

Department of Vermont Health Access Pharmacy Benefit Management Program

EFFECTIVE

Version

Updated: 06/02/2023

Vermont Preferred Drug List and Drugs Requiring Prior Authorization

(includes clinical criteria)

The Commissioner for Office of Vermont Health Access shall establish a pharmacy best practices and cost control program designed to reduce the cost of providing prescription drugs, while maintaining high quality in prescription drug therapies. The program shall include:

"A preferred list of covered prescription drugs that identifies preferred choices within therapeutic classes for particular diseases and conditions, including generic alternatives"

From Act 127 passed in 2002

The following pages contain:

- The therapeutic classes of drugs subject to the Preferred Drug List, the drugs within those categories and the criteria required for Prior Authorization (P.A.) of non-preferred drugs in those categories. The therapeutic classes of drugs which have clinical criteria for Prior Authorization may or may not be subject to a preferred agent.
- Within both categories there may be drugs or drug classes that are subject to Quantity Limit Parameters.

Therapeutic class criteria are listed alphabetically. Within each category the Preferred Drugs are noted in the left-hand columns. Representative non-preferred agents have been included and are listed in the right-hand column. Any drug not listed as preferred in any of the included categories requires Prior Authorization. Approval of non-preferred brand name products may require trial and failure of at least 2 different generic manufacturers. Drugs used for weight loss, drugs used to promote fertility, and drugs used for cosmetic purposes or hair growth are excluded from coverage under the Vermont Medicaid Pharmacy program.

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This is not an all-inclusive list of available covered drugs and includes only managed categories. Unless otherwise stated, the listing of a particular brand or generic name includes all dosage forms of that drug. NR indicates a new drug that has not yet been reviewed by the P&T Committee.

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PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
	ACNE AGENTS	
ORAL AGENTS		
AMNESTEEM (isotretinoin) capsules CLARAVIS (isotretinoin) capsules MYORISAN (isotretinoin) capsules ZENATANE (isotretinoin) capsules	Absorica® (isotretinoin) capsules Isotretinoin capsules	Absorica, Isotretinoin: patient has had a documented side effect, allergy, or treatment failure with at least two isotretinoin preferred products.
TOPICAL AGENTS		
BENZOYL PEROXIDE PRODUCTS BENZOYL PEROXIDE 2.5%, 5%, 10%G; 3%, 5%, 10% CL; 5.3%, 9.8% F CLINDAMYCIN PRODUCTS CLINDAMYCIN 1% S, G, L, P (compare to Cleocin-T) ERYTHROMYCIN PRODUCTS	Benzol Peroxide 5%, 10%L Clindacin (clindamycin) 1% CL, P, Swab Clindamycin 1%F Clindamycin 1%G (compare to Clindagel) 75mL bottle Cleocin-T® (clindamycin) 1% L	 Single ingredient products: patient has had a documented side effect, allergy, or treatment failure with two preferred products including one from the same sub-category, if there is one available. If a product has an AB rated generic, there must have been a trial of the generic. Benzaclin, Benzamycin: patient must have a documented intolerance to the generic equivalent. Sodium Sulfacetamide Products: patient has had a documented side effect, allergy or treatment failure with two preferred products, one of which must be Klaron lotion. Clindamycin/Benzoyl peroxide pump, Onexton: there must be a clinically
ERYTHROMYCIN 2% S, G MINOCYCLINE PRODUCTS All Products Require PA SODIUM SULFACETAMIDE PRODUCTS KLARON® (sodium sulfacetamide 10% L)	Erygel® (erythromycin 2% G) Ery (erythromycin 2%) P Amzeeq® (minocycline) 4% foam Sodium Sulfacetamide 10% L Sodium Sulfacetamide/Sulfur CL, C, P, E Sodium Sulfacetamide/Sulfur W Sumaxin ® (sulfacetamide/sulfur L, P, W)	compelling reason why clindamycin/benzoyl peroxide gel cannot be used. Limitations: Kits with non-drug products are not covered
COMBINATION PRODUCTS ERYTHROMYCIN / BENZOYL PEROXIDE CLINDAMYCIN/BENZOYL PEROXIDE (compare to Benzaclin®) G	Benzaclin® (clindamycin/benzoyl peroxide) Benzamycin® (erythromycin/benzoyl peroxide) Clindamycin/Benzoyl Peroxide Pump Onexton® (clindamycin/benzoyl peroxide)	
<u>OTHER</u>	Dapsone 5%, 7.5% G	
C=cream, CL=cleanser, E=emulsion, F=Foam, G=gel, L=lotion, O=ointment, P=pads,	All other brands any topical acne anti-infective medication	

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
S=solution, W =wash, B =bar		
TOPICAL – ANDROGEN RECEPTOR INHIB	ITORS	
All products require PA	Winlevi® (clascoterone) 1% C	Winlevi: patient has had a documented side effect, allergy, or treatment failure with two preferred topical acne agents.
TOPICAL - RETINOIDS		
AVITA [®] (tretinoin) ADAPALENE 0.1% G, 0.3% G DIFFERIN® (adapalene) 0.1% G RETIN-A® (tretinoin) 0.025%, 0.05%, 0.1% C; 0.01%, 0.025% G C= cream, G=gel, L=lotion	Adapalene (compare to Differin®) 0.1% C Adapalene/Benzoyl Peroxide 0.1-2.5% G Altreno™ (tretinoin) 0.05% L Arazlo® (tazarotene) 0.045% L Atralin® (tretinoin) 0.05% G Clindamycin/tretinoin 1.2-0.025% G Fabior® (tazarotene) 0.1% F Plixda® (adapalene) 0.1% F Plixda® (adapalene) 0.1% swabs Retin-A Micro® (tretinoin microsphere) 0.04%, 0.06%, 0.08%, 0.1% G Tazarotene (compare to Tazorac®) 0.1% C Tretinoin (compare to Retin-A®) 0.025%, 0.05%, 0.1% C; 0.01%, 0.025% G Tretinoin microsphere (compare to Retin-A Micro®) 0.1%, 0.04% Twyneo® (tretinoin/benzoyl peroxide) 0.1%-3% C	Altreno, Atralin, Retin-A Micro, Tretinoin, Tretinoin microsphere: diagnosis or indication is acne vulgaris, actinic keratosis, or rosacea AND patient has had a documented side effect, allergy, or treatment failure with a preferred topical tretinoin product (Avita or Retin-A®). Adapalene Cream: patient has had a documented side effect, allergy, or treatment failure with adapalene gel. Arazlo, Fabior, Tazarotene: patient has had a documented side effect or treatment failure with a preferred topical tretinoin product and adapalene. Adapalene/benzoyl peroxide gel, Clindamycin/tretinoin gel, Twyneo: patient has had a documented side effect or treatment failure on combination therapy with the separate ingredients of the combination product Plixda: patient has had a documented side effect, allergy, or treatment failure with brand Differin AND a generic adapalene product. Limitations: Coverage of topical retinoid products will not be approved for cosmetic use (wrinkles age spots, etc.) (i.e. Avage, Renova, Tri-Luma).
TOPICAL - ROSACEA		
FINACEA [®] (azelaic acid) 15% <i>G</i> , F METRONIDAZOLE 0.75% <i>C</i> , <i>G</i> , <i>L C=cream</i> , <i>F=Foam</i> , <i>G=gel</i> , <i>L=lotion</i>	All brand metronidazole products (MetroCream 0.75% <i>C</i> , Metrogel® 1% <i>G</i> , MetroLotion® 0.75% <i>L</i> , Noritate® 1% <i>C</i> etc.) Epsolay® (benzoyl peroxide) 5% C Ivermectin (compare to Soolanta®) 1% C Metronidazole 1% <i>G</i> Rhofade® (oxymetazoline) 1% C Zilxi® (minocycline) 1.5% F	Brand name metronidazole products, Metronidazole 1% gel (generic): diagnosis or indication is rosacea AND patient has had a documented side effect, allergy, or treatment failure with a preferred generic topical metronidazole product. If a product has an AB rated generic, there must have also been a trial of the generic formulation. Epsolay, Ivermectin, Rhofade: the patient has had a documented side effect, allergy, or treatment failure with 2 preferred topical rosacea agents. Zilxi: diagnosis or indication is rosacea AND patient has had a documented side effect, allergy, or treatment failure with a preferred generic topical metronidazole product and Finacea. Limitations: The use of Mirvaso (brimonidine topical gel) for treating skin redness is considered cosmetic. Medications used for cosmetic purposes are excluded from coverage. Mirvaso topical gel has not been shown to improve any other symptom of rosacea (e.g. pustules, papules, flushing, etc.) or to alter the course of the disease.

(No PA required unless otherwise noted)

NON-PREFERRED AGENTS

(PA required)

PA CRITERIA

ADHD AND NARCOLEPSY CATAPLEXY MEDICATIONS

SHORT/INTERMEDIATE ACTING STIMULANTS

AMPHETAMINE/DETROAMPHETAMINE (compare to Adderall $^{(8)}$)

DEXMETHYLPHENIDATE (compare to Focalin[®])
METHYLIN[®] (compare to Ritalin[®]) solution

METHYLPHENIDATE (compare to Ritalin $^{(8)}$) tablets, solution

METHYLPHENIDATE SR (compare to Ritalin® SR)

PROCENTRA® (dextroamphetamine sulfate) 1 mg/ml oral solution

Adderall® (amphetamine/dextroamphetamine)

Amphetamine Sulfate (compare to Evekeo)

Desoxyn[®] (methamphetamine)

Dextroamphetamine sulfate 1 mg/ml oral solution Dextroamphetamine IR (Zenzedi 5 or 10 mg, formerly

Dexedrine®)
Evekeo® (amphetamine sulfate)

Evekeo® ODT (amphetamine sulfate)

Focalin[®] (dexmethylphenidate)

Methamphetamine (compare to Desoxyn[®])
Methylphenidate (compare to Ritalin ®) chewable tablets

Ritalin® (methylphenidate)

Zenzedi[®] (dextroamphetamine IR) 2.5 mg, 7.5 mg, 15 mg, 20 mg, 30 mg tablets

Clinical Criteria for ALL non-preferred drugs: patient has a diagnosis of ADD, ADHD or narcolepsy AND patient has been started and stabilized on the requested medication. (Note: samples are not considered adequate justification for stabilization.) OR patient meets additional clinical criteria outlined below.

Focalin, Adderall, Ritalin: the patient must have had a documented intolerance to the preferred generic equivalent.

Methamphetamine and Desoxyn: Given the high abuse potential of methamphetamine and Desoxyn, the patient must have a diagnosis of ADD, ADHD or narcolepsy and have failed all preferred treatment alternatives. In addition, for approval of brand name Desoxyn, the patient must have had a documented intolerance to generic methamphetamine.

Methylphenidate chewable tablets: patient has a documented intolerance to methylphenidate and Methylin solution.

Evekeo ODT, Dextroamphetamine oral solution: patient has a medical necessity for a non-solid oral dosage form. (e.g. swallowing disorder). AND the patient has a documented intolerance Procentra oral solution.

Amphetamine Sulfate, Dextroamphetamine IR, Zenzedi, Evekeo: the patient has had a documented side-effect, allergy, or treatment failure of at least 2 preferred agents (If a product has an AB rated generic, there must have been a trial of the generic.)

LONG ACTING STIMULANTS METHYLPHENIDATE PRODUCTS ORAL

CONCERTA® (methylphenidate SA OSM IR/ER, 22:78%)

DEXMETHYLPHENIDATE SR 24 HR IR/ER, 50:50% (compare to Focalin XR®)

METHYLPHENIDATE CR, IR/ER, 30:70% (compare to Metadate CD®)

METHYLPHENIDATE SR 24 HR, IR/ER, 50:50% (compare to Ritalin LA®)

QUILLICHEW ER TM (methylphenidate IR/ER, 30:70%) chewable tablets

RITALIN LA® (methylphenidate SR 24 HR, IR/ER, 50:50%)

Adhansia ® XR (methylphenidate IR/ER 20:80%)

OTY LIMIT: 1 capsule/day

Aptensio® XR (methylphenidate DR 24HR IR/ER, 40:60%)

 $Az starys^{TM} (ser dex methyl phenidate / dex methyl phenidate) \\$

Cotempla® XR (methylphenidate IR/ER 25:75%) ODT Focalin® XR (dexmethylphenidate SR 24 HR

Jornay PMTM (methylphenidate ER) capsules *QTY LIMIT*: 1 capsule/day

Methylphenidate DR 24HR IR/ER, 40:60% (compare to Aptensio®XR)

Methylphenidate SA OSM IR/ER, 22:78% (compare to Concerta®)

 $Relexxii @ \ (methylphenidate \ ER\ OSM)\ IR/ER,\ 22:78\%$

Clinical criterial for ALL non-preferred drugs: the patient has a diagnosis of ADD, ADHD or narcolepsy AND has been started and stabilized on the requested medication. (Note: samples are not considered adequate justification for stabilization) OR meets the additional clinical criteria outlined below.

Azstarys, Adhasia XR, Cotempla XR ODT, Jornay PM: patient has had a documented side-effect, allergy, or treatment failure on 3 preferred long-acting Methylphenidate products.

Aptensio XR, Methylphenidate DR 40:60: patient has had a documented side effect, allergy, or treatment failure on two preferred long-acting Methylphenidate products. For approval of Methylphenidate DR 40:60, the patient must also have a documented intolerance to brand Aptensio XR.

Focalin XR: the patient must have had a documented intolerance to the preferred generic equivalent.

Methylphenidate SA OSM: the patient must have a documented intolerance to

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
ORAL SUSPENSION QUILLIVANT XR [®] (methylphenidate IR/ER, 20:80%) QTY LIMIT: 1 bottle/Rx (60ml, 120ml, 150ml) 2 bottles/Rx (180ml)		brand Concerta. Relexxi: Both Concerta and methylphenidate SA OSM must be on a long-term backorder and unavailable from the manufacturer.
AMPHETAMINE PRODUCTS ORAL ADDERALL XR® (amphetamine/dextroamphetamine SR 24 HR, IR/ER, 50:50%) AMPHETAMINE/DEXTROAMPHETAMINE SR 24 HR, IR/ER, 50:50% (compare to Adderall XR®) VYVANSE® (lisdexamfetamine) capsule QTY LIMIT: 1 cap/day	Daytrana [®] (methylphenidate patch) <i>QTY LIMIT</i> : 1 patch/day Methylphenidate patch (compare to Daytrana®) <i>QTY LIMIT</i> : 1 patch/day Adzenys XR® ODT (amphetamine SR 24 HR, IR/ER, 50:50%) <i>QTY LIMIT</i> : 1 cap/day Adzenys ER TM suspension (amphetamine SR 24 HR, IR/ER, 50:50%) Dyanavel TM suspension (amphetamine/dextroamphetamine SR) <i>QTY LIMIT</i> : 240ml/30days Dexedrine CR® (dextroamphetamine 24 HR SR) Dextroamphetamine 24 HR SR (compare to Dexedrine CR®) Mydayis® (mixed amphetamine salts) extended-release capsules	 Daytrana patch, Methylphenidate patch: patient has a documented medical necessity for a specialty non-oral dosage form AND for approval of generic Methylphenidate patch, the patient must have a documented intolerance to be Daytrana. Adzenys XR ODT, Adzenys ER suspension, Dyanavel XR suspension, Vyva Chew: patient must be unable to tolerate Adderall XR sprinkled onto applesauce Vyvanse mixed with yogurt, water, or orange juice. Dexedrine CR, Dextroamphetamine SR, Mydayis: patient must have a documented intolerance to two preferred amphetamine products. For approf brand Dexedrine CR, the patient must also have a documented intolerant to the generic equivalent.
MISCELLANEOUS ARMODAFINIL (compare to Nuvigil®)	Vyvanse® (lisdexamfetamine) chewable tablet QTY LIMIT: 1 tab/day	Intuniv, Nuvigil, Provigil, Strattera: patient must have a documented
ARMODAFINE (compare to Nivigit) QTY LIMIT: 50 mg = 2 tabs/day 150 mg/200 mg/250 mg = 1 tab/day, Max days supply = 30 days ATOMOXETINE (compare to Strattera®) QTY LIMIT: 10, 18, 25 and 40 mg = 2 capsules/day 60, 80 and 100 mg = 1 capsule/day FDA maximum recommended dose = 100 mg/day CLONIDINE ER QTY LIMIT: 4 tabs/day	Intuniv [®] (guanfacine extended release) tablet QTY LIMIT: 1 tablet/day Nuvigil [®] (armodafinil) QTY LIMIT: 50 mg = 2 tablets/day; 150 mg/200 mg/250 mg = 1 tablet/day, Max days supply = 30 days Provigil [®] (modafinil) QTY LIMIT: 100 mg = 1.5 tablets/day; 200 mg = 2 tablets/day Maximum Daily Dose = 400 mg, Max day supply = 30 days	intolerance to the generic equivalent. Qelbree: the patient has had a documented side effect, allergy, or treatment failuratomoxetine. Sunosi: patient has had a documented side effect, allergy, or treatment failure to preferred agents (may be stimulant or non-stimulant) Wakix patient has no known risk factors for increased QT prolongation (e.g. car arrhythmias, symptomatic bradycardia, hypokalemia, or congenital prolongation of the QT interval) AND medication is not being used in combination with o drugs known to prolong the QT interval (e.g. antipsychotics, erythromycin,

30 days

GUANFACINE ER (Intuniv®)

drugs known to prolong the QT interval (e.g. antipsychotics, erythromycin, $\,$

tricyclic antidepressants) AND patient has had a documented side effect, allergy,

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
MODAFINIL (compare to Provigil®) QTY LIMIT: 100 mg = 1.5 tablets/day; 200 mg = 2 tablets/day Maximum Daily Dose = 400 mg, Max day supply = 30 days Preferred After Clinical Criteria Are Met QELBREE® (viloxazine hydrochloride) ER capsule QTY LIMIT: 100 mg = 1 capsule/day 150 mg = 2 capsules/day 200 mg = 3 capsules/day FDA maximum recommended dose = 600 mg/day	Strattera [®] (atomoxetine) QTY LIMIT: 10, 18, 25 and 40 mg = 2 capsules/day 60, 80 and 100 mg = 1 capsule/day FDA maximum recommended dose = 100 mg/day Sunosi® (solriamfetol) tablet QTY LIMIT: 1 tablet/day FDA maximum recommended dose = 150 mg/day Wakix® (pitolisant) tablet QTY LIMIT: 2 tablets/day FDA maximum recommended dose = 35.6 mg/day Xyrem® (sodium oxybate) oral solution QTY LIMIT: 540 ml/30 days Xywav TM (calcium, magnesium, potassium, and sodium oxybates) solution QTY LIMIT: 9 g (18 mL)/day	or treatment failure to at least 3 agents (may be preferred or non-preferred; may be stimulant or non-stimulant), one of which must be Sunosi. Xyrem, Xywav: patient has had a documented side effect, allergy, or treatment failure to 2 preferred agents (may be stimulant or non-stimulant) and Sunosi AND patient has been enrolled in the REMS program AND for approval of Xywav, the patient must have a documented intolerance to Xyrem.
	ATTEDCEN IMMUNOTIC	ED A DV

ALLERGEN IMMUNOTHERAPY

Oralair:

Oralair®

All products require PA

QTY LIMIT: Palforzia® (pea	tablet/day • Patient age ≥10 years and ≤65 years AND • Treatment must start 12 weeks before expected onset of pollen season and only after confirmed by positive skin test or in vitro testing for pollen-specific IgE antibodies for any of the 5 grass species contained in Oralair AND
	Patient must have an auto-injectable epinephrine on-hand Palforzia:
	 Patient age ≥ 4 years and ≤ 17 years for initial dose escalation or ≥ 4 years for up-dosing and maintenance The prescriber is an allergist or immunologist Prescriber must provide the testing to show that the patient is allergic to peanuts Patient must not have a recent history of uncontrolled asthma, eosinophilic esophagitis, or other eosinophilic GI disease. Prescriber, pharmacy, and patient must be registered with the REMS program Patient must have an auto-injectable epinephrine on-hand Initial approval will be granted for 6 months and includes approval for initial dose escalation and Up Dosing. Approval for Up Dosing may be extended if the patient was unable to tolerate all the dose levels at 2-week intervals. For approval of Maintenance Dosing (300mg daily), pharmacy records will be evaluated to assess compliance with once daily therapy and ensure

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
		no level was missed during Up Dosing. Documentation must be provided attesting that the patient has not experienced any treatment restricting adverse events (e.g. systemic allergic reactions, severe anaphylaxis).
	ALPHA1-PROTEINASE INHI	BITORS
All products require PA	Aralast NP [®] Glassia [®] Prolastin-C [®] Zemaira [®] **Maximum days supply per fill for all drugs is 14 days**	Criteria for Approval: The indication for use is treatment of alpha1 -proteinase inhibitor deficiency-associated lung disease when all of the following criteria are met: Patient's alpha1 -antitrypsin (ATT) concentration < 80 mg per dl [or < 11 micromolar] AND patient has obstructive lung disease as defined by a forced expiratory volume in one second (FEV1) OF 30 - 65% of predicted or a rapid decline in lung function defined as a change in FEV1 of > 120 mL/year. AND medication is being administered intravenously (inhalation administration will not be approved) AND patient is a non-smoker OR patient meets above criteria except lung function has deteriorated beneath above limits while on therapy.
	ALZHEIMER'S MEDICAT	IONS
CHOLINESTERASE INHIBITORS		
DONEPEZIL (compare to Aricept [®]) tablet 5 mg and 10 mg QTY LIMIT: 1 tablet/day DONEPEZIL ODT (compare to Aricept® ODT) QTY LIMIT: 1 tablet/day GALANTAMINE tablet RIVASTIGMINE (compare to Exelon®) capsule QTY LIMIT: 2 capsules/day	Aricept [®] (donepezil) Tablet <i>QTY LIMIT</i> : 1 tablet/day Donepezil (compare to Aricept ®) Tablet 23 mg Galantamine ER capsule (compare to Razadyne® ER) Razadyne ER [®] (galantamine) capsule	 Donepezil 23mg Tablet, Galantamine ER Capsule, Razadyne ER Capsule: the patient has been started and stabilized on the requested medication (Note: samples are not considered adequate justification for stabilization) OR patient had a documented side effect, allergy, or treatment failure to a preferred cholinesterase inhibitor. Adlarity: medical necessity for a specialty dosage form has been provided AND the patient had a documented side effect, allergy, or treatment failure to Exelon patch. Aricept: the patient has a documented intolerance to the generic product. Galantamine Oral Solution, Rivastigmine patch: medical necessity for a specialty dosage form has been provided. AND for approval of rivastigmine
SOLUTION All products require PA	Galantamine (compare to Razadyne®) Oral Solution	patch the patient has a documented intolerance to brand Exelon patch.
TRANSDERMAL EXELON® (rivastigmine transdermal) Patch QTY LIMIT: 1 patch/day	Adlarity® (donzepezil) patch <i>QTY LIMIT:</i> 12 patches/84 days Rivastigmine (compare to Exelon®) patch <i>QTY LIMIT:</i> 1 patch/day	

IMMUNOGLOBULIN GAMMA 1 (IgG1) MONOCLONAL ANTIBODY

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
All products require PA NMDA RECEPTOR ANTAGONIST	Aduhelm® (aducanumab-avwa) IV solution	 Patient is 50 years of age or older Prescriber has assessed and documented baseline disease severity utilizing an objective measure/tool (e.g., MMSE, Alzheimer's Disease Assessment Scale-Cognitive Subscale [ADAS-Cog-13], Alzheimer's Disease Cooperative Study-Activities of Daily Living Inventory-Mild Cognitive, Impairment version [ADCS-ADL-MCI], Clinical Dementia Rating-Sum of Boxes [CDR-SB]). Patient has mild cognitive impairment (MCI) due to Alzheimer's Disease or mild Alzheimer's dementia as evidenced by the following: Clinical Dementia Rating (CDR) Global Score of 0.5 Objective evidence of cognitive impairment at screening MMSE score between 24 and 30 PET scan is positive for amyloid beta plaque OR Cerebrospinal fluid (CSF) test is positive for amyloid Patient has had a recent (within 1 year) brain MRI prior to initiating treatment and prescriber attests to a repeat brain MRI prior to the 7th infusion (first dose of 10mg/kg) and 12th infusion (sixth dose of 10mg/kg) Patient does not have any of the following within 1 year of treatment initiation: pretreatment localized superficial siderosis, 10 or more brain microhemorrhages, or brain hemorrhage >1 cm Patient has had a documented treatment failure, as defined by significant disease progression after 1 year of therapy, with a preferred cholinesterase inhibitor, unless contraindicated. For re-approval, the patient must have responded to therapy compared to pre-treatment baseline as evidenced by improvement, stabilization, or slowing in cognitive or functional impairment AND patient has not progressed to moderate or severe disease (there is insufficient evidence in moderate or severe AD).
ACTA CANADA TO THE CONTROL OF THE CO	.	
MEMANTINE Tablets	Memantine oral solution Memantine XR (compare to Namenda® XR) Oral capsule QTY LIMIT: 1 capsule/day Namenda® (memantine) tablet Namenda® XR (memantine ER) Oral Capsule QTY LIMIT: 1 capsule/day	 Namenda: Patient has a documented intolerance to the generic. Memantine XR, Namenda XR: Patient has not been able to tolerate twice daily dosing of immediate release memantine, resulting in significant clinical impact. Memantine Oral Solution: medical necessity for a specialty dosage form has been provided.
CHOLINESTERASE INHIBITOR/NMDA COM	MBINATION	
All products require PA	Namzaric [®] (donepezil/memantine) Capsule QTY LIMIT: 1 capsule/day	Namzaric: Clinically compelling reason why the individual ingredients of donepezil and memantine cannot be used.

PREFERRED AGENTS
(No PA required unless otherwise noted)
MISCELLANEOUS: TOPICAL AND TRANSDER
LIDOCAINE 3% Cream
LIDOCAINE 4% OTC Patch
LIDOCAINE 4% cream
LIDOCAINE 5% Ointment
LIDOCAINE 5% patch
QTY LIMIT: 3 patches/day
LIDOCAINE/PRILOCAINE 2.5-2.5% Cream
OPIOIDS: SHORT ACTING
OFIGIDS: SHORT ACTING
ACETAMINOPHEN W/CODEINE (compare to
Tylenol® w/codeine) (age >12 years)
BUTALBITAL COMP. W/ CODEINE (age >12 years)
CODEINE SULFATE (age >12 years)
ENDOCET®
(oxycodone w/ acetaminophen)
HYDROCODONE (plain, w/acetaminophen, or
w/ibuprofen) (some exceptions apply) QTY LIMIT: Hydrocodone/APAP 12 tablets/day
HYDROMORPHONE tablets (compare to Dilaudid [®])
MORPHINE SULFATE OXYCODONE (plain)
OXYCODONE (w/acetaminophen, w/aspirin or
w/ibuprofen)
QTY LIMIT: Oxycodone/APAP 12 tablets/day
TRAMADOL
QTY LIMIT: 8 tablets/day (Age \geq 16)
TRAMADOL/APAP
<i>QTY LIMIT</i> : 8 tablets/day (Age \geq 18)
**NOTE: As of 5/1/21 a completed safety

OROMORPHONE tablets (compare to Dilaudid [®]) RPHINE SULFATE	Butorph QTY Demero
YCODONE (plain) YCODONE (w/acetaminophen, w/aspirin or w/ibuprofen) PTY LIMIT: Oxycodone/APAP 12 tablets/day MADOL PTY LIMIT: 8 tablets/day (Age ≥ 16)	Dilaudid Dilaudid Fentany Fentora
AMADOL/APAP QTY LIMIT: 8 tablets/day (Age ≥18)	Hydrom Meperid
NOTE: As of 5/1/21, a completed safety checklist must be submitted for new patients exceeding 90 MME per day, and existing patients exceeding 120 MME per day (applies to any combination of short and/or long acting opioids)	QTI Nucynta Oxycodd Oxymor Pentazod Seglentis Tramado

NON-PREFERRED AGENTS (PA required)

PA CRITERIA

ANALGESICS

RMAL PATCH

Outenza® Patch (capsaicin 8 %) OTY LIMIT: 4 patches/90 days ZtlidoTM Patch (lidocaine 1.8%) QTY LIMIT: 3 patches/day

(Note: Please refer to Analgesics: COX IIs and NSAIDs for topical NSAIDS)

Qutenza, **Ztlido:** diagnosis or indication is post-herpetic neuralgia AND patient has had a documented side effect, allergy, treatment failure or contraindication to 2 drugs in the tricyclic antidepressant (TCA) class and/or anticonvulsant class as well as Lidocaine 5% patch. OR patient has a medical necessity for transdermal formulation (ex. dysphagia, inability to take oral medications) AND patient has had a documented side effect, allergy, treatment failure or contraindication to lidocaine 5% patch.

Acetaminophen w/hydrocodone: all branded products *OTY LIMIT:* = 12 tablets/day

Acetaminophen w/oxycodone: all branded products *QTY LIMIT:* = 12 tablets/day

Actiq® (fentanyl lozenge on a stick: 200 mcg, 400 mcg, 600 mcg, 800 mcg, 1200 mcg, 1600 mcg)

Apadaz® (benzhydrocodone/APAP) OTY LIMIT: 12 tablets/day

Benzhydrocodone/APAP (compare to Apadaz®)

OTY LIMIT: 12 tablets/day nanol Nasal Spray Y LIMIT: 2 bottles/month

ol (meperidine)

d[®](hydromorphone) tablets

id-5[®](hydromorphone) oral solution

yl citrate transmucosal (compare to Actiq[®])

(fentanyl citrate buccal tablets)

norphone oral solution (compare to Dilaudid- $5^{\textcircled{R}}$)

Y LIMIT: 30 tablets/5-day supply per 30 days

a® (tapentadol) lone (plain) capsules

rphone (compare to Opana®)

cine w/naloxone

is® (celecoxib/tramadol) oral tablet

lol oral solution 5mg/ml

Note: The initial fill for all short-acting opiates will be limited to 50 Morphine Milligram Equivalents (MME) and 7-day supply for patients ≥ 18 years of age OR 24 MME and 3-day supply for patients \leq 17 years of age.

Butorphanol Nasal Spray: documented site effect, allergy, treatment failure, or contraindication to codeine, hydrocodone, morphine, & oxycodone (all 4 generic entities) as single or combination products. OR is unable to use tablet or liquid formulations.

Actiq, Fentanyl transmucosal, Fentora: indication of cancer breakthrough pain AND patient is opioid tolerant AND is on a long acting opioid formulation AND is 18 years of age or older (Actiq 16 years of age or older) AND prescriber is registered in the Transmucosal Immediate Release Fentanyl (TIRF) Risk Evaluation and Mitigation Strategy (REMS) Access program AND member has had a documented treatment failure with or intolerance to 2 of the following 3 immediate release treatment options: morphine. hydromorphone or oxycodone. OR is unable to use tablet or liquid formulations AND if the request is for brand name Actiq, member has a documented intolerance to generic fentanyl transmucosal.

Dilaudid - 5 Oral Solution, Hydromorphone Oral Solution: member has had a documented side effect, allergy or treatment failure with oxycodone oral solution and morphine oral solution OR has been started and stabilized on another dosage form of hydromorphone AND if the request is for the branded product, patient has a documented intolerance to the generic product.

Oxycodone (generic) Capsules: member has a documented intolerance to generic oxycodone tablets.

Seglentis: The patient has a documented side effect, allergy, or treatment failure with two or more preferred agents AND the patient is unable to take the individual components separately

Tramadol Oral Solution: patient has a medical necessity for a non-solid oral dosage form. (e.g. swallowing disorder).

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
Note: The FDA restricts the use of prescription codeine pain and cough medicines in children. Prior authorization is required for patients <12 years of age.		Ultracet: member has a documented intolerance to the generic formulation Other Short acting Opioids: member has had a documented side effect, allerg or treatment failure to at least 3 medications not requiring prior approval. (If product has an AB rated generic, one trial must be the generic.) PA requests to exceed daily cumulative MME limits:
		 Non-Opioid alternatives (up to a maximum dose recommended by the FDA) and Non-Pharmacological Treatments have been considered, an any appropriate treatments are documented in the patient's medical records. Such treatments may include, but are not limited to: NSAIDs, Acetaminophen, Acupuncture, Chiropractic, Physical Therapy. Vermont Prescription Monitoring System (VPMS) has been queried. Patient education and informed consent have been obtained, and a Controlled Substance Treatment Agreement is included in the patient's medical record. A reevaluation of the effectiveness and safety of the patient's pain management plan, including an assessment of the patient's adherence to the treatment regimen is completed no less than once every 90 days. Patient has a valid prescription for or states they are in possession of naloxone. Patients in nursing homes, receiving or eligible for hospice services, or those with chronic pain associated with cancer or cancer treatment are exempt from these requirements. Limitations: APAP containing products: daily doses that result in > 4 grams of acetaminophen/day will reject for PA
OPIOIDS: LONG ACTING		
TRANSDERMAL BUTRANS (buprenorphine) TRANSDERMAL SYSTEM QTY LIMIT: 4 patches/28 days (Maximum 28-day fill) FENTANYL PATCH (compare to Duragesic®) QTY LIMIT: 12 mcg/hr, 25 mcg/hr, 50 mcg/hr = 15 patches/30 days, 75 mcg/hr, 100 mcg/hr = 30 patches/30 days	Buprenorphine patch (compare to Butrans®) <i>QTY LIMIT:</i> 4 patches/28 days (Maximum 28-day Fill) Fentanyl patch 37.5 mcg/hr, 62.5 mcg/hr, 87.5 mcg/hr	CLINICAL CONSIDERATIONS: Long acting opioid dosage forms are intended for use in opioid tolerant patients only. These tablet/capsule/topical medication strengths may cause fatal respiratory depression when administered to patients not previously exposed to opioids. LA opioids should be prescribed for patients with a diagnosis or condition that requires a continuous, around-the-clock analgesic. LA opioids should be reserved for use in patients for whom alternative treatment options (e.g., non-opioid analgesics or immediate-release opioids) are ineffective, not tolerated, or would be otherwise inadequate to provide sufficient management of pain. LA opioids are NOT intended for use as 'prn' analgesic. LA opioids are NOT intended for use as 'prn' analgesic.
BUCCAL All products require PA ORAL MORPHINE SULFATE CR 12 hr tablet (compare to MS Contin®) QTY LIMIT: 90 tablets/strength/30 days	Belbuca® (buprenorphine hcl buccal film) <i>QTY LIMIT:</i> 56 films/28 days (Maximum 28-day fill) Conzip® (tramadol ER biphasic release) capsule <i>QTY LIMIT:</i> 1 capsule/day Hydromorphone XR tablet <i>QTY LIMIT:</i> 30 tablets/30 days (8 mg, 12 mg, 16 mg tabs)	indicated for pain in the immediate post-operative period (the first 12-24 hours following surgery) or if the pain is mild, or not expected to persist for an extended period of time. LA opioids are not intended to be used in a dosage frequency other than FDA approved regimens. Patients should not be using other extended release opioids prescribed by another physician. Prescribers should consult the VPMS (Vermont Prescription Monitoring System) to review a patient's Schedule II - IV medication use before prescribing long acting opioids. Belbuca Films, Buprenorphine Patch: the patient has had a documented

Methadone5 mg, 10 mg tablets

intolerance to Butrans patches

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
ORAL, ABUSE-DETERRENT FORMULATION XTAMPZA ER® (oxycodone ER) QTY LIMIT: 60 caps/strength/30days **NOTE: As of 5/1/21, a completed safety checklist must be submitted for new patients exceeding 90 MME per day, and existing patients exceeding 120 MME per day (applies to any combination of short and/or long acting opioids)**	Oxycodone ER (compare to OxyContin [®]) QTY LIMIT: 90 tablets/strength/30 days OxyContin [®] (Oxycodone ER) QTY LIMIT: 90 tablets/strength/30 days	Fentanyl patches 37.5mcg/hr, 62.5mcg/hr, 87.5mcg/hr: provider must submit clinical rationale detailing why the patient is unable to use a combination of the preferred strengths. Methadone Tablet: patient has had a documented side effect, allergy, or treatment failure to morphine sulfate CR 12 hr tablets AND the initial methadone daily dose does not exceed 30mg (Note: Methadone products, when used for treatment of opioid addiction in detoxification or maintenance programs, shall be dispensed ONLY by certified opioid treatment programs as stipulated in 42 CFR 8.12, NOT retail pharmacy.) Methadone Liquid: Patient must have a medical necessity for an oral liquid (i.e. swallowing disorder, inability to take oral medications) AND the initial daily dose does not exceed 30mg OR patient has been started and stabilized on the requested oral liquid medication. (Note: Methadone products, when used for treatment of opioid addiction in detoxification or maintenance programs, shall be dispensed ONLY by certified opioid treatment programs as stipulated in 42 CFR 8.12, NOT retail pharmacy.) Conzip, Tramadol ER biphasic-release Capsule, Tramadol ER biphasic-release Tablet, Tramadol ER/SR: member has had a documented treatment failure to a preferred short-acting tramadol product. In addition, for approval of tramadol ER biphasic-release capsule or tablet or the patient must have a documented intolerance to generic tramadol ER/SR. Oral Non-Preferred (except methadone & tramadol containing products): the patient has had a documented side effect, allergy, or treatment failure to morphine sulfate CR 12hr tablet (generic) AND generic fentanyl patch. (If a product has an AB rated generic, there must have been a trial of the generic). AND the patient must have a documented side effect, allergy, or treatment failure to the preferred abuse deterrent formulation (Xtampza ER) before OxyContin will be approved. Hysingla ER: Available with PA for those unable to tolerate any preferred medications. All requests will go to the DVHA Medical

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
		Patients in nursing homes, receiving or eligible for hospice services, or those with chronic pain associated with cancer or cancer treatment are exempt from these requirements. Limitations: Methadone 40mg dispersible tablet not approved for retail dispensing.
NSAIDS		
ORAL SINGLE AGENT DICLOFENAC POTASSIUM DICLOFENAC SODIUM ETODOLAC FLURBIPROFEN IBUPROFEN INDOMETHACIN INDOMETHACIN ER KETOPROFEN KETOROLAC QTY LIMIT: 20 doses/5 day supply every 90 day MECLOFENAMATE SODIUM MEFANAMIC ACID capsules MELOXICAM tabs NABUMETONE NAPROXEN 250 mg, 375 mg, 500 mg NAPROXEN ENTERIC COATED 375 mg, 500 mg NAPROXEN SODIUM 275mg, 550mg NAPROXEN SODIUM OTC 220 mg OXAPROZIN (compare to Daypro®) PIROXICAM (compare to Feldene®) SULINDAC	Cambia [®] (diclofenac potassium) packet for oral solution <i>QTY LIMIT</i> : 9 packets/month Daypro [®] (oxaprozin) Etodolac ER Feldene [®] (piroxicam) Fenoprofen 400 mg cap Fenoprofen 600 mg tab Indocin [®] (indomethacin) suspension Ketoprofen ER Lofena TM (diclofenac) tablet Meloxicam capsule (compare to Vivlodex®) Nalfon [®] (fenoprofen) 400 mg capsules Naproxen oral suspension Naproxen oral suspension Naproxen sodium ER Naproxen suspension 125mg/5ml Relafen® DS (nabumetone) Zipsor [®] (diclofenac potassium) Zorvolex [®] (diclofenac) Capsules <i>QTY LIMIT</i> : 3 capsules/day	 Arthrotec, diclofenac/misoprostol, Duexis: patient has a documented side effect or treatment failure to 2 or more preferred generic NSAIDs OR patient is not a candidate for therapy with a preferred generic NSAID mono-therapy due to one of the following: patient is 60 years of age or older, Patient has a history of GI bleed, Patient is currently taking an oral corticosteroid, Patient is currently taking methotrexate AND for approval of diclofenac/misoprostol, the patient must have a documented intolerance to brand Arthrotec Cambia: drug is being prescribed for treatment of acute migraine attacks AND patient has had a documented side effect or treatment failure to 2 or more preferred generic NSAIDs, one of which must be generic diclofenac OR drug is being prescribed for treatment of acute migraine attacks AND patient has a requirement for an oral liquid dosage form (i.e. swallowing disorder, inability to take oral medications) AND patient has had a documented side effect or treatment failure with the generic ibuprofen suspension. Celebrex: patient has had a documented intolerance to generic celecoxib. Pennsaid: patient has had a documented side effect or inadequate response to Diclofenac Patch, Licart: patient has had a documented side effect or inadequate response to Diclofenac gel or topical solution. Diclofenac Patch, Licart: patient has had a documented side effect or inadequate response to Diclofenac gel or topical solution AND patient has a documented intolerance to brand Flector Patch. Duexis, Ibuprofen/famotidine, naproxen/esomeprazole, Vimovo: patient is unable to take the individual components separately AND for approval of ibuprofen/famotidine or naproxen/esomeprazole, the patient must have a documented intolerance to the brand name equivalent.
ORAL COX-II Selective CELECOXIB QTY LIMIT: 2 caps/day INJECTABLE KETOROLAC Injection (formerly Toradol®)	Celebrex® (celecoxib) capsule QTY LIMIT: 2 caps/day Elyxyb TM (celecoxib) oral solution	Elyxyb: drug is being prescribed for treatment of acute migraine attacks AND patient has had a documented side effect or treatment failure to 2 or more preferred generic NSAIDs, one of which must be generic celecoxib OR drug is being prescribed for treatment of acute migraine attacks AND patient has a requirement for an oral liquid dosage form (i.e. swallowing disorder, inability to take oral medications) AND patient has had a documented side effect or treatment failure with the generic ibuprofen suspension. Lofena, Zipsor, Zorvolex: patient has had a documented side effect, allergy, or

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
NASAL SPRAY All products require PA TOPICAL DICLOFENAC (compare to Voltaren®) gel 1% DICLOFENAC 1.5 % Topical Solution	Sprix [®] (ketorolac) Nasal Spray <i>QTY LIMIT</i> : 5 bottles/5 days – once every 90 days Pennsaid® (diclofenac) 2% Topical Solution	treatment failure to 4 or more preferred generic NSAIDs, one of which must be generic diclofenac. Meloxicam Capsule: patient has had a documented side effect, allergy, or treatment failure to 4 or more preferred generic NSAIDs, one of which must be generic meloxicam tablet. Naproxen suspension: patient has a requirement for an oral liquid dosage form (i.e. swallowing disorder, inability to take oral medications) AND patient has had a documented side effect or treatment failure with generic ibuprofen suspension. Relafen DS: patient has had a documented side effect, allergy, or treatment failure to 4 or more preferred generic NSAIDs, one of which must be generic
TRANSDERMAL Flector® (diclofenae) 1.3 % Patch QTY LIMIT: 2 patches/day	Diclofenac (compare to Flector®) 1.3% Patch QTY LIMIT: 2 patches/day Licart® (diclofenac epolamine) 1.3% Patch QTY LIMIT: 1 patch/day	nabumetone. Sprix: indication or diagnosis is moderate to moderately severe pain. AND patient has had a documented inadequate response or intolerance to generic ketorolac tablets. OR patient has a documented medical necessity for the specialty dosage form (i.e. inability to take medication orally (NPO)). All other PA requiring NSAIDs: patient has had a documented side effect or treatment failure to 2 or more preferred generic NSAIDS. (If a product has an
NSAID/ANTI-ULCER All products require PA	Arthrotec [®] (diclofenac sodium w/misoprostol) Diclofenac sodium w/misoprostol (compare to Arthrotec [®])	AB rated generic, one trial must be the generic.) AND if the request is for a non-preferred extended release formulation, the patient has not been able to adhere to the dosing schedule of the immediate release formulation resulting in significant clinical impact.
Note: Please refer to "Dermatological: Actinic Keratosis Therapy" for Solaraze [®] or Diclofenac 3% Gel	Duexis [®] (ibuprofen/famotidine) QTY LIMIT: 3 tablets/day Ibuprofen/famotidine (compare to Duexis®) QTY LIMIT: 3 tablets/day Naproxen/esomeprazole (compare to Vimovo®) Vimovo® (naproxen/esomeprazole) QTY LIMIT: 2 tablets/day	

ANKYLOSING SPONDYLITIS: INJECTABLES

Length of Authorization: Initial PA 3 months; 12 months thereafter

INJECTABLE AVSOLA® (infliximab-axxq) biosimilar to Remicade® ENBREL® (etanercept) QTY LIMIT:50 mg = 4 syringes/28 days,

Preferred After Clinical Criteria Are Met

25 mg = 8 syringes/28 days HUMIRA[®] (adalimumab) *QTY LIMIT*:2 syringes/28 days Cimzia® (certolizumab pegol)

QTY LIMIT: 1 kit/28 days (starter X 1, then regular) Cosentyx® (secukinumab) Subcutaneous

Remicade[®] (infliximab) Renflexis[™] (infliximab-abda) biosimilar to Remicade[®] Rinvoq ® (upadactinib) extended release tablet

QTY LIMIT: 1 tablet/day

Clinical Criteria:

For all drugs: patient has a diagnosis of ankylosing spondylitis (AS) and has already been stabilized on the medication being requested. OR patient has a confirmed diagnosis of AS, and conventional NSAID treatment and DMARD therapy (e.g. methotrexate therapy) resulted in an adverse effect, allergic reaction, inadequate response, or treatment failure. If methotrexate is contraindicated, another DMARD should be tried.

Additional criteria for Taltz, Xeljanz, Xeljanz XR: the patient had a trial and failure or contraindication to a preferred TNF Inhibitor

Additional criteria for Cimzia, Cosentyx, Simponi: the prescriber must provide a clinically valid reason why at least 2 preferred agents cannot be

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
INFLECTRA® (infliximab-dyyb) biosimilar to Remicade® TALTZ® (ixekizumab) QTY LIMIT: 80 mg prefilled syringe or autoinjector = 2/28 days for the first month and 1/28 days subsequently ORAL XELJANZ® (tofacitinib) tablet QTY LIMIT: 2 tablets/day XELJANZ® XR (tofacitinib) tablet QTY LIMIT: 1 tablet/day Maximum 30 days supply	Simponi [®] (golimumab) Subcutaneous <i>QTY LIMIT</i> : 50 mg prefilled syringe or autoinjector = 1/28 days	used. Note: Patient must be ≥ 18 years of age for Simponi approval as safety and efficacy has not been established in pediatric patients. Additional criteria for Remicade, Renflexis: the prescriber must provide a clinically valid reason why at least 2 preferred agents cannot be used, and the patient must be unable to use Avsola or Inflectra. Additional Criteria for Rinvoq: the prescriber must provide a clinically valid reason why at least 2 preferred agents cannot be used, one of which must be Xeljanz or Xeljanz XR. * Patients with documented diagnosis of active axial involvement should have a trial with two NSAIDs, but a trial with DMARD is not required. If no active axial skeletal involvement, then NSAID trial and a DMARD trial are required (unless otherwise contraindicated).
	ANTI-ANXIETY: ANXIOL	YTICS

DESTA	ODIA	ZEDD	NT C
BENZ	ODIA	WAKIPII	NH.

CHLORDIAZEPOXIDE (formerly Librium[®])
CLONAZEPAM (compare to Klonopin[®])

QTY LIMIT: 4 tabs/day except 2 mg.
2 mg = 3 tabs/day
CLONAZEPAM ODT

QTY LIMIT: 4 tabs/day except 2 mg.
2 mg = 3 tabs/day
DIAZEPAM (compare to Valium[®])
LORAZEPAM (compare to Ativan[®])

QTY LIMIT: 4 tablets/day
OXAZEPAM

OTY LIMIT: 4 tablets/day Alprazolam ER, Alprazolam XR[®] (compare to Xanax XR[®]) QTY LIMIT: 2 tablets/day Alprazolam ODT OTY LIMIT: 3 tablets/day Alprazolam Intensol® (alprazolam concentrate) Ativan® (lorazepam) *QTY LIMIT:* 4 tablets/day Clorazepate tabs (compare to Tranxene T[®]) Diazepam Intensol® (diazepam concentrate) Klonopin[®] (clonazepam) QTY LIMIT: 4 tabs/day except 2 mg. 2 mg = 3 tabs/dayLorazepam Intensol® (lorazepam concentrate) Loreev XRTM (lorazepam extended release) Tranxene T[®] (clorazepate tablets) Valium® (diazepam) Xanax[®] (alprazolam) OTY LIMIT: 4 tablets/day

Alprazolam (compare to Xanax[®])

Non-preferred Benzodiazepines (except for Alprazolam ODT, Intensol Products, and Loreev XR): patient has a documented side effect, allergy, or treatment failure to at least 2 preferred benzodiazepine medications. (If a product has an AB rated generic, there must also be a trial of the generic formulation.)
Alprazolam ODT: patient has a documented side effect, allergy, or treatment failure to at least 2 preferred benzodiazepine medications. (If a product has an AB rated generic, there must also be a trial of the generic formulation). OR patient has a medical necessity for disintegrating tablet administration (i.e. inability to swallow tablets) AND patient has a documented

side effect, allergy or treatment failure to clonazepam ODT. **Alprazolam Intensol, Diazepam Intensol, and Lorazepam Intensol:** patient has

a medical necessity for the specialty dosage form (i.e. swallowing disorder). AND the medication cannot be administered by crushing oral tablets.

Loreev XR: The patient is receiving a stable dose of lorazepam tablets, evenly divided, three times daily AND medical reasoning for use beyond convenience or enhanced compliance is provided.

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
	Xanax XR [®] (alprazolam XR) <i>QTY LIMIT</i> : 2 tablets/day	
NON-BENZODIAZEPINE		
BUSPIRONE (formerly Buspar [®]) HYDROXYZINE HYDROCHLORIDE (formerly Atarax [®]) HYDROXYZINE PAMOATE (compare to Vistaril [®]) (all strengths except 100 mg) MEPROBAMATE	Hydroxyzine Pamoate (100 mg strength ONLY) (compare to Vistaril®) Vistaril® (hydroxyzine pamoate)	Hydroxyzine Pamote 100mg strength ONLY: patient is unable to use generic 50 mg capsules.Vistaril: patient has a documented intolerance to the generic formulation.
	ANTICOAGULANTS	
ORAL		
WARFARIN (compare to Coumadin [®]) DIRECT THROMBIN INHIBITOR PRADAXA [®] (dabigatran etexilate) QTY LIMIT: 2 capsules/day FACTOR XA INHIBITOR ELIQUIS [®] (apixaban) QTY LIMIT: 2 tablets/day QTY LIMIT: 5 mg = 4 tablets/day for 7 days if indication is treatment of DVT or PE (followed by 5 mg twice daily) XARELTO [®] (rivaroxaban) QTY LIMIT: 10 mg = 1 tablet/day QTY LIMIT: 15 mg and 20 mg = 1 tablet/day QTY LIMIT: 15 mg = 2 tablets/day for 21 days if indication is treatment of DVT or PE (followed by 20mg once daily)	Savaysa® (edoxaban) QTY LIMIT: 1 tablet/daily	Savaysa: creatinine clearance is documented to be < 95 ml/min AND prescribed has provided another clinically valid reason why generic warfarin, Pradaxa, Xarelto or Eliquis cannot be used. A yearly creatinine clearance is required with renewal of PA request
QTY LIMIT: Starter Pack (15 mg/20 mg) = 51 tablets/30days Preferred After Clinical Criteria Are Met XARELTO® (rivaroxaban) 2.5 mg QTY LIMIT: 2 tablets/day)		Xarelto 2.5 mg: Patient has a diagnosis of chronic coronary artery disease (CAD) peripheral artery disease (PAD) AND medication is being used concurrently waspirin.
INJECTABLE		

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
UNFRACTIONATED HEPARIN INJECTABLE HEPARIN LOW MOLECULAR WEIGHT HEPARINS INJECTABLE ENOXAPARIN (compare to Lovenox®) QTY LIMIT: 2 syringes/day calculated in ml volume SELECTIVE FACTOR XA INHIBITON INJECTABLE	Fragmin [®] (dalteparin) Lovenox [®] (enoxaparin) <i>QTY LIMIT</i> : 2 syringes/day calculated in ml volume Arixtra [®] (fondaparinux) Fondaparinux (compare to Arixtra®)	Arixtra, Fondaparinux, Lovenox and Fragmin: patient has a documented intolerance to generic enoxaparin AND if the request is for brand Arixtra, the patient must also have a documented intolerance to generic fondaparinux.
All products require PA	, , ,	

ANTICONVULSANTS

ORAL

Aptiom[®] (eslicarbazepine acetate) CARBAMAZEPINE tablets (compare to Tegretol[®]) OTY LIMIT: 200, 400 = 1 tab/dayCARBAMAZEPINE capsules (compare to Carbatrol®) CARBAMAZEPINE extended release (compare to 600 mg, 800 mg = 2 tabs/dayTegretol XR®) Banzel® (rufinamide) CELONTIN® (methsuxamide) tabs/dav CLOBAZAM (compare to Onfi®) *QTY LIMIT*: 10 mg = 3 tabs/day, 20 mg = 2tabs/day, oral suspension = 16mL/day (40mg/day) CLONAZEPAM (compare to Klonopin®) OTY LIMIT: 4 tablets/day CLONAZEPAM ODT (formerly Klonopin Wafers®) OTY LIMIT: 4 tablets/day DIAZEPAM (compare to Valium^(R)) DILVALPROEX SODIUM capsules (compare to Diacomit® (stiripentol) Depakote Sprinkles®) DIVALPROEX SODIUM (compare to Depakote®) suspension DIVALPROEX SODIUM ER (compare to Depakote ER®) EPITOL (carbamazepine) Felbamate (compare to Felbatol[®]) ETHOSUXAMIDE (compare to Zarontin[®]) Fintepla® (fenfluramine) oral solution GABAPENTIN 100 mg, 300 mg, 400 mg capsules. Felbatol® (felbamate) 600 mg, 800 mg tablets, 250 mg/5 ml oral solution

QTY LIMIT: 400 mg = 8 tabs/day, 200 mg = 16Banzel® (rufinamide) oral suspension OTY LIMIT: 80 ml/day (3,200 mg/day) Briviact® (brivaracetam) tablets, oral suspension Carbatrol® (carbamazepine) capsules Clorazepate (compare to Tranxene-T®) tablets Depakote[®] (divalproex sodium) Depakote ER[®] (divalproex sodium) Depakote Sprinkles[®] (divalproex sodium caps) Dilantin® (phenytoin) chewable tablets, capsules, ElepsiaTM (levetiracetam) extended release EprontiaTM (topiramate) oral solution

Criteria for approval of ALL non-preferred drugs: patient has been started and stabilized on the requested medication (Note: samples are not considered adequate justification for stabilization.) OR patient meets additional criteria outlined below.

Aptiom: the diagnosis is adjunctive therapy of partial-onset seizures and the patient has had a documented side effect, allergy, treatment failure/inadequate response or a contraindication to at least TWO preferred anticonvulsants, one of which is oxcarbazepine.

Banzel, Rufinamide: diagnosis or indication is treatment of Lennox-Gastaut Syndrome. AND patient has had a documented side effect, allergy, treatment failure/inadequate response or a contraindication to at least TWO preferred anticonvulsants used for the treatment of Lennox-Gastaut syndrome (topiramate, lamotrigine, valproic acid) AND for approval of the oral suspension, patient must have medical necessity for a specialty dosage form AND for approval of generic rufinamide, the patient must have a documented intolerance to brand Banzel.

Briviact: the diagnosis is adjunctive therapy of partial-onset seizures and the patient has had a documented side effect, allergy, treatment failure/inadequate response, or a contraindication to at least TWO preferred anticonvulsants, one of which is levetiracetam.

Carbatrol, Depakote, Depakote ER, Depakote Sprinkles, Dilantin, Keppra tablets or oral solution, Klonopin, Klonopin Wafers, Lamictal tablets or chew tablets, Lyrica, Mysoline, Neurontin capsules, tablets, solution, Onfi, Phenytek, Tegretol tablets, Tegretol XR (200 mg & 400 mg). Topamax tabs, Topamax sprinkles, Trileptal tablets, Trileptal oral suspension, Vimpat, Zarontin: patient has had a documented intolerance to

PREFERRED AGENTS	NON-PREFERRED AGENTS
(No PA required unless otherwise noted)	(PA required)
(compare to Neurontin®) GABITRIL® (tiagabine) LACOSAMIDE (compare to Vimpat®) tabs, solution LAMOTRIGINE chew tabs (compare to Lamictal® chew tabs) LAMOTRIGINE tabs (compare to Lamictal® tabs) LEVETIRACETAM tabs (compare to Keppra® tabs) LEVETIRACETAM oral solution (compare to Keppra® oral solution) LEVETIRACETAM ER (compare to Keppra XR®) OXCARBAZEPINE tablets (compare to Trileptal®) OXCARBAZEPINE oral suspension (compare to Trileptal®) PHENYTOIN (compare to Dilantin®) PHENYTOIN EX cap (compare to Phenytek®) PREGABALIN capsules (compare to Lyrica) QTY LIMIT:3 capsules/day PRIMIDONE (compare to Mysoline®) TEGRETOL® (carbamazepine) suspension TEGRETOL XR® (carbamazepine) 100 mg ONLY TOPIRAMATE tabs (compare to Topamax® tabs) TOPIRAMATE sprinkle caps (compare to Topamax® Sprinkles) VALPROIC ACID ZONISAMIDE Preferred After Clinical Criteria Are Met EPIDIOLEX® (cannabidiol) oral solution	Fycompa® (perampanel) tablets QTY IJMIT: 1 tablet/day Keppra®* (levetiracetam) tablets, oral solution Keppra XR® (levetiracetam extended release) Klonopin® (clonazepam) QTY IJMIT: 4 tablets/day Lamictal® tabs (lamotrigine tabs) Lamictal ODT® (lamotrigine orally disintegrating tablets Lamictal XR® tablets (lamotrigine extended release) Lamotrigine ER (compare to Lamictal XR®) Lamotrigine ODT (compare to Lamictal ODT®) Lyrica® (pregabalin) capsules QTY IJMIT: 3 capsules/day Lyrica® (pregabalin) oral solution Mysoline® (primidone) Neurontin® (gabapentin) capsules, tablets and solution Onfi® (clobazam) Oral Suspension 2.5 mg/ml QTY IJMIT: 16 ml/day Onfi® (clobazam) Tablets QTY IJMIT: 10 mg = 3 tabs/day, 20 mg = 2 tabs/day Oxtellar® XR (oxcarbazapine ER) tablet Pregabalin oral solution (compare to Lyrica®) Qudexy® XR (topiramate) capsules Sabril® (vigabatrin) Spritam® (levetiracetam) tablets for oral suspension Sympazan® (clobazam) films Rufinamide (compare to Banzel®) tablet, oral suspension QTY LIMIT: 400 mg = 8 tabs/day, 200 mg = 16 tabs/day, oral suspension = 80 ml/day (3200 mg/day) Tegretol® (carbamazepine) tablets

Topamax[®] (topiramate) Sprinkle Capsules

PA CRITERIA

the generic equivalent of the requested medication.

Clorazepate, Fycompa, Tranxene-T: diagnosis is adjunctive therapy of partialonset seizures OR diagnosis is adjunctive therapy for primary generalized tonic-clonic seizures (Fycompa only) AND the patient has had a documented side effect, allergy, treatment failure, inadequate response, or a contraindication to at least TWO preferred anticonvulsants. AND for approval of Tranxene-T the patient must have a documented intolerance to the generic equivalent.

Diacomit: Diagnosis or indication is treatment of Dravet Syndrome AND neutrophil and platelet counts have been obtained prior to starting therapy and are monitored periodically thereafter AND Patient is unable to tolerate or has had an inadequate response to valproate and clobazam AND medication will used concurrently with clobazam. **Note:** There are no clinical data to support the use of Diacomit as monotherapy.

Eprontia: The patient has a medical necessity for a specialty dosage form.
Epidiolex: The patient is unable to tolerate or has had an inadequate response to at least 2 of the following medications: clobazam, levetiracetam, valproate, lamotrigine, topiramate, rufinamide, or felbamate Note: This is processed via automated (electronic step therapy)

Felbamate, Felbatol: patient information/consent describing aplastic anemia and liver injury has been completed AND diagnosis is adjunctive therapy of partial-onset seizures or Lennox-Gastaut seizures and the patient has had a documented side effect, allergy, treatment failure/inadequate response or a contraindication to at least THREE preferred anticonvulsants. Additionally, if brand is requested, the patient has a documented intolerance to the generic product.

Fintepla: Diagnosis or indication is treatment of Dravet Syndrome AND patient has had a documented side effect, allergy, treatment failure/inadequate response or contraindication to at least two preferred anticonvulsants and Epidiolex AND prescriber, pharmacy and patient are registered with the REMS programs AND for reapproval, the patient must have a documented decrease from baseline in seizure frequency per 28 days.

Elepsia XR, Keppra XR, Lamictal XR, Lamotrigine ER, Oxtellar XR, Qudexy XR, Topiramate ER, Trokendi XR: patient has been unable to be compliant with or tolerate twice daily dosing of the immediate release product. Additionally, if brand Elepsia XR, Keppra XR or Lamictal XR is requested, the patient has a documented intolerance to the generic product. If topiramate ER sprinkle caps are requested, the patient must have a documented intolerance to Qudexy XR.

Lamictal ODT, Lamotrigine ODT: medical necessity for a specialty dosage form has been provided AND lamotrigine chewable tabs cannot be used. For approval of brand Lamictal ODT, the patient must have a documented intolerance to the generic equivalent.

Lyrica oral solution, Pregabalin oral solution: the patient is unable to use pregabalin capsules (i.e. swallowing disorder). For approval of brand Lyrica oral solution, the patient must have a documented intolerance to the generic

PREFERRED AGENTS	NON-PREFERRED AGENTS	
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	Topiramate ER sprinkle capsules (compare to Qudexy® XR) Tranxene-T® (clorazepate) tablets Trileptal® tablets (oxcarbazepine) Trileptal® oral suspension (oxcarbazepine) Trokendi XR® (topiramate SR 24hr) capsules QTY LIMIT: 200 mg = 2 caps/day, all other strengths = 1 cap/day Vigabatrin (compare to Sabril®) Vimpat® (lacosamide) tablets, oral solution Xcopri® (cenobamate) tablets QTY LIMIT: 200 mg = 2 tabs/day, all other strengths = 1 tab/day Zarontin® (ethosuximide) Ztalmy® (ganaxolone) suspension QTY LIMIT: 36 mL/day	equivalent. Spritam: medical necessity for a specialty dosage form has been provided AND patient must have a documented intolerance to levetiracetam oral solution. Sympazan: diagnosis or indication is adjunctive treatment of refractory epilepsy (may include different types of epilepsy) AND patient has had a documented side effect, allergy, treatment failure, inadequate response or a contraindication to at least TWO preferred anticonvulsants AND prescriber must provide a clinically compelling reason why the patient is unable to use Clobazam tablets AND Clobazam suspension Tiagabine generic: patient has had a documented intolerance to the brand name product. Sabril, Vigabatrin: prescriber and patient are registered with the REMS program AND diagnosis is infantile spasms OR patient is > 16 years old and the indication is adjunctive therapy in refractory complex partial seizures and failure of THREE other preferred anticonvulsants. Xcopri: the diagnosis is adjunctive therapy of partial-onset seizures AND the patient is ≥ 18 years of age AND the patient has had a documented side effect, allergy, treatment failure/inadequate response or a contraindication to at least TWO preferred anticonvulsants AND for reapproval, the patient must have a documented decrease from baseline in seizure frequency per 28 days. Ztalmy: Diagnosis or indication is for the treatment of seizures associated with cyclin-dependent kinase-like 5 (CDKL5) deficiency disorder (CDD) confirmed by genetic testing (results must be submitted) AND patient has had a documented side effect, allergy, treatment failure, inadequate response or a contraindication to at least TWO preferred anticonvulsants AND for reapproval, the patient must have a documented decrease from baseline in seizure frequency per 28 days. PA Requests to Exceed QTY LIMIT for clonazepam/clonazepam ODT or Klonopin: all requests will be referred to the DVHA Medical Director for review unless the patient has been started and stabilized on the requested quantity for treatment of a se
NASAL		qualitity for deadment of a serzure disorder.
NAYZILAM® (midazolam) nasal spray (age ≥ 12 years) QTY LIMIT: 10 units/30 days VALTOCO® (diazepam) nasal spray (age ≥ 6 years) QTY LIMIT: 20 units/30 days		
RECTAL		
DIAZEPAM (compare to Diastat®) rectal gel	Diastat® (diazepam) rectal gel	Diastat: patient has had a documented intolerance to the generic equivalent

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PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
	ANTIDEPRESSANTS	S
MAO INHIBITORS		
PHENELZINE SULFATE (compare to Nardil [®]) FDA maximum recommended dose = 90 mg/day TRANYLCYPROMINE FDA maximum recommended dose = 60 mg/day	Emsam [®] (selegiline) QTY LIMIT: 1 patch/day Marplan [®] (isocarboxazid) Nardil [®] (phenylzine) FDA maximum recommended dose = 90 mg/day	 Marplan: patient has been started and stabilized on the requested medication (Note: samples are not considered adequate justification for stabilization). OR patient has had a documented side effect, allergy, or treatment failure to phenelzine and tranylcypromine. Nardil: patient has had a documented intolerance to generic equivalent product. Emsam: patient has had a documented side effect, allergy, or treatment failure with at least 3 antidepressants from 2 of the major antidepressant classes (Miscellaneous, SNRIs, SSRIs, and Tricyclic Antidepressants). OR patient is unable to tolerate oral medication.
MISCELLANEOUS		
BUPROPION SR (compare to Wellbutrin SR®) FDA maximum recommended dose = 400mg/day BUPROPION XL (compare to Wellbutrin XL®) 150 mg, 300 mg FDA maximum recommended dose = 450 mg/day BUPROPION FDA maximum recommended dose = 450 mg/day MAPROTILINE FDA maximum recommended dose = 225 mg/day MIRTAZAPINE (compare to Remeron®) FDA maximum recommended dose = 45 mg/day MIRTAZAPINE RDT (compare to Remeron Sol-Tab®) FDA maximum recommended dose = 45 mg/day TRAZODONE HCL (formerly Desyrel®) FDA maximum recommended dose = 600 mg/day VIIBRYD® (vilazodone) Tablet (Age ≥ 18 years) FDA maximum recommended dose = 40 mg/day	Aplenzin® (bupropion hydrobromide) ER tablets QTY LIMIT: 1 tablet/day Auvelity™ (bupropion/dextromethorphan) QTY LIMIT: 2 tablets/day Bupropion XL 450mg (compare to Forfivo XL®) QTY LIMIT: 1 tablet/day FDA maximum recommended dose = 450 mg/day Forfivo XL® (bupropion SR 24hr) 450 mg tablet QTY LIMIT: 1 tablet/day FDA maximum recommended dose = 450 mg/day Nefazodone FDA maximum recommended dose = 600 mg/day Remeron® (mirtazapine) FDA maximum recommended dose = 45 mg/day Remeron Sol Tab® (mirtazapine RDT) FDA maximum recommended dose = 45 mg/day Spravato® (esketamine) nasal spray QTY LIMIT: not to exceed FDA recommended dose and frequency for corresponding timeframe Trintellix® (vortioxetine) Tablet QTY LIMIT: 1 tablet/day Vilazodone (compare to Viibryd®) QTY LIMIT: 1 tablet/day FDA maximum recommended dose = 40 mg/day Wellbutrin SR® (bupropion SR) FDA maximum recommended dose = 400 mg/day Wellbutrin XL® (bupropion XL)	Criteria for approval for ALL non-preferred drugs: The patient has been started and stabilized on the requested medication. (Note: samples are not considered adequate justification for stabilization.) OR The patient meets additional criteria as outlined below. Aplenzin, Auvelity: The patient is ≥ 18 years of age AND The patient has had a documented side effect, allergy, or inadequate response to at least 3 different antidepressants from the SSRI, SNRI and/or Miscellaneous Antidepressant categories (may be preferred or non-preferred), one of which must be bupropion. Bupropion XL 450mg, Forfivo XL: The patient is unable to take the equivalent dose as generic bupropion XL (150mg & 300mg) AND for approval of brand, the patient must have a documented intolerance to the generic equivalent. Nefazodone: The patient is ≥ 18 years of age AND The patient has had a documented side effect, allergy, or inadequate response to at least 3 different antidepressants from the SSRI, SNRI and/or Miscellaneous Antidepressant categories (may be preferred or non-preferred) Remeron, Remeron SolTab, Wellbutrin SR, and Wellbutrin XL: The patient has had a documented intolerance to the generic formulation of the requested medication. Spravato: Diagnosis is treatment resistant depression: the patient is ≥ 18 years of age AND medication is being used as adjunct treatment with an oral antidepressant AND the patient has a documented treatment failure (defined by at least 8 weeks of therapy) with at least 2 different antidepressants from the SSRI, SNRI, and/or Miscellaneous Antidepressant categories (may be preferred or non-preferred) AND the healthcare site and patient are enrolled in the Spravato® REMS program. Initial approval will be granted for 3 months. For re-approval after 3 months, the patient must have documented improvement in symptoms. Diagnosis is Major Depressive Disorder (MDD) with Acute Suicidal Ideation or Behavior: the patient is ≥ 18 years of age AND the medication is being used as

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PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
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SNRI	FDA maximum recommended dose = 450 mg/day Zulresso TM (brexanolone) intravenous solution	adjunct treatment with an oral antidepressant AND the healthcare site and patient are enrolled in the Spravato® REMS program. Approval will be granted for 4 weeks. Trintellix: The patient is ≥ 18 years of age AND The diagnosis or indication is MDD AND The patient has had a documented side effect, allergy, or inadequate response (defined by at least 8 weeks of therapy) to at least 2 different antidepressants from the SSRI, SNRI, and/or Miscellaneous Antidepressant categories (may be preferred or non-preferred). Vilazodone: Patient is ≥ 18 years of age AND The patient has had a documented intolerance to brand Viibryd. Zulresso: Patient is ≥ 18 years of age and ≤ 6 months postpartum AND patient has a diagnosis of postpartum depression (PPD) with documented onset of symptoms occurring in the third trimester or within 4 weeks of delivery AND the patient has a documented treatment failure (defined by at least 8 weeks of therapy) with two different oral antidepressants unless contraindicated or documentation shows that the severity of depression would place the health of the mother or infant at significant risk AND the pharmacy, patient, and healthcare facility are enrolled in the REMS program. Note: Zulresso™ will be approved as a medical benefit ONLY and will NOT be approved if billed through pharmacy point of sale. Note: After a 4-month lapse in use of a non-preferred agent for a mental health indication, or if there is a change in therapy, a lookback through claims information will identify the need to re-initiate therapy following the PDL and clinical criteria.
DULOXETINE (compare to Cymbalta [®]) capsule <i>QTY LIMIT</i> : 2 capsules/day FDA maximum recommended dose = 120 mg/day (MDD and GAD), 60 mg/day all others VENLAFAXINE ER capsule (compare to Effexor XR [®]) <i>QTY LIMIT</i> : 37.5 mg and 75 mg = 1 capsule/day FDA maximum recommended dose = 225 mg/day VENLAFAXINE IR tablet FDA maximum recommended dose = 225 mg/day	Cymbalta [®] (duloxetine) capsule <i>QTY LIMIT</i> : 2 capsules/day FDA maximum recommended dose = 120 mg/day (MDD and GAD), 60 mg/day all others Desvenlafaxine base SR <i>QTY LIMIT</i> : 50 mg tablet only = 1 tablet/day FDA maximum recommended dose = 400 mg/day Desvenlafaxine succinate ER (compare to Pristiq®) <i>QTY LIMIT</i> : 50 mg tablet only = 1 tablet/day FDA maximum recommended dose = 400 mg/day Drizalma® (duloxetine) sprinkle capsule QTY LIMIT: 2 capsules/day FDA maximum recommended dose = 120 mg/day (MDD and GAD), 60 mg/day all others Effexor XR [®] (venlafaxine XR) capsule <i>QTY LIMIT</i> : 37.5 mg and 75 mg = 1 capsule/day FDA maximum recommended dose = 225 mg/day	Criteria for approval of ALL non-preferred drugs: The patient has been started and stabilized on the requested medication. (Note: samples are not considered adequate justification for stabilization.) OR The patient meets additional criteria as outlined below. Venlafaxine ER tablet (generic), Effexor XR Capsule (brand), Desvenlafaxine ER succinate, Pristiq: The patient has had a documented intolerance to generic venlafaxine ER caps AND if the request is for Pristiq, the patient has a documented intolerance to the generic. Desvenlafaxine SR (base), Fetzima: The patient has had a documented side effect, allergy, or inadequate response to at least 2 different antidepressants AND The patient has had a documented intolerance with generic desvenlafaxine succinate ER. Cymbalta, Drizalma: There must be a clinically compelling reason why the dosing needs cannot be accomplished with generic duloxetine. Note: After a 4-month lapse in use of a non-preferred agent for a mental health indication, or if there is a change in therapy, a lookback through claims information will identify the need to re-initiate therapy following the PDL and clinical criteria.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
	Fetzima [®] (levomilnacipran ER) capsule <i>QTY LIMIT</i> : 1 capsule/day FDA maximum recommended dose = 120 mg/day Fetzima [®] (levomilnacipran ER) capsule titration pack <i>QTY LIMIT</i> : 1 pack per lifetime FDA maximum recommended dose = 120 mg/day Pristiq [®] (desvenlafaxine succinate SR) <i>QTY LIMIT</i> : 50 mg tablet only = 1 tablet/day FDA maximum recommended dose = 400 mg/day Venlafaxine ER [®] tablet <i>QTY LIMIT</i> : 37.5 mg and 75 mg = 1 tablet/day FDA maximum recommended dose = 225 mg/day	
SSRIs		
CITALOPRAM (compare to Celexa®) tablets, solution FDA maximum recommended dose = 40 mg/day ESCITALOPRAM (compare to Lexapro®) tablets FDA maximum recommended dose = 20mg/day FLUOXETINE (compare to Prozac®) capsules, tablets, solution FDA maximum recommended dose = 80 mg/day FLUVOXAMINE FDA maximum recommended dose = 300 mg/day PAROXETINE hydrochloride tablet (compare to Paxil®) FDA maximum recommended dose = 60 mg/day SERTRALINE (compare to Zoloft®) tablet, solution FDA maximum recommended dose = 200 mg/day,	Brisdelle (paroxetine mesylate) QTY LIMIT: 1 capsule/day Celexa (citalopram) FDA maximum recommended dose = 40 mg/day Escitalopram solution FDA maximum recommended dose = 20 mg/day Fluoxetine 90 mg FDA maximum recommended dose = 90 mg/week Fluvoxamine CR QTY LIMIT: 2 capsules/day FDA maximum recommended dose = 300 mg/day Lexapro (escitalopram) QTY LIMIT: 5 mg and 10 mg tablets = 1.5 tabs/day FDA maximum recommended dose = 20mg/day Paroxetine mesylate (compare to Brisdelle (compare to Brisdelle)) QTY LIMIT: 1 capsule/day Paroxetine CR (compare to Paxil CR (compare) FDA maximum recommended dose = 75 mg/day Paxil (paroxetine) FDA maximum recommended dose = 60 mg/day Paxil CR (paroxetine) FDA maximum recommended dose = 75 mg/day Paxil CR (paroxetine CR) FDA maximum recommended dose = 75 mg/day Pexeva (paroxetine) FDA maximum recommended dose = 60 mg/day Prozac (fluoxetine)	Celexa, Fluvoxamine CR, Lexapro, Paxil tablet, Pexva, Paroxetine CR, Paxil CR, Prozac, Zoloft: The patient had a documented side effect, allergy, or treatment failure with 2 preferred SSRIs. One trial must be the generic formulation or IR formulation if CR formulation requested. Brisdelle, Paroxetine mesylate: The indication for use is moderate to severe vasomotor symptoms (VMS) associated with menopause. AND The patient has tried and failed generic paroxetine hydrochloride. Paxil suspension, Escitalopram solution: The patient has a requirement for an oral liquid dosage form. AND The patient had a documented side effect, allergy, or treatment failure with 2 preferred liquid SSRI formulations. Fluoxetine 90mg: The patient has been started and stabilized on the requested medication. (Note: samples are not considered adequate justification for stabilization.) OR The prient failed and is not a candidate for daily fluoxetine. AND The prescriber provides clinically compelling rationale for once-weekly dosing. Sertraline capsules: Prescriber must provide a clinically compelling reason why the patient is unable to use tablets. Note: After a 4-month lapse in use of a non-preferred agent for a mental health indication, or if there is a change in therapy, a lookback through claims information will identify the need to re-initiate therapy following the PDL and clinical criteria.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
TRICYCLICS AMITRIPTYLINE FDA maximum recommended dose = 300 mg/day AMOXAPINE DOXEPIN capsules, solution IMIPRAMINE FDA maximum recommended dose = 300 mg/day NORTRIPTYLINE (compare to Pamelor®) NORTRIPTYLINE Oral Solution	FDA maximum recommended dose = 80 mg/day Sertraline capsule 150 mg, 200 mg QTY LIMIT: 1 capsule/day Zoloft® (sertraline) QTY LIMIT: 25 mg and 50 mg tablets = 1.5 tabs/day FDA maximum recommended dose = 200 mg/day Anafranil® (clomipramine) Clomipramine (compare to Anafranil®) Imipramine Pamoate capsules Desipramine (compare to Norpramin®) Norpramin® (desipramine) Pamelor® (nortriptyline) Protriptyline Trimipramine (compare to Surmontil®)	Criteria for approval of ALL non-preferred drugs: patient has been started and stabilized on the requested medication. (Note: samples are not considered adequate justification for stabilization.) OR the patient meets additional criteria as outlined below. Imipramine Pamoate: The patient has had a documented side effect, allergy, or treatment failure to 3 preferred TCAs, one of which must be imipramine tablets. Desipramine: The patient has had a documented side effect, allergy, or treatment failure to nortriptyline. Clomipramine: The patient has had a documented side effect, allergy, or treatment failure to 2 or more preferred TCAs OR patient has a diagnosis of obsessive-compulsive disorder AND has had a documented side effect, allergy, or treatment failure to 2 SSRIs. All other non-preferred agents: The patient has had a documented side effect, allergy, or treatment failure to 2 or more preferred TCAs. One trial must be the AB rated generic formulation if available Limitation: Chlordiazepoxide/amitriptyline and amitriptyline/perphenazine combinations are not covered. Generic agents may be prescribed separately.
	ANTI-DIABETICS	
ALPHA-GLUCOSIDASE INHIBITORS		
ACARBOSE (compare to Precose [®]) MIGLITOL BIGUANIDES & COMBINATIONS	Precose [®] (acarbose)	Precose: patient must have a documented intolerance to generic acarbose
SINGLE AGENT		
METFORMIN (compare to Glucophage [®]) METFORMIN XR (compare to Glucophage XR [®])	Fortamet [®] (metformin ER Osmotic) Glumetza [®] (metformin ER modified release) Metformin ER modified release (compare to Glumetza) Metformin oral solution (compare to Riomet®) Metformin ER Osmotic (compare to Fortamet [®]) Riomet [®] (metformin oral solution)	Fortamet, Glumetza, Metformin ER mod release, Metformin ER osmotic: patient has had a documented intolerance to generic metformin XR (if product has an AB rated generic, there must have been a trial of the generic) Metformin oral solution, Riomet: prescriber provides documentation of medical necessity for the specialty dosage form (i.e. inability to swallow tablets, dysphagia)

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
COMBINATION GLIPIZIDE/METFORMIN GLYBURIDE/METFORMIN		
DIPEPTIDYL PEPTIDASE (DPP-4) INHIBITOR	S	
Preferred After Clinical Criteria Are Met SINGLE AGENT JANUVIA® (sitagliptin) QTY LIMIT: 1 tab/day TRADJENTA® (linagliptin) QTY LIMIT: 1 tab/day COMBINATION JANUMET® (sitagliptin/metformin) QTY LIMIT: 2 tabs/day JANUMET XR® (sitagliptin/metformin ER) QTY LIMIT: 50/500 and 100/1000 mg = 1 tab/day, 50/1000 mg = 2 tabs/day JENTADUETO® (linagliptin/metformin) QTY LIMIT: 2 tabs/day	Non-Preferred After Clinical Criteria Are Met Alogliptan (compare to Nesina®) QTY LIMIT: 1 tab/day Nesina® (alogliptin) QTY LIMIT: 1 tab/day Onglyza® (saxagliptin) QTY LIMIT: 1 tab/day Jentadueto XR (linagliptan/metformin ER) QTY LIMIT: 1 tab/day Kazano® (alogliptin/metformin) QTY LIMIT: 1 tab/day Kombiglyze XR® (saxagliptin/metformin ER) QTY LIMIT: 1 tab/day Oseni® (alogliptin/pioglitazone) QTY LIMIT: 1 tab/day	 Januvia, Tradjenta: patient has had a documented side effect, allergy, contraindication OR treatment failure with metformin Alogliptan, Nesina, Onglyza: patient has had a documented side effect, allergy, contraindication OR treatment failure with metformin AND patient has had a documented side effect, allergy OR treatment failure with at least one preferred DPP-4 agent. Janumet, Janumet XR: patient has had an inadequate response with Januvia Of Metformin/Metformin XR monotherapy OR patient has been started and stabilized on Januvia and Metformin/Metformin XR combination therapy. Kazano, Kombiglyze XR: patient has had a documented side effect, allergy OR treatment failure with at least one preferred DPP-4 combination agent. Jentadueto XR: patient is unable to take Tradjenta in combination with Metformin XR as the individual separate agents. Jentadueto: patient has had an inadequate response with Tradjenta OR Metformin monotherapy OR patient has been started and stabilized on Tradjenta and Metformin combination therapy. Oseni: patient is unable to take Nesina and Actos (pioglitazone) as the individual separate agents (after meeting clinical criteria for each individual agent)
HYPOGLYCEMIA TREATMENTS		
BAQSIMI® (glucagon nasal powder) 3mg QTY LIMIT: 2 devices/28 days GLUCAGEN® HYPOKIT® (glucagon for injection) 1mg ZEGALOGUE® (dasiglucagon SC injection) 0.6 mg QTY LIMIT: 2 prefilled syringes or auto- injectors/28 days	Glucagon emergency kit Gvoke TM (glucagon SC injection) prefilled syringe, auto- injector 0.5mg, 1mg	Glucagon Emergency Kit, Gvoke: Patient has recurrent episodes of symptomatic or severe hypoglycemia (<55 mg/dL) requiring the assistance of another individual AND the preferred formulations would not be suitable alternatives.
INSULINS		
RAPID-ACTING INJECTABLE HUMALOG® (insulin lispro) INSULIN ASPART (compare to Novolog® NOVOLOG® (insulin aspart)	Admelog® (insulin lispro) Afrezza ® Inhaled (insulin human) Apidra® (insulin glulisine) Fiasp® (insulin aspart) Insulin Aspart (compare to Novolog®) Insulin Lispro (compare to Humalog®)	 Admelog, Fiasp, Insulin Lispro, Lyumjev: Both Humalog and Novolog must be on a long-term backorder and unavailable from the manufacturer. Apidra, Humulin R (U-100), Novolin R: patient has been started and stabilized on the requested medication. (Note: samples are not considered adequate justification for stabilization.) OR patient has had a documented side effect, allergy OR treatment failure to Novolog or Humalog. Humulin N, Novolin N: patient has been started and stabilized on the requested

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
SHORT-ACTING INJECTABLE HUMULIN R® U-500 INTERMEDIATE-ACTING INJECTABLE All products require PA LONG-ACTING ANALOGS INJECTABLE LANTUS® (insulin glargine) LEVEMIR® (insulin detemir) TOUJEO® (insulin glargine) TRESIBA® (insulin degludec) MIXED INSULINS INJECTABLE NOVOLOG MIX 70/30® (Protamine/Aspart) HUMALOG MIX 50/50® (Protamine/Lispro) HUMALOG MIX 75/25® (Protamine/Lispro)	Lyumjev® (insulin lispro-aabc) Humulin R® (Regular) U-100 Novolin R® (Regular) U-100 Humulin N® (NPH) Novolin N® (NPH) Basaglar® (insulin glargine) Semglee® (insulin glargine) Toujeo® Max (insulin glargine) Insulin Aspart Protamine/Aspart 70/30 (compare to Novolog Mix 70/30®) Humulin 70/30® (NPH/Regular) Novolin 70/30® (NPH/Regular)	medication. (Note: samples are not considered adequate justification for stabilization.) OR patient has a documented treatment failure to at least one preferred long-acting agent (Lantus or Levemir). Humulin 70/30, Insulin Aspart Protamine/Aspart 70/30, Novolin 70/30: patient has been started and stabilized on the requested medication. (Note: samples are not considered adequate justification for stabilization.) OR patient has had a documented side effect, allergy or treatment failure to Novolog Mix or Humalog Mix. Toujeo Max: The patient is currently using insulin glargine 300 units/mL AND the dose exceeds 160 units. Basaglar, Semglee: Diagnosis of diabetes mellitus AND Lantus must be on a long-term backorder and unavailable from the manufacturer. AFREZZA INHALED INSULIN: Baseline PFT with FEV1 ≥ 70 % predicted Patient does not have underlying lung disease (Asthma, COPD) Patient is a non-smoker or has stopped smoking more than six months prior to starting Afrezza Patient is currently using a long-acting insulin Patient has failed to achieve HbA1c goal (defined as ≤ 7%) on a shortacting insulin in combination with a long-acting insulin Initial approval is for 3 months and improved glycemic control must be documented for further approvals
MEGLITINIDES SINGLE AGENT NATEGLINIDE REPAGLINIDE PEPTIDE HORMONES: GLP-1 RECEPTOR AGENTAL AGENTAL AGENTAL AGENT	GONISTS	
Preferred After Clinical Criteria Are Met SINGLE AGENTS OZEMPIC® (semaglutide) QTY LIMIT: 9mL/84 days TRULICITY® (dulaglutide) QTY LIMIT: 12 pens/84 days VICTOZA® (liraglutide) QTY LIMIT: 9 pens/90 days	Adlyxin® (lixisenatide) Bydureon® BCise TM (exenatide extended-release) QTY LIMIT: 12 pens/84 days Byetta® (exenatide) QTY LIMIT: 3 pens/90 days Mounjaro TM (tirzepatide) QTY LIMIT: 4 pens/28 days Rybelsus® (semaglutide) tablets QTY LIMIT: 1 tablet/day	Clinical criteria for all drugs: patient has a diagnosis of Type 2 Diabetes Mellitus Additional criteria for Adlyxin/Byetta/Bydureon BCise, Mounjaro: patient has a documented side effect, allergy, contraindication, or treatment failure with two preferred GLP-1 Receptor Agonists. Treatment failure is defined as < 1% reduction in HbA1c after 12 weeks at the maximally tolerated dose. Additional criteria for Rybelsus patient has a documented side effect, allergy, contraindication, or treatment failure with one preferred SGLT2 inhibitor AND patient has a documented side effect, allergy, contraindication, or treatment failure with two preferred GLP-1 Receptor Agonists, one of which must be Ozempic, or has a clinically valid reason for being unable to administer an injection (e.g. visual impairment, impaired dexterity). Treatment failure is

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PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
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		defined as < 1% reduction in HbA1c after 12 weeks at the maximally tolerated
COMBINATION AGENTS	Soliqua [®] (insulin glargine/lixisenatide)	dose. Soliqua/Xultophy: patient has a documented side effect, allergy, contraindication,
All products require PA	QTY LIMIT: 3 pens/25 days	or treatment failure with at least one preferred GLP-1 Receptor Agonist used in
	Xultophy® (insulin degludec/liraglutide)	combination with Lantus or Levemir. Treatment failure is defined as $<1\%$
AMYLINOMIMETICS AND ADDRESS OF THE PARTY OF	Symlin [®] (pramlintide)	reduction in HbA1c after 12 weeks at the maximally tolerated dose.
All products require PA		Symlin: patient is at least 18 years of age AND patient is on insulin.
SODIUM-GLUCOSE CO-TRANSPORTER 2 (SG	LT2) INHIBITORS AND COMBINATIONS	
SINGLE AGENTS		Steglatro: Patient has a documented side effect, allergy, or contraindication to
FARXIGA® (dapagliflozin)	Steglatro® (ertugliflozin)	two preferred SGLT2 inhibitors.
QTY LIMIT: 1 tab/day	QTY LIMIT: 1 tab/day	Invokamet XR/Segluromet/ Synjardy XR additional criteria: The patient has documentation of a failure of therapy with a preferred SGLT2 inhibitor used
INVOKANA® (canagliflozin)		in combination with metformin/metformin XR.
QTY LIMIT: 1 tab/day		Glyxambi/Qtern/Steglujan additional criteria: The patient has documentation
JARDIANCE (empagliflozin) <i>QTY LIMIT</i> : 1 tab/day		of a failure of therapy with the combination of a preferred SGLT2 inhibitor plus a preferred DPP-4 inhibitor
Q11 121/111. 1 uto/daty		Trijardy XR: patient has documentation of a failure of therapy with a preferred
COMBINATIONS AGENTS		SGLT2 inhibitor, a preferred DDP-4 inhibitor and metformin/metformin XR
INVOKAMET® (canagliflozin/metformin) QTY LIMIT: 1 tab/day	Glyxambi® (empagliflozin/ linagliptin) QTY LIMIT: 1 tab/day	used in combination.
SYNJARDY® (empagliflozin/metformin)	Invokamet® XR (canagliflozin/metformin ER)	
QTY LIMIT: 2 tabs/day	Qtern® (dapagliflozin/saxagliptin)	
XIGDUO XR® (dapagliflozin & metformin ER)	Segluromet® (ertugliflozin/metformin)	
QTY LIMIT: $5/1000 \text{ mg} = 2/\text{day}$, all other strengths	QTY LIMIT: 2 tabs/day Steglujan® (ertugliflozin/sitagliptin)	
= 1/day	QTY LIMIT: 1 tab/day	
	Synjardy [®] XR (empagliflozin/metformin ER) <i>QTY LIMIT</i> : 1 tab/day	
	Trijardy® XR (empagliflozin/linagliptin/metformin ER)	
SULFONYLUREAS 2 ND GENERATION		
GLIMEPIRIDE (compare to Amaryl)	Amaryl [®] (glimepiride)	
GLIPIZIDE (compare to Amaryl) GLIPIZIDE (compare to Glucotrol®)	Glucotrol® (glipizide)	Criteria for Approval: Patient must have a documented side effect, allergy or treatment failure to two preferred sulfonylureas. If a product has an AB rated
GLIPIZIDE (compare to Glucotrol ") GLIPIZIDE ER (compare to Glucotrol XL®)	Glucotrol XL [®] (glipizide ER)	generic, one trial must be the generic.
GLYBURIDE	Glynase [®] (glyburide micronized)	-
GLYBURIDE MICRONIZED		
THIAZOLIDINEDIONES & COMBINATIONS		
Preferred After Clinical Criteria Are Met		Actos, Pioglitazone: Patient has been started and stabilized on the requested

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
PIOGLITAZONE (compare to Actos [®]) COMBINATION All products require PA	Actos [®] (pioglitazone) Actoplus Met [®] (pioglitazone/metformin) Duetact [®] (pioglitazone/glimepiride) <i>QTY LIMIT</i> : 1 tablet/day Pioglitazone/Glimepiride (compare to Duetact®) <i>QTY LIMIT</i> : 1 tablet/day Pioglitazone/Metformin (Compare to Actoplus Met)	medication OR patient has had a documented side effect, allergy, contraindication OR treatment failure with metformin AND for approval of Actos, the patient has a documented intolerance to the generic equivalent. Actoplus Met, Duetact, Pioglitazone/Metformin, Pioglitazone/Glimepiride: patient is unable to take as the individual separate agents AND if the request is for Actoplus Met or Duetact, the patient has had a documented intolerance to the generic equivalent.

ANTI-EMETICS

5HT3 ANTAGONISTS: Length of Authorization: 6 months for chemotherapy or radiotherapy; 3 months for hyperemesis gravadarum, 1 time for prevention of post-op nausea/vomiting: see clinical criteria. Monthly quantity limits apply, PA required to exceed.

ONDANSETRON tablet OYY LIMIT: 3 tabs/day, maximum of 30 days per fill ONDANSETRON ODT OYY LIMIT: 3 tabs/day, maximum of 30 days per fill ONDANSETRON oral solution 4mg/5mL Akynzeo® (nutupitant/palonosetron) Granisetron 1 mg OYY LIMIT: 6 tabs/28 days Granisetron injectable Sancuso® 3.1 mg/24 hr transdermal patch OYY LIMIT: 4 patches/28 days Sustol® (granisetron) injection 10 mg/0.4t OYY LIMIT: 4 mj = 12 tabs/28 days Zofran® (ondansetron) oral tablets OYY LIMIT: 4 mg = 12 tabs/28 days Zuplenz® (ondansetron) oral soluble film OYY LIMIT: 4 mg = 12 films/28 days films/28 days	chemotherapy. AND patient has had a documented side effect, allergy, or treatment failure to generic ondansetron. Zofran: patient must have a documented intolerance to generic formulation. Sancuso: patient has a diagnosis of nausea and vomiting associated with cancer chemotherapy. AND prescriber provides documentation of medical necessity for the transdermal formulation. OR patient has had a documented side effect, allergy, or treatment failure with generic ondansetron.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
MISCELLANEOUS (PREGNANCY) DICLEGIS® (10 mg doxylamine succinate and 10 mg pyridoxine hydrochloride) DR tablet QTY LIMIT: 4 tablets/day	Bonjesta® (20 mg doxylamine succinate and 20 mg pyridoxine hydrochloride ER tablet) QTY LIMIT: 2 tablets/day Doxylamine succinate/pyridoxine hydrochloride DR tablet (compare to Diclegis®) QTY LIMIT: 4 tablets/day	for each day of chemotherapy and 2 tablets for 2 days after completion of chemotherapy may be approved. OR For nausea and vomiting associated with radiation therapy, 2 tablets for each day of radiation may be approved. Sancuso: For nausea and vomiting associated with chemotherapy, 1 patch for each chemotherapy cycle may be approved. Limitations: Aloxi is not considered an outpatient medication and is not covered in the pharmacy benefit. Bonjesta, Doxylamine/Pyridoxone: patient has a documented intolerance to Diclegis.
NK1 ANTAGONISTS		
CINVANTI® (aprepitant) injection EMEND® (fosaprepitant) injection Preferred After Clinical Criteria Are Met EMEND® (aprepitant) 80 mg QTY LIMIT: 2 caps/28 days EMEND® (aprepitant) Tri-fold Pack QTY LIMIT: 1 pack/28 days	Aprepitant (compare to Emend®) 40 mg QTY LIMIT: 1 cap/28 days Aprepitant (compare to Emend®) 80 mg QTY LIMIT: 2 caps/28 days Aprepitant (compare to Emend®) 125 mg QTY LIMIT: 1 cap/28 days Emend® (aprepitant) oral suspension Varubi® (rolapitant) QTY LIMIT: 4 tabs/28 days	Aprepitant, Emend (aprepitant): medication will be prescribed by an oncology practitioner. AND patient requires prevention of nausea and vomiting associated with moderate to highly emetogenic cancer chemotherapy. AND The requested quantity does not exceed one 125 mg and two 80 mg capsules OR one Tri-Fold Pack per course of chemotherapy. Patients with multiple courses of chemotherapy per 28 days will be approved quantities sufficient for the number of courses of chemotherapy. For approval of generic aprepitant, the patient must have a documented intolerance to brand Emend. Emend oral suspension: medication will be prescribed by an oncology practitioner AND patient requires prevention of nausea and vomiting associated with moderate to highly emetogenic cancer chemotherapy AND patient has a documented medical necessity for the specialty dosage form (e.g. swallowing disorder) Varubi: Medication will be prescribed by an oncology practitioner AND patient requires prevention of nausea and vomiting associated with moderate to highly emetogenic cancer chemotherapy AND the patient has had a documented side effect, allergy, or treatment failure with Emend®.
THC DERIVATIVES		
All products require PA	Dronabinol (compare to Marinol [®]) Marinol [®] (dronabinol)	Pharmacology: Marinol® is a schedule III cannabinoid agent containing the same active ingredient, tetrahydrocannabinol, as marijuana. While its exact mechanism of action is unknown, it is speculated to inhibit medullary activity

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
	Cesamet [®] (nabilone)	as well as suppress prostaglandin and endorphin synthesis. Cesamet® is a schedule II synthetic cannabinoid that acts by activating the endocannabinoid receptors, CB1 and CB2, which are involved in nausea/vomiting regulation. Both Marinol® and Cesamet® are FDA-approved for use in chemotherapy associated nausea and vomiting refractory to conventional antiemetics. In addition, Marinol® is indicated for patients with HIV/AIDS-related anorexia or wasting syndrome. Dronabinol/Marinol: patient has a diagnosis of chemotherapy-induced nausea/vomiting AND patient has had a documented side effect, allergy, or treatment failure to at least 2 antiemetic agents, of which, one must be a preferred 5HT3 receptor antagonist. If the request is for Marinol, the patient must additionally have a documented intolerance to generic dronabinol. OR patient has a diagnosis of HIV/AIDS associated anorexia. AND patient has had an inadequate response, adverse reaction, or contraindication to megestrol acetate. If the request is for Marinol, the patient must additionally have a documented intolerance to generic dronabinol. Cesamet: patient has a diagnosis of chemotherapy-induced nausea/vomiting AND patient has had a documented side effect, allergy, or treatment failure to at least 2 antiemetic agents, of which, one must be a preferred 5HT3 receptor antagonist.
ACE INHIBITORS	ANTI-HYPERTENS	SIVES
BENAZEPRIL (compare to Lotensin [®]) ENALAPRIL (compare to Vasotec [®]) tablet ENALAPRIL oral solution (age ≤ 12 years old) FOSINOPRIL LISINOPRIL (compare to Zestril®) QUINAPRIL (compare to Accupril [®]) RAMIPRIL (compare to Altace [®]) TRANDOLAPRIL	Accupril® (quinapril) Altace® (Ramipril) Captopril Enalapril oral solution (age > 12 years old) Epaned® (enalapril) oral solution Lotensin® (benazepril) Moexepril Perindopril Qbrelis® (Lisinopril) 1mg/ml solution Vasotec® (enalapril) Zestril® (lisinopril)	 Enalapril (Patients > 12 years old), Epaned Oral Solution: patient has a requirement for an oral liquid dosage form (i.e. swallowing disorder) AND for approval of Epaned, the patient must have a documented intolerance to the generic equivalent. Qbrelis Oral Solution: patient has a requirement for an oral liquid dosage form (i.e. swallowing disorder, inability to take oral medications) AND has a side effect, allergy, or treatment failure to Epaned oral solution. Other ACE Inhibitors: patient has had a documented side effect, allergy, or treatment failure to all available preferred generic ACEI. If a medication has an AB rated generic, there must have been a trial of the generic formulation.
ACE INHIBITOR W/ HYDROCHLOROTHIA	ZIDE	
$\label{eq:BENAZEPRIL/HYDROCHLOROTHIAZIDE} \\ (compare to Lotensin HCT^{\textcircled{R}})$	Accuretic [®] (quinapril/HCTZ) Lotensin HCT [®] (benazepril/HCTZ)	ACE Inhibitor/Hydrochlorothiazide combinations: patient has had a documented side effect, allergy, or treatment failure to all available preferred generic ACEI/Hydrochlorothiazide combination. If a medication has an AB

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
ENALAPRIL/HYDROCHLOROTHIAZIDE (compare to Vaseretic [®]) FOSINOPRIL/HYDROCHLOROTHIAZIDE LISINOPRIL/HYDROCHLOROTHIAZIDE (compare to Zestoretic [®]) QUINAPRIL/HYDROCHLOROTHIAZIDE (compare to Accuretic [®])	Vaseretic [®] (enalapril/HCTZ) Zestoretic [®] (lisinopril/HCTZ)	rated generic, there must have been a trial of the generic formulation.
ACE INHIBITOR W/CALCIUM CHANNEL BLOC	CKER	
AMLODIPINE/BENAZEPRIL (compare to Lotrel®)	Lotrel [®] amlodipine/(benazepril) Trandolapril/Verapamil ER	 Lotrel: The patient has had a documented side effect, allergy, or treatment failure to the generic formulation. Trandolapril/Verapamil ER: The patient has had a documented side effect, allergy, or treatment failure to amlodipine/benazepril AND the patient is unable to take as the individual separate agents.
ANGIOTENSIN RECEPTOR BLOCKERS (ARBs)		
CANDESARTAN IRBESARTAN (compare to Avapro [®]) LOSARTAN (compare to Cozaar [®]) OLMESARTAN (compare to Benicar [®]) TELMISARTAN (compare to Micardis®) VALSARTAN (compare to Diovan [®])	Avapro [®] (irbesartan) Benicar [®] (olmesartan) Cozaar [®] (losartan) Diovan [®] (valsartan) Edarbi [®] (azilsartan) Tablet QTY LIMIT:1 tablet/day Micardis® (telmisartan)	Avapro, Benicar, Cozaar, Diovan, Edarbi, and Micardis: Patient has had a documented side effect, allergy, or treatment failure with TWO preferred Angiotensin Receptor Blocker (ARB) or ARB combinations. AND If brand name product with generic available, the patient has had a documented intolerance with the generic product.
ANGIOTENSIN RECEPTOR BLOCKER/DIURET	TC COMBINATIONS	
IRBESARTAN/HYDROCHLOROTHIAZIDE (compare to Avalide®) LOSARTAN/HYDROCHLOROTHIAZIDE (compare to Hyzaar®) OLMESARTAN/HYDOCHLOROTHIAZIDE (compare to Benicar HCT®) TELMISARTAN/HYDROCHLOROTHIAZIDE (compare to Micardis HCT®) VALSARTAN/HYDROCHLOROTHIAZIDE (compare to Diovan HCT®)	Avalide [®] (irbesartan/hydrochlorothiazide) Benicar HCT [®] (olmesartan/hydrochlorothiazide) Candesartan/hydrochlorothiazide Diovan HCT® (valsartan/hydrochlorothiazide) Edarbyclor [®] (azilsartan/chlorthalidone) Tablet <i>QTY LIMIT:</i> 1 tablet/day Hyzaar [®] (losartan/hydrochlorothiazide) Micardis HCT [®] (telmisartan/hydrochlorothiazide)	Avalide, Benicar HCT, Candesartan/HCTZ, Diovan HCT, Edarbyclor, Hyzaar, Micardis HCT and Telmisartan/HCTZ: patient has had a documented side effect, allergy, or treatment failure with a preferred ARB/Hydrochlorothiazide combination AND If brand name product with generic available, the patient has had a documented intolerance with the generic product.

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
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OLMESARTAN/AMLODIPINE (compare to Azor®) VALSARTAN/AMLODIPINE (compare to Exforge®) QTY LIMIT: 1 tablet/day	Azor [®] (olmesartan/amlodipine) QTY LIMIT: 1 tablet/day Amlodipine/telmisartan QTY LIMIT: 1 tablet/day Exforge [®] (valsartan/amlodipine) QTY LIMIT: 1 tablet/day	Azor, Amlodipine/Telmisartan, Exforge, Olmesartan/amlodipine: The patient has had a documented side effect, allergy, or treatment failure to a preferred ARB/CCB combination product AND if brand name product with generic available, the patient has had a documented intolerance with the generic equivalent.
ANGIOTENSIN RECEPTOR BLOCKER/CALCI	UM CHANNEL BLOCKER/HCTZ COMBO	
VALSARTAN/AMLODIPINE/HCTZ (compare to Exforge HCT®) QTY LIMIT: 1 tablet/day	Exforge HCT® (amlodipine/valsartan/hydrochlorothiazide) QTY LIMIT: 1 tablet/day Olmesartan/amlodipine/hydrochlorothiazide (compare to Tribenzor®) QTY LIMIT: 1 tablet/day Tribenzor® (amlodipine/olmesartan/hydrochlorothiazide) QTY LIMIT: 1 tablet/day	Exforge HCT, Olmesartan/amlodipine/HCTZ, Tribenzor: patient has had a documented side effect, allergy, or treatment failure to Valsartan/amlodipine/HCTZ.
BETA BLOCKERS		
SINGLE AGENT ACEBUTOLOL ATENOLOL (compare to Tenormin®) BISOPROLOL FUMARATE BYSTOLIC® (nebivolol) CARVEDILOL (compare to Coreg®) LABETALOL METOPROLOL TARTRATE (compare to Lopressor®) METOPROLOL SUCCINATE XL (compare to Toprol XL®) NADOLOL NEBIVOLOL (compare to Bystolic®) PINDOLOL PROPRANOLOL PROPRANOLOL ER (compare to Inderal LA®) SOTALOL (compare to Betapace®, Betapace AF®) Preferred After Clinical Criteria Are Met HEMANGEOL® oral solution (propranolol)	Betapace (sotalol) Betapace AF (sotalol) Betaxolol Carvedilol CR (compare to Coreg)	Non-preferred drugs (except as noted below) patient has had a documented side effect, allergy, or treatment failure to at least three preferred drugs. (If a medication has an AB rated generic, one trial must be the generic formulation.) Carvedilol CR, Coreg CR: Indication: Heart Failure: patient has been started and stabilized on the medication. (Note: Samples are not considered adequate justification for stabilization.) OR patient has had a documented side effect, allergy, or treatment failure to metoprolol SR or bisoprolol. AND patient has been unabl to be compliant with or tolerate twice daily dosing of carvedilol IR. Indication: Hypertension: patient has been started and stabilized on the medication. (Note: Samples are not considered adequate justification for stabilization.) OR patient has had a documented side effect, allergy, or treatment failure to 3 (three) preferred anti-hypertensive beta-blockers. Hemangeol: indication for use is the treatment of proliferating infantile hemangioma Kapspargo: patient is unable to take a solid oral dosage form and has a treatment failure with an immediate release oral solution or crushed tablets.

PREFERRED AGENTS	NON-PREFERRED AGENTS	DA CDITEDIA
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
BETA-BLOCKER/DIURETIC COMBINATION ATENOLOL/CHLORTHALIDONE (compare to Tenoretic®) BISOPROLOL/HYDROCHLOROTHIAZIDE (compare to Ziac®) METOPROLOL/HYDROCHLOROTHIAZIDE	Nadolol/bendroflumethiazide Tenoretic [®] (atenolol/chlorthalidone) Ziac [®] (bisoprolol/HCTZ)	
CALCIUM CHANNEL BLOCKERS		
SINGLE AGENT DIHYDROPYRIDINES AMLODIPINE (compare to Norvasc®) FELODIPINE ER NIFEDIPINE IR (compare to Procardia®) NIFEDIPINE SR osmotic (compare to Procardia® XL) NIFEDIPINE SR (compare to Adalat® CC)	Isradipine Katerzia [®] (amlodipine) oral suspension Levamlodipine Nicardipine Nimodipine Norliqva® (amlodipine) oral solution Nisoldipine ER (compare to Sular [®]) Norvasc [®] (amlodipine) Nymalize [®] (nimodipine) Oral Solution Procardia [®] (nifedipine IR)	 Criteria for approval (except as noted below:) patient has had a documented side effect, allergy, or treatment failure to at least three preferred drugs. (If a medication has an AB rated generic, one trial must be the generic formulation.) Katerzia: patient has a medical necessity for a specialty dosage form (i.e. dysphagia, swallowing disorder). Norliqva, Nymalize: patient has a medical necessity for a specialty dosage form (i.e. dysphagia, swallowing disorder) and the patient has a had a documented side effect, allergy, or treatment failure to Katerzia.
NON-DIHYDROPYRIDINES CARTIA® XT (diltiazem SR, compare to Cardizem® CD) DILT-XR® (diltiazem SR) DILTIAZEM (compare to Cardizem®) DILTIAZEM ER 24-hour capsules (compare to Tiazac®) DILTIAZEM SR 24-hour capsules (compare to Cardizem®CD) DILTIAZEM SR 24-hour tablets TAZTIA® XT (diltiazem ER, compare to Tiazac®) VERAPAMIL (compare to Calan®)	Procardia (nifedipine IR) Procardia XL (nifedipine SR osmotic) Sular (nisoldipine) Calan SR (verapamil CR) Cardizem (diltiazem) Cardizem CD (diltiazem SR) Cardizem LA (diltiazem SR) Diltiazem ER 12-hour capsules Diltiazem ER/Matzin LA (compare to Cardizem LA) Tiazac (diltiazem ER) Verapamil SR 100 mg, 200 mg, 300mg (compare to Verelan PM®) Verelan (verapamil SR 120 mg, 180 mg, 240 mg and 360 mg)	

NON-PREFERRED AGENTS	
	PA CRITERIA
(i A required)	TACRITERIA
Verelan [®] PM (100 mg, 200 mg and 300 mg)	
Made 11 are Tellar	
ivietnyidopa Tablets	Methyldopa: The patient has a documented side effect, allergy, or
	contraindication to two preferred central alpha agonists.
Vecamyl [®] (mecamylamine) tablet	Vecamyl tabs: Patient has a diagnosis of moderately severe or severe hypertension AND patient has tried and failed, intolerant to, or contraindicated to at least THREE different antihypertension therapies of different mechanism of actions.
SINGLE AGENT Aliskiren (compare to Tekturna®) QTY LIMIT: 1 tablet/day Tekturna® (aliskiren) QTY LIMIT: 1 tablet/day COMBINATIONS Tekturna HCT® (aliskiren/hydrochlorothiazide) OTY LIMIT: 1 tablet/day	 Aliskiren, Tekturna: patient is NOT a diabetic who will continue on therapy with an ACEI or ARB AND patient has a diagnosis of hypertension. AND patient has had a documented side effect, allergy, or treatment failure with an angiotensin Receptor Blocker (ARB). Tekturna HCT: the patient must meet criteria as listed above for Tekturna and is unable to use the individual separate agents.
	Methyldopa Tablets Vecamyl [®] (mecamylamine) tablet SINGLE AGENT Aliskiren (compare to Tekturna®) QTY LIMIT: 1 tablet/day Tekturna® (aliskiren) QTY LIMIT: 1 tablet/day COMBINATIONS

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA		
ANTI-INFECTIVES ANTIBIOTICS				
AMINOGLYCOSIDES				
NEOMYCIN SULFATE	Arikayce® (amikacin inhalation suspension) QTY LIMIT: 28 vials (235.2 mL)/28 days	Arikayce: Patient is ≥ 18 years of age AND indication for use is treatment of <i>Mycobacterium avium complex</i> (MAC) lung disease AND patient has not achieved negative sputum cultures after a minimum of 6 consecutive months of a multidrug background regimen therapy (e.g. macrolide, rifampin, & ethambutol) within the past 12 months. Note: Initial approval will be granted for 6 months. For re-approval, the patient must have documentation of clinical improvement AND 3 consecutive monthly negative sputum cultures.		
CEPHALOSPORINS 1ST GENERATION				
CAPSULES/TABLETS CEFADROXIL capsules CEPHALEXIN capsules (compare to Keflex®) SUSPENSION CEFADROXIL suspension CEPHALEXIN suspension IV drugs are not managed at this time	Cefadroxil tablets Cephalexin tablets	Cephadroxil tabs: patient has had a documented intolerance to cefadroxil generic capsules. Cephalexin Tabs: patient has had a documented intolerance to cephalexin generic capsules.		
CEPHALOSPORINS 2 ND GENERATION				
CAPSULES/TABLETS CEFACLOR capsule CEFPROZIL tablet CEFUROXIME tablet	Cefaclor [®] ER tablet	Cefaclor ER Tabs: patient has had a documented intolerance to cefaclor capsules. Cefaclor Suspension: patient has a documented side effect, allergy, or treatment failure to Cefprozil suspension.		
SUSPENSION CEFPROZIL suspension	Cefaclor suspension			
IV drugs are not managed at this time				
CEPHALOSPORINS 3 RD GENERATION				
CAPSULES/TABLETS CEFDINIR CAPSULE CEFPODOXIME TABLET	Suprax [®] (cefixime) chewable tablets	 Suprax, chewable tablet: patient is completing a course of therapy which was initiated in the hospital. OR patient has had a documented side effect or treatment failure to cefdinir or cefpodoxime. Cefpodoxime Proxetil Susp, Cefixime Susp, Suprax Susp: patient is 		

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
SUSPENSION	Cefixime suspension	completing a course of therapy which was initiated in the hospital. OR patient
CEFDINIR suspension	Cefpodoxime proxetil suspension Suprax [®] (cefixime) suspension	has had a documented side effect or treatment failure to cefdinir suspension.
	Suprax (certaine) suspension	
IV drugs are not managed at this time		
CLINDAMYCIN DERIVATIVES		
CLINDAMYCIN (compare to Cleocin®) capsules	Cleocin (clindamycin) Capsules	Cleocin: the patient has a documented intolerance to the generic equivalent.
CLINDAMYCIN (compare to Cleocin®) oral solution	Cleocin® Ped (clindamycin) oral solution	
MACROLIDES		
AZITHROMYCIN tabs, liquid (≤5-day supply) (compare to Zithromax®)	Azithromycin tablets and liquid (if > 5-day supply) (compare to Zithromax [®])	Non-preferred agents (except as below): patient has a documented side-effect,
Maximum 10 days therapy/30 days	Azithromycin packet (compare to Zithromax [®])	allergy, or treatment failure to at least two of the preferred medications. (If a
	QTY LIMIT: 2 grams/fill	product has an AB rated generic, one trial must be the generic.) OR patient is
	Zithromax [®] (azithromycin) tablets and liquid	completing a course of therapy with the requested medication that was initiated in the hospital.
	QTY LIMIT: 5 days supply/RX, maximum 10 days, therapy/30 days Zithromax (azithromycin) packet	Azithromycin/Zithromax packets: A clinically valid reason why the dose
		cannot be obtained using generic azithromycin tablets or suspension AND If
	QTY LIMIT: 2 grams/fill	the request is for brand Zithromax, the patient has a documented intolerance
		to the generic product.
		Azithromycin > 5-day supply (criteria for approval based on indication):
		Lyme Disease: patient has had a documented side effect, allergy, or treatment failure to at least two of the following: doxycycline, amoxicillin, or a 2nd
		generation cephalosporin. For early Lyme disease, without neurologic or
CLARITHROMYCIN tablets	Clarithromycin SR	rheumatologic (arthritis) complications, the length of authorization is up to 10
	Clarithromycin suspension	days. For neurologic or rheumatologic Lyme disease, the length of
	E.E.S.® (erythromycin ethylsuccinate) ERY-TAB® (erythromycin base, delayed release)	authorization is up to 28 days
	ERYTHROMYCIN BASE	Cystic Fibrosis: length of authorization up to 12 months
	Erythromycin base, delayed release (compare to Ery-	HIV/immunocompromised status: azithromycin is being used for MAC or Toxoplasmosis treatment or prevention. (length of authorization up to 6
	tab [®]) ERYTHROMYCIN ETHYLSUCCINATE (compare to	months)
	E.E.S.®)	Bacterial Sinusitis: patient has had a documented side effect, allergy, or treatment
	Eryped [®] (erythromycin ethylsuccinate)	failure to penicillin, amoxicillin, or sulfamethoxazole/trimethoprim (Bactrim).
	Erythrocin (erythromycin stearate)	(length of authorization up to 10 days)
	Dificid [®] (fidaxomicin) tablet	Severe Bronchiectasis or COPD with frequent exacerbations: length of
	QTY LIMIT: 2 tablets per day, 10-day supply per 30	authorization up to 1 year (There is no safety or efficacy data for long-term therapy beyond one year)
	days	Babesiosis: blood smear or PCR is positive (results must be submitted; positive
		serology is not sufficient) AND patient is symptomatic (length of authorization

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
(NOTA required unless otherwise noted)	(1 A required)	TACKITEKIA
IV drugs are not managed at this time		up to 10 days) Difficid: patient's diagnosis or indication is Clostridium difficile associated diarrhea (CDAD) AND patient has had a side-effect, allergy, treatment failure or contraindication to oral vancomycin.
NITROFURANTOIN DERIVATIVES		
NITROFURANTOIN MACROCRYSTALLINE capsules (compare to Macrodantin®) NITROFURANTOIN MONOHYDRATE MACROCYSTALLINE capsules (compare to Macrobid®) NITROFURANTOIN SUSPENSION (age ≤ 12 yrs)	Macrobid® (nitrofurantoin monohydrate macrocrystalline) capsules Macrodantin® (nitrofurantoin macrocrystalline) capsules	 Macrobid, Macrodantin: the patient has a documented intolerance to the generic equivalent. Nitrofurantoin susp (age > 12 yrs): patient must have medical necessity for a liquid formulation (i.e. swallowing disorder)
OXAZOLIDINONES		
IV form of this medication not managed at this time	Linezolid (compare to Zyvox®) <i>QTY LIMIT</i> :56 tablets per 28 days Linezolid (compare to Zyvox®) suspension <i>QTY LIMIT</i> :60 ml/day, maximum 28 days supply Sivextro® (tedizolid) <i>QTY LIMIT</i> :1 tab/day Zyvox® (linezolid) <i>QTY LIMIT</i> :56 tablets per 28 days Zyvox® (linezolid) suspension <i>QTY LIMIT</i> : 60 ml/day, maximum 28 days supply	Criteria for Approval: patient has been started on intravenous or oral linezolid or tedizolid in the hospital and will be finishing the course of therapy in an outpatient setting OR patient has a documented blood, tissue, sputum, or urine culture that is positive for Vancomycin-Resistant Enterococcus (VRE) species. OR patient has a documented blood, sputum, tissue, or urine culture that is positive for Methicillin-Resistant Staphylococcus species AND patient has had a documented treatment failure with trimethoprim/sulfamethoxazole, clindamycin, doxycycline, or minocycline OR there is a clinically valid reason that the patient cannot be treated with one of those agents AND for approval of Zyvox or Sivextro the patient has an intolerance to generic linezolid.
PLEUROMUTILINS		
All products require PA IV form of this medication not managed at this time PENICILLINS (ORAL)	Xenleta® (lefamulin acetate) QTY LIMIT: 2 tabs/day	Xenleta: patient is completing a course of therapy which was initiated in the hospital OR patient is ≥ 18 years of age AND has a confirmed diagnosis of community-acquired bacterial pneumonia (CABP) AND culture and sensitivity (C&S) report shows isolated pathogen is a susceptible to lefamulin (If obtaining a C&S report is not feasible, provider must submit documentation.) AND patient has a documented treatment failure, intolerance, or contraindication to 2 preferred antibiotics AND patient has no known risk factors for increased QT prolongation (e.g. cardiac arrhythmias, symptomatic bradycardia, hypokalemia, or congenital prolongation of the QT interval) AND medication is not being used in combination with other drugs known to prolong the QT interval (e.g. antipsychotics, erythromycin, tricyclic antidepressants). If use of Xenleta® cannot be avoided in these patients, baseline EKG and plan for ongoing monitoring must be documented.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
SINGLE ENTITY AGENTS NATURAL PENICILLINS PENICILLIN V POTASSIUM tablets, oral solution PENICILLINASE-RESISTANT PENICILLINS DICLOXACILLIN Capsules AMINOPENICILLINS AMOXICILLIN capsules, tablets, chewable tablets, suspension AMPICILLIN capsules, suspension COMBINATION PRODUCTS AMOXICILLIN/CLAVULANATE tablets, chewable tablets, suspension	Amoxicillin/clavulanate ER tablets	Amoxicillin/Clavulanate ER: prescriber must provide a clinically valid reason for the use of the requested medication.
QUINOLONES		
CIPROFLOXACIN (compare to Cipro®) tabs CIPRO® (ciprofloxacin) oral suspension LEVOFLOXACIN (compare to Levaquin®) tabs, solution MOXIFLOXACIN tabs IV drugs are not managed at this time	Baxdela [™] (delafloxacin) Cipro [®] (ciprofloxacin) tabs Levaquin [®] (levofloxacin) tabs, solution Ofloxacin	 Cipro, Levaquin: the patient has had a documented intolerance to the generic equivalent. Baxdela: patient is completing a course of therapy with the requested medication that was initiated in the hospital OR patient is ≥ 18 years of age AND has a confirmed diagnosis of acute bacterial skin and skin structure infection (ABSSSI) AND current culture and sensitivity (C&S) report shows isolated pathogen is a grampositive or gram-negative organism susceptible to delafloxacin (If obtaining a C&S report is not feasible, provider must submit documentation.) AND member has a documented treatment failure, intolerance or contraindication to 2 preferred antibiotics, one of which must be a fluoroquinolone AND duration of therapy does not exceed 14 days. Ofloxacin: patient has had a documented side effect, allergy, or treatment failure with two preferred fluoroquinolones
RIFAMYCINS		
All products require PA	Aemcolo® (rifamycin) delayed release tablets **QTY LIMIT: 12 tablets, max of 3 days** Xifaxan ® (rifaximin) 200 mg tablets	Aemcolo: patient has a diagnosis of traveler's diarrhea caused by noninvasive strains of Escherichia coli AND Patient has had a documented side effect, allergy, treatment failure or contraindication with a fluoroquinolone or azithromycin.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
	QTY LIMIT: depends on indication	Xifaxan: Criterial for Approval Based on Indication:
	Xifaxan [®] (rifaximin) 550 mg tablets	Hepatic Encephalopathy (Xifaxan 550 mg Tablets Only): patient has a
	QTY LIMIT: depends on indication	diagnosis of hepatic encephalopathy. AND Patient has had a documented side effect, allergy, treatment failure or contraindication to lactulose. AND
		Quantity limit is 2 tablets/day (550 mg tablets only).
		Traveler's Diarrhea (Xifaxan 200 mg Tablets Only): patient has a diagnosis
		of traveler's diarrhea caused by noninvasive strains of Escherichia coli. AND
		Patient has had a documented side effect, allergy, treatment failure or contraindication with a fluoroquinolone or azithromycin. AND Quantity limit
		is 9 tablets/RX (200 mg tablets only).
		Small Intestinal Bacterial Overgrowth (Xifaxan 550 mg or 200 mg Tablets:
		patient has a diagnosis of SIBO AND Quantity limit is 1,200 mg to
		1,650mg/day for 14 days; maximum of 3 courses will be approved.
		Irritable Bowel Syndrome (Xifaxan 550 mg or 200 mg Tablets): patient has a diagnosis of irritable bowel syndrome without constipation or with symptoms
		of bloating. Quantity limit is 1,200 mg to 1,650 mg/day for 14 days;
		maximum of 3 courses will be approved.
		Inflammatory Bowel Disease: Crohn's Disease (Xifaxan 550 mg or 200 mg
		Tablets): patient has a diagnosis of Crohn's Disease. AND Patient has had a
		documented side effect, allergy, treatment failure or contraindication to two of the following: 6-mercaptopurine, azathioprine, corticosteroids, or
		methotrexate. AND Quantity limit is 600 mg to 1,600 mg/day.
		Clostridium difficile Diarrhea (Xifaxan 200 mg Tablets): patient has a
		diagnosis of C. difficile diarrhea. AND Patient has had a documented side
		effect, allergy, treatment failure or contraindication to vancomycin AND
		Quantity limit is 1200mg/day.
TETRACYCLINES		
DOVVOVCI INTE MONOTRYDD ATE 50 MC	Demeclocycline 150mg, 300mg tabs	Non-preferred doxycycline/minocycline products (except as listed below):
DOXYCYCLINE MONOHYDRATE 50 MG, 100 MG capsules, tablets	Doryx (doxycycline hyclate) delayed release tabs	patient has had a documented side effect, allergy, or treatment failure with a preferred doxycycline/minocycline. If a product has an AB rated generic, the
DOXYCYCLINE HYCLATE 20MG tablets	Doxycycline hyclate delayed release tabs	trial must be the generic formulation.
DOXYCYCLINE HYCLATE 100 MG	Doxycycline monohydrate 40mg cap Doxycycline 75mg, 150mg caps, tabs	Nuzyra: patient has been started on intravenous or oral omadacycline in the hospital
capsules, tablets	Minolira® ER (minocycline extended release) tablet	and will be finishing the course of therapy in an outpatient setting OR the patient
DOCYCYCLINE HYCLATE 50MG capsules	QTY LIMIT: 1 tablet/day	has a diagnosis of community-acquired bacterial pneumonia (CABP) or acute bacterial skin and skin structure infections (ABSSSI) AND the patient has had a
DOXYCYCLINE MONOHYDRATE suspension 25 MG/5ML	Minocycline 50 mg, 75 mg, 100 mg tabs Nuzyra® (omadacycline) tabs	documented treatment failure with two preferred antibiotics (from any class) OR
MINOCYCLINE 50 MG, 100 MG capsules	QTY LIMIT: Max 14-day supply	the provider submits clinical rationale as to why the preferred agents would not be appropriate for the patient.
,	Solodyn®(minocycline) tabs ER	Oracea: patient has a diagnosis of Rosacea AND patient has had a documented side
	Tetracycline 250 mg, 500 mg cap Vibramycin® (doxycycline hyclate) cap, suspension	effect, allergy, or treatment failure with both a preferred doxycycline and
	. Island only of the nyolate, suspension	

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
	Vibramycin® (doxycycline calcium) syrup Ximino® (minocycline) caps ER All other brands	 minocycline. Minolira ER/Solodyn/Ximino: patient is ≥ 12 years of age AND indication is to treat non-nodular inflammatory lesions of acne vulgaris AND patient has had a documented side effect, allergy, or treatment failure with a preferred minocycline. Note: no effect has been demonstrated on non-inflammatory acne lesions. Vibramycin Suspension, Syrup: patient has a medical necessity for a liquid dosage form AND a documented failure of preferred doxycycline suspension. Tetracycline: patient has had a documented side effect, allergy, or treatment failure with at least two preferred products OR the indication for use is the treatment of H. Pylori infection.
VANCOMYCIN		
All products require PA IV vancomycin products are not managed at this time	Firvanq TM (vancomycin HCl) powder for oral solution <i>QTY LIMIT:</i> 1 bottle (150ml) per course of therapy. If more than 150ml is required, use of 300ml bottle is required. Vancocin [®] Vancomycin (compare to Vancocin [®]) capsules, oral solution	 Firvanq, Vancomycin oral solution: The patient has a diagnosis or indication of Clostridium difficile associated diarrhea (CDAD) or staphylococcus enterocolitis AND for approval of Vancomycin oral solution, the patient has a documented intolerance to Firvanq. Vancocin, Vancomycin capsules: The patient has a diagnosis or indication of Clostridium difficile associated diarrhea (CDAD) or staphylococcus enterocolitis AND for approval of Vancocin, the patient has a documented intolerance to generic vancomycin capsules.
	ANTI-INFECTIVES ANTIFU	INGAL
ALLYLAMINES		
TERBINAFINE tabs (compare to Lamisil®) QTY LIMIT: 30 tablets/month (therapy limit of 90 days) GRISEOFULVIN MICROSIZE Suspension	Griseofulvin Microsize Tablets Griseofulvin Ultramicrosize Tablets	Griseofulvin Microsize Tabs/Griseofulvin Ultramicrosize: patient has had a documented side effect, allergy, or treatment failure with terbinafine tablets and a preferred formulation of griseofulvin.
AZOLES		
FLUCONAZOLE (compare to Diflucan®) tabs, suspension CLOTRIMAZOLE Troche (compare to Mycelex®) IV drugs are not managed at this time.	Cresemba [®] (isavuconazonium) caps Diflucan [®] (fluconazole) tabs, suspension Itraconazole (compare to Sporanox [®]) caps, solution Ketoconazole tabs Noxafil [®] (posaconazole) oral suspension	Cresemba: patient is completing a course of therapy that was initiated in the hospital OR patient has a diagnosis of mucormycosis OR patient has a diagnosis of invasive aspergillosis and has had a documented side effect, allergy, contraindication, or treatment failure with voriconazole. Ketoconazole/Itraconazole 100mg cap/Itraconzaole Solution/Sporanox patient has a documented side-effect, allergy, or treatment failure to at least

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
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	Noxafii [®] (posaconazole) DR Tablets	ONE of the preferred medications OR patient is completing a course of therapy that was initiated in the hospital. For approval of Sporanox® capsules, the patient must have a documented intolerance to generic itraconazole. For approval of Itraconazole solution, the patient must have a medical necessity for a liquid dosage form. Limitations: Coverage of Onychomycosis agents will NOT be approved solely for cosmetic purposes. Tokura: patient has a diagnosis of aspergillosis intolerant of or refractory to Amphotericin B therapy AND patient has a documented intolerance to both generic itraconazole and voriconazole OR patient has a diagnosis of blastomycosis or histoplasmosis AND the patient has a documented intolerance to itraconazole capsules and solution. Voriconazole/Vfend: Patient has a diagnosis of invasive aspergillosis. OR patient is completing a course of therapy with the requested medication that was initiated in the hospital. OR patient has a documented side-effect, allergy, or treatment failure to ONE of the preferred medications AND itraconazole. AND For approval of Vfend®, the patient must have a documented intolerance to generic voriconazole. AND For approval of voriconazole suspension, the patient must have a medical necessity for a liquid dosage form. Noxafil, Posaconazole: patient has a diagnosis of HIV/immunocompromised status (neutropenia secondary to chemotherapy, hematopoietic stem cell transplant recipients) AND medication is being used for the prevention of invasive Aspergillosis/ Candida infections. OR patient is completing a course of therapy with the requested medication that was initiated in the hospital. OR For Oral Suspension ONLY the patient has a documented side-effect, allergy, or treatment failure to one of the preferred medications and itraconazole AND the patient is being treated for oropharyngeal candidiasis. Diflucan (brand): For approval of Diflucan brand name product, the patient must have a documented intolerance to generic fluconazole. Oravig: The indication for use is t
TRITERPENOIDS		
All products require PA	Brexafemme® (ibrexafungerp) tablets	Brexafemme: The patient is not pregnant and has been counseled to use effective contraception during treatment and for 4 days after the last dose (if applicable) AND the patient has recurrent yeast infections despite a treatment course of 7-14

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
		days with a preferred vaginal azole AND a longer course of oral fluconazole (e one dose every 3 days for a total of 3 doses)
	ANTI-INFECTIVES AN	TIMALARIALS
ATOVAQUONE/PROGUANIL (compare to Malarone®) CHLOROQUINE COARTEM® (artemether/lumefantrine) DARAPRIM® (pyrimethamine) HYDROXYCHLOROQUINE SULFATE MEFLOQUINE PRIMAQUINE QUINIDINE SULFATE Preferred After Clinical Criteria Are Met KRINTAFEL® (tafenoquine succinate)	Malarone® (atovaquone/proguanil) Pyrimethamine (compare to Daraprim®) Quinine Sulfate (compare to Qualquin®) Qualaquin® (quinine sulfate)	 Krintafel: the patient is ≥ 16 years of age AND is receiving concurrent antimalar therapy Malarone: patient has a documented intolerance to the generic equivalent Pyrimethamine: patient has a documented intolerance to brand Daraprim Quinine sulfate, Qualaquin: diagnosis or indication is for the treatment of malaria. (Use for leg cramps not permitted.) AND If the request is for bran Qualaquin, the patient has a documented intolerance to the generic equival
	ANTI-PARAS	ITICS
ALBENDAZOLE (compare to Albenza®) BILTRICIDE® (praziquantel) IVERMECTIN (compare to Stromectol®)	Albenza® (albendazole) Benznidazole Emverm® (mebendazole) Lampit (nifurtimox) Stromectol® (ivermectin)	 Benznidazole, Lampit: patient must be between 2-12 years of age (Benznidazole) or ≤ 18 years (Lampit) AND patient has a diagnosis of Cha Disease (American trypanosomiasis) AND length of therapy does not exce 60 days. Emverm: patient has a documented side effect, allergy, treatment failure, or contraindication to albendazole OR indication for use is hookworm infection (e.g. ancyclostomiasis, necatoriasis, uninariasis). Albenza, Stromectol: patient has a documented intolerance to the generic product.
	ANTI-INFECTIVES A	NTI-VIRALS
HERPES SIMPLEX VIRUS MEDICATIONS (ORAL)	
ACYCLOVIR (compare to Zovirax®) tablets, capsules ACYCLOVIR suspension (age ≤ 12 yrs) VALACYCLOVIR (compare to Valtrex®)	Famciclovir (compare to Famvir [®]) Sitavig [®] (acyclovir) Buccal Tablet <i>QTY LIMIT:</i> 2 tablets/30 days	 Acyclovir suspension (age > 12 yrs), Zovirax suspension: patient has a med necessity for a non-solid oral dosage form AND for approval of brand Zovirax, the patient has a documented intolerance to generic acyclovir suspension. Famciclovir: patient has a documented side effect, allergy, or treatment failured.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
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	Zovirax [®] (acyclovir) tablets, capsules, suspension	 Sitavig: patient has a diagnosis of recurrent herpes labialis (cold sores), having at least 4 episodes in the previous year AND patient has a documented side effect or treatment failure with acyclovir AND valacyclovir. Valtrex, Zovirax (tabs, caps): patient has a documented intolerance to the generic equivalent.
INFLUENZA MEDICATIONS		
OSELTAMIVIR (compare to Tamiflu®) QTY LIMIT: 45 and 75 mg caps =10 caps/30 days, 30 mg caps = 20 caps/30 days, 6 mg/ml suspension = 180ml/30 days RELENZA® (zanamivir) QTY LIMIT: 20 blisters/30 days	Tamiflu® (oseltamivir) QTY LIMIT: 45 and 75 mg caps = 10 caps/30 days, 30 mg caps = 20 capsule /30 days, 6 mg/ml suspension = 180 ml/30 days Xofluza TM (baloxavir marboxil)	 Tamiflu: Patient has a documented intolerance to generic Oseltamivir Xofluza: Patient is ≥ 12 years of age AND there is a clinical, patient-specific reason the patient cannot use a preferred agent. Note: A maximum of one single dose per 30 days will be approved based on the patient's body weight: 40mg (2 x 20mg tablets) for patients weighing between 40kg and 80kg or 80mg for patients weighing at least 80kg. Limitations: Amantadine and rimantadine are not CDC recommended for use in influenza treatment or chemoprophylaxis at this time and are not covered for this indication. For information regarding amantadine see "Parkinson's Medications".
CYTOMEGALOVIRUS (CMV) INFECTION MEI	DICATIONS	
VALGNCICLOVIR (compare to Valctye®) tablet	Livtencity TM (maribavir) tablets Prevymis® (letermovir) Valcyte® tablets, solution Valganciclovir (compare to Valcyte®) solution	 Livtencity: Indication is for the treatment of CMV infection in a recipient of a hematopoietic stem cell or solid organ transplant AND infection is refractory to ganciclovir, valganciclovir, cidofovir, or foscarnet (as defined by >1 log₁₀ increase in CMV DNA levels in blood or serum after at least 14 days of therapy) AND medication will not be administered with ganciclovir or valganciclovir. For re-approval beyond 12 weeks, documentation must be submitted detailing continued medical necessity. Prevymis: Indication is for the prophylaxis of cytomegalovirus (CMV) infection in adult CMV-seropositive recipients [R+] of an allogenic hematopoietic stem cell transplant AND therapy is initiated between day 0 and day 28 post-transplantation AND therapy will continue through day 100 post-transplantation AND for approval of injection, the patient must be unable to take oral medications. Valcyte: the patient has a documented intolerance to generic valganciclovir AND for approval of solution, the patient has a medical necessity for a non-solid oral dosage form. Valganciclovir solution: the patient has a medical necessity for a non-solid oral dosage form.
INFLUENZA VACCINES		

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
SEASONAL INFLUENZA VACCINE INJECTION INACTIVATED INFLUENZA VACCINE, QUADRIVALENT (IIV4), STANDARD DOSE (EGG BASED) AFLURIA® QUADRIVALENT Injection FLUARIX® QUADRIVALENT Injection FLULAVAL® QUADRIVALENT Injection FLUZONE® QUADRIVALENT Injection	ADJUVANTED INACTIVATED INFLUENZA VACCINE, QUADRIVALENT (IIV4), STANDARD DOSE (EGG BASED) Fluad TM Injection INACTIVATED INFLUENZA VACCINE, QUADRIVALENT (IIV4), HIGH DOSE (EGG BASED) Fluzone High-Dose® Injection RECOMBINANT INFLUENZA VACCINE, QUADRIVALENT (RIV4) (EGG FREE) Flublok® Injection INACTIVATED INFLUENZA VACCINE, QUADRIVALENT (ccIIV4), STANDARD DOSE (CELL CULTURE BASED) (NOT EGG FREE) Flucelvax Quadrivalent® Injection LIVE ATTENUATED INFLUENZA VACCINE, QUADRIVALENT (LAIV4) (EGG BASED) Flumist® Quadrivalent Intranasal	 Flucelvax Quadrivalent: Patient must have a documented severe reaction to egg based influenza vaccine OR Prescriber provides clinical rationale why one of the preferred influenza vaccines cannot be used. Flublok: Patient is ≥ 65 years old OR Patient must have a documented severe reaction to egg based influenza vaccine AND the patient is unable to use Flucelvax. Flumist: Flumist is being requested for influenza prophylaxis during flu season AND The patient is between the ages of 19 and 49 years old, AND Prescriber provides documentation of a contraindication to an intramuscular injection (e.g., currently on warfarin; history of thrombocytopenia) or other compelling information to support the use of this dosage form. Fluzone High Dose, Fluad: Patient is ≥ 65 years old OR Prescriber provides clinical rationale why one of the preferred influenza vaccines cannot be used. Note: the CDC and its Advisory Committee on Immunization Practices (ACIP) have not expressed a preference for any flu vaccine formulation for this age group.
Preferred After Age Limit Is Met GARDASIL SHINGRIX		 Gardasil: Covered for 19 years old to 45 years old (those under 19 should be referred to their pediatrician or PCP for state-supplied vaccine) Shingrix: Covered if ≥ 50 years of age Vaccines on the Advisory Committee on Immunization Practices (ACIP) list of recommended vaccines for children ≤ 18 years of age are supplied through the Vaccines for Children program administered by the Vermont Department of Health, and are not available through DVHA's pharmacy Programs. • Vaccines on the ACIP list of recommended vaccines for adults ≥ 19 years of age are available at many primary care provider offices and through the pharmacy programs. Vaccines are subject to the same limitations as the ACIP guideline recommendations. Providers who participate in the Blueprint for

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
		Health initiative must enroll in the Vaccines for Adults program administered by the Vermont Department of Health. The ACIP guidelines and information about enrollment in these programs can be found at http://healthvermont.gov/hc/imm/provider.aspx •Vaccines not on the recommended list may require Prior Authorization.

MIGRAINE THERAPY: PREVENTATIVE TREATMENTS

Calcitonin gene-related peptide (CGRP) Inhibitors: Initial approval is 6 months; renewals are 1 year

AIMOVIG® (erenumab-aooe)

QTY LIMIT: 1 injection (1mL) per 30

days

AJOVY® (fremanezumab-vfrm)

QTY LIMIT: 225 mg (1 injection) per 30 days or 675 mg (3 injections) every 90 days

675 mg (3 injections) every 90 days

EMGALITY® (galcanezumab-gnlm) 120 mg/mL QTY LIMIT: 240 mg (2 injections) for the first 30 days followed by 120 mg (1 injection) per 30 days

Emgality ® (galcanezumab-gnlm) 100 mg/mL QTY LIMIT: 300 mg (3 injections) per 30 days, maximum of 6 months per year approved

Nurtec® ODT (rimegepant)

OTY LIMIT: 16 tablets/30 days

OuliptaTM (atogepant)

QTY LIMIT: 30 tablets/30 days

Vyepti® (eptinezumab-jjmr)

Note: Please refer to "Botulinum Toxins" for Botox

Aimovig, Ajovy, Emgality 120mg/mL: The patient is 18 years of age or older AND patient has a diagnosis of episodic migraine (4-14 headache days per month with migraine lasting 4 hours or more) or chronic migraine (≥ 15 headache days per month, of which ≥ 8 are migraine days, for at least 3 months) AND patient has failed or has a contraindication to an adequate trial (≥ 60 days) of at least TWO medications for migraine prophylaxis from at least 2 different classes (tricyclic antidepressants, SNRI's, beta-blockers, or anticonvulsants). Initial approval will be granted for 6 months. For reapproval after 6 months, the patient must have documentation of a decrease in the number of headache days per month or decreased use of acute migraine medications such as triptans. Pharmacy claims will also be evaluated to assess compliance with the medication. Clinical justification must be provided if there is an increase in triptan use noted in the patient's profile.

Nurtec ODT, Quilipta, Vyepti: The patient is 18 years of age or older AND The patient must have a documented side effect, allergy, or treatment failure to two preferred CGRP Inhibitors. Initial approval will be granted for 6 months. For reapproval after 6 months, the patient must have documentation of a decrease in the number of headache days per month or decreased use of acute migraine medications such as triptans. Pharmacy claims will also be evaluated to assess compliance with the medcation. Clinical justification must be provided if there is an increase in triptan use noted in the patient's profile.

Emgality 100mg/mL:

- Patient is 18 years of age or older AND
- Patient has a diagnosis of episodic cluster headache as defined by the following:
 - Severe to very severe unilateral pain felt in the orbital, supraorbital, and/or temporal regions lasting 15-180 minutes (when untreated)
 - Pain is accompanied by a sense of restlessness or agitation OR at least one of the following signs or symptoms, ipsilateral to the headache:
 - Conjunctival injection and/or lacrimation
 - Evelid edema

NON-PREFERRED AGENTS	
(PA required)	PA CRITERIA
	 Miosis and/or ptosis Nasal congestion and/or rhinorrhea Forehead and facial sweating Patient has ≥ 2 active cluster periods lasting 7 days to 1 year, separated by remission for periods lasting ≥ 3 months AND Patient has not achieved satisfactory response to adequate doses of corticosteroids (≥ 30mg prednisone or ≥ 16mg dexamethasone daily) started promptly at the start of the cluster period (Failure is defined as the need to use acute/abortive medications (oxygen, triptans, ergotamines, lidocaine) at least once daily for at least 2 days/week after the first full week of steroid therapy) AND Patient has not achieved satisfactory response to adequate doses of verapamil (480mg/day, titrated up as needed to a max of 960mg/day) given for at least 3 weeks (Failure is defined as the need to use acute/abortive medications (oxygen, triptans, ergotamines, lidocaine) at least once daily for at least 2 days/week after 3 weeks of adequately dosed verapamil) Note: this requirement will be waived if the patient's 2 most recent active cluster periods were less than 3 weeks in duration.
MIGRAINE THERAPY: ACUTE T	REATMENTS
Ubrelvy® (ubrogepant) QTY LIMIT: 10 tablets/30 days	Nurtec ODT: Patient has a documented side effect, allergy, or treatment failure with 2 preferred triptans, unless contraindicated. Ubrelvy: Patient has a documented side effect, allergy, or treatment failure with 2 preferred triptans, unless contraindicated AND patient has a documented side effect, allergy, or treatment failure with Nurtec ODT.
Dihydroergotamine mesylate nasal spray (compare to Migranal®) **QTY LIMIT: 8 units/30 days** Trudhesa TM (dihydroergotamine mesylate) nasal spray **QTY LIMIT: 8 units/30 days*	Dihydroergotamine, Trudhesa: The patient has a documented intolerance to Migranal nasal spray.
Reyvow® (lasmiditan) QTY LIMIT: 8 tablets/30 days	Reyvow: Patient has a documented side effect, allergy, or treatment failure with 2 preferred triptans, unless contraindicated AND patient has a documented side effect, allergy, or treatment failure with Nurtec ODT AND counseling has been documented regarding the risks of driving impairment
	WIGRAINE THERAPY: ACUTE T Ubrelvy® (ubrogepant) QTY LIMIT: 10 tablets/30 days Dihydroergotamine mesylate nasal spray (compare to Migranal®) QTY LIMIT: 8 units/30 days Trudhesa TM (dihydroergotamine mesylate) nasal spray QTY LIMIT: 8 units/30 days Reyvow® (lasmiditan)

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
TRIPTANS		
SINGLE AGENT ORAL ELETRIPTAN (compare to Relpax®) OTY LIMIT: 12 tablets/30 days FROVATRIPTAN (compare to Frova®) 2.5 mg QTY LIMIT: 9 tablets/30 days NARATRIPTAN OTY LIMIT: 9 tablets/30 days SUMATRIPTAN (compare to Imitrex®) OTY LIMIT: 25 mg = 18 tablets/30 days, 50 and 100 mg = 9 tablets/30 days RIZATRIPTAN (compare to Maxalt®) OTY LIMIT: 12 tablets/30 days RIZATRIPTAN ODT (compare to Maxalt-MLT®) OTY LIMIT: 12 tablets/30 days ZOLMITRIPTAN (compare to Zomig®) tablets OTY LIMIT: 2.5 mg = 12 tablets/30 days, 5 mg = 6 tablets/30 days	Almotriptan 6.25 mg, 12.5 mg QTY LIMIT: 12 tablets/30 days Frova® (frovatriptan) 2.5 mg QTY LIMIT: 9 tablets/30 days Imitrex® (sumatriptan) QTY LIMIT: 25 mg = 18 tablets/30 days, 50 and 100 mg = 9 tablets/30 days Maxalt® (rizatriptan) 5 mg, 10 mg tablet QTY LIMIT: 12 tablets/30 days Maxalt-MLT® (rizatriptan ODT) QTY LIMIT: 12 tablets/30 days Relpax® (eletriptan) 20 mg, 40 mg QTY LIMIT: 12 tablets/30 days Zomig® (zolmitriptan) tablets QTY LIMIT: 2.5 mg = 12 tablets/30 days, 5 mg = 6 tablets/30 days Zomig® ZMT (zolmitriptan ODT) QTY LIMIT: 2.5 mg = 12 tablets/30 days, 5 mg = 6 tablets/30 days Zolmitriptan ODT (compare to Zomig® ZMT) QTY LIMIT: 2.5 mg = 12 tablets/30 days, 5 mg = 6 tablets/30 days	Non-preferred single agents: The patient has had a documented side effect, allerg or treatment failure with at least two preferred triptans. If a product has an AB rated generic, there must have also been a trial of the generic formulation. Sumatriptan/naproxen, Treximet: patient has had a documented side effect, allergy or treatment failure with 2 preferred Triptans, AND patient is unable to take the individual components separately. Zolmitriptan Nasal Spray, Zomig Nasal Spray, Imitrex Nasal Spray, Onzetra Xsail, Tosymra: patient has had a documented side effect, allergy, o treatment failure with Sumatriptan Nasal Spray. For Zolmitriptan Nasal Spray, the patient must also have a documented intolerance to the brand Zomig Nasal Spray. Imitrex Injection, Zembrace: patient has had a documented intolerance to generic sumatriptan injection. To exceed quantity limits: patient is taking a medication for migraine prophylaxis.
<u>NASAL SPRAY</u>	Imitrex [®] (sumatriptan) <i>QTY LIMIT</i> : 5 mg nasal spray = 12 units/30 days, 20 mg nasal spray = 6 units/30 days Tosymra® (sumatriptan) <i>QTY LIMIT</i> : 6 units/30 days	
SUMATRIPTAN (compare to Imitrex ^(B)) QTY LIMIT: 5 mg nasal spray = 12 units/30 days, 20 mg nasal spray = 6 units/30 days	Zomig [®] (zolmitriptan) <i>QTY LIMIT:</i> 2.5 and 5 mg nasal spray = 12 units/30 days	

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
	Zolmitriptan (compare to Zomig®) QTY LIMIT: 2.5 and 5 mg nasal spray = 12 units/30 days	
	Onzetra Xsail® (sumatriptan succinate) QTY LIMIT: 8 doses/30 days	
NASAL POWDER All products require PA INJECTABLE	Imitrex [®] (sumatriptan) <i>QTY LIMIT</i> : 4 and 6 mg injection = 8 injections (4ml)/30 days Zembrace [®] SymTouch (sumatriptan) 3 mg/5ml <i>QTY LIMIT</i> : 4 injections/ 30 days	
SUMATRIPTAN (compare to Imitrex [®]) <i>QTY LIMIT:</i> 4 and 6 mg injection = 8 injections (4ml)/30 days	Sumatriptan/Naproxen (compare to Treximet®) QTY LIMIT: 9 tablets/30 days	
COMBINATION PRODUCT ORAL All products require PA	Treximet [®] (sumatriptan/naproxen) QTY LIMIT: 9 tablets/30 days	

ANTI-PSYCHOTIC ATYPICAL & COMBINATIONS (CHILDREN < 18 YEARS OLD)

Preferred After Clinical Criteria Are Met Target symptoms or Diagnosis that will be accepted for approval: Target Abilify® (aripiprazole) TABLETS/CAPSULES Symptoms - Grandiosity/euphoria/mania; Obsessions/compulsions; Psychotic ARIPIPRAZOLE (compare to Abilify®) FDA maximum recommended dose = 30 mg/day symptoms; Tics (motor or vocal). Diagnosis- Autism with Aggression and/or FDA maximum recommended dose = 30 mg/day Asenapine (compare to Saphris®) irritability; Disruptive Mood Dysregulation Disorder; Bipolar Disorder; LURASIDONE (compare to Latuda®) OTY LIMIT: 2 tabs/day FDA maximum recommended dose = FDA maximum recommended dose = 80 mg/day 20 mg/day OLANZAPINE (compare to Zyprexa[®]) Clozapine (compare to Clozaril®) FDA maximum recommended dose = 20 mg/day FDA maximum recommended dose = 900 mg/day RISPERIDONE (compare to Risperdal[®]) Clozaril[®] (clozapine) FDA maximum recommended dose = 16 mg/day FDA maximum recommended dose = 900 mg/day PALIPERIDONE (compare to Invega®) Geodon® (ziprasidone) FDA maximum recommended dose = FDA maximum recommended dose = 160 mg/day

Invega® (paliperidone)

12 mg/day

QUETIAPINE (compare to Seroquel®)

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
· · · · · · · · · · · · · · · · · · ·		
FDA maximum recommended dose = 800 mg/day QUETIAPINE ER (compare to Seroquel® XR) FDA maximum recommended dose = 800 mg/day ZIPRASIDONE (compare to Geodon®) FDA maximum recommended dose = 160 mg/day	QTY LIMIT: 3 and 9 mg = 1 tab/day, 6 mg = 2 tabs/day FDA maximum recommended dose = 12 mg/day Latuda® (lurasidone) FDA maximum recommended dose = 80 mg/day Risperdal® (risperidone) FDA maximum recommended dose = 16 mg/day Seroquel® (quetiapine) FDA maximum recommended dose = 800 mg/day Saphris® (asenapine) QTY LIMIT: 2 tabs/day FDA maximum recommended dose = 20 mg/day Seroquel XR® (quetiapine XR) FDA maximum recommended dose = 800 mg/day Zyprexa® (olanzapine) FDA maximum recommended dose = 20 mg/day	Asenapine, Saphris: patient has had a documented side effect, allergy or treatment failure with at least two preferred products (typical or atypical antipsychotics) one of which is risperidone. Abilify, Clozaril, Geodon, Invega, Latuda, Risperdal, Seroquel, Seroquel XR, Zyprexa: patient has a documented intolerance to the generic equivalent. Clozapine: patient has had a documented side effect, allergy or treatment failure with at least three other antipsychotic medications (typical or atypical antipsychotics), two of which must be preferred agents. Aripiprazole Oral Solution: patient has had a documented side effect, allergy or treatment failure with risperidone oral solution OR prescriber feels that risperidone would not be an appropriate alternative for the patient because of pre-existing medical conditions such as obesity or diabetes. Versacloz Oral Solution: patient has had a documented side effect, allergy or treatment failure with at least three other antipsychotic medications (typical or atypical antipsychotics). AND patient is unable to use clozapine orally disintegrating tablets. Aripiprazole ODT, Olanzapine ODT, Risperidone ODT, Zyprexa Zydis: Medical necessity for a specialty dosage form has been provided AND if the request is for Zyprexa Zydis, the patient has a documented intolerance to the generic equivalent. Clozapine ODT: Medical necessity for a specialty dosage form has been provided AND patient has had a documented side effect, allergy or treatment failure with at least three other antipsychotic medications (typical or atypical antipsychotics)
Preferred After Clinical Criteria Are Met ORAL SOLUTIONS RISPERIDONE (compare to Risperdal®) oral solution FDA maximum recommended dose = 16 mg/day	Aripiprazole oral solution FDA maximum recommended dose = 25 mg/day Risperdal [®] (risperidone) oral solution FDA maximum recommended dose = 16 mg/day Versacloz [®] (clozapine) Oral Suspension <i>QTY LIMIT</i> : 18ml/day	

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
ORALLY DISINTEGRATING TABLETS All products require PA	Aripiprazole orally disintegrating tablets QTY LIMIT: 10 and 15 mg = 2 tabs/day FDA maximum recommended dose = 30 mg/day Clozapine orally disintegrating tablets FDA maximum recommended dose = 900 mg/day Olanzapine orally disintegrating tablets (compare to Zyprexa Zydis [®]) QTY LIMIT: 5 and 10 mg = 1.5 tabs/day FDA maximum recommended dose = 20 mg/day Risperidone ODT FDA maximum recommended dose = 16 mg/day Zyprexa Zydis [®] (olanzapine orally disintegrating tablets) QTY LIMIT: 5 and 10 mg = 1.5 tabs/day FDA maximum recommended dose = 20 mg/day	

ANTI-PSYCHOTIC ATYPICAL & COMBINATIONS (ADULTS ≥ 18 YEARS OLD)

TABLETS/CAPSULES

ARIPIPRAZOLE (compare to Abilify®)

FDA maximum recommended dose = 30 mg/day

CLOZAPINE (compare to Clozaril®)

FDA maximum recommended dose = 900 mg/day

LURASIDONE (compare to Latuda®)

FDA maximum recommended dose =

160 mg/day

OLANZAPINE (compare to Zyprexa[®])

FDA maximum recommended dose = 20 mg/day

PALIPERIDONE (compare to Invega®)

FDA maximum recommended dose = 12 mg/day

RISPERIDONE (compare to Risperdal®)

FDA maximum recommended dose = 16 mg/day

QUETIAPINE (compare to Seroquel[®])

FDA maximum recommended dose = 800 mg/day

QUETIAPINE ER (compare to Seroquel® XR)

Abilify® (aripiprazole)

FDA maximum recommended dose = 30 mg/day Abilify® Mycite (aripiprazole tablets with sensor)

QTY LIMIT: 1 tab/day

FDA maximum recommended dose=30mg/day

Asenapine sublingual tablet (compare to Saphris®)

FDA maximum recommended dose = 20 mg/day

Clozaril® (clozapine)

FDA maximum recommended dose = 900 mg/day Caplyta® (lumateperone)

OTY LIMIT: 1 capsule/day

FDA maximum recommended dose = 42 mg/day Fanant (iloperidone)

QTY LIMIT: 2 tablets/day

FDA maximum recommended dose = 24 mg/day

Geodon® (ziprasidone)

FDA maximum recommended dose = 160 mg/day Invega® (paliperidone)

QTY LIMIT: 3 and 9 mg = 1 tab/day, 6 mg = 2

Criteria for approval of ALL non-preferred drugs: patient has been started and stabilized on the requested medication (Note: samples are not considered adequate justification for stabilization.) OR patient meets additional criteria outlined below.

Caplyta:

Indication for use is schizophrenia/schizoaffective disorder: The patient has had a documented side effect, allergy or treatment failure with at least three preferred products (typical or atypical antipsychotics).

Indication for use is Bipolar Depression: the patient has had a documented side effect, allergy, or treatment failure with two preferred products (typical or atypical antipsychotics). If the prescriber feels that neither quetiapine or olanzapine/fluoxetine combination would be appropriate alternatives for the patient because of pre-existing conditions such as obesity or diabetes, the patient must have a documented side effect, allergy, or treatment failure with lurasidone.

Fanapt: The indication for use is the treatment of schizophrenia/schizoaffective disorder or bipolar disorder. AND The patient has had a documented side effect, allergy, or treatment failure with at least three preferred products (typical or atypical antipsychotics).

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
FDA maximum recommended dose = 800 mg/day ZIPRASIDONE (compare to Geodon®) FDA maximum recommended dose = 160 mg/day	tabs/day FDA maximum recommended dose = 12 mg Latuda [®] (lurasidone) FDA maximum recommended dose = 160 mg/day Nuplazid [™] (primavaserin) QTY LIMIT: 2 tablets/day FDA maximum recommended dose = 34 mg Rexulti [®] (brexpiprazole) FDA maximum recommended dose = 3 mg (adjunct of MDD) or 5 mg (schizophrenia) Risperdal [®] (risperidone) FDA maximum recommended dose = 16 mg/day Saphris [®] (asenapine) sublingual tablet FDA maximum recommended dose = 20 mg/day Seroquel [®] (quetiapine) FDA maximum recommended dose = 800 mg/day Seroquel XR [®] (quetiapine XR) FDA maximum recommended dose = 800 mg/day Vraylar [®] (cariprazine) QTY LIMIT: 1 capsule/day FDA maximum recommended dose = 6 mg/day Zyprexa [®] (olanzapine) FDA maximum recommended dose = 20 mg/day	Asenapine, Saphris: The indication for use is the treatment of schizophrenia/schizoaffective disorder or bipolar disorder AND The patient has had a documented side effect, allergy, or treatment failure with at least two preferred products (typical or atypical antipsychotics), one of which is risperidone. Note: Prior therapy with injectable Invega Sustenna® is not considered to be started and stabilized for oral Invega. Patients transferring to oral therapy from Invega Sustenna® should transition to oral risperidone (unless patient previously failed such treatment). Abilify, Clozaril, Geodon, Invega, Latuda, Risperdal, Seroquel, Seroquel X and Zyprexa: patient has a documented intolerance to the generic equivalen Abilify Mycite: The patient has not been able to be adherent to aripiprazole tablets resulting in significant clinical impact (documentation of measures aimed at improving compliance is required) AnD there is a clinically compelling reason why Abilify Maintena or Aristada cannot be used. Initial approval will be granted for 3 months. For renewal, documentation supportin use of the tracking software must be provided and pharmacy claims will be evaluated to assess compliance with therapy. Vraylar: Indication for use is schizophrenia/schizoaffective disorder: the patient has had a documented side effect, allergy or treatment failure with three preferred product (typical or atypical antipsychotics) OR Indication for use is Bipolar I depression: the patient has had a documented side effect, allergy, or treatment failure with two preferred products (typical or atypical antipsychotics). If the prescriber feels that neither quetiapine or olanzapine/fluoxetine combination would be appropriate alternatives for the patient because of pre-existing conditions such as obesity or diabetes, the patient must have a documented side effect, allergy, or treatment failure with two preferred atypical antipsychotics in the patient has had a documented inadequate response to at least 3 different antidepressants from two d
ORAL SOLUTIONS	Aripiprazole oral solution FDA maximum recommended dose =	acting opioids. Nuplazid: The diagnosis or indication is the treatment of hallucinations/delusion

PREFERRED AGENTS	NON-PREFERRED AGENTS	
		PA CRITERIA
(No PA required unless otherwise noted)	(PA required)	PACRITERIA
	Risperdal [®] (risperidone) oral solution	Rexulti:
	FDA maximum recommended dose = 16 mg/day Versacloz [®] (clozapine) Oral Suspension QTY LIMIT: 18ml/day FDA maximum recommended dose = 900 mg/day	Indication for use is schizophrenia: the patient has had a documented side effect, allergy or treatment failure with at least three preferred products (typical or atypical antipsychotics), one of which must be aripiprazole OR Indication for use is adjunct treatment of Major Depressive Disorder (MDD): the
		patient has had a documented inadequate response to at least 3 different antidepressants from two different classes AND the patient has had a documented side effect, allergy or treatment failure with two preferred atypical antipsychotic products being used as adjunctive therapy, one of which must be aripiprazole
SHORT-ACTING INJECTABLE PRODUCTS		Aripiprazole Oral Solution: the patient has had a documented side effect, allergy, or treatment failure with preferred risperidone oral solution.
GEODON [®] IM (ziprasidone intramuscular injection) FDA maximum recommended dose = 40 mg/day		Risperdal Oral Solution: The patient has a documented intolerance to the generic product risperidone.
OLANZAPINE IM (compare to Zyprexa® IM) FDA maximum recommended dose = 30 mg/day		Versacloz Oral Solution: The patient has a medical necessity for a non-solid oral dosage form and is unable to use clozapine orally disintegrating tablets.
ZYPREXA® IM (olanzapine intramuscular		Invega Hafyera: The patient is started and stabilized on the medication OR The
injection) FDA maximum recommended dose = 30 mg/day		patient has been adequately treated with Invega Sustenna (paliperidone palmitate 1-month) for at least four months or Invega Trinza (paliperidone palmitate 3-
		month) following at least one 3-month injection cycle. Invega Trinza: The patient is started and stabilized on the medication OR telegrability has been established with Invest Statemen for at least 4 months.
LONG-ACTING INJECTABLE PRODUCTS		tolerability has been established with Invega Sustenna for at least 4 months. Note: This is processed via automated (electronic) step therapy.
ABILIFY MAINTENA® (aripiprazole monohydrate)		ORALLY DISINTEGRATING TABLETS: Medical necessity for a specialty
QTY LIMIT: 1 vial/28 days FDA maximum recommended dose = 400		dosage form has been provided AND If the request is Zyprexa Zydis, the patient has a documented intolerance to the generic equivalent.
mg/month		COMBINATION PRODUCTS: The patient has had a documented side effect,
ARISTADA® (aripiprazole lauroxil) QTY LIMIT: 441, 662, and 882 mg = 1 syringe/28		allergy, or treatment failure with two preferred products OR The prescriber provides a clinically valid reason for the use of the requested medication.
days, 1064 mg = 1 syringe/60 days ARISTADA Initio TM (aripiprazole lauroxil)		Secuado: The indication for use is the treatment of schizophrenia/schizoaffective disorder AND The patient has had a documented side effect, allergy or
INVEGA SUSTENNA (paliperidone palmitate) FDA maximum recommended dose = 234 mg/ month		treatment failure with at least three preferred products (typical or atypical antipsychotics) and Saphris OR The indication for use is the treatment of
PERSERIS® (risperidone)		schizophrenia/schizoaffective disorder AND the patient is unable to take oral
QTY LIMIT: 1 syringe/28 days FDA maximum recommended dose = 120		medications AND the patient has had a documented side effect, allergy or treatment failure with a preferred long-acting injectable.
mg/month		a cament famile with a preferred folig-acting injectable.
RISPERDAL [®] CONSTA (risperidone microspheres) FDA maximum recommended dose = 50 mg/14		
days ZYPREXA RELPREVV® (olanzapine pamoate)		

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
QTY LIMIT: 405 mg = 1 vial/month, 210 and 300 mg = 2 vials/month FDA maximum recommended dose = 600 mg/month Preferred After Clinical Criteria Are Met INVEGA HAFYERA TM (paliperidone palmitate) FDA maximum recommended dose = 1560 mg/6 months INVEGA TRINZA [®] (paliperidone palmitate) FDA maximum recommended dose = 819 mg/3 months ORALLY DISINTEGRATING TABLETS All products require PA	Aripiprazole ODT <i>QTY LIMIT</i> : 10 and 15 mg = 2 tabs/day FDA maximum recommended dose = 30 mg/day Clozapine orally disintegrating tablets FDA maximum recommended dose = 900 mg/day Olanzapine orally disintegrating tablets (compare to Zyprexa Zydis®) <i>QTY LIMIT</i> : 5 and 10 mg = 1.5 tabs/day FDA maximum recommended dose = 20 mg/day Risperidone ODT FDA maximum recommended dose = 16 mg/day Zyprexa Zydis® (olanzapine orally disintegrating tablets) <i>QTY LIMIT</i> : 5 and 10 mg = 1.5 tabs/day FDA maximum recommended dose = 20 mg/day	
COMBINATION PRODUCTS All products require PA	Lybalvi® (olanzapine/samidorphan) <i>QTY LIMIT:</i> 1 tablet/day FDA maximum recommended dose = 20mg/10mg (per day) Olanzapine/fluoxetine FDA maximum recommended dose = 18 mg/75 mg (per day) Secuado (asenapine) transdermal patch <i>QTY LIMIT:</i> 1 patch/day FDA maximum recommended dose = 7.6 mg/day	

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
TRANSDERMAL PRODUCTS All products require PA		
	ANTI-PSYCHOTI	C: TYPICALS
ORAL HALOPERIDOL LOXAPINE PERPHENAZINE PIMOZIDE	Chlorpromazine Fluphenazine Molindone Thioridazine	Chlorpromazine: patient has a diagnosis of acute intermittent porphyria or intractable hiccups OR patient has had a documented side effect, allergy or treatment failure with at least three preferred products (may be typical or atypical anti-psychotics).

LONG ACTING INJECTABLE PRODUCTS

TRIFLUOPERAZINE

FLUPHENAZINE DECANOATE HALOPERIDOL DECANOATE (compare to Haldol® decanoate)

Haldol® decanoate (haloperidol decanoate)

Thiothixene

Fluphenazine Oral Solution: patient has a requirement for an oral liquid dosage form (i.e. swallowing disorder, inability to take oral medications)

Fluphenazine tablets: patient is transitioning to the decanoate formulation or requires supplemental oral dosing in addition to decanoate OR patient has had a documented side effect, allergy or treatment failure with at least three preferred products (may be typical or atypical anti-psychotics).

All other oral medications: patient has had a documented side effect, allergy or treatment failure with at least three preferred products (may be typical or atypical anti-psychotics). If a product has an AB rated generic, one trial must be the generic.

Long Acting Injectable Products: for approval of Haldol decanoate, the patient has a documented intolerance to the generic product.

ANTIRETROVIRAL THERAPY HUMAN IMMUNODEFICIENCY VIRUS (HIV)

SINGLE PRODUCT REGIMENS

Tablets (STRs)

BIKTARVY® (bictegravir/emtricabine/tenofovir AF)

COMPLERA® (emtricitabine/relpivirine/tenofovir)

DELSTRIGO® (doravirine/lamivudine/tenofovir)

DOVATO® (dolutegravir/lamivudine)

EFAVIRENZ/EMTRICITABINE/TENOFOVIR

GENVOYA® (elvitegravir/cobicistat/

emtricitabine/tenofovir AF)

ODEFSEY® (emtricitabine/relpivirine/

tenofovir AF)

SYMFITM (efavirenz/lamivudine/tenofovir)

SYMFITM LO (efavirenz/lamivudine/tenofovir)

TRIUMEQ® (abacavir/lamivudine/dolutegravir)

Juluca® (dolutegravir/rilpivirine)

Stribild® (elvitegravir/cobicistat/

emtricitabine/tenofovir)

Symtuza® (darunavir/cobicistat/emtricitabine/tenofovir AF)

Cabenuva: The patient has been started and stabilized on the requested medication (Note: samples are not considered adequate justification for stabilization.) OR patient is virologically suppressed (HIV-1 RNA < 50 copies per mL) on a stable oral antiretroviral regimen with no history of treatment failure AND medical reasoning beyond convenience or enhanced compliance over preferred agents is provided. Note: oral lead-in with Vocabria® (cabotegravir) and Edurant® (rilpivirine) are provided at no charge and sent directly to the prescriber or patient by a specialty distributor and should be dispensed ONLY for those with prior approval for Cabenuva.

Juluca: The patient has been started and stabilized on the requested medication (Note: samples are not considered adequate justification for stabilization.) OR patient is virologically suppressed (HIV-1 RNA < 50 copies per mL) on a stable oral antiretroviral regimen for at least 6 months AND medical reasoning beyond

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
(No FA required unless otherwise noted)	(FA required)	FA CRITERIA
TRIUMEQ® PD tablets for oral suspension		convenience or enhanced compliance over preferred agents is provided.
(abacavir/lamivudine/dolutegravir)		Stribild:
Long-Acting Injectables All products require PA	Cabenuva® (cabotegravir/rilpivirine) Kit	 The patient has been started and stabilized on the requested medication. (Note: samples are not considered adequate justification for stabilization.) OR Genotype testing supporting resistance to other regimens OR Intolerance or contraindication to preferred combination of drugs AND
		 • Intolerance or contraindication to preferred combination of drugs AND • Medical reasoning beyond convenience or enhanced compliance over preferred agents AND • CrCl > 70mL/min to initiate therapy OR CrCl > 50mL/min to continue
		therapy
		Symtuza: The patient has been started and stabilized on the requested
		medication. (Note: samples are not considered adequate justification for stabilization.) OR Medical reasoning beyond convenience or enhanced
COMPINATION PROPLICITY AIRTH.		compliance over preferred agents (Prezcobix & Descovy)
COMBINATION PRODUCTS - NRTIS ABACAVIR/LAMIVUDINE (compare to Epzicom®)	Combivir® (lamivudine/zidovudine)	Combining Englands nations must have a decumented intelegence to the generic
ABACAVIR/LAMIVUDINE/ZIDOVUDINE ABACAVIR/LAMIVUDINE/ZIDOVUDINE	Epzicom® (abacavir/lamivudine)	Combivir, Epzicom: patient must have a documented intolerance to the generic equivalent
(compare to Trizivir®) LAMIVUDINE/ZIDOVUDINE (compare to Combivir®)	Trizivir® (abacavir/lamivudine/zidovudine)	Trizivir: The patient has been started and stabilized on the requested medication. (Note: samples are not considered adequate justification for stabilization.) OR The prescriber must provide a clinically compelling reason for the use of the requested medication including reasons why any of the preferred products would not be suitable alternatives
COMBINATION PRODUCTS - NUCLEOSIDE & 1	NUCLEOTIDE ANALOG RTIs	
CIMDUOTM (lamivudine/tenofovir) DESCOVY® (emtricitabine/tenofovir AF) EMTRICITABINE/TENOFOVIR (compare to Truvada®)	Truvada® (emtricitabine/tenofovir)	Truvada: patient must have a documented intolerance to the generic equivalent
COMBINATION PRODUCTS – PROTEASE INHII	RITORS	
KALETRA® (lopinavir/ritonavir)	Lopinavir/ritonavir (compare to Kaletra®)	Lopinavir/ritonavir: patient must have a documented intolerance to brand Kaletra
IMMUNOLOGIC THERAPIES		
Preferred After Clinical Criteria Are Met		Rukobia, Trogarzo: The patient must meet ALL of the following criteria:
TROGARZO™ (ibalizumab-uiyk)		• ≥ 18 years of age
QTY LIMIT: 10 vials (2000 mg) x 1 dose then 4 vials (800 mg) every 14 days thereafter		 Prescription is written by or in consultation with an infectious disease specialist
		 Viral Load is ≥ 1,000 copies/mL (results must be submitted) Patient has been compliant but has had an inadequate response to at

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
		least 6 months of treatment with anti-retroviral therapy (ART), including recent failure within the last 8 weeks • Patient has multi-drug resistant HIV-1 infection including documented resistance to at least one medication from each of the following classes: • Protease Inhibitor (PI) • Nucleoside Reverse Transcriptase Inhibitor (NRTI) • Non-nucleoside Reverse Transcriptase Inhibitor (NNRTI) • Medication will be used in combination with ART that includes at least one drug to which the individual's virus is susceptible • Initial approval will be granted for 6 months. For continuation of therapy, there must be a decrease in viral load from baseline AND the patient must continue to be compliant with the optimized background regiment of ART.
GP120 DIRECTED ATTACHMENT INHIBITOR		
Preferred After Clinical Criteria Are Met RUKOBIA® (fostemsavir) QTY LIMIT = 2 tablets per day		
INTEGRASE STRAND TRANSFER INHIBITORS		
ISENTRESS® (raltegravir potassium) ISENTRESS HD (raltegravir potassium)		
TIVICAY® (dolutegravir sodium)		
TIVICAY® PD (dolutegravir sodium)		
NUCLEOSIDE REVERSE TRANSCRIPTASE INH		
ABACAVIR SULFATE (compare to Ziagen®) EMTRIVA® (emtricitabine)	Epivir® (lamivudine) Retrovir® (zidovudine)	Epivir, Retrovir, Viread 300mg, Ziagen: patient must have a documented intolerance to the generic equivalent
LAMIVUDINE (compare to Epivir®)	Stavudine	Stavudine: The patient has been started and stabilized on the requested
TENOFOVIR DISOPROXIL FUMARATE (compare	Viread® (tenofovir disoproxil fumarate) 300mg tablet	medication. (Note: samples are not considered adequate justification for
to Viread®) 300mg	Ziagen® (abacavir sulfate) tablet	stabilization.) OR The prescriber must provide a clinically compelling reason
VIREAD® (tenofovir disoproxil fumarate) 150mg,		for the use of the requested medication including reasons why any of the
200mg, 250mg tablet, 40mg/gm powder		preferred products would not be suitable alternatives.
ZIAGEN® (abacavir sulfate) ZIDOVUDINE (compare to Retrovir®)		
•		
NON-NUCLEOSIDE REVERSE TRANSCRIPTASI		
EDURANT® (rilpivirine)	Etravirine (compare to Intelence®)	Etravirine: patient must have a documented intolerance to brand Intelence.
EFAVIRENZ (compare to Sustiva®) INTELENCE® (etravirine)	Nevirapine (compare to Viramune®) Nevirapine ER (compare to Viramune® ER)	Sustiva: patient must have a documented intolerance to the generic equivalent Nevirapine, Nevirapine ER, Viramune ER: The patient has been started and
PIFELTRO (doravirine)	Sustiva® (efavirenz)	stabilized on the requested medication. (Note: samples are not considered
- ()	Viramune® ER (nevirapine ER)	adequate justification for stabilization.) OR The prescriber must provide a

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
(NO FA required unless otherwise noted)	(FA required)	TA CRITERIA
		clinically compelling reason for the use of the requested medication including
		reasons why any of the preferred products would not be suitable alternatives.
PHARMACOENHANCER-CYTOCHROME P450	INHIBITOR	
All products require PA	Tybost® (cobicistat)	Tybost: The patient has been started and stabilized on the requested medication.
		(Note: samples are not considered adequate justification for stabilization.)
		OR a clinically valid reason beyond compliance or convenience is given for
		not using a preferred combination drug or a ritonavir- based regimen with similar components
PRE-EXPOSURE PROPHYLAXIS (PrEP) AGENT	8	similar components
Apretude® (cabotegravir extended-release)	Truvada® (Emtricitabine/Tenofovir DF) 200mg/300 mg	Truvada: The patient has a documented intolerance to the generic equivalent.
600mg/3mL IM injection	tablet	
Descovy® (emtricitabine/tenofovir AF) 200mg/25mg		
tablet		
Emtricitabine/Tenofovir DF (compare to Truvada®)		
200mg/300mg tablet		
PROTEASE INHIBITORS (PEPTICIC)		
ATAZANAVIR (compare to Reyataz®)	Fosemprenavir (compare to Lexiva®)	Fosemprenavir, Invirase, Lexiva, Viracept: The patient has been started and
EVOTAZ® (atazanavir/cobicistat) NORVIR® (ritonavir)	Invirase® (saquinavir mesylate) Lexiva® (fosemprenavir)	stabilized on the requested medication. (Note: samples are not considered adequate justification for stabilization.) OR The prescriber must provide a
RITONAVIR (compare to Norvir®)	Reyataz® (atazanavir)	clinically compelling reason for the use of the requested medication including
rational vine (compare to restaure)	Viracept® (nelfinavir)	reasons why any of the preferred products would not be suitable alternatives.
		Reyataz: patient must have a documented intolerance to the generic equivalent
PROTEASE INHIBITORS (NON-PEPTIDIC)		
PREZCOBIX® (darunavir/cobicistat)	Aptivus® (tipranavir)	Aptivus: The patient has been started and stabilized on the requested medication.
PREZISTA® (darunavir ethanolate)		(Note: samples are not considered adequate justification for stabilization.) OR The prescriber must provide a clinically compelling reason for the use of the
		requested medication including reasons why any of the preferred products
		would not be suitable alternatives.
ENTRY INHIBITORS-CCR5 CO-RECEPTOR AN	TAGONISTS	
All products require PA	Selzentry® (maraviroc)	Selzentry: The patient has been started and stabilized on the requested
		medication. (Note: samples are not considered adequate justification for
		stabilization.) OR The prescriber must provide a clinically compelling reason
		for the use of the requested medication including reasons why any of the
ENTRY INHIBITORS-FUSION INHIBITORS		preferred products would not be suitable alternatives.
All products require PA	Fuzeon® (enfuvirtide)	Fuzeon: The patient has been started and stabilized on the requested medication.
An products require 1 A	1 uzcone (cinuviruoc)	(Note: samples are not considered adequate justification for stabilization.) OR
		(1.00.) samples are not considered adequate justification for submittation, of

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
		The prescriber must provide a clinically compelling reason for the use of the requested medication including reasons why any of the preferred products would not be suitable alternatives.
	BILE SALTS AND BIL	JARY AGENTS
URSODIOL capsules	Bylvay™ (odevixibat) Chenodal® (chendiol) Cholbam® (cholic acid) Livmarli® (maralixibat) Ocaliva® (obeticholic acid) Urso® (Urosiol) Ursodiol tablets Urso® Forte (ursodiol)	Bylvay: The patient is experiencing moderate to severe pruritis associated with a diagnosis of progressive familial intrahepatic cholestasis (PFIC) confirmed by molecular genetic testing AND the patient does not have a ABCB11 variant resulting in non-functional or complete absence of the bile salt export pump protein (BSEP-3) AND the patient does not have a history of liver transplant or clinical evidence of decompensated cirrhosis AND baseline liver function tests and fat-soluble vitamin (A, D, E, and K) levels have been completed and will be monitored periodically during treatment AND patient has had an inadequate response or contraindication to cholestyramine and ursodiol. For re-approval, there must be documented clinical improvement (e.g. reduced serum bile acid or decreased pruritis). Chenodal: The indication for use is with radiolucent stones in well-opacifying gallbladders, in whom selective surgery would be undertaken except for the presence of increased surgical risk due to systemic disease or age AND the patient does not have any of the following contraindications to therapy: women who are pregnant or may become pregnant, known hepatocyte dysfunction or bile ductal abnormalities such as intrahepatic cholestasis, primary biliary cirrhosis or sclerosing cholangitis. Cholbam: The indication for use is the treatment of bile acid synthesis disorders due to single enzyme defects OR for the adjunctive treatment of peroxisomal disorders, including Zellweger spectrum disorders, AND the patient exhibits manifestations of liver disease, steatorrhea, or complications from decreased fat-soluble vitamin absorption AND the prescriber is a hepatologist or gastroenterologist. Initial approval will be granted for 3 months. For reapproval after 3 months, there must be documented clinical benefit. Livmarli: The patient is experiencing moderate to severe pruritis associated with a diagnosis of Alagille Syndrome (ALGS) AND baseline liver function tests and fat-soluble vitamin (A, D, E, and K) levels have been completed

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	ursodiol. Urso, Ursodiol tablets, Urso Forte: The patient must have a documented treatment limiting side effect to generic ursodiol capsules.
	BONE RESORPTION INH	IIBITORS
ORAL BISPHOSPHONATES TABLETS/CAPSULES ALENDRONATE (compare to Fosamax®) tablets INJECTABLE BISPHOSPHONATES ZOLEDRONIC ACID Injection (compare to Reclast®) 5 mg/100mL QTY LIMIT: 5 mg (one dose)/year ZOLEDRONIC ACID Injection 4mg/5mL concentrate and 4 mg/100mL IV solution	Actonel® (risedronate) Alendronate oral solution Atelvia (risedronate) Delayed Release Tablet QTY LIMIT: 4 tablets/28 days Boniva® (ibandronate) QTY LIMIT: 150 mg = 1 tablet/28 days Fosamax® (alendronate) Fosamax Plus D® (alendronate/vitamin D) Ibandronate (compare to Boniva®) QTY LIMIT: 150 mg = 1 tablet/28 days Risedronate (compare to Actonel®) Boniva® Injection (ibandronate) QTY LIMIT: 3 mg/3 months (four doses)/year Ibandronate Injection (compare to Boniva®) QTY LIMIT: 3 mg/3 months (four doses)/year Reclast® Injection (zoledronic acid) QTY LIMIT: 5 mg (one dose)/year Evista® (raloxifene) Tablet QTY LIMIT: 1 tablet/day Prolia® Injection (denosumab) QTY LIMIT: 60 mg/6 months (two doses)/year Xgeva® (denosumab) QTY LIMIT: 120 mg/28 days	Actonel, Atelvia, Boniva (oral), Ibandronate (oral), Risedronate patient has had a documented side effect, allergy, or treatment failure (at least a sixmonth trial) to generic alendronate tablets AND if the request is for brand, the patient has also had a documented intolerance to generic equivalent. Alendronate Oral Solution: prescriber provides documentation of medical necessity for the specialty dosage form (i.e. inability to swallow tablets, dysphagia). Evista, Fosamax, Reclast: patient has a documented intolerance to the generic formulation. Calcitonin Nasal: patient is started and stabilized on the requested medication. Note: Calcitonin Nasal Spray (brand and generic) no longer recommended for osteoporosis. Miacalcin Injection: patient has a diagnosis/indication of Paget's Disease Fosamax Plus D: there is a clinical reason why the patient is unable to take generic alendronate tablets and vitamin D separately. Forteo, Teriparatide: patient has a diagnosis/indication of postmenopausal osteoporosis in females, primary or hypogonadal osteoporosis in males or glucocorticoid induced osteoporosis AND patient has had a documented side effect, allergy, or treatment failure** to an oral bisphosphonate. AND prescriber has verified that the patient has been counseled about osteosarcoma risk AND for approval for Forteo the patient has had a documented intolerance to generic Teriparatide. Tymlos: patient has a diagnosis/indication of postmenopausal osteoporosis in females AND patient has had a documented side effect, allergy, or treatment failure ** to an oral bisphosphonate and teriparatide AND prescriber has verified that the patient has been counseled about osteosarcoma risk. Boniva Injection, Ibandronate Injection: patient has a diagnosis/indication of postmenopausal osteoporosis AND patient has had a documented side effect or treatment failure** to a preferred bisphosphonate.
ESTROGEN AGONIST/ANTAGONIST		used for another FDA approved indication

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
(1.6.1.24 required unless otherwise noted)	(1711cquireu)	111 CATIBAIN
RALOXIFENE (compare to Evista (B)) Tablet QTY LIMIT: 1 tablet/day INJECTABLE RANKL INHIBITOR All products require PA INJECTABLE SCLEROSTIN INHIBITOR All products require PA CALCITONIN NASAL SPRAY All products require PA CALCITONIN INJECTION All products require PA PARATHYROID HORMONE INJECTION All products require PA	Evenity® (romosozumab-aqqg) injection QTY LIMIT: 210 mg (2 syringes)/month (Lifetime max duration = 12 months) Calcitonin Nasal Spray (compare to Miacalcin®) Miacalcin® (calcitonin) Injection Forteo® (teriparatide) QTY LIMIT: 1 pen (2.4ml/30 days) (Lifetime max duration of treatment = 2 years) Teriparatide (compare to Forteo®) QTY LIMIT: 1 pen/30 days (Lifetime Max duration of treatment = 2 years) Tymlos™ (abaloparatide) injection QTY LIMIT: 1 pen (1.56ml)/30 days (Lifetime max duration of treatment = 2 years)	Xgeva Injection: diagnosis or indication is bone metastases from solid tumors (e.g. prostate, breast, thyroid, non-small lung cancer), multiple myeloma, hypercalcemia of malignancy, or giant cell tumor of bone. Evenity Injection: diagnosis or indication is postmenopausal osteoporosis AND patient has no history of stroke or MI within the previous year AND patient has had a documented side effect or treatment failure** to a preferred bisphosphonate and Teriparatide. **Treatment failure is defined as documented continued bone loss or fracture after one or more years despite treatment with an oral bisphosphonate.
	BOTULINUM TOXIN	S
All products require PA	Botox® (onabotulinumtoxinA) Dysport® (abobotulinumtoxinA) Myobloc® (rimabotulinumtoxinB) Xeomin® (incobotulinumtoxinA)	Criteria for approval of ALL drugs: The medication is being prescribed for an FDA approved indication AND the patient's age is FDA approved for the given indication AND the patient meets the following additional criteria (if applicable). Initial approval will be granted for 3 months unless otherwise noted. For re-approval, the patient must have

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
		documented improvement in symptoms. Additional criteria for Severe Axillary Hyperhidrosis (Botox only): the patient failed an adequate trial of topical therapy. Additional criteria for Overactive bladder or detrusor overactivity (Botox only): the patient failed an adequate trial of at least TWO urinary antispasmodics (either short- or long-acting formulations) Additional criteria for Chronic migraine (Botox only): the patient has ≥ 15 headache days per month, of which ≥ 8 are migraine days, for at least 3 months AND the member has failed or has a contraindication to an adequate trial (≥ 60 days) of at least TWO medications for migraine prophylaxis from at least two different classes (tricyclic antidepressants, SNRI's, beta-blockers, or anticonvulsants). Initial approval will be granted for 6 months. For re-approval after 6 months, the patient must have documentation of a decrease in the number of headache days per month or decreased use of acute migraine medications such as triptans. Additional criteria for chronic sialorrhea (Myobloc and Xeomin): the patient has a documented side effect, allergy, treatment failure, or contraindication to at least two anticholinergic agents (e.g. scopolamine, glycopyrrolate). LIMITATIONS: Coverage of botulinum toxins will not be approved for cosmetic use (e.g., glabellar lines, vertical glabellar eyebrow furrows, facial rhytides, horizontal neck rhytides, etc.). (BOTOX Cosmetic (onabotulinumtoxinA) is not covered)
	BPH AGENTS	
ALPHA BLOCKERS ALFUZOSIN ER QTY LIMIT: 1 tablet/day DOXAZOSIN (compare to Cardura®) TAMSULOSIN (compare to Flomax®) QTY LIMIT: 2 capsules/day TERAZOSIN ANDROGEN HORMONE INHIBITORS DUTASTERIDE (compare to Avodart®) QTY LIMIT: 1 capsule/day FINASTERIDE (compare to Proscar®) QTY LIMIT: 1 tablet/day	Cardura [®] (doxazosin) Cardura XL [®] (doxazosin) <i>QTY LIMIT</i> : 1 tablet/day Flomax [®] (tamsulosin) <i>QTY LIMIT</i> : 2 capsules/day Rapaflo [®] (silodosin) <i>QTY LIMIT</i> : 1 capsule/day Silodosin (compare to Rapaflo ®) <i>QTY LIMIT</i> : 1 tablet/day Avodart [®] (dutasteride) <i>QTY LIMIT</i> : 1 capsule/day Proscar [®] (finasteride) <i>QTY LIMIT</i> : 1 tablet/day	Cardura, Cardura XL: The patient has had a documented side effect, allergy or treatment failure with two alpha blockers, one of which must be generic doxazosin. Cialis, Tadalafil: The patient has a diagnosis of BPH (benign prostatic hypertrophy) AND the patient has a documented treatment failure/inadequate response to a preferred alpha blocker AND the patient has a documented treatment failure/inadequate response to a preferred 5-alpha reductase inhibitor AND for approval of Cialis, the patient must have a documented intolerance to the generic equivalent. Approval will be limited to 5mg daily for a maximum of 26 weeks. Entadfi: The patient has a diagnosis of BPH (benign prostatic hypertrophy) AND the patient has a documented treatment failure/inadequate response to a preferred alpha blocker AND the patient has a documented treatment failure/inadequate response to a preferred 5-alpha reductase inhibitor AND the patient has a documented treatment failure/inadequate response to a maximum of 26 weeks.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
PDE-5 INHIBITORS All products require PA COMBINATION PRODUCT All products require PA	Cialis® (tadalafil) QTY LIMIT: 1 tablet/day Tadalafil (compare to Cialis®) QTY LIMIT: 1 tablet/day Dutasteride/tamsulosin (compare to Jalyn®) QTY LIMIT: 1 capsule/day Entadfi™ (finasteride/tadalafil) QTY LIMIT: 1 capsule/day Jalyn® (dutasteride/tamsulosin) QTY LIMIT: 1 capsule/day	Flomax: The patient has had a documented side effect, allergy or treatment failure with two preferred alpha blockers, one of which must be generic tamsulosin. Rapaflo, Silodosin: The patient has had a documented side effect, allergy or treatment failure with two preferred alpha blockers Avodart, Proscar: The patient has a documented intolerance to the generic equivalent. Dutasteride/tamsulosin, Jalyn: The patient has a diagnosis of BPH (benign prostatic hypertrophy) AND the patient has a documented treatment failure/inadequate response to combination therapy with generic tamsulosin and finasteride AND is unable to take tamsulosin and dutasteride as the individual separate agents AND for approval of Jalyn, the patient must have a documented intolerance to generic dutasteride/tamsulosin. LIMITATIONS: Coverage of androgen hormone inhibitors will not be approved for cosmetic use in men or women (male-pattern baldness/alopecia or hirsutism). (This includes Propecia (finasteride) 1mg and its generic equivalent whose only FDA approved indication is for treatment of male pattern hair loss.).
BULK POWDERS		
https://dvha.vermont.gov/sites/dvha/files/doc_library/Covered%20Compounding%20Products.pdf		
	CARDIAC GLYCOSIDI	ES
DIGOXIN DIGOXIN Oral Solution		
CUSHING'S DISEASE		
All products require PA	Isturisa® (osilodrostat) tablets Korlym® tablets (mifepristone) QTY LIMIT: 4 tablets/day Signifor® (pasireotide) Ampules QTY LIMIT: all strengths = 2 ml (2 amps)/day Maximum day supply = 30 days	Korlym: Patient is ≥18 years of age AND Patient has a diagnosis of endogenous Cushing's syndrome AND Patient is diagnosed with type 2 diabetes mellitus or glucose intolerance AND Patient has hyperglycemia secondary to hypercortisolism AND Patient has failed or is not a candidate for surgery AND Patient has a documented side effect, allergy, treatment failure or contraindication to at least 2 adrenolytic medications (e.g. ketoconazole, etomidate) AND Patient does not have any of the following contraindications to Korlym: Pregnancy

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
		(pregnancy must be excluded before the initiation of therapy or if treatment is interrupted for >14 days in females of reproductive potential. Nonhormonal contraceptives should be used during and one month after stopping treatment in all women of reproductive potential) OR Patient requires concomitant treatment with systemic corticosteroids for serious medical conditions/illnesses (immunosuppression for organ transplant) OR Patient has a history of unexplained vaginal bleeding OR Patient has endometrial hyperplasia with atypia or endometrial carcinoma OR Patient is concomitantly taking simvastatin, lovastatin, or a CYP3A substrate with a narrow therapeutic index (e.g., cyclosporine, dihydroergotamine, ergotamine, fentanyl, pimozide, quinidine, sirolimus, or tacrolimus). Isturisa, Signifor: Patient has a diagnosis of (pituitary) Cushing's disease AND Patient is 18 years of age or older AND Pituitary surgery is not an option or has not been curative Note: Re-approval requires confirmation that the patient has experienced an objective response to therapy (i.e., clinically meaningful reduction in 24-hour urinary free cortisol levels and/or improvement in signs or symptoms of the disease).

GASTROINTESTINAL AGENTS: BOWEL PREP AGENTS, CONSTIPATION/DIARRHEA, IRRITABLE BOWEL SYNDROME-CONSTIPATION (IBS-C), IRRITABLE BOWEL SYNDROME-DIARRHEA (IBS-D), SHORT BOWEL SYNDROME, OPIOID INDUCED CONSTIPATION

Constipation: Chronic, IBS-C, or Opioid-Induced: Length of approval for non-preferred agents: Initial PA of 3 months and & 12 months thereafter		
BULK-PRODUCING LAXATIVES PSYLLIUM OSMOTIC LAXATIVES LACTULOSE POLYETHYLENE GLYCOL 3350 (PEG) STIMULANT LAXATIVE BISACODYL SENNA STOOL SOFTENER DOCUSATE MISCELLANEOUS DICYCLOMINE		 Linzess 72mcg: The patient has a diagnosis of chronic idiopathic constipation (CIC) AND the patient is unable to tolerate the 145 mcg dose Lubiprostone: The patient is 18 years of age or older has had a documented intolerance to brand name Amitiza Relistor Tablets, Symproic: The patient is current using an opiate for at least 4 weeks AND has documented opioid-induced constipation AND has had a documented side effect, allergy, or treatment failure to Amitiza and Movantik. Relistor Injection: The patient must have documented opioid-induced constipation and be receiving palliative care AND the patient must have had documented treatment failure to a 1 week trial of 2 preferred laxatives from 2 different laxative classes used in combination. Ibsrela, Motegrity: The patient is 18 years of age or older. AND the patient has had a documented side effect, allergy, or treatment failure to Amitiza and either Linzess or Trulance.
GUANYLATE CYCLASE-C AGONIST LINZESS® (linaclotide) 145 mcg and 290 mcg (age ≥ 6 years)	Linzess [®] (linaclotide) 72mcg <i>QTY LIMIT</i> : 1 capsule/day	

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
QTY LIMIT: 1 capsule/day TRULANCE® (plecanatide) (age ≥ 6 years) QTY LIMIT: 1 tablet/day	Lubiprostone (compare to Amitiza®) QTY LIMIT: 2 capsules/day	
Note: Linzess® and Trulance® are contraindicated in patients less than 6 years of age due to the risk of serious dehydration.		
CIC-2 CHLORIDE CHANNEL ACTIVATORS AMITIZA® (lubiprostone) (age ≥ 18 years) QTY LIMIT: 2 capsules/day	Relistor [®] (methylnaltrexone) tablets <i>QTY LIMIT:</i> 3 tablets/day Relistor [®] (methylnatrexone) injection Symproic [®] (naldemedine) <i>QTY LIMIT:</i> 1 tablet/day	
OPIOID ANTAGONISTS MOVANTIK® (naloxegol) QTY LIMIT: 1 tablet/day 5-HT4 RECEPTOR ANTAGONISTS All products require PA NHE3 INHIBITORS All products require PA	Motegrity® (prucalopride) QTY LIMIT: 1 tablet/day Ibsrela® (tenapanor) QTY LIMIT: 2 tablets/day	
Short Bowel Syndrome (SBS): Length of appro	val: 6 Months	
All products require PA	Gattex [®] (teduglutide) Vials Maximum day supply = 30 days	Gattex: Patient has a diagnosis of short bowel syndrome AND Patient is receiving specialized nutritional support administered intravenously (i.e. parenteral nutrition) AND Patient does not have an active gastrointestinal malignancy (gastrointestinal tract, hepatobiliary, pancreatic), colorectal cancer, or small bowel cancer. Note: Re-approval requires evidence of decreased parenteral nutrition support from baseline.
Antidiarrheal: HIV/AIDs: Length of approval:	Initial approval 3 months, subsequent 1 year	
DIPHENOXYLATE/ATROPINE LOPERAMIDE	Mytesi® (crofelemer) 125 mg DR Tablets QTY LIMIT: 2 tablets/day	Mytesi: Patient has HIV/AIDS and is receiving anti-retroviral therapy AND Patient is at least 18 years of age AND Patient requires symptomatic relief of noninfectious diarrhea AND Infectious diarrhea (e.g. cryptosporidiosis, c. difficile, etc.) has been ruled out AND Patient has tried and failed at least one anti-diarrheal medication (i.e. loperamide or atropine/diphenoxylate)

PREFERRED AGENTS	NON-PREFERRED AGENTS		
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA	
Antidiarrheal: IBS-D: Length of approval: Init	ial approval 3 months; subsequent 1 year		
All products require PA	Alosetron (compare to Lotronex®) Lotronex® (alosetron) Viberzi® (eluxadoline) Xermelo™ (telotristat ethyl) QTY LIMIT: 3 tablets/day	 Lotronex/alosetron: The patient is a woman and has a diagnosis of severe diarrhea-predominant irritable bowel syndrome (IBS) with symptoms lasting 6 months or longer AND has had anatomic or biochemical abnormalities of the GI tract excluded AND has not responded adequately to conventional therapies such as loperamide and TCA's. For approval of generic alosetron, the patient must have documented intolerance to brand Lotronex. Viberzi: The patient has a diagnosis of IBS-D AND does not have any of the following contraindications to therapy A) known or suspected biliary duct obstruction, or sphincter of Oddi disease or dysfunction B) alcoholism, alcohol abuse, alcohol addiction, or drink more than 3 alcoholic beverages/day C) a history of pancreatitis; structural diseases of the pancreas D) severe hepatic impairment (Child-Pugh Class C) AND has not responded adequately to conventional therapies such as loperamide and TCA's. Xermelo: The patient has a diagnosis of carcinoid syndrome diarrhea AND had an inadequate treatment response (defined as 4 or more bowel movements per day) despite use of a long-acting somatostatin analog for at least 3 consecutive months AND the medication will be used in combination with a long-acting somatostatin analog therapy. For reauthorization, documentation showing a decrease in the number of bowel movements per day is required. Note: Xermelo will not be approved in treatment naïve patients or as monotherapy. 	
BOWEL PREP AGENTS		N. C. I. C.	
CLENPIQ® GAVILTYE-G, GAVILYTE-H, GAVILYTE-N MOVIPREP PEG-3350	Gavilyte-C Golytely Nulytely Plenvu® Suprep® Sutab®	Non-preferred agents: The patient has a documented intolerance or treatment failure of at least one preferred agent (defined by failure to complete cleansing of the colon as a preparation for colonoscopy) AND if the product has an AB rated generic, there must have been a trial with the generic formulation.	
	CONTINUOUS GLUCOSE MONITORS		
Initial approval will be granted for 6 months;	renewals up to 1 year thereafter		
Preferred After Clinical Criteria Are Met DEXCOM G6 Initial prescription: 1 receiver, 1 wireless	Medtronic Guardian™ Connect Initial Prescription: 1 transmitter, 5 sensors	Patient has a diagnosis of Diabetes Mellitus AND • 2 years of age or older for Dexcom G6, ≥ 14 years for Medtronic	

PREFERRED AGENTS NON-PREFERRED AGENTS (PA required) (No PA required unless otherwise noted) PA CRITERIA transmitter, and 1 3-pack of sensors **Refill Quantity Limits:** 1 transmitter every year, 1 Guardian, or ≥ 4 years for Freestyle Libre 2, or ≥ 18 for Freestyle Libre **Refill Quantity Limits:** 1 transmitter every 3 sensor every 7 days (maximum of 5 sensors every months, 1 sensor every 10 days (maximum of 9 35 days) Patient requires multiple daily injections of a rapid/short acting insulin or sensors every 90 days) Medtronic 670G Guardian Link 3 is on an insulin pump. FREESTYLE LIBRE PRO (10-DAY SENSORS) Approval of non-preferred products will be limited to cases where the **Initial Prescription:** 1 transmitter, 5 sensors CGM is directly integrated with the patient's insulin pump. The make and **Initial Prescription:** 1 reader, 3 sensors Refill Quantity Limits: 1 transmitter every year, 1 **Refill Quantity Limits:** 1 sensor every 10 days model of pump must be documented on the prior authorization. sensor every 7 days (maximum of 5 sensors every (maximum of 9 sensors every 90 days) 35 days) FREESTYLE LIBRE 14 DAY (14-DAY SENSORS) **Re-authorization:** Medtronic 770G Guardian Link 3 **Initial Prescription:** 1 reader, 2 sensors There is documented evidence of compliance to CGM (log data and/or **Initial Prescription:** 1 transmitter, 5 sensors **Refill Quantity Limits:** 1 sensor every 14 days office visit notes required). (maximum of 6 sensors every 84 days) Refill Quantity Limits: 1 transmitter every year 1 Replacement will be considered when medically necessary and not for FREESTYLE LIBRE 2 (14-DAY SENSORS) sensor every 7 days (maximum of 5 sensors every recent technology upgrades (device must be malfunctioning and out of **Initial Prescription:** 1 reader, 2 sensors 35 days) warranty). **Refill Quantity Limits:** 1 sensor every 14 days Medtronic MiniLink (includes Enlite Serter) (maximum of 6 sensors every 84 days) **Initial Prescription:** 1 transmitter, 5 sensors FREESTYLE LIBRE 3 (14-DAY SENSORS) Refill Quantity Limits: 1 transmitter every year, 1 **Initial Prescription:** 1 reader, 2 sensors sensor every 7 days (maximum of 5 sensors every **Refill Quantity Limits:** 1 sensor every 14 days (maximum of 6 sensors every 84 days) 35 days) CONTRACEPTIVES **SELECT PRODUCTS:** Length of approval: 1 year MONOPHASIC AGENTS: **Non-preferred agents:** Trial with at least three preferred contraceptive products Due to the extensive list of products, any Beyaz (drospirenone/ethinyl estradiol/levomefol) monophasic BCP not listed as non-preferred is including the preferred formulation of the requested non-preferred agent Blisovi FE 24 (norethindrone/ethinyl estradiol/FE) considered preferred. Drospirenone/ethinyl estradiol/levomefol Kaitlib (norethindrone/ethinyl estradiol/FE) Layolis FE (norethindrone/ethinyl estradiol/FE) Lo-Estrin (norethindrone/ethinyl estradiol) Lo-Estrin FE (norethindrone/ ethinyl estradiol/FE)

Melodetta FE (drospirenone/ethinyl estradiol/levomefol) Mibelis FE (norethindrone/ethinyl

Nexstellis (drospirenone/estetrol)

Noretin-Eth Estra-Ferros Fum Tab Chew 0.8-25(24)

(norethindrone/ethinyl estradiol/FE)

estradiol/FE)

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
(110 111 required amoss outer wise noted)	(1111equilled)	
	Noretin-Eth Estra-Ferros Fum Tab Chew 1MG-20(24)	
BIPHASIC AGENTS		
AZURETTE (desogestrel/ethinyl estradiol) BEKYREE (desogestrel/ethinyl estradiol) DESOGESTREL/ETHINYL ESTRADIOL KARIVA (desogestrel/ethinyl estradiol) KIMIDESS (desogestrel/ethinyl estradiol) NORETHIDRONE/ETHINYL ESTRADIOL 0.5/1-35 PIMTREA (desogestrel/ethinyl estradiol) SIMLIYA (desogestrel/ethinyl estradiol) VIORELE (desogestrel/ethinyl estradiol) VOLNEA (desogestrel/ethinyl estradiol)	Lo Loestrin FE (norethindrone/ ethinyl estradiol/FE) Mircette (desogestrel/ ethinyl estradiol)	Non-preferred agents: Trial with at least three preferred contraceptive products including the preferred formulation of the requested non-preferred agent
TRIPHASIC AGENTS		
ALYACEN (norethindrone ethinyl estradiol) ARANELLE (norethindrone/ethinyl estradiol) CAZIANT (desogestrel/ ethinyl estradiol) CYCLAFEM (norethindrone/ethinyl estradiol) DASETTA (norethindrone/ethinyl estradiol) ENPRESSE (levonorgestrel/ ethinyl estradiol) LEENA (norethindrone/ethinyl estradiol) LEVONEST (levonorgestrel/ ethinyl estradiol) NATAZIA (dienogest/estradiol valerate) NORGESTIMATE/ETHINYL ESTRADIOL NORTREL 7/7/7 (norethindrone/ethinyl estradiol) PIRMELLA (norethindrone/ethinyl estradiol) TRI-ESTARYLLA (norgestimate/ ethinyl estradiol) TRI-FEMYNOR (norgestimate/ ethinyl estradiol) TRI-LINYAH (norgestimate/ ethinyl estradiol) TRI-LO-ESTARYLLA (norgestimate/ethinyl estradiol)	Estrostep FE (norethindrone/ethinyl estradiol/FE) Tilia FE (norethindrone/ethinyl estradiol/FE) Tri-Legest FE (norethindrone/ethinyl estradiol/FE) estradiol/FE)	Non-preferred agents: Trial with at least three preferred contraceptive products including the preferred formulation of the requested non-preferred agent

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PREFERRED AGENTS	NON-PREFERRED AGENTS	DA CIDITEDIA
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
TRI-LO-MARZIA (norgestimate/ethinyl estradiol) TRI-LO-SPRINTEC (norgestimate/ethinyl estradiol) TRI-PREVIFEM (norgestimate/ ethinyl estradiol) TRI-SPRINTEC (norgestimate/ ethinyl estradiol) TRI-VYLIBRA (norgestimate/ ethinyl estradiol) TRI-VYLIBRA LO (norgestimate/ ethinyl estradiol) TRIVORA (levonorgestrel/ ethinyl estradiol) VELIVET (desogestrel/ ethinyl estradiol) EXTENDED CYCLE		
EATERDED CICLE		
AMETHIA (levonorgestrel/ ethinyl estradiol) AMETHIA LO (levonorgestrel/ ethinyl estradiol) AMETHYST (levonorgestrel/ ethinyl estradiol) ASHLYNA (levonorgestrel/ ethinyl estradiol) CAMRESE (levonorgestrel/ ethinyl estradiol) CAMRESE LO (levonorgestrel/ ethinyl estradiol) DAYSEE (levonorgestrel/ ethinyl estradiol) INTROVALE (levonorgestrel/ ethinyl estradiol) JAIMIESS (levonorgestrel/ ethinyl estradiol) JOLESSA (levonorgestrel/ ethinyl estradiol) JOLESSA (levonorgestrel/ ethinyl estradiol) LEVONORGESTREL/ETHINYL ESTRADIOL TBDSPK 3 month LO-SEASONIQUE (levonorgestrel/ ethinyl estradiol) SIMPESSE (levonorgestrel/ ethinyl estradiol) SEASONIQUE (levonorgestrel/ ethinyl estradiol) SETLAKIN (levonorgestrel/ethinyl estradiol)	Fayosim (levonorgestrel/ ethinyl estradiol) Quartette (levonorgestrel/ ethinyl estradiol) Rivelsa (levonorgestrel/ ethinyl estradiol)	Non-preferred agents: Trial with at least three preferred contraceptive products including the preferred formulation of the requested non-preferred agent
PROGESTIN ONLY CONTRACEPTIVES		
CAMILA (norethindrone) DEBLITANE (norethindrone) ERRIN (norethindrone) HEATHER (norethindrone) INCASSIA (norethindrone) JENCYCLA (norethindrone) JOLIVETTE (norethindrone) LYZA (norethindrone) NORA-BE (norethindrone) NORETHINDRONE 0.35MG NORLYNDA (norethindrone)	Slynd® (drospirenone)	Non-preferred agents: Trial with at least three preferred contraceptive products including the preferred formulation of the requested non-preferred agent.

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
(NO FA required unless otherwise noted)	(FA required)	TA CRITERIA
SHAROBEL (norethindrone) TULANA (norethindrone)		
INJECTABLE CONTRACEPTIVES		
MEDROXYPROGESTERONE ACETATE 150MG (IM) VIAL/SYRINGE DEPO-PROVERA 104 (SUB-Q) SYRINGE (medroxyprogesterone acetate)	Depo-Provera (IM) (medroxyprogesterone acetate) 150 mg Susp vial/syringe	Depo-Provera IM: Patient must have a documented intolerance to medroxyprogesterone acetate 150mg.
VAGINAL RING		
NUVARING® (etonogestrel/ethinyl estradiol vaginal ring)	Annovera® (segesterone acetate/ethinyl estradiol vaginal ring) QTY LIMIT: 1 ring/year Eluryng (etonogestrel/ethinyl estradiol vaginal ring) Etonogestrel/ethinyl estradiol vaginal ring	Non-preferred agents: Trial with at least three preferred contraceptive products including the preferred formulation of the requested non-preferred agent.
LONG ACTING REVERSIBLE CONTRACEPTIV	YES (LARCs)	
KYLEENA (levonorgestrel) IUD LILETTA (levonorgestrel) IUD MIRENA (levonorgestrel) IUD PARAGARD (copper) IUD SKYLA (levonorgestrel) IUD NEXPLANON (etonogestrel) Implant		
TOPICAL CONTRACEPTIVES		
TWIRLA® (levonorgestrel/ethinyl estradiol) patch XULANE PATCH (norelgestromin/ethinyl estradiol)	Zafemy (norelgestromin/ethinyl estradiol) patch	Zafemy: Trial with at least three preferred contraceptive products including the preferred formulation of the requested non-preferred agent.
VAGINAL CONTRACEPTIVES		
Please refer to the DVHA website for covered OTC spermicidal gels https://dvha.vermont.gov/sites/dvha/files/documents/providers/Pharmacy/OTCWebList.pdf	Phexxi TM (lactic acid, citric acid, and potassium bitartrate) vaginal gel	Phexxi: Use of hormonal contraceptives is contraindicated AND the patient has a documented side effect or allergy to nonoxynol-9
EMERGENCY CONTRACEPTIVES		

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
AFTERA (levonorgestrel) ECONTRA EZ (levonorgestrel) LEVONORGESTREL MY CHOICE (levonorgestrel) MY WAY (levonorgestrel) NEW DAY (levonorgestrel) OPCICON ONE-STEP (levonorgestrel) OPTION 2 (levonorgestrel)		
CORO	NARY VASODILATORS/ANTIANGINALS	S/SINUS NODE INHIBITORS
ORAL		
ISOSORBIDE DINITRATE tablet (compare to Isordil®) ISOSORBIDE DINITRATE ER tablet ISOSORBIDE MONONITRATE tablet ISOSORBIDE MONONITRATE ER tablet ISOSORBIDE MONONITRATE ER tablet NITROGLYCERIN SPRAY LINGUAL (compare to Nitrolingual Pump Spray®) NITROSTAT® (nitroglycerin SL tablet) RANOLAZINE SR 12 HR (compare to Ranexa®) QTY LIMIT: 500 mg = 3 tablets/day, 1000 mg = 2 tablets/day	Aspruzyo Sprinkle TM (ranolazine) granule <i>QTY LIMIT</i> : 500 mg = 3 packets/day, 1000 mg = 2 packets/day BiDil [®] (isosorbide dinitrate/hydralazine) Dilatrate-SR [®] (isosorbide dinitrate SR capsule) Isosorbide dinitrate SL tablet Isordil [®] (isosorbide dinitrate tablet) Nitrolingual Pump Spray [®] Ranexa [®] (ranolazine) <i>QTY LIMIT</i> : 500 mg = 3 tablets/day, 1000 mg = 2 tablets/day	 Aspruzyo: the patient has medical necessity for a non-solid oral dosage form. Dilatrate-SR, Isosorbide dinitrate SL tablet, Isordil: the patient has had a side effect, allergy, or treatment failure to at least two preferred agents. Nitrolingual Pump Spray: the patient has had a side effect, allergy, or treatment failure to Nitroglycerin spray lingual. Bidil: The prescriber provides a clinically valid reason why the patient cannot use isosorbide dinitrate and hydralazine as separate agents. Ranexa: the patient has a documented intolerance to the generic equivalent.
TOPICAL		
NITRO-BID [®] (nitroglycerin ointment) NITROGLYCERIN TRANSDERMAL PATCHES (compare to Nitro-Dur [®])	Nitro-Dur [®] (nitroglycerin transdermal patch)	Nitro-Dur: patient has had a side effect, allergy, or treatment failure to generic nitroglycerin transdermal patches.
SINUS NODE INHIBITORS		
All products require a PA	Corlanor® (ivabradine)	Corlanor Clinical Criteria:

Diagnosis of stable, symptomatic heart failure:

• Resting heart rate ≥ 70 bpm AND

• In sinus rhythm AND

• Left ventricular ejection fraction of $\leq 35\%$ AND

QTY LIMIT: 60 tabs/30 days

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
		 Patient has persisting symptoms despite maximally tolerated doses of beta blockers or who have contraindication to beta blocker therapy Diagnosis of Inappropriate Sinus Tachycardia: Patient has persisting symptoms despite maximally tolerated doses of beta blockers or there is a contraindication to beta blocker therapy. Diagnosis of Postural Orthostatic Tachycardia Syndrome (POTS) The patient has a documented side effect, allergy, or treatment failure with at least 2 of the following medications: fludrocortisone, midodrine, beta blocker (metoprolol or propranolol), or pyridostigmine.
CORTICOSTEROIDS: ORAL		
DEXAMETHASONE tablets, elixir, intensol, solution DEXPAK [®] tabs (dexamethasone taper pack) HYDROCORTISONE tab (compare to Cortef [®]) MEDROL [®] (methylprednisolone) 2mg tablets METHYLPREDNISOLONE (compare to Medrol [®]) tabs METHYLPREDNISOLONE DOSE PACK (compare to Medrol Dose Pack [®]) tabs PREDNISOLONE 3 mg/ml oral solution, syrup PREDNISOLONE SODIUM PHOSPHATE 3 mg/ml oral solution (compare to Orapred [®]) PREDNISOLONE SOD PHOSPHATE ORAL SOLUTION 6.7mg/5ml (5mg/5ml base) (compare to Pediapred [®]) PREDNISONE intensol, solution, tablets	Alkindi® Sprinkle (hydrocortisone) granule Cortef [®] (hydrocortisone) tablets Hemady® (dexamethasone) tablets Medrol [®] (methylprednisolone) tablets Medrol Dose Pak [®] (methylprednisolone) tabs Prednisolone sodium phosphate oral solution 25 mg/5ml Rayos [®] (prednisone) Delayed Release Tablet <i>QTY LIMIT</i> : 1 tablet/day Tarpeyo TM (budesonide) delayed release capsule	 Rayos: The patient has had a trial of generic immediate release prednisone and has documented side effects that are associated with the later onset of activity of immediate release prednisone taken in the morning. Tarpeyo: The patient has a diagnosis of Immunoglobulin A Nephropathy (IgAN) confirmed by biopsy AND eGFR ≥ is 35ml/min/1.73m² AND The patient meets one of the following: Proteinuria ≥ 1g/day or Urine protein-to-creatinine ratio (UPCR) ≥ 1.5 g/g AND The patient is on a stable dose of maximally tolerated ACE-I or ARB therapy for a minimum of 3 months AND The patient's kidney function has continued to decline despite treatment with a preferred oral corticosteroid AND Duration of therapy does not exceed 9 months All Others: The patient has a documented side effect, allergy, or treatment failure to at least two preferred medications. If a product has an AB rated generic, one trial must be the generic formulation.
	COUGH AND COLD PREPAR	ATIONS
Please refer to the DVHA website for covered OTC cough & cold products https://dvha.vermont.gov/sites/dvha/files/documents/providers/Pharmacy/OTCWebList.pdf All RX generics Note: The FDA restricts the use of prescription	Hydrocodone/chlorpheniramine (compare to Tussionex®) QTY LIMIT: 60 ml/RX Tussionex® (hydrocodone/chlorpheniramine) QTY LIMIT: 60 ml/RX TussiCaps® (hydrocodone/chlorpheniramine) OTY LIMIT: 12 capsules/RX	Tussionex, TussiCaps, Hydrocodone/chlorpheniramine suspension (generic): The patient has had a documented side effect, allergy, or treatment failure to two of the following generically available cough or cough/cold products: hydrocodone/homatropine (compare to Hycodan), promethazine/codeine (previously Phenergan with Codeine), guaifenesin/codeine (Cheratussin AC) or benzonatate. AND patient is 6 years old of age or greater. AND The

QTY LIMIT: 12 capsules/RX

Note: The FDA restricts the use of prescription

quantity requested does not exceed 60 ml (Tussionex) or 12 capsules

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
codeine pain and cough medicines in children. Prior authorization is required for patients <12 years of age.	All other brands	 (TussiCaps). AND If the request is for Tussionex, the patient has a documented intolerance to generic hydrocodone/chlorpheniramine suspension. All Other Brands: The prescriber must provide a clinically valid reason for the use of the requested medication including reasons why any of the generically available preparations would not be a suitable alternative.

CYSTIC FIBROSIS MEDICATIONS

Preferred After Clinical Criteria Are Met

KITABIS® (tobramycin sol)

QTY LIMIT: 56 vials/56 days; maximum day supply = 56 days (2 vials/day for 28 days, then 28 days off)

 $\mathsf{TOBI}^{\circledR}$ PODHaler (tobramycin capsules for inhalation)

QTY LIMIT: 224 capsules/56 days; maximum day supply = 56 days (4 capsules twice daily for 28 days, then 28 days off)

TOBRAMYCIN inhalation solution (compare to Tobi®) 300mg/5mL

QTY LIMIT: 56 vials/56 days; maximum day supply = 56 days (2 vials/day for 28 days, then 28 days off) Bethkis® (tobramycin) inhalation solution *QTY LIMIT*: 56 vials/56 days; maximum day supply = 56 days (2 vials/day for 28 days, then 28 days off)

Bronchitol® (mannitol) capsules for inhalation QTY LIMIT: 560 capsules/28 days; maximum day supply = 28 days

Cayston® (aztreonam) inhalation solution

QTY LIMIT: 84 vials/56 days; maximum day supply = 56 days (3 vials/day for 28 days, then 28 days off)

Kalydeco® (ivacaftor) tablets

QTY LIMIT: 2 tablets/day, maximum day supply = 30 days

Kalydeco® (ivacaftor) packets

QTY LIMIT: 2 packets/day; maximum day supply = 30 days

Orkambi® (lumacaftor/ivacaftor)

QTY LIMIT: 120/30 days; maximum day supply=30 days

Pulmozyme® (dornase alfa) inhalation solution

QTY LIMIT: 60/30 days; maximum day supply=30 days

Symdeko® (tezacaftor/ivacaftor and ivacaftor)

QTY LIMIT: 56/28 days; maximum day supply = 28 days

Tobi® (tobramycin) inhalation solution

QTY LIMIT: 56 vials/56 days; maximum day supply = 56 days (2 vials/day for 28 days, then 28 days off)

Kitabis, Tobramycin inhalation solution (300mg/5mL), Pulmozyme: diagnosis or indication is cystic fibrosis

Bethkis, TOBI, tobramycin inhalation solutions (300mg/4mL): Diagnosis or indication is cystic fibrosis and the patient has a documented failure or intolerance to two preferred formulations of tobramycin inhalation solution.

Bronchitol: Diagnosis or indication is cystic fibrosis AND the patient is 18 years of age or older AND the patient has a documented inadequate response or contraindication to hypertonic saline and Pulmozyme AND the patient has passed the Bronchitol Tolerance Test (BTT) AND the patient has been counseled to use a short-acting beta agonist (SABA) 5-15 minutes prior to each dose.

Cayston: diagnosis or indication is cystic fibrosis and the patient has had a documented failure, intolerance or inadequate response to inhaled tobramycin therapy alone

Kalydeco: The patient has a diagnosis of Cystic Fibrosis AND Patient has a mutation on at least one allele in the cystic fibrosis transmembrane conductance regulator gene (CFTR gene) shown to be responsive to Kalydeco per FDA approval (documentation provided). AND The patient is ≥ 1 month old. Note: Renewal of Prior Authorization will require documentation of member response.

TOBI PODHALER: allowed after a trial of another form of inhaled tobramycin **Orkambi/Symdeko/Trikafta:** The patient has a diagnosis of Cystic Fibrosis AND

Initial Criteria

- Patient age is FDA approved for the requested medication AND
- Patient must have a confirmed mutation in the CFTR gene shown to be responsive to the requested medication per FDA approval (documentation provided) AND
- If the patient is under the age of 18, they must have undergone a baseline ophthalmic examination to monitor for lens opacities/cataracts AND

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
((Crossiquests)	
	Tobramycin inhalation solution 300mg/4mL QTY LIMIT: 56 vials/56 days; maximum day supply = 56 days (2 vials/day for 28 days, then 28 days off) Trikafta® (elexacaftor/tezacaftor/ivacaftor) QTY LIMIT: 84/28 days; maximum day supply = 28 days	 Prescriber is a CF specialist or pulmonologist Ongoing Approval Criteria Patient has clinically documented improvement in lung function (will be applied to the first renewal request only; requirement waived on subsequent renewals) Patient has LFTs/bilirubin monitored every 3 months for the first year of therapy and annually after the first year ALT or AST ≤ 5 X the upper limit of normal or ALT/AST ≤ 3 X the upper limits of normal and bilirubin is ≤ 2 X the upper limit of normal For patients under the age of 18, have follow up ophthalmic exam at least annually
	DERMATOLOGICAL AGE	NTS
ACTINIC KERATOSIS THERAPY		
CARAC [®] (fluorouracil) 0.5% cream FLUOROURACIL (compare to Efudex®) 5% cream IMIQUIMOD 5% Cream C=cream, F=foam, G=gel, L=lotion, O=ointment, S=solution	Aldara [®] (imiquimod) 5 % Cream Diclofenac Sodium 3 % Gel (compare to Solaraze [®]) QTY LIMIT: 1 tube/30 days Efudex® (fluorouracil) 5% cream Fluorouracil 5%, 2% solution Fluorouracil (compare to CARAC [®]) 0.5% cream Zyclara (imiquimod) 3.75 % Cream QTY LIMIT: 56 packets/6 weeks Zyclara (imiquimod) 2.5%, 3.75 % Cream Pump QTY LIMIT: 2 pumps/8 weeks	 Aldara: the patient has a documented intolerance to generic imiquimod 5% cream Efudex cream, Fluorouracil solution: The patient has a documented intolerance to fluorouracil 5% cream. Fluorouracil 0.5% cream: The patient has a documented intolerance to brand Carac. Diclofenac Gel: The diagnosis or indication is actinic keratosis AND The patient has had a documented side effect, allergy, contraindication or treatment failure with a preferred topical fluorouracil product. Zyclara Cream: The diagnosis or indication is actinic keratosis on the face or scalp AND The patient has had a documented side effect, allergy, or treatment failure with 5-fluorouracil and imiquimod 5% cream. OR The treatment area is greater than 25 cm2 on the face or scalp. AND The patient has had a documented side effect, allergy, or treatment failure with 5-fluorouracil.
ANTIBIOTICS TOPICAL		
SINGLE AGENT BACITRACIN MUPIROCIN OINTMENT (compare to Bactroban®) COMBINATION PRODUCTS BACITRACIN-POLYMYXIN NEOMYCIN-BACITRACIN-POLYMYXIN	Centany [®] Ointment (mupirocin) Gentamicin Cream or Ointment Mupirocin cream (compare to Bactroban [®]) Xepi cream (ozenoxacin)	Mupirocin cream, Centany Ointment, Xepi cream: The patient has had a documented intolerance with generic mupirocin ointment Gentamicin Cream or Ointment: The patient has had a documented side-effect, allergy, or treatment failure with at least one preferred generic topical antibiotic

NON-PREFERRED AGENTS (PA required)	PA CRITERIA
Ciclodan® (ciclopirox 8% solution) Kerydin® (tavaborole 5% solution) Jublia® (efinaconazole 10% solution) QTY LIMIT: 48 weeks treatment	Ciclodan, Jublia, Kerydin: The patient meets at least 1 of the following criteria: Pain to affected area that limits normal activity, Diabetes Mellitus, Patient is immunocompromised, Patient has diagnosis of systemic dermatosis, Patient has significant vascular compromise AND Documented intolerance to generic ciclopirox 8% solution. LIMITATIONS: Coverage of Onychomycosis agents will NOT be approved solely for cosmetic purposes. Kits with multiple drug products or non-drug items not covered.
Butenafine (compare to Mentax®) 1% C Ciclodan® (ciclopirox) C Econazole 1% C Ertaczo® (sertaconazole) 2% C Extina® (ketoconazole) 2% F Ketoconazole (compare to Extina®) 2 % Foam Luliconazole 1% C Luzu® (luliconazole) 1% Cream Mentax® 1% C Naftifine (compare to Naftin®) 1% & 2% C, 1% G Naftifine (compare to Naftin®) 1% & 2% C, 1% G Naftin® (naftifine) 1% C, 1%, 2% G Nystop®, Nyamyc® (nystatin) P Oxistat® (oxiconazole) 1% C Miconazole w/zinc oxide (compare to Vusion®) O QTY LIMIT: 50 g/30 days Vusion® (miconazole w/zinc oxide) O QTY LIMIT: 50 g/30 days All other branded products Note: Please refer to "Dermatological: Antifungals: Onychomycosis" for ciclopirox solution	 All Non-Preferred Agents (except Vusion): The patient has had a documented side effect, allergy, or treatment failure to at least TWO different preferred generic topical antifungal agents. (If a product has an AB rated generic, one trial must be the generic equivalent of the requested product.) OR The patient has a contraindication that supports the need for a specific product or dosage form of a brand topical antifungal. Miconazole w/ Zinc Oxide, Vusion: The patient has a diagnosis of diaper dermatitis complicated by documented candidiasis AND The patient is at least 4 weeks of age. AND The patient has had two trials (with two different preferred antifungal agents) used in combination with a zinc oxide diaper rash product resulting in documented side effects, allergy, or treatment failures.
	Ciclodan® (ciclopirox 8% solution) Kerydin® (tavaborole 5% solution) Jublia® (efinaconazole 10% solution) QTY LIMIT: 48 weeks treatment Butenafine (compare to Mentax®) 1% C Ciclodan® (ciclopirox) C Econazole 1% C Ertaczo® (sertaconazole) 2% C Extina® (ketoconazole) 2% F Ketoconazole (compare to Extina®) 2 % Foam Luliconazole 1% C Luzu® (luliconazole) 1% Cream Mentax® 1% C Naftifine (compare to Naftin®) 1% & 2% C, 1% G Naftifine (compare to Naftin®) 1% & 2% C, 1% G Nystop®, Nyamyc® (nystatin) P Oxistat® (oxiconazole) 1% C Miconazole w/ zinc oxide (compare to Vusion®) O QTY LIMIT: 50 g/30 days Vusion® (miconazole w/zinc oxide) O QTY LIMIT: 50 g/30 days All other branded products Note: Please refer to "Dermatological: Antifungals:

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
ACYCLOVIR (compare to Zovirax®) 5 % O ZOVIRAX® (acyclovir) 5% C C=cream, O=ointment	Acyclovir (compare to Zovirax [®]) 5 % O Denavir [®] (penciclovir) 1% C Docosanol 10% C Xerese® (acyclovir 5%/hydrocortisone 1%) C Zovirax [®] (acyclovir) 5% O	Acyclovir cream: The patient has a documented intolerance to brand Zovirax cream. Denavir, Docosanol, Xerese: The patient has a treatment failure with a preferred topical acyclovir product. Zovirax ointment: The patient has a documented intolerance to generic acyclovir ointment
AXILLARY HYPERHIDROSIS THERAPY		
Xerac-AC (aluminum chloride) 6.25% Solution	Qbrexza TM (glycopyrronium) 2.4% single use pads <i>QTY LIMIT</i> : 30 pads/month	Qbrexza : the patient has had a documented side effect, allergy, or treatment failure with Xerac-AC
CORTICOSTEROIDS: LOW POTENCY		
ALCLOMETASONE 0.05% C, O DESONIDE 0.05% C, O FLUOCINOLONE 0.01% C, S, oil (compare to Derma-Smoothe, Synalar®) HYDROCORTISONE 0.5%, 1%, 2.5% C; 2.5% L, 0.5%, 1%, 2.5% O C=cream, F=foam, G=gel, L=lotion, O=ointment, S=solution	Derma-Smoothe [®] (fluocinolone 0.01%) oil Desonide 0.05% L Synalar [®] (fluocinolone) 0.01% S All other brands	CRITERIA FOR APPROVAL (NON-PREFERRED AGENTS): The patient has a documented side effect, allergy, or treatment failure to at least two different preferred agents of similar potency. (If a product has an AB rated generic, one trial must be the generic.)
CORTICOSTEROIDS: MEDIUM POTENCY		
BETAMETHASONE DIPROPIONATE 0.05% C, L, O BETAMETHASONE VALERATE 0.1% C, L, O BETAMETHASONE VALERATE 0.12% (compare to Luxiq®) F FLUOCINOLONE 0.025% C, O (compare to Synalar®) FLUTICASONE 0.05% C; 0.005% O HYDROCORTISONE VALERATE 0.2% C, O MOMETASONE FUROATE 0.1% C, L, O, S TRIAMCINOLONE ACETONIDE 0.025%, 0.1% C, L, O	Clocortolone 0.1% C (compare to Cloderm®) Cloderm® (clocortolone) 0.1% C Desoximetasone 0.05% C, O (compare to Topicort®) Flurandrenolide C, L, O Fluticasone 0.05%, L Hydrocortisone Butyrate 0.1% C, O, S Kenalog® (triamcinolone) Aerosol Spray Luxiq® (betamethasone valerate) F Prednicarbate 0.1% C, O Synalar® (fluocinolone) 0.025% C, O Topicort® (desoximetasone) 0.05% C, O Triamcinolone Aerosol Spray Trianex® (triamcinolone) 0.05% O	CRITERIA FOR APPROVAL (NON-PREFERRED AGENTS): The patient has a documented side effect, allergy, or treatment failure to at least two different preferred agents of similar potency. (If a product has an AB rated generic, one trial must be the generic.)
C=cream, F=foam, G=gel, L=lotion, O=ointment, S=solution	All other brands	

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
CORTICOSTEROIDS: HIGH POTENCY		
AUGMENTED BETAMETHASONE 0.05% C, L (compare to Diprolene® AF) BETAMETHASONE VALERATE 0.1% C, O DESOXIMETASONE 0.25% C, O (compare to Topicort®) FLUOCINONIDE 0.05% C, G, O, TRIAMCINOLONE ACETONIDE 0.5% C, O C=cream, F=foam, G=gel, L=lotion, O=ointment, S=solution	Apexicon E [®] (diflorasone) 0.05% C Desoximetasone 0.05% G Diflorasone diacetate 0.05% C, O (compare to Apexicon E [®]) Halcinonide 0.1% C Halog [®] (halcinonide) all products Topicort [®] (desoximetasone) 0.05% G; 0.25% C, O, Spray All other brands	CRITERIA FOR APPROVAL (NON-PREFERRED AGENTS): The patient has a documented side effect, allergy, or treatment failure to at least two different preferred agents of similar potency. (If a product has an AB rated generic, one trial must be the generic.)
CORTICOSTEROIDS: VERY HIGH POTENCY		
CONTRODIENCIDO, VENT HIGH FOTENCI		
AUGMENTED BETAMETHASONE 0.05% C, L, O (compare to Diprolene®) 0.05% G CLOBETASOL PROPIONATE 0.05%, C, F, G, L, O, S, Shampoo, Spray HALOBETASOL PROPIONATE (compare to Ultravate®) 0.05% C, O C=cream, F=foam, G=gel, L=lotion, O=ointment, S=solution	Bryhali® (halobetasol propionate) L Clobetasol propionate emulsion (compare to Olux E®) 0.05% F Diprolene® (augmented betamethasone) 0.05% L, O Fluocinonide (compare to Vanos®)0.1% C Halobetasol (compare to Lexette TM) 0.05% F Impeklo TM (clobetasol propionate) 0.05% L Lexette TM (halobetasol) 0.05% F Olux®/Olux E® (clobetasol propionate) 0.05% F Tovet® (clobetasol propionate aerosol) 0.05% F Vanos® (fluocinonide) 0.1% C All other brands	CRITERIA FOR APPROVAL (NON-PREFERRED AGENTS): The patient has a documented side effect, allergy, or treatment failure to at least two different preferred agents of similar potency. (If a product has an AB rated generic, one trial must be the generic.)
GENITAL WART THERAPY		
IMIQUIMOD 5 % (compare to Aldara [®]) cream PODOFILOX SOLUTION (compare to Condylox [®])	Aldara® (imiquimod) 5% cream Condylox® Gel (podofilox gel) Imiquimod (compare to Zyclara®) 3.75% Cream QTY Limit: 56 packets/8 weeks Imiquimod (compare to Zyclara®) 3.75% Cream Pump QTY LIMIT: 2 pumps/ 8 weeks Veregan® (sinecatechins ointment) QTY LIMIT: 15 grams (1 tube)/30 days Zyclara® (imiquimod 3.75%) Cream QTY LIMIT: 56 packets/8 weeks Zyclara® (imiquimod 2.5%, 3.75%) Cream Pump QTY LIMIT: 2 pumps/8 weeks	 Aldara cream, Zyclara cream: The patient has had a documented intolerance to generic imiquimod Condylox gel, Veregan: The patient has had a documented side effect, allergy, or treatment failure with imiquimod. Imiquimod pump, Zyclara pump: The patient has had a documented intolerance to generic imiquimod cream and Zyclara cream.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
IMMUNOMODULATORS		
ELIDEL® (pimecrolimus) for ages ≥ 2 TACROLIMUS 0.03% Ointment for ages ≥ 2 TACROLIMUS 0.1% Ointment for ages ≥ 16 Preferred After Clinical Criteria Are Met ADBRY (tralokinumab-ldrm) subcutaneous injection QTY LIMIT: 6 syringes the first 28 days then 4 syringes every 28 days thereafter DUPIXENT® (dupilumab) subcutaneous injection QTY LIMIT: 4 syringes/pens the first 28 days then 2 Syringes/pens every 28 days thereafter Note: please refer to Dermatological Agents: Corticosteroids category for preferred topical corticosteroids.	Cibinqo® (abrocitinib) tablets QTY LIMIT: 1 tab/day Maximum 30 days supply Eucrisa® (crisaborole) Ointment Opzelura® (ruxolitinub) cream Pimecrolimus cream (compare to Elidel®) Rinvoq ® (upadactinib) extended-release tablet QTY LIMIT: 1 tablet/day Maximum 30 days supply	 Eucrisa: The patient has a diagnosis of mild-moderate atopic dermatitis (eczema) AND the patient has had a documented side effect, allergy, or treatment failure (defined as daily treatment for at least one month) with at least one preferred topical calcineurin inhibitor AND the quantity requested does not exceed 60 grams/fill and 180 grams/ 6 months. Trial of calcineurin inhibitor will be waived for patients ≥ 3 months through < 2 years of age. Opzelura: The patient is ≥ 12 years of age AND The patient has a diagnosis of mild-moderate atopic dermatitis (eczema) AND The patient has had a documented side effect, allergy, or treatment failure (defined as daily treatment for at least one month) with at least one moderate to high potency topical corticosteroid within the last 6 months, unless contraindicated AND The patient has had a documented side effect, allergy, or treatment failure (defined as daily treatment for at least one month) of a preferred topical calcineurin inhibitor and crisabarole ointment AND Patient is not receiving Opzelura in combination with another biologic medication (e.g. dupilumab), oral JAK inhibitor (e.g. upadactinib), or systemic immunosuppressant (e.g. cyclosporine) AND The quantity requested does not exceed 60 grams/fill; maximum of 8-weeks of continuous use. Pimecrolimus: The patient has a documente intolerance to brand Elidel. Adbry, Cibingo, Dupixent, Rinvoq: The patient has a diagnosis of moderate to severe atopic dermatitis AND The patient has a diagnosis of moderate to severe atopic dermatitis AND The patient has had a documented side effect, allergy, or treatment failure (defined as daily treatment for at least one month) with at least one moderate to high potency topical corticosteroid and one

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
		allergy, or treatment failure with Adbry or Dupixent AND the patient has a had a documented side effect, allergy, or treatment failure with Rinvoq. Rinvoq additional criteria: The patient has a had a documented side effect, allergy, or treatment failure with Adbry or Dupixent.
SCABICIDES AND PEDICULOCIDES		
PERMETHRIN 5 % (compare to Elimite [®]) C PERMETHRIN 1 % CR, L PIPERONYL BUTOXIDE AND PYRETHRINS G, S, Sh NATROBA [®] (spinosad 0.9 %) Ss C=cream, CR=crème rinse, G=gel, L=lotion, S=solution, Sh=shampoo, Sp=spray, Ss=suspension	Ivermectin 0.5% L Lindane Sh Malathion L (compare to Ovide®) Ovide® (malathion) L Spinosad (compare to Natroba) Ss Vanalice® (piperonyl butoxide/pyrethrins) G	Non-preferred Scabicides: The patient has had a documented side effect or allergy to permethrin cream or treatment failure with two treatments of permethrin cream. Non-Preferred Pediculicides: The patient has had a documented side effect or allergy to OTC permethrin and piperonyl butoxide and pyrethrins and one treatment of Natroba OR treatment failure with two treatments of OTC permethrin and/or piperonyl butoxide and pyrethrins and one treatment of Natroba. For approval of Ovide® Lotion, the patient must also have a documented intolerance to the generic equivalent product.

DESMOPRESSIN: INTRANASAL/ORAL

INTRANASAL All products require PA ORAL DESMOPRESSIN	DDAVP [®] (desmopressin) Nasal Solution or Spray 0.01% Desmopressin Nasal Solution or Spray 0.01 % (compare to DDAVP [®]) Noctiva [™] (desmopressin) Nasal Spray Stimate [®] (desmopressin) Nasal Solution 1.5 mg/ml Nocdurna [®] (desmopressin) SL tablets QTY LIMIT: 1 tablet/day DDAVP [®] (desmopressin) tablets	 CRITERIA FOR APPROVAL: Intranasal (except as indicated below): The diagnosis or indication for the requested medication is (1) Diabetes Insipidus, (2) hemophilia type A, or (3) Von Willebrand disease AND If the request is for brand DDAVP, the patient has a documented intolerance to generic desmopressin spray or solution. Oral: The diagnosis or indication for the requested medication is (1) Diabetes Insipidus and/or (2) primary nocturnal enuresis AND The patient has had a documented intolerance to generic desmopressin tablets Nocdurna, Noctiva: Patient is ≥18 years of age (Nocdurna) or ≥ 50 years of age (Noctiva) AND the indication for use is the treatment of nocturia due to nocturnal polyuria (defined as nighttime urine production exceeding 1/3 of the 24-hour urine production) causing patient to awaken more than 2 times per night to void for at least 6 months AND patient has eGFR > 50ml/min/1.73m2 AND patient does not have increased risk of severe hyponatremia (e.g. concomitant use of loop diuretics or corticosteroids, diagnosis of CHF, or uncontrolled hypertension) AND serum sodium concentrations are normal before starting therapy AND patient has had a documented intolerance to generic desmopressin tablets. LIMITATIONS: Desmopressin intranasal formulations will not be approved for
		the treatment of primary nocturnal enuresis (PNE) due to safety risks of hyponatremia. Oral tablets may be prescribed for this indication.

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
	DIABETIC TESTING SUP	PLIES
Please refer to the DVHA website for covered Diabetic testing supplies. Test strips are subject to a quantity limit of 200 strips per 30 days. https://dvha.vermont.gov/sites/dvha/files/doc_library/Vermont% 20PDSL% 20January% 202023_update d% 201.20.23.pdf		CRITERIA FOR APPROVAL: The prescriber demonstrates that the patient has a medical necessity for clinically significant features that are not available on any of the preferred meters/test strips. CRITERIA FOR APPROVAL to Exceed QTY LIMIT: Chart notes must be provided documenting medical necessity. LIMITATIONS: Talking monitors are not covered under the pharmacy benefit.
	ENDOMETRIOSIS/UTERINE FIBE	ROIDS AGENTS
LUPRON DEPOT® (leuprolide acetate for depot suspension) QTY LIMIT: 3.75 mg kit/month or 11.25 mg kit/3 months SYNAREL® (nafarelin acetate) nasal solution Preferred After Clinical Criteria are Met MYFEMBREE® (relugolix/estradiol/norethindrone) tablet QTY LIMIT: 1 tab/day ORIAHNN® (elagolix and elagolix/estradiol/norethindrone) capsules QTY LIMIT: 2 tabs/day ORILISSA® (elagolix) tablets QTY LIMIT: 200mg dose = 2 tabs/day; maximum of 6 months; 150mg = 1 tab/day	Lupaneta Pack™ (leuprolide acetate for depot suspension and norethindrone acetate tablets) QTY LIMIT: 3.75 mg kit/month or 11.25 mg kit/3 months	Lupaneta Pack: patient has a documented intolerance to Lupron Depot and norethindrone tablets used in combination. Myfembree, Orilissa, Oriahnn: Patient has a documented side effect, allergy, or treatment failure to at least TWO medications from at least 2 different classes (oral contraceptives, NSAIDs, progestins). Note: Use of GnRH receptor antagonists will be limited to 2 years.
EPINEPHRINE: SELF-ADMINISTERED		
EPIPEN-JR INJ 0.15mg EPIPEN INJ 0.3mg EPINEPHRINE INJ (compare to EpiPen-Jr®) (authorized generic, Mylan labeler code 49502 is the only preferred form) 0.15mg	Auvi-Q® Inj 0.1mg Auvi-Q® Inj 0.15mg Auvi-Q® Inj 0.3mg Epinephrine Inj 0.15 mg Epinephrine Inj 0.3 mg	Non-preferred Agents (0.15mg, 0.3mg): The patient must have a documented intolerance to a preferred epinephrine product. Auvi-Q 0.1mg: Patient weight is 7.5kg to 15kg (16.5 to 33 lbs).

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
EPINEPHRINE INJ (compare to EpiPen®) (authorized generic, Mylan labeler code 49502 is the only preferred form) 0.3mg	Symjepi® Inj 0.15mg Symjepi® Inj 0.3mg	
	ESTROGENS: VAGIN	AL
ESTRADIOL ESTRACE VAGINAL® Cream ESTRING® Vaginal Ring VAGIFEM® Vaginal Tablets CONJUGATED ESTROGENS PREMARIN VAGINAL® Cream ESTRADIOL ACETATE FEMRING® Vaginal Ring		
CACTDOINTECTINAL		

GASTROINTESTINAL

INFLAMMATORY BOWEL DISEASE BIOLOGICS: Initial approval is 3 months; renewals are 1 year

<u>Preferred After Clinical Criteria Are Met INJECTABLE</u>

AVSOLA ® (infliximab-axxq) biosimilar to Remicade®

HUMIRA® (adalimumab)

QTY LIMIT: 6 syringes/28 days for the first month (Crohn's starter kit);2 syringes/28 days subsequently

INFLECTRA® (infliximab-dyyb) biosimilar to Remicade®

Cimzia[®] (certolizumab pegol) *QTY LIMIT:* 1 kit/28 days

Entyvio® (vedolizumab)

QTY LIMIT: 300 mg X 3/42 days, 300 mg X 1 every

56 days thereafter Remicade® (infliximab)

RenflexisTM (infliximab-abda) biosimilar to Remicade®

Simponi® (golimumab) SC

QTY LIMIT: 3 of 100 mg prefilled syringe or autoinjector X 1, then 100 mg/28days

Skyrizi® (risankizumab-rzaa)

Clinical Criteria for approval of ALL drugs (Crohn's Disease): Patient has a diagnosis of moderate to severe Crohn's disease and has already been stabilized on the medication OR patient meets additional criteria outlined below:

Avsola, Humira, Inflectra: The patient has had a treatment failure with at least one conventional agent (e.g. methotrexate, corticosteroids) OR there is evidence of severely active disease and early introduction of a biologic without prior medication trials is medically necessary.

Cimzia, Entyvio, Simponi, Stelara, Tysabri: The patient never responded to a 12-week course of anti-TNFα therapy (primary nonresponse) OR the patient previously responded to infliximab (secondary nonresponse) and has a documented side effect, allergy, or treatment failure with adalimumab. Note: Initial IV dose for Stelara will be approved through the medical benefit. All subsequent subcutaneous doses may be approved through the pharmacy benefit

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
	QTY LIMIT: 360 mg (2.4ml)/56 days after initial IV loading dose Stelara® (ustekinumab) QTY LIMIT: 90mg (1 mL)/56 days after initial IV loading dose Tysabri® (natalizumab)	with quantity limit of 90mg every 8 weeks. Remicade, Renflexis: The prescriber must provide a clinically compelling reason why Avsola or Inflectra would not be suitable alternatives. Skyrizi: The patient has a documented side effect, allergy, or treatment failure to a 12-week course of therapy with a preferred TNF inhibitor AND the patient has a documented side effect, allergy, or treatment failure to a 12-week course of therapy with either Entyvio or Stelara.
ORAL XELJANZ® (tofacitinib) tablet QTY LIMIT: 2 tablets/day XELJANZ® XR (tofacitinib) tablet QTY LIMIT: 1 tablet/day	Rinvoq ® (upadactinib) extended-release tablet QTY LIMIT: 1 tablet/day Maximum 30 days supply Zeposia® (ozanimod) capsule QTY LIMIT: 1 capsule/day	Clinical Criteria for approval of ALL drugs (Ulcerative Colitis): Patient has a diagnosis of moderate to severe Ulcerative Colitis and has already been stabilized on the medication OR patient meets additional criteria outlined below: Avsola, Humira, Inflectra: The patient has had a treatment failure with at least one conventional agent (e.g. 5-ASA, corticosteroids) OR there is evidence of severely active disease and early introduction of a biologic without prior medication trials is medically necessary. Entyvio, Simponi, Stelara, Zeposia: The patient has had a treatment failure with at least one conventional agent (e.g. 5-ASA, corticosteroids) AND the patient has a documented side effect, allergy, or treatment failure with at least one preferred biologic. Rinvoq: The patient has had a treatment failure with at least one conventional agent (e.g. 5-ASA, corticosteroids) AND the patient has a documented side effect, allergy, or treatment failure with a preferred TNF inhibitor AND the patient has a documented side effect, allergy, or treatment failure with Xeljanz or Xeljanz XR. Xeljanz, Xeljanz XR: The patient has had a treatment failure with at least one conventional agent (e.g. 5-ASA, corticosteroids) AND the patient has a documented side effect, allergy, or treatment failure with at least one conventional agent (e.g. 5-ASA, corticosteroids) AND the patient has a documented side effect, allergy, or treatment failure with at least one preferred TNF Inhibitor. Note: Induction of Xeljanz 10mg twice daily or XR 22mg once daily will be limited to 16 weeks. Treatment should be discontinued after 16 weeks if adequate therapeutic response is not achieved. For patients with loss of response during maintenance treatment with 5mg twice daily or XR 11mg once daily, approval of 10mg twice daily or XR 22mg once daily will be considered and limited to the shortest duration possible.
H. PYLORI COMBINATION THERAPY		
LANSOPRAZOLE, AMOXICILLIN, CLARITHROMYCIN QTY LIMIT: 112 caps & tabs/14 days PYLERA® (bismuth subcitrate, metronidazole, tetracycline) capsules QTY LIMIT: 120 caps/10 days	Omeclamox-Pak® (omeprazole, clarithromycin, amoxicillin) QTY LIMIT: 80 caps & tabs/10 days Talicia® (omeprazole, amoxicillin, rifabutin) delayed release capsules QTY LIMIT: 168 caps/14 days	CRITERIA FOR APPROVAL: The patient has a documented treatment failure with Lansoprazole, amoxicillin, clarithromycin combo package or Pylera used in combination with a PPI.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
H-2 BLOCKERS		
FAMOTIDINE (compare to Pepcid [®]) tablet	Cimetidine (compare to Tagamet®) tablet Nizatidine capsule Pepcid® (famotidine) tablet	Cimetidine tablet, Nizatidine capsule, Pepcid tablet: The patient has had a documented side effect, allergy, or treatment failure to famotidine.
SYRUPS AND SPECIAL DOSAGE FORMS FAMOTIDINE oral suspension (compare to Pepcid®) age ≤ 12 years	Cimetidine oral solution Famotidine (compare to Pepcid [®]) oral suspension (age >12 years) Nizatidine Oral Solution	Cimetidine Oral Solution, Nizatidine oral solution: Patient has a medical necessity for a liquid dosage form AND the patient has had a documented side effect, allergy, or treatment failure to famotidine oral suspension. Famotidine Oral Suspension (Age >12): Patient has a medical necessity for a liquid dosage form
INFLAMMATORY BOWEL AGENTS (ORAL &	RECTAL PRODUCTS)	
MESALAMINE PRODUCTS ORAL APRISO® (mesalamine capsule extended release) LIALDA® (mesalamine tablet extended release) PENTASA ER® (mesalamine cap CR)	Delzicol® (mesalamine capsule delayed-release) QTY LIMIT: 6 capsules/day Mesalamine capsule delayed release (compare to Delzicol®) QTY LIMIT: 6 capsules/day Mesalamine capsule extended release 0.375gm (compare to Apriso®) Mesalamine tablet delayed release (compare to Asacol® HD) Mesalamine tablet extended release 1.2 g (compare to Lialda®)	 Azulfidine, Colazal: patient has had a documented intolerance to the generic equivalent of the requested medication. Budesonide ER 9mg, Ortikos: the patient has a documented intolerance to brandname Uceris. Delzicol, Mesalamine capsule DR, Mesalamine tablet DR, Mesalamine tablet ER: The patient has had a documented side effect, allergy, or treatment failure to 2 preferred oral mesalamine products. sfRowasa, Uceris Rectal Foam: The patient has had a documented intolerance to mesalamine enema or suppositories. LIMITATIONS: Kits with non-drug products are not covered.
RECTAL MESALAMINE ENEMA (compare to Rowasa®) MESALAMINE SUPPOSITORY	sfRowasa [®] (mesalamine enema sulfite free)	
CORTICOSTEROIDS ORAL BUDESONIDE 24HR QTY LIMIT: 3 capsules/day UCERIS® (budesonide) ER Tablet QTY LIMIT = 1 tablet/day	Budesonide ER 9 mg tablet (compare to Uceris®) QTY LIMIT: 1 tablet/day Ortikos® (budesonide) ER capsule QTY LIMIT: 1 capsule/day	

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
(110 171 required unless other wise noted)	(i i required)	TH CRITERIA
RECTAL All products require PA	Uceris® Rectal Foam (budesonide)	
OTHER BALSALAZIDE (compare to Colazal [®]) DIPENTUM [®] (olsalazine) SULFAZINE SULFAZINE EC SULFASALAZINE (compare to Azulfidine [®]) SULFASALAZINE DR	Azulfidine [®] (sulfasalazine) Colazal [®] (balsalazide)	
PROKINETIC AGENTS		
TABLETS METOCLOPRAMIDE tabs (compare to Reglan®) ORAL SOLUTION METOCLOPRAMIDE oral solution	Reglan [®] (metoclopramide)	Reglan: The patient has had a documented intolerance to generic metoclopramide tablets. Gimoti: The patient has a documented intolerance to metoclopramide tablets and oral solution.
NASAL SPRAY All products require PA	Gimoti [™] (metoclopramide) nasal spray	
PROTON PUMP INHIBITORS		
ORAL CAPSULES/TABLETS ESOMEPRAZOLE (compare to Nexium®) QTY LIMIT: 1 cap/day LANSOPRAZOLE generic RX capsules (compare to Prevacid®) QTY LIMIT: 1 cap/day OMEPRAZOLE RX capsules (compare to Prilosec®) QTY LIMIT: 1 cap/day PANTOPRAZOLE tablets (compare to Protonix®) QTY LIMIT: 1 tab/day	Aciphex [®] (rabeprazole) tablets QTY LIMIT: 1 tab/day Dexilant [®] (dexlansoprazole) capsules QTY LIMIT: 1 cap/day Nexium [®] (esomeprazole) capsules QTY LIMIT: 1 cap/day Omeprazole generic OTC tablets QTY LIMIT: 1 tab/day Omeprazole magnesium generic OTC 20 mg capsules QTY LIMIT: 1 cap/day Omeprazole/sodium bicarb capsules RX (compare to Zegerid [®]) QTY LIMIT: 1 cap/day Prevacid [®] RX (lansoprazole) capsules QTY LIMIT: 1 cap/day Prevacid [®] 24 hr OTC (lansoprazole) capsules QTY LIMIT: 1 cap/day	Nexium powder for suspension (for patients ≥ 12 years old): The patient has a requirement for a non-solid oral dosage form (e.g. an oral liquid, dissolving tablet or sprinkle). Aciphex Sprinkle, Prevacid Solutabs, Prilosec packet, and Protonix packet: The patient has a requirement for a non-solid oral dosage form (e.g. an oral liquid, dissolving tablet or sprinkle). AND the member has had a documented side effect, allergy or treatment failure to Nexium powder for suspension. Other non-preferred medications: The member has had a documented side effect, allergy, or treatment failure to ALL preferred PPIs AND if the product has an AB rated generic, there must be a trial of the generic. CRITERIA FOR APPROVAL (twice daily dosing): Gastroesophageal Reflux Disease (GERD) – If member has had an adequate trial (e.g. 8 weeks) of standard once daily dosing for GERD, twice daily dosing may be approved. Note: Approval of twice daily dosing for GERD is limited to 12 weeks. For continuation after 12 weeks, there must be a documented attempt to taper to once daily dosing of a PPI with an adjunctive H2 Blocker. The dosing of long-term PPI's should be periodically reevaluated so that the lowest effective dose can be prescribed to manage the

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
SUSPENSION & SPECIAL DOSAGE FORMS NEXIUM [®] (esomeprazole) powder for suspension	Protonix [®] (pantoprazole) tablets <i>QTY LIMIT</i> : 1 tab/day Rabeprazole (compare to Aciphex [®]) tablets <i>QTY LIMIT</i> : 1 tab/day Zegerid RX [®] (omeprazole/sodium bicarb) caps, oral, suspension <i>QTY LIMIT</i> : 1 cap/day	condition. Zollinger-Ellison (ZE) syndrome – Up to triple dose PPI may be approved. Hypersecretory conditions (endocrine adenomas or systemic mastocytosis) – Double dose PPI may be approved. Erosive Esophagitis, Esophageal stricture, Barrett's esophagitis (complicated GERD) – Double dose PPI may be approved. Treatment of ulcers caused by H. Pylori – Double dose PPI may be approved for up to 2 weeks. Laryngopharyngeal reflux – Double dose PPI may be approved. LIMITATIONS: First-Lansoprazole® and First-Omeprazole Suspension Kits are not covered as Federal Rebate is no longer offered.
(age < 12 years) QTY LIMIT: 1 packet/day	Aciphex [®] Sprinkle (rabeprazole) DR Capsule <i>QTY LIMIT</i> : 1 cap/day Lansoprazole ODT (compare to Prevacid Solutab®) <i>QTY LIMIT</i> : 1 tab/day Nexium [®] (esomeprazole) powder for suspension (age ≥ 12 years) <i>QTY LIMIT</i> : 1 packet/day Prevacid Solutabs [®] (lansoprazole) <i>QTY LIMIT</i> : 1 tab/day	
	Prilosec [®] (omeprazole magnesium) packet QTY LIMIT: 2 packets/day Protonix [®] (pantoprazole) packet QTY LIMIT: 1 packet/day GAUCHER'S DISEASE MEDIC	ATIONS
All products require PA	Cerezyme® (imiglucerase for injection) Cerdelga® (eliglustat)	 CRITERIA FOR APPROVAL: The diagnosis or indication is Gaucher disease (GD) type I. AND The diagnosis has been confirmed by molecular or enzymatic testing. Age Limits Elelyso, Vpriv: for patients ≥ 4 years old Cerezyme: for patients ≥ 2 years old
	Miglustat (compare to Zavesca®) QTY LIMIT: 3 caps/day Zavesca® (miglustat) OTY LIMIT: 3 caps/day	Cerdelga, Miglustat, Zavesca: for patients ≥ 18 years old Cerezyme/Vpriv additional criteria: Failure, intolerance or other
	QTY LIMIT:3 caps/day	contraindication to enzyme replacement therapy with Elelyso
	Maximum days supply per fill for all drugs is 14 days	Cerdelga additional criteria:

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA • Testing to verify if CYP2D6 extensive metabolizer (EM), intermediate metabolizer (IM), poor metabolizer (PM), or if CYP2D6 genotype cannot be determined ○ Dose max: 84mg twice/day if EM or IM ○ Dose max: 84mg/day if PM ○ Case by case determination if CYP2D6 cannot be determined Miglustat, Zavesca additional criteria: • For whom enzyme replacement therapy is not a therapeutic option (e.g. due to allergy, hypersensitivity, or poor venous access) AND for
	COUT ACENTS	approval of miglustat, the patient must have a documented intolerance to brand Zavesca.
	GOUT AGENTS	
ALLOPURINOL (compare to Zyloprim [®]) COLCHICINE tablets (compare to Colcrys®) COLCHICINE/PROBENECID PROBENECID	Colcrys [®] (colchicine) tablet QTY LIMIT: 3 tablets/day (gout) or 4 tablets/day (FMF) Colchicine capsules Febuxostat (compare to Uloric®) QTY LIMIT: 40 mg tablets = 1 tablet/day Mitigare® (colchicine) capsule QTY LIMIT: 2 capsules/day Uloric® (febuxostat) QTY LIMIT: 40 mg tablets = 1 tablet/day Zyloprim® (allopurinol)	 Colchicine capsules, Colcrys, Mitgare: the patient has a documented intolerance to generic colchicine tablets. Febuxostat, Uloric: The diagnosis or indication is treatment of gout AND The patient has had a documented side effect, allergy, treatment failure or a contraindication to allopurinol. NOTE: Treatment failure is defined as inability to reduce serum uric acid levels to < 6 mg/dl with allopurinol doses of 600 mg/day taken consistently. Additionally, renal impairment is not considered a contraindication to allopurinol use. Zyloprim: The patient has had a documented intolerance to generic allopurinol
	GROWTH STIMULATING A	AGENTS
ACHONDROPLASIA TREATMENTS All products require PA	Voxzogo TM (vosoritide)	Voxzogo: The patient must have a diagnosis of achondroplasia confirmed with
		genetic testing AND the medication must be prescribed by a pediatric endocrinologist AND Confirmation of non-closure of epiphyseal plates (x-ray determining bone age) must be provided for females > age 12 and males > age 14 AND Voxzogo will not be used in combination with growth hormone (e.g. somatropin), growth hormone analogs (e.g. somapacitan), or insulin-like growth factor (IGF-1) (e.g. mecasermin) AND patient's standing height, weight, BMI, and upper to lower body ratio will be measured at baseline and monitored throughout therapy. For re-approval, the patient must have an improvement in growth velocity compared to pre-treatment baseline.

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
Preferred After Clinical Criteria Are Met GENOTROPIN® NORDITROPIN®	Nutropin® AQ Omnitrope® Saizen® Skytrofa® (lonapegsomatropin-tcgd) Zomacton® Specialized Indications – See Specific Criteria Increlex® (mecasermin) Serostim® Zorbtive®	Criteria for Approval Pediatric: 1) The patient must have one of the following indications for growth hormone: □ Turner syndrome confirmed by genetic testing. □ Prader-Willi Syndrome confirmed by genetic testing. □ Growth deficiency due to chronic renal failure. □ Patient who is Small for Gestational Age (SGA) due to Intrauterine Growth Retardation (IUGR)and catch up growth not achieved by age 2 (Birth weight less than 2500g at gestational age of <37 weeks or a birth weight or length below the 3rd percentile for gestational age). OR □ Pediatric Growth Hormone Deficiency confirmed by results of two provocative growth hormone stimulation tests (insulin, arginine, levodopa, propranolol, clonidine, or glucagon) showing results (peak level) <10ng/ml. 2) The requested medication must be prescribed by a pediatric endocrinologist (or pediatric nephrologist if prescribed for growth deficiency due to chronic renal failure). 3) Confirmation of non-closure of epiphyseal plates (x-ray determining bone age) must be provided for females > age 12 and males > age 14. 4) Initial requests can be approved for 6 months. Subsequent requests can be approved for up to 1 year with documentation of positive response to treatment with growth hormone. Criteria for Approval Adult: The patient must have one of the following indications for growth hormone: Panhypopituitarism due to surgical or radiological eradication of the pituitary. OR Adult Growth Hormone Deficiency confirmed by one growth hormone stimulation test (insulin, arginine, levodopa, propranolol, clonidine, or glucagon) showing results (peak level) <5ng/ml. Growth hormone deficient children must be retested after completion of growth. LIMITATIONS: Coverage of Growth Hormone products will not be approved for patients who have Idiopathic Short Stature. NUTROPIN AQ, OMNITROPE, SAIZEN, SKYTROFA, ZOMACTON: The patient has a documented side effect, allergy, or treatment failure to both preferred agents. Increlex: Member has growth hormone, OR primary insulin-like growth factor

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PREFERRED AGENTS	NON-PREFERRED AGENTS	DA CDITEDIA
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
All products require PA	hattr treatment Amvuttra TM (vutrisiran) 25mg/0.5ml injection for subcutaneous use <i>QTY LIMIT</i> : 1 syringe (0.5ml) every 3 months Onpattro® (patisiran) 10 mg/5ml intravenous injection Weight < 100kg (0.3 mg/kg every 3 weeks) Weight ≥ 100kg (30 mg every 3 weeks) Tegsedi® (inotersen) 284 mg/1.5ml injection for subcutaneous use <i>QTY LIMIT</i> : 4 syringes/28 days Vyndamax® (tafamidis) <i>QTY LIMIT</i> : 1 capsule/day Vyndaqel® (tafamidis meglumine) <i>QTY LIMIT</i> : 4 capsules/day	Amvuttra, Onpattro, Tegsedi: • The patient is ≥ 18 years of age with a diagnosis of polyneuropathy of heredity transthyretin mediated (hATTR) amyloidosis (Documentation of TTR mutation by genetic testing or the presence of amyloid deposits via tissue biopsy has been submitted) AND • The medication is being prescribed by or in consultation with a neurologist AND • Clinical signs and symptoms of the disease (e.g., peripheral/autonomic neuropathy, motor disability, cardiovascular dysfunction, renal dysfunction) are present and other causes of neuropathy have been excluded AND • Patient is receiving vitamin A supplementation AND • Initial approval will be granted for 3 months. For re-approval, the patient must have documentation of clinical improvement or slower progression of the disease than would otherwise be expected. Vyndamax, Vyndaqel: • The patient is ≥ 18 years of age with a diagnosis of cardiomyopathy of wild type transthyretin-mediated amyloidosis or heredity transthyretin mediated (hATTR) amyloidosis AND • The presence of amyloid deposits showing cardiac involvement via tissue biopsy or imaging has been submitted AND • The medication is being prescribed by or in consultation with a cardiologist AND • Initial approval will be granted for 6 months. For re-approval, the patient must have a decrease in the frequency of cardiovascular-related hospitalizations or slower progression of the disease than would otherwise be expected.
	HEART FAILURE	
ANGIOTENSIN RECEPTOR – NEPRILYSIN INH	IIBITOR (ARNI)	
ENTRESTO® (valsartan/sacubitril)		
QTY LIMIT: 2 tablets/day		
CARDIAC MYOSIN INHIBITORS		
All procuts require PA	Camzyos® (mavacamten)	Camzyos:

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
(NOTA required unless officerwise noted)	QTY LIMIT: 1 capsule/day	 The diagnosis or indication is symptomatic New York Heart Association (NYHA) class II-III obstructive hypertrophic cardiomyopathy (HCM) AND LVEF ≥ 55% AND Valsalva LVOT peak gradient ≥50mmHg at rest or with provocation AND The patient has a documented side effect, allergy, or treatment failure at a maximally tolerated dose to at least two of the following: Non-vasodilating beta blocker (e.g., atenolol, bisoprolol, metoprolol, nadolol, propranolol), Nondihydropyridine calcium channel blocker (i.e., diltiazem, verapamil), and Disopyramide AND The medication will not be used concurrently with disopyramide, ranolazine, verapamil with a beta blocker, or diltiazem with a beta blocker. Approval will be granted for 12 months. For reapproval, there must be a documented positive clinical response as supported by one of the following: Stable or reduction in New York Heart Association (NYHA) class AND Patient has a left ventricular ejection fraction of greater than or equal to 50%
		•
SODIUM-GLUCOSE CO-TRANSORTER 2 (SGI	LT2) INHIBITORS	
FARXIGA [®] (dapagliflozin) <i>QTY LIMIT:</i> 1 tab/day		
SOLUBLE GUANYLATE CYCLASE (sGC) STIN	MULATORS	
All products require PA	Verquvo® (vericiguat) tablet <i>QTY LIMIT</i> : 1 tablet/day	 Verquvo: The diagnosis or indication is symptomatic heart failure (HF) with ejection fraction < 45% AND the patient has been hospitalized for HF within the previous 6 months or required the use of IV diuretics within the past 3 months AND the patient is not pregnant AND the patient is concurrently receiving the maximum tolerated dose of one agent from each of the following classes, unless contraindicated: ARNI, ACE-I, or ARB Beta Blocker (metoprolol, carvedilol, or bisoprolol) Aldosterone antagonist if LVEF ≤ 35% or LVEF ≤ 40% with diabetes mellitus or post myocardial infarction (MI) with HF symptoms
	HEMATOPOIETICS	
Colony Stimulating Factors		
Eflapegrastim Products	Rolvedon TM (eflapegrastim-xnst) Syringe	Granix, Leukine, Nivestym, Releuko, Zarxio: The prescriber must provide a

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
All products require PA Filgrastim Products NEUPOGEN® (filgrastim) Vial, Syringe Pegfilgrastim Products FULPHILA™ (pegfilgrastim-jmdb) Syringe NEULASTA® (pegfilgrastim) Syringe NEULASTA® Onpro® (pegfilgrastim) kit ZIEXTENZO® (pegfilgrastim-bmez)	Granix® (tbo-filgrastim) Vial, Syringe Leukine® (sargramostim) Nivestym™ (figrastim-aafi) Vial, Syringe Releuko™ (filgrastim-ayow) Zarxio® (filgrastim-sndz) Syringe Fylnetra® (pegfilgrastim-pbbk) Nyvepria (pegfilgrastim-apgf) Stimufend® (pegfilgrastim-fpgk) Udenyca™ (pegfilgrastim-cbqv)	clinically compelling reason for the use of the requested medication including reasons Neupogen would not be a suitable alternative. Fylnetra, Nyvepria, Rolvedon, Stimufend, Udenyca: The prescriber must provide a clinically compelling reason for the use of the requested medication including reasons why any of the preferred pegfilgrastim products would not be suitable alternatives.
Erythropoietic Stimulating Agents		
Preferred After Clinical Criteria Are Met EPOGEN® (epoetin alpha) MIRCERA® (methoxypolyethylene glycolepoetin beta)	Aranesp® (darbepoetin alfa) Procrit® (epoetin alpha) Retacrit® (epoetin alpha-epbx)	Aranesp, Procrit, Epogen, Retacrit: diagnosis or indication for the requested medication is anemia due to one of the following: Chronic kidney disease/renal failure, Post-renal transplant, use of zidovudine for the treatment of human immunodeficiency virus (HIV) (other causes of anemia, such as iron/folate/vitamin B12 deficiency have been eliminated), Surgery patients at high risk for perioperative blood loss, Cancer chemotherapy, Use of ribavirin or interferon therapy for Hepatitis C, Myelodysplastic syndrome. Hemoglobin level at initiation of therapy is <10 g/dL OR for patients currently maintained on therapy, hemoglobin level is < 11 g/dL in dialysis patients with chronic kidney disease, < 10 g/dL in non-dialysis patients with chronic kidney disease, or < 12 g/dL in patients treated for other indications AND for approval of Aranesp or Procrit, or Retacrit the patient has had a documented side effect, allergy, or treatment failure to Epogen. Mircera: The diagnosis or indication for the requested medication is anemia due to chronic kidney disease/renal failure AND Hemoglobin level at initiation of therapy is <10g/dl OR For patients currently maintained on therapy, hemoglobin level is ≤11 g/dL in dialysis patients with chronic kidney disease, ≤10 g/dL in non-dialysis patients with chronic kidney disease, ≤10 g/dL in non-dialysis patients with chronic kidney disease.
HEMOPHILIA FACTORS		
AHF-Factor VII		
All products require PA	Novoseven® RT	Novoseven RT: Medication is being used for the treatment of acute bleeding

PREFERRED AGENTS (No PA required unless otherwise noted) NON-PREFERRED AGENTS (PA required) Sevenfact® PA CRITERIA episodes in a patient with Hemophilia A or B with inhibitors congenital Factor VII deficiency. Sevenfact: Medication is being used for the treatment of acute I patient with Hemophilia A or B with inhibitors AND there i	
congenital Factor VII deficiency. Sevenfact: Medication is being used for the treatment of acute l	
congenital Factor VII deficiency. Sevenfact: Medication is being used for the treatment of acute l	
Sevenfact: Medication is being used for the treatment of acute l	s OR Patient has
natient with Hemonhilia A or B with inhibitors AND there i	
compelling reason why Novoseven RT cannot be used.	s a clinically
AHF-Factor VIII	
ADVATE® Adynovate® All Non-Preferred Products: The prescriber must provide a	clinically
AFSTYLA® compelling reason for the use of the requested medication	
HEMLIBRA® (emicizumab-kxwh) Esperoct® why any of the preferred products would not be suitable a	
HEMOFIL® M approval of Adynovate, Eloctate, or Esperoct, documentation	
JIVI® the member is unable to use the preferred extended half-life	concentrate Jivi.
KOATE®-DVI NOVOEIGHT®	
OBIZUR®	
RECOMBINATE®	
XYNTHA®	
AHF-Factor IX	
ALPHANINE® SD Kcentra® All Non-Preferred Products: The prescriber must provide a	clinically
ALPROLIX® Rebinyn® compelling reason for the use of the requested medication	-
BENEFIX® why any of the preferred products would not be suitable a	
IDELVION® approval of Rebinyn, documentation must include why the	
IXINITY® to use a preferred extended half-life concentrate Alprolix	or Idelvion.
PROFILNINE®	
RIXUBIS®	
AHF-Von Willebrand Factor	
All Non-Preferred Products: The prescriber must provide a	clinically
ALPHANATE® Compelling reason for the use of the requested medication	
HUMATE-P® why any of the preferred products would not be suitable a	lternatives.
WILATE®	
AHF-Anti-Inhibitor Coagulation Complex	
All products require PA	
Feiba® Feiba: medication is being used for the treatment of acute bleed	ling episodes or

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA routine prophylaxis in a patient with Hemophilia A or B with inhibitors.
		Toutine prophytaxis in a patient with Hemophina A of B with inhibitors.
	HEPATITIS B AGEN	TS
ENTECAVIR (compare to Baraclude®) VIREAD® (tenofovir disoproxil fumarate) tablet	Adefovir (compare to Hepsera®) Baraclude® (entecavir) tablet, solution Lamivudine HBV (compare to Epivir-HBV®) Vemlidy® (tenofovir alafenamide fumarate) Viread® (tenofovir disoproxil fumarate) powder	Adefovir, Lamivudine HBV, Epivir-HBV: The prescriber must provide a clinically compelling reason for the use of the requested medication including reasons why any of the preferred products would not be suitable alternatives Note: AASLD and WHO guidelines recommend these not be utilized first line due to potential for the development of resistance. Baraclude tabs: the patient has a documented intolerance to generic entecavir. Baraclude suspension, Viread Powder: the patient has a medical necessity for a non-solid oral dosage form. Vemlidy: the patient must have a diagnosis of osteoporosis, renal insufficiency (CrCl < 60ml/min), or other contraindication to Viread such as chronic steroid use.
HEPATITIS C AGENTS		
Initial PA: 3 months; subsequent maximun	n 3 months	
RIBAVIRIN 200 mg tablets	Ribavirin 200 mg capsules	Non-preferred Ribavirin Brands/strengths: The patient is unable to use generic ribavirin 200 mg tablets
PEGINTERFERON PRODUCTS PEGASYS® (peginterferon alfa-2a) QTY LIMIT: 4 vials or syringes/28 days		
DIRECT ACTING ANTIVIRALS Preferred After Clinical Criteria Are Met MAVYRET™ (glecaprevir/pibrentasvir) SOFOSBUVIR/VELPATASVIR (compare to Epclusa®)	Epclusa® (sofosbuvir/velpatasvir) Harvoni® (ledipasvir/sofosbuvir) Ledipasvir/sofosbuvir (compare to Harvoni®) Sovaldi® (sofosbuvir) Vosevi® (sofosbuvir/velpatasvir/voxilaprevir) Zepatier® (elbasvir/grazoprevir)	 Direct Acting Agents: Epclusa, Harvoni, Ledipasvir/sofosbuvir, Mavyret, Sofosbuvir/velpatasvir, Sovaldi, Vosevi, Zepatier: Hep C PA form must be completed, and clinical documentation supplied Combination therapy will be either approved or denied in its entirety. Prescriber is, or has consulted with, a hepatologist, gastroenterologist or infectious disease specialist. Consult must be within the past year with documentation of recommended regimen. Specialist requirement will NOT apply for patients meeting all the following: treatment naïve, non-cirrhotic, HBV negative, HIV negative, no prior liver transplantation, an not pregnant. See PA form for detailed requirements and for documentation required For approval of a non-preferred agent, the provider must submit clinical

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA documentation detailing why the patient is not a candidate for a preferred direct acting agent regimen.
	HEREDITARY ANGIOEDEMA M	MEDICATIONS
TREATMENT		
Preferred After Clinical Criteria are Met BERINERT® (human C1 inhibitor) ICATIBANT (compare to Firazyr®) QTY LIMIT: 3 syringes (9 ml)/fill	Firazyr® (icatibant) QTY LIMIT: 3 syringes (9 ml)/fill Kalbitor® (escallantide) QTY LIMIT: 6 vials (2 packs) per fill Ruconest® (recombinant C1 esterase inhibitor) QTY LIMIT: 4 vials/fill	Berinert, Firazyr, Icatibant: The diagnosis or indication is treatment of an acute Hereditary Angioedema (HAE) attack AND for approval of Firazyr, the patient must have a documented intolerance to generic Icatibant. (Approval may be granted so that 2 doses may be kept on hand for Berinert and 3 doses for Icatibant/Firazyr). Kalbitor, Ruconest: The diagnosis or indication is treatment of an acute Hereditary Angioedema (HAE) attack AND the patient has a documented side effect, allergy, treatment failure or contraindication to a preferred agent (Approval may be granted so that 2 doses may be kept on hand.)
PROPHYLACTIC		
Preferred After Clinical Criteria are Met CINRYZE® (human C1 inhibitor) QTY LIMIT: 20 vials/30days HAEGARDA® (human C1 inhibitor) ORLADEYO™ (berotralstat) QTY LIMIT: 1 capsule/day TAKHZYRO™ (lanadelumab-flyo) QTY LIMIT: 2 vials/28 days		Cinryze, Haegarda, Orladeyo, Takhzyro: The diagnosis or indication is prophylaxis of Hereditary Angioedema (HAE) attacks.

HIDRADENITIS SUPPURATIVA

BIOLOGICS: Initial approval is 3 months; renewals are 1 year

Preferred After Clinical Criteria Are Met INJECTABLE

HUMIRA® (adalimumab) QTY LIMIT: 6 syringes/28 days for the first month

(HS starter kit);4 syringes/28 days subsequently

- The patient has a diagnosis of moderate-severe hidradenitis suppurative (Hurley Stage II-III) AND
- The medication is being prescribed by, or in consultation with, a dermatologist AND
- The patient has not responded to a 12-week course of standard antibiotic therapy with an oral tetracycline (e.g. Doxycycline) or clindamycin plus

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA rifampin, unless contraindicated.
	HYPERKALEMIA AGE	NTS
Lokelma TM (sodium zirconium cyclosilicate) SPS® (sodium polystyrene sulfonate) suspension	Veltassa® (patiromer sorbitex calcium) powder packets <i>QTY LIMIT</i> : 1 packet/day	Veltassa: The patient requires therapy for the treatment of non-emergent hyperkalemia AND where clinically appropriate, medications known to cause hyperkalemia (e.g. ACE inhibitors, ARBs, aldosterone antagonists, NSAIDs) have been discontinued or reduced to the lowest effective dose AND where clinically appropriate, a loop or thiazide diuretic has failed for potassium removal, AND the patient has been counseled to follow a low potassium diet (\leq 3 grams/day).
	IDIOPATHIC PULMONARY FIE	BROSIS (IPF)
All products require PA	Esbriet® (pirfenidone) QTY LIMIT: 267 mg tablets = 270 tabs/month, 801 mg tablets = 90 tabs/month Ofev® (nintedanib) QTY LIMIT: 60 tabs/month	 Clinical Criteria: Esbriet, Ofev Age ≥ 18 Diagnosis of idiopathic pulmonary fibrosis (Esbriet and Ofev) OR chronic fibrosing interstitial lung disease or systemic sclerosis associated interstitial lung disease (Ofev Only) May not be used in combination The prescriber is a pulmonologist. Clinical documentation that the member is a non-smoker or has not smoked in 6 weeks. FVC≥ 50% of predicted Reauthorization Criteria: Documentation the patient is receiving clinical benefit to Esbrit® or Ofev® therapy as evidenced by < 10% decline in percent predicted FVC or < 200mL decrease in FVC AND There is clinical documentation that the member has remained tobaccofree.
	IMMUNOLOGIC THERAPIES FO	OR ASTHMA
Initial 3 months, Renewal 1 year Preferred After Clinical Criteria are Met DUPIXENT® (dupilumab) subcutaneous injection, pre-filled syringe, and auto-injector pen QTY LIMIT: 4 syringes/pens the first	Cinqair® (reslizumab) Intravenous injection Nucala® (mepolizumab) subcutaneous injection, vial, pre-filled syringe, and auto-injector pen	 Xolair: Diagnosis of moderate to severe persistent asthma: The patient must be 6 years of age or older AND

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
28 days then 2 syringes/pens every 28 days thereafter FASENRA® (benralizumab) subcutaneous Injection, pre-filled syringe and auto-injector pen QTY LIMIT: 1 mL every 28 days for 3 doses then 1 mL every 56 days XOLAIR® (omalizumab) subcutaneous injection vial, prefilled syringe QTY LIMIT: 900 mg every 28 days	QTY LIMIT: 1mL every 28 days Tezspire™ (tezepelumab-ekko) subcutaneous injection, pre-filled syringe QTY LIMIT: 1.91 mL every 28 days	 • The patient has a history of uncontrolled asthma symptoms (symptoms occurring almost daily or waking at night with asthma at least once a week) or 2 or more exacerbations in the previous year despite regular use of medium-high dose ICS/LABA for a minimum of 3 consecutive months, with or without oral corticosteroids. Pharmacy claims will be evaluated to assess compliance with therapy. AND • The prescriber is a pulmonologist, allergist, or immunologist AND • Patient has tested positive to at least one perennial aeroallergen by skin or blood test (i.e.: RAST, CAP, intracutaneous test) AND • Patient has an IgE level ≥ 30 and ≤ 700 IU/ml (ages 12 and older) OR IgE level ≥ 30 and ≤ 1300 IU/ml (ages 6-11) prior to beginning therapy with Xolair. • For continuation of therapy after the initial 6-month authorization, the patient must continue to receive therapy with an ICS/LABA AND have either a decreased frequency of exacerbations, decreased use of maintenance oral corticosteroids, reduction in the signs and symptoms of asthma, or an increase in predicted FEV1 from baseline. Diagnosis of chronic idiopathic urticaria: • The patient must be 12 years of age or older AND • The patient must be 12 years of age or older AND • The patient has a therapeutic failure or contraindication to an H1 antihistamine (e.g. cetirizine, fexofenadine) at double the daily dose • For continuation of therapy after the initial 6-month authorization, the patient must have documented clinical improvement in symptoms. Diagnosis of Chronic Rhinosinusitis with Nasal Polyps: • Patient is 18 years of age or older AND • Patient has had an inadequate response to at least a 3-month trial of 2 different nasal corticosteroids AND • Patient has had an inadequate response to at least a 10-14 day course of oral corticosteroid AND there must be documented improvement in nasal symptoms. Limitations: Xolair use will not be appr

(No PA required unless otherwise noted) PA CRITERIA * The patient must have a diagnosis of severe persistent asthma with an existing probability of the patient probability of patie	PREFERRED AGENTS	NON-PREFERRED AGENTS	
 The patient most have a diagnosis of severe persistent asthma with an eosinophilic phenotype as defined by pre-treatment blood eosinophil count of ≥ 150 cells per mcL. within the previous 6 weeks or ≥ 300 cells per mcL within the previous 6 weeks or ≥ 300 cells per mcL within 12 months prior to initiation of therapy AND The patient has a history of uncontrolled asthma symptoms (symptoms occurried annotation) or wasting at right with asthma at least once a week) or 2 or more exacerbations in the previous year despite regular use of medium-high dose CESALABA for a minimum of 2 consecutive months, with or without oral corticosteroids. Pharmacy claims will be evaluated to assess compliance with therapy. AND The prescriber is an altergist, immunologist, or pulmonologist, AND For approval of Cinapir or Nucala, the patient must have a documented side effect, allergy, or treatment failure with Dujskent or Fasenn. For continuation of therapy after the initial 6-month authorization, the patient must accommand the patient association of the patient and accurate of maintenance or all corticosteroids, reduction in the signs and symptoms of asthma, or an increase in predicted FFV, from baseline. Diagnosis of hyperecoinophile: yandrome (Nucada only): Patient must be 12 years of age or older AND The patient must have a blood eosinophile count of ≥ 1,000 cells per mcl. AND The patient must have a blood eosinophile count of ≥ 1,000 cells per mcl. AND The patient is on a subtle dose of background HFS therapy (chronic or episodic corticosteroids, immunosuppressive, or cytoxic therapy) for at least 4 weeks prior to to entered initiation AND The patient is an allergist or Kryl specialist AND Presente Khinosimositis with Nasal Pohyps (Nucala Only): Pratient has had an inadequate response to at least a 10-14-day course of oral corticosteroids AND Patient has had an inadequate response to at least a 10-14-day c			PA CRITERIA
eosinophilic phenotype as defined by gre-treatment blood eosinophilic count of ≥ 150 cells per mcl. within the previous 6 weeks or ≥ 300 cells per mcl. within the previous 6 weeks or ≥ 300 cells per mcl. within the previous 6 weeks or ≥ 300 cells per mcl. within the previous 6 weeks or ≥ 300 cells per mcl. within 12 months prior to initiation of therapy AND • The patient has a history of uncontrolled asthma symptoms occurring almost daily or waking at night with asthma at least once a week) or ≥ 0 more exacerbations in the previous year despite regular use of medium-high dose ICSL/ABA for a minimum of ≥ consecutive months, with or without oral corticosteroids. Pharmacy claims will be evaluated to assess compliance with therapy. AND • The prescriber is an alkergist, immunologist, or pulmonologist, AND • For approval of Cinqair or Nucala, the patient must have a documented side effect, allergy, or treatment failure with Dupixent or Fassana. • For continuation of therapy after the initial 6 month authorization, the patient must continue to reserve therapy with an ICSL-ABA AND have either a decreased frequency of exacerbations, decreased use of maintenance oral corticosteroids, reduction in the signs and symptoms of asthma, or an increase in predicted FFV ₁ from baseline. Diagnosts of hypereosinophilic symptome (Nucala and symptoms of asthma, on hypereosinophilic symptome (Nucala and ymptoms) of asthma, on hypereosinophilic symptome (Nucala and ymptoms). • Patient must be 12 years of age or older AND • The patient has bad at least 2 HES flares within the past 12 months AND • The patient has bad at least 2 HES flares within the past 12 months AND • The prescriber is an allergist in to an allergist, hematologist, or pulmonologist. Diagnosis of Chromic Rhinosinustiis with Nasul Polyps (Nucala Only): • Patient is 18 years of age or older AND • Prescriber is an allergist is earl Polyps (Nucala Only): • Patient has had an inadequate response to at least a 3-month trial of 2 different sast a corticosteroids	(100 171 required unless outer wise noted)	(1717equireu)	TH CHILDRIN
			eosinophilic phenotype as defined by pre-treatment blood eosinophil count of ≥ 150 cells per mcL within the previous 6 weeks or ≥ 300 cells per mcL within 12 months prior to initiation of therapy AND • The patient has a history of uncontrolled asthma symptoms (symptoms occurring almost daily or waking at night with asthma at least once a week) or 2 or more exacerbations in the previous year despite regular use of medium-high dose ICS/LABA for a minimum of 3 consecutive months, with or without oral corticosteroids. Pharmacy claims will be evaluated to assess compliance with therapy. AND • The prescriber is an allergist, immunologist, or pulmonologist. AND • For approval of Cinqair or Nucala, the patient must have a documented side effect, allergy, or treatment failure with Dupixent or Fasenra. • For continuation of therapy after the initial 6-month authorization, the patient must continue to receive therapy with an ICS/LABA AND have either a decreased frequency of exacerbations, decreased use of maintenance oral corticosteroids, reduction in the signs and symptoms of asthma, or an increase in predicted FEV₁ from baseline. Diagnosis of hypereosinophilic syndrome (Nucala only): • Patient must be 12 years of age or older AND • The patient must have a blood eosinophil count of ≥ 1,000 cells per mcl AND • The patient has had at least 2 HES flares within the past 12 months AND • The patient is on a stable dose of background HES therapy (chronic or episodic corticosteroids, immunosuppressive, or cytotoxic therapy) for at least 4 weeks prior to treatment initiation AND • The prescriber is an allergist, hematologist, immunologist, or pulmonologist Diagnosis of Chronic Rhinosinusitis with Nasal Polyps (Nucala Only): • Patient has had an inadequate response to at least a 3-month trial of 2 different nasal corticosteroids AND • Patient has had an inadequate response to at least a 10–14-day course of oral corticosteroids AND • Patient has had an inadequate response to at least a 10–14-day course of oral cortic

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
		corticosteroid AND there must be documented improvement in nasal
		symptoms.
		Dupixent:
		Diagnosis of moderate to severe persistent asthma:
		 The patient must be 6 years of age or older AND
		• The patient must have an eosinophilic phenotype as defined by pre-treatment
		blood eosinophil count of ≥ 150 cells per mcL within the previous 6 weeks
		or ≥ 300 cells per mcL within 12 months prior to initiation of therapy OR
		the patient is dependent on oral corticosteroids. • The patient has a history of uncontrolled asthma symptoms (symptoms
		occurring almost daily or waking at night with asthma at least once a
		week) or 2 or more exacerbations in the previous year despite regular use
		of medium-high dose ICS/LABA for a minimum of 3 consecutive months,
		with or without oral corticosteroids. Pharmacy claims will be evaluated to
		assess compliance with therapy. AND
		 The prescriber is an allergist, immunologist, or pulmonologist AND For continuation of therapy after the initial 6 month authorization, the patient
		must continue to receive therapy with an ICS/LABA AND have either a
		decreased frequency of exacerbations OR decreased use of maintenance
		oral corticosteroids OR reduction in the signs and symptoms of asthma
		OR an increase in predicted FEV1 from baseline.
		Diagnosis of Chronic Rhinosinusitis with Nasal Polyps:
		• Patient is 18 years of age or older AND
		Prescriber is an allergist or ENT specialist AND
		 Patient has had an inadequate response to at least a 3-month trial of 2 different nasal corticosteroids AND
		• Patient has had an inadequate response to at least a 10–14-day course of oral
		corticosteroids AND
		 Patient will use Dupixent concurrently with an intranasal corticosteroid
		• For continuation of therapy after the initial 6-month authorization, the patient
		must continue to receive therapy with an intranasal corticosteroid AND
		there must be documented improvement in nasal symptoms. Diagnosis of Eosinophilic Esophagitis:
		Patient is 12 years of age or older, weighing at least 40kg AND Prescriber is an altergist or gostscenteral oriet AND.
		Prescriber is an allergist or gastroenterologist AND Diagnosis is confirmed by andoscopia scaphogoal biopsy showing > 15
		 Diagnosis is confirmed by endoscopic esophageal biopsy showing ≥ 15 intraepithelial eosinophils per high-power field AND
		 Symptoms of esophageal dysfunction are present (e.g. pain while
		• symptoms of esophagear dystunction are present (e.g. pain white swallowing, sensation of food being stuck in the throat or chest) AND
		• The patient has had an inadequate response after a minimum trial of 8
		• The patient has had an inadequate response after a minimum trial of 8 weeks to at least one of the following: swallowed topical
		corticosteroids (e.g. Budesonide) or high-dose proton inhibitor.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
		• For continuation of therapy after the initial 6-month authorization, there
		must be documented improvement in EoE symptoms.
		Diagnosisis is Prurigo Nodularis:The patient must be 18 years of age or older AND
		 Diagnosis is confirmed based on the following: chronic pruritis lasting ≥ 6
		weeks, history and/or signs of repeated scratching, and multiple
		localized or generalized pruriginous skin lesions (e.g. whitish or pink
		papules, nodules and/or plaques) AND
		• The patient has had a documented side effect, allergy, or treatment failure
		(defined as daily treatment for at least one month) with at least one moderate to high potency topical corticosteroid and one preferred
		topical calcineurin inhibitor within the last 6 months
		• For continuation of therapy after the initial 6-month authorization, there
		must be documented improvement in PN symptoms.
		Limitations: Dupixent®, Fasenra®, Nucala® and Cinqair® will not be considered
		in patients who are currently smoking or in combination with omalizumab or Tezepelumab.
		Tezspire:
		• The patient must be 12 years of age or older AND
		• The patient has a history of uncontrolled asthma symptoms (symptoms
		occurring almost daily or waking at night with asthma at least once a
		week) or 2 or more exacerbations in the previous year despite regular use of medium-high dose ICS/LABA for a minimum of 3 consecutive
		months, with or without oral corticosteroids. Pharmacy claims will be
		evaluated to assess compliance with therapy. AND
		• The prescriber is an allergist, immunologist, or pulmonologist AND
		• If the patient has an eosinophilic phenotype (as defined by pretreatment
		blood eosinophil count of ≥ 150 cells per mcL within the previous 6 weeks or ≥ 300 cells per mcL within 12 months prior to initiation of
		therapy), there must have been a documented side effect, allergy, or
		treatment failure with Dupixent or Fasenra AND
		• For continuation of therapy after the initial 6 month authorization, the
		patient must continue to receive therapy with an ICS/LABA AND have
		either a decreased frequency of exacerbations OR decreased use of maintenance oral corticosteroids OR reduction in the signs and
		symptoms of asthma OR an increase in predicted FEV1 from baseline.
		Limitations: Tezspire will not be considered in patients who are currently
		smoking or in combination with anti-IgE, anti-IL4, or anti-IL5 monoclonal
		antibodies.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
	IMMUNOSUPPRESANTS,	ORAL
AZATHIOPRINE tablet CYCLOSPORINE capsule CYCLOSPORINE MODIFIED MYCOPHENOLATE MOFETIL tablet, capsule, suspension MYCOPHENOLIC ACID delayed release tablet SIROLIMUS tablet TACROLIMUS capsule	Astagraf® XL (tacrolimus) capsule Azasan® (azathioprine) tablet Cellcept® (mycophenolate mofetil) tablet, capsule, suspension Envarsus® XR (tacrolimus) tablet Everolimus (compare to Zortress®) tablet Gengraf® (cyclosporine modified) capsule, solution Imuran® (azathioprine) tablet Lupkynis™ (voclosporin) capsule Myfortic® (mycophenolic acid) delayed release tablet Neoral® (cyclosporine modified) capsule, solution Prograf® (tacrolimus) capsule, granules for suspension Rapamune® (sirolimus) tablet, solution Rezurock™ (belumosudil) tablet Sandimmune® (cyclosporine) capsule, solution Zortress® (everolimus) tablet	 Criteria (except Lupkynis and Rezurock): The patient has been started and stabilized on the requested product OR the patient has a documented side effect, allergy, or treatment failure to a preferred agent (if a product has and AB rated generic, there must be a trial of the generic formulation). Lupkynis: The patient has a diagnosis of Systemic Lupus Erythematosus (SLE) ANI The patient has active Lupus Nephritis confirmed by urine/blood tests or kidney biopsy AND The patient is ≥ 18 years of age AND Medication is prescribed by, or in consultation with, a nephrologist or rheumatologist AND The patient has clinical progression (e.g. worsening of proteinuria or serum creatinine) after 3 months of induction therapy with corticosteroids plus cyclophosphamide or mycophenolate mofetil OR failure to respond after 6 months of induction therapy with corticosteroids plus cyclophosphamide or mycophenolate mofetil AND Medication will be used in combination with background immunosuppressive therapy (e.g. mycophenolate mofetil and systemic corticosteroids) AND The patient has a documented intolerance or treatment failure with Benlysta Rezurock: The patient has a diagnosis of Chronic Graft-versus-host disease AND The patient has had a treatment failure with at least 2 prior courses of systemic immunosuppressant therapy (e.g. Corticosteroids, rituximab) AND The prescriber attests to monthly monitoring of liver function tests (total bilirubin, AST, and ALT)
CRYOPYRIN ASSO	OCIATED PERIODIC SYNDROMES (CAPS) A	ND PERIODIC FEVER SYNDROME (PFS)
All Products Require PA	Arcalyst [®] (rilonacept) **QTY LIMIT: 2 vials for loading dose, then 1 vial per Week **Ilaris® (canakinumab)	Ilaris: The diagnosis is Cryopyrin-Associated Periodic Syndrome (CAPS) OR The diagnosis is Familial Cold Autoinflammatory Syndrome (FCAS), Familial Mediterranean Fever (FMF), Hyper-IgD periodic fever syndrome (HIDS), Muckle-Wells Syndrome (MWS), or Tumor Necrosis Factor Recentor Associated Periodic Syndrome (TRAPS), AND, The patient is > 4

Receptor Associated Periodic Syndrome (TRAPS) AND The patient is > 4

Arcalyst: The diagnosis is Cryopyrin-Associated Periodic Syndrome (CAPS)

years old

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA OR The diagnosis is Familial Cold Autoinflammatory Syndrome (FCAS) OR The diagnosis is Muckle-Wells Syndrome (MWS) AND The patient is > 12 years old Note: Medical Records to support the above diagnosis must accompany the Prior Authorization request. Authorization for continued use shall be reviewed at least every 12 months to confirm patient has experienced disease stability or improvement while on therapy.
	IRON CHELATING AGE	
DEFERASIROX tablet	Deferasirox dispersible tablet, granule pack Deferiprone tablet Exjade® (defarasirox) dispersible tablet Ferripirox® (deferiprone) tablet, solution Jadenu® (deferasirox) tablet, granule pack	 Deferasirox dispersible tablet, Exjade dispersible tablet: The patient has a medical necessity for a non-solid oral dosage form AND for approval of Exjade, the patient has a documented intolerance to generic deferasirox dispersible tablets. Deferiprone tablet, Ferriprox tablet, Jadenu tablet: the patient has a documented intolerance to generic deferasirox tablets Deferasirox granule pack, Ferripirox solution, Jadenu granule pack: The patient has a medical necessity for a non-solid oral dosage form AND The patient has a documented intolerance to generic deferasirox dispersible tablets.
	LIPOTROPICS	
BILE ACID SEQUESTRANTS		
CHOLESTYRAMINE powder (compare to Questran®) CHOLESTYRAMINE LIGHT powder (compare to Questran Light®) COLESTIPOL tablets (compare to Colestid®) WELCHOL® (colesevelam) tablets, powder packets	Colesevelam (compare to Welchol®) Colestid® tablets, granules (colestipol) Colestipol granules, packets Prevalite powder (cholestyramine light) Questran® powder (cholestyramine) Questran Light® powder (cholestyramine light)	 Colesevelam: The patient has had a documented intolerance to the brand name equivalent. Colestipol granules, packets: The patient has a documented side effect, allery, or treatment failure with two preferred bile acid sequestrants. Prevalite, Questran, Questran Light, Colestid: The patient has had a documented intolerance to the preferred generic formulation.
FIBRIC ACID DERIVATIVES		
GEMFIBROZIL (compare to Lopid [®]) 600 mg FENOFIBRATE MICRONIZED CAPSULE (compare to Lofibra® capsules) 67 mg, 134 mg, 200 mg FENOFIBRATE NANOCRYSTALIZED (compare to Tricor [®]) 48 mg, 145 mg tablets FENOFIBRATE TABLETS (compare to Lofibra®	Antara [®] (fenofibrate micronized) 30 mg, 43 mg, 90 mg, 130 mg F Fenofibrate capsule (compare to (Lipofen [®]) 50 mg, 150 mg	Lopid: The patient has had a documented intolerance to generic gemfibrozil. All other non-preferred medications: The patient has had a documented side effect, allergy, or treatment failure with two preferred fibric acid derivatives (If a product has an AB rated generic, there must have been a trial with the generic formulation.)

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
tablets) 54 mg, 160 mg	Fenofibrate micronized (compare to Antara [®]) 43 mg, 130 mg Fenofibric acid (compare to Trilipix) 45 mg, 135 mg delayed release capsule Fenofibric acid 35 mg, 105 mg <i>QTY LIMIT:</i> 1 capsule/day Fenoglide [®] (fenofibrate MeltDose) 40 mg, 120 mg Lipofen [®] (fenofibrate) 50 mg, 150 mg Lopid [®] (gemfibrozil) 600 mg Tricor [®] (fenofibrate nanocrystallized) 48 mg, 145 mg Trilipix (fenofibric acid) 45 mg, 135 mg delayed release capsule	
MISC. HOMOZYGOUS FAMILIAL HYPERCHO	LESTEROLEMA (HoFH) AGENTS	
All products require PA	Evkeeza TM (evinacumab-dgnb) intravenous solution Juxtapid [®] (lomitapide) Capsule <i>QTY LIMIT</i> : 5 and 10 mg caps = 1/day, 20 mg cap = 3/day Maximum day supply per fill is 28 days	 CRITERIA FOR APPROVAL: Total cholesterol levels > 290mg/dL or LDL-C > 190mg/dL (adults) OR Total cholesterol levels > 260mg/dL or LDL-C > 155mg/dL (children < 16 years) and TG within reference range or Confirmation of diagnosis by gene testing AND Documented adherence to prescribed lipid lowering medications for the previous 90 days AND Recommended or prescribed by a lipidologist or Cardiologist AND Inability to reach goal LDL-C despite a trial of 2 or more maximum tolerated dose of statins (one of which must be atorvastatin or rosuvastatin), ezetimibe 10mg daily, and Repatha
NICOTINIC ACID DERIVATIVES		
NIACIN NIACIN extended release		
STATINS		
ATORVASTATIN (compare to Lipitor®) LOVASTATIN PRAVASTATIN ROSUVASTATIN (compare to Crestor®) SIMVASTATIN (compare to Zocor®) Note: All preferred agents have a quantity limit of 1 tablet/day except Lovastatin 40mg which has a quantity limit of 2 tablets/day	Altoprev® (lovastatin SR) Crestor® (rosuvastatin) Ezallor ® (rosuvastatin) sprinkle capsule Fluvastatin Fluvastatin ER (compare to Lescol® XL) Lescol® XL (fluvastatin ER) Lipitor® (atorvastatin) Livalo® (pitavastatin) Zocor® (simvastatin) Zypitamag™ (pitavastatin)	Non-preferred agents (except as noted below): The patient must have a documented side effect, allergy, or treatment failure to 3 preferred statins. If the product has an AB rated generic, one trial must be the generic formulation. Ezallor: medical necessity for a specialty dosage form has been provided Zypitamag: The patient must have a documented side effect, allergy, or treatment failure to 3 preferred statins AND clinical justification is provided documenting why the patient is unable to use Livalo.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
	Note: All non-preferred agents have a quantity limit of 1 tablet/day except fluvastatin IR which has a quantity limit of 2 tablets/day.	LIMITATIONS: Simvastatin 80 mg: initiation of simvastatin 80 mg or titration to 80 mg is not recommended by the FDA due to the increased risk of myopathy, including rhabdomyolysis. Patients may only continue on this dose when new to Medicaid if the patient has been taking this dose for 12 or more months without evidence of muscle toxicity. If the request is for Zocor 80 mg, the patient must have met the prior treatment length requirement and have a documented intolerance to the generic equivalent
MISCELLANEOUS/COMBOS		
Ezetimibe (compare to Zetia®) QTY LIMIT: 1 tab/day Omega-3-acid ethyl esters (compare to Lovaza®) Ezetimibe/simvastatin (compare to Vytorin®) 10/10 mg, 10/20mg, and 10/40mg QTY LIMIT: 1 tab/day	Amlodipine/atorvastatin (compare to Caduet®) QTY LIMIT: 1 tab/day Caduet® (atorvastatin/amlodipine) QTY LIMIT: 1 tab/day Ezetimibe/simvastatin (compare to Vytorin®) 10/80mg strength only Icosapent Ethyl (compare to Vascepa®) QTY LIMIT: 4 caps/day Lovaza® (omega-3-acid ethyl esters) Omega-3-acid ethyl esters (compare to Lovaza®) Nexletol® (bempedoic acid) QTY LIMIT: 1 tab/day Nexlizet® (bempedoic acid/ezetimibe) QTY LIMIT: 1 tab/day Vascepa® (icosapent ethyl) QTY LIMIT: 4 caps/day Vytorin® (ezetimibe/simvastatin) QTY LIMIT: 1 tab/day Zetia® (ezetimibe) QTY LIMIT: 1 tab/day	 Lovaza, Vytorin, Zetia: patient must have a documented intolerance to the generic equivalent. Icosapent Ethyl, Vascepa: Indication for use is severe hypertriglyceridemia: The patient has pre-treatment triglyceride levels > 500 mg/dL AND The patient has a documented contraindication, side effect, allergy, or treatment failure to Omega-3-acid ethyl esters. Indication for use is cardiovascular risk reduction: The patient has pre-treatment triglyceride levels > 150 mg/dL AND The patient is receiving adjunct therapy with a maximally tolerated high intensity statin AND For approval of icosapent ethyl, the patient has had a documented intolerance to brand Vascepa Amlodipine/atorvastatin, Caduet: The patient is unable to take the individual separate agents AND for approval of Caduet, the patient must have also had a documented intolerance to the generic equivalent. Nexletol, Nexlizet: The patient has had an inadequate response to a 3-month trial of atorvastatin or rosuvastatin OR Patient has demonstrated statin intolerability as defined by statin-related rhabdomyolysis or skeletal related muscle symptoms AND Patient (if eligible) will continue adjunct therapy with maximally tolerated high intensity statin. If patient is using simvastatin, dose should not exceed 20 mg/day; if patient is using pravastatin, dose should not exceed 40 mg/day Ezetimibe/simvastatin (10/80): the patient has been taking this dose for 12 or more months without evidence of muscle toxicity.
PCSK9 INHIBITORS		more mental mande endence of massic contents.
Preferred After Clinical Criteria Are Met PRALUENT® (alirocumab) QTY LIMIT: 2ml (75 mg injection every 2 weeks	Leqvio® (inclisiran) prefilled syringe	 Criteria for approval: The patient's age is FDA approved for the given indication AND Concurrent use with statin therapy AND

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
or 300 mg every month)/28 days Max 28-day supply REPATHA® (evolocumab) Sureclick, prefilled syringe QTY LIMIT: 2ml (2 injections)/28 days Max 28-day supply REPATHA® (evolocumab) Pushtronix TM QTY LIMIT: 3.5ml (One single-use infusor and prefilled cartridge)/28 days, Max 28-day supply		 Documented adherence to prescribed lipid lowering medications for the previous 90 days AND Inability to reach goal LDL-C despite a trial of 2 or more maximum tolerated dose of statins (one of which must be atorvastatin or rosuvastati For approval of Leqvio, the patient must have a documented side effect, allergy, or treatment failure (defined as inability to get within 10% of stated LDL-C goal, not to exceed guideline recognized goals) with a minimum 12-week trial of both Praluent and Repatha.
	MISCELLANEOUS	
KUVAN® (sapropterin) 100mg, 500mg powder PYRIDOSTIGMINE BROMIDE (Compare to Mestinon) SAPROPTERIN 100mg powder TRANEXAMIC ACID (compare to Lysteda®) QTY LIMIT: 30 tablets/28 days FENSOLVI® (leuprolide acetate) subcutaneous injection QTY LIMIT: 1 vial every 6 months Preferred After Clinical Criteria Are Met CARGLUMIC ACID (compare to Carbaglu®) dispersible tablets CRYSVITA® (burosumab-twza)	Brineura [™] (cerliponase alfa) <i>QTY LIMIT</i> : 1 package per 14 days (Brineura Injection, 2 vials of 150mg/5ml, and Intraventricular Electrolytes Injection, 1 vial of 5ml) Carbaglu® dispersible tablets (carglumic acid) Elaprase [®] (idursulfase) <i>QTY LIMIT</i> : calculated dose/week Firdapse® (amifampridine) <i>QTY LIMIT</i> : 8 tablets/day Galafold [™] (migalastat) <i>QTY LIMIT</i> : 14 caps/28 days Maximum day supply = 28 days Gamifant® (emapalumab-lzsg)	 Patient is 3 years of age or older AND The diagnosis or indication is late infantile neuronal ceroid lipofuscinosis type 2 (CLN2), confirmed by deficiency of the lysosomal enzyme tripeptidyl peptidase-1 (TPP1) (results of genetic testing must be submitted AND The prescriber is a neurologist or other physician specializing in intraventricular administration Note: Bineura will be approved as a medical benefit ONLY and will NOT be approved if billed through pharmacy point of sale. Initial approval will be granted for 3 months. Renewal may be granted for up to 12 months. For therapy continuation, clinical documentation must be submitted documenting improvement or maintenance of motor ability OR slower progression of

Hydroxyprogesterone caproate 250 mg/ml vial

Luxturna® (voretigine neparvovec-rzyl) suspension for

QTY LIMIT: one injection per eye per lifetime

Myalept® (metreleptin) vial for subcutaneous injection

Maximum day supply per fill = 30 days

(intramuscular injection)

Lysteda® tablets (tranexamic acid)

QTY LIMIT: one vial/day

QTY LIMIT: 30 tablets/28 days

subretinal injection

Mestinon®

Crysvita:

Patient is ≥ 1 year of age AND

equivalent of the requested medication.

Patient has a diagnosis of X-linked hypophosphatemia AND

Carbaglu, Carglumic Acid: The diagnosis or indication for the requested

deficiency, propionic acidemia, or methylmalonic acidemia AND The

Carbaglu, the patient has had a documented intolerance to the generic

medication is hyperammonemia due to N-acetylglutamate synthetase (NAGS)

prescriber is in consultation with a specialist AND for approval of brand name

prescriber is a specialist in metabolic disorders (e.g., medical geneticist) or

Medication is prescribed by or in consultation with an endocrinologist or

DD FEED DED. 1 GELVING	NOW PREPER PER 1 GENERAL	
PREFERRED AGENTS	NON-PREFERRED AGENTS	DA COMPENIA
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
	Oxlumo TM (pegvaliase-pqpz) Ruzurgi® (amifampridine) OTY LIMIT: 10 tablets/day Sapropterin (compare to Kuvan®) tablets, 500mg powder Thyquidity TM (levothyroxine sodium) oral solution Tirosint®-Sol (levothyroxine sodium) oral solution Vyvgart® (efgartigimod alfa-fcab) IV solution Xatmep TM (methotrexate) oral solution Zinplava TM (Bezlotoxumab) injection Zokinvy® (lonafarnib) capsule	 Patient has not received oral phosphate or vitamin D analogs within 1 week prior to starting therapy AND Baseline fasting serum phosphorous level is below the lower limit of the laboratory normal reference range AND Patient does not have severe renal impairment, defined as a GFR of < 30mL/min AND Dose does not exceed 90mg every 14 days (pediatrics) or 90mg every 28 days (adults) Note: Initial approval will be granted for 6 months. Renewal may be granted for up to 1 year. For therapy continuation, patient must have disease response as indicated by one of the following: Increased serum phosphate levels, not exceeding the upper limit of the laboratory normal range. A reduction in serum total alkaline phosphatase activity. Improvement in symptoms (e.g. skeletal pain, linear growth, etc.). Improvement in radiographic imaging of Rickets/osteomalacia. Elaprase (Hunter's Syndrome Injectable): The diagnosis or indication for the requested medication is Hunter's Syndrome Fabrazyme: Diagnosis or indication is Fabry Disease. Firdapse, Ruzurgi: patient has a diagnosis of Lambert-Eaton myasthenic syndrome (LEMS) AND prescription is initiated by or in consultation with a neurologist AND patient does not have a history of seizures AND for approval of Firdapse, the patient must have a documented intolerance to Ruzurgi. Initial approval will be granted for 3 months with documentation of the patient's baseline clinical muscle strength assessment using a standardized rating scale. For re-approval after 3 months, the patient must have improved, or stable symptoms documented with the appropriate standardized rating scale Galafold: Patient is ≥ 18 years of age AND Diagnosis or indication is Fabry Disease with an amenable galactosidase alpha (GLA) gene variant for treatment (result

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
		therapy. Initial approval will be granted for 3 months. For re-approval, there must be documented reduction in the size and redness of angiofibromas from baseline. Korsuva: The patient has a diagnosis of moderate-to-severe pruritis associated with chronic kidney disease AND the patient is receiving hemodialysis AND the patient has a documented side effect, allergy, or treatment failure with at least 1 topical and 1 systemic pruritis treatment (e.g. antihistamines, corticosteroids, gabapentin, pregabalin, capsaicin) Kuvan tabs, Sapropterin tabs: patient has a documented intolerance to the powder formulation. Luxturna: patient must have inherited retinal dystrophy due to mutations in both copies of the RPE65 gene (results of genetic testing must be submitted) AND patient has sufficient viable retinal cells as determined by the treating physician(s) AND Luxturna will be administered by a retinal specialist/surgeon experienced in performing intraocular surgery and associated with an Ocular Gene Therapy Treatment Center. Lysteda the patient has had a documented intolerance to the generic product. Myalept: Patient has a diagnosis of congenital or acquired generalized lipodystrophy AND Patient has one or more of the following metabolic abnormalities AND is refractory to current standards of care for lipid and diabetic management: Insulin resistance (defined as requiring > 200 units per day), Hypertriglyceridemia, Diabetes AND Prescription is written by or in consultation with an endocrinologist AND The prescriber is registered in the MYALEPT REMS program. Reauthorization for continued use criteria: Patient has experienced an objective response to therapy • Sustained reduction in hemoglobin A1c (HbA1c) level from baseline OR • Sustained reduction in triglyceride (TG) levels from baseline. Oxlumo: The patient has a diagnosis of Primary Hyperoxaluria Type I (PHI) confirmed via genetic testing (identification of alanine: glyoxylate aminotransferase gene (AGXT) mutation) AND urinary oxalate excretion > 0.5mmol/1.73 m

NON-PREFERRED AGENTS	
(PA required)	PA CRITERIA
	Palynziq has a black box warning for anaphylaxis which can occur at any time during treatment. Patients, pharmacies, and physicians must be enrolled in the Palynziq REMS program AND concurrent auto-injectable epinephrine must be prescribed. Note: Initial approval will be granted for 4 loading doses (the first 3 loading doses should be administered at 14-day intervals; the 4th loading dose should be administered 30 days after the 3th dose). Renewal may be granted for up to 12 months with a maximum of 3 doses approved per year (12mg(5ml) every 4 months). For therapy continuation, clinical documentation must be submitted documenting improvement or maintenance of motor ability OR slower progression of disease than would otherwise be expected. Sapropterin 500mg powder: patient has a documented intolerance to brand Kuvan Thyquidity, Tirosint-Sol: The patient has a medical necessity for a non-solid oral dosage form and the medication cannot be administered by crushing oral tablets AND for approval of Tirosint-Sol, the patient must have a documented intolerance to Thyquidity. Vyvgart: • Patient is ≥ 18 years of age AND • Patient has a diagnosis of generalized Myasthenia Gravis with Myasthenia Gravis Foundation of America (MGFA) clinical classification class II to IV AND • Patient is anti-acetylcholine receptor (AChR) positive AND • Patient has IgG levels of at least 6g/L AND • Patient has IgG levels of at least 6g/L AND • Patient has IgG levels of at least 6g/L AND • Patient has IgG levels of at least 6g/L AND • Patient has IgG levels of at least 6g/L AND • Patient has IgG levels of at least 6g/L AND • Patient has IgG levels of at least 6g/L AND • Patient has IgG levels of at least 6g/L AND • Patient has IgG levels of at least 6g/L AND • Patient has IgG levels of at least 6g/L AND • Patient has IgG levels of at least 6g/L AND • Patient has IgG levels of at least 6g/L AND • Patient has IgG levels of at least 6g/L AND • Patient has IgG levels of at least 6g/L AND • Patient has IgG levels of at leas
	NON-PREFERRED AGENTS (PA required)

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
		fidaxomicin) AND • The patient is at high risk for recurrence based on at least one of the following: ○ Age ≥ 65 years ○ Two or more episodes of CDI within the past 6 months ○ The patient is immunocompromised ○ The patient has clinically severe CDI (e.g. fever, abdominal tenderness, WBC ≥ 15,000 cells/mm³, albumin <30g/L, or renal failure) Zokinvy: The patient meets FDA approved age and BSA AND the patient has a diagnosis of Hutchinson-Gilford Progeria Syndrome (HGPS) OR the patient has a diagnosis of processing-deficient Progeroid Laminopathies with documentation of either Heterozygous LMNA mutation with progerin-like protein accumulation or Homozygous or compound heterozygous ZMPSTE24 mutations. Note: A single-dose of 10mg/kg will be approved per active CDI. A repeat dose will not be approved for recurrence of the same active infection.
AMYOTROPHIC LATERAL SCLEROSIS (ALS)		
RILUZOLE (Compare to Rilutek®)	Exservan TM (riluzole) film Radicava [®] (edaravone) IV injection Relyvrio TM (sodium phenylbutyrate/taurursodiol) powder for suspension OTY LIMIT: 2 packets/day Rilutek® (riluzole) Tiglutik TM (riluzole) suspension	 Exservan, Tiglutik: patient must be unable to take whole or crushed Riluzole tablets Radicava: The diagnosis is amyotrophic lateral sclerosis (ALS) AND Disease duration is ≤ 2 years AND Patient has functionally retained most activities of daily living AND Patient has normal respiratory function (defined as a % predicted forced vital capacity of ≥ 80%) AND Patient does not have a sulfite allergy AND Initial approval will be granted for 14 doses/28 days and all subsequent approvals will be for 10 doses/28 days Relyvrio: The diagnosis is amyotrophic lateral sclerosis (ALS) AND Disease duration is ≤ 18 months AND The patient has a slow vital capacity (SVC) spirometry test of greater than 60% of predicted at screening AND Baseline ALS Functional Rating Scale-Revised (ALSFRS-R) total score has been completed AND Initial approval will be granted for 6 months. For reapproval, clinical notes must indicate there has been improved or maintained baseline functional ability as measured by ALSFRS-R scale. Rilutek: patient must have a documented intolerance with riluzole.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
COMPLEMENT INHIBITORS		
All products require PA	Enjaymo [™] (sutimlimab-jome) Empaveli [™] (pegcetacoplan) subcutaneous solution <i>QTY LIMIT:</i> 8 vials/28 days Soliris® (eculizumab) vial Ultomiris® (ravulizumab-cwvz)	Enjaymo: The patient has a diagnosis of cold agglutinin syndrome (CAD) AND the patient does not have an active chronic systemic infection (e.g. Hepatitis B, Hepatitis C, HIV) AND the medication is prescribed by, or in consultation with, a hematologist AND the patient has had at least one blood transfusion in the 6 months prior to starting Enjaymo AND the patient has received the pneumococcal, Haemophilus influenzae, and meningococcal vaccines at least 2 weeks prior to therapy initiation. Authorization for continued use shall be reviewed to confirm that the patient has experienced an objective response to the therapy (e.g. stabilization of hemoglobin levels, reductions in transfusions, improvement in hemolysis, etc.) Empaveli: The patient has a diagnosis of paroxysmal nocturnal hemoglobinuria (PNH) documented by flow cytometry AND The patient has received the meningococcal vaccine at least 2 weeks prior to therapy initiation. Authorization for continued use shall be reviewed to confirm that the patient has experienced an objective response to the therapy (e.g. stabilization of hemoglobin levels, reductions in transfusions, improvement in hemolysis, etc.). Note: For patients switching from eculizumab, an additional 4 weeks of eculizumab will be approved before continuing monotherapy with Empaveli. For patients switching from ravulizumab, Empaveli will be initiated no more than 4 weeks after the last dose of ravulizumab. Ongoing combination therapy of complement inhibitors will not be approved.
		Soliris: Indication for use is Atypical Hemolytic Uremic Syndrome: Dose requested must be within the FDA parameters for loading and maintenance dose Indication for use is paroxysmal nocturnal hemoglobinuria (PNH): Diagnosis is documented by flow cytometry AND The patient has received the meningococcal vaccine at least 2 weeks prior to therapy initiation. Authorization for continued use shall be reviewed to confirm that the patient has experienced an objective response to the therapy (e.g. stabilization of hemoglobin levels, reductions in transfusions, improvement in hemolysis, etc.) Indication for use is Myasthenia Gravis: The patient is anti-aceytlcholine receptor (AchR) antibody positive AND the patient has a documented side effect, allergy, or treatment failure with at least 2 immunosuppressive therapies (e.g. corticosteroids, azathioprine, cyclosporine, mycophenolate, etc.). Ultomiris: The patient has a diagnosis of Atypical Hemolytic Uremic Syndrome or a diagnosis of paroxysmal nocturnal hemoglobinuria (PNH) documented by flow cytometry AND The patient has received the meningococcal vaccine at least 2 weeks prior to therapy initiation. Authorization for continued use shall be reviewed to confirm that the patient has experienced an objective response to the

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
		therapy (e.g. stabilization of hemoglobin levels, reductions in transfusions,
		improvement in hemolysis, etc.) Note: Dose requested must be within the weight-based parameters for loading and maintenance dose
GLYCOPYRROLATE		weight-based parameters for loading and maintenance dose
GLYCOPYRROLATE 1 mg, 2 mg tablets (compare	Cuvposa ^(g) oral solution (glycopyrrolate)	Cuvposa, Glycopyrrolate oral solution: The patient has medical necessity for a
to Robinul [®] , Robinul Forte [®])	Maximum days supply per fill is 30 days Dartisla ODT TM (glycopyrrolate)	non-solid oral dosage form OR the dose cannot be obtained from the tablet formulation.
	QTY LIMIT = 4 tabs/day	Dartisla ODT: The patient has been established on the 2mg dosage strength of
	Glycopyrrolate 1mg/5ml oral solution (compare to Cuvposa)	another form of glycopyrrolate AND the patient has a documented intolerance to glycopyrrolate tablets and solution.
	Robinul® (glycopyrrolate) 1mg	Robinul, Robinul Forte: The patient has a documented intolerance to
	Robinul® Forte (glycopyrrolate) 2mg	glycopyrrolate tablets.
INJECTABLE METHOTREXATE		
METHOTREXATE 25 MG/ML solution for injection	Otrexup® or Rasuvo® Single-dose auto-injector for subcutaneous use (methotrexate) QTY LIMIT: 4 syringes/28 days RediTrex® Prefilled syringe for subcutaneous use (methotrexate)	Otrexup, Rasuvo, Reditrex: The patient has a diagnosis of rheumatoid arthritis (RA), polyarticular juvenile idiopathic arthritis (pJIA) or psoriasis. AND The patient has been intolerant to oral methotrexate AND The patient has been unable to be compliant with a preferred form of injectable methotrexate (includes difficulty with manual dexterity)
	QTY LIMIT: 4 syringes/28 days	• • • • • • • • • • • • • • • • • • • •
MINERALOCORTICOID RECEPTOR ANTAG		
EPLERENONE	Aldactone® (spironolactone)	Aldactone, Inspra: The patient has a documented intolerance to the generic
SPIRONOLACTONE	CaroSpir® (spironolactone) oral suspension	formulation
	Inspra® (eplerenone) Kerendia® (finerenone)	Carospir: patient has a medical necessity for a specialty dosage form (i.e. dysphagia, swallowing disorder).
		Kerendia: The patient has a diagnosis of chronic kidney disease (CKD) associated
		with Type II Diabetes AND the estimated glomerular filtration rate at baseline is ≥ 25 mL/min/1.73m2 AND the urine albumin-to-creatinine ratio is ≥ 30mg/g
		AND the patient is currently receiving, or has a contraindication to, an ACE inhibitor or angiotensin receptor blocker (ARB)
NEUROMYELITIS OPTICA SPECTRUM DISO	RDFRS (NMOSD)	minibilor of angiotensin receptor blocker (AKB)
All Products Require PA	Enspryng® (satralizumab-mwge)	Enspryng, Soliris, Uplizna:
7 Trouves Acquire 171	prefilled syringe QTY LIMIT = 3/28 days for the first month then 1/28 days thereafter Soliris® (eculizumab) vial Uplizna® (inebilizumab-cdon) vial QTY LIMIT = 300mg x 2 doses for the first 2 weeks then 300mg every 6 months thereafter	 The patient is ≥ 18 years AND Diagnosis or indication is the treatment of neuromyelitis optica spectrum disorder (NMOSD) AND Patient is anti aquaporin-4 (AQP4) antibody positive AND Patient has a history of one or more relapses that required rescue therapy within the year prior to screening, or 2 or more relapses that required rescue therapy in 2 years prior to screening AND Patient must have a documented side effect, allergy, treatment failure, or contraindication to rituximab. Initial approval will be granted for 6 months. Renewal requires documentation of improvement or stabilization of neurologic symptoms

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
(1vo 1 A required unless otherwise noted)	(i A required)	TACKITERIA
		such as a decrease in acute relapses, reduced hospitalization, or reduction in plasma exchange treatments. Soliris, Uplizna additional criteria: The patient must have a documented side effect, allergy, treatment failure or contraindication to Enspryng.
SOMATOSTATIN ANALOGS OCTREOTIDE ACETATE solution for injection	D of '- @ (Described Conductation de materiale en la constalación de Cons
SANDOSTATIN® (octreotide acetate) LAR Depot	Bynfezia® (octreotide) pen Mycapssa® (octreotide) capsule QTY LIMIT: 4 caps/day Sandostatin® (octreotide) solution for injection Somatuline® Depot Injection (lanreotide) QTY LIMIT: 60 mg syringe = 0.2 ml/28 days, 90 mg syringe = 0.3 ml/28 days, 120 mg = 0.5 ml/28 days	 Bynfezia, Sandostatin: the patient has a documented intolerance to Octreotide injection. Mycapssa: the diagnosis or indication is long-term maintenance treatment of acromegaly AND the patient has already responded to and tolerated treatement with an injectable somatostatin alalog AND there is a clinically valid reason why the patient is unable to use Sandostatin LAR Depot. Somatuline: the patient has a documented side effect, allergy, treatment failure, or contraindication to Sandostatin LAR Depot.
SPINAL MUSCULAR ATROPHY		
Preferred After Clinical Criteria Are Met		Evrysdi:
ZOLGENSMA® (onasemnogene abeparvovec-xioi) intravenous suspension	Evrysdi® (risdiplam) oral solution Spinraza (nusinersen) injection 12mg/5ml single-dose vial	 The diagnosis is spinal muscular atrophy (SMA) AND Patient is 2 months of age or older AND Medication is prescribed per the dosing guidelines in the package insert AND A negative pregnancy test is obtained for females of reproductive potential prior to initiating therapy and patient has been advised to use effective contraception during treatment and for at least 1 month after her last dose AND A patient who has been started on Spinraza will not be approved for Evrysdi until at least 3 months after the fifth dose (i.e. nine months after the first loading dose, three months after the fifth dose). Concurrent use will not be approved. Note: For therapy continuation, clinical documentation must be submitted documenting improvement or maintenance of motor ability OR slower disease progression than would otherwise be expected. Spinraza: The diagnosis is spinal muscular atrophy (SMA) type 1,2, or 3 (results of genetic testing must be submitted) AND The patient has at least 2 copies of the SMN2 gene AND The need for invasive or noninvasive ventilation (if applicable) does not exceed more than 16 hours per 24 hour period AND Baseline motor ability has been established using one of the following exams:

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
(No FA required unless otherwise noted)	(FA required)	TA CKITEKIA
		 Prior to starting therapy, and prior to each dose, the following laboratory tests will be conducted: Platelet count, prothrombin time (PT), activated partial thromboplastin time (aPTT), and quantitative spot urine protein Concurrent use with Evrysdi will not be approved. Note: Initial approval will be granted for 4 loading doses (the first 3 loading doses should be administered at 14-day intervals; the 4th loading dose should be administered 30 days after the 3rd dose). Renewal may be granted for up to 12 months with a maximum of 3 doses approved per year (12mg(5ml) every 4 months). For therapy continuation, clinical documentation must be submitted documenting improvement or maintenance of motor ability OR slower progression of disease than would otherwise be expected. Zolgensma: The patient is less than 2 years of age AND The diagnosis is spinal muscular atrophy (SMA) AND The patient has bi-allelic mutations of the SMN1 gene AND The patient does not have advanced SMA (e.g. complete paralysis of limbs or permanent ventilator dependence) AND Medication is prescribed per the dosing guidelines in the package insert (recommended dose is 1.1 x 10⁴ vector genomes per kilogram) AND Baseline anti-AAV9 antibodies are less than 1:50 AND Prior to starting therapy and periodically for at least 3 months, the following laboratory tests will be conducted: Liver function (AST, ALT, total bilirubin, prothrombin time), platelet counts, and troponin-I Note: The safety and effectiveness of repeat administration has not been evaluated. Approval is limited to a single intravenous infusion.
SYSTEMIC LUPUS ERYTHEMATOSUS (SLE)		
	Benlysta® (belimumab) Maximum days supply per fill = 28 days Saphnelo TM (anifrolumab-fnia)	 Benlysta: Indication for use is Systemic Lupus Erythematosus (SLE): The patient is positive for autoantibodies (anti-nuclear antibody (ANA) and/or anti-double-stranded DNA (anti-dsDNA) AND The patient has had a documented inadequate response or intolerance to at least TWO of the following agents: NSAIDs, hydroxychloroquine, corticosteroids, azathioprine, methotrexate, mycophenolate mofetil AND Initial approval will be granted for 3 months. For therapy continuation, clinical documentation must be submitted documenting stable disease activity OR reduction in disease activity or corticosteroid dose. Note: The efficacy of Benlysta® has not been evaluated in patients with severe active central nervous system lupus. Benlysta has not been studied in combination with other biologics or intravenous cyclophosphamide. Use of Benlysta is not recommended in these situations. Indication for use is Active Lupus Nephritis: Diagnosis has been confirmed by urine/blood tests or kidney biopsy AND The patient is ≥ 18 years of age AND

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	 Medication is prescribed by, or in consultation with, a nephrologist or rheumatologist AND The patient has clinical progression (e.g. worsening of proteinuria or serum creatinine) after 3 months of induction therapy with corticosteroids plus cyclophosphamide or mycophenolate mofetil OR failure to respond after 6 months of induction therapy with corticosteroids plus cyclophosphamide or mycophenolate mofetil AND Medication will be used in combination with background immunosuppressive therapy (e.g. mycophenolate mofetil and systemic corticosteroids) AND Initial approval will be granted for 3 months. For therapy continuation, clinical documentation must be submitted documenting stable disease activity OR reduction in disease activity. Saphnelo: The patient has a diagnosis of moderate-severe Systemic Lupus Erythematosus AND The patient is ≥ 18 years of age AND Medication is prescribed by, or in consultation with, a nephrologist or rheumatologist AND The patient has had a documented inadequate response or intolerance to at least TWO of the following agents: hydroxychloroquine, corticosteroids, azathioprine, methotrexate, mycophenolate mofetil AND The patient has had a documented intolerance or treatment failure with Benlysta Initial approval will be granted for 3 months. For therapy continuation, clinical documentation must be submitted documenting stable disease
		 Initial approval will be granted for 3 months. For therapy continuation, clinical documentation must be submitted documenting stable disease activity OR reduction in disease activity or corticosteroid dose. Note: The efficacy of Saphnelo has not been evaluated in patients with severe active lupus nephritis or severe active central nervous system lupus. Saphnelo has not been studied in combination with other biologics or intravenous cyclophosphamide. Use of Saphnelo is not recommended in these situations.
	MOOD STABILIZE	iks ————————————————————————————————————
LITHIUM CARBONATE (formerly Eskalith®) LITHIUM CARBONATE SR (compare to Lithobid®, formerly Eskalith CR®) LITHIUM CITRATE SYRUP	Equetro [®] (carbamazepine SR) Lithobid [®] (lithium carbonate SR)	 Lithobid: The patient has had a documented side effect, allergy, or treatment failure with the generic equivalent of the requested medication. Equetro: The patient has had a documented side effect, allergy, or treatment failure with a carbamazepine product from the anticonvulsant therapeutic drug category

PREFERRED AGENTS

(No PA required unless otherwise noted)

NON-PREFERRED AGENTS

(PA required)

PA CRITERIA

MOVEMENT DISORDERS

Preferred After Clinical Criteria Are Met

AUSTEDO® (deutetrabenazine) tablets

QTY LIMIT: 48 mg/day

Maximum 1-month supply per fill

INGREZZA® (valbenazine tosylate) capsules

QTY LIMIT: 80 mg/day

Maximum 1-month supply per fill

TETRABENAZINE (compare to Xenazine®)

QTY LIMIT: 50 mg/day at initial approval (12.5

mg tablets ONLY), up to 100 mg/day at

subsequent approvals (12.5 mg or 25 mg tablets)

Maximum 1-month supply per fill

Xenazine® (tetrabenazine) tablets

QTY LIMIT: 50 mg/day at initial approval (12.5 mg tablets ONLY), up to 100 mg/day at subsequent

approvals (12.5 mg or 25 mg tablets)

Maximum 1-month supply per fill

Austedo: The diagnosis or indication for the requested medication is Huntington's Disease (HD) with chorea or Tardive Dyskinesia (TD) AND the results of an Abnormal Involuntary Movement Scale (AIMS) exam have been submitted AND the patient is ≥18 years of age. For re-approval, there must be documented clinical improvement.

Ingrezza: The diagnosis or indication for the requested medication is Tardive Dyskinesia (TD) AND the results of an Abnormal Involuntary Movement Scale (AIMS) exam have been submitted AND the patient is ≥18 years of age. For re-approval, there must be documented clinical improvement.

Tetrabenazine, Xenazine: The diagnosis or indication for use is Tourette Syndrome OR the diagnosis or indication for use is Huntington's Disease (HD) with Chorea or Tardive Dyskinesia (TD) AND the patient is ≥18 years of age AND for approval of Xenazine, the patient must have a documented intolerance to tetrabenazine.

Note: Austedo, Tetrabenazine, and Xenazine are contraindicated in patients with Huntington's Disease who are suicidal or with untreated/inadequately treated depression.

MULTIPLE SCLEROSIS MEDICATIONS

INJECTABLES

INTERFERONS

AVONEX® (interferon B-1a)
BETASERON® (interferon B-1b)
REBIF® (interferon B-1a)
REBIF® REBIDOSE (interferon B-1a)

OTHER

COPAXONE® 20 mg (glatiramer acetate) *QTY LIMIT*: 1 kit/30 days

Preferred After Clinical Criteria are Met

Extavia[®] (interferon beta-1b)

Copaxone[®] 40 mg (glatiramer)

QTY LIMIT: 12 syringes (12 ml)/28 days

QTY LIMIT: 1 kit/30days

Glatiramer Acetate (compare to Copaxone®) 40 mg

QTY LIMIT: 12 syringes (12 ml)/28 days

Glatopa® 20 mg (glatiramer acetate)

QTY LIMIT: 1 carton (30 syringes/30 days

Glatopa® 40 mg (glatiramer)

QTY LIMIT: 12 syringes (12 ml)/28 days

Ampyra, Tecfidera: patient must have a documented intolerance to the generic equivalent

Bafiertam, Vumerity: Patient is ≥ 18 years AND has a diagnosis of relapsing forms of Multiple Sclerosis AND the patient has a documented side effect, allergy, treatment failure, or contraindication to at least two preferred drugs, one of which must be Dimethyl fumarate.

Copaxone 40 mg Syringe: The patient is unable to tolerate or be compliant with Copaxone 20 mg daily dosing.

Extavia: Patient has a diagnosis of multiple sclerosis. AND The provider provides a clinical reason why Betaseron cannot be prescribed.

Glatiramer, Glatopa: Patient is ≥ 18 years AND diagnosis of relapsing forms of Multiple Sclerosis AND the provider provides a clinical reason why

PREFERRED AGENTS	NON-PREFERRED AGENTS		
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA	
ORAL AUBAGIO® (teriflunamide) tablet QTY LIMIT: 1 tablet/day Maximum 30-day supply per fill DALFAMPRIDINE ER tablet (compare to Ampyra®) QTY LIMIT: 2 tablets/day Maximum 30-day supply per fill DIMETHYL FUMARATE QTY LIMIT: 2 capsules/day Maximum 30-day supply per fill GILENYA® (fingolimod) capsule QTY LIMIT: 1 capsule/day Maximum 30-day supply per fill	Kesimpta® (ofatumumab) Lemtrada® (alemtuzumab) intravenous Ocrevus® (ocrelizumab)	 Copaxone cannot be prescribed. Mavenclad: Patient is ≥ 18 years AND has a diagnosis of relapsing-remitting MS (RRMS) or active secondary progressive MS (SPMS) AND Documentation is provided showing ≥ 1 relapse within the past year AND baseline CBC w/ diff (including lymphocyte count), liver function tests, and MRI (within the past 3 months) have been completed AND the patient is negative for HIV, Hepatitis B, and Hepatitis C infections AND the patient is not pregnant AND patient has a documented side effect, allergy, treatment failure or contraindication to at least three preferred drugs AND dosing does not exceed any of the following: 2 tablets per day, 10 tablets per cycle, 2 treatment cycles per course, 1 course per year. Following the administration of 2 treatment courses, Mavenclad may not be administered during the next 2 years. Mayzent, Ponvory, Zeposia: Diagnosis of relapsing-remitting MS, Clinical Isolated Syndrome, or Active Secondary Progressive MS (SPMS): Patient is ≥ 18 years AND Patient CYP2C9 variant status has been tested to determine genotyping (Mayzent only; required for dosing; therapy is contraindicated in CYP2C9*3/*3) AND Baseline CBC, electrocardiogram (ECG), and ophthalmic evaluation have been completed AND Patient has a documented side effect, allergy, treatment failure or contraindication to at least two preferred drugs, one of which must be Gilenya Kesimpta, Lemtrada, Ocrevus: Patient is ≥18 years AND has a diagnosis of relapsing multiple sclerosis AND has a documented side effect, allergy, treatment failure or contraindication to at least two preferred drugs, one of which must be Gilenya or Tysabri, unless contraindicated. OR Patient is ≥18 years AND has a diagnosis of relapsing form of Multiple Sclerosi AND has a documented side effect, allergy, treatment failure or contraindication to at least three prefe	
MUSCLE RELAXANTS, SKELETAL			
	MOSGLE RELIGION TO, SIXELLINE		
MUSCULOSKELETAL AGENTS		Amrix, Cyclobenzaprine 7.5 mg, Fexmid: The prescriber must provide a	

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
SINGLE AGENTS CYCLOBENZAPRINE 5 mg, 10 mg tablets (compare to Flexeril®) QTY LIMIT: 5 mg = 6 tablets/day, 10 mg = 3 tablets/day METHOCARBAMOL tablets (compare to Robaxin®) QTY LIMIT: 8 tablets/day ORPHENADRINE CITRATE ER 100 mg tablet QTY LIMIT: 2 tablets/day COMBINATION PRODUCT All products require PA ASA = aspirin	Amrix (cyclobenzaprine sustained-release) capsule <i>QTY LIMIT:</i> 1 capsule/day Carisoprodol tablets <i>QTY LIMIT:</i> 8 tablets/day Chlorzoxazone tablets <i>QTY LIMIT:</i> 4 tablets/day Cyclobenzaprine 7.5 mg tab (compare to Fexmid) <i>QTY LIMIT:</i> 3 tablets/day Fexmid (cyclobenzaprine) 7.5 mg tablet <i>QTY LIMIT:</i> 3 tablets/day Lorzone (chlorzoxazone) tablets <i>QTY LIMIT:</i> 4 tablets/day Metaxalone (compare to Skelaxin) tablets <i>QTY LIMIT:</i> 4 tablets/day Skelaxin (metaxalone) tablets <i>QTY LIMIT:</i> 4 tablets/day Soma (carisoprodol) tablets <i>QTY LIMIT:</i> 4 tablets/day Carisoprodol, ASA, codeine <i>QTY LIMIT:</i> 4 tablets/day	clinically valid reason why a preferred generic cyclobenzaprine 5mg or 10mg cannot be used. For approval of Fexmid, the patient must also have a documented intolerance to the generic equivalent. Baclofen oral solution Fleqsuvy: Patient has a medical necessity for a non-solid oral dosage form AND the patient has a documented intolerance to Lyvispah. Carisoprodol, Carisoprodol/ASA/codeine, Chlorzoxazone, Lorzone, Soma, Metaxalone, Skelaxin: The patient has had a documented side effect, allergy or treatment failure with two different preferred musculoskeletal agents. Additionally, if a brand name product is requested where an AB rated generic exists, the patient must also have had a documented intolerance to the generic product. Dantrium, Zanaflex tablets: The patient must have a documented intolerance with the AB rated generic product. Lyvispah: Patient has a meical necessity for the non-solid oral dosage form. Tizanidine capsules, Zanaflex capsules: The prescriber must provide a clinically valid reason why generic tizanidine tablets cannot be used. AND If the request is for Zanaflex capsules, the patient must have a documented intolerance to generic tizanidine capsules
ANTISPASTICITY AGENTS BACLOFEN tablets DANTROLENE (compare to Dantrium [®]) TIZANIDINE (compare to Zanaflex [®]) tablets	Baclofen oral solution Dantrium [®] (dantrolene) Fleqsuvy [™] (baclofen) oral suspension Lyvispah [™] (baclofen) oral granule packet Tizanidine (compare to Zanaflex [®]) capsules Zanaflex [®] (tizanidine) capsules Zanaflex [®] (tizanidine) tablets	
	MUSCULAR DYSTROPHY	AGENTS
All products require PA	Amondys®45 (casimersen) Emflaza TM (deflazacort) Maximum 30-day supply per fill Exondys 51 TM (eteplirsen) Viltepso® (viltorsen) Vyondys 53 TM (golodirsen)	 Emflaza: The patient must be ≥ 2 years of age AND The patient must have a diagnosis of Duchenne Muscular Dystrophy AND There is documented improvement in muscle function or strength with

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
		use of prednisone, but the patient has experienced weight gain >10% of body weight within 3 months or >25% within 1 year. Amondys, Exondys, Viltepso, Vyondys: • The patient must have a diagnosis of Duchenne Muscular Dystrophy with a confirmed mutation of the DMD gene that is amenable to exon 45 skipping (for Amondys) or exon 53 skipping (for Cyllepso, Vyondys) (results of genetic testing must be submitted) AND • The prescriber is, or has consulted with, a neuromuscular disorder specialist AND • The dose does not exceed 30mg/kg once weekly (for Amondys, Exondys, Vyondys) or 80mg/kg once weekly (for Viltepso) AND • The patient is currently on a stable corticosteroid dose for at least 6 months. AND • Baseline documentation of the members voluntary motor and cardiac function has been provided and results have shown member retains meaningful voluntary motor function: Optional • 6-minute walk test (6MWT) or other timed fuctions tests (e.g time to stand [TTSTAND], tiem to run/walk 10 meters [TTRW], time to climb 4 stairs [TTCLIMB]) • Brooks Upper Extremity Test • North Star Ambulatory Assessment (NSAA) Required • Forced Vital Capacity (FVC) percent predicted • Ejection Fraction Percentage Note: Initial approval will be granted for 6 months. For re-approval after 6 months, the patient must demonstrate a response to therapy compared to baseline as evidenced by stable, improved, or slowed rate of either motor function or cardiac function degradation. Evidence may include one or more of the following (not all-inclusive): • 6MWT or other timed function tests (e.g., time to stand [TTSTAND], time to run/walk 10 meters [TTRW], time to climb 4 stairs [TTCLIMB]) • Brooks Upper Extremity Test • North Star Ambulatory Assessment (NSAA) • Forced Vital Capacity (FVC) percent predicted • Ejection Fraction Percentage

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
	NEUROGENIC ORTHOSTATIC H	IYPOTENSION
FLUDROCORTISONE MIDODRINE	Northera®	 Quantity Limits: Initial 2 weeks approval Continued therapy approvals based on documentation of continued benefit clinically and as evidenced by positional blood pressure readings Clinical Criteria: diagnosis of neurogenic orthostatic hypotension caused by primary autonomic failure (Parkinson's disease, multiple system atrophy, or pure autonomic failure), dopamine beta-hydroxylase deficiency, or non-diabetic autonomic neuropathy, AND the presentation of symptoms including dizziness, lightheadedness, and the feeling of "blacking out" AND Failure of multiple non-pharmacologic measures as appropriate (e.g. removal of offending medications, compression stockings, increased fluid and salt intake) AND Failure, intolerance or contra-indication to fludrocortisone AND midodrine
	NEUROPATHIC PAIN & FIBROMY	ALGIA AGENTS
Oral DULOXETINE (compare to Cymbalta®) QTY LIMIT: 2 capsules/day PREGABALIN (compare to Lyrica®) capsules QTY LIMIT: 3 capsules/day SAVELLA® (milnacipran) tablet, titration pack QTY LIMIT: 2 tablets/day	Cymbalta® (duloxetine) <i>QTY LIMIT:</i> 2 capsules/day Gralise® (gabapentin) tablet, starter pack <i>QTY LIMIT:</i> 3 tablets/day Maximum 30-day supply per fill Horizant® (gabapentin enacarbil) ER Tablet FDA maximum recommended dose = 1200 mg/day Lyrica® (pregabalin) capsules <i>QTY LIMIT:</i> 3 capsules/day Lyrica® CR (pregabalin, extended release) FDA maximum recommended dose = 330 mg/day (DPN), 660 MG/day (PHN) Lyrica® (pregabalin) solution	Cymbalta, Lyrica: the patient has had a documented intolerance with the generic equivalent. Gralise, Horizant: The patient has a diagnosis of post-herpetic neuralgia (PHN) AND The patient has had a documented side effect, allergy, contraindication of treatment failure with at least one drug from the tricyclic antidepressant class AND The patient has had an inadequate response to the generic gabapentin immediate-release. Pregabalin ER, Lyrica CR: The patient has a diagnosis of post-herpetic neuralgi (PHN) or diabetic peripheral neuropathy (DPN) AND patient has not been able to adherent to a twice daily dosing schedule of pregabalin immediate release resulting in a significant clinical impact AND for approval of pregabalin ER, the patient had documented intolerance to brand Lyrica CR. Note: The efficacy of Lyrica® CR had not been established for the management of fibromyalgia or as adjunctive therapy adult patients with partial onset seizures.

adult patients with partial onset seizures.

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required) Pregabalin (compare to Lyrica®) solution Pregabalin extended release (compare to Lyrica® CR) FDA maximum recommended dose =	Pregabalin solution, Lyrica solution: the patient is unable to use Lyrica capsules (e.g. Swallowing disorder) AND for approval of brand Lyrica oral solution, the patient must have a documented intolerance to the generic equivalent.
	330 mg/day (DPN), 660 mg/day (PHN)	
	NUTRITIONALS, LIQUID ORAL SU	UPPLEMENTS
All products require PA	Note: Nutritional supplements administered via tube feeds may be provided through the Medical Benefit	 EleCare, EleCare Jr: The patient is an infant or child who needs an amino acid-based medical food or who cannot tolerate intact or hydrolyzed protein. AND The product is being requested for the dietary management of protein maldigestion, malabsorption, severe food allergies, short-bowel syndrome, eosinophilic GI disorders, GI-tract impairment, or other conditions for which an amino acid-based diet is required. All Others: Requested nutritional supplement will be administered via tube feeding. OR Patient has one of the following conditions where feeding is difficult or malabsorption or maldigestion occurs: AIDS, Cancer, Cerebral Palsy, Cystic Fibrosis, Dementia resulting in loss of motor skills, Neuromuscular Disease, Short Gut. OR Patient has experienced unplanned weight loss or is extremely low weight (see further definitions below) OR Patient has demonstrated nutritional deficiency identified by low serum protein levels (albumin or pre-albumin levels to be provided) (albumin <3.5 g/dL/pre-albumin <15 mg/dL) Unplanned Weight Loss/Low Weight Table: Adult: Involuntary loss of > 10 % of body weight within 6 months OR Involuntary loss of > 5% of body weight within 1 month OR Loss of > 2% of body weight within one week OR BMI of <18.5 kg/m2 Elderly: (>65): Involuntary loss of > 10 % of body weight within 6 months OR Involuntary loss of > 5 % of body weight within 3 months OR Loss of > 2% of body weight within one month OR BMI of <18.5 kg/m2 Children: Anatomic causes for malnutrition have been evaluated and treated AND clinical diagnosis and documentation supports the need for enteral nutrition (See Below) Members weight is below the 5th percentile for sex and corrected age AND weight-to-length ratio is below the 10th percentile OR Sustained decrease in growth velocity as demonstrated by weight-forage or weight-for-length fall by two major percentiles (percentile markers 95, 90, 75, 50, 25, 10, and 5) over time (defined by the WHO for children l

DD FEED DED A CENTER	MOM PREFERRED A COMPAG	
PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
(10171 required timess otherwise noted)	(i / required)	TACKILKI
		than 2 years of age)
		Limitations: Approvals will be based on medical necessity for supplemental
		nutrition. Approval will NOT be granted for individuals whose need is nutritional rather than medical, including an unwillingness to consume solid or pureed
		foods. For nonmedical needs contact WIC at 800-464-4343
	ONCOLOGY: DRUGS (se	lect)
		Clinical Criteria: Medication is being used for an FDA approved indication AND
		age, dose, duration, required concurrent therapy, and past treatment failures (if
		applicable) are consistent with prescribing information AND the patient does not have any contraindications prohibiting use of the medication OR medication is
		being used in accordance with the National Comprehensive Cancer Network®
		(NCCN®) Clinical Practice Guidelines. Requests outside of these parameters
		require medical director review. This includes all cell and gene therapies, including CAR-T therapies, regardless of site of administration. For physician-
		administered drugs, please refer to the Fee Schedule for which codes require a
		PA: http://vtmedicaid.com/#/feeSchedule/hcpcs
	OPHTHALMICS	
ANTIBIOTICS		
		Single and Combination Agents (except noted below): The patient has had a
QUINOLONES BESIVANCE® (besifloxacin) suspension	Ciloxan® (ciprofloxacin) solution	documented side effect, allergy or treatment failure with at least TWO
CILOXAN® ointment	Gatifloxacin 0.5% solution (compare to Zymaxid [®]) Levofloxacin 0.5% solution	preferred ophthalmic antibiotics or ophthalmic antibiotic combination agents,
CIPROFLOXACIN HCL (compare to Ciloxan [®]) solution	Moxifloxacin 0.5% (compare to Moxeza®)	one of which must be in the same therapeutic class. (If a product has an AB
MOXIFLOXACIN 0.5% solution (compare to	(preservative free) solution	rated generic, there must have also been a trial of the generic formulation.)
Vigamox®)	Ocuflox [®] (ofloxacin) solution Ofloxacin (compare to Ocuflox [®]) solution	
	Vigamox [®] (moxifloxacin 0.5%) (preservative free)	
ALL CDOVEDES	solution	
MACROLIDES ERYTHROMYCIN ointment	Zymaxid [®] (gatifloxacin 0.5%) solution	
AMINOGLYCOSIDES		
SINGLE AGENT	Azasite [®] (azithromycin) solution	
	Azasne (aziuromycin) solution	

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
AK-TOB (tobramycin) solution GARAMYCIN® (gentamicin) ointment, solution GENTAK (gentamicin) ointment GENTAMICIN solution	All other brands	
TOBRAMYCIN solution (compare to Tobrex [®]) COMBINATION TOBRADEX [®] (tobramycin/dexamethasone) suspension, ointment ZYLET [®] (tobramycin/loteprednol) suspension	Tobrex [®] ointment, solution (tobramycin) Tobramycin w/Dexamethasone (compare to Tobradex [®]) suspension Tobradex ST [®] (tobramycin/dexamethasone) suspension Pred-G [®] S.O.P. (gentamicin/prednisolone) ointment Pred-G® (gentamicin/prednisolone) ointment, suspension	
MISCELLANEOUS SINGLE AGENT All products require PA	Bacitracin ointment Bleph- $10^{\textcircled{R}}$ (sulfacetamide) solution Sulfacetamide sodium (compare to Bleph- $10^{\textcircled{R}}$) solution Sulfacetamide sodium ointment	
Combination BACITRACIN ZINC W/POLYMYXIN B ointment NEOMYCIN/BACITRACIN/POLYMYXIN ointment NEOMYCIN/POLYMYXIN W/DEXAMETHASONE (compare to Maxitrol®) ointment, suspension NEOMYCIN/POLYMYXIN/BACITRACIN/ HYDROCORTISONE ointment POLYMYXIN B W/TRIMETHOPRIM (compare to Polytrim®) solution SULFACETAMIDE W/PREDNISOLONE SOD PHOSPHATE solution	Blephamide (sulfacetamide/prednisolone acetate) suspension Blephamide S.O.P. (sulfacetamide/prednisolone acetate) ointment Maxitrol (neomycin/polymyxin/dexamethasone) suspension, ointment Neomycin/Polymyxin W/Gramicidin solution Neomycin/Polymyxin w/Hydrocortisone ointment, suspension Polytrim (polymyxin B/trimethoprim) soln	
ANTIHISTAMINES AZELASTINE QTY LIMIT: 1 bottle/month KETOTIFEN 0.025 % QTY LIMIT: 1 bottle/month OLOPATADINE 0.1%, 0.2% QTY LIMIT: 1 bottle/month	Bepotastine (compare to Bepreve®) Bepreve® (bepotastine besilate) Epinastine QTY LIMIT: 1 bottle/month Lastacaft® (alcaftadine) QTY LIMIT: 1 bottle/month	Bepotastine, Bepreve, Epinastine: The patient has had a documented side effect, allergy, or treatment failure to a preferred ophthalmic antihistamine AND for approval of Bepotastine, the patient must have a documented intolerance to brand Bepreve. Lastacaft: The patient is pregnant, and the diagnosis is allergic conjunctivitis OR The patient has had a documented side effect, allergy, or treatment failure to a

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
(No 1 A required timess other wise noted)	(i A required)	TACRITICAL
	Zerviate® (cetirizine 0.24%) QTY LIMIT:60 vials/30 days	preferred ophthalmic antihistamine. Zerviate: The patient has had a documented side effect, allergy, or treatment failure to TWO preferred ophthalmic antihistamines.
CORTICOSTEROIDS: TOPICAL		
ALREX® (loteprednol) 0.2% suspension DEXAMETHASONE sodium phosphate 0.1% solution DUREZOL® (difluprednate) 0.05% emulsion FLAREX® (fluorometholone acetate) 0.1% suspension FML Forte® (fluorometholone) 0.25% suspension FLUOROMETHOLONE 0.1% suspension FML® (fluorometholone) 0.1% ointment LOTEMAX® (loteprednol) 0.5% suspension, ointment MAXIDEX® (dexamethasone) suspension PRED MILD® (prednisolone acetate) 0.12% suspension PREDNISOLONE ACETATE 1% suspension PREDNISOLONE SODIUM PHOSPHATE 1% solution	Difluprednate (compare to Durezol®) FML Liquifilm® (fluorometholone) 0.1% suspension Inveltys™ (loteprednol) suspension Lotemax® (loteprednol) 0.5% gel Lotemax SM (loteprednol) 0.038% gel drops Loteprednol suspension Pred Forte® (prednisolone acetate) 1% suspension All other brands	Non-preferred agents: The patient has had a documented side effect, allergy, or treatment failure with TWO preferred ophthalmic corticosteroids. (If a product has an AB rated generic, there must have been a trial of the generic formulation)
CYSTEAMINE		
All products require PA		
	Cystadrops® (cysteamine) 0.37% ophthalmic solution QTY LIMIT: 4 bottles (20 ml)/28 days Maximum day supply/Rx = 28 days Cystaran® (cysteamine) 0.44% ophthalmic solution <i>QTY LIMIT:</i> 4 bottles (60 ml)/ 28 days Maximum day supply/RX = 28 days	Cystadrops, Cystaran: The indication for use is corneal cystine accumulation in patients with cystinosis.
DRY EYE SYNDROME		
OCULAR LUBRICANTS Please refer to the DVHA website for covered OTC ocular lubricants https://dvha.vermont.gov/sites/dvha/files/documents/providers/Pharmacy/OTCWebList.pdf IMMUNOMODULATORS	Cequa TM (cyclosporine ophthalmic solution) 0.09% Cyclosporin ophthalmic emulsion 0.05% droperette (compare to Restasis®) QTY LIMIT: 180 vials per 90 days Eysuvis® (loteprednol etabonate ophthalmic suspension) 0.25%	 Cequa: The patient has a diagnosis of Dry Eye Disease AND has a documented side effect, allergy, or treatment failure to two ophthalmic immunomodulators, one of which must be Restasis. Cyclosporin emulsion, Tyrvaya, Xiidra: The patient has a diagnosis of Dry Eye Disease AND has a documented side effect, allergy or treatment failure to Restasis. Eysuvis: The patient has a diagnosis of Dry Eye Disease AND has failed at least a
RESTASIS® (cyclosporine ophthalmic emulsion)	Restasis [®] (cyclosporine ophthalmic emulsion) 0.05%	14-day course of a preferred OTC ocular lubricant AND has a documented side

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
0.05% droperette (NDC 00023916330 and 00023916360 are the only preferred NDC's) QTY LIMIT: 180 vials per 90 days GLAUCOMA AGENTS/MIOTICS	multidose bottle QTY LIMIT: 1 bottle (5.5ml) per 25 days Tyrvaya TM (varenicline) nasal spray QTY LIMIT: 2 bottles (8.4 ml) per 30 days Verkazia® (cyclosporine ophthalmic emulsion) 0.1% single dose vials Xiidra® (lifitegrast) solution QTY LIMIT: 60 vials per 30 days	effect, allergy, or treatment failure with 2 preferred ophthalmic corticosteroids, one of which must be a formulation of loteprednol. Restasis Multidose: Both package sizes of the droperettes must be on a long-term backorder and unavailable from the manufacturer. Verkazia: The patient has a diagnosis of vernal keratoconjunctivitis (VKC) AND the patient has had a documented side effect, allergy, or treatment failure with a mast cell stabilizer (e.g. cromolyn sodium) or a dual acting antihistamine/mast cell stabilizer (e.g. olopatadine, azelastine)
		AV DVA A A DD DVADO GALLA GALL
ALPHA-2 ADRENERGIC SINGLE AGENT ALPHAGAN P® 0.1 %, 0.15 % (brimonidine tartrate) BRIMONIDINE TARTRATE 0.2 %	Apraclonidine (compare to Iopidine [®]) Brimonidine tartrate 0.15 % (compare to Alphagan P [®]) Iopidine [®] (apraclonidine)	ALPHA 2 ADRENERGIC AGENTS: Single Agent: The patient has had a documented side effect, allergy, or treatment failure with at least one preferred ophthalmic alpha 2 adrenergic agent. If the request is for brimonidine tartrate 0.15%, the patient must have a documented intolerance of brand name Alphagan P 0.15%.
COMBINATION COMBIGAN® (brimonidine tartrate/timolol maleate) SIMBRINZA® (brinzolamide 1% and brimonidine 0.2%) Suspension	Brimonidine tartrate/timolol maleate (compare to Combigan®)	Brimonidine/timolol: the patient must have a documented intolerance to brand Combigan. BETA BLOCKERS: The patient has had a documented side effect, allergy or treatment failure with at least one preferred ophthalmic beta blocker.
BETA BLOCKER CARTEOLOL HCL LEVOBUNOLOL HCL TIMOLOL MALEATE (compare to Timoptic®)	Betaxolol HCl solution Betoptic S [®] (betaxolol suspension) Istalol [®] (timolol) Timoptic (timolol maleate) Timoptic XE [®] (timolol maleate gel) Timolol maleate gel (compare to Timotic XE [®])	PROSTAGLANDIN INHIBITORS Bimatoprost, Travoprost, Vyzulta, Xalatan, Xelpros, Zioptan: The patient has had a documented side effect, allergy or treatment failure with at least 2 preferred prostaglandin inhibitors. Durysta: The patient has had a documented side effect, allergy, or treatment failure with at least 2 preferred prostaglandin inhibitors OR the patient is not a candidate for topical drop therapy AND the patient does not have any of the
PROSTAGLANDIN INHIBITORS LATANOPROST (compare to Xalatan®) LUMIGAN® (bimatoprost) TRAVATAN Z® (travoprost) (BAK free)	Bimatoprost 0.03% (Lumigan [®]) Durysta® (bimatoprost) 10 mcg implant Travoprost BAK Free (compare to Travatan Z®) Vyzulta® (latanoprostene bunod) Xelpros [®] (latanoprost) (BAK free) Zioptan [®] (tafluprost)	 following contraindications: History of prior corneal transplantation or endothelial cell transplants (e.g. Descemet's Stripping Automated Endothelial Keratoplasty) Diagnosis of corneal endothelial dystrophy (e.g. Fuchs' Dystrophy) Absent or ruptured posterior lens capsule Approval will be limited to a single implant per eye without retreatment. CARBONIC ANHYDRASE INHIBITORS
RHO KINASE INHIBITORS SINGLE AGENT RHOPRESSA® (netarsudil)		Trusopt: The patient has had a documented intolerance to the generic equivalent product.Cosopt PF: The patient has had a documented intolerance to the preservatives in the generic combination product.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
COMBINATION ROCKLATAN® (netarsudil/latanoprost)		Miscellaneous: The patient has had a documented side effect, allergy or treatment failure with a preferred miscellaneous ophthalmic agent. If a product has an AB rated generic, there must have also been a trial of the
CARBONIC ANHYDRASE INHIBITOR SINGLE AGENT AZOPT® (brinzolamide 1%) DORZOLAMIDE 2 % (compare to Trusopt®)	Trusopt [®] (dorzolamide 2 %)	generic formulation)
COMBINATION DORZOLAMIDE w/TIMOLOL (compare to Cosopt®)	Cosopt PF [®] (dorzolamide w/timolol) (pres-free)	
MISCELLANEOUS ISOPTO® CARPINE (pilocarpine) PILOCARPINE HCL PHOSPHOLINE IODIDE® (echothiophate)	Miochol-E [®] (acetylcholine)	
MAST CELL STABILIZERS		
CROMOLYN SODIUM	Alocril [®] (nedocromil sodium) Alomide [®] (lodoxamide)	Criteria for Approval: The patient has had a documented side effect, allergy, or treatment failure with generic cromolyn sodium
NEUROTROPHIC KERATITIS		
All products require PA	Oxervate TM (cenegermin-bkbj) ophthalmic solution 0.002% <i>QTY LIMIT</i> : 1 vial (1mL) per eye per day Maximum of 8 weeks therapy	Oxervate: Medication is being prescribed by, or in consultation with, an ophthalmologist AND Patient has a diagnosis of Stage 2 or 3 neurotrophic keratitis (in one or both eyes) as evidenced by persistent epithelial defect or corneal ulceration AND patient has evidence of decreased corneal sensitivity in at least one corneal quadrant AND patient has failed one or more conventional non-surgical treatments such as artificial tears, gels, or ointments.
NON-STEROIDAL ANTI-INFLAMMATORY DR	UGS (NSAIDs)	
DICLOFENAC 0.1% ophthalmic solution KETOROLAC 0.4 % ophthalmic solution (compare to Acular LS®) KETOROLAC 0.5 % ophthalmic solution (compare to Acular®) NEVANAC® ophthalmic suspension (nepafenac 0.1%)	Acular [®] (ketorolac 0.5% ophthalmic solution) Acular LS [®] (ketorolac 0.4% ophthalmic solution) Acuvail (ketorolac 0.45 %) Ophthalmic Solution <i>QTY LIMIT</i> : 30-unit dose packets/15 days Bromfenac 0.09 % ophthalmic solution BromSite [™] (bromfenac 0.075%) solution Flurbiprofen 0.03% ophthalmic solution Ilevro® ophthalmic suspension (nepafenac 0.3%)	 Acuvail: The patient has had a documented side effect, allergy, or treatment failure to Acular OR ketorolac 0.5% OR The patient has a documented hypersensitivity to the preservative benzalkonium chloride. All other non-preferred agents: The patient has had a documented side effect, allergy, or treatment failure to TWO preferred agents. In addition, if a product has an AB rated generic, there must have also been a trial of the generic formulation.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
	Prolensa [®] ophthalmic solution (bromfenac 0.07%)	
PRESBYOPIA AGENTS		
All products require PA	Vuity TM (pilocarpine) 1.25% solution	Vuity: The patient has a diagnosis of presbyopia AND the patient is between the ages of 40-55 at the time of therapy initiation AND the medication is being prescribed by or in consultation with an optometrist or ophthalmologist AND the patient has failed corrective eyeglasses or contact lenses, unless contraindicated.
	OTIC ANTI-INFECTIVES/ANTI-INFL	AMMATORIES
ANTI-INFECTIVE SINGLE AGENT OFLOXACIN 0.3% Otic solution	Ciprofloxacin 0.2% otic solution QTY LIMIT: 14-unit dose packages/ 7 days	 Anti-infective single and combination agents: The patient has had a documented side effect, allergy, or treatment failure to two preferred products. DermOtic, Flac Oil: the patient has a documented intolerance to generic fluocinolone oil.
ANTI-INFECTIVE/CORTICOSTEROID COMBINATION CIPRODEX® (ciprofloxacin 0.3%/dexamethasone 0.1%) otic suspension CIPRO-HC® (ciprofloxacin 0.2%/hydrocortisone 1%) otic suspension NEOMYCIN/POLYMYXIN B SULFATE/HYDROCORTISONE SOLUTION, SUSPENSION	Cortisporin-TC® (neomycin/colistin/thonzium/hydrocortisone) Ciprofloxacin/Dexamethasone (compare to Ciprodex®) otic suspension Ciprofloxacin/Fluocinolone otic solution QTY LIMIT: 28-units dose packages/7days	
CORTICOSTEROID FLUOCINOLONE OIL 0.01%	DermOtic® Oil (fluocinolone acetonide) 0.01% Flac® Oil (fluocinolone acetonide) 0.01%	
MISCELLANEOUS AGENTS ACETIC ACID Otic solution	Acetic Acid/Hydrocortisone Otic Solution	

OVER THE COUNTER (OTC) MEDICATIONS

Please refer to the DVHA website for covered OTC categories not already managed on the PDL. Many categories limited to generics ONLY and other categories not covered. No PA process for non-covered OTCs.

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
https://dvha.vermont.gov/sites/dvha/files/docum	ents/providers/Pharmacy/OTCWebList.pdf	
	PANCREATIC ENZYME	PRODUCTS
CREON [®] DR Capsule ZENPEP [®] DR Capsule	Pertzye [®] DR Capsule Viokace [®] DR Capsule	Pertzye, Viokace: The patient has been started and stabilized on the requested product. OR The patient has had treatment failure or documented intolerance with both Creon and Zenpep.
	PARATHYROID A	GENTS
CALCITRIOL (compare to Rocaltrol®) CINACALCET (compare to Sensipar®) ERGOCALCIFEROL (compare to Drisdol®) PARICALCITOL (compare to Zemplar®)	Doxercalciferol (compare to Hectoral®) Drisdol® (ergocalciferol) Hectoral® (doxercalciferol) Natpara® (parathyroid hormone) QTY LIMIT: 2 cartridges per 28 days Parsabiv™ (etelcalcetide) Rayaldee® (calcifediol ER) Rocaltrol® (calcitriol) Sensipar® (cinacalcet) Zemplar® (paricalcitol)	Doxercalciferol, Drisdol, Hectoral, Rayaldee, Rocaltrol, Zemplar: The patient must have a documented side effect, allergy, or treatment failure to two preferred agents. If a product has an AB rated generic, one trial must be the generic formulation. Natpara: Natpara: Natpara: diagnosis of hypocalcemia secondary to hypoparathyroidism (but NOT acute post-surgical hypoparathyroidism within 6 months of surgery) AND Natpara PA form must be completed and clinical and lab documentation supplied AND Must be prescribed by an endocrinologist AND Must be documented by ALL of the following: History of hypoparathyroidism >18 months AND Concomitant serum intact parathyroid hormone (PTH) concentrations below the lower limit of the normal laboratory reference range on 2 test dates at least 21 days apart within the past 12 months AND No history of the following: mutation in CaSR gene OR pseudohypoparathyroidism OR a condition with an increased risk of osteosarcoma AND Hypocalcemia is not corrected by calcium supplements and preferred active forms of vitamin D alone AND Patients must be taking vitamin D metabolite/analog therapy with calcitriol ≥0.25 µg per day OR equivalent AND Must be taking supplemental oral calcium treatment ≥ 1000 mg per day over and above normal dietary calcium intake AND Serum calcium must be ≥ 7.5 mg/dl prior to starting Natpara AND Serum thyroid function tests and serum magnesium levels must be within normal limits AND Documentation of creatinine clearance > 30 mL/min on two separate

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
		measurements OR creatinine clearance > 60 mL/min AND serum creatinine < 1.5 mg/dL
		Parsabiv: indication is for the treatment of secondary hyperparathyroidism in a
		patient with Chronic Kidney Disease (CKD) receiving hemodialysis AND the
		patient has a documented side effect, allergy, or treatment failure with Sensipar.
		Note: treatment failure is defined as < 30% reduction from baseline in mean pre- dialysis PTH concentrations.
		Sensipar: the patient has a documented intolerance to the generic equivalent.
	PARKINSON'S MEDIC	ATIONS
DOPAMINE PRECURSOR		Inbrija: The patient has a diagnosis of Parkinson's disease with intermittent
CARBIDOPA/LEVODOPA (compare to Sinemet [®])	Inbrija® (levodopa capsule for inhalation)	presence
CARBIDOPA/LEVODOPA ER (compare to	QTY LIMIT: 10 caps/day	of OFF episodes AND the patient is currently taking Carbidopa/Levodopa AND
Sinemet [®] CR)	Rytary® (carbidopa/levodopa ER caps)	the patient has had a documented side effect, allergy, or treatment failure with Apokyn®
CARBIDOPA/LEVODOPA ODT	Sinemet [®] (carbidopa/levodopa)	Comtan, Sinemet, Parlodel, Stalevo: The patient has had a documented
		intolerance to the generic product.
DODAMINE A CONICTE (OD AL)	Mirapex ER [®] (pramipexole ER)	Ongentys: The diagnosis or indication is Parkinson's disease AND the patient
DOPAMINE AGONISTS (ORAL)	QTY LIMIT: 1 tab/day	has had a documented side effect, allergy, or treatment failure with
BROMOCRIPTINE (compare to Parlodel [®])	Pramipexole ER (compare to Mirapex ER®)	entacapone.
PRAMIPEXOLE (compare to Mirapex [®])	QTY LIMIT: 1 tab/day	Rytary: The patient has a diagnosis of Parkinson's disease,
ROPINIROLE (compare to Requip [®])	Ropinirole XL	post-encephalitic parkinsonism, or parkinsonism following intoxication from
	QTY LIMIT: $12 \text{ mg} = 2 \text{ tabs/day}$, All other strengths = 1 tab/day	carbon monoxide or manganese AND the prescriber is a neurologist AND the
	All other strengths = 1 tab/day	patient is having breakthrough symptoms despite a combination of concurrent
		IR and ER formulations of carbidopa/levodopa Azilect, Rasagiline: The diagnosis or indication is Parkinson's disease. AND
		The patient has had a documented side effect, allergy, or treatment failure
	Neupro® (rotigotine) transdermal patch	with selegiline. AND The dose requested does not exceed 1 mg/day
	QTY LIMIT: $2, 4, 6, \text{ and } 8 \text{ mg} = 1 \text{ patch/day}$	Gocovri: diagnosis or indication is for the treatment of dyskinesia in a patient
		with Parkinson's Disease AND the patient is currently receiving levodopa-
DOPAMINE AGONISTS (TRANSDERMAL)	Comtan® (entacapone)	based therapy (with or without concomitant dopaminergic medications) AND
All products require PA	Ongentys® (opicapone)	the patient has a documented side effect, allergy, or treatment failure with
	Tasmar [®] (tolcapone)	immediate release amantadine. Note: treatment failure is defined by a
COMT INHIBITORS	Tolcapone (compare to Tasmar®)	decrease in effectiveness despite attempts to increase dosage to 300mg/day or
		by temporarily discontinuing amantadine for several weeks and restarting
ENTACAPONE (compare to Comtan [®])	Azilect [®] (rasagiline)	therapy.
	QTY LIMIT: 1 mg/day	Kynmobi: The patient has a diagnosis of Parkinson's disease with intermittent
	Rasagiline (compare to Azilect®)	presence of OFF episodes AND the patient is receiving concomitant levodopa which has been at a stable dose for a minimum of 4 weeks AND the patient is
MAO-B INHIBITORS	QTY LIMIT: 1 mg/day	not taking a 5HT3 antagonist (e.g ondansetron, alosetron) concurrently AND
SELEGILINE	Xadago [®] (safinamide) <i>QTY LIMIT:</i> 1 tab/day	the patient has had a documented side effect, allergy or treatment failure with
	Q11 LIMI1. 1 tau/day	and patient has had a documented side effect, affer by or deathern failule with

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
ANTICHOLINERGICS BENZTROPINE TRIHEXYPHENIDYL ADENSOSINE RECEPTOR AGONIST All products require PA OTHER APOKYN® (apomorphine) AMANTADINE syrup AMANTADINE capsules, tablets (PA required for ≤ 10-day supply) CARBIDOPA/LEVODOPA/ENTACAPONE (compare to Stalevo®)	Zelapar [®] (selegiline ODT) QTY LIMIT: 2.5 mg/day Nourianz (istradefylline) QTY LIMIT: 1 tab/day Gocovri TM (amantadine extended release) QTY LIMIT: 2 tabs/day Kynmobi® (apomorphine) sublingual film Osmolex® ER (amantadine extended-release) QTY LIMIT: 1 tablet/strength/day Stalevo® (carbidopa/levodopa/entacapone)	Apokyn. Mirapex ER, Pramipexole ER, Ropinirole XL: The diagnosis or indication is Parkinson's disease. Requests will not be approved for Restless Leg Syndrome (RLS) AND The patient has had an inadequate response (i.e. wearing off effect or "off" time) with the immediate release product. OR The patient has not been able to be adherent to a three times daily dosing schedule of the immediate release product resulting in a significant clinical impact. AND If the requested product has an AB rated generic, the patient has a documented intolerance to the generic product. Neupro: The patient has a medical necessity for a specialty dosage form. Nourianz: The patient has a diagnosis of Parkinson's disease with intermittent presence of OFF episodes AND the patient is currently taking Carbidopa/Levodopa AND the patient has had a documented side effect, allergy, or treatment failure with TWO preferred medications being used as adjunct therapy. Osmolex ER: patient has not been able to be adherent to the dosing schedule of amantadine immediate release resulting in a significant clinical impact. Tasmar, Tolcapone: The diagnosis or indication is Parkinson's disease. AND The patient has had a documented side effect, allergy, or treatment failure with entacapone AND patient has provided written acknowledgement of risks per the package insert. For approval of brand Tasmar, the patient must have documented intolerance to the generic equivalent. Xadago: The diagnosis or indication is Parkinson's disease AND The patient is on current therapy with levodopa/carbidopa AND The patient has had a documented side effect, allergy, or treatment failure with selegiline. Note: Xadago will not be approved for monotherapy. Zelapar: The diagnosis or indication is Parkinson's disease. AND The patient is on current therapy with levodopa/carbidopa. AND Medical necessity for disintegrating tablet administration is provided (i.e. inability to swallow tablets or drug interaction with oral selegiline). AND the dose requested does not exceed 2.5 mg
	PLATELET INHIBITO	DRS
AGGREGATION INHIBITORS BRILINTA® (ticagrelor) Tablet	Effient [®] (prasugrel) Tablet	Agrylin, Effient, Plavix: The patient has had a documented intolerance to the generic formulation of the medication.

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
QTY LIMIT: 2 tablets/day CILOSTAZOL CLOPIDOGREL 75 mg (compare to Plavix [®]) PRASUGREL (compare to Effient®) OTHER ANAGRELIDE (compare to Agrylin [®]) ASPIRIN DIPYRIDAMOLE DIPYRIDAMOLE/ASPIRIN	QTY LIMIT: 1 tablet/day Plavix [®] 75 mg (clopidogrel bisulfate) Zontivity [®] (vorapaxar) Tablet QTY LIMIT: 1 tablet/day Agrylin [®] (anagrelide)	Zontivity: The patient is started and stabilized on the medication. (Note: samples are not considered adequate justification for stabilization.) OR The patient has a history of myocardial infarction (MI) or peripheral arterial disease (PAD) AND The indication for use is reduction of thrombotic cardiovascular events. AND The medication is being prescribed in combination with aspirin and/or clopidogrel. Limitations: Plavix/clopidogrel 300 mg is not an outpatient dose and is not covered in the pharmacy benefit.

OTHER ANAGRELIDE (compare to Agrylin [®]) ASPIRIN DIPYRIDAMOLE DIPYRIDAMOLE/ASPIRIN	Agrylin [®] (anagrelide)	combination with aspirin and/or clopidogrel. Limitations: Plavix/clopidogrel 300 mg is not an outpatient dose and is not covered in the pharmacy benefit.
	PLATELET STIMULATING	G AGENTS
Preferred After Clinical Criteria Are Met PROMACTA® (eltrombopag)	Doptelet® (avatrombopag) Mulpleta® (lusutrombopag) Nplate® (romiplostim) Tavalisse™ (fostamatinib disodium hexahydrate)	Doptelet: Indication for use is chronic immune (idiopathic) thrombocytopenic purpura (ITP): The patient's platelet count is less than 30,000/μL (< 30 x 10°/L) or the patient is actively bleeding AND The patient has an insufficient response or documented intolerance to corticosteroids, immunoglobulins, or splenectomy AND Promacta. Indication for use is thrombocytopenia in a patient with chronic liver disease scheduled to undergo an elective surgical or dental procedure: The patient is at least 18 years of age AND the patient's platelet count is less than 50,000/μL (< 50 x 10°/L) AND approval will be limited to a maximum of 5 days' supply per procedure Mulpleta: The patient is at least 18 years of age AND the diagnosis is thrombocytopenia in a patient with chronic liver disease scheduled to undergo an elective surgical or dental procedure AND the patient's platelet count is less than 50,000/μL (< 50 x 10°/L) AND approval will be limited to a maximum of 7 days' supply per procedure. AND patient has had a documented side effect, allergy, contraindication, or treatment failure to Doptelet. Nplate: The diagnosis or indication is chronic immune (idiopathic) thrombocytopenic purpura (ITP). AND The patient's platelet count is less than 30,000/μL (< 30 x 10°/L) or the patient is actively bleeding. AND The patient has an insufficient response or documented intolerance to corticosteroids, immunoglobulins, or splenectomy AND Promacta. Promacta: Indication for use is chronic immune thrombocytopenia (ITP): The patient's platelet count is less than 30,000/μL (< 30 x 10°/L) or the patient is actively bleeding, AND the patient has had an insufficient response or documented intolerance to corticosteroids, immunoglobulins, or splenectomy. Indication for use is chronic Hepatitis-C associated thrombocytopenia: The patient is at least 18 years of age AND medication is used to initiate or maintain interferon-based therapy. Indication for use is Severe Aplastic Anemia: patient has had an inadequate response

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
		to standard immunosuppressive therapy (e.g. cyclosporine). Tavalisse: The patient is at least 18 years of age AND The diagnosis is chronic immune thrombocytopenia (ITP) AND The patient's platelet count is less than < 30 x 10°/L AND The patient has had a documented side effect, allergy, treatment failure or a contraindication to therapy with corticosteroids AND the patient has failed at least one of the following additional treatments: immunoglobulins, rituximab, splenectomy, or a thrombopoietin receptor agonist (e.g. eltrombopag, romiplostim, etc.). Note: Initial approval will be granted for 12 weeks. For therapy continuation, the patient must have achieved and maintained a platelet count of at least 50 x 10°/L and/or have a documented decrease in rescue treatment(s) with platelet transfusions.
	PSEUDOBULBAR AFFECT A	GENTS
All products require PA	Nuedexta® capsules (dextromethorphan/quinidine) QTY LIMIT: 2 capsules/day	Nuedexta: The patient must have a diagnosis of pseudobulbar affect (PBA) secondary to a neurological condition AND the patient has had a trial and therapy failure at a therapeutic dose with a tricyclic antidepressant (TCA) or an SSRI AND the patient has documentation of a current EKG (within the past 3 months) without QT prolongation AND initial authorizations will be approved for 6 months with a baseline Center for Neurologic Studies Lability Scale (CNS-LS) questionnaire AND subsequent prior authorizations will be considered at 6 month intervals with documented efficacy as seen in an improvement in the CNS-LS questionnaire
	PSORIASIS	
BIOLOGICS: Initial approval is 3 months, renewals	s are 1 year	
Preferred After Clinical Criteria Are Met INJECTABLE AVSOLA® (infliximab-axxq) biosimilar to Remicade® ENBREL® (etanercept) QTY LIMIT: 50 mg = 8 syringes/28 days for the first 3 months; then 4 syringes/28 days 25 mg = 8 syringes/28 days subsequently HUMIRA® (adalimumab) QTY LIMIT: 4 syringes/28 days for one month; 2 syringes/28 days subsequently INFLECTRA® (infliximab-dyyb) biosimilar	Cimzia® (certolizumab pegol) <i>QTY LIMIT:</i> 1 kit/28 days (starter X 1, then regular) Cosentyx® (secukinumab) Ilumya™ (tildrakizumab-asmn) <i>QTY LIMIT:</i> 2 ml (2 syringes) for the first month then 1 ml (1 syringe)/84 days subsequently Remicade® (infliximab) Renflexis™ (infliximab-abda) biosimilar to Remicade® Siliq™ (brodalumab) injection <i>QTY LIMIT:</i> 6 ml (4 syringes) for the first month then 3 ml (2 syringes)/28 days subsequently	Clinical Criteria: For all drugs (except Spevigo): The prescription must be written by a dermatologist or rheumatologist AND The patient has a documented diagnosis of moderate to severe plaque psoriasis and has already been stabilized on the drug being requested OR The prescription must be written by a dermatologist or rheumatologist AND The patient has a documented diagnosis of moderate to severe plaque psoriasis affecting > 10% of the body surface area (BSA) and/or has involvement of the palms, soles, head and neck, or genitalia and has had a documented side effect, allergy, inadequate treatment response, or treatment failure to at least 2 different categories of therapy [i.e. at least 2 topical agents and at least 1 oral systemic agent, (unless otherwise contraindicated)] from the following categories: Topical agents:

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
to Remicade® TALTZ® (ixekizumab) QTY LIMIT: 3 syringes/28 days for the first month, 2 syringes/28 days months 2 and 3 and 1 syringe/28 days subsequently ORAL OTEZLA® tablet (apremilast) QTY LIMIT: Starter Pack = 55 tablets/28 days, 30 mg = 2 tablets/day	Skyrizi TM (risankizumab-rzaa) <i>QTY LIMIT</i> : 150 mg/28 days for the first month and 150mg/84 days thereafter Spevigo® (spesolimab-sbzo) <i>QTY LIMIT</i> : 900 mg (15 ml) per dose Stelara® (ustekinumab) <i>QTY LIMIT</i> : 45 mg (0.5 ml) or 90 mg (1 ml) per dose (90mg dose only permitted if patient weight > 100kg) One dose/28 days for the first month and one dose/84 days thereafter Tremfya® (guselkumab) <i>QTY LIMIT</i> : 2 syringes/28 days for the first month, then 1 syringe every 56 days thereafter Sotyktu® (deucravacitinib) <i>QTY LIMIT</i> : 1 tablet/day	emollients, keratolytics, corticosteroids, calcipotriene, tazarotene, etc. Systemic agents: methotrexate, sulfasalazine, azathioprine, cyclosporine, tacrolimus, mycophenolate mofetil, etc. Phototherapy: ultraviolet A and topical psoralens (topical PUVA), ultraviolet A and oral psoralens (systemic PUVA, narrow band ultraviolet B (NUVA), etc. Additional Criteria for Taltz: The prescriber must provide evidence of a trial and failure or contraindication to a preferred TNF Inhibitor. Additional Criteria for Cimzia, Cosentyx, Ilumya, Siliq, Skyrizi, Sotyktu, Stelara, Tremfya: The prescriber must provide a clinically valid reason why both a preferred TNF Inhibitor and Taltz [®] cannot be used. Note: Cosentyx approvals for 300mg dose(s) must use "300DOSE" package (containing 2x150mg pens or syringes) Approval will not be granted for 2 separate 150mg packages. Additional Criteria for Remicade, Renflexis: The prescriber must provide a clinically valid reason why Humira®, Taltz®, and Avsola/Inflectra cannot be used. Spevigo: • The patient is experiencing a moderate-to-severe intensity flare of generalized pustular psoriasis (GPP) as defined by: • A Generalized Pustular Psoriasis Physician Global Assessment (GPPPGA) total score of at least 3 (moderate) or greater AND • The presence of fresh pustules (new appearance or worsening of pustules) AND At least 5% of body surface area (BSA) covered with erythema and the presence of pustules AND • The patient will not use concomitantly with other systemic immunosuppressants or topical agents AND Approval will be granted for a maximum of two 900mg doses, given 7 days apart.
NON-BIOLOGICS		
ORAL ACITRETIN capsules CYCLOSPORINE (generic) METHOTREXATE (generic) TOPICAL CALCIPOTRIENE Cream, Ointment, Solution	Methoxsalen (compare to Oxsoralen-Ultra [®]) Oxsoralen-Ultra [®] (methoxsalen) Calcitriol (compare to Vectical [®]) Ointment QTY LIMIT: 200 g (2 tubes)/week Calcipotriene Foam (compare to Sorilux®) Calcipotriene/betamethasone ointment (compare to Taclonex [®]) QTY LIMIT: Initial fill = 60 grams	 Duobrii lotion: the patient has had an inadequate response to at least 2 different preferred high or very high potency corticosteroids AND tazarotene cream. Enstilar, Taclonex or Calcipotriene/betamethasone dipropionate Ointment or Scalp Suspension: The patient has had an inadequate response to a trial (defined as daily treatment for at least one month) of a betamethasone dipropionate product and Dovonex (or generic calcipotriene), simultaneously. Calcipotriene Foam, Calcitriol Ointment, Sorilux, Tazarotene, Vtama, Zoryve: The patient has a diagnosis of mild-to-moderate plaque psoriasis AND The patient has demonstrated inadequate response, (defined as daily treatment for at least one month), adverse reaction, or contraindication to a preferred formulation of calcipotriene.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
	Duobrii TM (halobetasol propionate/tazarotene) lotion Enstilar® (calcipotriene/betamethasone) foam Sorilux [®] (calcipotriene) foam Taclonex [®] (calcipotriene/betamethasone ointment/scalp suspension) QTY LIMIT: Initial fill = 60 grams Tazarotene Cream, Gel Vtama® (tapinarof) cream Zoryve® (roflumilast) Cream	Methoxsalen, Oxsoralen Ultra: The patient has a documented diagnosis of moderate to severe psoriasis affecting > 10% of the body surface area (BSA) and/or has involvement of the palms, soles, head and neck, or genitalia and has had a documented side effect, allergy, inadequate treatment response, or treatment failure to at least 2 topical agents and at least 1 oral systemic agent, unless otherwise contraindicated. Limitations: Kits with non-drug or combinations of 2 drug products are not covered.
	PULMONARY AGENT	S
ANTICOLINERGICS: INHALED		
SHORT-ACTING BRONCHODILATORS ATROVENT HFA® (ipratropium) COMBIVENT® RESPIMAT (ipratropium/albuterol) QTY LIMIT: 3 inhalers (12 grams)/90 days IPRATROPIUM NEBULIZER SOLN IPRATROPIUM/ALBUTEROL NEBULIZER SOLN LONG-ACTING BRONCHODILATORS (LAMA) INCRUSE ELLIPTA® (umeclidinium bromide) QTY LIMIT: 1 inhaler/30 days SPIRIVA® HANDIHALER (tiotropium) QTY LIMIT: 1 capsule/day SPIRIVA® RESPIMAT (tiotropium) QTY LIMIT: 3 inhalers/90 days	Lonhala® Magnair (glycopyrollate) inhalation solution <i>QTY LIMIT</i> : 60 vials/30 days Tudorza® Pressair® (aclidinium bromide) <i>QTY LIMIT</i> : 3 inhalers/90 days Yupelri TM (revefenacin) inhalation solution <i>QTY LIMIT</i> : 300 vials/30 days	 Tudorza: The patient has had documented side effect, allergy or treatment failure with a preferred LAMA. Bevespi Aerosphere, Duaklir Pressiar: The patient has a documented side effect, allergy, or treatment failure to TWO preferred LAMA/LABA combinations. Lonhala Magnair, Yupelri: patient has a diagnosis of COPD (not FDA approved for asthma) AND has a failure of nebulized ipratropium solution AND at least 3 inhaled LAMAs. Breztri: patient has a diagnosis of COPD (not FDA approved for asthma) AND patient has a treatment failure of at least 2 different combinations of a preferred Inhaled Corticosteroid, LABA, and LAMA used in combination for a minimum of 30 consecutive days AND patient has a documented side effect, allergy, treatment failure, or contraindication with Trelegy Ellipta. Trelegy Ellipta: patient has a treatment failure of at least 2 different combinations of a preferred Inhaled Corticosteroid, LABA, and LAMA used in combination for a minimum of 30 consecutive days.
COMBINATION LONG-ACTING BRONCHODILATORS (LAMA & LABA) ANORO® ELLIPTA (umeclidinium/vilanterol) QTY LIMIT: 3 inhalers (180 blisters)/90 days STIOLTO® RESPIMAT (tiotropium/olodaterol) QTY LIMIT: 3 inhalers/90 days	Bevespi Aerosphere® (glycopyrrolate/formoterol) <i>QTY LIMIT:</i> 3 inhalers/90 days Duaklir® Pressair (aclidinium bromide/ formoterol fumarate) QTY LIMIT: 3 inhalers/90 days	
LAMA/LABA/ICS COMBINATION	Breztri® Aerosphere	

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
(No FA required unless otherwise noted)	(FA required)	FACRITERIA
All products require PA	(budesonide/glycopyrrolate/formoterol fumarate) QTY LIMIT: 1 inhaler (120 blisters)/30 days Trelegy® Ellipta (fluticasone/umeclidinium/vilanterol) QTY LIMIT: 1 inhaler (60 blisters)/30 days	
ANTIHISTAMINES: INTRANASAL		
SINGLE AGENT		
AZELASTINE 0.1% Nasal Spray <i>QTY LIMIT:</i> 1 bottle (30 ml)/25 days	Azelastine 0.15 % Nasal Spray <i>QTY LIMIT</i> : 1 bottle (30 ml)/25 days Olopatadine 0.6% (compare to Patanase®) Nasal Spray <i>QTY LIMIT</i> : 1 bottle (31 gm)/30 days Patanase® (olopatadine 0.6%) Nasal Spray <i>QTY LIMIT</i> : 1 bottle (31 gm)/30 days	Azelastine 0.15%, Olopatadine, Patanase: The patient has a documented side effect, allergy, or treatment failure to Azelastine 0.1% Azelastine/Fluticasone, Ryaltris: The patient has a documented side effect, allergy, or treatment failure to azelastine 0.1% AND The patient has a documented side effect, allergy, or treatment failure to a preferred nasal corticosteroid OR the patient has a documented intolerance to Dymista.
COMBO WITH CORTICOSTEROID DYMISTA® (azelastine/fluticasone) Nasal Spray QTY LIMIT: 1 bottle (23 gm)/30 days	Azelastine/fluticasone (compare to Dymista®) Nasal Spray QTY LIMIT: 1 bottle (23 gm)/30 days Ryaltris® (olopatadine/mometasone)	
CVI.	QTY LIMIT: 1 bottle (29 gm)/30days	
ANTIHISTAMINES: 1 ST GENERATION		
All generic antihistamines All generic antihistamine/decongestant combinations	All brand antihistamines (example: Benadryl [®]) All brand antihistamine/decongestant combinations (example: Deconamine SR [®] , Rynatan [®] , Ryna-12 [®])	CRITERIA FOR APPROVAL: The prescriber must provide a clinically valid reason for the use of the requested medication including reasons why any of the generically available products would not be a suitable alternative.
ANTIHISTAMINES: 2 ND GENERATION		
SINGLE AGENT TABLET CETIRIZINE OTC 5 mg, 10 mg tablets FEXOFENADINE tablets LEVOCETIRIZINE 5 mg tablets LORATADINE	Clarinex [®] (desloratadine) 5 mg tablet Desloratadine (compare to Clarinex [®]) 5 mg tablet All other brands	Clarinex tablets, Desloratadine tablets: The patient has had a documented side effect, allergy, or treatment failure to 2 preferred second generation antihistamines, at least one of which must be loratadine AND If the request is for Clarinex, the patient must also have a documented intolerance to the generic equivalent tablets.
COMBINATION WITH PSEUDOEPHEDRINE CETIRIZINE/PSEUDOEPHEDRINE SR 12hr 5 mg/120 mg FEXOFENADINE/PSEUDOEPHEDRINE SR 12hr 60mg/120 mg FEXOFENADINE/PSEUDOEPHEDRINE SR 24hr 180mg/240 mg LORATADINE/PSEUDOEPHEDRINE SR 12hr 5 mg/120 mg LORATADINE/PSEUDOEPHEDRINE SR 24hr 10	Clarinex-D [®] 12 hr (desloratadine/pseudoephedrine 2.5 mg/120 mg)	Cetirizine chewable tablets, Desloratadine ODT: The patient has had a documented side effect, allergy, or treatment failure to cetirizine oral solution and one of the following loratadine formulations: chewable tablet, rapidly disintegrating tablet, or oral solution. Fexofenadine suspension, Levocetirizine solution: the patient has had a documented side effect, allergy, or treatment failure to loratadine syrup AND cetirizine syrup. Clarinex-D: The patient has had a documented side effect, allergy, or treatment failure to loratadine-D and cetirizine-D. LIMITATIONS: Many Allegra® and Zyrtec® brand products as well as Claritin capsules are not covered as no Federal Rebate is offered.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
(110 111 required amess outer wise noted)	(FITTEQuined)	TH CHILDREN
mg/240 mg		
SINGLE AGENT ORAL LIQUID CETIRIZINE syrup	Fexofenadine (compare to Allegra®) suspension Levocetirizine Solution	
LORATADINÉ syrup		
CHEWABLE/ORALLY DISINTEGRATING TABLET LORATADINE chewable tablet 5mg LORATADINE rapidly disintegrating tablet (RDT)	Certirizine OTC Chewable Tablets 5 mg, 10 mg Desloratadine ODT (compare to Clarinex Reditabs [®]) 2.5 mg, 5 mg	
10 mg	All other brands	
BETA-ADRENERGIC AGENTS		
DEIN-ADREMENTO AGENTO		
METERED-DOSE INHALERS (SHORT-ACTING) PROAIR® Respiclick (albuterol) PROVENTIL® HFA (albuterol) VENTOLIN® HFA (albuterol)	Albuterol HFA (compare to Proventil® HFA, ProAir® HFA, Ventolin® HFA) Levalbuterol Aerosol (compare to Xopenex ® HFA) ProAir® Digihaler (albuterol)	Albuterol HFA, Levalbuterol (aerosol), Xopenex HFA: patient has a documented side effect, allergy, or treatment failure to two preferred short acting metered dose inhalers. AND for approval of levalbuterol aerosol, the patient must have a documented intolerance to brand Xopenex HFA. ProAir Digihaler: Preferred albuterol metered dose inhalers and Xopenex HFA are on a long-term backorder and unavailable from the manufacturer
METERED-DOSE INHALERS (LONG-ACTING)	Xopenex [®] HFA (levalbuterol)	Serevent: The patient has a diagnosis of asthma and is prescribed an inhaled corticosteroid (pharmacy claims will be evaluated to assess compliance with long term controller therapy) OR the patient has a diagnosis of COPD.Striverdi: The patient has a diagnosis of COPD (not FDA approved for asthma).
Preferred After Clinical Criteria Are Met SEREVENT® DISKUS (salmeterol xinafoate) QTY LIMIT: 1 inhaler (60 blisters)/30 days	Strivenski Degnimet® (eledetenel)	AND The patient has a documented side effect, allergy, or treatment failure to Serevent. Levalbuterol, Xopenex nebulizer solution (age > 12 years): The patient must
NEBULIZER SOLUTIONS (SHORT-ACTING)	Striverdi Respimat® (olodaterol)	have had a documented side effect, allergy, or treatment failure to albuterol nebulizer. AND for approval of brand Xopenex, the patient must have had a documented intolerance to the generic.
ALBUTEROL neb solution (all strengths) LEVALBUTEROL neb solution (age ≤ 12 years)	Levalbuterol neb solution (compare to Xopenex [®]) (age > 12 years)	Xopenex (age <12 years): The patient must have a documented intolerance to generic levalbuterol nebulizer solution
NEBULIZER SOLUTIONS (LONG-ACTING)	Xopenex [®] neb solution (all ages)	Arformoterol, Brovana, Formoterol, Perforomist Nebulizer Solution: The
All products require PA		patient must have a diagnosis of COPD. AND The patient must be unable to use a non-nebulized long-acting bronchodilator or anticholinergic (Serevent or
	Arformoterol (compare to Brovana®)	Spiriva) due to a physical limitation
	QTY LIMIT: 2 vials/day	Terbutaline tablets: The medication is not being prescribed for the

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
TABLETS/SYRUP (SHORT-ACTING) ALBUTEROL tablets/syrup TABLETS (LONG-ACTING) ALBUTEROL ER tablets	Brovana® (arformoterol) QTY LIMIT: 2 vials/day Formoterol (compare to Perforomist®) QTY LIMIT: 2 vials/day Perforomist® (formoterol) QTY LIMIT: 2 vials/day Terbutaline tablets	prevention/treatment of preterm labor.
CORTICOSTEROIDS/COMBINATIONS: INHA	LED	
METERED DOSE INHALERS (SINGLE AGENT) ASMANEX® (mometasone furoate) QTY LIMIT: 3 inhalers/90 days FLOVENT® DISKUS (fluticasone propionate) QTY LIMIT: 3 inhalers/90 days FLOVENT® HFA (fluticasone propionate) QTY LIMIT: 3 inhalers (36 gm)/90 days PULMICORT FLEXHALER® (budesonide) QTY LIMIT: 6 inhalers/90 days	Armonair® Digihaler (fluticasone propionate) QTY LIMIT = 3 inhalers/90 days Alvesco® (ciclesonide) QTY LIMIT: 80 mcg = 3 inhalers/90 days Arnuity Ellipta 100 or 200 mcg/inh (fluticasone furoate) QTY LIMIT: 90 blisters/90 days Asmanex® (mometasone furoate) HFA QTY LIMIT: 3 inhalers (39 gm)/90 days Fluticasone propionate HFA (compare to Flovent® HFA) QTY LIMIT: 3 inhalers (36 gm)/90 days Qvar® Redihaler™ 40mcg/inh QTY LIMIT: 2 inhalers (21.2 gm)/90 days Qvar® Redihaler™ 80mcg/inh QTY LIMIT: 3 inhalers (31.8 gm)/90 days	 Metered-dose inhalers (single agent): The patient has had a documented side effect, allergy, or treatment failure to at least two preferred agents AND for approval of Asmanex HFA, there must be a clinically compelling reason the patient is unable to use Asmanex. Advair HFA (age < 12 years): The patient has had a documented side effect, allergy, or treatment failure to Dulera or Symbicort. AirDuo Digihaler, AirDuo Respiclick, Breo Ellipta, Fluticasone/Salmeterol (non-authorized generics): The patient has had a documented side effect, allergy, or treatment failure to any 2 of the following: Advair HFA, Advair Diskus, Dulera, or Symbicort. Budesonide/formoterol: the patient has a documented intolerance to brand Symbicort. Budesonide Inh Suspension: Medical necessity for the use of a nebulized solution has been provided AND if the dose is 1mg, the patient must be unable to use two 0.5 mg vials Fluticasone/salmeterol powder (authorized generic), Wixela Inhub: A clinically compelling reason must be provided detailing why the patient is unable to use Advair HFA or Advair Diskus.
METERED DOSE INHALERS (COMBINATION PRODUCT) ADVAIR® DISKUS (fluticasone/salmeterol) (Age ≥ 4 years) QTY LIMIT: 3 inhalers/90 days ADVAIR® HFA (fluticasone/salmeterol) (Age ≥ 12 years) QTY LIMIT: 3 inhalers (36 gm)/90 days DULERA® (mometasone/formoterol) QTY LIMIT: 9 inhalers (39 gm)/90 days SYMBICORT® (budesonide/formoterol) QTY LIMIT: 9 inhalers (91.8gm)/90 days	AirDuo® Digihaler (fluticasone/salmeterol) QTY LIMIT: 3 inhalers/90 days AirDuo Respiclick® (fluticasone/salmeterol) QTY LIMIT: 3 inhalers/90 days Breo Ellipta® (fluticasone furoate/vilanterol) QTY LIMIT: 3 inhalers (180 blisters) 90 days Budesonide/formoterol (compare to Symbicort®) QTY LIMIT: 9 inhalers (91.8gm)/90 days Fluticasone/salmeterol (compare to AirDuo Respiclick®) QTY LIMIT: 3 inhalers/90 days Fluticasone/salmeterol inhalation Powder (compare to	Pulmicort Respules: medical necessity for the use of a nebulized solution has been provided AND if the dose is 1 mg, the patient must be unable to use two 0.5 mg vials AND the patient has a documented intolerance to the generic.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
NEBULIZER SOLUTIONS BUDESONIDE INH SUSPENSION 0.25mg, 0.5mg (Age ≤ 12 yrs)	Advair® Diskus) QTY LIMIT: 3 inhalers/90 days Wixela TM Inhub TM (fluticasone/salmeterol inhalation powder) (compare to Advair® Diskus) QTY LIMIT: 3 inhalers/90 days Budesonide Inh Suspension 1mg (all ages), 0.25mg and 0.5mg (age >12 years) Pulmicort Respules® (budesonide)	
	Pulmicort Respules (budesonide)	
CORTICOSTEROIDS: INTRANASAL		
BUDESONIDE QTY LIMIT: 1 inhaler (8.43 ml)/30 days FLUTICASONE PROPIONATE QTY LIMIT: 1 inhaler (16 gm)/30 days OMNARIS® (ciclesonide) QTY LIMIT: 1 inhaler (12.5 gm)/30 days TRIAMCINOLONE QTY LIMIT: 1 inhaler (16.9 ml)/30 days ZETONNA® (ciclesonide) QTY LIMIT: 1 inhaler (6.1 gm)/30 days	Beconase AQ [®] (beclomethasone) QTY LIMIT: 2 inhalers (50 gm)/30 days Flunisolide 25 mcg/spray QTY LIMIT: 2 inhalers (50 ml)/30 days Mometasone (compare to Nasonex [®]) QTY LIMIT: 1 inhaler (17 gm)/30 days QNASL [®] (beclomethasone dipropionate) QTY LIMIT: 1 inhaler (10.6 gm)/30 days Xhance TM (fluticasone propionate) QTY LIMIT: 1 inhaler (16 ml)/30 days	 Beconase AQ, Flunisolide 25 mcg/spray, Mometasone, QNASL: The patient has had a documented side effect, allergy, or treatment failure of two preferred nasal glucocorticoids. If a product has an AB rated generic, one trial must be the generic. Xhance: The patient has had a documented side effect, allergy, or treatment failure of three preferred nasal glucocorticoids, one of which must be fluticasone. Limitations: Nasacort Allergy OTC and Flonase are not covered as no Federal Rebate is offered.
LEUKOTRIENE MODIFIERS		
Preferred After Age Criteria Are Met MONTELUKAST SODIUM (compare to Singulair®) tablets, 10mg for ages ≥ 15 MONTELUKAST SODIUM (compare to Singulair®) chews, 4 mg for ages 2-5, 5 mg for age 6-14 MONTELUKAST SODIUM (compare to Singulair®) granules, ages 6 months-23 months	Accolate [®] (zafirlukast) QTY LIMIT: 2 tablets/day Singulair [®] (montelukast sodium) tablets, chew tabs, granules QTY LIMIT: 1 tablet or packet per day Zafirlukast (compare to Accolate [®]) Zileuton ER (compare to Zyflo CR®) QTY LIMIT: 4 tablets/day Zyflo (zileuton) QTY LIMIT: 4 tablets/day	 Montelukast: Clinical rationale must be provided for prescribing a dose and formulation that differs from age recommendations AND If the request is for brand Singulair, the patient has a documented intolerance to the generic equivalent montelukast preparation. Zafirlukast, Accolate: The diagnosis or indication for the requested medication is asthma. AND If the request is for Accolate, the patient has a documented intolerance to generic zafirlukast. Zileuton ER, Zyflo: The diagnosis or indication for the requested medication is asthma. AND The patient has had a documented side effect, allergy, or treatment failure to Accolate/Zafirlukast or Singulair/Montelukast
PHOSPHODIESTERASE-4 (PDE-4) INH	BITORS	
All products require PA	Daliresp® tablet (roflumilast) QTY LIMIT: 1 tablet/day	Daliresp: The indication for the requested medication is treatment to reduce the risk of COPD exacerbations in patients with severe COPD associated with

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
	* Maximum days' supply per fill = 30 *	chronic bronchitis and a history of exacerbations. AND The patient has had a documented side effect, allergy, treatment failure, or a contraindication to at least one inhaled long-acting anticholinergic AND at least one inhaled long-acting beta-agonist AND at least one inhaled corticosteroid.
SYNAGIS		
	SYNAGIS® (palivizumab) QTY LIMIT: 50 mg = 1 vial/month, 100 mg = 2 vials/month	 CRITERIA FOR APPROVAL: Infants born at 28 weeks of gestation or earlier (i.e., ≤ 28 weeks, 6 days) and under twelve months of age at the start of the RSV season (maximum 5 doses). Infants born at 29-32 weeks (i.e., between 29 weeks, 0 days and 31 weeks, 6 days) of gestation and under 1 year of age at the start of the RSV season who develop chronic lung disease of prematurity defined as a requirement for >21% oxygen for at least the first 28 days after birth (maximum 5 doses). Children under 24 months of age with chronic lung disease of prematurity defined as born at 31 weeks, 6 days or less who required >21% oxygen for at least the first 28 days after birth and continue to require medical support (chronic corticosteroid therapy, diuretic therapy, or supplemental oxygen) during the 6-month period before the start of the second RSV season (maximum 5 doses). Children under 12 months of age with hemodynamically significant congenital heart disease (CHD) (dosing continues in the RSV season through the end of the month the infant reaches 12 months old maximum 5 doses): Acyanotic heart disease and receiving medication to control congestive heart failure and will require cardiac surgical procedures, Moderate to severe pulmonary hypertension, Cyanotic heart disease and recommended for Synagis therapy by Pediatric Cardiologist Infants under 12 months of age with either: (dosing continues in the RSV season through the end of the month the infant reaches 12 months old -maximum 5 doses) Congenital abnormalities of the airways that impairs the ability to clear secretions from the upper airway because of ineffective cough, Neuromuscular condition that impairs the ability to clear secretions from the upper airway because of ineffective cough Infants and children less than 24 months of age who are profoundly immunocompromised during the RSV season Infants and children less than 24 months of age who are profoundly immunocompromised during the RSV season (e.

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
		EXCLUDED FROM APPROVAL: Infants and children with hemodynamically insignificant heart disease. Infants with cardiac lesions adequately corrected by surgery, unless they continue to require medication for congestive heart failure. Infants with mild cardiomyopathy who are not receiving medical therapy. Breakthrough hospitalization for RSV disease (Synagis therapy should be discontinued for the season once hospitalization for RSV has occurred). Infants and children with Down syndrome unless other indications above are present. Infants and children with cystic fibrosis unless other specific conditions are present This drug must be obtained and billed through a DVHA enrolled specialty pharmacy and processed through the DVHA POS prescription processing system using NDC values. Under no circumstances will claims processed through the medical benefit be accepted.
	PULMONARY ARTERIAL HYPERT	ENSION MEDICATIONS
ENDOTHELIN RECEPTOR ANTAGONISTS AMBRISENTAN (compare to Letairis®) QTY LIMIT: 1 tablet/day BOSENTAN (compare to Tracleer) QTY LIMIT: 2 tablets/day	Letairis® (ambrisentan) Tablet QTY LIMIT: 1 tablet/day Opsumit® (macitentan) Tablet QTY LIMIT: 1 tablet/day	Adempas: The patient has a diagnosis of pulmonary arterial hypertension (PAH) with New York Heart Association (NYHA) Functional Class II or III. OR The patient has a diagnosis of chronic thromboembolic pulmonary hypertension (CTEPH, WHO Group 4) AND the patient has persistent or recurrent disease after surgical treatment (e.g., pulmonary endarterectomy) or has CTEPH that

PROSTACYCLIN AGONISTS INJECTION

EPOPROSTENOL (compare to Flolan®) $\begin{array}{l} REMODULIN^{\textcircled{R}} \ (treprostinil \ sodium \ injection) \\ VELETRI^{\textcircled{R}} \ (epoprostinil) \end{array}$

INHALATION

All products require PA

Tracleer[®] tablets for oral suspension (32 mg) Tracleer® (bosentan) tablet (62.5 mg, 125 mg)

QTY LIMIT: 2 tablets/day

Flolan® (epoprostenol) Treprostinil sodium injection (compare to Remodulin®)

Tyvaso® (Treprostinil) inhalation solution Tyvaso® DPI (treprostinil) powder for inhalation Ventavis® (iloprost) inhalation solution

is inoperable AND The patient is 18 years of age or older AND The patient will not use Adempas concomitantly with the following: Nitrates or nitric oxide donors (such as amyl nitrate) in any form. Phosphodiesterase (PDE) inhibitors, including specific PDE-5 inhibitors (such as sildenafil, tadalafil, or vardenafil) or non-specific PDE inhibitors (such as dipyridamole or theophylline) AND The patient is not pregnant AND Female patients are enrolled in the Adempas REMS Program

Flolan, Letairis, Tracleer: patient has a documented intolerance to the generic equivalent.

Tracleer tablets for oral suspension: Patient has a diagnosis of PAH with NYHA Functional Class II or III AND patient is ≤ 12 years of age and <40kg. **Opsumit:** Patient has a diagnosis of PAH with NYHA Functional Class II or III

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
ORAL ORENITRAM® (treprostinil) ER Tablet SGC STIMULATOR All products require PA **Maximum days supply for all drugs is 30 days**	Uptravi® (selexipag) tablets QTY LIMIT: 200 mcg = 140 tablets/30 days for the first 2 months, then 2 tablets/day thereafter All other strengths = 2 tablets/day Adempas® (riociguat) Tablets QTY LIMIT: 3 tablets/day	AND Patient is not pregnant AND Female patients have been enrolled in the REMS Program AND the patient has a documented side effect, allergy, or treatment failure with Tracleer or Letairis. Treprostinil: Patient has a diagnosis of pulmonary arterial hypertension AND The patient has had a documented intolerance to the brand Remodulin. Tyvaso, Ventavis: The patient has a diagnosis of pulmonary arterial hypertension (PAH) with New York Heart Association (NYHA) Functional Class II or III heart failure AND the patient is unable to tolerate or has failed 2 different preferred medications. Uptravi: The patient has a diagnosis of pulmonary arterial hypertension (PAH) with New York Heart Association (NYHA) Functional Class II or III heart failure AND the patient is unable to tolerate or has failed 2 different preferred medications, one of which must be Orenitram
PHOSPHODIESTERASE-5 (PDE-5) INHIBITORS Effective 7/1/06, phosphodiesterase-5 (PDE-5) inhibitors are no longer a covered benefit for all Vermont Pharmacy Programs for the treatment of erectile dysfunction. This change is resultant from changes set into effect January 1, 2006 and as detailed in Section 1903 (i)(21)(K) of the Social Security Act (the Act), precluding Medicaid Federal Funding for outpatient drugs used for the treatment of sexual or erectile dysfunction. Sildenafil will remain available for coverage via prior-authorization for the treatment of Pulmonary Arterial Hypertension.		

Preferred After Clinical Criteria Are Met

SILDENAFIL CITRATE (compare to Revatio®) tablet

QTY LIMIT: 3 tablets/day TADALAFIL (compare to Adcirca®)

QTY LIMIT: 2 tablets/day

Adcirca® (tadalafil)

QTY LIMIT: 2 tablets/day

Revatio® (sildenafil) tabs OTY LIMIT: 3 tablets/day

Revatio® (sildenafil citrate) suspension

Revatio® (sildenafil citrate) vial

QTY LIMIT: 3 vials/day

Maximum 14-day supply per fill

Sildenafil (compare to Revatio ®) suspension

Sildenafil (compare to Revatio®) vial

Tadliq® (tadalafil) suspension

Sildenafil, Tadalafil: Clinical Diagnosis of Pulmonary Hypertension
 Adcirca (tadalafil) 20 mg, Revatio (sildenafil citrate) 20 mg: Clinical diagnosis of pulmonary hypertension AND No concomitant use of organic

nitrate-containing products AND patient has a documented intolerance to the generic equivalent.

Revatio Suspension, Sildenafil Suspension: Clinical diagnosis of pulmonary hypertension AND medical necessity for a liquid formulation is provided OR the patient is unable to tolerate a 20 mg dose AND for approval of Revatio, the patient must have a documented intolerance to the generic equivalent.

Revatio IV, Sildenafil IV: Clinical diagnosis of pulmonary hypertension AND No concomitant use of organic nitrate-containing products AND The patient has a requirement for an injectable dosage form. AND Arrangements have been made for IV bolus administration outside of an inpatient hospital setting.

Tadliq: Clinical diagnosis of pulmonary hypertension AND medical necessity for a liquid formulation is provided AND the patient has a documented side effect, allergy, or treatment failure with sildenafil suspension.

PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
RENAL DISEASE: PHOSPHATE BINDERS		
CALCIUM ACETATE capsule	Auryxia® (ferric citrate) <i>QTY LIMIT</i> : 12/day	Renvela Oral Suspension Packet, Sevelamer Packet: The patient has a requirement for a liquid dosage form.

CALCIUM ACETATE tablet SEVELAMER CARBONATE (compare to Renvela®) tablets

ORAL SOLUTIONS

PHOSLYRA® (calcium acetate) oral solution

Fosrenol® (lanthanum carbonate) Lanthanum carbonate (compare to Fosrenol) Renagel® (sevelamer) Renvela® (sevelamer carbonate) Oral Suspension Packet

QTY LIMIT: 0.8 g = 2 packs/dayRenvela® (sevelamer carbonate) tablets Sevelamer carbonate Oral Suspension Packet (compare to Renvela®)

QTY LIMIT: 0.8 g = 2 packs/daySevelamer hydrochloride (compare to Renagel®) Velphoro® (sucroferric oxyhydroxide) Chew Tablet Auryxia, lanthanum carbonate, Renagel, Renvela tablets, sevelamer hydrochloride tablets, Velphoro Chew Tablet: The patient must have a documented side effect, allergy, or inadequate response to one preferred phosphate binder.

RESTLESS LEG SYNDROME MEDICATIONS

DOPAMINE AGONISTS (ORAL)

PRAMIPEXOLE (compare to Mirapex[®]) ROPINIROLE (compare to Requip[®])

DOPAMINE AGONISTS (TRANSDERMAL) All products require PA

GAMMA-AMINOBUTYRIC ACID ANALOG GABAPENTIN IR

Mirapex® (pramipexole)

Neupro® (rotigotine) transdermal patch QTY LIMIT: 1, 2, and 3 mg ONLY = 1 patch/day

Horizant® (gabapentin enacarbil) ER Tablet OTY LIMIT: 1 tablet/day

Mirapex: The patient has had a documented intolerance to the generic product. **Neupro:** The patient has a medical necessity for a specialty dosage form. Horizant: The patient has a diagnosis of restless legs syndrome (RLS). AND The patient has had a documented side effect, allergy, contraindication or treatment failure to two preferred dopamine agonists AND gabapentin IR. Limitations: Requests for Mirapex ER and Requip XL will not be approved for Restless Leg Syndrome (RLS).

RHEUMATOID, JUVENILE & PSORIATIC ARTHRITIS: IMMUNOMODULATORS

Preferred After Clinical Criteria Are Met INJECTABLE

AVSOLA® (infliximab-axxq) biosimilar to Remicade®

ENBREL® (etanercept) QTY LIMIT: 50 mg = 4 syringes/28 days, 25 mg = 8 syringes/28 days INFLECTRA® (infliximab-dyyb) biosimilar

Actemra® (tocilizumab) Intravenous Infusion QTY LIMIT: 80 mg vial = 4 vials/28 days, 200 mg vial = 3 vials/28 days, 400 mg vial = 2 vials/28 daysActemra® (tocilizumab) Subcutaneous Prefilled Syringe QTY LIMIT: 4 prefilled syringes (3.6ml)/28 days Actemra® (tocilizumab) ACTPen OTY LIMIT: 4 pens (3.6ml)/28 days

Clinical Criteria for all drugs: Patient has a diagnosis of rheumatoid arthritis (RA), juvenile idiopathic arthritis* or psoriatic arthritis and has already been stabilized on the drug being requested OR Diagnosis is RA, juvenile idiopathic arthritis or psoriatic arthritis, and methotrexate therapy resulted in an adverse effect, allergic reaction, inadequate response, or treatment failure. If methotrexate is contraindicated, another DMARD should be tried prior to approving therapy. Other DMARDs include leflunomide, sulfasalazine, gold, antimalarials, minocycline, D-penicillamine, azathioprine, cyclophosphamide

PREFERRED AGENTS **NON-PREFERRED AGENTS** (No PA required unless otherwise noted) (PA required) PA CRITERIA to Remicade® Cimzia[®] (certolizumab pegol) and cyclosporine *QTY LIMIT:* 1 kit/28 days KINERET® (anakinra) **Taltz, Xeljanz, Xeljanz XR additional criteria:** patient must be ≥ 18 years of Cosentyx® (secukinumab) QTY LIMIT: 1 syringe/day age AND the prescriber must provide evidence of a trial and failure or Kevzara® (sarilumab) contraindication to a preferred TNF Inhibitor. Note: Xeljanz 10mg BID and HUMIRA® (adalimumab) QTY LIMIT: 2 syringes/28 days XR 22mg are NOT recommended for Rheumatoid Arthritis or Psoriatic OTY LIMIT: 4 syringes/28 days Ilaris® (canakinumab) Orencia® (abatacept) Subcutaneous Injection TALTZ® (ixekizumab) Arthritis. Please refer to Gastrointestinal: Inflammatory Bowel Disease QTY_LIMIT: 4 syringes/28 days Biologics for Ulcerative Colitis criteria. QTY LIMIT: 80 mg prefilled syringe or Orencia (abatacept) Intravenous Infusion Remicade (infliximab) Renflexis (Infliximab-abda) biosimilar to Remicade Actemra, Cimzia, Cosentyx, Kevzara, Orencia, Simponi (subcutaneous), autoinjector = 2/28 days for the first month Skyrizi, Stelara, and Tremfya additional criteria: The prescriber must and 1/28 days subsequently Simponi[®] (golimumab) Subcutaneous provide clinically valid reason why at least 2 preferred agents cannot be used. *QTY LIMIT:* 50 mg = 1 prefilled syringe or **Ilaris:** The diagnosis is systemic juvenile idiopathic arthritis (sJIA) with active autoinjector/28 days systemic features and varying degrees of synovitis with continued disease Simponi Aria^(R) (golimumab) 50 mg/4 ml Vial for activity after initial therapy (initial therapy defined as 1 month of anakinra Intravenous Infusion (Kineret), 2 weeks of glucocorticoid monotherapy (oral or IV) or one month SkyriziTM (risankizumab-rzaa) of NSAIDs). AND patient is > 2 years of age. QTY LIMIT: 150 mg/28 days for the **Remicade, Renflexis additional criteria:** The prescriber must provide a first month and 150mg/84 days clinically valid reason why at least 2 preferred agents cannot be used AND the thereafter patient must be unable to use Avsola or Inflectra. Stelara[®] (ustekinumab) **Simponi Aria additional criteria:** The patient has not responded adequately to *OTY LIMIT*: 45 mg (0.5 ml) or 90 mg (1 ml) per dose Simponi subcutaneous. AND The prescriber must provide a clinically valid (90 mg dose only permitted for pt weight > 100 kg) One reason why at least 2 preferred agents cannot be used. dose/28 days for the first month and one dose/84 Olumiant, Rinvoq additional criteria: The patient must be ≥ 18 years of age AND days thereafter The prescriber must provide a clinically valid reason why at least two preferred Tremfya® (guselkumab) agents cannot be used, one of which must be Xeljanz or Xeljanz XR. OTY LIMIT: 1 syringe/28 days for the Note: Patients with systemic juvenile arthritis (SJRA/SJIA) and fever are not first month, then 1 syringe every 56 required to have a trial of a DMARD, including methotrexate. Patients with ORAL days thereafter systemic juvenile arthritis without fever should have a trial of methotrexate, OTEZLA® tablet (apremilast) but a trial of another DMARD in the case of a contraindication to OTY LIMIT: Starter Pack = 55 tablets/28 methotrexate is not required. * Patients with psoriatic arthritis with a days, 30 mg = 2 tablets/dayOlumiant® (baricitinib) tablets documented diagnosis of active axial involvement should have a trial of Maximum 30 days supply OTY LIMIT: 1 tablet/day NSAID therapy, but a trial with DMARD is not required before a TNF-Maximum 30 days supply XELJANZ® (tofacitinib) 5 mg tablet Rinvoq ® (upadactinib) extended release tablet blocker is approved. If no active axial skeletal involvement, then an NSAID OTY LIMIT: 2 tablets/day OTY LIMIT: 1 tablet/day trial and a DMARD trial are required (unless otherwise contraindicated). Maximum 30 days supply Maximum 30 days supply XELJANZ® XR (tofacitinib) tablet

QTY LIMIT: 1 tablet/day

SICKLE CELL DISEASE THERAPIES

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
DROXIA® (hydroxyurea) 200 mg, 300 mg, 400 mg cap HYDROXYUREA (compare to Hydrea®) 500 mg cap	Adakveo® (crizanlizumab-tmca) Endari (L-glutamine powder for oral solution) QTY LIMIT: maximum of 30-day supply Hydrea® (hydroxyurea) 500 mg cap Oxbryta® (voxelotor) 500 mg tablet QTY LIMIT: 3 tablets/day Oxbryta® 300mg tablets for oral suspension Siklos® (hydroxyurea) 100 mg, 1000 mg tablet	Adakveo: Patient has a diagnosis of Sickle Cell Disease AND patient is at least 16 years of age or older AND patient has had an inadequate response to a 6-month trial of hydroxyurea dosed at 15-35 mg/kg/day, unless contraindicated AND patient has experienced at least 2 vaso-occlusive crises in the previous 12 months despite compliance with hydroxyurea. Initial approval will be granted for 6 months. For re-approval, the patient must have a decrease in the frequency or severity of VOC compared to baseline. Note: Adakveo will not be approved in conjunction with Oxbryta. Endari: Indication for use is to reduce the acute complications of Sickle Cell Anemia AND medication will be approved with quantity limits based on patient weight (<30kg = 2 packets/day, 30-65kg = 4 packets/day, > 65kg = 6 packets/day). Hydrea: Patient has had a documented intolerance to the generic equivalent. Oxbryta: Patient has a diagnosis of Sickle Cell Disease AND patient is at least 4 years of age or older AND patient has a baseline hemoglobin (Hb) ≤10.5 g/dL AND patient has had an inadequate response to a 6-month trial of hydroxyurea dosed at 15-35 mg/kg/day, unless contraindicated AND patient has experienced at least 2 vaso-occlusive crises in the previous 12 months despite compliance with hydroxyurea. Initial approval will be granted for 6 months. For re-approval, the patient must have a decrease in the frequency or severity of VOC compared to baseline. Note: Oxbryta will not be approved in conjunction with Adakveo. Siklos: Patient has a diagnosis of Sickle Cell Anemia with recurrent moderate to severe pain crises AND the required dose is < 200mg OR Patient has a diagnosis of Sickle Cell Anemia with recurrent moderate to severe pain crises AND the a documented docrease in vaso-occlusive episodes, acute chest syndrome, SCD related hospitalizations, or blood transfusions.
	SEDATIVE/HYPNOTIC	cs control of the second of th
BENZODIAZEPINE		
TEMAZEPAM 7.5mg, 15 mg, 30 mg (compare to Restoril [®])	Estazolam Flurazepam Halcion [®] (triazolam) Restoril [®] (temazepam) Temazepam 22.5 mg (compare to Restoril [®]) Triazolam (compare to Halcion [®])	Criteria for Approval: The patient has had a documented side effect, allergy, or treatment failure with Temazepam. If a product has an AB rated generic, one trial must be the generic.
NON BENZODIAZEPINE, NON BARBITURATE		
ESZOPICLONE (compare to Lunesta) QTY LIMIT: 1 tab/day	Ambien [®] (zolpidem) <i>QTY LIMIT</i> : 1 tab/day	Ambien, Ambien CR, Lunesta: The patient has had a documented intolerance to the generic equivalent.

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
ZALEPLON QTY LIMIT: 5 mg = 1 cap/day, 10 mg = 2 caps/day ZOLPIDEM (compare to Ambien®) QTY LIMIT: 1 tab/day ZOLPIDEM CR (compare to Ambien CR®) QTY LIMIT: 1 tab/day	Ambien CR [®] (zolpidem) QTY LIMIT: 1 tab/day Belsomra [®] (suvorexant) QTY LIMIT: 1 tab/day Dayvigo® (lemborexant) tablet QTY LIMIT: 1 tab/day Doxepin 3mg tablets (compare to Silenor) QTY LIMIT: 1 tab/day Edluar [®] (zolpidem) sublingual tablet QTY LIMIT: 1 tab/day Hetlioz® (tasimelteon) 20 mg oral capsule QTY LIMIT: 1 capsule/day Maximum days supply per fill is 30 days Lunesta [®] (eszopiclone) QTY LIMIT: 1 tab/day Quviviq TM (daridorexant) QTY LIMIT: 1 tab/day Ramelteon (compare to Rozerem®) QTY LIMIT: 1 tab/day Rozerem [®] (ramelteon) QTY LIMIT: 1 tab/day Silenor [®] (doxepin) QTY LIMIT: 1 tab/day Zolpidem sublingual tablet QTY LIMIT: 1 tab/day	Belsomra: The patient has had a documented side effect, allergy, or treatment failure to one preferred sedative/hypnotic. Dayvigo, Quviviq: The patient has had a documented side effect, allergy, or treatment failure to two preferred sedative/hypnotics and Belsomra. Edluar, Zolpidem sublingual: The patient has a medical necessity for a disintegrating tablet formulation (i.e. swallowing disorder). Hetlioz: Patient has documentation of Non-24-Hour Sleep-Wake Disorder (Non24) or Insomnia due to Smith-Magenis Syndrome AND Patient has had a documented side effect, allergy or treatment failure with Rozerem and at least one OTC melatonin product. Ramelteon, Rozerem: The patient has had a documented side effect, allergy, contraindication, or treatment failure to one preferred sedative/hypnotic OR the patient has had a treatment failure after a minimum 2-week trial of melatonin. OR There is a question of substance abuse with the patient or family of the patient. If the request is for Ramelteon, there must also have been a documented intolerance to brand Rozerem. Silenor: The patient has had a documented side effect, allergy, contraindication, or treatment failure to two preferred sedative/hypnotics AND The patient has had a documented intolerance with a preferred generic doxepin formulation.
	SMOKING CESSATION	THERAPIES

NICOTINE REPLACEMENT: maximum duration is 16 weeks (2 x 8 weeks)/365 days for non-preferred. For approval of therapy beyond the established maximum duration, the prescriber must provide evidence that the patient is engaged in a smoking cessation counseling program.

NICOTINE GUM NICOTINE LOZENGE NICOTINE PATCH OTC	Nicotrol Inhaler®	Nicotrol Inhaler: The patient has had a documented treatment failure with nicotine patch used in combination with nicotine gum or lozenge.
NICOTROL® (nicotine) NASAL SPRAY		*Smoking Cessation Counseling is encouraged with the use of smoking cessation
		therapies*
		*The combined prescribing of long acting (patch) and faster acting (gum or
ORAL THERAPY		lozenge) nicotine replacement therapy is encouraged for greater likelihood of quit
		success*

DDEEEDDED A CENTS	NON DECEMBED ACENTS	
PREFERRED AGENTS (No PA required unless otherwise noted)	NON-PREFERRED AGENTS (PA required)	PA CRITERIA
BUPROPION SR (compare to Zyban®) CHANTIX® (varenicline) (Limited to 18 years and older) QTY LIMIT: 2 tabs/day Max duration 24 weeks (2x12 weeks)/365 days) VARENICLINE (Limited to 18 years and older) QTY LIMIT: 2 tabs/day Max duration 24 weeks (2x12 weeks)/365 days)		Vermont QUIT LINE (available free to all patients) 1-800-QUIT-NOW (1-800-784-8669) https://802quits.org/ GETQUIT™ Support Plan available free to all Chantix® patients 1-877-CHANTIX (242-6849) https://www.get-quit.com/
	SUBSTANCE USE DISORDER TR	EATMENTS
ALCOHOL USE DISORDER		
ACAMPROSATE DISULFIRAM NALTREXONE VIVITROL® (naltrexone for extended-release injectable suspension) QTY LIMIT: 1 injection (380 mg) per 28 days		
OPIOID USE DISORDER		
NALTREXONE tablet BUPRENORPHINE/NALOXONE TABLET QTY LIMIT: 8 mg = 2 tablets/day, 2mg N/A (Maximum Daily Dose = 16 mg/day, PA required for over 16 mg) SUBOXONE® sublingual FILM (buprenorphine/naloxone) QTY LIMIT: 8 mg = 2 films per day, 4 and 12 mg = 1 film per day, 2mg N/A (Maximum daily Dose = 16 mg/day, PA required for over 16 mg) *Maximum days supply for Suboxone Films, Buprenorphine/naloxone tablets is 30 days* VIVITROL® (naltrexone for extended-release injectable suspension) QTY LIMIT: 1 injection (380 mg) per 28 days	Buprenorphine sublingual TABLET QTY LIMIT: 2 mg = 3 tablets per day, 8 mg = 2 tablets/day Maximum Daily Dose = 16 mg/day Buprenorphine/naloxone (compare to Suboxone®) sublingual FILM QTY LIMIT: 8 mg = 2 films per day, 4 and 12 mg = 1 film per day, 2mg N/A Maximum daily Dose = 16 mg/day Zubsolv® (buprenorphine/naloxone) sublingual tablet QTY LIMIT: 1 tablet per day of all strengths **Maximum days supply for oral buprenorphine/naloxone films or buprenorphine is 30 days** Sublocade® (buprenorphine extended-release) injection QTY LIMIT: Maximum 30-day supply	CLINICAL CONSIDERATIONS: These products are not FDA approved for alleviation of pain. For this indication, please refer to the Opioid Analgesics PDL category. Note: As of 1/1/23, a completed Buprenorphine safety checklist must be submitted with all PA requests. Buprenorphine/naloxone films, Zubsolv: Clinical justification explaining why the member cannot use the preferred buprenorphine formulations has been provided AND the Buprenorphine Safety Checklist has been completed (see PA form for detailed requirements). Buprenorphine: Patient is pregnant and is experiencing an intolerance to a preferred combination product that cannot be resolved or mitigated through alternative efforts (duration of PA will be 90 days post anticipated delivery date). Other requests will be considered with clinical justification explaining why the member cannot use the preferred buprenorphine formulations AND the Buprenorphine Safety Checklist has been completed (see PA form for detailed requirements). Requests to exceed quantity limits or maximum daily dose: documentation must be submitted detailing medical necessity for requested dosage regimen AND the Buprenorphine Safety Checklist has been completed (see PA form for detailed requirements).). Requests for treatment of pain AND opioid use disorder: The Buprenorphine Safety Checklist has been completed (see PA form for detailed requirements and
Note: Methadone for opioid use disorder can only be prescribed through a Methadone Maintenance Clinic		for documentation required) AND other non-opioid medications and pain management modalities have been trialed prior to increasing the buprenorphine

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
OPIOID WITHDRAWAL TREATMENT		dose for pain AND split dosing (multiple daily administrations) on current dose have been trialed for pain control as recommended in the ASAM 2020 practice guidelines AND clinical rationale has been provided if the request is for a dose increase > 25% the current daily dose. Sublocade: Diagnosis of opiate use disorder confirmed (will not be approved for alleviation of pain) AND patient has been stabilized (clinically controlled cravings and withdrawal symptoms) on a steady dose of 8mg to 24mg of a transmucosal buprenorphine product for at least 7 days AND clinical justification must be provided detailing why the member cannot use a more cost effective buprenorphine formulation. Note: Approval will be granted for 300mg monthly for the first 2 months followed by a maintenance dose of 100mg thereafter for a total length of approval not to exceed 6 months. A maintenance dose increase to 300mg will be considered for those patients who are able to tolerate the 100mg dose but do not demonstrate a satisfactory clinical response (including supplemental oral buprenorphine dosing, documentation of self-reported illicit opioid use, or urine drug screens positive for illicit opioid use). Once the patient is established on a maintenance dose, concurrent use of Sublocade and supplemental oral buprenorphine dosing will not be permitted. Sublocade must be dispensed directly to a healthcare provider and will not be approved for dispensing to the patient.
Central Alpha Agonists CLONIDINE IR tablets (compare to Catapres®) Note: Methadone for opiate dependency or withdrawal can only be prescribed through a Methadone Maintenance Clinic	Lucemyra® (lofexidine) Maximum length of therapy = 14 days	Lucemyra: Indication for use is the mitigation of opioid withdrawal symptoms to facilitate abrupt opioid discontinuation AND the patient is ≥ 18 years of age AND the patient is unable to tolerate clonidine due to significant side effects.
OVERDOSE TREATMENT		
NALOXONE HCL Prefilled luer-lock needleless syringe plus intranasal mucosal atomizing device (Rescue kit) NARCAN® (naloxone hcl) 4mg Nasal Spray QTY LIMIT: 4 single-use sprays/28days	Kloxxado TM (naloxone HCl) 8mg Nasal Spray <i>QTY LIMIT</i> : 4 single-use sprays/28days Naloxone HCl (compare to Narcan® 4 mg Nasal Spray) <i>QTY LIMIT</i> : 4 single-use sprays/28days Zimhi TM (naloxone HCl) 5mg Prefilled Syringe	 Kloxxado: The prescriber must provide a clinically compelling reason why Narcan cannot be used. Naloxone Nasal Spray: Narcan must be on a backorder and unavailable from the manufacturer. Zimhi: The prescriber must provide a clinically compelling reason why the preferred agents would not be suitable alternatives. Limitations: Effective 4/1/17, Evzio® is not classified as a covered outpatient drug and is therefore not covered by Vermont Medicaid.

PREFERRED AGENTS	NON-PREFERRED AGENTS			
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA		
	TERCTO CITED ON EDENIA CEMEN			
TESTOSTERONE REPLACEMENT THERAPY				
TOPICAL				
ANDRODERM® Transdermal 2 mg, 4 mg (testosterone patch) QTY LIMIT: 1 patch/day/strength TESTOSTERONE 1.62% Gel Packets QTY LIMIT: 1.25 gm packet (1.62%) = 1 packet/day, 2.5 gm packet (1.62%) = 2 packets/day TESTOSTERONE 1.62% Gel Pump (compare to Androgel®) QTY LIMIT: 2 bottles/30 days TESTOSTERONE 1% Gel Packets (compare to Androgel®, Vogelxo®) QTY LIMIT: 2.5 gm packet = 1 packet/day, 5 gm packet = 2 packets/day	Androgel [®] pump 1.62% (testosterone pump bottles) <i>QTY LIMIT</i> : 2 bottles/30 days Fortesta [®] (testosterone 2 % Gel) 60 gm Pump Bottle <i>QTY LIMIT</i> : 2 bottles/30 days Testim [®] Gel 5 gm (testosterone 1% gel tube) <i>QTY LIMIT</i> : 2 tubes/day Testosterone 1% gel tube (compare to Testim [®] Gel 5 gm, Vogelxo [®] , Androgel [®]) <i>QTY LIMIT</i> : 2 tubes/day Testosterone 1% Gel Pump (Vogelxo [®]) <i>QTY LIMIT</i> : 4 bottles/30 days Testosterone 2% gel 60 gm pump bottle (compare to Fortesta [®]) <i>QTY LIMIT</i> : 2 bottles/30 days Testosterone 2% solution 90ml Pump Bottle <i>QTY LIMIT</i> : 2 bottles/30 days Vogelxo [®] 1% (testosterone 1%) gel, pump <i>QTY LIMIT</i> : 2 tubes/day (5 gm gel tubes), 4 bottles/30 days (gel pump bottle)	Non-preferred agents: The patient has a documented side effect, allergy, or treatment failure to at least two preferred topical products.		
NASAL				
	Natesto® (testosterone) nasal gel QTY LIMIT: 3 bottles/30 days	Natesto: The patient has had a documented side effect, allergy, or treatment failure to TWO preferred testosterone products (topical and/or injectable formulations)		
ORAL				
All products require PA	Methitest (methyltesterone) tablet 10 mg Methyltestosterone capsule 10 mg Jatenzo (testosterone undecanoate) capsule Tlando (testosterone undecanoate) capsule	Oral non-preferred agents: The patient has had a documented side effect, allergy, or treatment failure to TWO preferred testosterone products (topical and/or injectable formulations) AND if the request is for Methitest or methyltestosterone, the patient has had a documented side effect, allergy, or treatment failure with Jatenzo.		
	Maximum day supply all products is 30 days			
INJECTABLE				

PREFERRED AGENTS	NON-PREFERRED AGENTS	
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA
TESTOSTERONE CYPIONATE IM (compare to Depo®-Testosterone) TESTOSTERONE ENANTHATE IM	Aveed® (testosterone undecanote) IM Depo®-Testosterone (testosterone cypionate) IM Testopel® (testosterone) implant pellets Xyosted™ (testosterone enanthate) SC	 Depo-Testosterone: The patient has a documented intolerance to generic testosterone cypionate. Aveed, Testopel, Xyosted: The patient has had a documented side effect, allergy, or treatment failure to TWO preferred testosterone products, one of which must be an injectable formulation. Treatment failure is defined as inability to achieve testosterone values in the 300-1,000ng/dL range despite adjustments to dose and frequency of injection.
	URINARY ANTISPASM	IODICS
SHORT-ACTING AGENTS OXYBUTYNIN	Detrol [®] (tolterodine) Flavoxate Tolterodine (compare to Detrol [®]) Trospium	Darifenacin, Detrol, Ditropan XL, tolterodine (generic), tolterodine SR (generic), trospium (generic), trospium ER (generic), Vesicare: The patient has had a documented side effect, allergy, or treatment failure with two preferred long-acting agents. If a medication has an AB rated
LONG-ACTING AGENTS OXYBUTYNIN XL (compare to Ditropan® XL) QTY LIMIT: 1/day SOLIFENACIN (compare to Vesicare®) QTY LIMIT: 1/day TOVIAZ® (fesoterodine) QTY LIMIT: 1/day	Darifenacin ER (compare to Enablex®) Ditropan XL® (oxybutynin XL) Tolterodine SR (compare to Detrol LA®) Trospium ER Vesicare® (solifenacin) Vesicare LS™ (solifenacin) oral suspension	generic, there must have also been a trial of the generic formulation. Gelnique 10%, Oxytrol: The patient is unable to swallow a solid oral formulation (e.g. patients with dysphagia) OR The patient is unable to be compliant with solid oral dosage forms. Gemtesa: The patient has had a documented side effect, allergy, treatment failure, or contraindication with one preferred long-acting urinary antimuscarinic agent and Myrbetriq. Myrbetriq Granules, Vesicare LS: The patient has a diagnosis of neurogenic
TRANSDERMAL/TOPICAL All products require PA	Gelnique 10% [®] (oxybutynin topical gel) <i>QTY LIMIT</i> : 1 sachet/day Oxytrol [®] (oxybutinin transdermal) <i>QTY LIMIT</i> : 8 patches/28 days Gemtesa® (vibegron) tablet	 detrusor overactivity AND the patient has a documented side effect, allergy, or treatment failure with oxybutynin or Toviaz AND for patients ≥ 18 years of age, medical necessity has been provided for a liquid formulation. Limitations: Oxytrol (for Women) OTC not covered. Oxytrol RX is available but subject to prior authorization.
BETA-3 ADRENERGIC AGONISTS MYRBETRIQ® (mirabegron) ER Tablet QTY LIMIT: 1 tablet/day	QTY LIMIT: 1 tablet/day Myrbetriq® ER Granules for Suspension	
	VAGINAL ANTI-INFEC	TIVES
CLEOCIN [®] Vaginal Ovules (clindamycin vaginal suppositories) CLINDAMYCIN VAGINAL (clindamycin vaginal	Cleocin [®] (clindamycin vaginal cream 2%) Gynazole-1® (butoconazole vaginal cream 2%) Nuvessa TM (metronidazole 1.3% Vaginal Gel)	Cleocin, Xaciato: The patient has had a documented side effect, allergy, or treatment failure to a preferred clindamycin vaginal cream.

PREFERRED AGENTS	NON-PREFERRED AGENTS			
(No PA required unless otherwise noted)	(PA required)	PA CRITERIA		
cream 2%) CLINDESSE [®] (clindamycin vaginal cream 2%) CLOTRIMAZOLE Vaginal cream MICONAZOLE Nitrate Vaginal cream, suppositories MICONAZOLE 1 Vaginal Kit MICONAZOLE 3 Vaginal Kit, cream MICONAZOLE 7 Vaginal cream, suppositories METRONIDAZOLE VAGINAL GEL 0.75%	Solosec [™] (secnidazole) oral granules packet Terconazole (compare to Terazol®) vaginal cream 0.4%, 0.8%, vaginal suppositories 80 mg Vandazole (metronidazole vaginal 0.75%) Xaciato [™] (clindamycin vaginal gel 2%)	 Nuvessa, Vandazole: The patient has had a documented side effect, allergy, or treatment failure to preferred metronidazole vaginal gel. Solosec: The patient has had a documented side effect, allergy, or treatment failure to a preferred topical anti-infective and oral metronidazole. Gynazole, Terconazole: The patient has a documented side effect, allergy, or treatment failure to a preferred miconazole or clotrimazole formulation. 		
VASOPRESSIN RECEPTOR ANTAGONIST				
	Jynarque® tablets (tolvaptan) QTY LIMIT: 56 tablets/28 days Samsca® tablets (tolvaptan) QTY LIMIT: 15 mg = 1 tablet/day, 30 mg 2 tablets/day	 Jynarque: The patient must be ≥ 18 years of age AND the patient is at risk of rapidly progressing Autosomal Polycystic Kidney Disease (ADPKD) AND the patient has normal serum sodium concentrations before starting the medication (results must be submitted) AND the patient and provider are enrolled in the Jynarque® REMS program Samsca: The agent is being used for the treatment of euvolemic or hypervolemic hyponatremia AND Despite optimal fluid restriction, the patient's serum sodium < 120 mEq/L or the patient is symptomatic with a serum sodium < 125 mEq/L. AND The treatment will be initiated or is being reinitiated in a hospital setting where serum sodium can be monitored 		
VITAMINS: PRENATAL MULTIVITAMINS				
C-NATE DHA M-NATAL PLUS NIVA-PLUS PRENATAL PLUS IRON PRENATAL VITAMINS PLUS SE-NATAL CHEW WESTAB PLUS	All others	All Non-Preferred: The prescriber must provide a clinically valid reason for the use of the requested medication including reasons why any of the preferred products would not be a suitable alternative.		