Drug classes not included on this list are not managed through a Preferred Drug List (PDL). HOWEVER, THIS EXCLUSION IS NOT A GUARANTEE OF PAYMENT OR COVERAGE. Dosage limits and other requirements may apply.

Drugs new to market are non-preferred until a clinical review has been completed. PA criteria will apply to both the pediatric population, as well as the adult population for those plans where PA/PDL limits are allowed

Yellow highlighted items below indicate new changes to the PDL. Red font indicates quantity/dosage limits apply. *Indicates BRAND is Preferred. May Use DAW 5. Contact the OptumRx PA Helpdesk @ 877-207-1126 for prior authorization if client has primary insurance that will not cover the brand name medication.

THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIES THE UST 5 NOT ALL INCLUSIVE PLASE CONTACT CONTINUE WITH ANY QUESTIONS
ADDICTION	BUPRENORPHINE	COMBINATIONS buprenorphine/naloxone tablets SUBOXONE FILM*	Client must have a diagnosis of opioid dependence or abuse. This is not to be used for the treatment of chronic pain. Prior authorization will be required before any narcotic, benzodiazepine, or carisoprodol prescription will be allowed between fills. Prior authorization will be required before any short-acting stimulant prescription from any doctor other than the prescriber of buprenorphine or suboxone, will be allowed between fills.	buprenorphine (oral) buprenorphine/naloxone film BRAND IS PREFERRED) ZUBSOLV
			Oral buprenorphine will be approved for clients with a documented allergy to naloxone.	
			Please submit PA requests on the "Oral Buprenorphine/Naloxone or Oral Buprenorphine" PA form available at www.wymedicaid.org.	
		OXONE	Dosage limits apply Prior authorization will be required for doses >24mg Kloxxado, naloxone products, and Narcan nasal spray will be limited to one fill per 180 days	OPVEE
	KLOXXADO naloxone nasal spray NARCAN	UXUNE	NOXAOU, naloxone products, and Marcan nasa spray will be inniced to one nil per 100 days without nrice authorization. Naloxone formulations available in quantities of 10ml will require prior authorization.	REXTOVY ZIMHI
	NALT	REXONE naltrexone	Client must have a diagnosis of alcohol or opioid dependance.	topiramate*
			Prior authorization will be required before any narcotic, carisoprodol, or benzodiazepine prescription will be allowed between fills. Prior authorization will be required before a short- acting stimulant prescription from any doctor other than the prescriber of naltrexone or Vivitrol will be allowed between fills. *Topiramate requires 4 week trial and failure of naltrexone or acamprosate in AUD	
LERGY / ASTHMA / COPD	ANTIHISTAMINES cetirizine fexofenadine loratadine	, MINIMALLY SEDATING	Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent.	desloratadine CLARINEX RDT/SYRUP levocetirizine
	ANTIHISTAMINE/DECO cetirizine/pseudoephedrine fexofenadine/pseudoephedrine loratadine/pseudoephedrine	NGESTANT COMBINATIONS	Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent.	CLARINEX-D
		C BRONCHODILATORS	Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent.	TIOTROPIUM BROM (<i>use brand</i>) TUDORZA YUPELRI
	SPIRIVA RESPIMAT	OMBINATION AGENTS	Spiriva 5 day STARTER package will be allowed one (1) time per recipient. Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent.	BEVESPI BREZTRI DUAKLIR TRELEGY
	LEUKOTRIE montelukast	NE MODIFIERS	**Will also require the diagnosis of COPD. Trial and failure of preferred agent greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent.	zafirlukast
	LONG ACTING BR arformoterol SEREVENT STRIVERDI	ONCHODILATORS	Trial and failure of two (2) preferred agents greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent.	BROVANA
		THISTAMINES	Trial and failure of preferred agent greater than or equal to 90 days in the last 12 months will be required before approval can be given for a non-preferred agent.	azelastine 0.15% DYMISTA (<i>use separate agents</i>) olopatadine 0.6% RYALTRIS
	NASAL budesonide flunisolide fluticasone mometasone	STEROIDS	Trial and failure of two (2) preferred agents greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent. Budesonide will be approved for pregnancy.	DYMISTA (use separate agents) OMNARIS QNASL XHANCE ZETONNA
		NCHODILATORS - INHALERS	Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent. Prior authorization will be required after a total of 12 albuterol inhalers are dispensed within 365 days. Minimum day supply of 16 days is required.	levalbuterol (BRAND IS PREFERRED) PROAR DIGHALER PROVENTIL HFA XOPENEX HFA
	STEROID AIRDUO RESPICLICK ARNUITY ELLIPTA ASMANEX TWISTHALER budesonide suspension PULMICORT FLEXHALER	INHALANTS	Trial and failure of two (2) preferred agents greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent. *Fluticasone HFA and Asmanex HFA will be approved for pediatric clients 8 years of age or younger. Alvesco will be approved for a history of oral thrush with steroid inhalants.	AIRDUO DIGIHALER AIRSUPRA ALVESCO ARMONAIR ASMANEX HFA* fluticasone HFA*
	ADVAIR (HFA, Diskus) BREO ELLIPTA** DULERA	BINATION AGENTS	Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent.	QVAR REDIHALER fluticasone/vilanterol (use preferred agent fluticasone/salmeterol 55-14/113-14/232- fluticasone/salmeterol 100-50/250-50/500 (BRAND IS PREFERRED)
	SYMBICORT*	EPHRINE	**Will also require the diagnosis of COPD or uncontrolled asthma. Advair 7 and 14-day STARTER package will be allowed one (1) time per recipient.	TRELEGY WIXELA AUVI-Q (use preferred agent)
	epinephrine auto-injector pen EPI-PEN EOSINOPHILIC	ASTHMA AGENTS	Approval for these agents will require additional clinical criteria which can be found on the	FASENRA
		DUPIXENT XOLAIR	Additional Therapeutic Criteria Chart.	NUCALA TEZSPIRE

THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLI THIS UST IS NOT ALL INCLUSIVE PLASE CONTACT Optimile. WITH ANY OUTSTIDIES
THRITIS		MODULATORS SPONDYLITIS (AS)	Client must have diagnosis of AS prior to approval of a preferred agent. To receive a non- preferred agent, client must have diagnosis of AS and 56-day trial and failure of two preferred	CIMZIA** COSENTYX
		ENBREL	agente	REMICADE
		HUMIRA TALTZ	**Cimzia will be allowed for clients that are pregnant or breast-feeding	RINVOQ SIMPONI
			Quantity Limits apply for all diagnoses:	XELJANZ/XR
			Enbrel 25mg - limited to 10 per month	
			Enbrel 50mg - limited to 5 per month Humira 20mg - limited to 10 per month	
			Humira 40mg - limited to 5 per month	
	JUVENILE IDIOPA	THIC ARTHRITIS (JIA)	Client must have diagnosis of JIA prior to approval of a preferred agent. To receive a non-	ACTEMRA
		ENBREL HUMIRA	preferred agent, client must have a diagnosis of JIA and a 56-day trial and failure of both	ILARIS ORENCIA
			preferred agents.	XELJANZ/XR
	PSORIATIC	ARTHRITIS (PA)	Client must have diagnosis of PA prior to approval of a preferred agent. To receive a non-	CIMZIA**
		ENBREL HUMIRA	preferred agent, client must have a diagnosis of PA and a 56-day trial and failure of two preferred agents.	COSENTYX ORENCIA
		OTEZLA*	preferred agents.	REMICADE
		TALTZ		RINVOQ SIMPONI
				SKYRIZI
			*Otezla starter pack is non-preferred	STELARA
			**Cimzia will be allowed for clients that are pregnant or breast-feeding	TREMFYA XELJANZ/XR
	RHEUMATOID	ARTHRITIS (RA)	Client must have diagnosis of RA and a 56-day trial and failure of methotrexate prior to approval	ACTEMRA
		ENBREL HUMIRA	of a preferred agent. To receive a non-preferred agent, client must have a diagnosis of RA and a	CIMZIA* KEVZARA
		HOWINA	56-day trial and failure of both preferred agents.	KINERET
				OLUMIANT
			*Cimzia will be allowed for clients that are pregnant or breast-feeding	ORENCIA REMICADE
				RINVOQ**
			**See Dermatology criteria for Atopic Dermatitis approval	RITUXAN SIMPONI
				SIMPONI XELIANZ/XR
VULSIONS		REOTYPIC SEIZURE EPISODES	*Nayzilam will be allowed for patients 12 years of age and older	
	diazepam gel NAYZILAM*			
	VALTOCO			
		CONVULSANTS	Preferred agents with clinical criteria will be limited to FDA approved indications related to	APTIOM
	carbamazepine divalproex	BANZEL (tablets only) clonazepam	seizures and epilepsy. Non-preferred agents require 30 day trial and failure of two preferred agents prior to approval.	BRIVIACT clobazam**
	FELBAMATE	EPIDIOLEX	agents prior to approval.	DIACOMIT**
	fosphenytoin lacosamide (tablets)	FYCOMPA gabapentin	For indications not related to seizures and epilepsy, please refer to the Additional Therapeutic	FINTEPLA** levetiracetam ER
	lamotrigine/XR	pregabalin*	Criteria chart at www.wymedicaid.org.	LIBERVANT
	levetiracetam	topiramate/ER sprinkle caps	to see the first of the second for the second s	OXTELLAR
	oxcarbazepine phenytoin		*Pregabalin will also be allowed for diagnoses of restless leg syndrome or anxiety **Please refer to the Additional Therapeutic Criteria chart at www.wymedicaid.org for specific	TROKENDI XR XCOPRI
	subvenite		requirements.	VIMPAT (tablets)
	valproate/valproic acid VIMPAT (suspension)			zonisamide oral susp.
	zonisamide			
HN'S	IMMUNO	MODULATORS HUMIRA	Client must have diagnosis of Crohn's prior to approval of the preferred agent. To receive a non-	CIMZIA** ENTYVIO*
		HOWINA	preferred agent, client must have a diagnosis of Crohn's and a 56-day trial and failure of the preferred agent.	REMICADE
				RINVOQ
			* Refer to Additional Therapeutics Clinical Criteria Chart for more info	SKYRIZI STELARA
			**Cimzia will be allowed for clients that are pregnant or breast-feeding	TYSABRI (additional criteria applies)
MATOLOGY	BENZOYL PEROXIDE	/CLINDAMYCIN COMBOs clindamycin/benzoyl peroxide 1-5%	Clients must be 12 to 20 years of age. Requires prior authorization for clients less than 12 years	ACANYA ONEXTON
		clindamycyin/benzoyl peroxide	of age. Acne combinations are limited to clients under the age of 21.	ONEXTON
		1.2-5% (Refrig)		
	AMNESTEEM	RETINOIN	Clients must be 12 to 20 years of age.	ABSORICA
	CLARAVIS			
	isotretinoin ZENATANE			
	CORTICOSTEROI	DS - STEP 1 AGENTS		
		N; O=OINTMENT; S=SOLUTION	Web and follows of two conferences are a set of the set	DANOSI
	alclometasone	POTENCY	Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days.	PANDEL TEXACORT 2.5% (S)
	desonide*			
	fluocinolone 0.01% hydrocortisone butyrate 0.1% (C)		*Cream, ointment, and lotion formulations of desonide are preferred.	
	hydrocortisone 1%, 2.5% (C,L,O)			
	MEDIUI	M POTENCY	Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days.	Clocortolone Pivalate
	betamethasone valerate desoximetasone 0.05%, 0.25% (C)			flurandrenol fluticasone 0.05% (L)
	fluocinolone 0.025%			hydrocortisone butyrate 0.1% (O)
	fluticasone 0.05% (C) mometasone			triamcinolone 0.05% (O)
	SYNALAR 0.025% (C, O)			
	triamcinolone 0.025%, 0.1%	DOTENOV	Web and follows of two conferences are the second	
	HIGH betamethasone dipropionate	POTENCY	Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days.	APEXICON 0.05% (C) amcinonide 0.1% (C,L,O)
	clobetasol/E 0.05% (C,G,O,S)			augmented betamethasone 0.05% (G,L,C
	diflorasone 0.05% (O) fluocinonide			clobetasol 0.05% (L) desoximetasone 0.05%, 0.25% (G,O)
	flucinonide flurandrenolide			desoximetasone 0.05%, 0.25% (G,O) diflorasone 0.05% (C)
	fluticasone 0.005% (O)			fluocinonide 0.1% (C)
	halobetasol TOPICORT 0.025% (C)			halcinonide 0.1% (C) HALOG 0.1% (O)
	triamcinolone 0.5%			11ALUG 0.1/0 (U)
	ULTRAVATE 0.05% (C,O)			
	IMMUNOMODU	LATORS - STEP 2 AGENTS ELIDEL	To receive a step 2 agent: Trial and failure of a preferred medium or high potency topical	pimecrolimus (brand preferred)
		tacrolimus	corticosteroid greater than or equal to a 21 day trial in the last 90 days.	
			Exceptions will be made for application to the face and for clients age 12 and under, a trial and	
			failure of a preferred low potency corticosteroid greater than or equal to a 21 day trial in the	
		•		1
		4 INHIBITOR - STEP 3 AGENT	last 90 days will be required. To receive a step 3 agent: Trial and failure of a preferred step 2 agent	EUCRISA

THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIES THIS LIST IS NOT ALL INCLUSIVE PLEASE CONTACT OPTIMINE WITH ANY QUESTIONS
ERMATOLOGY (continued)	ATOPIC	DERMATITIS ADBRY	Dupixent requires member be at least 6 months of age or older, Adbry requires member be at least 12 years of age or older. No high-potency steroid trial will be necessary. For clients with	CIBINQO** NEMLUVIO
(continueu)		DUPIXENT*	>20% BSA, no immunomodulator trial and failure will be necessary for preferred agent(s).	OPZELURA**
			Trial and failure of all criteria to receive a step 3 agent as defined above including	RINVOQ ZORYVE
			medium/high potency topical corticosteroid, preferred step 2 immunomodulator AND 56-day	
			trial and failure of a preferred biologic for Atopic Dermatitis in Step 3 will be required for	
			approval of the non-preferred agents.	
	PLAQUE P	SORIASIS (PP) ENBREL	Client must have diagnosis of PP prior to approval of a preferred agent. To receive a non- preferred agent, client must have a diagnosis of PP and a 56-day trial and failure of two of the	CIMZIA** COSENTYX
		HUMIRA	preferred agents.	ILUMYA
		OTEZLA SOTYKTU*	*Sotyktu requires diagnosis of moderate to severe PP and 56 day trial and failure of Humira.	REMICADE SILIQ
		TALTZ	**Cimzia will be allowed for clients that are pregnant or breast-feeding	SKYRIZI
		ZORYVE***	***Zoryve will be allowed for PP after a 21-day trial and failure of a high-potency corticosteroid OR a mild-potency corticosteroid if using in intertriginous areas.	STELARA TREMFYA
	SCABICIDES	/PEDICULICIDES	Trial and failure of a preferred agent in the last 12 months.	malathion lotion
	permethrin VANALICE			NATROBA spinosad (BRAND IS PREFERRED)
ABETES		ES AGENTS		metformin SR 24H osm (use preferred ager
		JANIDES		metformin SR 24H mod (use preferred age
	metformin/ER			
	GLUCOSIDASE	INHIBITORS	Trial and failure of metformin and a preferred agent greater than or equal to a 90 day supply in the last 12 months will be required before approval can be given for a non-preferred agent.	miglitol
		LITINIDES	Trial and failure of metformin and a preferred agent greater than or equal to a 90 day supply in	repaglinide
	nateglinide		the last 12 months will be required before approval can be given for a non-preferred agent.	- choPulling
		IDINEDIONES	Trial and failure of metformin and a preferred agent greater than or equal to a 90 day supply in	ACTOPLUS MET (use separate agents)
	pioglitazone		the last 12 months will be required before approval can be given for a non-preferred agent.	
	SULFO	NYLUREAS	Trial and failure of metformin and a preferred agent greater than or equal to a 90 day supply in	
	glimepiride/ER		the last 12 months will be required before approval can be given for a non-preferred agent.	
	glipizide/ER glyburide/ER			
		ASE 4 (DPP-4) INHIBITORS	Trial and failure of metformin greater than or equal to a 90 day supply in the last 12 months will	alogliptin
		JANUVIA ONGLYZA	be required before approval can be given for a preferred agent. A 90 day trial and failure of the preferred agent is required before approval can be given for a non-preferred agent.	GLYXAMBI (use separate preferred agents) QTERN (use separate preferred agents)
		TRADJENTA		STEGLUJAN (use separate preferred agents
	DPP-4 INHIBITOR	JANUMET/XR	Trial and failure of metformin greater than or equal to a 90 day supply in the last 12 months will be required before approval can be given for a preferred agent. A 90 day trial of failure of the	alogliptin/metformin alogliptin/pioglitazone (use separate prefer
		JENTADUETO	preferred agent is required before approval can be given for a non-preferred agent.	agents)
		KOMBIGLYZE/XR		JENTADUETO XR saxagliptin/metformin (BRAND IS PREFERR
				sitagliptin/metformin (BRAND IS PREFERRE
	INCRETIN MIMETICS (C	SLP-1 RECEPTOR AGONISTS)	Trial and failure of metformin greater than or equal to a 90 day supply in the last 12 months will	BYDUREON
		BYETTA RYBELSUS	be required before approval can be given for a preferred agent unless ASCVD or risk factors are present, in which case the trial of metformin is waived. A 90 day trial of failure of the preferred	liraglutide (use brand) MOUNJARO
		TRULICITY	agent is required before approval can be given for a non-preferred agent.	OZEMPIC*
		VICTOZA	Dosage Limits Apply:	SOLIQUA XULTOPHY (use separate preferred agents)
			Ozempic: 2mg/week Victoza: 1.8mg/day	
	50173	NHIBITORS	Trial and failure of metformin greater than or equal to a 90 day supply in the last 12 months will	GLYXAMBI (use separate preferred agents)
	30112	FARXIGA	be required before approval can be given for a preferred agent unless there is a diagnosis of	QTERN (use separate preferred agents)
		JARDIANCE SYNJARDY	ASCVD, CKD, or heart failure, in which case the trial of metformin will be waived. A 90 day trial	INVOKAMET INVOKANA
		XIGDUO XR	and failure of a preferred agent is required before approval can be given for a non-preferred agent.	SEGLUROMET (use separate preferred ager
			65-11-1 	STEGLATRO STEGLUJAN (use separate preferred agents)
				SYNJARDY XR (use separate preferred agen
	EAST-AC	TING INSULIN	Prior authorization will be required when using two different delivery forms of the same type of	TRIJARDY XR (use separate preferred agent. ADMELOG (use preferred agent)
	HUMALOG		insulin concurrently.	FIASP (use preferred agent)
	HUMALOG 75/25 HUMALOG JR.			insulin lispro (use preferred agents) LYUMJEV
	HUMALOG MIX			LIONDEV
	NOVOLOG MIX			
	LONG-AC LANTUS SOLOSTAR*	TING INSULIN	Prior authorization will be required when using two different delivery forms of the same type of insulin concurrently.	BASAGLAR (use preferred agent) Insulin Glargine (use preferred agent)
	LANTUS vial		/	Insulin Degludec
				SOLIQUA TOUJEO (use preferred agent)
				TRESIBA* (use preferred agent)
	DIABETIC ME	TERS/TEST STRIPS	Quantity limits apply:	XULTOPHY (use separate preferred agents) ALL OTHER METERS AND TEST STRIPS
	FREESTYLE (strips only)		Insulin Dependent Clients: 10 strips/day	
	FREESTYLE FREEDOM FREESTYLE FREEDOM LITE		Non-Insulin Dependent Clients: 4 strips/day	
	FREESTYLE INSULINX FREESTYLE PRECISION NEO B		Clients are limited to 1 meter/365 days	
	FREESTYLE SIDEKICK II			
	ONE TOUCH ULTRA II ONE TOUCH ULTRA MINI			
	ONE TOUCH ULTRA BLUE ONE TOUCH VERIO			
	ONE TOUCH VERIO FLEX			
	ONE TOUCH VERIO REFLECT ONE TOUCH VERIO IQ			
	PRECISION XTRA			
	EXTERNAL DIA OMNIPOD DASH	ABETIC DEVICES	4	OMNIPOD GO
	OMNIPOD 5			
	OMNIPOD G5 FSL 2 PLUS G6		Reference to a standard second second for the second second for the second second second second second second s	CHADDIAN
	CONTINUOUS BLOC	D GLUCOSE MONITORS DEXCOM G6	Prior authorization will be required to verify if the client is injecting insulin daily. Monitors will also be limited to the labeled age.	GUARDIAN MINIMED
		DEXCOM G7		
		FREESTYLE LIBRE FREESTYLE LIBRE 2		
		FREESTYLE LIBRE 3/PLUS		
	ACUTE HYPOG BAQSIMI	GLYCEMIA AGENTS		GVOKE (use preferred agent)
	ZEGALOGUE (autoinjector)	1		1

THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	red font indicates quantity/dose limits apply), and Wyoming Medicaid Pro CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIES THE LIST IS NOT ALL INCLUSIVE PLASE CONTACT OPIDIME WITH ANY QUESTION
IBROMYALGIA	FIBRO amitriptyline cyclobenzaprine duloxetine	MYALGIA gabapentin	Trial and failure of a preferred agent greater than or equal to six (6) weeks in the last 12 months is required prior to approval of a non-preferred agent Clients will not be allowed to take gabapentin and pregabalin concurrently	pregabalin SAVELLA tablets (savella titration pak will not be covered)
ASTROINTESTINAL		EVACUANTS	Crients win not de anowed to take gadapentin and pregadami concurrentiy	GAVILYTE H (use preferred agents) POLY-PREP (use preferred agents) SUTAB
		THIC CONSTIPATION AMITIZA LINZESS TRULANCE	Client must have a diagnosis of chronic idiopathic constipation to receive a preferred agent. To receive a non-preferred agent, the client must have a diagnosis of chronic idiopathic constipation and a 30-day trial and failure of a preferred agent within the last 12 months.	MOTEGRITY
	CREON ZENPEP	VE ENZYMES PERTZYE* DROME WITH CONSTIPATION	Prior authorization required. *Pertzye will be preferred for members diagnosed with cystic fibrosis. Client must have a diagnosis of Irritable Bowel Syndrome (IBS) with constipation.	VIOKACE
		AMITIZA LINZESS TRULANCE		
	MESA APRISO* LIALDA* mesalamine 400mg DR capsule mesalamine enema PENTASA	LAMINE	Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent.	mesalamine DR tab 800mg, 1.2g mesalamine ER cap 0.375gm mesalamine Sup 1000mg SFROWASA
	OPIOID-INDUCED	CONSTIPATION AGENTS AMITIZA	Client must have a diagnosis of opioid-induced constipation and a three (3) month trial and failure of a stool softener to receive the preferred agent. To receive the non-preferred agent, the client must have a diagnosis of opioid-induced constipation, a three (3) month trial and failure of a secretory agent, and a three (3) month trial and failure of the preferred agent. *Movantik will be approved for a diagnosis of cancer or for clients in hospice or palliative care.	MOVANTIK* RELISTOR SYMPROIC
		ED NAUSEA/VOMITING		BONJESTA
	DICLEGIS PROTON PU lansoprazole capsules/ODT omeprazole capsules/ODT pantoprazole	MP INHIBITORS	Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent.	amox/clarith/lanso pack DEXILANT dexlansoprazole esomeprazole 20.6mg capsules omeprazole 20.6mg capsules omeprazole fodium bicarbonate OMECLAMOX (use separate agents) PREVACID solutals (use preferred agents) rabeprazole TALICIA (use separate agents) VIMOVO (use separate agents)
	POTASSIUM COMPE	TITIVE ACID REDUCERS	Voquezna will require trial and failure of two proton pump inhibitors twice daily at max dose for 30 days	
оит	COL colchicine (tablets)	CHICINE	30 08/5	MITIGARE (use preferred agent)
	XANTHINE OXIDASE		Trial and failure of the preferred agent greater than or equal to a 60 day supply in the last 12	ULORIC*
		AND ORATI INHIBITORS		
MATOLOGY	allopurinol LOW MOLECULAR W	VEIGHT HEPARIN (LMWH)	Prior authorization will be required before approval can be given for a non-preferred agent. Prior authorization will be required for the 300mg/3ml strength.	FRAGMIN (use preferred agent)
MATOLOGY	allopurinol LOW MOLECULAR W enoxaparin		months will be required before approval can be given for a non-preferred agent.	
IMATOLOGY	allopurinol LOW MOLECULAR W enoxaparin DIRECT THROI SELECTIVE FAC ELIQUIS XARELTO (10/15/20mg, starter)	VEIGHT HEPARIN (LMWH) MBIN INHIBITOR PRADAXA TOR XA INHIBITOR XARELTO 2.5mg*	months will be required before approval can be given for a non-preferred agent. Prior authorization will be required for the 300mg/3ml strength. Client must have diagnosis of non-valvular atrial fibrillation and relative contraindication to warfarin for approval, treatment for deep vein thrombosis (DVT) or pulmonary embolism (PE), or for the reduction in the risk of recurrence of DVT and PE after initial therapy. *To receive Xarelto 2.5mg, client must have a diagnosis of chronic coronary artery disease or peripheral artery disease with the need to reduce risk of major cardiovascular events	FRAGMIN (use preferred agent)
MATOLOGY	allopurinol LOW MOLECULAR W enoxaparin DIRECT THROI SELECTIVE FAC ELIQUIS XARELTO (10/15/20mg, starter) CPTP DI	/EIGHT HEPARIN (LMWH) MBIN INHIBITOR PRADAXA TOR XA INHIBITOR	months will be required before approval can be given for a non-preferred agent. Prior authorization will be required for the 300mg/3ml strength. Client must have diagnosis of non-valvular atrial fibrillation and relative contraindication to warfarin for approval, treatment for deep vein thrombosis (DVT) or pulmonary embolism (PE), or for the reduction in the risk of recurrence of DVT and PE after initial therapy. "To receive Xarelto 2.5mg, client must have a diagnosis of chronic coronary artery disease or peripheral artery disease with the need to reduce risk of major cardiovascular events Client must have a diagnosis of acute coronary syndrome, history of myocardial infarction, or history of stroke and transient ischemic attack.	FRAGMIN (use preferred agent) enoxaparin 300MG/3ML ELIQUIS (starter pack)
EMATOLOGY	allopurinol LOW MOLECULAR W enoxaparin DIRECT THROJ ELIQUIS XARELTO (10/15/20mg, starter) CPTP DI PAR-1 AL ADVATE ADVATE ASTYLA ELOCTATE ESPEROCT HEMOFIL M HEMLIBRA JIVI KOATE FS/BIO-SET NOVOEIGHT NUWQ OBIZUR RECOMBINATE XONTIA SOLOFUSE	VEIGHT HEPARIN (LMWH) MBIN INHIBITOR PRADAXA ETOR XA INHIBITOR XARELTO 2.5mg* ERIVATIVES BRILINTA TRAGONIST ZONTIVITY	months will be required before approval can be given for a non-preferred agent. Prior authorization will be required for the 300mg/3ml strength. Client must have diagnosis of non-valvular atrial fibrillation and relative contraindication to warfarin for approval, treatment for deep vein thrombosis (DVT) or pulmonary embolism (PE), or for the reduction in the risk of recurrence of DVT and PE after initial therapy. *To receive Xarelto 2.5mg, client must have a diagnosis of chronic coronary artery disease or peripheral artery disease with the need to reduce risk of major cardiovascular events Client must have a diagnosis of acute coronary syndrome, history of myocardial infarction, or history of stroke and transient ischemic attack.	FRAGMIN (use preferred agent) enoxaparin 300MG/3ML ELIQUIS (starter pack)
EMATOLOGY	Allopurinol COMPARIANCE CULAR W A Constant of the second	IPEIGHT HEPARIN (LMWH) VEIGHT HEPARIN (LMWH) PRADAXA TOR XA INHIBITOR XARELTO 2.5mg* RIVATVES BRLINTA NTAGONIST ZONTIVITY ILIC FACTOR VIII ILIC FACTOR VIII	months will be required before approval can be given for a non-preferred agent. Prior authorization will be required for the 300mg/3ml strength. Client must have diagnosis of non-valvular atrial fibrillation and relative contraindication to warfarin for approval, treatment for deep vein thrombosis (DVT) or pulmonary embolism (PE), or for the reduction in the risk of recurrence of DVT and PE after initial therapy. *To receive Xarelto 2.5mg, client must have a diagnosis of chronic coronary artery disease or peripheral artery disease with the need to reduce risk of major cardiovascular events Client must have a diagnosis of acute coronary syndrome, history of myocardial infarction, or history of stroke and transient ischemic attack. Client must have diagnosis of reduction of thrombotic cardiovascular events with a history of myocardial infarction (MI) or with peripheral arterial disease (PAD). Must be used in	FRAGMIN (use preferred agent) enoxaparin 300MG/3ML ELIQUIS (starter pack) SAVAYSA (use preferred agent) ALTUVIIIO
EMATOLOGY	Allopurinol COMPARIANCE CULAR W A Constant of the second	VEIGHT HEPARIN (LMWH) MBIN INHIBITOR PRADAXA ETOR XA INHIBITOR XARELTO 2.5mg* ERIVATIVES BRILINTA TRAGONIST ZONTIVITY	months will be required before approval can be given for a non-preferred agent. Prior authorization will be required for the 300mg/3ml strength. Client must have diagnosis of non-valvular atrial fibrillation and relative contraindication to warfarin for approval, treatment for deep vein thrombosis (DVT) or pulmonary embolism (PE), or for the reduction in the risk of recurrence of DVT and PE after initial therapy. *To receive Xarelto 2.5mg, client must have a diagnosis of chronic coronary artery disease or peripheral artery disease with the need to reduce risk of major cardiovascular events Client must have a diagnosis of acute coronary syndrome, history of myocardial infarction, or history of stroke and transient ischemic attack. Client must have diagnosis of reduction of thrombotic cardiovascular events with a history of myocardial infarction (MI) or with peripheral arterial disease (PAD). Must be used in	FRAGMIN (use preferred agent) enoxaparin 300MG/3ML ELIQUIS (starter pack) SAVAYSA (use preferred agent) ALTUVIIIO
EMATOLOGY	allopurinol	IPEIGHT HEPARIN (LMWH) VEIGHT HEPARIN (LMWH) PRADAXA TOR XA INHIBITOR XARELTO 2.5mg* RIVATVES BRLINTA NTAGONIST ZONTIVITY ILIC FACTOR VIII ILIC FACTOR VIII	months will be required before approval can be given for a non-preferred agent. Prior authorization will be required for the 300mg/3ml strength. Client must have diagnosis of non-valvular atrial fibrillation and relative contraindication to warfarin for approval, treatment for deep vein thrombosis (DVT) or pulmonary embolism (PE), or for the reduction in the risk of recurrence of DVT and PE after initial therapy. *To receive Xarelto 2.5mg, client must have a diagnosis of chronic coronary artery disease or peripheral artery disease with the need to reduce risk of major cardiovascular events Client must have a diagnosis of acute coronary syndrome, history of myocardial infarction, or history of stroke and transient ischemic attack. Client must have diagnosis of reduction of thrombotic cardiovascular events with a history of myocardial infarction (MI) or with peripheral arterial disease (PAD). Must be used in	FRAGMIN (use preferred agent) enoxaparin 300MG/3ML ELIQUIS (starter pack) SAVAYSA (use preferred agent) ALTUVIIIO

THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLI THIS LIST IS NOT ALL INCLUSIVE PLASE CONTACT Optimities WITH ANY QUESTIONS
PATITIS C	DIRECT ACTIN		Limited to FDA approved indication. Prior authorization will be required prior to use of	EPCLUSA (use preferred agent)
		sofosbuvir/velpatasvir MAVYRET	preferred agents. Please submit PA requests on the Hepatitis C PA form available at www.wymedicaid.org.	HARVONI SOVALDI
			riedse submit i A requests on the reparties of A form available at www.wynieuteat.org.	VOSEVI** ZEPATIER
RADENITIS SUPPURATIVA	IMMUNO	MODULATORS	Humira will not be covered as a first line agent for the diagnosis for hidradenitis suppurativa.	COSENTYX
RMONES	GnBH AN	HUMIRA	*Please refer to the Additional Therapeutic Criteria chart at www.wymedicaid.org for specific	ORIAHNN
	MYFEMBREE		requirements.	
	ORILISSA GROWTH	HORMONE		HUMATROPE
		GENOTROPIN		NGENLA
		NORDITROPIN NUTROPIN AQ		SAIZEN SEROSTIM
		SKYTROFA		SOGROYA
	TESTOSTERON	IE TOPICAL GELS	Testosterone agents are only allowed for diagnosis of hypogonadism or insufficient testosterone	ZOMACTON ANDRODERM (use preferred agent)
		TESTIM GEL	production.	FORTESTA (use preferred agent)
			Other testosterone dosage form products will require a diagnosis of hypogonadism or	JATENZO (use preferred agent) TESTOPEL (use preferred agent)
			insufficient testosterone production (not outlined on PDL).	testosterone gel (use preferred agent)
				testosterone solution (use preferred agen XYOSTED (use preferred agent)
		HORMONES	Ermeza will be covered with confirmed diagnosis of dysphagia.	THYQUIDITY
	ARMOUR THYROID LEVOXYL	ERMEZA		TIROSINT
	levothyroxine (tablets)			
	LEVO-T liothyronine			
	SYNTHROID			
	UNITHROID	ACEPTIVES		alyacen 1-35, 7/7/7
	afirmelle		1	aranelle
	altavera amethia			BALCOLTRA balziva
	amethyst			briellyn
	apri ashlyna			drospir/ethinyl estradiol/levomefolate enpresse
	aubra/EQ			ethynodiol/ethinyl estradiol
	aurovela 1-20/FE 1-20, 1-35			FALESSA KIT
	aviane ayuna			fayosim FEMLYV
	azurette			kaitlib FE chew
	blisovi 1-20 FE, 1.5-30 FE bekyree			layolis FE chew levonest
	beyaz			levonorgest/ethinyl estradiol/LO (84-7)
	camila camrese/LO			levonorgest/ethinyl estradiol 0.15- MERZEE
	chateal/EQ			MINASTRIN FE chew*
	CHARLOTTE 24 FE chew cyred			NEXSTELLIS noreth/ethinyl estradiol/FE chew 0.8/25
	dasetta 1-35, 7/7/7			nortrel
	daysee deblitane			OPILL PHEXXI
	deso/ethinyl estradiol			philith
	drospir/ethinyl estradiol elinest			rivelsa QUARTETTE
	emzahh			SAFYRAL
	enskyce errin			SLYND TAYSOFY
	estarylla			TAYTULLA
	falmina finzala FE chew			tilia FE tri-legest FE
	gianvi			TRIVORA
	hailey FE 1/20, 1/35 heather			TWIRLA TYBLUME
	iclevia			tydemy
	incassia introvale			vyfemla wera
	isibloom			wymzya FE chew
	jaimiess jencycla			ZAFEMY
	jolessa			
	juleber junel 1-20/FE, 1.5-30/FE			
	kalliga			
	kariva kelnor			
	kurvelo			
	larin 1-20/FE, 1.5-30/FE Jeena			
	leena lessina			
	levora lo loestrin			
	loestrin FE			
	loryna LOSEASONIQUE*			
	low-ogestrel			
	lutera			
	marlissa melodetta			
	mibelas FE chew			
	microgestin 1-20/FE, 1.5-30/FE mili			
	mono-linyah			
	natazia NECON 0.5/35, 1/35, 1/50, 7/7/7,			
	nikki			
	nora-be noreth/ethinyl estradiol/FE chw			
	noreth/ethinyl estradiol 1-20/FE			
	norgest/ethinyl estradiol/LO			
	norethindrone norlynda			
	nylia	1		1

		PREFERRED AGENTS NON-PREFERRED AGENTS				
THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIES THE LIST IS NOT ALL INCLUSIVE PLEASE CONTACT OPLIMITIK WITH ANY QUESTIONS		
IORMONES; CONTRACEPTIVES	ocella pimtrea					
ontinued)	portia					
	previfem					
	reclipsen					
	safyral					
	SEASONIQUE*					
	setlakin					
	sharobel					
	simliya simpesse					
	sprintec					
	sronyx					
	syeda					
	tri-estarylla/LO					
	tri-femynor					
	tri-linyah					
	tri-marzia LO tri-mili/LO					
	tri-sprintec/LO					
	tri-nymyo					
	tri-vylibra					
	velivet					
	vestura					
	vienva					
	viorele					
	volnea					
	vylibra					
	yasmin-28 YAZ					
	YAZ zumandimine					
PERLIPIDEMIA		EQUESTRANT	Trial and failure of ALL preferred agents greater than or equal to six (6) months in the last 12	WELCHOL		
	cholestyramine/light		months will be required before approval can be given for a non- preferred agent.			
	colestipol					
	STATINS, L	OW POTENCY	Trial and failure of a preferred agent greater than or equal to a 90 day supply in the last 12	fluvastatin/ER		
	lovastatin		months will be required before approval can be given for a non-preferred agent.			
	pravastatin					
			If client's current medication therapy is contraindicated with the preferred statin(s) due to a			
			drug-drug interaction, a non-preferred agent may be obtained with a prior authorization.			
			Prior authorization will be required for clients under the age of 10.			
	CTATING I	IGH POTENCY	Trial and failure of a preferred agent greater than or equal to a 90 day supply in the last 12	EZALLOR		
	atorvastatin		months will be required before approval can be given for a non-preferred agent.	LIVALO		
	rosuvastatin		montris will be required before approval can be given for a non-preferred agent.	ZYPITAMAG		
	simvastatin		10 - Provide a construction of the state of			
			If client's current medication therapy is contraindicated with the preferred statin(s) due to a			
			drug-drug interaction, a non-preferred agent may be obtained with a prior authorization.			
			Prior authorization will be required for clients under the age of 10.			
		MBINATIONS	Trial and failure of a preferred agent greater than or equal to a 90 day supply in the last 12	ezetimibe/simvastatin (BRAND IS PREFERR		
	amlodipine/atorvastatin		months will be required before approval can be given for a non-preferred agent.			
	VYTORIN*					
			Prior authorization will be required for clients under the age of 10.			
	PCSK9-REL	ATED AGENTS	Client must have a diagnosis of homozygous familial hypercholesterolemia; have a diagnosis of	LEQVIO		
		PRALUENT	heterozygous familial hypercholesterolemia or atherosclerotic cardiovascular disease AND not			
		REPATHA	at goal with a maximum dose statin; or be intolerant to statin therapy. Approval for a non-			
	1		preferred agent requires trial and failure of a preferred agent.			
	1					
	TRIGLYCERIDE LC		Tetal and failure of a professed event assets they are smaller a 60 day small by 2 - 1 - 1 - 2	for of brie original		
	TRIGLYCERIDE LC	WERING AGEN IS	Trial and failure of a preferred agent greater than or equal to a 90 day supply in the last 12	fenofibric acid fenofibrate (43/50/120/130/150mg)		
			fenofibrate months will be required before approval can be given for a non-preferred agent.			
	fenofibrate		months will be required before approval can be given for a non-preferred agent.			
	fenofibrate gemfibrozil		months will be required before approval can be given for a non-preferred agent.	icosapent LIPOFEN		
	fenofibrate gemfibrozil omega-3-acid			icosapent LIPOFEN VASCEPA		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeza-3-acid ANGIOTENSIN RECE	PTOR BLOCKERS (ARBs)	months will be required before approval can be given for a non-preferred agent. Non-preferred ARBs will require a history of ALL preferred ARBs before approval	icosapent LIPOFEN VASCEPA candesartan		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeza-3-acid ANGIOTENSIN RECE EDARBI			icosapent LIPOFEN VASCEPA		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeea-3-acid EDARBI Irbesartan			icosapent LIPOFEN VASCEPA candesartan		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeza-3-acid EDARBI Irbesartan Iosartan Iosartan			icosapent LIPOFEN VASCEPA candesartan		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeea-3-acid EDARBI Irbesartan			icosapent LIPOFEN VASCEPA candesartan		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeza-3-acid EDARBI Irbesartan Iosartan olmesartan telmisartan valsartan	YTOR BLOCKERS (ARBs)		icosapent UPOFEN VASCEPA Candesartan eprosartan 600mg		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan olmesartan telmisartan valsartan Valsartan ARBs AN			icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan elemisartan valsartan valsartan EDARBYCLOR	YTOR BLOCKERS (ARBs)	Non-preferred ARBs will require a history of ALL preferred ARBs before approval	icosapent UPOFEN VASCEPA Candesartan eprosartan 600mg		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeza-3-acid EDARBI Irbesartan Iosartan olmesartan valsartan Valsartan EDARBYCLOR Irbesartan HCTZ	YTOR BLOCKERS (ARBs)	Non-preferred ARBs will require a history of ALL preferred ARBs before approval	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan telmisartan valsartan EDARBYCLOR irbesartan HCTZ losartan HCT	YTOR BLOCKERS (ARBs)	Non-preferred ARBs will require a history of ALL preferred ARBs before approval	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan elemisartan valsartan EDARBVCLOR irbesartan HCTZ losartan HCTZ olmesartan HCTZ	YTOR BLOCKERS (ARBs)	Non-preferred ARBs will require a history of ALL preferred ARBs before approval	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan elmisartan telmisartan telmisartan Valsartan EDARBYCLOR irbesartan HCTZ losartan HCTZ olmesartan HCTZ valsartan HCTZ	PTOR BLOCKERS (ARBs)	Non-preferred ARBs will require a history of ALL preferred ARBs before approval	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan elemisartan valsartan EDARBYCLOR irbesartan HCTZ losartan HCTZ valsartan HCTZ valsartan HCTZ	YTOR BLOCKERS (ARBs)	Non-preferred ARBs will require a history of ALL preferred ARBs before approval	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan olmesartan telmisartan valsartan EDARBYCLOR irbesartan HCTZ losartan HCTZ valsartan HCTZ valsartan HCTZ valsartan HCTZ valsartan HCTZ clonidine	PTOR BLOCKERS (ARBs)	Non-preferred ARBs will require a history of ALL preferred ARBs before approval	icosapent LIPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan elmisartan telmisartan valsartan EDARBYCLOR irbesartan HCTZ losartan HCTZ olmesartan HCTZ valsartan HCTZ clonidine clonidine TD patches	PTOR BLOCKERS (ARBs)	Non-preferred ARBs will require a history of ALL preferred ARBs before approval Non-preferred ARB/diuretic combinations will require a history of ALL preferred	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ telmisartan HCTZ		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan elmisartan telmisartan valsartan EDARBYCLOR irbesartan HCTZ losartan HCTZ olmesartan HCTZ valsartan HCTZ clonidine clonidine TD patches	PTOR BLOCKERS (ARBs) D DIURETICS BLOCKERS	Non-preferred ARBs will require a history of ALL preferred ARBs before approval Non-preferred ARB/diuretic combinations will require a history of ALL preferred Client must be greater than one year of age and have a diagnosis of Congestive Heart Failure	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ telmisartan HCTZ telmisartan HCTZ		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan elmisartan telmisartan valsartan EDARBYCLOR irbesartan HCTZ losartan HCTZ olmesartan HCTZ valsartan HCTZ clonidine clonidine TD patches	PTOR BLOCKERS (ARBs) D DIURETICS BLOCKERS ON PRODUCTS	Non-preferred ARBs will require a history of ALL preferred ARBs before approval Non-preferred ARB/diuretic combinations will require a history of ALL preferred Client must be greater than one year of age and have a diagnosis of Congestive Heart Failure (CHF) NYHA Class II-IV. Angiotensin converting enzyme (ACE) inhibitors and angiotensin receptor	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ telmisartan HCTZ		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan elmisartan telmisartan valsartan EDARBYCLOR irbesartan HCTZ losartan HCTZ olmesartan HCTZ valsartan HCTZ clonidine clonidine TD patches	PTOR BLOCKERS (ARBs) D DIURETICS BLOCKERS ON PRODUCTS	Non-preferred ARBs will require a history of ALL preferred ARBs before approval Non-preferred ARB/diuretic combinations will require a history of ALL preferred Client must be greater than one year of age and have a diagnosis of Congestive Heart Failure	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ telmisartan HCTZ telmisartan HCTZ		
	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan olmesartan telmisartan EDARBYCLOR irbesartan HCTZ losartan HCTZ valsartan HCTZ valsartan HCTZ valsartan HCTZ clonidine clonidine TD patches COMBINAT	PTOR BLOCKERS (ARBs) D DIURETICS BLOCKERS ON PRODUCTS ENTRESTO	Non-preferred ARBs will require a history of ALL preferred ARBs before approval Non-preferred ARB/diuretic combinations will require a history of ALL preferred Client must be greater than one year of age and have a diagnosis of Congestive Heart Failure (CHF) NYHA Class II-IV. Angiotensin converting enzyme (ACE) inhibitors and angiotensin receptor blockers (ARBs) will not be allowed in combination with Entresto.	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ telmisartan HCTZ telmisartan HCTZ ENTRESTO SPRINKLES VERQUVO		
	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan olmesartan telmisartan EDARBYCLOR irbesartan HCTZ losartan HCTZ valsartan HCTZ valsartan HCTZ valsartan HCTZ clonidine clonidine TD patches COMBINAT	PTOR BLOCKERS (ARBs) D DIURETICS BLOCKERS ON PRODUCTS	Non-preferred ARBs will require a history of ALL preferred ARBs before approval Non-preferred ARB/diuretic combinations will require a history of ALL preferred Client must be greater than one year of age and have a diagnosis of Congestive Heart Failure (CHF) NYHA Class II-IV. Angiotensin converting enzyme (ACE) inhibitors and angiotensin receptor blockers (ARBs) will not be allowed in combination with Entresto. Please refer to the Additional Therapeutic Criteria Chart located at	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ telmisartan HCTZ telmisartan HCTZ		
	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan olmesartan telmisartan Valsartan EDARBVCLOR irbesartan HCTZ losartan HCTZ valsartan HCTZ valsartan HCTZ valsartan HCTZ conidine clonidine TD patches COMBINAT	PTOR BLOCKERS (ARBs) D DIURETICS BLOCKERS ON PRODUCTS ENTRESTO	Non-preferred ARBs will require a history of ALL preferred ARBs before approval Non-preferred ARB/diuretic combinations will require a history of ALL preferred Client must be greater than one year of age and have a diagnosis of Congestive Heart Failure (CHF) NYHA Class II-IV. Angiotensin converting enzyme (ACE) inhibitors and angiotensin receptor blockers (ARBs) will not be allowed in combination with Entresto.	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ telmisartan HCTZ telmisartan HCTZ		
	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan telmisartan valsartan EDARBYCLOR irbesartan HCT losartan HCT olmesartan HCTZ valsartan HCTZ clonidine clonidine TD patches COMBINAT ciprofloxacin levofloxacin	PTOR BLOCKERS (ARBs) D DIURETICS BLOCKERS ON PRODUCTS ENTRESTO OLONES	Non-preferred ARBs will require a history of ALL preferred ARBs before approval Non-preferred ARB/diuretic combinations will require a history of ALL preferred Client must be greater than one year of age and have a diagnosis of Congestive Heart Failure (CHF) NYHA Class II-IV. Angiotensin converting enzyme (ACE) inhibitors and angiotensin receptor blockers (ARBs) will not be allowed in combination with Entresto. Please refer to the Additional Therapeutic Criteria Chart located at	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ telmisartan HCTZ telmisartan HCTZ ENTRESTO SPRINKLES VERQUVO moxifloxacin (use preferred agents)		
	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan telmisartan valsartan EDARBYCLOR irbesartan HCT losartan HCT olmesartan HCTZ valsartan HCTZ clonidine clonidine TD patches COMBINAT ciprofloxacin levofloxacin	PTOR BLOCKERS (ARBs) D DIURETICS BLOCKERS ON PRODUCTS ENTRESTO	Non-preferred ARBs will require a history of ALL preferred ARBs before approval Non-preferred ARB/diuretic combinations will require a history of ALL preferred Client must be greater than one year of age and have a diagnosis of Congestive Heart Failure (CHF) NYHA Class II-IV. Angiotensin converting enzyme (ACE) inhibitors and angiotensin receptor blockers (ARBs) will not be allowed in combination with Entresto. Please refer to the Additional Therapeutic Criteria Chart located at	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ telmisartan HCTZ telmisartan HCTZ ENTRESTO SPRINKLES VERQUVO		
	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan telmisartan valsartan EDARBYCLOR irbesartan HCT losartan HCT olmesartan HCTZ valsartan HCTZ clonidine clonidine TD patches COMBINAT ciprofloxacin levofloxacin	PTOR BLOCKERS (ARBs) D DIURETICS BLOCKERS ON PRODUCTS ENTRESTO OLONES	Non-preferred ARBs will require a history of ALL preferred ARBs before approval Non-preferred ARB/diuretic combinations will require a history of ALL preferred Client must be greater than one year of age and have a diagnosis of Congestive Heart Failure (CHF) NYHA Class II-IV. Angiotensin converting enzyme (ACE) inhibitors and angiotensin receptor blockers (ARBs) will not be allowed in combination with Entresto. Please refer to the Additional Therapeutic Criteria Chart located at	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ telmisartan HCTZ telmisartan HCTZ ENTRESTO SPRINKLES VERQUVO moxifloxacin (use preferred agents)		
	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan telmisartan valsartan EDARBYCLOR irbesartan HCTZ losartan HCTZ olmesartan HCTZ valsartan HCTZ clonidine TD patches comBINAT clorofloxacin levofloxacin ofloxacin oxycycline MINC	PTOR BLOCKERS (ARBs) D DIURETICS BLOCKERS ON PRODUCTS ENTRESTO OLONES	Non-preferred ARBs will require a history of ALL preferred ARBs before approval Non-preferred ARB/diuretic combinations will require a history of ALL preferred Client must be greater than one year of age and have a diagnosis of Congestive Heart Failure (CHF) NYHA Class II-IV. Angiotensin converting enzyme (ACE) inhibitors and angiotensin receptor blockers (ARBs) will not be allowed in combination with Entresto. Please refer to the Additional Therapeutic Criteria Chart located at	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ telmisartan HCTZ telmisartan HCTZ ENTRESTO SPRINKLES VERQUVO moxifloxacin (use preferred agents)		
	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan olmesartan telmisartan valsartan EDARBVCLOR irbesartan HCTZ losartan HCTZ valsartan HCTZ valsartan HCTZ valsartan HCTZ clonidine TD patches COMBINAT	PTOR BLOCKERS (ARBs) DIURETICS BLOCKERS ENTRESTO DLONES CYCLINE	Non-preferred ARBs will require a history of ALL preferred ARBs before approval Non-preferred ARB/diuretic combinations will require a history of ALL preferred Client must be greater than one year of age and have a diagnosis of Congestive Heart Failure (CHF) NYHA Class II-IV. Angiotensin converting enzyme (ACE) inhibitors and angiotensin receptor blockers (ARBs) will not be allowed in combination with Entresto. Please refer to the Additional Therapeutic Criteria Chart located at	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ telmisartan HCTZ telmisartan HCTZ ENTRESTO SPRINKLES VERQUVO moxifloxacin (use preferred agents) DORYX (use preferred agent) minocycline 65mg and 115mg ER (use preferred agent)		
	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan olmesartan telmisartan valsartan EDARBVCLOR irbesartan HCTZ losartan HCTZ olmesartan HCTZ valsartan HCTZ clonidine TD patches clonidine TD patc	PTOR BLOCKERS (ARBs) D DIURETICS BLOCKERS ON PRODUCTS ENTRESTO OLONES CYCLINE CYCLINE	Non-preferred ARBs will require a history of ALL preferred ARBs before approval Non-preferred ARB/diuretic combinations will require a history of ALL preferred Client must be greater than one year of age and have a diagnosis of Congestive Heart Failure (CHF) NYHA Class II-IV. Angiotensin converting enzyme (ACE) inhibitors and angiotensin receptor blockers (ARBs) will not be allowed in combination with Entresto. Please refer to the Additional Therapeutic Criteria Chart located at	icosapent LIPOFEN VASCEPA Candesartan eprosartan 600mg Candesartan HCTZ telmisartan HCTZ telmisartan HCTZ ENTRESTO SPRINKLES VERQUVO moxifloxacin (use preferred agent) DORYX (use preferred agent) minocycline 65mg and 115mg ER (use preferred agent)		
PERTENSION/ CARDIOLOGY	fenofibrate gemfibrozil omeza-3-acid EDARBI irbesartan losartan elmisartan valsartan EDARBVCLOR irbesartan HCTZ losartan HCTZ olmesartan HCTZ valsartan HCTZ clonidine TD patches clonidine TD patches	PTOR BLOCKERS (ARBs) DIURETICS BLOCKERS ENTRESTO DLONES CYCLINE	Non-preferred ARBs will require a history of ALL preferred ARBs before approval Non-preferred ARB/diuretic combinations will require a history of ALL preferred Client must be greater than one year of age and have a diagnosis of Congestive Heart Failure (CHF) NYHA Class II-IV. Angiotensin converting enzyme (ACE) inhibitors and angiotensin receptor blockers (ARBs) will not be allowed in combination with Entresto. Please refer to the Additional Therapeutic Criteria Chart located at	icosapent UPOFEN VASCEPA candesartan eprosartan 600mg candesartan HCTZ telmisartan HCTZ telmisartan HCTZ ENTRESTO SPRINKLES VERQUVO moxifloxacin (use preferred agents) DORYX (use preferred agent) minocycline 65mg and 115mg ER (use preferred agent)		

Please refer to the Addi	tional Therapeutic Criteria (Chart, Dosage Limitation List (red font indicates quantity/dose limits apply), and Wyoming Medicaid Pro	vider Manual for additional criteria
THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIES THIS UST IS NOT ALL INCLUSIVE PLASE CONTACT Optimulie WITH ANY QUESTIONS
NFECTIOUS DISEASE (continued)	ANTI-RI BIRTARVY CIMDUO DELSTRIGO DOVATO EVOTAZ GENVOYA ODEFSEY PIFELTRO PREZCOBIX ritonavir tablets SYMFI/LO TRIUMEQ TROGARZO	HOVIRALS CABENUVA* DESCOV* TRUVADA*	*Please refer to the Additional Therapeutic Criteria chart at www.wymedicaid.org for specific requirements. **Rukobia approval requires documentation of multi-drug resistance defined as failure of two medications from different classes.	JULUCA NORVIR RUKOBIA** STRIBILD (use separate agents) SUNLENCA SYMTUZA (use separate preferred agents)
INFLAMMATION	N Celecoxib diclofenac tablets etodolac FLECTOR* flurbiprofen indomethacin ketoprofen ketoprofen ketorolac meclofenamate meloxicam nabumetone naproxen oxaprozin piroxicam sulindac	SAIDs	Trial and failure of two (2) preferred agents each greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non- preferred agent. Dosing and quantity limits apply for ketorolac (limit Sdays/34 days; max dose 40mg/day for oral tablets).	CALDOLOR (use preferred agent) diclofenac 1.3% patch (BRAND IS PREFERRED diclofenac 3.5% gel fenoprofen mefenamic acid NEOPROFEN (use preferred agent)
		TICOSTEROIDS		CELESTONE (use preferred agent) EMFLAZA
INSOMNIA	NON-BEN BELSOMRA eszopicione zalepion zolnidem zolpidem ER	COLIAZEPINES	Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. Prior Authorization will be required for clients under the age of 18 *Quviviq requires trial and failure of two preferred agents with different mechanisms of action **Rozerem is non-preferred without a history of substance abuse Prior authorization will be required when a client is taking more than one insomnia agent concurrently. Dosage limits apply: zalepion: 30mg/day zolpidem: 15mg/day	EDLUAR (additional criteria applies) DAYVIGO QUVYIQ* ROZEREM** zolpidem sublingual (additional criteria applies)
MENTAL HEALTH		ER'S AGENTS donepezil/ODT galantamine/ER memantine tablets/solution	Client must have a diagnosis of dementia.	donepezil 23mg (use preferred agent) memantine ER NAMZARIC (use separate agents) rivastigmine capsules/patches
	NORADRENERGIC/SPEC mirtazapine tablets NOREPINEPHRINE/DOPAI bupropion ER/SR/XL SELECTIVE SEROTONIN citalopram escitalopram fluoxetine capsules paroxetine IR/CR settraline	PRESSANTS FIC SEROTONERGICS (NaSS) AINE REUPTAKE INHIBITORS (NDRI) REUPTAKE INHIBITORS (SSRI)	Trial and failure of two (2) prefered agents greater than or equal to six (6) weeks <u>WITHIN THE</u> LAST 2 YEARS will be required before approval can be given for a non-preferred agent. One of the trials of preferred agents must be in the same class (NaSS, NDRI, SSRI, or SNRI) as the requested non-preferred agent. Trazodone, buspirone, fluvoxamine, MAO inhibitors, TCA's, bupropion IR, and venlafaxine IR do not require prior authorization but will not count towards meeting preferred therapy requirements. Clients will not be allowed to be on more than one antidepressant, including fluvoxamine, bupropion IR, and venlafaxine IR, at one time with the exception of mirtazapine or bupropion with a SSRI or SNRI. ***Trintellix requires trial and failure of two preferred agents in anv class Clients five (5) years of age and younger will require prior authorization before approval. Dosage limits apply: bupropion EK/SR/XL: 450mg/day citalopram < 60 years of age: 60mg/day divextien < 18 years of age: 90mg/day fluvoxetine < 18 years of age: 90mg/day mirtazapine: 67.5mg/day paroxetine IR/CR < 18 years of age: 90mg/day paroxetine IR/CR < 18 years of age: 90mg/day paroxetine IR/CR < 18 years of age: 90mg/day venlafaxine EK = 112.5mg/day sertraline: 300mg/day venlafaxine EX 337.5mg/day	NaSS mirtazapine rapid discolve tablets (use preferred agent) NDRI APLENZIN AUVELITY FORFIVO XL* Citalopram capsules fluoxetine tablets VIBRYD FETZIMA venlafaxine ER tablets (use preferred agent) OTHER TRINTELLIX***

THERAPEUTIC CLASS		PREFERRED AGENTS REQUIRING CLINICAL		NON-PREFERRED AGENTS
		CRITERIA	CLINICAL CRITERIA	GENERIC MANDATORY POLICY APPLIE THIS LIST IS NOT ALL INCLUSIVE PLEASE CONTACT Optumike WITH ANY QUESTIONS
initial de la companya de la company	ABILIFY MAINTENA	NTIPSYCHOTICS	*Quetiapine doses less than 100mg will require prior authorization <u>without</u> a diagnosis of mood disorder or major depressive disorder. For titration doses, contact the OptumRx Pharmacy Help	
	ABILIFY ASIMTUFII		Desk for an override.	GEODON 20MG INJ
	aripiprazole tab/solution/ODT ARISTADA		**Clients nine (9) years of age and younger will require a prior authorization to receive approval	LYBALVI (additional criteria applies) NUPLAZID
	asenapine FANAPT**		of lurasidone and asenapine. Clients eighteen (18) years of age and younger will require a prior	olanzapine 10mg Inj SAPHRIS (use preferred agent)
	paliperidone INVEGA HAFYERA		authorization to receive approval of Fanapt.	SECUADO REXULTI***
INVEGA SUSTENNA INVEGA TRINZA	INVEGA SUSTENNA			ZYPREXA RELPREVV
	INVEGA TRINZA lurasidone**		***Rexulti approval for MDD treatment requires concurrent antidepressant therapy as well as a trial and failure of aripiprazole or other preferred atypical antipsychotic indicated for adjunct	
	olanzapine		MDD treatment.	
PERSERIS quetiapine* quetiapine ER RISPERDAL CONSTA risperidone		Trial and failure of two (2) preferred agents greater than or equal to 30 days in the last 12		
		months will be required before approval can be given for a non-preferred agent unless		
	risperidone		otherwise specified.	
	RYKINDO UZEDY			
	VRAYLAR ziprasidone		Prior authorization will be required for any client five (5) years of age or younger, or for any	
	ziprasidone		client taking both an injectable and oral dosage form of the same medication concurrently.	
			Dosage limits apply:	
			aripiprazole <13 years of age: 15mg/day; <a>>13 years of age: 30mg/day asenapine: 20mg/day	
			ABILIFY MAINTENA: 400mg per 26 days	
			ARISTADA 441/662/882mg: 1 injection per 28 days; 1064mg: 1 injection per 56 days ARISTADA INITIO: 1 injection per 365 days	
			FANAPT: 24mg/day	
			INVEGA HAFYERA: 1 injection per 6 months INVEGA SUSTENNA: 1 injection per 28 days	
			INVEGA TRINZA: 1 injection per 84 days	
			lurasidone 10-17 years of age: 80mg/day; >17 years of age: 160mg/day olanzapine <13 years of age: 10mg/day; ≥13 years of age: 20mg/day	
			paliperidone: 12mg/day	
			PERSERIS: 1 injection per 28 days	
			<pre>quetiapine <13 years of age: 400mg/day; 13-17 years of age: 600mg/day; >17 years of age: 800mg/day</pre>	
			risperidone <10 years of age: 3mg/day; 10-17 years of age: 6mg/day; >17 years of age:	
			16mg/day RISPERDAL CONSTA: 2 injections per 28 days	
			ziprasidone ≤17 years of age: 120mg/day; >17 years of age: 200mg/day	
	SPECIAL ATYPICA	L ANTIPSYCHOTICS	Dosage limits apply: 900mg/day	VERSACLOZ Suspension (use preferred ag
	clozapine/ODT	ETAMINES	Clients over the age of 17 must have a diagnosis for ADD, ADHD (see ADD/ADHD criteria below),	AMPHETAMINES
		AMPHETAMINES	narcolepsy, obstructive sleep apnea, shift work sleep disturbance, MS fatigue (see MS Fatigue	ADZENYS XR ODT
		ADDERALL XR* amphetamine salts combo XR	criteria below), or refractory depression (see refractory depression criteria below).	DYANAVEL XR EVEKEO/ODT
		dextroamphetamine CR caps VYVANSE CAPSULES**		MYDAYIS PROCENTRA
	IMMEDIATE RELE	ASE AMPHETAMINES	For clients over the age of 17, they must meet the DSM-5 criteria for diagnosis of ADHD. These	VYVANSE CHEWABLES
		amphetamine salts combo dextroamphetamine tablets	 criteria include: Five or more symptoms of inattention, present for at least 6 months, 	ZENZEDI 2.5 AND 7.5MG TABLETS
		PHENIDATES ETHYLPHENIDATES	inappropriate for developmental level.	METHYLPHENIDATES APTENSIO XR
	Long Acting Mi	CONCERTA*	OR Five or more symptoms of hyperactivity and impulsivity, present for at least 6 months, to an 	AZSTARYS
		dexmethylphenidate ER methylphenidate ER tablets	extent that is disruptive and inappropriate for developmental level.	COTEMPLA XR DAYTRANA
			AND • Sumptoms must be present in two or more settings (home, school or work):	FOCALIN XR JORNAY PM
			 Symptoms must be present in two or more settings (home, school or work); There must be clear evidence that the symptoms interfere or reduce the quality of social, 	methylphenidate ER osmotic release
	IMMEDIATE RELEAS	E METHYLPHENIDATES dexmethylphenidate	school or work functioning; and • The symptoms must not be better explained by another mental disorder.	(BRAND IS PREFERRED) methylphenidate ER/CR/SR capsules
		methylphenidate chewables	 The symptoms must not be better explained by another mental disorder. 	(METADATE CD/RITALIN LA, APTENSIO XF
		methylphenidate solution methylphenidate tablets		RELEXXII QUILLICHEW ER
			Diagnosis of MS fatigue will require a fatigue severity scale score of 5.0, a 60-day trial of	QUILLIVANT
			amantadine and discontinuation of medications that may contribute to drowsiness and fatigue. Diagnosis of refractory depression will require a 6-week trial and failure of an antidepressant	
			(monotherapy) and continued concomitant use of an antidepressant with the stimulant. **Vyvanse will be approved for the diagnosis of binge-eating disorder for clients 18 years of age	
			and older. Authorizations will be approved for 12 weeks. After initiation a one-year prior authorization of Vyvanse for this diagnosis will require additional documentation prior to approved	
			approval. Claims will require Prior Authorization if client is under the age of 4, or has a history of the	
			following: glaucoma, cardiac arrhythmias, arteriosclerosis, untreated hypertension, untreated	
			hyperthyroidism, substance abuse, or current MAO inhibitor use. Trial and failure of two (2) preferred agents (each from a different class: methylphenidate and	
			amphetamine) greater than or equal to a 30 day supply in the last 12 months will be required	
			before approval can be given for a non-preferred agent. Two or more long-acting agents will not be allowed concurrently.	
			Dosage limits apply:	
			amphetamine salts combo XR: 60mg/day amphetamine salts combo: 60mg/day	
			amphetamine salts combo (narcolepsy): 90mg/day	
			DAYTRANA: 45mg/9 hour patch/day	
			DAYTRANA: 45mg/9 hour patch/day dextroamphetamine: 90mg/day dextroamphetamine CR: 90mg/day dexmethylphenidate: 30mg/day	
			DAYTRANA: 45mg/9 hour patch/day dextroamphetamine: 90mg/day dextroamphetamine CR: 90mg/day dexmethylphenidate: 30mg/day FOCALIN XR < 13 years of age: 45mg/day	
			DAYTRANA: 45mg/9 hour patch/day dextroamphetamine: 90mg/day dextroamphetamine CR: 90mg/day dexmethylphenidate: 30mg/day	

THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA		NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIE: THIS LIST IS NOT ALL INCLUSIVE PLASE CONTACT OPLICING WITH ANY OUTSTICKS
ENTAL HEALTH ontinued	clonidine, clonidine ER guanfacine, guanfacine ER	A-ADRENERGIC AGONIST	Client must must have a diagnosis of ADD or ADHD. Prior authorization will be required for clients under the age of 4.	ONYDA XR
	SELECTIVE NOREPINEP	HRINE REUPTAKE INHIBITOR atomoxetine	Atomoxetine: Clients must have a diagnosis for ADD, ADHD, narcolepsy, obstructive sleep apnea, shift work sleep disturbance, or refractory depression (see refractory depression criteria below).	QELBREE
			Diagnosis of refractory depression will require a 6-week trial and failure of an antidepressant (monotherapy) and continued concomitant use of an antidepressant with the stimulant.	
			Prior Authorization required for clients under the age of 4.	
			Claims will require Prior Authorization if clients have a history of the following: glaucoma, cardiac arrhythmias, arteriosclerosis, untreated hypertension, untreated hyperthyroidism, substance abuse, or current MAO inhibitor use.	
			Qelbree: Clients must have a 30-day trial and failure of a preferred non-stimulant. Approval will be limited to a max dose of 400mg in clients 6-17 years of age, and 600mg for clients 18 years of age or older. Dosage limits apply: atomoxetine: 100mg/day	
GRAINE		PROPHYLAXIS	Trial and failure of both an anticonvulsant and a beta blocker (Step 1 agents) greater than or	NURTEC
	STEP : beta blockers	1 AGENTS divalproex topiramate	equal to three (3) months will be required before approval can be given for the step 2 agents. Concurrent use of Botox will not be approved. Nurter will be limited to 16 tabs/30 days.	
	STEP	2 AGENTS AIMOVIG*	*Starting dose will be limited to 70mg **Approval for non-preferred agents requires trial and failure of a preferred agent along with	QULIPTA**
		AIMOVIG AJOVY EMGALITY AINE TREATMENT	TApprovanion non-preferred agents requires that and ratio to a preferred agent along with the trial and failures described with Step 1 Agents' criteria above.	
		AINE TREATMENT 1 AGENTS	Trial and failure of two preferred agents will be required for approval of a non- preferred agent.	almotriptan ELYXYB
	naratriptan RELPAX* sumatriptan rizatriptan		Rizatriptan will be limited to clients 6 years of age or older Quantity limits apply: naratriptan 1mg: 25 tabs/34 days	Sumatriptan-Naproxen Sodium TOSYMRA (use preferred agent) ZEMBRACE (use preferred agent) ZAVZPRET zolmitriptan
			naratriptan 2.5mg: 10 tabs/34 days RELPAX 20mg: 20 tabs/34 days RELPAX 40mg: 14 tabs/34 days rizatriptan 5mg: 27 doses/34 days rizatriptan 10mg: 14 doses/34 days	20111111
			sumatriptan usus: 2 vials/34 days sumatriptan nasal 20mg: 6 bottles/34 days; 5mg: 12 bottles/34 days sumatriptan 25mg: 41 tabs/34 days sumatriptan 50mg: 20 tabs/34 days	
	STED	2 AGENTS	sumatriptan 100mg: 10 tabs/34 days Trial and failure of two triptan agents required for Step 2 Agent approval	REYVOW
	STEP.	NURTEC UBRELVY	Trial and failure of two original agents required for step 2 Agent approval Trial and failure of two originered triptan agents AND Nurtec OR Ubrelvy will be required for approval of a non-preferred agent. Quantity limits apply: NURTEC 75mg: limited to 8 tabs/30 days REVVOW: 200mg/day or 1 tab/day, 4 tab/30 days	RETVOW
OVEMENT DISORDERS		INHIBITORS	Quantity limits apply:	
	AUSTEDO/XR* INGREZZA* TETRABENAZINE		AUSTEDO: limited to 4 tabs/dav INGREZZA: limited to 4 tabs/day	
JLTIPLE SCLEROSIS		AGENTS GILENYA KESIMPTA	*Please refer to the Additional Therappeutic Criteria chart at www.wymedicaid.org Gilenya, Kesimpta, Lemtrada, Ocrevus, and Tysabri will be approved for highly active disease, please refer to the ATCC for additional information.	AUBAGIO BAFIERTAM BRIUMVI
	COPAXONE 20MG/ML* dimethyl fumarate REBIF teriflunomide	LEMTRADA OCREVUS TYSABRI	Trial and failure of two preferred agents for at least 56 days (each from a separate class) will be required before approval can be given for a non-preferred agent.	EXTAVIA glatiramer (BRAND IS PREFERRED) GLATOPA (use preferred agent) MAVENCLAD
	VUMERITY		For Mavenclad, in addition to the above criteria, approval will be granted on a case- by-case basis.	MAYZENT PLEGRIDY PONVORY TECFIDERA ZEPOSIA
RCOLEPSY	STIN	IULANTS modafinil NUVIGIL*	Modafinil and Nuvigii: Client must be > 16 years of age. Client must have a diagnosis of narcolepsy, obstructive sleep apnea, shift work sleep disorder, Multiple Sclerosis (MS) Fatigue, or ADD/ADH with a concurrent diagnosis of substance abuse.	
	NON-ST	TIMULANTS	Diagnosis of MS will require a fatigue severity scale score of 5.0, a 60 day trial of amantadine AND discontinuation of medications that may contribute to drowsiness or fatigue.	SUNOSI WAKIX XYREM
			Clients will not be allowed to take two or more agents in this class concurrently	
UROPATHIC PAIN		APENTIN gabapentin pregabalin	Clients will not be allowed to take gabapentin and pregabalin concurrently Prior authorizations for perioperative pain will be approved for gabapentin OR pregabalin for less than or equal to 14 day supplies	
	TOPICAI Lidocaine Patches	LIDOCAINE		ZTLIDO
		NAL AGENTS	Trial and failure of a tricyclic antidepressant greater than or equal to a 12 week supply AND trial and failure of gabapentin at a dose of 3600mg per day OR pregabalin for greater than or equal to a 12 week supply in the last 12 months will be required before approval can be given for a non-preferred agent.	carbamazepine imipramine (capsules) oxcarbazepine valproic acid
STRUCTIVE SLEEP APNEA		Agonists ZEPBOUND	Client must have diagnosis of moderate to severe obstructive sleep apnea. Will be approved for obese adults with an AHI (Apnea-Hypopnea Index) of greater than 15. Prior authorization will be required again at 6 months to show at least 5% weight loss. Prior authorization will be required again at 12 months to demonstrate improvement in obstructive sleep apnea.	

THERAPEUTIC CLASS			t (red font indicates quantity/dose limits apply), and Wyoming Medicaid Provider Manual for additi	
	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIES THIS LIST IS NOT ALL INCLUSIVE PLEASE CONTACT Optimits WITH ANY QUESTIONS
OPHTHALMICS		I-ALLERGICS	Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will	ALOCRIL
	ALREX azelastine		be required before approval can be given for a non-preferred agent.	ALOMIDE bepotastine
	BEPREVE*		Alomide and Alocril will be approved for pregnancy. Alomide will be approved for children under the age of 3.	epinastine
	cromolyn 0.4%	l	-	ZERVIATE
	OPANTIBIOT ciprofloxacin	ICS- QUINOLONES	Trial and failure of a preferred agent greater than or equal to 5 days in the last 12 months will	gatifloxacin ZYMAXID
	BESIVANCE		be required before approval can be given for a non-preferred agent.	
	gentamlcin			
	moxifloxacin 0.5% ofloxacin			
	tobramycin			
		IFLAMMATORY	Trial and failure of ALL preferred agents each greater than or equal to 5 day supply in the last 12	
	flurbiprofen diclofenac		months will be required before approval can be given for a non- preferred agent.	ACUVAIL bromfenac 0.9%
	LOTEMAX*			BROMSITE
	ketorolac			DUREZOL
	NEVANAC			ILEVRO INVELTYS
				LOTEMAX SM
				loteprednol 0.5% (BRAND IS PREFERRED)
	00.057		T	PROLENSA
	betaxolol	A-BLOCKERS	Trial and failure of three (3) preferred agents each greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non- preferred agent.	BETIMOL BETOPTIC S*
	carteolol		*Betoptic S will be approved for those with heart and lung conditions.	
	levobunolol	1		
	timolol OPCARBONIC A	NHYDRASE INHIBITOR	Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will	brinzolamide (BRAND IS PREFERRED)
	AZOPT		be required before approval can be given for a non-preferred agent.	
	dorzolamide	<u> </u>		
		BO PRODUCTS		dorzolamide/timolol (BRAND IS PREFERRED)
	COMBIGAN* ROCKLATAN			
	SIMBRINZA			
		EYE AGENTS	Trial and failure of the preferred agent greater than or equal to 12 weeks will be required	CEQUA
	RESTASIS* XIIDRA		before approval can be given for the non-preferred agent.	cyclosporine (BRAND IS PREFERRED) EYSUVIS
	AIIDKA			MIEBO
				RESTASIS MULTIDOSE (see preferred)
		L		TYRVAYA
	OPPROST latanoprost	TAGLANDINS	Trial and failure of ALL preferred agents each greater than or equal to 30 days in the last 12	bimatoprost IYUZEH
	LUMIGAN		months will be required before approval can be given for a non- preferred agent.	tafluprost
	TRAVATAN Z			
	XALATAN ZIOPTAN			
	OPRHO KINA			
	RHOPRESSA			
		THOMIMETICS	Trial of a preferred agent greater than or equal to 30 days in the last 12 months will be required	brimonidine 0.15% (BRAND IS PREFERRED)
	ALPHAGAN P 0.1% ALPHAGAN P 0.15%*		before approval can be given for a non-preferred agent.	
	brimonidine 0.2%			
OSTEOPOROSIS		PHONATES	Trial and failure of a preferred agent greater than or equal to 12 months will be required before	
	alendronate ibandronate		approval can be given for a non-preferred agent.	FORTEO*** FOSAMAX-D
	risedronate			TYMLOS***
		1	Fosamax liquid will be approved for clients that have difficulty swallowing.	
			**Evinity will only be allowed for a maximum of 12 months of treatment, will not be allowed	
			**Evinity will only be allowed for a maximum of 12 months of treatment, will not be allowed with any concurrent osteoporosis treatment, and will be limited to approved indication	
			with any concurrent osteoporosis treatment, and will be limited to approved indication	
		ALCITONIN	with any concurrent osteoporosis treatment, and will be limited to approved indication	
זדוכ	calcitonin-salmon	ALCITONIN OID COMBINATION	with any concurrent osteoporosis treatment, and will be limited to approved indication	ciprofloxacin 0.2% (use preferred agent)
	calcitonin-salmon ANTIBIOTIC/STER ciprofloxacin/dexamethasone		with any concurrent osteoporosis treatment, and will be limited to approved indication	CIPRO HC (use preferred agent)
	calcitonin-salmon ANTIBIOTIC/STER ciprofloxacin/dexamethasone Neo/Poly/HC suspension/solution		with any concurrent osteoporosis treatment, and will be limited to approved indication	CIPRO HC (use preferred agent) CORTISPORIN-TC (use preferred agent)
	calcitonin-salmon ANTIBIOTIC/STER ciprofloxacin/dexamethasone		with any concurrent osteoporosis treatment, and will be limited to approved indication	CIPRO HC (use preferred agent)
	calcitonin-salmon ANTIBIOTIC/STER ciprofloxacin/dexamethasone Neo/Poly/HC suspension/solution ofloxacin tobramycin/dexamethasone OVERACTIVE B		with any concurrent osteoporosis treatment, and will be limited to approved indication ***Will be limited to 2 years of use Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12	CIPRO HC (use preferred agent) CORTISPORIN-TC (use preferred agent) FLUOCINOLONE ACET OIL 0.01% (use preferred agent) darifenacin
	calcitonin-salmon ANTIBIO TIC/STER ciprofloxacin/dexamethasone Neo/Poly/HC suspension/solution ofloxacin tobramycin/dexamethasone OVERACTIVE B MYRBETRIQ	OID COMBINATION	with any concurrent osteoporosis treatment, and will be limited to approved indication ***Will be limited to 2 years of use Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent.	CIPRO HC (use preferred agent) CORTISPORIN-TC (use preferred agent) FLUOCINOLONE ACET OIL 0.01% (use preferred agent) darifenacin GELNIQUE GEL 10%
	calcitonin-salmon ANTIBIOTIC/STER ciprofloxacin/dexamethasone Neo/Poly/HC suspension/solution ofloxacin tobramycin/dexamethasone OVERACTIVE B MYRBETRIQ oxybuttynin /ER	OID COMBINATION	with any concurrent osteoporosis treatment, and will be limited to approved indication ***Will be limited to 2 years of use Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12	CIPRO HC (use preferred agent) CORTISPORIN-TC (use preferred agent) FLUOCINOLONE ACET OIL 0.01% (use preferred agent) darifenacin GELNIQUE GEL 10% GELNIQUE GEL 10%
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OVERACTIVE BLADDER	calcitonin-salmon ANTIBIOTIC/STER ciprofloxacin/dexamethasone Neo/Poly/HC suspension/solution ofloxacin tobramycin/dexamethasone OVE:ACTIVE B MYRBETRIQ oxybutynin /ER solifenacin	OID COMBINATION	with any concurrent osteoporosis treatment, and will be limited to approved indication ***Will be limited to 2 years of use Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. Oxytrol will be approved for clients that have an inability to swallow. Trial and failure of a preferred agent(s) greater than or equal to a 14 day supply in the last 12	CIPRO NC (use preferred agent) CORTISPORINATC (use preferred agent) FLUOCINOLONE ACET OIL 0.01% (use preferred agent) darifenacin GELNIQUE GEL 10% GEMTESA mirabegron (BRAND IS PREFERRED) OXYTROL DIS totlerodine/ER TOVIAZ trospium fentanyl patches hydrocodone ER hydrocodone ER
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OVERACTIVE BLADDER	calcitonin-salmon ANTIBIOTIC/STER ciprofloxacin/dexamethasone Neo/Poly/HC suspension/solution ofloxacin tobramycin/dexamethasone OVE:ACTIVE B MYRBETRIQ oxybutynin /ER solifenacin	OID COMBINATION	with any concurrent osteoporosis treatment, and will be limited to approved indication ***Will be limited to 2 years of use Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. Oxytrol will be approved for clients that have an inability to swallow. Trial and failure of a preferred agent(s) greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. C-IIIs and C-IVs that are not included on the PDL and are available without prior authorization	CIPRO NE (use preferred agent) CORTISPORIN-TC (use preferred agent) FLUOCINOLONE ACET OL 0.01% (use preferred agent) darifenacin GELNIQUE GEL 10% GEMTESA mirabegron (BRAND IS PREFERRED) OXYTROL DIS tolterodine/ER TOVIAZ trospium fentanyl patches hydrocodone ER hydromorphone ER HYSINGLA ER METHADONE morphine ER cansules (use preferred agents)
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OTIC OVERACTIVE BLADDER PAIN	calcitonin-salmon ANTIBIOTIC/STER ciprofloxacin/dexamethasone Neo/Poly/HC suspension/solution ofloxacin tobramycin/dexamethasone OVE:ACTIVE B MYRBETRIQ oxybutynin /ER solifenacin	OID COMBINATION	with any concurrent osteoporosis treatment, and will be limited to approved indication ***Will be limited to 2 years of use Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. Oxytrol will be approved for clients that have an inability to swallow. Trial and failure of a preferred agent(s) greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. Oxytrol will be approved for clients that have an inability to swallow. Trial and failure of a preferred agent(s) greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non- preferred agent. C-IIIs and C-IVs that are not included on the PDL and are available without prior authorization with the exception of Butrans (generic substitution is mandatory). Concurrent therapy with a benzodiazepine and a narcotic medication or with duplicate benzodiazepines is not covered by Wyoning Medicaid. A single medication will continue to be allowed to process unless another benzodiazepine (or narcotic) is billed to Wyoming Medicaid. Fentanyl: SOmcg. 1 strength at a time, 1 patch every 3 days Hydromorphone ER: 30mg/day HysingIa ER: 120mg/day	CIPRO NE (<i>use preferred agent</i>) CORTISPORIN-TC (<i>use preferred agent</i>) FLUOCINOLONE ACET OIL 0.01% (<i>use preferred agent</i>) darifenacin GELNIQUE GEL 10% GEMTESA mirabegron (BRAND IS PREFERRED) OXYTROL DIS totterodine/ER TOVIAZ trospium fentanyl patches hydrocodone ER hydrocodone ER HYSINGLA ER METHADONE morphine ER capsules (<i>use preferred agents</i>) axymorphone ER
OVERACTIVE BLADDER	calcitonin-salmon ANTIBIOTIC/STER ciprofloxacin/dexamethasone Neo/Poly/HC suspension/solution ofloxacin tobramycin/dexamethasone OVE:ACTIVE B MYRBETRIQ oxybutynin /ER solifenacin	OID COMBINATION	with any concurrent osteoporosis treatment, and will be limited to approved indication ***Will be limited to 2 years of use Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. Oxytrol will be approved for clients that have an inability to swallow. Trial and failure of a preferred agent(s) greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. Oxytrol will be required before approval can be given for a non-preferred agent. Crills and C-IVs that are not included on the PDL and are available without prior authorization with the exception of Butrans (generic substitution is mandatory). Concurrent therapy with a benzodiazepine and a narcotic medication or with duplicate benzodiazepines is not covered by Wyoming Medicaid. A single medication will continue to be allowed to process unless another benzodiazepine (or narcotic) is billed to Wyoming Medicaid. Fentanyl: SOncg. 1 strength at a time, 1 patch every 3 days Hydromorphone ER: 30mg/day Hysingla ER: 120mg/day Morphine ER: 90mg/day	CIPRO NE (<i>use preferred agent</i>) CORTISPORIN-TC (<i>use preferred agent</i>) FLUOCINOLONE ACET OIL 0.01% (<i>use preferred agent</i>) darifenacin GELNIQUE GEL 10% GEMTESA mirabegron (BRAND IS PREFERRED) OXYTROL DIS totterodine/ER TOVIAZ trospium fentanyl patches hydrocodone ER hydrocodone ER HYSINGLA ER METHADONE morphine ER capsules (<i>use preferred agents</i>) axymorphone ER
OVERACTIVE BLADDER	calcitonin-salmon ANTIBIOTIC/STER ciprofloxacin/dexamethasone Neo/Poly/HC suspension/solution ofloxacin tobramycin/dexamethasone OVE:ACTIVE B MYRBETRIQ oxybutynin /ER solifenacin	OID COMBINATION	with any concurrent osteoporosis treatment, and will be limited to approved indication ***Will be limited to 2 years of use Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. Oxytrol will be approved for clients that have an inability to swallow. Trial and failure of a preferred agent(s) greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. Oxytrol will be required before approval can be given for a non-preferred agent. Crills and C-IVs that are not included on the PDL and are available without prior authorization with the exception of Butrans (generic substitution is mandatory). Concurrent therapy with a benzodiazepine and a narcotic medication or with duplicate benzodiazepines is not covered by Wyoming Medicaid. A single medication will continue to be allowed to process unless another benzodiazepine (or narcotic) is billed to Wyoming Medicaid. Fentanyl: SOncg. 1 strength at a time, 1 patch every 3 days Hydromorphone ER: 30mg/day Hysingla ER: 120mg/day Morphine ER: 90mg/day	CIPRO NE (<i>use preferred agent</i>) CORTISPORIN-TC (<i>use preferred agent</i>) FLUOCINOLONE ACET OIL 0.01% (<i>use preferred agent</i>) darifenacin GELNIQUE GEL 10% GEMTESA mirabegron (BRAND IS PREFERRED) OXYTROL DIS totterodine/ER TOVIAZ trospium fentanyl patches hydrocodone ER hydrocodone ER HYSINGLA ER METHADONE morphine ER capsules (<i>use preferred agents</i>) axymorphone ER

Please refer to the Addit	tional Therapeutic Criteria C	hart, Dosage Limitation List (r	ed font indicates quantity/dose limits apply), and Wyoming Medicaid Pro	vider Manual for additional criteria.
THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIES THIS LIST IS NOT ALL INCLUSIVE PLASE CONTACT Optimum WITH ANY OUTSTONS
PAIN continued	codeine sulfate hvdrocodone/APAP hydrocodone/IBU	CTING C-IIs	Trial and failure of three (3) preferred agents greater than or equal to a 6 day supply in the last 90 days will be required before approval can be given for a non-preferred agent.	levorphanol oxvmorohone ROXYBOND
	hydromorphone meoeridine morphine oxycodone oxycodone/APAP		Concurrent therapy with a benzodiazepine and a narcotic medication or with duplicate benzodiazepines is not covered by Wyoming Medicaid. A single medication will continue to be allowed to process unless another benzodiazepine (or narcotic) is billed to Wyoming Medicaid.	
			All short-acting narcotics, after 42 days of consecutive use of any combination of short-acting narcotics, will be limited to 4 tablets per day (liquids have specific dosing limits per medication - please refer to dosage limitation chart at www.wymedicaid.org)	
			Clients will be limited to one short-acting narcotic at a time	
	C-III/C- BUTRANS tramadol	V AGENTS	Trial and failure of a preferred agent(s) greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non- preferred agent. Quantity and dosage limits apply (max 8 tabs/day).	BELBUCA tramadol/apap tramadol ER capsules/tablets
			Concurrent therapy with a benzodiazepine and a narcotic medication or with duplicate benzodiazepines is not covered by Wyoming Medicaid. A single medication will continue to be allowed to process unless another benzodiazepine (or narcotic) is billed to Wyoming Medicaid.	
PARKINSON'S DISEASE		TING AGENTS		
	amantadine benztropine tablets carbidopa/levodopa pramipexole ropinirole			
	LONG-ACT ropinirole ER RYTARY	ING AGENTS	**Non-preferred Parkinson's agents will require a 30 day trial and failure of 2 preferred medications including at least one short-acting agent and one long-acting agent	APOKYN benztropine injectables GOCOVRI INBRIJA
			Neupro will be approved for clients with difficulty swallowing	NEUPRO ONGENTYS pramipexole ER XADAGO
PHOSPHATE BINDERS	PHOSPHA calcium acetate	TE BINDERS	Prior authorization required for non-preferred agents.	AURYXIA lanthanum sevelamer VELPHORO
PROSTATE	5-ALPHA-REDU finasteride	ICTASE INHIBITORS	Trial and failure of a preferred agent greater than or equal to a 30 day supply in the last 12 months will be required before approval can be given for a non-preferred agent.	dutasteride dutasteride/tamsulosin (<i>use separate agents</i>)
	ALPHA doxazosin tamsulosin terazosin	BLOCKERS	Trial and failure of a preferred agent greater than or equal to a 30 day supply in the last 12 months will be required before approval can be given for a non-preferred agent.	alfuzosin dutasteride/tamsulosin <i>(use separate agents)</i> silodosin
PULMONARY ANTIHYPERTENSIVES		ICTASE INHIBITORS ALYQ sildenafil suspension sildenafil (A/B rated generics)	Prior authorization required. Client must have a diagnosis of pulmonary hypertension.	OPSYNVI REVATIO (suspension)
	ENDOTHELIN REC	EPTOR ANTAGONISTS LETAIRIS TRACLEER TABS*	Prior authorization required. Client must have a diagnosis of pulmonary hypertension.	bosentan (BRAND IS PREFERRED) OPSUMIT (use preferred agent) TRACLEER TABS FOR ORAL SUSP (use preferred agent)
	GUANYLATE C	YCLASE INHIBITORS	Prior authorization required.	WINREVAIR ADEMPAS (use preferred agent)
		E VASODILATORS	Prior authorization required. Client must have a diagnosis of pulmonary hypertension.	
		ORENITRAM		
	PROSTACYCLINE F	RECEPTOR AGONIST	Prior authorization required.	UPTRAVI (use preferred agent)
RESTLESS LEG SYNDROME	RESTLESS LE pramipexole ropinirole	G SYNDROME gabapentin pregabalin	Client must have a diagnosis of Restless Leg Syndrome (RLS). Trial and failure of gabapentin greater than or equal to 60 days <u>and</u> a trial and failure of a dopamine agonist greater than or equal to 60 days in the last 12 months will be required before approval can be given for a non- preferred agent. "Neupro will be approved for clients with difficulty swallowing or for clients with a diagnosis of Parkinson's Disease.	HORIZANT NEUPRO*
			Clients will not be allowed to take gabapentin and pregabalin concurrently	
SKELETAL MUSCLE RELAXANTS	MUSCLE baclofen (5, 10, 20mg tablets) cyclobenzaprine tizanidine tablets	RELAXANTS	Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent.	carisoprodol chlorzoxazone cyclobenzaprine ER LYVISPAH metaxalone
			Cyclobenzaprine will require a prior authorization for clients concurrently taking a tricylic antidepressant. Carisoprodol is limited to 84 tabs/365 days	methocarbamol orphenadrine tizanidine capsules (use preferred agent)
	IMMUNON	IODULATORS HUMIRA	Client must have diagnosis of UC prior to approval of a preferred agent. To receive a non- preferred agent, client must have a diagnosis of UC and a 56-day trial and failure of the preferred agent.	ENTYVIO* REMICADE RINVOQ
ULCERATIVE COLITIS			preferred ugent.	SIMPONI
ULCERATIVE COLITIS			* Refer to Additional Therapeutics Clinical Criteria Chart for more information	SIMPONI SKYRIZI STELARA TREMFYA XELJANZ/XR