) ≥ 30 kg/m² Prescribed by or in consultation with an endocrinologist or geneticist Member has Bardet-Biedl syndrome (BBS) Member has proopiomelanocortin (POMC), proprotein convertase 	All medications: Renewals will no longer be granted once a member reaches a BMI < 25
Imcivree Saxenda SQ	subtilisin/kexin type 1 (PCSK1), or leptin receptor (LEPR) deficiency, as confirmed by a genetic test	Benzphetamine, diethylpropion, phendimetrazine, phentermine:
Wegovy SQ Zepbound	 Member's genetic variants are interpreted as pathogenic, likely pathogenic, or of uncertain significance (VUS) 	If member achieves at least a 10-lb. weight loss during initial 3 months of therapy, an additional 3-month PA may

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- Wegovy (min age 12 years), Saxenda (min age 12 years), and Zepbound (min age 18 years):
 - o Member meets one of the following:
 - Body mass index (BMI) > 40 kg/m² if no applicable risk factors
 - BMI > 37 kg/m² with one or more of the following risk factors: dyslipidemia, hypertension, type II diabetes
 - Member meets one of the following:
 - Member has tried and failed one of the non-Glucagon-Like Peptide-1 (GLP-1) weight loss medications*
 - Member is intolerant to all non-GLP1 weight-loss medications*, member is not concurrently on another GLP-1 receptor agonist, and the member has tried and failed* the selected product as indicated on the preferred drug list (Saxenda)
 - Note: definitions of accepted drug trials are as follows:
 - Benzphetamine, diethylpropion, phendimetrazine, phentermine: 3-month trial without a weight loss of 10lbs
 - Orlistat: 6-month trial without a weight loss of 10lbs
 - GLP-1 receptor agonist: 6-month trial without a body weight reduction of 5%

Initial request requirements:

- No contraindications to use (i.e. uncontrolled hypertension, hyperthyroidism etc for stimulant based products)
- No malabsorption syndromes, cholestasis, pregnancy and/or lactation (for orlistat)
- No history of an eating disorder (for example, anorexia, bulimia)
- No acute pancreatitis, acute suicidal behavior/ideation, personal or family history of medullary thyroid cancer or multiple endocrine neoplasia 2 syndrome (if requesting

be granted. Maximum length of continuous drug therapy = 6 months (waiting period of 6 months before next request)

Orlistat:

 If member achieves at least a 10-lb. weight loss, an additional 6-month PA may be granted. Maximum length of continuous drug therapy = 24 months (waiting period of 6 months before next request)

Imcivree:

If the member has experienced ≥ 5% reduction in body weight (or ≥ 5% of baseline BMI in those with continued growth potential), an additional 1-year SA may be granted.

GLP-1 Receptor agonists:

 If the member achieves a weight loss of ≥ 5% in body weight compared to the most recent authorization, an additional 6-month PA may be granted.

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	 a GLP-1 Receptor Agonist) Qualifying criteria (excluding Imcivree): Participation in nutritional counseling 	
	 Participation in physical activity program, unless medically contraindicated Commitment to continue weight-loss treatment plan The provider attests that the member's obesity is disabling and life threatening (i.e. puts the member at risk for high morbidity conditions) 	
	Following documentation must be included in medical records:	
	 Current medical status and weight loss plan. An individualized weight-loss program sho include a specific reduced-calorie meal plan, recommended routine physical activity and behavioral intervention, including lifestyle modification as needed to improve adherence and outcomes Note: Providers should also summarize details of previous weight-loss treatment plans to include diet and exercise plans, in addition to submitting a copy of the plan 	y,
	Current height and weight measurements	
Xifaxan	 Clinical Criteria for Xifaxan: Xifaxan®: 200 mg tabs: Treatment of travelers' diarrhea caused by noninvasive strains of E. coli Member is 12 years of age or older 	Approvals: Xifaxan 200 mg tabs: 1 month Xifaxan 550 mg tabs: 3 months
	 Xifaxan®550 mg tabs: - Reduction in risk of overt hepatic encephalopathy recurrence Member is greater than or equal to 18 years of age OR Treatment of irritable bowel syndrome with diarrhea (IBS-D) 	 Quantity Level Limits: Xifaxan 200 mg tabs: 9 tabs per claim Xifaxan 550 mg tabs: Hepatic encephalopathy: 2 tablets per day

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	o Member is greater than or equal to 18 years of age	 Irritable bowel syndrome with diarrhea (IBS-D): 3 times a day for 14 days (42 tablets per 14 days); can be retreated up to two times with the same regimen. Max 126 tablets per 365 days
Zolgensma	One dose of Zolgensma per lifetime will be approved – medication will not be renewed	Approval: 1 month
	Clinical criteria for Zolgensma:	
	Medication is prescribed by a Pediatric Neuromuscular Neurologist with expertise in SMA	Quantity Level Limits: One dose/kit per lifetime
	 Member has a diagnosis of 5q spinal muscular atrophy confirmed by either bi-allelic deletion or dysfunctional point mutation of the SMN1 gene, with 4 or fewer copies of SMN2 	
	Member is less than 24 months of age	
	Member is not ventilator-dependent, defined as requiring invasive ventilation (tracheostomy) or respiratory assistance for 16 or more hours per day (including noninvasive ventilator support) continuously for 21 or more days in the absence of an acute reversible event	
	Member has baseline anti-AAV9 antibody titer of less than or equal to 1:50 measured by ELISA	
	Member has LFTs less than 2X the upper limit of normal determined by certified laboratory	
	Member has received NO treatment with immunosuppressive therapy in the 3 months prior to starting Zolgensma treatment (e.g., corticosteroids, cyclosporine, tacrolimus, methotrexate, cyclophosphamide, intravenous immunoglobulin, rituximab)	

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Member does NOT have advanced disease (e.g., complete limb paralysis, permanent
ventilation support)
Member does NOT have symptoms of active viral infection
Member does NOT have concomitant illness that may create unnecessary risks for
gene transfer
Member has had NO prior treatment with Zolgensma
Member will NOT revceive the requested treatment in combination with Spinraza
(nusinersen) or Evrysdi (risdiplam)

Otezla References

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