Ohio Medicaid

Pharmacy Benefit Management Program



Unified Preferred Drug List

Medicaid Fee-for-Service and Managed Care Plans

Effective July 1, 2025

Helpful Links

Prior Authorization (PA)

Prior Authorization (PA) Information | medicaid.ohio.gov

- General Prior Authorization Requirements
- PA and Step Therapy Frequently Asked Questions (FAQ)

Unified Preferred Drug List (UPDL)

Ohio Unified Preferred Drug List | medicaid.ohio.gov

Unified Preferred Drug List (UPDL)

General Information

- The Statewide UPDL is not an all-inclusive list of drugs covered by the Ohio Department of Medicaid (ODM). Non UPDL drugs receive PA in accordance with the Gainwell SPBM medical necessity policy as posted on the Gainwell SPBM website.
- Medications that are new to the market will be non-preferred, PA required, until reviewed by the ODM Pharmacy and Therapeutics (P&T) Committee.
- The UPDL document is organized by therapeutic class. Brand name drugs are listed in CAPITAL letters; generic drug names are listed in lower case letters. In most cases, when a generic of a brand-name drug is available, the generic drug will be preferred and appear on the UPDL while the brand name will be non-preferred but not appear on the UPDL. The <u>Drug Search tool</u> is a handy reference to check the status of a drug. Some generic drugs may require a specific labeler, or the brand to be dispensed.
- ODM will only cover drugs that are part of the Medicaid Drug Rebate Program, with limited exceptions. This document may not reflect the most current rebate status of a drug (i.e., a drug may be listed on the document but is non-rebateable and therefore non-payable).
- Some therapeutic categories are deemed 'legacy' categories. These categories are denoted with an "*" and LEGACY CATEGORY listed next to their title on the table on contents and their place within the criteria document. Legacy is defined as: Patients who have a claim for a non-preferred drug in the previous 120 days will be automatically approved to continue the drug. Patients who have taken the drug previously, but do not have claims history (e.g., new to Medicaid), will need to submit a prior authorization to continue coverage.
- ALL authorizations must be prescribed in accordance with FDA approved labeling or listed on a CMS-supported compendia.
- For ALL authorizations, there must be a trial and failure of preferred strengths prior to authorization of non-preferred strengths (if available).

- For ALL non-preferred authorizations, there must be documentation of medical necessity beyond convenience for why the patient cannot be changed to a preferred drug form (i.e., allergies, drug-drug interactions, contraindications, or intolerances). Must have had an inadequate clinical response of preferred individual components for any combination non-preferred product.
- For any nonsolid oral dosage formulation, there must be documentation of medical necessity for why patient cannot be changed to a solid oral dosage formulation (if available).
- For non-preferred extended-release formulations, there must be documentation of an inadequate clinical response with its immediate release formulation (if available).
- For non-preferred brand names that have preferred generics, there must be documentation of an inadequate clinical response or allergy to two or more generic labelers (if available).
- For ALL subsequent authorizations, there must be documentation of patient's clinical response to treatment and ongoing safety monitoring unless otherwise stated.
- Some therapeutic categories have sub-sections to divide the medications by their mechanism of action, route of administration, or duration of action. References to 'sub-section' in the Clinical Criteria shall be defined as the separate groupings that appear in that category's drug placement columns.
- Some therapeutic categories may have quantity limits on specific drugs. For a list of the quantity limits on specific drugs, please reference the Quantity Limit Document found here: <u>Quantity Limits Document | spbm.medicaid.ohio.gov</u>

Terminology/Abbreviations:

AR (Age Restriction) – An edit allowing claims for members within a defined age range to be covered without PA

BvG (Brand Preferred Over the Generic) – The brand name drug is preferred over the generic equivalent

PA (Clinical Prior Authorization) - PA is required before the drug will be covered

ST (Step Therapy) – Drug requires a trial with one or more preferred drugs before being covered

- With a few exceptions, the clinical criteria have a cumulative top-to-bottom format.

Example Category		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
Example Drug	Example Drug	LENGTH OF AUTHORIZATIONS: X days or Initial: X days; Subsequent: X days (if different)
		LEGACY*: Patients who have a claim for a non-preferred drug in the previous 120 days will be automatically approved to continue the drug. Patients who have taken the drug previously, but do not have claims history (e.g. new to Medicaid), will need to submit a prior authorization in order to continue coverage.
		CLINICAL PA CRITERIA (if applicable):
		<u>"DRUG" CRITERIA (if applicable):</u>
		 STEP THERAPY CRITERIA: Must have had an inadequate clinical response of at least <u>X days</u> with at least <u>X preferred</u> drugs
		 NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>X days</u> with <u>X</u> preferred drugs
		ADDITIONAL "DRUG" CRITERIA (if applicable):
		ADDITIONAL INFORMATION (if applicable):
		 SUBSEQUENT AUTHORIZATION CRITERIA: Must provide documentation of patient's response to treatment from baseline and/or attestation of clinical stabilization AR – a PA is required for patients X years and older OR younger than X years

Interpretation of UPDL Format

- The UPDL criteria is designed to have a cumulative approach from top-to-bottom. The following scenarios will aid in illustrating this point:

Scenario 1: Clinical PA drug

- All Authorizations
- Clinical PA Criteria

Scenario 2: Clinical PA drug with drug-specific criteria

- All Authorizations
- Drug-Specific Criteria

Scenario 3: Step-therapy drug

- All Authorizations
- Clinical PA Criteria (if applicable)
- Step Therapy Criteria

Scenario 4: Non-preferred drug

- All Authorizations
- Clinical PA Criteria (if applicable)
- Step Therapy Criteria (if applicable)
- Non-Preferred Criteria

Scenario 5: Non-preferred drug with drug-specific criteria

- All Authorizations
- Clinical PA Criteria (if applicable)
- Step Therapy Criteria (if applicable)
- Non-Preferred Criteria
- Additional Drug-Specific Criteria

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Infectious Disease Agents: Antivirals – Herpes	
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Analgesic Agents: Gout			
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA	
allopurinol 100, 300mg colchicine tab	allopurinol 200mg febuxostat	LENGTH OF AUTHORIZATIONS: 365 days	
probenecid probenecid/colchicine	MITIGARE ^{BVG}	 NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>one preferred</u> drug in this UPDL category and indicated for diagnosis 	
		 ADDITIONAL COLCHICINE CAPSULE (MITIGARE) CRITERIA: Must have had an inadequate clinical response of <u>30 days</u> with colchicine tablets 	

Analgesic Agents: NSAIDS			
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA	
celecoxib diclofenac sodium DR, ER, gel 1% etodolac IR, ER flurbiprofen ibuprofen indomethacin IR, ER cap ketorolac mefenamic acid meloxicam tab nabumetone naproxen IR naproxen susp ^{AR} oxaprozin piroxicam sulindac	diclofenac/misoprostol diclofenac patch 1.3%; soln 1.5%, 2% diclofenac potassium ELYXYB fenoprofen ibuprofen/famotidine indomethacin supp, susp ketoprofen IR, ER meclofenamate meloxicam cap naproxen EC, ER naproxen/esomeprazole RELAFEN DS	 LENGTH OF AUTHORIZATIONS: 365 days NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>two preferred</u> drugs in this UPDL category and indicated for diagnosis AR – naproxen susp: a PA is required for patients 12 years old and older 	

Analgesic Agents: Opioids			
PREFERRED AGENTS NON-PREFERRED AGENTS		PA CRITERIA	
SHORT-ACTING APAP/codeine AR APAP/caffeine/ dihydrocodeine		**Ohio law requires prescribers to request and review an OARRS report before initially prescribing or personally furnishing any	
but/APAP/caff/cod ^{AR} 50/325/40/30mg but/ASA/caff/cod ^{AR} butorphanol	but/APAP/caff/cod ^{AR} 50/300/40/30mg hydrocodone/APAP 5, 7.5, 10-300mg hydrocodone/ibuprofen	controlled substance, such as an opioid analgesic or a benzodiazepine, and gabapentin**	
codeine ^{AR} hydrocodone/APAP hydromorphone IR morphine IR	levorphanol meperidine oxymorphone IR pentazocine/naloxone	LENGTH OF AUTHORIZATIONS: Initial short-acting and long-acting requests may only be authorized for up to 90 days. For reauthorization, up to 180 days.	
oxycodone IR cap, soln, tab oxycodone/APAP tramadol IR ^{AR} 50mg tramadol/APAP ^{AR}	PROLATE ROXYBOND SEGLENTIS ^{AR} tramadol IR ^{AR} soln, 25, 75, 100mg tab	 BUPRENORPHINE TOPICAL (BUTRANS) CRITERIA: For doses greater than 5 mcg/hour must provide documentation of an inadequate clinical response with at least one opioid formulation taken for at least 30 of the last 	
LONG-	ACTING	60 days	
fentanyl hydrocodone bitartrate ER 12H	buprenorphine TD patch weekly fentanyl hydrocodone bitartrate ER 12HR cap hydrocodone bitartrate ER 24HR tab hydromorphone ER methadone morphine ER 24HR cap	 MORPHINE SULFATE ER (MS CONTIN) CRITERIA: Unless receiving for cancer pain, palliative care, or end-of-life/hospice care, must provide documentation of an inadequate clinical response with at least one opioid formulation taken for at least 30 of the last 60 days Must also meet LONG-ACTING OPIOID CRITERIA 	
	oxymorphone ER	 MON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>7</u> <u>days</u> of at least <u>two</u> preferred drugs with different active ingredients of the same duration of action (SHORT-ACTING or LONG-ACTING) Must also meet applicable SHORT-ACTING or LONG-ACTING OPIOID CRITERIA 	
		 ADDITIONAL SHORT-ACTING OPIOIDS CRITERIA: The system defines an "initial request" as having no opioid 	

claims in the previous 90 days Initial short-acting requests can be authorized up to 90 days ٠ Length of authorization is dependent on indication, 0 previous patient utilization, and requested length of therapy (could be more restrictive) To exceed acute opioid limits, documentation of the 0 following must be provided: Diagnosis code which must be for somatic type pain Prescriber attestation that the benefits and risks of opioid therapy have been discussed with patient Exemptions to the additional criteria: 0 Patients receiving short-acting opioids for active cancer treatment, palliative care, and end-of-life/hospice care, sickle cell, severe burn, traumatic crushing of tissue, amputation, major orthopedic surgery Prescriber attestation that patient is opioid tolerant (i.e., new to Medicaid or was on higher dose in hospital) Subsequent short-acting requests can be authorized up to 180 days • Documentation of the following must be provided: Current treatment plan Demonstrated adherence to treatment plan through progress notes, including pain and function scores, random urine screening results reviewed, concerns addressed, and no serious adverse outcomes observed Exemptions to the additional criteria: 0 Patients receiving short-acting opioids for cancer pain, palliative care, or end-oflife/hospice care Patients residing in LTC facilities are

exempted from urine drug screening requirements

- Dose escalation requests can be authorized up to 180 days
 - Documentation of the following must be provided:
 - Prescriber attestation that dose escalation is likely to result in improved function or pain control
 - Requests for a cumulative daily dose > 80 MED must be prescribed by or in consultation with a pain specialist, specialist in the area of the body affected by pain, or anesthesiologist

Patients with initial prescriptions for opioid therapy, defined as no rx claims for opioids in the last 90 days, will be limited to 30 MED per day and a maximum of 7 days per prescription. Prior authorization will be required to exceed these limits.

ADDITIONAL LONG-ACTING OPIOIDS CRITERIA:

 The system defines an "initial long-acting request" as having no opioid claims in the previous 90 days Initial long-acting requests can be authorized up to 90 days Documentation of the following must be provided: Request is a daily dose equivalent of ≤ 80 MED Inadequate clinical response to both non-opioid pharmacologic and non-pharmacologic treatments Current use of opioids for ≥ 30 of the last 60 days Treatment plan including risk 	

Pain and function scores at each visit Opioid contract required to be in place and submitted with PA form Exemptions to the additional criteria: 0 Patients receiving long-acting opioids for cancer pain, palliative care, or end-oflife/hospice care Patients residing in LTC facilities are exempted from urine drug screening and opioid contract requirements Subsequent long-acting requests can be authorized up to 180 days Documentation of the following must be provided: 0 Current treatment plan Demonstrated adherence to treatment plan through progress notes, including pain and function scores, random urine screening results reviewed, concerns addressed, and no serious adverse outcomes observed Exemptions to the additional criteria: 0 Patients receiving long-acting opioids for cancer pain, palliative care, or end-oflife/hospice care Patients residing in LTC facilities are exempted from urine drug screening and opioid contract requirements **Dose escalation requests** can be authorized up to 180 days ٠ Documentation of the following must be 0 provided: Prescriber attestation that dose escalation is likely to result in improved function or pain control Requests for a cumulative daily dose > 80

	consultation	e prescribed by or in a with a pain specialist, specialist of the body affected by pain, or ogist
ADI	DITIONAL TRANSMUCOSAL FENTAN	YL CRITERIA:
	 Must be prescribed by an one hospice/palliative prescriber 	cologist, pain specialist, or
	 Must be concurrently taking therapeutic dose of any of th <u>days</u> without adequate pain 	e following for at least <u>7</u>
	≥ 60 mg oral morphine/day	≥ 8 mg oral hydromorphone/day
	≥ 25 mcg/hr transdermal fentanyl	≥ 25 mg oral oxymorphone/day
	≥ 30 mg oral oxycodone/day	Equianalgesic dose of another opio
BUF	PRENORPHINE BUCCAL FILM (BELBU	CA) CRITERIA:
	Must meet ADDITIONAL LON	IG-ACTING OPIOID Criteria
AR	R – All codeine and tramadol containi	
	for patients younger than 12 years	010

Blood Formation, Coagulation, and Thrombosis Agents: Colony Stimulating Factors			
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA	
FULPHILA PA	FYLNETRA	LENGTH OF AUTHORIZATIONS: 30 days or duration of chemotherapy regimen	
NEUPOGEN PA	GRANIX		
NIVESTYM PA	LEUKINE	CLINICAL PA CRITERIA:	
NYVEPRIA PA	NEULASTA	Must provide documentation of diagnosis, patient's weight (for weight-based	
	RELEUKO	dosed medications only), and duration of treatment	
	ROLVEDON		
	STIMUFEND	NON-PREFERRED CRITERIA:	
	UDENYCA	 Must have had an inadequate clinical response of at least <u>14 days</u> with at 	
	ZARXIO	least one preferred drug in this UPDL category and indicated for diagnosis	
	ZIEXTENZO		

Blood Formation, Coagulation, and Thrombosis Agents: Hematopoietic Agents		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
EPOGEN ^{PA} RETACRIT ^{PA}	ARANESP MIRCERA PROCRIT	 LENGTH OF AUTHORIZATIONS: 180 days; except 365 days for patients with chronic renal failure CLINICAL PA CRITERIA: Must provide documentation of baseline hemoglobin level NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>14 days</u> with at least <u>one preferred</u> drug in this UPDL category and indicated for diagnosis
		SUBSEQUENT AUTHORIZATION CRITERIA:
		Provide current hemoglobin lab result

Blood Formation, Coagulation, and Thrombosis Agents: Hemophilia A, von Willebrand Disease, and Factor XIII Deficiency* LEGACY CATEGORY		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
ADVATE PA	HYMPAVZI	LENGTH OF AUTHORIZATIONS: 365 Days
ADYNOVATE PA	NUWIQ INJ	
AFSTYLA PA	OBIZUR	CLINICAL PA CRITERIA:
ALPHANATE PA	SEVENFACT	 Must provide documentation of patient's body weight (for weight-based
ALTUVIIIO PA	VONVENDI	dosed medications only)
CORIFACT PA		
ELOCTATE PA		NON-PREFERRED CRITERIA:
ESPEROCT PA		 Must have had an inadequate clinical response of at least <u>14 days</u> with at
FEIBA PA		least one preferred drug in this UPDL category and indicated for
HEMLIBRA PA		diagnosis
HEMOFIL M PA		
HUMATE-P PA		ADDITIONAL EXTENDED HALF-LIFE FACTOR CRITERIA
JIVI PA		 Must provide attestation that the patient is not a suitable candidate for
KOATE PA		treatment with a shorter-acting half-life drug
KOGENATE FS PA		
KOVALTRY PA		ADDITIONAL HYMPAVZI (MARSTACIMAB-HNCQ) CRITERIA
NOVOEIGHT PA		 Must have had an inadequate clinical response of at least <u>30 days</u> with
NOVOSEVEN RT PA		HEMLIBRA
NUWIQ KIT PA		
RECOMBINATE PA		
WILATE PA		
XYNTHA PA		

Blood Formation, Coagulation, and Thrombosis Agents: Hemophilia B* LEGACY CATEGORY		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
ALPHANINE SD PA	HYMPAVZI	LENGTH OF AUTHORIZATIONS: 365 Days
ALPROLIX PA	SEVENFACT	
BENEFIX PA		CLINICAL PA CRITERIA:
FEIBA PA		 Must provide documentation of patient's body weight (for weight-based
IDELVION PA		dosed medications only)
IXINITY PA		
NOVOSEVEN RT PA		NON-PREFERRED CRITERIA:
PROFILNINE PA		 Must have had an inadequate clinical response of at least <u>14 days</u> with at
REBINYN PA		least one preferred drug in this UPDL category and indicated for
RIXUBIS PA		diagnosis
		ADDITIONAL EXTENDED HALF-LIFE FACTOR CRITERIA
		 Must provide attestation that the patient is not a suitable candidate for treatment with a shorter-acting half-life drug

Blood Formation, Coagulation, and Thrombosis Agents: Heparin-Related Preparations			
PREFERRED AGENTS NON-PREFERRED AGENTS PA CRITERIA			
enoxaparin	fondaparinux FRAGMIN	 LENGTH OF AUTHORIZATIONS: 35 days; except 365 days for patients with cancer, pregnancy, or unable to be converted to an oral anticoagulant NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>14 days</u> with at least <u>one preferred</u> drug in this UPDL category and indicated for diagnosis 	

Blood Formation, Coagulation, and Thrombosis Agents: Oral Anticoagulants			
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA	
ELIQUIS PRADAXA CAP ^{BvG} , PELLET PAK ^{AR} warfarin XARELTO ^{BvG} SUSP ^{AR} , TAB	dabigatran cap rivaroxaban tab SAVAYSA	LENGTH OF AUTHORIZATION: 365 days NON-PREFERRED CRITERIA: • Must have had an inadequate clinical response of at least 14 days with at	
		least <u>two preferred</u> drugs in this UPDL category and indicated for diagnosis	
		AR – PRADAXA PELLET PAK, XARELTO SUSP: a PA is required for patients 12 years and older	

Blood Formation, Coagulation, and Thrombosis Agents: Oral Antiplatelet			
PREFERRED AGENTS NON-PREFERRED AGENTS PA CRITERIA			
aspirin IR, ER aspirin/dipyridamole ER BRILINTA ^{BvG} clopidogrel prasugrel	ticagrelor	 LENGTH OF AUTHORIZATION: 365 days NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>one preferred</u> drug in this UPDL category and indicated for diagnosis 	

Cardiovascular Agents: Angina, Hypertension and Heart Failure		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
ACE INHIBITORS/DIUR	ETICS/COMBINATIONS	LENGTH OF AUTHORIZATIONS: 365 days except nimodipine: 21 days
amlodipine/benazepril benazepril benazepril/HCTZ captopril captopril/HCTZ enalapril soln, tab enalapril/HCTZ fosinopril fosinopril/HCTZ lisinopril lisinopril/HCTZ moexipril	QBRELIS	 PROPRANOLOL ORAL SOLN (HEMANGEOL) CRITERIA: Must provide documentation of the patient's weight NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>30 days</u> of at least <u>two preferred</u> drugs within the same sub-section classification in this UPDL category and indicated for diagnosis ADDITIONAL APROCITENTAN (TRYVIO) CRITERIA: Must have had an inadequate clinical response of at least <u>30 days</u> of at least <u>four</u> different classes of antihypertensive medications <u>concurrently</u> without adequate blood pressure control
quinapril quinapril/HCTZ ramipril trandolapril trandolapril/verapamil ARBs/DIURETICS amlodipine/olmesartan amlodipine/valsartan amlodipine/valsartan/HCTZ candesartan	/COMBINATIONS EDARBI EDARBYCLOR valsartan soln	 ADDITIONAL FINERENONE (KERENDIA) CRITERIA: Must be on a maximally tolerated dose of an angiotensin-converting enzyme inhibitor or angiotensin receptor blocker AND Must provide documentation of an inadequate clinical response to a SGLT2 Inhibitor OR provide documentation of medical necessity beyond convenience for why the patient cannot try a SGLT2 inhibitor (i.e., chronic kidney disease diagnosis)
candesartan/HCTZ		ADDITIONAL MAVACAMTEN (CAMZYOS) CRITERIA:
irbesartan irbesartan/HCTZ losartan losartan/HCTZ olmesartan olmesartan/amlodipine/HCTZ olmesartan/HCTZ telmisartan		 Must be prescribed by or in consultation with a cardiologist Must provide documentation of NYHA Class II-III symptoms and left ventricular ejection fraction ≥55% ADDITIONAL SOTAGLIFLOZIN (INPEFA) CRITERIA: Must provide documentation of an inadequate clinical response to at least two SGLT2 Inhibitors (refer to Endocrine Agents: Diabetes – Non-Insulin class for complete list)

telmisartan/HCTZ		ADDITIONAL VERICIGUAT (VERQUVO) CRITERIA:
valsartan tab		Must provide documentation of ejection fraction
valsartan/HCTZ		
BETA BLOCKERS/COMBINATIONS		 Must have been hospitalized for the treatment of heart failure in the previous 180 days or needs treatment with an outpatient intravenous
acebutolol bisoprolol 2.5mg		diuretic in the previous 90 days
atenolol	carvedilol ER	
atenolol/chlorthalidone	INNOPRAN XL	 Must be treated with an agent from ALL the following unless contraindicated:
betaxolol	KAPSPARGO	
bisoprolol 5, 10mg	labetalol 400mg	 Angiotensin-converting enzyme inhibitor, angiotensin II receptor blocker, OR an angiotensin receptor neprilysin inhibitor
bisoprolol/HCTZ	SOTYLIZE AR	 Beta-blocker
carvedilol IR	0011111	 Aldosterone antagonist and/or SGLT2 inhibitor as appropriate for
		renal function
labetalol 100, 200, 300mg		
metoprolol succ		AR – SOTYLIZE SOLN: a PA is required for patients 6 years and older
metoprolol tart		
metoprolol/HCTZ		
nadolol		
nebivolol		
propranolol IR, ER		
sotalol		
timolol		
CALCIUM CHAI	NNEL BLOCKERS	
amlodipine	diltiazem 24HR ER tabs	
cartia XT	isradipine	
diltiazem IR	KATERZIA	
diltiazem 12HR ER cap	nimodipine	
diltiazem 24HR ER cap	nisoldipine	
felodipine ER	NORLIQVA	
levamlodipine	NYMALIZE	
nicardipine	verapamil ER (gen of VERELAN PM)	
nifedipine IR, ER		
verapamil IR, ER, SR		
	HER	
amiloride	aliskiren	
amiloride/HCTZ	ASPRUZYO SPRINKLE	
clonidine IR, patch	CAMZYOS	

doxazosin	clonidine ER (gen of NEXICLON XR)
ENTRESTO TAB BVG	CORLANOR SOLN
eplerenone	ENTRESTO SPRINKLE CAP
guanfacine IR, ER	INPEFA
hydralazine	ivabradine tab (gen of CORLANOR)
methyldopa	KERENDIA
minoxidil	spironolactone susp
ranolazine	TRYVIO
spironolactone tab	VERQUVO
spironolactone/HCTZ	
terazosin	

Cardiovascular Agents: Antiarrhythmics		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
amiodarone	quinidine IR, ER	LENGTH OF AUTHORIZATIONS: 365 Days
disopyramide		
dofetilide		NON-PREFERRED CRITERIA:
flecainide		• Must have had an inadequate clinical response of at least <u>30 days</u> with at
mexiletine		least one preferred drug in this UPDL category and indicated for
MULTAQ		diagnosis
NORPACE CR		
propafenone IR, ER		

Cardiovascular Agents: Lipotropics			
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA	
BILE ACID SE	QUESTRANTS	LENGTH OF AUTHORIZATIONS: See below	
cholestyramine light, regular colesevelam tab	colesevelam packet colestipol granules	JUXTAPID (Initial)	180 days
colestipol tab		icosapent ethyl cap, LOVAZA, ACL inhibitors (Initial)	84 days
prevalite		All others (Initial and Subsequent)	365 days
FIBRIC ACID	DERIVATIVES		
fenofibrate 48, 54, 145, 160mg tab gemfibrozil	fenofibrate IR, DR cap fenofibrate 40, 120mg tab fenofibric acid	 CLINICAL PA CRITERIA: Must provide baseline labs AND have adherence to <u>90</u> lipid lowering medications 	days of preferred
PCSK9 IN	HIBITORS	 Must have had an inadequate clinical response of at le 	ast 90 days AND
PRALUENT ^{PA} REPATHA ^{PA}		unable to reach goal LDL-C (see below) despite treatment with maxima tolerated or high-potency statin (or a clinical reason that these drugs	
STATINS/CO	MBINATIONS	cannot be utilized)	
atorvastatin ezetimibe/simvastatin lovastatin pravastatin rosuvastatin	ALTOPREV amlodipine/atorvastatin ATORVALIQ EZALLOR fluvastatin IR, ER	 Must have had an inadequate clinical response of at least 90 days ANI unable to reach goal LDL-C (see below) despite treatment with ezetim OR documentation that LDL is >25% above goal despite current statin therapy 	ent with ezetimibe
simvastatin	pitavastatin ZYPITAMAG	NON-PREFERRED CRITERIA:	aast 20 days (or 90
01	HER	 Must have had an inadequate clinical response of at least <u>30 days</u> (or <u>90</u> days for fibrates) with at least one preferred drug within the same sub- 	
ezetimibe niacin IR, ER OTC	icosapent ethyl cap JUXTAPID	section classification in this UPDL category and indicat	
omega-3-acid ethyl esters	NEXLETOL NEXLIZET niacin ER tab	ADDITIONAL LOVASTATIN ER (ALTOPREV), PITAVASTATIN (LIV FLUVASTATIN (LESCOL) CRITERIA: • Must have had an inadequate clinical response of at le	
		two preferred drugs in the same drug class ADDITIONAL COLESEVELAM (WELCHOL) CRITERIA:	
		Must provide documentation of a Type 2 Diabetes diag	gnosis
		ADDITIONAL ICOSAPENT ETHYL CRITERIA:	

 Must provide documentation of baseline labs indicating triglyceride levels ≥500mg/dL after an inadequate clinical response to fibrates, niacin, and diet/exercise ADDITIONAL LOMITAPIDE (JUXTAPID) & ATP CITRATE LYASE (ACL) INHIBITOR CRITERIA: Must provide documentation of baseline labs AND have documented adherence to <u>90 days</u> of prescribed lipid lowering medications Must have had inadequate clinical response of at least <u>90 days</u> AND unable to reach goal LDL-C with high-potency statin, ezetimibe and PCSK9 inhibitor (or a clinical reason that these drugs cannot be utilized)
ADDITIONAL INFORMATION:
 High potency statins: atorvastatin (LIPITOR) 40-80mg & rosuvastatin (CRESTOR) 20-40mg
 LDL goals for Familial Hypercholesterolemia (includes Heterozygous & Homozygous FH): LDL ≤ 100mg/dL for adults or LDL ≤ 110mg/dL for those < 18 years of age
 LDL goals for Clinical Atherosclerotic Cardiovascular Disease (ASCVD) not at very high risk: LDL ≤ 70mg/dL
 LDL goals for Clinical Atherosclerotic Cardiovascular Disease (ASCVD) at very high risk: LDL ≤ 55mg/dL
 Must provide documentation of multiple major ASCVD events or 1 major ASCVD event and multiple high-risk conditions if citing goal LDL ≤ 55mg/dL

Cardiovascular Agents: Pulmonary Arterial Hypertension* LEGACY CATEGORY		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
ENDOTHELIN RECEPTOR ANTAGONISTS		LENGTH OF AUTHORIZATIONS: 365 Days
ambrisentan PA	OPSUMIT	
bosentan PA	TRACLEER SUSP	CLINICAL PA CRITERIA:
F	PDE5 INHIBITORS	 Must provide documentation of NYHA Functional Class symptoms for
sildenafil PA		Pulmonary Hypertension experienced by patient
sildenafil susp ^{AR PA}		
tadalafil ^{PA}		NON-PREFERRED CRITERIA:
TADLIQ AR PA		 Must have had an inadequate clinical response of at least <u>30 days</u> with at
Р	ROSTAGLANDINS	least two preferred drugs in this UPDL category and indicated for
epoprostenol	ORENITRAM	diagnosis, if available, <u>one</u> of which must be a phosphodiesterase-5
	treprostinil	inhibitor
	TYVASO	
	VENTAVIS	ADDITIONAL INFORMATION:
OTHER		 Patients who have class III or IV symptoms defined by the NYHA
	ADEMPAS	Functional Class for Pulmonary Hypertension may be authorized for
	OPSYNVI	inhalation or intravenous agents
	UPTRAVI	
	WINREVAIR	AR – sildenafil susp: a PA is required for patients 18 years and older
		AR – TADLIQ: a PA is required for patients younger than 18 years

Central Nervous System (CNS) Agents: Alzheimer's Agents* LEGACY CATEGORY		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
donepezil ^{AR} galantamine IR tab, ER cap ^{AR}	ADLARITY ^{AR} galantamine soln ^{AR}	LENGTH OF AUTHORIZATIONS: 365 Days
memantine IR, ER tab ^{AR} rivastigmine cap ^{AR} rivastigmine patch ^{AR}	memantine/donepezil cap ^{AR} 14-10, 21-10, 28-10mg memantine soln ^{AR} NAMZARIC ^{AR}	 MON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>two preferred</u> drugs in this UPDL category and indicated for diagnosis
		AR – All drugs: a PA is required for patients younger than 40 years

Central Nervous System (CNS) Agents: Anti-Migraine Agents, Acute		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
CGRP INHIBITORS		LENGTH OF AUTHORIZATIONS: 180 Days
NURTEC ODT ST	ZAVZPRET	
UBRELVY ST		STEP THERAPY CRITERIA:
TRIPTANS/COMBINATIONS		 Must have had an inadequate clinical response of at least <u>14 days</u> with at
IMITREX NASAL SPRAY	almotriptan	least two preferred drugs in this UPDL category OR documentation why
naratriptan	eletriptan	patient is unable to take product not requiring step therapy
rizatriptan	frovatriptan	
sumatriptan inj, nasal spray, tab	sumatriptan/naproxen	NON-PREFERRED CRITERIA:
	TOSYMRA	 Must have had an inadequate clinical response of at least <u>14 days</u> with at
	zolmitriptan	least <u>one</u> preferred drug and <u>one</u> step therapy drug in this UPDL category
OTHER		and indicated for diagnosis, if available
	dihydroergotamine	
	MIGERGOT	ADDITIONAL INFORMATION:
	REYVOW	NURTEC has a maximum quantity of 8 tablets per month for acute
		migraines

Central Nervous System (CNS) Agents: Anti-Migraine Agents, Cluster Headache		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
verapamil IR, ER	EMGALITY 100mg/ml	LENGTH OF AUTHORIZATIONS: 180 days
		 Must have had an inadequate clinical response of at least <u>60 days</u> to at least <u>one preferred</u> drug in this UPDL category and indicated for diagnosis
		ADDITIONAL INFORMATION:
		 An inadequate clinical response to verapamil is defined as a titration to at least 480mg daily or maximally tolerated dose based on blood pressure or heart rate and maintained for at least 60 days

	Central Nervous System (Cl	NS) Agents: Anti-Migraine Agents, Prophylaxis
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
AIMOVIG ST AJOVY ST	NURTEC ODT QULIPTA	LENGTH OF AUTHORIZATIONS: Initial: 180 days; Subsequent: 365 days
EMGALITY 120mg/ml st	QUEITA	STEP THERAPY CRITERIA:
EMGALITY 120mg/ml ³¹ Cardiovascular Agents: Beta- Blockers CNS Agents: Anticonvulsants CNS Agents: Serotonin- Norepinephrine Reuptake Inhibitors CNS Agents: Tricyclic Antidepressants		 Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>two preferred</u> controller migraine drugs. For patients already established on a serotonergic medication, only <u>one</u> preferred controller migraine drugs will be required Must include objective documentation of severity, frequency, type of migraine, and number of headache days per month Controller migraine drug classes include beta-blockers, anticonvulsants, serotonin-norepinephrine reuptake inhibitors, or tricyclic
		 antidepressants <u>ERENUMAB (AIMOVIG) CRITERIA:</u> Must have had an inadequate clinical response of at least <u>60 days</u> with the 70mg dose to request a dose increase
		FREMANEZUMAB (AJOVY) CRITERIA:
		 Must have demonstrated efficacy for at least <u>90 days</u> before quarterly administration will be authorized
		NON-PREFERRED CRITERIA:
		 Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>three preferred</u> controller migraine drugs AND <u>one step therapy</u> drug in this UPDL category
		ADDITIONAL INFORMATION:
		 NURTEC has a maximum quantity of 16 tablets per month for migraine prophylaxis
		SUBSEQUENT AUTHORIZATION CRITERIA:
		 Must provide documentation of patient's clinical response to treatment (Objective documentation of severity, frequency, and number of headache days per month).

Central Nervous System (CNS) Agents: Anticonvulsants* LEGACY CATEGORY		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
BANZEL TAB ^{BVG}	APTIOM	LENGTH OF AUTHORIZATIONS: 365 days except EPIDIOLEX and DIACOMIT – Initial:
BRIVIACT SOLN ^{AR} , TAB		180 days
carbamazepine IR, ER	clonazepam ODT	
clobazam	ELEPSIA XR	STEP THERAPY CRITERIA:
clonazepam	felbamate	 Must have had an inadequate clinical response of at least <u>30 days</u> with at
	FINTEPLA	least <u>one preferred</u> drug in this UPDL category
divalproex DR, ER	lamotrigine ER	
EPIDIOLEX PA	levetiracetam ER tab	CANNABIDIOL (EPIDIOLEX) CRITERIA
EPRONTIA AR	MOTPOLY XR	Must have had an inadequate clinical response of at least <u>30 days</u> with any
ethosuximide	oxcarbazepine susp	two of the following anticonvulsants: clobazam, levetiracetam, valproic
FYCOMPA ST	OXTELLAR XR ^{BVG}	acid, lamotrigine, topiramate, rufinamide, or felbamate within the past <u>365 days</u> (members who meet this criterion will not require a PA)
gabapentin	QUDEXY XR ^{BvG}	<u>565 days</u> (members who meet this chieffort will not require a PA)
lacosamide	rufinamide tab, soln	STIRIPENTOL (DIACOMIT) CRITERIA
lamotrigine chew, IR, ODT	SPRITAM ^{BVG}	Exempt from Legacy rules
levetiracetam IR tab, soln	SYMPAZAN	 Must be prescribed by or in consultation with a neurologist
oxcarbazepine IR tab	tiagabine	 Must be presended by or in consultation with a hearologist Must be concomitantly taking clobazam (ONFI)
phenobarbital	topiramate sprinkle cap	 Must be conconntantly taking clobazari (ONT) Must provide documentation of addressed comorbidities and baseline
phenytoin IR, ER	TROKENDI XR BVG	hematologic testing (CBC)
pregabalin IR	vigabatrin	 Patients with phenylketonuria (PKU) must provide evidence of tota
primidone	vigabatrin powder AR	daily amount of phenylalanine
topiramate IR	VIGAFYDE AR	• Prescribers must include management plans for patients with
TRILEPTAL SUSP BVG	XCOPRI	neutrophil counts <1,500 cells/mm ³ or platelet count <150,000/µl
valproic acid	ZONISADE SUSP	 Must provide documentation of patient's weight
zonisamide cap	ZTALMY	 Maximum daily dose does not exceed: 50 mg/kg/day or 3,000mg/day
		NON-PREFERRED CRITERIA:
		 Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>two preferred</u> drugs in this UPDL category and indicated for diagnosis

 Prescriptions submitted from a prescriber who is credentialed as a neurology specialty with Ohio Medicaid AND for drugs that are used only for seizures, there must have been an inadequate clinical response of at least <u>30 days</u> with <u>one preferred</u> drug. This provision applies only to the standard tablet/capsule dosage form.
AR – BRIVIACT SOLN: a PA is required for patients 12 years and older
AR – EPRONTIA SOLN: a PA is required for patients 12 years and older
AR – vigabatrin powder: a PA is required for patients 2 years and older
AR – VIGAFYDE SOLN: a PA is required for patients 2 years and older
Central Nervous System (CNS) Agents: Anticonvulsants Rescue
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PREFERRED AGENTS
diazepam gel LIBERVANT ^{AR} NAYZILAM ^{AR} VALTOCO ^{AR}

	Central Nervous System (CNS) Age	nts: Antidepressants* LEGACY CATEGORY
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
NE	DRIs	LENGTH OF AUTHORIZATIONS: 365 Days except 14 days with no renewal for
bupropion	APLENZIN	ZURZUVAE
bupropion SR (gen of WELLBUTRIN SR)	bupropion XL (gen of FORFIVO XL)	
bupropion XL (gen of WELLBUTRIN XL)		PSYCHIATRIST EXEMPTION:
	IRIs	Prescribers (as identified below) are exempt from prior authorization of
desvenlafaxine succ ER (gen of PRISTIQ) duloxetine 20, 30, 60mg venlafaxine IR tab, ER cap	desvenlafaxine ER (gen of KHEDEZLA) DRIZALMA SPRINKLE duloxetine 40mg FETZIMA venlafaxine ER tab	any non-preferred antidepressant, or step therapy of any preferred drug, in the standard tablet/capsule dosage forms. Other dosage forms may still require prior authorization. The exemption will be processed by the claims system when the pharmacy has submitted the prescriber on the claim using the individual national provider identifier (NPI) for the prescriber. Prescribers are defined as: Physicians with a specialty in
SS	Ris	psychiatry, nurse practitioners certified in psychiatric mental health, or
citalopram tab, soln escitalopram	citalopram cap fluoxetine IR 60mg, DR	clinical nurse specialists certified in psychiatric mental health, who are credentialed with the Ohio Department of Medicaid.
fluoxetine IR 10, 20, 40mg	fluvoxamine ER	
fluoxetine soln	paroxetine ER tab	CLINICAL PA CRITERIA:
fluvoxamine IR	sertraline cap	 Must have a diagnosis of moderate to severe Post-Partum Depression (PPD) no earlier than the 3rd trimester OR within 12 months of pregnancy
paroxetine IR tab, soln		delivery
sertraline tab		
	HER AUVELITY	STEP THERAPY CRITERIA:
mirtazapine nefazodone tranylcypromine	CAPLYTA clomipramine	 Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>two preferred</u> drugs in this UPDL category
trazodone 50, 100, 150mg vilazodone VRAYLAR ST ZURZUVAE ^{PA}	EMSAM MARPLAN phenelzine REXULTI trazodone 300mg	 NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>two preferred</u> drugs in this UPDL category and indicated for diagnosis
	TRINTELLIX	ADDITIONAL DEXTROMETHORPHAN/BUPROPION (AUVELITY) CRITERIA:
		 Must have an inadequate clinical response of at least <u>30 days</u> with ALL of the following: <u>ONE</u> norepinephrine/dopamine reuptake inhibitor (NDRI)

	 <u>ONE</u> serotonin and norepinephrine reuptake inhibitor (SNRI)
	 <u>TWO</u> selective serotonin reuptake inhibitors (SSRIs) (<u>ONE</u> of
	which must be either vilazodone (VIIBRYD) OR vortioxetine
	(TRINTELLIX))

Central Nervous System (CNS) Agents: Attention Deficit Hyperactivity Disorder Agents		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
NON-STI	MULANTS	LENGTH OF AUTHORIZATIONS: 365 days
NON-STIL atomoxetine cap ^{AR} clonidine ER guanfacine ER ONYDA XR SUSP ^{AR} QELBREE ST		
		AR – methylphenidate soln/susp/chewable tab: a PA is required for patients 12 years and older
		AR – ONYDA XR SUSP: a PA is required for patients 12 years and older

Central Nervous System (CNS) Agents: Atypical Antipsychotics* LEGACY CATEGORY		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
PREFERRED AGENTSABILIFY ASIMTUFII, MAINTENAaripiprazoleARISTADAARISTADA INITIOasenapine STclozapineFANAPT STGEODONINVEGA HAFYERA ER PAINVEGA TRINZAlurasidoneolanzapinepaliperidone tab		
PERSERIS quetiapine IR, ER RISPERDAL CONSTA ^{BvG} risperidone RYKINDO UZEDY VRAYLAR ST ziprasidone	SECOADO VERSACLOZ ZYPREXA RELPREVV	 PALIPERIDONE PALMITATE (INVEGA HAFYERA) CRITERIA: Must have had 4 months of treatment with INVEGA SUSTENNA or 3 months with INVEGA TRINZA STEP THERAPY CRITERIA: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>one preferred</u> drug in this UPDL category MUST have had an inadequate clinical response of at least <u>30 days</u> with at least <u>one preferred</u> drug in this UPDL category MUST have had an inadequate clinical response of at least <u>30 days</u> with at least <u>two preferred</u> drugs in this UPDL category and indicated for diagnosis ADDITIONAL ARIPIPRAZOLE (ABILIFY MYCITE) CRITERIA: Must be prescribed by or in consultation with a psychiatrist following an aripiprazole serum blood level draw indicating need for further investigation of adherence

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ADDITIONAL OLANZAPINE/SAMIDORPHAN (LYBALVI) CRITERIA:
 Must provide documentation that patient is not using opioids or
undergoing acute opioid withdrawal
ADDITIONAL PIMAVANSERIN (NUPLAZID) CRITERIA:
For Parkinson-related Hallucinations & Delusions ALL of the following
must be met:
 Psychotic symptoms are severe and frequent enough to warrant treatment with an antipsychotic AND are not related to dementia or delirium
 The patient's other Parkinson's Disease drugs have been
reduced or adjusted and psychotic symptoms persist OR patient
is unable to tolerate adjustment of these other drugs
 Must have been inadequate clinical response or contraindication to at least <u>30 days</u> of either quetiapine or clozapine
• An exemption to the criteria will be authorized for prescribers with a
neurology specialty to a patient with a history of the related condition
ADDITIONAL INFORMATION:
 Long-acting injectable antipsychotics may be billed by the pharmacy if
they are not dispensed directly to the patient. If not administered by the
pharmacist, the drug must be released only to the administering
provider or administering provider's staff, following all regulations for a
Prescription Pick-Up Station as described by the Ohio Board of Pharmacy

Central Nervous System (CNS) Agents: Fibromyalgia Agents			
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA	
pregabalin IR SAVELLA		All products are covered without a PA	

Central Nervous System (CNS) Agents: Medication Assisted Treatment of Opioid Addiction		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
BRIXADI buprenorphine/naloxone	buprenorphine LUCEMYRA ^{BvG}	LENGTH OF AUTHORIZATIONS: 180 days except 14 days for LUCEMYRA
clonidine IR, ER		ADDITIONAL LOFEXIDINE (LUCEMYRA) CRITERIA
SUBLOCADE SUBOXONE VIVITROL ZUBSOLV		 May be authorized if ALL of the following criteria are met: Must provide medical justification supporting why an opioid taper (such as with buprenorphine or methadone) cannot be used Must have had an inadequate clinical response or contraindication to clonidine
		 Must provide documentation that the drug was initiated in an inpatient setting to be exempt from the above criteria
		BUPRENORPHINE SAFETY EDITS AND DRUG UTILIZATION REVIEW CRITERIA:
		 Prescribing for buprenorphine products must follow the requirements of Ohio Administrative Code rule 4731-33-03 Office based treatment for opioid addiction. In favor of eliminating prior authorization for all forms of oral short acting buprenorphine- containing products, ODM and the Managed Care Plans will implement safety edits and a retrospective drug utilization review process for all brand and generic forms of oral short acting buprenorphine-containing products. Safety edits are in place for dosages over 24mg of buprenorphine equivalents/day. buprenorphine sublingual tablets (generic SUBUTEX) will be restricted to pregnancy, breastfeeding, or allergy/contraindication to preferred products
		 ADDITIONAL INFORMATION VIVITROL, SUBLOCADE, and BRIXADI may be billed by the pharmacy if it is not dispensed directly to the patient. If not administered by the pharmacist, the drug must be released only to the administering provider or administering provider's staff, following all regulations for a Prescription Pick-Up Station as described by the Ohio Board of Pharmacy.

Central Nervous System (CNS) Agents: Movement Disorders		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
AUSTEDO IR, XR ^{PA ST} INGREZZA ^{PA ST}		LENGTH OF AUTHORIZATIONS: 365 Days
tetrabenazine		CLINICAL PA CRITERIA:
		 Must be prescribed by or in consultation with a neurologist or psychiatrist
		STEP THERAPY CRITERIA:
		 Must have an inadequate clinical response of at least <u>90 days</u> to a maximally tolerated dose of tetrabenazine for Huntington's Disease only

Central Nervous System (CNS) Agents: Multiple Sclerosis* LEGACY CATEGORY		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
AVONEX	BAFIERTAM	LENGTH OF AUTHORIZATIONS: 365 Days
BETASERON	glatiramer	
	glatopa	NON-PREFERRED CRITERIA:
dalfampridine	MAVENCLAD	 Must have had an inadequate clinical response of at least <u>30 days</u> with at
dimethyl fumarate	MAYZENT	least one preferred drug in this UPDL category and indicated for
fingolimod	OCREVUS	diagnosis
GILENYA	PLEGRIDY	
KESIMPTA	PONVORY	ADDITIONAL OCRELIZUMAB (OCREVUS) CRITERIA:
REBIF	TASCENSO ODT	 Must provide documentation of diagnosis of primary progressive
teriflunomide	VUMERITY	multiple sclerosis OR must have had an inadequate clinical response of
	ZEPOSIA	at least <u>30 days</u> with at least <u>one preferred</u> drug in this UPDL category
		ADDITIONAL SIPONIMOD (MAYZENT) CRITERIA:
		 Must provide documentation of CYP2C9 genotype

Central Nervous System (CNS) Agents: Narcolepsy		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
amphetamine/ dextroamphetamine IR/ER ^{AR} armodafinil dextroamphetamine ER ^{AR} methylphenidate ER methylphenidate tab modafinil	SUNOSI WAKIX XYREM ^{BvG} XYWAV	 LENGTH OF AUTHORIZATIONS: 365 days NON-PREFERRED CRITERIA: Must have had an inadequate clinical response with at least two preferred drugs - either at least <u>30 days</u> of armodafinil or modafinil; OR at least <u>7 days</u> of a preferred amphetamine or methylphenidate drug in this UPDL category and indicated for diagnosis
		 ADDITIONAL OXYBATE SALTS (XYWAV) CRITERIA: Must have documented adherence to sodium restricted diet AR –amphetamine/dextroamphetamine: a PA is required for patients younger than 3 years AR –amphetamine/dextroamphetamine XR, dextroamphetamine ER: a PA is required for patients younger than 6 years

	Central Nervous Syste	em (CNS) Agents: Neuropathic Pain
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
DIBENZA	ZEPINES	LENGTH OF AUTHORIZATIONS: 365 Days
carbamazepine IR, ER	oxcarbazepine susp	
oxcarbazepine tab		STEP THERAPY CRITERIA:
TRILEPTAL SUSP BVG		• Must have had an inadequate clinical response of at least <u>30 days</u> with
GAPAPEN	ITINOIDS	generic lidocaine patch
gabapentin IR	gabapentin ER	
GRALISE ^{BVG}		NON-PREFERRED CRITERIA:
HORIZANT		• Must have had an inadequate clinical response of at least <u>30 days</u> with at
TRICYCLIC ANT	IDEPRESSANTS	least two preferred drugs within the same sub-section classification in
amitriptyline		this UPDL category and indicated for diagnosis
desipramine		
doxepin 10, 25, 50, 75, 100, 150mg		
doxepin soln		
imipramine		
nortriptyline		
OTHER		
duloxetine 20, 30, 60mg	duloxetine 40mg	
lidocaine patch	pregabalin ER	
pregabalin IR		
ZTLIDO ST		

	Central Nervous System	m (CNS) Agents: Parkinson's Agents
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
COMT IN	IHIBITORS	LENGTH OF AUTHORIZATIONS: 365 Days
entacapone	ONGENTYS	
	tolcapone	NON-PREFERRED CRITERIA:
DOPAMIN	E AGONISTS	 Must have had an inadequate clinical response of at least <u>30 days</u> with at
pramipexole IR	apomorphine	least one preferred drug within the same sub-section classification in this
ropinirole IR, ER	КҮММОВІ	UPDL category and indicated for diagnosis
	NEUPRO	
	pramipexole ER	ADDITIONAL APOMORPHINE (APOKYN/KYNMOBI), LEVODOPA INHALATION
MAO-B II	NHIBITORS	(INBRIJA), & ISTRADEFYLLINE (NOURIANZ) CRITERIA:
selegiline	rasagiline	• Must have had inadequate clinical response to at least <u>30 days</u> with one
	XADAGO	other drug for the treatment of "off episodes" (COMT inhibitor, dopamine
	ZELAPAR	agonist, or MAO-B inhibitor)
ОТ	HER	
amantadine cap, tab	amantadine soln	
carbidopa	carbidopa/levodopa dispersible	
carbidopa/levodopa IR, ER	carbidopa/levodopa/entacapone	
	CREXONT	
	GOCOVRI	
	INBRIJA	
	NOURIANZ	
	OSMOLEX ER	
	RYTARY	
	VYALEV	

Central Nervous System (CNS) Agents: Restless Legs Syndrome			
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA	
HORIZANT pramipexole IR ropinirole IR, ER	NEUPRO	 LENGTH OF AUTHORIZATIONS: 365 Days NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>one preferred</u> drug in this UPDL category and indicated for diagnosis 	

Central Nervous System (CNS) Agents: Sedative-Hypnotics, Non-Barbiturate			
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA	
BELSOMRA	DAYVIGO	LENGTH OF AUTHORIZATIONS: 180 Days	
estazolam	doxepin 3, 6mg		
eszopiclone	EDLUAR	NON-PREFERRED CRITERIA:	
ramelteon	flurazepam	Must have had an inadequate clinical response of at least <u>7 days</u> with at	
temazepam	quazepam	least two preferred drugs in this UPDL category and indicated for	
triazolam	QUVIVIQ	diagnosis	
zaleplon	zolpidem cap, SL		
zolpidem tab ER, IR	ZOLPIMIST	ADDITIONAL INFORMATION	
		 Non-controlled medications may be authorized if the prescriber indicates the patient has a history of addiction 	

Central Nervous System (CNS) Agents: Skeletal Muscle Relaxants, Non-Benzodiazepine			
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA	
baclofen susp ^{AR} , tab	baclofen soln	LENGTH OF AUTHORIZATIONS: 365 Days	
chlorzoxazone 500mg	carisoprodol		
cyclobenzaprine IR	chlorzoxazone 250, 375, 750mg	NON-PREFERRED CRITERIA:	
dantrolene	cyclobenzaprine ER	Must have had an inadequate clinical response of at least <u>30 days</u> with at	
metaxalone 800mg	FLEQSUVY AR	least two preferred drugs in this UPDL category and indicated for	
methocarbamol 500, 750mg	LYVISPAH	diagnosis	
orphenadrine	metaxalone 400mg		
tizanidine	methocarbamol 1000mg	ADDITIONAL CARISOPRODOL (SOMA) CRITERIA:	
	orphenadrine/ASA/caffeine	 Must provide medical justification that no other muscle relaxant or agent to treat fibromyalgia, or any musculoskeletal condition would serve the clinical needs of the patient 	
		AR – FLEQSUVY (baclofen susp): a PA is required for patients 12 years and older	

Central Nervous System (CNS) Agents: Smoking Deterrents			
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA	
bupropion SR		All products are covered without a PA	
CHANTIX			
nicotine			
varenicline			

Dermatologic Agents: Oral Acne Products			
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA	
amnesteem ^{PA} claravis ^{PA}	ABSORICA ABSORICA LD	LENGTH OF AUTHORIZATIONS: 150 days	
isotretinoin ^{PA} zenatane ^{PA}		 <u>CLINICAL PA CRITERIA:</u> Must have had an inadequate clinical response of at least <u>90 days</u> with at least <u>one preferred</u> topical AND <u>one preferred</u> oral antibiotic for acne NON-PREFERRED CRITERIA: 	
		 Must have had an inadequate clinical response of at least <u>90 days</u> with at least <u>two preferred</u> drugs in this UPDL category and indicated for diagnosis 	
		 ADDITIONAL INFORMATION Authorization length will be for no more than 150 days at a time then must take 56 days off 	

Dermatologic Agents: Topical Acne Products		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
NON-RE	TINOIDS	LENGTH OF AUTHORIZATIONS: 365 Days
azelaic acid gel	CLINDACIN KIT	
benzoyl peroxide	clindamycin foam	NON-PREFERRED CRITERIA:
clindamycin gel, lot, soln, swabs	clindamycin/benz perox 1.2-3.75%	 Must have had an inadequate clinical response with at least two
clind/benz perox 1-5%, 1.2-2.5%, 1.2-5%	dapsone gel	preferred drugs within the same sub-section classification in
erythromycin	FINACEA FOAM	this UPDL category. Trials must be 30 days for preferred non-
erythromycin/benzoyl peroxide	NEUAC	retinoids and 90 days for preferred retinoids.
ONEXTON GEL BVG	sodium sulfacetamide/sulfur	
sodium sulfacetamide gel, liq	sodium sulfacetamide pads	ADDITIONAL CLINDAMYCIN/ADAPALENE/BENZOYL PEROXIDE
	WINLEVI	(CABTREO) CRITERIA
	ZMA CLEAR SUSP	 Must provide documentation for patient's inability to use the
RETINOIDS/CC	OMBINATIONS	individual drugs in this UPDL category
adapalene gel ^{AR} 0.1%, 0.3%	adapalene cream ^{AR}	
adapalene/benzoyl peroxide AR	ARAZLO AR	ADDITIONAL INFORMATION
ALTRENO AR	CABTREO GEL AR	All retinoids - May be authorized with a diagnosis of skin cancer
RETIN-A MICRO AR BVG 0.04%, 0.1%	clindamycin/tretinoin AR	• tazarotene (TAZORAC) - May be authorized with a diagnosis of
tretinoin ^{AR} cream, gel	RETIN-A MICRO ^{AR} 0.06%, 0.08% ^{BvG}	psoriasis
	tazarotene AR cream, foam, gel 0.1%	
	tretinoin micro ^{AR} 0.04%, 0.1%	AR - All topical retinoids: a PA is required for patients 24 years and older

Endocrine Agents: Androgens			
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA	
PREFERRED AGENTS depo-testosterone ^{AR PA} testosterone cypionate ^{AR PA} testosterone gel 1% packet ^{AR PA} testosterone gel 1.62% pump ^{AR PA}			
		 SUBSEQUENT AUTHORIZATION CRITERIA: Must provide documentation of patient's clinical response to treatment and ongoing safety monitoring (i.e., testosterone and hematocrit) AR: All drugs: a PA is required for patients younger than 18 years 	

Endocrine Agents: Diabetes – Hypoglycemia Treatments			
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA	
BAQSIMI glucagon emerg kit [labeler 00548] GVOKE ZEGALOGUE	glucagon emerg kit [labeler 00378, 63323]	LENGTH OF AUTHORIZATIONS: 365 Days NON-PREFERRED CRITERIA: • Must have had an inadequate clinical response of at least one preferred drug in this UPDL category and indicated for diagnosis OR the inability of the member and/or caregiver to administer a preferred glucagon product in a timely fashion	
		 SUBSEQUENT AUTHORIZATION CRITERIA: Renewal will be allowed for expired/unused products WITHOUT documentation of patient's clinical response to treatment 	

	Endocrine A	nts: Diabetes – Insulin
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
RAPI	D-ACTING	LENGTH OF AUTHORIZATIONS: 365 Days
APIDRA HUMALOG U-100 KWIKPEN, VIAL insulin aspart insulin lispro	ADMELOG AFREZZA FIASP HUMALOG U-100 TEMPO PEN HUMALOG U-200	 STEP THERAPY CRITERIA: Must have had an inadequate clinical response (defined as the inability to reach target A1C) after at least <u>120 days</u> with at least <u>one preferred</u> drug having a similar duration of action in this UPDL category
	LYUMJEV NOVOLOG U-100	NON-PREFERRED CRITERIA:
SHOP HUMULIN R U-500	RT-ACTING HUMULIN R U-100 NOVOLIN R U-100	 Must have had an inadequate clinical response (defined as the inability to reach target A1C) after at least <u>120 days</u> with at least <u>two</u> <u>preferred</u> drugs having a similar duration of action in this UPDL
INTERME	DIATE-ACTING	category and indicated for diagnosis, if available
	HUMULIN N U-100 NOVOLIN N U-100	ADDITIONAL TEMPO PEN CRITERIA
LON	G-ACTING	Must have had an inadequate clinical response or documentation of
LANTUS ^{BVG} LEVEMIR TOUJEO ^{BVG}	BASAGLAR insulin degludec insulin glargine	medical necessity beyond convenience for why the patient cannot use the corresponding FlexPens or Kwikpens
TRESIBA ^{BVG ST}	REZVOGLAR	ADDITIONAL INHALED INSULIN (AFREZZA) CRITERIA:
	SEMGLEE ^{BVG}	 Must provide documentation of spirometry testing prior to initiation
	D INSULIN	with a predicted FEV1 ≥70% - Will not be authorized for patients with
HUMALOG 50-50 HUMALOG 75-25 HUMULIN 70-30	NOVOLIN 70-30 NOVOLOG 70-30	 asthma or COPD Must provide documentation of being nicotine-free for at least 180 days
insulin aspart pro/insulin aspart		ADDITIONAL INFORMATION
		 An inadequate clinical response is defined as the inability to reach A1C goal after at least 120 days of current regimen with documented adherence and appropriate dose escalation. Must include a patient specific A1C goal if less than 7% Must include current A1C (within last 6 months) Requests may be authorized for patients with a condition that is difficult to control (i.e., prone to ketoacidosis, hypoglycemia)

SUBSEQUENT AUTHORIZATION CRITERIA:
Must provide documentation of patient's clinical response to treatment
and ongoing safety monitoring
 Must include a patient specific A1C goal if less than 7%
 Must include current A1C (within last 6 months)

	Endocrine Agents: Dia	betes – Non-Insulin
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
DPP4 INHIBITC	RS/COMBINATIONS	LENGTH OF AUTHORIZATIONS: 365 Days
JANUMET JANUMET XR JANUVIA JENTADUETO KOMBIGLYZE XR ^{BVG} ONGLYZA ^{BVG} TRADJENTA	alogliptin alogliptin/metformin alogliptin/pioglitazone JENTADUETO XR saxagliptin saxagliptin/metformin sitagliptin/metformin (gen of ZITUVIMET) ZITUVIMET XR ZITUVIO ^{BvG}	 NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>120 days</u> with at least <u>three preferred</u> drugs in this UPDL category and indicated for diagnosis, if available ADDITIONAL SITAGLIPTIN (ZITUVIO) CRITERIA Must have had a trial of at least 120 days with JANUVIA OR must provide documentation of medical necessity for patient's inability
GLP-1 RECEPTOR AG	CONISTS/COMBINATIONS	to use JANUVIA
metformin ER (gen of GLUCOPHAGE XR)	BYDUREON BCISE liraglutide MOUNJARO OZEMPIC RYBELSUS SOLIQUA XULTOPHY FORMIN metformin ER (gen of FORTAMET, GLUMETZA)	 ADDITIONAL INFORMATION An inadequate clinical response is defined as the inability to reach A1C goal after at least 120 days of current regimen, with use of two or more drugs concomitantly per ADA guidelines, documented adherence, and appropriate dose escalation (must achieve maximum recommended dose or document that maximum recommended dose is not tolerated or is clinically inappropriate).
metformin IR 500, 850, 1000mg	metformin IR 625, 750mg metformin soln	 For non-preferred drugs that have preferred drugs in the same drug
SGLT2 INHIBITO	DRS/COMBINATIONS	 class: must provide documentation that there was at least <u>one</u> inadequate clinical response with a drug in same drug class SUBSEQUENT AUTHORIZATION CRITERIA: Must provide documentation of patient's clinical response to treatment and ongoing safety monitoring Must include a patient specific A1C goal if less than 7% Must include current A1C (within last 6 months)
FARXIGA ^{BVG} JARDIANCE SYNJARDY XIGDUO XR ^{BVG}	dapagliflozin dapagliflozin/metformin ER GLYXAMBI INVOKAMET INVOKANA QTERN SEGLUROMET STEGLATRO STEGLUJAN SYNJARDY XR	

	TRIJARDY XR
SULFONYLURE	AS/COMBINATIONS
glimepiride 1, 2, 4mg	glimepiride 3mg
glipizide IR, ER	glimepiride/pioglitazone
glipizide/metformin	
glyburide	
glyburide/metformin	
	DTHER
acarbose	SYMLINPEN
miglitol	
nateglinide	
pioglitazone	
pioglitazone/metformin	
repaglinide	

Endocrine Agents: Endometriosis		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
danazol ST DEPO-SUBQ PROVERA 104 ST LUPRON DEPOT ST 3.75, 11.25mg MYFEMBREE ST ORILISSA ST	SYNAREL	 <u>LENGTH OF AUTHORIZATIONS</u>: 365 Days <u>STEP THERAPY CRITERIA</u>: Must have had an inadequate clinical response of at least <u>84 days</u> with at least one preferred NSAID and one oral contraceptive
		 MON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>84 days</u> with at least <u>one preferred</u> step-therapy drug in this UPDL category and indicated for diagnosis
		 A total lifetime duration of therapy of 730 days between ORILISSA and MYFEMBREE or 365 days for LUPRON DEPOT will be authorized

Endocrine Agents: Estrogenic Agents		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
	ORAL	LENGTH OF AUTHORIZATIONS: 365 Days
ANGELIQ estradiol tab ethinyl estradiol/norethindrone PREMARIN TAB PREMPHASE PREMPRO	DUAVEE estradiol/norethindrone MENEST	 MON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>two preferred</u> drugs in this UPDL category within the same subsection classification and indicated for diagnosis
	OPICAL	
DIVIGEL ^{BVG} ELESTRIN estradiol cream	estradiol gel 0.06% (gen of ESTROGEL)	
TRA	NSDERMAL	
CLIMARA COMBIPATCH dotti estradiol patch lyllana MINIVELLE VIVELLE -DOT	EVAMIST MENOSTAR	
v	AGINAL	
ESTRING PREMARIN CREAM	estradiol 10mcg vag tab FEMRING	

Endocrine A		e Agents: Growth Hormone	
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA	
	DAILY-DOSING	LENGTH OF AUTHORIZATIONS: Initial: 180 days; Subsequent: 365 days	
GENOTROPIN ^{PA} NORDITROPIN ^{PA}	HUMATROPE NUTROPIN OMNITROPE SEROSTIM ZOMACTON	 <u>CLINICAL PA CRITERIA:</u> Pediatric Approvals (under 18 years of age): Must be treated and followed by a pediatric endocrinologist, nephrologist, clinical geneticist, endocrinologist, or gastroenterologist 	
	WEEKLY-DOSING	(or as appropriate for diagnosis)	
WEEKLY-DOSING SKYTROFA PAST NGENLA SOGROYA SOGROYA	 Must provide documentation to justify criteria being met, including height, weight, bone age (children), date and results of most current x-ray, stimulus test results, IGF-1 levels, and a growth chart (children) Must not be used in combination with another somatropin agent Adult Approvals (18 years of age or older): Must be treated and followed by an endocrinologist Must provide documentation of growth hormone deficiency by means of a negative response to an appropriate stimulation test (clonidine test is not acceptable for adults) STEP THERAPY CRITERIA: Must have had an inadequate clinical response of at least <u>90 days</u> with at least <u>one preferred</u> daily-dosed growth hormone formulation 		
		NON-PREFERRED CRITERIA:	
		 Must have had an inadequate clinical response of at least <u>90 days</u> with at least <u>one preferred</u> drug within the same sub-section classification in this UPDL category and indicated for diagnosis 	
		SUBSEQUENT AUTHORIZATION CRITERIA:	
		 Must provide documentation of patient's clinical response to treatment and ongoing safety monitoring (i.e., height, weight gain, improved body composition) For adults: must provide documentation by endocrinologist that 	
		discontinuing agent would have a detrimental effect on body composition or other metabolic parameters	

	Endocrine Agents: Osteo	oporosis – Bone Ossification Enhancers
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
B	ISPHOSPHONATES	LENGTH OF AUTHORIZATIONS: 365 Days
alendronate tab ibandronate	alendronate soln BINOSTO FOSAMAX PLUS D risedronate zoledronic acid	 CLINICAL PA CRITERIA: Must have had an inadequate clinical response of at least <u>365 days</u> with <u>one</u> bisphosphonate A total lifetime duration of therapy of 730 days will be authorized
OTHER BONE RESORPTIO	ON SUPPRESSION AND RELATED AGENTS	between any parathyroid analog
calcitonin-salmon FORTEO ^{BVG PA}	EVENITY PROLIA	NON-PREFERRED CRITERIA:
raloxifene	teriparatide TYMLOS	 Must have had an inadequate clinical response of at least <u>365 days</u> with at least <u>one preferred</u> drug within the same sub-section classification in this UPDL category and indicated for diagnosis <u>ADDITIONAL "OTHER BONE RESORPTION SUPPRESSION AND RELATED AGENTS"</u> <u>CRITERIA:</u> Must have had an inadequate clinical response of at least <u>365 days</u> with <u>one</u> bisphosphonate A total lifetime duration of therapy of 730 days will be authorized between any parathyroid analog A total lifetime duration of therapy of 365 days will be authorized for EVENITY
		 ADDITIONAL INFORMATION Patients should only be on ONE of the therapeutic classes (bisphosphonates, calcitonin-salmon)

Endocrine Agents: Progestin Agents		
PREFERRED AGENTS NON-PREFERRED AGENTS PA CRITERIA		
medroxyprogesterone acetate tab megestrol norethindrone acetate progesterone progesterone in oil		All products are covered without a PA

Endocrine Agents: Uterine Fibroids		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
LUPRON DEPOT ^{PA} 3.75, 11.25mg MYFEMBREE ^{PA}		LENGTH OF AUTHORIZATIONS: Up to 180 Days
ORIAHNN PA		CLINICAL PA CRITERIA:
		 Must have had an inadequate clinical response of at least 90 days with at least one oral contraceptive
		ADDITIONAL INFORMATION:
		 A total lifetime duration of therapy of 730 days between MYFEMBREE and ORIAHNN or 365 days for LUPRON DEPOT will be authorized

	Gastrointestinal	Agents: Anti-Emetics
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
5-HT3 ANT	AGONISTS	LENGTH OF AUTHORIZATIONS: 365 Days
granisetron tab	ondansetron 16mg	
ondansetron 4, 8mg	SANCUSO	CLINICAL PA CRITERIA:
ANTICHOL	INERGICS	 dronabinol is only covered for nausea and vomiting associated with
scopolamine		chemotherapy in adult patients who failed at least <u>3 days</u> with at least
ANTIHISTAMINES and ANTIH	IISTAMINE COMBINATIONS	one preferred drug in this UPDL category.
dimenhydrinate	BONJESTA	
diphenhydramine		NON-PREFERRED CRITERIA:
doxylamine/pyridoxine		 Must have had an inadequate clinical response of at least <u>3 days</u> with at least <u>one preferred</u> drug in this UPDL category within the same sub-
meclizine		section classification and indicated for diagnosis
trimethobenzamide		section classification and indicated for aldenosis
PHENOTH	IIAZINES	
prochlorperazine		
promethazine		
SUBSTANCE P/NEUROKININ 1 (NK-1) ANTAGONISTS		
aprepitant 40mg, tripac	aprepitant 80, 125mg	
EMEND 125mg SUSP		
OTHER		
dronabinol PA		
metoclopramide		

Gastrointestinal Agents: Bowel Preparations		
PREFERRED AGENTS NON-PREFERRED AGENTS PA CRITERIA		
CLENPIQ	PLENVU	LENGTH OF AUTHORIZATIONS: 365 Days
GAVILYTE -C	SUFLAVE	
GAVILYTE -G	SUTAB	NON-PREFERRED CRITERIA:
GAVILYTE -N		Must have had an inadequate clinical response with at least one
GOLYTELY		preferred drug in this UPDL category and indicated for diagnosis
MOVIPREP		
sod sulf-potass sulf-mag sulf soln		

Gastrointestinal Agents: Crohn's Disease		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
azathioprine 50mg budesonide ER mercaptopurine tab methotrexate sulfasalazine IR, DR	azathioprine 75, 100mg	 LENGTH OF AUTHORIZATIONS: 365 days NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>two preferred</u> drugs in this UPDL category and indicated for diagnosis

Gastrointestinal Agents: Hepatic Encephalopathy		
PREFERRED AGENTS NON-PREFERRED AGENTS PA CRITERIA		
lactulose XIFAXAN st		LENGTH OF AUTHORIZATIONS: 365 Days
		 STEP THERAPY CRITERIA: Must have had an inadequate clinical response of at least <u>14 days</u> with at least <u>one preferred</u> drug in this UPDL category

Gastrointestinal Agents: Irritable Bowel Syndrome (IBS) with Diarrhea		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
diphenoxylate/atropine loperamide	alosetron VIBERZI	LENGTH OF AUTHORIZATIONS: 365 Days
XIFAXAN ST		 STEP THERAPY CRITERIA: Must have had an inadequate clinical response of at least <u>14 days</u> with at least <u>one preferred</u> drug in this UPDL category
		 NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>14 days</u> with at least <u>one preferred</u> drug and one <u>step therapy</u> drug in this UPDL category and indicated for diagnosis
Gastrointestinal Agents: Pancreatic Enzymes		
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PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
CREON PERTZYE ST	VIOKACE	LENGTH OF AUTHORIZATIONS: 365 Days
ZENPEP		 STEP THERAPY CRITERIA: For a diagnosis of Cystic Fibrosis, no trials required For all other diagnoses, must have had an inadequate clinical response of at least <u>14 days</u> with at least <u>one preferred</u> drug in this UPDL category
		 NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>14 days</u> with at least <u>two preferred</u> drugs in this UPDL category and indicated for diagnosis

Gastrointestinal Agents: Proton Pump Inhibitors		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
esomeprazole	DEXILANT ^{BVG}	LENGTH OF AUTHORIZATIONS: 180 days, except as listed under additional
lansoprazole cap	esomeprazole granules	criteria
NEXIUM GRANULES BVG	KONVOMEP	
omeprazole	lansoprazole ODT	NON-PREFERRED CRITERIA:
pantoprazole tab	omeprazole/sodium bicarbonate	Must have had an inadequate clinical response of at least <u>30 days</u> with
PROTONIX PAK AR BVG	pantoprazole packet AR	at least two preferred drugs in this UPDL category and indicated for
rabeprazole	PRILOSEC SUSP	diagnosis
		ADDITIONAL CRITERIA FOR PPI DOSES GREATER THAN ONCE DAILY
		 Must have had an inadequate clinical response of at least <u>30 days</u> of once daily dosing with the requested drug OR
		 For H. Pylori diagnosis: Must provide documentation of diagnosis Authorization length: 30 days
		 For any of the following diagnoses: carcinoma of GI tract, COPD, Crest Syndrome, dyspepsia, esophageal varices, gastritis, gastroparesis, scleroderma, symptomatic uncomplicated Barret's Esophagus, systemic mastocytosis, or Zollinger Ellison Syndrome: Must provide documentation of diagnosis AND must have failed once-daily dosing of the requested drug
		 Authorization length: 365 days
		ADDITIONAL INFORMATION
		 Request may be authorized If the drug was initiated in the hospital for the treatment of a condition such as a GI bleed or the presence of a gastrostomy and/or jejunostomy (G, GJ, J-tube)
		AR – PROTONIX PAK/pantoprazole packet: a PA is required for patients 6 years and older

Gastrointestinal Agents: Ulcerative Colitis		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
OR	AL CONTRACTOR OF CONTRACTOR	LENGTH OF AUTHORIZATIONS: 365 Days; except UCERIS FOAM – 90 days
balsalazide disodium	DIPENTUM	
budesonide ER tab	mesalamine DR tab 800mg	NON-PREFERRED CRITERIA:
mesalamine DR cap, tab 1.2gm	VELSIPITY	Must have had an inadequate clinical response of at least <u>30 days</u> with at
mesalamine ER cap 0.375gm, 500mg	ZEPOSIA	least two preferred drugs in this UPDL category within the same sub-
PENTASA 250mg		section classification and indicated for diagnosis, if available
sulfasalazine IR, DR		
RECT	TAL CONTRACT	ADDITIONAL BUDESONIDE (UCERIS) CRITERIA:
mesalamine enema, supp	mesalamine enema kit SF ROWASA	 Must have had a documented side effect, allergy, or treatment failure of at least <u>30 days</u> with mesalamine enema or suppository
	UCERIS FOAM ^{BVG}	ADDITIONAL OZANIMOD (ZEPOSIA) AND ETRASIMOD (VELSIPITY) CRITERIA:
		 Must have had a documented side effect, allergy, or treatment failure of at least <u>90 days</u> with at least <u>one preferred Systemic Immunomodulator</u> <u>indicated for Ulcerative Colitis</u> (refer to Immunomodulator Agents: Systemic Inflammatory Disease class for complete list)

Gastrointestinal Agents: Unspecified GI		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
bisacodyl	AEMCOLO	LENGTH OF AUTHORIZATIONS: 365 days except 3 days for AEMCOLO
dicyclomine	AMITIZA	
diphenoxylate/atropine	GATTEX	STEP THERAPY CRITERIA:
lactulose	IBSRELA	 Must have had an inadequate clinical response to at least <u>14 days</u>
LINZESS	MYTESI	with at least two preferred drugs in this UPDL category, if
loperamide	polyethylene glycol oral powder packet	indicated for diagnosis
lubiprostone st	prucalopride	
MOVANTIK ST	RELISTOR	NON-PREFERRED CRITERIA:
polyethylene glycol oral powder bottle	SYMPROIC	 Must have had an inadequate clinical response of at least <u>14 days</u>
senna		with <u>one step therapy</u> drug this UPDL category and indicated for
TRULANCE ST		diagnosis
XIFAXAN ST		
		ADDITIONAL METHYLNALTREXONE (RELISTOR) AND NALDEMEDINE
		(SYMPROIC) CRITERIA:
		 Must have a history of chronic pain requiring continuous opioid therapy for ≥84 days
		ADDITIONAL RIFAMYCIN DELAYED-RELEASE (AEMCOLO) CRITERIA:
		 Must have the inability to take, or failure of ALL of the following: azithromycin, ciprofloxacin, levofloxacin, ofloxacin, or rifaximin
		SUBSEQUENT AUTHORIZATION CRITERIA:
		 Must provide documentation of patient's clinical response to
		treatment and ongoing safety monitoring (i.e., decreased
		frequency of specialized nutrition support or improvement in
		symptoms)

LENGTH OF AUTHORIZATIONS: 365 Days TADALAFIL (CIALIS) CRITERIA:
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TADALAFIL (CIALIS) CRITERIA:
 Must have had an inadequate clinical response of at least <u>30 days</u> with a
least <u>one</u> alpha-1 adrenergic blocker. If prostate volume of > 30cc on
imaging, a prostate specific antigen (PSA) > 1.5ng/dL, or palpable prosta
enlargement on digital rectal exam (DRE), then a trial of at least <u>90 days</u>
finasteride is required.
NON-PREFERRED CRITERIA:
least two preferred drugs, with at least one preferred within the same
sub-section classification and indicated for diagnosis, if available

Genitourinary Agents: Electrolyte Depleter Agents		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
CALCIUM BASED		LENGTH OF AUTHORIZATIONS: 365 Days
calcium acetate, carbonate		
IRON BASED		STEP THERAPY CRITERIA:
VELPHORO ST	ferric citrate tab	Must have had an inadequate clinical response of at least <u>7 days</u> with at
OTHER		least <u>one preferred</u> drug in this UPDL category
sevelamer	FOSRENOL POWDER lanthanum carbonate XPHOZAH	 NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>14 days</u> with at least <u>one preferred</u> step therapy drug in this UPDL category and indicated for diagnosis, if available

Genitourinary Agents: Urinary Antispasmodics		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
ANTIMUSCARINICS		LENGTH OF AUTHORIZATIONS: 365 Days
fesoterodine	darifenacin	
oxybutynin IR, ER	tolterodine IR, ER	NON-PREFERRED CRITERIA:
OXYTROL	VESICARE LS AR	 Must have had an inadequate clinical response of at least <u>30 days</u> with at
solifenacin		least two preferred drugs in this UPDL category and indicated for
trospium IR, ER		diagnosis, one of which must be within the same sub-section classification,
BETA-3 AGONISTS		if available
MYRBETRIQ TAB ^{BVG}	GEMTESA mirabegron tab	AR – MYRBETRIQ GRANULES: a PA is required for patients younger than 3 years old AND 5 years and older
	MYRBETRIQ GRANULES AR	AR – VESICARE LS: a PA is required for patients younger than 2 years old AND 5 years and older

Hyperkalemia Agents: Potassium Binders		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
LOKELMA	kionex susp sodium polystyrene sulfonate VELTASSA	 LENGTH OF AUTHORIZATIONS: 365 Days NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>30 days with at least one preferred</u> drug in this UPDL category and indicated for diagnosis

nts: Systemic Inflammatory Disease
bsequent: 365 days
g/maintenance) will be based
d loading and maintenance dosing
st prior to initiation of biologic
oonse of at least 90 days with at
ed for diagnosis in this UPDL
sponse of at least <u>90 days</u> with at
tegory that are not biosimilars of
ed for diagnosis
llators: must provide
nical response to its preferred
in this UPDL category and
ailable
with a specialist (i.e.
with a specialist (i.e.,
equate clinical response of at

	ZYMFENTRA	ADDITIONAL ATOPIC DERMATITIS CRITERIA:
	OTHER	 Must have at least 10% body surface area (BSA) involvement with an
OTEZLA PA	ENTYVIO	inadequate clinical response of at least <u>90 days</u> with <u>two</u> of the
	ORENCIA	following: topical corticosteroids or topical calcineurin inhibitors [e.g.,
	SOTYKTU	ELIDEL] unless atopic dermatitis is severe and involves >25% BSA
		 ADDITIONAL PRURIGO NODULARIS CRITERIA: Must be prescribed by or in consultation with a specialist (i.e., dermatologist, rheumatologist) Must provide documentation of an inadequate clinical response of at least 90 days with a topical steroid

Infectious Disease Agents: Antibiotics – Cephalosporins		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
cefaclor IR cefaclor susp ^{AR} cefadroxil cefdinir cefprozil cefprozil susp ^{AR} cefuroxime cephalexin cap 250, 500mg cephalexin susp ^{AR}	cefaclor ER cefixime cap cefixime susp ^{AR} cefpodoxime cephalexin cap 750mg, tab	 LENGTH OF AUTHORIZATIONS: Based on indication NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>3 days</u> with at least <u>one preferred</u> drug in this UPDL category and indicated for diagnosis ADDITIONAL INFORMATION Requests may be authorized if the patient is completing a course of therapy that was started in the hospital or other similar location or was started before Medicaid eligibility, only the remaining course will be authorized SUBSEQUENT AUTHORIZATION CRITERIA: Must provide documentation of patient's clinical response to treatment, ongoing safety monitoring, AND medical necessity for continued use AR – cefaclor susp: a PA is required for patients 12 years and older AR – cefprozil susp: a PA is required for patients 12 years and older AR – cephalexin susp: a PA is required for patients 12 years and older

Infectious Disease Agents: Antibiotics – Inhaled		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
tobramycin 300mg/5ml neb soln ^{PA} tobramycin inj	ARIKAYCE CAYSTON TOBI PODHALER tobramycin 300mg/4ml neb soln	 LENGTH OF AUTHORIZATIONS: Initial: 180 days; Subsequent: 365 days CLINICAL PA CRITERIA: Must provide documentation of cultures demonstrating drug is prescribed in alignment with approved indication NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>28 days</u> with at least <u>one preferred</u> drug in this UPDL category and indicated for diagnosis SUBSEQUENT AUTHORIZATION CRITERIA: Must provide documentation of patient's clinical response to treatment and ongoing safety monitoring (i.e., culture conversion, symptom improvement)

Infectious Disease Agents: Antibiotics – Macrolides		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
azithromycin clarithromycin IR, susp ^{AR}	clarithromycin ER erythromycin IR, ER	 LENGTH OF AUTHORIZATIONS: Based on indication NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>3 days</u> with at least <u>one preferred</u> drug in this UPDL category and indicated for diagnosis
		 ADDITIONAL INFORMATION Requests may be authorized if the patient is completing a course of therapy that was started in the hospital or other similar location or was started before Medicaid eligibility, only the remaining course will be authorized
		 SUBSEQUENT AUTHORIZATION CRITERIA: Must provide documentation of patient's clinical response to treatment, ongoing safety monitoring, AND medical necessity for continued use
		AR – clarithromycin susp: a PA is required for patients 12 years and older

Infectious Disease Agents: Antibiotics – Quinolones			
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA	
CIPRO ORAL SUSP ^{AR} ciprofloxacin ciprofloxacin susp ^{AR} levofloxacin soln ^{AR} , tab	BAXDELA ofloxacin	 <u>LENGTH OF AUTHORIZATIONS</u>: Based on indication <u>NON-PREFERRED CRITERIA:</u> Must have had an inadequate clinical response of at least <u>3 days</u> with at 	
moxifloxacin		least <u>one preferred</u> drug in this UPDL category and indicated for diagnosis ADDITIONAL INFORMATION	
		 Requests may be authorized if the patient is completing a course of therapy that was started in the hospital or other similar location or was started before Medicaid eligibility, only the remaining course will be authorized 	
		SUBSEQUENT AUTHORIZATION CRITERIA:	
		 Must provide documentation of patient's clinical response to treatment, ongoing safety monitoring, AND medical necessity for continued use 	
		AR – ciprofloxacin susp: a PA is required for patients 12 years and older	
		AR – CIPRO ORAL SUSP: a PA is required for patients 12 years and older	
		AR – levofloxacin oral soln: a PA is required for patients 12 years and older	

Infectious Disease Agents: Antibiotics – Tetracyclines			
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA	
doxycycline 20, 50, 100mg doxycycline susp ^{AR} minocycline IR tetracycline	demeclocycline doxycycline 75, 150mg doxycycline DR minocycline ER MINOLIRA NUZYRA	LENGTH OF AUTHORIZATIONS: Based on indication for acute infections or 365 days for acne NON-PREFERRED CRITERIA: • Must have had an inadequate clinical response of at least <u>3 days</u> with at least <u>one preferred</u> drug for acute infections OR at least <u>90 days</u> with at least <u>one preferred oral</u> drug for acute infections OR at least <u>90 days</u> with at least <u>one preferred oral</u> drug for acne in this UPDL category and indicated for diagnosis ADDITIONAL INFORMATION • Requests may be authorized if the patient is completing a course of therapy that was started in the hospital or other similar location or was started before Medicaid eligibility, only the remaining course will be authorized SUBSEQUENT AUTHORIZATION CRITERIA: • Must provide documentation of patient's clinical response to treatment, ongoing safety monitoring, AND medical necessity for continued use AR – doxycycline susp: a PA is required for patients 12 years and older	

Infectious Disease Agents: Antifungals		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
clotrimazole	BREXAFEMME	LENGTH OF AUTHORIZATIONS: Based on indication
fluconazole	CRESEMBA	
griseofulvin	flucytosine	NON-PREFERRED CRITERIA:
itraconazole cap	itraconazole soln	 Must have had an inadequate clinical response of at least <u>3 days</u> with at
ketoconazole	NOXAFIL PAK	least two preferred drugs in this UPDL category and indicated for
nystatin	ORAVIG	diagnosis
terbinafine voriconazole susp ^{AR} , tab	posaconazole TOLSURA	ADDITIONAL OTESECONAZOLE (VIVJOA) CRITERIA:
	VIVJOA	 Must provide documentation of at least three symptomatic episodes of vulvovaginal candidiasis in the past 12 months
		 Must provide documentation of non-reproductive potential (i.e., post- menopausal)
		 Must have had an inadequate clinical response of at least <u>180-day</u> maintenance course with oral fluconazole shown by documentation of more than <u>one</u> breakthrough infection
		ADDITIONAL INFORMATION:
		 posaconazole can be approved for aspergillosis treatment and prophylaxis without trials of preferred agents
		 Requests may be authorized if the patient is completing a course of therapy that was started in the hospital or other similar location or was started before Medicaid eligibility, only the remaining course will be authorized
		SUBSEQUENT AUTHORIZATION CRITERIA:
		• Must provide documentation of patient's clinical response to treatment, ongoing safety monitoring, AND medical necessity for continued use
		AR – voriconazole susp: a PA is required for patients 12 years and older

Infectious Disease Agents: Antivirals – Hepatitis C Agents		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
MAVYRET PA PEGASYS PA ribavirin PA sofosbuvir/velpatasvir PA	HARVONI ledipasvir/sofosbuvir SOVALDI VOSEVI ZEPATIER	LENGTH OF AUTHORIZATIONS: Dependent upon authorized course CLINICAL PA CRITERIA: • Only regimens recommended by the American Association for the Study of Liver Diseases (AASLD) will be authorized • Please see the Hepatitis C Direct Acting Antiviral Prior Authorization Form for criteria
		 NON-PREFERRED CRITERIA: Must have had an inadequate clinical response defined as not achieving sustained virologic response (SVR) with guideline-recommended preferred drugs in this UPDL category and indicated for diagnosis

Infectious Disease Agents: Antivirals – Herpes		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
acyclovir valacyclovir	famciclovir SITAVIG	LENGTH OF AUTHORIZATIONS: For the duration of the prescription (up to 180 days)
		 NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>3 days</u> with at least <u>one preferred</u> drug in this UPDL category and indicated for diagnosis

	Infectious Disease Agents: An	ntivirals – HIV* LEGACY CATEGORY
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
INTEGRASE STRAND	TRANSFER INHIBITORS	LENGTH OF AUTHORIZATIONS: 365 Days
APRETUDE	VOCABRIA	
ISENTRESS		ABACAVIR/DOLUTEGRAVIR/LAMIVUDINE (TRIUMEQ PD) CRITERIA:
ISENTRESS CHEW TAB AR		Must provide documentation of patient's weight (only authorized for
TIVICAY		those 6 – 25 kg)
TIVICAY PD		
NUCLEOSIDE REVERSE TE	RANSCRIPTASE INHIBITORS	NON-PREFERRED CRITERIA:
abacavir	abacavir soln	• Must have had an inadequate clinical response of at least <u>30 days</u> with
emtricitabine	EMTRIVA SOLN	at least <u>one preferred</u> drug in this UPDL category and indicated for
entecavir	lamivudine tab	diagnosis. If applicable, the request must address the inability to use
lamivudine soln ^{AR}	VIREAD 250, 300mg TAB	the individual components.
tenofovir dis fum 300mg		AD ISENITRESS CHEVAARIE TARIET, a RA is required for notionts 12 years
VIREAD 150, 200mg TAB, POWDER		AR – ISENTRESS CHEWABLE TABLET: a PA is required for patients 12 years and older
zidovudine		AR – lamivudine soln: a PA is required for patients 3 years and older
NON-NUCLEOSIDE REVERSE	TRANSCRIPTASE INHIBITORS	AR – nevirapine soln: a PA is required for patients 3 years and older
efavirenz	EDURANT	
nevirapine soln ^{AR}	etravirine	
PIFELTRO	nevirapine IR, ER tab	
PROTEASE	INHIBITORS	
atazanavir	APTIVUS	
darunavir 600, 800mg tab	fosamprenavir	
EVOTAZ	NORVIR POWDER	
PREZCOBIX	PREZISTA SUSP, 75, 150mg TAB	
REYATAZ POWDER	VIRACEPT	
ritonavir tab		
OTHER SINGLE INGREDIENT PRODUCTS		
RUKOBIA	FUZEON	
	SELZENTRY ^{BVG}	
	SUNLENCA	
	TYBOST	
COMBINATIO	ON PRODUCTS	

abacavir/lamivudine	CIMDUO
BIKTARVY	lamivudine/zidovudine
CABENUVA	STRIBILD
COMPLERA	SYMFI ^{BVG}
DELSTRIGO	SYMFI LO ^{BVG}
DESCOVY	
DOVATO	
efavirenz/emtricitabine/tenofovir	
emtricitabine/tenofovir dis fum	
GENVOYA	
JULUCA	
lopinavir/ritonavir	
ODEFSEY	
SYMTUZA	
TRIUMEQ	
TRIUMEQ PD PA	

Ophthalmic Agents: Antibiotic and Antibiotic-Steroid Combination Drops and Ointments		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
bacitracin-polymyxin	AZASITE	LENGTH OF AUTHORIZATIONS: 30 days
CILOXAN	bacitracin	
ciprofloxacin	BESIVANCE	NON-PREFERRED CRITERIA:
erythromycin	gatifloxacin	 Must have had an inadequate clinical response of at least <u>3 days</u> with at
gentamicin	moxifloxacin (gen of MOXEZA)	least two preferred drugs in this UPDL category and indicated for
moxifloxacin	neo/poly/hydrocortisone	diagnosis
neo/poly/bacitracin	sulfacetamide sodium oint 10%	
neo/poly/bacitracin/hydrocortisone	TOBRADEX ST	ADDITIONAL INFORMATION
neo/poly/dexamethasone	ZYLET	• Requests may be authorized if the patient is completing a course of
neo/poly/gramicidin		therapy that was started in the hospital or other similar location or was
ofloxacin		started before Medicaid eligibility, only the remaining course will be
polymyxin/trimethoprim		authorized
sulfacetamide sodium soln 10%		
sulfacetamide/prednisolone		
TOBRADEX OINT		
tobramycin		
tobramycin/dexameth 0.3/0.1%		
TOBREX OINT		

Ophthalmic Agents: Antihistamines & Mast Cell Stabilizers		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
azelastine BEPREVE ^{BVG}	alomide bepotastine	LENGTH OF AUTHORIZATIONS: 365 Days
cromolyn ketotifen olopatadine	epinastine ZERVIATE	 NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>7 days</u> with at least <u>two preferred</u> drugs in this UPDL category and indicated for diagnosis

Ophthalmic Agents: Dry Eye Treatments		
NON-PREFERRED AGENTS	PA CRITERIA	
CEQUA cyclosporine EYSUVIS MIEBO RESTASIS MULTI-DOSE TYRVAYA VEVYE	 LENGTH OF AUTHORIZATIONS: 14 days for EYSUVIS; 365 days for all other drugs STEP THERAPY CRITERIA: Must have had an inadequate clinical response of at least <u>14 days</u> with at least <u>one</u> preferred drug in this UPDL category in the previous 120 days NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>14 days</u> with at least <u>one</u> preferred drug in this UPDL category in the previous 120 days 	
	NON-PREFERRED AGENTS CEQUA cyclosporine EYSUVIS MIEBO RESTASIS MULTI-DOSE TYRVAYA	

	Ophthalmic Age	nts: Glaucoma Agents
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
ALPHA-2 A	AGONISTS	LENGTH OF AUTHORIZATIONS: 365 Days
ALPHAGAN P 0.1% ^{BvG} ALPHAGAN P 0.15% ^{BvG} brimonidine 0.2% BETA BL betaxolol carteolol levobunolol timolol gel, soln CARBONIC ANHYD	BETIMOL 0.25% BETOPTIC S timolol hemihydrate soln 0.5% timolol maleate once daily, PF	 STEP THERAPY CRITERIA: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>one preferred</u> drug in the same sub-section classification in this UPDL category and indicated for diagnosis, if available NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>two preferred</u> drugs within the same sub-section classification in this UPDL category and indicated for diagnosis of at least <u>30 days</u> with at least <u>two preferred</u> drugs within the same sub-section classification in this UPDL category and indicated for diagnosis
AZOPT ^{BVG ST} dorzolamide PROSTAG	brinzolamide	
latanoprost TRAVATAN Z ^{BVG ST}	bimatoprost IYUZEH LUMIGAN tafluprost travoprost VYZULTA XELPROS	
OTH	IER	
COMBIGAN ^{BVG ST} dorzolamide/timolol RHOPRESSA ROCKLATAN SIMBRINZA	brimonidine/timolol	

Ophthalmic Agents: NSAIDs		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
diclofenac	ACUVAIL	LENGTH OF AUTHORIZATIONS: 30 days
flurbiprofen	bromfenac	
ketorolac	ILEVRO	NON-PREFERRED CRITERIA:
NEVANAC		Must have had an inadequate clinical response of at least <u>3 days</u> with at
		least two preferred drugs in this UPDL category and indicated for
		diagnosis

Ophthalmic Agents: Ophthalmic Steroids		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
	INVELTYS	LENGTH OF AUTHORIZATIONS: 30 days
dexamethasone sodium phosphate	LOTEMAX SM	
difluprednate	loteprednol	NON-PREFERRED CRITERIA:
DUREZOL		• Must have had an inadequate clinical response of at least <u>7 days</u> with at
FLAREX		least two preferred drugs in this UPDL category and indicated for
fluorometholone		diagnosis
FML FORTE		
MAXIDEX		
PRED FORTE		
PRED MILD		
prednisolone acetate		
prednisolone sodium phosphate		

Otic Agents: Antibacterial and Antibacterial/Steroid Combinations		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
CIPRO HC ciprofloxacin/dexamethasone	ciprofloxacin ciprofloxacin/fluocinolone	LENGTH OF AUTHORIZATIONS: 30 days
CORTISPORIN-TC neomycin/poly B/hydrocortisone ofloxacin		 NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>3 days</u> with at least <u>two preferred</u> drugs in this UPDL category and indicated for
		diagnosis

Respiratory Agents: Antihistamines – Second Generation		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
cetirizine cap, syr, tab cetirizine/pseudoephedrine	cetirizine chewable ^{AR} CLARINEX-D	LENGTH OF AUTHORIZATIONS: 365 Days
desloratadine fexofenadine levocetirizine loratadine rapid dissolve loratadine syr, tab	loratadine chewable AR fexofenadine/pseudoephedrine	 MON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>7 days</u> with at least <u>two preferred</u> drugs in this UPDL category and indicated for diagnosis
loratadine/pseudoephedrine		AR – cetirizine chewable, loratadine chewable: a PA is required for patients 6 years and older

Respiratory Agents: Cystic Fibrosis		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
ALYFTREK ^{PA} KALYDECO ^{PA} ORKAMBI ^{PA} PULMOZYME ^{PA} SYMDEKO ^{PA} TRIKAFTA ^{PA} PAK ^{AR} , TAB	BRONCHITOL	 LENGTH OF AUTHORIZATIONS: Initial: 90 days; Subsequent: 365 days CLINICAL PA CRITERIA: Must be prescribed by or in consultation with a pulmonologist or infectious disease specialist Must provide documentation of the specific Cystic Fibrosis Transmembrane Conductance Regular (CFTR) genetic mutation NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>one preferred</u> drug in this UPDL category and indicated for diagnosis ADDITIONAL BRONCHITOL CRITERIA: Must be used as an add-on maintenance therapy Must provide documentation of a completed BRONCHITOL Tolerance Test
		AR – TRIKAFTA PAK: a PA is required for patients 6 years and older

Respiratory Agents: Epinephrine		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
epinephrine (labeler 49502) EPIPEN	AUVI-Q epinephrine (labeler 00093, 00115)	LENGTH OF AUTHORIZATIONS: 365 Days
EPIPEN JR	NEFFY	 NON-PREFERRED CRITERIA: Must have had an inadequate clinical response to at least <u>one preferred</u> drug in this UPDL category and indicated for diagnosis
		 SUBSEQUENT AUTHORIZATION CRITERIA: Subsequent reauthorizations for expired epinephrine auto-injectors are allowable

Respiratory Agents: Hereditary Angioedema		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
	ACUTE	LENGTH OF AUTHORIZATIONS: Acute: 30 days; Prophylaxis: 180 Days
BERINERT PA	KALBITOR	
icatibant acetate PA	RUCONEST	CLINICAL PA CRITERIA:
	PROPHYLAXIS	Acute Treatment
TAKHZYRO ^{PA}	CINRYZE HAEGARDA ORLADEYO	 Must provide documentation that diagnosis is verified by a C4 level below the lower limit of normal as defined by laboratory testing AND one of the following: C1 inhibitor (C1-INH) antigenic level below the lower limit of normal as defined by laboratory testing; OR C1-INH functional level below the lower limit of normal as defined by laboratory testing Prophylactic Treatment Must provide documentation that diagnosis is verified by a C4 level below the lower limit of normal as defined by laboratory testing Prophylactic Treatment Must provide documentation that diagnosis is verified by a C4 level below the lower limit of normal as defined by laboratory testing AND one of the following: C1 inhibitor (C1-INH) antigenic level below the lower limit of normal as defined by laboratory testing; OR C1 inhibitor (C1-INH) antigenic level below the lower limit of normal as defined by laboratory testing; OR C1-INH functional level below the lower limit of normal as defined by laboratory testing; OR C1-INH functional level below the lower limit of normal as defined by laboratory testing; OR Presence of a known HAE-causing C1-INH mutation All indications History of moderate or severe attacks such as airway swelling, severe abdominal pain, facial swelling, nausea and vomiting, or painful facial distortion
		NON-PREFERRED CRITERIA:
		 Must have had an inadequate clinical response of at least <u>3 days</u> with at least <u>one preferred</u> acute drug in this UPDL category and indicated for diagnosis to request a non-preferred acute drug. Must have had an inadequate clinical response of at least <u>14 days</u> with at least <u>one preferred</u> prophylaxis drug in this UPDL category and indicated for diagnosis to request a non-preferred prophylaxis drug in this UPDL category and indicated for diagnosis to request a non-preferred prophylaxis drug.

	Respiratory Age	ents: Inhaled Agents
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
ANTICHOLINERGIC BRONCHODILATORS/COMBINATIONS		LENGTH OF AUTHORIZATIONS: 365 Days
ANORO ELLIPTA ^{BVG} ATROVENT HFA COMBIVENT RESPIMAT INCRUSE ELLIPTA ipratropium/albuterol neb soln SPIRIVA ^{BVG} STIOLTO ADRENERGIC BRO albuterol HFA albuterol neb 0.021% (0.63mg/3mL), 0.042% (1.25mg/3mL) ^{AR} albuterol neb 0.083% (2.5mg/3mL) albuterol neb 0.5% (5mg/mL) conc arformoterol neb SEREVENT DISKUS STRIVERDI RESPIMAT VENTOLIN HFA XOPENEX HFA ^{BVG}	BEVESPI AEROSPHERE DUAKLIR PRESSAIR tiotropium inhaled caps TUDORZA umeclidinium/vilanterol YUPELRI	 <u>NON-PREFERRED CRITERIA:</u> Must have had an inadequate clinical response of at least <u>14 days</u> with at least <u>two preferred</u> drugs in this UPDL category within the same subsection classification and indicated for diagnosis <u>ADDITIONAL STEROID-CONTAINING INHALER CRITERIA:</u> Must have had an inadequate clinical response of at least <u>14 days</u> with at least one preferred steroid-containing drug <u>ADDITIONAL BUDESONIDE/ALBUTEROL (AIRSUPRA) CRITERIA:</u> Must have had an inadequate clinical response of at least <u>14 days</u> with at least one preferred steroid-containing drug <u>ADDITIONAL BUDESONIDE/ALBUTEROL (AIRSUPRA) CRITERIA:</u> Must have had an inadequate clinical response of at least <u>14 days</u> with either DULERA or SYMBICORT AR – albuterol nebulizer soln 0.021% (0.63mg/3mL), 0.042% (1.25mg/3mL): a PA is required for patients 13 years and older AR – budesonide nebulizer soln: a PA is required for patients 13 years and older
BRONCHODILATOR/GLUCOC	ORTICOID COMBINATIONS	
ADVAIR DISKUS ^{BvG} ADVAIR HFA ^{BvG} DULERA SYMBICORT ^{BvG} GLUCOCO	AIRDUO DIGIHALER AIRSUPRA BREO ELLIPTA ^{BVG} BREYNA BREZTRI AEROSPHERE budesonide/formoterol fluticasone/salmeterol TRELEGY ELLIPTA WIXELA INHUB	

ARNUITY ELLIPTA	ALVESCO
ASMANEX TWISTHALER	ARMONAIR DIGIHALER
budesonide neb susp ^{AR}	ASMANEX HFA
FLOVENT	
fluticasone propionate	
PULMICORT FLEXHALER	
QVAR	
OTHER	
cromolyn neb soln	

Respiratory Agents: Leukotriene Receptor Modifiers & Inhibitors		
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
montelukast zafirlukast ^{s⊤}	zileuton ER ZYFLO	LENGTH OF AUTHORIZATIONS: 365 Days
		 STEP THERAPY CRITERIA: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>one preferred</u> drug in this UPDL category
		 NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>two preferred</u> drugs in this UPDL category and indicated for diagnosis

	Respiratory Agents:	Monoclonal Antibodies-Anti-IL/Anti-IgE
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA
DUPIXENT ^{PA} FASENRA ^{PA} XOLAIR ^{PA}	CINQAIR NUCALA TEZSPIRE	 LENGTH OF AUTHORIZATIONS: Initial: 180 days; Subsequent: 365 days CLINICAL PA CRITERIA: Must be prescribed by or in consultation with an applicable specialist (i.e., allergist/ immunologist, pulmonologist, or otolaryngologist) For Asthma – Must have had uncontrolled asthma symptoms and/or exacerbations despite at least <u>30 days</u> with: Medium dose preferred ICS/LABA inhaler for 6 years and older OR medium dose preferred ICS/LABA inhaler with tiotropium or high dose ICS/LABA inhaler if 12 years and older For Chronic Rhinosinusitis with Nasal Polyposis – Must have had an inadequate clinical response of at least <u>30 days</u> to at least <u>one oral</u> corticosteroid AND <u>one nasal</u> corticosteroid spray For Chronic Urticaria – Must have had an inadequate clinical response to at least <u>14 days</u> with at least <u>two different</u> second-generation antihistamines at 4 times standard dose For Chronic Obstructive Pulmonary Disease (COPD): The patient must have an eosinophilic count of greater than or equal to 300 cells per mcL within 12 months prior to initiation of therapy AND The patient has a history of uncontrolled disease, as indicated by greater than or equal to 2 COPD exacerbations or greater than or equal to 1 COPD exacerbation resulting in a hospitalization despite being on standard of care, defined as triple therapy (LAMA+LABA+ICS) for at least 3 months prior to request, and at a stable dose for at least 1 month prior.
		 NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>90 days</u> with at least <u>one preferred</u> drug in this UPDL category and indicated for diagnosis
		 SUBSEQUENT AUTHORIZATION CRITERIA: Must provide documentation of patient's clinical response to treatment and ongoing safety monitoring (i.e., PFT improvement, reduced affected BSA)

Respiratory Agents: Nasal Preparations			
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA	
GLUCOCORTI	COIDS/COMBINATIONS	LENGTH OF AUTHORIZATIONS: 365 days	
flunisolide fluticasone (gen of FLONASE)	azelastine/fluticasone spray BECONASE AQ mometasone OMNARIS QNASL RYALTRIS XHANCE ZETONNA	 Mon-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>14 days</u> with at least <u>two preferred</u> drugs in this UPDL category within the same sub-section classification and indicated for diagnosis 	
OTHER			
azelastine			
ipratropium			
olopatadine			

Respiratory Agents: Pulmonary Fibrosis				
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA		
OFEV PA	pirfenidone	LENGTH OF AUTHORIZATIONS: 365 Days		
		CLINICAL PA CRITERIA:		
		Must be prescribed by or in consultation with a pulmonologist		
		NON-PREFERRED CRITERIA:		
		Must have had an inadequate clinical response of at least <u>30 days</u> with		
		at least one preferred drug in this UPDL category and indicated for		
		diagnosis		

Topical Agents: Antifungals				
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA		
ALEVAZOL	ciclopirox kit	LENGTH OF AUTHORIZATIONS: 365 days		
butenafine	JUBLIA			
ciclopirox	ketoconazole foam	NON-PREFERRED CRITERIA:		
clotrimazole	luliconazole	• Must have had an inadequate clinical response of at least 14 days		
clotrimazole/betamethasone	miconazole/zinc/white petrolatum oint	with at least two preferred drugs in this UPDL category and		
econazole	naftifine	indicated for diagnosis		
ketoconazole	oxiconazole			
miconazole	OXISTAT	ADDITIONAL EFINACONAZOLE (JUBLIA) CRITERIA:		
nystatin	tavaborole	Must have had an inadequate clinical response of at least 48 week		
nystatin/triamcinolone	tolnaftate soln	of ciclopirox AND 6 weeks of oral terbinafine (if fingernail) OR 12		
terbinafine		weeks of oral terbinafine (if toenail)		
tolnaftate cream, powder				

Topical Agents: Antiparasitics				
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA		
NATROBA ^{BvG} permethrin piperonyl butoxide/pyrethrins VANALICE	CROTAN ivermectin lot malathion spinosad	 <u>LENGTH OF AUTHORIZATIONS</u>: 14 Days <u>NON-PREFERRED CRITERIA:</u> Must have had an inadequate clinical response of at least <u>14 days</u> with at least <u>one preferred</u> drug in this UPDL category and indicated for diagnosis 		

Topical Agents: Corticosteroids				
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA		
LOW POTENCY		LENGTH OF AUTHORIZATIONS: 365 days		
desonide cream, oint	alclometasone			
fluocinolone acetonide 0.01%	desonide lotion	NON-PREFERRED CRITERIA:		
hydrocortisone	TEXACORT	Must have had an inadequate clinical response of at least <u>14</u>		
MEDIUM	POTENCY	days with at least two preferred drugs within the same sub-		
betamethasone valerate	betamethasone val aerosol foam	section classification in this UPDL category and indicated for		
flurandrenolide	clocortolone pivalate	diagnosis, if available		
fluticasone propionate cream, oint	fluocinolone acetonide 0.025%			
prednicarbate	fluticasone propionate lotion			
triamcinolone cream, lotion, oint	hydrocortisone butyrate, valerate			
	PANDEL			
	triamcinolone spray			
HIGH P	OTENCY			
betamethasone dip/calcipotriene oint	betamethasone dip			
fluocinonide 0.05%	betamethasone dip/calcipotriene susp			
mometasone furoate	desoximetasone			
	diflorasone diacetate			
	ENSTILAR			
	halcinonide			
ULTRA HIGH POTENCY				
clobetasol propionate	APEXICON E			
	BRYHALI			
	fluocinonide 0.1%			
	halobetasol propionate			
	ULTRAVATE			

Topical Agents: Immunomodulators				
PREFERRED AGENTS	NON-PREFERRED AGENTS	PA CRITERIA		
ELIDEL ^{AR} pimecrolimus ^{AR} [labeler 68682] tacrolimus ^{AR}	EUCRISA HYFTOR OPZELURA pimecrolimus ^{AR} [labeler 00591, 68462] VTAMA ZORYVE CREAM, FOAM	 LENGTH OF AUTHORIZATIONS: 365 Days NON-PREFERRED CRITERIA: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>one preferred</u> drug in this UPDL category and indicated for diagnosis ADDITIONAL ROFLUMILAST (ZORYVE) CRITERIA: 0.15% CREAM: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>one preferred</u> topical corticosteroid OR topical calcineurin inhibitor 0.3% CREAM: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>one preferred</u> topical corticosteroid OR topical calcipotriene FOAM: Must have had an inadequate clinical response of at least <u>30 days</u> with at least <u>one preferred</u> agent indicated for Seborrheic Dermatitis (such as a topical antifungal, topical calcineurin inhibitor, or topical corticosteroid) AR – ELIDEL, pimecrolimus, and tacrolimus: a PA is required for patients younger than 2 years old 		