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- Prior authorization for a non-preferred agent in any class will be given only if there has been a trial of the preferred brand/generic
 equivalent or preferred formulation of the active ingredient, at a therapeutic dose, that resulted in a partial response with a documented
 intolerance.
- Prior authorization of a non-preferred isomer, pro-drug, or metabolite will be considered with a trial of a preferred parent drug
 of the same chemical entity, at a therapeutic dose, that resulted in a partial response with documented intolerance or a previous
 trial and therapy failure, at a therapeutic dose, with a preferred drug of a different chemical entity indicated to treat the
 submitted diagnosis. The required trial may be overridden when documented evidence is provided that the use of these
 preferred agent(s) would be medically contraindicated.
- Unless otherwise specified, the listing of a particular brand or generic name includes all legend forms of that drug. OTC drugs are not covered unless specified.
- PA criteria for non-preferred agents apply in addition to general Drug Utilization Review policy that is in effect for the entire pharmacy program, including, but not limited to, appropriate dosing, duplication of therapy, etc.
- The use of pharmaceutical samples will not be considered when evaluating the members' medical condition or prior prescription history for drugs that require prior authorization.
- Quantity limits may apply. Refer to the Limits List on the BMS Website by clicking the hyperlink.
- Unless otherwise indicated, non-preferred combination products require medical reasoning beyond convenience or enhanced compliance as to why the clinical need cannot be met with a preferred agent or combination of preferred single-ingredient agents.
- Acronyms
 - CL Requires clinical PA. For detailed clinical criteria, please go to the PA criteria page by clicking the hyperlink.
 - NR Denotes a new drug which has not yet been reviewed by the P & T Committee. These agents are available only on appeal to the BMS Medical Director.
 - o AP Non-preferred and selected preferred drugs, where indicated, are subject to auto-PA criteria. See PA criteria column.



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CLASSES CHANGING	Status Changes	PA Criteria Changes	New Drugs
ANALGESICS, NARCOTIC SHORT ACTING			XXX
ANTICONVULSANTS			XXX
ANTIFUNGALS, ORAL			XXX
ANTIPSYCHOTICS, ATYPICAL			XXX
BLADDER RELAXANT PREPARATIONS			XXX
BRONCHODILATORS, BETA-AGONISTS			XXX
MULTIPLE SCLEROSIS AGENTS			XXX
OPHTHALMICS FOR ALLERGIC CONJUNCTIVITIS			XXX
OPHTHALMICS, ANTI-INFLAMMATORIES			XXX
PITUITARY SUPPRESSIVE AGENTS, LHRH	XXX		XXX
STIMULANTS AND RELATED AGENTS			XXX



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THERAPEUTIC DRUG CLASS				
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA		
ACNE AGENTS, TOPICALAP	ACNE AGENTS, TOPICALAP			
CLASS PA CRITERIA: Non-preferred agents r subclasses, including the generic version of the present.	require a thirty (30) day trial of one (1) preferred retinor requested non-preferred product, before they will be	oid and two (2) unique chemical entities in two (2) other approved, unless one (1) of the exceptions on the PA form is		
In cases of pregnancy, a trial of retinoids will <i>no</i> Acne kits are non-preferred.	t be required. For members eighteen (18) years of a	ge or older, a trial of retinoids will not be required.		
Specific Criteria for sub-class will be listed to 30-day trial of all preferred agents in that sub-	class.	sub-class are available only on appeal and require at least a		
	ANTI-INFECTIVE			
CLINDAGEL (clindamycin) clindamycin lotion, medicated swab, solution erythromycin gel, solution	AMZEEQ FOAM (minocycline) CLEOCIN-T (clindamycin) CLINDACIN ETZ kit, medicated swab (clindamycin) CLINDACIN P (clindamycin) CLINDACIN PAC (clindamycin) clindamycin gel, foam dapsone ERYGEL (erythromycin) erythromycin medicated swab EVOCLIN (clindamycin) FABIOR (tazarotene) KLARON (sulfacetamide) OVACE/PLUS (sulfacetamide) sodium sulfacetamide 10% cleansing gel sulfacetamide			
	RETINOIDS			
DIFFERIN (adapalene) RETIN-A (tretinoin) RETIN-A MICRO (tretinoin)	adapalene AKLIEF CREAM (trifarotene) ALTRENO LOTION (tretinoin) ARAZLO (tazarotene) ATRALIN (tretinoin) AVITA (tretinoin) tazarotene cream tretinoin cream, gel tretinoin gel micro	In addition to the Class Criteria: PA required for members eighteen (18) years of age or older.		
	KERATOLYTICS			



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benzoyl peroxide cleanser Rx & OTC, 10% cream OTC, gel Rx & OTC, lotion OTC, wash OTC	BENZEFOAM benzoyl peroxide) BP 10-1 (benzoyl peroxide) BPO (benzoyl peroxide)	
PANOXYL-4 OTC (benzoyl peroxide)	COMBINATION AGENTS	
ACANYA (clindamycin phosphate/benzoyl	adapalene-benzoyl peroxide*	In addition to the Class Criteria: Non-preferred combination
peroxide) BENZAMYCIN PAK (benzoyl peroxide/ erythromycin) benzoyl peroxide/clindamycin gel (generic DUAC only) EPIDUO (adapalene/benzoyl peroxide)* EPIDUO FORTE (adapalene/benzoyl peroxide)* ONEXTON (clindamycin phosphate/benzoyl	AVAR/-E/LS (sulfur/sulfacetamide) BENZACLIN GEL (benzoyl peroxide/ clindamycin) benzoyl peroxide/clindamycin gel (all generics other than DUAC) benzoyl peroxide/urea clindamycin phosphate/benzoyl peroxide (generic Acanya) clindamycin-tretinoin gel*	agents require thirty (30) day trials of the corresponding preferred single agents before they will be approved. *PA required for combination agents with Retinoid products for members eighteen (18) years of age or older.
peroxide) sulfacetamide sodium/sulfur suspension ZIANA (clindamycin/tretinoin)*	erythromycin/benzoyl peroxide NEUAC (clindamycin phosphate/benzoyl peroxide) SSS 10-4 (sulfacetamide /sulfur) SSS 10-5 foam (sulfacetamide /sulfur) sulfacetamide sodium/sulfur cloths, lotion, pads sulfacetamide/sulfur wash/cleanser sulfacetamide/sulfur wash kit sulfacetamide sodium/sulfur/ urea SUMADAN/XLT (sulfacetamide/sulfur) SUMAXIN/TS (sulfacetamide sodium/sulfur)	
	ROSACEA AGENTS	
FINACEA GEL (azelaic acid) MIRVASO GEL (brimonidine) metronidazole cream metronidazole gel 0.75% (NDCs 00115-1474- 46, 00168-0275-45, 51672-4116-06, 66993-0962-45 only)	azelaic acid gel FINACEA FOAM (azelaic acid) ivermectin METROCREAM (metronidazole) METROGEL GEL (metronidazole) metronidazole lotion metronidazole gel (all other NDCs) NORITATE CREAM (metronidazole) RHOFADE (oxymetazoline) ROSADAN (metronidazole) SOOLANTRA CREAM (ivermectin) ZILXI (minocycline) foam	Subclass criteria : Non-preferred agents are available only on appeal and require evidence of 30-day trials of all chemically-unique preferred agents in the sub-class.
ALZHEIMER'S AGENTSAP		

ALZHEIMER'S AGENTSAP

CLASS PA CRITERIA: Non-preferred agents require a thirty (30) day trial of a preferred agent in the same sub-class before they will be approved, unless one (1) of the exceptions on the PA form is present.

Prior authorization is required for members up to forty-five (45) years of age if there is no diagnosis of Alzheimer's disease.



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	CHOLINESTERASE INHIBITORS	
donepezil 5 and 10 mg donepezil ODT	ARICEPT (donepezil) donepezil 23 mg* EXELON PATCH (rivastigmine) galantamine galantamine ER RAZADYNE ER (galantamine) Rivastigmine	*Donepezil 23 mg tablets will be authorized if the following criteria are met: 1. There is a diagnosis of moderate-to-severe Alzheimer's Disease and 2. There has been a trial of donepezil 10 mg daily for at least three (3) months and donepezil 20 mg daily for an additional one (1) month.
	NMDA RECEPTOR ANTAGONIST	()
memantine	memantine ER memantine solution NAMENDA (memantine) NAMENDA XR (memantine)*	*Namenda XR requires ninety (90) days of compliant therapy with Namenda.
CHOLINE	STERASE INHIBITOR/NMDA RECEPTOR ANTAG	
	NAMZARIC (donepezil/memantine)	Combination agents require thirty (30) day trials of each corresponding preferred single agent.
ANALGESICS, NARCOTIC LONG	ACTING (Non-parenteral) ^{AP}	
the requested non-preferred agent (if available) to for the requested non-preferred brand agent, the	pefore they will be approved, unless one (1) of the exc n another generic non-preferred agent must be trialed	et preferred agents AND a six (6) day trial of the generic form of eptions on the PA form is present. If no generic form is available instead. NOTE: All long-acting opioid agents require a prior indication and specify previous opioid and non-opioid therapies *Belbuca prior authorization requires manual review. Full PA criteria may be found on the PA Criteria page by clicking the hyperlink. **Methadone will be authorized without a trial of the preferred agents if a diagnosis of cancer is submitted. ***Tramadol ER (generic Conzip) requires a manual review and may be authorized for ninety (90) days with submission of a detailed treatment plan including anticipated duration of treatment and scheduled follow-ups with the prescriber. ****Nucynta requires six (6) day trials of three (3) chemically distinct preferred agents



BUREAU FOR MEDICAL SERVICES WEST VIRGINIA MEDICAID REFERENCE DELICATION CRITERIA

PREFERRED DRUG LIST WITH PRIOR AUTHORIZATION CRITERIA

This is not an all-inclusive list of available covered drugs and includes only

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ANALGESICS, NARCOTIC SHORT ACTING (Non-parenteral) AP

CLASS PA CRITERIA: Non-preferred agents require six (6) day trials of at least four (4) chemically distinct preferred agents (based on the narcotic ingredient only), including the generic formulation of the requested non-preferred agent, before they will be approved, unless one (1) of the exceptions on the PA form is present.

NOTE: All tramadol and codeine products require a prior authorization for children under 18 years of age. Requests must be for an FDA approved age and indication and specify non-opioid therapies attempted.

managed categories. Refer to cover page for complete list of rules governing this PDL.

APAP/codeine

butalbital/APAP/caffeine/codeine

codeine

hydrocodone/APAP 2.5/325 mg, 5/325 mg,

7.5/325 mg,10/325 mg hydrocodone/APAP solution hydrocodone/ibuprofen hydromorphone tablets

LORTAB SOLUTION

LOK TAB SOLUTION

(hydrocodone/acetaminophen)

morphine

oxycodone tablets, concentrate, solution

oxycodone/APAP oxycodone/ASA pentazocine/naloxone

tramadol

tramadol/APAP

ABSTRAL (fentanyl) ACTIQ (fentanyl)

butalbital/ASA/caffeine/codeine

butorphanol

DEMEROL (meperidine) dihydrocodeine/ APAP/caffeine

DILAUDID (hydromorphone)

fentanyl

FENTORA (fentanyl)

FIORICET W/ CODEINE

(butalbital/APAP/caffeine/codeine)

FIORINAL W/ CODEINE

(butalbital/ASA/caffeine/codeine)

hydrocodone/APAP 5/300 mg, 7.5/300 mg,

10/300 mg

hydromorphone liquid, suppositories

levorphanol

LORCET (hydrocodone/APAP)

LORTAB (hydrocodone/APAP)

meperidine

NORCO (hydrocodone/APAP)

NUCYNTA (tapentadol) oxycodone capsules oxycodone/ibuprofen

oxymorphone

PÉRCOCET (oxycodone/APAP)

QDOLO SOLÙTION (tramadol)

ROXICODONE (oxycodone)
ULTRACET (tramadol/APAP)

VICOPROFEN (hydrocodone/ibuprofen)

Fentanyl buccal, nasal and sublingual products will only be authorized for a diagnosis of cancer and as an adjunct to a long-acting agent. These dosage forms will not be authorized for monotherapy.

Limits: Unless the patient has escalating cancer pain or another diagnosis supporting increased quantities of short-acting opioids, all short acting solid forms of the narcotic analgesics are limited to 120 tablets per thirty (30) days. Longer-acting medications should be maximized to prevent unnecessary breakthrough pain in chronic pain therapy.

Immediate-release tramadol is limited to 240 tablets per thirty (30) days.

ANDROGENIC AGENTS

CLASS PA CRITERIA: A non-preferred agent will only be authorized if one (1) of the exceptions on the PA form is present.

ANDRODERM (testosterone)

ANDROGEL (testosterone)

ANDROGEL (testosterone)

ANDROBEL (testosterone)

METHITEST (methyltestosterone)

JATENZO (testosterone undecanoate)

testosterone cypionate vial^{CL} methyltestosterone capsule testosterone enanthate vial^{CL} NATESTO (testosterone)

TESTIM (testosterone)
TESTRED (methyltestosterone)

testosterone gel

VOGELXO (testosterone)



managed categories. Refer to cover page for complete list of rules governing this PDL.

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	XYOSTED (testosterone enanthate)	
ANESTHETICS, TOPICALAP		
•		ent before they will be approved, unless one (1) of the exceptions on the
PA form is present.	a agents require terr (10) day thats of each preferred age	ent before they will be approved, unless one (1) of the exceptions on the
lidocaine	lidocaine/hydrocortisone	
lidocaine/prilocaine	LIDOTRAL CREAM (lidocaine)	
xylocaine	LIDOZION LOTION (lidocaine)	
ANGIOTENSIN MODULATO	SYNERA (lidocaine/tetracaine)	
	d agents require fourteen (14) day trials of each prefer d, unless one (1) of the exceptions on the PA form is pr	red agent in the same sub-class, with the exception of the Direct Renin esent.
	ACE INHIBITORS	
benazepril	ACCUPRIL (quinapril)	*Epaned will be authorized with a diagnosis of hypertension,
captopril	ALTACE (ramipril)	symptomatic heart failure or asymptomatic left ventricular
enalapril	EPANED (enalapril)*	dysfunction provided that the patient is less than seven (7)
fosinopril	LOTENSIN (benazepril)	years of age OR is unable to ingest a solid dosage form due
lisinopril quinapril	moexipril perindopril	to documented oral-motor difficulties or dysphagia.
ramipril	PRINIVIL (lisinopril)	**Qbrelis solution may be authorized for children ages 6-10
Tampin	QBRELIS SOLUTION (lisinopril)**	who are unable to tolerate a solid dosage form. Qbrelis may
	trandolapril	also be authorized for older patients with clinical
	VASOTEC (enalapril)	documentation indicating oral-motor difficulties or
	ZESTRIL (lisinopril)	dysphagia.
	ACE INHIBITOR COMBINATIO	
benazepril/amlodipine	ACCURETIC (quinapril/HCTZ)	
benazepril/HCTZ	LOTENSIN HCT (benazepril/HCTZ)	
captopril/HCTZ	LOTREL (benazepril/amlodipine)	
enalapril/HCTZ	TARKA (trandolapril/verapamil)	
fosinopril/HCTZ	trandolapril/verapamil	
lisinopril/HCTZ	VASERETIC (enalapril/HCTZ)	
quinapril/HCTZ	ZESTORETIC (lisinopril/HCTZ) ANGIOTENSIN II RECEPTOR BLOG	CKERS (ARRs)
irbesartan	ATACAND (candesartan)	ONENO (ANDS)
losartan	AVAPRO (irbesartan)	
valsartan	BENICAR (olmesartan)	
olmesartan	candesartan	
	COZAAR (losartan)	
	DIOVAN (valsartan)	
	EDARBI (azilsartan)	
	MICARDIS (telmisartan)	
	telmisartan	
	ARB COMBINATIONS	
ENTRESTO (valsartan/sacubitril) ^{AP*}	ATACAND-HCT (candesartan/HCTZ)	*Entresto will only be authorized for patients 18 years of age
	AVALIDE (irbesartan/HCTZ)	or older who are diagnosed with chronic heart-failure.



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PREFERRED DRUG LIST WITH PRIOR AUTHORIZATION CRITERIA

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irbesartan/HCTZ losartan/HCTZ olmesartan/amlodipine olmesartan/HCTZ TRIBENZOR (olmesartan/amlodipine/HCTZ) valsartan/amlodipine valsartan/amlodipine/HCTZ valsartan/HCTZ	AZOR (olmesartan/amlodipine) BENICAR-HCT (olmesartan/HCTZ) candesartan/HCTZ DIOVAN-HCT (valsartan/HCTZ) EDARBYCLOR (azilsartan/chlorthalidone) EXFORGE (valsartan/amlodipine) EXFORGE HCT (valsartan/amlodipine/HCTZ) HYZAAR (losartan/HCTZ) MICARDIS-HCT (telmisartan/HCTZ) olmesartan/amlodipine/HCTZ telmisartan/amlodipine telmisartan HCTZ	
	DIRECT RENIN INHIBITORS	
	aliskiren TEKTURNA (aliskiren) TEKTURNA HCT (aliskiren/HCTZ)	Substitute for Class Criteria: Tekturna requires a thirty (30) day trial of one (1) preferred ACE, ARB, or combination agent, at the maximum tolerable dose, before it will be authorized unless one (1) of the exceptions on the PA form is present. Amturnide, Tekamlo, Tekturna HCT or Valturna will be authorized if the criteria for Tekturna are met and the patient also needs the other agents in the combination.
ANTIANGINAL & ANTI-ISCHEMIC		ŭ de la
as single agents or a combination agent contain		also taking a calcium channel blocker, a beta blocker, or a nitrite
ranolazine ^{AP}	RANEXA	
ANTIBIOTICS, GI & RELATED AG	ENTS	
CLASS PA CRITERIA: Non-preferred agents		efore they will be approved, unless one (1) of the exceptions on
the PA form is present.		
FIRVANQ (vancomycin) metronidazole tablet neomycin tinidazole	DIFICID (fidaxomicin)* FLAGYL (metronidazole) metronidazole capsule paromomycin VANCOCIN (vancomycin) vancomycin XIFAXAN (rifaximin)*	*Full PA criteria may be found on the PA Criteria page by clicking the hyperlink.
ANTIBIOTICS, INHALED		
CLASS PA CRITERIA: Non-preferred agents a approved, unless one (1) of the exceptions on the exceptions of the exception of the exceptions of the exceptions of the exception of the ex		nt and documentation of therapeutic failure before they will be
BETHKIS (tobramycin) KITABIS PAK (tobramycin)	CAYSTON (aztreonam) TOBI (tobramycin) TOBI PODHALER (tobramycin) tobramycin	
ANTIBIOTICS, TOPICAL		

CLASS PA CRITERIA: Non-preferred agents require ten (10) day trials of at least one preferred agent, including the generic formulation of the requested non-preferred agent, before they will be approved, unless one (1) of the exceptions on the PA form is present.



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bacitracin (Rx, OTC) gentamicin sulfate mupirocin ointment	CENTANY (mupirocin) CORTISPORIN (bacitracin/neomycin/polymyxin/HC) mupirocin cream neomycin/polymyxin/pramoxine XEPI CREAM (ozenoxacin)	
ANTIBIOTICS, VAGINAL		
will be approved, unless one (1) of the exception	ns on the PA form is present.	nt at the manufacturer's recommended duration, before they
CLEOCIN OVULE (clindamycin) CLINDESSE (clindamycin) metronidazole gel NUVESSA (metronidazole)	CLEOCIN CREAM (clindamycin) clindamycin cream METROGEL (metronidazole) SOLOSEC (secnidazole) VANDAZOLE (metronidazole)	
ANTICOAGULANTS		
CLASS PA CRITERIA: Non-preferred agents represent.	equire a trial of each preferred agent in the same sub	o-class, unless one (1) of the exceptions on the PA form is
	INJECTABLECL	
enoxaparin	ARIXTRA (fondaparinux) fondaparinux FRAGMIN (dalteparin) LOVENOX (enoxaparin)	
	ORAL	
ELIQUIS (apixaban) PRADAXA (dabigatran) warfarin XARELTO (rivaroxaban)	SAVAYSA (edoxaban)	

ANTICONVULSANTS

CLASS PA CRITERIA: For a diagnosis of seizure disorder, non-preferred agents require a fourteen (14) day trial of a preferred agent in the same sub-class before they will be approved, unless one (1) of the exceptions on the PA form is present; patients currently on established therapies shall be grandfathered.

For all other diagnoses, non-preferred agents require a thirty (30) day trial of a preferred agent in the same sub-class before they will be approved, unless one (1) of the exceptions on the PA form is present.

In situations where AB-rated generic equivalent products are available, "Brand Medically Necessary" must be hand-written by the prescriber on the prescription for the brand name product to be reimbursed.



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ADJUVANTS			
carbamazepine carbamazepine ER divalproex divalproex Sprinkle EPITOL (carbamazepine) GABITRIL (tiagabine) lamotrigine levetiracetam IR levetiracetam IR suspension oxcarbazepine suspension and tablets TEGRETOL SUSPENSION (carbamazepine) topiramate IR topiramate ER* valproic acid VIMPAT (lacosamide) zonisamide	APTIOM (eslicarbazepine) BANZEL (rufinamide) BRIVIACT (brivaracetam) carbamazepine oral suspension CARBATROL (carbamazepine) DEPAKOTE (divalproex) DEPAKOTE ER (divalproex) DEPAKOTE SPRINKLE (divalproex) DIACOMIT CAPSULE/POWDER PACK (stripentol)** ELEPSIA XR (levetiracetam) EQUETRO (carbamazepine) felbamate FELBATOL (felbamate) FINTEPLA (fenfluramine) SOLUTION**** FYCOMPA (perampanel) KEPPRA SOLUTION (levetiracetam) KEPPRA XR (levetiracetam) LAMICTAL (lamotrigine) LAMICTAL CHEWABLE (lamotrigine) LAMICTAL CHEWABLE (lamotrigine) LAMICTAL XR (lamotrigine) lamotrigine dose pack lamotrigine dose pack lamotrigine ODT OXTELLAR XR (oxcarbazepine) QUDEXY XR (topiramate ER)**** rufinamide oral suspension, tablets SABRIL (vigabatrin) SPRITAM (levetiracetam) TEGRETOL TABLETS (carbamazepine) TEGRETOL XR (carbamazepine) tiagabine TOPAMAX (topiramate) TRILEPTAL SUSPENSION and TABLETS (oxcarbazepine) TROKENDI XR (topiramate)*** vigabatrin tablet/powder pack	*Topiramate ER will be authorized after a thirty (30) day trial of topiramate IR. **Diacomit may only be approved as adjunctive therapy for diagnosis of Dravet Syndrome when prescribed by, or in consultation with, a neurologist AND requires a thirty (30) day trial of valproate and clobazam unless one (1) of the exceptions on the PA form is present. Diacomit must be used concurrently with clobazam. ***Qudexy XR and Trokendi XR are only approvable on appeal. ****Full PA criteria for Fintepla may be found on the PA Criteria page by clicking the hyperlink.	
	XCOPRI (cenobamate)		
	BARBITURATES ^{AP}		
phenobarbital	MYSOLINE (primidone)		
primidone	mi ocania (primidono)		
F	BENZODIAZEPINESAP		
	DENZUDIAZEPINEOAP		



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clonazepam diazepam rectal gel diazepam tablets NAYZILAM NASAL SPRAY (midazolam) VALTOCO NASAL SPRAY (diazepam)	clobazam* clonazepam ODT DIASTAT (diazepam rectal) KLONOPIN (clonazepam) ONFI (clobazam)* ONFI SUSPENSION (clobazam)* SYMPAZAN (clobazam film)* CANNABINOIDS	*Onfi shall be authorized as adjunctive therapy for treatment of Lennox-Gastaut Syndrome and Dravet Syndrome without further restrictions. All other indications require an appeal to the Medical Director. NOTE: generic clobazam is preferred over brand ONFI.
	EPIDIOLEX SOLUTION (cannabidiol)*	* Full PA criteria may be found on the PA Criteria page by
	HYDANTOINS ^{AP}	clicking the hyperlink.
DILANTIN (phenytoin sodium, extended)	DILANTIN INFATABS (phenytoin)	
PEGANONE (ethotoin) phenytoin capsules, chewable tablets, suspension	PHENYTEK (phenytoin)	
	SUCCINIMIDES	
CELONTIN (methsuximide) ethosuximide capsules ethosuximide syrup	ZARONTIN (ethosuximide) capsules ZARONTIN (ethosuximide) syrup	
ANTIDEPRESSANTS, OTHER		
CLASS PA CRITERIA: See below for indivi	dual sub-class criteria. MAOIs^{AP}	
	MARPLAN (isocarboxazid) NARDIL (phenelzine) phenelzine tranylcypromine	Patients stabilized on MAOI agents will be grandfathered.
	SNRISAP	
duloxetine capulses venlafaxine ER capsules	CYMBALTA (duloxetine) desvenlafaxine ER desvenlafaxine fumarate ER EFFEXOR XR (venlafaxine) FETZIMA (levomilnacipran) PRISTIQ (desvenlafaxine) venlafaxine IR venlafaxine ER tablets (venlafaxine)	Non-preferred agents require separate thirty (30) day trials of a preferred agent in this sub-class AND an SSRI before they will be approved, unless one (1) of the exceptions on the PA form is present.
	SECOND GENERATION NON-SSRI,	
bupropion IR bupropion SR bupropion XL mirtazapine trazodone	APLENZIN (bupropion hbr) EMSAM (selegiline) FORFIVO XL (bupropion) nefazodone REMERON (mirtazapine) TRINTELLIX (vortioxetine) VIIBRYD (vilazodone HCI) WELLBUTRIN SR (bupropion)	Non-preferred agents require separate thirty (30) day trials of a preferred agent in this sub-class AND an SSRI before they will be approved, unless one (1) of the exceptions on the PA form is present.



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	WELLBUTRIN XL (bupropion)	
	SELECTED TCAs	
imipramine HCI	imipramine pamoate	Non-preferred agents require a twelve (12) week trial of imipramine HCl before they will be approved, unless one (1) of the exceptions on the PA form is present.
ANTIDEPRESSANTS, SSRISAP		
CLASS PA CRITERIA: Non-preferred agent exceptions on the PA form is present.	s require thirty (30) day trials of at least two (2) prefe	erred agents before they will be approved, unless one (1) of the
Upon hospital discharge, patients admitted wir continue that drug.	h a primary mental health diagnosis who have been st	tabilized on a non-preferred SSRI will receive an authorization to
citalopram escitalopram tablets fluoxetine capsules, solution fluvoxamine paroxetine sertraline	BRISDELLE (paroxetine) CELEXA (citalopram) escitalopram solution fluoxetine tablets fluvoxamine ER LEXAPRO (escitalopram) paroxetine 7.5 mg capsules paroxetine ER PAXIL (paroxetine) PAXIL CR (paroxetine) PEXEVA (paroxetine) PROZAC (fluoxetine) SARAFEM (fluoxetine) ZOLOFT (sertraline)	
ANTIEMETICSAP		
CLASS PA CRITERIA: See below for sub-cla	ss criteria.	
	5HT3 RECEPTOR BLOCKERS	
granisetron ondansetron ODT, solution, tablets	ondansetron vials SANCUSO (granisetron) SUSTOL (granisetron) ZOFRAN (ondansetron) ZUPLENZ (ondansetron) CANNABINOIDS	Non-preferred agents require a three (3) day trial of a preferred agent before they will be approved, unless one (1) of the exceptions on the PA form is present.
	dronabinol*	*Dronabinol will only be authorized for:
	MARINOL (dronabinol)*	The treatment of anorexia associated with weight loss in patients with AIDS or cancer and unresponsive to megestrol or The prophylaxis of chemotherapy induced nausea and vomiting unresponsive to three (3) day trials of ondansetron or promethazine for patients from eighteen (18) up to sixty-five (65) years of age.



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	SUBSTANCE P ANTAGONISTS	
EMEND (aprepitant)	aprepitant VARUBI (rolapitant)	Non-preferred agents require a three (3) day trial of a preferred agent before they will be approved, unless one (1) of the exceptions on the PA form is present.
	COMBINATIONS	Non-restaurational annual control
	AKYNZEO (netupitant/palonosetron) BONJESTA (doxylamine/pyridoxine) DICLEGIS (doxylamine/pyridoxine) doxylamine/pyridoxine (generic Diclegis)	Non-preferred agents will only be approved on appeal.
ANTIFUNGALS, ORAL		
CLASS PA CRITERIA: Non-preferred agent	s will only be authorized if one (1) of the exceptions on	the PA form is present.
clotrimazole fluconazole* nystatin terbinafine ^{CL}	ANCOBON (flucytosine)CRESEMBA (isovuconazonium)CL** DIFLUCAN (fluconazole) flucytosine griseofulvi** itraconazole ketoconazole**** MYCELEX (clotrimazole) NOXAFIL (posaconazole) ORAVIG (miconazole) posaconazole tablet SPORANOX (itraconazole) TOLSURA (itraconazole) VFEND (voriconazole) voriconazole suspension voriconazole tablets	**PA is required when limits are exceeded. **Full PA criteria may be found on the PA Criteria page by clicking the hyperlink. ***PA is not required for griseofulvin suspension for children up to eighteen (18) years of age for the treatment of tinea capitis. ****Ketoconazole will be authorized if the following criteria are met: 1. Diagnosis of one of the following fungal infections: blastomycosis, coccidioidomycosis, histoplasmosis, chromomycosis, or paracoccidioidomycosis and 2. Documented failure or intolerance of all other diagnosis-appropriate antifungal therapies, i.e. itraconazole, fluconazole, flucytosine, etc and 3. Baseline assessment of the liver status including alanine aminotransferase (ALT), aspartate aminotransferase (AST), total bilirubin, alkaline phosphatase, prothrombin time, and international normalized ratio (INR) before starting treatment and 4. Weekly monitoring of serum ALT for the duration of treatment (If ALT values increase to a level above the upper limit of normal or 30% above baseline, or if the patient develops symptoms of abnormal liver function, treatment should be interrupted and a full set of liver tests be obtained. Liver tests should be repeated to ensure normalization of values.) and 5. Assessment of all concomitant medications for potential



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Ketoconazole will not be authorized for treatment for

fungal infections of the skin and nails.

ANTIFUNGALS, TOPICALAP		
		gents before they will be approved, unless one (1) of the y trial of one (1) preferred product (i.e. ketoconazole shampoo) is
	ANTIFUNGALS	
econazole ketoconazole cream, shampoo MENTAX (butenafine) miconazole (OTC) nystatin	CICLODAN (ciclopirox) ciclopirox ERTACZO (sertaconazole) EXELDERM (sulconazole) EXTINA (ketoconazole) GYNAZOLE 1 CREAM (butoconazole) JUBLIA (efinaconazole) ketoconazole foam KERYDIN (tavaborole) KETODAN (ketoconazole) LOPROX (ciclopirox) LUZU (luliconazole) miconazole/petrolatum/zinc oxide NAFTIN GEL (naftifine) OXISTAT (oxiconazole)* tavaborole 5% topical solution VUSION (miconazole/petrolatum/zinc oxide)	*Oxistat cream will be authorized for children up to thirtee (13) years of age for tinea corporis, tinea cruris, tinea pedis and tinea (pityriasis) versicolor.
clotrimazole/betamethasone cream	ANTIFUNGAL/STEROID COMBINATI	ONS
ciotiimazoie/betamethasone cream	clotrimazole/betamethasone lotion nystatin/triamcinolone	
ANTIHEMOPHILIA FACTOR A	-	
CLASS PA CRITERIA: All agents will requal preferred product.	uire prior-authorization, and non-preferred agents require grandfathered with documentation of adherence to thera	e medical reasoning explaining why the need cannot be met using the second seco
	FACTOR VIII	
ADVATE AFSTYLA ALPHANATE HEMOFIL M HUMATE-P KOATE KOGENATE FS NOVOEIGHT NUWIQ WILATE XYNTHA	ADYNOVATE ELOCTATE ESPEROCT JIVI KOVALTRY RECOMBINATE VONVENDI	



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XYNTHA SOLOFUSE				
FACTOR VII				
	NOVOSEVEN ^{NR} SEVENFACT ^{NR}			
	FACTOR IX			
ALPHANINE SD ALPROLIX BENEFIX IXINITY MONONINE PROFILNINE RIXUBIS	IDELVION REBINYN			
	FACTOR IXa/IX			
	HEMLIBRA (emicizumab-kxwh)*	*Hemlibra shall be approved without further restriction for patients with Hemophilia A with documented presence of Factor VIII inhibitors.		
ANTIHYPERTENSIVES, SYMPATH	HOLYTICS			
CLASS PA CRITERIA: Non-preferred agents rebe approved, unless one (1) of the exceptions o		hemical entity in the corresponding formulation before they will		
CATAPRES-TTS (clonidine) clonidine patch clonidine tablets	CATAPRES TABLETS (clonidine)			
ANTIHYPERURICEMICS				
	equire a thirty (30) day trial of one (1) of the preferred oil) before they will be approved, unless one (1) of the			
	ANTIMITOTICS			
COLCRYS (colchicine) tablets	colchicine capsules colchicine tablets MITIGARE (colchicine) GLOPERBA (colchicine)*	In the case of acute gouty attacks, a ten (10) day supply (twenty (20) units) of the preferred agent(s) in this subclass will be authorized per ninety (90) days. *Gloperba may only be authorized for those who are unable to ingest solid dosage forms due to documented oralmotor difficulties or dysphagia.		
ANTIMITOTIC-URICOSURIC COMBINATION				
colchicine/probenecid				
URICOSURIC				
probenecid				
XANTHINE OXIDASE INHIBITORS				



BUREAU FOR MEDICAL SERVICES WEST VIRGINIA MEDICAID

PREFERRED DRUG LIST WITH PRIOR AUTHORIZATION CRITERIA

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allopurinol	febuxostat tablets ULORIC (febuxostat) ZYLOPRIM (allopurinol)	
ANTIMIGRAINE AGENTS, PRO	PHYLAXISCL	
		d on the PA Criteria page by clicking the hyperlink. Non-preferred
agents require a 90-day trial of all preferred		*F
AIMOVIG (erenumab) AJOVY (fremanezumab)	EMGALITY (galcanezumab) 120mg/mL EMGALITY (galcanezumab) 300mg/3 mL*	*Emgality 300 mg/3 mL requires review by the Medical Director and is available only on appeal.
ANTIMIGRAINE AGENTS, ACU	TEAP	
	ts require three (3) day trials of each preferred unique available), before they will be approved, unless one (1	chemical entity as well as a three (3) day trial using the same route) of the exceptions on the PA form is present.
	TRIPTANS	
naratriptan	almotriptan	*In addition to the Class Criteria: Onzetra Xsail and
rizatriptan ODT rizatriptan tablet	AMERGE (naratriptan) eletriptan	Tosymra require three (3) day trials of each preferred oral,
sumatriptan injection ^{CL}	FROVA (frovatriptan)	nasal and injectable forms of sumatriptan.
sumatriptan nasal spray	frovatriptan	
sumatriptan tablets	IMITREX NASAL SPRAY (sumatriptan)	
Sumatriplan tablets	IMITREX tablets (sumatriptan)	
	MAXALT MLT (rizatriptan)	
	MAXALT (rizatriptan)	
	ONZETRA XSAIL (sumatriptan)*	
	RELPAX (eletriptan)	
	TOSYMRA NASAL SPRAY (sumatriptan)*	
	ZEMBRACE SYMTOUCH (sumatriptan)	
	zolmitriptan	
	zolmitriptan ODT	
	ZOMIG (zolmitriptan)	
	ZOMIG ZMT (zolmitriptan)	
	TRIPTAN COMBINATIONS TREXIMET (sumatriptan/naproxen sodium)	
	TREATIVET (Sumamplan/naproxen socium)	
	OTHER	
NURTEC ODT (rimegepant)*	CAMBIA (diclofenac)	*Nurtec ODT For a diagnosis of Migraine treatment:
	UBRELVY (ubrogepant)**	requires three (3) day trials of two (2) preferred chemically
	REYVOW (lasmiditan)**	distinct triptans before it may be approved, unless one (1) of
		the exceptions on the PA form is present. Maximum Quantity
		limit of 8 tablets per 30 days.
		*Nurtec ODT For a diagnosis of Migraine prophylaxis:
		In addition to the Class Criteria for CGRP PA Criteria, a
		90-day trial of each preferred agent for antimigraine
		prophylaxis is required, unless one (1) of the exceptions on
		propriyianio io required, drilego erio (1) or the exceptions on



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		the PA form is present. Maximum Quantity limit of 16 tablets per 32 days.		
		**Ubrelvy and Reyvow require three (3) day trials of two (2) preferred chemically distinct triptans as well as a three (3) day trial of Nurtec ODT before they may be approved, unless one (1) of the exceptions on the PA form is present.		
ANTIPARASITICS, TOPICALAP				
CLASS PA CRITERIA: Non-preferred agone (1) of the exceptions on the PA form is		age and weight appropriate) before they will be approved, unless		
NATROBA (spinosad) permethrin 5% cream pyrethrins-piperonyl butoxide OTC	ELIMITE CREAM (permethrin) EURAX (crotamiton) ivermectin 0.5% lotion LICE EGG REMOVER OTC (benzalkonium chloride) lindane malathion OVIDE (malathion) SKLICE (ivermectin) spinosad VANALICE (piperonyl/pyrethin)			
ANTIPARKINSON'S AGENTS	, , , , ,			
CLASS PA CRITERIA: Patients starting to before a non-preferred agent will be authorally to the control of the c		d allergy to all preferred agents in the corresponding sub-class,		
	ANTICHOLINERGICS			
benztropine trihexyphenidyl				
	COMT INHIBITORS			
entacapone	COMTAN (entacapone) ONGENTYS (opicapone) TASMAR (tolcapone) tolcapone	COMT Inhibitor agents will only be approved as add-on therapy to a levodopa-containing regimen for treatment of documented motor complications.		
DOPAMINE AGONISTS				
APOKYN (apomorphine) PEN bromocriptine pramipexole ropinirole	KYNMOBI (apomorphine) FILM MIRAPEX ER (pramipexole)* NEUPRO (rotigotine) pramipexole ER ropinirole ER	*Mirapex ER will be authorized for a diagnosis of Parkinsonism without a trial of preferred agents.		
	OTHER ANTIPARKINSON'S AG	ENTS		
amantadine*AP carbidopa/levodopa levodopa/carbidopa/entacapone	AZILECT (rasagiline) carbidopa GOCOVRI ER (amantadine)	*Amantadine will not be authorized for the treatment or prophylaxis of influenza.		

INBRIJA (levodopa)

levodopa/carbidopa ODT

selegiline



BUREAU FOR MEDICAL SERVICES WEST VIRGINIA MEDICAID REFERENCE DELICATION CRITERIA

PREFERRED DRUG LIST WITH PRIOR AUTHORIZATION CRITERIA

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LODOSYN (carbidopa)
NOURIANZ (istradefylline)
OSMOLEX ER (amantadine)
PARLODEL (bromocriptine)
rasagiline
RYTARY (levodopa/carbidopa)
SINEMET (levodopa/carbidopa)
STALEVO (levodopa/carbidopa/entacapone)
XADAGO (safinamide)
ZELAPAR (selegiline)

ANTIPSORIATICS, TOPICAL

CLASS PA CRITERIA: Non-preferred agents require thirty (30) day trials of two (2) preferred unique chemical entities before they will be approved, unless one (1) of the exceptions on the PA form is present.

TACLONEX (calcipotriene/ betamethasone) VECTICAL (calcitriol)

calcipotriene cream
calcipotriene ointment
calcipotriene solution
calcipotriene/betamethasone ointment,
suspension
calcitriol
DOVONEX (calcipotriene)
ENSTILAR (calcipotriene/betamethasone)
SORILUX (calcipotriene)
tazarotene cream

ANTIPSYCHOTICS, ATYPICAL

CLASS PA CRITERIA: All antipsychotic agents require prior authorization for children up to eighteen (18) years of age. All PA requests for antipsychotics for children 6 years of age and younger will be reviewed by Medicaid's consultant psychiatrist.

Non-preferred agents require thirty (30) day trials of two (2) preferred agents, including the generic formulation of the requested agent (if available), before they will be approved unless one (1) of the exceptions on the PA form is present. All trials must be at the maximum recommended dose for the diagnosis provided before they would be considered a failure unless an adverse reaction is documented necessitating a change in therapy.

Patients shall be grandfathered onto their existing therapy, provided the requested agent is being used according to the manufacturer label. Continuation of therapy for an off-label indication or non-standard dosage may be granted a thirty (30) day prior-authorization while the Medical Director reviews the request.

SINGLE INGREDIENT



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ABILIFY MAINTENA (aripiprazole)^{CL} aripiprazole tablets ARISTADA (aripiprazole)^{CL} ARISTADA INITIO (aripiprazole)CL clozapine INVEGA SUSTENNA (paliperidone)CL INVEGA TRINZA (paliperidone)* CL olanzapine olanzapine ODT PERSERIS (risperidone)CL quetiapine ER quetiapine** AP for the 25 mg Tablet Only RISPERDAL CONSTA (risperidone)CL risperidone ODT risperidone solution, tablet ziprasidone

ZYPREXA RELPREVV (olanzapine)

ABILIFY MYCITE (aripiprazole) ABILIFY TABLETS (aripiprazole) ADASUVE (loxapine) aripiprazole solution asenapine sublingual tablets CAPLYTA (lumateperone) clozapine ODT CLOZARIL (clozapine) FANAPT (iloperidone) GEODON (ziprasidone) GEODON IM (ziprasidone) INVEGA ER (paliperidone) LATUDA (lurasidone)*** LYBALVI (olanzapine and samidorphan)^{NR} NUPLAZID (pimavanserin) **** olanzapine IMCL paliperidone ER REXULTI (brexipiprazole) RISPERDAL (risperidone) SAPHRIS (asenapine) SECUADO (asenapine) SEROQUEL (quetiapine) SEROQUEL XR (quetiapine) VERSACLOZ (clozapine) VRAYLAR (capriprazine)***** VRAYLAR DOSE PAK (capriprazine)***** ZYPREXA (olanzapine) ZYPREXA IM (olanzapine)CL

•

The following criteria exceptions apply to the specified products:

*Invega Trinza will be authorized after four months' treatment with Invega Sustenna

- **Quetiapine 25 mg will be authorized:
 - 1. For a diagnosis of schizophrenia or
 - 2. For a diagnosis of bipolar disorder **or**
 - When prescribed concurrently with other strengths of Seroquel in order to achieve therapeutic treatment levels.

Quetiapine 25 mg will not be authorized for use as a sedative hypnotic.

- *** Latuda will be be authorized for the indication of Bipolar Depression with documentation of the diagnosis. All other indications require class criteria to be followed.
- ****Nuplazid may only be authorized for the treatment of Parkinson Disease Induced Psychosis after documented treatment failure with quetiapine.
- ***** Vraylar may be authorized for the indication of Bipolar Depression only after failure of a 30-day trial of Latuda and a 30-day trial of either quetiapine OR a combination of olanzapine + fluoxetine. All other indications require class criteria to be followed.

ATYPICAL ANTIPSYCHOTIC/SSRI COMBINATIONS

olanzapine/fluoxetine

ANTIRETROVIRALS^{AP}

CLASS PA CRITERIA: Non-preferred drugs require medical reasoning beyond convenience or enhanced compliance as to why the clinical need cannot be met with a preferred agent or combination of preferred agents. <u>NOTE</u>: Regimens consisting of preferred agents will result in no more than one additional unit per day over equivalent regimens composed of non-preferred agents. Patients already on a non-preferred regimen shall be grandfathered.

SINGLE TABLET REGIMENS

BIKTARVY (bictegravir/emtricitabine/
tenofovir alafenamide)

COMPLERA (emtricitabine/rilpivirine/tenofovir)

DELSTRIGO (doravirine/lamivudine/
tenofovir df)

GENVOYA (elvitegravir/cobicistat/
emtricitabine/tenofovir)

ATRIPLA (efavirenz/emtricitabine/tenofovir)
DOVATO (dolutegravir/lamivudine)
efavirenz/emtricitabine/tenofovir
JULUCA (dolutegravir/rilpivirine)
SYMTUZA (darunavir/cobicistat/
emtricitabine/tenofovir alafenamide)
STRIBILD (elvitegravir/cobicistat/

- *Stribild requires medical reasoning beyond convenience or enhanced compliance as to why the medical need cannot be met with the the preferred agent Genvoya.
- **Triumeq requires medical reasoning beyond convenience or enhanced compliance as to why the medical need cannot be met with the preferred agents Epzicom and Tivicay.



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ODEFSEY (emtricitabine/rilpivirine/tenofovir) SYMFI (efavirenz/lamivudine/tenofovir) SYMFI LO (efavirenz/lamivudine/tenofovir)	emtricitabine/tenofovir)* TRIUMEQ (abacavir/lamivudine/ dolutegravir)**	
	INTEGRASE STRAND TRANSFER INHIBI	TORS
ISENTRESS (raltegravir potassium) TIVICAY (dolutegravir sodium) TIVICAY PD (dolutegravir sodium)	ISENTRESS HD (raltegravir potassium)	
	NUCLEOSIDE REVERSE TRANSCRIPTASE INHIE	BITORS (NRTI)
abacavir sulfate tablet EMTRIVA (emtricitabine) EPIVIR SOLUTION (lamivudine) lamivudine tenofovir disoproxil fumarate	abacavir sulfate solution didanosine DR capsule emtricitabine capsule EPIVIR TABLET (lamivudine) RETROVIR (zidovudine)	
VIREAD ORAL POWDER (tenofovir disoproxil fumarate) ZIAGEN SOLUTION (abacavir sulfate) zidovudine	stavudine VIDEX EC (didanosine) VIDEX SOLUTION (didanosine) VIREAD TABLETS (tenofovir disoproxil fumarate) ZIAGEN TABLET (abacavir sulfate)	
NO	DN-NUCLEOSIDE REVERSE TRANSCRIPTASE IN	HIBITOR (NNRTI)
SUSTIVA (efavirenz)	EDURANT (rilpivirine) efavirenz etravirine INTELENCE (etravirine) nevirapine nevirapine ER PIFELTRO (doravirine) VIRAMUNE ER 24H (nevirapine) VIRAMUNE SUSPENSION (nevirapine) PHARMACOENHANCER – CYTOCHROME P456	
TYBOST (cobicistat)	PHARMACOENHANCER - CTTOCHROME P430	UINNIBITOR
112001 (oobioistat)	DDOTEASE INDIDITORS (DEPTIDIO	
atazanavir	PROTEASE INHIBITORS (PEPTIDIC	
EVOTAZ (atazanavir/cobicistat) NORVIR (ritonavir) REYATAZ POWDER PACK (atazanavir)	fosamprenavir LEXIVA (fosamprenavir) REYATAZ CAPSULE (atazanavir) ritonavir tablet VIRACEPT (nelfinavir mesylate) PROTEASE INHIBITORS (NON-PEPTIL	DIC)
PREZCOBIX (darunavir/cobicistat)	APTIVUS (tipranavir)	
PREZISTA (darunavir ethanolate)		
	ENTRY INHIBITORS - CCR5 CO-RECEPTOR AN	ITAGONISTS
	SELZENTRY (maraviroc)	
	ENTRY INHIBITORS – FUSION INHIBIT	ORS
	FUZEON (enfuvirtide)	
	COMBINATION PRODUCTS - NRTI:	
abacavir/lamivudine	abacavir/lamivudine/zidovudine	



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CIMDUO (lamivudine/tenofovir) lamivudine/zidovudine	COMBIVIR (lamivudine/zidovudine) EPZICOM (abacavir/lamivudine) TEMIXYS (lamivudine/tenofovir) TRIZIVIR (abacavir/lamivudine/zidovudine)				
COM	IBINATION PRODUCTS - NUCLEOSIDE & NUCLE	OTIDE ANALOG RTIS			
DESCOVY (emtricitabine/tenofovir)	TRUVADA (emtricitabine/tenofovir)* emtricitabine/tenofovir	*Truvada shall be treated as preferred when prescribed for PrEP in members assigned female at birth. Truvada may also be approved over Descovy where guidelines clearly indicate superiority over Descovy (documentation may be required to support the request for PA).			
	COMBINATION PRODUCTS - PROTEASE IN				
KALETRA (lopinavir/ritonavir)	lopinavir/ritonavir				
	GP 120 DIRECTED ATTACHMENT INHIE	BITORS			
RUKOBIA (fostemsavir tromethamine) TABLETS					
ANTIVIRALS, ORAL					
CLASS PA CRITERIA: Non-preferred agents of the exceptions on the PA form is present.	require five (5) day trials of each preferred agent in the	ne same sub-class before they will be approved, unless one (1)			
	ANTI HERPES				
acyclovir valacyclovir	famciclovir SITAVIG (acyclovir) VALTREX (valacyclovir) ZOVIRAX (acyclovir)				
	ANTI-INFLUENZA				
oseltamivir RELENZA (zanamivir) TAMIFLU (oseltamivir)	FLUMADINE (rimantadine) rimantadine XOFLUZA (baloxavir)	In addition to the Class Criteria: The anti-influenza agents will be authorized only for a diagnosis of influenza.			
ANTIVIRALS, TOPICALAP					
•	require a five (5) day trial of the preferred agent before	re they will be approved, unless one (1) of the exceptions on the			
ZOVIRAX CREAM (acyclovir) ZOVIRAX OINTMENT (acyclovir)	acyclovir ointment docosanol cream DENAVIR (penciclovir)				
BETA BLOCKERSAP					
	require fourteen (14) day trials of three (3) chemically will be approved, unless one (1) of the exceptions on	distinct preferred agents, including the generic formulation of the PA form is present.			
	BETA BLOCKERS				
acebutolol atenolol betaxolol bisoprolol CORGARD (nadolol)	BETAPACE (sotalol) BYSTOLIC (nebivolol) HEMANGEOL (propranolol)* INDERAL LA (propranolol) INDERAL XL (propranolol)	*Hemangeol will be authorized for the treatment of proliferating infantile hemangioma requiring systemic therapy.			



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metoprolol metoprolol ER pindolol propranolol SORINE (sotalol) sotalol timolol atenolol/chlorthalidone bisoprolol/HCTZ metoprolol/HCTZ	INNOPRAN XL (propranolol) KAPSPARGO SPRINKLE (metoprolol) LOPRESSOR (metoprolol) nadolol propranolol ER** TENORMIN (atenolol) TOPROL XL (metoprolol) BETA BLOCKER/DIURETIC COMBINA nadolol/bendroflumethiazide TENORETIC (atenolol/chlorthalidone) ZIAC (bisoprolol/HCTZ)	**Propranolol ER shall be authorized for patients with a diagnosis of migraines. Existing users will be grandfathered for use in migraine prophylaxis. ATION DRUGS
propranolol/HCTZ	DETA AND ALBUA DI COV	FD0
	BETA- AND ALPHA-BLOCK	ERS
carvedilol labetalol	carvedilol ER capsule COREG (carvedilol) COREG CR (carvedilol)	
	d agents require thirty (30) day trials of each chemically d	distinct preferred agent before they will be approved, unless one (1) of
the exceptions on the PA form is pres		
GELNIQUE (oxybutynin) oxybutynin IR oxybutynin ER solifenacin TOVIAZ (fesoterodine)	darifenacin ER tablet DETROL (tolterodine) DITROPAN XL (oxybutynin) ENABLEX (darifenacin) flavoxate GEMTESA (vibegron)* MYRBETRIQ (mirabegron) OXYTROL (oxybutynin) tolterodine tolterodine ER trospium trospium ER VESICARE (solifenacin) VESICARE LS (solifenacin)	*In addition to class criteria, a 30-day trial of Myrbetriq (mirabegron) is required prior to Gemtesa approval, unless one (1) of the exceptions on the PA form is present.
BONE RESORPTION SUPP	PRESSION AND RELATED AGENTS	

BONE RESORPTION SUPPRESSION AND RELATED AGENTS

CLASS PA CRITERIA: See below for class criteria.

RI	SPF	201	DЦ	ON	۸Т	EC
- DI	огг	ıvə	пп	OI4	~ .	

alendronate tablets	ACTONEL (risedronate)	Non-preferred agents require thirty (30) day trials of each
ibandronate	alendronate solution	preferred Bisphosphonate agent before they will be approved,
	ATELVIA (risedronate)	unless one (1) of the exceptions on the PA form is present.



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	BINOSTO (alendronate) BONIVA (ibandronate) FOSAMAX TABLETS (alendronate) FOSAMAX PLUS D (alendronate/vitamin D) Risedronate	
	OTHER BONE RESORPTION SUPPRESSION ANI	O RELATED AGENTS
	calcitonin EVISTA (raloxifene)* FORTEO (teriparatide) MIACALCIN (calcitonin) raloxifene* teriparatide TYMLOS (abaloparatide)	Non-preferred agents require a thirty (30) day trial of a preferred Bisphosphonate agent before they will be approved, unless one (1) of the exceptions on the PA form is present. *Raloxifene will be authorized for postmenopausal women with osteoporosis who are at high risk for invasive breast cancer.
BPH TREATMENTS		
	erred agents require thirty (30) day trials of at least two (2) cherent before they will be approved, unless one (1) of the exceptio	nically distinct preferred agents, including the generic formulation ns on the PA form is present.
	5-ALPHA-REDUCTASE (5AR) INHIBITORS A	ND PDE-5 AGENTS
finasteride	AVODART (dutasteride) CIALIS 5 mg (tadalafil) dutasteride PROSCAR (finasteride)	
	ALPHA BLOCKERS	
alfuzosin doxazosin tamsulosin terazosin	CARDURA (doxazosin) CARDURA XL (doxazosin) FLOMAX (tamsulosin) RAPAFLO (silodosin) silodosin	
	5-ALPHA-REDUCTASE (5AR) INHIBITORS/ALPHA	BLOCKER COMBINATION
	dutasteride/tamsulosin JALYN (dutasteride/tamsulosin)	Substitute for Class Criteria : Concurrent thirty (30) day trials of dutasteride and tamsulosin are required before the non-preferred agent will be authorized.
BRONCHODILATORS, B	ETA AGONISTAP	
· ·	erred agents require thirty (30) day trials of each chemically dist	tinct preferred agent in their corresponding sub-class unless one (1)
	INHALATION SOLUTION	
albuterol	arformoterol BROVANA (arformoterol) formoterol levalbuterol metaproterenol PERFOROMIST (formoterol) XOPENEX (levalbuterol)*	*Xopenex Inhalation Solution will be authorized for twelve (12) months for a diagnosis of asthma or COPD for patients on concurrent asthma controller therapy (either oral or inhaled) with documentation of failure on a trial of albuterol or documented intolerance of albuterol, or for concurrent diagnosis of heart disease.



BUREAU FOR MEDICAL SERVICES WEST VIRGINIA MEDICAID

PREFERRED DRUG LIST WITH PRIOR AUTHORIZATION CRITERIA

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INHALERS, LONG-ACTING				
SEREVENT (salmeterol)	STRIVERDI RESPIMAT (olodaterol)			
	INHALERS, SHORT-ACTING			
PROAIR HFA (albuterol) PROAIR RESPICLICK (albuterol) VENTOLIN HFA (albuterol)	albuterol HFA PROAIR DIGIHALER (albuterol) PROVENTIL HFA (albuterol) XOPENEX HFA (levalbuterol)			
	ORAL			
	albuterol ER albuterol IR metaproterenol terbutaline			
CALCIUM CHANNEL DI OCKEDSAP				

CALCIUM CHANNEL BLOCKERS AP

CLASS PA CRITERIA: Non-preferred agents require fourteen (14) day trials of each preferred agent within the corresponding sub-class before they will be approved, unless one (1) of the exceptions on the PA form is present.

	LONG-ACTING	
amlodipine diltiazem ER felodipine ER nifedipine ER verapamil ER	ADALAT CC (nifedipine) CALAN SR (verapamil) CARDIZEM CD, LA (diltiazem) diltiazem LA KATERZIA SUSPENSION (amlodipine)* MATZIM LA (diltiazem) nisoldipine NORVASC (amlodipine) PROCARDIA XL (nifedipine) SULAR (nisoldipine) TIAZAC (diltiazem) verapamil ER PM VERELAN/VERELAN PM (verapamil)	*Katerzia will be authorized for children who are 6-10 years of age who are unable to ingest solid dosage forms. Katerzia may also be authorized for older patients with clinical documentation indicating oral-motor difficulties or dysphagia.
	SHORT-ACTING	
diltiazem verapamil	CARDIZEM (diltiazem) isradipine nicardipine nifedipine nimodipine NYMALIZE SOLUTION (nimodipine) PROCARDIA (nifedipine)	
CEDUAL OCDODING AN	ID DEL ATED ANTIDIOTICS	

CEPHALOSPORINS AND RELATED ANTIBIOTICS

CLASS PA CRITERIA: Non-preferred agents require a five (5) day trial of a preferred agent within the corresponding sub-class before they will be approved, unless one (1) of the exceptions on the PA form is present.



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BETA LACTAMS AND BETA LACTAM/BETA-LACTAMASE INHIBITOR COMBINATIONS			
amoxicillin/clavulanate IR	amoxicillin/clavulanate ER AUGMENTIN (amoxicillin/clavulanate)		
	CEPHALOSPORINS		
cefaclor capsule cefadroxil capsule, tablet cefdinir cefuroxime tablet cephalexin capsule, suspension	cefaclor suspension cefaclor ER tablet cefadroxil suspension cefixime cefpodoxime cefprozil cefuroxime suspension cephalexin tablet KEFLEX (cephalexin) SUPRAX (cefixime)		
COPD AGENTS			
CLASS PA CRITERIA: Non-preferred agents unless one (1) of the exceptions on the PA form		rom the corresponding sub-class before they will be approved,	
	ANTICHOLINERGICAP		
ATROVENT HFA (ipratropium) ipratropium nebulizer solution SPIRIVA (tiotropium) SPIRIVA RESPIMAT (tiotropium) TUDORZA (aclidinium)	INCRUSE ELLIPTA (umeclidinium) LONHALA MAGNAIR (glycopyrrolate) YUPELRI SOLUTION (revefenacin)		
	ANTICHOLINERGIC-BETA AGONIST COMBIN		
ANORO ELLIPTA (umeclidinium/vilanterol) albuterol/ipratropium nebulizer solution BEVESPI (glycopyrrolate/formoterol) COMBIVENT RESPIMAT (albuterol/ipratropium)	DUAKLIR PRESSAIR (aclidinium/formoterol)* STIOLTO RESPIMAT (tiotropium/olodaterol)**	*In addition to the Class PA criteria, Duaklir Pressair requires sixty (60) day trials of each long acting preferred agent, as well as a 60-day trial of Stiolto Respimat. **In addition to the Class PA criteria, Stiolto Respimat requires a sixty (60) day trial of a long acting preferred agent.	
ANTI	CHOLINERGIC-BETA AGONIST-GLUCOCORTICO	ID COMBINATIONS	
	TRELEGY ELLIPTA (fluticasone/umeclidinium/vilanterol)* BREZTRI AEROSPHERE (budesonide/glycopyrrolate/formoterol)**	* Trelegy Ellipta may be prior authorized for patients currently established on the individual components for at least 30 days. **Breztri may be prior authorized for patients currently established on the individual components for at least 30 days.	
	PDE4 INHIBITOR		
	DALIRESP (roflumilast)*	*Daliresp will be authorized if the following criteria are met: 1. Patient is forty (40) years of age or older and 2. Diagnosis of severe chronic obstructive pulmonary disease (COPD) associated with chronic bronchitis and multiple exacerbations requiring systemic glucocorticoids in the preceding six (6) months and	



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		 Concurrent therapy with an inhaled corticosteroid and long-acting bronchodilator and evidence of compliance and No evidence of moderate to severe liver impairment (Child-Pugh Class B or C) and No concurrent use with strong cytochrome P450 inducers (rifampicin, phenobarbital, carbamazepine or phenytoin)
CROHNS DISEASE ORAL STERO	DIDS	
	ORAL	
budesonide ER capsule (generic Entocort EC)	ENTOCORT EC (budesonide)* ORTIKOS (budesonide)*	*Please see the following PDL classes for PDL status of additional agents used for induction and remission (Cytokine and CAM Antagonists/ Immunosuppressives, Oral/ Ulcerative Colitis Agents)
		*Entocort EC and Ortikos may only be authorized if the patient has a documented allergy or intolerance to the generic budesonide 3mg 24-hour capsules.
CYTOKINE & CAM ANTAGONIST	S cl	
exceptions on the PA form is present. Patients	stabilized for at least 6-months on their existing non-	which are indicated for the diagnosis, unless one (1) of the -preferred regimen shall be grandfathered (provided the current II PA criteria may be found on the PA Criteria page by
ENDDEL (storovent)		*For all required the reset and effective alternative will be
ENBREL (etanercept) HUMIRA (adalimumab)	AVSOLA (infliximab)* CIMZIA (certolizumab pegol) INFLECTRA (infliximab)* REMICADE (infliximab)* RENFLEXIS (infliximab)* SIMPONI subcutaneous (golimumab)	*For all requests, the most cost-effective alternative will be approved. Should the provider request a different infliximab product, documentation of contraindication or allergy to the required agent must be provided. As of 10/1/2021, Avsola is the most cost-effective alternative.
	OTHERS	
TALTZ (ixekizumab)* XELJANZ (tofacitinib)**	ACTEMRA subcutaneous (tocilizumab) COSENTYX (secukinumab) ENTYVIO (vedolizumab) ILARIS (canakinumab) ILUMYA (tildrakizumab) KEVZARA (sarilumab) KINERET (anakinra) OLUMIANT (baricitinib)	*Taltz will be authorized for treatment of plaque psoriasis, psoriatic arthritis, and ankylosing spondylitis only after inadequate response to a ninety (90) day trial of one preferred agent. **Xeljanz will only be preferred for the treatment of rheumatoid arthritis and ulcerative colitis. For all other indications it is non preferred. Full PA criteria may be found on the PA Criteria



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SKYRIZI (risankizumab) STELARA subcutaneous (ustekinumab) TREMFYA (guselkumab) XELJANZ XR (tofacitinib)

EPINEPHRINE, SELF-INJECTED

CLASS PA CRITERIA: A non-preferred agent may be authorized with documentation showing the patient's inability to follow the instructions, or the patient's failure to understand the training for the preferred agent(s).

epinephrine (labeler 49502 only)

epinephrine (all labelers except 49502)

EPIPEN (epinephrine)
EPIPEN JR (epinephrine)
SYMJEPI (epinephrine)

ERYTHROPOIESIS STIMULATING PROTEINSCL

CLASS PA CRITERIA: Non-preferred agents require a thirty (30) day trial of a preferred agent before they will be approved, unless one (1) of the exceptions on the PA form is present.

EPOGEN (rl	HuEPO)
RETACRIT ((epoetin alfa)

ARANESP (darbepoetin)
MIRCERA (methoxy PEG-epoetin)
PROCRIT (rHuEPO)

Erythropoiesis agents will be authorized if the following criteria are met:

- Hemoglobin or Hematocrit less than 10/30 respectively. For renewal, hemoglobin or hematocrit levels greater than 12/36 will require dosage reduction or discontinuation. Exceptions will be considered on an individual basis after medical documentation is reviewed. (Lab oratory values must be dated within six (6) weeks of request.) and
- Transferrin saturation ≥ 20%, ferritin levels ≥100 mg/ml, or on concurrent therapeutic iron therapy. (Laboratory values must be dated within three (3) weeks of request. For re-authorization, transferrin saturation or ferritin levels are not required if the patient has been responsive to the erythropoietin agent and
- 3. For HIV-infected patients, endogenous serum erythropoietin level must be \leq 500mU/ml to initiate therapy **and**
- No evidence of untreated GI bleeding, hemolysis, or Vitamin B-12, iron or folate deficiency.



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FLUOROQUINOLONES (Oral)AP			
CLASS PA CRITERIA: Non-preferred agents form is present.	require a five (5) day trial of a preferred agent before the	hey will be approved, unless one (1) of the exceptions on the PA	
CIPRO SUSPENSION (ciprofloxacin) ciprofloxacin levofloxacin tablet	BAXDELA (delafloxacin) CIPRO TABLETS (ciprofloxacin) ciprofloxacin suspension levofloxacin solution moxifloxacin ofloxacin		
GLUCOCORTICOIDS, INHALEDAP			
CLASS PA CRITERIA: Non-preferred agents require thirty (30) day trials of each chemically unique preferred agent before they will be approved, unless one (1) of the exceptions on the PA form is present.			
	GLUCOCORTICOIDS		
ASMANEX TWISTHALER (mometasone) budesonide nebulizer 0.5 mg/2 ml & 0.25 mg/2 ml solution* FLOVENT DISKUS (fluticasone) FLOVENT HFA (fluticasone) PULMICORT FLEXHALER (budesonide)	ARMONAIR DIGIHALER (fluticasone) ALVESCO (ciclesonide) ARNUITY ELLIPTA (fluticasone) ASMANEX HFA (mometasone) budesonide nebulizer 1 mg/2ml solution PULMICORT NEBULIZER SOLUTION (budesonide) QVAR REDIHALER (beclomethasone)	*Budesonide Respules are only preferred for children up to nine (9) years of age. For patients nine (9) and older, prior authorization is required and will be approved only for a diagnosis of severe nasal polyps.	
	GLUCOCORTICOID/BRONCHODILATOR COM	BINATIONS	
ADVAIR DISKUS (fluticasone/salmeterol) ADVAIR HFA (fluticasone/salmeterol) DULERA (mometasone/formoterol) SYMBICORT(budesonide/formoterol)	AIRDUO DIGIHALER (fluticasone/salmeterol) AIRDUO RESPICLICK (fluticasone/salmeterol) budesonide/formoterol BREO ELLIPTA (fluticasone/vilanterol) fluticasone/salmeterol WIXELA (fluticasone/salmeterol)		
GUANYLATE CYCLASE STIMULATORSCL			
	ADEMPAS (riociguat)* VERQUVO (vericiguat)**	*Adempas requires a thirty (30) day trial of a preferred agent from any other PAH Class before it may be approved, unless one (1) of the exceptions on the PA form is present.	
		**Full PA criteria for Verquvo may be found on the <u>PA Criteria</u> page by clicking the hyperlink.	

GROWTH HORMONECL

CLASS PA CRITERIA: Non-preferred agents require three (3) month trials of each preferred agent before they will be approved, unless one (1) of the exceptions on the PA form is present.



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manage	d categories. Refer to cover page for complete list of	rules governing this PDL.
GENOTROPIN (somatropin) NORDITROPIN (somatropin)	INCRELEX (mecasermin) NUTROPIN AQ (somatropin) OMNITROPE (somatropin) SAIZEN (somatropin) SEROSTIM (somatropin) ZOMACTON (somatropin) ZORBTIVE (somatropin)	Patients already on a non-preferred agent will receive authorization to continue therapy on that agent for the duration of the existing PA.
H. PYLORI TREATMENT		
		d components of the requested non-preferred agent and must hey will be approved, unless one (1) of the exceptions on the
Please use individual components: preferred PPI (omeprazole or pantoprazole) amoxicillin tetracycline metronidazole clarithromycin bismuth PYLERA (bismuth/metronidazole/tetracycline)	HELIDAC (bismuth/metronidazole/tetracycline) lansoprazole/amoxicillin/clarithromycin OMECLAMOX-PAK (omeprazole/amoxicillin/clarithromycin) TALICIA (omeprazole/amoxicillin/rifabutin)	
HEPATITIS B TREATMENTS		
	require ninety (90) day trials of each preferred agent b	before they will be approved, unless one (1) of the exceptions on
BARACLUDE SOLUTION (entecavir) * entecavir lamivudine HBV	adefovir BARACLUDE TABLET (entecavir) EPIVIR HBV (lamivudine) HEPSERA (adefovir) VEMLIDY (tenofovir alafenamide fumarate)	*Baraclude <u>solution</u> will be authorized only for patients with documentation of dysphagia.
HEPATITIS C TREATMENTSCL		
CLASS PA CRITERIA: For patients starting to require medical reasoning why a preferred reg		I on the PA Criteria page. Requests for non-preferred regimens
MAVYRET (pibrentasvir/glecaprevir)* ribavirin sofosbuvir/velpatasvir (labeler 72626)* ZEPATIER (elbasvir/grazoprevir)*	EPCLUSA (sofosbuvir/velpatasvir)* HARVONI (ledipasvir/sofosbuvir)* ledipasvir/sofosbuvir* PEGASYS (pegylated interferon) PEG-INTRON (pegylated interferon) RIBASPHERE RIBAPAK (ribavirin) RIBASPHERE 400 mg, 600 mg (ribavirin) SOVALDI (sofosbuvir)* VIEKIRA XR (dasabuvir/ombitasvir/ paritaprevir/ritonavir)* VOSEVI (sofosbuvir/velpatasvir/voxilaprevir)	*Full PA criteria may be found on the PA Criteria page by clicking the hyperlink.



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HYPERPARATHYROID AGENTSAP

CLASS PA CRITERIA: Non-preferred agents require thirty (30) day trials of each preferred agent before they will be approved, unless one (1) of the exceptions on the PA form is present.

paricalcitol capsule cinacalcet

doxercalciferol

HECTOROL (doxercalciferol)

paricalcitol injection RAYALDEE (calcifediol) SENSIPAR (cinacalcet) ZEMPLAR (paricalcitol)

HYPOGLYCEMICS, BIGUANIDES

CLASS PA CRITERIA: Non-preferred agents require a ninety (90) day trial of a preferred agent of similar duration before they will be approved, unless one (1) of the exceptions on the PA form is present.

metformin

metformin ER (generic Glucophage XR)

FORTAMET (metformin ER)
GLUCOPHAGE XR (metformin ER)

GLUMETZA (metformin ER)*
metformin solution (generic Riomet)

metformin ER (generic Glumetza & Fortamet)

RIOMET (metformin)

*Glumetza will be approved only after a 30-day trial of Fortamet.

HYPOGLYCEMICS, DPP-4 INHIBITORS

CLASS PA CRITERIA: Non-preferred agents are available only on appeal.

NOTE: DPP-4 inhibitors will NOT be approved in combination with a GLP-1 agonist.

JANUMET (sitagliptin/metformin)

JANUMET XR (sitagliptin/metformin) JANUVIA (sitagliptin)

JENTADUETO (linagliptin/metformin)

TRADJENTA (linagliptin)

alogliptin

alogliptin/metformin alogliptin/pioglitazone

JENTADUETO XR (linagliptin/metformin)

KAZANO (alogliptin/metformin)

KOMBIGLYZE XR (saxagliptin/metformin)

NESINA (alogliptin)
ONGLYZA (saxagliptin)
OSENI (alogliptin/pioglitazone)

HYPOGLYCEMICS, GLP-1 AGONISTSCL

CLASS PA CRITERIA: Non-preferred agents will only be approved (in 6-month intervals) if ALL of the following criteria has been met:

- 1) Current A1C must be submitted. Agents in this class will not be approved for patients with a starting A1C of less than (<) 7%.
- 2) Documentation demonstrating 90 days of compliance on all current diabetic therapies is provided.
- Documentation demonstrating treatment failure with all unique preferred agents in the same class.

Re-authorizations will require documentation of <u>continued</u> compliance on all diabetic therapies and A1C levels must reach goal, (either an A1C of ≤8%, or demonstrated continued improvement).



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NOTE: GLP-1 agents will NOT be approved in combination with a DPP-4 inhibitor.

OZEMPIC (semaglutide) ADLYXIN (lixisenatide) TRULICITY (dulaglutide) BYETTA (exenatide)

VICTOZA (liraglutide)

BYDUREON BCISE (exenatide)

RYBELSUS (semaglutide)

HYPOGLYCEMICS, INSULIN AND RELATED AGENTS

CLASS PA CRITERIA: Non-preferred agents require a ninety (90) day trial of a pharmacokinetically similar agent before they will be approved, unless one (1) of the exceptions on the PA form is present.

APIDRA (insulin gluisine)AP*

FIASP (insulin aspart)

HUMALOG (insulin lispro)

HUMALOG JR KWIKPEN (insulin lispro)

HUMALOG KWIKPEN U-100 (insulin lispro)

HUMALOG MIX PENS (insulin lispro/lispro protamine)

protamine)

HUMALOG MIX VIALS (insulin lispro/lispro protamine)

HUMULIN N VIAL (insulin)

HUMULIN R U-500 VIAL (insulin)

HUMULIN R U-500 KWIKPEN (insulin)

LANTUS (insulin glargine)

LEVEMIR (insulin detemir)

NOVOLOG (insulin aspart)

NOVOLOG MIX (insulin aspart/aspart

protamine)

TOUJEO SOLOSTAR (insulin glargine)

TOUJEO MAX SOLOSTAR (insulin glargine

ADMELOG (insulin lispro)

AFREZZA (insulin)CL

BASAGLAR (insulin glargine)

HUMALOG KWIKPEN U-200 (insulin lispro)

HUMULIN PENS (insulin)

HUMULIN R VIAL (insulin)

HUMULIN 70/30 (insulin)

insulin aspart

insulin aspart/aspart protamine

insulin lispro

LYUMJEV (insulin lispro)

NOVOLIN (insulin)

SEMGLEE (insulin glargine)

SOLIQUA (insulin glargine/lixisenatide)**

TRESIBA (insulin degludec)***

TRESIBA FLEXTOUCH (insulin degludec)***
XULTOPHY (insulin degludec/liraglutide)**

*Apidra will be authorized if the following criteria are met:

- 1. Patient is four (4) years of age or older; and
- 2. Patient is currently on a regimen including a longer acting or basal insulin, **and**
- Patient has had a trial of a similar preferred agent, Novolog or Humalog, with documentation that the desired results were not achieved..
- ** Non-preferred insulin combination products require that the patient must already be established on the individual agents at doses not exceeding the maximum dose achievable with the combination product, and require medical reasoning beyond convenience or enhanced compliance as to why the clinical need cannot be met with a combination of preferred single-ingredient agents.
- ***Patients stabilized on Tresiba may be grandfathered <u>at the</u> request of the prescriber, if the prescriber considers the preferred products to be clinically inappropriate.
- *** Tresiba U-100 may be approved only for: Patients who have demonstrated at least a 6-month history of compliance on a preferred long-acting insulin and who continue to have regular incidents of hypoglycemia.
- ***Tresiba U-200 may be approved only for: Patients who require once-daily doses of at least 60 units of long-acting insulin and have demonstrated at least a 6-month history of compliance on preferred long-acting insulin and who continue to have regular incidents of hypoglycemia.



HYPOGLYCEMICS MEGLITINIDES

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CLASS PA CRITERIA: Non-preferred agents are available only on appeal.			
	MEGLITINIDES		
nateglinide repaglinide	PRANDIN (repaglinide) STARLIX (nateglinide)		
	MEGLITINIDE COMBINATIONS		
	repaglinide/metformin		
HYPOGLYCEMICS, MISCELLANE	OUS AGENTS		
CLASS PA CRITERIA: Welchol will be authoriz agent.	zed for add-on therapy for type 2 diabetes when there	is a previous history of a thirty (30) day trial of an oral diabetic	
WELCHOL (colesevelam) ^{AP}	colesevelam SYMLIN (pramlintide)*	*Symlin will be authorized with a history of bolus insulin utilization in the past ninety (90) days with no gaps in insulin therapy greater than thirty (30) days.	
HYPOGLYCEMICS, SGLT2 INHIBI	TORS		
•	ill only be approved (in 6-month intervals) if ALL of the	e following criteria has been met:	
 Current A1C must be submitted. Agents in this class will not be approved for patients with a starting A1C of less than (<) 7%. Documentation demonstrating 90 days of compliance on all current diabetic therapies is provided. Documentation demonstrating treatment failure with all unique preferred agents in the same class. Re-authorizations will require documentation of continued compliance on all diabetic therapies and A1C levels must reach goal, (either an A1C of ≤8%, or demonstrated continued improvement).			
*Preferred SGLT2 inhibitors and combinations may be approved for a diagnosis of Heart Failure with Reduced Ejection Fraction (HFrEF) with or without Type II			
DM, Chronic Kidney Disease (CKD) with or without Type II DM, or Atherosclerotic Cardiovascular Disease (ASCVD) with Type II DM without further restrictions.			
SGLT2 INHIBITORS			
FARXIGA (dapagliflozin)* INVOKANA (canagliflozin)* JARDIANCE (empagliflozin)*	STEGLATRO (ertugliflozin)		
INIVOLANTE (consciilo zin/mottormin)*	SGLT2 COMBINATIONS		
INVOKAMET (canagliflozin/metformin)* SYNJARDY (empagliflozin/metformin)*	GLYXAMBI (empagliflozin/linagliptin) INVOKAMET XR (canagliflozin/metformin) SEGLUROMET (ertugliflozin/metformin STEGLUJAN (ertugliflozin/sitagliptin) SYNJARDY XR (empagliflozin/metformin)		



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TRIJARDY XR (empagliflozin/linagliptin/metformin) QTERN (dapagliflozin/saxagliptin) XIGDUO XR (dapagliflozin/metformin) HYPOGLYCEMICS, TZD CLASS PA CRITERIA: Non-preferred agents are available only on appeal. **THIAZOLIDINEDIONES** ACTOS (pioglitazone) pioglitazone AVANDIA (rosiglitazone) **TZD COMBINATIONS** ACTOPLUS MET (pioglitazone/ metformin) Patients are required to use the components of Actoplus Met DUETACT (pioglitazone/glimepiride) and Duetact separately. Exceptions will be handled on a casepioglitazone/glimepiride by-case basis. pioglitazone/ metformin IMMUNOMODULATORS, ATOPIC DERMATITIS CLASS PA CRITERIA: Non-preferred agents require 30-day trial of a medium to high potency topical corticosteroid AND all preferred agents in this class unless one (1) of the exceptions on the PA form is present. Requirement for topical corticosteroids may be excluded with involvement of sensitive areas such as the face and skin folds. ELIDEL (pimecrolimus) DUPIXENT (dupilumab)* *Full PA criteria for Dupixent may be found on the PA Criteria EUCRISA (crisaborole)AP** PROTOPIC (tacrolimus) page by clicking the hyperlink pimecrolimus cream tacrolimus ointment **Eucrisa requires a 30-day trial of Elidel OR a medium to high potency corticosteroid unless contraindicated. **IMMUNOMODULATORS, GENITAL WARTS & ACTINIC KERATOSIS AGENTS** CLASS PA CRITERIA: Non-preferred agents require thirty (30) day trials of each preferred agent before they will be approved, unless one (1) of the exceptions on the PA form is present. CONDYLOX GEL (podofilox) ALDARA (imiguimod) *Zyclara will be authorized for a diagnosis of actinic keratosis. CARAC (fluorouracil) EFUDEX (fluorouracil) imiquimod diclofenac 3% gel fluorouracil 0.5% cream fluorouracil 5% cream podofilox TOLAK (fluorouracil 4% cream) VEREGEN (sinecatechins) ZYCLARA (imiquimod)*

IMMUNOSUPPRESSIVES, ORAL

CLASS PA CRITERIA: Non-preferred agents require a fourteen (14) day trial of a preferred agent before they will be approved, unless one (1) of the exceptions on the PA form is present.



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azathioprine cyclosporine cyclosporine, modified mycophenolate mofetil sirolimus tacrolimus capsule	ASTAGRAF XL (tacrolimus) AZASAN (azathioprine) CELLCEPT (mycophenolate mofetil) ENVARSUS XR (tacrolimus) IMURAN (azathioprine) LUPKYNIS (voclosporin)* mycophenolic acid mycophenolic mofetil suspension MYFORTIC (mycophenolic acid) NEORAL (cyclosporine, modified) PROGRAF (tacrolimus) RAPAMUNE (sirolimus) SANDIMMUNE (cyclosporine) ZORTRESS (everolimus)	*Lupkynis requires a ninety (90) day trial of Benlysta prior to approval. Full PA criteria for Lupkynis may be found on the PA Criteria page by clicking the hyperlink.
INTRANASAL RHINITIS AGENTS	SAP	
CLASS PA CRITERIA: See below for individ	ual sub-class criteria.	
	ANTICHOLINERGICS	
ipratropium	ATROVENT (ipratropium)	Non-preferred agents require thirty (30) day trials of one (1) preferred nasal anti-cholinergic agent, AND one (1) preferred antihistamine AND one (1) preferred intranasal corticosteroid agent before they will be approved, unless one (1) of the exceptions on the PA form is present.
	ANTIHISTAMINES	
azelastine	olopatadine PATANASE (olopatadine)	Non-preferred agents require thirty (30) day trials of one (1) preferred antihistamine AND one (1) preferred intranasal corticosteroid before they will be approved, unless one (1) of the exceptions on the PA form is present.
	COMBINATIONS	
	azelastine/fluticasone DYMISTA (azelastine / fluticasone)	Dymista requires a concurrent thirty (30) day trial of each preferred component before it will be approved, unless one (1) of the exceptions on the PA form is present.
	CORTICOSTEROIDS	
fluticasone propionate OMNARIS (ciclesonide) QNASL HFA (beclomethasone) ZETONNA (ciclesonide)	BECONASE AQ (beclomethasone) flunisolide mometasone NASONEX (mometasone)	Non-preferred agents require thirty (30) day trials of each preferred agent in this sub-class before they will be approved, unless one (1) of the exceptions on the PA form is present
IRRITABLE BOWEL SYNDROME/SHORT BOWEL SYNDROME/SELECTED GI AGENTS CL		
CLASS PA CRITERIA: All agents are approvable only for patients age eighteen (18) and older. See below for additional sub-class criteria.		
	CONSTIPATION	
AMITIZA (lubiprostone) MOVANTIK (naloxegol) LINZESS 145 and 290 mcg (linaclotide)	LINZESS 72 mcg (linaclotide) lubiprostone capsule MOTEGRITY (prucalopride)	All agents in this subclass require documentation of the current diagnosis and evidence that the patient has failed to



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	RELISTOR INJECTION (methylnaltrexone) RELISTOR TABLET (methylnaltrexone) SYMPROIC (naldemedine) TRULANCE (plecanatide) ZELNORM (tegaserod maleate)	find relief with dietary modification and a fourteen (14) day trial of an osmotic laxative. No agent shall be approved to treat opioid induced constipation (OIC) without evidence of at least 90-days of opioid use preceding the request. Continuation of coverage shall be granted with evidence of continuous and concurrent opioid use. Agents may be authorized only for their FDA-approved labeled indication. The following agent-specific criteria shall also apply, unless one (1) of the exceptions on the PA form is present: Motegrity requires a 30-day trial of both Amitiza and Linzess. Relistor and Symproic are indicated for OIC and require thirty (30) day trials of both Movantik and Amitiza. Trulance requires thirty (30) day trials of both Amitiza and Linzess, however for the indication of IBS-C in males, a trial of Amitiza is not required. Linzess 72mcq may only be approved for a diagnosis of chronic idiopathic constipation (CIC) AND for those who cannot tolerate the 145mcg dose. Zelnorm is indicated for females < 65 years of age diagnosed with irritable bowel syndrome with constipation (IBS-C) AND requires thirty (30) day trials of Amitiza and Linzess. Lubiprostone may only be authorized with a documented allergy or intolerance to Amitiza.	
DIARRHEA			
	Alosetron MYTESI (crofelemer) LOTRONEX (alosetron) VIBERZI (eluxadoline)	Full PA criteria may be found on the PA Criteria page by clicking the hyperlink	
LAXATIVES AND CATHARTICS	LAXATIVES AND CATHARTICS		
CLASS PA CRITERIA: Non-preferred agents require thirty (30) day trials of each preferred agent before they will be approved, unless one (1) of the exceptions on the PA form is present			
COLYTE GOLYTELY NULYTELY peg 3350	CLENPIQ (sodium picosulfate, magnesium oxide, citric acid) MOVIPREP OSMOPREP		



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	SUPREP SUTAB (magnesium sulfate, potassium sulfate, sodium sulfate)		
LEUKOTRIENE MODIFIERS			
CLASS PA CRITERIA: Non-preferred agents the PA form is present.	require thirty (30) day trials of each preferred agent be	efore they will be approved, unless one (1) of the exceptions on	
montelukast zafirlukast	ACCOLATE (zafirlukast) SINGULAIR (montelukast) zileuton ZYFLO (zileuton)		
LIPOTROPICS, OTHER (Non-stati	ns)		
CLASS PA CRITERIA: Non-preferred agents require a twelve (12) week trial of a preferred agent before they will be approved, unless one (1) of the exceptions on the PA form is present.			
	BILE ACID SEQUESTRANTSAP		
cholestyramine colestipol tablets	COLESTID (colestipol) colesevelam colestipol granules QUESTRAN (cholestyramine) WELCHOL (colesevelam)*	*Welchol will be authorized for add-on therapy for type 2 diabetes when there is a previous history of a thirty (30) day trial of an oral agent (metformin, sulfonylurea or thiazolidinedione (TZD)). See HYPOGLYCEMICS, MISCELLANEOUS.	
	CHOLESTEROL ABSORPTION INHIBIT	ORS	
ezetimibe	ZETIA (ezetimibe)		
	FATTY ACIDSCL		
omega-3 acid ethyl esters VASCEPA (icosapent ethyl)*	icosapent ethyl capsules LOVAZA (omega-3-acid ethyl esters)	 CLAII agents in this subclass require a prior authorization and an initial triglyceride level ≥ 500 mg/dL. *Additionally, Vascepa may be approved if the following criteria is met: 1. The patient has an initial triglyceride level of ≥ 150 mg/dL prior to start of therapy; AND 2. The patient has established cardiovascular disease or diabetes; AND 3. The patient is concomitantly receiving a statin. 	
FIBRIC ACID DERIVATIVES ^{AP} ANTARA (fonefibrate)			
fenofibrate 54 and 160 mg fenofibrate micronized 67mg, 134mg & 200mg fenofibrate nanocrystallized 48 mg, 145 mg gemfibrozil	ANTARA (fenofibrate) FENOGLIDE (fenofibrate) FIBRICOR (fenofibric acid) fenofibrate 40 mg tablet fenofibrate 150 mg capsules fenofibrate 43, 50, 120 and 130 mg fenofibric acid LIPOFEN (fenofibrate)		



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	LOPID (gemfibrozil) TRICOR (fenofibrate nanocrystallized) TRILIPIX (fenofibric acid)	
	MTP INHIBITORS	
	JUXTAPID (lomitapide)*	*Full PA criteria may be found on the PA Criteria page by clicking the hyperlink.
	NIACIN	
niacin niacin ER (OTC) NIASPAN (niacin)	niacin ER (Rx)	
	PCSK-9 INHIBITORS/BEMPEDOIC	ACID
	PRALUENT (alirocumab)* REPATHA (evolocumab)* NEXLETOL (bempedoic acid)* NEXLIZET (bempedoic acid/ezetimibe)*	*Full PA criteria may be found on the PA Criteria page by clicking the hyperlink.
LIPOTROPICS, STATINSAP		
CLASS PA CRITERIA: See below for individua	ll sub-class criteria.	
	STATINS	
atorvastatin lovastatin pravastatin rosuvastatin simvastatin*	ALTOPREV (Iovastatin) CRESTOR (rosuvastatin) EZALLOR (rosuvastatin) ^{NR} EZALLOR SPRINKLE (rosuvastatin)* fluvastatin fluvastatin ER LESCOL XL (fluvastatin) LIPITOR (atorvastatin) LIVALO (pitavastatin) PRAVACHOL (pravastatin) ZOCOR (simvastatin)** ZYPITAMAG (pitavastatin)	Non-preferred agents require twelve (12) week trials of two (2) preferred agents, including the generic formulation of the requested non-preferred agent, before they will be approved, unless one (1) of the exceptions on the PA form is present. *Ezallor SPRINKLE will only be authorized for those who are unable to ingest solid dosage forms due to documented oralmotor difficulties or dysphagia. **Zocor/simvastatin 80mg tablets will require a clinical PA.
STATIN COMBINATIONS		
	amlodipine/atorvastatin CADUET (atorvastatin/amlodipine) ezetimibe/simvastatin VYTORIN (simvastatin/ezetimibe)*	Non-preferred agents require thirty (30) day concurrent trials of the corresponding preferred single agents before they will be approved, unless one (1) of the exceptions on the PA form is present.
		*Vytorin will be authorized only after an insufficient response to a twelve (12) week trial of the maximum tolerable dose of atorvastatin or rosuvastatin, unless one (1) of the exceptions on the PA form is present.
		Vytorin 80/10mg tablets will require a clinical PA.



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MABS, ANTI-IL/IgE
CLASS PA CRITERIA: For FDA-appr

roved indications, non-preferred agents require a ninety (90) day trial of Xolair. Full PA Criteria may be found on the PA Criteria page by clicking the hyperlink.

XOLAIR (omalizumab)

DUPIXENT (dupilumab) FASENRA (benralizumab) FASENRA PEN (benralizumab)

NUCALA SYRINGE/VIAL (mepolizumab) NUCALA AUTO INJECTOR (mepolizumab)

MACROLIDES

CLASS PA CRITERIA: Non-preferred agents require a five (5) day trial of each preferred agent before they will be approved, unless one (1) of the exceptions on the PA form is present.

MACROLIDES

azithromycin clarithromycin tablets erythromycin base clarithromycin ER clarithromycin suspension

E.E.S. (erythromycin ethylsuccinate) ERYPED (erythromycin ethylsuccinate)

ERY-TAB (erythromycin)

ERYTHROCIN (erythromycin stearate)

ervthromycin estolate ZITHROMAX (azithromycin)

MULTIPLE SCLEROSIS AGENTSCL

CLASS PA CRITERIA: All agents require a prior authorization and documented diagnosis of multiple sclerosis. Preferred oral agents require a ninety (90) day trial of any preferred injectable agent. Non-preferred agents require ninety (90) day trials of two (2) chemically unique preferred agents (in the same sub-class) before they will be approved, unless one (1) of the exceptions on the PA form is present.

INTERFERONS^{AP}

AVONEX (interferon beta-1a) EXTAVIA KIT (interferon beta-1b) AVONEX PEN (interferon beta-1a) EXTAVIA VIAL (interferon beta-1b) BETASERON (interferon beta-1b) PLEGRIDY (peginterferon beta-1a)

REBIF (interferon beta-1a)

REBIF REBIDOSE (interferon beta-1a)

NON-INTERFERONS

AUBAGIO (teriflunomide)* AMPYRA (dalfampridine)** dalfampridine ER**

COPAXONE 20 mg (glatiramer)

GILENYA (fingolimod)

TECFIDERA (dimethyl fumarate)***

BAFIERTAM CAPSULES (monomethyl fumarate)

COPAXONE 40 mg (glatiramer)****

dimethyl fumerate*** alatiramer

GLATOPA (glatiramer)

KESIMPTA INJECTION (ofatumumab)

MAYZENT (siponimod)***** MAVENCLAD (cladribine)

In addition to class PA criteria, the following conditions and criteria may also apply:

*Aubagio requires the following additional criteria to be met:

- 1. Diagnosis of relapsing multiple sclerosis and
- 2. Measurement of transaminase and bilirubin levels within the (6) months before initiation of therapy and ALT levels at least monthly for six (6) months after initiation of therapy and



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PONVORY ((ponesimod)
VUMERITY	(diroximel)
7FPOSIA (o	zanimod)

- Complete blood cell count (CBC) within six (6) months before initiation of therapy and
- Female patients must have a negative pregnancy test before initiation of therapy and be established on a reliable method of contraception if appropriate and
- 5. Patient is between eighteen (18) up to sixty-five (65) years of age **and**
- Negative tuberculin skin test before initiation of therapy

**Dalfampridine ER and Ampyra require the following additional criteria to be met:

- 1. Diagnosis of multiple sclerosis and
- 2. No history of seizures and
- 3. No evidence of moderate or severe renal impairment.

***Dimethyl fumerate and Tecfidera require the following additional criteria to be met:

- 1. Diagnosis of relapsing multiple sclerosis and
- Complete blood count (CBC) within six (6) months of initiation of therapy and six (6) months after initiation and
- 3. Complete blood count (CBC) annually during therapy.

****Copaxone 40mg will only be authorized for documented injection site issues.

*****Mayzent may be authorized with no additional requirement beyond the diagnosis for patients with documented secondary progressive MS.

NEUROPATHIC PAIN

CLASS PA CRITERIA: Non-preferred agents require a thirty (30) day trial of a preferred agent in the corresponding dosage form (oral or topical) before they will be approved, unless one (1) of the exceptions on the PA form is present.

capsaicin OTC duloxetine gabapentin lidocaine patch 5% pregabalin capsule ZTLIDO PATCH (lidocaine) CYMBALTA (duloxetine)
DRIZALMA SPRINKLE (duloxetine)*
GRALISE (gabapentin)**
HORIZANT (gabapentin)
lidocaine patch 4%
LIDODERM (lidocaine)
LYRICA CR (pregabalin)***
LYRICA SOLUTION (pregabalin)***
NEURONTIN (gabapentin)^{AP}
pregabalin ER tablet (generic Lyrica CR)
QUTENZA (capsaicin)

*Drizalma SPRINKLE will only be authorized for those who are unable to ingest solid dosage forms due to documented oral-motor difficulties or dysphagia.

**Gralise will be authorized only if the following criteria are met:

- 1. Diagnosis of post herpetic neuralgia and
- Trial of a tricyclic antidepressant for a least thirty (30) days and
- 90-day trial of gabapentin immediate release formulation (positive response without adequate duration) and



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	SAVELLA (milnacipran)**** LYRICA CAPSULE (pregabalin)	4. Request is for once daily dosing with 1800 mg maximum daily dosage. ***Lyrica CR and Lyrica Solution require medical reasoning beyond convenience as to why the need cannot be met using preferred pregabalin capsules. ****Savella will be authorized for a diagnosis of fibromyalgia only after a 90-day trial of one preferred agent
NSAIDS ^{AP}		
CLASS PA CRITERIA: See below for sub-cl	ass PA criteria.	
	NON-SELECTIVE	
diclofenac (IR, SR)	DAYPRO (oxaprozin)	Non-preferred agents require thirty (30) day trials of each
flurbiprofen ibuprofen (Rx and OTC) INDOCIN SUSPENSION (indomethacin) indomethacin ketoprofen ketorolac meloxicam tablet nabumetone naproxen sodium tablet naproxen suspension EC-naproxen DR tablet piroxicam sulindac	diflunisal DUEXIS (famotidine/ibuprofen) etodolac IR etodolac SR FELDENE (piroxicam) fenoprofen INDOCIN SUPPOSITORIES (indomethacin) indomethacin ER ketoprofen ER meclofenamate mefenamic acid meloxicam submicronized capsule (generic Vivlodex) meloxicam suspension MOBIC TABLET (meloxicam) NALFON (fenoprofen) NAPRELAN (naproxen) naproxen CR oxaprozin RELAFEN DS (nabumetone) SPRIX (ketorolac) TIVORBEX (indomethacin) tolmetin VIVLODEX (meloxicam) VOLTAREN (diclofenac) ZIPSOR (diclofenac potassium) ZORVOLEX (diclofenac)	preferred agent before they will be approved, unless one (1) of the exceptions on the PA form is present.

NSAID/GI PROTECTANT COMBINATIONS



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Non-preferred agents are only available on appeal and require

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ARTHROTEC (diclofenac/misoprostol)

diclofenac/misoprostol naproxen/esomeprazole VIMOVO (naproxen/esomeprazole)	medical reasoning beyond convenience as to why the need cannot be met with the combination of preferred single agents.
COX-II SELECTIVE	
CELEBREX (celecoxib) celecoxib	COX-II Selective agents require thirty (30) day trials of each preferred Non-Selective Oral NSAID, UNLESS the following criteria are met:
	Patient has a history or risk of a serious GI complication; OR Agent is requested for treatment of a chronic condition and 1. Patient is seventy (70) years of age or older, or 2. Patient is currently on anticoagulation therapy.
TOPICAL	
diclofenac patch diclofenac solution LICART PATCH (diclofenac) PENNSAID (diclofenac)	*Flector patches are limited to two per day. **diclofenac gel will be limited to 100 grams per month. Non-preferred agents require a thirty (30) day trial of the preferred Topical agent and thirty (30) day trials of each preferred oral NSAID before they will be approved, unless one(1) of the exceptions on the PA form is present.
require three (3) day trials of each preferred agent be	efore they will be approved, unless one (1) of the exceptions on
AZASITE (azithromycin) bacitracin BLEPH-10 (sulfacetamide) BESIVANCE (besifloxacin)* CILOXAN (ciprofloxacin) gatifloxacin moxifloxacin** NATACYN (natamycin) neomycin/polymyxin/gramicidin OCUFLOX (ofloxacin) POLYTRIM (polymyxin/trimethoprim) sulfacetamide drops sulfacetamide ointment TOBREX (tobramycin) VIGAMOX (moxifloxacin) ZYMAXID (gatifloxacin) OID COMBINATIONSAP	*Prior authorization of any fluoroquinolone agent requires three (3) day trials of all other preferred agents unless definitive laboratory cultures exist indicating the need to use a fluoroquinolone.
	diclofenac/misoprostol naproxen/esomeprazole VIMOVO (naproxen/esomeprazole) COX-II SELECTIVE CELEBREX (celecoxib) celecoxib TOPICAL diclofenac patch diclofenac solution LICART PATCH (diclofenac) PENNSAID (diclofenac) require three (3) day trials of each preferred agent be AZASITE (azithromycin) bacitracin BLEPH-10 (sulfacetamide) BESIVANCE (besifloxacin)* CILOXAN (ciprofloxacin) gatifloxacin moxifloxacin** NATACYN (natamycin) neomycin/polymyxin/gramicidin OCUFLOX (ofloxacin) POLYTRIM (polymyxin/trimethoprim) sulfacetamide drops sulfacetamide ointment TOBREX (tobramycin) VIGAMOX (moxifloxacin) ZYMAXID (gatifloxacin)

OPHIHALMIC ANTIBIOTIC/STEROID COMBINATIONS

CLASS PA CRITERIA: Non-preferred agents require three (3) day trials of each preferred agent before they will be approved, unless one (1) of the exceptions on the PA form is present.



BUREAU FOR MEDICAL SERVICES **WEST VIRGINIA MEDICAID**

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neomycin/polymyxin/dexamethasone sulfacetamide/prednisolone TOBRADEX OINTMENT (tobramycin/ dexamethasone) TOBRADEX SUSPENSION (tobramycin/ dexamethasone) ZYLET (loteprednol/tobramycin)

BLEPHAMIDE (prednisolone/sulfacetamide) MAXITROL ointment (neomycin/polymyxin/ dexamethasone) MAXITROL suspension (neomycin/polymyxin/ dexamethasone) neomycin/bacitracin/polymyxin/ hydrocortisone neomycin/polymyxin/hydrocortisone PRED-G (prednisolone/gentamicin) TOBRADEX ST (tobramycin/ dexamethasone) tobramycin/dexamethasone suspension

OPHTHALMICS FOR ALLERGIC CONJUNCTIVITISAP

CLASS PA CRITERIA: Non-preferred agents require thirty (30) day trials of three (3) preferred chemically unique agents before they will be approved, unless one (1) of the exceptions on the PA form is present.

ALAWAY (ketotifen) ALREX (loteprednol) BEPREVE (bepotastine) cromolyn

ketotifen LASTACAFT (alcaftadine) olopatadine 0.1% (Generic PATANOL labeler 61314 only)

ZADITOR OTC (ketotifen)

ALOCRIL (nedocromil) ALOMIDE (lodoxamide) azelastine

bepotastine

Epinastine

LUMIFY (brimonidine)

olopatadine 0.1% (all formulations except Generic

PATANOL labeler 61314) olopatadine 0.2% (all labelers) PATANOL (olopatadine) ZERVIATE (cetirizine)

OPHTHALMICS, ANTI-INFLAMMATORIES-IMMUNOMODULATORSCL

CLASS PA CRITERIA: All agents require a prior authorization. Non-preferred agents require a 60-day trial of the preferred agent(s).

RESTASIS (cyclosporine)

CEQUA (cyclosporine) EYSUVIS (loteprednol) RESTASIS MULTIDOSE (cyclosporine)* XIIDRA (lifitegrast)

*Restasis Multidose is approvable only on appeal and requires medical reasoning as to why the clinical need cannot be met with the preferred product (Restasis).

All agents must meet the following prior-authorization criteria:

- 1.) Patient must be sixteen (16) years of age or greater;
- 2.) Prior Authorization must be requested by an ophthalmologist or optometrist; AND
- 3.) Clinically diagnosed tear deficiency due to ocular inflammation in patients with keratoconjunctivitis sicca or dry eye syndrome (also known as dry eye); AND
- 4.) Patient must have a functioning lacrimal gland; AND
- 5.) Patient using artificial tears at least four (4) times a day over the last thirty (30) days; AND Patient must not have an active ocular infection



BUREAU FOR MEDICAL SERVICES WEST VIRGINIA MEDICAID REFERENCE DELICATION CRITERIA

PREFERRED DRUG LIST WITH PRIOR AUTHORIZATION CRITERIA
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OPHTHALMICS, ANTI-INFLAMMATORIES

CLASS PA CRITERIA: Non-preferred agents require five (5) day trials of at least two (2) preferred agents before they will be approved, unless one (1) of the exceptions on the PA form is present. Trials must include at least one agent with the same mechanism of action as the requested non-preferred agent.

dexamethasone ACULAR (ketorolac) diclofenac ACULAR LS (ketorolac) ACUVAIL (ketorolac tromethamine) DUREZOL (difluprednate) fluorometholone bromfenac FML FORTE (fluorometholone) BROMSITE (bromfenac) FML S.O.P. (fluorometholone) FLAREX (fluorometholone) ketorolac flurbiprofen LOTEMAX DROPS, OINTMENT (loteprednol) FML (fluorometholone) MAXIDEX (dexamethasone) ILEVRO (nepafenac) NEVANAC (nepafenac) INVELTYS (loteprednol) PRED MILD (prednisolone) LOTEMAX GEL (loteprednol) prednisolone acetate loteprednol drops, gel OMNIPRED (prednisolone) prednisolone sodium phosphate OZURDEX (dexamethasone) PRED FORTE (prednisolone) PROLENSA (bromfenac) RETISERT (fluocinolone) TRIESENCE (triamcinolone)

OPHTHALMICS, GLAUCOMA AGENTS

CLASS PA CRITERIA: Non-preferred agents will only be authorized if there is an allergy to all preferred agents in the corresponding sub-class.

CLASS PA CRITERIA: Non-preferred agents v	will only be authorized if there is an allergy to all prefer	red agents in the corresponding sub-class.
	COMBINATION AGENTS	
COMBIGAN (brimonidine/timolol) dorzolamide/timolol SIMBRINZA (brinzolamide/brimonidine)	COSOPT PF (dorzolamide/timolol)	
	BETA BLOCKERS	
BETOPTIC S (betaxolol) carteolol levobunolol timolol drops	betaxolol ISTALOL (timolol) timolol gel TIMOPTIC (timolol)	
CARBONIC ANHYDRASE INHIBITORS		
AZOPT (brinzolamide) dorzolamide	brinzolamide TRUSOPT (dorzolamide)	
PARASYMPATHOMIMETICS		
PHOSPHOLINE IODIDE (echothiophate iodide)	pilocarpine	
PROSTAGLANDIN ANALOGS		
latanoprost TRAVATAN-Z (travoprost)	bimatoprost LUMIGAN (bimatoprost) travoprost VYZULTA (latanoprostene)*	*Vyzulta – prior authorization requires failure on a 3-month trial of at least one preferred prostaglandin eye drop used in combination with an agent from another subclass.



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	XALATAN (latanoprost) XELPROS (latanoprost) ZIOPTAN (tafluprost)	
	RHO-KINASE INHIBITORS	
RHOPRESSA (netarsudil) ROCKLATAN (netarsudil/latanoprost)		
huine anidine 0.20/	SYMPATHOMIMETICS	
brimonidine 0.2%	ALPHAGAN P 0.1% Solution (brimonidine) ALPHAGAN P 0.15% Solution (brimonidine) apraclonidine IOPIDINE (apraclonidine)	
OPIATE DEPENDENCE TREATME		
CLASS PA CRITERIA: Bunavail and Zubsolv rablets.		or allergy to Suboxone strips AND buprenorphine/naloxone Buprenorphine Coverage Policy and Related Forms
buprenorphine/naloxone tablets* naloxone NARCAN NASAL SPRAY (naloxone) SUBLOCADE (buprenorphine soln) ^{CL} SUBOXONE FILM (buprenorphine/naloxone)* VIVITROL (naltrexone)	BUNAVAIL (buprenorphine/naloxone) buprenorphine tablets buprenorphine/naloxone film LUCEMYRA (lofexidine) ZUBSOLV (buprenorphine/naloxone)	* Full PA criteria may be found on the PA Criteria page by clicking the hyperlink.
OTIC ANTIBIOTICSAP		
CLASS PA CRITERIA: Non-preferred agents re PA form is present.	equire five (5) day trials of each preferred agent befo	re they will be approved, unless one (1) of the exceptions on the
CIPRO HC (ciprofloxacin/hydrocortisone) CIPRODEX (ciprofloxacin/dexamethasone) ofloxacin CORTISPORIN-TC (colistin/hydrocortisone/ neomycin)	ciprofloxacin ciprofloxacin/dexamethasone) ciprofloxacin/fluocinolone neomycin/polymyxin/HC solution/suspension OTOVEL (ciprofloxacin/fluocinolone)	
PAH AGENTS - ENDOTHELIN RE	CEPTOR ANTAGONISTS ^{CL}	
CLASS PA CRITERIA: Non-preferred agents r PA form is present.	equire a thirty (30) day trial of a preferred agent before	re they will be approved, unless one (1) of the exceptions on the
LETAIRIS (ambrisentan) TRACLEER TABLET (bosentan)	ambrisentan bosentan OPSUMIT (macitentan) TRACLEER SUSP (bosentan)	
PAH AGENTS - PDE5scl		



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CLASS PA CRITERIA: Non-preferred agents require a thirty (30) day trial of a preferred agent before they will be approved, unless one (1) of the exceptions on the PA form is present.

Patients stabilized on non-preferred agents will be grandfathered.

sildenafil tablets

ADCIRCA (tadalafil) REVATIO IV (sildenafil)

REVATIO SUSPENSION (sildenafil) REVATIO TABLETS (sildenafil) sildenafil suspension (generic Revatio)

PAH AGENTS - PROSTACYCLINSCL

CLASS PA CRITERIA: Non-preferred agents require a thirty (30) day trial of a preferred agent, including the preferred generic form of the non-preferred agent (if available), before they will be approved, unless one (1) of the exceptions on the PA form is present.

epoprostenol (generic Flolan)

VENTAVIS (iloprost)*

epoprostenol (generic Veletri) FLOLAN (epoprostenol) ORENITRAM ER (treprostinil) REMODULIN (treprostinil sodium)

TYVASO (treprostinil) UPTRAVI (selexipag) VELETRI (epoprostenol) *Ventavis will only be authorized for the treatment of pulmonary artery hypertension (WHO Group 1) in patients with NYHA Class III or IV symptoms.

PANCREATIC ENZYMESAP

CLASS PA CRITERIA: Non-preferred agents require a thirty (30) day trial of a preferred agent before they will be approved, unless one (1) of the exceptions on the PA form is present.

For members with cystic fibrosis, a trial of a preferred agent will not be required.

CREON PANCREAZE ZENPEP PERTZYE VIOKACE

PHOSPHATE BINDERSAP

CLASS PA CRITERIA: Non-preferred agents require a thirty (30) day trial of at least two (2) preferred agents before they will be approved, unless one (1) of the exceptions on the PA form is present.

calcium acetate

CALPHRON (calcium acetate)

MAGNEBIND RX (calcium carbonate, folic acid, magnesium carbonate)

PHOSLYRA (calcium acetate) sevelamer carbonate

AURYXIA (ferric citrate) FOSRENOL (lanthanum) lanthanum chewable RENAGEL (sevelamer)

RENVELA (sevelamer carbonate) sevelamer carbonate powder packet VELPHORO (sucroferric oxyhydroxide)

PITUITARY SUPPRESSIVE AGENTS, LHRHCL

CLASS PA CRITERIA: Unless otherwise noted, non-preferred agents are available only on appeal.



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LUPANETA (leuprolide) LUPRON DEPOT KIT (leuprolide) LUPRON DEPOT-PED KIT (leuprolide) ORILISSA (elagolix)* ORIAHNN (elagolix-estradiol-norethindrone)* SYNAREL (nafarelin) TRELSTAR (triptorelin) TRIPTODUR (triptorelin) VANTAS (histrelin) ZOLADEX (goserelin)	leuprolide MYFEMBREE (relugolix, estradiol, norethindrone) SUPPRELIN LA KIT (histrelin)	* Full PA criteria may be found on the PA Criteria page by clicking the hyperlink.
PLATELET AGGREGATION INHIB	SITORS	
CLASS PA CRITERIA: Non-preferred agents reparted pagents in the present.	equire a thirty (30) day trial of a preferred agent before	e they will be approved, unless one (1) of the exceptions on the
BRILINTA (ticagrelor) clopidogrel dipyridamole prasugrel	clopidogrel kit dipyridamole/aspirin EFFIENT (prasugrel) PLAVIX (clopidogrel) ZONTIVITY (vorapaxar)	
PROGESTATIONAL AGENTS		
CLASS PA CRITERIA: Full PA criteria may be	found on the PA Criteria page by clicking the hyperlin	k.
MAKENA (hydroxyprogesterone caproate) AUTO INJECTOR MAKENA (hydroxyprogesterone caproate) VIAL	hydroxyprogesterone caproate	
PROGESTINS FOR CACHEXIA		
CLASS PA CRITERIA: Non-preferred agents re PA form is present.	equire a thirty (30) day trial of a preferred agent before	e they will be approved, unless one (1) of the exceptions on the
Megestrol		
PROTON PUMP INHIBITORSAP		
		d pantoprazole at the maximum recommended dose*, inclusive ved, unless one (1) of the exceptions on the PA form is present.
NEXIUM PACKETS (esomeprazole)** omeprazole (Rx) pantoprazole PROTONIX GRANULES (pantoprazole)**	ACIPHEX (rabeprazole) ACIPHEX SPRINKLE (rabeprazole) DEXILANT (dexlansoprazole) esomeprazole magnesium lansoprazole Rx NEXIUM (esomeprazole) omeprazole/sodium bicarbonate (Rx) PREVACID CAPSULES (lansoprazole) PREVACID SOLUTABS (lansoprazole)** PRILOSEC Rx (omeprazole)	*Maximum recommended doses of the PPIs and H2-receptor antagonists may be located at the BMS Pharmacy PA criteria page titled "Max PPI and H2RA" by clicking on the hyperlink. **Prior authorization is required for members nine (9) years of age or older for these agents.



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	PROTONIX DR TABLETS (pantoprazole) rabeprazole ZEGERID Rx (omeprazole/sodium bicarbonate)	
SEDATIVE HYPNOTICSAP		
of the exceptions on the PA form is present. A	require thirty (30) day trials of all preferred agents in Ell agents except melatonin will be limited to fifteen (15) without a PA. Melatonin labeler code 51645 is preferred BENZODIAZEPINES	BOTH sub-classes before they will be approved, unless one (1) tablets in a thirty (30) day period. NOTE: WV Medicaid covers ed if available, however all NDCs are payable.
temazepam 15, 30 mg	estazolam	
Tomas paint of the same	flurazepam HALCION (triazolam) RESTORIL (temazepam) temazepam 7.5, 22.5 mg triazolam	
	OTHERS	
melatonin ROZEREM (ramelteon) zolpidem 5, 10 mg	AMBIEN (zolpidem) AMBIEN CR (zolpidem) BELSOMRA (suvorexant) DAYVIGO (lemborexant) EDLUAR (zolpidem) eszopiclone HETLIOZ (tasimelteon) ^{CL*} LUNESTA (eszopiclone) ramelteon SILENOR (doxepin) zaleplon zolpidem ER 6.25, 12.5 mg	Strengths of zolpidem that are non-preferred (6.25 and 12.5 mg) must be created by combining or splitting the preferred doses (5 and 10 mg) of zolpidem, if appropriate. For treatment naïve female patients, zolpidem and zolpidem ER maximum dosages will be limited to 5 mg and 6.25 mg respectively per day. *Full PA criteria may be found on the PA Criteria page by clicking the hyperlink.
SKELETAL MUSCLE RELAXANT	TS AP	
CLASS PA CRITERIA: See below for individ	ual sub-class criteria.	
	ACUTE MUSCULOSKELETAL RELAXANT	AGENTS
chlorzoxazone (generic PARAFON FORTE) cyclobenzaprine IR 5, 10 mg methocarbamol	AMRIX (cyclobenzaprine) carisoprodol* carisoprodol/ASA* carisoprodol/ASA/codeine* chlorzoxazone (generic LORZONE) cyclobenzaprine ER cyclobenzaprine IR 7.5 mg FEXMID (cyclobenzaprine) LORZONE (chlorzoxazone) metaxalone orphenadrine	Non-preferred agents require thirty (30) day trials of each preferred agent before they will be approved, unless one (1) of the exceptions on the PA form is present, with the exception of carisoprodol. *Carisoprodol requires thirty (30) day trials of each of the preferred acute musculoskeletal relaxants and Skelaxin before it will be approved.



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	orphenadrine ER ROBAXIN (methocarbamol) SKELAXIN (metaxalone) SOMA (carisoprodol)	
M	USCULOSKELETAL RELAXANT AGENTS USED F	OR SPASTICITY
baclofen tizanidine tablets	DANTRIUM (dantrolene) dantrolene tizanidine capsules ZANAFLEX (tizanidine)	Non-preferred agents require thirty (30) day trials of each preferred agent before they will be approved, unless one (1) of the exceptions on the PA form is present.

STEROIDS, TOPICAL

CLASS PA CRITERIA: Non-preferred agents require five (5) day trials of one (1) form of EACH preferred unique active ingredient in the corresponding potency

	VERY HIGH & HIGH POTENCY
group before they will be approved, unless on betamethasone dipropionate cream betamethasone valerate cream betamethasone valerate lotion betamethasone valerate oint clobetasol propionatecream, gel, ointment, solution clobetasol emollient clobetasol propionate shampoo fluocinonide gel triamcinolone acetonide cream, ointment triamcinolone acetonide lotion	

TOPICORT SPRAY (desoximetasone)



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	TOVET FOAM (clobetasol) ULTRAVATE (halobetasol propionate) ULTRAVATE PAC cream VANOS (fluocinonide)	
	MEDIUM POTENCY	
fluticasone propionate cream, ointment mometasone furoate triamcinolone acetonide 0.025% and 0.1% cream	BESER LOTION (fluticasone) betamethasone valerate foam CLODERM (clocortolone pivalate) clocortolone cream CORDRAN (flurandrenolide) CUTIVATE (fluticasone propionate) fluocinolone acetonide cream, ointment, solution fluticasone propionate lotion hydrocortisone butyrate cream hydrocortisone butyrate ointment, solution hydrocortisone valerate LOCOID (hydrocortisone butyrate) LOCOID LIPOCREAM (hydrocortisone butyrate/emollient) LUXIQ (betamethasone valerate) PANDEL (hydrocortisone probutate) prednicarbate	
	LOW POTENCY	
DERMA-SMOOTHE FS (fluocinolone acetonide) hydrocortisone acetate (Rx, OTC) hydrocortisone cream (Rx, OTC) hydrocortisone lotion OTC hydrocortisone ointment (Rx, OTC) hydrocortisone solution OTC hydrocortisone-aloe cream OTC hydrocortisone-aloe ointment OTC	alclometasone dipropionate AQUA GLYCOLIC HC (hydrocortisone) CAPEX (fluocinolone acetonide) DESONATE (desonide) desonide cream, ointment desonide lotion fluocinolone oil hydrocortisone/mineral oil/petrolatum hydrocortisone acetate/urea hydrocortisone lotion hydrocortisone/aloe gel SCALPICIN OTC (hydrocortisone) SYNALAR (fluocinolone) TEXACORT (hydrocortisone)	

STIMULANTS AND RELATED AGENTS

CLASS PA CRITERIA: A PA is required for adults eighteen (18) years of age or older.

Non-preferred agents require a thirty (30) day trial of at least one preferred agent in the same subclass and with a similar duration of effect and mechanism of action, unless one (1) of the exceptions on the PA form is present. **NOTE**: Non-preferred agents will NOT be "grandfathered" for adults. Children under the age of 18 may continue their current therapy until the end of the school year after which they will be required to switch to a preferred agent.

AMPHETAMINES



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amphetamine salt combination ER amphetamine salt combination IR dextroamphetamine ER dextroamphetamine IR VYVANSE CHEWABLE (lisdexamfetamine) VYVANSE CAPSULE (lisdexamfetamine)

ADDERALL (amphetamine salt combination)
ADDERALL XR (amphetamine salt combination)
ADZENYS XR ODT (amphetamine)
ADZENYS ER SUSP (amphetamine)
amphetamine tablets
DESOXYN (methamphetamine)
DEXEDRINE ER (dextroamphetamine)
dextroamphetamine solution
DYANAVEL XR SUSP (amphetamine)
EVEKEO (amphetamine)
EVEKEO ODT (amphetamine)
methamphetamine
MYDAYIS (dextroamphetamine/amphetamine

In addition to the Class Criteria: Thirty (30) day trials of at least three (3) antidepressants are required before amphetamines will be authorized for depression.

*Mydayis requires a 30-day trial of at least one long-acting preferred agent in this subclass and a trial of Adderall XR.

NON-AMPHETAMINE

atomoxetine
CONCERTA (methylphenidate)
clonidine IR
dexmethylphenidate IR
FOCALIN XR (dexmethylphenidate)
guanfacine ER
guanfacine IR
methylphenidate IR
methylphenidate ER tablet (generic RITALIN
SR)
methylphenidate solution
QUILLICHEW ER (methylphenidate)
QUILLIVANT XR (methylphenidate)

ADHANSIA XR (methylphenidate) APTENSIO XR (methylphenidate)

(dextroamphetamine)ZENZEDI

AZSTARYS

salt)*

PROCENTRA solution

(dextroamphetamine)

(dexmethylphenidate;serdexmethylphenidate)NF

clonidine ER

COTEMPLA XR ODT (methylphenidate)

DAYTRANA (methylphenidate)

dexmethylphenidate XR

FOCALIN IR (dexmethylphenidate)

INTUNIV (guanfacine extended-release)

JORNAY PM (methylphenidate)

METHYLIN SOLUTION (methylphenidate)

methylphenidate CD

methylphenidate chewable tablets

methylphenidate ER 24 tablet (generic

CONCERTA)

methylphenidate ER capsule methylphenidate LA capsule

QELBREE (viloxazine)**

RITALIN (methylphenidate)
RITALIN LA (methylphenidate)
STRATTERA (atomoxetine)*

* Strattera is limited to a maximum of 100 mg per day.

**Full PA criteria may be found on the <u>PA Criteria</u> page by clicking the hyperlink.

NARCOLEPTIC AGENTS

armodafinil^{CL} modafinil^{CL} NUVIGIL (armodafinil) PROVIGIL (modafinil) SUNOSI (solriamfetol)* * Sunosi is approvable only with documentation of treatment failure after 30-day trials of both armodafinil and modafinil.



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	WAKIX (pitolisant)**	**Wakix is approvable only with documentation of treatment failure after 30-day trials of armodafinil, modafinil and Sunosi.
TETRACYCLINES		·
CLASS PA CRITERIA: Non-preferred age PA form is present.	nts require ten (10) day trials of each preferred agent befo	ore they will be approved, unless one (1) of the exceptions on the
doxycycline hyclate capsules doxycycline hyclate 100 mg tablets doxycycline monohydrate 50, 100 mg capsules minocycline capsules	demeclocycline* DORYX (doxycycline hyclate) doxycycline hyclate 75, 150 mg tablets doxycycline hyclate tablet DR 75, 100, 150, 200 mg doxycycline hyclate tablet DR 50 mg doxycycline monohydrate 40, 75, 150 mg capsule doxycycline monohydrate tablet doxycycline monohydrate suspension MINOCIN (minocycline) minocycline ER capsules minocycline tablets MINOLIRA ER (minocycline) MORGIDOX KIT (doxycycline) ORACEA (doxycycline monohydrate) SOLODYN (minocycline) tetracycline VIBRAMYCIN CAPSULES, SUSPENSION, SYRUP (doxycycline) XIMINO (minocycline)	*Demeclocycline will be authorized for conditions caused by susceptible strains of organisms designated in the product information supplied by the manufacturer. A C&S report must accompany this request. Demeclocycline will also be authorized for SIADH.
ULCERATIVE COLITIS AGENT	SAP	

CLASS PA CRITERIA: Non-preferred agents require thirty (30) day trials of each preferred dosage form or chemical entity before the corresponding non-preferred agent of that dosage form or chemical entity will be approved, unless one (1) of the exceptions on the PA form is present.

ORAL.		
APRISO (mesalamine) ASACOL HD (mesalamine) balsalazide PENTASA (mesalamine) 250 mg PENTASA (mesalamine) 500 mg sulfasalazine	AZULFIDINE (sulfasalazine) budesonide ER tablet COLAZAL (balsalazide) DELZICOL (mesalamine) DIPENTUM (olsalazine) LIALDA (mesalamine) mesalamine UCERIS (budesonide) ZEPOSIA (ozanimod)	
RECTAL		
mesalamine	DELZICOL DR (mesalamine) mesalamine kit ROWASA (mesalamine) SF ROWASA (mesalamine) UCERIS (budesonide)	



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VASODILATORS, CORONARY

CLASS PA CRITERIA: Non-preferred agents require thirty (30) day trials of each preferred dosage form before they will be approved, unless one (1) of the exceptions on the PA form is present.

SUBLINGUAL NITROGLYCERIN

nitroglycerin spray (generic NITROLINGUAL) nitroglycerin sublingual

NITROSTAT SUBLINGUAL (nitroglycerin)

GONITRO SPRAY POWDER (nitroglycerin) nitroglycerin spray (generic NITROMIST) NITROLINGUAL SPRAY (nitroglycerin) NITROMIST (nitroglycerin)

MISCELLANEOUS COVERED AGENTS

This category contains covered agents which either did not easily fit into a single PDL category or had criteria that was too lengthy to cite within the PDL itself. Full criteria for the agents listed below may be found by following this hyperlink: (https://dhhr.wv.gov/bms/BMS%20Pharmacy/Pages/PA-Criteria.aspx). Please note that some agents may be available only by billing the appropriate HCPCS code noted in the criteria.

Afinitor

Albenza and Emverm

Ampyra

Antifungal Agents

Austedo

Belbuca

Benlysta

Botox

Cabenuva

Carbaglu

CGRP Receptor Antagonists

Continuous Glucose Monitors

Corlanor

Cresemba

Cuvposa

Cytokine & CAM Antagonists

Diclegis

Dificid

Dojolvi

Droxidopa

Duavee

Dupixent

Epidiolex

Emflaza

Enspryng

Esbriet

Evrysdi

ExJade

Exondys 51



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Fasenra
Ferriprox
Firazyr
Fuzeon
Gattex
Gralise
Growth Hormone for Adults
Growth Hormone for Children
Hepatitis C PA Criteria
Hereditary Angioedema Agents
Hetlioz
Home Infusion Drugs and Supplies
Horizant
HP Acthar
HyQvia
Increlex
Ingrezza
Jublia
Juxtapid
Kalydeco
Ketoconazole
Korlym
Kuvan
Kymriah
Kynamro
Lucemyra
Lutathera
Lupkynis
Luxturna
Makena
Max PPI an H2RA
Mozobil
Myalept
Mytesi
Natpara
Nexletol and Nexlizet
Non-Sedating Antihistamines
Nuvigil
Nucala
OFEV
Oforta
Omnipod
Orilissa
Oralair
Oriahnn

Orkambi Osphena



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Oxlumo
Palforzia
Palynziq
PCSK9 Inhibitor
Provigil
Qbrexza
Qelbree
Rectiv
Regranex
Restasis
Rilutek
Riluzole
Risperdal Consta
Ruconest
Sirturo
Spinraza
Spravato
Sprycel
Suboxone Policy
Symdeko
Synagis
Testosterone
Thalomid
Tobacco Cessation Policy
Trikafta
V-Go
Viberzi and Lotronex
Verquvo
Vyondys 53
Xanax XR
Xenazine
Xhance
Xifaxan
Xolair Xonana and Youngain
Xyrem and Xywav
Yescarta
Zolgensma
Zulresso
Zurampic

Zyvox