

WYOMING MEDICAID
Preferred Drug List (PDL) - September 10, 2013

Drug classes not included on this list are not managed through a Preferred Drug List (PDL). HOWEVER, THIS EXCLUSION IS NOT A GUARANTEE OF PAYMENT OR COVERAGE. Dosage limits and other requirements may apply. Drugs new to market are non-preferred until a clinical review has been completed. PA criteria will apply to both the pediatric population, as well as the adult population for those plans where PA/PDL limits are allowed.

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

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

Please refer to the Additional Therapeutic Criteria Chart, Dosage Limitation List (red font indicates quantity/dosage limits apply), Epocrates, and the Wyoming Medicaid Provider Manual at <http://wyomedicaid.org> for additional criteria.

THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIES
ALLERGY / ASTHMA	ANTIHISTAMINES, MINIMALLY SEDATING		Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent.	desloratadine CLARINEX RDT/SYRUP levocetirizine
	cetirizine fexofenadine loratadine			
	ANTIHISTAMINE/DECONGESTANT COMBINATIONS		Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent.	CLARINEX-D
	cetirizine/pseudoephedrine fexofenadine/pseudoephedrine loratadine/pseudoephedrine			
	ANTICHOLINERGIC BRONCHODILATORS		Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent. Spiriva 5 day STARTER package will be allowed one (1) time per recipient.	ATROVENT HFA TUDORZA
	ipratropium SPIRIVA			
	CORTICOSTEROID / BRONCHODILATOR COMBO'S		Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent. Advair 7 and 14-day STARTER package will be allowed one (1) time per recipient.	
	ADVAIR/HFA DULERA SYMBICORT			
	LEUKOTRIENE MODIFIERS		Trial and failure of preferred agent greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent.	zafirlukast ZYFLO
	montelukast			
	NASAL ANTIHISTAMINES		Trial and failure of preferred agent greater than or equal to 90 days in the last 12 months will be required before approval can be given for a non-preferred agent.	ASTEPRO 0.15% DYMISTA (use separate agents) PATANASE
	azelastine			
	NASAL STEROIDS		Trial and failure of two (2) preferred agents greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent. Rhinocort will be approved for pregnancy.	BECONASE AQ DYMISTA (use separate agents) flunisolide OMNARIS QNASL RHINOCORT triamcinolone (BRAND IS PREFERRED) VERAMYST ZETONNA
	fluticasone NASACORT AQ* NASONEX			
	SHORT ACTING BRONCHODILATORS - INHALERS		Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent.	VENTOLIN HFA XOPENEX HFA
PROAIR HFA PROVENTIL HFA				
SHORT ACTING BRONCHODILATORS - NEBULIZERS		Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent.	levulbuterol (BRAND IS PREFERRED)	
albuterol neb XOPENEX neb*				
STERIOD INHALANTS		Trial and failure of three (3) preferred agents greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent. Alvesco will be approved for a history of oral thrush with steroid inhalants.	AEROBID/AEROBID-M ALVESCO ASMANEX PULMICORT SUSPENSION	
budesonide FLOVENT HFA/DISK PULMICORT FLEXHALER QVAR				
ALZHEIMERS	ALZHEIMER AGENTS		Client must have a diagnosis of dementia.	donepezil 23mg (use preferred) donepezil ODT (use preferred) NAMENDA XR
		donepezil EXELON PATCH/SOLUTION galantamine/ER NAMENDA rivastigmine capsules		
ANALGESICS	BUPRENORPHINE COMBINATIONS		Client must have a diagnosis of opioid dependence or abuse. This is not to be used to for the treatment of chronic pain. Only one (1) narcotic prescription will be allowed between fills. Prescriber must have a XDEA number. Subutex will be approved for clients pregnant or nursing or with a documented allergy to naloxone. Dosage limits apply (Max Dose: 24mg/day). Client is limited to two (2) years of Suboxone or buprenorphine use. Please submit PA requests on the Suboxone and buprenorphine PA form available at wyequalitycare.org .	buprenorphine/naloxone tablets (use preferred) SUBUTEX
	SUBOXONE FILM			
	LONG-ACTING		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. Fentanyl patches are limited to one patch every 72 hours. C-III's and C-IV's are not included and are available without prior authorization with the exception of Butrans (generic substitution is mandatory). **Butrans requires a trial of morphine sulfate or low dose trial of fentanyl patch. ***Nucynta ER will be allowed for diabetic peripheral neuropathy or slients with significant gastrointestinal concerns with other CII narcotics.	AVINZA BUTRANS** EXALGO KADIAN (10mg/200mg) morphines sulfate ER capsules NUCYNTA ER*** OPANER ER (5mg/10mg/20mg/30mg/40mg) oxymorphone ER (7.5mg/15mg) OXYCONTIN/CR
	fentanyl patch morphine sulfate ER tablets			
	SHORT-ACTING C-III's		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. **In addition to above criteria, Embeda and Oxecta require a diagnosis of drug/substance abuse. ***Nucynta will be allowed for diabetic peripheral neuropathy or slients with significant gastrointestinal concerns with other CII narcotics.	EMBEDA** levorphanol NUCYNTA*** OXECTA** oxymorphone oxycodone/IBU
codeine sulfate hydromorphone morphine sulfate oxycodone oxycodone/APAP oxycodone/ASA				
C-III/C-V AGENTS		Trial and failure of a preferred agent(s) greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. Quantity and dosage limits apply (max 8 tabs/day). **Butrans will require a 14 day trial and failure of tramadol IR and a 14 day trial and failure of tramadol ER prior to approval	BUTRANS** CONZIP RYBIX ODT tramadol/apap tramadol ER	
tramadol				

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ANDROGENS	TESTOSTERONE TOPICAL GELS		Testosterone agents are only allowed for diagnosis of hypogonadism or insufficient testosterone production. <i>Other testosterone dosage form products will require a diagnosis of hypogonadism or insufficient testosterone production (not outlined on PDL).</i>	FORTESTA (use preferred) TESTIM GEL (use preferred)	
		ANDROGEL			
ANTIBIOTICS	QUINOLONES			AVELOX FACTIVE NOROXIN PROQUIN ADDOX (use preferred) DORYX (use preferred) ORACEA (use preferred) SOLODYN (use preferred)	
	ciprofloxacin/ER levofloxacin ofloxacin				
	DOXYCYCLINE				
	doxycycline				
	MINOCYCLINE				
	minocycline/ER				
ANTICOAGULANTS	LOW MOLECULAR WEIGHT HEPARIN (LMWH)		Client must have diagnosis of non-valvular atrial fibrillation and relative contraindication to warfarin for approval. Client must have diagnosis of non-valvular atrial fibrillation and relative contraindication to warfarin for approval. Client must have diagnosis of non-valvular atrial fibrillation, deep vein thrombosis (DVT), pulmonary embolism (PE), reduction in risk of recurrence DVT or PE, or prophylaxis of deep vein thrombosis which can lead to pulmonary embolism in clients undergoing hip or knee replacement.	enoxaparin (BRAND IS PREFERRED) FRAGMIN (use preferred) LOVENOX 300MG/3ML (use preferred)	
	LOVENOX*				
	DIRECT THROMBIN INHIBITOR				
		PRADAXA			
	SELECTIVE FACTOR XA INHIBITOR				
	ELIQUIS				
		XARELTO			
ANTICONVULSANTS	DIAZEPAM RECTAL GEL			diazepam gel (BRAND IS PREFERRED)	
	DIASTAT*				
	LACOSAMIDE		Client must have a diagnosis of partial onset seizures.		
	VIMPAT				
ANTIDEPRESSANTS	ANTIDEPRESSANTS		Trial and failure of two (2) preferred agents greater than or equal to six (6) weeks 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. 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. 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. Trial and failure of an ACE Inhibitor greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for preferred ARB. Non-preferred ARBs and ARB/diuretic combinations also require a history of ALL preferred ARBs before approval can be given. Trial and failure of an ACE Inhibitor greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for preferred ARB. Non-preferred ARBs and ARB/diuretic combinations also require a history of ALL preferred ARBs before approval can be given. ARB COMBINATIONS EXFORGE/EXFORGE-HCTZ ALPHA-BLOCKERS CATAPRES PATCHES* clonidine		
	NORADRENERGIC/SPECIFIC SEROTONINICS (NaSS)				
	mirtazapine 15, 30, and 45mg				NaSS mirtazapine 7.5mg and rapid dissolve tablets (use preferred)
	NOREPINEPHRINE/DOPAMINE REUPTAKE INHIBITORS (NDRI)				NDRI
	bupropion ER/SR/XL			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.	APLENZIN FORFIVO XL
	SELECTIVE SEROTONIN REUPTAKE INHIBITORS (SSRI)				SSRI
	citalopram escitalopram fluoxetine capsules paroxetine IR/CR sertraline			*Cymbalta will be approved for clients with a diagnosis of osteoarthritis of the knee or chronic low back pain. Dosage limits apply: bupropion ER/SR/XL: 450mg/day citalopram ≤ 60 years of age: 60mg/day citalopram > 60 years of age: 30mg/day escitalopram: 30mg/day fluoxetine ≤ 18 years of age: 90mg/day fluoxetine > 18 years of age: 120mg/day mirtazapine: 67.5mg/day paroxetine IR/CR ≤ 18 years of age: 75mg/day paroxetine IR > 18 years of age: 90mg/day paroxetine CR > 18 years of age: 112.5mg/day sertraline: 300mg/day venlafaxine ER: 337.5mg/day	fluoxetine tablets (use preferred) VIBRYD
	SEROTONIN/NORPINEPHRINE REUPTAKE INHIBITORS (SNRI)				SNRI
	venlafaxine ER capsules				CYMBALTA* PRISTIQ venlafaxine ER tablets (use preferred)
	ANTIHYPERTENSIVES	ACE INHIBITORS			
benazepril captopril enalapril fosinopril lisinopril moexipril perindopril quinapril ramipril trandolapril					
ACE INHIBITORS AND DIURETICS					
benazepril/HCTZ captopril/HCTZ enalapril/HCTZ fosinopril/HCTZ lisinopril/HCTZ moexipril/HCTZ quinapril/HCTZ					
ANGIOTENSIN RECEPTOR BLOCKERS (ARBs)					
		AVAPRO* BENICAR DIOVAN losartan		candesartan EDARBI eprosartan 600mg irbesartan (BRAND IS PREFERRED) MICARDIS TEVETEN 400mg	
ARBs AND DIURETICS					
		AVALIDE* BENICAR HCTZ DIOVAN HCTZ losartan HCTZ		candesartan HCTZ EDARBYCLOR irbesartan HCTZ (BRAND IS PREFERRED) MICARDIS HCTZ TEVETEN HCTZ	
ARB COMBINATIONS					
		EXFORGE/EXFORGE-HCTZ		AZOR TWYNSTA (use separate agents) TRIBENZOR (use separate agents)	
ALPHA-BLOCKERS					
	CATAPRES PATCHES* clonidine		clonidine patch (BRAND IS PREFERRED) NEXICLON XR (use preferred)		

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ANTIPSYCHOTICS	ATYPICAL ANTIPSYCHOTICS		<p>**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 GHS Pharmacy Help Desk for an override.</p> <p><i>Typical antipsychotics do <u>not</u> require prior authorization.</i></p> <p>Dosage limits apply: ABILIFY <13 years of age: 23mg/day ABILIFY ≥13 years of age: 45mg/day FANAPT: 36mg/day INVEGA: 18mg/day LATUDA: 240mg/day Risperidone ≤ 17 years of age: 5mg/day Risperidone > 17 years of age: 24mg/day SAPHRIS: 30mg/day Olanzapine < 13 years of age: 15mg/day Olanzapine > 13 years of age: 30mg/day Quetiapine <13 years of age: 600mg/day Quetiapine 13-17 years of age: 900mg/day Quetiapine > 17 years of age: 1200mg/day ziprasidone < 17 years of age: 180mg/day ziprasidone > 17 years of age: 300mg/day</p>	SEROQUEL XR (use preferred)
	SPECIAL ATYPICAL ANTIPSYCHOTICS		Dosage limits apply: 1350mg/day	
CHOLESTEROL	BILE ACID SEQUESTRANT		Trial and failure of ALL preferred agents greater than or equal to six (6) months in the last 12 months will be required before approval can be given for a non-preferred agent.	WELCHOL
	STATINS, LOW POTENCY		Trial and failure of a preferred agent greater than or equal to a 90 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. If client's current medication therapy is contraindicated with the preferred statin(s) due to a drug-drug interaction, a non-preferred agent may be obtained with a prior authorization.	ALTOPREV fluvastatin/ER
	STATINS, HIGH POTENCY		Trial and failure of a preferred agent greater than or equal to a 90 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. If client's current medication therapy is contraindicated with the preferred statin(s) due to a drug-drug interaction, a non-preferred agent may be obtained with a prior authorization.	CRESTOR LIVALO
	STATIN COMBINATIONS		Trial and failure of a preferred agent greater than or equal to a 90 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. Zetia monotherapy will require PA.	ADVICOR (use separate agents) amlodipine/atorvastatin (BRAND IS PREFERRED) CHOLESTIN PRAVIGARD SIMCOR ZETIA* (use preferred)
	TRIGLYCERIDE LOWERING AGENTS		Trial and failure of a preferred agent greater than or equal to a 90 day supply in the last 12 months will be required before approval can be given for a non-preferred agent.	ANTARA fenofibric fenofibrate LOVAZA VASCEPA
CONTRACEPTIVES	ORAL CONTRACEPTIVES			amethia (BRAND IS PREFERRED) aranelle (BRAND IS PREFERRED) azurette (BRAND IS PREFERRED) BEYAZ (PA required) camila (use preferred) camrese (BRAND IS PREFERRED) caziant (use preferred) cesia (use preferred) cyclafem (BRAND IS PREFERRED) FEMCON FE (PA required) GENERESS FE CHW (PA required) gianvi (BRAND IS PREFERRED) heather (use preferred) introvale (use preferred) karva (BRAND IS PREFERRED) leena (BRAND IS PREFERRED) LO LOESTRIN (PA required) loryna (BRAND IS PREFERRED) NATAZIA (PA required) neon 0.5/35, 1/35, 7/7/7 (BRAND IS PREFERRED) NECON 1/50 (use preferred) norethindrone/ethinyl estradiol chew (PA required) norethindrone (use preferred) NORINYL 1/35 (use preferred) nortrel (BRAND IS PREFERRED) ocella (BRAND IS PREFERRED) ORTHO-NOVUM 1/50 (use preferred) quasense (use preferred) SAFYRAL (PA required) syeda (BRAND IS PREFERRED) tilia FE (BRAND IS PREFERRED) tri-legest FE (BRAND IS PREFERRED) tri-lo-sprintec (BRAND IS PREFERRED) viorele (BRAND IS PREFERRED) zarah (BRAND IS PREFERRED) zenchent FE chewable (PA required) zeosa chewable (PA required)

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CORTICOSTEROIDS	ORAL CORTICOSTEROIDS			CELESTONE (use preferred)
	budesonide cortisone acetate dexamethasone/intensol hydrocortisone methylprednisone prednisolone prednisone			
DIABETES	DIABETES AGENTS			FORTAMET (use preferred) GLUMETZA (use preferred) RIOMET (use preferred)
	BIGUANIDES			GLYSET
	α-GLUCOSIDASE INHIBITORS			
	acarbose		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.	
	MEGLITINIDES			
	STARLIX*		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.	nateglinide (BRAND IS PREFERRED) repaglinide
	THIAZOLIDINEDIONES			
	pioglitazone		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.	ACTOSPLUS MET (use separate agents) AVANDIA AVANDAMET (use separate agents)
	SULFONYLUREAS			
	glimepiride/ER glipizide/ER glyburide/ER		Trial and failure of metformin and a preferred agent greater than or equal to a 90 day supply in the last 12 months will be required before approval can be given for a non-preferred agent.	
	DIPEPTIDYL PEPTIDASE 4 (DPP-4) INHIBITORS			
		JANUVIA ONGLYZA	Trial and failure of metformin greater than or equal to a 90 day supply in the last 12 months will be required before approval can be given for a preferred agent. A 90 day trial of failure of the preferred agent is required before approval can be give for a non-preferred agent.	NESINA TRADJENTA
	DIPEPTIDYL PEPTIDASE 4 (DPP-4) INHIBITOR COMBO AGENTS			
		JANUMET JUVISYNC KOMBIGLYZE	Trial and failure of metformin greater than or equal to a 90 day supply in the last 12 months will be required before approval can be given for a preferred agent. A 90 day trial of failure of the preferred agent is required before approval can be give for a non-preferred agent.	JANUMET XR JENTADUETO KAZANO OSENI
	INCRETIN MIMETICS (GLP-1 RECEPTOR AGONISTS)			
		BYETTA	Trial and failure of metformin greater than or equal to a 90 day supply in the last 12 months will be required before approval can be given for a preferred agent. A 90 day trial of failure of the preferred agent is required before approval can be give for a non-preferred agent.	BYDUREON VICTOZA
	SGLT2 INHIBITORS			
		INVOKANA	Trial and failure of metformin greater than or equal to a 90 day supply in the last 12 months will be required before approval can be given for a preferred agent.	
	INTERMEDIATE-ACTING INSULIN			
	HUMULIN N HUMULIN 70/30 NOVOLOG N NOVOLOG 70/30			
	LONG-ACTING INSULIN			
	LANTUS <small>via</small>			LANTUS OPTICLIK/SOLOSTAR (use preferred) LEVEMIR (use preferred)
RAPID-ACTING INSULIN				
APIDRA HUMALOG NOVOLOG				
DIABETIC METERS/TEST STRIPS				
		Quantity limit applies (1 meter/365days).	ALL OTHER METERS AND TEST STRIPS	
	FREESTYLE INSULINX FREESTYLE LITE FREESTYLE FREEDOM LITE ONE TOUCH ULTRA ONE TOUCH ULTRA 2 ONE TOUCH ULTRA MINI ONE TOUCH ULTRASMART PRECISION XTRA			
EAR	ANTIBIOTIC/STEROID COMBINATION			CETRAXAL (use preferred) CIPRODEX (use preferred) CIPRO HC (use preferred) COLY-MYCIN S (use preferred) CORTISPORIN-TC (use preferred) dexamethasone sodium phosphate (use preferred) FLUOCINOLONE ACET OIL 0.01% (use preferred) <small>Necomycin/Polymyxin B Sulfates/Hydrocortisone 201/202 (BRAND 6 PREPARED)</small>
	CORTISPORIN SOL 1% OTIC* <small>Necomycin/Polymyxin B Sulfates/Hydrocortisone 201/202</small> ofloxacin			
FIBROMYALGIA	FIBROMYALGIA STEP 1			
	amitriptyline cyclobenzaprine			
	FIBROMYALGIA STEP 2			
		SAVELLA	Trial and failure of a Step 1 agent greater than or equal to six (6) weeks in the last 12 months is required for approval of a Step 2 agent.	
FIBROMYALGIA STEP 3				
	CYMBALTA LYRICA	Trial and failure of a Step 1 agent and a Step 2 agent greater than or equal to six (6) weeks in the last 12 months is required for approval of a Step 3 agent.		

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GASTROINTESTINAL	DIGESTIVE ENZYMES		Prior authorization required.	PANCREAZE pancrelipase (BRAND IS PREFERRED) PERTZYE TRI-PASE ULTRESA VIOKASE
	CREON 3000, 6000, 12000, 24000 UNIT ZENPEP*			
	PROTON PUMP INHIBITORS		Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. Lansoprazole solutabs will be approved for children less than or equal to 8 years of age. Lansoprazole capsules will be approved for children less than 1 year of age.	ACIPHEX lansoprazole NEXIUM omeprazole <u>tablets</u> (use preferred) omeprazole bicarbonate OMECLAMOX (use separate agents) PREVPAC (use separate agents) VIMOVO (use separate agents)
	MESALAMINE		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.	APRISO ASACOL/HD CANASA LIALDA PENTASA 500MG (use preferred) ROWASA
	DEXILANT omeprazole <u>capsules</u> pantoprazole			
	mesalamine enema PENTASA 250MG ONLY			
GROWTH HORMONE	GROWTH HORMONE		PA is required for use outside of FDA-approved indications. Evaluation by an endocrinologist is preferred. Clinical evidence of improved growth will be required on a yearly basis to support ongoing utilization. Clinical evidence of need for growth hormone will be required for adult growth hormone deficiency and pediatric growth failure due to inadequate endogenous growth hormone. Trial and failure of two (2) preferred agents within the last 12 months will be required for the following indications: Pediatric: Growth failure due to inadequate endogenous growth hormone, Prader-Willi syndrome, children born small for gestation. Turner syndrome. Adult: Replacement for those with growth hormone deficiency.	HUMATROPE OMNITROPE SAIZEN SEROSTIM TEV-TROPIN ZORBTIVE
		GENOTROPIN NORITROPIN NUTROPIN AQ		
HEPATITIS C	INTERFERON		Trial and failure of preferred agent greater than or equal to 30 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. Peg-intron will be approved for pediatric patients (aged 18 and under), for retreatment, and for dosage adjustments that cannot be achieved with Pegasys.	PEG-INTRON
	PEGASYS			
	PROTEASE INHIBITOR		Prior authorization required for non-preferred agent.	INCIVEK
	VICTRELIS			
IMMUNOMODULATORS	IMMUNOMODULATORS		Client must have diagnosis prior to approval for preferred agents (outlined below): Enbrel : Ankylosing Spondylitis (AS), Juvenile Idiopathic Arthritis (JIA), Plaque Psoriasis (PP), Psoriatic Arthritis (PA), Rheumatoid Arthritis (RA)** Humira : AS, Crohn's, JIA, PP, PA, Ulcerative Colitis (UC), RA** **56-day trial and failure of methotrexate required prior to approval of Enbrel or Humira for diagnosis of Rheumatoid Arthritis (RA) For non-preferred agents , 56-day trial and failure of a preferred agent is required and client must have diagnosis prior to approval (outlined below): Actemra : RA (60-day trial of methotrexate is required) Amevive : PP Cimzia : Crohn's, RA Kineret : RA Orencia : JIA, RA Remicade : AS, Crohn's, PP, PA, RA, UC Rituxan : RA Simponi : AS, PA, RA Stelara : PP Tysabri : Crohn's (additional PA criteria applies)	ACTEMRA AMEVIVE CIMZIA KINERET ORENCIA RAPTIVA REMICADE RITUXAN SIMPONI STELARA TYSABRI (additional criteria applies)
		ENBREL HUMIRA		
INSOMNIA	NON-BENZODIAZEPINES		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. Rozerem is non-preferred without a history of substance abuse Dosage limits apply: zaleplon: 30mg/day zolpidem: 15mg/day	EDLUAR (additional criteria applies) INTERMEZZO (additional criteria applies) ROZEREM zolpidem ER ZOLPIMIST (additional criteria applies)
	zaleplon zolpidem			

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MIGRAINE	TRIPTANS		Trial and failure of all preferred agents will be required for approval of a non-preferred agent. Quantity limits apply: naratriptan 1mg: 25 tabs/34 days naratriptan 2.5mg: 10 tabs/34 days sumatriptan kit: 3 kits/34 days sumatriptan vials: 2 vials/34 days sumatriptan nasal: 6 bottles/34 days sumatriptan 25mg: 41 tabs/34 days sumatriptan 50mg: 20 tabs/34 days sumatriptan 100mg: 10 tabs/34 days	AXERT FROVA RELPAK rizatriptan TREXIMET zomig triptan
	naratriptan sumatriptan			
MULTIPLE SCLEROSIS	IMMUNOMODULATOR (GLATIRAMER INJECTION)		Trial and failure of preferred interferon agent AND failure of Copaxone before approval can be given for a non-preferred agent. For Tysabri, in addition to the above criteria, additional prior authorization criteria applies.	AUBAGIO EXTAVIA BETASERON GILENYA REBIF TECFIDERA TYSABRI (additional criteria applies)
	COPAXONE			
NEUROPATHIC PAIN	INTERFERON		For the diagnosis of neuropathic pain, 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 for greater than or equal to a 12 week supply in the last 12 months will be required before approval can be given for a non-preferred agent.	CYMBALTA LYRICA
	AVONEX			
NSAIDS	TRICYCLIC ANTIDEPRESSANTS		For the diagnosis of neuropathic pain, 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 for greater than or equal to a 12 week supply in the last 12 months will be required before approval can be given for a non-preferred agent.	CYMBALTA LYRICA
		amitriptyline imipramine nortriptyline		
NSAIDS	GABAPENTIN		Trial and failure of two (2) preferred agents each greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. Dosing and quantity limits apply for ketorolac (limit 5days/34 days; max dose 40mg/day for oral tablets).	CALDOLOR CAMBIA POWDER CELEBREX FLECTOR (additional criteria applies) mefenamic acid NAPRELAN NEOPROFEN PENNSAID (additional criteria applies) SOLARAZE (additional criteria applies) SPRIX (additional criteria applies) VOLTAREN (additional criteria applies) ZIPSOR
		gabapentin		
OPHTHALMICS	NSAIDs		Trial and failure of two (2) preferred agents each greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. Dosing and quantity limits apply for ketorolac (limit 5days/34 days; max dose 40mg/day for oral tablets).	CALDOLOR CAMBIA POWDER CELEBREX FLECTOR (additional criteria applies) mefenamic acid NAPRELAN NEOPROFEN PENNSAID (additional criteria applies) SOLARAZE (additional criteria applies) SPRIX (additional criteria applies) VOLTAREN (additional criteria applies) ZIPSOR
	diclofenac etodolac fenoprofen flurbiprofen ibuprofen indomethacin ketoprofen ketorolac meclofenamate meloxicam nabumetone naproxen oxaprozin sulindac tolmetin			
OPHTHALMICS	OP. -ANTI-ALLERGICS		Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent. Emadine, Alomide, and Alocril will be approved for pregnancy. Alomide will be approved for children under the age of 3.	ALAMAST ALOCRIL ALOMIDE ALREX BEPREVE ELESTAT EMADINE ketotifen LASTACRAFT
	azelastine cromolyn PATADAY PATANOL			
OPHTHALMICS	OP. -ANTIBIOTICS- QUINOLONES		Trial and failure of a preferred agent greater than or equal to 5 days in the last 12 months will be required before approval can be given for a non-preferred agent. Azasite will be approved for pregnancy.	AZASITE BESIVANCE IQUIX levofloxacin ZYMAR ZYMAXID
	ciprofloxacin ofloxacin MOXEZA VIGAMOX			
OPHTHALMICS	OP. -ANTI-INFLAMMATORY- NSAIDS		Trial and failure of ALL preferred agents each greater than or equal to 5 day supply in the last 12 months will be required before approval can be given for a non-preferred agent.	ACULAR/LS/PP (use preferred) ACUVAIL BROMDAY bromfenac NEVANAC
	flurbiprofen diclofenac ketorolac			
OPHTHALMICS	OP. -BETA-BLOCKERS		Trial and failure of three (3) preferred agents each greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent. Betoptoc S will be approved for those with heart and lung conditions.	BETIMOL BETOPTIC S ISTALOL
	betaxolol carteolol levobunolol metipranolol timolol			
OPHTHALMICS	OP. -CARBONIC ANHYDRASE INHIBITOR		Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent.	AZOPT
	dorzolamide			
OPHTHALMICS	OP. -CARBONIC ANHYDRASE INHIBITOR COMBO		Trial and failure of a preferred agent greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent.	
	dorzolamide/timolol			
OPHTHALMICS	OP. -PROSTAGLANDINS		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.	LUMIGAN ZIOPTAN
	latanoprost TRAVATAN Z			
OPHTHALMICS	OP. -SYMPATHOMIMETICS		Trial of a preferred agent greater than or equal to 30 days in the last 12 months will be required before approval can be given for a non-preferred agent.	ALPHAGAN P 0.1% brimonidine 0.15% (BRAND IS PREFERRED) COMBIGAN (use separate agents)
	ALPHAGAN P 0.15%* brimonidine 0.2%			
OSTEOPOROSIS	BISPHOSPHONATES		Trial and failure of a preferred agent greater than or equal to 12 months will be required before approval can be given for a non-preferred agent. Fosamax liquid will be approved for clients that have difficulty swallowing.	ACTONEL ATELVIA FOSAMAX-D ibandronate
	alendronate			
OSTEOPOROSIS	NASAL CALCITONIN			
	calcitonin-salmon fortical			
OVERACTIVE BLADDER	OVERACTIVE BLADDER AGENTS		Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12 months will be required before approval can be given for a non-preferred agent. Oxytrol will be approved for clients that have an inability to swallow.	DETROL LA ENBLEX GELNIQUE GEL 10% MYRBETRIQ OXYTROL DIS SANCTURA XR tolterodine trospium
	oxybutynin /ER TOVIAZ VESICARE			
PHOSPHATE BINDERS	PHOSPHATE BINDERS		Prior authorization required for non-preferred agents.	calcium acetate tabs (BRAND IS PREFERRED) FOSRENOL PHOSLYRA RENAGEL 800MG (use preferred) RENEVELA
	calcium acetate capsules ELIPHOS* RENAGEL 400MG ONLY			

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THERAPEUTIC CLASS	PREFERRED AGENTS	PREFERRED AGENTS REQUIRING CLINICAL CRITERIA	CLINICAL CRITERIA	NON-PREFERRED AGENTS GENERIC MANDATORY POLICY APPLIES
PLATELET AGGREGATE INHIBITORS	THIENOPYRIDINE DERIVATIVES		Prior authorization required for clients on antiplatelet therapy greater than one (1) year.	
	clopidogrel EFFIENT ticlopidine	BRIILTA		
PROSTATE	CYCLOPENTYLTRIAZOLOPYRIMIDINE (CPTP) Derivatives		Client must have diagnosis of acute coronary syndrome to reduce thrombotic cardiovascular events.	AVODART JALYN (use separate agents)
	5-ALPHA-REDUCTASE INHIBITORS			
	finasteride			
	ALPHA 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 JALYN (use separate agents) RAPAFLO
	doxazosin tamsulosin terazosin			
PULMONARY ANTIHYPERTENSIVES	5-ALPHA-REDUCTASE INHIBITORS		Client must have a diagnosis of pulmonary hypertension.	
	ENDOTHELIN RECEPTOR ANTAGONISTS			
		ADICIRCA sildenafil (Revatio A/B rated generic) LETAIRS		
RESTLESS LEG SYNDROME	RESTLESS LEG SYNDROME		Client must have a diagnosis of Restless Leg Syndrome (RLS). Trial and failure of gabapentin greater than or equal to 60 days and a trial and failure of a dopamine agonist greater than or equal to 60 days in the last 12 months will be required before approval can be given for a non-preferred agent. *Neupro will be approved for clients with difficulty swallowing or for clients with a diagnosis of Parkinson's Disease.	HORIZANT NEUPRO*
		gabapentin pramipexole ropinirole		
SKELETAL MUSCLE RELAXANTS	MUSCLE RELAXANTS		Trial and failure of a preferred agent greater than or equal to a 14 day supply in the last 12 months, along with a medical diagnosis of muscle spasticity will be required before approval can be given for a non-preferred agent. Cyclobenzaprine will require a prior authorization for clients concurrently taking a tricyclic antidepressant.	carisoprodol chlorzoxazone cyclobenzaprine ER metaxalone methocarbamol orphenadrine tizanidine capsules (use preferred) Carisoprodol is limited to 84 tabs/365 days.
	baclofen cyclobenzaprine tizanidine tablets			
STIMULANT	AMPHETAMINES		Clients must have a diagnosis for ADD, ADHD, narcolepsy, obstructive sleep apnea, shift work sleep disturbance, MS fatigue (see MS Fatigue criteria below), or refractory depression (see refractory depression criteria below). Diagnosis of MS fatigue will require a fatigue severity scale score of 5.0, a 60-day trial of amantadine and discontinuation of medications that may contribute to drowsiness and fatigue. Diagnosis of refractory depression will require a 6-week trial and failure of an antidepressant (monotherapy) and continued concomitant use of an antidepressant with the stimulant. Prior Authorization will be required for clients under the age of 4. 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 MAOI 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. Dosage limits apply: ADDERALL XR: 60mg/day amphetamine salts combo: 60mg/day amphetamine salts combo (narcolepsy): 90mg/day CONCERTA: 135mg/day DAYTRANA: 45mg/9 hour patch dextroamphetamine: 90mg/day dextroamphetamine CR: 90mg/day dexmethylphenidate: 30mg/day FOCALIN XR < 13 years of age: 45mg/day FOCALIN XR > 13 years of age: 60mg/day methyl/methylphenidate: 135mg/day methyl/methylphenidate ER/CR/SR: 135mg/day VYVANSE: 105mg/day	AMPHETAMINES: ZENEDI 2.5 AND 7.5MG TABLETS METHYLPHENIDATES: methylphenidate ER/CR/SR capsules (METADATE CD/RITALIN LA) QUILLIVANT XR SUSPENSION
	LONG ACTING AMPHETAMINES			
		amphetamine salts combo XR VYVANSE dextroamphetamine CR		
	IMMEDIATE RELEASE AMPHETAMINES			
		amphetamine salts combo* dextroamphetamine		
	METHYLPHENIDATES			
	LONG ACTING METHYLPHENIDATES			
		DAYTRANA FOCALIN XR methylin ER methylphenidate ER/CR/SA/SR tablets		
	IMMEDIATE RELEASE METHYLPHENIDATES			
		FOCALIN methylin tablets methylphenidate		

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STIMULANT-LIKE AGENTS	SELECTIVE ALPHA-ADRENERGIC AGONIST		Trial and benefit of clonidine IR will be required prior to approval of the non-preferred agent.	KAPVAY
	CLONIDINE AGENTS			
	clonidine			
	GUANFACINE AGENTS		To obtain the non-preferred agent , client must meet the following criteria: Client must have a diagnosis of ADHD or ADD. Prior authorization will be required for clients under the age of 5. Client must have a trial and failure of a stimulant greater than or equal to a 14 day trial OR a trial and failure of Strattera greater than or equal to a 30 day supply AND trial and benefit of guanfacine (Tenex) in the previous 12 months OR a contraindication to ADHD medications (including stimulant and non-stimulant) OR a TIC disorder associated with stimulants (trial of stimulant required).	INTUNIV
guanfacine				
SELECTIVE NOREPINEPHRINE REUPTAKE INHIBITOR		Clients must have a diagnosis for ADD, ADHD, narcolepsy, obstructive sleep apnea, shift work sleep disturbance, MS fatigue (see MS Fatigue criteria below), or refractory depression (see refractory depression criteria below). Diagnosis of MS fatigue will require a fatigue severity scale score of 5.0, a 60-day trial of amantadine and discontinuation of medications that may contribute to drowsiness and fatigue. Diagnosis of refractory depression will require a 6-week trial and failure of an antidepressant (monotherapy) and continued concomitant use of an antidepressant with the stimulant. Prior Authorization will be required for clients under the age of 5. 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. Strattera is limited to 1 tablet/day; unless the dose is greater than 40mg/day or unable to achieve a prescribed dose with 1 tablet. <small>Dosage limits apply: STRATTERA: 150mg/day</small>		
STRATTERA				
TOPICAL AGENTS	IMPETIGO ANTIBIOTICS		Trial and failure of ALL preferred agents greater than or equal to 7 days in the past 90 days. Use smallest size appropriate for 7 day trial.	ALTABAX
	gentamicin mupirocin			
	BENZOYL PEROXIDE/CLINDAMYCIN COMBOS		Clients must be 12 to 20 years of age and have a diagnosis of acne vulgaris. Requires prior authorization for clients less than 12 years of age. Acne combinations are limited to clients under the age of 21.	ACANYA benzoyl peroxide/clindamycin (BRAND IS PREFERRED)
	BENZACLIN* clindamycin/benzoyl peroxide 1.2 (1)-5% (Refrig)			
	CORTICOSTEROIS <small>CREAM, GEL, LOTION, OINTMENT</small>		Trial and failure of ALL preferred agents greater than or equal to 14 days in the last 90 days.	PANDEL
	LOW POTENCY			
	alclometasone desonide fluocinolone 0.01% hydrocortisone butyrate 0.1% (C) hydrocortisone 1%, 2.5% (C,L,O) prednicarbate			
	MEDIUM POTENCY		Trial and failure of ALL preferred agents greater than or equal to 14 days in the last 90 days.	CLODERM CORDRAN/SP TOPICORT LP TRIANEX
	betamethasone valerate desoximetasone 0.05% (C) fluocinolone 0.025% fluticasone 0.05% (C) hydrocortisone butyrate 0.1% (O) hydrocortisone probutate 0.1% (C) mometasone triamcinolone 0.025%, 0.1%			
	HIGH POTENCY		Trial and failure of ALL preferred agents greater than or equal to 14 days in the last 90 days.	APEXICON HALOG
	amcinonide betamethasone dipropionate clobetasol desonimetasone 0.25%, 0.05% (G) diflorasone flucinonide flurandrenolide fluticasone 0.005% (O) halobetasol triamcinolone 0.5%			
	IMMUNOMODULATORS		Trial and failure of a preferred medium potency topical corticosteroid greater than or equal to a 21 day trial <u>and</u> a trial and failure of a preferred high potency topical corticosteroid greater than or equal to a 21 day trial in the last 90 days. For clients less than two (2) years of age, a trial and failure of a preferred low potency corticosteroid greater than or equal to a 21 day trial <u>and</u> a trial and failure of a preferred medium potency topical corticosteroid greater than or equal to a 21 day trial in the last 90 days.	
	EUIDEL PROTOPIC			
	SALICYLIC ACID			All other topical salicylic acid formulations.
alliclen shampoo 6% salacyn cream/lotion 6% Salicylic Acid Shampoo 6%				
SCABICIDES/PEDICULICIDES		Trial and failure of a preferred agent in the last 12 months.	NATROBA OVIDE SKLICE ULESFIA	
permethrin LINDANE				
UREA			All other topical urea formulations.	
Kerafoam Aerosol 30% Remeven Cream 50% urea hydration aerosol 35% urea emulsion 50% urea nail suspension 40% urea suspension 50% X-Viate Cream 40%				