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.

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 OptumRx 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/dose limits apply), and Wyoming Medicaid Provider 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 PLEASE CONTACT OPLIMITAL WITH ANY QUESTIONS
ADDICTION	BUPRENORPHINE	COMBINATIONS	Client must have a diagnosis of opioid dependence or abuse. This is not to be used for the	buprenorphine (oral)
		buprenorphine/naloxone tablets	treatment of chronic pain. Prior authorization will be required before any narcotic,	buprenorphine/naloxone film
		SUBOXONE FILM*	benzodiazepine, or carisoprodol prescription will be allowed between fills. Prior authorization	BRAND IS PREFERRED)
			will be required before any short-acting stimulant prescription from any doctor other than the	ZUBSOLV
			prescriber of buprenorphine or Suboxone, will be allowed between fills.	
			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.	
			To the available at www.wymedicaid.org.	
			Dosage limits apply Prior authorization will be required for doses >24mg	
	NAL	OXONE	Kloxxado, naloxone products, and Narcan nasal spray will be limited to one fill per 180 days	OPVEE
	KLOXXADO		without prior authorization.	REXTOVY
	naloxone nasal spray NARCAN		Naloxone formulations available in quantities of 10ml will require prior authorization.	ZIMHI
	NALT	REXONE	Client must have a diagnosis of alcohol or opioid dependance.	topiramate*
		naltrexone VIVITROL		
	ĺ		Prior authorization will be required before any narcotic, carisoprodol, or benzodiazepine	
	ĺ		prescription will be allowed between fills. Prior authorization will be required before a short-	
	ĺ		acting stimulant prescription from any doctor other than the prescriber of naltrexone or Vivitrol will be allowed between fills.	ĺ
	ĺ		111111111111111111111111111111111111111	
			*Topiramate requires 4 week trial and failure of naltrexone or acamprosate in AUD	
LLERGY / ASTHMA / COPD		, MINIMALLY SEDATING	Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12	desloratadine
	cetirizine fexofenadine		months will be required before approval can be given for a non-preferred agent.	CLARINEX RDT/SYRUP levocetirizine
	loratadine			
	ANTIHISTAMINE/DECO	NGESTANT COMBINATIONS	Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12	CLARINEX-D
	cetirizine/pseudoephedrine		months will be required before approval can be given for a non-preferred agent.	
	fexofenadine/pseudoephedrine			
	loratadine/pseudoephedrine			
	ANTICHOLINERGI ATROVENT HFA	C BRONCHODILATORS	Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will	TIOTROPIUM BROM (use brand)
	INCRUSE ELLIPTA		be required before approval can be given for a non-preferred agent.	TUDORZA YUPELRI
	ipratropium			TOT EEM
	SPIRIVA HANDIHALER			
	SPIRIVA RESPIMAT		Spiriva 5 day STARTER package will be allowed one (1) time per recipient.	
		OMBINATION AGENTS	Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will	BEVESPI
	ANORO ELLIPTA** COMBIVENT		be required before approval can be given for a non-preferred agent.	BREZTRI DUAKLIR
	STIOLTO			TRELEGY
			**Will also require the diagnosis of COPD.	
		NE MODIFIERS	Trial and failure of preferred agent greater than or equal to 30 days in the last 12 months will be	zafirlukast
	montelukast		required before approval can be given for a non-preferred agent.	
	LONG ACTING BR	ONCHODILATORS	Trial and failure of two (2) preferred agents greater than or equal to 30 days in the last 12	BROVANA
	arformoterol		months will be required before approval can be given for a non-preferred agent.	
	SEREVENT STRIVERDI			ĺ
		TIHISTAMINES	Trial and failure of preferred agent greater than or equal to 90 days in the last 12 months will be	azelastine 0.15%
	azelastine 0.1%		required before approval can be given for a non-preferred agent.	DYMISTA (use separate agents)
	ĺ		and the second second second second second second	olopatadine 0.6%
				RYALTRIS
		STEROIDS	Trial and failure of two (2) preferred agents greater than or equal to 30 days in the last 12	DYMISTA (use separate agents)
	budesonide flunisolide		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 BRO	NCHODILATORS - INHALERS	Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will	levalbuterol (BRAND IS PREFERRED)
	albuterol HFA		be required before approval can be given for a non-preferred agent. Prior authorization will be	PROAIR DIGIHALER
	PROAIR RESPICLICK		required after a total of 12 albuterol inhalers are dispensed within 365 days.	PROVENTIL HFA
	VENTOLIN HFA		Minimum day supply of 16 days is required.	
	STEROID	INHALANTS	Trial and failure of two (2) preferred agents greater than or equal to 30 days in the last 12	AIRDUO DIGIHALER
	AIRDUO RESPICLICK		months will be required before approval can be given for a non-preferred agent.	ALVESCO
	ARNUITY ELLIPTA		*Fluticasone HFA and Asmanex HFA will be approved for pediatric clients 8 years of age or	ARMONAIR
	ASMANEX TWISTHALER		younger.	ASMANEX HFA*
	budesonide suspension PULMICORT FLEXHALER		Alvesco will be approved for a history of oral thrush with steroid inhalants.	fluticasone HFA* QVAR REDIHALER
		BINATION AGENTS	Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will	QVAR REDIHALER fluticasone/vilanterol (use preferred agent)
	ADVAIR (HFA, Diskus)	BINAHON AGENTS	be required before approval can be given for a non-preferred agent.	fluticasone/vilanterol (use preferred agent) fluticasone/salmeterol 55-14/113-14/232-14
	BREO ELLIPTA**		be required before approval can be given for a non-preferred agent.	fluticasone/salmeterol 100-50/250-50/500-5
	DULERA			(BRAND IS PREFERRED)
	SYMBICORT*		**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.	WIXELA
		PHRINE		AUVI-Q (use preferred agent)
	epinephrine auto-injector pen			
	EPI-PEN FOSINORHILIC	ASTHMA AGENTS	*Approval for these agents will require additional clinical criteria which can be found on the	FASENRA*
	EOSINOPHILIC	DUPIXENT	Approval for these agents will require additional clinical criteria which can be found on the	PASENRA* NUCALA*
	ĺ	XOLAIR	Additional Therapeutic Citiena Chart	TEZSPIRE

THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPL THIS LIST IS NOT ALL INCLUSIVE PLEASE CONTACT Optumins WITH ANY QUESTION
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 a diagnosis of AS and a 56-day trial and failure of both	CIMZIA** COSENTYX
		ENBREL	nreferred agents	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	
ļ	ILIVENII E IDIOPA	ATHIC ARTHRITIS (JIA)	Client must have diagnosis of JIA prior to approval of a preferred agent. To receive a non-	ACTEMRA
	70741114-12107	ENBREL HUMIRA	preferred agent, client must have a diagnosis of JIA and a 56-day trial and failure of both preferred agents.	ILARIS ORENCIA
	PSORIATIC	ARTHRITIS (PA)	Client must have diagnosis of PA prior to approval of a preferred agent. To receive a non-	XELJANZ/XR CIMZIA**
ļ		ENBREL HUMIRA	preferred agent, client must have a diagnosis of PA and a 56-day trial and failure of two of the	COSENTYX ORENCIA
ļ		OTEZLA*	three preferred agents.	REMICADE
ļ		TALTZ		RINVOQ
ļ			*Otezla starter pack is non-preferred	SIMPONI STELARA
ļ			Ocean Starter pack is non-preferred	TREMFYA
ļ			**Cimzia will be allowed for clients that are pregnant or breast-feeding	XELJANZ/XR
l	RHEUMATOII	ARTHRITIS (RA) ENBREL	Client must have diagnosis of RA and a 56-day trial and failure of methotrexate prior to approval of a preferred agent. To receive a non-preferred agent, client must have a diagnosis of RA and a	
ļ		HUMIRA	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.	KEVZARA
ļ				KINERET
l		Ī		OLUMIANT ORENCIA
ļ			*Cimzia will be allowed for clients that are pregnant or breast-feeding	REMICADE
l		Ī	**Coo Dormatology critoria for Atonia Dormatikia anavorral	RINVOQ**
ļ			**See Dermatology criteria for Atopic Dermatitis approval	RITUXAN SIMPONI
				XELJANZ/XR
/ULSIONS		REOTYPIC SEIZURE EPISODES	*Nayzilam will be allowed for patients 12 years of age and older	
l	diazepam gel NAYZILAM*	Ī		
l	VALTOCO	<u> </u>		<u> </u>
ļ		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**
ļ	FELBAMATE	EPIDIOLEX	agents prior to approvai.	DIACOMIT**
ļ	fosphenytoin	FYCOMPA	For indications not related to seizures and epilepsy, please refer to the Additional Therapeutic	FINTEPLA**
ļ	lacosamide (tablets) lamotrigine/XR	gabapentin pregabalin	Criteria chart at www.wymedicaid.org.	levetiracetam ER LIBERVANT
ļ	levetiracetam	topiramate/ER sprinkle caps		OXTELLAR
ļ	oxcarbazepine		**Please refer to the Additional Therapeutic Criteria chart at www.wymedicaid.org for specific	TROKENDI XR XCOPRI
ļ	phenytoin subvenite		requirements.	VIMPAT (tablets)
	valproate/valproic acid			zonisamide oral susp.
	VIMPAT (suspension) zonisamide			
IN'S		MODULATORS	Client must have diagnosis of Crohn's prior to approval of the preferred agent. To receive a non-	
ļ		HUMIRA	preferred agent, client must have a diagnosis of Crohn's and a 56-day trial and failure of the	ENTYVIO* REMICADE
			preferred agent.	RINVOQ
ļ			* Refer to Additional Therapeutics Clinical Criteria Chart for more info	SKYRIZI
			**Cimzia will be allowed for clients that are pregnant or breast-feeding	STELARA TYSABRI (additional criteria applies)
MATOLOGY	BENZOYL PEROXIDE	/CLINDAMYCIN COMBOs	Clients must be 12 to 20 years of age. Requires prior authorization for clients less than 12 years	ACANYA
ļ		clindamycin/benzoyl peroxide 1-5%	of age. Acne combinations are limited to clients under the age of 21.	ONEXTON
ļ		clindamycyin/benzoyl peroxide 1.2-5% (Refrig)		
	ISOT	1.2-5% (Retrig) RETINOIN	Clients must be 12 to 20 years of age.	ABSORICA
İ]	
	AMNESTEEM			
	CLARAVIS			
	CLARAVIS isotretinoin ZENATANE			
	CLARAVIS isotretinoin ZENATANE CORTICOSTERO	DS - STEP 1 AGENTS		
	CLARAVIS isotretinoin ZENATANE CORTICOSTERO C=CREAM; G=GEL; L=LOTIC	N; O=OINTMENT; S=SOLUTION	Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days	PANDEL
	CLARAVIS isotretinoin ZENATANE CORTICOSTERO C=CREAM; G=GEL; L=LOTIC LOW alclometasone		Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days.	PANDEL TEXACORT 2.5% (S)
	CLARAVIS isotretinoin ZENATANE CORTICOSTERO C=CREAM; G=GEL; L=LOTIC LOW alclometasone desonide*	N; O=OINTMENT; S=SOLUTION		
	CLARAVIS isotretinoin ZENATANE CORTICOSTERO C=CREAM; G=GEL; L=LOTIC LOW alclometasone	N; O=OINTMENT; S=SOLUTION	Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days. *Cream, ointment, and lotion formulations of desonide are preferred.	
	CLARAVIS isotretinoin ZENATANE CORTICOSTERO C=CREAM; G=GEL; L=LOTIC IOW alclometasone desonide* fluocinolone 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone 1%, 2.5% (C,LO)	N; O=OINTMENT; S=SOLUTION POTENCY	*Cream, ointment, and lotion formulations of desonide are preferred.	TEXACORT 2.5% (S)
	CLARAVIS isotretinoin ZENATANE CORTICOSTERO C=CREAM; G=GEI; L=LOTIC LOW alclometasone desonide* fluccinolnoe 0.01% hydrocortisone butvrate 0.1% (C, hydrocortisone 1%, 2.5% (C,L,O) MEDIU	N; O=OINTMENT; S=SOLUTION		TEXACORT 2.5% (S) Clocortolone Pivalate
	CLARAVIS isotretinoin ZENATANE CORTICOSTERO C=CREAM; G=GEL; L=LOTIC IOW alclometasone desonide* fluocinolone 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone 1%, 2.5% (C,LO)	N; O=OINTMENT; S=SOLUTION POTENCY	*Cream, ointment, and lotion formulations of desonide are preferred.	TEXACORT 2.5% (S)
	CLARAVIS isotretinoin ZENATANE CORTICOSTERO C=CREAM; G=GEI; L=LOTIC LOW alciometasone desonide* fluocinolone 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone 1%, 2.5% (C,L.O) MEDIU betamethasone valerate desoximetasone 0.05%, 0.25% (C) fluocinolone 0.025%	N; O=OINTMENT; S=SOLUTION POTENCY	*Cream, ointment, and lotion formulations of desonide are preferred.	TEXACORT 2.5% (5) Clocortolone Pivalate flurandrenol fluticasone 0.05% (L) hydrocortisone butyrate 0.1% (O)
	CLARAVIS isotretinoin ZENATANE CORTICOSTERO C=CREAM; G=GEL; L=LOTIC LOW alclometasone desonide* fluocinolone 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone butyrate 0.1% (C, L, O) MEDIU betamethasone valerate desoximetasone 0.05%, 0.25% (C) fluocinolone 0.025% (T) fluiciasone 0.05% (C)	N; O=OINTMENT; S=SOLUTION POTENCY	*Cream, ointment, and lotion formulations of desonide are preferred.	TEXACORT 2.5% (S) Clocortolone Pivalate flurandrenol fluticasone 0.05% (L)
	CLARAVIS isotretinoin ZENATANE CORTICOSTERO CECREAM; GEGEL; LELOTIC LOW alclometasone desonide* fluocinolone 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone butyrate 0.1% (C) hydrocortisone volume on the company of the co	N; O=OINTMENT; S=SOLUTION POTENCY	*Cream, ointment, and lotion formulations of desonide are preferred.	TEXACORT 2.5% (5) Clocortolone Pivalate flurandrenol fluticasone 0.05% (L) hydrocortisone butyrate 0.1% (O)
	CLARAVIS isotretinoin ZENATANE CORTICOSTERO C=CREAM; G=GEL; L=LOTIC LOW alciometasone desonide* fluocinolone 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone butyrate 0.1% (C) hydrocortisone 1%, 2.5% (C,L,O) MEDIU betamethasone valerate desoximetasone 0.05%, 0.25% (C) fluocinolone 0.025% (Iluocinolone 0.025% (C) onometasone SYNALAR 0.025% (C, O) triamcinolone 0.025%, 0.1%	M. O-COINTMENT; S=SOLUTION POTENCY M. POTENCY	*Cream, ointment, and lotion formulations of desonide are preferred. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days.	Clocortolone Pivalate flurandrenol flutiasone 0.05% (L) hydrocortisone butyrate 0.1% (O) triamcinolone 0.05% (O)
	CLARAVIS isotretinoin ZENATANE CCREAM; G=GEI; L=LOTIC LOW alciometasone desonide* fluocinolone 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone butyrate 0.1% (C) hydrocortisone valerate desoximetasone valerate desoximetasone 0.05% (C) fluocinolone 0.025% (C) fluorinoine 0.025% (C) mometasone SYNALAR 0.025% (C, O) triamcinolone 0.025%, 0.1%	N; O=OINTMENT; S=SOLUTION POTENCY	*Cream, ointment, and lotion formulations of desonide are preferred.	TEXACORT 2.5% (5) Clocortolone Pivalate flurandrenol fluticasone 0.05% (L) hydrocortisone butyrate 0.1% (O) triamcinolone 0.05% (O) APEXICON 0.05% (C)
	CLARAVIS isotretinoin ZENATANE C-CREAM; G-GEI; LI-OTIC LOW alciometasone desonide* fluorinoine 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone butyrate 0.1% (C) hydrocortisone butyrate 0.1% (C) hydrocortisone 0.05% (C, LO) betamethasone valerate desoximetasone 0.05% (C) fluorinoine 0.025% (L) guidentinoine 0.05% (C) mometasone SYNALAR 0.025% (C, O) triamcinolone 0.025% (L) hydrocortisone 0.05% (C) hydrocortiso	M. O-COINTMENT; S=SOLUTION POTENCY M. POTENCY	*Cream, ointment, and lotion formulations of desonide are preferred. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days.	TEXACORT 2.5% (S) Clocortolone Pivalate flurandrenol fluticasone 0.05% (L) hydrocortisone butyrate 0.1% (O) triamcinolone 0.05% (C) APEXICON 0.05% (C) amcinonide 0.1% (C, L, O) augmented betamethasone 0.05% (G, L, augmented betamethasone 0.05% (G, L,
	CLARAVIS isotretinoin ZENATANE CORTICOSTERO C=CREAM; G=GEL; L=LOTIC LOW alcometasone desonide* fluocinolone 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone butyrate 0.1% (C) hydrocortisone 1%, 2.5% (C,L,O) MEDIU betamethasone valerate desoximetasone 0.05%, 0.25% (C) fluocinolone 0.025% fluticasone 0.05% (C) mometasone SYNALAR 0.025% (C, O) triamcinolone 0.025%, 0.1% HIGH betamethason/E 0.05% (C, G,O,S) diflorasone 0.05% (O)	M. O-COINTMENT; S=SOLUTION POTENCY M. POTENCY	*Cream, ointment, and lotion formulations of desonide are preferred. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days.	TEXACORT 2.5% (\$) Clocortolone Pivalate flurandrenol fluticasone 0.05% (L) hydrocortisone butyrate 0.1% (O) triamcinolone 0.05% (C) amcinonide 0.1% (C,L,O) augmented betamethasone 0.05% (G,L) clobetasol 0.05% (L)
	CLARAVIS isotretinoin ZENATANE CORTICOSTERO C=CREAM; G=GEL; L=LOTIC LOW alciometasone desonide* fluocinolone 0.01% hydrocritisone butyrate 0.1% (C) hydrocritisone butyrate 0.1% (C) hydrocritisone 1%, 2.5% (C,L,O) MEDIU betamethasone valerate desoximetasone 0.05% (D, 2.5% (C) fluocinolone 0.025% (C) mometasone SYNALAR 0.025% (C, O) triamcinolone 0.025% (C, G) triamcinolone 0.025% (C, G) diflorasone 0.05% (C) fluocinoide desoximetasone 0.05% (C, G,O,S) diflorasone 0.05% (O) fluocinoide fluocinoide	M. O-COINTMENT; S=SOLUTION POTENCY M. POTENCY	*Cream, ointment, and lotion formulations of desonide are preferred. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days.	Clocortolone Pivalate flurandrenol flutiasone 0.05% (L) hydrocortisone butyrate 0.1% (O) triamcinolone 0.05% (C) APEXICON 0.05% (C) amcinonide 0.1% (C,L,O) augmented betamethasone 0.05% (G,L,I clobetasol 0.05% (L) desoximetasone 0.05%, 0.25% (G,O)
	CLARAVIS isotretinoin ZENATANE CORTICOSTERO C=CREAM; G=GEL; L=LOTIC LOW alcometasone desonide* fluocinolone 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone butyrate 0.1% (C) hydrocortisone 1%, 2.5% (C,L,O) MEDIU betamethasone valerate desoximetasone 0.05%, 0.25% (C) fluocinolone 0.025% fluticasone 0.05% (C) mometasone SYNALAR 0.025% (C, O) triamcinolone 0.025%, 0.1% HIGH betamethason/E 0.05% (C, G,O,S) diflorasone 0.05% (O)	M. O-COINTMENT; S=SOLUTION POTENCY M. POTENCY	*Cream, ointment, and lotion formulations of desonide are preferred. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days.	TEXACORT 2.5% (5) Clocortolone Pivalate flurandrenol fluticasone 0.05% (L) hydrocortisone butyrate 0.1% (0) triamcinolone 0.05% (O) APEXICON 0.05% (C) amcinonide 0.1% (C,L,O) augmented betamethasone 0.05% (G,L,ciobetasol 0.05% (L)
	CLARAVIS isotretinoin 2ENATANE CCREAM; G=GEI; L=LOTIC 10W alciometasone desonide* fluorinoine 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone butyrate 0.1% (C) hydrocortisone 1%, 2.5% (C,LO) MEDIU betamethasone valerate desoximetasone 0.05%, 0.25% (C) fluorinoine 0.025% (C) mometasone SYNALAR 0.025% (C, O) triamcinoione 0.05% (C) G,O,S) diflorasone 0.05% (O) fluorinoide fluorinoi	M. O-COINTMENT; S=SOLUTION POTENCY M. POTENCY	*Cream, ointment, and lotion formulations of desonide are preferred. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days.	TEXACORT 2.5% (5) Clocortolone Pivalate flurandrenol fluticasone 0.05% (L) hydrocortisone butyrate 0.1% (O) triamcinolone 0.05% (C) amcinonide 0.1% (C, L, O) augmented betamethasone 0.05% (G, L, clobetasol 0.05% (L) desoximetasone 0.05% (O, 25% (G, O) diflorasone 0.05% (C) fluocinonide 0.1% (C) halcinonide 0.1% (C)
	CLARAVIS isotretinoin ZENATANE CORTICOSTERO CECREAM; GEGEL; LELOTIC LOW alcometasone desonide* fluocinolone 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone butyrate 0.1% (C) hydrocortisone 0.05%, 0.25% (C, L) betamethasone valerate desoximetasone 0.05%, 0.25% (C) fluocinolone 0.025% fluticasone 0.05% (C) mometasone SYNALAR 0.025% (C, O) triamcinolone 0.025%, 0.1% HIGH betamethasone dipropionate clobetaso/f E 0.05% (C, G, O, S) diflorasone 0.05% (O) fluocinoinide fluticasone 0.005% (O) halobetasol TOPICORT 0.025% (C)	M. O-COINTMENT; S=SOLUTION POTENCY M. POTENCY	*Cream, ointment, and lotion formulations of desonide are preferred. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days.	TEXACORT 2.5% (5) Clocortolone Pivalate flurandrenol fluticasone 0.05% (L) hydrocortisone butyrate 0.1% (O) triamcinolone 0.05% (C) APEXICON 0.05% (C) augmented betamethasone 0.05% (G, L, clobetasol 0.05% (L) desoximetasone 0.05% (0.5%, 0.25% (G, O) difforasone 0.05%, 0.25% (G, O) fluocinonide 0.1% (C)
	CLARAVIS isotretinoin 2ENATANE CCREAM; G=GEI; L=LOTIC 10W alciometasone desonide* fluorinoine 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone butyrate 0.1% (C) hydrocortisone 1%, 2.5% (C,LO) MEDIU betamethasone valerate desoximetasone 0.05%, 0.25% (C) fluorinoine 0.025% (C) mometasone SYNALAR 0.025% (C, O) triamcinoione 0.05% (C) G,O,S) diflorasone 0.05% (O) fluorinoide fluorinoi	M. O-COINTMENT; S=SOLUTION POTENCY M. POTENCY	*Cream, ointment, and lotion formulations of desonide are preferred. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days.	TEXACORT 2.5% (S) Clocortolone Pivalate flurandrenol fluticasone 0.05% (L) hydrocortisone butyrate 0.1% (O) triamcinolone 0.05% (C) amcinonide 0.1% (C, L, O) augmented betamethasone 0.05% (G, L, clobetasol 0.05% (L) desoximetasone 0.05% (O, 25% (G, O) diflorasone 0.05% (C) fluocinonide 0.1% (C) halcinonide 0.1% (C)
	CLARAVIS isotretinoin 2ENATANE CORTICOSTERO C=CREAM; G=GEI; L=LOTIC LOW alcometasone desonide* fluocinolone 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone butyrate 0.1% (C) hydrocortisone 1%, 2.5% (C,L,O) MEDIU betamethasone valerate desoximetasone 0.05%, 0.25% (C) fluocinolone 0.025%, 0.25% (C) fluocinolone 0.025%, 0.25% (C) mometasone SYNALAR 0.025% (C, O) triamcinolone 0.05% (C,G,O,S) difforasone 0.05% (C,G,O,S) difforasone 0.05% (C) fluocinonide fluriacsone 0.05% (O) halobetasol fluriacsone 0.05% (C) triamcinolone 0.5% (C) triamcinolone 0.5% (C) triamcinolone 0.5% (C) tutrancinolone 0.5% (C) ULTRAVATE 0.05% (C,O)	M POTENCY M POTENCY POTENCY ATORS - STEP 2 AGENTS	*Cream, ointment, and lotion formulations of desonide are preferred. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days.	TEXACORT 2.5% (5) Clocortolone Pivalate flurandrenol fluticasone 0.05% (L) hydrocortisone butyrate 0.1% (O) triamcinolone 0.05% (O) APEXICON 0.05% (C) augmented betamethasone 0.05% (G,L, clobetasol 0.05% (L) desoximetasone 0.05% (O, 25% (G,O) diflorasone 0.05% (C) fluocinonide 0.1% (C) halcinonide 0.1% (C)
	CLARAVIS isotretinoin 2ENATANE CORTICOSTERO C=CREAM; G=GEI; L=LOTIC LOW alcometasone desonide* fluocinolone 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone butyrate 0.1% (C) hydrocortisone 1%, 2.5% (C,L,O) MEDIU betamethasone valerate desoximetasone 0.05%, 0.25% (C) fluocinolone 0.025%, 0.25% (C) fluocinolone 0.025%, 0.25% (C) mometasone SYNALAR 0.025% (C, O) triamcinolone 0.05% (C,G,O,S) difforasone 0.05% (C,G,O,S) difforasone 0.05% (C) fluocinonide fluriacsone 0.05% (O) halobetasol fluriacsone 0.05% (C) triamcinolone 0.5% (C) triamcinolone 0.5% (C) triamcinolone 0.5% (C) tutrancinolone 0.5% (C) ULTRAVATE 0.05% (C,O)	POTENCY POTENCY ATORS - STEP 2 AGENTS ELIDEL	*Cream, ointment, and lotion formulations of desonide are preferred. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days.	TEXACORT 2.5% (S) Clocortolone Pivalate flurandrenol flutiasone 0.05% (L) hydrocortisone butyrate 0.1% (O) triamcinolone 0.05% (C) amcinonide 0.1% (C,LO) augmented betamethasone 0.05% (G,L) clobetasol 0.05% (L) diflorasone 0.05% (C) fluocinonide 0.1% (C,LO) halcinonide 0.1% (C)
	CLARAVIS isotretinoin 2ENATANE CORTICOSTERO C=CREAM; G=GEI; L=LOTIC LOW alcometasone desonide* fluocinolone 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone butyrate 0.1% (C) hydrocortisone 1%, 2.5% (C,L,O) MEDIU betamethasone valerate desoximetasone 0.05%, 0.25% (C) fluocinolone 0.025%, 0.25% (C) fluocinolone 0.025%, 0.25% (C) mometasone SYNALAR 0.025% (C, O) triamcinolone 0.05% (C,G,O,S) difforasone 0.05% (C,G,O,S) difforasone 0.05% (C) fluocinonide fluriacsone 0.05% (O) halobetasol fluriacsone 0.05% (C) triamcinolone 0.5% (C) triamcinolone 0.5% (C) triamcinolone 0.5% (C) tutrancinolone 0.5% (C) ULTRAVATE 0.05% (C,O)	M POTENCY M POTENCY POTENCY ATORS - STEP 2 AGENTS	*Cream, ointment, and lotion formulations of desonide are preferred. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days. To receive a step 2 agent: Trial and failure of a preferred medium or high potency topical corticosteroid greater than or equal to a 21 day trial in the last 90 days.	TEXACORT 2.5% (5) Clocortolone Pivalate flurandrenol flutiasone 0.05% (L) hydrocortisone butyrate 0.1% (O) triamcinolone 0.05% (C) amcinonide 0.1% (C,LO) augmented betamethasone 0.05% (G,L, clobetasol 0.05% (L) diflorasone 0.05% (C) fluocinonide 0.1% (C,LO) halcinonide 0.1% (C) HALOG 0.1% (O) HALOG 0.1% (O)
	CLARAVIS isotretinoin 2ENATANE CORTICOSTERO C=CREAM; G=GEI; L=LOTIC LOW alcometasone desonide* fluocinolone 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone butyrate 0.1% (C) hydrocortisone 1%, 2.5% (C,L,O) MEDIU betamethasone valerate desoximetasone 0.05%, 0.25% (C) fluocinolone 0.025%, 0.25% (C) fluocinolone 0.025%, 0.25% (C) mometasone SYNALAR 0.025% (C, O) triamcinolone 0.05% (C,G,O,S) difforasone 0.05% (C,G,O,S) difforasone 0.05% (C) fluocinonide fluriacsone 0.05% (O) halobetasol fluriacsone 0.05% (C) triamcinolone 0.5% (C) triamcinolone 0.5% (C) triamcinolone 0.5% (C) tutrancinolone 0.5% (C) ULTRAVATE 0.05% (C,O)	POTENCY POTENCY ATORS - STEP 2 AGENTS ELIDEL	*Cream, ointment, and lotion formulations of desonide are preferred. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days. To receive a step 2 agent: Trial and failure of a preferred medium or high potency 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	TEXACORT 2.5% (5) Clocortolone Pivalate flurandrenol fluticasone 0.05% (L) hydrocortisone butyrate 0.1% (O) APEXICON 0.05% (C) amcinonide 0.1% (C,L,O) augmented betamethasone 0.05% (G,L, clobetasol 0.05% (L) desoximetasone 0.05% (D, 0.25% (G,O) difforasone 0.05% (C) fluocinonide 0.1% (C) halcinonide 0.1% (C) halcinonide 0.1% (C) HALOG 0.1% (O)
	CLARAVIS isotretinoin 2ENATANE CORTICOSTERO CECREAM; GEGEI; LEJOTIC LOW alcometasone desonide* fluocinolone 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone butyrate 0.1% (C) hydrocortisone 1%, 2.5% (C,L,O) MEDIU betamethasone valerate desoximetasone 0.05%, 0.25% (C) fluocinolone 0.025% fluticasone 0.05% (C) mometasone SYNALAR 0.025% (C, O) triamcinolone 0.05% (D) fluocinolone (I) florasone 0.05% (C,G,O,S) difforasone 0.05% (O) fluocinoinide fluticasone 0.05% (O) halobetasol fluradrenolide fluticasone 0.05% (C) triamcinolone 0.5% (C) triamcinolone 0.5% (C) triamcinolone 0.5% (C) tutrancinolone 0.5% (C,O)	POTENCY POTENCY ATORS - STEP 2 AGENTS ELIDEL	*Cream, ointment, and lotion formulations of desonide are preferred. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days. Trial and failure of two preferred agents greater than or equal to 14 days in the last 90 days. To receive a step 2 agent: Trial and failure of a preferred medium or high potency topical corticosteroid greater than or equal to a 21 day trial in the last 90 days.	TEXACORT 2.5% (S) Clocortolone Pivalate flurandrenol flutiasone 0.05% (L) hydrocortisone butyrate 0.1% (O) triamcinolone 0.05% (C) amcinonide 0.1% (C,LO) augmented betamethasone 0.05% (G,L) clobetasol 0.05% (L) diflorasone 0.05% (C) fluocinonide 0.1% (C,LO) halcinonide 0.1% (C)

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 Optuming WITH ANY QUESTIONS
ERMATOLOGY (continued)	ATOPIC	DERMATITIS ADBRY	*Trial and failure of a preferred step 2 agent (immunomodulator) greater than or equal to a 21 day trial within the last 30 days required. Dupixent requires member be at least 6 months of age	CIBINQO** OPZELURA**
(continued)		DUPIXENT*	or older. No high-potency steroid trial will be necessary.	RINVOQ**
			**Trial and failure of all criteria to receive a step 3 agent as defined above including	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	
	PLAQUE P	SORIASIS (PP)	approval of the non-preferred agents. Client must have diagnosis of PP prior to approval of a preferred agent (Enbrel, Humira, or	CIMZIA**
		ENBREL HUMIRA	Otezla). To receive a non-preferred agent, client must have a diagnosis of PP and a 56-day trial	COSENTYX ILUMYA
		OTEZLA	and failure of two of the preferred agents.	REMICADE
		SOTYKTU* TALTZ	*Sotyktu requires diagnosis of moderate to severe PP and 56 day trial and failure of	SILIQ SKYRIZI
		IALIZ	Humira.	STELARA
			**Cimzia will be allowed for clients that are pregnant or breast-feeding	TREMFYA
	SCABICIDES	/PEDICULICIDES	Trial and failure of a preferred agent in the last 12 months.	malathion lotion
	permethrin]	NATROBA
BETES	VANALICE DIABET	ES AGENTS		spinosad (BRAND IS PREFERRED) metformin SR 24H osm (use preferred age
		JANIDES		metformin SR 24H mod (use preferred age
	metformin/ER			
	GLUCOSIDASE	INHIBITORS		miglitol
	acarbose	LITINUDES	the last 12 months will be required before approval can be given for a non-preferred agent.	zonogliui do
	nateglinide	LITINIDES	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
		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	1		
		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	GLYXAMBI (use separate preferred agents QTERN (use separate preferred agents)
		TRADJENTA	preferred agent is required before approval can be given for a non-preferred agent.	STEGLUJAN (use separate preferred agents)
	DPP-4 INHIBITOR	COMBO AGENTS	Trial and failure of metformin greater than or equal to a 90 day supply in the last 12 months will	alogliptin/metformin
		JANUMET/XR JENTADUETO	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 given for a non-preferred agent.	alogliptin/pioglitazone (use separate prefe agents)
		KOMBIGLYZE/XR	preferred agent is required before approval can be given for a non-preferred agent.	JENTADUETO XR
				saxagliptin/metformin (BRAND IS PREFER) sitagliptin/metformin (BRAND IS PREFERR)
	INCRETIN MIMETICS (G	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	liraglutide (use brand) MOUNJARO
		TRULICITY	present, in which case the trial of metformin is waived. A 90 day trial of failure of the preferred agent is required before approval can be given for a non-preferred agent.	OZEMPIC*
		VICTOZA	, , , , , , , , , , , , , , , , , , ,	SOLIQUA XULTOPHY (use separate preferred agents
			Dosage Limits Apply: Ozempic: 2mg/week	Action in face separate preferred agents
			Victoza: 1.8mg/day	
	SGLT2 I	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)
		FARXIGA JARDIANCE	be required before approval can be given for a preferred agent unless there is a diagnosis of ASCVD, CKD, or heart failure, in which case the trial of metformin will be waived. A 90 day trial	QTERN (use separate preferred agents) INVOKAMET
		SYNJARDY	and failure of a preferred agent is required before approval can be given for a non-preferred	INVOKANA
		XIGDUO XR	agent.	SEGLUROMET (use separate preferred age STEGLATRO
				STEGLUJAN (use separate preferred agent
				SYNJARDY XR (use separate preferred age TRIJARDY XR (use separate preferred agen
		TING INSULIN	Prior authorization will be required when using two different delivery forms of the same type of	ADMELOG (use preferred agent)
	HUMALOG HUMALOG 75/25	<u> </u>	insulin concurrently.	FIASP (use preferred agent) insulin lispro (use preferred agents)
	HUMALOG JR.	1		LYUMJEV
	HUMALOG MIX NOVOLOG MIX	1		
	LONG-AC	TING INSULIN	Prior authorization will be required when using two different delivery forms of the same type of	
	LANTUS SOLOSTAR* LANTUS vial	1	insulin concurrently.	Insulin Glargine (use preferred agent) Insulin Degludec
		1		SOLIQUA
		Ī		TOUJEO (use preferred agent) TRESIBA* (use preferred agent)
		<u> </u>		XULTOPHY (use separate preferred agents
	FREESTYLE (strips only)	TERS/TEST STRIPS	Quantity limits apply: Insulin Dependent Clients: 10 strips/day	ALL OTHER METERS AND TEST STRIPS
	FREESTYLE FREEDOM	1	Non-Insulin Dependent Clients: 10 strips/day Non-Insulin Dependent Clients: 4 strips/day	
	FREESTYLE FREEDOM LITE FREESTYLE INSULINX	Ī		
	FREESTYLE PRECISION NEO B		Clients are limited to 1 meter/365 days	
	FREESTYLE SIDEKICK II ONE TOUCH ULTRA II	1		
	ONE TOUCH ULTRA MINI ONE TOUCH ULTRA BLUE	Ī		
	ONE TOUCH VERIO ONE TOUCH VERIO FLEX	Ī		
	ONE TOUCH VERIO REFLECT	1		
	ONE TOUCH VERIO IQ PRECISION XTRA	Ī		
	EXTERNAL DIA	ABETIC DEVICES		OMNIPOD GO
	OMNIPOD DASH OMNIPOD 5			
		D GLUCOSE MONITORS	Prior authorization will be required to verify if the client is injecting insulin daily. Monitors will	GUARDIAN
		DEXCOM G6	also be limited to the labeled age.	MINIMED
		DEXCOM G7 FREESTYLE LIBRE		
		FREESTYLE LIBRE 2		
	ACUTE HYPOG	FREESTYLE LIBRE 3/PLUS GLYCEMIA AGENTS		GVOVE (use professed access)
	BAQSIMI ACUTE HYPOC	SET CEIVITA AGENTS	1	GVOKE (use preferred agent)
	ZEGALOGUE (autoinjector)	<u> </u>		

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 THIS LIST IS NOT ALL INCLUSIVE PLEASE CONTACT Optimility. WITH ANY QUESTIONS
IBROMYALGIA	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)
SASTROINTESTINAL		evacuants		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
	DIGESTIV CREON ZENPEP	PERTYZE*	Prior authorization required. *Pertyze will be preferred for members diagnosed with cystic fibrosis.	VIOKACE
		DROME WITH CONSTIPATION AMITIZA LINZESS TRULANCE	Client must have a diagnosis of Irritable Bowel Syndrome (IBS) with constipation.	
	MESA APRISO* LIALDA* mesalamine 400mg DR capsule mesalamine enema PENTASA	ALAMINE	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 (BRAND IS PREFERRED) mesalamine ER cap 0.375gm (BRAND IS PREFERRED) mesalamine sup 1000mg SFROWASA
		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 omeprazole 20.6mg capsules omeprazole tablets omeprazole tablets omeprazole by the composition of the c
GOUT	colchicine (tablets)	CHICINE		MITIGARE (use preferred agent)
	XANTHINE OXIDASE	AND URAT1 INHIBITORS	Trial and failure of the preferred agent greater than or equal to a 60 day supply in the last 12	ULORIC*
IEMATOLOGY	allopurinol	/EIGHT HEPARIN (LMWH)	months 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)
LIMATOLOGI	enoxaparin	MBIN 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.	enoxaparin 300MG/3ML
	SELECTIVE FACE ELIQUIS XARELTO (10mg, 15mg, 20mg, and starter pack)	CTOR XA INHIBITOR XARELTO 2.5mg*	*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	ELIQUIS (starter pack) SAVAYSA (use preferred agent)
	CPTP DE	BRILINTA	Client must have a diagnosis of acute coronary syndrome, history of myocardial infarction, or history of stroke and transient ischemic attack.	
		NTAGONIST ZONTIVITY	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 conjunction with aspirin or clopidogrel.	
	ANTIHEMOPH ADVATE ADVNOVATE AFSTVIA ELOCTATE ESPEROCT HEMOFIL M HEMLIBRA JIVI KOATE/KOATE-DVI KOGENATE FS/BIO-SET	ILIC FACTOR VIII		ALTUVIIIO KOVALTRY
	NOVOEIGHT NUWIQ OBIZUR RECOMBINATE XYNTHA/SOLOFUSE			
	NUWIQ OBIZUR RECOMBINATE XYNTHA/SOLOFUSE COAGULAT ALPHANINE SD ALPROILX BENEFIX IDELVION IXINITY REBINYN RIXUBIS	ION FACTOR IX		
	NUWIQ OBIZUR RECOMBINATE XYNTHA/SOLOFUSE COAGULAT ALPHANINE SD ALPROILX BENEFIX IDELVION IXINITY REBINYN RIXUBIS ANTIHEMOPH ALPHANATE HUMATE-P VONVENDI WILATE	ILIC FACTOR/VWF		
	NUWIQ OBIZUR RECOMBINATE XYNTHA/SOLOFUSE COAGULAT ALPHANINE SD ALPROLIX BEENEFIX IDELVION IXINITY REBINYN RIXUBIS ANTIHEMOPH ALPHANATE HUMATE-P VONVENDI WILATE EPOGEN MIRCERA RETACRIT			ARANESP PROCRIT

HERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLI THIS LIST IS NOT ALL INCLUSIVE PLEASE CONTACT OPLIMING WITH ANY QUESTIONS
ATITIS 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 VOSEVI** ZEPATIER
ADENITIS SUPPURATIVA	IMMUNO	MODULATORS	Humira will not be covered as a first line agent for the diagnosis for hidradenitis suppurativa.	ZELATEN
MONES		HUMIRA ITAGONISTS	*Please refer to the Additional Therapeutic Criteria chart at www.wymedicaid.org for specific	ORIAHNN
	MYFEMBREE ORILISSA		requirements.	
	GROWTH	HORMONE GENOTROPIN		HUMATROPE 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) JATENZO (use preferred agent)
			Other testosterone dosage form products will require a diagnosis of hypogonadism or	TESTOPEL (use preferred agent)
			insufficient testosterone production (not outlined on PDL).	testosterone gel (use preferred agent) testosterone solution (use preferred age
	THYROID	HORMONES	Ermeza will be covered with confirmed diagnosis of dysphagia.	XYOSTED (use preferred agent) THYQUIDITY
	ARMOUR THYROID	ERMEZA	Ethicza wiii de covered with commined diagnosis of dysphagia.	TIROSINT
I	LEVOXYL levothyroxine (tablets)			
	LEVO-T liothyronine			
	SYNTHROID UNITHROID			
	CONTR	ACEPTIVES		alyacen 1-35, 7/7/7
	afirmelle altavera			aranelle BALCOLTRA
	amethia amethyst			balziva briellyn
	apri			drospir/ethinyl estradiol/levomefolate
	ashlyna aubra/EQ			enpresse ethynodiol/ethinyl estradiol
ä	aurovela 1-20/FE 1-20, 1-35 aviane			FALESSA KIT fayosim
	ayuna			FEMLYV
a	azurette blisovi 1-20 FE, 1.5-30 FE			kaitlib FE chew layolis FE chew
	bekyree beyaz			levonest levonorgest/ethinyl estradiol/LO (84-7)
	camila			levonorgest/ethinyl estradiol 0.15-
	camrese/LO chateal/EQ			MERZEE MINASTRIN FE chew*
	CHARLOTTE 24 FE chew cyred			NEXSTELLIS noreth/ethinyl estradiol/FE chew 0.8/25
	dasetta 1-35, 7/7/7			nortrel OPILL
	daysee deblitane			PHEXXI
	deso/ethinyl estradiol drospir/ethinyl estradiol			philith rivelsa
	elinest emzahh			QUARTETTE SAFYRAL
	enskyce			SLYND
	errin estarylla			TAYSOFY TAYTULLA
	falmina finzala FE chew			tilia FE tri-legest FE
	gianvi			TRIVORA
	hailey FE 1/20, 1/35 heather			TWIRLA TYBLUME
	iclevia incassia			tydemy vyfemla
	introvale isibloom			wera wymzya FE chew
	jaimiess			XULANE
	jencycla jolessa			ZAFEMY
	juleber junel 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 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	1	1	I
	norlynda			

THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIES HASSES SON ACCOUNTED WITH ACCOUNTED TO A COUNTED TO A C
ORMONES; CONTRACEPTIVES	ocella	CRITERIA		PLEASE CONTACT OptumRx WITH ANY QUESTIONS
ontinued)	pimtrea			
	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			1
	viorele			1
	volnea			1
	vylibra			1
	yasmin-28 YAZ			1
	zumandimine			1
/PERLIPIDEMIA		EQUESTRANT	Trial and failure of ALL preferred agents greater than or equal to six (6) months in the last 12	WELCHOL
T ENEIT IDENIIA	cholestyramine/light	EQUESTIVANT	months will be required before approval can be given for a non- preferred agent.	WEEGIOE
	colestipol		months will be required before approval can be given for a non- preferred agent.	
		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		months will be required before approval can be given for a non-preferred agents.	
			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.	
			arag arag interaction, a non-preferred agent may be obtained with a prior authorization.	
			Prior authorization will be required for clients under the age of 10.	
		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			ZYPITAMAG
	simvastatin		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.	
	CTATINI CO	MBINATIONS		
	amlodipine/atorvastatin	WIBINATIONS	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
	VYTORIN*		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.	
	PCSK9-REL	ATED AGENTS	Client must have a diagnosis of homozygous familial hypercholesterolemia; have a diagnosis of	LEQVIO
		PRALUENT REPATHA	heterozygous familial hypercholesterolemia or atherosclerotic cardiovascular disease AND not	
		REPAIRA	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.	
	TRIGLYCERIDE LO	WERING AGENTS	Trial and failure of a preferred agent greater than or equal to a 90 day supply in the last 12	fenofibric acid
	fenofibrate		months will be required before approval can be given for a non-preferred agent.	fenofibrate (43/50/120/130/150mg)
	gemfibrozil			icosapent
	ĺ			LIPOFEN
	ĺ			omega-3-acid VASCEPA
PERTENSION/ CARDIOLOGY	ANGIOTENSIN RECE	PTOR BLOCKERS (ARBs)	Non-preferred ARBs will require a history of ALL preferred ARBs before approval	candesartan
	EDARBI			eprosartan 600mg
	irbesartan			1
	losartan			1
	olmesartan			1
	telmisartan			1
	valsartan	DUIDETICS	Non-aveferred ADD/disposite combinations will residue to bishows of ALL aveferred	anderestan UCT7
	EDARBYCLOR ARBs ANI	DIURETICS	Non-preferred ARB/diuretic combinations will require a history of ALL preferred	candesartan HCTZ telmisartan HCTZ
	irbesartan HCTZ			termisal tarr rie12
	losartan HCT			1
	olmesartan HCTZ			1
	valsartan HCTZ			<u> </u>
		BLOCKERS		
	clonidine			1
	clonidine TD patches			<u> </u>
	COMBINATI	ON PRODUCTS	Client must be greater than one year of age and have a diagnosis of Congestive Heart Failure	ENTRESTO SPRINKLES
	ĺ	ENTRESTO	(CHF) NYHA Class II-IV. Angiotensin converting enzyme (ACE) inhibitors and angiotensin receptor	VERQUVO
	ĺ		blockers (ARBs) will not be allowed in combination with Entresto.	1
				100
FECTIOUS DISEASE		OLONES	Please refer to the Additional Therapeutic Criteria Chart located at	moxifloxacin (use preferred agents)
	ciprofloxacin		http://www.wymedicaid.org/additional-therapeutic-criteria for Baxdela criteria.	1
	levofloxacin			I
	ofloxacin	CVCLINE		DORYY (use preferred asset)
	doxycycline	CYCLINE	-	DORYX (use preferred agent)
		CYCLINE	<u> </u>	minocycline 65mg and 115mg ER (use
	minocycline/ER	CTCLINE	7	
				preferred agent) SOLODYN (use preferred agent)
	INILIALED	ORRAMYCIN		
	INHALED 1 KITABIS	OBRAMYCIN	*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.	BETHKIS inhaled tobramycin

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 Optimile WITH ANY QUESTIONS
NFECTIOUS DISEASE continued)	ANTI-RE APRETUDE BIKTARVY CIMDUO DELSTRIGO DOVATO EVOTAZ GENVOYA ODEFSEY PIFELTRO PREZCOBIX ritonavir tablets SYMFI/LO TRIUMEQ TROGARZO	TROVIRALS 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.	JULIUCA NORVIR RUKOBIA** STRIBILD (use separate agents) SUNLENCA SYMTUZA (use separate preferred agents)
NFLAMMATION		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 PREFERREI diclofenac 3.5% soln. diclofenac 3% gel fenoprofen mefenamic acid NEOPROFEN (use preferred agent)
	ORAL COR budesonide cortisone acetate dexamethasone/intensol hydrocortisone methylprednisolone prednisolone prednisolone prednisone	TICOSTEROIDS		CELESTONE (use preferred agent) EMFLAZA
usomnia	BELSOMRA eszopicione zalepion zolpidem zolpidem ER	CODIAZEPINES	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: zaleplon: 30mg/day zolpidem: 15mg/day	EDLUAR (additional criteria applies) DAYVIGO QUVIVIQ* ROZEREM* zolpidem sublingual (additional criteria applies)
MENTAL HEALTH	NORADRENERGIC/SPECI mirtazapine tablets NOREPINEPHRINE/DOPAR bupropion ER/SR/XL SELECTIVE SEROTONIN citalopram escitalopram fluoxetine capsules paroxetine IR/CR sertraline	RES AGENTS donepezi/ODT galantamine/ER memantine tablets/solution PRESSANTS FIC SEROTONERGICS (NASS) INNE REUPTAKE INHIBITORS (NDRI) REUPTAKE INHIBITORS (SSRI)	Client must have a diagnosis of dementia. Trial and failure of two (2) preferred agents greater than or equal to six (6) weeks WITHIN THE LAST_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 reouries trial and failure of two preferred agents in any class Clients five (5) years of age and younger will require prior authorization before approval. Dosage limits apply: bupropion ER/SR/XL: 450mg/day citalopram < 60 years of age: 30mg/day fluoxetine < 18 years of age: 120mg/day mirtazapine: 67.5mg/day paroxetine IR > 18 years of age: 120mg/day paroxetine IR > 18 years of age: 175mg/day paroxetine IR > 18 years of age: 112.5mg/day paroxetine IR > 18 years of age: 112.5mg/day sertraline: 300mg/day glood of the properties of the	donepeil 23mg (use preferred agent) memantine ER NAMZARIC (use separate agents) rivastigmine capsules/patches NASS mirtazapine rapid dissolve tablets (use preferred agent) NDRI APLENZIN AUVELITY FORFIVO XL* SSRI citalopram capsules fluoxetine tablets VIIBRYD SNRI desvenlafaxine FETZIMA venlafaxine ER tablets (use preferred agent) OTHER TRINTELLIX***

WYOMING MEDICAID

Preferred Drug List (PDL) January 1, 2025

	Thomas merapeutic Citteria	situite) Bosage Elimeation Elst	(red font indicates quantity/dose limits apply), and Wyoming Medicaid Prov	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 PLEASE CONTACT Optimile. WITH ANY QUESTIONS
MENTAL HEALTH (continued)	ATYPICAL A ABILIFY MAINTENA ABILIFY ASIMTUFII	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 MYCITE (use preferred agent) CAPLYTA GEODON 20MG INJ
	aripiprazole tab/solution/ODT		Desk for an override.	LYBALVI (additional criteria applies)
	ARISTADA asenapine		**Clients nine (9) years of age and younger will require a prior authorization to receive approval of lurasidone and asenapine. Clients eighteen (18) years of age and younger will require a prior	NUPLAZID olanzapine 10mg Inj
	FANAPT** paliperidone		authorization to receive approval of Fanapt.	SAPHRIS (use preferred agent) SECUADO
	INVEGA HAFYERA			REXULTI***
	INVEGA SUSTENNA INVEGA TRINZA		***Rexulti approval for MDD treatment requires concurrent antidepressant therapy as well as a	UZEDY ZYPREXA RELPREVV
	lurasidone** olanzapine		trial and failure of aripiprazole or other preferred atypical antipsychotic indicated for adjunct MDD treatment.	
	PERSERIS		Trial and failure of two (2) preferred agents greater than or equal to 30 days in the last 12	
	quetiapine* quetiapine ER		months will be required before approval can be given for a non-preferred agent unless	
	RISPERDAL CONSTA risperidone		otherwise specified.	
	RYKINDO VRAYLAR	•		
	ziprasidone		Prior authorization will be required for any client five (5) years of age or younger, or for any	
			client taking both an injectable and oral dosage form of the same medication concurrently. Dosage limits apply:	
			aripiprazole <13 years of age: 15mg/day; ≥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	
			quetiapine <13 years of age: 400mg/day; 13-17 years of age: 600mg/day; >17 years of age: 800mg/day	
			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	
			asenapine: 20mg/day	
			ziprasidone ≤17 years of age: 120mg/day; >17 years of age: 200mg/day	
	SPECIAL ATYPICA clozapine/ODT	AL ANTIPSYCHOTICS	Dosage limits apply: 900mg/day	VERSACLOZ Suspension (use preferred agent)
	AMPH	S AMPHETAMINES	Clients over the age of 17 must have a diagnosis for ADD, ADHD (see ADD/ADHD criteria below),	AMPHETAMINES ADZENYS XR ODT
	LONG ACTING	ADDERALL XR*	narcolepsy, obstructive sleep apnea, shift work sleep disturbance, MS fatigue (see MS Fatigue criteria below), or refractory depression (see refractory depression criteria below).	DYANAVEL XR
		amphetamine salts combo XR dextroamphetamine CR caps		EVEKEO/ODT MYDAYIS
	IMMEDIATE REI	VYVANSE CAPSULES** EASE AMPHETAMINES	For clients over the age of 17, they must meet the DSM-5 criteria for diagnosis of ADHD. These	PROCENTRA VYVANSE CHEWABLES
	ININIEDIATE REE	amphetamine salts combo	criteria include:	ZENZEDI 2.5 AND 7.5MG TABLETS
		dextroamphetamine tablets PHENIDATES	Five or more symptoms of inattention, present for at least 6 months, inappropriate for developmental level.	METHYLPHENIDATES
	LONG ACTING M	IETHYLPHENIDATES CONCERTA*	OR OR	APTENSIO XR AZSTARYS
		dexmethylphenidate ER methylphenidate ER tablets	 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 DAYTRANA
		methylphemidate Ex tablets		
			AND	FOCALIN XR
			AND 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,	
	IMMEDIATE RELEAS	SE METHYLPHENIDATES devmethylphenidate	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, school or work functioning; and	FOCALIN XR JORNAY PM methylphenidate ER osmotic release (BRAND IS PREFERRED)
	IMMEDIATE RELEAS	dexmethylphenidate methylphenidate chewables	 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, 	FOCALIN XR JORNAY PM methylphenidate ER osmotic release (BRAND IS PREFERRED) methylphenidate ER/CR/SR <u>capsules</u> [METADATE CD/RITALIN LA APTENSIO XR]
	IMMEDIATE RELEA:	dexmethylphenidate	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, school or work functioning; and	FOCALIN XR JORNAY PM methylphenidate ER osmotic release (BRAND IS PREFERRED) methylphenidate ER/CR/SR capsules (METADATE CD/RITALIN LA, APTENSIO XR) RELEXXII QUILLICHEW ER
	IMMEDIATE RELEAS	dexmethylphenidate methylphenidate chewables methylphenidate solution	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, school or work functioning; and	FOCALIN XR JORNAY PM methylphenidate ER osmotic release (BRAND IS PREFERRED) methylphenidate ER/CR/SR <u>capsules</u> (METADATE CD/RITALIN LA, APTENSIO XR) RELEXXII
	IMMEDIATE RELEAS	dexmethylphenidate methylphenidate chewables methylphenidate solution	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, school or work functioning; and The symptoms must not be better explained by another mental disorder. 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.	FOCALIN XR JORNAY PM methylphenidate ER osmotic release (BRAND IS PREFERRED) methylphenidate ER/CR/SR capsules (METADATE CD/RITALIN LA, APTENSIO XR) RELEXXII QUILLICHEW ER
	IMMEDIATE RELEAS	dexmethylphenidate methylphenidate chewables methylphenidate solution	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, school or work functioning; and The symptoms must not be better explained by another mental disorder. Diagnosis of MS fatigue will require a fatigue severity scale score of 5.0, a 60-day trial of	FOCALIN XR JORNAY PM methylphenidate ER osmotic release (BRAND IS PREFERRED) methylphenidate ER/CR/SR capsules (METADATE CD/RITALIN LA, APTENSIO XR) RELEXXII QUILLICHEW ER
	IMMEDIATE RELEAS	dexmethylphenidate methylphenidate chewables methylphenidate solution	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, school or work functioning; and The symptoms must not be better explained by another mental disorder. 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	FOCALIN XR JORNAY PM methylphenidate ER osmotic release (BRAND IS PREFERRED) methylphenidate ER/CR/SR capsules (METADATE CD/RITALIN LA, APTENSIO XR) RELEXXII QUILLICHEW ER
	IMMEDIATE RELEAS	dexmethylphenidate methylphenidate chewables methylphenidate solution	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, school or work functioning; and The symptoms must not be better explained by another mental disorder. 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. "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	FOCALIN XR JORNAY PM methylphenidate ER osmotic release (BRAND IS PREFERRED) methylphenidate ER/CR/SR c <u>apsules</u> (METADATE CD/RITALIN LA, APTENSIO XR) RELEXXII QUILLICHEW ER
	IMMEDIATE RELEAS	dexmethylphenidate methylphenidate chewables methylphenidate solution	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, school or work functioning; and The symptoms must not be better explained by another mental disorder. 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. "Vyvanse will be approved for the diagnosis of binge-eating disorder for clients 18 years of age	FOCALIN XR JORNAY PM methylphenidate ER osmotic release (BRAND IS PREFERRED) methylphenidate ER/CR/SR capsules (METADATE CD/RITALIN LA, APTENSIO XR) RELEXXII QUILLICHEW ER
	IMMEDIATE RELEAS	dexmethylphenidate methylphenidate chewables methylphenidate solution	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, school or work functioning; and The symptoms must not be better explained by another mental disorder. 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. **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 client is under the age of 4, or has a history of the	FOCALIN XR JORNAY PM methylphenidate ER osmotic release (BRAND IS PREFERRED) methylphenidate ER/CR/SR capsules (METADATE CD/RITALIN LA, APTENSIO XR) RELEXXII QUILLICHEW ER
	IMMEDIATE RELEAS	dexmethylphenidate methylphenidate chewables methylphenidate solution	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, school or work functioning; and The symptoms must not be better explained by another mental disorder. 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. "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.	FOCALIN XR JORNAY PM methylphenidate ER osmotic release (BRAND IS PREFERRED) methylphenidate ER/CR/SR capsules (METADATE CD/RITALIN LA, APTENSIO XR) RELEXXII QUILLICHEW ER
	IMMEDIATE RELEAS	dexmethylphenidate methylphenidate chewables methylphenidate solution	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, school or work functioning; and The symptoms must not be better explained by another mental disorder. 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. **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 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	FOCALIN XR JORNAY PM methylphenidate ER osmotic release (BRAND IS PREFERRED) methylphenidate ER/CR/SR capsules (METADATE CD/RITALIN LA, APTENSIO XR) RELEXXII QUILLICHEW ER
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	IMMEDIATE RELEAS	dexmethylphenidate methylphenidate chewables methylphenidate solution	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, school or work functioning; and The symptoms must not be better explained by another mental disorder. 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. **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 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.	FOCALIN XR JORNAY PM methylphenidate ER osmotic release (BRAND IS PREFERRED) methylphenidate ER/CR/SR capsules (METADATE CD/RITALIN LA, APTENSIO XR) RELEXXII QUILLICHEW ER
	IMMEDIATE RELEAS	dexmethylphenidate methylphenidate chewables methylphenidate solution	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, school or work functioning; and The symptoms must not be better explained by another mental disorder. 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. "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 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	FOCALIN XR JORNAY PM methylphenidate ER osmotic release (BRAND IS PREFERRED) methylphenidate ER/CR/SR capsules (METADATE CD/RITALIN LA, APTENSIO XR) RELEXXII QUILLICHEW ER
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	IMMEDIATE RELEAS	dexmethylphenidate methylphenidate chewables methylphenidate solution	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, school or work functioning; and The symptoms must not be better explained by another mental disorder. 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. **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 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 agent (seach 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 KR: 60mg/day amphetamine salts combo: 60mg/day amphetamine salts combo (narcolepsy): 90mg/day dextroamphetamine: 90mg/day dextroamphetamine CR: 90mg/day	FOCALIN XR JORNAY PM methylphenidate ER osmotic release (BRAND IS PREFERRED) methylphenidate ER/CR/SR capsules (METADATE CD/RITALIN LA, APTENSIO XR) RELEXXII QUILLICHEW ER

MIGRAINE bet fron nar REI Sur	MIGRAINE STEP 1 ACUTE MIGRA	PROPHYLAXIS AGENTS divaloroex topiramate AMOVIG* AJOVY EMGALITY INE TREATMENT AGENTS	Client must must have a diagnosis of ADD or ADHD. Prior authorization will be required for clients under the age of 4. 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). 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 MAD 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 Trial and failure of both an anticonvulsant and a beta blocker (Step 1 agents) 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 will be limited to 16 tabs/30 days. **Starting dose will be limited to 70mg **Approval for non-preferred agents requires trial and failure of a preferred agent along with the trial and failures described with Step 1 Agents' criteria above.	QELBREE NURTEC QULIPTA** almotriptan
fro nar REI	MIGRAINE STEP 1 ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1 PARTITION TO THE MIGRA STEP 1 ACUTE MIGRA STEP 1	PROPHYLAXIS AGENTS divaloroex topiramate AGENTS AMMOVIG* AJOVY EMGALITY INE TREATMENT	apnea, shift work sleep disturbance, or refractory depression (see refractory depression criteria below). 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 Trial and failure of both an anticonvulsant and a beta blocker (Step 1 agents) 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. Nurtee will be limited to 16 tabs/30 days. *Starting dose will be limited to 70mg *Approval for non-preferred agents requires trial and failure of a preferred agent along with the trial and failures described with Step 1 Agents' criteria above. Trial and failure of two preferred agents will be required for approval of a non- preferred agent.	NURTEC QULIPTA**
fro nar REI sur	ACUTE MIGRA ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1 PARTITION PARTITION STEP 1 ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1	AGENTS divaloroex topiramate AGENTS AIMOVIG* AJOVY EMGALITY INETERATMENT	(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. Qeibree: 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 Trial and failure of both an anticonvulsant and a beta blocker (Step 1 agents) 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 will be limited to 16 tabs/30 days. *Starting dose will be limited to 70mg *Approval for non-preferred agents requires trial and failure of a preferred agent along with the trial and failures described with Step 1 Agents' criteria above. Trial and failure of two preferred agents will be required for approval of a non-preferred agent.	QULIPTA**
fro nar REI sur	ACUTE MIGRA ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1 PARTITION PARTITION STEP 1 ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1	AGENTS divaloroex topiramate AGENTS AIMOVIG* AJOVY EMGALITY INETERATMENT	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. Qeibree: 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 Trial and failure of both an anticonvulsant and a beta blocker (Step 1 agents) 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 will be limited to 16 tabs/30 days. *Starting dose will be limited to 70mg *Approval for non-preferred agents requires trial and failure of a preferred agent along with the trial and failures described with Step 1 Agents' criteria above. Trial and failure of two preferred agents will be required for approval of a non-preferred agent.	QULIPTA**
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fro nar REI sur	ACUTE MIGRA ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1 PARTITION PARTITION STEP 1 ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1	AGENTS divaloroex topiramate AGENTS AIMOVIG* AJOVY EMGALITY INETERATMENT	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 Trial and failure of both an anticonvulsant and a beta blocker (Step 1 agents) 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 will be limited to 16 tabs/30 days. *Starting dose will be limited to 70mg *Approval for non-preferred agents requires trial and failure of a preferred agent along with the trial and failures described with Step 1 Agents' criteria above. Trial and failure of two preferred agents will be required for approval of a non-preferred agent.	QULIPTA**
fro nar REI sur	ACUTE MIGRA ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1 PARTITION PARTITION STEP 1 ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1	AGENTS divaloroex topiramate AGENTS AIMOVIG* AJOVY EMGALITY INETERATMENT	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 will be limited to 16 tabs/30 days. *Starting dose will be limited to 70mg *Approval for non-preferred agents requires trial and failure of a preferred agent along with the trial and failures described with Step 1 Agents' criteria above. Trial and failure of two preferred agents will be required for approval of a non-preferred agent.	QULIPTA**
fro nar REI sur	ACUTE MIGRA STEP 1 ACUTE MIGRA STEP 1 PARTITION PARTIT	divaloroex topiramate AGENTS AIMOVIG* AJOVY EMGALITY INETRATMENT	Concurrent use of Botox will not be approved. Nurtec will be limited to 16 tabs/30 days. *Starting dose will be limited to 70mg *Approval for non-preferred agents requires trial and failure of a preferred agent along with the trial and failures described with Step 1 Agents' criteria above. Trial and failure of two preferred agents will be required for approval of a non-preferred agent.	
nar REI sur	ACUTE MIGRA STEP 1 ovatriptan ratriptan LPAX* matriptan	AIMOVIG* AJOVY EMGALITY INE TREATMENT	*Starting dose will be limited to 70mg **Approval for non-preferred agents requires trial and failure of a preferred agent along with the trial and failures described with Step 1 Agents' criteria above. Trial and failure of two preferred agents will be required for approval of a non- preferred agent.	
nar REI sur	ACUTE MIGRA STEP 1 ovatriptan ratriptan LPAX* matriptan	AIMOVIG* AJOVY EMGALITY INE TREATMENT	**Approval for non-preferred agents requires trial and failure of a preferred agent along with the trial and failures described with Step 1 Agents' criteria above. Trial and failure of two preferred agents will be required for approval of a non- preferred agent.	
nar REI sur	STEP 1 ovatriptan rratriptan ELPAX* matriptan			almotriptan
nar REI sur	ovatriptan Iratriptan ELPAX* matriptan	AGENIS		aimotriptan
REI sur	ELPAX* matriptan			ELYXYB
rizz	atriptan		Rizatriptan will be limited to clients 6 years of age or older Quantity limits apply:	Sumatriptan-Naproxen Sodium TOSYMRA (use preferred agent) ZEMBRACE (use preferred agent)
			naratriptan 1.mg; 25 tabs/34 days naratriptan 2.5mg; 10 tabs/34 days REIPAX 20mg; 20 tabs/34 days	ZAVZPRET zolmitriptan
			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	AGENTS NURTEC	Trial and failure of two triptan agents required for Step 2 Agent approval Trial and failure of two preferred triptan agents AND Nurtec will be required for approval of a non-preferred agent.	REYVOW UBRELVY
			Quantity limits apply: NURTEC 75mg: limited to 8 tabs/30 days REVIOW: 200mg/day or 1 tab/day, 4 tab/30 days	
OVEMENT DISORDERS		INHIBITORS	Quantity limits apply:	
INC TET	JSTEDO/XR* GREZZA* TRABENAZINE		AUSTEDO: limited to 4 tabs/day INGREZZA: limited to 4 tabs/day *Please refer to the Additional Therapeutic Criteria chart at www.wymedicaid.org	
BET	/ONEX :TASERON DPAXONE 20MG/ML*	AGENTS GILENYA KESIMPTA LEMTRADA	Gilenya, Kesimpta, Lemtrada, Ocrevus, and Tysabri will be approved for highly active disease, please refer to the ATCC for additional information.	AUBAGIO BAFIERTAM BRIUMVI EXTAVIA
din REE teri	methyl fumarate :BIF riflunomide	OCREVUS TYSABRI	Trial and failure of two preferred agents for at least 56 days (each from a separat class) will be required before approval can be given for a non-preferred agent.	glatiramer (BRAND IS PREFERRED) GLATOPA (use preferred agent) MAVENCLAD
VU	JMERITY		For Mavenclad, in addition to the above criteria, approval will be granted on a case- by-case basis.	MAYZENT PLEGRIDY PONVORY TECFIDERA ZEPOSIA
RCOLEPSY	STIM	ULANTS 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 (MS) Fatigue,	ZEPUSIA
	NON-ST	NUVIGIL* IMULANTS	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 amantadine AND discontinuation of medications that may contribute to drowsiness or fatigue.	SUNOSI WAKIX
			Clients will not be allowed to take two or more agents in this class concurrently	XYREM
UROPATHIC PAIN	GABA	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	
Lid	TOPICAL docaine Patches	LIDOCAINE		ZTLIDO
am		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	carbamazepine imipramine (capsules) oxcarbazepine

riedse reier to the Addi	tional inerapeutic Criteria C	PREFERRED AGENTS	red font indicates quantity/dose limits apply), and Wyoming Medicaid Pro	NON-PREFERRED AGENTS	
THERAPEUTIC CLASS	PREFERRED AGENTS	REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	GENERIC MANDATORY POLICY APPLIES THE UST IS NOT ALL INCLUSIVE PLEASE CONTACT OPTUMBA WITH ANY QUESTIONS	
PHTHALMICS	OPANTI	I-ALLERGICS	Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will	ALOCRIL ALOMIDE	
	azelastine		be required before approval can be given for a non-preferred agent. Alomide and Alocril will be approved for pregnancy. Alomide will be approved for children	bepotastine	
	BEPREVE* cromolyn 0.4%		under the age of 3.	epinastine ZERVIATE	
		ICS- QUINOLONES	Trial and failure of a preferred agent greater than or equal to 5 days in the last 12 months will	gatifloxacin	
	ciprofloxacin		be required before approval can be given for a non-preferred agent.	ZYMAXID	
	BESIVANCE gentamicin				
	moxifloxacin 0.5%				
	ofloxacin tobramycin				
	OPANTI-IN	FLAMMATORY	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 NEVANAC			DUREZOL ILEVRO	
	neviune			INVELTYS	
				LOTEMAX SM loteprednol 0.5% (BRAND IS PREFERRED)	
				PROLENSA	
		A-BLOCKERS	Trial and failure of three (3) preferred agents each greater than or equal to 30 days in the last 12	BETIMOL BETOPTIC S*	
	betaxolol carteolol		months will be required before approval can be given for a non-preferred agent. *Betoptic S will be approved for those with heart and lung conditions.		
	levobunolol		betoptic 5 will be approved for those with heart and tang conditions.		
	timolol OPCARBONIC AI	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 OR -COMB	O PRODUCTS		dorzolamide/timolol (BRAND IS PREFERRE	
	COMBIGAN*	O TRODUCTS	1	aoi zoiamide) diffuiul (BRAND IS PREFERRE	
	ROCKLATAN				
	SIMBRINZA OPDRY E	EYE AGENTS	Trial and failure of the preferred agent greater than or equal to 12 weeks will be required	CEQUA	
	RESTASIS*		before approval can be given for the non-preferred agent.	cyclosporine (BRAND IS PREFERRED)	
	XIIDRA			EYSUVIS MIEBO	
				RESTASIS MULTIDOSE (see preferred)	
	00.0000	TAGLANDINS	Trial and fail are fall and an about a short and a second a 20 days in the last 42	TYRVAYA	
	latanoprost	AGLANDINS	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-preferred agent.	bimatoprost IYUZEH	
	LUMIGAN			tafluprost	
	TRAVATAN Z XALATAN				
	ZIOPTAN				
	OPRHO KINA RHOPRESSA	SE INHIBITOR			
	OPSYMPAT	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%				
STEOPOROSIS		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***	
			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 with any concurrent osteoporosis treatment, and will be limited to approved indication		
			***Will be limited to 2 years of use		
		ALCITONIN			
ric	calcitonin-salmon ANTIBIOTIC/STER	OID COMBINATION		ciprofloxacin 0.2% (use preferred agent)	
-	ciprofloxacin/dexamethasone		1	CIPRO HC (use preferred agent)	
	Neo/Poly/HC suspension/solution ofloxacin			CORTISPORIN-TC (use preferred agent) FLUOCINOLONE ACET OIL 0.01%	
	tobramycin/dexamethasone			(use preferred agent)	
/ERACTIVE BLADDER	OVERACTIVE B	LADDER AGENTS	Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12	darifenacin	
	MYRBETRIQ oxybutynin /ER		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.	GELNIQUE GEL 10% GEMTESA	
	solifenacin		, ,	mirabegron (BRAND IS PREFERRED)	
				OXYTROL DIS tolterodine/ER	
				TOVIAZ	
MN	LONG-A	CTING C-IIs	Trial and failure of a preferred agent(s) greater than or equal to a 14 day supply in the last 12	trospium fentanyl patches	
	morphine ER tablets		months will be required before approval can be given for a non- preferred agent.	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 agen NUCYNTA ER**	
			(0-11-11-11-11-11-11-11-11-11-11-11-11-11	oxymorphone ER OXYCONTIN	
			Concurrent therapy with a benzodiazepine and a narcotic medication or with duplicate	XTAMPZA ER (additional criteria applies)	
			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.		
			**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		
	1		Methadone: Limited to 3 tablets per day		
		4	Morphine ER: 90mg/day	i	
			Nucynta ER: 327mg/day		
			Nucynta ER: 327mg/day Oxycontin: 80mg/day		
			Oxycontin: 80mg/day Oxymorphone: 40mg/day		
			Oxycontin: 80mg/day		

				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 PLEASE CONTACT Optumits: WITH ANY QUESTIONS
PAIN continued	codeine sulfate hydrocodone/APAP hydrocodone/IBU hydromorphone	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 NUCYNTA* oxymorphone ROXYBOND
	meperidine morphine oxvcodone oxycodone/APAP		*Nucynta will be allowed for diabetic peripheral neuropathy or clients with significant gastrointestinal concerns with other CII narcotics. 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)	
	C 111/C	V AGENTS	Clients will be limited to one short-acting narcotic at a time	DELDUCA
	BUTRANS tramadol	VAGENIS	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-AC amantadine	TING AGENTS		
	benztropine tablets carbidopa/levodopa pramipexole ropinirole			
		TING 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-REDI finasteride	JCTASE 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 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 (use separate agents silodosin
PULMONARY ANTIHYPERTENSIVES		JCTASE INHIBITORS ALYQ sildenafil suspension	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) WINNEVAIR
	GUANYLATE C	YCLASE INHIBITORS	Prior authorization required.	ADEMPAS (use preferred agent)
	PROSTACYCLII	NE VASODILATORS ORENITRAM	Prior authorization required. Client must have a diagnosis of pulmonary hypertension.	
	PROSTACYCLINE	RECEPTOR AGONIST	Prior authorization required.	UPTRAVI (use preferred agent)
RESTLESS LEG SYNDROME	RESTLESS LI	EG SYNDROME	Client must have a diagnosis of Restless Leg Syndrome (RLS). Trial and failure of gabapentin	HORIZANT
	pramipexole ropinirole	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	NEUPRO*
			Parkinson's Disease. Clients will not be allowed to take gabapentin and pregabalin concurrently	
SKELETAL MUSCLE RELAXANTS		RELAXANTS	Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12	carisoprodol
	baclofen (5, 10, 15mg tablets) cyclobenzaprine tizanidine tablets		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	chlorzoxazone cyclobenzaprine ER LYVISPAH metaxalone methocarbamol
ULCERATIVE COLITIS	IMMUNO	MODULATORS HUMIRA	antidepressant. Carisonradol is limited to 84 tabs/365 days Client must have diagnosis of UC prior to approval of a preferred agent. To receive a non- referred once tighen trust have a diagnosis of UC and a 56 day trial and failure of the	orphenadrine tizanidine capsules (use preferred agent) ENTYVIO* REMICADE
DECEMATIVE COLITIS	1	ITOWINA	preferred agent, client must have a diagnosis of UC and a 56-day trial and failure of the preferred agent.	REMICADE RINVOQ
OCCLIATIVE COLITS				SIMPONI SKYRIZI
SCEIMINE COLINS			* Refer to Additional Therapeutics Clinical Criteria Chart for more information	