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

Unless otherwise noted on the PDL, generic substitution is mandatory.

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 Change Healthcare PA Helpdesk @ 877-207-1126 for prior authorization if client has primary insurance that will not cover the brand name medication.

Please refer to the Additional Therapeutic Criteria Chart, Dosage Limitation List (red font indicates quantity/dosage limits apply), and the Wyoming Medicaid Provider

	Manual at http://wymedicaid.org 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 PLEASE CONTACT GHANGE HEALTHCARE WITH ANY QUESTION:	
DDICTION	BUPRENORPHINE CO		Client must have a diagnosis of opioid dependence or abuse. This is not to be used	buprenorphine (oral)	
		buprenorphine/naloxone tablets SUBOXONE FILM*	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/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.		
			Dosage limits apply Prior authorization will be required for doses > 24mg		
	NALOXO KLOXXADO naloxone NARCAN NASAL SPRAY	NE	Kloxxado, naloxone products, and Narcan nasal spray will be limited to one fill per 180 days without prior authorization.	naloxone nasal spray	
			Naloxone formulations available in quantities of 10ml will require prior authorization.		
	NALTREXO	naltrexone	Client must have a diagnosis of alcohol or opioid dependance.		
		NITROL	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.		
LLERGY / ASTHMA / COPD	ANTIHISTAMINES, MINI	MALLY SEDATING		desloratadine	
	cetirizine fexofenadine loratadine		last 12 months will be required before approval can be given for a non-preferred agent.	CLARINEX RDT/SYRUP levocetirizine	
	ANTIHISTAMINE/DECONGES cetirizine/pseudoephedrine fexofenadine/pseudoephedrine loratadine/pseudoephedrine		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	
	ANTICHOLINERGIC BRC ATROVENT HFA INCRUSE ELLIPTA ipratropium SPIRIVA HANDIHALER SPIRIVA RESPINAT	INCHODILATORS	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 (use brand) TUDORZA YUPELRI	
			Spiriva 5 day STARTER package will be allowed one (1) time per recipient		
	ANTICHOLINERGIC COMI ANORO ELLIPTA** COMBIVENT STIOLTO	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.	BEVESPI BREZTRI DUAKLIR TRELEGY	
	LEUKOTRIENE M	IODIEIERS	**Will also require the diagnosis of COPD. Trial and failure of preferred agent greater than or equal to 30 days in the last 12	zafirlukast	
	montelukast	COPPERS	Irial and failure of preterred 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.	Lammukast	
	LONG ACTING BRON arformoterol SEREVENT STRIVERDI	CHODILATORS	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	
	NASAL ANTIHIS azelastine 0.1%	TAMINES	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 (use separate agents) olopatadine 0.6% RYALTRIS	

	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIE PLEASE CONTACT CHARGE HEATHCOSE WITH ANY QUESTI
RGY / ASTHMA / COPD	NASAL STEP	ROIDS	Trial and failure of two (2) preferred agents greater than or equal to 30 days in the	DYMISTA (use separate agents)
itinued	budesonide flunisolide		last 12 months will be required before approval can be given for a non-preferred agent.	OMNARIS QNASL
	fluticasone			XHANCE
	mometasone		Budesonide will be approved for pregnancy.	ZETONNA
	SHORT ACTING BRONCHOL	DILATORS - INHALERS	Trial and failure of a preferred agent greater than or equal to 30 days in the last 12	levalbuterol (BRAND IS PREFERRED)
	albuterol HFA PROAIR RESPICLICK VENTOLIN HFA		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.	PROAIR DIGIHALER PROVENTIL HFA
	XOPENEX HFA*		Minimum day supply of 16 days is required	
	STEROID INH	ALANTS	Trial and failure of two (2) preferred agents greater than or equal to 30 days in the	AIRDUO DIGIHALER
	AIRDUO RESPICLICK ARNUITY ELIPTA		last 12 months will be required before approval can be given for a non-preferred agent.	ALVESCO ARMONAIR
	ASMANEX TWISTHALER budesonide suspension		*Fluticasone HFA and Asmanex HFA will be approved for pediatric clients 8 years of	ASMANEX HFA* fluticasone HFA*
	PULMICORT FLEXHALER		age or younger. Alvesco will be approved for a history of oral thrush with steroid inhalants.	QVAR REDIHALER
	STEROID COMBINA BREO ELLIPTA**	TION 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.	fluticasone/vilanterol (use preferred agent, fluticasone/salmeterol 55-14/113-14/232-1
	DULERA SYMBICORT*			fluticasone/salmeterol 100-50/250-50/500- (BRAND IS PREFERRED)
	WIXELA		**Will also require the diagnosis of COPD or uncontrolled asthma.	TRELEGY
			Advair 7 and 14-day STARTER package will be allowed one (1) time per recipient.	
	EPINEPHR epinephrine auto-injector pen	RINE		AUVI-Q (use preferred agent) EPI-PEN (use preferred agent)
	EOSINOPHILIC ASTI	HMA AGENTS	*Approval for these agents will require additional clinical criteria which can be	FASENRA*
	EGSINOT TIETE AST	DUPIXENT XOLAIR	found on the Additional Therapeutic Criteria Chart	NUCALA* TEZSPIRE
HRITIS	IMMUNOMOD		Client must have diagnosis of AS prior to approval of a preferred agent. To receive a	
HRITIS	ANKYLOSING SPON	NDYLITIS (AS)	non-preferred agent, client must have a diagnosis of AS and a 56-day trial and	COSENTYX
		ENBREL HUMIRA	failure of both preferred agents.	REMICADE RINVOQ
			**Cimzia will be allowed for clients that are pregnant or breast-feeding	SIMPONI TALTZ XELJANZ/XR
			Quantity Limits apply for all diagnoses:	ALEXALLY AND
			Enbrel 25mg - limited to 10 per month	
			Enbrel 50mg - limited to 5 per month Humira 20mg - limited to 10 per month	
	JUVENILE IDIOPATHIC	ARTHRITIS (JIA)	Humira 40mg - limited to 5 per month Client must have diagnosis of JIA prior to approval of a preferred agent. To receive	ACTEMRA
		ENBREL HUMIRA	a non-preferred agent, client must have a diagnosis of JIA and a 56-day trial and failure of both preferred agents.	ILARIS ORENCIA
				XELJANZ/XR
	PSORIATIC ARTH	ENBREL	Client must have diagnosis of PA prior to approval of a preferred agent. To receive a non-preferred agent, client must have a diagnosis of PA and a 56-day trial and	CIMZIA** COSENTYX
		HUMIRA	failure of two of the three preferred agents.	ORENCIA
		OTEZLA*		REMICADE RINVOQ
				SIMPONI
			*Otezla starter pack is non-preferred	STELARA TALTZ
			**Cimzia will be allowed for clients that are pregnant or breast-feeding	TREMFYA
	RHEUMATOID AR		Client must have diagnosis of RA and a 56-day trial and failure of methotrexate	XELJANZ/XR ACTEMRA
		ENBREL HUMIRA	prior to approval of a preferred agent. To receive a non-preferred agent, client must have a diagnosis of RA and a 56-day trial and failure of both preferred agents.	CIMZIA* KEVZARA
				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 XELJANZ/XR
VULSIONS	INTERMITTENT, STEREOTYF	PIC SEIZURE EPISODES	*Nayzilam will be allowed for patients 12 years of age and older	ALLIMINZ/AR
	diazepam gel NAYZILAM*			
	VALTOCO ORAL ANTICON	/I II CANTS	Professed agents with clinical criteria will be limited to 500	ADTIONA (uso preferred as 11)
	ORAL ANTICON carbamazepine	BANZEL (tablets only)	Preferred agents with clinical criteria will be limited to FDA approved indications related to seizures and epilepsy. Non-preferred agents require 30 day trial and	APTIOM (use preferred agent) BRIVIACT (use preferred agent)
	divalproex FELBAMATE	clonazepam EPIDIOLEX	failure of two preferred agents prior to approval.	clobazam** DIACOMIT**
	fosphenytoin	gabapentin	For indications not related to seizures and epilepsy, please refer to the Additional	FINTEPLA**
	lacosamide (tablets)	pregabalin	Therapeutic Criteria chart at www.wymedicaid.org.	levetiracetam ER
	lamotrigine/XR levetiracetam	topiramate/ER sprinkle caps	**Please refer to the Additional Therapeutic Criteria chart at www.wymedicaid.org	OXTELLAR (use preferred agent) TROKENDI XR (use preferred agent)
	oxcarbazepine		for specific requirements.	XCOPRI
!	phenytoin subvenite			VIMPAT (tablets) zonisamide oral susp. (use preferred agen
	valproate/valproic acid			action action and a supply (use preferred agent
			The state of the s	İ.
	VIMPAT (suspension) zonisamide			
	VIMPAT (suspension)			
	VIMPAT (suspension)			

THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIES
		CLINICAL CRITERIA		THIS LIST IS NOT ALL INCLUSIVE PLEASE CONTACT CHANGE HEALTHCARE WITH ANY QUESTION
OHN'S	IMMUNOMOD	JLATORS HUMIRA	Client must have diagnosis of Crohn's prior to approval of the preferred agent. To receive a non-preferred agent, client must have a diagnosis of Crohn's and a 56-day	CIMZIA** ENTYVIO*
			trial and failure of the preferred agent.	REMICADE
			* Refer to Additional Therapeutics Clinical Criteria Chart for more information	RINVOQ STELARA
			**Cimzia will be allowed for clients that are pregnant or breast-feeding	TYSABRI (additional criteria applies)
RMATOLOGY	BENZOYL PEROXIDE/CLIN		Clients must be 12 to 20 years of age. Requires prior authorization for clients less	ACANYA (use preferred agent)
		clindamycin/benzoyl peroxide 1-5% clindamycyin/benzoyl peroxide	than 12 years of age. Acne combinations are limited to clients under the age of 21.	ONEXTON (use preferred agent)
		1.2-5% (Refrig)		
	AMNESTEEM ISOTRETIN	OIN		ABSORICA (use preferred agents)
	CLARAVIS			
	isotretinoin			
	ZENATANE			
	CORTICOSTEROIDS - S		Trial and failure of two preferred agents greater than or equal to 14 days in the last	
	C=CREAM; G=GEL; L=LOTI		90 days.	TEXACORT 2.5% (S)
	alclometasone desonide*		*Cream, ointment, and lotion formulations of Desonide are preferred.	
	fluocinolone 0.01%		Gream, omathent, and lodion formulations of Desortide are preferred.	
	hydrocortisone butyrate 0.1% (C)			
	hydrocortisone 1%, 2.5% (C,L,O)			
	MEDIUM PO	TENCY	Trial and failure of two preferred agents greater than or equal to 14 days in the last	Clocortolone Pivalate
	betamethasone valerate		90 days.	flurandrenol
	desoximetasone 0.05%, 0.25% (C) fluocinolone 0.025%			fluticasone 0.05% (L) hydrocortisone butyrate 0.1% (O)
	fluticasone 0.05% (C)			triamcinolone 0.05% (O)
	mometasone SYNALAR 0.025% (C, O)			
	triamcinolone 0.025%, 0.1%			
	HIGH POTE betamethasone dipropionate	NCY	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,O)
	diflorasone 0.05% (O) fluocinonide			clobetasol 0.05% (L) desoximetasone 0.05%, 0.25% (G,O)
	flurandrenolide			diflorasone 0.05% (C)
	fluticasone 0.005% (O) halobetasol			fluocinonide 0.1% (C) halcinonide 0.1% (C)
	TOPICORT 0.025% (C)			HALOG 0.1% (O)
	triamcinolone 0.5% ULTRAVATE 0.05% (C,O)			
	IMMUNOMODULATORS			pimecrolimus (brand preferred)
		ELIDEL tacrolimus	topical 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 last 90 days will be required.	
	PHOSPHODIESTERASE 4 INHI	BITOR - STEP 3 AGENT	To receive a step 3 agent: Trial and failure of a preferred step 2 agent	EUCRISA
			(immunomodulator) greater than or equal to a 21 day trial within the last 30 days.	
	ATOPIC DERM	IATITIS DUPIXENT*	*Trial and failure of a preferred step 2 agent (immunomodulator) greater than or	ADBRY** CIBINOO**
		DOT MENT	lequal to a 21 day trial within the last 30 days required. Dupixent requires member be at least 6 months of age or older. No high-potency steroid trial will be necessary.	OPZELURA**
				RINVOQ**
			**Trial and failure of all criteria to receive a step 3 agent as defined above including	
			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 PSORI		Client must have diagnosis of PP prior to approval of a preferred agent (Enbrel,	CIMZIA**
		ENBREL	Humira, or Otezla). To receive a non-preferred agent, client must have a diagnosis of PP and a 56-day trial and failure of two of the three preferred agents.	COSENTYX
		HUMIRA OTEZLA	or Pr and a 36-day trial and failure of two of the three preferred agents.	ILUMYA REMICADE
		SOTYKTU*		SILIQ
			*Sotyktu requires diagnosis of moderate to severe PP and 56 day trial and failure of Humira.	SKYRIZI STELARA
			**Cimzia will be allowed for clients that are pregnant or breast-feeding	TALTZ
	SCABICIDES/PED	CULICIDES	Trial and failure of a preferred agent in the last 12 months.	TREMFYA malathion lotion
				NATROBA
	permethrin VANALICE			spinosad (BRAND IS PREFERRED)

Please refer to the A	Additional Therapeutic Criteria (ist (red font indicates quantity/dosage limits apply), and the dicaid.org for additional criteria.	e Wyoming Medicaid Provider
THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIES THIS SELECT OF THE PROPERTY OF THE PROPE
NABETES	DIABETES AC BIGUANII metformin/ER			metformin SR 24HR osmotic release(use preferred agent) metformin SR 24HR modified release (use preferred agent)
	α-GLUCOSIDASE acarbose	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
	MEGLITINI nateglinide	DES	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.	repaglinide
	THIAZOLIDINE pioglitazone	DIONES	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.	ACTOPLUS MET (use separate agents)
	SULFONYLU glimepiride/ER glipizide/ER glyburide/ER	REAS	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.	
	DIPEPTIDYL PEPTIDASE 4	DPP-4) INHIBITORS JANUVIA ONGLYZA TRADJENTA	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 preferred agent is required before approval can be give for a non-preferred agent.	GLYXAMBI (use separate preferred agents)
	DPP-4 INHIBITOR CO	JANUMET/KR JANUMET/KR JENTADUETO KOMBIGLYZE/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 preferred agent is required before approval can be give for a non-preferred agent.	alogliptin/metformin alogliptin/pioglitazone (use separate preferred agents) IENTADUETO XR saxagliptin/metformin (use brand)
	INCRETIN MIMETICS (GLP-1	RECEPTOR AGONISTS) BYETTA TRULICITY VICTOZA	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 unless ASCVD or risk factors are present, in which case the trial of metform is waived. A 90 day trial of failure of the preferred agent is required before approval can be give for a non-preferred agent. *Rybelsus requires documentation of inability to use injectable agents. Dosage Limits Apply: Ozempic: 2mg/week Victoza: 1.8mg/day	MOUNJARO OZEMPIC*
	SGLT2 INHIB	TORS FARXIGA INVOKAMET INVOKANA JARDIANCE SYNJARDY XIGDUO 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 unless there is a diagnosis of ASCVQ. KCN, or heart failure, in which case the trial of metformin will be waived. A 90 day trial and failure of a preferred agent is required before approval can be given for a non-preferred agent.	QTERN (use separate preferred agents) SEGLUROMET (use separate preferred agents)
	FAST-ACTING HUMALOG HUMALOG 75/25 HUMALOG JR. HUMALOG MIX	INSULIN	Prior authorization will be required when using two different delivery forms of the same type of insulin concurrently.	ADMELOG (use preferred agent) FIASP (use preferred agent) insulin lispro (use preferred agents) LYUMJEV
	NOVOLOG MIX LANTUS SOLOSTAR* LANTUS vial LEVEMIR	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) Insulin Degludec SOLIQUA TOUIEO (use preferred agent) TRESIBA* (use preferred agent) XULTOPHY (use separate preferred agent)
	DIABETIC METERS/ FREESTYLE (Strips only) FREESTYLE FREEDOM FREESTYLE FREEDOM LITE FREESTYLE FREEDOM LITE FREESTYLE INSULINX FREESTYLE SIDEMCK II ONE TOUCH ULTRA MINI ONE TOUCH ULTRA MINI ONE TOUCH VERIO ONE TOUCH VERIO ONE TOUCH VERIO ONE TOUCH VERIO FLEX ONE TOUCH VERIO FLEX ONE TOUCH VERIO REFLECT ONE TOUCH VERIO IQ PRECISION XTRA	TEST STRIPS	Quantity limits apply: Insulin Dependent Clients: 10 strips/day Non-Insulin Dependent Clients: 4 strips/day Clients are limited to 1 meter/365 days	ALL OTHER METERS AND TEST STRIPS
	EXTERNAL DIABET OMNIPOD CLASSIC OMNIPOD 5 CONTINUOUS BLOOD GL		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
BROMYALGIA	ACUTE HYPOGLYCE BAQSIMI ZEGALOGUE (autoinjector) FIBROMYA	MIA AGENTS	Trial and failure of a preferred agent greater than or equal to six (6) weeks in the	GVOKE (use preferred agent) pregabalin
I ALGIA	amitriptyline cyclobenzaprine duloxetine	gabapentin	last 12 months is required prior to approval of a non-preferred agent	pregabatin SAVELLA tablets (savella titration pak will not be covered)
			Clients will not be allowed to take gabapentin and pregabalin concurrently	

				NON-PREFERRED AGENTS
THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	GENERIC MANDATORY POLICY APPLIE THIS LIST IS NOT ALL INCLUSIVE PLEASE CONTACT CHANGE HEALTHCARE WITH ANY QUESTS
STROINTESTINAL	BOWEL EVA	CUANTS		CLENPIQ (use preferred agents) GAVILYTE H (use preferred agents)
	GOLYTELY MOVIPREP PEG 3350 SOLUTION			POLY-PREP (use preferred agents) SUFLAVE SUTAB
	SUPREP			
	CHRONIC IDIOPATHI	C CONSTIPATION AMITIZA	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	MOTEGRITY
		LINZESS TRULANCE	of chronic idiopathic constipation and a 30-day trial and failure of a preferred agent within the last 12 months.	
	DIGESTIVE E	NZYMES	Prior authorization required.	PERTZYE VIOKACE
	ZENPEP			
	IRRITABLE BOWEL SYNDROI	ME WITH CONSTIPATION AMITIZA	Client must have a diagnosis of Irritable Bowel Syndrome (IBS) with constipation.	
		LINZESS TRULANCE		
	MESALAI APRISO* LIALDA* mesalamine 400mg DR capsule mesalamine enema PENTASA	VINE	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 (BRAND IS PREFERRED) mesalamine DR tab 1.2gm (BRAND IS PREFERRED) mesalamine ER cap 0.375gm (BRAND IS
				PREFERRED) mesalamine sup 1000mg SFROWASA
	OPIOID-INDUCED CON	AMITIZA	Client must have a diagnosis of opioid-induced constipation and a three (3) month trial and failure of a stool softner 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* RELISTOR SYMPROIC
			*Movantik will be approved for a diagnosis of cancer or for clients in hospice or palliative care.	
	PREGNANCY INDUCED BONJESTA DICLEGIS	NAUSEA/VOMITING		
	PROTON PUMP	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	
	lansoprazole <u>capsules</u> omeprazole <u>capsules</u> pantoprazole		agent. PREVACID solutabs will be approved for children less than or equal to 8 years of age.	DEXILANT dexlansoprazole esomeprazole omeprazole 20.6mg capsules (use preferre omeprazole tablets (use preferred agent) omeprazole/sodium bicarbonate
				OMECLAMOX (use separate agents) PREVACID solutabs* rabeprazole TALICIA (use separate agents) VIMOVO (use separate agents)
	COLCHI	CINE		MITIGARE (use preferred agent)
	colchicine (tablets) XANTHINE OXIDASE ANI	D URAT1 INHIBITORS	Trial and failure of the preferred agent greater than or equal to a 60 day supply in	ULORIC*
	allopurinol		the last 12 months will be required before approval can be given for a non- preferred agent.	
TOLOGY	LOW MOLECULAR WEIG	HT HEPARIN (LMWH)	Prior authorization will be required for the 300mg/3ml strength.	FRAGMIN (use preferred agent) enoxaparin 300MG/3ML
	DIRECT THROME	IN INHIBITOR PRADAXA	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.	
	SELECTIVE FACTOR	XA INHIBITOR XARELTO 2.5mg* (use preferred)		ELIQUIS (starter pack) SAVAYSA (use preferred agent)
	XARELTO 10mg, 15mg, 20mg, and starter pack		*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	

THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIE THIS LIST IS NOT ALL INCLUSIVE
IATOLOGY	СРТР С	ERIVATIVES	Client must have a diagnosis of acute coronary syndrome, history of myocardial	PLEASE CONTACT CHANGE HEALTHCARE WITH ANY QUESTI
tinued		BRILINTA	infarction, or history of stroke and transient ischemic attack.	
	PAR-1	NTAGONIST	Client must have diagnosis of reduction of thrombotic cardiovascular events with a	
		ZONTIVITY	history of myocardial infarction (MI) or with peripheral arterial disease (PAD). Must be used in conjunction with aspirin or clopidogrel.	
	ANTIHEMOP	HILIC FACTOR VIII		ALTUVIIIO
	ADVATE ADVNOVATE			KOVALTRY
	AFSTYLA			
	ELOCTATE ESPEROCT			
	HEMOFIL M HEMLIBRA			
	JIVI KOATE/KOATE-DVI			
	KOGENATE FS/BIO-SET			
	NOVOEIGHT NUWIQ			
	OBIZUR RECOMBINATE			
	XYNTHA/SOLOFUSE			
		ION FACTOR IX		
	ALPHANINE SD ALPROLIX			
	BENEFIX			
	IDELVION IXINITY			
	REBINYN RIXUBIS			
	ANTIHEMOPP ALPHANATE	ILIC FACTOR/VWF		
	HUMATE-P VONVENDI			
	WILATE			
	ERYTHROPOIESIS EPOGEN	STIMULATING AGENTS		ARANESP PROCRIT
	MIRCERA			T NOCHT
	RETACRIT SICKLE	ELL ANEMIA		
	DROXIA SIKLOS			
PATITIS C	DIRECT ACT	ING ANTIVIRALS	Limited to FDA approved indication. Prior authorization will be required prior to	EPCLUSA (use preferred agent)
		sofosbuvir/velpatasvir MAVYRET	use of preferred agents.	HARVONI SOVALDI
		WWW.		VOSEVI**
			Please submit PA requests on the Hepatitis C PA form available at	ZEPATIER
			www.wymedicaid.org.	
PRADENITIS SUPPURATIVA	INAMIINO	MODULATORS	Humira will not be covered as a first line agent for the diagnosis for hidradenitis	
RADENITIS SOFFORATIVA	IIVIIVIONO	HUMIRA	suppurativa.	
RMONES		TAGONISTS	*Please refer to the Additional Therapeutic Criteria chart at www.wymedicaid.org	ORILISSA
	MYFEMBREE ORIAHNN		for specific requirements.	
	GROWT	HORMONE GENOTROPIN	-	HUMATROPE NGENLA
		NORDITROPIN		OMNITROPE
		NUTROPIN AQ		SAIZEN SEROSTIM
				SKYTROFA SOGROYA
				ZOMACTON
	TESTOSTERO	NE TOPICAL GELS ANDROGEL*	Testosterone agents are only allowed for diagnosis of hypogonadism or insufficient	ANDRODERM (use preferred agent) FORTESTA (use preferred agent)
		TESTIM GEL		JATENZO (use preferred agent)
				TESTOPEL (use preferred agent) testosterone gel (use preferred agent)
				testosterone solution (use preferred agent
				XYOSTED (use preferred agent)
	THYROID ARMOUR THYROID	HORMONES ERMEZA	Ermeza will be covered with confirmed diagnosis of dysphagia.	THYQUIDITY TIROSINT
	LEVOXYL	ENIVIEZA		TINGOINT
	levothyroxine (tablets) LEVO-T			
	liothyronine			
	SYNTHROID			
	UNITHROID			

Please refer to the Additional Therapeutic Criteria Chart, Dosage Limitation List (red font indicates quantity/dosage limits apply), and the Wyoming Medicaid Provider Manual at http://wymedicaid.org for additional criteria. NON-PREFERRED AGENTS
GENERIC MANDATORY POLICY APPLIES PREFERRED AGENTS REQUIRING CLINICAL CRITERIA THERAPEUTIC CLASS PREFERRED AGENTS CLINICAL CRITERIA HORMONES continued alyacen 1-35, 7/7/7 aranelle afirmelle BALCOLTRA altavera amethia amethysi balziva briellyn drospir/ethinyl estradiol/levomefolate enpresse ethynodiol/ethinyl estradiol FALESSA KIT ashlvna aubra/EQ aurovela 1-20/FE 1-20, 1-35 aviane favosim kaitlib FE chew layolis FE chew levonest ayuna azurette blisovi 1-20 FE, 1.5-30 FE levonorgest/ethinyl estradiol/LO (84-7) levonorgest/ethinyl estradiol 0.15-MINASTRIN FE chew* bekyree beyaz camrese/LO chateal/EQ CHARLOTTE 24 FE chew noreth/ethinyl estradiol/FE chew 0.8/25 nortrel philith cyred dasetta 1-35, 7/7/7 philith rivelsa QUARTETTE SAFYRAL TAYTULLA tilia FE tri-legest FE TRIVORA daysee deso/ethinyl estradiol drospir/ethinyl estradiol elinest enskyce estarylla falmina finzala FE chew TWIRLA tydemy gianvi hailey FE 1/20, 1/35 iclevia vyfemla wera wymzya FE chew introvale isibloom jaimiess iunel 1-20/FE. 1.5-30/FE kalliga kariva kelnor kurvelo larin 1-20/FE, 1.5-30/FE 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 miii mono-linyah natazia NECON 0.5/35, 1/35, 1/50, 7/7/7, 10/11 nikki noreth/ethinyl estradiol/FE chw 0.4/35, 1/20 noreth/ethinyl estradiol 1-20/FE norgest/ethinyl estradiol/LO norethindrone nylia nymyo ocella pimtrea portia previfem reclipsen safyral SEASONIQUE* setlakin simliva simpesse sprintec sronyx syeda tri-estaryll/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

THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIE THIS USY IS NOT ALL INCLUSIVE. PLEASE CONTACT CHANGE HEALTHCARE WITH ANY QUEST
LIPIDEMIA	BILE ACID SEQUI cholestyramine/light colestipol	STRANT	Trial and failure of ALL preferred agents greater than or equal to six (6) months in the last 12 months will be required before approval can be given for a non- preferred agent.	WELCHOL
	STATINS, LOW P lovastatin pravastatin	OTENCY	Trial and failure of 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. 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.	fluvastatin/ER
ro	STATINS, HIGH F atorvastatin rosuvastatin simvastatin	OTENCY	Prior authorization will be required for clients under the age of 10. Trial and failure of 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.	EZALLOR LIVALO ZYPITAMAG
			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.	
	STATIN COMBIN amlodipine/atorvastatin VYTORIN*	IATIONS	Trial and failure of 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. Prior authorization will be required for clients under the age of 10.	ezetimibe/simvastatin (BRAND IS PREFERI
	PCSK9-RELATED	AGENTS PRALUENT	Client must have a diagnosis of homozygous familial hypercholesterolemia; have a diagnosis of heterozygous familial hypercholesterolemia or atherosclerotic cardiovascular disease AND not at goal with a maximum dose statin, or be intolerant to statin therapy. Approval for a non-preferred agent requires trial and failure of a preferred agent.	LEQVIO REPATHA
	TRIGLYCERIDE LOWE fenofibrate 40/48/54/67/134/145/160/200mg gemfibrozil	RING AGENTS	Trial and failure of 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.	fenofibric fenofibrate (43/50/120/130/150mg) icosapent LIPOFEN omega-3-acid

Please refer to the Additional Therapeutic Criteria Chart, Dosage Limitation List (red font indicates quantity/dosage limits apply), and the Wyoming Medicaid Provider Manual at http://wymedicaid.org for additional criteria.				
THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIES THE LIST IS NOT TALLINGUISME PLEASE CONTACT CHANGE HEALTHCASE WITH ANY QUESTION
YPERTENSION/ CARDIOLOGY	ANGIOTENSIN RECEPTOR BLOCKERS (ARBs) EDARBI irbesartan losartan olmesartan telmisartan valsartan		Non-preferred ARBs will require a history of ALL preferred ARBs before approval	candesartan eprosartan 600mg
	ARBS AND DIURETICS EDARBYCLOR irbesartan HCTZ losartan HCT olmesartan HCTZ valsartan HCTZ valsartan HCTZ ALPHA-BLOCKERS clonidine		Non-preferred ARB/diuretic combinations will require a history of ALL preferred	candesartan HCTZ telmisartan HCTZ
	COMBINATION PRODUCTS	ENTRESTO	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.	VERQUVO
FECTIOUS DISEASE	QUINOLO ciprofloxacin levofloxacin ofloxacin	NES	Please refer to the Additional Therapeutic Criteria Chart located at http://www.wymedicaid.org/additional-therapeutic-criteria for Baxdela criteria.	moxifloxacin (use preferred agents)
	DOXYCYCI doxycycline MINOCYCI minocycline/ER			DORYX (use preferred agent) minocycline 65mg and 115mg ER (use prefeagent)
	INHALED TOBR	AMYCIN	*Tobi Podhaler requires a 28 day trial of a preferred agent, as well as 28 days off of that same preferred agent prior to approval.	SOLODYN (use preferred agent) BETHKIS inhaled tobramycin
			Minimum day supply of at 56 days is required	TOBI PODHALER (use preferred agent)
	ANTI-RETRO APRETUDE BIKTARVY CIMDUO DELSTRIGO	VIRALS CABENUVA* DESCOVY* TRUVADA*	*Please refer to the Additional Therapeutic Criteria chart at www.wymedicaid.org for specific requirements.	JULUCA NORVIR RUKOBIA** STRIBILD (use separate agents) SUNLENCA
	DOVATO EVOTAZ GENVOYA ODEFSEY PIFELTRO PREZCOBIX ritonavir tablets SYMFI/LO TRIUMEQ TROGARZO		**Rukobia aproval requires documentation of multi-drug resistance defined as failure of two medications from different classes.	SYMTUZA (use separate preferred agents)
FLAMMATION	NSAID: celecoxib diclofenac tablets etodolac FLECTOR* flurbiprofen ibuprofen indomethacin ketoprofen ketorolac meclofenamate meloxicam nabumetone		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 5days/34 days; max dose 40mg/day for oral tablets).	CALDOLOR (use preferred agent) diclofenac 1.3% patch (BRAND IS PREFERREI diclofenac 1.5% soln. diclofenac 3% gel fenoprofen mefenamic acid NEOPROFEN (use preferred agent)
	naproxen oxaprozin piroxicam sulindac ORAL CORTICO:	STEROIDS		CELESTONE (use preferred agent)
	budesonide cortisone acetate dexamethasone/intensol hydrocortisone methylprednisolone prednisolone prednisolone			
SOMNIA	BELSOMRA eszopiclone zalepion zolpidem zolpidem ER	ZZEPINES	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	EDLUAR (additional criteria applies) DAYVIGO QUVIVIQ ROZEREM* zolpidem sublingual (additional criteria appli
			Prior authorization will be required when a client is taking more than one insomnia agent concurrently. Dosage limits apply: zaleplon: 30mg/day zolpidem: 15mg/day	

THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLII
ENTAL HEALTH	ALZHEIMER'S	donepezil/ODT galantamine/ER	Client must have a diagnosis of dementia.	donepezil 23mg (use preferred agent) memantine ER NAMZARIC (use separate agents)
	ANTIDEPRES	memantine tablets/solution SANTS	Trial and failure of two (2) preferred agents greater than or equal to six (6) weeks	rivastigmine capsules/patches
	NORADRENERGIC/SPECIFIC S	EROTONERGICS (NaSS)	WITHIN THE LAST 2 YEARS will be required before approval can be given for a non-	NaSS
	mirtazapine tablets		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.	mirtazapine rapid dissolve tablets (use preferred agent)
	NOREPINEPHRINE/DOPAMINE RE	UPTAKE INHIBITORS (NDRI)		NDRI
	bupropion ER/SR/XL			APLENZIN AUVELITY
	SELECTIVE SEROTONIN REUPT	AKE INHIBITORS (SSRI)	Trazodone, buspirone, fluvoxamine, MAO inhibitors, TCA's, bupropion IR, and	FORFIVO XL*
	citalopram escitalopram		venlafaxine IR do not require prior authorization but will not count towards meeting preferred therapy requirements.	SSRI citalopram capsules
	fluoxetine capsules			fluoxetine tablets
	paroxetine IR/CR sertraline		Clients will not be allowed to be on more than one antidepressant, including	VIIBRYD
	Scradine		fluvoxamine, bupropion IR, and venlafaxine IR, at one time with the exception of	
	SEROTONIN/NORPINEPHRINE RE	UPTAKE INHIBITORS (SNRI)	mirtazapine or bupropion with a SSRI or SNRI.	SNRI
	duloxetine venlafaxine ER capsules			desvenlafaxine FETZIMA
	Vermanality Err capsules		***Trintellix requires trial and failure of two preferred agents in any class	venlafaxine ER tablets (use preferred ager
			Clients five (5) years of age and younger will require prior authorization before	
			approval.	OTHER
			Dosage limits apply:	TRINTELLIX***
			bupropion ER/SR/XL: 450mg/day	
			citalopram < 60 years of age: 60mg/day citalopram > 60 years of age: 30mg/day	
			escitalopram: 30mg/day	
			fluoxetine < 18 years of age: 90mg/day fluoxetine > 18 years of age: 120mg/day	
			mirtazapine: 67.5mg/day	
			paroxetine IR/CR < 18 years of age: 75mg/day paroxetine IR > 18 years of age: 90mg/day	
			paroxetine CR > 18 years of age: 112.5mg/day	
			sertraline: 300mg/day venlafaxine ER: 337.5mg/day	
	ATYPICAL ANTIPS	SYCHOTICS	*Quetiapine doses less than 100mg will require prior authorization without a	ABILIFY MYCITE (use preferred agent)
	ABILIFY MAINTENA	Teno nes	diagnosis of mood disorder or major depressive disorder. For titration doses,	CAPLYTA
	ABILIFY ASIMTUFII aripiprazole tab/solution/ODT		contact the Change Healthcare Pharmacy Help Desk for an override.	GEODON 20MG INJ (use preferred agent) LYBALVI (additional criteria applies)
	ARISTADA		**Clients nine (9) years of age and younger will require a prior authorization to	NUPLAZID
	FANAPT**		receive approval of Latuda and Saphris. Clients eighteen (18) years of age and younger will require a prior authorization to receive approval of Fanapt.	olanzapine 10mg Inj (use preferred agent
	paliperidone INVEGA HAFYERA/SUSTENNA/TRINZA		younger will require a prior authorization to receive approval of Fanapt.	SECUADO REXULTI***
	lurasidone		***Rexulti approval for MDD treatment requires concurrent antidepressant	RYKINDO
	olanzapine PERSERIS		therapy as well as a trial and failure of aripiprazole or other preferred atypical antipsychotic indicated for the adjunct treatment of MDD.	UZEDY ZYPREXA RELPREVV
	quetiapine*			
	quetiapine ER RISPERDAL CONSTA		Trial and failure of two (2) preferred agents greater than or equal to 30 days in the	
	risperidone		last 12 months will be required before approval can be given for a non-preferred	
	SAPHRIS** VRAYLAR		agent unless otherwise specified.	
	ziprasidone			
			Prior authorization will be required for any client five (5) years of age or younger, or for any cleint taking both an injectable and oral dosage form of the same	
			medication concurrently.	
			Dosage limits apply:	
			aripiprazole <13 years of age: 15mg/day	
			aripiprazole ≥13 years of age: 30mg/day ABILIFY MAINTENA: 400mg per 26 days	
			ARISTADA 441/662/882mg: 1 injection per 28 days ARISTADA 1064mg: 1 injection per 56 days	
			ARISTADA 1064mg: 1 injection per 36 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	
		İ	INVEGA TRINZ: 1 injection per 84 days	
				1
			LATUDA 10-17 years of age: 80mg/day LATUDA >17 years of age: 160mg/day	
			LATUDA >17 years of age: 160mg/day olanzapine <13 years of age: 10mg/day	
			LATUDA >17 years of age: 160mg/day olanzapine <13 years of age: 10mg/day olanzapine ≥13 years of age: 20mg/day paliperidone: 12mg/day	
			LATUDA >17 years of age: 160mg/day olanzapine <13 years of age: 10mg/day olanzapine ≥13 years of age: 20mg/day paliperidone: 12mg/day PERSERIS: 1 injection per 28 days	
			LATUDA >17 years of age: 160mg/day olanzapine <13 years of age: 10mg/day olanzapine <13 years of age: 20mg/day paliperidone: 12mg/day peliperidone: 12mg/day persersis: 1 injection per 28 days quetiapine <13 years of age: 400mg/day quetiapine 13-17 years of age: 600mg/day	
			LATUDA >17 years of age: 160mg/day olanzapine <13 years of age: 10mg/day olanzapine >13 years of age: 20mg/day paliperidone: 12mg/day PERSERIS: 1 injection per 28 days quetiapine <13 years of age: 400mg/day quetiapine <13 years of age: 600mg/day quetiapine 137 years of age: 800mg/day quetiapine >17 years of age: 800mg/day	
			LATUDA >17 years of age: 160mg/day olanzapine <13 years of age: 10mg/day olanzapine ≥13 years of age: 20mg/day paliperidone: 12mg/day PERSERIS: 11njection per 28 days quetiapine <13 years of age: 400mg/day quetiapine <13 years of age: 600mg/day quetiapine 13-17 years of age: 600mg/day risperidone <10 years of age: 800mg/day risperidone <10-17 years of age: 3mg/day risperidone <10-17 years of age: 6mg/day	
			LATUDA >17 years of age: 160mg/day olanzapine <13 years of age: 10mg/day olanzapine >13 years of age: 20mg/day paliperidone: 12mg/day PERSERIS: 1 injection per 28 days quetiapine <13 years of age: 400mg/day quetiapine 13-17 years of age: 600mg/day quetiapine 17 years of age: 800mg/day risperidone <10 years of age: 3mg/day risperidone 10-17 years of age: 6mg/day risperidone 10-17 years of age: 6mg/day	
			LATUDA >17 years of age: 160mg/day olanzapine <13 years of age: 10mg/day olanzapine >13 years of age: 20mg/day paliperidone: 12mg/day PERSERIS: 1 injection per 28 days quetiapine <13 years of age: 400mg/day quetiapine <13 years of age: 600mg/day quetiapine =17 years of age: 600mg/day risperidone <10 years of age: 80mg/day risperidone <10 years of age: 6mg/day risperidone <10 years of age: 6mg/day risperidone >17 years of age: 16mg/day RISPERDAL CONSTA: 2 injections per 28 days SAPHRIS: 20mg/day	
			LATUDA >17 years of age: 160mg/day olanzapine <13 years of age: 10mg/day olanzapine ≥13 years of age: 20mg/day paliperidone: 12mg/day PERSERIS: 11njection per 28 days quetiapine <13 years of age: 400mg/day quetiapine <13 years of age: 600mg/day quetiapine >17 years of age: 800mg/day quetiapine >17 years of age: 800mg/day risperidone <10 years of age: 3mg/day risperidone <17 years of age: 6mg/day risperidone >17 years of age: 16mg/day RISPEROLA CONSTA: 2 injections per 28 days	
			LATUDA >17 years of age: 160mg/day olanzapine <13 years of age: 10mg/day olanzapine >13 years of age: 20mg/day paliperidone: 12mg/day PERSERIS: 1 injection per 28 days quetiapine <13 years of age: 400mg/day quetiapine <13 years of age: 600mg/day quetiapine 13-17 years of age: 800mg/day risperidone <10 years of age: 3mg/day risperidone <10 years of age: 6mg/day risperidone <10 years of age: 6mg/day RISPERDAL CONSTA: 2 injections per 28 days SAPHRIS: 20mg/day ziprasidone <17 years of age: 120mg/day ziprasidone <17 years of age: 120mg/day	

THED ADELITIC CLASS		PREFERRED AGENTS REQUIRING	CHNICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIES	
THERAPEUTIC CLASS	PREFERRED AGENTS	CLINICAL CRITERIA	CLINICAL CRITERIA	THIS LIST IS NOT ALL INCLUSIVE PLEASE CONTACT CHANGE HEALTHCARE WITH ANY QUEST	
TAL HEALTH	AMPHETAN		Clients over the age of 17 must have a diagnosis for ADD, ADHD, (see ADD/ADHD	AMPHETAMINES	
tinued	LONG ACTING AME	PHETAMINES ADDERALL XR	criteria below), narcolepsy, obstructive sleep apnea, shift work sleep disturbance,	ADZENYS XR ODT DYANAVEL XR	
		amphetamine salts combo XR	MS fatigue (see MS Fatigue criteria below), or refractory depression (see refractory depression criteria below).	EVEKEO/ODT	
		dextroamphetamine CR caps	depression enteria belowy.	MYDAYIS	
		VYVANSE CAPSULES**		PROCENTRA	
	IMMEDIATE RELEASE A	AMPHETAMINES	For clients over the age of 17, they must meet the DSM-5 criteria for diagnosis of	VYVANSE CHEWABLES ZENZEDI 2.5 AND 7.5MG TABLETS	
		amphetamine salts combo	ADHD. These criteria include:		
		dextroamphetamine tablets	Five or more symptoms of inattention, present for at least 6 months,		
	METHYLPHEN LONG ACTING METHY		inappropriate for developmental level.	METHYLPHENIDATES APTENSIO XR	
		CONCERTA*	OR	AZSTARYS	
		dexmethylphenidate ER	 Five or more symptoms of hyperactivity and impulsivity, present for at least 6 months, to an extent that is disruptive and inappropriate for developmental level. 	COTEMPLA XR	
		methylphenidate ER tablets	AND	DAYTRANA FOCALIN XR	
		metry premate Extables	Symptoms must be present in two or more settings (home, school or work); and	JORNAY PM	
			There must be clear evidence that the symptoms interfere or reduce the quality of social, school or work functioning; and	methylphenidate ER osmotic release	
	IMMEDIATE RELEASE ME	THYLPHENIDATES dexmethylphenidate	The symptoms must not be better explained by another mental disorder.	(BRAND IS PREFERRED) methylphenidate ER/CR/SR capsules	
		methylphenidate chewables		(METADATE CD/RITALIN LA, APTENSIO	
		methylphenidate solution		QUILLICHEW ER	
		methylphenidate tablets		QUILLIVANT	
			Diagnosis of MS fatigue 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 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.		
			Prior Authorization will be required for clients under the age of 4.		
			·		
			**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 approval.		
			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.		
			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: 60mg/day		
			amphetamine salts combo: 60mg/day amphetamine salts combo (narcolepsy): 90mg/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		
			FOCALIN XR > 13 years of age: 60mg/day JORNAY PM: 100mg/day		
			methylin/methylphenidate/ER: 90mg/day		
			VYVANSE: 105mg/day		
	SELECTIVE ALPHA-ADRE	NERGIC AGONIST	To obtain the non-preferred agent, client must meet the following criteria:	clonidine ER	
	clonidine		Client must have a diagnosis of ADD or ADUD		
			Client must have a diagnosis of ADD or ADHD		
			Prior authorization will be required for clients under the age of 4.		
			To receive clonidine ER, clients must have completed a 14 day trial of clonidine IR		
			with benefit in the previous 12 months.		
	1	l .		i	

THERAPEUTIC CLASS PREFERRED AGENTS PREFERRED AGENTS REQUIRING			CLINICAL CRITERIA	NON-PREFERRED AGENTS
THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	GENERIC MANDATORY POLICY APPLIES THIS USY IS NOT ALL INCLUSIVE PLEASE CONTACT CHANGE HEALTHCARE WITH ANY QUESTIC
ENTAL HEALTH continued	SELECTIVE NOREPINEPHRINE	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	
MIGRAINE	MIGRAINE PROI	PHYLAXIS	Trial and failure of both an anticonvulsant and a beta blocker (Step 1 agents)	NURTEC
	STEP 1 AGI	NTS	greater than or 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. Nurtec	
	beta blockers	divalproex topiramate	will be limited to 16 tabs/30 days.	
	STEP 2 AGI	AIMOVIG*	*Starting dose will be limited to 70mg **Approval for non-preferred agents requires trial and failure of a preferred agent	QULIPTA**
		AJOVY	along with the trial and failures described with Step 1 Agents' criteria above.	
	ACUTE MIGRAINE	EMGALITY TREATMENT		
	STEP 1 AGI		Trial and failure of two preferred agents will be required for approval of a non-	almotriptan
	frovatriptan naratriptan RELPAX* sumatriotan		preferred agent. Rizatriptan will be limited to clients 6 years of age or older	ELYXYB Sumatriptan-Naproxen Sodium TOSYMRA (use preferred agent) TROKENDI XR
	rizatriptan		Quantity limits apply:	ZEMBRACE (use preferred agent) zolmitriptan
			naratriptan 1mg: 25 tabs/34 days 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 sumatriptan vials: 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	
			sumatriptan 100mg: 10 tabs/34 days	
	STEP 2 AGE	NTS	Trial and failure of two triptan agents is required for approval of a Step 2 Agent.	REYVOW
		NURTEC		UBRELVY
			Trial and failure of two preferred triptan agents AND Nurtec will be required for approval of a non-preferred agent.	
			Quantity limits apply:	
			NURTEC 75mg: limited to 8 tabs/30 days REYVOW: 200mg/day or 1 tab/day, 4 tab/30 days	
OVEMENT DISORDERS	VMAT 2 INF	HIBITORS	Quantity limits apply:	
	AUSTEDO/XR* INGREZZA*		AUSTEDO: limited to 4 tabs/day INGREZZA: limited to 4 tabs/day	
	TETRABENAZINE		*Please refer to the Additional Therapeutic Criteria chart at www.wymedicaid.org for specific requirements.	
IULTIPLE SCLEROSIS	MS AGEN		Kesimpta, Lemtrada, Ocrevus, and Tysabri will be approved for highly active	AUBAGIO
	AVONEX BETASERON	GILENYA KESIMPTA	disease, please refer to the ATCC for additional information.	BAFIERTAM BRIUMVI
	COPAXONE 20MG/ML* dimethyl fumarate	LEMTRADA OCREVUS	Trial and failure of two preferred agents for at least 56 days (each from a separate	EXTAVIA glatiramer (BRAND IS PREFERRED)
	REBIF	TYSABRI	class) will be required before approval can be given for a non-preferred agent.	GLATOPA (use preferred agent)
	teriflunomide VUMERITY			MAVENCLAD MAYZENT
			For Mavenclad, in addition to the above criteria, approval will be granted on a case-	
			by-case basis.	PONVORY TECFIDERA ZEPOSIA
ARCOLEPSY	CTIMILII ANITC		Modefinil and Nurisil Client must be a 15 cores of a 15 cores	EL SSIA
MANCOLEPST	STIMULANTS	modafinil	Modafinil and Nuvigil: Client must be > 16 years of age. Client must have a diagnosis of narcolepsy, obstructive sleep apnea, shift work sleep disorder, Multiple Sclerosis	
		NUVIGIL*	(MS) Fatigue, or ADD/ADHD with a concurrent diagnosis of substance abuse. Diagnosis of MS will require a fatigue severity scale score of 5.0, a 60 day trial of	
	NON-STIMULANTS		amantadine AND discontinuation of medications that may contribute to drowsiness or fatigue.	SUNOSI
			Clients will not be allowed to take two or more agents in this class concurrently	WAKIX XYREM
			sector with not be anowed to take two or more agents in this class concurrently	principal (
EUROPATHIC PAIN	GABAPEN		Clients will not be allowed to take gabapentin and pregabalin concurrently	
		gabapentin pregabalin	Prior authorizations for perioperative pain will be approved for gabapentin OR	
			pregabalin for less than or equal to 14 day supplies	
	TOPICAL LIDO Lidocaine Patches			ZTLIDO
	ADDITIONAL / amitriptyline	AGENTS	Trial and failure of a tricyclic antidepressant greater than or equal to a 12 week	carbamazepine imipramine (capsules)
	desipramine		supply AND trial and failure of gabapentin at a dose of 3600mg per day OR	oxcarbazepine
	imipramine (tablets) nortriptyline		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.	valproic acid
	nor a ipcynnic	İ		İ

		PREFERRED AGENTS REQUIRING		NON-PREFERRED AGENTS
THERAPEUTIC CLASS	PREFERRED AGENTS	CLINICAL CRITERIA	CLINICAL CRITERIA	GENERIC MANDATORY POLICY APPLIE: THIS LIST IS NOT ALL INCLUSIVE PLEASE CONTACT CHANGE HEALTHCARE WITH ANY QUESTIX
HTHALMICS	OPANTI-ALL	ERGICS	Trial and failure of a preferred agent greater than or equal to 30 days in the last 12	ALOCRIL ALOMIDE
	azelastine		months will be required before approval can be given for a non-preferred agent.	bepotastine
	BEPREVE*		Alomide and Alocril will be approved for pregnancy.	epinastine
	cromolyn 0.4%		Alomide will be approved for children under the age of 3.	ZERVIATE
	OPANTIBIOTICS- C	QUINOLONES	Trial and failure of a preferred agent greater than or equal to 5 days in the last 12 months will be required before approval can be given for a non-preferred agent.	gatifloxacin ZYMAXID
	BESIVANCE gentamicin			
	moxifloxacin 0.5% ofloxacin			
	tobramycin OPANTI-INFLAN	MMATORY	Trial and failure of ALL preferred agents each greater than or equal to 5 day supply	ACULAR/LS/PF (use preferred agent)
	flurbiprofen		in the last 12 months will be required before approval can be given for a non-	ACUVAIL
	diclofenac LOTEMAX*		preferred agent.	bromfenac 0.9% BROMSITE
	ketorolac			DUREZOL
	NEVANAC			ILEVRO INVELTYS
				LOTEMAX SM
				loteprednol 0.5% (BRAND PREFERRED) PROLENSA
	OPBETA-BLC	OCKERS		BETIMOL
	betaxolol carteolol		in the last 12 months will be required before approval can be given for a non- preferred agent.	BETOPTIC S*
	levobunolol timolol		*Betoptic S will be approved for those with heart and lung conditions.	
	OPCARBONIC ANHYD	RASE INHIBITOR	Trial and failure of a preferred agent greater than or equal to 30 days in the last 12	brinzolamide (BRAND PREFERRED)
	A7OPT dorzolamide		months will be required before approval can be given for a non-preferred agent.	
	OPCOMBO PR	ODUCTS		dorzolamide/timolol (BRAND PREFERRED)
	COSOPT* ROCKLATAN			
	SIMBRINZA OPDRY EYE A	AGENTS	Trial and failure of the preferred agent greater than or equal to 12 weeks will be	CEQUA
	RESTASIS* XIIDRA		required before approval can be given for the non-preferred agent.	cyclosporine (BRAND PREFERRED) EYSUVIS MIEBO RESTASIS MULTIDOSE (use preferred agent TYRVAYA
	OPPROSTAGL	ANDINS	Trial and failure of ALL 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-	bimatoprost IYUZEH
	LUMIGAN TRAVATAN Z XALATAN ZIOPTAN		preferred agent.	tafluprost
	OPRHO KINASE	INHIBITOR		
	RHOPRESSA			
	OPSYMPATHOI ALPHAGAN P 0.1% ALPHAGAN P 0.15%*	MIMETICS	Trial 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.	brimonidine 0.15% (BRAND IS PREFERRED)
TEOPOROSIS	brimonidine 0.2% BISPHOSPHO	NATES	Trial and failure of a preferred agent greater than or equal to 12 months will be	EVENITY**
	alendronate ibandronate		required before approval can be given for a non-preferred agent.	FORTEO*** FOSAMAX-D
	risedronate		Fosamax liquid will be approved for clients that have difficulty swallowing.	TYMLOS***
			**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	
			***Will be limited to 2 years of use	
	NASAL CALCI calcitonin-salmon	TONIN		
С	ANTIBIOTIC/STEROID	COMBINATION		ciprofloxacin 0.2% (use preferred agent)
	ciprofloxacin/dexamethasone Neo/Poly/HC Suspension and Solution			CIPRO HC (use preferred agent) CORTISPORIN-TC (use preferred agent)
	ofloxacin tobramycin/dexamethasone			FLUOCINOLONE ACET OIL 0.01% (use preferred agent)
ERACTIVE BLADDER	OVERACTIVE BLADI	DER AGENTS	Trial and failure of a preferred agent greater than or equal to a 14 day supply in the	darifenacin GELNIQUE GEL 10%
	oxybutynin /ER		last 12 months will be required before approval can be given for a non-preferred agent.	GEMTESA
	solifenacin			OXYTROL DIS
	TOVIAZ		Oxytrol will be approved for clients that have an inability to swallow.	tolterodine/ER trospium

Please refer to the Additional Therapeutic Criteria Chart, Dosage Limitation List (red font indicates quantity/dosage limits apply), and the Wyoming Medicaid Provider Manual at http://wymedicaid.org 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 PLEASE CONTACT CHANGE HEALTHCASE WITH ANY QUESTION.		
ZAIN	LONG-ACTIN morphine ER <u>tablets</u>	G C-IIs	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.	fentanyi patches hydrocodone ER hydromorphone ER HYSINGLA ER METHADONE		
			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).	morphine ER capsules (use preferred agents) NUCYNTA ER** oxymorphone ER OXYCONTIN		
			Concurrent therapy with a benzodiazepine and a narcotic medication or with duplicate benzodiazepines is not cowered by Wyoming Medicaid. A single medication will continue to be allowed to process unless another benzodiazepine (or narcotic) is billed to Wyoming Medicaid.	XTAMPZA ER (additional criteria applies)		
			**Nucynta ER will be allowed for diabetic peripheral neuropathy or clients with significant gastrointestinal concerns with other CII narcotics.			
			Fentanyl: 50mcg, 1 strength at a time, 1 patch every 3 days Hydromorphone ER: 30mg/day Hysingla ER: 120mg/day Methadone: Limited to 3 tablets per day Morphine ER: 90mg/day			
			Nucynta ER: 327mg/day Oxycontin: 80mg/day Oxymorphone: 40mg/day Xtampza ER: 80mg/day			
			Clients will be limited to one long-acting narcotic at a time			
	short-Actin codeine sulfate hydrocodone/APAP hydrocodone/IBU hydromorphone	G C-lls	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 NUCYNTA* oxymorphone ROXYBOND		
	meperidine morphine oxycodone oxycodone/APAP		*Nucynta will be allowed for diabetic peripheral neuropathy or clients with significant gastrointestinal concerns with other CII narcotics.			
	5A, 55C 6A, 7. 1. 1.		Concurrent therapy with a benzodiazepine and a narcotic medication or with duplicate benzodiazepines is not cowered 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-V AG BUTRANS tramadol	ENTS	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.			
ARKINSON'S DISEASE	SHORT-ACTING amantadine	AGENTS				
	benztropine tablets carbidopa/levodopa pramipexole ropinirole					
	LONG-ACTING ropinirole ER RYTARY	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		
HOSPHATE BINDERS	PHOSPHATE B	INDERS	Prior authorization required for non-preferred agents.	AURYXIA		
	calcium acetate			lanthanum sevelamer VELPHORO		

Manual at http://wymedicaid.org 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 PLEASE CONTACT GHANGE HEALTHCARE WITH ANY QUESTION:			
PROSTATE	5-ALPHA-REDUCTAS finasteride	E 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 (use separate agents)			
	ALPHA BLOC doxazosin tamsulosin terazosin	KERS	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 (use separate agents) silodosin			
PULMONARY ANTIHYPERTENSIVES	5-ALPHA-REDUCTAS	E INHIBITORS ALYQ tadalafil REVATIO SUSPENSION* sildenafil (Revatio A/B rated generic)	Prior authorization required. Client must have a diagnosis of pulmonary hypertension.	sildenafil suspension (BRAND IS PREFERRED)			
	ENDOTHELIN RECEPTO	R 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 CYCLAS	E INHIBITORS	Prior authorization required.	ADEMPAS (use preferred agent)			
	PROSTACYCLINE VA	SODILATORS ORENITRAM	Prior authorization required. Client must have a diagnosis of pulmonary hypertension with documented right-heart catheterization validating the diagnosis.				
	PROSTACYCLINE RECE	PTOR AGONIST	Prior authorization required.	UPTRAVI (use preferred agent)			
ESTLESS LEG SYNDROME	RESTLESS LEGSY pramipexole ropinirole	NDROME gabapentin	Client must have a diagnosis of Restless Leg Syndrome (RLS). Trial and failure of gabapentin greater than or equal to 60 days and 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				
KELETAL MUSCLE RELAXANTS	MUSCLE RELA baclofen (tablets) cyclobenzaprine tizanidine tablets	XANTS	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. Cyclobenzaprine will require a prior authorization for clients concurrently taking a tricylic antidepressant. Carisoprodol is limited to 84 tabs/365 days	cartsproded chlorzoxazone cyclobenzaprine ER LVVISPAH metaxalone methocarbamol orohenadrine tizanidine capsules (use preferred agent)			
ICERATIVE COLITIS	IMMUNOMODI	JLATORS 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 SIMPONI STELARA			
			* Refer to Additional Therapeutics Clinical Criteria Chart for more information	XELJANZ/XR			
IVEITIS	IMMUNOMODU	HUMIRA	Client must have diagnosis of non-infectious intermediate, posterior, or panuveitis in adult patients				