

CATEGORY	Coverage Indicator	Step Order	PREFERRED DRUGS	Coverage Indicator	Step Order	NON-PREFERRED DRUGS PA Required	Criteria	
PDL Effective April 4, 2025								
*PLEASE NOTE: For a search box hit Ctrl F								
* PLEASE NOTE: All <i>cost effective</i> generics applicable to DEL are considered PREFERRED Drugs. "BASIC" Covered Drugs are bolded with the Coverage Indicator of "MC / DEL".								
General Criteria for all PDL categories- For more information or help using the PDL, providers may call 1-888-445-0497; members should call 1-866-796-2463. To access PDL and PA materials via the internet: www.mainearepdl.org								
A: Preferred Drugs- Unless otherwise specified, preferred drugs are available without prior authorization. Step order may apply for preferred drugs in some drug categories as indicated on the PDL. (See item "D" below for explanation of step order.)								
B: <u>Requests for Non-preferred Drugs</u> - Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.								
C: <u>Adequate Drug Trials</u> - 1. The minimum trial period for each preferred and step order drug is two weeks, unless otherwise stated within specific PDL drug categories; trials with less than a two week duration will be reviewed on a case-by-case basis; 2. A trial will not be considered valid if preferred or non-preferred products were readily available (by override, individual purchase, samples, etc.); 3. Certain drug trials, such as with controlled substances, may require evidence that the preferred drugs were actually tried (example: with random pill counts and with random urine drug tests, using the methods of GC/MS with no lower threshold); 4. Adequate trials require documentation of attempts to titrate dose of preferred agents toward desired clinical response. 5. Adequate trials include prevention/treatment of common adverse effects associated with preferred agents (example: antinausea, antipruritic, etc.)								
D: <u>Step Order</u> - When numbers appear in the "step order" column, it means drugs in this category must be used in the order specified, with the lower numbers having preference over the higher numbers. Chart notes should be provided to confirm drug trials that do not appear in the member's MaineCare drug profile.								
E. The Department will institute strategies to ensure cost effectiveness through the use of an enhanced Drug Benefit Preferred brand drugs will no longer be preferred in any PDL drug category where preferred generic drugs are also available. It is expected that preferred generics will be used prior to any preferred brands. This will be operated as a form of step care. Preferred brands in these categories will require prior authorization for these high utilization / high cost members.								
F: <u>Brand Name Medication Requests</u> - (Must be submitted on the Brand Name PA request form)- According to MaineCare Benefits Manual Chapter II (80.07-5), when medically necessary covered brand-name drugs have an A-rated generic equivalent available, the most cost effective medically necessary version will be approved and reimbursed, since the brand-name and A-rated generic drugs have been determined by the FDA to be chemically and therapeutically equivalent. The Bureau does not make determinations as to whether or not a generic drug is clinically inferior or inequivalent to its brand version. This is the proper role of the FDA. Physicians should submit their reports of generic inequivalence directly to the FDA via the MEDWATCH.								
G: <u>PA requests for non- FDA Approved Indications</u> - Decisions will be made on a case-by-case basis until the DUR committee is able to review the evidence and make a recommendation. Interim approvals and DUR recommendations for approval of a drug for a non- FDA approved indication will require a minimum of two published, peer reviewed, non contradicted, double- blind, placebo-controlled randomized clinical studies establishing both safety and efficacy.								
H: <u>Dose Consolidation Requirements</u> - Some drugs may also be affected by dose consolidation requirements. Please see Dose Consolidation List and/or Splitting Tables provided in the PDL.								
I: <u>Trials from Multiple Drug Classes</u> - Trial/failure/intolerance to preferred agents from multiple classes within the same category or other categories of drugs may be required prior to the approval of non-preferred agents (e.g., Cymbalta, Zofran, Elidel and others).								
J: <u>Drug-specific PA Forms</u> - Drug-specific PA forms contain medical necessity documentation requirements and/or criteria that may not be repeated in the PDL. Drug-specific PA forms may be obtained on the web at www.mainearepdl.org .								
K: <u>PA Exemptions for Prescribers</u> - According to MaineCare Benefits Manual Chapter II (80.07-4), providers may receive a three (3) month exemption from prior authorization requirement for certain categories of drugs when they demonstrate high compliance with the Department's PDL. The Department will notify providers in writing which drug categories are included and what dates apply to the exemption. If a provider loses his/ her exemption, members who previously were not required to obtain a PA while the prescriber was exempt will be required to do so, and criteria for approval of that medication will need to be met.								
L: <u>Drug-Drug Interactions (DDI)</u> - The DUR Committee has implemented new drug-drug interaction edits requiring prior authorization. Several drug-drug combinations and PDL drug categories are affected by new PA requirements. These will be indicated in the PDL with DDI notation. Please see the DDI document provided in the PDL.								
ASSORTED ANTIBIOTICS								
BETA-LACTAMS / CLAVULANATE COMBO'S	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC MC/DEL MC MC MC/DEL		AMOXICILLIN AMOXICILLIN/POTASSIUM CLA CHEW AMOXICILLIN/POTASSIUM CLA SUSR AMOXICILLIN/POTASSIUM CLA TABS AMPICILLIN BICILLIN L-A SUSP DICLOXACILLIN SODIUM CAPS OXACILLIN SODIUM SOLR PENICILLIN V POTASSIUM TIMENTIN SOLR UNASYN SOLR ZOSYN	MC/DEL MC/DEL		AUGMENTIN ³ AUGMENTIN XR TB12 ⁴	3. Chewable 125mg & 250mg and Solution 125mg/5ml and 250mg/5ml available without PA. 4. Use preferred generic amoxicillin/clavulanate potassium alternatives. Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. DDI: Ampicillin will now be non-preferred and require prior authorization if it is currently being used in combination with either Prevacid, pantoprazole, Prilosec, or any currently non preferred PPI.
CEPHALOSPORINS	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		CEFADROXIL HEMIHYDRATE CEFAZOLIN SODIUM SOLR CEFENDINIR CEFEPIME CEFPODOXIME CEFPODOXIME PROXETIL SUS CEFPODOXIME PROXETIL TAB CEFIXIME 400MG ² CAP	MC MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC		CEDAX CEFACTOR ¹ CEFADROXIL MONOHYDRATE TABS CEFIXIME SUS CEPHALEXIN TABS CEPHALEXIN 750MG CAPS CEFTIN DAXBIA	1. Both brand and generic are clinically non-preferred. 2. Dosing limits apply, please see Dosage Consolidation List. 3. Approvals will only be considered for patients 18 years of age or older who	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.

			MC/DEL		ZEMDRI ²		Current users of Tobi Nebu and Tobramycin Soln will be allowed a grace period until 10/1/15 to transition to preferred Kitabis. Arikayce will require clinical PA to confirm MAC lung disease and for use in adults who have limited or no alternative treatment options. Zemdri will be reserved for patients with limited or no alternative treatment of care.
ANTI-MYCOBACTERIALS / ANTI-TUBERCULOSIS	MC/DEL MC/DEL MC/DEL MC/DEL		ETHAMBUTOL HCL TABS MYAMBUTOL TABS RIFABUTIN CAPS RIFAMPIN	MC/DEL MC/DEL MC MC/DEL	MYCOBUTIN CAPS PRETOMANID RIFADIN CAPS	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Pretomanid is indicated as part of a combination regimen with bedaquiline and linezolid for the treatment of adults with pulmonary extensively drug resistant (XDR) or treatment-intolerant or non-responsive multidrug-resistant (MDR) tuberculosis (TB). Approval of this indication is based in limited clinical safety and efficacy data. This drug is indicated for use in a limited and specific population of patients. DDI: Preferred rifampin will be non-preferred and require prior authorization if it is currently being used in combination with either Pradaxa or Latuda.
ANTIMALARIAL AGENTS	MC/DEL MC MC/DEL MC/DEL		DARAPRIM TABS KRINTAFEL ² MEFLOQUINE HCL TABS QUININE SULFATE	MC MC/DEL MC MC MC/DEL	ARALEN TABS CHLOROQUINE PHOSPHATE TABS ³ HYDROXYCHLOROQUINE TABS ³ ISONARIF ¹ MALARONE TABS PLAQUENIL TABS	Use PA Form# 20420 1. Ingredients available as preferred without PA. 2. Krintafel is preferred for ≥ 16 years of age. 3. Established users will be grandfathered	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. DDI: Avoid coadministration of Krintafel® with Organic Cation Transporter 2 (OCT2) and Multidrug and Toxin Extrusion (MATE) substrates (e.g. dofetilide, metformin).
ANTHELMINTICS	MC/DEL MC/DEL MC/DEL		ALBENDAZOLE PRAZICANTEL TAB STROMECTOL TABS	MC MC MC/DEL	ALBENZA TABS EMVERM BILTRICIDE TABS	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
ANTIBIOTICS - MISC.	MC MC MC MC/DEL MC MC/DEL MC MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL MC		AZACTAM SOLR COLY-MYCIN-M SOLR COLISTIMETHATE SODIUM SOLR FIRVANQ ⁴ FUROXONE TABS METRONIDAZOLE ¹ PENTAMIDINE ISETHIONATE SOLR SOLOSEC TRIMETHOPRIM TABS VANCOMYCIN 5GM INJ. VANCOMYCIN CAPS XIFAXAN 200mg	MC MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL MC MC/DEL MC MC MC/DEL MC MC	AEMCOLO COLISTIMETHATE SODIUM SOLR CAYSTON ³ FLAGYL CAPS FLAGYL TABS FLAGYL ER TBCR KETEK LIKMEZ METRONIDAZOLE 375MG CAPS ¹ METRONIDAZOLE 750MG TABS ¹ NEBUPENT SOLR REBYOTA ⁵ TINDAMAX VANCOMYCIN 10GM INJ. ² XENLETA XIFAXAN VOWST ⁵	1. 375mg caps and 750mg tabs are non-preferred. Please use available preferred strengths(250mg & 500mg tabs) to obtain required dose without PA. 2. Please use multiple 5gm which are preferred to obtain dose without PA. 3. Clinical PA is required to establish CF diagnosis and medical necessity. Prior trail and failure of preferred Tobi before approval will be granted. 4. Quantity limit of one per 150ml bottle. 5. For the treatment of patients 18 years of age and older.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. 1. For macrolide resistant infections when quinolones inappropriate DDI: Ketek is non-preferred but with any prior authorization requests, the member's drug profile will also be monitored for concurrent use with either Enablex 15mg or carbamazepine. Cayston is only indicated to improve respiratory symptoms in CF patients with Pseudomonas aeruginosa. Dosing limits, as should be given TID X28 days (followed by 28 days OFF Cayston therapy). A bronchodilator should be used before administration of Cayston. Xenleta will be considered for the treatment of adults with community-acquired bacterial pneumonia (CABP) caused by the following susceptible microorganisms: Streptococcus pneumoniae, Staphylococcus aureus (methicillin-susceptible isolates), Hemophilus influenzae, Legionella pneumophila, Mycoplasma pneumoniae, and Chlamydia pneumoniae. Vowst: To prevent the recurrence of C.difficile infection (CDI) in individuals 18 years of age and older following antibacterial treatment for recurrent CDI (rCDI). Likmez: patient has a medical necessity for a non-solid oral dosage form. Rebyota: For the prevention of recurrence of C. difficile infection (CDI) in individuals 18 years of age and older following antibiotic treatment for recurrent CDI. The limitation of use is that Rebyota® is not indicated for treatment of CDI.
CARBAPENEMS				MC	INVANZ SOLR	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on

				MC MC/DEL MC/DEL		MERREM SOLR PRIMAXIN RECARBRIO		the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
LINCOSAMIDES / OXAZOLIDINONES / LEPROSTATICS	MC/DEL MC/DEL MC/DEL MC MC/DEL		CLEOCIN SOLN CLEOCIN SUSR CLINDAMYCIN HCL 150CAPS DAPSONE TABS LINEZOLID 600mg TABS ²	MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL	8 8 8 8 9 9	CLEOCIN CAPS CLINDAMYCIN HCL 300CAPS ¹ SIVEXTRO VIBATIV ZYVOX SUSR ZYVOX TABS	1. Use multiple 150's for Clindamycin instead of 300's. 2. Quantity limit of 14 days supply within a 60day period. Use PA Form# 30820 for Zyvox & Vibativ Use PA Form# 20420 for all others	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. For Zyvox or Vibativ, please see the criteria listed in the Antibacterial Antibiotics PA form.
ANTI INFECTIVE COMBO'S - MISC.	MC/DEL MC/DEL MC/DEL MC/DEL		ERYTHROMYCIN/SULF SUSR SEPTRA/DS TABS SULFAMETHOXAZOLE/TRIMETH TRIMETHOPRIM/SULFAMETHOXA	MC MC		BACTRIM DS TABS VABOMERE ¹	Use PA Form# 20420 1. For the treatment of patients ≥ 18 years of age.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
ANTIPROTOZOALS	MC/DEL MC/DEL		BENZNIDAZOLE ² LAMPIT ²	MC		ALINIA ¹	1. Alina is preferred for children less than 12 years of age. 2. Clinical PA required for appropriate diagnosis. Use PA Form# 20420	Benznidazole is indicated for pediatric patients 2 to 12 years of age for the treatment of Chagas disease (American trypanosomiasis) caused by Trypanosoma cruzi.
ANTI - FUNGALS								
ANTIFUNGALS - ASSORTED	MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		ANCOBON CAPS FLUCONAZOLE ¹ KETOCONAZOLE TABS ⁷ NYSTATIN TERBINAFINE TABS ⁴ VORICONAZOLE TABS	MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL MC	6 6 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	LAMISIL TABS ⁴ ITRACONAZOLE BREXAFEMME CRESEMBA ⁹ GRIFULVIN V TABS GRISEOFULVIN SUSP GRISEOFULVIN ULTRAMICROSI TABS GRIS-PEG TABS REZZAYO ⁹ SPORANOX SOLN ² SPORANOX PULSEPAK CAPS ³ SPORANOX CAPS ³ DIFLUCAN ERAXIS INJ ⁶ GRIFULVIN SUSP ONMEL NOXAFIL ⁵ TOLSURA VFEND TABS VIVJOA	See quantity limit table. Non-preferred products must be used in specified step order. Continue to use Anti-Fungal PA form for non-preferred products. 1. QL--1/every 7-day period (150mg only). 2. Sporanox QL 300cc/month with PA. See quantity limit table. 3. Sporanox QL 30/month with PA. 4. Quantity limit of one tablet daily. Please see dosage consolidation list. 5. Approved if immuno suppressed/ HIV or if the member has failed a 7 day trial of a preferred antifungal therapy. 6. Eraxis will be approved if submitting with documentation that it was initiated during a hospitalization and this request is to finish the hospital course.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. The other criteria are listed on the Antifungal PA form including the required proof of a non-cosmetic fungal infection. DDI: Any Griseofulvin will now be non-preferred and require prior authorization if it is currently being used in combination with either Prevacid, pantoprazole, Prilosec, or any currently non preferred PPI. DDI: Sporanox is non-preferred but with any prior authorization requests, the member's drug profile will also be monitored for current use with Enablex 15mg, Vesicare 10mg, Prandin, Prevacid, pantoprazole, Prilosec, or any currently non preferred PPI, due to a significant drug-drug interaction. DDI: Vfend is non-preferred but with any prior authorization requests, the member's drug profile will also be monitored for concurrent use with Warfarin. DDI: Fluconazole (except 150mg strength) will now be non-preferred and require prior authorization if it is currently being used with glimepiride (Amaryl), Enablex 15mg, or Vesicare 10mg. Diflucan is non-preferred but with any prior authorization requests, the member's drug profile will also be monitored for concurrent use with either glimepiride (Amaryl), Enablex 15mg, or Vesicare 10mg. DDI: Fluconazole will require prior authorization if being used in combination with Plavix or Warfarin. DDI: Ketoconazole will now be non-preferred and require prior authorization if they are currently being used in combination with any of the following medications: Prevacid, Pantoprazole, Plavix, Onglyza, Enablex 15mg, Vesicare 10mg, Latuda, Cometriq, Tafinlar or Omeprazole. Rezzayo: In patients 18 years of age or older who have limited or no alternative options for the treatment of candidemia and invasive candidiasis.

7. Quantity limits allowing 30 day supply without PA. PA will be required if using > 30 days.

8. For children < 18, quantity limits allows 8 weeks supply without PA. PA will be required if using > than 8 weeks. If 18 and older PA will be required for any quantity. Not approving for Onychomycosis indication.

9. For patients ≥ 18years of age

[Use PA Form# 10120](#)

ANTI - VIRALS

ANTIRETROVIRALS

MC/DEL	MC	MC/DEL	MC/DEL	8	ABACAVIR SOL
MC		ABACAVIR TABS	MC/DEL	8	ABACAVIR SOL
MC/DEL		APRETUDE	MC/DEL	8	APTIVUS
MC		ATAZANAVIR	MC	8	ATRIPLA ¹
MC		BIKTARVY	MC/DEL	8	CIMDUO
MC		CABENUVA	MC/DEL	8	COMBIVIR TABS
MC		COMPLERA ¹	MC/DEL	8	EDURANT
MC/DEL		DELSTRIGO	MC/DEL	8	EPZICOM ¹
MC		DESCOVY ¹	MC/DEL	8	FUZEON
MC		DIDANOSINE	MC/DEL	8	INTELENCE
MC/DEL		DOVATO	MC/DEL	8	ISENTRESS ³
MC		EFAVIRENZ TAB	MC/DEL	8	ISENTRESS HD
MC/DEL		EFAVIRENZ CAP	MC	8	JULUCA
MC		EFAVIRENZ-EMTRICITABINE-TENOFOVIR DF TAB	MC	8	KALETRA
MC		EMTRICITABINE-TENOFOVIR	MC/DEL	8	LAMIVUDINE SOLN
MC		EMTRIVA ¹	MC/DEL	8	LEXIVA
MC		EPIVIR SOL	MC/DEL	8	NEVIRAPINE
MC/DEL		EVOTAZ ¹	MC	8	NORVIR
MC		GENVOYA ^{1,4}	MC/DEL	8	PIFELTRO
MC/DEL		ISENTRESS 400MG ⁵	MC	8	RETROVIR
MC/DEL		ISENTRESS CHEW ³	MC	8	REYATAZ
MC/DEL		ISENTRESS POWDER	MC/DEL	8	SELZENTRY
MC/DEL		LAMIVUDINE TABS	MC	8	STAVUDINE
MC/DEL		LAMIVUDINE/ZIDOVUDINE	MC	8	STRIBILD ¹
MC/DEL		LOPINAVIR-RITONAVIR SOL	MC/DEL	8	SYMFI ⁴
MC		LOPINAVIR-RITONAVIR TAB	MC/DEL	8	SYMFI LO ⁴
MC		ODEFSEY ¹	MC/DEL	8	SYM TUZA
MC/DEL		PREZCOBIX	MC/DEL	8	TRIZIVIR TABS
MC		PREZISTA ²	MC	8	TRUVADA ¹
MC/DEL		RITONAVIR TAB 100MG	MC/DEL	8	VIRACEPT TABS
MC		RUKOBIA ⁴	MC	8	VITEKTA
MC		SUNLENCA ⁴	MC	8	ZERIT
MC		SUSTIVA ¹	MC	8	VIDEX EC
MC		TIVICAY	MC	8	VIREAD TABS ¹
MC		TIVICAY PD	MC/DEL	8	ZIAGEN TABS
MC		TRIUMEQ ¹	MC/DEL	8	ZIAGEN SOL
MC		TROGARZO ⁴	MC/DEL	9	VIRAMUNE XR

[Use PA Form# 20420](#)

1. Quantity limit of one per day

2. Only preferred if Norvir script is in member's profile within the past 30 days of filling Prezista

3. Isentress Chewable will only be approved if between the age of 2-12 years old

4. Clinical PA required.

5. Only preferred for post-exposure prophylaxis.

Fuzeon: Prescriber is either an HIV specialist provider or has consulted with one. Documentation of genotype testing is supplied and shows that there is no other potent, appropriate two or three drug oral regimen available, AND patient has a positive HIV viral load within past 6 months while on his/her current antiretroviral regimen AND the drug will be prescribed with at least two other drugs that are likely to be active based on the genotype testing.

DDI: Reyataz requires prior authorization if it is currently being used in combination with either Prevacid, pantoprazole, Prilosec, or any currently non preferred PPI .

DDI: Norvir requires prior authorization if it is currently being used in combination with either Enblex 15mg or Vesicare 10mg.

DDI: Preferred Crixivan caps requires prior authorization if it is currently being used in combination with either Enblex 15mg or Vesicare 10mg.

DDI: The concomitant use of the following drugs with **Descovy**® is not recommended: tipranavir/ritonavir, St. John's wort, and the antimycobacterials rifabutin, rifampin, or rifapentine.

DDI: Administration with the following drugs: the anticonvulsants carbamazepine, oxcarbazepine, phenobarbital, and phenytoin; the antimycobacterials rifampin and rifapentine; proton pump inhibitors such as dexlansoprazole, esomeprazole, lansoprazole, omeprazole, pantoprazole, rabeprazole; systemic dexamethasone (more than a single dose); and St. John's wort with **Odefsey** is contraindicated.

Stribild: PA required; must provider rationale as to why the member's medical need cannot be met with preferred agents, particularly Genvoya or combinations of preferred and agents AND must be antiretroviral treatment-naïve or virologically controlled on current therapy (HIV-1RNA < copies/ml) AND be HBV negative AND not be combined with other anti-retroviral agents.

DDI: Tivicay will require prior authorization is used with nevirapine, oxcarbazepine, phenytoin, phenobarbital, carbamazepine, and St. John's wort.

DDI:Aatazanavir or darunavir and the following drugs are contraindicated (due to potential for serious and/or life-threatening events or loss of therapeutic effect): alfuzosin, dronedarone, rifampin, irinotecan, dihydroergotamine, ergotamine, methylergonovine, cisapride, St. John's wort, lovastatin, simvastatin, pimozone, nevirapine, sildenafil (when given as Revatio® for treatment of PAH), indinavir, triazolam, or PO midazolam will be non-preferred and require prior authorization if it is currently being used in combination with Tybost.

DDI: Combined P-gp, UGT1A1 and strong CYP3A inhibitors may significantly increase plasma concentrations of Sunlenca®. Concomitant administration of Sunlenca® with these inhibitors is not recommended.

Sunlenca: In combination with other antiretroviral(s) for the treatment of HIV-1 infection in heavily treatment-experienced adults with multidrug resistant HIV-1 infection failing their current

	MC MC MC/DEL		TYBOST VIREAD POW ZIDOVUDINE				antiretroviral regimen due to resistance, intolerance, or safety considerations.
CYTO-MEGALOVIRUS AGENTS	MC MC MC/DEL MC/DEL		CIDOFOVIR FOSCARNET SODIUM GANCICLOVIR VALGANCICLOVIR	MC MC/DEL MC/DEL MC/DEL		VALCYTE TABS FOSCAVIR LIVTENCITY ¹ PREVYMIS	<p>Use PA Form# 20420</p> <p>1. Must show failure or contraindication to all the following ganciclovir, valganciclovir, cidofovir and foscarnet before Livtency will be approved.</p> <p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>Prevymis: Documentation that member is high-risk for CMV reactivation as defined by transplant guidelines or that there has been significant myelosuppression by one of the preferred agents.</p> <p>DDI: Livtency is a substrate of CYP3A4. Coadministration of Livtency® with strong inducers of CYP3A4 is not recommended, except for selected anticonvulsants.</p>
HERPES AGENTS	MC/DEL MC/DEL		ACYCLOVIR VALACYCLOVIR HCL	MC/DEL MC MC/DEL MC MC/DEL	8 8 8 8 9	FAMCICLOVIR ¹ SITAVIG ZOVIRAX ¹ VALTREX TABS ¹ FAMVIR TABS ¹	<p>1. Must fail Acyclovir and Valacyclovir before non-preferred products in step order.</p> <p>Use PA Form# 20420</p> <p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved (in step order), unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p>
INFLUENZA AGENTS	MC MC MC/DEL		AMANTADINE CAPS RELENZA DISKHALER AEPB OSELTAMIVIR ¹	MC MC MC MC/DEL MC/DEL MC/DEL MC/DEL		AMANTADINE TABS FLUMADINE TABS FLUMIST RIMANTADINE HCL TABS TAMIFLU ¹ TAMIFLU SUS XOFLUZA	<p>1. Tamiflu and Oseltamivir 10 caps or 60cc's per month. Will be audited for presence of positive influenza tests in patient or family member.</p> <p>Use PA Form# 20420 for all others</p> <p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved (in step order), unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p>
IMMUNE SERUMS							
IMMUNE SERUMS	MC		HYPERRHO INJ				
HEPATITIS AGENTS							
HEPATITIS C AGENTS	MC MC MC/DEL MC/DEL MC/DEL MC MC/DEL		SOFOSBUVIR/VELPATASVIR ² (Authorized generic labeler 72626 Asegua Therapeutics) MAVYRET ² PEGASYS KIT ¹ PEGASYS SOLN PEG-INTRON KIT ¹ RIBAVIRIN RIBASPHERE	MC/DEL MC/DEL MC MC/DEL MC MC MC MC MC/DEL		COPEGUS TABS DAKLINZA EPCLUSA ² HARVONI ² REBETOL CAPS RIBAPAK SOVALDI ² VIEKIRA PAK ² VIEKIRA XR ² VOSEVI ZEPATIER ⁴	<p>1. Dosing limits apply, please see dosage consolidation list.</p> <p>2. Approvals will require clinical PA. Please see the Hepatitis PA form for criteria</p> <p>Use PA Form #10700</p> <p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved (in step order), unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>DDI: Olysio will require a prior authorization if it is currently being used in combination with drugs known to be significant CYP3A4 inhibitors (ketoconazole, itraconazole, clarithromycin, indinavir, nefazodone, nelfinavir, ritonavir, atazanavir, saquinavir and telithromycin).</p>
HEPATITIS AGENTS - MISC.				MC		ACTIMMUNE	<p>Use PA Form# 20420</p> <p>Approved for chronic granulomatous disease, osteopetrosis and idiopathic pulmonary fibrosis.</p>
HEPATITIS B ONLY	MC/DEL MC		ENTECAVIR TENOFIVIR	MC MC MC MC		BARACLUDE HEPSERA TABS TYZEKA VEMLIDY	<p>Use PA Form# 20420</p> <p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved (in step order), unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>Baraclude is indicated for treatment of chronic Hep B virus (HBV) in adults with: evidence of active viral replication AND either evidence of persistent elevation in serum aminotransferases (ALT or AST) or histologically active disease. Patient is 16 years of age or older. Boxed warning: Use not recommended for those co-infected with HIV and HBV who are not also receiving highly active antiretroviral therapy (HAART).</p>

							<p>Treating provider agrees to do a quantitative spot urine protein test before each dose</p> <p>Concomitant use of Spinraza and Zolgensma is investigational and will not be approved AND Use of Spinraza after gene replacement therapy, including Zolgensma is investigational and will not be approved</p> <p>Note: Initial approval will be granted for 4 loading doses (the first 3 loading doses should be administered at 14-day intervals; the 4th loading dose should be administered 30 days after the 3rd dose). Renewal may be granted for up to 12 months with a maximum of 3 doses approved per year (12mg (5ml) every 4 months). For therapy continuation, clinical documentation must be submitted documenting improvement or maintenance of motor ability OR slower progression of disease than would otherwise be expected.</p> <p>Use PA Form# 20420</p>	
NEUROLOGICS- RETT SUNDROME				MC		DAYBUE ¹²	<p>1. Clinical PA required for appropriate diagnosis</p> <p>2. For the treatment of patients 2 years of age and older.</p> <p>Use PA Form# 20420</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved (in step order), unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p>
ALS DRUGS	MC/DEL		RILUZOLE	MC MC MC MC MC		<p>EXSERVAN</p> <p>QALSODY</p> <p>RILUTEK TABS</p> <p>RADICAVA¹</p> <p>RELYVRIO¹</p> <p>TIGLUTIK</p>	<p>1. Clinical PA for indication required</p> <p>Use PA Form# 20420</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved (in step order), unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>Qalsody: For the treatment of amyotrophic lateral sclerosis (ALS) in adults who have a mutation in the superoxide dismutase 1 (SOD1) gene. Continued approval for this indication may be contingent upon verification of clinical benefit in confirmatory trial(s).</p>
MOVEMENT DISORDERS	MC MC MC MC		<p>AUSTEDO¹</p> <p>AUSTEDO XR¹</p> <p>INGREZZA¹</p> <p>TETRABENAZINE¹</p>	MC/DEL		XENAZINE	<p>1. Clinical PA required for appropriate diagnosis</p> <p>Use PA Form# 20420</p> <p>Use PA Form# 20710 for Xenazine</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved (in step order), unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>DDI: Avoid concomitant use of Ingrezza® with MAO inhibitors (e.g. isocarboxazid, phenelzine, or selegiline). Concomitant use with strong CYP3A4 inducers (e.g. rifampin, carbamazepine, phenytoin, St. John's wort) is not recommended</p>
MUSCULAR DYSTROPHY AGENTS	MC		EMFLAZA ^c	MC MC MC MC MC MC MC		<p>AGAMREE^c</p> <p>AMONDYS 45¹</p> <p>DEFLAZACORT</p> <p>ELEVIDYS³</p> <p>DUVYZAT</p> <p>EXONDYS 51¹</p> <p>VILTEPSO³</p> <p>VYONDYS 53</p>	<p>1. Clinical prior authorization to verify diagnosis and use of stable dose of corticosteroid for at least 6 months.</p> <p>2. For the treatment of Duchenne muscular dystrophy (DMD) in patients 2 years of age and older and a documented intolerance of oral corticosteroid.</p> <p>3. Clinical prior authorization to verify diagnosis and use of stable dose of corticosteroid</p> <p>4. For the treatment of Duchenne muscular dystrophy (DMD) in patients 2 years of age and older</p> <p>Use PA Form# 20420</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved (in step order), unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>Amondys 45, Exondys 51 and Vyondys 53: • The prescriber is, or has consulted with, a neuromuscular disorder specialist AND • The dose does not exceed 30mg/kg once weekly AND • The patient is currently on a stable corticosteroid dose for at least 6 months (at least 3 months for Elevidy).</p> <p>Amondys 45, Exondys 51, Vyondys 53 Note: Initial approval will be granted for 6 months. For re-approval after 6 months, the patient must demonstrate a response to therapy</p> <p>Duvyzat: The patient must meet the FDA approved age AND have a diagnosis of Duchenne Muscular Dystrophy confirmed with a confirmed mutation of the DMD gene AND</p> <ul style="list-style-type: none"> • The prescriber is, or has consulted with, a neuromuscular disorder specialist • The patient is ambulatory AND • The patient is currently on a stable corticosteroid dose for at least 6 months. AND • Baseline platelet counts are > 150 x 109/L and baseline triglycerides are < 300 mg/dL <p>Elevidys and Viltepsos: The prescriber is, or has consulted with, a neuromuscular disorder specialist AND • The dose does not exceed dosing AND • The patient is currently on a stable corticosteroid dose for at least 3 months.</p> <p>Viltepsos: For Duchenne muscular dystrophy (DMD) in patients who have a confirmed mutation of the DMD gene that is amenable to exon 53 skipping. Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial.</p>
MYASTHENIA GRAVIS	MC		<p>PYRIDOSTIGMINE</p>	MC MC		<p>MESTINON</p> <p>VYVGART¹</p>	<p>1. For the treatment of generalized myasthenia gravis (AMG) in adult</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved (in step order), unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between</p>

ESTROGENS - PATCHES / TOPICAL	MC MC/DEL MC/DEL		EVAMIST MINIVELLE PATCH VIVELLE-DOT PTTW	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	5 8 8 8 8	ESTRADIOL PTWK DIVIGEL ¹ CLIMARA PTWK ELESTRIN ¹ MENOSTAR PATCH	1. Step order drugs must be used in specified step order. Use PA Form# 20420	Approved for failures on multiple oral estrogen agents after 90 day trials or if unable to swallow any oral medication.
ESTROGENS - TABS	MC/DEL MC/DEL		ESTRADIOL PREMARIN TABS	MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL MC		ENJUVIA ESTRADIOL-NORETHINDRONE ESTRACE TABS ESTRATAB TABS MENEST TABS NORETHINDRON-ETHINYL ORTHO-EST TABS	Must fail preferred products before non-preferred products. Use PA Form# 20420	Preferred drugs must be tried for at least 90 days and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
ESTROGEN COMBO'S	MC/DEL MC/DEL MC/DEL		ANGELIQ COMBIPATCH PTTW PREMPHASE TABS PREMPRO TABS	MC/DEL MC/DEL MC MC/DEL MC/DEL		FEMHRT 1/5 TABS ¹ FYAVOLV LOPREEZA TAB ORTHO-PREFEST TABS ¹ SYNTEST H.S. TABS ¹	1. Must fail Premphase and Prempro products before non preferred products. Use PA Form# 20420	Preferred drugs must be tried for at least 90 days and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
PROGESTINS	MC/DEL MC/DEL MC MC		MEDROXYPROGESTERONE ACETA ¹ NORETHINDRONE ACETATE TABS ¹ 17-ALPHA HYDROXYPROGESTERONE PWDR PROGESTERONE CAPS	MC/DEL MC MC MC/DEL MC/DEL		AYGESTIN TABS CYCRIN TABS PROGESTERONE POWD PROMETRIUM CAPS PROVERA TABS	1. Must fail Medroxyprogesterone and Norethindrone products before non-preferred Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
ENDOMETRIOSIS								
CENTRAL PRECOCIOUS PUBERTY AGENTS	MC		FENSOLVI ¹				1. For pediatric patients 2 years of age and older with central precocious puberty (CPP). Use PA Form# 20420	
ENDOMETRIOSIS- NASAL	MC/DEL		SYNAREL (NASAL) SPRAY				Use PA Form# 20420	Synarel is also indicated for central precocious puberty
ENDOMETRIOSIS/ UTERINE FIBROIDS- ORAL	MC/DEL MC		ORLISSA ¹ MYFEMBREE ^{1,2}	MC		ORIAHNN ¹	1. Prior treatment of NSAID and hormonal contraceptives required 2. Limited to 24 months due to the risk of continued bone loss, which may not be reversible. Use PA Form# 20420	
ENDOMETRIOSIS- INJECTABLE	MC/DEL		DEPO-SUBQ PROVERA 104					

CONTRACEPTIVES

CONTRACEPTIVES - PROGESTIN ONLY	<p>MC/DEL MC/DEL MC MC MC/DEL MC/DEL</p>	<p>CAMILA TABS ERRIN INCASSIA TAB HEATHER TAB NORETHINDRONE ACETATE 0.35MG TABS SLYND</p>	<p>MC/DEL MC/DEL MC</p>	<p>JOLIVETTE NORA-BE TABS ORTHO MICRONOR TABS</p>	<p>Use PA Form# 20420</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>If member experienced adverse reactions, consider using Oral Contraceptives from other groups.</p> <p>DDI: Preferred Oral Contraceptives will now be non-preferred and require prior authorization if it is currently being used in combination with Tracleer.</p>
CONTRACEPTIVES - INJECTABLE	<p>MC/DEL</p>	<p>MEDROXYPROGESTERONE ACETATE 150mg IM</p>	<p>MC/DEL</p>	<p>DEPO-PROVERA 150 mg SUSP</p>	<p>Use PA Form# 20420</p>	<p>The preferred drug must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p>
CONTRACEPTIVE - EMERGENCY	<p>MC/DEL MC MC MC MC MC/DEL MC MC/DEL MC/DEL</p>	<p>ELLA ENCONTRA ONE STEP ECONTRA EZ NEW DAY OPCION OPTION 2 MY CHOICE MY WAY LEVONORGESTREL NEXT CHOICE¹</p>			<p>1. Allowed 2 tablets per 30 days without PA</p> <p>Use PA Form# 20420</p>	<p>Due to the extensive list of products, any covered emergency contraceptive product preferred is and available without a PA.</p>
CONTRACEPTIVES - PATCHES/ VAGINAL PRODUCTS	<p>MC MC MC MC/DEL</p>	<p>ELURYN¹ NUVARING RING¹ TWIRLA XULANE²</p>	<p>MC MC MC</p>	<p>ANNOVERA PHEXXI ZAFEMY</p>	<p>Use PA Form# 20420</p> <p>1. Quantity limit allowing 1 every 28 days with out PA.</p> <p>2. Dose limits apply allowing 3 patches per 28 days supply.</p>	<p>Approved if adequate clinical reason given why patient unable to comply with other preferred agents including long acting injectable.</p>
CONTRACEPTIVES- LONG ACTING REVERSIBLE	<p>MC/DEL</p>	<p>MIRENA</p>	<p>MC/DEL MC MC MC/DEL MC/DEL</p>	<p>KYLEENA LILETTA NEXPLANON PARAGARD SKYLA</p>		
CONTRACEPTIVES - MONOPHASIC COMBINATION O/C'S	<p>MC/DEL MC/DEL MC/DEL MC MC/DEL MC MC/DEL MC/DEL MC MC/DEL MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL</p>	<p>APRI TABS AVIANE TABS BALZIVA CRYSSELLE-28 TABS DESODEN TABS ESTARYLLA TAB HAILEY FE TAB ISIBLOOM TAB JUNEL FE TAB LARIN FE TAB LESSINA TAB LEVORA-28 TAB MILI TAB NORGESTIMATE-ETHINYL ESTRADIOL TAB MIBELAS 24 FE TAB MICROGESTIN FE TAB RECLIPSEN SAFYRAL TAB SPRITEC 28 TABS</p>	<p>MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL</p>	<p>BEYAZ BREVICON-28 TABS LESSINA-28 TABS LEVORA LOESTRIN FE 1/20 TABS LOESTRIN 1.5/30-21 TABS MICROGESTIN FE TABS LOESTRIN 1/20-21 TABS LO/OVRAL 21 TABS LO/OVRAL 28 TABS NEXTSTELLIS NORDETTE-28 TABS NORTREL OCELLA OVRAL PORTIA-28 TABS SAFYRAL</p>	<p>Use PA Form# 20420</p> <p>If member experienced adverse reactions, consider using Oral Contraceptives from other groups.</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>If member experienced adverse reactions, consider using Oral Contraceptives from other groups.</p> <p>DDI: Preferred Oral Contraceptives will now be non-preferred and require prior authorization if it is currently being used in combination with Tracleer.</p>

	<p>MC</p> <p>MC</p> <p>MC</p> <p>MC</p> <p>MC</p> <p>MC</p> <p>MC</p> <p>MC</p> <p>MC</p> <p>MC/DEL</p>	<p>HUMALOG JUNIOR KWIKPEN 100/ML</p> <p>HUMALOG MIX 75/25</p> <p>HUMALOG 50/50 VIAL</p> <p>HUMULIN INJ 70/30 KWIKPEN</p> <p>HUMULIN INJ 70/30</p> <p>HUMULIN R INJ U-500</p> <p>INSULIN ASPART PROT MIX 70-30</p> <p>INSULIN ASPART</p> <p>INSULIN LISPRO</p> <p>LANTUS SOLN</p>	<p>MC/DEL</p> <p>MC</p> <p>MC</p> <p>MC</p> <p>MC</p> <p>MC</p> <p>MC/DEL</p> <p>MC/DEL</p> <p>MC/DEL</p> <p>MC/DEL</p> <p>MC</p>	<p>AFREZZA¹</p> <p>BASAGLAR</p> <p>HUMALOG KWIKPEN U-200</p> <p>HUMULIN INJ 50/50</p> <p>HUMULIN N INJ U-100</p> <p>HUMULIN R U-100</p> <p>INSULIN DEGLUDEC</p> <p>LYUMJEV</p> <p>NOVOLIN</p> <p>NOVOLOG</p> <p>NOVOLOG MIX</p> <p>NOVOLOG MIX 70/30 FLEXPEN</p> <p>RELION</p>	<p>monotherapy. Obtain lab values of pulmonary function and recent smoking history</p> <p>2. For the treatment of patients ≥3 years of age</p>	
DIABETIC - PENFILLS	<p>MC</p> <p>MC</p> <p>MC</p> <p>MC</p> <p>MC/DEL</p> <p>MC</p> <p>MC</p> <p>MC</p> <p>MC/DEL</p> <p>MC/DEL</p> <p>MC/DEL</p>	<p>HUMALOG MIX KWIK 50/50</p> <p>HUMALOG MIX INJ 75/25 KWP</p> <p>HUMALOG KWIK INJ 100/ML</p> <p>HUMALOG KWIK INJ 200/ML</p> <p>HUMULIN R U-500 KWP</p> <p>INSULIN ASPART PROT MIX 70-30 PEN</p> <p>INSULIN ASPART PEN</p> <p>INSULIN LISPRO KWIKPEN U-100</p> <p>LANTUS SOLOSTAR</p> <p>TOUJEO MAX SOLOSTAR</p> <p>TOUJEO SOLOSTAR</p>	<p>MC</p> <p>MC/DEL</p> <p>MC/DEL</p> <p>MC/DEL</p> <p>MC/DEL</p> <p>MC</p> <p>MC/DEL</p>	<p>APIDRA OPTICLIK PEN</p> <p>NOVOLIN 70/30 PEN</p> <p>NOVOLOG MIX PENFILL</p> <p>NOVOLOG PENFILL SOLN</p> <p>NOVOLOG FLEXPEN</p> <p>NOVOLOG MIX 70/30 VIAL</p> <p>REZVOGLAR KWIKPEN</p> <p>TRESIBA</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved (in step order), unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>Use PA Form# 20420</p>	
DIABETIC - DPP- 4 ENZYME INHIBITOR	<p>MC/DEL</p> <p>MC/DEL</p>	<p>JANUVIA^{1,2}</p> <p>TRADJENTA²</p>	<p>MC/DEL</p> <p>MC/DEL</p> <p>MC/DEL</p> <p>MC</p>	<p>NESINA</p> <p>ONGLYZA²</p> <p>QTERN</p> <p>ZITUVIO</p>	<p>1. Preferred if therapeutic doses of metformin are seen in members drug profile for at least 60 days within the past 18 months or if phosphate binder is currently seen in the members drug profile.</p> <p>2. Dosing limits apply. Please refer to Dose consolidation list.</p> <p>Use PA Form# 20420</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved (in step order), unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>DDI: Onglyza 5mg will require a prior authorization if it is currently being used in combination with drugs known to be significant CYP3A4 inhibitors (ketoconazole, itraconazole, clarithromycin, indinavir, nefazodone, nelfinavir, ritonavir, atazanavir, saquinavir and telithromycin).</p>
DIABETIC - DPP- 4 ENZYME INHIBITOR-COMBO	<p>MC/DEL</p> <p>MC/DEL</p> <p>MC/DEL</p>	<p>JANUMET^{1,2}</p> <p>JANUMET XR^{1,2}</p> <p>JENTADUETO¹</p>	<p>MC/DEL</p> <p>MC/DEL</p> <p>MC</p> <p>MC/DEL</p> <p>MC</p> <p>MC</p>	<p>JENTADUETO XR</p> <p>KAZANO</p> <p>KOMBIGLYZE XR</p> <p>OSENI</p> <p>ZITUVIMET</p> <p>ZITUVIMET XR</p>	<p>1. Preferred if therapeutic doses of metformin are seen in members drug profile for at least 60 days within the past 18 months or if phosphate binder is currently seen in the members drug profile.</p> <p>2. Dosing limits apply. Please refer to Dose consolidation list.</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved (in step order), unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>Zituvimet/ Zituvimet XR: Approvals will require trial of preferred sitagliptin/metformin products or other preferred diabetic agents.</p>

DIABETIC - LANCET-LANCET DEVICE						Use PA Form# 20420		Use PA Form# 20420	Please refer to the MaineCare Preferred Diabetic Supply List available at www.mainearepdl.org
DIABETIC - SYRINGES-NEEDLES								Use PA Form# 20420	Please refer to the MaineCare Preferred Diabetic Supply List available at www.mainearepdl.org
DIABETIC - OTHER				MC/DEL MC		CYCLOSET SYMLIN		Use PA Form #20420 for all others	
SGLT 2 INHIBITORS	MC/DEL MC/DEL		FARXIGA JARDIANCE	MC/DEL MC/DEL		INVOKANA ¹ STEGLATRO		Use PA Form# 20420	1.Dosing limits apply please refer to Dose Consolidation List Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
SGLT 2 INHIBITOR COMBINATIONS	MC/DEL MC/DEL MC/DEL		SYNJARDY SYNJARDY XR XIGDOU XR	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		GLYXAMBI INVOKAMET INVOKAMET XR SEGLUROMET STEGLUJAN TRIJARDY XR		Use PA Form# 20420	Preferred drugs must be tried for at least 3 months at full therapeutic doses and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Glyxambi /Xigduo XR- Verify prior trials and failures or intolerance of preferred treatments from other diabetic categories Synjardy® XR is not recommended for patients with type 1 DM or for the treatment of diabetic ketoacidosis.
DIABETIC MONITOR	MC MC MC		ONE TOUCH ULTRA 2 KIT ONE TOUCH ULTRA MINI KIT TRUE METRIX	MC MC MC MC MC MC MC MC MC MC		ACCUCHECK ASCENSIA ASSURE CONTOUR BREEZE Z EXACTECH FREESTYLE INSULINX FREESTYLE LITE SYSTEM KIT ONE TOUCH ULTRA SMART KIT PRECISION XTRA METER PRODIGY TRUETRACK		Use PA Form# 20420	Effective October 25th 2007, approvals for all non preferred meters/ test strips will require medical necessity documenting clinically significant features that are not available on any of the preferred meters.
DIABETIC TEST STRIPS	MC MC		ONE TOUCH ULTRA ¹ TRUE METRIX	MC MC MC MC MC MC MC MC MC MC		ACCUCHECK ASCENSIA ASSURE CONTOUR BREEZE Z EXACTECH FREESTYLE FREESTYLE LITE FREESTYLE INSULINX ONE TOUCH DELICA PRECISION XTRA PRODIGY TRUETRACK		Use PA Form# 20420	1. Only 50 ct & 100 ct package size. Effective October 25th 2007, approvals for all non preferred meters/ test strips will require medical necessity documenting clinically significant features that are not available on any of the preferred meters.
INCRETIN MIMETIC	MC/DEL MC MC/DEL		RYBELSUS TRULICITY VICTOZA	MC/DEL MC/DEL MC/DEL	5 8 8	OZEMPIC ADLYXIN BYDUREON BCISE			Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved (in step order), unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.

			MC MC/DEL MC/DEL	8 8 8	MOUNJARO SOLIQUA XULTOPHY		Soliqua must try both insulin and a preferred incretin mimetic and have a medical necessity for use that is not based on convenience or simply due to the fact that one injection is needed instead of two.
DIABETIC - ORAL SULFONYLUREAS	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL		AMARYL TABS DIABETA TABS GLUCOTROL TABS GLUCOTROL XL TBCR GLYNASE TABS MICRONASE TABS	Use PA Form# 20420 Use PA Form# 20420 1. Pa required for members ≥65. Glyburide has a greater risk of severe prolonged hypoglycemia in older adults.	Preferred drugs must be tried for at least 3 months at full therapeutic doses and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. DDI: All sulfonylureas (except glyburide) will now be non-preferred and require prior authorization if it is currently being used with either ranitidine or cimetidine. DDI: Glimepiride will now be non-preferred and require prior authorization if it is currently being used with either fluconazole (except 150mg strength) or fluvoxamine. Amaryl is non-preferred but with any prior authorization requests, the member's drug profile will also be monitored for concurrent use with either fluconazole or fluvoxamine.
DIABETIC -ORAL BIGUANIDES	MC/DEL MC/DEL		MC MC MC MC/DEL		GLUCOPHAGE TABS GLUCOPHAGE XR TB24 FORTAMET METFORMIN ER OSMOTIC	Use PA Form# 20420	Preferred drug must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
DIABETIC - THIAZOL / BIGUANIDE COMBO			MC/DEL MC MC		ACTOPLUS MET ¹ ACTOPLUS MET XR AVANDARYL ¹ AVANDAMET TABS ¹	Use PA Form# 20420 1. Requires use of Actos, Metformin, or other preferred anti-diabetics.	DDI: Actos, Avandia, or any combination product with Actos or Avandia will now be non-preferred and require prior authorization if it is currently being used with gemfibrozil.
DIABETIC - / THIAZOL	MC/DEL		MC/DEL MC		ACTOS TABS ³ AVANDIA TABS ²	1. Pioglitazone HCL is non-preferred as monotherapy. Pioglitazone HCL is preferred if therapeutic doses of metformin, sulfonylurea or insulin are seen in members drug profile for at least 60 days within the past 18 months. 2. Current users of Avandia who have tried Actos will be able to continue use of Avandia. 3. Dosing limits apply please refer to Dose Consolidation List Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. DDI: Actos, Avandia, or any combination product with Actos or Avandia will now be non-preferred and require prior authorization if it is currently being used with gemfibrozil.
DIABETIC - ALPHAGLUCOSIDASE	MC/DEL		MC		PRECOSE TABS		Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Use PA Form# 20420
DIABETIC - SULFONYLUREA / BIGUANIDE	MC/DEL		MC MC MC/DEL		GLUCOVANCE TABS ¹ METAGLIP TABS ¹ DUETACT ²	1. Use individual ingredients. 2. Use Actos with generic glimepiride. Use PA Form# 20420	Approved for patients failing to achieve good diabetic control with maximal doses of individual components.
DIABETIC - MEGLITINIDES	MC		MC/DEL MC/DEL		PRANDIN TABS STARLIX TABS	Use PA Form# 20420	Preferred drugs from other diabetic sub-categories must be tried and failed due to lack of inadequate diabetic control or intolerable side effects before non-preferred drug will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.

DDI: Prandin is non-preferred but with any prior authorization requests, the member's drug profile will also be monitored for current use with both Sporanox and gemfibrozil, due to a significant drug-drug interaction.

GLUCOSE ELEVATING AGENTS

GLUCOSE ELEVATING AGENTS	MC/DEL MC/DEL		BAQSIMI ¹ GVOKE ²	MC MC MC		GLUCAGON DIAGNOSTIC KIT ZEGALOGUE ³	Use PA Form# 20420 1. For the treatment of patients ≥ 4 years of age. 2. For the treatment of patients ≥ 2 years of age. 3. For the treatment of patients ≥ 6 years of age.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved (in step order), unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
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THYROID

THYROID EYE DISEASE				MC		TEPEZZA	Use PA Form# 20420	
THYROID HORMONES	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		ARMOUR THYROID TABS CYTOMEL TABS ERMEZA ¹ LEVOTHROID TABS LEVOTHYROXINE SODIUM TABS LEVOXYL TABS UNITHROID TABS	MC MC/DEL MC MC/DEL		LEVOTHYROXINE SODIUM SOLR LIOTHYRONINE SYNTHROID TABS THYQUIDITY	Use PA Form# 20420 1. Clinical PA is required to confirm diagnosis of dysphagia.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
ANTITHYROID THERAPIES	MC/DEL MC/DEL		METHIMAZOLE TABS PROPYLTHIOURACIL TABS	MC/DEL		TAPAZOLE TABS	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.

CUSHING DISEASE AGENTS

CUSHING DISEASE AGENTS				MC MC		ISTURISA ¹ RECORLEV	1. For the treatment of adult patients with Cushing's disease for whom pituitary surgery is not an option or has not been curative. Use PA Form #20420	Recorlev® is associated with dose-related QT interval prolongation. QT interval prolongation may lead to life-threatening ventricular dysrhythmias such as Torsade de pointes.
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OSTEOPOROSIS / BONE AGENTS

OSTEOPOROSIS	MC/DEL		ALENDRONATE	MC/DEL MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC MC/DEL		ACTONEL TABS AREIDIA SOLR BINOSTO BONIVA INJECTION KIT BONIVA TABS ^{2,4} CALCITONIN NS DUAVEE DIDRONEL TABS EVISTA TABS ¹ EVENTY ² FORTEO FORTICAL	Use PA Form# 20420 1. Approval only requires failure of Alendronate. 2. Quantity limits apply, please see dosage consolidation list. 3. Please use Alendronate and Vitamin D. 4. Please use other preferred agents.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Binosto use preferred generic alendronate tablets Eventy® should be limited to 12 monthly doses Sohonos: For the reduction in volume of new heterotopic ossification in adults and pediatric patients aged 8 years and older for females and 10 years and older for males with fibrodysplasia ossificans progressive (FOP).
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				MC/DEL MC MC MC MC MC/DEL		FOSAMAX TABS AND PLUS D ³ PROLIA SOHONOS ⁶ STRENSIQ ⁵ TYMLOS XGEVA ZOMETA	5. Obtain baseline ophthalmology exams and renal ultrasounds and then periodically during treatment 6. Clinical PA for indication required.	
FIBROBLAST GROWTH FACTOR 23 INHIBITORS	MC		CRYSVITA ¹				1. Preferred for patients <21 years for the treatment of X-linked hypophosphatemia. Use PA Form #20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
CALCIMIMETIC AGENTS								
CALCIMIMETIC AGENTS				MC MC		PARSABIV SENSIPAR	Use PA Form# 30115	For Sensipar baseline PTH, Ca, and phosphorous levels are required and initial approvals will be limited to 3 months. Subsequent approvals will require additional levels being done to assess changes. Will not approve if baseline Ca is less than 8.4. Parsabiv is for the treatment of secondary hyperparathyroidism (HPT) in adults with chronic kidney disease (CKD) on hemodialysis. Parsabiv® has not been studied in adults with parathyroid carcinoma, primary hyperparathyroidism, or with chronic kidney disease who are not on hemodialysis and is not recommended for use in these populations.
GROWTH HORMONE								
GROWTH HORMONE	MC/DEL MC/DEL MC		GENOTROPIN ¹ NORDITROPIN SOLN ¹ SKYTROFA ^{1,2}	MC MC MC/DEL MC MC MC/DEL MC/DEL	8 8 8 8 8 8 8	HUMATROPE SOLR INCRELEX NUTROPIN NGENLA OMNITROPE SAIZEN SOLR SOGROYA TEV-TROPIN	Use PA Form# 10710 1. Clinical PA is required to establish diagnosis and medical necessity. 2. Preferred after single step therapy of short acting growth hormone.	See Growth Hormone PA form for criteria. Step-order will still apply unless clinical contraindication supplied. Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
ACHONDROPLASIA TREATMENT				MC		VOXZOGO ¹	1. Pediatric patients with achondroplasia who are 5 years of age and older with open epiphyses. Use PA Form# 20420	Voxzogo: To increase linear growth in pediatric patients with achondroplasia who are 5 years of age and older with open epiphyses. This indication is approved under accelerated approval based on an improvement in annualized growth velocity. Continued approval for this indication may be contingent upon verification and description of clinical benefit in confirmatory trial(s).
SOMATOSTATIC AGENTS				MC/DEL MC MC MC/DEL MC	7 8 8 8 8	OCTREOTIDE INJ ¹ BYNFEZIA ¹ MYCAPSSA ¹ SANDOSTATIN ¹ SOMATULINE ¹	Use PA Form# 10710 1. Non-preferred products must be used in specified step order.	
GROWTH HORMONE ANTAGONISTS								
GH ANTAGONISTS				MC		SOMAVERT	Use PA Form# 10710	Approved for acromegaly patients failing surgery/radiation/drug therapy including bromocriptine and sandostatin.
VASOPRESSIN RECEPTOR ANTAGONIST								
VASOPRESSIN RECEPTOR ANTAGONIST				MC MC/DEL		JYNARQUE ¹ SAMSCA	Use PA Form# 20420 1. Clinical PA required for appropriate diagnosis	Samsca Drug Warning- Avoid use in patients with underlying liver disease, including cirrhosis, because the ability to recover from liver injury may be impaired. Limit duration of therapy to 30 days to minimize the risk of liver injury. DDI: Jynarque- Concomitant use with strong CYP3A inhibitors is contraindicated. Avoid concomitant use of Jynarque® with OATP1B1/B3 and OAT3 substrates (e.g. statins, bosentan, glyburide, nateglinide, repaglinide, methotrexate, furosemide).
URINARY INCONTINENCE								

VASOPRESSINS	MC/DEL MC/DEL		DESMOPRESSIN TABS DDAVP SOLN	MC/DEL MC/DEL MC MC/DEL MC MC/DEL	5 6 8 8 8 8	DDAVP TABS DESMOPRESSIN SPRAY ¹ DESMOPRESSIN ACETATE SOLN ¹ NOCDURNA ¹ NOCTIVA ¹ STIMATE SOLN ^{1,2}	1. Products must be used in specified step order. Nocturnal enuresis patients will be encouraged to periodically attempt stopping DDAVP. 2. Patients with a diagnosis of hemophilia or Von Willebrand's disease will be exempt from prior authorization. Use PA Form# 20420	Approved for central diabetes insipidus and for nocturnal enuresis. For nocturnal enuresis- must be over 6 years old, must fail an adequate trial of alarm training (higher success rate, lower relapse rate) and must periodically attempt wearing (at 6 month intervals).	
ANTISPASMODICS	MC/DEL MC/DEL		OXYBUTYNIN TOLTERODINE	MC/DEL MC/DEL MC/DEL MC/DEL	8 8 8	DARIFENACIN ER TAB DITROPAN FLAVOXATE HCL TAB	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.	
ANTISPASMODICS - LONG ACTING	MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		FESOTERODINE GELNIQUE GEL PACKET MYRBETRIQ OXYBUTYNIN ER TABS OXYTROL SOLIFENACIN SUCCINATE TAB TROSPIUM	MC MC/DEL MC MC/DEL MC/DEL MC MC	8 8 8 8 8 8 8	DITROPAN XL TBCR ENABLEX ^{1,2} GEMTESA ² TOLTERODINE TAB TOVIAZ VESICARE ¹ VESICARE ³ LS	Use PA Form# 20420 1. See Criteria Section. 2. Use a preferred long acting antispasmodic. 3. For the treatment of patients ≥ 2 years of age.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. 1. Vesicare 5mg and Enablex 7.5mg maximum doses if given with drugs known to be significant CYP3A4 inhibitors.(Ketoconazole, Sporanox, Erythromycin, Fluconazole, Nefazodone, Nelfinavir, and Ritonavir) DDI: Enablex 15mg and Vesicare 10mg will now be non-preferred and require prior authorization if they are currently being used in combination with any of the following medications: clarithromycin, erythromycin, Ketek, Crixivan, Norvir, ketoconazole, fluconazole (except 150mg strength), Sporanox, nefazodone, or diltiazem.	
CHOLINERGIC	MC/DEL		BETHANECHOL	MC/DEL		URECHOLINE	Use PA Form# 20420		
HYPERAMMONIA TREATMENTS	MC		CARGLUMIC ACID TABS	MC		CARBAGLU TABS	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.	
UREA CYCLE DISORDER	MC MC		BUPHENYL TABLET PHEBURANE GRANULES	MC MC MC MC/DEL MC/DEL		BUPHENYL POWDER RAVICTI LIQUID OLPRUVA SODIUM PHENYL BUTYRATE POWDER SODIUM PHENYL BUTYRATE TAB	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Olpruva: As adjunctive therapy to standard of care, which includes dietary management, for the chronic management of adult and pediatric patients weighing 20kg or greater and with a body surface area (BSA) of 1.2m ² or greater, with urea cycle disorders (UCDs) involving deficiencies of carbamylphosphate synthetase (CPS), ornithine transcarbamylase (OTC), or argininosuccinic acid synthetase (AS).	
METABOLIC MODIFIER									
HERED. TYROSINEMIA				MC		ORFADIN	Use PA Form# 20420	Approved for Type 1 hereditary tyrosinemia patients. Must include laboratory evidence of dx at first PA.	
FABRY DISEASE AGENTS				MC MC MC/DEL		ELFABRIO ¹ FABRAZYME ² GALAFOLD ¹	1.Clinical PA to verify appropriate diagnosis. 2.For the treatment of patients 2 years of age and older. Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Elfabrio and Galfold: For the treatment of adults with confirmed Fabry disease.	
ANTIHYPERTENSIVES / CARDIAC									
CARDIAC GLYCOSIDES	MC/DEL		DIGITEK TABS				Use PA Form# 20420		

	MC/DEL MC/DEL	DIGOXIN LANOXIN					
CARDIAC MYOSIN INHIBITORS				MC		CAMZYOS	<p>Use PA Form# 20420</p> <p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>Camzyos: For the treatment of adults with symptomatic New York Heart Association (NYHA) class II-III obstructive hypertrophic cardiomyopathy (HCM) to improve functional capacity and symptoms.</p> <p>DDI: Concomitant use of Camzyos® with a moderate to strong CYP2C19 inhibitor or a strong CYP3A4 inhibitor is contraindicated.</p>
CARDIAC - SINUS NODE INHIBITORS				MC		CORLANOR	<p>Use PA Form#20420</p> <p>In patients with stable, symptomatic chronic heart failure with left ventricular ejection fraction ≤35%, who are in sinus rhythm with resting heart rate ≥70 beats per minute (bpm) and either</p>
CARDIAC- ERAs				MC		TRYVIO	<p>Use PA Form#20420</p> <p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>Tryvio: In combination with other antihypertensive drugs, is indicated for the treatment of resistant hypertension, to lower blood pressure (BP) in adult patients who are not adequately controlled on other drugs. Resistant HTN is defined as a patient who takes at least 3 different class antihypertensive medications with complementary mechanisms including thiazide, ACE inhibitor, ARB, long-acting calcium channel blocker, with a trial of spironolactone, unless contra-indicated</p>
CARDIAC- SOLUBLE GUANYLATE CYCLASE STIMULATORS				MC/DEL		VERQUVO	<p>Use PA Form# 20420</p>
CARDIAC RISK REDUCTION- SGLT2/GLP-1				MC MC MC/DEL		INPEFA ¹ LODOCO WEGOVY	<p>1. To reduce the risk of cardiovascular death, hospitalization for heart failure, and urgent heart failure visit in adults with: Heart failure or Type 2 diabetes mellitus, chronic kidney disease, and other cardiovascular risk factors.</p> <p>Other Preferred SGLT inhibitors must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>Lodoco: Patient must have tried and failed generic colchicine due to lack of efficacy or intolerable side effects</p> <p>Wegovy: Patient has BMI > 27 kg/m2, and is not being used for weight loss only Patient has history of at least one of the following: o Stroke o Myocardial Infarction o Symptomatic peripheral arterial disease</p> <p>Patient does not have diagnosis of diabetes, end stage renal disease/dialysis, or HFrEF (EF < 45%)</p>
ANTIANGINALS--Isosorbide Di-nitrate/ Mono-Nitrates	MC/DEL MC/DEL		ISOSORBIDE MONONITRATE TABS ISOSORBIDE MONONITRATE ER	MC MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC		DILATRATE SR CPCR ISORDIL TABS ISORDIL TITRADOSE TABS ISOSORBIDE DINITRATE SUBL ISOSORBIDE DINITRATE TABS ISOSORBIDE DINITRATE CR TBCR ISOSORBIDE DINITRATE ER TBCR ISOSORBIDE DINITRATE TD TBCR IMDUR TB24 ISMO TABS MONOKET TABS	<p>Use PA Form# 20420</p> <p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p>

NITRO - OINTMENT/CAP/CR	MC/DEL MC/DEL MC MC		NITROBID OINT NITROGLYCERIN CPR NITROL OINT NITRO-TIME CPR			Use PA Form# 20420	
NITRO - PATCHES	MC/DEL MC/DEL	1 1	NITROGLYCERIN PT24 ¹ NITRO-DUR PT 24 0.8MG ¹	MC MC/DEL		NITRODISC PT24 NITRO-DUR PT24	1. At least 2 step 1's and step 3 of the preferred products must be used in specified order or PA will be required. Use PA Form# 20420 Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
NITRO - SUBLINGUAL/ SPRAY	MC/DEL		NITROSTAT SUBL	MC/DEL MC MC		NITROQUICK SUBL NITROLINGUAL SOLN NITROLINGUAL TABS	Use PA Form# 20420 Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
BETA BLOCKERS - NON SELECTIVE	MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL		CARVEDILOL LEVATOL TABS NADOLOL TABS PINDOLOL TABS PROPRANOLOL HCL SOLN ¹ PROPRANOLOL HCL TABS ¹ PROPRANOLOL HCL 60MG TABS PROPRANOLOL LA CAPS RANOLAZINE ER TABS SOTALOL AF SOTALOL HCL TABS TIMOLOL MALEATE TABS	MC MC/DEL MC MC MC/DEL MC/DEL MC/DEL MC MC MC MC		ASPRUZYO BETAPACE TABS BETAPACE AF TABS COREG CR ³ COREG TABS CORGARD TABS INDERAL TABS HEMANGEOL SOL INDERAL XL CAP INDERAL LA CPR INNOPRAN XL RANEXA	1. Recommend using BID since its effects do not last 24 hours. 2. Please use other strengths in combination to obtain this dose. 3. Dosing limits still apply. Please see dose consolidation list Use PA Form# 20420 Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. DDI: Concomitant use of Ranolazine products with strong CYP3A inhibitors, including ketoconazole, itraconazole, clarithromycin, nefazodone, nelfinavir, ritonavir, indinavir, and saquinavir, is contraindicated.
BETA BLOCKERS - CARDIO SELECTIVE	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		ACEBUTOLOL HCL CAPS ATENOLOL TABS ¹ BETAXOLOL HCL TABS BISOPROLOL FUMARATE TABS BYSTOLIC METOPROLOL TARTRATE TABS ¹ METOPROLOL ER NEBIVOLOL HCL TAB	MC MC/DEL MC MC/DEL MC/DEL MC/DEL		KERLONE TABS LOPRESSOR TABS SECTRAL CAPS TENORMIN TABS TOPROL XL TB24 ZEBETA TABS	1. Recommend using Atenolol (and metoprolol) BID since its effects do not last 24 hours. Use PA Form# 20420 Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
BETA BLOCKERS - ALPHA / BETA	MC/DEL		LABETALOL HCL TABS	MC		TRANDATE TABS	Use PA Form# 20420 Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
BETA BLOCKERS & DURECTIC COMBOS	MC/DEL		METOPROLOL-HYDROCHLOROTHIAZIDE TAB	MC/DEL		DUTOPROL	Use PA Form# 20420
CALCIUM CHANNEL BLOCKERS-- Amlodipine, Bepridil, Diltiazem, Felodipines, Isradipines, Nifedipines, Nisoldipine, and Verapamil	MC/DEL		AMLODIPINE ¹	MC/DEL MC MC/DEL		KATERZIA NORLIQVA NORVASC TABS ¹	1. Dosing limits apply, please see dose consolidation list. Use PA Form# 20420
	MC MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL		DILTIA XT CP24 DILTIAZEM HCL ER CP24 DILTIAZEM HCL XR CP24 DILTIAZEM CD 300MG CP24 DILTIAZEM CD 360MG CP24 CARTIA XT CP24 ¹ DILTIAZEM CD CP24 ¹ DILTIAZEM HCL ER CP24 ¹ DILTIAZEM XR CP24 ¹ TIAZAC CP24 ¹	MC/DEL MC/DEL MC MC MC MC MC/DEL MC/DEL MC/DEL	5 6 8 8 8 8 8 8 8	DILACOR XR CP24 ¹ TAZTIA ¹ CARDIZEM TABS ¹ CARDIZEM CD CP24 ¹ CARDIZEM LA TB24 ¹ CARDIZEM SR CP12 ¹ DILTIAZEM HCL TABS ¹ DILTIAZEM HCL ER CP12 ¹ DILTIAZEM HCL ER CP12 ¹	1. Products must be used in specified order or PA will be required. Just write "Diltiazem 24-hour" and the pharmacy will use a preferred long acting diltiazem that does not require PA. Use PA Form# 20420 Preferred drugs must be tried and failed (in step-order) due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. DDI: All preferred diltiazem will now be non-preferred and require prior authorization if they are currently being used in combination with either Enablex 15mg or Vesicare 10mg. All non-preferred diltiazem require prior authorization, but with any prior authorization request, the member's drug profile will also be monitored for current use with Enablex 15mg or Vesicare 10mg.
				MC/DEL		PLENDIL TB24	Use PA Form# 20420 Other Preferred calcium channel blockers must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable

			MC/DEL		FELODIPINE		clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.		
			MC MC		DYNACIRC CAPS DYNACIRC CR TBCR ¹	Use PA Form# 20420 1. Established users will be grandfathered	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.		
			MC MC		CARDENE SR CPCR NICARDIPINE HCL CAPS	Use PA Form# 20420	Other Preferred calcium channel blockers must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.		
			MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		ADALAT CC TBCR ¹ NIFEDIPINE CAPS PROCARDIA CAPS PROCARDIA XL TBCR	1. Established users of Adalat CC are grandfathered. Use PA Form# 20420	Preferred drug must be tried and failed in step order due to lack of efficacy or intolerable side effects before non-preferred drugs in step order will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.		
			MC MC		SULAR TB24 SULAR CR ¹	1. Established users of 10MG and 20MG strengths are grandfathered. Use PA Form# 20420			
		1 1 1	MC/DEL MC/DEL MC/DEL		VERAPAMIL HCL CR TBCR VERAPAMIL HCL ER TBCR VERAPAMIL HCL SR TBCR	MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	CALAN TABS CALAN SR TBCR COVERA-HS TBCR ISOPTIN-SR VERAPAMIL HCL ER CP24 VERAPAMIL HCL SR CP24 VERAPAMIL HCL TABS VERELAN CP24 VERELAN PM CP24 Use PA Form# 20420	Products must be used in specified order or PA will be required. Just write "Verapamil 24-hour" and the pharmacy will use a preferred long acting generic that does not require PA. Preferred drugs must be tried and failed (in step-order) due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.	
ANTIARRHYTHMICS			MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL		AMIODARONE HCL DISOPYRAMIDE FLECAINIDE MEXILETINE HCL PROCAINAMIDE PROPAFENONE QUINAGLUTE QUINIDINE GLUCONATE QUINIDINE SULFATE	MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL MC MC/DEL	CORDARONE DISOPYRAMIDE MULTAQ NORPACE PACERONE QUINIDEX TAMBOCOR TIKOSYN ¹ RYTHMOL SR RYTHMOL	1. Prescription must be written by Cardiologist. Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. DDI: Amiodarone will now be non-preferred and require prior authorization if it is currently being used in combination with either Lovastatin (doses greater than 40mg/day) or Lipitor (doses greater than 20mg/day) or Levofloxacin or Gemifloxacin, or Moxifloxacin, or Ofloxacin. DDI: Multaq will be preferred unless the following medications are seen in the member's drug profile within the last 35 days for brand name medications or 90 days for generic medications: Erythromycin, Amiodarone and other antiarrhythmics, TCA's, Phenothiazine, Ketoconazole, Itraconazole, Voriconazole, Cyclosporine, Telithromycin, Clarithromycin, Nefazodone, Ritonavir.
ACE INHIBITORS			MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		BENAZEPRIL HCL CAPTOPRIL TABS ENALAPRIL MALEATE TABS FOSINOPRIL SODIUM LISINAPRIL TABS RAMIPRIL QUINAPRIL HCL	MC MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL MC MC MC/DEL MC MC/DEL	5 MAVIK TABS 5 ACCUPRIL TABS 8 ACEON TABS ¹ 8 ALTACE CAPS ¹ 8 EPANED 8 LOTENSIN TABS ¹ 8 MOEXIPRIL HCL ¹ 8 MONOPRIL HCT TABS ¹ 8 PRINIVIL TABS ¹ 8 QBRELIS 8 UNIVASC ¹ 8 VASOTEC TABS ¹ 8 ZESTRIL TABS ¹	1. Non-preferred products must be used in specified order. Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs (in step-order) will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Non-preferred products are subject to step-order requirements unless clinical circumstances warrant exception.
ANGIOTENSIN RECEPTOR BLOCKER			MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		AMLODIPINE-OLMESARTAN TAB ³ IRBESARTAN ¹ LOSARTAN ¹ MICARDIS TABS ³ OLMESARTAN ¹ TELMISARTAN ¹	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	8 ATACAND TABS 8 AVAPRO 8 BENICAR TABS 8 COZAAR 8 DIOVAN 8 EDARBI	Use PA Form# 20420 1. Dosing limits apply, please see dose consolidation list. 2. Use preferred active ingredients which are available at PA	Per best practices patient should have trialed prior therapy of ACE inhibitor or currently on a diabetic therapy

			MC	8	TEVETEN TABS	available without PA. 3. Preferred without a PA only if patient on a diabetic therapy or prior ACE therapy.
DIRECT RENIN INHIBITOR			MC/DEL MC/DEL MC/DEL		AMTURNIDE TEKTURNA ¹ TEKAMLO	1. Must show failure of single and combination therapy from all preferred antihypertensive categories. Use PA Form# 20420
ANTIHYPERTENSIVES - CENTRAL	MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL		MC/DEL MC/DEL MC MC MC/DEL MC MC/DEL		CLONIDINE HCL TABS GUANFACINE HCL TABS HYDRALAZINE HCL TABS HYLOREL TABS METHYLDOPA TABS MINOXIDIL TABS PRAZOSIN HCL CAPS RESERPINE TABS CLONIDINE PATCH CLONIDINE TTS GUANABENZ ACETATE TABS ISMELIN TABS MINIPRESS CAPS NEXICLON TENEX TABS	Use PA Form# 20420 Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
ACE INHIBITORS AND CA CHANNEL BLOCKERS			MC/DEL MC MC MC/DEL	8 8 8 9	AMLODIPINE/BENAZEPRIL PRESTALIA ¹ TARKA TBCR LOTREL CAPS	1. Prestalia will only be approved for patients ≥ 18 years of age. Use individual preferred generic medications. Use PA Form# 20420
ACE AND THIAZIDE COMBO'S	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		MC/DEL MC MC/DEL MC/DEL MC MC/DEL		ACCURETIC TABS MONOPRIL HCT TABS PRINZIDE TABS UNIRETIC TABS VASERETIC TABS ZESTORETIC TABS BENAZEPRIL HCL/HYDROCHLOR CAPTOPRIL/HYDROCHLOROTHIA ENALAPRIL MALEATE/HCTZ TABS LISINAPRIL-HCTZ TABS LOTENSIN HCT TABS	Use PA Form# 20420 Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
BETA BLOCKERS AND DIURETIC COMBO'S	MC/DEL MC/DEL MC/DEL		MC/DEL MC/DEL MC MC MC/DEL		CORZIDE TABS LOPRESSOR HCT TABS TENORETIC TIMOLIDE 10/25 TABS ZIAC TABS ATENOLOL/CHLORTHALIDONE BISOPROLOL FUMARATE/HCTZ PROPRANOLOL/HCTZ	Use PA Form# 20420 Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
ARB'S AND CA CHANNEL BLOCKERS	MC/DEL MC/DEL MC/DEL		MC/DEL MC MC/DEL MC/DEL		AZOR BYVALSON EXFORGE EXFORGE HCT AMLODIPINE/VALSARTAN AMLODIPINE/VALSARTAN HCT TRIBENZOR	DDI: Byvalson will be non-preferred and require a prior authorization if it is currently being used in combination with drugs known to be significant CYP2D6 inhibitors (e.g. quinidine, propafenone, fluoxetine, paroxetine). Per best practices patient should have trialed prior therapy of ACE inhibitor or currently on a diabetic therapy Use PA Form# 20420
ARB'S AND DIURETICS	MC/DEL MC/DEL MC/DEL MC/DEL		MC/DEL MC/DEL MC MC/DEL MC MC/DEL	7 8 8 8 8 8	IRBESARTAN HYDROCHLOROTHIAZIDE ATACAND HCT TABS AVALIDE TABS ¹ DIOVAN HCT TABS ¹ HYZAAR TABS TEVETEN HCT TABS	1. Dosing limits apply, please see dose consolidation list. Use PA Form# 20420 Per best practices patient should have trialed prior therapy of ACE inhibitor or currently on a diabetic therapy
ANGIOTENSIN MODULATORS-ARB COMBINATION	MC		MC/DEL MC		EDARBYCLOL ENTRESTO SPRINKLES ENTRESTO	Use PA Form# 20420
ARB'S AND DIRECT RENIN INHIBITOR COMBINATION			MC/DEL		VALTURNIA	Use PA Form# 20420
DIURETICS	MC/DEL MC/DEL MC/DEL		MC/DEL MC/DEL MC/DEL		ALDACTAZIDE TABS ALDACTONE TABS AMILORIDE HCL ACETAZOLAMIDE TABS BUMETANIDE CHLOROTHIAZIDE TABS	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.

	MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC	CHLORTHALIDONE TABS EDECRIN TABS EDECRIN TABS HYDROCHLOROTHIAZIDE INDAPAMIDE TABS METHAZOLAMIDE TABS METHYLCLOTHIAZIDE TABS SPIRONOLACTONE SPIRONOLACTONE/HYDRO TORSEMIDE TABS TRIAMTERENE/HCTZ ZAROXOLYN TABS	MC/DEL MC/DEL MC MC MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC	BUMEX TABS DEMADEX TABS DIAMOX DIURIL DYAZIDE CAPS CAROSPIR ENDURON TABS FUROSCIX INSPIRA KERENDIA KEVEYIS LASIX TABS MAXZIDE MICROZIDE CAPS MIDAMOR TABS NAQUA TABS		Furoscix: The indication for use is the treatment of congestion due to fluid overload in adults with NYHA Class II or Class III chronic heart failure AND the medication is being prescribed by or in consultation with a cardiologist AND the patient is experiencing symptoms despite compliance with oral loop diuretic therapy AND oral loop diuretic therapy will be resumed as soon as practical AND medical reasoning beyond convenience is provided for not pursuing therapy in an outpatient infusion setting. PA approval will be authorized for 1 month. DDI: The concomitant use of Keveysi® with high dose aspirin is contraindicated. Kerendia: Patient must be on max tolerated preferred ACE-I/ARB and SGLT-2
CCB / LIPID			MC/DEL	CADUET	Use PA Form# 20420	
NEUROGENIC ORTHOSTATIC HYPOTENSION						
NEUROGENIC ORTHOSTATIC HYPOTENSION			MC	NORTHERA	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
LIPID DRUGS						
CHOLESTEROL - BILE SEQUESTRANTS	MC/DEL MC/DEL	CHOLESTYRAMINE COLESTIPOL HCI	MC/DEL MC/DEL MC MC/DEL	COLESTID PREVALITE QUESTRAN WELCHOL TABS	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
CHOLESTEROL - FIBRIC ACID DERIVATIVES	MC/DEL MC/DEL MC/DEL	FENOFIBRATE TAB GEMFIBROZIL TABS NIACIN ER	MC MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL MC MC	ANTARA LOPID FENOFIBRATE 120mg TAB FENOFIBRATE CAP FIBRICOR LIPOFEN LOFIBRA NIASPAN ER TRICOR TRIGLIDE	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. DDI: Fenofibrate is preferred but will require a prior authorization requests if used concurrent with Warfarin. DDI: Gemfibrozil will now be non-preferred and require prior authorization if it is currently being used with any of the following medications: Prandin, Actos, Avandia, any Avandia/Actos combination product, any HMG-COA Reductase Inhibitors (statins), or Warfarin.
CHOLESTEROL - HMG COA + ABSORB INHIBITORS MORE POTENT DRUGS/COMBINATIONS	MC/DEL MC/DEL MC MC/DEL	ATORVASTATIN EZETIM/SIMVA TAB ROSUVASTATIN SIMVASTATIN ¹	MC MC/DEL MC/DEL MC MC/DEL MC MC/DEL MC/DEL MC MC	ATORVALIQ CRESTOR EZALLOR SPRINKLES ³ FLOLIPID LIPITOR LIPTRUZET ZOCOR SIMVASTATIN 80MG ^{1,2} VYTORIN	1. Dosing limits apply, please see dosage consolidation list. 2. Current users grandfathered. 3. For the treatment of patients ≥ 18 years of age. Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. DDI: Lipitor (doses greater than 20mg/day) will now be non-preferred and require prior authorization if they are currently being used in combination cyclosporine. DDI: Lipitor (doses greater than 20mg/day) will now be non-preferred and require prior authorization if it is currently being used in combination with Amiodarone. DDI: All preferred statins will now be non-preferred and require prior authorization if it is currently being used in combination with Gemfibrozil.
CHOLESTEROL - HMG COA + ABSORB INHIBITORS LESS POTENT DRUGS/COMBINATIONS	MC/DEL MC/DEL MC/DEL	EZETIMIBE TABS LOVASTATIN TABS ² PRAVASTATIN ²	MC MC/DEL MC/DEL	8 8 8	ALTOPREV TB24 FLUVASTATIN TAB ER LESCOL XL TB24	2. Dosing limits apply, please see dosage consolidation list. Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Zetia will be approved for patients unable to tolerate all other therapies or unable to achieve cholesterol goal with maximally tolerated dose of most potent statins.

				MC	8	LIVALO			
				MC/DEL	8	MEVACOR TABS			DDI: Lescol will now be non-preferred and require prior authorization if it is currently being used in combination with diclofenac.
				MC	8	NEXLETOL			DDI: Lovastatin (doses greater than 40mg/day) will now be non-preferred and require prior authorization if it is currently being used in combination with Amiodarone.
				MC	8	NEXLIZET			
				MC/DEL	8	PRAVACHOL TABS			DDI: Lovastatin (doses greater than 20mg per day) will now be non-preferred and require prior authorization if it is currently being used in combination cyclosporine.
				MC/DEL	8	PRAVIGARD			
				MC	8	ZETIA TABS	Use PA Form# 20420		DDI: All preferred statins will now be non-preferred and require prior authorization if it is currently being used in combination with Gemfibrozil.
CHOLESTEROL - HMG COA + ABSORB INHIBITORS STATIN/ NIACIN COMBO	MC		SIMCOR	MC		ADVICOR TBCR	Use PA Form# 20420		
FAMILIAL HYPERCHOLESTEROLEMIA	MC MC		PRALUENT (LABLER 72733) PEN ^{1,2,3} REPATHA ^{1,2,3}	MC MC MC MC		EVKEEZA™ JUXTAPID KYNAMRO ¹ LEQVIO	1. Clinical PA required for appropriate diagnosis 2. Quantity limits apply 3. Documented adherence to lipid lowering medications and abstinence from tobacco for previous 90 days 4. For the treatment of patients ≥ 12 years of age. 5. Approval of Praluent NDC's with labeler code 00024 will be considered only if labeler code 72733 NDC's are on a long-term backorder and unavailable from the manufacturer.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists Juxtapid is contraindicated with strong CYP3A4 inhibitors. Juxtapid dosage should not exceed 30mg daily when it is used concomitantly with weak CYP3A4 inhibitors. Kynamro requires an appropriate lab testing prior to starting (ALT<AST), Alkaline phosphatase and total bilirubin, monthly liver-related tests for the first year, then every three months. Repatha and Praluent Criteria for approval: The patient's age is FDA approved for the given indication AND • Concurrent use with statin therapy AND • Documented adherence to prescribed lipid lowering medications for the previous 90 days AND • Recommended or prescribed by a lipidologist or cardiologist AND • Inability to reach goal LDL-C despite a trial of 2 or more maximum tolerated dose of statins (one of which must be atorvastatin or rosuvastatin) and ezetimibe 10mg daily	
							Use PA Form# 20420		Additional criteria for the diagnosis of heterozygous familial hypercholesterolemia (HeFH): (both are required): Total cholesterol > 290 mg/dL OR LDL-C > 190 mg/dL AND one of the following • Presence of tendon xanthomas OR • In 1st or 2nd degree relative-documented tendon xanthomas, MI at age ≤ 60 years or TC > 290 mg/dL . Additional criteria for the diagnosis of clinical atherosclerotic cardiovascular disease: History of MI, angina, coronary or other arterial revascularization, stroke, TIA, or PVD of atherosclerotic origin. Additional criteria for the diagnosis of homozygous familial hypercholesterolemia (Repatha only): Total cholesterol levels > 290mg/dL or LDL-C > 190mg/dL (adults) OR Total cholesterol levels > 260mg/dL or LDL-C > 155mg/dL (children < 16 years) and TG within reference range OR Confirmation of diagnosis by gene testing.
PULMONARY ANTI-HYPERTENSIVES									
PULMONARY ANTI-HYPERTENSIVES	MC MC/DEL MC/DEL MC		EPOPROSTENOL INJ ^{3,6} SILDENAFIL TADALAFIL VENTAVIS ³	MC/DEL MC MC/DEL MC MC MC MC MC/DEL MC MC		ADEMPAS™ ³ ADCIRCA ⁴ ALYQ TAB FLOLAN ³ LIQREV OPSUMIT ^{1,2} OPSYNVI ⁴ ORENITRAM REMODULIN ³ REVATIO ⁴ TADLIQ ⁴ TYVASO	1. Requires previous trials/failure of multiple preferred medications. 2. Dosing limits apply, please see the dose consolidation list. 3. Require WHO Group 1 diagnosis of primary PAH (Primary Pulmonary Hypertension) and NYHA functional class 3 or 4. 4. Require WHO Group 1	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Sildenafil will be preferred with clinical PA for treatment of pulmonary arterial hypotension (WHO Group 1) in adults to improve exercise ability and delay clinical worsening. Avoid concomitant use of Sildenafil with moderate or strong Cyp3A inhibitors DDI: Upravi will require a prior authorization if it is currently being used in combination with strong inhibitors of CYP2C8 (gemfibrozil) DDI: Opsumit will require a prior authorization if it is currently being used in combination with drugs known to be significant CYP3A inhibitors (ketoconazole, itraconazole, clarithromycin, indinavir, nefazodone, nelfinavir, ritonavir, atazanavir, saquinavir and telithromycin). DDI: Adempas will require a prior authorization if it is currently being used in combination with drugs known to be PDE inhibitors should be avoided (including dipyrindamole, adcirca and	

	MC/DEL MC	LORATADINE TAVIST ND (OTC)	MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	5 5 5 8 8 8 8 8 8 8 9	FEXOFENADINE ¹ ZYRTEC ¹ ZYRTEC SYR ^{1,2} ALLEGRA ³ CLARITIN ³ DESLORATADIN LORATADINE ODT ⁴ LEVOCETIRIZINE ⁴ XYZAL ³	<p>benzine before moving to non-preferred step order drugs.</p> <p>2. Clarinex and Zyrtec syrup <6 yr w/o PA.</p> <p>3. Must fail all step 5 drugs (Clarinex, Fexofenadine and Zyrtec) before moving to next step product.</p> <p>4. All OTC versions of loratadine ODT are now non-preferred.</p> <p>5. Pa's for Clarinex RediTabs will only be approved if between the ages of 6-11 years old.</p> <p>Use PA Form# 20530</p>	<p>and the preferred drug(s) exists. No combination product with decongestant will be approved since pseudoephedrine available without PA.</p> <p>Pseudoephedrine is available with prescription.</p>
ANTI-HISTAMINES - OTHER	MC/DEL MC/DEL MC/DEL	CLEMASTINE CHLORPHENIRAMINE DIPHENHYDRAMINE				<p>Use PA Form# 20530</p>	
ALLERGY / ASTHMA THERAPIES							
ANAPHYLACTIC DEVICES	MC/DEL MC/DEL MC/DEL	EPINEPHRINE EPIPEN EPIPEN JR	MC MC MC		AUVI-Q NEFFY TWINJECT		<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>Use PA Form# 20420</p>
ALLERGEN IMMUNOTHERAPY			MC MC MC MC MC		ODACTRA ORALAIR ¹ PALFORZIA RAGWITEK GRASTEK	<p>Use PA Form# 20420</p> <p>1. See criteria section</p>	<p>Prescriber must provide the testing to show that the patient is allergic to the components in the prescribed therapy and must provide a clinically valid rationale why single agent sublingual therapy is being chosen over subcutaneous therapy</p> <p>Palforzia® is approved for use in patients with a confirmed diagnosis of peanut allergy. Initial dose escalation may be administered to patients aged 4 through 17 years. Up-dosing and maintenance may be continued in patients 4 years of age and older.</p> <p>Odactra® is approved for use in persons 12 through 65 years of age. Note that Odactra® is not indicated for the immediate relief of allergic symptoms.</p> <p>Treatment must start 12 weeks before expected onset of pollen season and only after confirmed by positive skin test or in vitro testing for pollen-specific IgE antibodies for any of the 5 grass species contained in Oralair</p> <p>Oralair: Patient age ≥10 years and ≤65 years</p> <p>Have an auto-injectable epinephrine on-hand</p>
ANTI-ASTHMATIC - ANTICHOLINERGICS - INHALER	MC MC/DEL MC/DEL	INCRUSE ELLIPTA ³ SPIRIVA HANDIHALER ^{1,2} SPIRIVA RESPIMAT	MC MC/DEL		LONHALA MAGNAIR TUDORZA	<p>Use PA Form# 20420</p> <p>1. Quantity limit of 1 inhalation daily (1 capsule for inhalation daily) See criteria section</p> <p>2. We ask physicians to write "asthma" on the prescription whenever Spiriva is primarily being used for that condition.</p> <p>3. Quantity limit of 1 inhalation daily</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p>

ANTIASTHMATIC - PHOSPHODIESTERASE 4 INHIBITORS	MC/DEL		ROFLUMILAST	MC/DEL MC		DALIRESP OHTUVAYRE ¹	Use PA Form# 20420 1. For the maintenance treatment of chronic obstructive pulmonary disease (COPD) in adult patients	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
ANTIASTHMATIC - ANTICHOLINERGICS - NEBULIZER	MC/DEL		IPRATROPIUM BROMIDE SOLN	MC MC/DEL		ATROVENT SOLN YUPELRI	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
ANTIASTHMATIC - ANTIINFLAMMATORY AGENTS	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		CROMOLYN SODIUM NEBU DUPIXENT ^{2,4} FASENRA ² FASENRA ² AUTO INJCT XOLAIR ¹	MC MC MC		CINQAIR ³ NUCALA ² TEZSPIRE ⁵	1. Need max inhaled steroids and written by pulmonary or allergy specialist. Must have elevated IgE and ≥ to age 6. 2. For patients with severe asthma aged 12 years or older and eosinophilia. 3. For patients ≥ 18 years of age with eosinophilia. 4. Clinical PA required. 5. For adult and pediatric patients aged 12 years and older with severe asthma. Use PA Form# 20420	All will require suboptimal response to maximal doses of inhaled steroid as evidenced by asthmatic ER/Hospital admissions and Allergy/Pulmonary specialist management. Dupixent limited to patient with asthma not controlled on high dose ICS-LABA who have eosinophil greater than or equal to 150 cells or the patient is depend on an oral corticosteroid Fasenra, Nucala and Cinqair are not indicated for treatment of other eosinophilic conditions and are not indicated for the relief of acute bronchospasm or status asthmaticus.
ANTIASTHMATIC - NASAL STEROIDS	MC/DEL MC/DEL MC MC/DEL MC/DEL MC		BUDESONIDE SPRAY FLUTICASONE SPR ³ OLOPATADINE SPRAY OMNARIS SPR ³ TRIAMCINOLONE NS QNASL	MC MC/DEL MC/DEL MC/DEL MC MC/DEL MC MC MC/DEL MC MC/DEL	5 8 8 8 8 8 8 8 8 8 8	BECONASE AQ INHA ^{1,3} DYMISTA FLONASE SUSP ^{2,3} FLUNISOLIDE SOLN ^{1,3} NASONEX SUSP RHINOCORT AERO ^{2,3} RHINOCORT AQUA SUSP ^{2,3} RYALTRIS ⁴ TRI-NASAL SOLN ^{2,3} VANCENASE POKKETHALER AERS ^{2,3} VERAMYST ^{2,3} XHANCE ² ZETONNA ³	Use PA Form# 20420 1. All preferred drugs must be tried before moving to non preferred steps. 2. All step 5 medications need to be tried before moving to step 8's. 3. Dosing limits apply to whole category, please see dosage consolidation list. 4. Use of individual ingredients or other preferred agents.	Preferred drugs and step therapy must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Xhance will be considered for the treatment of nasal polyps in patients 18 years of age or older. The patient has had a documented side effect, allergy, or treatment failure of two preferred nasal glucocorticoids, one of which must be fluticasone.
ANTIASTHMATIC - NASAL MISC.	MC/DEL MC/DEL MC		AZELASTINE CROMOLYN NASAL 4% IPRATROPIUM NASAL SOL ¹	MC/DEL MC/DEL MC	8 8	ASTEPRO ² PATANASE	Use PA Form# 20420 1. Ipratropium will be approved if submitted with documentation supporting use of CPAP machine.	Approved if patient fails on non-sedating antihistamines and steroid nasal sprays. Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.

	MC/DEL MC/DEL MC/DEL		THEOPHYLLINE SOLN THEOPHYLLINE ER CP12 THEOPHYLLINE ER TB12					
ANTIASTHMATIC - STEROID INHALANTS	MC MC/DEL MC/DEL MC/DEL MC/DEL MC		ARNUITY ELLIPTA ASMANEX TWISTHALER ^{3,4} ASMANEX HFA ³ BUDESONIDE NEB 0.25MG & 0.5MG ¹ PULMICORT FLEXHALER ³ QVAR AERS ³	MC MC/DEL MC MC/DEL MC/DEL	8 8 8 8 8	AEROSPAN ALVESCO ³ ARMONAIR DIGIHALER BUDESONIDE NEB 1MG PULMICORT SUSP	1. Budesonide Neb 0.25mg & 0.5mg will be preferred for members under the age of 8 years old. PA will be required for members 8 years of age and older, please consider other preferred options. 2. All preferred must be tried before moving to non preferred steps. 3. Dosing limits apply, please see dosage consolidation list. 4. Asmanex 110mcg will be limited to member between the ages of 4-11years old. 5. Asmanex HFA will be preferred for members under the age of 6 years old. PA will be required for members 6 years of age and older, please consider other preferred options. Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
ANTIASTHMATIC - 5-Lipoxygenase Inhibitors				MC		ZYFLO CR TABS	Use PA Form# 20420	Other Preferred asthma controller drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
ANTIASTHMATIC - LEUKOTRIENE RECEPTOR ANTAGONISTS	MC/DEL MC/DEL MC/DEL		MONTELUKAST GRANULE ¹ MONTELUKAST SODIUM TAB MONTELUKAST SODIUM CHEW TAB	MC/DEL MC/DEL MC/DEL	8 8 8	ACCOLATE TABS SINGULAIR ² SINGULAIR GRANULES	Use PA Form# 20420 1.Montelukast Granules will only be approved if between ages of 6months-24 months. 2.Singulair Chewable 4mg from 2years-5years and Singulair Chewable 5mgs from 6years-14years old.	
ANTIASTHMATIC - ALPHA-PROTEINASE INHIBITOR				MC MC/DEL MC MC	8 8 8 8	ARALAST ZEMAIRA GLASSIA PROLASTIN SUSR	Use PA Form# 20420	Prolastin and Azemaira will be approved for members with A1AT deficiency and clinically demonstrable panacinar emphysema.
ANTIASTHMATIC - HYDRO-LYTIC ENZYMES				MC/DEL		PULMOZYME SOLN	Use PA Form# 20420	Will be approved for cystic fibrosis patients.

	MC/DEL MC/DEL MC/DEL MC/DEL		PAMINE TABS PROPANTHELINE BROMIDE TABS SODIUM BICARBONATE TABS TUMS	MC MC MC		OSCIMIN ROBINUL INJ ROBINUL TABS		Mytesi requires a diagnosis of non-infectious diarrhea in patients with HIV/AIDS on anti-retroviral therapy, prior trials of preferred, more cost effective anti-diarrheal.
GI- BILE ACID				MC		CHOLBAM	Use PA Form# 20420	Indication of bile acid synthesis disorders due to single enzyme defects (SEDs) AND for adjunctive treatment of peroxisomal disorders (PDs)
GI- EOSINOPHILIC ESOPHAGITIS	MC		EOHILIA ¹				Use PA Form# 20420 1. Approvals will not be longer than 12 weeks of treatment in adult and pediatric patients 11 years of age and older	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved (in step order), unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Eohilia: Dietary modification, PPIs, and topical glucocorticoids are required as initial therapy.
GI- H2-ANTAGONISTS	MC MC/DEL MC/DEL		ACID REDUCER TABS CIMETIDINE FAMOTIDINE	MC MC MC/DEL MC/DEL MC		AXID CAPS AXID AR TABS NIZATIDINE CAPS PEPCID PEPCID AC	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. DDI: Cimetidine will now be non-preferred and require prior authorization if it is currently being used with any sulfonylurea (except for glyburide). DDI: Cimetidine will require prior authorization if being used in combination with Plavix.
GI- IBAT INHIBITORS				MC MC		BYLVAY ^{1,2} LIVMARLI ^{1, 2}	Use PA Form# 20420 1. For the treatment of patients ≥ 3months of age 2. Clinical PA required for appropriate diagnosis	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Certain drugs require specific diagnoses for approval.
GI- PROTON PUMP INHIBITOR	MC/DEL MC/DEL MC/DEL		OMEPRAZOLE CAPS ² PANTOPRAZOLE ² LANSOPRAZOLE CAPS ²	MC/DEL MC/DEL MC MC MC/DEL MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	6 6 7 7 8 8 8 8 8 8 8 8 8	NEXIUM CPDR ³ NEXIUM SUS ⁵ PRILOSEC OTC ³ ACIPHEX TBEC ³ DEXILANT (KAPIDEX) ² KONVOME ² OMEPRAZOLE-SODIUM BICARBONATE CAPS OMEPRAZOLE MAGNESIUM PREVACID CPDR ³ PREVACID SOLUTABS ^{1,4} PRILOSEC CPDR PROTONIX INJ PROTONIX ² VOQUEZNA TABS	1. Prevacid Solutabs available without PA for children less than 9 years old. 2. Dosing limits apply, please see dosage consolidation list. 3. All preferred and step therapy must be tried and failed. 4. Payment for Prevacid SoluTabs for patients 9 and older will be considered for those patients who cannot tolerate a preferred solid oral dosage form. 5. Nexium sus available without PA if member is < 12 yrs of age and ≤ 1 pack per day. Use PA Form# 20720	All preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs (in step-order) will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Please refer to the PPI PA form for additional criteria on Non-Preferred PPIs DDI: Omeprazole will require prior authorization if being used in combination with Plavix. DDI: Lansoprazole will require prior authorization if being used in combination with Plavix. DDI: Prevacid, Omeprazole and pantoprazole will now be non-preferred and require prior authorization if they are currently being used in combination with any of the following medications: Ampicillin, B-12, Fe salts, Griseofulvin, Sporanox, Ketoconazole, Reyataz, or Vantin. DDI: All non-preferred PPIs require prior authorization, but with any prior authorization request, the member's drug profile will also be monitored for current use with ampicillin, B-12, Fe salts, griseofulvin, itraconazole, ketoconazole, Reyataz or Vantin due to a significant drug-drug interaction.
GI- ULCER ANTI-INFECTIVE	MC MC		PYLERA TALICIA			VOQUEZNA DUAL PAK VOQUEZNA TRIPLE PAK	Use PA Form# 20420	

GI - PROSTAGLANDINS	MC		MISOPROSTOL TABS	MC/DEL		CYTOTEC TABS		Preferred drug must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drug will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Use PA Form# 20420
GI - DIGESTIVE ENZYMES	MC/DEL MC		CREON ¹ ZENPEP ¹	MC/DEL MC/DEL MC/DEL		PERTZYE ULTRESA VIOKACE		Non -Preferred drugs must be tried and failed in step-order due to lack of efficacy or intolerable side effects before other non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Use PA Form# 20420 1. Clinical PA is required to establish CF diagnosis and medical necessity. In all cases except cystic fibrosis patients, objective evidence of pancreatic insufficiency (fat malabsorption test etc...) must be supplied.
GI - ANTI - FLATULENTS / GI STIMULANTS	MC/DEL MC MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL		AMITIZA CALULOSE SYRP CONSTULOSE SYRP ENULOSE SYRP GASTROCROM CONC GENERLAC SYRP LACTULOSE SYRP METOCLOPRAMIDE HCL	MC MC/DEL MC MC/DEL		CEPHULAC SYRP INFANTS GAS RELIEF SUSP GIMOTI SPRAY REGLAN TABS		Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Certain drugs require specific diagnoses for approval. Use PA Form# 20420
GI - INFLAMMATORY BOWEL AGENTS	MC MC/DEL MC MC MC/DEL MC/DEL		APRISO BALSALAZIDE MESALAMINE ENMA KIT PENTASA SULFAZINE EC TBEC SULFASALAZINE TABS	MC/DEL MC/DEL MC MC MC/DEL MC MC MC/DEL MC/DEL MC MC/DEL MC MC		ASACOL 800MG HD AZULFIDINE EN-TABS TBEC AZULFIDINE TABS COLAZAL CAPS DELZICOL DIPENTUM CAPS GIAZO LIALDA TABS ¹ MESALAMINE TAB ROWASA ENEM SFROWASA UCERIS RECTAL FOAM ² UCERIS TABS ²		Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Use PA Form# 20420 1. Current users grandfathered. 2. Diagnosis required Giazto is only indicated for males, as the safety,efficacy for use in females has not been established. Prior trials of preferred products. Uceris Rectal Foam or Tab- Concomitant use with CYP3A inhibitors (e.g. ketoconazole, itraconazole, ritonavir, indinavir, saquinavir, erythromycin, cyclosporine, and grapefruit juice) should be avoided. Verify prior trials and failures or intolerance of preferred treatments
GI - IRRITABLE BOWEL SYNDROME AGENTS	MC/DEL		LOTROXEN TABS	MC		VIBERZI		Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Use PA Form# 20420
GI- SHORT BOWL SYNDROME				MC		GATTEX		Gattex requires a diagnosis of adult SBS who are dependent on parenteral support. Appropriate colonoscopy and lab assessments 6months prior to starting Use PA Form #20420
GI- NASH				MC		REZDIFRA		Rezdiffra: The patient must have a diagnosis of NASH with fibrosis Stage 2 or 3 and utilizing imaging and scanning test such as fibro scan, MRI or ultra sound AND the patient does not have evidence of decompensated cirrhosis Use PA Form #20420
MISCELLANEOUS GI								
GI - MISC.	MC/DEL MC/DEL MC		BISAC-EVAC SUPP BISACODYL BISCOLAX SUPP	MC/DEL MC MC/DEL		ACTIGALL CAPS BENEFIBER CARAFATE		1. PA required to confirm FDA approved indication. 2. For the treatment of Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Certain drugs require specific diagnoses for approval.

MC	CINOBAC CAPS	MC/DEL	CLEARLAX POW
MC/DEL	CITRATE OF MAGNESIA SOLN	MC/DEL	COLACE CAPS
MC/DEL	CITRUCEL	MC	DIOCTO-C SYRP
MC/DEL	CLENPIQ SOL	MC	DOC SOD /CAS CAP
MC/DEL	COLYTE	MC	DOC-Q-LAX CAPS
MC/DEL	DIOCTO SYRP	MC/DEL	DOCUSATE SODIUM/CAS CAPS
MC	DOCUSATE CALCIUM CAPS	MC/DEL	DOK PLUS
MC/DEL	DOCUSATE SODIUM	MC/DEL	DULCOLAX SUPP
MC/DEL	FIBER LAXATIVE TABS	MC	ENEMEEZ
MC	FLEET	MC	FIBER CON TABS
MC/DEL	GENFIBER POWD	MC/DEL	FIBER-LAX TABS
MC/DEL	GLYCERIN	MC/DEL	GAVILYTE-H
MC	HIPREX TABS	MC	GOLYTELY SOLR
MC/DEL	KRISTALOSE PACK	MC	IBSRELA
MC/DEL	LINZESS	MC	IQIRVO
MC	MAALOX	MC/DEL	LINZESS 72mcg ⁴
MC/DEL	MILK OF MAGNESIA SUSP	MC	LIVDELZI
MC	MINERAL OIL	MC	MALTSUPEX
MC	MIRALAX BULK POWD (BRAND)	MC	MIRALAX PACKETS
MC/DEL	MOVANTIK	MC/DEL	MOTEGRITY
MC/DEL	MOVIPREP POWD PACK	MC	OCALIVA ¹
MC	NULYTELY SOLR	MC	PEG-ELECTROLYTES SOLR
MC	PEG 3350- ELECTROLYTE SOL	MC	PEG 3350 PACKETS
MC	PEG 3350 POWDER	MC	PREPOPIK PAK
MC/DEL	SENNA	MC	RELISTOR TABS
MC/DEL	SEKOT GRAN	MC/DEL	SEKONX TABS
MC/DEL	SEKOT SYRP	MC/DEL	SEKOT TABS
MC/DEL	SEKOT CHILDRENS SYRP	MC	SEKOT S TABS
MC	SEKOT XTRA TABS	MC/DEL	SORBITOL
MC/DEL	STOOL SOFTENER CAPS	MC	STOOL SOFTENER PLUS CAPS
MC/DEL	SUCRALFATE TABS	MC	SUFLAVE
MC/DEL	SUPREP SOL	MC	SUTAB
MC	TRULANCE ²	MC/DEL	SYMPROIC ³
MC	UNI-EASE CAPS	MC/DEL	UNI-CENNA TABS
MC	URSO FORTE	MC	UNI-EASE PLUS CAPS
MC/DEL	URSODIOL	MC	V-R NATURAL SENNA LAXATIV TABS
		MC	URSO 250
		MC	XERMELO ⁴

carcinoid syndrome diarrhea in combination with somatostatin analog (SSA) therapy in adults inadequately controlled by SSA therapy	
Linzess is preferred for adults as treatment of IBS-Constipation AND treatment of chronic idiopathic constipation in adults.	
3. For the treatment of Opioid Induced Constipation(OIC)	Trulance should be avoided in pediatric patients less than 18 years of age.
4. Established users will be grandfathered	
Iqirvo: For the treatment of primary biliary cholangitis (PBC) in combination with ursodeoxycholic acid (UDCA) in adults who have had an inadequate response to UDCA, or as monotherapy in patients unable to tolerate UDCA. Continued approval may be contingent upon verification and description of clinical benefit in confirmatory trial(s).	
Livdelzi: Clinical PA is required for the treatment of primary biliary cholangitis (PBC) in combination with ursodeoxycholic acid (UDCA) in adults who have had an inadequate response to UDCA, or as monotherapy in patients unable to tolerate UDCA. Patients who do not have a diagnosis of decompensated cirrhosis.	
	Use PA Form# 20420

MISC. UROLOGICAL

UROLOGICAL - MISC.	MC	ACETIC ACID 0.25% SOLN	MC	CITRIC ACID/SODIUM CITRAT SOLN
	MC	CYTRA-K SOLN	MC/DEL	CYTRA-2 SOLN
	MC	FOSFOMYCIN (NDC 82036427401 ONLY)	MC/DEL	ELMIRON CAPS ¹
	MC	K-PHOS MF TABS	MC	FURADANTIN SUSP
	MC/DEL	METHENAMINE MANDELATE TABS	MC/DEL	MACROBID CAPS
	MC/DEL	NEOSPORIN GU IRRIGANT SOLN	MC/DEL	MACRODANTIN CAPS
	MC/DEL	NITROFURANTOIN MONO CAPS	MC/DEL	NITROFURANTOIN MACR SUSP
	MC/DEL	PHENAZOPYRIDINE HCL TABS	MC	POTASSIUM CITRATE/CITRIC SOLN
	MC/DEL	PHENAZOPYRIDINE PLUS	MC/DEL	PYRIDIUM PLUS TABS
	MC	POT CITRATE TAB	MC	PYRIDIUM TABS
	MC/DEL	PROSED/DS TABS	MC/DEL	RENACIDIN SOLN
	MC	TRICITRATES SYRP	MC	UROCID-K
	MC/DEL	URELIEF PLUS		
	MC	UREX TABS		
	MC/DEL	URISED TABS		
	MC/DEL	UROQID #2 TABS		

1. Elmiron requires adequate proof of Dx with supportive testing.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
Use PA Form# 20420	

PHOSPHATE BINDERS						
PHOSPHATE BINDERS	MC/DEL MC/DEL MC/DEL MC MC/DEL		CALCIUM ACETATE CAP ¹ FOSRENOL CHEW ¹ MAGNEBIND - 400 ¹ PHOSLYRA ¹ REVELA ¹	MC MC/DEL MC/DEL MC MC		AURYXIA ¹ CALCIUM ACETATE TAB ¹ ELIPHOS ¹ FOSRENOL PWDR ¹ VELPHORO ¹ XPHOZAH
						Use PA Form# 20420 1. Diag required.
						Preferred drugs must be tried and failed in step-order due to lack of efficacy or intolerable side effects before less preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Xphozah to reduce serum phosphorus in adults with chronic kidney disease (CKD) on dialysis as add-on therapy in patients who have an inadequate response to phosphate binders or who are intolerant of any dose of phosphate binder therapy.
INTRA-VAGINALS						
VAGINAL - ANTIBACTERIALS	MC/DEL MC/DEL MC MC/DEL MC/DEL		CLEOCIN CREA CLEOCIN SUPP CLINDESSE CREA METRONIDAZOLE VAGINAL GEL ¹ NUVESSA	MC/DEL MC/DEL MC MC/DEL MC/DEL		METROGEL VAGINAL GEL ¹ VANDAZOLE XACIATO
						1. Dosing limits apply, please see Dosage Consolidation List. Use PA Form# 20420
						Preferred drugs must be tried and failed in step-order due to lack of efficacy or intolerable side effects before less preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
VAGINAL - ANTI FUNGALS	MC/DEL MC/DEL MC/DEL MC MC MC/DEL MC/DEL MC MC/DEL MC MC		CLOTRIMAZOLE CREA CLOTRIMAZOLE-3 CREA GYNE-LOTRIMIN CREA MICONAZOLE CREA MICONAZOLE 3 KIT CREA OTC MICONAZOLE 7 CREA MICONAZOLE NITRATE CREA NYSTATIN TABS TERCONAZOLE CREAM VAGITROL V-R MICONAZOLE-7 CREA	MC MC MC MC MC/DEL MC/DEL MC MC MC/DEL MC MC		AVC CREA CLOTRIMAZOLE 3 DAY CREA GYNAZOLE-1 CREA GYNE-LOTRIMIN 3 TABS MICONAZOLE 3 COMBO PACK KIT ¹ MICONAZOLE 3 SUPP TERAZOL 3 CREA TERAZOL 7 CREA TERCONAZOLE SUPP
						1. Quantity limit: 1/script/2 weeks Use PA Form# 20420
						Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. DDI: Miconazole will require prior authorization if being used in combination with Warfarin.
VAGINAL - CONTRACEPTIVES						
						Preferred drug must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drug will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Use PA Form# 20420
VAGINAL - ESTROGENS	MC/DEL MC/DEL		ESTRING RING PREMARIN CREA	MC/DEL MC/DEL		ESTRACE CREA ¹ VAGIFEM TABS ¹
						1. Must fail all preferred products before non-preferred. Use PA Form# 20420
						Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
VAGINAL - OTHER	MC/DEL MC MC		ACID JELLY GEL ACI-JEL GEL CERVICAL AMINO ACID CREA	MC		AMINO ACID CERVICAL CREA
						Use PA Form# 20420
						Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
BENIGN PROSTATIC HYPERPLASIA (BPH)						
BPH	MC/DEL MC/DEL MC/DEL MC/DEL		DOXAZOSIN MESYLATE TABS FINASTERIDE ¹ 5mg TERAZOSIN HCL CAPS TAMSULOSIN HCL	MC/DEL MC/DEL MC MC/DEL MC MC MC/DEL MC/DEL	5 8 8 8 8 8 8 8	FLOMAX CP24 ALFUZOSIN AVODART ^{2,4} CARDURA TABS ⁴ ENTADF ^{5,6} JALYN ^{3,4} PROSCAR TABS ⁴ RAPAFLO ⁴ UROXATRAL ⁴
						1. There will be dosing limits of 1 tab per day with out PA. 2. Prior use of preferred agent prior to any approvals. 3. Use of preferred (tamsulosin and finasteride) and (tamsulosin and non-preferred Avodart). 4. Non-preferred products must be used in specified order. 5. Use of individual
						Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs (in step-order) will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Approval of a non-preferred 5-alpha reductase inhibitor requires objective clinical evidence of a very enlarged prostate rather than just the presence of obstructive urinary outflow symptoms along with adequate trial of preferred Proscar.

							Ingredients preferred (Finasteride and tadalafil).	
							6. Entadfi® is not recommended for more than 26 weeks	
							Use PA Form# 20420	
ANXIOLYTICS								
ANXIOLYTICS - BENZODIAZEPINES	MC/DEL		ALPRAZOLAM TABS	MC/DEL	8	ALPRAZOLAM ER	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
	MC/DEL		CHLORDIAZEPOXIDE HCL CAPS	MC/DEL	8	ATIVAN		
	MC/DEL		CLORAZEPATE DIPOTASSIUM TABS	MC	8	LOREEV XR		
	MC/DEL		DIAZEPAM	MC/DEL	8	NIRAVAM		
	MC/DEL		LORAZEPAM	MC/DEL	8	SERAX		
	MC/DEL		OXAZEPAM CAPS	MC/DEL	8	TRANXENE		
				MC/DEL	8	XANAX TABS		
				MC/DEL	9	XANAX XR		
ANXIOLYTICS - MISC.	MC/DEL		BUSPIRONE HCL TABS	MC		BUSPAR TABS	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
	MC		HYDROXYZINE HCL SOLN	MC		DROPERIDOL SOLN	1. Dosing limits apply, please refer to Dose consolidation list.	
	MC		HYDROXYZINE HCL SYRP	MC/DEL		DROPERIDOL SOLN		
	MC/DEL		HYDROXYZINE HCL TABS ¹	MC/DEL		DROPERIDOL SOLN		
	MC/DEL		HYDROXYZINE PAMOATE CAPS					
	MC/DEL		MEPROBAMATE TABS					
ANTI-DEPRESSANTS								
ANTIDEPRESSANTS - MAO INHIBITORS	MC/DEL		NARDIL TABS	MC/DEL		TRANLYCYPROMIINE	Use PA Form# 20420	
ANTIDEPRESSANTS - MAO INHIBITORS TOPICAL				MC/DEL		EMSAM ¹	1. Dosing limits apply, please refer to Dose consolidation list.	Preferred drugs (including a preferred SSRI, a non-SSRI, and Venlafaxine ER) must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
							Use PA Form# 20420	
ANTIDEPRESSANTS - SELECTED SSRI's AND OTHERS	MC/DEL		BUPROPION HCL TABS	MC/DEL	8	APLENZIN ⁴	1. Strong caution with pediatric population. 2. Max daily dose allowed is 120mg. Combination of multiple strengths require PA 4. Dosing limits allowing 2 tabs/day and a max daily limit of 200mg / day applies. Please see dose consolidation list. 5. Dosing limits apply, please refer to Dose consolidation list and max daily dose applies. Max daily dose allowed is 375mg. 6. Non-preferred products must be used in specified step order. 7. Requires previous trials/failure of multiple preferred medications. Dosing limits apply, please see the dose consolidation list. Max daily dose of 80mg if used concomitantly with strong CYP3A4 inhibitor.	Preferred drugs (including failure of at least one preferred SSRI, one SNRI and one non-SSRI/SNRI) must be tried for at least 4 weeks each and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
	MC/DEL		BUPROPION SR	MC	8	AUVELITY ¹¹		
	MC/DEL		BUPROPION XL 150mg and 300mg	MC/DEL	8	BUPROPION XL 450mg		
	MC/DEL		CITALOPRAM	MC/DEL	8	CELEXA		
	MC/DEL		DULOXETINE ^{2,9}	MC	8	CYMBALTA ²		
	MC/DEL		ESCITALOPRAM	MC/DEL	8	DRIZALMA SPRINKLES		
	MC/DEL		FLUOXETINE 10mg AND 20mg AND 40mg CAPS	MC/DEL	8	EFFEXOR TABS		
	MC/DEL		FLUOXETINE HCL LIQD	MC/DEL	8	EFFEXOR XR CP24		
	MC/DEL		FLUVOXAMINE MALEATE TABS	MC/DEL	8	FETZIMA ⁷		
	MC/DEL		MIRTAZAPINE	MC/DEL	8	FLUOXETINE 10mg AND 20mg AND 60mg TABS		
	MC/DEL		NEFAZODONE	MC	8	FORFIVO XL		
	MC/DEL		PAROXETINE ¹	MC/DEL	8	IRENKA		
	MC/DEL		SERTRALINE HCL	MC/DEL	8	KHEDEZLA		
	MC/DEL		TRAZODONE HCL TABS	MC/DEL	8	LEXAPRO TABS		
	MC/DEL		VENLAFAXINE ER CAPS ⁵	MC	8	LUVOX TABS		
	MC/DEL		VENLAFAXINE TABS ⁵	MC	8	MAPROTILINE HCL TABS		
				MC/DEL	8	MIRTAZAPINE ODT		
				MC	8	OLEPTRO		
				MC/DEL	8	PAROXETINE CR ¹		
				MC/DEL	8	PAXIL ¹		
				MC/DEL	8	PAXIL CR ¹		
				MC/DEL	8	PRISTIQ		
				MC	8	PROZAC		
				MC	8	PROZAC CAPS		
							Zulresso® is available only through a restricted program under a Risk Evaluation and Mitigation Strategy (REMS) called the Zulresso® REMS.	

ANTI-PSYCHOTICS

ANTI-PSYCHOTICS - ATYPICALS						
MC		ABILIFY ASIMTUFII	MC/DEL	8	ABILIFY DISC TAB, INJ and SOL ¹	<p>If prescribing 2 or more antipsychotics, PA will be required for both drugs, except if one is Clozapine. This also includes combination of Seroquel with Seroquel XR.</p> <p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Non preferred atypicals will be approved for patients with FDA-approved indications, and for specific conditions supported by at least two published peer-reviewed double-blinded, placebo-controlled randomized trials that are not contradicted by other studies of similar quality and as long as all first line preferred therapies have been tried and failed at full therapeutic doses for adequate durations (at least two weeks).</p> <p>Prescriptions for quetiapine are limited to a maximum daily dose of 800mg.</p> <p>Uzedy: Establish tolerability with oral risperidone prior to initiating Uzedy</p> <p>Atypicals: Prior Authorization will be required for preferred medication to assure indication is in accordance with FDA approved or literature supported evidence-based best practices. The approved indications are:</p> <ul style="list-style-type: none"> • schizophrenia • bipolar disorder • agitation related to autism • adjunct in major depressive disorder <p>If prescribing 2 or more antipsychotics, PA will be required for both drugs, except if one is Clozapine. This also includes combination of Seroquel with Seroquel XR.</p> <p>DDI: It is recommended to reduce the Vraylar® dose if it is used concomitantly with a strong CYP3A inhibitor (such as itraconazole, ketoconazole). The concomitant use of Vraylar® with a CYP3A4 inducer (such as rifampin, carbamazepine) is not recommended.</p> <p>DDI: The concomitant use of Nuplazid with other drugs known to prolong the QT interval (e.g. Class IA antiarrhythmics, Class 3 antiarrhythmics, antipsychotics, and antibiotics such as gatifloxacin and moxifloxacin).</p> <p>Lybalvi: Step through aripiprazole and Latuda. If criteria is met then initial approval for 3 months. Subsequent approvals will be based on evidence of not gaining >= 10 % baseline body weight for ongoing approval. If weight gain >= 10 % of initial body weight, then criteria for ongoing use not met.</p> <p>Cobenfy: Patient must be 18–65 years old AND meet criteria for the diagnosis of schizophrenia, AND Trial of 2 prior preferred second generation antipsychotics showing minimal response in control of symptoms of schizophrenia OR Trial of SGA that have yielded side effects of weight gain which has not been responsive to lifestyle & medication augmentation AND Patient must have baseline tests including heart rate, liver enzymes, kidney function tests, and bilirubin prior to starting treatment.</p> <p>Invega Hafyera: The patient is started and stabilized on the medication OR The patient has been adequately treated with Invega Sustenna (paliperidone palmitate 1-month) for at least four months or Invega Trinza (paliperidone palmitate 3- month) following at least one 3-month injection cycle.</p>
MC		ABILIFY MAINTENA	MC	8	ABILIFY TABS ²	
MC/DEL		ARIPIPRAZOLE TAB ³	MC/DEL	8	ARIPIPRAZOLE SOL	
MC		ARISTADA	MC/DEL	8	ARIPIPRAZOLE ODT	
MC		ARISTADA INITIO	MC	8	CAPLYTA	
MC/DEL		OLANZAPINE ^{2,3}	MC	8	COBENFY	
MC/DEL		OLANZAPINE ^{2,3} ODT	MC	8	ERZOFRI	
MC/DEL		INVEGA HAFYERA	MC	8	FANAPT	
MC		INVEGA SUSTENNA	MC/DEL	8	GEODON	
MC/DEL		INVEGA TRINZA INJ	MC	8	INVEGA	
MC/DEL		LURASIDONE TAB	MC	8	IGALMI	
MC/DEL		PALIPERIDONE ER	MC	8	LATUDA	
MC/DEL		PERSERIS	MC	8	LYBALVI	
MC		RISPERDAL CONSTA	MC	8	NUPLAZID	
MC/DEL		RISPERIDONE ODT	MC	8	OPIPZA	
MC/DEL		RISPERIDONE TAB ^{2,3}	MC	8	REXULTI	
MC/DEL		RISPERIDONE SOLN ²	MC	8	RISPERDAL TAB	
MC		RYKINDO	MC	8	RISPERDAL M TAB ¹	
MC/DEL		QUETIAPINE ^{2,3}	MC	8	RISPERDAL SOLN	
MC/DEL		QUETIAPINE XR	MC/DEL	8	SAPHRIS ¹	
MC		VRAYLAR ⁴	MC	8	SECUADO	
MC/DEL		ZIPRASIDONE ^{2,3}	MC/DEL	8	SEROQUEL TABS	
			MC	8	UZEDY	
			MC	8	ZYPREXA TABS	
			MC		ZYPREXA RELPREVV	
			MC	8	ZYPREXA ZYDIS TBDP ¹	
			MC/DEL	9	SEROQUEL XR	
					1. Established users of single therapy atypicals were grandfathered.	
					2. Prior Authorization will be required for preferred medications for members under the age of 5.	
					3. Dosing limits apply please refer to the dose consolidation list.	
					4.Requires step through 1 preferred drug for all indications except AMDD. AMDD requires insufficient response from two antidepressants	
ANTI-PSYCHOTICS - SPECIAL ATYPICALS						
MC/DEL		CLOZAPINE TABS	MC/DEL		CLOZAPINE ODT	<p>Preferred generic drug must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred brand will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Patients previously stabilized on brand name drug will be approved.</p>
			MC/DEL		CLOZARIL TABS	
			MC/DEL		VERSACLOZ SUSP	

ANTIPSYCHOTICS - TYPICAL	MC/DEL MC/DEL MC/DEL MC MC/DEL MC MC MC/DEL MC/DEL MC MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL	CHLORPROMAZINE HCL FLUPHENAZINE DECANOATE FLUPHENAZINE HCL HALDOL HALOPERIDOL HALOPERIDOL DECANOATE SOLN HALOPERIDOL LACTATE SOLN LOXAPINE SUCCINATE CAPS LOXITANE-C CONC MOBAN TABS PERPHENAZINE PROCHLORPERAZINE SERENTIL THIORIDAZINE HCL THIOTHIXENE TRIFLUOPERAZINE HCL TABS	MC/DEL MC/DEL MC MC/DEL MC MC/DEL MC MC	COMPAZINE COMPRO SUPP FLUPHENAZINE HCL CONC HALDOL DECANOATE LOXITANE CAPS MELLARIL NAVANE CAPS PROLIXIN STELAZINE TABS	Use PA Form# 20420 If prescribing 2 or more antipsychotics, PA will be required for both drugs, except if one is Clozapine.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. If prescribing 2 or more antipsychotics, PA will be required for both drugs, except if one is Clozapine.
LITHIUM						
LITHIUM	MC/DEL MC/DEL	LITHIUM CARBONATE LITHIUM CITRATE SYRP	MC/DEL MC/DEL	ESKALITH CAPS ESKALITH CR TBCR	Use PA Form# 20420	
COMBINATION - PSYCHOTHERAPEUTIC						
PSYCHOTHERAPEUTIC COMBINATION			MC/DEL MC/DEL	CHLORDIAZEPOXIDE/AMITRIPT PERPHENAZINE/AMITRIPTYLIN	Use PA Form# 20420	
STIMULANTS						
STIMULANT - AMPHETAMINES -SHORT ACTING	MC/DEL MC/DEL MC	AMPHETAMINE SALT COMBO ^{1,4} DEXTROAMPHET SULF TABS PROCENTRA	MC/DEL MC MC/DEL MC	ADDERALL TABS EVEKEO METHAMPHETAMINE HCL ZENZEDI	Use PA Form# 20420 1. Preferred stimulants will be available without PA if diagnosis of ADHD or Narcolepsy. 2. As per recent FDA alert, Adderall & Dexedrine should not be used in patients with underlying heart defects since they may be at increased risk for sudden death. 3. Dosing limits apply, please see dosing consolidation list. 4. Max daily dose of 50mg.	
STIMULANT - LONG ACTING AMPHETAMINES SALT	MC/DEL MC MC	AMPHETAMINE/DEXTROAMPHET ER ^{3,4,7} ADDERALL XR CP24 ^{1,3,4,7} VYVANSE ^{2,3,4}	MC MC MC	MYDAYIS ⁵ VYVANSE CHEW ^{4,6} XELSTRYM ⁸	Use PA Form# 20420 1. As per recent FDA alert, Adderall should not be used in patients with underlying heart defects since they may be at increased risk for sudden death.	DDI: The concomitant use of Mydayis® is contraindicated with monoamine oxidase inhibitors (MAOIs) or within 14 days after discontinuing MAOI treatment, as concomitant use can increase hypertensive crisis.

				MC/DEL		RITALIN	please see dosing consolidation list. Maximum daily doses are as follows: 72mg daily for methylphenidate and 36mg daily for dexamethylphenidate.	
STIMULANT - METHYLPHENIDATE - LONG ACTING	MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC MC MC/DEL		CONCERTA TBCR DEXMETHYLPHENIDATE CAP ER 50/50 FOCALIN XR METHYLPHENIDATE LA CAPS METHYLPHENIDATE ER CAPS 50/50 METHYLPHENIDATE ER CAPS 40/60 METHYLPHENIDATE CD CAPS 30-70 QUILLICHEW ER ^{5,1} QUILLIVANT XR SUS ^{1,5} RITALIN LA ⁴	MC MC/DEL MC MC MC MC/DEL MC/DEL MC/DEL	5 8 8 8 8 8 8 8 8	METADATE CD CPR ADHANSIA XR ^{2,6} APTENSIO XR ² AZSTARYS ⁶ COTEMPLA XR ² COTEMPLA XR ODT ² DAYTRANA ^{2,3} JORNAY PM ^{2,6} METHYLPHENIDATE ER CAPS ^{2,4}	1. Preferred stimulants will be available without PA if diagnosis of ADHD. 2. Non-preferred products must be used in specified step order. 3. FDA approval currently only for ages 6-16. Limit of one patch daily. Max dose of 30MG daily. 4. Dosing limits apply, please see dosing consolidation list. 5. Quillivant XR and Quillichew ER are only indicated for use in patients 6 years of age and older. 6. For the treatment of patients ≥ 6 years of age. Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
STIMULANT - STIMULANT LIKE	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC		ATOMOXETINE HCL ARMODAFINIL CLONIDINE ER GUANFACINE ER MODAFINIL TABS QELBREE ^{6,7}	MC/DEL MC MC MC/DEL MC MC MC/DEL MC MC	7 7 8 8 8 8 8 8 8	PROVIGIL TABS ³ STRATTERA ^{1,2} CAFCIT SOLN ³ INTUNIV KAPVAY ONYDA XR ⁶ SUNOSI WAKIX XYREM SOL XYWAV ⁵ NUVIGIL ³ DESOXYN TABS ³ DESOXYN CR ³	1. Failure of both an amphetamine and methylphenidate is required for consideration for approval of Strattera, unless history of substance abuse without current use of abusable medication(s). Additionally, for patients <17 years of age, a trial of guanfacine is required before approval of Strattera. 2. Strattera currently has dosing limitations allowing one tablet per day for all strengths if obtain approval. Max daily dose of Strattera is 100mg. Please see dosing consolidation list. 3. Non-preferred products must be used in specified 4. Please use generic Guanfacine. 5. For patients 7 years of age and older with narcolepsy 6. For pediatric patients 6 years of age or older	Provigil requests require diagnosis of Narcolepsy, ADHD, or Obstructive Sleep Apnea. Previous failures of methylphenidate and amphetamine is required for Narcolepsy and ADHD diagnosis, with additional Strattera trial needed with ADHD diagnosis. Please refer to detailed criteria on Provigil PA form Sunsosi is non-preferred and is indicated for to improve wakefulness in adult patients with excessive daytime sleepiness associated with narcolepsy or obstructive sleep apnea (OSA). Wakix is non-preferred and is indicated for the treatment of excessive daytime sleepiness (EDS) in adults with narcolepsy DDI: Sunosi® is contraindicated with MAO inhibitors or within 14 days after discontinuing the MAO inhibitor. Xywav: Diagnosis of cataplexy associated with narcolepsy OR excessive daytime sleepiness associated with narcolepsy. Diagnosis must be confirmed by submission of supporting documentation to include the specialist's interpretation of the Polysomnography (PSG) and Multiple Sleep Latency Test (MSLT) results FDA reminded healthcare professionals and patients that the combined use of Xyrem (sodium oxalate) with alcohol or central nervous system (CNS) depressant drugs can markedly impair consciousness and may lead to severe breathing problems (respiratory depression) DDI: Concomitant use of Qelbree® with an MAO inhibitor or within 2 weeks after discontinuing an MAO inhibitor is contraindicated DDI: Concomitant use of Qelbree® significantly increases the total exposure, but not peak exposure, of sensitive CYP1A2 substrates, which may increase the risk of adverse reactions

							7. Preferred with a trial and fail either Atomoxetine OR any 2 preferred ADHD agents.	<p>Other combination use of selective serotonin reuptake inhibitors, but not paroxetine, or serotonin 2-1A reuptake inhibitors, which may increase the risk of serotonin toxicity associated with these CYP1A2 substrates. Coadministration of Qelbree® with sensitive CYP1A2 substrates or CYP1A2 substrates with a narrow therapeutic range (e.g. alosetron, duloxetine, ramelteon, tasimelteon, tizanidine, theophylline), is contraindicated.</p> <p>Use PA Form# 20710 for Proviqil, Nuviqil and Xyrem</p> <p>Use PA Form# 20420 for all others</p>
ANTI-CATALECTIC AGENTS								
PSYCHOTHERAPEUTIC AGENTS - MISC.				MC MC			<p>NUDEXTA XENAZINE</p> <p>Use PA Form# 20710 for Xenazine</p>	
WEIGHT LOSS								
WEIGHT LOSS							No longer covered: PHENTERMINE, XENICAL, DIDREX, and MERIDIA	Weight loss drugs are not covered as permitted by Federal Medicaid regulations and Maine Medicaid (MaineCare) Policy.
ALZHEIMER DISEASE								
ALZHEIMER - Cholinimetics/Others	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		<p>DONEPEZIL HYDROCHLORIDE TABS¹</p> <p>DONEPEZIL HYDROCHLORIDE ODT¹</p> <p>EXELON DIS¹</p> <p>GALANTAMINE CAPS¹</p> <p>GALANTAMINE TAB¹</p> <p>MEMANTINE¹</p> <p>RIVASTIGMINE TARTRATE CAPS¹</p>	MC MC MC/DEL MC MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL MC MC	6 6 7 8 8 8 8 8 8 8 8 8 8 8 8 9	<p>ARICEPT TABS²</p> <p>ARICEPT ODT²</p> <p>DONEPEZIL HYDROCHLORIDE TABS 23MG</p> <p>ADLARITY³</p> <p>EXELON CAP</p> <p>GALANTAMINE HYDROBROMIDE SOL</p> <p>KISUNLA</p> <p>LEQEMBI^{1,2}</p> <p>MEMANTINE HCL SOL</p> <p>NAMENDA</p> <p>NAMENDA XR CAPS</p> <p>NAMZARIC</p> <p>RAZADYNE²</p> <p>COGNEX CAPS²</p>	<p>1. PA is required to establish dementia diagnosis and baseline mental status score.</p> <p>2. Must fail all preferred products before moving to non-preferred.</p> <p>3. Approvals will require trials and failure or clinical rationale why preferred patches cant be used.</p> <p>Use PA Form# 20420</p>	<p>Preferred drug must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs (in step-order) will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>Kisunla and Leqembi: Testing to rule out reversible causes of dementia (CBC, CMP, TSH, B12, urine drug screen, RPR/VDRL, (folate (if alcohol abuse is present), HIV (if risk present) and an assessment including a review of current medications as a cause of intellectual decline</p> <p>- Prescribed by or in consultation with a neurologist or geriatrician or geriatric psychiatrist. Diagnosis of Alzheimer's disease defined as:</p> <ul style="list-style-type: none"> •Confirmed presence of amyloid pathology and mild cognitive impairment or mild dementia stage of disease, consistent with Stage 3 and Stage 4 Alzheimer's disease OR •Confirmed presence of amyloid pathology and prodromal or mild dementia stage of disease, consistent with Stage 3 and Stage 4 Alzheimer's disease <p>-Testing:</p> <ul style="list-style-type: none"> •Clinical Dementia Rating (CDR) global score of 0.5 or 1.0 OR •Repeatable Battery for Assessment of Neuropsychological Status (RBANS) delayed memory index score ≤ 85 OR •Mini-Mental State Examination (MMSE) score of 20-30 OR •Montreal Cognitive Assessment (MoCA) score ≤ 22 <p>- Member is age 50 or older</p> <p>- Obtain recent (within one year) brain magnetic resonance imaging (MRI) prior to initiating treatment</p> <p>- Provider attestation to obtain MRIs prior to the 7th infusion (first dose of 10 mg/kg) and 12th infusion (sixth dose of 10 mg/kg)</p> <p>- Member does NOT have history or increased risk of amyloid related imaging abnormalities-edema (ARIA-E), which includes brain edema or sulcal effusions and amyloid related imaging abnormalities hemosiderin deposition (ARIA-H), which includes microhemorrhage and superficial siderosis</p> <p>- Member does NOT have hypersensitivity to any components of these drugs</p> <p>- Failure of or inability to tolerate at least two other preferred Alzheimer therapies for at least four months each, one of which should include a combination of a cholinesterase inhibitor with memantine</p> <ul style="list-style-type: none"> •If the initial drug utilized is the combination of a cholinesterase inhibitor and memantine, then only that single trial of two drugs is required
SMOKING CESSATION								
NICOTINE PATCHES / TABLETS	MC/DEL MC/DEL MC/DEL MC/DEL		<p>CHANTIX TAB¹</p> <p>CHANTIX STARTER PACK</p> <p>NICOTINE DIS PT24¹</p> <p>VARENICLINE TAB</p>	MC/DEL			<p>NICODERM CQ PT24¹</p> <p>Use PA Form# 20420</p> <p>1. See criteria section for exemptions</p>	<p>As of July 1, 2014 per MaineCare policy, smoking cessation products will be covered without a copay(including MEDEL). No annual or lifetime limits, must follow FDA approved indications and therapy guidelines.</p> <p>Preferred drug must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs (in step-order) will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>Note: MaineCare policy, smoking cessation product were "not covered" except for during pregnancy between 9/1/12 and 1/1/14, between 1/1/2014 and 7/1/14 smoking cessation products were covered with limitations.</p>

								Patients may qualify for the medication through The Maine Tobacco Helpline if they do not have MaineCare or MEDEL. Patients are encouraged to call The Maine Tobacco helpline at 1-800-207-1230.
NICOTINE REPLACEMENT - OTHER	MC/DEL MC/DEL MC/DEL		NICOTINE POLACRILEX GUM ¹ NICOTINE LOZENGE MINI NICOTINE LOZENGE	MC/DEL MC/DEL MC/DEL MC	8 8 8 8	NICOTROL INHALER ^{1,2} NICOTROL NASAL SPRAY ^{1,2} NICORETTE GUM ^{1,2} NICORETTE LOZENGES	Use PA Form# 20420 1. See criteria section for exemptions 2. Must use non-preferred products in specified step order.	As of July 1, 2014 per MaineCare policy, smoking cessation products will be covered without a copay(including MEDEL). No annual or lifetime limits, must follow FDA approved indications and therapy guidelines. Preferred drug must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs (in step-order) will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Note: MaineCare policy, smoking cessation product were "not covered" except for during pregnancy between 9/1/12 and 1/1/14, between 1/1/2014 and 7/1/14 smoking cessation products were covered with limitations Patients may qualify for the medication through The Maine Tobacco Helpline if they do not have MaineCare or MEDEL. Patients are encouraged to call The Maine Tobacco helpline at 1-800-207-1230.
ALCOHOL DETERRENTS								
ALCOHOL DETERRENTS	MC/DEL MC MC MC/DEL		ACAMPROSATE ANTABUSE TABS DISULFIRAM TABS NALTREXONE HCL TABS	MC/DEL		ACAMPRO ¹	1. Should only be used in conjunction with formal structured outpatient detoxification program. Use PA Form# 20420	Preferred generic drug must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
MISCELLANEOUS ANALGESICS								
ANALGESICS - MISC.	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL		ACETAMINOPHEN ASPIRIN ASPRIN/ APAP/ CAFF TAB BUTAL/ASA/CAFF BUTALBITAL COMPOUND BUTALBITAL/ACET TABS BUTALBITAL/APAP CAPS BUTALBITAL/APAP/CAFFEINE TABS CHOLINE MAGNESIUM TRISALI DIFLUNISAL TABS EXCEDRIN SALSALATE TABS	MC MC/DEL MC/DEL MC MC MC/DEL MC MC MC MC MC MC		AXOCET CAPS ESGIC-PLUS FIORICET TABS FIORINAL CAPS FIORTAL CAPS FORTABS TABS PHRENILIN TABS PHRENILIN FORTE CAPS TRILISATE LIQD TRILISATE TABS ZEBUTAL CAPS ZORPRIN TBCR	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
LONG ACTING NARCOTICS								
NARCOTICS - LONG ACTING	MC/DEL MC/DEL MC/DEL		FENTANYL PATCH ⁴ BUTRANS ⁴ MORPHINE SULFATE ER TB12	MC MC MC MC MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL MC MC MC/DEL MC MC/DEL MC MC	8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	ARYMO ER AVINZA BELBUCA EXALGO HYSINGLA ER KADIAN METHADONE METHADOSE MORPHABOND ER MORPHINE SULFATE ER CAP MORPHINE SULFATE SUPP MS CONTIN TB12 OPANA ER ORAMORPH SR TB12 OXYCONTIN TB12 ¹ XARTEMIS ER ZOHYDRO ER	Use PA Form# 20510 Use PA form #10300 for PAS over the opiate limit 1. Oxycontin will be available without PA for patients treated for or dying from cancer or hospice patients. CA (cancer) or HO (hospice) diag code may be used but store must verify since all scripts will be audited and stores will be liable. 2. Established users are grandfathered. 3. Oxycodone ER allowed only 2 per day for all strengths except 80 mg, where 4 are allowed to	Preferred drugs (Fentanyl Patch, Morphine Sulfate ER tab, Butrans and Embeda) must be tried for at least 2 weeks each & failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug & the preferred drug(s) exists. Adequate trials include prevention/treatment of common adverse effects associated w/ narcotics (antinausea, antipruritic, etc.) as well as adequate equianalgesic dosing when converting from one narcotic to another. Also, adequate documentation of attempts to titrate dose of preferred agents to achieve adequate pain relief & desired clinical response must be provided. Member's drug regimen for additions &/or discontinuations of medications that may affect absorption &/or metabolism of preferred agents must be monitored. Approvals will not be granted if patient had access to either non-preferred products or high doses of short acting narcotics during the trial period. Non-preferred drugs will not be approved for patients showing evidence of usage patterns consistent w/ controlled substance abuse such as: 1.Frequent or persistent early refills of controlled drugs; 2.Multiple instances of early refill overrides due to reports of misplacement, stolen, dropped in toilet or sink, distant travel, etc.; 3.Breaches of narcotic contracts with any provider; 4.Failure to comply with patient responsibilities in attached opioid documentation (see PA form) including but not limited to failing to submit to and pass pill counts; 5.Failing to take or pass random drug testing; 6.Failing to provide old records regarding prior use of narcotics; 7.Receiving controlled substances from other prescribers that the provider submitting the PA is unaware of

				MC MC/DEL	8 9	OXYCODONECONC OXYCODONE ER ^{3,5}	4. Dosing limits apply. Please see dose consolidation list. 5. Non-preferred products must be used in specific order. 6. Methadone will be available without PA for patients treated for or dying from cancer or hospice patients or similar conditions as supported by clinical documentation. CA (cancer) or HO (hospice) diag code may be used but store must verify since all scripts will be audited and stores will be liable.	8.Documented history of substance abuse. Substance abuse evaluations may be required for patients with medical records displaying documented substance abuse or potential signs of narcotic misuse and abuse such as chronic early refills, short dosing intervals, frequent dose increases, multiple lost/stolen etc scripts and intolerance or "allergy" to all products but Oxycontin. 9.Circumventing MaineCare prior authorization requirements for narcotics by paying cash for affected narcotics (prescribers failed to submit prior authorization prior to cash narcotic scripts being filled by member). 10.Requests for any Brand name controlled substance, considered by authorities to be highly abused and diverted (Oxycontin, Percocet, Tylox, Vicodin, Dilaudid, Ultracet...) with an available AB rated generic equivalent will be denied unless it will be provided in a setting that virtually eliminates the risk of diversion. 11.Allergic reactions to any product within a specific narcotic class will justify and preclude use of any other product in the same class due to the risk of cross-hypersensitivity. Hysingla ER- Concomitant use should be avoided with mixed agonist/antagonist analgesics, partial agonist analgesics, and MAOIs. Verify prior trials and failures or intolerance of preferred treatments Methadone – Established users must have a trial and failure of at least 2preferred drugs for least 2 weeks. Otherwise they will be allowed 180 days to transition to a preferred product.
NARCOTICS - SELECTED	MC/DEL MC/DEL	TRAMADOL HCL TABS TRAMADOL/APAP TABS	MC/DEL MC MC/DEL MC MC MC MC MC MC	7 8 8 8 8 8 8 8 8 9		RYZOLT BUPRENEX SOLN BUTORPHANOL NALBUPHINE HCL SOLN QDOLO SOLN SEGLENTIS ¹ STADOL NS SOLN TRAMADOL ER ULTRACET TABS ¹ ULTRAM ER	Use PA Form# 20420 Use PA form #10300 for PA's over the opiate limit 1. Only available if component ingredients are unavailable.	Preferred drugs from this and other narcotic classes must be tried for at least 2 weeks each and failed due to lack of efficacy or intolerable side effects before non-preferred drugs from this class will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Approvals will not be granted if patient had access to either non-preferred products or high doses of short acting narcotics during the trial period. Substance abuse evaluations may be required for patients with medical records displaying potential signs of narcotic misuse and abuse such as chronic early refills, short dosing intervals, frequent dose increases, multiple lost/stolen etc scripts and intolerance or "allergy" to all products but desired product. Allergic reactions to any product within a specific narcotic class will justify and preclude use of any other product in the same class due to the risk of cross-hypersensitivity. Non-preferred drugs will not be approved for patients showing evidence of usage patterns consistent with controlled substance abuse such as: 1.frequent or persistent early refills of controlled drugs; 2.multiple instances of early refill overrides due to reports of misplacement, stolen, dropped in toilet or sink, distant travel; 3.breaches of narcotic contracts with any provider; 4.failure to comply with patient responsibilities in attached opioid documentation (see PA form) including but not limited to failing to submit to and pass pill counts; 5.failing to take or pass random drug testing; 6.failing to provide old records regarding prior use of narcotics; 7.receiving controlled substances from other prescribers that the provider submitting the PA is unaware of. In Substance abuse evaluations may be required for patients with medical records displaying potential signs of narcotic misuse and abuse such as chronic early refills, short dosing intervals, frequent dose increases, multiple lost/stolen etc scripts and intolerance or "allergy" to all products but Oxycontin. Allergic reactions to any product within a specific narcotic class will justify and preclude use of any other product in the same class due to the risk of cross-hypersensitivity. Beginning January 2017, all current opiate users who are above the maximum combined daily dose of 100 MME must titrate their total daily dose of opioid medications below 300 MME. Also, the maximum daily supply of an opiate prescription for acute pain will be limited to 7-day supplies. The maximum day supply of an opiate prescription for chronic pain will be limited to 30-day supplies. As of July 1, 2017 all users of opioid medications must comply with the maximum combined daily dose of 100 MME. However, for MaineCare members, effective January 1, 2017, opioid prescription(s) for more than a 7-day supply and/or more than 30 MME/ day will require a prior authorization. Please note that MaineCare implemented a 30 MME limit January 1, 2013 that is still effective. Post-surgical members may receive prior authorizations for opiates up to a 60 days in length if medical necessity is provided by the surgical provider. An MME conversion chart is available at www.mainearepdl.org. Click on "General Pharmacy Info." Please see the Pain Management Policy tab for the complete criteria

MISCELLANEOUS NARCOTICS

MISCELLANEOUS NARCOTICS							
NARCOTICS - MISC.	MC/DEL		ACETAMINOPHEN/CODEINE	MC/DEL	8	ABSTRAL	<p>1. Fentanyl OT loz (Barr) and Capital and codeine suspension products require PA for users over 18 years of age. PA is not required if under 18 years of age.</p> <p>2. Oxycodone/acet 10/650 is 8 times more expensive. Use twice as many of oxycod/acet 5/325 instead. You can mix and match preferred strengths of oxycodone and oxycodone/acet to minimize acet. dose similar to certain non-preferred drugs.</p> <p>3. Only preferred manufacturer's products will be available without prior authorization.</p>
	MC/DEL		ASPIRIN/CODEINE TABS	MC/DEL	8	APADAZ	
	MC/DEL		BUTAL/ASA/CAFF/COD CAPS	MC/DEL	8	ASCOMP/CODEINE CAPS	
	MC		BUTALBITAL/ASPIRIN/CAFFEI CAPS	MC/DEL	8	BUTALBITAL/APAP/CAFFEINE/ CAPS	
	MC		CAPITAL AND CODEINE SUSP ¹	MC/DEL	8	BUTALBITAL COMPOUND- CODEINE CAP	
	MC		CAPITAL/CODEINE SUSP ¹	MC	8	DEMEROL	
	MC/DEL		CODEINE PHOSPHATE SOLN	MC/DEL	8	DILAUDID	
	MC/DEL		CODEINE SULFATE TABS	MC	8	DILAUDID-HP SOLN	
	MC/DEL		ENDOCET TABS ³	MC	8	FENTANYL CITRATE SOLN	
	MC/DEL		ENDODAN TABS	MC/DEL	8	FENTORA	
	MC/DEL		FENTANYL OT LOZ ¹	MC/DEL	8	FIORICET/CODEINE CAPS	
	MC/DEL		FENTANYL OT LOZ1	MC	8	FIORINAL/CODEINE #3 CAPS	
	MC/DEL		HYDROCODONE/ACETAMINOPHEN	MC	8	FIORTAL/CODEINE CAPS	
	MC/DEL		HYDROMORPHONE HCL ³	MC/DEL	8	HYDROCODONE/IBUPROFEN	
	MC		LORTAB ELX	MC/DEL	8	HYDROMORPHONE ER	
	MC/DEL		MEPERIDINE SOL	MC/DEL	8	HYDROMORPHONE RECTAL SUPP	
	MC/DEL		OXYCODONE TAB	MC	8	IBUDONE	
	MC/DEL		OXYCODONE/ACETAMINOPHEN ^{2,3}	MC/DEL	8	LEVORPHANOL TARTRATE TAB	
	MC/DEL		ROXICET	MC/DEL	8	LORCET	
	MC		ROXIPRIN TABS	MC	8	LORTAB	
				MC	8	MAXIDONE TABS	
				MC/DEL	8	MEPERIDINE TABS	
				MC/DEL	8	NORCO TABS	
				MC/DEL	8	ONSOLIS	
				MC/DEL	8	OXECTA	
				MC/DEL	8	OXYCODONE CAP	
				MC/DEL	8	OXYCODONE/APAP 10/650	
				MC/DEL	8	OXYCODONE/APAP 7.5/500	
				MC/DEL	8	PENTAZOCINE/ACET TABS	
				MC/DEL	8	PENTAZOCINE/NALOXONE TABS	
				MC	8	PERCOCET TABS	
				MC	8	PERCOCET TABS	
				MC	8	PHRENILIN W/CAFFEINE/CODE CAPS	
				MC/DEL	8	ROXICET 5/500 TABS	
				MC	8	ROXICODONE TABS	
				MC/DEL	8	ROXYBOND	
				MC	8	SYNALGOS-DC CAPS	
				MC	8	TALACEN TABS	
				MC	8	TREZIX	
				MC	8	TYLENOL/CODEINE #3 TABS	
			MC	8	TYLOX CAPS		
			MC	8	XOLOX		
			MC	8	VICODIN		
			MC	8	VICOPROFEN TABS		
			MC	8	ZYDONE TABS		
			MC	9	ACTIQ LPOP		
			MC	9	CONZIP		
			MC	9	OPANA		
OPIOID DEPENDENCE TREATMENTS	MC		SUBOXONE FILM ²	MC/DEL		BUPRENORPHINE ¹	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>Use PA Form #20420</p> <p>Use PA form #10300 for PAs over the opiate limit</p> <p>Use PA Form #20100</p>

	MC/DEL		BUPRENORPHINE/NALOXONE TABS ²	MC		ZUBSOLV	<p>1. Buprenorphine will only be approved for use during pregnancy.</p> <p>2. See Criteria Section</p>	<p>Members will continue to be required to follow the criteria listed below:</p> <p>1-Induction period for 30 days</p> <p>2-Max dose of 32 mg for induction</p> <p>3-Max dose of 24 mg for maintenance</p> <p>4-There is not more than one opioid fill in member's drug profile between current fill of buprenorphine and a prior buprenorphine fill within the past 90 days</p> <p>5- Should provide evidence of monthly monitoring including random pill counts, urine drug tests and use of Maine Prescription Monitoring Program reports.</p> <p>6- Buprenorphine monotherapy is preferred if member is pregnant and dose not > 24 mg day and pregnancy diagnosis is noted on the prescription.</p>
EXTENDED RELEASE BUPRENORPHINE	MC MC		BRIXADI ¹ SUBLOCADE ¹				<p>Use PA form #20200 for Extended Release Buprenorphine</p> <p>1. Clinical PA required.</p>	<p>Brixadi and Sublocade:</p> <p>The prescriber can attest (and medical record should document) that:</p> <ul style="list-style-type: none"> -member has a documented history of opioid use disorder (OUD), -XRB is being used for the treatment of OUD (rather than pain or any other non-FDA approved indication) and -member's total daily dose of sublingual buprenorphine is less than or equal to 24 mg daily. <p>AND at least one of the following is true:</p> <ul style="list-style-type: none"> -The member's previous use of sublingual buprenorphine has included misuse, overuse, or diversion. -The member is at high risk of overdose (e.g., individuals leaving incarceration or abstinence-based treatment programs; individuals who are unhoused; or those facing potential gaps in care due to delays in care or geographically limited treatment access). -The member has experienced significant medical complications of OUD and/or of injection drug use. Occurrence should be in the last 5 years, or it should be clearly documented that the risk indicated by this infection or complication is ongoing (Examples of medical complications of OUD include: threatened the function of organs or life or limb threatening and required medical and/or surgical therapy. Examples of medical complications of injection drug use include osteomyelitis, endocarditis, renal failure, joint infection or other serious medical complications directly related to OUD.) -The member has treatment-resistant OUD, including those with ongoing illicit substance use in the context of sublingual buprenorphine treatment as documented by positive urine drug screens or other clear objective evidence, and/or further functional decline with explicit documentation of the functional decline. -The member has a significant intolerance of, or documented allergy to, sublingual buprenorphine (either buprenorphine monotherapy or buprenorphine/naloxone combination therapy) that has resulted in the patient's inability to comply with continued treatment using the sublingual product. (A true allergy is usually accompanied by rash, respiratory symptoms, or anaphylaxis. Other complaints such as bad taste, mouth tingling, etc. do not constitute evidence of allergy or significant intolerance. Formulation preference or convenience are not, in and of themselves, indications for using XRB.) -The member is in ongoing treatment with XRB and would like to continue the medication.
OPIOID WITHDRAWAL AGENTS				MC		LUCEMYRA ¹	<p>1. Clinical PA for appropriate approved use and patient has documented contraindication to clonidine.</p> <p>Use PA Form#20420</p>	
NARCOTIC ANTAGONISTS								
NARCOTIC - ANTAGONISTS	MC/DEL MC MC MC MC		NALTREXONE HCL TABS NALOXONE INJ NARCAN NS NALOXONE SPRAY OTC VIVITROL INJ ZIMHI	MC MC MC/DEL		EVZIO OPVEE ² KLOXXADO REVIA TABS ¹	<p>Use PA Form# 20420</p> <p>1. Will only be approved for side effects experienced with generic that are not described in the literature as occurring with the brand version.</p> <p>2. For the treatment of adult and pediatric patients 12 years of age and older.</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p>
COX 2 / NSAIDS								
COX 2 INHIBITORS - SELECTIVE / HIGHLY SELECTIVE	MC/DEL MC/DEL		CELECOXIB ^{4,5} KETOROLAC TROMETHAMINE ^{2,3,5}	MC/DEL MC/DEL		CELEBREX CAPS ^{4,5} MELOXICAM CAPS ⁵	<p>Use PA Form# 20420</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p>

				MC		SB IBUPROFEN TABS		
				MC		SPRIX		
				MC		TIVORBEX		
				MC		TOLECTIN		
				MC		V-R IBUPROFEN TABS		
				MC		ZORVOLEX		

NSAID - PPI				MC		PREVACID NAPRA-PAC		1. Use a preferred NSAID and PPI separately.
				MC/DEL		VIMOVO ¹		Use PA Form# 20420

RHEUMATOID ARTHRITIS

RHEUMATOID ARTHRITIS	MC/DEL		ACTEMRA VIALS			ADALIMUMAB-AACF		Use PA Form# 20900	See criteria as listed on Rheumatoid Arthritis PA form.
	MC/DEL		ACTEMRA SYRINGES	MC		AMJEVITA			
	MC/DEL		ADALIMUMAB-FKJP ³	MC/DEL		ARAVA		1. Dosing limits apply. Please see dose consolidation list.	Preferred injectable products allowed without PA if trial of a preferred oral agents (azathioprine, hydroxychloroquine, leflunomide, methotrexate, sulfasalazine tabs) are seen in the members drug profile. Dosing limits apply.
	MC		AVSOLA	MC/DEL		CIMZIA			
	MC/DEL		AZATHIOPRINE	MC/DEL		CYLTEZO		2. Established users will be grandfathered.	
	MC		ENBREL ²	MC/DEL		ENTYVIO			
	MC		ENBREL SURECLICK ²	MC		HADLIMA		3. Clinical PA is required to establish diagnosis and medical necessity.	Xeljanz is limited to adults with moderate to severe RA and UC who have had an inadequate response or intolerance to methotrexate. Should not be used concomitantly with biologic DMARDs or potent immunosuppressants.
	MC		KINERET SOLN	MC/DEL		HULIO		4. Verification of age for appropriate indication.	Jylamvo will require using preferred methotrexate if unable please provide clinical rational as why inappropriate.
	MC/DEL		LEFLUNOMIDE	MC/DEL		HYDROXYCHLOROQUINE ²		5. Treatment failure or intolerance to other forms of preferred methotrexate	Zymfentra: In adults for maintenance treatment of: Moderately to severely active ulcerative colitis following treatment with an infliximab product administered intravenously. Moderately to severely active Crohn's disease following treatment with an infliximab product administered intravenously.
	MC/DEL		METHOTREXATE	MC/DEL		HYRIMOZ		6. See criteria section	
	MC		ORENCIA	MC		IDACIO			
	MC/DEL		SULFASALAZINE TABS	MC/DEL		ILARIS ^{1,3,4}			
	MC		SIMLANDI ³	MC/DEL		INFLECTRA			
	MC		SIMPONI PEN	MC		INFLIXIMAB VIAL			
	MC		SIMPONI AUTOINJECTOR	MC		JYLAMVO			
	MC/DEL		RINVOQ ³	MC/DEL		KEVZARA			
	MC		HUMIRA ^{1,2}	MC		OLUMIANT			
	MC/DEL		XELJANZ ^{3,6}	MC		OMVOH			
	MC/DEL		XELJANZ XR	MC		OTREXUP			
	MC/DEL		XELJANZ XR SOL	MC		RASUVO ⁷			
				MC		REDITREX			
				MC		REMICADE			
				MC/DEL		RENFLEXIS			
				MC		SIMLANDI			
				MC		TOFIDENCE			
				MC		VELSIPITY			
				MC		YUFLYMA			DDI: The concomitant use of Xeljanz® XR with biologic DMARDs or potent immunosuppressants such as azathioprine and cyclosporine are not recommended. The concomitant use of Xeljanz® XR with potent CYP3A4 inducers (e.g. rifampin) is not recommended
				MC		YUSIMRY			
				MC		XATMEP ⁵			
				MC		ZYMFENTRA			

ALOPECIA AREATA AGENTS

ALOPECIA AREATA AGENTS				MC	7	OLUMIANT			
				MC/DEL	8	LITFULO			Preferred drug must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
								Use PA Form# 20420	

MISCELLANEOUS ARTHRITIS

ARTHRITIS - MISC.	MC		RIDAURA CAPS	MC/DEL		ARTHROTEC ¹		1. The individual components of Arthrotec are available without PA.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. The individual components of Arthrotec are available without PA.
	MC		MYOCHRYSLINE SOLN					Use PA Form# 20420	

LUPUS-SLE

LUPUS-SLE				MC MC MC		BENLYSTA ¹ LUPKYNIS SAPHNELO	Use PA Form# 20420 1. Approvals will require previous trial of corticosteroids, antimalarials, NSAIDs and immunosuppressives.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved (in step order), unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. DDI: Lupkynis is a sensitive CYP3A4 substrate. Co-administration with strong or moderate CYP3A4 inhibitors increases voclosporin exposure, which may increase the risk of Lupkynis® adverse reactions. Co-administration of Lupkynis® with strong CYP3A4 inhibitors (e.g. ketoconazole, itraconazole, clarithromycin) is contraindicated. Reduce Lupkynis® dosage when co-administered with moderate CYP3A4 inhibitors (e.g. verapamil, fluconazole, diltiazem)
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PIK3CA-Related Overgrowth Spectrum (PROS)

PIK3CA-Related Overgrowth Spectrum (PROS)				MC		VIJOICE ¹	Use PA Form# 20420 1. PA required to confirm FDA approved indication.	Preferred drugs must be tried and failed, in step-order, due to lack of efficacy (failure to reach target IOP reduction) or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
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MIGRAINE THERAPIES

MIGRAINE - ERGOTAMINE DERIVATIVES				MC/DEL MC		D.H.E. 45 SOLN TRUDHESA	Use PA Form# 10110	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
MIGRAINE - CARBOXYLIC ACID DERIVATIVES	MC		DIVALPROEX ER TB24	MC		DEPAKOTE ER TB24	Use PA Form# 10110	
MIGRAINE - SELECTIVE SEROTONIN AGONISTS (5HT)--Tabs/Nasal	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	1 1 1 1 1 1 2	MIGRANAL NASAL SPRAY RELPAX ¹ RIZATRIPTAN ODT RIZATRIPTAN TABS SUMATRIPTAN TABS ¹ ZOLMITRIPTAN TAB ¹ NARATRIPTAN HCl TABS ¹	MC MC MC/DEL MC MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL		AMERGE TABS ^{1,2} AXERT TABS ^{1,2} FROVA TABS ^{1,2} IMITREX NASAL SPRAY ¹ IMITREX TABS ^{1,2} MAXALT ^{1,2,3} MAXALT MLT ^{1,2,3} ONZETRA XSAIL ² SUMATRIPTAN NASAL SPRAY ¹ ZOLMITRIPTAN ODT ZOLMITRIPTAN SPRAY ZOMIG TABS ^{1,2} ZOMIG NASAL SPARY ^{1,2} ZOMIG ZMT TBP ^{1,2}	1. All drugs in this category have dosing limits. Please refer to dose consolidation table. 2. Must fail all preferred products before non-preferred. 3. Established users will be grandfathered Use PA Form# 10110	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Quantity limit exceptions will require ongoing therapy with therapeutic doses of highly effective prophylactic medication as listed on the Triptan PA form.
MIGRAINE - SELECTIVE SEROTONIN AGONISTS (5HT)--Injectables	MC MC/DEL MC/DEL		IMITREX CARTRIDGE ¹ SUMATRIPTAN SYRINGE ¹ SUMATRIPTAN PEN INJCTR ¹	MC/DEL MC MC		TOSYMRA ZEMBRACE ¹ IMITREX PEN INJCTR ¹	Use PA Form# 10110 1. Dosing limits apply. Please refer to the dose consolidation table.	
MIGRAINE - SELECTIVE SEROTONIN AGONISTS (5HT)--Combinations				MC/DEL		TREXIMET ^{1,2}	Use PA Form# 10110 1. Dosing limits apply. Please see dose consolidation list. 2. Use preferred Sumatriptan and Naproxen separately. Treximet only available if component ingredients of sumatriptan and naproxen are unavailable.	
MIGRAINE - PREVENTATIVE TREATMENT	MC MC/DEL MC/DEL		AIMOVI ¹ AJOVY ¹ AJOVY AUTO INJCT ¹	MC MC MC		NURTEC ODT ² QULIPTA VYEPTI ²	Use PA Form# 10110 1. See criteria section 2. Dosing limits apply,	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.

	MC/DEL MC/DEL		EMGALITY SYRINGE ¹ 200mg/ml EMGALITY PEN ¹				please see the dose consolidation list.	Aimovig, Ajovy and Emgality: The patient is 18 years of age or older AND patient has a diagnosis of episodic migraine (4-14 headache days per month with migraine lasting 4 hours or more) or chronic migraine (≥ 15 headache days per month, of which ≥ 8 are migraine days, for at least 3 months) AND patient has failed or has a contraindication to an adequate trial (≥ 60 days) of at least 2 medications for migraine prophylaxis from at least 2 different classes. Ubrelyv is non-preferred and is indicated for the acute treatment of migraine with or without aura in adults. This is not indicated for the preventive treatment of migraine. Nurtec ODT will be preferred after 2 adequate trials of at least two preferred triptans
MIGRAINE - ACUTE TREATMENT	MC MC/DEL		NURTEC ODT ¹ SPASTRIN TABS	MC MC/DEL MC/DEL MC MC MC/DEL		BELCOMP-PB SUPP ELYXYB MIGRAZONE CAPS MIGERGOT SUP REYVOW UBRELVY ZAVZPRET	1. Dosing limits apply, please see the dose consolidation list. Use PA Form# 10110	Reyvow is non-preferred and is indicated for the acute treatment of migraine with or without aura in adults. Reyvow® is not indicated for the preventive treatment of migraine. Zavzpret: The patient must have a documented side effect, allergy, or treatment failure to preferred oral CGRP Inhibitor and two non-preferred oral CGRP Inhibitors. Ubrelyv is non-preferred and is indicated for the acute treatment of migraine with or without aura in adults. This is not indicated for the preventive treatment of migraine. Nurtec ODT will be preferred after 2 adequate trials of at least two preferred triptans
GOUT								
GOUT	MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL		ALLOPURINOL TABS COLCHICINE TAB FEBUXOSTAT TAB MITIGARE PROBENECID TABS PROBENECID/COLCHICINE TABS	MC/DEL MC MC MC/DEL MC		COLCHICINE CAP COLCRYS GLOPERBA ULORIC ¹ ZYLOPRIM TABS	Use PA Form# 20420 1. Failure of therapeutic (300mg) dose of Allopurinol (failure define as not being able to get uric acid levels below 6mg/dl) or severe renal disease.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. DDI: The concomitant use of Gloperba® and CYP3A4 inhibitors (e.g. clarithromycin, ketoconazole, grapefruit juice, erythromycin, verapamil, etc.) should be avoided due to the potential for serious and life-threatening toxicity.
MISC.								
ACID SPHINGOMYELINASE DEFICIENCY (ASMD)				MC		XENPOZYME ^{1,2}	1.For treatment of non-central nervous system manifestations of acid sphingomyelinase deficiency (ASMD) in adult and pediatric patients 2. Clinical PA required for appropriate diagnosis and clinical parameters.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
ANESTHETICS - MISC.	MC MC MC		BUPIVACAINE HCL SOLN LIDOCAINE HCL SOLN MARCAINE SOLN	MC MC/DEL MC		SENSORCAINE-MPF SOLN SYNVISC INJ XYLOCAINE SOLN	Use PA Form# 30130	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
COLD AGGLUTININ DISEASE (CAD)				MC		ENJAYMO ¹	1. Indicated to decrease the need for red blood cell transfusion due to hemolysis in adults with cold agglutinin disease (CAD).	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
CONGENITAL ADRENAL HYPERPLASIA				MC		CRENESSITY	Use PA Form# 20420	Crenessity - As adjunctive treatment to glucocorticoid replacement to control androgens in adults and pediatric patients 4 years of age and older with classic congenital adrenal hyperplasia (CAH)
PRIMARY HYPEROXALURIA TYPE 1 (PH1)						OXLUMO ¹ RIVFLOZA	1. PA is required to establish diagnosis and medical Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Rivfloza: The patient has a diagnosis of Primary Hyperoxaluria Type I (PH1) confirmed via genetic testing (identification of alanine: glyoxylate aminotransferase gene (AGXT) mutation) AND urinary oxalate excretion > 0.5mmol/1.73 m2 or urinary oxalate: creatinine ratio is above the upper limit of normal for age AND is at least 9 years of age AND medication is being

SICKLE CELL DISEASE	MC MC/DEL MC		DROXIA HYDROXYUREA LYFGENIA ^{2,3}	MC MC MC MC/DEL		ADAKVEO CASGEVY ^{2,3} ENDARI ¹ SIKLOS	1.Evidence of other preferred L-glutamine products utilization and reason for failure. 2. For the treatment of patients ≥ 12 years of age. 3. PA required to confirm FDA approved indication. Use PA Form# 20420	prescribed by, or in consultation, with a nephrologist or urologist Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
HUTCHINSON- GILFORD PROGERIA SYNDROME (HGPS)				MC		ZOKINVY ^{1,2}	1.In patients 12 months of age and older with a body surface area (BSA) of 0.39m2 and above 2. PA required to confirm FDA approved indication. Use PA Form# 20420	ZOKINVY: To reduce the risk of mortality in Hutchinson-Gilford Progeria Syndrome (HGPS). For the treatment of processing-deficient Progeroid Laminopathies with either: Heterozygous LMNA mutation with progerin-like protein accumulation OR Homozygous or compound heterozygous ZMPSTE24 mutations
OBSTRUCTIVE SLEEP APNEA				MC		ZEPBOUND	Use PA Form# 20420	Zepbound for adults with a BMI ≥ 30 mg/kg2 and diagnosis of moderate to severe OSA, confirmed by sleep study within the last 3 years documenting AHI ≥ 15, AND in which CPAP is ineffective (AHI > 5 during therapeutic section of sleep study) or patient is unable to tolerate CPAP for at least 90 days AND for whom lifestyle modifications have been attempted for at least 3 months with failure to achieve weight loss. Note: Not for patients with T1DM, T2DM
VACCINES	MC/DEL MC MC/DEL MC/DEL		ABRYSVO AREXVY GARDASIL 9 SHINGRIX				Use PA Form# 20420	Gardasil 9 will be preferred by MaineCare for ages 19-45 for FDA approved indications. Under the Maine Immunization Program Gardasil 9 is covered under the Vaccine for Children Program for ages 9-18. Please contact 1-800-867-4775 or 207-287-3746 for assistance. Abrysvo will be a preferred vaccine indicated for active immunization for the prevention of lower respiratory tract disease (LRTD) caused by respiratory syncytial virus (RSV) in individuals 60 years of age and older. Active immunization of pregnant individuals at 32 through 36 weeks gestational age for the prevention of LRTD and severe LRTD caused by RSV in infants from birth through 6 months of age. Arexvy will be preferred for active immunization for the prevention of LRTD caused by respiratory syncytial virus (RSV) in individuals 60 years of age and older. SHINGRIX (>= 50yo) is preferred as of 11-20-20 with respective age edit.
APDS				MC		JOENJA ^{1,2,3}	Use PA Form# 20420 1.Clinical PA required for appropriate diagnosis 2. For the treatment of patients 2 years of age and older. 3. Avoid CYP3A drug interaction.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
ALPHA- MANNOSIDOSIS				MC		LAMZEDE	Use PA Form# 20420 1.Clinical PA required for appropriate diagnosis	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
ANTI-CONVULSANTS								
ANTICONSULSANTS	MC/DEL		BRIVIACT	MC	8	APTIOM	Use PA Form# 20420	

MC/DEL	CARBAMAZEPINE	MC	8	BANZEL	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
MC	CARBAMAZEPINE ER CAP	MC	8	CARBAMAZEPINE SUS	
MC/DEL	CARBATROL CP12	MC	8	DEPAKOTE	All non-preferred meds must be used in specified order
MC/DEL	CELONTIN CAPS	MC	8	DEPAKOTE ER	
MC/DEL	CLOBAZAM	MC	8	DIACOMIT	1. Quantity limit. 5/month
MC/DEL	CLONAZEPAM TABS	MC/DEL	8	DIVALPROEX SODIUM SPRINKLE CAPS	
MC	DEPAKOTE SPRINKLES CPSP	MC	8	ELEPSIA XR ⁹	2. Dosing limits apply, please see dose consolidation list.
MC/DEL	DIAZEPAM GEL ¹	MC	8	EPRONTIA SOLN ¹⁰	
MC/DEL	DILANTIN	MC/DEL	8	FELBATOL	Approvals will be for patients with a variety of drug-specific FDA-approved indications and for specific conditions supported by at least two published peer-reviewed double-blinded, placebo-controlled randomized trials that are not contradicted by other studies of similar quality after recommendation by the DUR Committee and as long as all first line therapies have been tried and failed at full therapeutic doses for adequate durations (at least two weeks).
MC/DEL	DIVALPROEX SODIUM	MC/DEL	8	FELBATOL SUS	
MC	DIVALPROEX SPRINKLE CAP	MC/DEL	8	FELBAMATE SUS	3. Dosing limits apply per strength as well as a maximum daily dose of 600mg. Please see dose consolidation list.
MC/DEL	EPIDIOLEX ⁷	MC	8	FINTEPLA ⁸	
MC/DEL	EPITOL TABS	MC	8	FYCOMPA ²	4. Adjunctive therapy 17 and older.
MC/DEL	ETHOSUXIMIDE SYRP	MC/DEL	8	HORIZANT	
MC/DEL	EQUETRO	MC	8	GRALISE	5. Max dose 2400mg
MC/DEL	GABAPENTIN ² CAP	MC/DEL	8	KEPPRA TABS	
MC/DEL	GABAPENTIN ² TAB	MC/DEL	8	KEPPRA SOLN	6. Clinical PA required for appropriate diagnosis
MC/DEL	GABAPENTIN SOL	MC/DEL	8	KLONOPIN TABS	
MC/DEL	GABITRIL TABS	MC	8	LAMICTAL IR	Topamax and Neurontin - Second line therapy for migraine prophylaxis after trial of at least three preferred preventive medications from Group 1 listed on page 2 of the Acute Migraine PA form.
MC/DEL	LACOSAMIDE SOL	MC	8	LAMICTAL ODT	
MC/DEL	LACOSAMIDE TAB	MC	8	LAMICTAL XR	7. Epidiolex is for the treatment of seizures associated with Lennox-Gastaut syndrome (LGS), Dravet syndrome (DS) or TS (Tuberous Sclerosis Complex) in patients 1 years of age and older.
MC	LAMICTAL CHEW	MC/DEL	8	LEVETIRACETAM INJ	
MC/DEL	LAMOTRIGINE ER ODT	MC	8	LIBERVANT	All non-preferred meds must be used in specified order.
MC/DEL	LAMOTRIGINE IR ²	MC/DEL	8	LYRICA CR	
MC/DEL	LAMOTRIGINE XR	MC/DEL	8	LYRICA SOL ³	Please use Drug-Drug Interaction PA form #10400 for this combination.
MC/DEL	LEVETIRACETAM SOLN	MC	8	MOTPOLY XR	
MC/DEL	LEVETIRACETAM TABS	MC/DEL	8	MYSOLINE TABS	8. For seizures associated with Dravet syndrome in patients 2 years of age and older
MC/DEL	LEVETIRACETAM ER TABS	MC	8	ONFI	
MC/DEL	LYRICA ³	MC/DEL	8	OXCARBAZEPINE SUS	Epidiolex Criteria for Lennox-Gastaut syndrome (LGS) and Dravet: a trial of two drugs (clobazam, levetiracetam, valproate derivatives, lamotrigine, topiramate, rufinamide, or felbamate).
MC/DEL	NAYZILAM ¹	MC	8	OXTELLAR XR ⁵	
MC/DEL	OXCARBAZEPINE	MC/DEL	8	PHENYTEK CAPS	Diacomit is for the treatment of seizures associated with Dravet syndrome (DS) in patients 6 months of age and older and weighing 7kg or more There are no clinical data to support the use of Diacomit® as monotherapy in DS.
MC/DEL	PREGABALIN CAPS	MC/DEL	8	POTIGA	
MC/DEL	PHENYTOIN	MC/DEL	8	PREGABALIN (ORAL) SOL	9. Adjunctive therapy 12 and older.
MC/DEL	PRIMIDONE TABS	MC	8	ROWEEPRA TAB	
MC/DEL	QUDEXY XR	MC	8	SABRIL	DDI: Concomitant use of Diacomit® with other CNS depressants, including alcohol, may increase the risk of sedation and somnolence. Concomitant use of strong inducers (CYP1A2, CYP3A4, or CYP2C19 inducers, such as rifampin, phenytoin, phenobarbital, and carbamazepine) should be avoided, or dosage adjustments should be made.
MC/DEL	TEGRETOL SUS	MC	8	SEZABY	
MC/DEL	TOPIRAMATE	MC	8	SPRITAM	DDI: Avoid concomitant use of Nayzilam® with moderate or strong CYP3A inhibitors.
MC/DEL	TOPIRAMATE SPRINKLE IR CAPS	MC	8	SYMPAZAN	
MC/DEL	TRILEPTAL SUS	MC/DEL	8	TEGRETOL TAB	10. Initial monotherapy for the treatment of partial-onset or primary generalized tonic-clonic seizures in patients 2 years of age and older.
MC/DEL	VALPROIC ACID TABS	MC/DEL	8	TIAGABINE	
MC/DEL	VALPROIC ACID SOL	MC	8	TOPAMAX	Adjunctive therapy for the treatment of partial-onset seizures, primary generalized tonic-clonic seizures, and seizures associated with Lennox-Gastaut syndrome in patients 2 years of age and older. The preventive treatment of migraine in patients 12 years and older.
MC	VALTOCO ²	MC/DEL	8	TOPIRAMATE ER CAPS	
MC/DEL	ZONISAMIDE	MC	8	TOPAMAX SPRINKLE ER CAPS ²	Motpoly XR: pediatric patient weight must be > 50kg and requires multiple preferred medication trials including generic lacosamide
		MC	8	TOPAMAX SPRINKLE IR CAPS ²	
		MC/DEL	8	TOPIRAMATE SPRINKLE ER CAPS ²	Libervant: For the acute treatment of intermittent, stereotypic episodes of frequent seizure activity (i.e., seizure clusters, acute repetitive seizures) that are distinct from a patient's usual seizure pattern in patients with epilepsy 2 to 5 years of age as long as all preferred therapies have been tried and failed at full therapeutic doses.
		MC	8	TROKENDI ^{2,6}	
		MC	8	VIGAFYDE	Vigafyde: Indicated as monotherapy for the treatment of infantile spasms in pediatric patients 1 month to 2 years of age for whom the potential benefits outweigh the potential risk of vision loss.
		MC/DEL	8	VIMPAT ⁴	
		MC/DEL	8	VIMPAT SOL ⁴	
		MC	8	XCOPRI	
		MC/DEL	8	ZARONTIN SYRP	
		MC/DEL	8	ZARONTIN CAP	
		MC/DEL	8	ZARONTIN SOL	
		MC	8	ZONISADE	
		MC	8	ZTALMY	

AGONISTS	MC/DEL		ROPINIROLE	MC MC/DEL MC/DEL	8 8 8	REQUIP TABS MIRAPEX ER NEUPRO PATCH	1. As of 12/08 users of Mirapex will be grandfathered if diagnosis is Parkinson's.	is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
PARKINSONS- MAOIS				MC		XADAGO		Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Use PA Form# 20420
PARKINSONS - DOPAMINERGICS/CARBII/ LEVO	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL		AMANTADINE HCLCAPS AMANTADINE HCL TABS BROMOCRIPTINE MESYLATE TABS BROMOCRIPTINE MESYLATE CAPS CARBIDOPA/LEVODOPA TABS ³ CARBIDOPA/LEVODOPA ER CARBIDOPA/LEVO/ENTACAPONE TAB LARODOPA TABS SELEGILINE CAPS HCL SELEGILINE TABS HCL	MC/DEL MC MC/DEL MC MC MC MC/DEL MC MC MC MC/DEL MC/DEL MC MC MC MC		APOKYN AZILECT ² CARBIDOPA/LEVODOPA RAPDIS CREXONT ⁴ ELDEPRYL CAPS GOCOVRI INBRIJA KYNMOBI LODOSYN TABS OSMOLEX ER PARLODEL CAPS PARLODEL TABS RYTARY SINEMET TABS SINEMET TBCR ZELAPAR ¹	1. Approvals will require concurrent therapy with Levodopa and failed trials of Selegiline, Comtan, and Stalevo. 2. Approvals will require trials of Carbidopa/Levodopa, Selegiline, Comtan, and Stalevo. 3. Only preferred manufacturer's products will be available without prior authorization. 4. Approvals will require trials of preferred medications including extended-release levodopa/carbidopa tablets Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Inbrija is recommended for the intermittent treatment of OFF episodes in patients with Parkinson's disease treated with carbidopa/levodopa.
PARKINSONS - COMBO.				MC/DEL MC		STALEVO ¹ CARBIDOPA/LEVODOPA/ENTACA ¹	Use PA Form# 20420	1.Clinical PA is required to establish diagnosis and medical necessity.
MUSCLE RELAXANTS								
MUSCLE RELAXANTS	MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL		BACLOFEN TABS CHLORZOXAZONE TABS CYCLOBENZAPRINE HCL 5mg & 10mg TABS LIORESAL INTRATHECAL KIT METHOCARBAMOL TABS TIZANIDINE HCL TABS	MC/DEL MC/DEL MC/DEL MC/DEL MC MC MC MC MC/DEL MC MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	7 8 8 8 8 8 8 8 8 8 8 8 8 9 9 9 9	ORPHENADRINE CITRATE CARISOPRODOL 350MG TABS AMRIX DANTRIUM CAPS FLEQSUVY LIORESAL TABS LORZONE LYVISPAP METAXALONE NORFLEX TBCR OZOBAX ROBAXIN-750 TABS VECUROMIUM INJ ZANAFLEX TABS CARISOPRODOL 250MG TABS CHLORZOXAZONE 250mg TABS SKELAXIN TAB SOMA TABS		Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. At least 4 preferred drugs (including tizanidine) must be tried for at least 2 weeks and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Elderly patients, over 65, will require written notice of the increased sedative risks and impaired driving. Prior Authorization will not be given for:1. frequent or persistent early refills of controlled drugs; 2. multiple instances of early refill overrides due to reports of misplacement, stolen, dropped in toilet or sink, distant travel, etc. Non-preferred products must be used in specified step order. Non-preferred drugs will not be approved if members circumventing MaineCare prior authorization requirements by paying (prescribers failed to submit prior authorization prior to cash narcotic scripts being filled by member).

				MC	9	TANLOR	Use PA Form# 20420	Lorzone is non preferred and requires at least 4 preferred drugs (including tizanidine) and step care therapy (orphenadrine), as well as reasons for why chlorzoxazone is not acceptable.
MUSCLE RELAXANT - COMBO.				MC/DEL MC/DEL MC MC/DEL MC/DEL MC		CARISOPRODOL/ASPIRIN TABS CARISOPRODOL/ASPIRIN/CODE NORGESIC TABS ORPHENADRINE COMPOUND ORPHENADRINE/ASA/CAFF ORPHENGESIC	Use PA Form# 20420	Individual components are available with PA described in the section above. 1. frequent or persistent early refills of non-controlled drugs; 2. multiple instances of early refill overrides due to reports of misplacement stolen, dropped in toilet or sink, distant travel, etc.
PARATHYROID HORMONE								
PARATHYROID HORMONE				MC MC		NATPARA ¹ YORVIPATH ¹	Use PA Form# 20420	1. Recommended only for those who cannot be well-controlled on calcium supplements and active forms of vitamin D alone. Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
VITAMINS								
VITAMINS	MC MC MC MC/DEL MC/DEL MC/DEL MC MC/DEL MC MC MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL MC MC		CYANOCOBALAMIN SOLN FERIVA CAP FERIVAFA CAP FOLIC ACID TABS MEPHYTON TABS NIACIN NIACOR TABS NICOTINIC ACID SR CPCR PYRIDOXINE HCL TABS TANDEM CAP THIAMINE HCL SOLN VITAMIN B-1 TABS VITAMIN B-12 VITAMIN B-6 TABS VITAMIN C VITAMIN E CAPS VITAMIN E/D-ALPHA CAPS VITAMIN K1 SOLN V-R VITAMIN E CAPS	MC MC MC MC MC MC MC MC MC MC MC MC MC MC MC MC MC MC		AQUASOL E SOLN AQUAVIT-E SOLN DHT SOLN FUSION PLUS CAP HEMOCYTE PLU CAP INTEGRA CAP INTEGRA F CAP INTEGRA PLUS CAP NASCOBAL GEL TANDEM PLUS CAP	Use PA Form# 20420 Please refer to OTC list for covered products. Click here for the OTC List	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Certain drugs require specific diagnoses for approval. Please refer to OTC list for covered products. DDI: B-12 will now be non-preferred and require prior authorization if it is currently being used in combination with either Prevacid, pantoprazole, Prilosec, or any currently non preferred PPI. Preferred products that used to require diag codes still require diag codes unless indicated otherwise.
VITAMIN D's	MC/DEL MC/DEL MC/DEL MC/DEL MC		CALCITRIOL CAPS ¹ ROCALTROL VITAMIN D2 ² VITAMIN D3 ² VITAMIN DROPS PARICALCITOL CAPS	MC MC/DEL MC/DEL MC/DEL MC/DEL MC MC MC		CALCIJEX DOXERCALCIF CAP DOXERCALCIF INJ PARICALCITROL CAP PARICALCITROL INJ HECTOROL (ORAL) HECTOROL (PARENTERAL) RAYALDEE ZEMPLAR INJ ZEMPLAR CAPS	Use PA Form# 20420	1. Diagnosis of dialysis (renal failure) required. 2. Only specific NDCs available Preferred products require dialysis/renal failure diagnosis. Rayaldee requires clinical PA to verify stage 3 or 4 CKD.
EMZYMES								
POMPE DISEASE AGENTS				MC MC MC MC		NEXVIAZYME ¹ LUMIZYME OPFOLDA POMBILITI		All preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs (in step-order) will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. 1. For patients 1 year of age and older with late-onset Pompe disease (lysosomal acid alpha-glucosidase [GAA] deficiency). Pombiliti and Opfolda are for the treatment of adult patients with late-onset Pompe disease (lysosomal acid alpha-glucosidase [GAA] deficiency) weighing ≥40kg and who are not improving on their current enzyme replacement therapy (ERT).

MISC MULTI-VITAMINS

VITAMINS - MISC.

MC	CENTRUM TABS	MC
MC	CENTRUM JR/IRON CHEW	MC/DEL
MC	CENTRUM-LUTEIN TABS	MC
MC	CEROVITE ADVANCED FO TABS	MC
MC/DEL	CHEWABLE MULTIVIT/FL CHEW	MC
MC	COD LIVER OIL CAPS	MC
MC/DEL	COMPLETE NATAL DHA (ORAL) COMBO PKG	MC
MC	COMPLETE SENIOR TABS	MC
MC	DAILY MULTI VIT/IRON	MC
MC/DEL	DIALYVITE 1MG	MC
MC/DEL	DIALYVITE 800MG	MC
MC/DEL	FULL SPECTRUM B	MC
MC	M.V.I.-12 INJ	MC
MC	MULTI-VIT/FLUORIDE	MC/DEL
MC/DEL	NATALCARE RX TABS	MC/DEL
MC/DEL	NEPHRONEX	MC/DEL
MC/DEL	NIVA-PLUS (ORAL) TABLET	MC
MC/DEL	ONE DAILY TABS	MC
MC/DEL	ONE-DAILY MULTIVITAMINS	MC/DEL
MC/DEL	ONE-TABLET-DAILY	MC
MC/DEL	POLY-VIT/IRON/FLUORID SOLN	MC/DEL
MC/DEL	POLY-VITAMIN/FLUORIDE SOLN	MC
MC/DEL	POLY-VITAMINS/IRON SOLN	MC
MC	PRENATA (ORAL) TAB CHEW	MC/DEL
MC/DEL	PRENATAL TABS ¹	MC/DEL
MC/DEL	PRENATAL FORMULA 3 TABS ¹	MC/DEL
MC/DEL	PRENATAL PLUS TABS ¹	MC
MC/DEL	PRENATAL PLUS NF TABS ¹	MC/DEL
MC	PRENATAL PLUS/27MG IRON ¹	MC/DEL
MC	PRENATAL PLUS/IRON TABS ¹	MC
MC	PRENATAL VITAMIN PLUS LOW IRON (ORAL) TAB	MC
MC/DEL	PRENATAL RX/BETA-CAROTENE ¹	MC
MC/DEL	PREPLUS (ORAL) TABLET	MC
MC/DEL	RENAL CAPS	MC
MC/DEL	RENAPHRO CAPS	MC
MC	STRESS TAB NF TABS	MC
MC	THERAPEUTIC-M TABS	MC
MC	THERAVITE LIQD	MC
MC/DEL	TRINATAL RX 1 (ORAL) TABLET	MC
MC/DEL	TRIVEEN-DUO DHA (ORAL) COMBO. PKG	MC/DEL
MC/DEL	TRI-VITAMIN/FLUORIDE SOLN	MC
MC	VITA CON FORTE CAPS	MC
MC	VITAPLEX PLUS TABS	MC/DEL

ADEKS
ADVANCED NATALCARE TABS
AQUADEKS
CENTRUM JR/EXTRA C CHEW
CENTRUM PERFORMANCE TABS
CENTRUM SILVER TABS
DALYVITE LIQD
EMBREX 600 MISC
FERRALET 90
IBERET
MATERNA TABS
MAXARON
MULTIRET FOLIC -500 TBCR
NATAFORT TABS
NATALCARE CFE 60 TABS ¹
NATALCARE GLOSS TABS ¹
NATALCARE PIC TABS ¹
NATALCARE PIC FORTE TABS ¹
NATALCARE PLUS TABS ¹
NATALCARE THREE TABS ¹
NATACHEW CHEW
NATALFIRST TABS
NATATAB RX TABS
NEPHPLEX RX TABS
NEPHROCAPS CAPS
NEPHRO-VITE TABS
NESTABS RX TABS
NIFEREX
OCUVITE TABS
POLY-VI-FLOR SOLN
POLY-VI-SOL SOLN
POLY-VI-SOL/IRON SOLN
POLY-VITAMIN DROPS SOLN
PRECARE
PREFERA OB
PREMESIS RX TABS
PRENATABS CBF TABS ¹
PRENATAL CARE TABS ¹
PRENATAL MR 90 TBCR ¹
PRENATAL MTR/SELENIUM TABS ¹
PRENATAL OPTIMA ADVANCE TABS ¹
PRENATAL PC 40 TABS ¹
PRENATAL RX TABS ¹
PRENATE ¹
PRENATE ELITE ¹
PRIMACARE MISC
PROTEGRA CAPS
STUARTNATAL PLUS 3 TABS ¹
TRI-VI-SOL SOLN
TRI-VI-SOL/IRON SOLN
ULTRA NATALCARE TABS
ULTRA-NATAL TABS ¹

1. Diag codes are no longer required on prenatal vitamins.

Please refer to OTC list.

Use PA Form# 20420

Click here for the OTC List

Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Certain drugs require specific diagnoses for approval.

Please refer to OTC list.

Preferred products that used to require diag codes still require diag codes unless indicated otherwise.

			MC	VICON FORTE CAPS		
			MC	VINATAL FORTE TABS ¹		
			MC	VINATE ¹		
			MC/DEL	VINATE ADVANCED TABS ¹		
MISCELLANEOUS MINERALS						
MINERALS	MC	CALCARB	MC	ANEMAGEN	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Certain drugs require specific diagnoses for approval.
	MC	CALCH-MIX CAPSULE CAPS	MC	CALCET TABS	Please refer to OTC list.	
	MC	CALCIQUID SYRP	MC/DEL	CALCIUM 600-D TABS		
	MC	CALCITRATE/VITAMIN D TABS	MC	CALCIUM/VITAMIN D TABS		
	MC/DEL	CALCIUM	MC	CALTRATE 600 PLUS/VIT D TABS	Click here for the OTC List	
	MC/DEL	CALCIUM CARBONATE	MC	CALTRATE PLUS TABS		DDI: Fe salts will now be non-preferred and require prior authorization if it is currently being used in combination with either Prevacid, pantoprazole, Prilosec, or any currently non preferred PPI.
	MC/DEL	CALCIUM CITRATE TABS	MC	CHROMAGEN		
	MC/DEL	CALCIUM GLUCONATE TABS	MC	CITRACAL PLUS TABS		
	MC/DEL	CALCIUM LACTATE TABS	MC	CONTRIN CAPS		Please refer to OTC list.
	MC	CALCIUM/MAGNESIUM TABS	MC	FEOGEN FORTE CAPS		
	MC/DEL	CALCIUM/VITAMIN D TABS	MC	FEROCON CAPS		Preferred products that used to require diag codes still require diag codes unless indicated otherwise.
	MC	CALTRATE 600 TABS	MC/DEL	FERREX 150 CAPS		
	MC/DEL	CHEWABLE CALCIUM CHEW	MC	FERRO-SEQUELS TBCR		
	MC	CITRACAL TABS	MC	FE-TINIC CAPS		
	MC	CITRACAL + D TABS	MC	FE-TINIC 150 FORTE CAPS		
	MC	CITRUS CALCIUM TABS	MC/DEL	FLUOR-A-DAY SOLN		
	MC	CITRUS CALCIUM 1500 + D TABS	MC	HEMOCYTE TABS		
	MC	EFFERVESCENT POTASSIUM TBEF	MC/DEL	K-DUR TBCR		
	MC/DEL	FEOSTAT CHEW	MC	KLOR-CON PACK		
	MC	FERATAB TABS	MC	K-LYTE		
	MC/DEL	FER-GEN-SOL SOLN	MC/DEL	K-PHOS TABS NEUTRAL		
	MC	FER-IRON SOLN	MC	K-TABS TBCR		
	MC	FERRONATE TABS	MC	K-VESCENT PACK		
	MC/DEL	FERROUS SULFATE	MC	MICRO-K 10 MEG CPCR		
	MC/DEL	FLUOR-A-DAY CHEW	MC	NU-IRON 150 CAPS		
	MC	FLUORIDE CHEW	MC/DEL	OYSTER SHELL CALCIUM/VITA TABS		
	MC	FLUORIDE SODIUM CHEW	MC/DEL	POLY-IRON 150 CAPS		
	MC	FLUORITAB CHEW	MC/DEL	POLYSACCHARIDE IRON CAPS		
	MC	HM CALCIUM TABS	MC/DEL	POTASSIUM BICARB/CHLORIDE		
	MC	K+ POTASSIUM PACK	MC/DEL	POTASSIUM CHLORIDE 10MEQ CAPS		
	MC	KAON ELIX	MC/DEL	POTASSIUM CHLORIDE 8MEQ CAPS		
	MC	KAON-CL-10 TBCR	MC	TUMS 500 CHEW		
	MC	KCL 0.075%/D5W/NACL 0.2% SOLN	MC	VIActiv CHEW		
	MC	K-EFFERVESCENT TBEF				
	MC	KLOR-CON				
	MC	KLOTRIX TBCR				
	MC/DEL	K-PHOS TABS				
	MC/DEL	K-VESCENT TBEF				
	MC/DEL	LURIDE CHEW				
	MC/DEL	MAGNESIUM GLUCONATE TABS				
	MC/DEL	MAGNESIUM SULFATE SOLN				
	MC	MAGTABS				
	MC	MICRO-K 8 MEG				
	MC/DEL	OS-CAL TABS				
	MC/DEL	OS-CAL 500 + D TABS				
	MC/DEL	OYSCO				
	MC/DEL	OYST-CAL TABS				
	MC/DEL	OYST-CAL D TABS				
	MC/DEL	OYST-CAL/VITAMIN D TABS				

	MC/DEL		OYSTER CALCIUM TABS				
	MC/DEL		OYSTER SHELL				
	MC		PHARMA FLUR				
	MC/DEL		PHOSPHA 250 NEUTRAL TABS				
	MC		POTASSIUM BICARBONATE TBEF				
	MC/DEL		POTASSIUM CHLORIDE 8MEQ				
	MC		POTASSIUM EFFERVESCENT				
	MC/DEL		SELENIUM TABS				
	MC		SLOW-MAG TBCR				
	MC/DEL		SODIUM FLUORIDE				
	MC		V-R CALCIUM				
	MC		V-R OYSTER SHELL CALCIUM				
	MC		ZINC SULFATE CAPS				

PHENYLKETONURIA (PKU) TREATMENT AGENTS

PHENYLKETONURIA (PKU) TREATMENT AGENTS- INJECTABLES				MC		PALYNZIQ ¹	1. For the treatment of patients ≥ 18 years of age. Use PA Form# 20420	Palynziq is not to be used in combination with Kuvan
PHENYLKETONURIA (PKU) TREATMENT AGENTS- ORAL				MC		KUVAN	Use PA Form# 20420	

MISC. ELECTROLYTES/NUTRITIONALS

ELECTROLYTES/ NUTRITIONALS	MC		INTRALIPID EMUL ¹	MC		BOOST ¹	1. This list of nutritional is incomplete. All nutritional still require a PA except for the miscellaneous products listed as preferred. SGA form required for nutritional unless member has a G/I tube. 2. Formerly known as Omacor. Use PA Form# 20420 & SGA Form	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Certain drugs require specific diagnoses for approval. Medical foods are not to be authorized solely for the purpose of enhancing nutrient intake or managing body weight if the participant is able to eat conventional foods adequately. Medical foods may be approved if the member has a medical condition which precludes or restricts the use of conventional foods and necessitates the use of a formula. Concurrent Stimulant therapy is not an acceptable medical reason/condition for use of medical foods for enhancing nutrient intake or managing body weight. For children under the age of 5, MaineCare will not provide milk- or soy-based standard infant formulas. Regular formulas may be sought through your nearest WIC office. MaineCare will continue to cover medical food for all participants in MaineCare when medical necessity is met. Vascepa requires adjunct therapy for specific indication to reduce TG in those with severe hypertriglyceridemia (500mg per deciliter or more). Proper indication per lab values is required before approval
	MC		P.T.E.-5 SOLN ¹	MC		CASEC POWD ¹		
	MC		SEA-OMEGA CAPS ¹	MC		CHOICE DM LIQD ¹		
				MC		DELIVER 2.0 LIQD ¹		
				MC		DOJOLVI		
				MC		ENFAMIL ¹		
				MC		ENSURE ¹		
				MC		GLUCERNA ¹		
				MC		ISOCAL LIQD ¹		
				MC		KINDERCAL TF LIQD ¹		
				MC		KINDERCAL TF/FIBER LIQD ¹		
				MC		L-CARNITINE CAPS ¹		
				MC		LIPISORB LIQD ¹		
				MC		LOVAZA ^{1,2}		
				MC		MODULEN IBD POWD ¹		
				MC		NUTRAMIGEN POWD ¹		
				MC		NUTREN ¹		
				MC		NUTRITIONAL SUPPLEMENT LIQD ¹		
				MC		NUTRIVENT 1.5 LIQD ¹		
				MC		PEPTAMEN ¹		
			MC		PHENYLADE ¹			
			MC		PHENYL-FREE ¹			
			MC		PKU 3 POWD ¹			
			MC		PREGESTIMIL POWD ¹			
			MC		PROBALANCE LIQD ¹			
			MC		PROSOBEE ¹			

MONOCLONAL ANTIBODY				<p>MC</p> <p>MC/DEL</p> <p>MC</p> <p>MC/DEL</p> <p>MC</p> <p>MC</p> <p>MC/DEL</p> <p>MC</p> <p>MC</p>	<p>EMPAVELI</p> <p>ENSPRYNG</p> <p>FABHALTA</p> <p>GAMIFANT</p> <p>PIASKY</p> <p>SOLIRIS</p> <p>ULTOMIRIS</p> <p>UPLIZNA</p> <p>VOYDEYA</p>	<p>Use PA Form# 20420</p>	<p>A diagnosis of Paroxysmal nocturnal hemoglobinuria (PNH) using the HAM test or flow cytometry is required. In addition, the patient must show evidence of having received a meningitis vaccine at least 2 weeks prior to the start of therapy.</p> <p>Gamifant is recommended for the treatment of adult and pediatric (newborn and older) patients with primary hemophagocytic lymphohistiocytosis (HLH) with refractory, recurrent, or progressive disease or intolerance with conventional HLH therapy.</p> <p>Fabhalta and Ultomiris are recommended for the treatment of adults with paroxysmal nocturnal hemoglobinuria (PNH).</p>
IMMUNE GLOBULIN	<p>MC</p> <p>MC/DEL</p> <p>MC</p> <p>MC</p> <p>MC/DEL</p> <p>MC/DEL</p> <p>MC</p>		<p>BIVIGAM¹</p> <p>CUTAQUIG¹</p> <p>GAMUNEX-C</p> <p>GAMMAGARD S-D¹</p> <p>HIZENTRA¹</p> <p>PANZYGA¹</p> <p>PRIVIGEN¹</p>	<p>MC</p> <p>MC</p> <p>MC/DEL</p> <p>MC</p> <p>MC/DEL</p> <p>MC</p> <p>MC/DEL</p>	<p>ALYGLO</p> <p>ASCENIV²</p> <p>CUVITRU</p> <p>GAMMAPLEX INJ</p> <p>HYQVIA</p> <p>OCTAGAM INJ¹</p> <p>XEMBIFY</p>	<p>Use PA Form# 20420</p> <p>1. Clinical PA required</p> <p>2. For the treatment of patients between 12 to 17 years of age.</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>Alyglo is indicated for treatment of primary humoral immunodeficiency in adults ages 17 or older.</p> <p>Cutaquig is indicated as replacement therapy for primary humoral immunodeficiency (PI) in adults.</p> <p>Xembify is indicated for treatment of primary humoral immunodeficiency (PI) in patients 2 years of age and older.</p> <p>Asceniv indicated for the treatment of primary humoral immunodeficiency (PI) in adults and adolescents (12 to 17 years of age). PI includes but is not limited to the humoral immune defect in congenital agammaglobulinemia, common variable immunodeficiency (CVID), X-linked agammaglobulinemia, Wiskott-Aldrich syndrome, and severe combined immunodeficiencies (SCID).</p>
HEREDITARY ANGIOEDEMA	<p>MC</p> <p>MC</p> <p>MC</p> <p>MC/DEL</p> <p>MC/DEL</p> <p>MC</p> <p>MC/DEL</p>		<p>PROPHYLAXIS</p> <p>CINRYZE¹</p> <p>HAEGARDA¹</p> <p>ORLADEYO^{1,2}</p> <p>TAKHZYRO¹</p> <p>TREATMENT</p> <p>BERINERT KIT¹</p> <p>FIRAZYR¹</p> <p>RUCONEST VIAL¹</p>		<p>PROPHYHLAXIS</p> <p>TREATMENT</p> <p>KALBITOR VIAL</p>	<p>1. Clinical PA is required to establish diagnosis and medical necessity.</p> <p>2. For the treatment of patients ≥ 12 years of age.</p> <p>Use PA Form# 20420</p>	<p>Haegarda is indicated for routine prophylaxis to prevent Hereditary Angioedema (HAE) attacks in adolescent and adult patients</p>
HEMATOLOGICAL AGENTS-THROMBOPOIETIN RECEPTOR AGONISTS	<p>MC</p> <p>MC</p>		<p>PROMACTA¹</p> <p>NPLATE¹</p>	<p>MC</p> <p>MC/DEL</p> <p>MC/DEL</p>	<p>ALVAIZ</p> <p>DOPLETEL</p> <p>MULPLETA</p>	<p>Use PA Form# 20420</p> <p>1. Clinical PA required. Must see prior trial with insufficient response to corticosteroids and immunoglobulins.</p>	<p>Doptelet and Mulpelta: For the treatment of thrombocytopenia in adults with chronic liver disease who are scheduled to undergo a procedure.</p>
HEMATOLOGICAL AGENTS-IgAN				<p>MC/DEL</p> <p>MC</p>	<p>FILSPARI¹</p> <p>TARPEYO</p>	<p>Use PA Form# 20420</p> <p>1. PA required to confirm FDA approved indication.</p>	<p>All preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs (in step-order) will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists</p>
ANEMIA- BETA THALASSEMIA				<p>MC</p> <p>MC</p>	<p>REBLOZYL</p> <p>ZYNTEGLO</p>	<p>Use PA Form# 20420</p>	<p>Reblozyl is indicated for the treatment of anemia in adult patients with beta thalassemia who require regular red blood cell (RBC) transfusion. It is not indicated for use as a substitute for RBC transfusions in patients who require immediate correction of anemia.</p> <p>Zynteglo is indicated for the treatment of adult and pediatric patients with β-thalassemia who require regular red blood cell (RBC) transfusions.</p>
HEMATOLOGIC DISORDER TREATMENT				<p>MC/DEL</p>	<p>CABLIVI</p>	<p>Use PA Form# 20420</p>	<p>Tavalisse is recommended for patients at risk of bleeding when one line of therapy (steroids, IVIG, splenectomy) has failed.</p>

AGENTS				MC		TAVALISSE		Cablivi is recommended for the treatment of adult patients with acquired thrombotic thrombocytopenic purpura (aTTP), in combination with plasma exchange and immunosuppressive therapy.
COMPLEMENT RECEPTOR ANTAGONIST				MC		TAVNEOS	Use PA Form# 20420	
WHIM SYNDROME AGENTS				MC		XOLREMDI	Use PA Form#20420	Xolremdi: In patients 12 years of age and older with WHIM syndrome (warts, hypogammaglobulinemia, infections, and myelokathexis) to increase the number of circulating mature neutrophils and lymphocytes.
HEMOSTATIC								
HEMOSTATIC	MC/DEL MC		AMICAR AMINOCAPROIC ACID	MC MC		FIBRYGA RIASTAP	Use PA Form# 20420	Fibryga and Riastap are indicated for the treatment of acute bleeding episodes in adults and adolescents with congenital fibrinogen deficiency, including afibrinogenemia and hypofibrinogenemia. Fibryga® is not indicated for dysfibrinogenemia.
ACUTE HEPATIC PORPHYRIA (AHP)								
ACUTE HEPATIC PORPHYRIA (AHP)				MC		GIVLAARI	Use PA Form# 20420	Givlaari is indicated for the treatment of adults with acute hepatic porphyria (AHP).
PYRUVATE KINASE DEFICIENCY AGENTS								
PYRUVATE KINASE DEFICIENCY AGENTS				MC		PYRUKYND ¹	Use PA Form# 20420 1.PA required to confirm FDA approved indication.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s).
OP. - ANTIBIOTICS								
OP. - ANTIBIOTICS	MC MC MC/DEL MC MC/DEL MC MC MC/DEL MC/DEL		AK-SPORE OINT BACITRACIN/NEOMYCIN/POLYM BACITRACIN/POLYMYXIN B OINT CHLOROPTIC SOLN ERYTHROMYCIN OINT NEOSPORIN SOLN POLYSPORIN TRIMETHOPRIM SULFATE/POLY TOBRAMYCIN SULFATE SOLN	MC MC MC MC MC/DEL MC MC MC/DEL MC/DEL MC MC MC/DEL MC/DEL MC/DEL MC		AK-POLY-BAC OINT AK-SULF OINT AK-TOB SOLN AZASITE BACITRACIN OINT BLEPH-10 SOLN GATIFLOXACIN DROPS GENTAMICIN SULFATE GENTAK ILOTYCIN OINT LEVOFLOXACIN DROPS NEOMYCIN/BACI/POLYM OINT NEOMYCIN/POLYMYXIN/GRAMIC NEOSPORIN OINT OCUSULF-10 SOLN OCUTRICIN SOLN POLYTRIM DROPS SULFACETAMIDE SODIUM DROPS SULFACETAMIDE SODIUM OINT TERAK OINT	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
OP. - ANTI-PARASITIC								
OP. - ANTI-PARASITIC				MC		XDEM ¹	Use PA Form# 20420 1. For the treatment of Demodex blepharitis.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
OP. - RHO KINASE INHIBITORS								
OP. - RHO KINASE INHIBITORS	MC		RHOPRESSA					the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s)
OP. - QUINOLONES								
OP. - QUINOLONES	MC/DEL		CILOXAN OINT	MC/DEL		BESIVANCE	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s).

	MC MC/DEL	SULFACETAMIDE/PREDNISOLONE ZYLET SUSP	MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC		RETISERT IMPLANT SULFACET SOD/PRED SOLN TRIESENCE VIAL TOBRADEX ST TOBRAMYCIN SUSP DEXAMETHASONE VASOCIDIN SOLN VEXOL SUSP XIPERE		
OP. - PROSTAGLANDINS	MC/DEL MC MC/DEL MC/DEL	LATANOPROST SOL 0.005% LUMIGAN SOLN ROCKLATAN TRAVATAN-Z	MC/DEL MC/DEL MC MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	7 8 8 8 8 8 8 8 8 8	ZIOPTAN BIMATOPROST 0.03% DROPS DURYSTA IYUZEH RESCULA ^{1,2,3} TRAVATAN SOLN TRAVOPROST VYZULTA XALATAN SOLN ¹ XELPROS	1. All preferred must be tried. 2. Dosing limits apply, please see dosing consolidation list. 3. Clinical PA is required to establish diagnosis and medical necessity. Use PA Form# 20420	Preferred drugs must be tried and failed, in step-order, due to lack of efficacy (failure to reach target IOP reduction) or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
OP. - CYCLOPLEGICS	MC MC/DEL MC/DEL MC/DEL	AK-PENTOLATE SOLN ATROPINE SULFATE CYCLOPENTOLATE HCL SOLN ISOPTO HYOSCINE SOLN	MC/DEL MC MC/DEL MC		CYCLOGYL SOLN ISOPTO ATROPINE SOLN ISOPTO HOMATROPINE SOLN MUROCOLL-2 SOLN	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
OP. - MIOTICS - DIRECT ACTING	MC/DEL MC MC MC/DEL MC/DEL	ISOPTO CARBACHOL SOLN ISOPTO CARPINE SOLN PILOCAR SOLN PILOCARPINE HCL SOLN PILOPINE HS GEL				Use PA Form# 20420	
OP. - SELECTIVE ALPHA ADRENERGIC AGONISTS	MC MC MC MC/DEL MC/DEL	ALPHAGAN SOLN ALPHAGAN P 0.1% SOLN ALPHAGAN P 0.15% SOLN BRIMONIDINE DROPS 0.2 % SIMBRINZA	MC/DEL MC/DEL		BRIMONIDINE TARTRATE DROPS 0.15 % IOPIDINE SOLN	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
OP. - ANTI-ALLERGENICS	MC/DEL MC MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL	AZELASTINE HCL DROPS BEPREVE CROMOLYN SODIUM DROPS KETOTIFEN FUMARATE DROPS LASTACRAFT OLOPATADINE HCL 0.1% OLOPATADINE HCL 0.2% ZADITOR SOLN	MC MC/DEL MC/DEL MC MC/DEL MC MC/DEL MC/DEL	8 8 8 8 8 8 8 9	ALOCRIOL SOLN ALOMIDE SOLN EMADINE SOLN OPTICROM SOLN PATANOL SOLN ZERVIAE EPINASTINE	Use PA Form# 20420	All preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
OP. ANTI-ALLERGENICS- MASTCELL STABILIZER CLASS			MC/DEL		ALAMAST SOLN	Use PA Form# 20420	
OP. - CARBONIC ANHYDRASE INHIBITORS/COMBO	MC/DEL MC MC/DEL MC/DEL	AZOPT SUSP COMBIGAN DORZOLAMIDE DORZOLAMIDE/TIMOLOL	MC/DEL		COSOPT SOLN PF	Use PA Form# 20420	
OP. - NSAID'S	MC MC/DEL	ACULAR SOLN ¹ DUREZOL	MC MC	8 8	ACULAR LS ¹ BROMSITE ¹	1. Must fail all preferred products before non-preferred	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.

				<p>MC BREVOXYL</p> <p>MC CABTREO GEL⁵</p> <p>MC/DEL CLEOCIN-T²</p> <p>MC CLINAC BPO GEL</p> <p>MC CLINDAGEL GEL</p> <p>MC/DEL CLINDAMYCIN PHOSPHATE CREAM²</p> <p>MC CLINDETS SWAB</p> <p>MC DESQUAM-E GEL</p> <p>MC DESQUAM-X</p> <p>MC DIFFERIN 0.3% GEL</p> <p>MC DIFFERIN</p> <p>MC EMGEL GEL</p> <p>MC EPIDUO</p> <p>MC EPSOLAY</p> <p>MC ERYCETTE PADS</p> <p>MC FINEVIN CREA</p> <p>MC/DEL KLARON LOTN</p> <p>MC METROCREAM CREA²</p> <p>MC METROGEL GEL²</p> <p>MC METROLOTION LOTN²</p> <p>MC NEOBENZ MICRO</p> <p>MC/DEL NORITATE CREA</p> <p>MC ONEXTON⁵</p> <p>MC/DEL PLIXDA</p> <p>MC RETIN-A GEL²</p> <p>MC RETIN-A CREA²</p> <p>MC RETIN-A MICRO GEL</p> <p>MC RHOFAGE</p> <p>MC/DEL SODIUM SULFACET/SULF LOTN</p> <p>MC SOOLANTRA⁴</p> <p>MC/DEL TRIAZ</p> <p>MC TWYNEO</p> <p>MC VELTIN</p> <p>MC WINLEVI⁵</p> <p>MC ZENCIA WASH</p> <p>MC ZETACET</p> <p>MC/DEL ZIANA</p> <p>MC ZILXI</p>		<p>6. For the treatment of patients ≥ 9 years of age.</p> <p>Use PA Form# 10220 for Brand Name requests</p> <p>Use PA Form# 20420 for all other requests</p>	
TOPICAL- ATOPIC DERMATITIS	<p>MC/DEL</p> <p>MC/DEL</p> <p>MC/DEL</p> <p>MC</p> <p>MC/DEL</p> <p>MC</p> <p>MC</p>	<p>1</p> <p>1</p> <p>1</p> <p>2</p> <p>2</p> <p>2</p> <p>2</p>	<p>ELIDEL CREA</p> <p>PIMECROLIMUS CRE (AUTH GENERIC LABELER 68682 Oceanside Pharmaceuticals)</p> <p>PROTOPIC OINT</p> <p>TACROLIMUS OINT</p> <p>ADBRY^{2,4}</p> <p>DUPIXENT^{1,2,4}</p> <p>EUCRISA^{2,4}</p> <p>OPZELURA^{2,3,4}</p>	<p>MC/DEL</p> <p>MC</p> <p>MC</p>	<p>CIBINQO</p> <p>EBGLYSS^{2,3}</p> <p>NEMLUVIO</p>	<p>1. Avoid live vaccines if treated with Dupixent</p> <p>2. Clinical PA required.</p> <p>3. For the treatment of patients ≥ 12 years of age.</p> <p>4. Preferred after a trial and failure of TCI.</p> <p>Use PA Form# 20420</p>	<p>Preferred drugs also indicated for this condition, including topical steroids, cyclosporin AND calcineurin inhibitors must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Note: If unable to use TCIs then a trial of Eucrisa could be recommended before Dupixent.</p>
TOPICAL - ANTIBIOTIC	<p>MC</p> <p>MC/DEL</p> <p>MC/DEL</p>		<p>BACIT/NEOMYCIN/POLYM OINT</p> <p>BACITRACIN OINT</p> <p>GENTAMICIN SULFATE</p>	<p>MC/DEL</p> <p>MC/DEL</p> <p>MC/DEL</p>	<p>CENTANY OINT 2%¹</p> <p>MUPIROCIIN CREA¹</p> <p>TRIPLE ANTIBIOTIC OINT</p>	<p>1. Dosing limits apply, please see dosing consolidation list.</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p>

	MC/DEL MC/DEL MC MC		<p style="text-align: center;">VERY HIGH POTENCY</p> AUGMENTED BETA DIP BETAMETHASONE VALERATE DIFLORASONE DIACETATE HALOBETASOL	MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC	BETAMETHASONE DIPPROPIONATE DESOXIMETASONE 0.25% CREA/OINT <p style="text-align: center;">VERY HIGH POTENCY</p> BRYHALI LOTN CLOBETASOL PROPINATE LOTN CLOBETASOL PROPINATE SHAMPOO 0.05% CORMAX DIPROLENE IMPEKLO ¹ LEXETTE OLUX FOAM PSORCON PSORCON E SERNIVO SPRAY ² TEMOVATE ULTRAVATE		
TOPICAL - STEROID LOCAL ANESTHETICS				MC	EPIFOAM FOAM	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
TOPICAL - STEROID COMBINATIONS	MC		DERMA-SMOOTH-FS SCALP	MC	CARMOL-HC CREA	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
TOPICAL - EMOLLIENTS	MC/DEL MC MC		AMMONIUM LACTATE CREA ¹ AMMONIUM LACTATE LOTN 12% ¹ VITAMIN A & D MEDICATED OINT	MC MC MC MC MC	LAC-HYDRIN CREA ¹ LAC-HYDRIN LOTN 12% MEDERMA GEL MIMYX RENOVA CREA	Use PA Form# 20420 1. Dosing limits still apply. Please see dose consolidation list.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
TOPICAL - ENZYMES / KERATOLYTICS / UREA				MC MC MC	CARMOL 40 CREA SALEX CREA SALEX LOTN	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Ziox, Panafil and Papain products have been removed from the PDL due to FDA safety concerns regarding drugs containing Papain.
TOPICAL - GENITAL WARTS	MC/DEL		IMIQUIMOD 5% ²	MC/DEL MC/DEL MC/DEL MC MC MC	5 PODOFILOX SOLN 8 CONDYLOX ¹ 8 ALDARA ¹ 8 PICATO 8 VEREGEN ¹ 8 ZYCLARA ¹	Use PA Form# 20420 1. Non-preferred products must be used in specified order. 2. Dosing limits still apply. Please see dose consolidation list.	
TOPICAL - LOCAL ANESTHETICS	MC MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL		AF CAPSICUM OLEORESIN CREA CAPSAICIN CREA CAPSAICIN PATCH DIBUCAINE OINT ELA-MAX ¹ LIDOCAINE/PRILOCAINE CREA ¹ LIDOCAINE CREAM LIDOCAINE GEL LIDOCAINE PTCH 5%	MC/DEL MC/DEL MC MC MC MC MC/DEL	EMLA PADS EMLA CREA LIDA MANTLE CREA PONTOCAINE SOLN SYNERA ZOSTRIX ZTLIDO ²	Use PA Form# 20420 1. Lidocaine/Prilocaine cream and Ela-Max products require PA for users over 18 years of age. 2. Dosing limits still apply. Please see dose consolidation list. Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
TOPICAL - DEPIGMENTING AGENTS				MC MC MC	8 ALUSTRA CREA 8 EPIQUIN MICRO 8 GLYQUIN CREA		As per Medicaid Policy, cosmetic drugs are not covered. Non-cosmetic clinical applications will be considered by prior authorization on a case by case basis.

				MC/DEL	8	HYDROQUINONE CREA	Use PA Form# 20420	
				MC/DEL	8	HYDROQUINONE/SUNSCREENS		
				MC	8	SOLAQUIN FORTE CREA		
				MC	8	TRI-LUMA CREA		
				MC	9	ELDOQUIN		
TOPICAL - SCABICIDES AND PEDICULICIDES	MC/DEL MC MC/DEL MC/DEL MC		ACTICIN CREA LICE KILLING SHAM LICE TREATMENT CREME RINS LIQD PERMETHRIN LOTN NATROBA'	MC MC MC/DEL MC MC MC/DEL		ELIMITE CREA EURAX LINDANE MALATHION OVIDE LOTN SPINOSAD SUSP	Use PA Form# 20420 1. Dosing limits apply, please refer to dosage consolidation list.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
TOPICAL - WOUND / DECUBITUS CARE				MC MC MC		FILSUIVEZ REGRANEX GEL VYJUVEK	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Regranex will be approved for diabetic patients in good control (hgba1c <8), who are not smoking, with a stage III or IV WOCN AND NPUAP lower extremity diabetic ulcer and with an adequate blood supply (TcP O2 >30, ABI>0.7 or ASP> 70), and where the underlying cause has been corrected. The wound must be free of infection and have been previously treated with preferred standard therapies for at least 2 months. Maximum approval for 20 weeks. Vyjuvek: For the treatment of wounds in patients 6 months of age and older with dystrophic epidermolysis bullosa (DEB) with mutation(s) in the collagen type VII alpha 1 chain (COL7A1) gene. Filsuvez: The patient has a diagnosis of dystrophic or junctional epidermolysis bullosa. The patient is at least 6 months old and does not have current evidence or history of squamous cell carcinoma or active infection in the area requiring Filsuvez application. The patient has used standard wound care treatments, including silicone or foam dressings without wound resolution Accuzyme and Ethezyme products have been removed from the PDL due to FDA concerns regarding drugs containing Papain.
TOPICAL - ASTRINGENTS / PROTECTANTS	MC		XERAC AC SOLN	MC MC MC MC		LOWILA BAR MOISTURIN DRY SKIN CREA PROSHIELD PLUS SKIN PROTE CREA SURGILUBE GEL	Use PA Form# 20420 1. Dosing limits apply, please refer to dosage consolidation list.	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
TOPICAL - ANTISEPTICS / DISINFECTANTS	MC/DEL		POVIDONE-IODINE SOLN	MC MC MC MC		BETADINE OINT FORMALYDE-10 AERS IODOSORB LAZERFORMALYDE SOLUTION SOLN	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
MISCELLANEOUS EYE								
OP. - EYE	MC MC MC MC MC MC/DEL		AK-DILATE SOLN EYE WASH SOLN NAPHAZOLINE HCL SOLN PHENYLEPHRINE HCL SOLN PONTOCAINE SOLN SODIUM CHLORIDE	MC MC/DEL MC		LENS PLUS REWETTING DROPS MURO 128 NEO-SYNEPHRINE SOLN	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
MISCELLANEOUS EAR								
EAR	MC/DEL MC MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL MC MC MC MC/DEL		A/B OTIC SOLN ACETASOL SOLN ACETASOL HC SOLN ACETIC ACID ACETIC ACID/HYDROCORTISON ALLERGEN SOLN CARBAMIDE PEROXIDE 6.5% OTIC SOLN. CIPRO HC SUSP CORTISPORIN-TC SUSP CORTOMYCIN COLY-MYCIN-S SUSP EAR DROPS SOLN EAR DROPS RX SOLN EAR WAX REMOVAL DROPS	MC MC MC/DEL MC/DEL MC MC MC/DEL MC MC MC MC/DEL		ANTIBIOTIC EAR SOLN ANTIBIOTIC EAR SUSP CIPRODEX CIPROFLOXACIN HCL DEBROX SOLN DERMOTIC FLOXIN OTIPRIO OTOVEL	Use PA Form# 20420	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.

	MC MC/DEL MC/DEL		FLUOCINOLONE ACETONIDE OIL DROPS 0.01% NEOMYCIN/POLYMYXIN/HC OFLOXACIN 0.3% OTIC				
MOUTH ANTISEPTICS							
MOUTH ANTI-INFECTIVES	MC MC/DEL		NILSTAT SUSP NYSTATIN SUSP	MC MC		MYCELEX TROC ORAVIG	Use PA Form# 20420 Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
MOUTH ANTISEPTICS	MC/DEL MC/DEL MC MC		CHLORHEXIDINE GLUCONATE LIDOCAINE VISCOUS SOLN TRIAMCINOLONE IN ORABASE PSTE TRIAMCINOLONE ORADENT PSTE	MC MC MC MC		APHTHASOL PSTE ¹ PERIOGARD SOLN ¹ TRIAMCINOLONE ACETONIDE PSTE ¹	Use PA Form# 20420 1. Must fail all preferred products before non-preferred. Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
DENTAL PRODUCTS							
DENTAL PRODUCTS	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC		ETHEDENT CREA GEL-KAM CONC GEL-KAM GEL 0.4% PHOS FLUR SOLN SF 5000 PLUS CREA SF GEL STANNOUS FLUORIDE ORAL RI CONC	MC/0MC MC/DEL MC/DEL MC		APF GEL DENTAGEL GEL PHOS-FLUR GEL THERA-FLUR-N GEL	Use PA Form# 20420 Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
ARTIFICIAL SALIVA/STIMULANTS							
ARTIFICIAL SALIVA/STIMULANTS	MC		SALIVA SUBSTITUTE SOLN	MC MC MC		EVOXAC CAPS RADIACARE SOLR SALAGEN TABS	Use PA Form# 20420 Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
MISCELLANEOUS ANORECTAL							
ANORECTAL - MISC.	MC MC MC/DEL MC/DEL MC/DEL		CORTENEMA ENEM ELA-MAX 5 CREA HYDROCORTISONE ENEM PROCTOSOL HC CREA PROCTOZONE-HC CREA	MC/DEL MC/DEL MC/DEL MC/DEL MC		ANUSOL-HC CREA CORTIFOAM FOAM PROCTOFOAM HC FOAM PROCTO-KIT CREA 2.5% RECTIV OINT	Use PA Form# 20420
T-CELL ACTIVATION INHIBITOR							
PSORIASIS BIOLOGICALS	MC MC MC MC MC/DEL MC		ADALIMUMAB-FKJP ENBREL ^{1,5} ENBREL SURECLICK ¹ HUMIRA ^{1,5} OTEZLA SIMLANDI SKYRIZI ⁶ TALTZ ²	MC MC/DEL MC MC/DEL MC/DEL MC/DEL MC MC/DEL MC MC MC MC MC		AMJEVITA BIMZELX ³ COSENTYX ⁴ CYLTEZO HADLIMA HULIO HYRIMOZ IDACIO ILUMYA ³ SOTYKTU SPEVIGO SILIQ STELARA TREMIFYA YUFLYMA YUSIMRY	1. Dosing limits apply, please refer to dosage consolidation list. 2. Clinical PA required and will be preferred for the indication of plaque psoriasis, psoriatic arthritis and ankylosing spondylitis. 3. For the treatment of adults with moderate-to-severe plaque psoriasis who are candidates for systemic therapy or phototherapy. 4. Please see criteria section 5. Will not require a PA if at least one systemic drug such as methotrexate, cyclosporine, methoxsalen Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Cosentyx approvals for 300mg dose(s) must use "300DOSE" package (containing 2 x 150mg pens or syringes). It is recommended to assess for TB infection prior to starting treatment with Taltz®. Stelara will require using preferred trial of Skyrizi if unable please provide clinical rational as why inappropriate.

or acitretin is in members drug profile.

6. Clinical PA required and will be preferred for the indication of plaque psoriasis, psoriatic arthritis, Crohn's disease and ulcerative colitis.

[Use PA Form# 20910](#)

ALTERNATIVE MEDICINES

ALTERNATIVE MEDICINES	MC MC		DIMETHYL SULFOXIDE SOLN MELATONIN	MC/DEL		CO-ENZYME Q-10	Use PA Form# 20420	Will only be approved for specific conditions supported by at least two double-blinded, placebo-controlled randomized trials that are not contradicted by other studies of similar quality.
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CHELATING AGENTS

CHELATING AGENTS	MC/DEL		CUPRIMINE CAPS	MC MC MC/DEL MC MC/DEL		CLOVIQUE DEPEN TITRATABS TABS EXJADE ¹ SYPRINE TRIENTINE CAPS	Use PA Form# 20420 1. FDA indication of treatment of chronic iron overload due to blood transfusions in members 2	Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved (in step order), unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists. Clovique® should be used when continued treatment with penicillamine is no longer possible because of intolerable or life endangering side effects.
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ANTILEPROTIC

ANTILEPROTIC				MC		THALOMID CAPS ¹	1. All PA requests for 150mg dosing will require use of Thalomid 100mg and 50mg capsules. Use PA Form# 20420	Approved for indications of leprosy, treatment-resistant multiple myeloma and AIDS.
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ANTINEOPLASTIC AGENTS

ANTINEOPLASTIC AGENTS - ANTIADNDROGENS	MC/DEL		BICALUTAMIDE	MC/DEL		CASODEX	Use PA Form# 20420	
ANTINEOPLASTIC AGENTS- LHRH ANALOGS	MC/DEL MC/DEL MC/DEL MC/DEL		LUPRON DEPOTSYPHNGEKIT ¹ LUPRON DEPOT- PED KIT ¹ (1-month) LUPRON DEPOT-PED SYRINGEKIT (3-month) TRIPTODUR VIAL	MC/DEL MC/DEL MC/DEL MC/DEL MC		LUPRON DEPOT SYRINGEKIT FIRMAGON ² SUPPRELIN LA (IMPLANT) KIT TRELSTAR VANTAS ²	1. Dosing limits apply, please refer to dosage consolidation list. 2. PA required to confirm FDA approved indication. Use PA Form# 20420	
ANTINEOPLASTIC AGENTS - TYROSINE KINASE INHIBITORS				MC MC/DEL MC		SPRYCEL ¹ TYKERB ² GLEEVEC ¹	Use PA Form# 20420 1. Verification of diagnosis is required. 2. PA required to confirm FDA approved indication and to monitor for potential drug-drug interactions.	
ANTINEOPLASTICS-MISCELLANEOUS	MC		AMIFOSTINE	MC		DOCEFREZ	Use PA Form# 20420	

MC	FOTIVDA
MC	FRUZAQLA
MC	GAVRETO
MC/DEL	GILOTRIF ^{4,5}
MC/DEL	IBRANCE
MC	ICLUSIG ³
MC/DEL	IDHIFA ³
MC	IMBRUVICA
MC	IMDELLTRA
MC/DEL	IMFINZI
MC/DEL	IMJUDO
MC	IMKELDI
MC	IMLYGIC
MC/DEL	INLYTA
MC/DEL	INREBIC
MC	INQOVI
MC	ITOVEBI
MC	IWILFIN
MC	JAKAFI
MC	JAYPIRCA ^{1,2}
MC	JEMPERLI
MC/DEL	KEYTRUDA ¹
MC	KIMMTRAK
MC	KISQALI ¹
MC/DEL	KOSELUGO
MC	KRAZATI ³
MC	KYMRIAH ^{3,9}
MC	KYPROLIS ¹
MC	LARTRUVO ¹
MC	LAZCLUZE
MC	LENVIMA
MC/DEL	LIBTAYO ¹
MC	LONSURF
MC/DEL	LORBRENA
MC	LOQTORZI
MC	LUMAKRAS
MC/DEL	LUMOXITI ¹
MC	LUNSUMIO ¹
MC	LYNPARZA ¹
MC	LYTGOBI
MC	NEXAVAR ¹
MC	NERLYNX ³
MC	NINLARO(PO)
MC/DEL	NUBEQA
MC	MARGENZA
MC/DEL	MEKINIST ^{3,4}
MC/DEL	MEKTOVI ¹
MC	MONJUVI
MC/DEL	MYLOTARG ³
MC/DEL	MVASI
MC	ODOMZO ^{1,2,5}
MC	OGSIVEO
MC	OJEMDA
MC	OJJAARA
MC	OMISIRGE

MC	ONUREG
MC/DEL	OPDIVO ³
MC	OPDIVO QVANTIG
MC	OPDUALAG
MC	ORGOVYX
MC	ORSERDU ^{2,3}
MC	PADCEV
MC	PEMAZYRE
MC	PEPAXTO
MC	PHESGO
MC/DEL	PIQRAY
MC	POLIVY
MC	POMALYST
MC	PORTRAZZA ³
MC	QINLOCK
MC	RETEVMO
MC	REZLIDHIA
MC/DEL	ROZLYTREK
MC	RUBRACA
MC	RITUXAN
MC	RYBREVANT
MC	RYDAPT
MC	RYLAZE
MC	RYTELO
MC/DEL	SARCLISA
MC	SCEMBLIX ¹
MC/DEL	STIVARGA
MC/DEL	SUTENT ^{1,2}
MC/DEL	SYLATRON
MC	TABRECTA
MC	TALVEY
MC/DEL	TAFINLAR ^{3,4,5,6}
MC	TAZVERIK
MC/DEL	TALZENNA ¹
MC/DEL	TAGRISO
MC	TECARTUS
MC	TECELRA
MC	TECENTRIQ ¹
MC	TECENTRIQ HYBREZA
MC	TEPMETKO
MC	TEVIMBRA
MC/DEL	TIBSOVO ¹
MC	TIVDAK
MC	TRODELVY
MC	TRUSELTIQ
MC/DEL	TRUXIMA
MC/DEL	TRUQAP
MC	TUKYSA
MC	UKONIQ
MC/DEL	VANFLYTA
MC	VEGZELMA
MC	VENCLEXTA ³
MC	VERZENIO ³
MC/DEL	VITRAKVI
MC/DEL	VIZIMPRO ¹

			MC	VONJO	
			MC	VORANIGO	
			MC/DEL	VYLOY	
			MC/DEL	WELIREG	
			MC/DEL	XALKORI	
			MC/DEL	XPOVIO	
			MC/DEL	XOSPATA	
			MC/DEL	XTANDI	
			MC/DEL	YERVOY	
			MC	YESCARTA ³	
			MC/DEL	ZALTRAP	
			MC	ZEJULA ¹	
			MC/DEL	ZELBORAF	
			MC	ZEPZELCA	
			MC	ZIHERA	
			MC	ZYDELIG	
			MC/DEL	ZYKADIA	
			MC	ZYNLONTA	
			MC	ZYNYZ ¹	
			MC	ZYTIGA	

IMMUNOSUPPRESSANTS

IMMUNOSUPPRESSANTS	MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		CYCLOSPORINE MODIFIED GENGRAF CAPS MYCOPHENOLATE MYFORTIC NEORAL SOL RAPAMUNE SANDIMMUNE TACROLIMUS CAPS	MC/DEL MC/DEL MC MC MC/DEL MC MC/DEL	CELLCEPT CYCLOSPORINE CAPS CYCLOSPORINE SOL. MODIFIED ENVARBUS XR MYHIBBIN ² NEORAL CAP PROGRAF CAPS REZUROCK ¹ ZORTRESS	<p>1. For the treatment of adult and pediatric patients 12 years and older with chronic graft-versus-host disease (chronic GVHD) after failure of at least 2 prior lines of systemic therapy</p> <p>2. Clinical PA is required.</p> <p>Use PA Form# 20420</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p> <p>DDI: Cyclosporine will now be non-preferred and require prior authorization if it is currently being used in combination with either Lipitor (doses greater than 20mg/day), Crestor, or lovastatin (doses greater than 20mg).</p> <p>DDI: Cyclosporine will require prior authorization when used with Livalo.</p> <p>Myhibbin: For the prophylaxis of organ rejection, in adult and pediatric recipients 3 months of age and older of allogeneic kidney, heart, or liver transplants, in combination with other immunosuppressants.</p> <p>DDI: All preferred immunosuppressants will require clinical PA for patients over 60 that are currently on fluoroquinolone therapy.</p>
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IMMUNOSUPPRESSANTS- Misc.				MC	HYFTOR ¹²	<p>1. For the treatment of patients ≥ 6 years of age.</p> <p>2. Clinical PA required for appropriate diagnosis and clinical parameters.</p> <p>Use PA Form# 20420</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p>
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PURINE ANALOG

PURINE ANALOG	MC MC/DEL		AZASAN TABS AZATHIOPRINE TABS	MC/DEL	IMURAN TABS	<p>Use PA Form# 20420</p>	<p>Preferred drugs must be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.</p>
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K REMOVING RESINS

K REMOVING RESINS	MC/DEL MC/DEL		LOKELMA SODIUM POLYSTYRENE SULFON	MC/DEL MC/DEL MC	SPS SUSP SPS 30GM/120ML ENEMA SUSP VELTASSA	<p>Use PA Form# 20420</p>	
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