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 Ctr	1 F									
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		• • • •				BASIC" Covered Drugs are bolded with t					
General Criteria for all PDL categories - Fo	or more informa	ation or help u	using the PDL, providers may call 1-888-445-0497	7; members sh	ould call	1-866-796-2463. To access PDL and PA materials via the	internet: www.mainecarepdl.c	org			
A: Preferred Drugs- Unless otherwise spe	cified, preferre	d drugs are a	vailable without prior authorization. Step order i	may apply for	preferred	drugs in some drug categories as indicated on the PDL.	(See item "D" below for expla	anation of step order.)			
B: Requests for Non-preferred Drugs- Pre the preferred drug(s) exists.	ferred drugs m	nust be tried a	nd failed due to lack of efficacy or intolerable sic	de effects befo	re non-pr	eferred drugs will be approved, unless an acceptable cli	ical exception is offered on th	he 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			
C: Adequate Drug Trials- 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 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: Step Order- When numbers appear in	the "step order	r" column, it n	neans drugs in this category must be used in the	e order specif	ed, with th	e lower numbers having preference over the higher nur	bers. 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 categories will require prior authorization				Preferred bra	ıd drugs v	vill no longer be preferred in any PDL drug category who	re preferred generic drugs are	e 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			
								d 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 per role of the FDA. Physicians should submit their reports of generic inequivalence directly to the FDA via the MEDWATCH.			
G: PA requests for non- FDA Approved In randomized clinical studies establishing I			made on a case-by-case basis until the DUR con	nmittee is able	to review	the evidence and make a recommendation. Interim app	rovals and DUR recommendat	tions 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			
H: <u>Dose Consolidation Requirements</u> - Sol	me drugs may	also be affect	ed by dose consolidation requirements. Please	see Dose Cor	solidatior	List and/or Splitting Tables provided in the PDL.					
I. <u>Trials from Multiple Drug Classes</u> - Tria	l/failure/intoler	ance to prefer	rred agents from multiple classes within the sam	e category or	other cate	gories of drugs may be required prior to the approval of	non-preferred agents (e.g., C	ymbalta, Zofran, Elidel and others).			
J. <u>Drug-specific PA Forms</u> - Drug-specific	PA forms con	tain medical n	necessity documentation requirements and/or cri	iteria that may	not be re	peated in the PDL. Drug-specific PA forms may be obtain	ned on the web at <u>www.maine</u>	ecarepdl.org.			
						nption from prior authorization requirement for certain of t will be required to do so, and criteria for approval of th		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 net.			
L. Drug Drug Interactions (DDI) The DHP	Committee has	a implemente	d now drug drug interaction adite requiring prior	r authorization	Soveral	drug drug combinations and DDL drug agterories are a	footed by now BA requiremen	ts. These will be indicated in the PDL with DDI notation. Please see the DDI document provided in the PDL.			
L. <u>brug-brug interactions (bbij</u> - The box	Committee nas	s implemente	a new drug-drug interaction earls requiring prior	autilorizatioi	. Several	drug-drug combinations and PDL drug categories are a	lected by new PA requirement	is. These will be indicated in the FDL with DDI hotation. Flease see the DDI document provided in the FDL.			
			ASSORTED ANT				_				
BETA-LACTAMS / CLAVULANATE COMBO'S	MC/DEL MC/DEL MC/DEL	A	MOXICILLIN MOXICILLIN/POTASSIUM CLA CHEW MOXICILLIN/POTASSIUM CLA SUSR MOXICILLIN/POTASSIUM CLA TABS	MC/DEL MC/DEL		AUGMENTIN <sup>3</sup> AUGMENTIN XR TB12 <sup>4</sup>	Chewable 125mg &     250mg and Solution     125mg/5ml and 250mg/5ml     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.			
	MC/DEL MC MC/DEL MC	E	MPICILLIN BICILLIN L-A SUSP DICLOXACILLIN SODIUM CAPS DXACILLIN SODIUM SOLR				Use preferred generic amoxicillin/clavulanate potassium alternatives.	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.			
	MC/DEL MC MC	F	PENICILLIN SOUR SOUR  TIMENTIN SOLR  JINASYN SOLR				Use PA Form# 20420				
	MC/DEL		OSYN								
CEPHALOSPORINS	MC/DEL		CEFADROXIL HEMIHYDRATE	MC		CEDAX	Both brand and generic	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/DEL		CEFAZOLIN SODIUM SOLR	MC/DEL		CEFACLOR <sup>1</sup>	are clinically non-preferred.	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			
	MC/DEL	C	CEFDINIR	MC/DEL		CEFADROXIL MONOHYDRATE TABS		preferred drug(s) exists.			
	MC/DEL	C	CEFEPIME	MC/DEL		CEFIXIME SUS	2. Dosing limits apply,				
	MC/DEL	C	CEFPODOXIME	MC/DEL		CEPHALEXIN TABS	please see Dosage				
	MC/DEL	c	CEFPODOXIME PROXETIL SUS	MC		CEPHALEXIN 750MG CAPS	Consolidation List.				
	MC/DEL	d	CEFPODOXIME PROXETIL TAB	MC/DEL		CEFTIN	3. Approvals will only be				
l	MC/DEL	C	CEFIXIME 400MG <sup>2</sup> CAP	MC		DAXBIA	considered for patients 18	I			
						Poo	o 1 of 77				

	MC/DEL	CEFPROZIL	MC	FETROJA <sup>3</sup>	have limited or no alternative	DDI: Vantin 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
	MC/DEL	CEPHALEXIN 250MG & 500MG CAPS	MC/DEL	FORTAZ	treatment options for the	PPI.
	MC	CEFTAZIDIME 6MG	MC/DEL	FORTAZ SOLN	treatment of complicated	
	MC/DEL	CEFTIN SUSP	MC	KEFLEX CAPS	urinary tract infections	
	MC/DEL	CEFTRIAXONE	MC	OMNICEF	(cUTIs)	As outlined in the US CDC Guidance on the Use of Expedited Partner Therapy (EPT) in the Treatment of Gonorrhea, MaineCare will cover a single 800 mg dose of cefixime for the
	MC/DEL	CEFUROXIME AXETIL TABS	MC/DEL	ROCEPHIN		treatment of gonorrhea as part of EPT.
	MC/DEL	CEPHALEXIN MONOHYDRATE	MC/DEL	SUPRAX <sup>2</sup>		1
	MC	FORTAZ SOLR	MC	TAZICEF SOLR		
	MC/DEL	SUPRAX CHEWABLE	MC/DEL	TEFLARO		
	MC	TAZICEF 6GM				
					Use PA Form# 20420	
MACROLIDES / ERYTHROMYCIN'S	MC/DEL	AZITHROMYCIN TABS	MC/DEL	AZITHROMYCIN POW	1. 7- Day supply per month	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/DEL	AZITHROMYCIN SUSP	MC/DEL	CLARITHROMYCIN SUSP	without PA.	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
	MC	E.E.S.	MC/DEL	CLARITHROMYCIN TABS		preferred drug(s) exists.
	MC	ERYPED 200 SUSR	MC	DIFICID		
	MC	ERYPED 400 SUSR	MC	PCE TBEC		DDI: Preferred erythromycin will now be non-preferred and require prior authorization if it is currently being used in combination with either Carbamazepine, Enablex 15mg or Vesicare
	MC	ERY-TAB TBEC	MC/DEL	ZITHROMAX TABS		10mg. Any non preferred formulation of erythromycin will require prior authorization and the member's drug profile will also be monitored for concurrent use with either Carbamazepine,
	MC	ERYTHROCIN STEARATE TABS	MC/DEL		H DA F# 00400	Enablex 15mg or Vesicare 10mg.
				ZITHROMAX 1GM PAK	Use PA Form# 20420	_ · · · · · · · · · · · · · · · · · · ·
	MC/DEL	ERYTHROMYCIN	MC/DEL	ZITHROMAX TRI-PAK		
			MC/DEL	ZITHROMAX SUSP		DDI: Preferred clarithromycin formulations (clarithromycin tablets) will now be non-preferred and require prior authorization if they are currently being used in combination with either
			MC/DEL	ZMAX		Carbamazepine, Onglyza 5mg, Enablex 15mg or Vesicare 10mg. Any non preferred formulation of clarithromycin will require prior authorization and the member's drug profile will also be
			MC/DEL	ZINPLAVA		monitored for concurrent use with either Carbamazepine, Onglyza 5mg, Enablex 15mg or Vesicare 10mg.
						Zinplava® will be non-preferred and require clinical prior authorization to verify it is prescribed or consulted by Gl or ID specialist, diagnosis, and concurrent use of an antibacterial agent
						as well as limiting its use to those who have recurrent C. diff disease that has recurred despite use of guideline recommended vancomycin taper or for whom this would be contraindicated.
						Contramidicated.
			L	DEGLOUNGHU TARO		
TETRACYCLINES	MC/DEL	DOXYCYCLINE MONOHYDRATE 100mg & 50mg	MC	DECLOMYCIN 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
		CAPS	l l		Use PA Form# 20420	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	MINOCYCLINE HCL CAPS	MC/DEL	DORYX CPEP		prototros staglej onicis.
	MC/DEL	TETRACYCLINE HCL CAPS	MC/DEL	DOXYCYCLINE HYCLATE	1. For the treatment of	1
			MC/DEL	DOXYCYCLINE MONOHYDRATE 150mg & 75mg CAPS	patients ≥ 8 years of age.	1
						1
			MC/DEL	DYNACIN CAPS	2. For the treatment of	1
			MC/DEL	MINOLIRA ER	patients ≥ 9 years of age.	1
			MC/DEL	NUZYRA <sup>1</sup>		1
			MC	ORACEA		1
			MC/DEL	PERIOSTAT		1
						1
			MC	SEYSARA <sup>2</sup>		1
			MC/DEL	SOLODYN ER		1
			MC	XIMINO		
FLUOROQUINOLONES	MC/DEL	CIPROFLOXACIN	MC	AVELOX SOLN		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/DEL	LEVOFLOXACIN	MC	AVELOX ABC PACK TABS	Use PA Form# 20420	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
	MC/DEL	OFLOXACIN	MC	BAXDELA	1. Dosing limits apply, see	preferred drug(s) exists.
			MC	CIPRO	Dosage Consolidation List.	DDI: Preferred ofloxacin will now be non-preferred and require prior authorization if they are currently being used in combination with amiodarone.
			MC	FACTIVE		DDI: Preferred levofloxacin will now be non-preferred and require prior authorization if they are currently being used in combination with amiodarone.
			MC	LEVAQUIN TABS SOLN/INJ		DDI: Preferred Avelox will now be non-preferred and require prior authorization if they are currently being used in combination with amiodarone.
			MC	LEVAQUIN TABS <sup>1</sup>		DDI: All preferred fluoroquinolones will require clinical PA for patients over 60 that are currently on immunosuppressants or steroid therapy.
			MC	NOROXIN TABS		
			MC	PROQUIN XR		DDI: Factive is non-preferred but with any prior authorization requests, the member's drug profile will also be monitored for concurrent use with amiodarone
			MC	PROQUIN XR		DDI: Factive is non-preferred but with any prior authorization requests, the member's drug profile will also be monitored for concurrent use with amiodarone.
AMINO GI YCOSIDES	MC	GENTAMICIN			Lico DA Essenti 20420	
AMINO GLYCOSIDES	MC	GENTAMICIN	MC/DEL	ARIKAYCE <sup>1,4</sup>	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
AMINO GLYCOSIDES	MC	KITABIS PAK	MC/DEL MC	ARIKAYCE <sup>1,2</sup> BETHKIS <sup>1</sup>	1. Clinical PA to verify	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
AMINO GLYCOSIDES	MC MC/DEL	KITABIS PAK NEOMYCIN SULFATE TABS	MC/DEL MC MC/DEL	ARIKAYCE <sup>1,2</sup> BETHKIS <sup>1</sup> TOBI PODHALER <sup>1</sup>	Clinical PA to verify appropriate diag	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.
AMINO GLYCOSIDES	MC	KITABIS PAK	MC/DEL MC MC/DEL MC	ARIKAYCE <sup>1,2</sup> BETHKIS <sup>1</sup>	1. Clinical PA to verify	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
AMINO GLYCOSIDES	MC MC/DEL	KITABIS PAK NEOMYCIN SULFATE TABS	MC/DEL MC MC/DEL	ARIKAYCE <sup>1,2</sup> BETHKIS <sup>1</sup> TOBI PODHALER <sup>1</sup>	Clinical PA to verify appropriate diag	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.

	I I	I	MC/DEL	ZEMDRI <sup>2</sup>		Current users of Tobi Nebu and Tobramycin Soln will be allowed a grace period until 10/1/15 to transition to preferred Kitabis.
						Address will associate disciplinate and the same of forces in adults who have limited as an alternative tradered and
						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-	MC/DEL	ETHAMBUTOL HCL TABS	MC/DEL	MYCOBUTIN 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
TUBERCULOSIS	MC/DEL	MYAMBUTOL TABS	MC/DEL	PRETOMANID		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	RIFABUTIN CAPS	MC	RIFADIN CAPS		profession unugla) exista.
	MC/DEL	RIFAMPIN				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	DARAPRIM TABS	MC	ARALEN 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
	MC	KRINTAFEL <sup>2</sup>	MC/DEL	CHLOROQUINE PHOSPHATE TABS <sup>3</sup>	1. Ingredients available as	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
	MC/DEL	MEFLOQUINE HCL TABS	MC/DEL	HYDROXYCHLOROQUINE TABS <sup>3</sup>	preferred without PA.	preferred drug(s) exists.
	MC/DEL	QUININE SULFATE	MC	ISONARIF <sup>1</sup>	<ol><li>Krintafel is preferred for ≥</li></ol>	
			MC	MALARONE TABS	16 years of age.	DDI: Avoid coadministration of Krintafel® with Organic Cation Transporter 2 (OCT2) and Multidrug and Toxin Extrusion (MATE) substrates (e.g. dofetilide, metformin).
			MC/DEL	PLAQUENIL TABS	Established users will be grandfathered	
ANTHELMINTICS	MC/DEL	ALBENDAZOLE	MC	ALBENZA 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
	MC/DEL	PRAZIQUANTEL TAB	MC	EMVERM	USE FA FUITH 20420	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
	MC/DEL MC/DEL	STROMECTOL TABS	MC/DEL	BILTRICIDE TABS		preferred drug(s) exists.
	IIIO/DEE	OTTOMESTOE TABO	IIIQ/DEL	SETTIONE TABO		
ANTIBIOTICS - MISC.	MC	AZACTAM SOLR	MC	AEMCOLO	1. 375mg caps and 750mg	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	COLY-MYCIN-M SOLR	MC	COLISTIMETHATE SODIUM SOLR	tabs are non-preferred.	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
	MC	COLISTIMETHATE SODIUM SOLR	MC	CAYSTON <sup>3</sup>	Please use available	preferred drug(s) exists.
	MC/DEL	FIRVANQ <sup>4</sup>	MC/DEL	FLAGYL CAPS	preferred strengths(250mg & 500mg tabs) to obtain	For macrolide resistant infections when quinolones inappropriate
	MC	FUROXONE TABS	MC/DEL	FLAGYL TABS	required dose without PA.	
	MC/DEL	METRONIDAZOLE <sup>1</sup>	MC/DEL	FLAGYL ER TBCR		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 Vesicare 10mg or
	MC	PENTAMIDINE ISETHIONATE SOLR	MC/DEL	КЕТЕК		carbamazepine.
	MC/DEL	SOLOSEC	MC	LIKMEZ		
	MC/DEL	TRIMETHOPRIM TABS	MC/DEL	METRONIDAZOLE 375MG CAPS <sup>1</sup>	which are preferred to obtain	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 acyston therapy). A bronchodilator should be used before administration of Cayston.
	MC/DEL	VANCOMYCIN 5GM INJ.	MC/DEL	METRONIDAZOLE 750MG TABS <sup>1</sup>	dose without PA.	
	MC/DEL	VANCOMYCIN CAPS	MC	NEBUPENT SOLR		
	MC	XIFAXAN 200mg	MC	REBYOTA <sup>5</sup>		Xenleta will be considered for the treatment of adults with community-acquired bacterial pneumonia (CABP) caused by the following susceptible microorganisms: Streptococcus
			MC	TINDAMAX	3. Clinical PA is required to	pneumoniae, Staphylococcus aureus (methicillin-susceptible isolates), Hemophilus influenzae, Legionella pneumophila, Mycoplasma pneumoniae, and Chlamydophila pneumoniae.
			MC/DEL	VANCOMYCIN 10GM INJ. <sup>2</sup>	establish CF diagnosis and	
			MC/DEL	XENLETA	medical necessity. Prior trail and failure of preferred Tobi	
			MC	XIFAXAN	before approval will be	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).
			MC	VOWST <sup>5</sup>	granted.	
					Quantity limit of one per     150ml bottle.	Likmez: patient has a medical necessity for a non-solid oral dosage form.
					5. For the treatment of patients 18 years of age and older.	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.
					Use PA Form# 20420	
CARBAPENEMS	1 1		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 MC/DEL	CLEOCIN SOLN CLEOCIN SUSR CLINDAMYCIN HCL 150CAPS DAPSONE TABS LINEZOLID 600mg TABS <sup>2</sup>	MC/DEL MC/DEL MC MC/DEL MC/DEL	8 8 8 9	CLEOCIN CAPS  CLINDAMYCIN HCL 300CAPS <sup>1</sup> SIVEXTRO VIBATIV ZYVOX SUSR ZYVOX TABS	Use multiple 150's for Clindamycin instead of 300's.      Quantity limit of 14 days supply within a 60day period.	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.
						Use PA Form# 30820 for Zyvox & Vibativ Use PA Form# 20420 for all others	
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 <sup>1</sup>	1. I of 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.
ANTIPROTOZOALS	MC/DEL MC/DEL	BENZNIDAZOLE <sup>2</sup> LAMPIT <sup>2</sup>	МС		ALINIA <sup>†</sup>	Alina is preferred for children less than 12 years of age.     Clinical PA required for appropriate diagnosis.	Benznidazole is indicated for pediatric patients 2 to 12 years of age for the treatment of Chagas disease (American trypanosomiasis) caused by Trypanosoma cruzi.
						Use PA Form# 20420_	
ANTIFUNGALS - ASSORTED	MC MC/DEL	ANCOBON CAPS FLUCONAZOLE <sup>1</sup>	MC/DEL MC/DEL	6	LAMISIL TABS <sup>4</sup> ITRACONAZOLE	See quantity limit table.  Non-preferred products must be used in specified step order.	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.
	MC/DEL MC/DEL MC/DEL MC/DEL	KETOCONAZOLE TABS <sup>7</sup> NYSTATIN TERBINAFINE TABS <sup>4</sup> VORICONAZOLE TABS	MC MC/DEL MC/DEL MC MC MC MC MC MC MC MC/DEL MC/DEL MC/DEL	8 8 8 8 8 8 8	BREXAFEMME CRESEMBA <sup>9</sup> GRIFULVIN V TABS GRISEOFULVIN SUSP GRISEOFULVIN ULTRAMICROSI TABS GRIS-PEG TABS REZZAYO <sup>9</sup> SPORANOX SOLN <sup>2</sup> SPORANOX PULSEPAK CAPS <sup>3</sup> SPORANOX CAPS <sup>3</sup>	QL1/every 7-day period (150mg only).     Sporanox QL	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
			MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL MC	8 8 8 8 8 8	DIFLUCAN ERAXIS INJ <sup>6</sup> GRIFULVIN SUSP ONMEL NOXAFIL <sup>5</sup> TOLSURA VFEND TABS VIVJOA	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.	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. 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. Diflucan is non-preferred but with any prior authorization if being used in combination with Plavix or Warfarin.
						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.	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 ≥ 18 years of age	
		l l	ı			U DA Farre# 40420	
		ANTI - VIRALS				Use PA Form# 10120	
ANTIRETROVIRALS	MC/DEL	ANTI - VIRALS ABACAVIR TABS	MC/DEL	8	ABACAVIR SOL		
	MC MC	APRETUDE	MC/DEL	8	APTIVUS	Use PA Form# 20420	
ll i	MC/DEL	ATAZANAVIR	MC	8	ATRIPLA <sup>1</sup>	Quantity limit of one per	
ll I	MC	BIKTARVY	MC/DEL	8	CIMDUO	day	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
	MC	CABENUVA	MC/DEL	8	COMBIVIR TABS	Only preferred if Norvir     perint is in member's prefile.	least two other drugs that are likely to be active based on the genotype testing.
l I	MC	COMPLERA <sup>1</sup>	MC/DEL	8	EDURANT	script is in member's profile within the past 30 days of	DDI: Reyataz requires prior authorization if it is currently being used in combination with either Prevacid, pantoprazole, Prilosec, or any currently non preferred PPI.
ll I	MC/DEL	DELSTRIGO DESCOVY <sup>1</sup>	MC/DEL	8	EPZICOM <sup>1</sup>	filling Prezista	
l I	MC MC	DIDANOSINE	MC/DEL MC/DEL	8 8	FUZEON INTELENCE	3.Isentress Chewable will	DDI: Norvir requires prior authorization if it is currently being used in combination with either Enablex 15mg or Vesicare 10mg.
	MC/DEL	DOVATO	MC/DEL	8	ISENTRESS <sup>3</sup>	only be approved if between	
	MC	EFAVIRENZ TAB	MC/DEL	8	ISENTRESS HD	the age of 2-12 years old	
	MC/DEL	EFAVIRENZ CAP	MC	8	JULUCA	4. Clinical PA required.	DDI: Preferred Crixivan caps requires prior authorization if it is currently being used in combination with either Enablex 15mg or Vesicare 10mg.
	MC	EFAVIRENZ-EMTRICITABINE-TENOFOVIR DF TAB	MC	8	KALETRA	5. Only preferred for post-	
	MC	EMTRICITABINE-TENOFOVIR	MC/DEL	8	LAMIVUDINE SOLN	exposure prophylaxis.	DDI: The concomitant use of the following drugs with <b>Descovy®</b> is not recommended: tipranavir/ritonavir, St. John's wort, and the antimycobacterials rifabutin, rifampin, or rifapentine.
	MC	EMTRIVA <sup>1</sup>	MC/DEL	8	LEXIVA		
	MC	EPIVIR SOL	MC/DEL	8	NEVIRAPINE		
	MC/DEL	EVOTAZ <sup>1</sup>	MC	8	NORVIR		DDI: Administration with the following drugs: the anticonvulsants carbamazepine, oxcarbazepine, phenobarbital, and phenytoin; the antimycobacterials rifampin and rifapentine; proton
	MC	GENVOYA <sup>1,4</sup>	MC/DEL	8	PIFELTRO		pump inhibitors such as dexlansoprazole, esomeprazole, lansoprazole, omeprazole, pantoprazole, rabeprazole; systemic dexamethasone (more than a single dose); and St. John's wort with <b>Odefsey</b> is contraindicated.
	MC/DEL	ISENTRESS 400MG <sup>5</sup>	MC	8	RETROVIR		· ·
	MC/DEL MC/DEL	ISENTRESS CHEW <sup>3</sup> ISENTRESS POWDER	MC MC/DEL	8	REYATAZ SELZENTRY		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
	MC/DEL	LAMIVUDINE TABS	MC MC	8	STAVUDINE		agents.
	MC/DEL	LAMIVUDINE/ZIDOVUDINE	MC	8	STRIBILD <sup>1</sup>		
	MC/DEL	LOPINAVIR-RITONAVIR SOL	MC/DEL	8	SYMFI <sup>4</sup>		DDI: Tivicay will require prior authorization is used with nevirapine, oxcarbazepine, phenytoin, phenobarbital, carbamazepine, and St. John's wort.
	MC	LOPINAVIR-RITONAVIR TAB	MC/DEL	8	SYMFI LO⁴		
	MC	ODEFSEY <sup>1</sup>	MC/DEL	8	SYMTUZA		
	MC/DEL	PREZCOBIX	MC/DEL	8	TRIZIVIR TABS		
	MC	PREZISTA <sup>2</sup>	MC	8	TRUVADA <sup>1</sup>		
	MC/DEL	RITONAVIR TAB 100MG	MC/DEL	8	VIRACEPT TABS		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,
	MC	RUKOBIA⁴	MC	8	VITEKTA		rifampin, irinotecan, dihydroergotamine, ergotamine, methylergonovine, cisapride, St. John's wort, lovastatin, simvastatin, pimozide, nevirapine, sildenafil (when given as Revatio® for
	MC	SUNLENCA <sup>4</sup>	MC	8	ZERIT		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.
	MC	SUSTIVA <sup>1</sup>	MC	8	VIDEX EC		
	MC	TIVICAY	MC MC/DEL	8	VIREAD TABS <sup>1</sup>		DDI: Combined P-gp, UGT1A1 and strong CYP3A inhibitors may significantly increase plasma concentrations of Sunlenca®. Concomitant administration of Sunlenca® with these
	MC	TIVICAY PD TRIUMEQ <sup>1</sup>	MC/DEL	8	ZIAGEN TABS		inhibitors is not recommended.
	MC	TRIGINEQ TROGARZO <sup>4</sup>	MC/DEL MC/DEL	8 9	ZIAGEN SOL VIDAMI INE YD		Suplanas: In combination with other antiratroviral/a) for the treatment of LIV 1 infection in bequily treatment experienced adults with multiday registrant LIV 1 infection failing their current
1	MC	INUGANZU	WIC/DEL	Э	VIRAMUNE XR	I	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

Ī	МС	TYBOST	Ī	Ī	I	ı	antiretroviral regimen due to resistance, intolerance, or satety considerations.
1	MC	VIREAD POW					
	MC/DEL	ZIDOVUDINE					
		1					
CYTO-MEGALOVIRUS AGENTS	MC	CIDOFOVIR	MC		VALCYTE 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
	MC	FOSCARNET SODIUM	MC/DEL		FOSCAVIR		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
	MC/DEL	GANCICLOVIR	MC/DEL		LIVTENCITY <sup>1</sup>		preferred drug(s) exists.
	MC/DEL	VALGANCICLOVIR	MC/DEL		PREVYMIS	1. Must show failure or	
						contraindication to all the	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
						following ganciclovir, valganciclovir, cidofovir and	agents.
						foscarnet before Livtencity	
						will be approved.	DDI: Livtencity is a substrate of CYP3A4. Coadministration of Livtencity® with strong inducers of CYP3A4 is not recommended, except for selected anticonvulsants.
HERPES AGENTS	MC/DEL	ACYCLOVIR	MC/DEL		5.110(0) 0.170 <sup>1</sup>	Must fail Acyclovir and	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
HERFES AGENTS	MC/DEL MC/DEL	VALACYCLOVIR HCL	MC	0	FAMCICLOVIR <sup>1</sup> SITAVIG	Valacyclovir before non-	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
	MO/DEE	VALACTOLOVIKTIOL	MC/DEL	8	ZOVIRAX <sup>1</sup>	preferred products in step	another drug and the preferred drug(s) exists.
			MC	8		order.	
			MC/DEL	9	VALTREX TABS <sup>1</sup> FAMVIR TABS <sup>1</sup>	Use PA Form# 20420	
INFLUENZA AGENTS	MC	AMANTADINE CAPS	MC	Ť	AMANTADINE TABS	CCC 1 7 (1 Offing 20720	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
	MC	RELENZA DISKHALER AEPB	MC		FLUMADINE TABS		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
	MC/DEL	OSELTAMIVIR <sup>1</sup>	MC		FLUMIST	Tamiflu and Oseltamivir	another drug and the preferred drug(s) exists.
			MC/DEL		RIMANTADINE HCL_TABS	10 caps or 60cc's per month.	
			MC/DEL		TAMIFLU <sup>1</sup>	Will be audited for presence	
			MC/DEL		TAMIFLU SUS	of positive influenza tests in patient or family member.	
			MC/DEL		XOFLUZA	patient of family member.	
						Use PA Form# 20420 for all	
						<u>others</u>	
IMMUNE SERUMS	MC	IMMUNE SERUMS HYPERRHO INJ	1	ı		1	
IMMONE SEROMS	WC	HEPATITIS AGENTS	1				
HEPATITIS C AGENTS	т т	SOFOSBUVIR/VELPATASVIR <sup>2</sup> (Authorized generic	MC/DEL	T T	COPEGUS TABS	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 (in step order), unless an acceptable clinical
	мс	labeler 72626 Asegua Therapeutics)				please see dosage	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
	MC	MAVYRET <sup>2</sup>	MC/DEL		DAKLINZA	consolidation list.	another drug and the preferred drug(s) exists.
	MC/DEL	PEGASYS KIT <sup>1</sup>	MC		EPCLUSA <sup>2</sup>		
	MC/DEL	PEGASYS SOLN	MC		HARVONI <sup>2</sup>	Approvals will require	
	MC/DEL	PEG-INTRON KIT <sup>1</sup>	MC/DEL		REBETOL CAPS	clinical PA. Please see the	DDL Olygip will require a prior outbookston if it is gurroutly being used in combination with drugs known to be significant CVD2AA inhibitors //etecanomals item
	MC	RIBAVIRIN	MC		RIBAPAK	Hepatitis PA form for criteria	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).
	MO/DEL	DIDACDUEDE	MC		SOVALDI <sup>2</sup>		
	MC/DEL	RIBASPHERE	MC				
					VIEKIRA PAK <sup>2</sup> VIEKIRA XR <sup>2</sup>		
			MC MC		VOSEVI		
					ZEPATIER <sup>2</sup>	H DAE #40700	
HEPATITIS AGENTS - MISC.			MC/DEL		ACTIMMUNE	Use PA Form #10700	Approved for chronic granulomatous disease, estephatosis and idiopathic nulmonary fibrosis
	MOIDEL	ENTERNANCE	MC			Use PA Form# 20420	Approved for chronic granulomatous disease, osteopetrosis and idiopathic pulmonary fibrosis.
HEPATITIS B ONLY	MC/DEL MC	ENTECAVIR TENOFOVIR	MC		BARACLUDE HEPSERA 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 (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
	IVIC	IENOFOVIK	MC MC		TYZEKA		another drug and the preferred drug(s) exists.
			MC		VEMLIDY		
			IVIC		VEINELD I		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
	I I	I	1		1		
1 1							aminotransferases (ALT of AST) of histologically active disease, Patient is To years of age of older. Boxed warning: Use not recommended for those co-infected with HIV and HBV who
							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).

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			'				Vemlidy® remain non-preferred and require prior authorization and be available to those who have evidence of bone loss or renal insufficiency or who are unable to tolerate or who have failed on preferred medications.
		RSV PROPHYLAXIS					
RSV PROPHYLAXIS			MC		SYNAGIS <sup>1</sup>	Use PA Form# 30120  1. MaineCare will approve Synagis PA's for start date of November 29, 2021 for infants who meet the guidelines. PA will be approved for max of 5 doses. Maximum 1 dose/30 days. MaineCare will start accepting PAs November 1, 2021."	Please see the criteria listed on the Synagis PA form.
		MS TREATMENTS					
MULTIPLE SCLEROSIS - INTERFERONS	MC MC/DEL MC	AVONEX KIT <sup>1</sup> BETASERON SOLR <sup>1</sup> REBIF SOLN <sup>1</sup>	MC MC/DEL		PLEGRIDY <sup>1</sup> EXTAVIA	establish diagnosis and	Non-Preferred drugs must be tried in step-order and failed due to lack of efficacy or intolerable side effects before lower ranked 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.
MILITIDI E SCI EROSIS - NON-	MC	CODAYONE		8	AMDVDA		Professed datas must be tried and failed due to lack of officacy or intolerable side effects before non-preferred drups will be approved (in step order), unless an acceptable clinical
MULTIPLE SCLEROSIS - NON- INTERFERONS	MC MC/DEL MC/DEL MC MC MC MC MC	COPAXONE  DALFAMPRIDINE ER  DIMETHYL FUMARATE CAP  FINGOLIMOD CAP <sup>2</sup> KESIMPTA <sup>2,5</sup> TERIFLUNOMIDE TAB <sup>2</sup> TYSABRI <sup>1,2</sup>	MC MC MC MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC	8 8 8 8 8 8 8 8 8	AMPYRA AUBAGIO BAFIERTAM BRIUMVI GILENYA GLATOPA MAVENCLAD³ MAYZENT  OCREVUS² OCREVUS ZUNOVO² PONVORY² TASCENSO ODT²⁴ TECFIDERA VUMERITY ZEPOSIA	Prescribing program, a restricted distribution program. Clinical PA is required to establish diagnosis and medical necessity.  2. Clinical PA is required to establish diagnosis and medical necessity.  3. Due to safety profile, use of Mavenclad® is generally recommended for patients who have had an inadequate response to, or are unable to tolerate, an alternate drug indicated for the treatment of MS  4. For the treatment of patients 10 years of age and older.  5. Approved after single step through preferred drugs.	Obtain an electrocardiogram (ECG) to determine whether pre-existing conduction abnormalities are present. In patients with certain pre-existing conditions, advice from a cardiologist should be sought and first-dose monitoring is recommended.  Obetermine whether patients are taking drugs that could slow heart rate of atrioventricular (AV) conduction.  *Liver Function Tests- Obtain recent (i.e. within the last 6 months) transaminase and bilirubin levels.  Ophthalmic Evaluation- Obtain an evaluation of the fundus, including the macula.  *Current or prior medications with immune system effects- If patients are taking anti-neoplastic, immunosuppressive, or immune-modulating therapies, or if there is a history of prior use of these drugs, consider possible unintended additive immunosuppressive effects before starting treatment with Ponvory®. Vaccinations- Test for antibodies to varicella zoster virus (VZV) before starting Ponvory®, VZV vaccination of antibody-negative patients is recommended prior to commencing treatment with Ponvory®. If live attenuated vaccine immunizations are required, administer at least 1 month prior to initiation of Ponvory®. If live attenuated vaccine immunizations are required, administer at least 1 month prior to commencing treatment with Ponvory®. If live attenuated vaccine immunizations are required, administer at least 1 month prior to initiation of Ponvory®.  Mayzent for Relapsing forms of MS: multiple trials of preferred agents, including an intravenous MS product.
<b>/  </b>			, j	1			Mayzent for Active secondary progressive disease: prior trials of two preferred agents are required.
<b>.</b> [	1 1	l	1 ,	1		Use PA Form# 20430	

MULTIPLE SCLEROSIS - MISC			MC		ZINBRYTA <sup>1</sup>	The safety and efficacy of use in children under the age of 17 years have not been established.	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 #20430	
		ASSORTED NEUROLOGICS					
NEUROLOGICS - MISC.	MC	BOTOX <sup>2,4</sup>	MC		DAXXIFY	1. Approval will be limited to	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
	MC	DYSPORT⁴	MC/DEL MC MC/DEL MC		FIRDAPSE MYOBLOC <sup>1</sup> RUZURGI <sup>3</sup> SKYSONA <sup>4,6</sup>	Cervical dystonia.  2. Please see botulinum PA form for additional criteria	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.  Failed/did not tolerate therapeutic trials of muscle relaxants, unless contraindicated, including but not limited to baclofen, cyclobenzaprine, orphenadrine, Skelaxin, and tizanidine.
			MC/DEL		XEOMIN <sup>2</sup>		Migraine: Consideration for Botox approvals will only be made after failures of required trials of the following preferred medications: tricyclic or venlafaxine, beta blocker, valproic acid
						patients between ages 6-16 years of age.	,topiramate.
						Clinical PA required.     For adult patients who are anti-acetylcholine receptor (AChR) antibody positive.	Firdapse is recommended for the treatment of Lambert-Eaton myasthenic syndrome (LEMS) in adults.
						6. For the treatment of patients between ages 4-17 years of age.	
						Use PA Form# 10210	Ruzurgi is recommended for the treatment of Lambert-Eaton myasthenic syndrome (LEMS) in patients 6 years to less than 17 years of age.
NEUROLOGICS- hATTR AGENTS			MC MC/DEL MC/DEL MC/DEL MC/DEL		AMVUTTRA <sup>1</sup> ONPATTRO <sup>1</sup> TEGSEDI <sup>1</sup> VYNDAMAX <sup>1</sup> VYNDAMAX <sup>1</sup> WAINUA <sup>1</sup>	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.  Tegsedi® should be non-preferred and approved for patients for whom other treatments, including Onpattro®, have been ineffective.  Vyndamax will be considered for the treatment of the cardiomyopathy of wild type or hereditary transthyretin-mediated amyloidosis (ATTR-CM) in adults to reduce cardiovascular mortality
					!	II DA E # 00400	and cardiovascular-related hospitalization
NEUDOLOGICS SMA		GENE			CENE	Use PA Form# 20420	Zalganama: The nation is location 2 years of ago AND the diagnacia is agind munaular stracky (CMA) AND the nation has highlight suitations of the CMAM
NEUROLOGICS- SMA	MC	ZOLGENSMA <sup>1</sup>			GENE	Clinical PA is required to establish diagnosis and medical necessity	Zolgensma: The patient is less than 2 years of age AND The diagnosis is spinal muscular atrophy (SMA) AND The patient has bi-allelic mutations of the SMN1 gene AND The patient does not have advanced SMA (e.g. complete paralysis of limbs or permanent ventilator dependence) AND Medication is prescribed per the dosing
		NON-GENE EVRYSDI <sup>1,2</sup>		ŀ	NON-GENE	2. For patients 2 months of age and older.	
	MC MC				!		Spinraza:
		SPINRAZA <sup>1</sup>					The diagnosis is spinal muscular atrophy (SMA) type 1, 2, or 3 (results of genetic testing must be submitted) AND
					 		The patient has at least 2 copies of the SMN2 gene AND
					 		The prescriber is a neurologist, pulmonologist, or other physician with expertise in treating SMA AND
					 		Baseline motor ability has been established using one of the following exams:
							Hammersmith Infant Neurological Exam (HINE)
					 		Hammersmith Functional Motor Scale Expanded (HFMSE)
							Upper Limb Module Test (non-ambulatory)
							Children's Hospital of Philadelphia Infant Test of Neuromuscular Disorders (CHOP INTEND) AND
							Prior to starting therapy, and prior to each dose, the following laboratory tests will be conducted:
					 		Treating provider attests the member has a platelet count > 50,000/ml or greater
1			l I	L	!	I	Treating provider agrees to do platelet count and coagulation test before each dose

								Treating provider agrees to do a quantitative spot urine protein test before each dose  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  will not be approved  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
							Use PA Form# 20420	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.
NEUROLOGICS- RETT SUNDROME		++		MC		DAYBUE <sup>1,2</sup>		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
						DATBUE	1.Clinical PA required for appropriate diagnosis     2. For the treatment of	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 2 years of age and older. Use PA Form# 20420	
ALS DRUGS	MC/DEL		RILUZOLE	MC	-	EXSERVAN	000 1 7 1 0 1 1 1 1 1 1 1 1 1 1 1 1 1 1	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
				MC MC		QALSODY RILUTEK TABS	Clinical PA for indication required	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		RADICAVA <sup>1</sup> RELYVRIO <sup>1</sup>		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).
<i>i</i> I			, ,	MC	ŀ	TIGLUTIK	Use PA Form# 20420	
MOVEMENT DISORDERS	MC MC MC	1	AUSTEDO <sup>1</sup> AUSTEDO XR <sup>1</sup> INGREZZA <sup>1</sup> TETRABENAZINE <sup>1</sup>	MC/DEL		XENAZINE	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 (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.
							Use PA Form# 20420 Use PA Form# 20710 for	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
MUSCULAR DYSTROPHY AGENTS	MC	Ē	EMFLAZA²	MC MC MC MC		AGAMREE* AMONDYS 45 <sup>1</sup> DEFLAZACORT ELEVIDYS <sup>3</sup> DUVYZAT	Clinical prior authorization     to verify diagnosis and use     of stable dose of     corticosteroid for at least 6     months.	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 MC		EXONDYS 51 <sup>1</sup> VILTEPSO <sup>3</sup> VYONDYS 53	For the treatment of Duchenne muscular dystrophy (DMD) in patients	Amondy 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).
							2 years of age and older and a documented intolerance of oral corticosteroid.	Amondy 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
								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  The prescriber is, or has consulted with, a neuromuscular disorder specialist  The patient is ambulatory AND
							to verify diagnosis and use of stable dose of	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
							corticosteroid	Elevidys and Viltepso: 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.
							For the treatment of Duchenne muscular dystrophy (DMD) in patients 2 years of age and older	Viltepso: 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.
			, ,	1	•	1	Use PA Form# 20420	
MYASTHENIA GRAVIS	MC	F	PYRIDOSTIGMINE	MC MC		MESTINON VYVGART <sup>1</sup>	For the treatment of generalized myasthenia gravis (gMG) in adult	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

FRIEDREICH'S ATAXIA AGENTS			MC MC	VYVGART HYTRULO <sup>1</sup> ZILBRYSQ <sup>1</sup> SKYCLARYS <sup>1,2</sup>	patients who are anti- acetylcholine receptor (AChR) antibody positive  Use PA Form# 20420  1. Clinical PA required for	another drug and the preferred drug(s) exists.  Zilbrysq recommended to vaccinate patients for meningococcal infection per current Advisory Committee on Immunization Practices (ACIP) recommendations at least 2 weeks prior to administering the first dose.  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.
		STEROIDS				
GLUCOCORTICOIDS/ MINERALOCORTICOIDS	MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC MC/DEL MC MC/DEL MC MC/DEL MC MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	BUDESONIDE EC 3mg DR CAPS CELESTONE SUSP CORTEF 5 CORTISONE ACETATE TABS DELTASONE TABS DEPO-MEDROL SUSP DEXAMETHASONE DEXPAK FLUDROCORTISONE ACETATE TABS HYDROCORTISONE KENALOG METHYLPREDNISOLONE TABS PREDNISOLONE PREDNISONE SOLU-CORTEF SOLR SOLU-MEDROL SOLR	MC MC MC/DEL MC MC/DEL MC	ALKINDI SPRINKLE CORTEF 10 and 20 TABS FLORINEF TABS HEMADY MEDROL TABS MEDROL DOSEPAK TABS MILLIPRED ORTIKOS ORAPRED SOLN PEDIAPRED LIQD PREDNISONE INTENSOL CONC STERAPRED TABS ZILRETTA	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: All preferred steroids will require clinical PA for patients over 60 that are currently on fluoroquinolone therapy.
		HORMONE REPLACEMENT THERA		<u></u>		
ANDROGENS / ANABOLICS	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	ANDRODERM PT24 ANDROGEL 1% ANDROGEL PUMP 1.62% DANAZOL CAPS TESTOSTERONE CYP	MC MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC MC/DEL MC MC/DEL MC MC/DEL MC MC/DEL MC MC MC/DEL MC	ANADROL-50 ANDRO LA 200 OIL ANDROGEL PACKETS 1.62% ANDROID CAPS AXIRON AZMIRO DELATESTRYL OIL DEPO-TESTOSTERONE OIL FORTESTA HALOTESTIN TABS JATENZO METHITEST TAB METHYLTESTOSTERONE CAP OXANDROLONE STRIANT MUC ER TESTIM TESTOSTERONE GEL PACKETS TESTOSTERONE SOL TESTRED CAPS TLANDO VOGELXO XYOSTED	Use PA Form# 20420	Preferred drugs must be tried and failed due to tack 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. Additionally, laboratory evidence of a testosterone deficiency must be supplied. One of each dosage form should be tried (tablet, injection, and topical)  Oxandrolone: Weight gain (adjunctive therapy): Adjunctive therapy to promote weight gain after weight loss following extensive surgery, chronic infections, or severe trauma, and in some patients who, without definite pathophysiologic reasons, fail to gain or to maintain normal weight. Other indications included in manufacturer labeling: Adjunctive therapy to offset protein catabolism with prolonged corticosteroid administration. Requirement for documentation of weight loss over two readings- Patient has involuntary weight loss of more than 10% of total body weight in less than four months) and, BMI < 18.5 (Normal BMI = 18.5 to 24.9)

ESTROGENS - PATCHES / TOPICAL	MC	EVAMIST	MC/DEL	5	ESTRADIOL PTWK		Approved for failures on multiple oral estrogen agents after 90 day trials or if unable to swallow any oral medication.
	MC/DEL	MINIVELLE PATCH	MC/DEL	8	DIVIGEL <sup>1</sup>	used in specified step order.	
	MC/DEL	VIVELLE-DOT PTTW	MC/DEL	8	CLIMARA PTWK		
			MC/DEL	8	ELESTRIN <sup>1</sup>		
			MC/DEL	8	MENOSTAR PATCH		
						Use PA Form# 20420	
ESTROGENS - TABS	MC/DEL	ESTRADIOL	MC/DEL		ENJUVIA	Must fail preferred products	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
	MC/DEL	PREMARIN TABS	MC/DEL		ESTRADIOL-NORETHINDRONE	before non-preferred products.	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		ESTRACE TABS	products.	all of the site product druggly cases.
			MC		ESTRATAB TABS		
			MC/DEL		MENEST TABS		
			MC/DEL		NORETHINDRON-ETHINYL		
			MC		ORTHO-EST TABS		
						Use PA Form# 20420	
ESTROGEN COMBO'S	MC/DEL	ANGELIQ	MC/DEL		FEMHRT 1/5 TABS <sup>1</sup>	Must fail Premphase and	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
	MC/DEL	COMBIPATCH PTTW	MC/DEL		FYAVOLV	Prempro products before non preferred products.	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	PREMPHASE TABS	MC		LOPREEZA TAB		anound dray and the protected dray(e) exists.
	MC/DEL	PREMPRO TABS	MC/DEL		ORTHO-PREFEST TABS <sup>1</sup>	Use PA Form# 20420	
			MC/DEL		SYNTEST H.S. TABS <sup>1</sup>		
PROGESTINS	MC/DEL	MEDROXYPROGESTERONE ACETA 1	MC/DEL		AYGESTIN TABS	Must fail     Madray prograterana and	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
	MC/DEL	NORETHINDRONE ACETATE TABS <sup>1</sup>	MC		CYCRIN TABS		preferred drug(s) exists.
	MC	17-ALPH HYDROXYPROGESTERONE PWDR	MC		PROGESTERONE POWD	before non-preferred	507
	MC	PROGESTERONE CAPS	MC/DEL		PROMETRIUM CAPS		
			MC/DEL		PROVERA TABS		
						Use PA Form# 20420	
		ENDOMETROSIS					
CENTRAL PRECOCIOUS PUBERTY	MC	FENSOLVI <sup>1</sup>					
AGENTS						1. For pediatric patients 2	
						years of age and older with central precocious puberty	
						(CPP).	
ENDOMETROSIS- NASAL	MC/DEL	SYNAREL (NASAL) SPRAY				,	Synarel is also indicated for central precocious puberty
		, ,					
						Use PA Form# 20420	
ENDOMETROSIS/ UTERINE FIBROIDS-	MC/DEL	ORILISSA <sup>1</sup>	MC		ORIAHNN'	Prior treatment of NSAID	
ORAL	MC	MYFEMBREE <sup>1,2</sup>				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	
ENDOMETROSIS- INJECTABLE	MC/DEL	DEPO-SUBQ PROVERA 104					
I	1	1	I		I	ı	I

1 '	1 1	· · · · · · · · · · · · · · · · · · ·	( I	ı	Use PA Form# 20420	
		CONTRACEPTIVES				
CONTRACEPTIVES - PROGESTIN ONLY	MC/DEL	CAMILA TABS	MC/DEL	JOLIVETTE	T	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
<b>!</b>	MC/DEL	ERRIN	MC/DEL	NORA-BE TABS		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
ļ , , , , , , , , , , , , , , , , , , ,	MC	INCASSIA TAB	MC	ORTHO MICRONOR TABS		preferred drug(s) exists.
ļ , , , , , , , , , , , , , , , , , , ,	MC	HEATHER TAB	1			If member experienced adverse reactions, consider using Oral Contraceptives from other groups.
ļ , , , , , , , , , , , , , , , , , , ,	MC/DEL	NORETHINDRONE ACETATE 0.35MG TABS	1	1		DDI: Preferred Oral Contraceptives will now be non-preferred and require prior authorization if it is currently being used in combination with Tracleer.
1 '	MC/DEL	SLYND	1	1	Use PA Form# 20420	
CONTRACEPTIVES - INJECTABLE	MC/DEL	MEDROXYPROGESTERONE ACETATE 150mg IM	MC/DEL	DEPO-PROVERA 150 mg SUSP	Use PA Form# 20420	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.
CONTRACEPTIVE - EMERGENCY	MC/DEL	ELLA	$\Box$			Due to the extensive list of products, any covered emergency contraceptive product preferred is and available without a PA.
· /	MC	ENCONTRA ONE STEP	1	1	days without PA	
ļ ,	MC	ECONTRA EZ	1	1		
, , , , , , , , , , , , , , , , , , ,	MC	NEW DAY	1	1		
, , , , , , , , , , , , , , , , , , ,	MC	OPCION	1	1		
· /	MC/DEL	OPTION 2	1			
ļ ,	MC	MY CHOICE	1	1		
· /	MC/DEL	MY WAY	1			
· /	MC	LEVONORGESTREL	1	1		
· /	MC/DEL	NEXT CHOICE <sup>1</sup>	1	1	Use PA Form# 20420	
CONTRACEPTIVES - PATCHES/ VAGINAL	MC	ELURYNG <sup>1</sup>	MC	ANNOVERA	Use PA Form# 20420	Approved if adequate clinical reason given why patient unable to comply with other preferred agents including long acting injectable.
PRODUCTS	MC	NUVARING RING <sup>1</sup>	MC	PHEXXI	1. Quantity limit allowing 1	
ļ ,	MC	TWIRLA	MC	ZAFEMY	every 28 days with out PA.	
, , , , , , , , , , , , , , , , , , ,	MC/DEL	XULANE <sup>2</sup>	1	1	2 Deser livrite apply allowin	
· /	1	,	1	1	<ol> <li>Dose limits apply allowing 3 patches per 28 days</li> </ol>	a a contract of the contract o
· /	1	,	1	1	supply.	
· /	1	,	1	1		
CONTRACEPTIVES- LONG ACTING	MC/DEL	MIRENA	MC/DEL	KYLEENA	†	
REVERSIBLE	1 1	,	MC	LILETTA		
· /	1	,	MC	NEXPLANON		
· /	1	,	MC/DEL	PARAGARD		
· /	1 1	,	MC/DEL	SKYLA		
· /	1	,	1	1		
CONTRACEPTIVES - MONOPHASIC	MC/DEL	APRI TABS	MC/DEL	BEYAZ	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
COMBINATION O/C'S	MC/DEL	AVIANE TABS	MC/DEL	BREVICON-28 TABS	If member experienced	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
.   '	MC/DEL	BALZIVA	MC/DEL	LESSINA-28 TABS	adverse reactions, consider	r preferred drug(s) exists.
ļ ,	MC/DEL	CRYSELLE-28 TABS	MC/DEL	LEVORA	using Oral Contraceptives	
ļ ,	MC	DESOGEN TABS	MC/DEL	LOESTRIN FE 1/20 TABS	from other groups.	
, <b> </b>	MC/DEL	ESTARYLLA TAB	MC/DEL	LOESTRIN 1.5/30-21 TABS		
, <b> </b>	MC	HAILEY FE TAB	1			
<b>!</b>	MC/DEL	ISIBLOOM TAB	MC/DEL	MICROGESTIN FE TABS		If member experienced adverse reactions, consider using Oral Contraceptives from other groups.
ļ ,	MC/DEL	JUNEL FE TAB	MC/DEL	LOESTRIN 1/20-21 TABS		
ļ ,	MC	LARIN FE TAB	1	1		
ļ ,	MC/DEL	LESSINA TAB	MC	LO/OVRAL 21 TABS		
· /	MC	LEVORA-28 TAB	MC/DEL	LO/OVRAL 28 TABS		
ļ ,	MC	MILI TAB	MC	NEXTSTELLIS		
· /	1 - 1	,	1	NORDETTE-28 TABS		
ļ ,	MC/DEL	NORGESTIMATE-ETHINYL ESTRADIOL TAB	MC/DEL			
· /	MC/DEL	MIBELAS 24 FE TAB	MC/DEL	NORTREL		DDI: Preferred Oral Contraceptives will now be non-preferred and require prior authorization if it is currently being used in combination with Tracleer.
<u> </u>	MC/DEL	MICROGESTIN FE TAB	MC/DEL	OCELLA		
.  '	MC/DEL	RECLIPSEN	MC/DEL	OVRAL		
, <b> </b>	MC/DEL	SAFYRAL TAB	MC/DEL	PORTIA-28 TABS		
,	MC/DEL	SPRINTEC 28 TABS	MC/DEL	SAFYRAL		
, II	MIC/DEL	OF MINIEO 20 TADO	MIC/DEL	Ohi HAL		I control of the cont

_	_		_	_	_	
<b>                                     </b>	MC/DEL	YASMIN 28 TABS	MC/DEL	ZOVIA		
	MC/DEL	YAZ	. [			
CONTRACEPTIVES - BI-PHASIC	MC/DEL	AZURETTE TAB	MC/DEL	LOSEASONIQUE	If member experienced	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
COMBINATIONS	MC/DEL	CAMRESE	. [		adverse reactions, consider	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
1	MC/DEL	CAMRESE LO	,	<b>I</b>	using Oral Contraceptives	preferred drug(s) exists.
1	MC	DESOGESTREL/ ETH/ ESTRAD 0.15/30mcg	.		from other groups.	If member experienced adverse reactions, consider using Oral Contraceptives from other groups.
	MC/DEL	KARIVA TABS	,	<b>I</b>		
•	MC/DEL	LO LOESTRIN FE	,	<b>I</b>		
		LO LOESTRIN FE PIMTREA TAB	,	<b>I</b>		
<b> </b>	MC/DEL MC	PIMTREA TAB NORETHINDRONE-ETH ESTRADIOL TAB 0.5-35/1-	.			
	WIG	35	,	<b>I</b>		
·	МС	SIMPESSE TBDSPK 3MO	. [			DDI: Preferred Oral Contraceptives will now be non-preferred and require prior authorization if it is currently being used in combination with Tracleer.
			,	<b>I</b>	Lloo DA Form# 20420	DDI. Freteried Oldi Contracepares wiii now be northerened and require prior adminization in it is containly being does in combination with macroscot.
CONTRACEPTIVES - TRI-PHASIC	MC/DEL	VIORELE TAB ENPRESSE	MOIDEL	NORTREL 7/7/7	Use PA Form# 20420	5. C.
CONTRACEPTIVES - TRI-PHASIC COMBINATIONS	MC/DEL		MC/DEL		If member experienced adverse reactions, consider	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
COMBINATIONS	MC/DEL	NORGESTIMATE-ETHINYL ESTRADIOL TAB	MC	ORTHO TRI-CYCLEN LO TABS	using Oral Contraceptives	preferred drug(s) exists.
<b> </b>	MC/DEL	TRIPHASIL 28 TABS	. [		from other groups.	and the diagraph states.
·	MC	TRI-LO-MILI TAB	. [			
<b> </b>	MC	TRI-LO-ESTARYLLA TAB	. [			
<b> </b>	MC	TRI-ESTARYLLA	. [			If member experienced adverse reactions, consider using Oral Contraceptives from other groups.
<b> </b>	MC/DEL	TRI-SPRINTEC TAB	. [			
<b> </b>	MC/DEL	TRI-LO-SPRINTEC	. [			
<b> </b>	MC	TRINESSA	. [			
<b> </b>		1	. [			
<b> </b>		1	. [			
<b>                                     </b>		1	. [			DDI: Preferred Oral Contraceptives will now be non-preferred and require prior authorization if it is currently being used in combination with Tracleer.
<b> </b>		1	. [			DDI. I loterioù Oldi Comitacoparco mil nom de noir-proteirea ana require prior administration in a content pound account combination mai ricolor.
<b>                                     </b>		1	. [		LL - DA 5# 00400	
CONTRACEDTIVES MULTIPLIACIO		4		511.74.71A	Use PA Form# 20420	
CONTRACEPTIVES - MULTI-PHASIC COMBINATIONS		1	MC	NATAZIA		
COMBINATIONS				<u></u>	Use PA Form# 20420	
		VASOMOTOR SYMPTOMS AGENTS				
VASOMOTOR SYMPTOMS AGENTS		1	MC/DEL	VEOZAH		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
'		1				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
<b> </b>		1	. [			preferred drug(s) exists.
<b> </b>		1	. [			
<b> </b>		1	. [			DDI: Avoid concomitant use of Veozah with drugs that are weak, moderate or strong CYP1A2 inhibitors.
<b> </b>		1	. [			
<b>                                     </b>		1	. [			
<b> </b>		1		1	Use PA Form# 20420	Veozah: Approval requires at least one preferred Hormone Replacement Therapy (HRT) and two preferred non-hormonal therapies (i.e., SSRIs, SNRIs, gabapentin, pregabalin, clonidine).
		DIABETES SUPPLIES			USE FA FUITH 20420	
					4.0	
DIABETIC- SUPPLIES		CONTINUOUS GLUCOSE MONITORING <sup>1</sup>	,	<b>I</b>	<ol> <li>Dosing limits apply.</li> <li>Please refer to Dose</li> </ol>	Please refer to the MaineCare Preferred Diabetic Supply List available at www.mainecarepdl.org
<b>                                     </b>		DIABETIC- LANCETS		1	consolidation list.	
<b>4 I</b> '		DIABETIC- LANCING DEVICES		1	001100110011111111111111111111111111111	<b>1</b>
<b>/  </b>		DIABETIC- LANCING DEVICES		1		Continuous Glucose Monitoring Criteria: Patient has a diagnosis of Diabetes Mellitus AND Practitioner feels patient has sufficient training to use CGM
<b>/                                    </b>		DIABETIC- PEN NEEDLES		1		• 2 years of age or older for Dexcom G6, Dexcom G7, Libre 3 Plus sensor, ≥ 14 years for Medtronic Guardian, or ≥ 4 years for Freestyle Libre 2 and Libre 3.
<b>/  </b>		DIABETIC- SYRINGES		1		At least one of the following are documented:
<b>/  </b>		DIABETIC- TEST STRIPS		1		o Hypoglycemic unawareness
<b>/  </b>		DIABETIC- METERS		1		o Treated with insulin (at least 1X day)
<b>                                     </b>		1	. [			o Has history of problematic hypoglycemia with documentation of at least one recurrent level 2 hypoglycemic events, or 1 level 3 hypoglycemic event
<b>                                     </b>		1		1		1
<b>                                     </b>		1	. [			• Approval of non-preferred products will be limited to cases where the CGM is directly integrated with the patient's insulin pump. The make and model of pump must be documented on
<b>                                     </b>		1	. [			the prior authorization.
<b> </b>		1	. [		Use PA Form#20420	
		DIABETES THERAPIES				
DIABETIC - INSULIN	MC/DEL	FIASP	MC/DEL	APIDRA	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
<b>/                                    </b>	MC		MC/DEL	ADMELOG	1. Not to be as a	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
41			WIC/DEL	ADMELOG	manatharany Obtain lab	preferred drua(s) exists.

	MC M	HUMALOG JUNIOR KWIKPEN 100/ML HUMALOG MIX 75/25 HUMALOG 50/50 VIAL HUMULIN INJ 70/30 KWIKPEN HUMULIN INJ 70/30 HUMULIN R INJ U-500 INSULIN ASPART PROT MIX 70-30 INSULIN ASPART INSULIN LISPRO LANTUS SOLN	MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	AFREZZA¹ BASAGLAR HUMALOG KWIKPEN U-200 HUMULIN INJ 50/50 HUMULIN N INJ U-100 HUMULIN R U-100 INSULIN DEGLUDEC LYUMJEV NOVOLIN NOVOLOG NOVOLOG MIX NOVOLOG MIX RELION	monomerapy. ∪onain iao ravalues of pulmonary function and recent smoking history  2. For the treatment of patients ≥3 years of age	
DIABETIC - PENFILLS	MC M	HUMALOG MIX KWIK 50/50 HUMALOG MIX INJ 75/25 KWP HUMALOG KWIK INJ 100/ML HUMALOG KWIK INJ 200/ML HUMULIN R U-500 KWP INSULIN ASPART PROT MIX 70-30 PEN INSULIN ASPART PEN INSULIN LISPRO KWIKPEN U-100 LANTUS SOLOSTAR TOUJEO MAX SOLOSTAR TOUJEO SOLOSTAR	MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	APIDRA OPTICLIK PEN NOVOLIN 70/30 PEN NOVOLOG MIX PENFILL NOVOLOG PENFILL SOLN NOVOLOG FLEXPEN NOVOLOG MIX 70/30 VIAL REZVOGLAR KWIKPEN TRESIBA	e	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.
DIABETIC - DPP- 4 ENZYME INHIBITOR	MC/DEL MC/DEL	JANUVIA <sup>12</sup> TRADJENTA <sup>2</sup>	MC/DEL MC/DEL MC/DEL MC	NESINA ONGLYZA <sup>2</sup> QTERN ZITUVIO	doses of metformin are seen e in members drug profile for at least 60 days within the	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: 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 tellithromycin).
DIABETIC - DPP- 4 ENZYME INHIBITOR- COMBO	MC/DEL MC/DEL MC/DEL	JANUMET <sup>1,2</sup> JANUMET XR <sup>1,2</sup> JENTADUETO <sup>1</sup>	MC/DEL MC/DEL MC MC/DEL MC MC	JENTADUETO XR KAZANO KOMBIGLYZE XR OSENI ZITUVIMET ZITUVIMET XR	doses of metformin are seen e in members drug profile for at least 60 days within the past 18 months or if phosphate binder is currently	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.  Zituvimet/ Zituvimet XR: Approvals will require trial of preferred sitagliptin/metformin products or other preferred diabetic agents.

						_	
						Use PA Form# 20420	
DIABETIC - LANCET-LANCET DEVICE	1 1					Use PA Form# 20420	Please refer to the MaineCare Preferred Diabetic Supply List available at www.mainecarepdl.org
'	1 1						·
DIABETIC - SYRINGES-NEEDLES	<del>                                      </del>	. —	+		<u> </u>	Use PA Form# 20420	Please refer to the MaineCare Preferred Diabetic Supply List available at www.mainecarepdl.org
DIABETIC - OTHER			MC/DEL		CYCLOSET	Use PA Form #20420 for all	1
'	1 1	,	MC		SYMLIN	<u>others</u>	]
AA: TA BUUDITADA	—		<del></del>		<b></b>		
SGLT 2 INHIBITORS	MC/DEL MC/DEL	FARXIGA	MC/DEL		INVOKANA <sup>1</sup>	1.Dosing limits apply please	
'	WIC/DEL	JARDIANCE	MC/DEL		STEGLATRO	refer to Dose Consolidation List	preferred drug(s) exists.
'	1 1						'
'	1 1						· [
	1				<u> </u>	Use PA Form# 20420	
SGLT 2 INHIBITOR COMBINATIONS	MC/DEL	SYNJARDY	MC/DEL		GLYXAMBI		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
'	MC/DEL	SYNJARDY XR	MC/DEL		INVOKAMET		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	XIGDOU XR	MC/DEL		INVOKAMET XR		
'	1 1		MC/DEL		SEGLUROMET		'
'	1 1		MC/DEL		STEGLUJAN		'
, <b> </b>	1 1		MC/DEL		TRIJARDY XR		Glyxambi /Xigduo XR- Verify prior trials and failures or intolerance of preferred treatments from other diabetic categories
, [	1 1						Synjardy® XR is not recommended for patients with type 1 DM or for the treatment of diabetic ketoacidosis.
DIABETIC MONITOR	MC	ONE TOUCH ULTRA 2 KIT	MC		ACCUCHECK	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
DIABETIC MONTON	MC	ONE TOUCH ULTRA MINI KIT	MC		ASCENSIA	Use PA Form# 20420	preferred meters.
, [	MC	TRUE METRIX	MC		ASSURE		
, <b> </b>		,	MC		CONTOUR BREEZE Z		
, <b> </b>			MC		EXACTECH		
, <b> </b>			MC		FREESTYLE INSULINX		
, <b> </b>			MC		FREESTYLE LITE SYSTEM KIT		
, [	1 1		MC		ONE TOUCH ULTRA SMART KIT		
, [	1 1		MC		PRECISION XTRA METER		
, [	1 1		MC		PRODIGY		
, <b> </b>	1 1		MC		TRUETRACK		
ı'							
DIABETIC TEST STRIPS	MC	ONE TOUCH ULTRA <sup>1</sup>	MC		ACCUCHECK	1. Only 50 ct & 100 ct	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
<b>/                                    </b>	MC	TRUE METRIX	MC		ASCENSIA	package size.	preferred meters.
<b>/                                    </b>	1 1		MC		ASSURE	Use PA Form# 20420	
<b>/                                    </b>	1 1		MC		CONTOUR BREEZE Z		
<b>/                                    </b>	1 1		MC MC		EXACTECH		
<b>/                                    </b>	1 1		MC		FREESTYLE		
<b>,                                    </b>	1 1		MC MC		FREESTYLE LITE		
<b>,                                    </b>	1 1		MC		FREESTYLE INSULINX ONE TOUCH DELICA		
, <b> </b>	1 1		MC		PRECISION XTRA		
, <b> </b>	1 1		MC		PRODIGY		
<b>/</b>	1 1		MC		TRUETRACK		
INCRETIN MIMETIC	MC/DEL	RYBELSUS	MC/DEL	5	OZEMPIC		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
,	MC	TRULICITY	MC/DEL	8	ADLYXIN		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
, <b> </b>	MC/DEL	VICTOZA	MC/DEL	8	BYDUREON BCISE		another drug and the preferred drug(s) exists.
ı <sup>-</sup>	•	- -	-		-	-	•

			MC MC/DEL MC/DEL	8 8 8	MOUNJARO SOLIQUA XULTOPHY	Use PA Form# 20420	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	CHLORPROPAMIDE TABS GLIMEPIRIDE GLIPIZIDE TABS GLIPIZIDE ER TABS GLYBURIDE MICRONIZED TABS GLYBURIDE TABS TOLAZAMIDE TABS TOLBUTAMIDE TABS	MC/DEL MC MC MC/DEL MC/DEL MC/DEL		AMARYL TABS DIABETA TABS GLUCOTROL TABS GLUCOTROL XL TBCR GLYNASE TABS MICRONASE TABS	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	METFORMIN HCL TABS METFORMIN ER	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/DEL MC MC		ACTOPLUS MET <sup>1</sup> ACTOPLUS MET XR AVANDARYL <sup>1</sup> AVANDAMET TABS <sup>1</sup>	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	PIOGLITAZONE HCL'	MC/DEL MC		ACTOS TABS <sup>3</sup> AVANDIA TABS <sup>2</sup>	Pioglitazone HCL is preferred if therapeutic doses of metformin, sulfonylurea or insulin are	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 or a significant potential drug interaction between another drug and the preferred drug or a significant potential drug interaction between another drug and the preferred drug or a significant potential drug interaction between another drug and the preferred drug or a significant potential drug interaction between another drug and the preferred drug or a significant potential drug interaction between another drug and the preferred drug or a significant potential drug interaction between another drug and the preferred drug or a significant potential drug interaction between another drug and the preferred drug or a significant potential drug interaction between another drug and the preferred drug or a significant potential drug interaction between another drug and the preferred drug or a significant potential drug interaction between another drug and the preferred drug or a significant potential drug interaction between another drug and the preferred drug or a significant potential drug interaction between another drug and the preferred drug or a significant potential drug interaction between another drug and the preferred drug or a significant potential drug interaction between another drug and the preferred drug or a significant potential drug interaction between another drug and the preferred drug or a significant potential drug interaction for a significant potent
						Current users of Avandia who have tried Actos will be able to continue use of Avandia.     Dosing limits apply please refer to Dose Consolidation List	
DIABETIC - ALPHAGLUCOSIDASE	MC/DEL		MC	<b></b> '	PRECOSE 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
DIABETTO - ALFTINGEOGGIDAGE	WIO/DEL			'	FREGORE TABO	<u>Use PA Form# 20420</u>	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 - SULFONYLUREA / BIGUANIDE	MC/DEL	GLYBURIDE/METFORMIN	MC MC MC/DEL		GLUCOVANCE TABS <sup>1</sup> METAGLIP TABS <sup>1</sup> DUETACT <sup>2</sup>	Use individual ingredients.     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	NATEGLINIDE	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.

		1		1		
						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.
		CHICOGE ELEVATINO ACENT			<u> </u>	
OLUGOOF FLEWATING AGENTS	MOIDEL	GLUCOSE ELEVATING AGENTS		OLUGA GON DIA ONGOTIO KIT		
GLUCOSE ELEVATING AGENTS	MC/DEL	BAQSIMI <sup>1</sup>	MC	GLUCAGON DIAGNOSTIC KIT	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 (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
	MC/DEL	GVOKE <sup>2</sup>	MC	ZEGALOGUE <sup>3</sup>	1. For the treatment of	another drug and the preferred drug(s) exists.
			MC		patients ≥ 4 years of age.	and the did give the protection of didgle) states.
					2. For the treatment of	
					patients ≥ 2 years of age.	
					3. For the treatment of	
					patients ≥ 6 years of age.	
					, ,	
		THYROID				
THYROID EYE DISEASE			MC	TEPEZZA	Use PA Form# 20420	
TITIKOID ETE DISEASE			IVIC	ILFLEZZA	USE PA FORM# 20420	
TINDOID HORMONES	MOIDEL	ARMOUR THYPOIR TARG		LEVOTING CONTINUE CONTINUE COLD		
THYROID HORMONES	MC/DEL	ARMOUR THYROID TABS	MC	LEVOTHYROXINE SODIUM 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 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
	MC/DEL	CYTOMEL TABS	MC/DEL	LIOTHYRONINE	1.Clinical PA is required to	preferred drug(s) exists.
	MC/DEL	ERMEZA <sup>1</sup>	MC	SYNTHROID TABS	confirm diagnosis of dysphagia.	promote diagram state.
	MC/DEL	LEVOTHROID TABS	MC/DEL	THYQUIDITY	dyspriagia.	
	MC/DEL	LEVOTHYROXINE SODIUM TABS				
	MC/DEL	LEVOXYL TABS				
	MC/DEL	UNITHROID TABS				
ANTITHYROID THERAPIES	MC/DEL	METHIMAZOLE 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
	MC/DEL	PROPYLTHIOURACIL TABS				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.
						preferred drug(s) exists.
_		CUSHING DISEASE AGENTS				
CUSHING DISEASE AGENTS			MC	ISTURISA <sup>1</sup>		
			MC	RECORLEV	1. For the treatment of adult	Recorlev® is associated with dose-related QT interval prolongation. QT interval prolongation may lead to life-threatening ventricular dysrhythmias such as Torsade de pointes.
					patients with Cushing's	
					disease for whom pituitary	
					surgery is not an option or has not been curative.	
					nas not been carative.	
					Use PA Form #20420	
		OSTEOPOROSIS / BONE AGENTS			0361 A 1 01111 #20420	
OSTEOPOROSIS	MC/DEL	OSTEOPOROSIS / BONE AGENTS ALENDRONATE	MC/DEL	ACTONEL TABS	H DA F # 00400	Destarted drugs must be tried and failed due to lack of afficacy or intelerable side affects before non professed drugs will be approved, unless an accordable aliased expension is affected as
OG I EUFURUSIS	MC/DEL	ALENDRONATE	MC/DEL		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
l I			MC	AREDIA SOLR	<ol> <li>Approval only requires failure of Alendronate.</li> </ol>	preferred drug(s) exists.
			MC	BINOSTO	idilate of Alcharonate.	
			MC/DEL	BONIVA INJECTION KIT		
			MC/DEL	BONIVA TABS <sup>2,4</sup>	Quantity limits apply,	Binosto use preferred generic alendronate tablets
l I			MC/DEL	CALCITONIN NS	please see dosage consolidation list.	
			MC/DEL	DUAVEE	consolidation list.	Evenity® should be limited to 12 monthly doses
			MC/DEL	DIDRONEL TABS	3. Please use Alendronate	
			MC	EVISTA TABS <sup>1</sup>	and Vitamin D.	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
			MC/DEL	EVENITY <sup>2</sup>		males with fibrodysplasia ossificans progressive (FOP).
			MC	FORTEO	4. Please use other preferred	
			MC/DEL	FORTICAL	agents.	
•                   •	ı	1		I. Sittler		

			MC/DEL MC MC MC MC MC MC			Obtain baseline ophthalmology exams and renal ultrasounds and then periodically during treatment     Clinical PA for indication required.	
FIBROBLAST GROWTH FACTOR 23 INHIBITORS	MC	CRYSVITA <sup>1</sup>				1.Preferred for patients <21 years for the treatment of X-linked hypophosphatemia. <u>Use PA Form #20420</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.
		CALCIMIMETIC AGENTS					
CALCIMIMETIC AGENTS			MC MC		PARSABIV SENSIPAR	<u>Use PA Form# 30115</u>	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 <sup>1</sup> NORDITROPIN SOLN <sup>1</sup> SKYTROFA <sup>1,2</sup>	MC MC/DEL MC/DEL MC MC MC MC MC/DEL	8 8 8 8 8	NGENLA OMNITROPE	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		VOXZOGO1	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
SOMATOSTATIC AGENTS			MC/DEL MC MC MC/DEL MC	8	OCTREOTIDE INJ <sup>1</sup> BYNFEZIA <sup>1</sup> MYCAPSSA <sup>1</sup> SANDOSTATIN <sup>1</sup> SOMATULINE <sup>1</sup>	Use PA Form# 10710  1. Non-preferred products must be used in specified step order.	
GH ANTAGONISTS		GROWTH HORMONE ANTAGONISTS					
GH ANTAGUNISTS		<b>!</b> !	WC	1 '	SOMAVERT	<u>Use PA Form# 10710</u>	Approved for acromegaly patients failing surgery/radiation/drug therapy including bromocriptine and sandostatin.
	_	VASOPRESSIN RECEPTOR ANTAGONI	IST				
VASOPRESSIN RECEPTOR ANTAGONIST			MC MC/DEL		JYNARQUE¹ 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	DESMOPRESSIN TABS	MC/DEL	5	DDAVP TABS	1. Products must be used in	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,
	MC/DEL	DDAVP SOLN	MC/DEL	6	DESMOPRESSIN SPRAY <sup>1</sup>	specified step order.	lower relapse rate) and must periodically attempt weaning (at 6 month intervals).
			MC	8	DESMOPRESSIN ACETATE SOLN <sup>1</sup>	Nocturnal enuresis patients will be encouraged to	
			MC/DEL	8	NOCDURNA <sup>1</sup>	periodically attempt stopping	
						DDAVP.	
					,		
			MC	8	NOCTIVA <sup>1</sup>		
			MC/DEL	8	STIMATE SOLN <sup>1,2</sup>	Patients with a diagnosis	
						of hemophilia or Von Willebrand's disease will be	
						exempt from prior	
						authorization.	
						Use PA Form# 20420	
ANTISPASMODICS	MC/DEL	OXYBUTYNIN	MC/DEL	8	DARIFENACIN ER 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
	MC/DEL	TOLTERODINE	MC/DEL	8	DITROPAN	OSCI AT OTHE 20420	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
	IIIO/DEE	TOPTEROBIRE	MC/DEL	8	FLAVOXATE HCL TAB		preferred drug(s) exists.
			MC/DEL	Ů	ENTONIE HOE IND		
			mo/DLL				
ANTISPASMODICS - LONG ACTING	MC	FESOTERODINE	MC	8	DITROPAN XL 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
	MC/DEL	GELNIQUE GEL PACKET	MC/DEL	8	ENABLEX <sup>1,2</sup>	See Criteria Section.	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
	MC/DEL	MYRBETRIQ	MC	8	GEMTESA <sup>2</sup>	2. Use a preferred long	preferred drug(s) exists.
	MC/DEL	OXYBUTYNIN ER TABS	MC/DEL	8	TOLTERODINE TAB	acting antispasmodic.	1. Vesicare 5mg and Enablex 7.5mg maximum doses if given with drugs known to be significant CYP3A4 inhibitors.(Ketoconazole, Sporanox, Erythromycin, Fluconazole, Nefazodone,
	MC/DEL	OXYTROL	MC/DEL	8	TOVIAZ	3. For the treatment of	Nelfinavir, and Ritonavir)
	MC/DEL	SOLIFENACIN SUCCINATE TAB	MC	8	VESICARE <sup>1</sup>	patients ≥ 2 years of age.	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:
	MC/DEL	TROSPIUM	MC	8	VESICARE <sup>3</sup> LS		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		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
ENGINIONIA INCAINICITO	IVIC	CARGLUMIC ACID TABS	ino		ONINDAGEO TADO		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	
UREA CYCLE DISORDER	MC	BUPHENYL TABLET	MC		BUPHENYL POWDER		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	PHEBURANE GRANULES	MC		RAVICTI LIQUID		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
			MC		OLPRUVA		preferred drug(s) exists.
			MC/DEL		SODIUM PHENYLBUTYRATE POWDER		
			MC/DEL		SODIUM PHENYLBUTYRATE TAB		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 1.2m2 or greater, with urea cycle disorders (UCDs) involving deficiencies of carbamylphosphate synthetase (CPS), omithine transcarbamylase (OTC), or
							argininosuccinic acid synthetase (AS).
						Use PA Form# 20420	
		METABOLIC MODIFIER			Tops and		
HERED. TYROSINEMIA			MC		ORFADIN	Use PA Form# 20420	Approved for Type 1 hereditary tyrosinemia patients. Must include laboratory evidence of dx at first PA.
FARRY DISEASE ASSESTS	1		1		CLEARDIO!	4 Oliminal DA 1	
FABRY DISEASE AGENTS			MC		ELFABRIO <sup>1</sup>	<ol> <li>Clinical PA to verify appropriate diagnosis.</li> </ol>	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
			MC MC/DEL		FABRAZYME <sup>2</sup>		preferred drug(s) exists.
			WIC/DEL		GALAFOLD <sup>1</sup>	2.For the treatment of patients 2 years of age and	
						older.	Elfabria and Calfald: For the treatment of adults with confirmed Febru discoses
							Elfabrio and Galfold: For the treatment of adults with confirmed Fabry disease.
						Lico DA Form# 20420	
		ANTIHYPERTENSIVES / CARDIAC				Use PA Form# 20420	
CARDIAC GLYCOSIDES	MC/DEL	DIGITEK TABS				Use PA Form# 20420	
		I				000 1 AT 0111# 20420	

1 !	MC/DEL	ı	DIGOXIN	1 1	I	I	ı	
	MC/DEL		LANOXIN		1 '	1		
CARDIAC MYOSIN INHIBITORS			<del> </del>	MC		CAMZYOS	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.
			· '			!		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.
			1		1 '	1		DDI: Concomitant use of Camzyos® with a moderate to strong CYP2C19 inhibitor or a strong CYP3A4 inhibitor is contraindicated.
CARDIAC - SINUS NODE INHIBITORS				MC		CORLANOR		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
			1		1 '	1	Use PA Form#20420	
CARDIAC- ERAS				MC		TRYVIO		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.
						!		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
			1		1 '	1	54.5 #00400	1
CARDIAC- SOLUBLE GUANYLATE CYCLASE STIMULATORS	<b> </b>		<u> </u>	MC/DEL	$\overline{}$	VERQUVO	Use PA Form#20420	<del> </del>
CTOLAGE STIMULATORS			1		1 '	1		
			1		<b>'</b>	1	Use PA Form# 20420	
CARDIAC RISK REDUCTION- SGLT2/GLP- 1				MC MC	$\Box$	INPEFA <sup>1</sup> LODOCO	To reduce the risk of cardiovascular death,	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
			1	MC/DEL	1 '	WEGOVY	hospitalization for heart failure, and urgent heart	another drug and the preferred drug(s) exists.
	'	1	1		1 '		failure visit in adults with: Heart failure or Type 2	Lodoco: Patient must have tried and failed generic colchicine due to lack of efficacy or intolerable side effects
			1	1 )	1 '		diabetes mellitus, chronic kidney disease, and other	Wegovy:
			1	1 )	1 '	1	cardiovascular risk factors.	Patient has BMI > 27 kg/m2, and is not being used for weight loss only
			1	1 )	1 '	1		Patient has history of at least one of the following:  o Stroke
		1	1	1 )	1 '	1		o Myocardial Infarction
		1	1	1 )	1 '	1		o Symptomatic peripheral arterial disease  Patient does not have diagnosis of diabetes, end stage renal disease/dialysis, or HFrEF (EF < 45%)
ANTIANGINALSIsosorbide Di-nitrate/ Mono-Nitrates	MC/DEL		ISOSORBIDE MONONITRATE TABS	MC		DILATRATE SR CPCR	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
MONO MARKES	MC/DEL	<u> </u>	ISOSORBIDE MONONITRATE ER	MC MC		ISORDIL TABS ISORDIL TITRADOSE TABS		preferred drug(s) exists.
	'	1 '	1	MC		ISOSORBIDE DINITRATE SUBL		
	1 '	1 '	1	MC/DEL MC/DEL		ISOSORBIDE DINITRATE TABS ISOSORBIDE DINITRATE CR TBCR		
	1 '	1 '	1	MC/DEL		ISOSORBIDE DINITRATE ER TBCR		
	1 '	1 '	1	MC/DEL		ISOSORBIDE DINITRATE TD TBCR		
	1 '	1 '	1	MC/DEL		IMDUR TB24		
	1 '	1 '	1	MC/DEL MC		ISMO TABS MONOKET TABS		
, L						WONORET TABO	<u> </u>	.1

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NITRO - OINTMENT/CAP/CR	MC/DEL	, '	NITROBID OINT	1 1			Use PA Form# 20420	1
l ,	MC/DEL	, '	NITROGLYCERIN CPCR	1 1	,	1		
l ,	MC	, '	NITROL OINT	1 1		1		
1	MC	, '	NITRO-TIME CPCR	1 1	,	1		
NITRO - PATCHES	MC/DEL	1	NITROGLYCERIN PT24 <sup>1</sup>	MC		NITRODISC PT24	1. At least 2 step 1's and	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/DEL	1	NITRO-DUR PT 24 0.8MG <sup>1</sup>	MC/DEL	,	NITRO-DUR PT24	step 3 of the preferred	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
		, '	1	1 1	,	1	products must be used in	preferred drug(s) exists.
		, '	1	1 1	,	1	specified order or PA will be required.	
		, '	1	1 1	,	1	lequileu.	
		, '	1	1 1	,		Use PA Form# 20420	
NITRO - SUBLINGUAL/ SPRAY	MC/DEL		NITROSTAT SUBL	MC/DEL	,	NITROQUICK SUBL	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
		, '	1	MC		NITROLINGUAL SOLN		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
		, '	1	MC	,	NITROLINGUAL TABS		preferred drug(s) exists.
BETA BLOCKERS - NON SELECTIVE	MC/DEL		CARVEDILOL	MC		ASPRUZYO	Recommend using BID	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
<b> </b>	MC	, '	LEVATOL TABS	MC/DEL	,	BETAPACE TABS	since its effects do not last	
<b> </b>	MC/DEL	, '	NADOLOL TABS	MC	,	BETAPACE AF TABS	24 hours.	preferred drug(s) exists.
	MC/DEL		PINDOLOL TABS	MC	,	COREG CR <sup>3</sup>	2. Please use other	
	MC/DEL		PROPRANOLOL HCL SOLN <sup>1</sup>	MC	,	COREG TABS	strengths in combination to	DDI: Concomitant use of Ranolazine products with strong CYP3A inhibitors, including ketoconazole, itraconazole, clarithromycin, nefazodone, nelfinavir, ritonavir, indinavir, and
	MC/DEL		PROPRANOLOL HCL SOLN PROPRANOLOL HCL TABS <sup>1</sup>	MC/DEL	,	CORGARD TABS	obtain this dose.	saquinavir, is contraindicated.
<b> </b>	MC/DEL		PROPRANOLOL HCL TABS PROPRANOLOL HCL 60MG TABS	MC/DEL	,	INDERAL TABS		
<b> </b>	MC/DEL		PROPRANOLOL LA CAPS	MC/DEL	,	HEMANGEOL SOL	Dosing limits still apply.	
	MC		RANOLAZINE ER TABS	MC	,	INDERAL XL CAP	Please see dose	
<b> </b>	MC/DEL			MC	,	INDERAL LA CPCR	consolidation list	
<b> </b>	MC/DEL		SOTALOL AF SOTALOL HCL TABS	MC	,	INNOPRAN XL		
<b> </b>	MC/DEL	, '	TIMOLOL MALEATE TABS	MC	,			
/ <b> </b>	WIC/DEL	, '	TIMOLOL WALEATE TABS	WIC	,	RANEXA		
		, '	1	1 1	,	1	Use PA Form# 20420	
DETA DI COVEDE CADDIO SEI ECTIVE	MO/DEL		ACCRUTOLOL HOL CARS	- MC		VEDLONIC TARE	1 Decemmend using	The first seed folial due to local of officers or intellegable side officers non professed drugs will be approved uplace an accordable clinical expension in efform on
BETA BLOCKERS - CARDIO SELECTIVE	MC/DEL		ACEBUTOLOL HCL CAPS	MC MC/DEL	,	KERLONE TABS  LOPRESSOR TABS	Recommend using     Atenolol (and metoprolol)	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
<b> </b>	MC/DEL MC/DEL		ATENOLOL TABS <sup>1</sup> BETAXOLOL HCL TABS	MC/DEL MC	,	SECTRAL CAPS	BID since its effects do not	
i <b>i</b>	MC/DEL		BISOPROLOL FUMARATE TABS	MC/DEL	,	TENORMIN TABS	last 24 hours.	
i <b>i</b>	MC/DEL		BYSTOLIC	MC/DEL	,	TOPROL XL TB24	U DA 5# 00400	
	MC/DEL			MC/DEL		ZEBETA TABS	Use PA Form# 20420	1
<b>                                   </b>	MC/DEL		METOPROLOL TARTRATE TABS <sup>1</sup> METOPROLOL ER	NIC/DEL	,	ZEBETA TADO		
11 7				1 1	,	1		
TTT DI COVERG AL DUA / DETA	MC/DEL		NEBIVOLOL HCL TAB	110		TO WOATE TARO		
BETA BLOCKERS - ALPHA / BETA	MC/DEL	i '	LABETALOL HCL TABS	MC		TRANDATE 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
11 7		, '	1	1 1	,	1		preferred drug(s) exists.
THE SOURCE A PUREATIO COMPOS	MO/DEL		THE STORY OF THE STORY OF TAR	· · · · · · · · · · · · · · · · · · ·		2000000	Use PA Form# 20420	professional diagrams and a second se
BETA BLOCKERS & DURECTIC COMBOS	MC/DEL	. !	METOPROLOL-HYDROCHLOROTHIAZIDE TAB	MC/DEL		DUTOPROL	Use PA Form# 20420	
CALCIUM CHANNEL BLOCKERS	MO/DEL		<u> </u>	$\longrightarrow$		<del></del>		<u></u>
CALCIUM CHANNEL BLOCKERS Amlodipine, Bepridil, Diltiazem,	MC/DEL	, '	AMLODIPINE <sup>1</sup>	1 1		1	<ol> <li>Dosing limits apply, please see dose</li> </ol>	
Felodipines, Isradipines, Nifedipines,		, '	1	MOIDEL	,	WATER TIA	consolidation list.	
Nisoldipine, and Verapamil		, '	1	MC/DEL	,	KATERZIA		
11 /		, '	1	MC/DEL	,	NORLIQVA		
11 /			4	MC/DEL		NORVASC TABS <sup>1</sup>	Use PA Form# 20420	
11 ,	MC		DILTIA XT CP24	MC/DEL	5	DILACOR XR CP24 <sup>1</sup>	Products must be used in specified order or PA will be	
11 7	MC/DEL		DILTIAZEM HCL ER CP24	MC/DEL	6	TAZTIA <sup>1</sup>	required. Just write	e exception is offered on the Prior Authorization form, such as the presence of a condition that prevents usage of the preferred drug of a significant potential drug interaction between another drug and the preferred drug(s) exists.
11 7	MC/DEL		DILTIAZEM HCL XR CP24	MC	8	CARDIZEM TABS <sup>1</sup>	"Diltiazem 24-hour"and the	0 1 0(7
11 /	MC/DEL		DILTIAZEM CD 300MG CP24	MC	8	CARDIZEM CD CP24 <sup>1</sup>	pharmacy will use a	
11 7	MC/DEL		DILTIAZEM CD 360MG CP24	MC	8	CARDIZEM LA TB24 <sup>1</sup>	preferred long acting	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-
11 /	MC		CARTIA XT CP24 <sup>1</sup>	MC	8	CARDIZEM SR CP12 <sup>1</sup>	diltiazem that does not require PA.	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.
11 7	MC/DEL		DILTIAZEM CD CP24 <sup>1</sup>	MC/DEL	8	DILTIAZEM HCL TABS <sup>1</sup>	lequile FA.	
41 /	MC/DEL		DILTIAZEM HCL ER CP241	MC/DEL		DILTIAZEM HCL ER CP12 <sup>1</sup>		· [
41 /	MC/DEL	, '	DILTIAZEM XR CP24 <sup>1</sup>	MC/DEL	8	DILTIAZEM HCL ER CP12 <sup>1</sup>		· <b> </b>
41 7	MC/DEL	!	TIAZAC CP24 <sup>1</sup>	4		<u> </u>	Use PA Form# 20420	
/   /			'	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

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				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.
1		Ī		MC		DYNACIRC 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
				MC		DYNACIRC CR TBCR <sup>1</sup>	Established users will be grandfathered	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.
							9	
				MC		CARDENE SR CPCR	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		NICARDIPINE HCL CAPS		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		AFEDITAB CR	MC/DEL		ADALAT CC TBCR1	Established users of	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
	MC/DEL		NIFEDIAC CC	MC/DEL		NIFEDIPINE CAPS	Adalat CC are	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
	MC/DEL		NIFEDICAL XL TBCR	MC/DEL		PROCARDIA CAPS	grandfathered.	between another drug and the preferred drug(s) exists.
	MC/DEL		NIFEDIPINE TBCR	MC/DEL		PROCARDIA XL TBCR	Use PA Form# 20420	
	MC/DEL		NIFEDIPINE ER TBCR				550 T T T GITTIN 20 120	
				MC		SULAR TB24	Established users of	
				MC		SULAR CR <sup>1</sup>	10MG and 20MG strengths are grandfathered.	
							ale granulathereu.	
							Use PA Form# 20420	
	MC/DEL	1	VERAPAMIL HCL CR TBCR	MC/DEL		CALAN TABS	Products must be used in	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
	MC/DEL		VERAPAMIL HCL CK TBCK VERAPAMIL HCL ER TBCR	MC/DEL		CALAN TABS CALAN SR TBCR	specified order or PA will be	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
	MC/DEL		VERAPAMIL HCL SR TBCR	MC/DEL		COVERA-HS TBCR	required. Just write	another drug and the preferred drug(s) exists.
				MC		ISOPTIN-SR	"Verapamil 24-hour" and the	
				MC/DEL		VERAPAMIL HCL ER CP24	pharmacy will use a preferred long acting generic	
				MC/DEL		VERAPAMIL HCL SR CP24	that does not require PA.	
				MC/DEL		VERAPAMIL HCL TABS		
				MC/DEL		VERELAN CP24		
				MC/DEL		VERELAN PM CP24	Use PA Form# 20420	
ANTIARRHYTHMICS	MC/DEL		AMIODARONE HCL	MC/DEL		CORDARONE	Prescription must be	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/DEL		DISOPYRAMIDE	MC/DEL		DISOPYRAMIDE	written by Cardiologist.	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
	MC/DEL		FLECAINIDE	MC/DEL		MULTAQ		preferred drug(s) exists.
	MC/DEL		MEXILETINE HCL	MC/DEL		NORPACE		
	MC/DEL		PROCAINAMIDE	MC/DEL		PACERONE		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
	MC/DEL		PROPAFENONE	MC		QUINIDEX	Use PA Form# 20420	(doses greater than 20mg/day) or Levofloxacin or Gemifloxacin, or Moxifloxacin, or Ofloxacin.
	MC		QUINAGLUTE	MC/DEL		TAMBOCOR		
	MC/DEL		QUINIDINE GLUCONATE	MC/DEL		TIKOSYN <sup>1</sup>		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
	MC/DEL		QUINIDINE SULFATE	MC		RYTHMOL SR		medications: Erythromycin, Amiodarone and other antiarrhythmics, TCA's, Phenothiazine, Ketoconazole, Itraconazole, Voriconazole, Cyclosporine, Telithromycin, Clarithromycin,
				MC/DEL		RYTHMOL		Nefazodone, Ritonavir.
ACE INHIBITORS	MC/DEL		BENAZEPRIL HCL	MC	5	MAVIK TABS	Non-preferred products	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
	MC/DEL		CAPTOPRIL TABS	MC/DEL	5	ACCUPRIL TABS	must be used in specified	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
	MC/DEL		ENALAPRIL MALEATE TABS	MC/DEL	8	ACEON TABS <sup>1</sup>	order.	another drug and the preferred drug(s) exists. Non-preferred products are subject to step-order requirements unless clinical circumstances warrant exception.
	MC/DEL		FOSINOPRIL SODIUM	MC/DEL	8	ALTACE CAPS <sup>1</sup>	Use PA Form# 20420	
	MC/DEL		LISINOPRIL TABS	MC	8	EPANED		
	MC/DEL		RAMIPRIL	MC/DEL	8	LOTENSIN TABS <sup>1</sup>		
	MC/DEL		QUINAPRIL HCL	MC/DEL	8	MOEXIPRIL HCL <sup>1</sup>		
				MC	8	MONOPRIL HCT TABS <sup>1</sup>		
				MC/DEL	8	PRINIVIL TABS <sup>1</sup>		
				MC	8	QBRELIS		
				MC/DEL	8	UNIVASC1		
				MC	8	VASOTEC TABS <sup>1</sup>		
				MC/DEL	8	ZESTRIL TABS <sup>1</sup>		
ANGIOTENSIN RECEPTOR BLOCKER	MC/DEL		AMLODIPINE-OLMESARTAN TAB <sup>3</sup>	MC/DEL	8	ATACAND TABS	Use PA Form# 20420	Per best practices patient should have trialed prior therapy of ACE inhibitor or currently on a diabetic therapy
	MC/DEL		IRBESARTAN <sup>1</sup>	MC/DEL	8	AVAPRO	Dosing limits apply,	
	MC/DEL		LOSARTAN <sup>1</sup>	MC/DEL	8	BENICAR TABS	please see dose	
	MC/DEL		MICARDIS TABS <sup>3</sup>	MC/DEL	8	COZAAR	consolidation list.	
	MC/DEL		OLMESARTAN <sup>1</sup>	MC/DEL	8	DIOVAN	2. Use preferred active	
	MC/DEL		TELMISARTAN <sup>1</sup>	MC/DEL	8	EDARBI	ingredients which are	
•	•	•			-	•	DA	•

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•	1		MC	8	TEVETEN TABS	avaliable without PA.	· [
1	1	1	J	1		Preferred without a PA	· [
1	1	1	J	1		only if patient on a diabetic	· [
1	1	1	J	1		therapy or prior ACE therapy.	· [
!	1		_ <b> </b>	1			· [
DIRECT RENIN INHIBITOR	+	<del></del>	MC/DEL		AMTURNIDE	Must show failure of single	
DINEVI NEIM III	1 1		MC/DEL	1	TEKTURNA <sup>1</sup>	and combination therapy	1
1	1 1		MC/DEL	1	TEKTURNA* TEKAMLO	from all preferred	·
•	1 1	1		1	TETOWIEG	antihypertensive categories.	·]
	1	1		1		Use PA Form# 20420	·
ANTIHYPERTENSIVES - CENTRAL	MC/DEL	CLONIDINE HCL TABS	MC/DEL		CLONIDINE PATCH	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/DEL	GUANFACINE HCL TABS	MC/DEL	1	CLONIDINE TTS		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
	MC/DEL	HYDRALAZINE HCL TABS	MC	1	GUANABENZ ACETATE TABS	•	preferred drug(s) exists.
	MC	HYLOREL TABS	MC	1	ISMELIN TABS	•	· [
	MC/DEL	METHYLDOPA TABS	MC/DEL	1	MINIPRESS CAPS	•	· [
•	MC/DEL	MINOXIDIL TABS	MC	1	NEXICLON	,	· [
•	MC/DEL	PRAZOSIN HCL CAPS	MC/DEL	1	TENEX TABS	,	· [
1	MC/DEL	RESERPINE TABS		1		•	· 1
CONTRICTORS AND SA CHANNEL	$\longleftarrow$					4 Dt-li- will only bo	<u> </u>
ACE INHIBITORS AND CA CHANNEL BLOCKERS	1 1	1	MC/DEL	8	AMLODIPINE/BENAZEPRIL	<ol> <li>Prestalia will only be approved for patients ≥ 18</li> </ol>	· [
BEOCKERG	1	1	MC	8	PRESTALIA <sup>1</sup>	years of age.	· [
<b> </b>	1	1	MC	8	TARKA TBCR		· [
	1	1	MC/DEL	9	LOTREL CAPS	Use individual preferred	· [
	1			1		generic medications.	· [
			ليلل			Use PA Form# 20420	
ACE AND THIAZIDE COMBO'S	MC/DEL	BENAZEPRIL HCL/HYDROCHLOR	MC/DEL	1	ACCURETIC 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
<b>   </b>	MC/DEL	CAPTOPRIL/HYDROCHLOROTHIA	MC	1	MONOPRIL HCT TABS		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.
<b>                                     </b>	MC/DEL	ENALAPRIL MALEATE/HCTZ TABS	MC/DEL	1	PRINZIDE TABS	,	profotod drug(s) states.
<b>                                     </b>	MC/DEL	LISINOPRIL-HCTZ TABS	MC/DEL	4	UNIRETIC TABS	•	· [
11	MC/DEL	LOTENSIN HCT TABS	MC	1	VASERETIC TABS	•	
/ L	4		MC/DEL		ZESTORETIC TABS		
BETA BLOCKERS AND DIURETIC	MC/DEL	ATENOLOL/CHLORTHALIDONE	MC/DEL	1	CORZIDE 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
COMBO'S	MC/DEL	BISOPROLOL FUMARATE/HCTZ	MC/DEL	1	LOPRESSOR HCT TABS		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
11	MC/DEL	PROPRANOLOL/HCTZ	MC	1	TENORETIC	•	preferred drug(s) exists.
<b>                                     </b>	1 1	1	MC	1	TIMOLIDE 10/25 TABS	,	
//	11		MC/DEL	1	ZIAC TABS		
ARB'S AND CA CHANNEL BLOCKERS	MC/DEL	AMLODIPINE/VALSARTAN	MC/DEL	<i></i>	AZOR		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,
11	MC/DEL	AMLODIPINE/VALSARTAN HCT	MC	1	BYVALSON		propafenone, fluoxetine, paroxetine).
11	MC/DEL	TRIBENZOR	MC/DEL	1	EXFORGE	,	
41	1		MC/DEL	1	EXFORGE HCT	<b>.</b>	Per best practices patient should have trialed prior therapy of ACE inhibitor or currently on a diabetic therapy
			/	<i></i>		Use PA Form# 20420	
ARB'S AND DIURETICS	MC/DEL	BENICAR HCT <sup>1</sup>		7	IRBESARTAN HYDROCHLOROTHIAZIDE		Per best practices patient should have trialed prior therapy of ACE inhibitor or currently on a diabetic therapy
11	MC/DEL	LOSARTAN HCT <sup>1</sup>	MC/DEL	8	ATACAND HCT TABS	please see dose consolidation list.	
<b>                                     </b>	MC/DEL	MICARDIS HCT TABS <sup>1</sup>	MC	8	AVALIDE TABS <sup>1</sup>	COHSUMATION NOT.	
<b>                                     </b>	MC/DEL	VALSARTAN-HCT <sup>1</sup>	MC/DEL	8	DIOVAN HCT TABS <sup>1</sup>	,	
11	1 1		MC/DEL	8	HYZAAR TABS	•	
	4		MC	8	TEVETEN HCT_TABS	<u>Use PA Form# 20420</u>	
ANGIOTENSIN MODULATORS-ARB COMBINATION	MC	ENTRESTO	MC/DEL	1	EDARBYCLOR		
	4		MC		ENTRESTO SPRINKLES	Use PA Form# 20420	
ARB'S AND DIRECT RENIN INHIBITOR COMBINATION			MC/DEL	1	VALTURNA	Use PA Form# 20420	
DIURETICS	MOIDEL	ACETAZOLAMIDE TABS	MOIDEL		TARA TARA		The second secon
DIURETICS	MC/DEL MC/DEL	ACETAZOLAMIDE TABS BUMETANIDE	MC/DEL MC/DEL	1	ALDACTAZIDE TABS ALDACTONE 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
41	MC/DEL MC/DEL	CHLOROTHIAZIDE TABS	MC/DEL		AMILORIDE HCL		preferred drug(s) exists.
<b>/</b>	MC/DEL	CHLUKUTHIAZIDE TADS	WIC/DEL		AMILORIDE HCL	l '	,

	MC/DEL MC MC/DEL	CHLORTHALIDONE TABS EDECRIN TABS EDECRIN TABS HYDROCHLOROTHIAZIDE INDAPAMIDE TABS METHAZOLAMIDE TABS METHYCLOTHIAZIDE TABS SPIRONOLACTONE SPIRONOLACTONE TORSEMIDE TABS TRIAMTERENE/HCTZ ZAROXOLYN TABS	MC/DEL MC/DEL MC/DEL MC MC MC MC MC MC MC MC MC MC/DEL		BUMEX TABS DEMADEX TABS DIAMOX DIURIL DYAZIDE CAPS CAROSPIR ENDURON TABS FUROSCIX INSPRA KERENDIA KEVEYIS LASIX TABS MAXZIDE MICROZIDE CAPS MIDAMOR TABS NAQUA TABS	<u>Use PA Form# 20420</u>	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 Keveyis® with high dose aspirin is contraindicated.  Kerendia: Patient must be on max tolerated preferred ACE-I/ARB and SGLT-2
CCB / LIPID	<del></del>		MC/DEL		CADUET	Use PA Form# 20420	<del> </del>
		NEUROGENIC ORTHOSTATIC HYPOTE			ONDOCI	OSCI AT OTHER 20420	
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			1		
CHOLESTEROL - BILE SEQUESTRANTS	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/DEL MC MC/DEL MC MC/DEL MC MC/DEL MC MC/DEL 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 <sup>1</sup>	MC MC/DEL MC/DEL MC MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL		EZALLOR SPRINKLES <sup>3</sup> FLOLIPID LIPITOR LIPTRUZET ZOCOR	Dosing limits apply, please see dosage consolidation list.      Current users grandfathered.      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	<b>EZETIMIBE TABS</b> LOVASTATIN TABS <sup>2</sup> PRAVASTATIN <sup>2</sup>		8	ALTOPREV TB24 FLUVASTATIN TAB ER LESCOL XL TB24	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.

							<b>,</b>
1	I	I	MC	8	LIVALO	I	1
•	l [		MC/DEL	8	MEVACOR TABS	1	DDI: Lescol will now be non-preferred and require prior authorization if it is currently being used in combination with diclofenac.
•	] [		MC	8	NEXLETOL	1	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.
, <b>I</b>	] [		MC	8	NEXLIZET	1	1
, <b> </b>	l I		MC/DEL	8	PRAVACHOL TABS		1
1			MC/DEL	8	PRAVIGARD		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	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	MC	SIMCOR	MC		ADVICOR TBCR	Use PA Form# 20420	
INHIBITORS STATIN/ NIACIN COMBO							
FAMILIAL HYPERCHOLESTEROLEMIA	MC	PRALUENT (LABLER 72733) PEN1,2,3,3	MC	<del></del>	EVKEEZA <sup>1,4</sup>	Clinical PA required for	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
Pumbirth	MC	REPATHA <sup>1,2,3</sup>	MC	1	JUXTAPID	appropriate diagnosis	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
<b>.</b>	IVIC	INLI ATTIA	MC MC	1	JUXTAPID KYNAMRO <sup>1</sup>	Quantity limits apply	preferred drug(s) exists
	] [		MC	1	LEQVIO	Qualitity liftilis apply     Documented adherence to	10
			IVIC	1	LEQVIO	lipid lowering medications	
/ <b> </b>			ļ	1		and abstinence from tobacco	Juxtapid is contraindicated with strong CYP3A4 inhibitors. Juxtapid dosage should not exceed 30mg daily when it is used concomitantly with weak CYP3A4 inhibitors.
				1		for previous 90 days	"/
/				1		4 F4- 4-patment of	Kynamro requires an appropriate lab testing prior to starting (ALT <ast), alkaline="" and="" bilirubin,="" every="" first="" for="" liver-related="" monthly="" months.<="" phosphatase="" td="" tests="" the="" then="" three="" total="" year,=""></ast),>
/				1		<ol> <li>For the treatment of patients ≥ 12 years of age.</li> </ol>	
			l l	1		pallerits ≤ 12 years or ago.	
<b>[ ]</b>			l l	1			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
<b>                                     </b>				1		5.Approval of Praluent	HIDIE HAXIIIUIII (Dielaleu uuse oi staniis (one oi winori must be atorvastanii oi rosuvastanii) aha ezenimbe romg uung
<b>                                     </b>				1		NDC's with labeler code	
				1		00024 will be considered only if labeler code 72733	
11				1		NDC's are on a long-term	
11				1		backorder and unavailable	
41				1		from the manufacturer.	
41 '				1			
41			ļ	1			
41			ļ	1			
41				1			
<b>                                     </b>			ļ	1			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
<b>                                     </b>			ļ	1			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.
<b>                                     </b>			ļ	1			of the following 11000100 of an activities a
/ /			ļ	1			
<b>                                     </b>				1			Additional criteria for the diagnosis of clinical atherosclerotic cardiovascular disease: History of MI, angina, coronary or other arterial revascularization, stroke, TIA, or PVD of
/			ļ	1			atherosclerotic origin.
11			ļ	1			
41			ļ	1			Additional criteria for the diagnosis of homozygous familial hypercholesterolemia (Repatha only): Total cholesterol levels > 290mg/dL or LDL-C > 190mg/dL (adults) OR Total
41 '			ļ	1			cholesterol levels > 260mg/dL or LDL-C > 155mg/dL (children < 16 years) and TG within reference range OR Confirmation of diagnosis by gene testing.
41			ļ	1			
41 '			ļ	1			· [
41 '				1		Use PA Form# 20420	·]
		PULMONARY ANTI-HYPER	RTENSIVES				
PULMONARY ANTI-HYPERTENSIVES	MC	EPOPROSTENOL INJ <sup>3,6</sup>	MC/DEL		ADEMPAS <sup>1,3</sup>	Requires previous	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
41 '	MC/DEL	SILDENAFIL	MC	1	ADCIRCA <sup>4</sup>	trials/failure of multiple	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
41 '	MC/DEL	TADALAFIL	MC/DEL	1	ALYQ TAB	preferred medications.	preferred drug(s) exists.
41 '	MC	VENTAVIS <sup>3</sup>	MC	1		Dosing limits apply,	
41 '	iii C	VENTAVIS		1	FLOLAN <sup>3</sup> LIQREV	please see the dose	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
41 '			MC	1	CIQREV  OPSUMIT <sup>1,2</sup>	consolidation list.	concomitant use of Sildenafil with moderate or strong Cyp3A inhibitors
41 '			MC	1			
41 '			MC	1	OPSYNVI <sup>4</sup>	3.Require WHO Group 1	DDI: Uptravi will require a prior authorization if it is currently being used in combination with strong inhibitors of CYP2C8 (gemfibrozil)
41 '			MC	1	ORENITRAM	diagnosis of primary PAH (Primary Pulmonary	
41 '			MC	1	REMODULIN <sup>3</sup>	Hypertension) and NYHA	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,
41 '			MC/DEL	1	REVATIO <sup>4</sup>	functional class 3 or 4.	indinavir, nefazodone, nelfinavir, ritonavir, atazanavir, saquinavir and telithromycin).
41 '			MC	1	TADLIQ⁴		
41 '			MC	1	TYVASO	4.Require WHO Group 1	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 dipyridamole, adcira and
<b>/ '</b>		Ī		4		•	DDI. Adempas will require a prior adminization in it is currently being used in combination with drugs known to be 1 DE minibilities should be avoided (including dipyridamole, addit a and

1	]	<i>i</i> I	MC	1	UPTRAVI	diagnosis of primary PAH	tadalafil) with adempas
	]	, I	MC	1	VELVETRi <sup>3</sup>	(Primary Pulmonary	
1	]	, I	MC/DEL	1	WINREVAIR <sup>4</sup>	Hypertension) and NYHA (WHO) functional class 2 or	Ligrev: treatment of pulmonary arterial hypertension (WHO Group 1) in adults to improve exercise ability and delay clinical worsening. Avoid concomitant use of Ligrev with moderate or
		,	4 1	1		3.	strong CYP3A inhibitors.
, <b> </b>	]	,	1 1	1			
		, <u> </u>	11	<i>1</i> '		Use PA Form# 20420	
ERA / ENDOTHELIN RECEPTOR	MC	LETAIRIS <sup>1,2</sup>	i			Providers must be	Tracleer approvals will require WHO Group 1 diagnosis of primary PAH (Primary Pulmonary Hypertension) and NYHA functional class 2 thru 4.
ANTAGONIST	MC	TRACLEER	1 1	1		registered with LEAP	
, <b> </b>	l j	,	1 1	1		Prescribing program, a restricted distribution	DDI: Preferred Oral Contraceptives will now be non-preferred and require prior authorization if it is currently being used in combination with Tracleer.
, [	l l	,	1 1	1		program.	
1	l j	,	( J	4			
	l l	,	( )	4		Clinical PA is required to     setablish diagnosis and	Letairis approvals will require WHO Group 1 diagnosis of primary PAH (Primary Pulmonary Hypertension) and functional class 2 or 3 symptoms.
	l l	,	( )	4		establish diagnosis and medical necessity.	
	l l	,	( )	4		modisal needed ,	
1	l l	,	( )	4			
11	]	, I	4 1	1		U DA C# 20420	
	Щ	WROTENOE ACENTS			<u> </u>	Use PA Form# 20420	<u></u>
		IMPOTENCE AGENTS				f l 1 2006 per	• · · · · · · · · · · · · · · · · · · ·
IMPOTENCE AGENTS	l l	,	( )	4		As of January 1, 2006, per CMS (federal govt.),	As of January 1, 2006, per CMS (federal govt.), impotence agents are no longer covered.
<b>/                                    </b>	l j	,	1 1	1		impotence agents are no	1
1	l l	,	( )	4		longer covered.	
					<u> </u>		<u> </u>
ANTIQUAL INFRAIG		ANTI-EMETOGENICS	- "				The second black of the se
ANTIEMETIC - ANTICHOLINERGIC / DOPAMINERGIC	MC	DOXYLAMINE SUCC-PYRIDOXINE HCL	MC	4	ANTIVERT 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
DOF AMIRE NOIS	MC/DEL	MECLIZINE HCL TABS	MC	1	BARHEMSYS		preferred drug(s) exists.
<b>/                                    </b>	MC	PROMETHAZINE SUPP	MC	1	BONJESTA		
<b>/                                    </b>	MC/DEL	PROMETHAZINE	MC	1	DICLEGIS		1
11	MC	TRANSDERM-SCOP PT72	MC	1	PHENERGAN SOLN		
	l j	,	MC	4	PROMETHECAN SURP		
	l l	,	MC MC	1	PROMETHEGAN SUPP TORECAN TABS		DDI: Concomitant use of MAOIs and Bonjesta® is contraindicated.
ANTIEMETIC - 5-HT3 RECEPTOR	MC/DEL	DRONABINOL CAPS	MC	8	AKYNZEO'	Approvals will require	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
ANTAGONISTS/ SUBSTANCE P	MC/DEL MC/DEL	GRANISETRON TAB	MC	8	APREPITANT	diagnosis of chemo-induced	Preferred drugs and step therapy must be tried and falled due to lack of efficacy of infolerable 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
NEUROKININ	MC/DEL	ONDANSETRON TAB	MC	8	ALOXI	nausea/vomiting and failed	another drug and the preferred drug(s) exists. * Ondansetron limits still apply as listed on the Ondansetron PA form for covered indications including chemotherapy, radiotherapy, post
<b>/                                    </b>	MC/DEL	ONDANSETRON ODT TBDP	MC	8	ANZEMET TABS	trials of all preferred anti-	operative nausea & vomiting and hyperemesis gravidarum. Other medical indications will be approved or denied on a case by case basis. Hyperemesis and other medical indications
<b>/                                    </b>	MC/DEL	ONDANSETRON SOL	MC	8	APONVIE <sup>4</sup>	emetics, including 5-HT3 class (Ondansetron) and	approved are still subject to failure of multiple preferred antiemesis drugs.
	MO/DII	J. J	MC	8	CESAMET <sup>1</sup>	Marinol.	1
	l l	,	MC	8	CINVANTI <sup>4</sup>		1
1	l j	, l	MC	8	EMEND <sup>2</sup>	1	Akynzeo- Concomitant use should be avoided in patients who are chronically using a strong CYP3A inducer such as rifampin.
<b>/                                    </b>	l j	, l	MC	8	FOCINVEZ <sup>1,2</sup>		Akylized- Collociticalit use stitutione avoluted in patients who are chronically using a strong of the stranger scenarios manipuls.
	l l	,	MC/DEL	8	KYTRIL	Clinical PA is required for	Varubi – Available to the few who are unable to tolerate or who have failed on preferred medications
	l l	,	MC/DEL	8	MARINOL CAPS	members on highly emetic	Valuation - Artificiation to this form white and animate to testing of the first testing of t
<b>/                                    </b>	l j	, l	MC	8	SANCUSO		Aponvie is for the prevention of postoperative nausea and vomiting (PONV) in adults.
<b>/                                    </b>	l j	, l	MC	8	SUSTOL		reputition to the provention of protoporative madeda and retimining ( ) in addition.
	l j	,	MC	8	SYNDROS	Dosing limits apply,	
1	l j	, l	MC	8	TRIMETHOBENZAMIDE CAP	please see Dosage	
1	l j	, l	MC	8	VARUBI	Consolidation List	
	l l	,	MC/DEL	8	ZOFRAN ODT TBDP <sup>3</sup>	4. Clinical PA required for	
1	l j	, l	MC/DEL	8	ZOFRAN TABS <sup>3</sup>	appropriate diagnosis	
<b>/                                    </b>	l j	, l	MC/DEL	8	ZOFRAN INJ <sup>3</sup>		
/	]	, I	MC	8	ZUPLENZ		
<b>/                                    </b>	]	,	1 1	1		Use PA Form# 20420	
		NON-SEDATING ANTIHISTAMINES / DECONGE	ESTANTS			550	
ANTIHISTIMINES - NON-SEDATING	MC	ALAVERT TABS	MC	5	CLARINEX TABS <sup>1,5</sup>	Must fail preferred drugs,	Preferred drug 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
	MC/DEL	CETIRIZINE TABS	MC		CLARINEX TABS CLARINEX SYR <sup>1,2</sup>	OTC loratadine and	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
<b>/</b> "		•		- 1	OD WINES. STA	antirizina hafara mavina ta	and the preferred develop exists. No combination product with deconsected will be consequed since accordance-belief existiable without DA

	MC/DEL MC	LORATADINE TAVIST ND (OTC)	MC/DEL MC/DEL MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL	5 5 8 8 8	FEXOFENADINE <sup>1</sup> ZYRTEC <sup>1</sup> ZYRTEC SYR <sup>1,2</sup> ALLEGRA <sup>3</sup> CLARITIN <sup>3</sup> DESLORATADIN LORATADINE ODT <sup>4</sup> LEVOCETIRIZINE <sup>4</sup> XYZAL <sup>3</sup>	non-preferred step order drugs.	
			11	<i></i> '		Use PA Form# 20530	
ANTIHISTIMINES - OTHER	MC/DEL	CLEMASTINE				Use PA Form# 20530	
	MC/DEL	CHLORPHENIRAMINE	1 1	1		'	
	MC/DEL	DIPHENHYDRAMINE			<u> </u>		<u></u>
ANAPHYLACTIC DEVICES	MC/DEL	ALLERGY / ASTHMA THERAPIES  EPINEPHRINE			AUVI- Q	1	
ANAPHTLACTIC DEVICES	MC/DEL	EPINEPHRINE EPIPEN	MC MC		NEFFY	·	
<b>[ ]</b>	MC/DEL	EPIPEN JR	MC	1	TWINJECT		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.
ALLERGEN IMMUNOTHERAPY			MC	1 '	ODACTRA ODALARI		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
		, <u>'</u>	MC		ORALAIR <sup>1</sup>		therapy is being chosen over subcutaneous therapy
		, <u>'</u>	MC		PALFORZIA	See criteria section	
		!	MC MC		RAGWITEK GRASTEK		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.
		!		1			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.
							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
		!		1			Oralair: Patient age ≥10 years and ≤65 years
	l			l'			Have an auto-injectable epinephrine on-hand
ANTIASTHMATIC - ANTICHOLINERGICS - INHALER	MC MC/DEL MC/DEL	INCRUSE ELLIPTA <sup>3</sup> SPIRIVA HANDIHALER <sup>1,2</sup> SPIRIVA RESPIMAT	MC MC/DEL		LONHALA MAGNAIR TUDORZA		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 - PHOSPHODIESTERASE 4 INHIBITORS	MC/DEL	ROFLUMILAST	MC/DEL MC		DALIRESP OHTUVAYRE <sup>1</sup>		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		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 <sup>2,4</sup> FASENRA <sup>2</sup> FASENRA <sup>2</sup> AUTO INJCT  XOLAIR <sup>1</sup>	MC MC		CINQAIR <sup>3</sup> NUCALA <sup>2</sup> TEZSPIRE <sup>5</sup>	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	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 MC/DEL MC/DEL MC/DEL	BUDESONIDE SPRAY  FLUTICASONE SPR <sup>3</sup> OLOPATADINE SPRAY  OMNARIS SPR <sup>3</sup> TRIAMCINOLONE NS  QNASL	MC MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC MC/DEL	8 8 8 8 8 8 8	BECONASE AQ INHA <sup>1,3</sup> DYMISTA FLONASE SUSP <sup>2,3</sup> FLUNISOLIDE SOLN <sup>1,3</sup> NASONEX SUSP RHINOCORT AERO <sup>2,3</sup> RHINOCORT AQUA SUSP <sup>2,3</sup> RYALTRIS <sup>4</sup> TRI-NASAL SOLN <sup>2,3</sup> VANCENASE POCKETHALER AERS <sup>2,3</sup> VERAMYST <sup>2,3</sup> XHANCE <sup>2</sup> ZETONNA <sup>3</sup>	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 the step 9th process.	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 <sup>1</sup>	MC/DEL MC/DEL		ASTEPRO <sup>2</sup> PATANASE	approved if submitted with	Approved if patient fails on nonsedating 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.

		_		_		
	1				O LUCE- MARKET	
1	1				<ol> <li>Utilize Multiple preferred, as well as step therapy</li> </ol>	
	l				Azelastine.	
ANTIASTHMATIC - BETA - ADRENERGICS	MC/DEL	ALBUTEROL NEB	MC/DEL	ACCUNEB NEBU	1. Xopenex users w/ prior	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/DEL	ALBUTEROL HFA (Teva labeler 00093 AND	MC/DEL	ALBUTEROL HFA	asthma hospitalization due	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
1	<i>i</i> 1	Sandoz 00781)	1 1			preferred drug(s) exists.
	MC	LEVALBUTEROL TARTRATE	MC/DEL	BRETHINE	will be grandfathered.	
1	MC/DEL	METAPROTERENOL	MC/DEL	PROAIR DIGIHALER⁴	ļ ,	
1	MC	PROAIR RESPICLICK	MC	VOLMAX TBCR	ļ ,	
	MC/DEL	PROVENTIL HFA	MC	VOSPIRE ER TB12	2. Quantity Limit: 12 cc/day.	
	MC	SEREVENT	MC	XOPENEX HFA <sup>3</sup>		
1	MC/DEL	STRIVERDI	MC	XOPENEX NEBU <sup>1,2</sup>	<ol><li>Dosing limits apply,</li></ol>	
1	MC/DEL	TERBUTALINE SULFATE TABS	11.0	NOTENEX NEED	please see dosage	
1	MC/DEL	ALBUTEROL 0.63mg/3ml	1 1		consolidation list.	
1	MC	VENTOLIN HFA AERS	1 1		4. For the treatment of	
. I	mo		1 1		patients ≥ 4 years of age.	
/ <b> </b>	<i>i</i> 1	1	1 1		<u> </u>	
	<i>l</i>	l	1 1			
1	<i>i</i> 1	1	1 1			
1	<i>i</i>	l	1 1		Use PA Form# 20420	
ANTIASTHMATIC - ADRENERGIC	MC	and a province	<del>+ + .</del>	AIRDUO DIGIHALER <sup>2</sup>		5 ( - 1 lives with a first and failed due to lock of efficiency or intolerable cide effects before non preferred drugs will be approved unless an acceptable clinical exception is offered on
ANTIASTHMATIC - ADRENERGIC COMBINATIONS	MC MC	ADVAIR DISKUS <sup>1</sup>	MC/DEL		Dosing limits apply, please see dosage	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
OG MENTATIONS		ADVAIR HFA <sup>1</sup>	MC/DEL	AIRSUPRA		preferred drug(s) exists.
	MC MC	AIRDUO RESPICLICK <sup>2</sup>	MC/DEL	BREZTRI AEROSPHERE	0.5	
11 7		BREO ELLIPTA <sup>1</sup>	MC	TRELEGY ELLIPTA <sup>1</sup>	<ol> <li>For patients ≥ 12 years and older.</li> </ol>	
11 7	MC/DEL	DULERA				
11 /	MC/DEL	FLUTICASONE-SALMETEROL			<u> </u>	AirDuo® Respiclick be non-preferred and require prior authorization and be available to those who are unable to tolerate or who have failed on preferred medications
11 7	MC/DEL	SYMBICORT			,	
11 7	l I				,	
11 7	l I				,	
11 ,	i					DDI: Avoid concomitant use of strong CYP3A4 inhibitors (e.g. ritonavir, atazanavir, clarithromycin, indinavir, itraconazole, nefazodone, nelfinavir, saquinavir, ketoconazole, telithromycin)
<b> </b>	l I				, , , , , , , , , , , , , , , , , , ,	with AirDuo® Respiclick is not recommended due to increased systemic corticosteroid and increased cardiovascular adverse effects
<b> </b>	l I				,	
<b> </b>	l I				,	
<b> </b>	l I				,	
11 7	l I				,	
11 7	l I				<u> </u>	
<b> </b>			<del></del>		Use PA Form# 20420	
ANTIASTHMATIC - ADRENERGIC ANTICHOLINERGIC	MC/DEL	ALBUTEROL/IPRATROPIUM NEB. SOLN	MC/DEL	BEVESPI AEROSPHERE <sup>2,3</sup>	Please use preferred individual ingredients	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
ANTICHOLINERGIC	MC	ANORO ELLIPTA	MC/DEL	DUAKLIR PRESSAIR		preferred drug(s) exists. DuoNeb components are available separately without PA.
11 /	MC/DEL	COMBIVENT RESPIMAT	MC/DEL	DUONEB SOLN <sup>1</sup>		provided druggly oxidite. Somponente de dramatie sopration, minister i in
11 /	MC/DEL	STIOLTO			2. Dosing limits apply,	
11 /	l I					DDI: Avoid concomitant use of Bevespi with other anticholinergic-containing drugs, due to an increased risk of anticholinergic adverse events. Bevespi® should be used with extreme
11 /	1 1		1 1			caution in patients being treated with MAO inhibitors, TCAs, or other drugs known to prolong the QTc interval.
11 /	l I				3. The safety and efficacy of	
11 7	l I				use in children under the age of 18 years have not been	
11 7	l I					Bevespi should be used with extreme caution in patients being treated with MAO inhibitors, TCAs, or other drugs known to prolong the QTc interval.
11 ,	i				,	
11 7	l I				,	
11 7	l I				Use PA Form# 20420	
ANTIASTHMATIC - XANTHINES	MC/DEL	AMINOPHYLLINE TABS	MC/DEL	THEO-24 CP24	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
ANTIAGITIMATIO - AARTTIMEG	MC/DEL	THEOCHRON TB12	MC	THEO-24 GF24 THEOLAIR TABS		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
41 /	MC/DEL	THEOLAIR-SR TB12	MC/DEL	UNIPHYL TBCR		preferred drug(s) exists.
41 /	MC/DEL	THEOPHYLLINE CR TB12	MO/DEL	ON THE TOOK	, l	
11 /	MC	THEOPHYLLINE ELIX			,	
<b>4</b> 1	/ "" I	THEOTHELINE LEIA	1 1	<u>I</u>	ı ,	<b>.</b>

1_		_	-	_	_	-	<u> </u>
1 ,	MC/DEL	THEOPHYLLINE SOLN		1	1	'	
1	MC/DEL	THEOPHYLLINE ER CP12	l ,	1		<u> </u>	·
	MC/DEL	THEOPHYLLINE ER TB12	'			'	
ANTIASTHMATIC - STEROID INHALANTS	MC	ARNUITY ELLIPTA	MC	8	AEROSPAN		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
1	MC/DEL	ASMANEX TWISTHALER 3,4	MC/DEL	8	ALVESCO <sup>3</sup>	& 0.5mg will be preterred for	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
	MC/DEL	ASMANEX HFA <sup>5</sup>	MC	8	ARMONAIR DIGIHALER	members under the age of 8 years old. PA will be	preferred drug(s) exists.
'	MC/DEL	BUDESONIDE NEB 0.25MG & 0.5MG <sup>1</sup>	MC/DEL	8	BUDESONIDE NEB 1MG	required for members 8	1
	MC/DEL	PULMICORT FLEXHALER 3	MC/DEL	8	PULMICORT SUSP	years of age and older,	1
'	MC	QVAR AERS <sup>3</sup>	_ <u> </u>	1		please consider other	1
'	1 1		_ <u> </u>	1		preferred options.	1
.	1 1		_ <u> </u>	1		<u>'</u>	1
'	1	1	_ <u> </u>	1		<b> </b>	· ·
	1 1		_ <u> </u>	1		2. All preferred must be tried	A
	1 1		ļ	1		before moving to non	1
	1 1		_ <u> </u>	1		preferred steps.	1
'	1 1		_ <u> </u>	1		<u> </u>	1
	1 1		_ <u> </u>	1		3. Dosing limits apply,	1
	1 1			1		please see dosage consolidation list.	1
	1 1			1		CONSONUATION IIST.	1
. [	1 1			1		4. Asmanex 110mcg will be	1
	1 1			1		limited to member between	1
	1 1			1		the ages of 4-11years old.	1
	1 1			1		<u> </u>	1
1	1	1	_ <u> </u>	1		<b> </b>	
, <b>[</b>	1 1			1		<u> </u>	1
/ l	1 1			1		5. Asmanex HFA will be	1
/ <b> </b>	1 1			1		preferred for members under	.1
/ <b> </b>	1 1			1		the age of 6 years old. PA	1
<b> </b>	1 1			1		will be required for members	A
<b> </b>	1 1			1		6 years of age and older,	1
<b> </b>	1 1			1		please consider other	1
<b> </b>	1 1			1		preferred options.	1
<b> </b>	1 1			1		<u> </u>	1
	<u>1L</u>		,			Use PA Form# 20420	
ANTIASTHMATIC - 5-Lipoxygenase			MC	1	ZYFLO CR TABS		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
Inhibitors	1	1	_ <u> </u>	1			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
	1 <u></u>	<u></u>		1 `	<u></u>	Use PA Form# 20420	between another drug and the preferred drug(s) exists.
ANTIASTHMATIC - LEUKOTRIENE	MC/DEL	MONTELUKAST GRANULE <sup>1</sup>	MC/DEL	8	ACCOLATE TABS		
RECEPTOR ANTAGONISTS	1 1			1		Use PA Form# 20420	1
/ <b> </b>	MC/DEL	MONTELUKAST SODIUM TAB	MC/DEL	8	SINGULAIR <sup>2</sup>	1.Montelukast Granules will	1
/ <b> </b>	MC/DEL	MONTELUKAST SODIUM CHEW TAB	MC/DEL		SINGULAIR GRANULES	only be approved if between	1
/	1		mo, see	1	ONGOLAN GRANGLEG	ages of 6months-24 months.	
/ <b> </b>	1 1			1		<u> </u>	1
<b> </b>	1 1			1		0.01   1.00	1
/ <b> </b>	1 1			1		2.Singulair Chewable 4mg from 2years-5years and	1
/	1 1			1		Singulair Chewable 5mgs	1
<b> </b>	1 1			1		from 6years-14years old.	1
<b> </b>	1 1		ļ	1			1
<b> </b>	1 1			1		<u> </u>	1
<b> </b>	1 1			1		<u> </u>	1
<b> </b>	1 1		ļ	1		<b>,</b>	1
<b> </b>	1 1			1		<u> </u>	1
ANTIASTHMATIC - ALPHA-PROTEINASE			+	<del></del>	TADALACT	II DA 5# 00400	Prolastin and Azemaira will be approved for members with A1AT deficiency and clinically demonstrable panacinar emphysema.
INHIBITOR	1 1		MC		ARALAST	Use PA Form# 20420	Profastin and Azemaira will be approved for Thembers with ATAT deliciency and clinically definitisticable particular emphysienta.
INTIBATION.	1 1		MC/DEL		ZEMAIRA	<b>,</b>	1
<b> </b>	1 1		MC		GLASSIA	<b>,</b>	1
<u> </u>	1 <u></u>	<u></u>	MC	8	PROLASTIN SUSR	'	
ANTIACTUMATIC HYDDO LYTIC				-	DULMOZVME COLNI		Will be approved for quetic fibracia nationts
ANTIASTHMATIC - HYDRO-LYTIC			MC/DEL	١	PULMOZYME SOLN		Will be approved for cystic fibrosis patients.
ENZYMES	Ţ		MC/DEL	•	POLMOZYME SOLN	Use PA Form# 20420	will be approved for cystic fibrosis patients.

ANTIASTHMATIC - MUCOLYTICS	MC/DEL	ACETYLCYSTEINE <sup>1</sup>	MC	MUCOMYST	Acetylcysteine is covered	
					with diagnosis of CF.	
					Use PA Form# 20420_	
ANTIASTHMATIC-CFTR POTENTIATOR AND COMBINATIONS			MC	ALYFTREK	For the treatment of	Alfytrek will be considered for the treatment of patients 6 years and older with at least one responsive mutation, including 31 additional mutations not responsive to other CFTR modulator
AND COMBINATIONS			MC	BRONCHITOL <sup>1</sup>	patients ≥18 years of age	therapies
			MC	KALYDECO	with CF.	Bronchitol will be considered as add-on maintenance therapy to improve pulmonary function in adult patients 18 years and older with cystic fibrosis (CF). Use Bronchitol® only for adults
			MC	ORKAMBI		who have passed the Bronchitol® Tolerance Test (BTT). (see Recommended Dosage section for further information
			MC	SYMDEKO		Kalydeco will be considered for patients with cystic fibrosis (CF) aged 1 month and older who have at least one mutation in the CFTR gene that is responsive to ivacaftor potentiation
		MC/DEL	TRIKAFTA		based on clinical and/or in vitro assay data. If the patient's genotype is unknown, an FDA-cleared CF mutation test should be used to detect the presence of a CFTR mutation followed by	
						verification with bi-directional sequencing when recommended by the mutation test instructions for use.
						Orkambi will be considered for patients with cystic fibrosis (CF) aged 1 year and older who are homozygous for the F508del mutation in the CFTR gene. If the patient's genotype is
						unknown, an FDA-cleared CF mutation test should be used to detect the presence of the F508del mutation on both alleles of the CFTR gene. The efficacy and safety of Orkambi have not
						been established in patients with CF other than those homozygous for the F508del mutation.
						Symdeko will be considered for patients with cystic fibrosis (CF) aged 6 years and older who are homozygous for the F508del mutation or who have at least one mutation in the cystic
						fibrosis transmembrane conductance regulator (CFTR) gene that is responsive to tezacaftor/ivacaftor based on in vitro data and/or clinical evidence. If the patient's genotype is unknown,
						an FDA-cleared CF mutation test should be used to detect the presence of a CFTR mutation followed by verification with bi-directional sequencing when recommended by the mutation test instructions for use.
						Trikafta will be considered for the treatment of cystic fibrosis (CF) in patients aged 2 years and older who have at least one F508del mutation in the cystic fibrosis transmembrane
						conductance regulator (CFTR) gene or mutation in the CFTE gene that is responsive based on in vitro data. If the patient's genotype is unknown, an FDA-cleared CF mutation test should
						be used to confirm the presence of at least one F508del mutation or a mutation that is responsive based on in vitro data.
					Use PA Form# 20420	
IDIOPATHIC PULMONARY FIBROSIS	MC/DEL	OFEV <sup>1</sup>	MC	ESBRIET <sup>1</sup>	1. Diagnosis required	
			MC	PIRFENIDONE		Ofev- Avoid concomitant use with P-gp and CYPA4 inducers (e.g. carbamazepine, phenytoin, and St. John's wort
						Esbriet- The concomitant use with strong CYP1A2 inhibitors (e.g. fluvoxamine, enoxacin) is not recommended
					Use PA Form# 20420	
		COUGH/COLD				
COUGH/COLD	MC/DEL	DEXTROMETHORPHAN CAPS <sup>1</sup>			1. All of cough cold	All non-preferred products are not covered as permitted by Federal Medicaid regulations and MaineCare Policy.
	MC/DEL	DEXTRO-GUAIF SYRP <sup>1</sup>			preparations are not covered	
	MC/DEL	GUAIFENESIN SYRP <sup>1</sup>			except these preferred products.	
	MC/DEL	PSEUDOEPHEDRINE <sup>1</sup>			pioducis.	
	MC	ROBITUSSIN DM SYRP <sup>1</sup>				
	MC	ROBITUSSIN SUGAR FREE SYRP <sup>1</sup>			Use PA Form# 20420	
		DIGESTIVE AIDS / ASSORTED GI				
GI - ANTIPERISTALTIC AGENTS	MC/DEL	DIPHENOXYLATE	MC/DEL	LOFENE 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
	MC/DEL	DIPHENOXYLATE/ATROPINE	MC	LONOX TABS	030 1 A 1 0111# 20420	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
	MC/DEL	LOPERAMIDE HCL CAPS/LIQ	MC	MOTOFEN TABS		preferred drug(s) exists. Certain drugs require specific diagnoses for approval.
	MC/DEL	OPIUM TINCTURE TINC				
	MC	PAREGORIC TINC				
GI - ANTI-DIARRHEAL/ ANTACID - MISC.	MC	ATROPINE SULFATE SOLN	MC/DEL	BELLADONNA ALKALOIDS & OP	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
1	MC/DEL	BISMATROL	MC/DEL	BENTYL TABS	1.Dosing limits apply please	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
	MC/DEL	BISMUTH SUBSALICYLATE	MC/DEL	BENTYL SYRP	refer to Dose Consolidation	preferred drug(s) exists. Certain drugs require specific diagnoses for approval.
	MC/DEL	CALCIUM CARBONATE (ANTACID) CHEW	MC	CUVPOSA	List	
	MC/DEL	DICYCLOMINE HCL	MC	DARTISLA ODT <sup>2</sup>	2. It is not indicated as	
	MC/DEL	GLYCOPYRROLATE TABS	MC	ED-SPAZ	monotherapy for treatment of	
	MC/DEL	HYOSCYAMINE CAPS & TABS	MC	MYTESI <sup>1</sup>	peptic ulcer because	
	MC/DEL	HYOSCYAMINE SULFATE	MC/DEL	GLYCOPYRROLATE INJ	effectiveness in peptic ulcer healing has not been	
	MC/DEL	KAOPECTATE	MC	LEVSIN TABS	established.	
	MC/DEL	MAGNESIUM OXIDE TABS	MC	LEVSIN/SL SUBL		Preferred products that used to require diag codes still require diag codes unless indicated otherwise.
	MC/DEL	MAG-OX 400 TABS	MC	NULEV TBDP		
, <b>-</b>		•		•		•

1							
	MC/DEL	PAMINE 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.
			MC		OSCIMIN		
	MC/DEL	PROPANTHELINE BROMIDE TABS	MC		ROBINUL INJ		
	MC/DEL	SODIUM BICARBONATE TABS	MC		ROBINUL TABS		
	MC/DEL	TUMS					
OL DILE ACID	<b>.</b>		110		CUOLDAM		lating this side with the distribution of the common defeat (CCD) AND for this best of a surface of the common defeat (CDD)
GI- BILE ACID			MC		CHOLBAM		Indication of bile acid synthesis disorders due to single enzyme defects (SEDs) AND for adjunctive treatment of peroxisomal disorders (PDs)
GI- EOSINOPHILIC ESOPHAGITIS		FOLIN IAT				Use PA Form# 20420	
GI- EOSINOPHILIC ESOPHAGITIS	MC	EOHILIA <sup>1</sup>				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 (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
						<ol> <li>Approvals will not be longer than 12 weeks of</li> </ol>	another drug and the preferred drug(s) exists.
						treatment in adult and	
						pediatric patients 11 years o	
						age and older	Eohilia: Dietary modification, PPIs, and topical glucocorticoids are required as initial therapy.
GI - H2-ANTAGONISTS	MC	ACID REDUCER TABS	MC		AXID 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
	MC/DEL	CIMETIDINE	MC		AXID AR TABS	03C 1 A 1 0111# 20420	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
	MC/DEL	FAMOTIDINE	MC/DEL		NIZATIDINE CAPS		preferred drug(s) exists.
			MC/DEL		PEPCID		
			MC		PEPCID AC		DDI: Cimetidine will now be non-preferred and require prior authorization if it is currently being used with any sulfonylurea (except for glyburide).
			iii O				
							DDI: Cimetidine will require prior authorization if being used in combination with Plavix.
GI- IBAT INHIBITORS			MC		BYLVAY <sup>1,2</sup>	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		LIVMARLI <sup>1,2</sup>		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.
						1. For the treatment of	preferred drug(s) exists. Certain drugs require specific diagnoses for approval.
						patients ≥ 3months of age	
						Clinical PA required for     prepariete diagnosis	
						appropriate diagnosis	
GI - PROTON PUMP INHIBITOR	MC/DEL	OMEPRAZOLE CAPS <sup>2</sup>	MC/DEL	6	NEXIUM CPDR <sup>3</sup>	Prevacid Solutabs	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
	MC/DEL	PANTOPRAZOLE <sup>2</sup>	MC/DEL	6	NEXIUM SUS⁵	available without PA for	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
	MC/DEL	LANSOPRAZOLE CAPS <sup>2</sup>	MC	7	PRILOSEC OTC <sup>3</sup>	children less than 9 years	another drug and the preferred drug(s) exists.
		ENNOUT WAZOLE ON O	MC	7	ACIPHEX TBEC <sup>3</sup>	old.	
			MC/DEL	8	DEXILANT (KAPIDEX) <sup>2</sup>	Dosing limits apply,	Please refer to the PPI PA form for additional criteria on Non-Preferred PPIs
			MC	8	KONVOMEP <sup>2</sup>	please see dosage	Today to to 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
			MC	8	OMEPRAZOLE-SODIUM BICARBONATE CAPS	consolidation list.	
			MC	8	OMEPRAZOLE MAGNESIUM		DDI: Omeprazole will require prior authorization if being used in combination with Plavix.
			MC/DEL	8	PREVACID CPDR <sup>3</sup>	All preferred and step	DDI: Lansoprazole will require prior authorization if being used in combination with Plavix.
			MC/DEL	0	PREVACID COUTABS <sup>1,4</sup>	therapy must be tried and	
			MC/DEL	0	PRILOSEC CPDR	4. Payment for Prevacid	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.
				8	PROTONIX INJ	SoluTabs for patients 9 and	
			MC/DEL MC/DEL	8		older will be considered for	
			WIC/DEL	·	PROTONIX <sup>2</sup>	those patients who cannot	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.
				8	VOQUEZNA TABS	tolerate a preferred solid ora	al Control of the Con
						dosage form.	
						5 Novium que available	
						<ol> <li>Nexium sus available without PA if member is &lt; 12</li> </ol>	2
						yrs of age and ≤ 1 pack per	
						dav.	
						Use PA Form# 20720	
						036 I A I UIII# 20120	
GI - ULCER ANTI-INFECTIVE	MC	PYLERA	+ +		VOQUEZNA DUAL PAK	Use PA Form# 20420	<del> </del>
	MC	TALICIA			VOQUEZNA TRIPLE PAK	030 1 A 1 0111# 20420	
1			1 1		- OGOLLINI III LE ITAN	ı	

I	1 1	Ī	1 1	ı	ı	1
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.
GI - DIGESTIVE ENZYMES	MOIDEL			DEDITA/E	Use PA Form# 20420	
GI - DIGESTIVE ENZYMES	MC/DEL MC	CREON <sup>1</sup> ZENPEP <sup>1</sup>	MC/DEL MC/DEL MC/DEL	PERTZYE ULTRESA VIOKACE	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.	between another drug and the preferred drug(s) exists.  is ce
GI - ANTI - FLATULENTS / GI	MC/DEL	AMITIZA	MC	CEPHULAC SYRP		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
STIMULANTS	MC	CALULOSE SYRP	MC/DEL	INFANTS GAS RELIEF SUSP		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
, <b>[</b>	MC/DEL	CONSTULOSE SYRP	MC	GIMOTI SPRAY		preferred drug(s) exists. Certain drugs require specific diagnoses for approval.
, <b>.</b>	MC/DEL	ENULOSE SYRP	MC/DEL	REGLAN TABS		·
. [	MC	GASTROCROM CONC		11.001		·
. [	MC/DEL	GENERLAC SYRP				·
. [	MC/DEL	LACTULOSE SYRP				·
, ]	MC/DEL	METOCLOPRAMIDE HCL	l i			·
<b>,                                    </b>	1				Use PA Form# 20420	·   · · · · · · · · · · · · · · · · · ·
GI - INFLAMMATORY BOWEL AGENTS	MC	APRISO	MC/DEL	ASACOL 800MG HD	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
<b>4</b> I	MC/DEL	BALSALAZIDE	MC/DEL	AZULFIDINE EN-TABS TBEC		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
4 l	MC	MESALAMINE ENMA KIT	MC	AZULFIDINE TABS	1. Current users	preferred drug(s) exists.
<b>,  </b>	MC	PENTASA	MC	COLAZAL CAPS	grandfathered.	·
<b>4</b>	MC/DEL	SULFAZINE EC TBEC	MC/DEL	DELZICOL	2. Diagnosis required	·
<b>, I</b>	MC/DEL	SULFASALAZINE TABS	MC	DIPENTUM CAPS		·
<b>4</b>	1	1	MC	GIAZO		Giazo is only indicated for males, as the safety.efficacy for use in females has not been established. Prior trials of preferred products.
<b>,  </b>	1 1	1	MC/DEL	LIALDA TABS <sup>1</sup>		·
<b>4</b>	1	1	MC/DEL	MESALAMINE TAB		Uceris Rectal Foam or Tab- Concomitant use with CYP3A inhibitors (e.g. ketoconazole, itraconazole, ritonavir, indinavir, saquinavir, erythromycin, cyclosporine, and grapefruit juice)
<b>4</b>	1	1	MC/DEL	ROWASA ENEM		should be avoided. Verify prior trials and failures or intolerance of preferred treatments
<b>4</b>	1	1	MC	SFROWASA		
<b>4</b>	1	1	MC	UCERIS RECTAL FOAM <sup>2</sup>		·
<b>/</b>	1	1	MC	UCERIS TABS <sup>2</sup>		·
GI - IRRITABLE BOWEL SYNDROME		LOTRONEX TABS	<del></del>	VIBERZI		The state of the s
AGENTS	MC/DEL	LUTRONEA TABO	MC	VIBERZI	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.
GI- SHORT BOWL SYNDROME	<del></del>	<del></del>	MC	GATTEX	<del></del>	Gattex requires a diagnosis of adult SBS who are dependent on parenteral support. Appropriate colonoscopy and lab assessments 6months prior to starting
<b>/</b>  '					<u>Use PA Form #20420</u>	
GI- NASH			МС	REZDIFFRA		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
<b>∡</b>	1	1		1	<u>Use PA Form #20420</u>	
		MISCELLANEOUS GI				
GI - MISC.	MC/DEL	BISAC-EVAC SUPP	MC/DEL	ACTIGALL CAPS		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
<b>∡</b>	MC/DEL	BISACODYL	MC	BENEFIBER	FDA approved indication.	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
<b>∡</b>	MC	BISCOLAX SUPP	MC/DEL	CARAFATE	2. For the treatment of	preferred drug(s) exists. Certain drugs require specific diagnoses for approval.
<b>⊿</b> "		•		•	•	•

	MC	CINOBAC CAPS	MC/DEL	CLEARLAX POW	carcinoid syndrome diarrhea	
	MC/DEL	CITRATE OF MAGNESIA SOLN	MC/DEL	COLACE CAPS	in combination with	
	MC/DEL	CITRUCEL	MC	DIOCTO-C SYRP	somatostatin analog (SSA)	
	MC/DEL	CLENPIQ SOL	MC	DOC SOD /CAS CAP	therapy in adults	
	MC/DEL	COLYTE	MC	DOC-Q-LAX CAPS	inadequately controlled by SSA therapy	
	MC/DEL	DIOCTO SYRP	MC/DEL	DOCUSATE SODIUM/CAS CAPS	OOA thorapy	
	MC		MC/DEL MC/DEL	DOK PLUS	2	Linzess is preferred for adults as treatment of IBS-Constipation AND treatment of chronic idiopathic constipation in adults.
		DOCUSATE CALCIUM CAPS			For the treatment of Opioid Induced	
	MC/DEL	DOCUSATE SODIUM	MC/DEL	DULCOLAX SUPP	Constipation(OIC)	L
	MC/DEL	FIBER LAXATIVE TABS	MC	ENEMEEZ		Trulance should be avoided in pediatric patients less than 18 years of age.
	MC	FLEET	MC	FIBER CON TABS	Established users will be	
	MC/DEL	GENFIBER POWD	MC/DEL	FIBER-LAX TABS	grandfathered	
	MC/DEL	GLYCERIN	MC/DEL	GAVILYTE-H		lqirvo: 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
	MC	HIPREX TABS	MC	GOLYTELY SOLR		monotherapy in patients unable to tolerate UDCA. Continued approval for this indication may be contingent upon verification and description of clinical benefit in confirmatory trial(s).
	MC/DEL	KRISTALOSE PACK	MC	IBSRELA		
	MC/DEL	LINZESS	MC	IQIRVO		
	MC	MAALOX	MC/DEL	LINZESS 72mcg <sup>4</sup>		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
	MC/DEL	MILK OF MAGNESIA SUSP	MC	LIVDELZI		UDCA, or as monotherapy in patients unable to tolerate UDCA. Patients who do not have a diagnosis of decompensated cirrhosis.
	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 <sup>1</sup>		
	MC 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	SENOKOT GRAN	MC/DEL	SENEXON TABS		
	MC/DEL	SENOKOT SYRP	MC/DEL	SENOKOT TABS		
	MC/DEL	SENOKOT CHILDRENS SYRP	MC	SENOKOT S TABS		
	MC	SENOKOT 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	Use PA Form# 20420	
	MC	TRULANCE <sup>2</sup>	MC/DEL	SYMPROIC <sup>3</sup>		
	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 <sup>2</sup>		
		MISC. UROLOGICAL				
UROLOGICAL - MISC.	MC I	ACETIC ACID 0.25% SOLN	MC	CITRIC ACID/SODIUM CITRAT SOLN	Elmiron requires	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
UNULUGICAL - IVIISC.	MC	CYTRA-K SOLN	MC/DEL	CYTRA-2 SOLN		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
	MC				supportive testing.	preferred drug(s) exists.
	MC	FOSFOMYCIN (NDC 82036427401 ONLY)	MC/DEL	ELMIRON CAPS <sup>1</sup>		
	MC	K-PHOS MF TABS	MC	FURADANTIN SUSP	Use PA Form# 20420	
	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		1
	MC/DEL	PHENAZOPYRIDINE HCL TABS	MC	POTASSIUM CITRATE/CITRIC SOLN		1
	MC/DEL	PHENAZOPYRIDINE PLUS	MC/DEL	PYRIDIUM PLUS TABS		1
	MC	POT CITRATE TAB	MC	PYRIDIUM TABS		
	MC/DEL	PROSED/DS TABS	MC/DEL	RENACIDIN SOLN		
	MC	TRICITRATES SYRP	MC	UROCIT-K		
	MC/DEL	URELIEF PLUS	I			1
	MC	UREX TABS				1
		URISED TABS				
	MC/DEL	UROQID #2 TABS				
	MC/DEL	ONOQID#Z IABO				
1	l I	I	ı I	I		I

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		PHOSPHATE BINDERS					
PHOSPHATE BINDERS	MC/DEL	CALCIUM ACETATE CAP <sup>1</sup>	MC		AURYXIA <sup>1</sup>	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
	MC/DEL	FOSRENOL CHEW <sup>1</sup>	MC/DEL		CALCIUM ACETATE TAB <sup>1</sup>	<ol> <li>Diag required.</li> </ol>	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	MAGNEBIND - 400 <sup>1</sup>	MC/DEL		ELIPHOS <sup>1</sup>		and the prefered drug(s) exists.
	MC	PHOSLYRA <sup>1</sup>	MC/DEL		FOSRENOL PWDR <sup>1</sup>		
	MC/DEL	RENVELA <sup>1</sup>	MC		VELPHORO <sup>1</sup>		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
			MC		XPHOZAH		who are intolerant of any dose of phosphate binder therapy.
		INTRA-VAGINALS					
VAGINAL - ANTIBACTERIALS	MC/DEL	CLEOCIN CREA	MC/DEL		METROGEL VAGINAL GEL <sup>1</sup>	Dosing limits apply,	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
	MC/DEL	CLEOCIN SUPP	MC/DEL		VANDAZOLE	please see Dosage	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
	MC	CLINDESSE CREA	MC		XACIATO	Consolidation List.	and the preferred drug(s) exists.
	MC/DEL		IVIC		ACIATO		
		METRONIDAZOLE VAGINAL GEL <sup>1</sup>					
	MC/DEL	NUVESSA				H DA F# 00400	
VACINAL ANTI FUNCAL C	MO/DEL	OLOTPINAZOLE ODEA	Mo			Use PA Form# 20420	
VAGINAL - ANTI FUNGALS	MC/DEL	CLOTRIMAZOLE CREA	MC		AVC CREA		
	MC/DEL	CLOTRIMAZOLE-3 CREA	MC		CLOTRIMAZOLE 3 DAY CREA	Quantity limit: 1/script/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
	MC/DEL	GYNE-LOTRIMIN CREA	MC		GYNAZOLE-1 CREA	weeks	preferred drug(s) exists.
	MC	MICONAZOLE CREA	MC		GYNE-LOTRIMIN 3 TABS	Use PA Form# 20420	pretented utugija) exista.
	MC	MICONAZOLE 3 KIT CREA OTC	MC/DEL		MICONAZOLE 3 COMBO PACK KIT <sup>1</sup>		
	MC/DEL	MICONAZOLE 7 CREA	MC/DEL		MICONAZOLE 3 SUPP		DDI: Miconazole will require prior authorization if being used in combination with Warfarin.
	MC/DEL	MICONAZOLE NITRATE CREA	MC		TERAZOL 3 CREA		
	MC	NYSTATIN TABS	MC		TERAZOL 7 CREA		
	MC/DEL	TERCONAZOLE CREAM	MC/DEL		TERCONAZOLE SUPP		
	MC	VAGITROL					
	MC	V-R MICONAZOLE-7 CREA					
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
						H DA F# 00400	preferred drug(s) exists.
VAGINAL - ESTROGENS	MC/DEL	ESTRING RING	MC/DEL		FOTDAGE ODEA1	Use PA Form# 20420  1. Must fail all 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
VACINAL - ESTROGENS	MC/DEL				ESTRACE CREA <sup>1</sup>	products before non-	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
	MO/DEE	PREMARIN CREA	MC/DEL		VAGIFEM TABS <sup>1</sup>	preferred.	preferred drug(s) exists.
						H DA F# 00400	
VAGINAL - OTHER	MC/DEL	ACID JELLY GEL	MC		AMINO ACID CERVICAL CREA	Use PA Form# 20420	Designed draws must be bried and failed due to local of efficacy or intelegable side effects before an explanate draw will be approved unless a constable disingle expension in effects and
VAGINAL - OTHER			IVIC		AWIINO ACID CERVICAL CREA	<u>Use PA Form# 20420</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
	MC	ACI-JEL GEL					preferred drug(s) exists.
	MC	CERVICAL AMINO ACID CREA					<u> </u>
DDU	MOIDEL	BENIGN PROSTATIC HYPERPLASIA			ITI OMAY ODGA		
BPH	MC/DEL	DOXAZOSIN MESYLATE TABS	MC/DEL	5	FLOMAX CP24		s 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
	MC/DEL	FINASTERIDE <sup>1</sup> 5mg	MC/DEL	8	ALFUZOSIN	or read per day with out r A.	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
	MC/DEL	TERAZOSIN HCL CAPS	MC	8	AVODART <sup>2,4</sup>		presence of obstructive urinary outflow symptoms along with adequate trial of preferred Proscar.
	MC/DEL	TAMSULOSIN HCL	MC/DEL	8	CARDURA TABS⁴	Prior use of preferred	
			MC	8	ENTADFI <sup>5,6</sup>	agent prior to any approvals.	
			MC	8	JALYN <sup>3,4</sup>		
			MC/DEL	8	PROSCAR TABS⁴	<ol><li>Use of preferred</li></ol>	
			MC/DEL	8	RAPAFLO <sup>4</sup>	(tamsulosin and finasteride)	
						and (tamsulosin and non- preferred Avodart).	
						preferred Avodart).	
	1 1						
			MC/DEL	8	UROXATRAL <sup>4</sup>	<ol> <li>Non-preferred products</li> </ol>	
						must be used in specified	
						order.	
						<ol><li>Use of individual</li></ol>	
<del>-</del>		•			-		-

		1 1	1 1	<u>İ</u>	1	ingredients preterred (Finasteride and tadalafil).	I I
			1	I	1	,	
			1	I	1	<ol><li>Entadfi® is not recommended for more than</li></ol>	
			1 1	l	1	26 weeks	
			1 1	l	1		
			1 1	l	1		
			ll	ı <u></u>		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
	MC/DEL	CHLORDIAZEPOXIDE HCL CAPS	MC/DEL	8	ATIVAN		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	CLORAZEPATE DIPOTASSIUM TABS	MC MC/DEL	8	LOREEV XR		
	MC/DEL MC/DEL	DIAZEPAM LORAZEPAM	MC/DEL MC/DEL	8	NIRAVAM SERAX		
	MC/DEL MC/DEL	LORAZEPAM OXAZEPAM CAPS	MC/DEL MC/DEL	8	SERAX TRANXENE		
	MO/DEL	OMALLI MINI OM O	MC/DEL	8	XANAX TABS		
<i>i</i>			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
<i>i</i> I	MC	HYDROXYZINE HCL SOLN	MC	İ	DROPERIDOL SOLN	Dosing limits apply,	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
	MC	HYDROXYZINE HCL SYRP	MC/DEL	İ	DROPERIDOL SOLN	please refer to Dose	preferred drug(s) exists.
	MC/DEL	HYDROXYZINE HCL TABS <sup>1</sup>	MC/DEL	I	DROPERIDOL SOLN	consolidation list.	
/ I	MC/DEL	HYDROXYZINE PAMOATE CAPS	1 1	İ			
	MC/DEL	MEPROBAMATE TABS					
WAS INTERESTED IN A SHURDITORS	***	ANTI-DEPRESSANTS	· · · · · · · · · · · · · · · · · · ·				
ANTIDEPRESSANTS - MAO INHIBITORS	MC/DEL	NARDIL TABS	MC/DEL	İ	TRANYLCYPROMINE	Use PA Form# 20420	
ANTIDEPRESSANTS - MAO INHIBITORS		j	MC/DEL		EMSAM <sup>1</sup>	Dosing limits apply,	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
TOPICAL			IIIO/DEE	I	EMSAM	please refer to Dose	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
ll i			1 1	İ		consolidation list.	potential drug interaction between another drug and the preferred drug(s) exists.
ll I		1	i j	I	1	Use PA Form# 20420	
ANTIDEPRESSANTS - SELECTED SSRI's	MC/DEL	BUPROPION HCL TABS	MC/DEL	8	APLENZIN⁴	Strong caution with	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
AND OTHERS	MC/DEL	BUPROPION SR	MC	8	AUVELITY <sup>11</sup>	pediatric population.	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
ll I	MC/DEL	BUPROPION XL 150mg and 300mg	MC/DEL	8	BUPROPION XL 450mg	Max daily dose allowed is	prevents usage of the preferred drug or a significant potential drug interaction between another drug and the preferred drug(s) exists.
ll I	MC/DEL	CITALOPRAM	MC/DEL	8	CELEXA	120mg, Combination of multiple strengths require	
ll i	MC/DEL	DULOXETINE <sup>2,9</sup>	MC	8	CYMBALTA <sup>2</sup>	PΔ	
ll i	MC/DEL MC/DEL	ESCITALOPRAM FLUOXETINE 10mg AND 20mg AND 40mg CAPS	MC/DEL MC/DEL	8 8	DRIZALMA SPRINKLES EFFEXOR TABS	<ol> <li>Dosing limits allowing 2 tabs/day and a max daily</li> </ol>	
ll i	WIC/DEL	FLUONETHINE TUHNY AND ZUHNY AND 40HINY OAL O	WIC/DEL	o I	EFFEXUR TADS	limit of 200mg / day applies.	
ll i	MC/DEL	FLUOXETINE HCL LIQD	MC/DEL	8	EFFEXOR XR CP24	Please see dose	CYMBALTA: Fibromyalgia diagnosis- prior use and failure of preferred generics (amitriptyline or cyclobenzaprine) and gabapentin prior to approval.
ll i	MC/DEL	FLUVOXAMINE MALEATE TABS	MC/DEL	8	FETZIMA <sup>7</sup>	consolidation list.	, <u>, , , , , , , , , , , , , , , , , , </u>
<b>[</b> ]	MC/DEL	MIRTAZAPINE	MC/DEL	8	FLUOXETINE 10mg AND 20mg AND 60mg TABS	<ol><li>Dosing limits apply,</li></ol>	
<b>[</b> ]	MC/DEL	NEFAZODONE	MC	8	FORFIVO XL	please refer to Dose	DDI: Fluvoxamine will now be non-preferred and require prior authorization if it is currently being used with glimepiride (Amaryl).
<b>[</b> ]	MC/DEL	PAROXETINE <sup>1</sup>	MC/DEL	8	IRENKA	consolidation list and max daily dose applies. Max	<u> </u>
ll i	MC/DEL	SERTRALINE HCL	MC/DEL	8	KHEDEZLA	daily dose allowed is 375mg.	
/ I	MC/DEL	TRAZODONE HCL TABS	MC/DEL	8	LEXAPRO TABS		DDI: Preferred nefazodone will now be non-preferred and require prior authorization if it is currently being used in combination with either Onglyza 5mg, Enablex 15mg or Vesicare 10mg.
ll I	MC/DEL MC/DEL	VENLAFAXINE ER CAPS <sup>5</sup>	MC MC	8	LUVOX TABS	<ol><li>Non-preferred products must be used in specified</li></ol>	DOLET OF THE STATE
11	WIC/DEL	VENLAFAXINE TABS <sup>5</sup>	MC/DEL	8	MAPROTILINE HCL TABS	step order.	DDI: Fluoxetine will require prior authorization if being used in combination with Plavix.  DDI: Fluoxamine will require prior authorization if being used in combination with Plavix.
<b>[</b> ]			MC MC	8	MIRTAZAPINE ODT OLEPTRO	7. Requires previous	DDI. Pluvoxallille will require prior additionazation it being used in combination with Lavix.
ll i			MC/DEL	8	PAROXETINE CR <sup>1</sup>	trials/failure of multiple	SAVELLA: Fibromyalgia diagnosis and trial of a preferred generic amitriptyline, cyclobenzaprine, duloxetine and gabapentin prior to approval.
<b>[</b> ]			MC/DEL	8	PAXIL <sup>1</sup>	preferred medications.	orteler i monifugu diagnoso and states a protestor gonoro annu prymor, systematical and general and general and a protestor gonoro annu prymor, systematical and a protestor gonoro annu prymor, systematical and general and general annual ann
<b>[</b> ]			MC/DEL	8	PAXIL CR <sup>1</sup>	Dosing limits apply, please see the dose consolidation	DDI: Drizalma Sprinkle avoid the concomitant use of duloxetine with potent CYP1A2 inhibitors (e.g. fluvoxamine, cimetidine, ciprofloxacin, enoxacin).
<b>[</b> ]			MC/DEL	8	PRISTIQ	list. Max daily dose of 80mg	
11			MC	8	PROZAC	if used concomitantly with	Zulresso® is available only through a restricted program under a Risk Evaluation and Mitigation Strategy (REMS) called the Zulresso® REMS.
ll i		1	MC	8	PROZAC CAPS	strong CYP3A4 inhibitor.	

				MC	8	PROZAC WEEKLY CPDR		
				MC/DEL	8	REMERON TABS	8. Psychiatry recommended.	Spravato: Treatment Resistant Depression
				MC/DEL	8	SARAFEM CAPS	Please see criteria section.	• Must be 18 years of age or older; and medication must be administered under the direct, on site, supervision of a licensed healthcare provider with post-administration observation of a
				MC/DEL	8	SPRAVATO <sup>8</sup>	9. Please use multiples of	minimum of least 2-hours. The medication must be prescribed by or in consultation with a psychiatrist and prescriber must be enrolled in the REMS program.
				MC/DEL	8	TRAZODONE HCL 300MG TABS	the 20mg, the 40mg is still	Approval is based upon failure of at least two antidepressants and failure of an antidepressant used adjunctively with one recognized augmentation strategy such as lithium, an atypical
				MC/DEL	8	TRINTELLIX	non-preferred.	antipsychotic, thyroid hormone, etc.
				MC	8	WELLBUTRIN TABS	10. For the treatment of	Ongoing use of Spravato beyond 3 months is based upon a positive response as evidenced by at least a 30 % reduction from baseline as measured by a standardized rating scale
				MC	8	WELLBUTRIN SR TBCR	patients ≥ 18 years of age.	
				MC	8	WELLBUTRIN XL	11. Use individual	Spravato: MDD with Suicidal Ideation
				MC/DEL	8	REMERON SOLTAB TBDP	ingredients separately.	Approval for this indication only if it is started in an inpatient unit, given adjunctively with an optimized antidepressant regimen, and with an 8-12 week initial approval with ongoing use
				MC/DEL	8	SAVELLA <sup>4</sup>	12. Approval will be limited	dependent upon documentation of ongoing benefit.
				MC/DEL	8	ZOLOFT	to a 14-day treatment course.	
				MC/DEL	8	ZULRESSO <sup>10</sup>	oouroo.	DDI: Reduce the Zurzuvae® dosage when used with a strong CYP3A4 inhibitor.
				MC	8	ZURZUVAE <sup>12</sup>		
				MC/DEL MC/DEL	8	VENLAFAXINE ER TABS <sup>5</sup>		
				MC/DEL	0	VIIBRYD <sup>6</sup>		
ANTIDERDEGGANTO, TRI OVOLIOO	MO/DEI				J	FLUOXETINE 90mg TABS <sup>6</sup>	Use PA Form# 20420	
ANTIDEPRESSANTS - TRI-CYCLICS	MC/DEL		AMITRIPTYLINE HCL TABS <sup>1</sup>	MC/DEL		AMOXAPINE TABS	<ol> <li>Users over the age of 65 require a pa.</li> </ol>	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
	MC/DEL		CLOMIPRAMINE HCL CAPS <sup>1</sup>	MC/DEL		ANAFRANIL CAPS	- dama arkan	preferred drug(s) exists.
	MC/DEL		DESIPRAMINE HCL TABS <sup>1</sup>	MC/DEL		DOXEPIN HCL 150 MG <sup>2</sup>		
	MC/DEL		DOXEPIN HCL <sup>1</sup> (not generic Silenor)	MC/DEL		DOXEPIN (generic Silenor)		
	MC/DEL		IMIPRAMINE HCL TABS <sup>1</sup>	MC/DEL		NORPRAMIN TABS	Use multiples of 50mg.	
	MC/DEL MC		NORTRIPTYLINE HCL <sup>1</sup>	MC/DEL MC		PAMELOR Tofranil		
	MC		PROTRIPTYLINE HCL TABS <sup>1</sup>	MC		VIVACTIL TABS	Use PA Form# 20420	
	WC		SURMONTIL CAPS <sup>1</sup>	WIC		VIVACTIL TABS	Use PA Form# 10220 for Brand Name requests	
							brand Name requests	
			SEDATIVE / HYPNOTICS					
SEDATIVE/HYPNOTICS - BARBITURATE	MC		SEDATIVE / HYPNOTICS	MC		LUMINAL SOLN	1 PA required for new users	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
SEDATIVE/HYPNOTICS - BARBITURATE	MC MC/DEL		BUTISOL SODIUM TABS <sup>1</sup>	MC MC/DEL		LUMINAL SOLN SOMNOTE CAPS	PA required for new users of preferred products if over	
SEDATIVE/HYPNOTICS - BARBITURATE	MC MC/DEL MC			MC MC/DEL				
SEDATIVE/HYPNOTICS - BARBITURATE			BUTISOL SODIUM TABS <sup>1</sup> CHLORAL HYDRATE SYRP <sup>1</sup>	MC MC/DEL			of preferred products if over	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
SEDATIVE/HYPNOTICS - BARBITURATE	MC		BUTISOL SODIUM TABS¹ CHLORAL HYDRATE SYRP¹ MEBARAL TABS¹	MC MC/DEL			of preferred products if over	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
SEDATIVE/HYPNOTICS -	MC/DEL		BUTISOL SODIUM TABS¹ CHLORAL HYDRATE SYRP¹ MEBARAL TABS¹	MC		SOMNOTE CAPS  HALCION TABS <sup>1</sup>	of preferred products if over 65 years. <u>Use PA Form# 20420</u> 1. Dosing limits apply,	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.  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/DEL MC/DEL MC/DEL		BUTISOL SODIUM TABS <sup>1</sup> CHLORAL HYDRATE SYRP <sup>1</sup> MEBARAL TABS <sup>1</sup> PHENOBARBITAL <sup>1</sup>	MC MC		SOMNOTE CAPS	of preferred products if over 65 years. <u>Use PA Form# 20420</u> 1. Dosing limits apply, please see dosing	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.  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
SEDATIVE/HYPNOTICS -	MC/DEL MC/DEL MC/DEL MC/DEL		BUTISOL SODIUM TABS¹ CHLORAL HYDRATE SYRP¹ MEBARAL TABS¹ PHENOBARBITAL¹  DORAL TABS¹ ESTAZOLAM TABS¹ FLURAZEPAM HCL CAPS¹	MC MC MC/DEL		SOMNOTE CAPS  HALCION TABS <sup>1</sup> MIDAZOLAM HCL SYRP RESTORIL CAPS <sup>1</sup>	of preferred products if over 65 years. <u>Use PA Form# 20420</u> 1. Dosing limits apply,	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.  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. Benzodiazepines do cause dependence with continued use and usage should be limited to 7-10 days at a time. Chronic intermittent use (2-3 Days per week
SEDATIVE/HYPNOTICS -	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		BUTISOL SODIUM TABS¹ CHLORAL HYDRATE SYRP¹ MEBARAL TABS¹ PHENOBARBITAL¹  DORAL TABS¹ ESTAZOLAM TABS¹ FLURAZEPAM HCL CAPS¹ TEMAZEPAM CAPS 15 & 30MG¹	MC MC		SOMNOTE CAPS  HALCION TABS <sup>1</sup> MIDAZOLAM HCL SYRP	of preferred products if over 65 years. <u>Use PA Form# 20420</u> 1. Dosing limits apply, please see dosing	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.  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
SEDATIVE/HYPNOTICS - BENZODIAZEPINES	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		BUTISOL SODIUM TABS¹ CHLORAL HYDRATE SYRP¹ MEBARAL TABS¹ PHENOBARBITAL¹  DORAL TABS¹ ESTAZOLAM TABS¹ FLURAZEPAM HCL CAPS¹ TEMAZEPAM CAPS 15 & 30MG¹ TRIAZOLAM TABS¹	MC MC MC/DEL MC/DEL		SOMNOTE CAPS  HALCION TABS <sup>1</sup> MIDAZOLAM HCL SYRP RESTORIL CAPS <sup>1</sup> TEMAZEPAM 7.5MG <sup>1</sup>	of preferred products if over 65 years.  Use PA Form# 20420  1. Dosing limits apply, please see dosing consolidation list.  Use PA Form# 30110	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.  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. Benzodiazepines do cause dependence with continued use and usage should be limited to 7-10 days at a time. Chronic intermittent use (2-3 Days per week max) is the standard of care
SEDATIVE/HYPNOTICS - BENZODIAZEPINES SEDATIVE/HYPNOTICS - Non-	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	1	BUTISOL SODIUM TABS¹ CHLORAL HYDRATE SYRP¹ MEBARAL TABS¹ PHENOBARBITAL¹  DORAL TABS¹ ESTAZOLAM TABS¹ FLURAZEPAM HCL CAPS¹ TEMAZEPAM CAPS 15 & 30MG¹ TRIAZOLAM TABS¹ MIRTAZAPINE	MC MC MC/DEL MC/DEL	7	SOMNOTE CAPS  HALCION TABS <sup>1</sup> MIDAZOLAM HCL SYRP RESTORIL CAPS <sup>1</sup> TEMAZEPAM 7.5MG <sup>1</sup> AMBIEN <sup>1</sup>	of preferred products if over 65 years. <u>Use PA Form# 20420</u> 1. Dosing limits apply, please see dosing consolidation list.	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.  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. Benzodiazepines do cause dependence with continued use and usage should be limited to 7-10 days at a time. Chronic intermittent use (2-3 Days per week max) is the standard of care  4 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
SEDATIVE/HYPNOTICS - BENZODIAZEPINES	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	1 1 1	BUTISOL SODIUM TABS¹ CHLORAL HYDRATE SYRP¹ MEBARAL TABS¹ PHENOBARBITAL¹  DORAL TABS¹ ESTAZOLAM TABS¹ FLURAZEPAM HCL CAPS¹ TEMAZEPAM CAPS 15 & 30MG¹ TRIAZOLAM TABS¹ MIRTAZAPINE TRAZODONE	MC MC MC/DEL MC/DEL	7 7	SOMNOTE CAPS  HALCION TABS <sup>1</sup> MIDAZOLAM HCL SYRP RESTORIL CAPS <sup>1</sup> TEMAZEPAM 7.5MG <sup>1</sup>	of preferred products if over 65 years.  Use PA Form# 20420  1. Dosing limits apply, please see dosing consolidation list.  Use PA Form# 30110  1. Quantity Limit of 12 per 34 days.	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.  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. Benzodiazepines do cause dependence with continued use and usage should be limited to 7-10 days at a time. Chronic intermittent use (2-3 Days per week max) is the standard of care
SEDATIVE/HYPNOTICS - BENZODIAZEPINES SEDATIVE/HYPNOTICS - Non-	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	1 1 1	BUTISOL SODIUM TABS¹ CHLORAL HYDRATE SYRP¹ MEBARAL TABS¹ PHENOBARBITAL¹  DORAL TABS¹ ESTAZOLAM TABS¹ FLURAZEPAM HCL CAPS¹ TEMAZEPAM CAPS 15 & 30MG¹ TRIAZOLAM TABS¹ MIRTAZAPINE	MC MC MC/DEL MC/DEL MC/DEL	7 7	SOMNOTE CAPS  HALCION TABS¹ MIDAZOLAM HCL SYRP RESTORIL CAPS¹ TEMAZEPAM 7.5MG¹  AMBIEN¹ ESZOPICLONE	of preferred products if over 65 years.  Use PA Form# 20420  1. Dosing limits apply, please see dosing consolidation list.  Use PA Form# 30110  1. Quantity Limit of 12 per 34 days.  2. Quantity limits will be	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.  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. Benzodiazepines do cause dependence with continued use and usage should be limited to 7-10 days at a time. Chronic intermittent use (2-3 Days per week max) is the standard of care  4 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
SEDATIVE/HYPNOTICS - BENZODIAZEPINES SEDATIVE/HYPNOTICS - Non-	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		BUTISOL SODIUM TABS¹ CHLORAL HYDRATE SYRP¹ MEBARAL TABS¹ PHENOBARBITAL¹  DORAL TABS¹ ESTAZOLAM TABS¹ FLURAZEPAM HCL CAPS¹ TEMAZEPAM CAPS 15 & 30MG¹ TRIAZOLAM TABS¹ MIRTAZAPINE TRAZODONE ZOLPIDEM²	MC MC MC/DEL MC/DEL	7 7 7 8	SOMNOTE CAPS  HALCION TABS¹ MIDAZOLAM HCL SYRP RESTORIL CAPS¹ TEMAZEPAM 7.5MG¹  AMBIEN¹ ESZOPICLONE  ZOLPIDEM ER	of preferred products if over 65 years. <u>Use PA Form# 20420</u> 1. Dosing limits apply, please see dosing consolidation list. <u>Use PA Form# 30110</u> 1. Quantity Limit of 12 per 34 days.  2. Quantity limits will be allowed up to 30/30, but intermittent therapy is	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.  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. Benzodiazepines do cause dependence with continued use and usage should be limited to 7-10 days at a time. Chronic intermittent use (2-3 Days per week max) is the standard of care  4 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.
SEDATIVE/HYPNOTICS - BENZODIAZEPINES SEDATIVE/HYPNOTICS - Non-	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		BUTISOL SODIUM TABS¹ CHLORAL HYDRATE SYRP¹ MEBARAL TABS¹ PHENOBARBITAL¹  DORAL TABS¹ ESTAZOLAM TABS¹ FLURAZEPAM HCL CAPS¹ TEMAZEPAM CAPS 15 & 30MG¹ TRIAZOLAM TABS¹ MIRTAZAPINE TRAZODONE	MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		SOMNOTE CAPS  HALCION TABS¹ MIDAZOLAM HCL SYRP RESTORIL CAPS¹ TEMAZEPAM 7.5MG¹  AMBIEN¹ ESZOPICLONE	of preferred products if over 65 years.  Use PA Form# 20420  1. Dosing limits apply, please see dosing consolidation list.  Use PA Form# 30110  1. Quantity Limit of 12 per 34 days.  2. Quantity limits will be allowed up to 30/30, but	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.  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. Benzodiazepines do cause dependence with continued use and usage should be limited to 7-10 days at a time. Chronic intermittent use (2-3 Days per week max) is the standard of care  4 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
SEDATIVE/HYPNOTICS - BENZODIAZEPINES SEDATIVE/HYPNOTICS - Non-	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		BUTISOL SODIUM TABS¹ CHLORAL HYDRATE SYRP¹ MEBARAL TABS¹ PHENOBARBITAL¹  DORAL TABS¹ ESTAZOLAM TABS¹ FLURAZEPAM HCL CAPS¹ TEMAZEPAM CAPS 15 & 30MG¹ TRIAZOLAM TABS¹ MIRTAZAPINE TRAZODONE ZOLPIDEM²	MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		SOMNOTE CAPS  HALCION TABS¹ MIDAZOLAM HCL SYRP RESTORIL CAPS¹ TEMAZEPAM 7.5MG¹  AMBIEN¹ ESZOPICLONE  ZOLPIDEM ER	of preferred products if over 65 years. <u>Use PA Form# 20420</u> 1. Dosing limits apply, please see dosing consolidation list. <u>Use PA Form# 30110</u> 1. Quantity Limit of 12 per 34 days.  2. Quantity limits will be allowed up to 30/30, but intermittent therapy is	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.  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. Benzodiazepines do cause dependence with continued use and usage should be limited to 7-10 days at a time. Chronic intermittent use (2-3 Days per week max) is the standard of care  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.  Ambien, Ambien CR, Lunesta, Sonata, Zaleplon and Zolpidem may cause dependence with continued use and as with benzodiazepines, usage should be limited to 7-10 days at a time. Chronic intermittent use (2-3 days per week max) is the standard of care. Please refer to Sedative/Hypnotic PA form.
SEDATIVE/HYPNOTICS - BENZODIAZEPINES SEDATIVE/HYPNOTICS - Non-	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL		BUTISOL SODIUM TABS¹ CHLORAL HYDRATE SYRP¹ MEBARAL TABS¹ PHENOBARBITAL¹  DORAL TABS¹ ESTAZOLAM TABS¹ FLURAZEPAM HCL CAPS¹ TEMAZEPAM CAPS 15 & 30MG¹ TRIAZOLAM TABS¹ MIRTAZAPINE TRAZODONE ZOLPIDEM²	MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	8	HALCION TABS¹ MIDAZOLAM HCL SYRP RESTORIL CAPS¹ TEMAZEPAM 7.5MG¹  AMBIEN¹ ESZOPICLONE  ZOLPIDEM ER AMBIEN CR¹	of preferred products if over 65 years.  Use PA Form# 20420  1. Dosing limits apply, please see dosing consolidation list.  Use PA Form# 30110  1. Quantity Limit of 12 per 34 days.  2. Quantity limits will be allowed up to 30/30, but intermittent therapy is recommended.  3. Only zolpidem trial/failure will be required to obtain	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.  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. Benzodiazepines do cause dependence with continued use and usage should be limited to 7-10 days at a time. Chronic intermittent use (2-3 Days per week max) is the standard of care  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.  Ambien, Ambien CR, Lunesta, Sonata, Zaleplon and Zolpidem may cause dependence with continued use and as with benzodiazepines, usage should be limited to 7-10 days at a time. Chronic intermittent use (2-3 days per week max) is the standard of care. Please refer to Sedative/Hypnotic PA form.
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		ANTI-PSYCHOTICS					
ANTIPSYCHOTICS - ATYPICALS	MC	ABILIFY ASIMTUFII	MC/DEL	8	ABILIFY DISC TAB, INJ and SOL <sup>1</sup>	If prescribing 2 or more	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	ABILIFY MAINTENA	MC	8	ABILIFY TABS <sup>2</sup>	antipsychotics, PA will be	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
	MC/DEL	ARIPIPRAZOLE TAB <sup>3</sup>	MC/DEL	8	ARIPIPRAZOLE SOL	required for both drugs,	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-
	MC	ARISTADA	MC/DEL	8	ARIPIPRAZOLE ODT	except if one is Clozapine.	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
	MC					This also includes combination of Seroquel with	and failed at full therapeutic doses for adequate durations (at least two weeks).
	MC/DEL	ARISTADA INITIO <b>OLANZAPINE<sup>2,3</sup></b>	MC MC	8	CAPLYTA COBENFY	Seroquel XR.	
		OLANZAPINE OLANZAPINE <sup>2,3</sup> ODT	MC	8	ERZOFRI	·	
	MC/DEL		MC	0	FANAPT		Prescriptions for quetiapine are limited to a maximum daily dose of 800mg.
	MC/DEL MC	INVEGA HAFYERA INVEGA SUSTENNA	MC/DEL	0	GEODON		rescriptions for quetraprine are limited to a maximum daily dose or ocoring.
				0			Handu Fatabilish telepahilih with and singgidana garagta initiating Handu
	MC/DEL	INVEGA TRINZA INJ	MC	8	INVEGA	Use PA form# 20440 for	Uzedy: Establish tolerability with oral risperidone prior to initiating Uzedy
	MC/DEL	LURASIDONE TAB	MC MC	8	IGALMI	Multiple Antipsychotic	
	MC/DEL	PALIPERIDONE ER	IVIC	8	LATUDA	<u>requests</u>	
	MO/DEL	PEROFERIO		•	LVDALV(	Use PA form# 10130 for non	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:
	MC/DEL MC	PERSERIS RISPERDAL CONSTA	MC	8	LYBALVI	preferred single therapy	"
			MC	8	NUPLAZID	<u>atypical requests</u>	• schizophrenia
	MC/DEL	RISPERIDONE ODT	MC	8	OPIPZA		• bipolar disorder
	MC/DEL	RISPERIDONE TAB <sup>2,3</sup>	MC	8	REXULTI		• agitation related to autism
	MC/DEL	RISPERIDONE SOLN <sup>2</sup>	MC	8	RISPERDAL TAB		adjunct in major depressive disorder
	MC	RYKINDO	MC	8	RISPERDAL M TAB <sup>1</sup>		
		0.15714.011.523	MC	8	RISPERDAL SOLN	Established users of single therepy at minels were	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.
	MC/DEL	QUETIAPINE <sup>2,3</sup>				single therapy atypicals were grandfathered.	
						grandatiorou.	
	MC/DEL	QUETIAPINE XR	MC/DEL	8	SAPHRIS <sup>1</sup>		
		VRAYLAR <sup>4</sup>	MC	0	SECUADO	Prior Authorization will be	
	MC	ZIPRASIDONE <sup>2,3</sup>		8		required for preferred	
	MC/DEL	ZIFRASIDONE	MC/DEL	8	SEROQUEL TABS	medications for members	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
			MC	8	UZEDY	under the age of 5.	a CYP3A4 inducer (such as rifampin, carbamazepine) is not recommended.
			MC MC	8	ZYPREXA TABS	2 Danier limite and colored	<u></u>
			IVIC		ZYPREXA RELPREVV	refer to the dose	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).
			MC	8	1	consolidation list.	gaunozadin and mozinozadin).
			MC/DEL	9	ZYPREXA ZYDIS TBDP <sup>1</sup> SEROQUEL XR		
			WIO/DEL	9	SENOQUEE XIV	4.Requires step through 1	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
						preferred drug for all	weight for ongoing approval. If weight gain >= 10 % of initial body weight, then criteria for ongoing use not met.
						indications except AMDD.	
						AMDD requires insufficient	
						response from two	
						antidepressants	
							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.
							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.
ANTIPSYCHOTICS - SPECIAL ATYPICALS	MC/DEL	CLOZAPINE TABS	MC/DEL		CLOZAPINE ODT	Use PA Form# 20420	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
			MC/DEL		CLOZARIL TABS		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.
			MC/DEL		VERSACLOZ SUSP		

ANTIPSYCHOTICS - TYPICAL	MC/DEL MC/DEL MC MC/DEL MC MC MC/DEL MC MC/DEL MC MC/DEL MC MC/DEL MC MC/DEL MC/DEL MC/DEL MC MC/DEL MC MC/DEL MC	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/DEL MC MC/DEL 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	LITHIUM CARBONATE	MC/DEL	ESKALITH CAPS	Use PA Form# 20420	
	MC/DEL	LITHIUM CITRATE SYRP	MC/DEL	ESKALITH CR TBCR		
		COMBINATION - PSYCHOTHERAP	EUTIC			
PSYCHOTHERPEUTIC COMBINATION			MC/DEL	CHLORDIAZEPOXIDE/AMITRIPT		
			MC/DEL	PERPHENAZINE/AMITRIPTYLIN		
	$\Box$				<u>Use PA Form# 20420</u>	
	MC/DEL	STIMULANTS  AMPHETAMINE SALT COMBO <sup>1,4</sup>	MC/DEL	ADDERALL TABS	Preferred stimulants will	
STIMULANT - AMPHETAMINES -SHORT	MC/DEL	DEXTROAMPHET SULF TABS	MC	EVEKEO	be available without PA if	
ACTING	MC	PROCENTRA	MC/DEL	METHAMPHETAMINE HCL	diagnosis of ADHD or Narcolepsy.	
			MC	ZENZEDI	ivarcolepsy.	
					0.4	
					<ol> <li>As per recent FDA alert,</li> <li>Adderall &amp; Dexedrine should</li> </ol>	
					not be used in patients with	
					underlying heart defects since they may be at	
					increased risk for sudden	
					death.	
					<ol><li>Dosing limits apply,</li></ol>	
					please see dosing consolidation list.	
					oonoonaa.on noo	
					4. Max daily dose of 50mg.	
					<u>Use PA Form# 20420</u>	
STIMULANT - LONG ACTING AMPHETAMINES SALT	MC/DEL	AMPHETAMINE/DEXTROAMPHET ER <sup>3,4,7</sup>	MC	MYDAYIS <sup>5</sup>	Use PA Form# 20420	
AMETICIANINES SALI	MC MC	ADDERALL XR CP24 <sup>1,3,4,7</sup> VYVANSE <sup>2,3,4</sup>	MC MC	VYVANSE CHEW <sup>4,6</sup> XELSTRYM <sup>8</sup>	<ol> <li>As per recent FDA alert,</li> <li>Adderall should not be used</li> </ol>	
	IVIC	VYVANSE	WIC	AELOTATIVI	in patients with underlying	
					heart defects since they may be at increased risk for	DDI: The concomitant use of Mydayis® is contraindicated with monoamine oxidase inhibitors (MAOIs) or within 14 days after discontinuing MAOI treatment, as
					sudden death.	concomitant use can increase hypertensive crisis.
	I I		l I	I		

1					
1	1	I	1 1	I	2. FDA approval is currently
ļ ,		1		1	for adults and children 6 or
<u>'</u>		1		1	older. Will be available
<u>'</u>		1		1	without PA for this age group
<u>'</u>		1		1	if within dosing limits. Limit
, <b>I</b>		1		1	of one capsule daily. Max
/ <b> </b>		1		1	dose of 70MG daily.
/ <b> </b>		1		1	
/ <b> </b>		1		1	
1 <b>1</b>		1		1	3. Preferred stimulants will
, <b> </b>		1		1	be available without PA if
, <b> </b>		1		1	diagnosis of ADHD.
, <b> </b>		1		1	4. Dosing limits apply,
<b>∤</b>		1		1	please see dosing
•		1		1	consolidation list.
<b> </b>		1		1	5. For the treatment of
<b> </b>		1		1	5. For the treatment or Attention Deficit
· •		1		1	Hyperactivity Disorder
<b>/  </b>		1		1	(ADHD) in patients 13 years
<b>/                                    </b>		1		1	and older
<b> </b>		1		l	
<b>/                                    </b>		1		1	6. Vyvanse chew grace
<b> </b>		1		1	period for current user
/		1		l	through June 2022.
<b> </b>		1		1	7. FDA approval is currently
<b> </b>		1		1	for adults and children 6 or
<b>/                                    </b>		1		1	older. Will be available
<b>/                                    </b>		1		1	without PA for this age group
<b>[ ]</b>		1		1	if within dosing limits. Max
<b>[ ]</b>					dose of 50MG daily without a
<b>/ </b>					PA.
<b>/                                    </b>					
<b>[ ]</b>					8. For the treatment of
<b>                                     </b>					patients 6 years of age and
<b>/                                    </b>					older.
LONG ACTING AMPHETAMINES	MC	DEXTROAMPHET SULF CPSR <sup>1,3</sup>	MC/DEL	ADZENYS ER <sup>3</sup>	<del>- </del>
Lone Advinto Aum 11211 mm 125	MC/DEL	DEXTROAMPHETAMINE ER	1110/1522	ADZERIO ER	Preferred stimulants will
<b>/  </b>	WIC/DLL	DEXTROAMIFRE LAMING LIX			be available without PA if
/I		1			diagnosis of ADHD.
/I			MC	ADZENYS XR- ODT	2. As per recent FDA alert,
<b>/                                    </b>	1				2 As not recent EDA plant
4 I				<b>.</b>	
<b>11</b>					Adderall & Dexedrine should
					Adderall & Dexedrine should not be used in patients with
					Adderall & Dexedrine should not be used in patients with underlying heart defects
					Adderall & Dexedrine should not be used in patients with underlying heart defects since they may be at increased risk for sudden
					Adderall & Dexedrine should not be used in patients with underlying heart defects since they may be at
					Adderall & Dexedrine should not be used in patients with underlying heart defects since they may be at increased risk for sudden
	МС	DYANAVEL XR SUS	мс	ADZENYS XR <sup>3</sup>	Adderall & Dexedrine should not be used in patients with underlying heart defects since they may be at increased risk for sudden death.
	мс	DYANAVEL XR SUS	MC MC	ADZENYS XR <sup>3</sup> DEXEDRINE CAP SR <sup>2,3</sup>	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,  DDI: The concomitant use of Adzenys® XR is contraindicated with monoamine oxidase inhibitors (MAOIs) or within 14 days after discontinuing MAOI treatment.
	мс	DYANAVEL XR SUS			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  DDI: The concomitant use of Adzenys® XR is contraindicated with monoamine oxidase inhibitors (MAOIs) or within 14 days after discontinuing MAOI treatment.
	MC	DYANAVEL XR SUS			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,  DDI: The concomitant use of Adzenys® XR is contraindicated with monoamine oxidase inhibitors (MAOIs) or within 14 days after discontinuing MAOI treatment.
	МС	DYANAVEL XR SUS		DEXEDRINE CAP SR <sup>2,3</sup>	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  DDI: The concomitant use of Adzenys® XR is contraindicated with monoamine oxidase inhibitors (MAOIs) or within 14 days after discontinuing MAOI treatment.
	МС	DYANAVEL XR SUS	MC		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.  DDI: The concomitant use of Adzenys® XR is contraindicated with monoamine oxidase inhibitors (MAOIs) or within 14 days after discontinuing MAOI treatment.
CTIMUS ANT METUVI DUEMINATE			MC	DEXEDRINE CAP SR <sup>2,3</sup> DYANAVEL XR TAB	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.  DDI: The concomitant use of Adzenys® XR is contraindicated with monoamine oxidase inhibitors (MAOIs) or within 14 days after discontinuing MAOI treatment.
STIMULANT - METHYLPHENIDATE	MC/DEL	DEXMETHYLPHENIDATE IR TABS	MC MC/DEL	DEXEDRINE CAP SR <sup>2,3</sup> DYANAVEL XR TAB  FOCALIN IR TABS	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.  DDI: The concomitant use of Adzenys® XR is contraindicated with monoamine oxidase inhibitors (MAOIs) or within 14 days after discontinuing MAOI treatment. please see dosing consolidation list.  Use PA Form# 20420  1. Preferred stimulants will 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 or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is or intolerable side effects before non-preferred drugs will be approved.
STIMULANT - METHYLPHENIDATE	MC/DEL MC/DEL	DEXMETHYLPHENIDATE IR TABS METHYLPHENIDATE SOL	MC MC/DEL MC/DEL	DEXEDRINE CAP SR <sup>2,3</sup> DYANAVEL XR TAB  FOCALIN IR TABS  METADATE ER	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.  DDI: The concomitant use of Adzenys® XR is contraindicated with monoamine oxidase inhibitors (MAOIs) or within 14 days after discontinuing MAOI treatment. please see dosing consolidation list.  Use PA Form# 20420  1. Preferred stimulants will be available without PA if 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
STIMULANT - METHYLPHENIDATE	MC/DEL MC/DEL MC/DEL	DEXMETHYLPHENIDATE IR TABS METHYLPHENIDATE SOL METHYLPHENIDATE TAB	MC  MC/DEL  MC/DEL  MC	DEXEDRINE CAP SR <sup>2,3</sup> DYANAVEL XR TAB  FOCALIN IR TABS  METADATE ER  METHYLPHENIDATE HCL CHEW	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.  DDI:: The concomitant use of Adzenys® XR is contraindicated with monoamine oxidase inhibitors (MAOIs) or within 14 days after discontinuing MAOI treatment. please see dosing consolidation list.  Use PA Form# 20420  1. Preferred stimulants will be available without PA if diagnosis of ADHD.  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 or 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 General Criteria category E.
STIMULANT - METHYLPHENIDATE	MC/DEL MC/DEL	DEXMETHYLPHENIDATE IR TABS METHYLPHENIDATE SOL	MC  MC/DEL  MC/DEL  MC  MC  MC	DEXEDRINE CAP SR <sup>2,3</sup> DYANAVEL XR TAB  FOCALIN IR TABS  METADATE ER  METHYLPHENIDATE HCL CHEW  METHYLIN CHEWABLES	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.  DDI: The concomitant use of Adzenys® XR is contraindicated with monoamine oxidase inhibitors (MAOIs) or within 14 days after discontinuing MAOI treatment. please see dosing consolidation list.  Use PA Form# 20420  1. Preferred stimulants will be available without PA if diagnosis of ADHD.  Use PA Form# 20420  Verently and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is or 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 General Criteria category E.
STIMULANT - METHYLPHENIDATE	MC/DEL MC/DEL MC/DEL	DEXMETHYLPHENIDATE IR TABS METHYLPHENIDATE SOL METHYLPHENIDATE TAB	MC  MC/DEL  MC/DEL  MC	DEXEDRINE CAP SR <sup>2,3</sup> DYANAVEL XR TAB  FOCALIN IR TABS  METADATE ER  METHYLPHENIDATE HCL CHEW	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.  DDI:: The concomitant use of Adzenys® XR is contraindicated with monoamine oxidase inhibitors (MAOIs) or within 14 days after discontinuing MAOI treatment. please see dosing consolidation list.  Use PA Form# 20420  1. Preferred stimulants will be available without PA if diagnosis of ADHD.  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 or 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 General Criteria category E.

			MC/DEL		RITALIN	please see dosing consolidation list. Maximum daily doses are as follows: 72mg daily for methylphenidate and 36mg daily for dexmethylphenidate.	
STIMULANT - METHYLPHENIDATE - LONG ACTING	MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC MC MC	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 <sup>5,1</sup> QUILLIVANT XR SUS <sup>1,5</sup> RITALIN LA <sup>4</sup>	MC MC/DEL MC MC MC MC MC MC MC/DEL	8	METADATE CD CPCR ADHANSIA XR <sup>26</sup> APTENSIO XR <sup>2</sup> AZSTARYS <sup>6</sup> COTEMPLA XR <sup>2</sup> COTEMPLA XR ODT <sup>2</sup> DAYTRANA <sup>23</sup> JORNAY PM <sup>2,6</sup> METHYLPHENIDATE ER CAPS <sup>2,4</sup>	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	ATOMOXETINE HCL  ARMODAFINIL  CLONIDINE ER  GUANFACINE ER  MODAFINIL TABS  QELBREE <sup>6,7</sup>	MC/DEL MC MC MC/DEL MC MC MC/DEL	8	PROVIGIL TABS <sup>3</sup> STRATTERA <sup>1,2</sup> CAFCIT SOLN <sup>3</sup> INTUNIV KAPVAY ONYDA XR <sup>6</sup> SUNOSI	1. Failure of both an amphetamine and methylphenidate is required for consideration for approva 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 in required before approval of Strattera.	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.
			MC MC	8 8	WAKIX XYREM SOL	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	s  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
			MC MC/DEL MC MC	-	XYWAV <sup>5</sup> NUVIGIL <sup>3</sup> DESOXYN TABS <sup>3</sup> DESOXYN CR <sup>3</sup>	must be used in specified 4. Please use generic Guanfacine. 5. For patients 7 years of age and older with Company 6. For pediatric patients 6 years of age or older	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 neak exposure of sensitive CYP1A2 substates, which may increase the risk of adverse reactions

PSYCHOTHERAPEUTIC AGENTS - MISC.		ANTI-CATAPLECTIC AGENTS	MC MC		NUEDEXTA XENAZINE	7. Preferred with a trial and fail either Atomoxetine OR any 2 preferred ADHD agents.  Use PA Form# 20710 for Provigil, Nuvigil and Xyrem  Use PA Form# 20420 for all others	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.
			IIIV		AEIVAZ IIVE	Use PA Form# 20710 for Xenazine	
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 - Cholinomimetics/Others	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	DONEPEZIL HYDROCHLORIDE TABS¹ DONEPEZIL HYDROCHLORIDE ODT¹ EXELON DIS¹ GALANTAMINE CAPS¹ GALANTAMINE TAB¹ MEMANTINE¹ RIVASTIGMINE TARTRATE CAPS¹	MC MC/DEL MC/DEL MC/DEL MC MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	6 6 7 8 8 8 8 8 8 8 8	ARICEPT TABS <sup>2</sup> ARICEPT ODT <sup>2</sup> DONEPEZIL HYDROCHLORIDE TABS 23MG ADLARITY <sup>3</sup> EXELON CAP  GALANTAMINE HYDROBROMIDE SOL KISUNLA LEQEMBI <sup>1,2</sup> MEMANTINE HCL SOL NAMENDA NAMENDA XR CAPS NAMZARIC RAZADYNE <sup>2</sup> COGNEX CAPS <sup>2</sup>		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.  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  - Prescribed by or in consultation with a neurologist or geniatrician or geniatric psychiatrist. Diagnosis of Alzheimer's disease defined as:  - 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  - Testing:  - 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  - Member is age 50 or older  - Obtain recent (within one year) brain magnetic resonance imaging (MRI) prior to initiating treatment  - 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)  - Member dose NOT have history or increased risk of amyloid related imaging abnormalities hemosiderin deposition (ARIA-H), which includes microhemorrhage and superficial siderosis  - Member dose NOT have hypersensitivity to any components of these drugs  - Failure of or inability to tolerate at least two other preferred Alzheimer th
		SMOKING CESSATION				Use PA Form# 20420	
NICOTINE PATCHES / TABLETS	MC/DEL MC/DEL MC/DEL MC/DEL	CHANTIX TAB <sup>1</sup> CHANTIX STARTER PACK NICOTINE DIS PT24 <sup>1</sup> VARENICLINE TAB	MC/DEL		NICODERM CQ PT24 <sup>1</sup>	Use PA Form# 20420  1. See criteria section for exemptions	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

				1		-	maia covalan miniminanone
							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 GUM¹ NICOTINE LOZENGE MINI NICOTINE LOZENGE	MC/DEL MC/DEL MC/DEL MC	8 8 8	NICOTROL INHALER <sup>12</sup> NICOTROL NASAL SPRAY <sup>12</sup> NICORETTE GUM <sup>12</sup> NICORETTE LOZENGES	Use PA Form# 20420  1. See criteria section for exemptions  2. Must use non-preferred products in specified step	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.
						order.	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	ACAMPROSATE ANTABUSE TABS DISULFIRAM TABS	MC/DEL		ACAMPRO <sup>1</sup>	<ol> <li>Should only be used in conjunction with formal structured outpatient detoxification program.</li> </ol>	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.
	MC/DEL	NALTREXONE HCL TABS				Use PA Form# 20420	
		MISCELLANEOUS ANALGESICS				USG 1 A 1 UIIII# 20420	
ANALGESICS - MISC.	MC/DEL	ACETAMINOPHEN	MC		AXOCET 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
	MC/DEL	ASPIRIN	MC/DEL		ESGIC-PLUS	00017110111111120120	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
	MC/DEL	ASPRIN/ APAP/ CAFF TAB	MC/DEL		FIORICET TABS		preferred drug(s) exists.
	MC/DEL	BUTAL/ASA/CAFF	MC		FIORINAL CAPS		
	MC/DEL	BUTALBITAL COMPOUND	MC		FIORTAL CAPS		
	MC/DEL	BUTALBITAL/ACET TABS	MC/DEL		FORTABS TABS		
	MC/DEL	BUTALBITAL/APAP CAPS	MC		PHRENILIN TABS		
	MC/DEL	BUTALBITAL/APAP/CAFFEINE TABS	МС		PHRENILIN FORTE CAPS		
	MC/DEL	CHOLINE MAGNESIUM TRISALI	MC		TRILISATE LIQD		
	MC/DEL	DIFLUNISAL TABS	MC		TRILISATE TABS		
	MC	EXCEDRIN	MC		ZEBUTAL CAPS		
	MC/DEL	SALSALATE TABS	MC		ZORPRIN TBCR		
		LONG ACTING NARCOTICS					
NARCOTICS - LONG ACTING	MC/DEL	FENTANYL PATCH⁴	MC	8	ARYMO ER	Use PA Form# 20510	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
	MC/DEL	BUTRANS⁴	MC	8	AVINZA	Use PA form #10300 for PA	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
	MC/DEL	MORPHINE SULFATE ER TB12	MC	8	BELBUCA	over the opiate limit	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
			MC	8	EXALGO	Oxycontin will be	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
			MC/DEL	8	HYSINGLA ER	available without PA for patients treated for or dying	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
			MC	8	KADIAN	from cancer or hospice	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:
			MC/DEL	8	METHADONE	patients. CA (cancer) or HO	
			MC/DEL	8	METHADOSE	(hospice) diag code may be	
			MC/DEL	8	MORPHABOND ER	used but store must verify since all scripts will be	
			MC/DEL	8	MORPHINE SULFATE SUPP	audited and stores will be	
			MC/DEL	8	MORPHINE SULFATE SUPP	liable.	1. Frequent or persistent early refills of controlled drugs;
			MC/DEL	8	MS CONTIN TB12		2.Multiple instances of early refill overrides due to reports of misplacement, stolen, dropped in toilet or sink, distant travel, etc.;
[ <b>]</b>			MC	8	OPANA ER		3.Breaches of narcotic contracts with any provider;
			MC/DEL MC/DEL	8	ORAMORPH SR TB12 OXYCONTIN TB12 <sup>1</sup>	Established users are	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;
						grandfathered.	
			MC	8	XARTEMIS ER	Oxycodone ER allowed only 2 per day for all attendable oxygent 90 mg.	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
<b> </b>	I I	I	MC	8	ZOHYDRO ER	strengths except 80 mg,	1

			MC MC/DEL	8 9	OXYCODONECONC OXYCODONE ER <sup>3,5</sup>	Please see dose consolidation list.	3. 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/APAP TABS TRAMADOL/APAP TABS	MC/DEL MC MC/DEL MC MC MC MC MC MC MC MC	7 8 8 8 8 8 8 8 9	RYZOLT BUPRENEX SOLN BUTORPHANOL NALBUPHINE HCL SOLN QDOLO SOLN SEGLENTIS¹ STADOL NS SOLN TRAMADOL ER ULTRACET TABS¹ ULTRAM ER	Use PA form #10300 for PAs over the opiate limit  1. Only available if component ingredients are unavailable.	Preferred drugs from this and other narcotic classes must be tired for at least 2 weeks each and failed due to lack of efficacy or inclorables 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. Carding the trait period. Substance abuse evaluations may be required for patients with medical records splaying colarists larging of narcotic missue 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, famility to stance 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 opicid documentation (see PA form) including but not limited to failing to submit to and pass pill counts;  5, 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 missae and base such as stronic early refills, short dosing int

		1	I	1	ſ	I	Ī	I
			MISCELLANEOUS NARCOTICS					
ARCOTICS - MISC.	MC/DEL	. 1	ACETAMINOPHEN/CODEINE	MC/DEL	8	ABSTRAL	Fentanyl OT loz (Barr)	
,	MC/DEL		ASPIRIN/CODEINE TABS	MC/DEL	8	APADAZ	and Capital and codeine	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
,	MC/DEL	_1 '	BUTAL/ASA/CAFF/COD CAPS	MC/DEL	8	ASCOMP/CODEINE CAPS	suspension products require PA for users over 18 years of	ire preferred drug(s) exists. Please refer to General Criteria category E.
,	MC		BUTALBITAL/ASPIRIN/CAFFEI CAPS	MC/DEL	8	BUTALBITAL/APAP/CAFFEINE/ CAPS	age. PA is not required if	1
,	MC	1 '	CAPITAL AND CODEINE SUSP <sup>1</sup>	MC/DEL	8	BUTALBITAL COMPOUND- CODEINE CAP	under 18 years of age.	
•	MC		CAPITAL/CODEINE SUSP <sup>1</sup>	MC	8	DEMEROL		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.
,	MC/DEL		CODEINE PHOSPHATE SOLN	MC/DEL		DILAUDID		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
, I	MC/DEL	_	CODEINE SULFATE TABS	MC		DILAUDID-HP SOLN	2. Oxycodone/acet 10/650	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.
,	MC/DEL	_	ENDOCET TABS <sup>3</sup>	MC	8	FENTANYL CITRATE SOLN	is 8 times more expensive.	
•	MC/DEL		ENDODAN TABS	MC/DEL	8	FENTORA	Use twice as many of oxycod/acet 5/325 instead.	
•	MC/DEL	_	FENTANYL OT LOZ <sup>1</sup>	MC/DEL	8	FIORICET/CODEINE CAPS	oxycod/acet 5/325 instead. You can mix and match	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
•	MC/DEL		FENTANYL OT LOZ1	MC	8	FIORINAL/CODEINE #3 CAPS	preferred strengths of	note that MaineCare implemented a 30 MME limit January 1, 2013 that is still effective.
i ,	MC/DEL	_	HYDROCODONE/ACETAMINOPHEN	MC	8	FIORTAL/CODEINE CAPS	oxycodone and	1
1	MC/DEL	_	HYDROMORPHONE HCL <sup>3</sup>	MC/DEL	8	HYDROCODONE/IBUPROFEN	oxycodone/acet to minimize	
1	MC/DEL	_	LORTAB ELX	MC/DEL		HYDROMORPHONE ER	acet. dose similar to certain non-preferred drugs.	Prost-Sulgical members may receive prior authorizations for opinios up to a so says
1	MC/DEL		MEPERIDINE SOL	MC/DEL		HYDROMORPHONE RECTAL SUPP	Horryrolonou alag	An MME conversion chart is available at www.mainecarepdl.org. Click on "General Pharmacy Info."
1	MC/DEL	_	OXYCODONE TAB	MC/DEL		IBUDONE		AN MME Currension chair is available at www.mainecarepoi.org. Onco. On Societary mainecy mic.
1	MC/DEL MC/DEL	-	OXYCODONE/ACETAMINOPHEN <sup>2,3</sup>	MC/DEL	8	LEVORPHANOL TARTRATE TAB		1
1	MC/DEL MC/DEL		OXYCODONE/ACETAMINOPHEN*** ROXICET	MC/DEL	8	LORCET	3. Only preferred	
.1	MC/DEL MC	_	ROXIDET ROXIPRIN TABS	MC/DEL MC	8	LORTAB	manufacturer's products will	
.1	MIC	1 ,	RUXIPRIIN TADO	MC		MAXIDONE TABS	be available without prior	
.1	י	1 ,	1	MC/DEL	8		authorization.	Please see the Pain Management Policy for the complete criteria
.1	ب	1 ,	1	MC/DEL		MEPERIDINE TABS NORCO TABS		Please see the Pain Management Policy for the complete Criteria
, <b>1</b>	י	1 ,	1	MC/DEL MC/DEL		NORCO TABS ONSOLIS		
.1	_{	1 '	1					1
<u>,                                    </u>		1 ,	1	MC/DEL		OXECTA  OXYCODONE CAP		1
<u>,                                    </u>	י	1 ,	1	MC/DEL		OXYCODONE (APAR 10/650		
<u>,                                    </u>		1 ,	1	MC/DEL		OXYCODONE/APAP 10/650		
<u>,                                    </u>		1 ,	1	MC/DEL		OXYCODONE/APAP 7.5/500		
, <b>1</b>		1 ,	1	MC/DEL		PENTAZOCINE/NALOVONE TARS		
<u>,                                    </u>		1 ,	1	MC/DEL		PENTAZOCINE/NALOXONE TABS		
.1	_{	1 '	1	MC		PERCOCET TABS		1
. <b>1</b>	_{	1 '	1	MC	8	PERCOCET TABS		1
<u>,                                    </u>	י	1 ,	1	MC	8	PHRENILIN W/CAFFEINE/CODE CAPS		
, <b>1</b>	י	1 ,	1	MC/DEL		ROXICET 5/500 TABS		
, <b>1</b>	י	1 ,	1	MC		ROXICODONE TABS		
<u>,                                    </u>	י	1 ,	1	MC/DEL		ROXYBOND		
<u>,</u>	י	1 ,	1	MC		SYNALGOS-DC CAPS		
<u>,                                    </u>	י	1 ,	1	MC	8	TALACEN TABS		
<u>,                                    </u>		1 ,	1	MC	8	TREZIX		
<u>,</u>	י	1 ,	1	MC	8	TYLENOL/CODEINE #3 TABS		
<u>,</u>	י	1 ,	1	MC	8	TYLOX CAPS		
<u>,                                    </u>	י	1 ,	1	MC	8	XOLOX	Use PA Form# 20420	
<u>,                                    </u>	י	1 ,	1	MC	8	VICODIN		
<u>,                                    </u>		1 ,	1	,	1		Use PA form #10300 for PAs	As
<u>,</u>	י	1 ,	1	MC		VICOPROFEN TABS	over the opiate limit	1
<b>4</b> [	י	1 ,	1	MC		ZYDONE TABS		
<u>,</u>	י	1 ,	1	MC	9	ACTIQ LPOP		
<b>4</b> [	י	1 ,	1	MC	9	CONZIP		
<b>4</b>	1 '	1 '		MC	9	OPANA		
OPIOID DEPENDENCE TREATMENTS	MC	1-	SUBOXONE FILM <sup>2</sup>	MC/DEL		BUPRENORPHINE <sup>1</sup>	+	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
<b>∡</b> 1	י ן	1 '		'	1			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
<b>∡</b> 1	י	1 ,	1	,	1		<u>Use PA Form #20100</u>	preferred drug(s) exists.
4	4	4	4	•		-	•	•

	MC/DEL	BUPRENORPHINE/NALOXONE TABS <sup>2</sup>	МС	ZUBSOLV	Buprenorphine will only be approved for use during pregnancy.     See Criteria Section	Members will continue to be required to follow the criteria listed below:  1-Induction period for 30 days  2-Max dose of 32 mg for induction  3-Max dose of 24 mg for maintenance  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  5- Should provide evidence of monthly monitoring including random pill counts, urine drug tests and use of Maine Prescription Monitoring Program reports.
						6- Buprenorphine monotherapy is preferred if member is pregnant and dose not > 24 mg day and pregnancy diagnosis is noted on the prescription.
EXTENDED RELEASE BUPRENORPHINE	MC MC	BRIXADI <sup>1</sup> SUBLOCADE <sup>1</sup>			Use PA form #20200 for Extended Release Buprenorphine  1. Clinical PA required.	Brixadi and Sublocade: The prescriber can attest (and medical record should document) that:
OPIOID WITHDRAWAL AGENTS			МС	LUCEMYRA <sup>1</sup>	Clinical PA for appropriate approved use and patient has documented contraindication to clonidine.     Use PA Form#20420	
		NARCOTIC ANTAGONISTS				
NARCOTIC - ANTAGONISTS	MC/DEL  MC  MC  MC  MC  MC	NALTREXONE HCL TABS  NALOXONE INJ NARCAN NS NALOXONE SPRAY OTC VIVITROL INJ ZIMHI  COX 2 / NSAIDS	MC MC MC MC/DEL	EVZIO  OPVEE <sup>2</sup> KLOXXADO  REVIA TABS <sup>1</sup>	1. Will only be approved for side effects experienced with generic that are not described in the literature as occurring with the brand version.  2. For the treatment of adult and pediatric patients 12 years of age and older.	
COV 2 INILIDITODS SELECTIVE /	MOUDEL	COX 2 / NSAIDS  CELECOXIB <sup>4,5</sup>	MC/DE!	OF FRREY 010045		Destarted drugs must be tried and failed due to look of officeas or intelegable side offices heavy and restarted drugs will be approved unless an accordable alliging to the control of th
COX 2 INHIBITORS - SELECTIVE / HIGHLY SELECTIVE	MC/DEL MC/DEL	KETOROLAC TROMETHAMINE <sup>2,3,5</sup>	MC/DEL MC/DEL	CELEBREX CAPS <sup>4,5</sup> MELOXICAM CAPS <sup>5</sup>	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

l =		-		•	-	Borrenes como se como
		NABUMETONE TABS <sup>5</sup>		MOBIC <sup>5</sup>		presented druggo) exists.
l j	MC/DEL	MELOXICAM TABS <sup>1,5</sup>	MC/DEL	MOBIC SUSP <sup>5</sup>	Meloxicam has dosing	<b>1</b>
l	]   '	1	MC/DEL		limits allowing one tablet	<u> </u>
l l	<b>                                     </b>	1	MC/DEL	QMIIZ ODT	daily of all strengths without	<b>1</b>
l j	<i>l</i> '	1		VIVLODEX	PA.	!
l I	]   '	1	1	1	Ketorolac Tromethamine	<u> </u>
l I	<b>                                     </b>	1	4	1	is indicated for the short term	.
l I	<b>                                     </b>	1	4		(up to 5 days) management	<b>1</b>
l I	<b>                                     </b>	1	4		of moderately severe acute	<b>1</b>
l I	<b>                                     </b>	1	4		pain that requires analgesic	<b>1</b>
l	<b>                                     </b>	1	4		at the opioid level in adults.  Not indicated for minor of	<b>1</b>
l I	l '	1	4 I		Not indicated for minor of chronic pain conditions.	<u> </u>
l I	]   '	1	1	1	Cilionic pain conditions.	<u> </u>
l I	]   '	1	1	1	<b>!</b>	<u> </u>
l	<b>                                     </b>	1	4	1	2 V-t has desing	<b>1</b>
l I	<b>                                     </b>	1	4	1	Ketorolac has dosing limits allowing 24 tablets for	<b>1</b>
l I	<b>                                     </b>	1	4		a 5 day supply every 30	<b>1</b>
l I	<b>                                     </b>	1	4		days.	<b>1</b>
l I	]   '	1	1		4. Dosing limits will be set at	<u> </u>
.l	]   '	1	1	1	a maximum of 400mg daily	<u> </u>
l I	]   '	1	1	1	5. The FDA has issued a	<u> </u>
l I	<b>                                     </b>	1	4		Public Health Advisory	<b>1</b>
l	]   '	1	1		warning of the potential for	<u> </u>
l I	<b>                                     </b>	1	4		increased cardiovascular risk & GI bleeding with NSAID	<b>1</b>
l	]   '	1	1	1	& GI bleeding with NoAiD	<u> </u>
l	]   '	1	1	1	use.	<u> </u>
.l	]   '	1	1	1	<b>!</b>	<u> </u>
, I	<i>l</i> '	1	<i>i</i> [	1	<b>!</b>	!
20100	***	SOUR BREWS IRVINDOFFN	<del></del>	7		
NSAIDS		CHILDRENS IBUPROFEN	MC			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
, I		DICLOFENAC POTASSIUM TABS	MC MC		warning of the potential for	preferred drug(s) exists.
, I		DICLOFENAC SODIUM TABS		ANAI NOX DO TADO		Approvals will be granted for other requests based on failure of at least one generic NSAID from at least 3 different NSAID classes as described in the COX-II PA form.
, I		DICLOFENAC SODIUM 1% GEL <sup>1</sup>	MC	CAMBIA	& GI bleeding with NSAID	
.l		ETODOLAC	MC/DEL	CATAFLAM TABS	use.	
. I		FENOPROFEN CALCIUM TABS	MC	CHILDRENS ADVIL SUSP	· [	
, I		FLURBIPROFEN TABS	MC	CHILD'S IBUPROFEN SUSP	· [	
, I		IBUPROFEN	MC/DEL	CHILDREN'S MOTRIN SUSP	Dosing limits apply,	
<i>i</i> I		INDOMETHACIN	MC/DEL		please see Dosage	<b>1</b>
, I		KETOPROFEN	MC/DEL		Consolidation List.	
.l		MECLOFENAMATE SODIUM CAPS	MC/DEL	DICLFENAC GEL	<b>!</b>	DDI: Diclofenac will now be non-preferred and require prior authorization if it is currently being used in combination with lescol.
, I		NAPROSYN SUSP		EC-NAPROSYN TBEC	<b>!</b>	
, I	MC/DEL	NAPROXEN SUSP	MC/DEL	ETODOLAC ER 600MG	Use PA Form# 20420	The FDA has issued a Public Health Advisory warning of the potential for increased cardiovascular risk & GI bleeding with NSAID use.
, I	MC/DEL	NAPROXEN TABS	MC	FELDENE CAPS	<b>!</b>	
, I	MC/DEL	NAPROXEN SODIUM TABS	MC/DEL	FLECTOR PATCH	<b>!</b>	
<i>i</i> I	MC/DEL	NAPROXEN SODIUM CAPS	MC/DEL	IBU-200	· [	<b> </b>
, I	MC/DEL	NAPROXEN DR TBEC	MC	INDOCIN	· [	
, I		OXAPROZIN TABS	MC	LICART	· [	
, I		SULINDAC TABS	MC/DEL	LODINE	· [	
, I	**	TOLMETIN SODIUM	MC	LOFENA	· [	
.l I		VOLTAREN GEL		MOTRIN	<b>!</b>	
, I	MIC/DEL	VOETAKEN GEE	MC	NALFON CAPS	<b>!</b>	
. I	<b>                                     </b>	1	MC/DEL	NAPRELAN TBCR	· [	
. I	<b>                                     </b>	1		NAPROSYN TABS	· [	
, I	]   '	1		NAPROSTN TABS NAPROXEN SODIUM TBCR	<b>!</b>	
.	/ I '	1	MC/DEL		<b>!</b>	
i l	7 I		9			
	1	1	MC	PENNSAID		
			MC/DEL	PIROXICAM CAPS		
			MC/DEL MC	PIROXICAM CAPS PONSTEL CAPS		
			MC/DEL MC	PIROXICAM CAPS		

	1 1		MC		SB IBUPROFEN TABS		
			MC		SPRIX		
			MC		TIVORBEX		
			MC		TOLECTIN		
			MC		V-R IBUPROFEN TABS		
			MC		ZORVOLEX		
NSAID - PPI	+		MC		PREVACID NAPRA-PAC	Use a preferred NSAID	
NOAID - FFI			MC/DEL		VIMOVO <sup>1</sup>	and PPI separately.	
			WIC/DEL		VIIIIOVO	Use PA Form# 20420	
		RHEUMATOID ARTHRITIS				036 1 A 1 0111# 20420	
RHEUMATOID ARTHRITIS	MC/DEL	ACTEMRA VIALS	1		ADALIMUMAB-AACF	Use PA Form# 20900	See criteria as listed on Rheumatoid Arthritis PA form.
Taribonist oils Attributio		ACTEMRA SYRINGES				USE PA FOITH 20900	de difficilità de la dece di l'information d'utilità d'information de la dece de la companya de la dece de la companya de la c
	MC/DEL	ADALIMUMAB-FKJP <sup>3</sup>	MC MC/DEL		AMJEVITA <b>ARAVA</b>	1 Desire limite and	
	MC/DEL					<ol> <li>Dosing limits apply.</li> <li>Please see dose</li> </ol>	Preferred injectable products allowed without PA if trial of a preferred oral agents (azathioprine, hydroxychloroquine, leflunomide, methotrexate, sulfasalazine tabs) are seen in the
	MC	AVSOLA	MC/DEL		CIMZIA	consolidation list.	members drug profile. Dosing limits apply.
	MC/DEL	AZATHIOPRINE	MC/DEL		CYLTEZO		
	MC	ENBREL <sup>2</sup>	MC/DEL		ENTYVIO	<ol><li>Established users will be grandfathered.</li></ol>	
	MC	ENBREL SURECLICK <sup>2</sup>	MC		HADLIMA	Ť	
	MC	KINERET SOLN	MC/DEL		HULIO	3.Clinical PA is required to	
	MC/DEL	LEFLUNOMIDE	MC/DEL		HYDROXYCHLOROQUINE <sup>2</sup>	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
	MC/DEL	METHOTREXATE	MC/DEL		HYRIMOZ		with biologic DMARDs or potent Immunosuppressants.
	MC	ORENCIA	MC		IDACIO	<ol><li>Verification of age for</li></ol>	
	MC/DEL	SULFASALAZINE TABS	MC/DEL		ILARIS <sup>1,3,4</sup>	appropriate indication.	Jylamvo will require using preferred methotrexate if unable please provide clinical rational as why inappropriate.
	MC	SIMLANDI <sup>3</sup>	MC/DEL		INFLECTRA	<ol><li>Treatment failure or</li></ol>	
-	MC	SIMPONI PEN	MC		INFLIXIMAB VIAL	intolerance to other forms of preferred methotrexate	Zymfentra: In adults for maintenance treatment of:
	MC	SIMPONI AUTOINJECTOR	MC		JYLAMVO	preferred methotrexate	Moderately to severely active ulcerative colitis following treatment with an infliximab product administered intravenously.
	MC/DEL	RINVOQ <sup>3</sup>	MC/DEL		KEVZARA		Moderately to severely active Crohn's disease following treatment with an infliximab product administered intravenously.
	MC	HUMIRA <sup>1,2</sup>	MC		OLUMIANT		
	MC/DEL	XELJANZ <sup>3,6</sup>	MC		OMVOH	6. See criteria section	
	MC/DEL	XELJANZ XR	MC		OTREXUP		
	MC/DEL	XELJANZ XR SOL	MC		RASUVO <sup>7</sup>		
			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
			MC		YUSIMRY		Xeljanz® XR with potent CYP3A4 inducers (e.g. rifampin) is not recommended
					XATMEP <sup>5</sup>		
			MC				
		11 005011 105171 105170	MC		ZYMFENTRA		l .
ALODEOIA ADEATA ACENTO		ALOPECIA AREATA AGENTS	1 110	7	OLUMIANT		
ALOPECIA AREATA AGENTS					OLUMIANT		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
			MC/DEL	8	LITFULO		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	
						330 17(1 OIIIIII 20420	
MISCELLANEOUS ARTHRITIS							
	MC	RIDAURA CAPS	MC/DEL	1	ADTUDOTEC <sup>1</sup>	The individual	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 expension is effected an
ARTHRITIS - MISC.	MC MC	MYOCHRYSINE SOLN	WIG/DEL		ARTHROTEC <sup>1</sup>		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
	WIC	IVITOGRATISINE SOLIV				available without PA.	preferred drug(s) exists. The individual components of Arthrotec are available without PA.
	1 1		J		I	Use PA Form# 20420	

				LUPUS-SLE				
LUPUS-SLE					MC MC MC	 BENLYSTA <sup>1</sup> LUPKYNIS SAPHNELO	Use PA Form# 20420  1. Approvals will require previous trial of corticosteroids, antimalarials	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.
							NSAIDS and immunosuppressives.	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)
				PIK3CA-Related Overgrowth Spectrum (P	PROS)			
PIK3CA-Related Overgrowth Spectrum (PROS)					MC	VIJOICE <sup>1</sup>	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.
				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		D	DIVALPROEX ER TB24	MC	DEPAKOTE ER TB24	Use PA Form# 10110	
MIGRAINE - SELECTIVE SEROTONIN AGONISTS (5HT)Tabs/Nasal  MIGRAINE - SELECTIVE SEROTONIN AGONISTS (5HT)Injectables	MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	1 1 1 1 1 1 1 2	1 R R R R S 1 Z N N	MIGRANAL NASAL SPRAY RELPAX¹ RIZATRIPTAN ODT RIZATRIPTAN TABS SUMATRIPTAN TABS¹ ZOLMITRIPTAN TAB¹ NARATRIPTAN HCI TABS¹  IMITREX CARTRIDGE¹ SUMATRIPTAN SYRINGE¹ SUMATRIPTAN PEN INJCTR¹	MC MC MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	AMERGE TABS <sup>1,2</sup> AXERT TABS <sup>1,2</sup> FROVA TABS <sup>1,2</sup> IMITREX NASAL SPRAY <sup>1</sup> IMITREX TABS <sup>1,2</sup> MAXALT **12,3  MAXALT **MLT <sup>1,2,3</sup> ONZETRA XSAIL*  SUMATRIPTAN NASAL SPRAY <sup>1</sup> ZOLMITRIPTAN ODT  ZOLMITRIPTAN SPRAY  ZOMIG TABS <sup>1,2</sup> ZOMIG TASS <sup>1,2</sup> ZOMIG ZMT TBDP <sup>1,2</sup> TOSYMRA  ZEMBRACE <sup>1</sup> IMITREX PEN INJCTR <sup>1</sup>	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  1. Dosing limits apply. Please refer to the dose consolidation table.	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)Combinations					MC/DEL	TREXIMET <sup>1,2</sup>	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/DEL MC/DEL		A	AIMOVIG¹ AJOVY¹ AJOVY AUTO INJCT¹	MC MC MC	NURTEC ODT <sup>2</sup> QULIPTA VYEPTI <sup>2</sup>	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 <sup>1</sup> 200mg/ml EMGALITY PEN <sup>1</sup>			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.  Ubrelvy 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 <sup>1</sup> SPASTRIN TABS	MC MC MC/DEL MC/DEL MC	BELCOMP-PB SUPP ELYXYB MIGRAZONE CAPS MIGERGOT SUP REYVOW UBRELVY	Dosing limits apply, please see the dose consolidation list.	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.  Ubrelvy 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
			MC/DEL	ZAVZPRET	Use PA Form# 10110	Natice OD 1 Will be professed after 2 acceptable strains of at least two professed after so
GOUT	MC/DEL MC/DEL MC/DEL MC MC/DEL MC 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 ULORIC¹ 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 <sup>1,2</sup>	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 <sup>1</sup>	I.Indicated to decrease the need for red blood cell transfusion due to hemolysis in adults with cold agglutinin disease (CAD).	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 spreferred 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)				OXLUMO¹ RIVFLOZA	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

	l I	Ī						prescribed by, or in consultation, with a nephrologist or urologist
SICKLE CELL DISEASE	MC MC/DEL MC		DROXIA HYDROXYUREA  LYFGENIA <sup>2,3</sup>	MC MC MC		ADAKVEO CASGEVY <sup>23</sup> ENDARI <sup>1</sup>		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		SIKLOS	2. For the treatment of patients ≥ 12 years of age.  3. PA required to confirm FDA approved indication. <u>Use PA Form# 20420</u>	
HUTCHINSON- GILFORD PROGERIA SYNDROME (HGPS)				МС		ZOKINVY <sup>12</sup>	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					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				МС		JOENJA <sup>1,2,3</sup>	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	1.Clinical PA required for	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					
ANTICONVULSANTS	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
MC	CARBAMAZEPINE ER CAP	MC	8	CARBAMAZEPINE SUS	7 til Holl protottod mode made	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
MC/DEL	CARBATROL CP12	MC	8	DEPAKOTE	be used in specified order	preferred drug(s) exists.
MC/DEL	CELONTIN CAPS	MC	8	DEPAKOTE ER		
MC/DEL	CLOBAZAM	MC	8	DIACOMIT		
MC/DEL	CLONAZEPAM TABS	MC/DEL	8	DIVALPROEX SODIUM SPRINKLE CAPS	1. Quantity limit. 5/month	
MC	DEPAKOTE SPRINKLES CPSP	МС	8	ELEPSIA XR <sup>9</sup>	2. Dosing limits apply,	
MC/DEL	DIAZEPAM GEL <sup>1</sup>	MC	8	EPRONTIA SOLN <sup>10</sup>	please see dose	
MC/DEL	DILANTIN	MC/DEL	8	FELBATOL	consolidation list.	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,
MC/DEL	DIVALPROEX SODIUM	MC/DEL	8	FELBATOL SUS	3. Dosing limits apply per	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
MC	DIVALPROEX SPRINKLE CAP	MC/DEL	8	FELBAMATE SUS	strength as well as a	been tried and failed at full therapeutic doses for adequate durations (at least two weeks).
MC/DEL	EPIDIOLEX <sup>7</sup>	MC		FINTEPLA <sup>8</sup>	maximum daily dose of	
MC/DEL	EPITOL TABS	MC	0	FYCOMPA <sup>2</sup>	600mg. Please see dose	
MC/DEL	ETHOSUXIMIDE SYRP	MC/DEL	ο ,	HORIZANT	consolidation list.	
MC/DEL	EQUETRO	MC	0	GRALISE	4. Adjunctive therapy 17 and	ATT OFF CHAPT AT END OF DOCUMENT
MC/DEL	GABAPENTIN <sup>2</sup> CAP	MC/DEL	0	KEPPRA TABS	older.	*** SEE CHART AT END OF DOCUMENT
	GABAPENTIN <sup>2</sup> TAB		0	KEPPRA SOLN	E May doos 2400mg	
MC/DEL		MC/DEL	ŏ		5. Max dose 2400mg	
MC/DEL	GABAPENTIN SOL	MC/DEL	8	KLONOPIN TABS	<ol><li>Clinical PA required for appropriate diagnosis</li></ol>	
MC/DEL	GABITRIL TABS	MC	8	LAMICTAL IR	appropriate diagnosis	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	
MC	LAMICTAL CHEW	MC/DEL	8	LEVETIRACETAM INJ	treatment of seizures	All non-preferred meds must be used in specified order.
MC/DEL	LAMOTRIGINE ER ODT	MC	8	LIBERVANT	associated with Lennox-	
MC/DEL	LAMOTRIGINE IR <sup>2</sup>	MC/DEL	8	LYRICA CR	Gastaut syndrome (LGS),	
MC/DEL	LAMOTRIGINE XR	MC/DEL	8	LYRICA SOL <sup>3</sup>	Dravet syndrome (DS) or TS (Tuberous Sclerosis	Please use Drug-Drug Interaction PA form #10400 for this combination.
MC/DEL	LEVETIRACETAM SOLN	MC	8	MOTPOLY XR	Complex) in patients 1	
MC/DEL	LEVETIRACETAM TABS	MC/DEL	8	MYSOLINE TABS	years of age and older.	
MC/DEL	LEVETIRACETAM ER TABS	MC	8	ONFI	8. For seizures associated	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	LYRICA <sup>3</sup>	MC/DEL	8	OXCARBAZEPINE SUS	with Dravet syndrome in	
MC/DEL	NAYZILAM <sup>1</sup>	МС	8	OXTELLAR XR⁵	patients 2 years of age and older	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
MC/DEL	OXCARBAZEPINE	MC/DEL	8	PHENYTEK CAPS	oldei	use of Diacomit® as monotherapy in DS.
MC/DEL	PREGABALIN CAPS	MC/DEL	8	POTIGA	9. Adjunctive therapy 12	
MC/DEL	PHENYTOIN	MC/DEL	0	PREGABALIN (ORAL) SOL	and older.	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,
MC/DEL	PRIMIDONE TABS		0	ROWEEPRA TAB		CYP3A4, or CYP2C19 inducers, such as rifampin, phenytoin, phenobarbital, and carbamazepine) should be avoided, or dosage adjustments should be made.
MC/DEL		MC MC	8 8	ROWEEPRA TAB SABRIL		
MC/DEL MC/DEL	QUDEXY XR TEGRETOL SUS		ŏ	SEZABY		DDI: Avaid appearitant use of Neurilana® with moderate or strong CVP2A inhibitors
MC/DEL MC/DEL	TOPIRAMATE	MC MC	o o	SEZABY SPRITAM	10. Initial monotherapy for	DDI: Avoid concomitant use of Nayzilam® with moderate or strong CYP3A inhibitors.
MC/DEL	TOPIRAMATE SPRINKLE IR CAPS	MC	0	SYMPAZAN	the treatment of partial-onset	
WC/DEL WC/DEL	TRILEPTAL SUS	MC/DEL	ο Ω	TEGRETOL TAB	or primary generalized tonic-	Xcopri criteria: History of trials with at least 4 AEDs (2 generic, 2 branded or Uncontrolled seizures on three AEDs; or Uncontrolled on 2 AEDs given along with VNS. Uncontrolled defined as 3 or more TC seizures per year (increases risk of SUDEP); > 6 disabling seizures per year. Any patient who has gone to the ED 2 or more times in the prior 12 months (who has also
MC/DEL	VALPROIC ACID TABS	MC/DEL	Ω	TIAGABINE	cionic scizures in patients z	tried and failed at least 3 other drugs). Ongoing use requires 50 percent reduction in seizure frequency after three months.
MC/DEL	VALPROIC ACID TABS	MC/DEL	8	TOPAMAX	years of age and older. Adjunctive therapy for the	· · · · · · · · · · · · · · · · · · ·
MC	VALTOCO <sup>2</sup>	MC/DEL	8	TOPIRAMATE ER CAPS	treatment of partial-onset	Motpoly XR: pediatric patient weight must be > 50kg and requires multiple preferred medication trials including generic lacosamide
MC/DEL	ZONISAMIDE	MC	8	TOPAMAX SPRINKLE ER CAPS <sup>2</sup>	seizures, primary	morpory zura: positiano patient maignit must de z dong and requires multiple preferied medicalitori trials including generic lacosamilide
	20110/ WIBE	MC	8	TOPAMAX SPRINKLE ER CAPS  TOPAMAX SPRINKLE IR CAPS <sup>2</sup>	generalized tonic-clonic	
		MC/DEL	8	TOPAMAX SPRINKLE IR CAPS <sup>2</sup> TOPIRAMATE SPRINKLE ER CAPS <sup>2</sup>	seizures, and seizures associated with Lennox	
		MC	8	TROKENDI <sup>2,6</sup>	Gastaut syndrome in	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
		MC	8	VIGAFYDE	patients 2 years of age and	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/DEL	8	VIMPAT*	older. The preventive	
		MC/DEL	8	VIMPAT SOL <sup>4</sup>	treatment of migraine in	
		MC	8	XCOPRI	patients 12 years and older.	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	ZARONTIN SYRP	Will require a step though topiramate.	VISIUI 1095.
		MC/DEL	8	ZARONTIN STRF ZARONTIN CAP	юрнанасе.	
		MC/DEL	8	ZARONTIN GAI ZARONTIN SOL		
		MC	8	ZONISADE		
		WIC	0	ZONIOADL		
					ī	

	<i>i</i>	, J		MC/DEL		NEURONTIN	·	
	<i>i</i>	, J		MC/DEL	9	TEGRETOL-XR TB12	'	1 <b>7</b>
	<i>i</i> ]	, J		1 1	( '	1	'	1
	<i>i</i>	, J		1 1	( '	1	SEE ANTICONVULSANT	1
	<i>i</i> ]	, J		1 '	( )	1	INDICATION CHART AT THE END OF THIS	1
	ı J	, J	. J	1 '	( )	BIPOLAR DISORDER: STEP ORDER	DOCUMENT	1
	ı J	, J	<u>.</u>	1 '	M ~ A	BIPOLAK DISOKDEK: STEP OKDEN	M= Monotherapy A= Adjunctive	1
	ı J	, J	. J			LAMICTAL	9= No Evidence	1
	ı J	, J		1 '	4 ~ 4	LITHIUM	The step orders show the relative strength of evidence	1
	ı J	, J	. J			Carbamazepine Valproate	for use in bi-polar and will	1
	ı J	, J	, J			VALPROATE ATYPICAL ANTIPSYCHOTICS EXC. CLOZAPINE	guide prior authorization determinations.	1
	ı J	, J	. J			TRILEPTAL	Step 4 drugs-no PA required.	1
	ı J	, J	. J	Ş	9 ~ 6	TOPAMAX	·   · · · · · · · · · · · · · · · · · ·	1
	ı J	, J	. J			KEPPRA TABS	·   · · · · · · · · · · · · · · · · · ·	1
	ı J	, J	<u>.</u>			GABITRIL TABS NEURONTIN	·  '	1
	ı J	, J	<u>.</u>	1 '	( '	1	·   '	1
	ı J	, J	<u>.</u>	1 '	( '	1	·   '	1
	ı J	, J	<u>.</u>	1 '	( '	1	·   · · · · · · · · · · · · · · · · · ·	1
	ı J	, J	<u>.</u>	1 '	( )	1	·  '	1
	ı J	, J	<u>.</u>	1 '	( )	1	·  '	1
	ı J	, J	. J	1 '	( )	PEDIATRIC BIPOLAR1 DISORDER: STEP ORDER	Two-step 1 preferred drugs must be tried before	1
	ı J	, J	. J	1 '	t !		Trileptal.	1
	ı J	, J	. J	1 "		(6-18 YEARS WITH OR WITHOUT PSYCHOSIS) LITHIUM	The step orders show the relative strength of evidence	1
	ı J	, J	. J	1 '		LTHIUM CARBAMAZEPINE	for use in bi-polar and will	1
	ı J	, J	. J	1 '	4 ~ 4	VALPROATE	guide prior authorization determinations.	
	ı J	, J	. J	1 '		ATYPICAL ANTIPSYCHOTICS EXC.CLOZAPINE	Step 4 drugs-no PA required.	.1
	ı J	, J	<u>.</u>	1 "		Lamictal Trilepta	·   · · · · · · · · · · · · · · · · · ·	
			ANTI-PARKINSON DRUGS		) ~ 0	IRILEPTA		
PARKINSONS - ANTICHOLINERGICS	MC/DEL		BENZTROPINE MESYLATE TABS				Use PA Form# 20420	
	MC		COGENTIN SOLN	1 '	( '	1	,	
PARKINSONS - ADENOSINE RECEPTOR	MC/DEL	<u> </u> '	TRIHEXYPHENIDYL	MC/DEL	<b>←</b>	NOURIANZ	<b></b> '	The form of the second field due to lock of officers or intelerable cities affects before non professed drups will be approved unless an appeniable clinical expension is offered on
ANTAGONIST	ı J	, J	<u>.</u>	MC/DEL	( )	NOURIANZ	1	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
	ı J	, J	. J	1 '	( )	1	·   · · · · · · · · · · · · · · · · · ·	preferred drug(s) exists.
	1 I	, J	. J	1 '	( )	1	·   · · · · · · · · · · · · · · · · · ·	
4 <b>1</b> ,	1			4	4 *	1		DDI: Avoid use of Nourianz® with strong CYP3A4 inducers (e.g. carbamazepine, rifampin, phenytoin, St. John's wort).
	'	` <u> </u>	`	١ ،	١ ،			· ·
				$^{f ar{L}}$	L		Use PA Form# 20420	
PARKINSONS - COMT INHIBITORS				MC/DEL		COMTAN TABS	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 Drieg Authorization form such as the presence of a condition that received the preferred drug as a similar product of the product of the preferred drug as a similar product of the product of the product of the product of the product of the product of the produ
PARKINSONS - COMT INHIBITORS				MC		ONGENTYS	Use PA Form# 20420	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
PARKINSONS - COMT INHIBITORS							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.
PARKINSONS - COMT INHIBITORS				MC		ONGENTYS	Use PA Form# 20420	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
PARKINSONS - COMT INHIBITORS				MC		ONGENTYS	Use PA Form# 20420	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
PARKINSONS - COMT INHIBITORS  PARKINSONS - SELECTED DOPAMIN	MC/DEL	P	PRAMIPEXOLE	MC MC/DEL		ONGENTYS	Use PA Form# 20420	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

AGUNISTS	MC/DEL	ROPINIROLE	MC/DEL MC/DEL	8 8 8	REQUIP TABS MIRAPEX ER NEUPRO PATCH	<ol> <li>As of 12/08 users of Mirapex will be grandfathered if diagnosis is Parkinson's.</li> </ol>	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	AMANTADINE HCLCAPS  AMANTADINE HCL TABS  BROMOCRIPTINE MESYLATE TABS  BROMOCRIPTINE MESYLATE CAPS	MC/DEL MC MC/DEL		APOKYN AZILECT <sup>2</sup> Carbidopa/Levodopa Rapdis	<ol> <li>Approvals will require concurrent therapy with Levodopa and failed trials of Selegiline, Comtan, and Stalevo.</li> </ol>	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 f preferred drug(s) exists.
					CREXONT <sup>4</sup>		
	MC/DEL MC/DEL MC/DEL MC/DEL MC	CARBIDOPA/LEVODOPA TABS <sup>3</sup> CARBIDOPA/LEVODOPA ER CARBIDOPA/LEVO/ENTACAPONE TAB LARODOPA TABS	MC MC MC MC/DEL MC		ELDEPRYL CAPS GOCOVRI INBRIJA KYNMOBI	Approvals will require trials of Carbidopa/Levodopa, Selegiline, Comtan, and Stalevo.	Inbrija is recommended for the intermittent treatment of OFF episodes in patients with Parkinson's disease treated with carbidopa/levodopa.
	MC/DEL MC/DEL	SELEGILINE CAPS HCL SELEGILINE TABS HCL	MC MC		LODOSYN TABS  OSMOLEX ER	Only preferred     manufacturer's products will     be available without prior     authorization.	
			MC/DEL MC/DEL MC MC		PARLODEL CAPS PARLODEL TABS RYTARY SINEMET TABS	Approvals will require trials of preferred medications including extended-release	
			MC MC		SINEMET TBCR ZELAPAR <sup>1</sup>	levodopa/carbidopa tablets  Use PA Form# 20420	
PARKINSONS - COMBO.			MC/DEL MC		STALEVO <sup>1</sup> CARBIDOPA/LEVODOPA/ENTACA <sup>1</sup>	Use PA Form# 20420 1.Clinical PA is required to establish diagnosis and medical necessity.	
		MUSCLE RELAXANTS					
MUSCLE RELAXANTS	MC/DEL	BACLOFEN TABS	MC/DEL	7	ORPHENADRINE CITRATE		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
	MC/DEL MC/DEL	CHLORZOXAZONE TABS CYCLOBENZAPRINE HCL 5mg & 10mg TABS	MC/DEL	8	CARISOPRODOL 350MG TABS AMRIX		preferred drug(s) exists.
	MC MC/DEL MC/DEL	LIORESAL INTRATHECAL KIT METHOCARBAMOL TABS TIZANIDINE HCL TABS	MC/DEL MC MC MC MC MC MC MC MC MC MC MC MC MC	8 8 8 8 8 8 8	DANTRIUM CAPS FLEQSUVY LIORESAL TABS LORZONE LYVISPAH METAXALONE NORFLEX TBCR OZOBAX ROBAXIN-750 TABS VECUROMIUM INJ ZANAFLEX TABS		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/DEL MC/DEL MC/DEL MC/DEL	9 9 9	CARISOPRODOL 250MG TABS CHLORZOXAZONE 250mg TABS SKELAXIN TAB SOMA TABS		

		Ī	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.	1 1	1	MC/DEL		CARISOPRODOL/ASPIRIN TABS	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
			MC/DEL		CARISOPRODOL/ASPIRIN/CODE		to reports of misplacement stolen, dropped in toilet or sink, distant travel, etc.
			MC		NORGESIC TABS		
			MC/DEL		ORPHENADRINE COMPOUND		
			MC/DEL		ORPHENADRINE/ASA/CAFF		
ı <b>İ</b>			MC		ORPHENGESIC		
		PARATHYROID	HORMONE		•		
PARATHYOID HORMONE	Т		MC		NATPARA <sup>1</sup>	1. Recommended only for	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		YORVIPATH <sup>1</sup>	those who cannot be well-	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
						controlled on calcium supplements and active	preferred drug(s) exists.
						forms of vitamin D alone.	
						Use PA Form# 20420	
		VITAMINS					
VITAMINS	MC	CYANOCOBALAMIN SOLN	MC		AQUASOL E 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
	MC	FERIVA CAP	MC		AQUAVIT-E SOLN	Please refer to OTC list for	r 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	FERIVAFA CAP	MC		DHT SOLN	covered products.	pretented drug(s) exists. Certain drugs require specific diagnoses for approval.
	MC/DEL	FOLIC ACID TABS	MC		FUSION PLUS CAP		
	MC/DEL	MEPHYTON TABS			HEMOCYTE PLU CAP		
			MC			Click here for the OTC List	
	MC/DEL	NIACIN	MC		INTEGRA CAP		
	MC	NIACOR TABS	MC		INTEGRA F CAP		
	MC/DEL	NICOTINIC ACID SR CPCR	MC		INTEGRA PLUS CAP		
	MC	PYRIDOXINE HCL TABS	MC		NASCOBAL GEL		
	MC	TANDEM CAP	MC		TANDEM PLUS CAP		
	MC/DEL	THIAMINE HCL SOLN					
	MC/DEL	VITAMIN B-1 TABS					Please refer to OTC list for covered products.
	MC/DEL	VITAMIN B-12					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
	MC	VITAMIN B-6 TABS					PPI. PPI.
	MC/DEL	VITAMIN C					
	MC/DEL	VITAMIN E CAPS					Preferred products that used to require diag codes still require diag codes unless indicated otherwise.
	MC/DEL	VITAMIN E/D-ALPHA CAPS					
	МС	VITAMIN K1 SOLN					
	MC	V-R VITAMIN E CAPS					
VITAMIN D's	MC/DEL	CALCITRIOL CAPS <sup>1</sup>	MC		CALCIJEX	Diagnosis of dialysis	Preferred products require dialysis/renal failure diagnosis.
	MC/DEL	ROCALTROL	MC/DEL		DOXERCALCIF CAP	(renal failure) required.	
	MC/DEL	VITAMIN D2 <sup>2</sup>	MC/DEL		DOXERCALCIF INJ	2. Only specific NDCs	
	MC/DEL	VITAMIN D3 <sup>2</sup>	MC/DEL		PARICALCITROL CAP	available	
	MC/DEL	VITAMIN DROPS	MC/DEL		PARICALCITROL INJ		
	MC	PARICALCITOL CAPS	MC/DEL		HECTOROL (ORAL)		
			MC/DEL		HECTOROL (PARENTERAL)		Rayaldee requires clinical PA to verify stage 3 or 4 CKD.
			MC		RAYALDEE		
			MC		ZEMPLAR INJ		
			MC		ZEMPLAR CAPS	Use PA Form# 20420	
		EMZYMES					
POMPE DISEASE AGENTS			MC		NEXVIAZYME <sup>1</sup>		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
			MC		LUMIZYME		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		OPFOLDA	For patients 1 year of age     and older with late enset	g another or any aria the previous analys) exists.
			MC		POMBILITI	and older with late-onset Pompe disease (lysosomal	
						acid alpha-glucosidase	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
						[GAA] deficiency).	improving on their current enzyme replacement therapy (ERT).
=		<del>-</del>	- '	=	=	-	-

					Use PA Form# 20420	
		MISC MULTI-VITAMINS				
MINS - MISC.	MC	CENTRUM TABS	MC	ADEKS	Diag codes are no longer	
	MC	CENTRUM JR/IRON CHEW	MC/DEL	ADVANCED NATALCARE TABS	required on prenatal	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
	MC	CENTRUM-LUTEIN TABS	MC	AQUADEKS	vitamins.	preferred drug(s) exists. Certain drugs require specific diagnoses for approval.
	MC	CEROVITE ADVANCED FO TABS	MC	CENTRUM JR/EXTRA C CHEW	Please refer to OTC list.	
	MC/DEL	CHEWABLE MULTIVIT/FL CHEW	MC	CENTRUM PERFORMANCE TABS		Please refer to OTC list.
	MC	COD LIVER OIL CAPS	MC	CENTRUM SILVER TABS	Use PA Form# 20420	
			MC	DALYVITE LIQD		Preferred products that used to require diag codes still require diag codes unless indicated otherwise.
	MC/DEL	COMPLETE NATAL DHA (ORAL) COMBO PKG				
	MC	COMPLETE SENIOR TABS	MC	EMBREX 600 MISC		
	MC	DAILY MULTI VIT/IRON				
			MC	FERRALET 90	Click here for the OTC List	
	MC/DEL	DIALYVITE 1MG	MC	IBERET		
	MC/DEL	DIALYVITE 800MG	MC	MATERNA TABS		
	MC/DEL	FULL SPECTRUM B	MC	MAXARON		
	MC	M.V.I12 INJ	MC	MULTIRET FOLIC -500 TBCR		
	MC	MULTI-VIT/FLUORIDE	MC/DEL	NATAFORT TABS		
	MC/DEL	NATALCARE RX TABS	MC/DEL	NATALCARE CFE 60 TABS <sup>1</sup>		
	MC/DEL	NEPHRONEX	MC/DEL	NATALCARE CHE 60 TABS  NATALCARE GLOSS TABS <sup>1</sup>		
	MC/DEL		MC			
	MC/DEL	NIVA-PLUS (ORAL) TABLET ONE DAILY TABS	MC	NATAL CARE PIC FORTE TARGE		
	MC/DEL	ONE-DAILY MULTIVITAMINS	MC/DEL	NATALCARE PIC FORTE TABS <sup>1</sup>		
	MC/DEL			NATALCARE PLUS TABS <sup>1</sup>		
		ONE-TABLET-DAILY	MC MC/DEL	NATACHEM CHEM		
	MC/DEL	POLY-VITAMIN/FLUORIDE SOLN	MC/DEL	NATALFIRET TARE		
	MC/DEL	POLY-VITAMIN/FLUORIDE SOLN	MC	NATALFIRST TABS		
	MC/DEL	POLY-VITAMINS/IRON SOLN	MC	NATATAB RX TABS		
	MC	PRENATA (ORAL) TAB CHEW	MC/DEL	NEPHPLEX RX TABS		
	MC/DEL	PRENATAL TABS <sup>1</sup>	MC/DEL	NEPHROCAPS CAPS		
	MC/DEL	PRENATAL FORMULA 3 <sub>2</sub> TABS <sup>1</sup>	MC/DEL	NEPHRO-VITE TABS		
	MC/DEL	PRENATAL PLUS TABS <sup>1</sup>	MC	NESTABS RX TABS		
	MC/DEL	PRENATAL PLUS NF TABS <sup>1</sup>	MC/DEL	NIFEREX		
	MC	PRENATAL PLUS/27MG IRON <sup>1</sup>	MC/DEL	OCUVITE TABS		
	MC	PRENATAL PLUS/IRON TABS <sup>1</sup>	MC	POLY-VI-FLOR SOLN		
	MC	PRENATAL VITAMIN PLUS LOW IRON (ORAL) TA		POLY-VI-SOL SOLN		
	MC/DEL	PRENATAL RX/BETA-CAROTENE <sup>1</sup>	MC	POLY-VI-SOL/IRON SOLN		
	MC/DEL	PREPLUS (ORAL) TABLET	MC	POLY-VITAMIN DROPS SOLN		
	MC/DEL	RENAL CAPS	MC	PRECARE		
	MC/DEL	RENAPHRO CAPS	MC	PREFERA OB		
	MC	STRESS TAB NF TABS	MC	PREMESIS RX TABS		
	MC	THERAPEUTIC-M TABS	MC	PRENATABS CBF TABS <sup>1</sup>		
	MC	THERAVITE LIQD	MC	PRENATAL CARE TABS <sup>1</sup>		
	MC/DEL	TRINATAL RX 1 (ORAL) TABLET	MC	PRENATAL MR 90 TBCR <sup>1</sup>		
	MC/DEL	TRIVEEN-DUO DHA (ORAL) COMBO. PKG	MC/DEL	PRENATAL MTR/SELENIUM TABS <sup>1</sup>		
	MC/DEL	TRI-VITAMIN/FLUORIDE SOLN	MC	PRENATAL OPTIMA ADVANCE TABS <sup>1</sup>		
	MC	VITA CON FORTE CAPS	MC	PRENATAL PC 40 TABS <sup>1</sup>		
	MC	VITAPLEX PLUS TABS	MC/DEL	PRENATAL RX TABS <sup>1</sup>		
			MC	PRENATE <sup>1</sup>		
			MC	PRENATE ELITE <sup>1</sup>		
		1		PREMATE ELITE PRIMACARE MISC		
			MC MC	PROTEGRA CAPS		
			MC	STUARTNATAL PLUS 3 TABS <sup>1</sup>		
			MC	TRI-VI-SOL SOLN		
		ı	MC	TRI-VI-SOL/IRON SOLN		
			MC/DEL	ULTRA NATALCARE TABS		
	1 1		MC	ULTRA-NATAL TABS <sup>1</sup>		

	1 1	Ī	MC	VICON FORTE CAPS	Ī	1
				VINATAL FORTE TABS <sup>1</sup>		
			MC			
			MC MC/DEL	VINATE 1		
		MISCELLANEOUS MINERAL		VINATE ADVANCED TABS <sup>1</sup>		
MINERALS		CALCARB		ANEMAGEN	I	
MINERALS	MC		MC	CALCET TABS	Use PA Form# 20420 Please refer to 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
	MC	CALCI-MIX CAPSULE CAPS	MC		Please refer to OTC list.	preferred drug(s) exists. Certain drugs require specific diagnoses for approval.
	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		produce ( ) .
	MC/DEL	CALCIUM GLUCONATE TABS	MC	CITRACAL PLUS TABS		n ( ) 070 ii
	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				1

	MC/DEL MC/DEL MC MC/DEL MC MC/DEL	OYSTER CALCIUM TABS OYSTER SHELL PHARMA FLUR PHOSPHA 250 NEUTRAL TABS POTASSIUM BICARBONATE TBEF POTASSIUM CHLORIDE 8MEQ				
	MC MC/DEL MC MC/DEL MC MC MC MC	POTASSIUM EFFERVESCENT SELENIUM TABS SLOW-MAG TBCR SODIUM FLUORIDE V-R CALCIUM V-R OYSTER SHELL CALCIUM ZINC SULFATE CAPS				
		PHENYLKETONURIA (PKU) TREATMENT AGENTS	5			
PHENYLKETONURIA (PKU) TREATMENT AGENTS- INJECTABLES	$\prod$		MC	PALYNZIQ <sup>1</sup>		Palynziq is not to be used in combination with Kuvan
<i>I</i> [	4				<u>Use PA Form# 20420</u>	<u> </u>
PHENYLKETONURIA (PKU) TREATMENT AGENTS- ORAL			MC	KUVAN		
/ <u> </u>		THE TATE OF THE CONTROL OF			<u>Use PA Form# 20420</u>	<u> </u>
ELECTROLYTES/ NUTRITIONALS	MC	MISC. ELECTROLYTES/NUTRITIONA	MC MC	To a part	This list of nutritionals is	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
ELECTROLITED/ NOTATIONALS	MC MC	INTRALIPID EMUL <sup>1</sup> P.T.E5 SOLN <sup>1</sup> SEA-OMEGA CAPS <sup>1</sup>	MC MC MC MC MC MC MC MC MC MC MC MC	BOOST <sup>1</sup> CASEC POWD <sup>1</sup> CHOICE DM LIQD <sup>1</sup> DELIVER 2.0 LIQD <sup>1</sup> DOJOLVI ENFAMIL <sup>1</sup> ENSURE <sup>1</sup> GLUCERNA <sup>1</sup> ISOCAL LIQD <sup>1</sup> KINDERCAL TF LIQD <sup>1</sup>	incomplete. All nutritionals to still require a PA except for the miscellaneous products listed as preferred. SGA form required for nutritionals unless member has a G/I tube.	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
			MC MC MC MC MC MC MC MC MC MC MC MC MC M	KINDERCAL TF/FIBER LIQD¹  L-CARNITINE CAPS¹  LIPISORB LIQD¹  LOVAZA¹²  MODULEN IBD POWD¹  NUTRAMIGEN POWD¹  NUTREN¹  NUTRITIONAL SUPPLEMENT LIQD¹  PEPTAMEN¹  PHENYLADE¹  PHENYLADE¹  PHENYL-FREE¹  PKU 3 POWD¹  PREGESTIMIL POWD¹  PROBALANCE LIQD¹	<u>Use PA Form# 20420</u> <u>&amp; SGA Form</u>	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

 		•					
	<b>l</b> [	· · · · · · · · · · · · · · · · · · ·	MC	1	SCANDISHAKE PACK <sup>1</sup>		
	l	'	1 1	1	1		
			MC		VASCEPA	<u> </u>	
ERYTHROPOEITINS	***	EPOGEN SOLN	- 110		Towns and	T. 54 5 # 40500	No. Dufaced design with the best and failed in step ander due to lock of effects are intelestable pide effects herein non-professed drugs will be entranged unless an accordable clinical
ERTIHKUPUEITINS	MC		MC MC	8	ARANESP SOLN <sup>1</sup>	Use PA Form# 10520  1. Clinical PA is required to	Non-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
	MC	MIRCERA SYRINGE	WIC	ı °	PROCRIT SOLN <sup>1</sup>	establish medical necessity	another drug and the preferred drug(s) exists. Please see the EPO PA form for other approval and renewal criteria.
	MC	RETACRIT	1 1	1	1	and that appropriate lab	
		GRANULOCYTE CSF				amonitoring is being done.	
GRANULOCYTE CSF	MC	FULPHILA	MC	8	FYLNETRA		See approval criteria detailed on Granulocyte Colony Stimulating Factor PA form.
	MC	NEUPOGEN SYRINGE	MC	8	GRANIX SYRINGE	step order.	<b>,</b>
<b>!</b>	MC	NEUPOGEN VIAL	MC	8	GRANIX VIAL		<b>!</b>
1	MC/DEL	NYVEPRIA SYRINGE	MC	8	LEUKINE		
1		,	MC/DEL	8	NIVESTYM		<b>!</b>
1		,	MC	8	ROLVEDON		
1		,	MC	8	STIMUFEND		<b>!</b>
1		' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' ' '	MC/DEL	8	ZARXIO		!
!		,	MC/DEL	1	ZIEXTENZO		!
. [		,	MC	9	NEULASTA <sup>1</sup>	Use PA Form# 20520	
		GAUCHER DISEASE			7 HE SECTION 11		
GAUCHER DISEASE			MC	$\overline{}$	CERDELGA <sup>1</sup>	Clinical PA for 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
. [		,	MC	1	YARGESA <sup>1</sup>	roquirod	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
.		,	1 1	1	THROES.		preferred drug(s) exists. Exceeding days supply limits for LMWH class requires PA.
		,	1 1	1	1		<b>!</b>
		,	1 1	1	1		Yargesa: As monotherapy for the treatment of adult patients with mild to moderate type 1 Gaucher disease for whom enzyme replacement therapy is not a therapeutic option (e.g., due to
		,	1 1	1	1		allergy, hypersensitivity, or poor venous access).
		,	1 1	1	1		
		,	1 1	1	1	Use PA Form# 20420	
		NIEMANN-PICK DISEASE AGENTS				USE FA FUITH 20420	
NIEMANN-PICK DISEASE AGENTS		NIEWANN-PICK DISEASE AGENTS	MC		ACMEUROA1	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
MEMANIFFIOR DIOLAGE AGENTO		,	MC	1	AQNEURSA <sup>1</sup> MIPLYFFA <sup>1</sup>	Clinical PA required for     appropriate diagnosis	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
		,	INIC	1	MIPLYFFA	appropriate diagnosis.	preferred drug(s) exists.
		,	1 1	1	1		
		,	1 1	1	1		
		,	1 1	1	1		
		,	1 1	1	1		
		,	1 1	1	1		
ı <u> </u>	<u> </u>		<u> </u>			Use PA Form# 20420	
		ANTICOAGULANTS / PLATELET AGENT					
ANTICOAGULANTS	MC	COUMADIN TABS	MC	1	ARIXTRA SOLN	Enoxaparin therapy	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
, <b> </b>	MC/DEL	ENOXAPARIN <sup>1</sup>	MC/DEL	1	FONDAPARINUX	durations greater than 7 days every 30 days require	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. Exceeding days supply limits for LMWH class requires PA.
. 1	MC	ELIQUIS	MC/DEL	1	FRAGMIN INJ	DΛ	preferred drugs) exists. Exceeding days supply littics for climant dass requires 1 7.
. 1	MC	ELIQUIS STARTER PACK	MC/DEL	1	FRAGMIN VIAL	2. Use other strengths	
. 1	MC	HEPARIN SODIUM/NACL 0.9% SOLN	MC/DEL	1	LOVENOX SOLN	available to obtain desired	
, <b> </b>	MC	HEP-LOCK SOLN	MC/DEL	1	LOVENOX 300 <sup>2</sup>	dose.	
, <b> </b>	MC	INNOHEP	MC/DEL	1	LOVENOX SUBQ SYRINGE	3. Diagnosis required	DDI: Warfarin will require prior authorization if being used in combination with fluconazole, miconazole, or voriconazole.
, <b> </b>	MC	HEPARIN LOCK SOLN	MC/DEL	1	PRADAXA ORAL PELLETS <sup>4</sup>		
, <b> </b>	MC/DEL	HEPARIN LOCK FLUSH SOLN	MC	1	IPRIVASK	4 F th- transment of	
, <b> </b>	MC/DEL	HEPARIN SODIUM SOLN	MC/DEL	1	SAVAYSAS <sup>3</sup>	For the treatment of patients aged 3 months to	DDI: Warfarin will require prior authorization if being used in conjunction with Gemfibrozil or Fenofibrate.
	MC/DEL	HEPARIN SODIUM LOCK FLUSH SOLN	1 1	1		less than 12 years of age.	
	MC/DEL	PRADAXA	1 1	1	1	1000 01011 12 ,00.0 0. 0.5.	
, <b> </b>			1 1	1	1		
, <b> </b>	MC/DEL	JANTOVEN	1 1	1	1		DBL Diference will remain a subhasination if hairs used in combination with Courses
, <b> </b>	MC/DEL	WARFARIN SODIUM TABS	1 1	1	1		DDI: Rifampin will require prior authorization if being used in combination with Savaysa
, 1	MC/DEL	XARELTO	1 1	1	1		1

ı	MC/DEL	XARELTO STARTER PACK	1 1	1	1	Ī	1
	MC/DEL	AARELTO STARTER FACK		1		· ·	
				1		Use PA form# 20420	
	l		\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	1			
ANTIHEMOPHILIC AGENTS	MC	ALPHANATE	MC/DEL	$\overline{}$	ADYNOVATE VIAL	Only if other products	Non-preferred will only be approved if other preferred products are unavailable.
	MC	ALPHANINE SD	MC	1	ADVATE <sup>1,2,5</sup>	unavailable.	
	MC/DEL	ALPROLIX VIAL	MC/DEL	1	ALHEMO		Begvez:FDA Approved Indication: An adeno-associated virus vector-based gene therapy indicated for the treatment of adults with moderate to severe hemophilia B (congenital factor IX
	MC/DEL	BEBULIN VIAL	MC	1	ALTUVIIIO⁴	2. Advate may be available	deficiency) who:
	MC/DEL	BENEFIX SOLR	MC/DEL	1	AFSTYLA	with PA in cases of large	· Currently use factor IX prophylaxis therapy, or · Have current or historical life-threatening hemorrhage, or
	MC/DEL	HELIXATE FS KIT	MC/DEL	1	BEQVEZ	volume dosing in patients with poor venous access.	Have repeated, serious spontaneous bleeding episodes, and,
	MC	HEMLIBRA	MC/DEL	1	ESPEROCT	with poor verious access.	Do not have neutralizing antibodies to adeno-associated virus serotype Rh74var (AAVRh74var) capsid as detected by an FDA- approved test.
<b> </b>	MC	HEMOFIL - M	MC/DEL	1	ELOCTATE	<b> </b>	
	MC	HUMATE-P SOLR	MC/DEL	1	HEMGENIX	<b> </b>	Hemgenix® is an adeno-associated viral vector-based gene therapy for IV infusion after dilution. For treatment of adults with Hemophilia B (congenital Factor IX deficiency) who: Currently
	MC/DEL	IXINITY VIAL	MC/DEL	1	HYMPAVZI	3. Not indicated for use in	use Factor IX prophylaxis therapy, or have current or historical life-threatening hemorrhage, or Have repeated, serious spontaneous bleeding episodes.
	MC/DEL	JIVI <sup>3</sup>	MC/DEL	1	IDELVION	children <12 years of age	
	MC	KOATE-DVI	MC/DEL	1	KOGENATE FS <sup>5</sup>	due to greater risk for	Altuviiio is a von Willebrand Factor (VWF) independent recombinant DNA-derived, Factor VIII concentrate indicated for use in adults and children with hemophilia A (congenital factor VIII
<b> </b>	MC	KONYNE - 80	MC		RECOMBINATE VIAL <sup>5</sup>	hypersensitivity reactions and is not indicated for use	deficiency) for: Routine prophylaxis to reduce the frequency of bleeding episodes, On-demand treatment and control of bleeding episodes, Perioperative management of bleeding.
<b> </b>	MC/DEL	KOVALTRY	MC	1	ROCTAVIAN⁴	in previously untreated	
<b> </b>	MC/DEL	REBINYN	MC	1	SEVENFACT	patients.	
	MC	MONARC - M	n.o	1	GEVENI AGT	<b> </b>	Roctavian: For the treatment of adults with severe hemophilia A (congenital factor VIII deficiency with factor VIII activity <1 IU/dL) without antibodies to adeno-associated virus serotype 5
		MOTALICO - IM		1			(AAV5) detected by an FDA-approved test.
	MC	MONOCLATE - P	1 1	1		<b> </b>	Inclusion:
<b> </b>	MC	MONONINE	<b>l</b> <i>l</i>	1		<b> </b>	Severe factor VIII deficiency (less than 1% native factor VIII).
	MC/DEL	NOVOEIGHT		1			Exclusion Criteria:
	MC	NOVOSEVEN SOLR		1			Antibodies to the virus AAV5
	MC	NUWIQ		1			
	MC/DEL	PROFILNINE	<b>l</b> <i>l</i>	1		16.11	Known significant fibrosis of cirrhosis of the liver, or unexplained elevated LFTs
	MC	RECOMBINATE SOLR		1		Ť	History of inadequate compliance with prophylaxis, or regular bleeds despite adequate prophylaxis
	MC	REFACTO		1			Conditions in which high-dose steroids are contraindicated.
	MC/DEL	RIXUBIS VIAL		1		<b> </b>	-Inability to abstain from alcohol for one year
	MC	WILATE INJ	<b>l</b> <i>l</i>	1		<b> </b>	Plan to impregnate a partner within 6 months of infusion
	MC/DEL	WILATE INJ XYNTHA	1 1	1		<b> </b>	-Hypersensitivity to mannitol
	WIC/DEL	XYNIDA		1		<b> </b>	-Active infections, either acute or uncontrolled chronic
	l		<b>l</b> <i>l</i>	1		Use PA Form# 20420	-HIV infection (limited information on use in this population)
	l			1		USE PA FUIII# 20420	-FITV fillection (illinited information on use in this population)
	l			1		<b> </b>	
	l			1		<b> </b>	
PLATELET AGGREGATION INHIBITORS	MC/DEI	ASPIRIN	MC/DEI	<del>-</del>	TICLOPIDINE HCL TABS		Destayed drives must be tried and failed due to look of officeau or intelerable side offices before non professed drives will be approved, unless an acceptable clinical expension is officed on
PLATELET AGGREGATION INFIDITIONS	MC/DEL		MC/DEL MC/DEL	8	BRILINTA 60mg	Use PA Form# 20715 for	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
	MC MC/DEL	ASPIRIN-DIPYRIDAMOLE ER CPMP 12HR			=	Flavix, Ellient & Brillinta	preferred drug(s) exists.
	MC/DEL	BRILINTA 90mg	MC	8	DURLAZA	Use PA form# 20420 for	
	MC/DEL	DIPYRIDAMOLE TABS	MC	8	EFFIENT DEDEANTINE TARE	other requests	A special PA may be obtained at the pharmacy for members scheduled for "stent" placement or have had placement if in the last 12months. Please indicate on prescription date of stent placement.
	MC/DEL	CLOPIDOGREL 75MG	MC/DEL	8	PERSANTINE TABS		placement.
H	MC/DEL	PRASUGREL HCL TAB	MC/DEL	8	PLAVIX TABS	Dosing limits apply,	
	l		MC/DEL	8	ZONTIVITY		DDI: Plavix will require prior authorization if being used in combination with omeprazole, esomeprazole, cimetidine, fluconazole, ketoconazole, intelence, fluoxetine, ticlopidine, and
	l		<b>l</b> <i>l</i>	1			fluvoxamine.
11	l			1			DDI: exists for using maintenance ASA dose >100mg, as it reduces the effectiveness of Brilinta
11	l			1		<b> </b>	Brilinta- Concomitant use with strong CYP3A4 inhibitors should be avoided (including ketoconazole, itraconazole, atazanavir, and telithromycin). Doses of simvastatin and lovastatin
	l			1		!	>40mg should be avoided.
				<b>—</b>			
PLATELET AGGR. INHIBITORS / COMBO'S - MISC.	MC/DEL	CILOSTAZOL	MC/DEL	1	AGRYLIN 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
COMBO 3 - MISC.	MC/DEL	PENTOXIFYLLINE ER TBCR	MC/DEL	1	ANAGRELIDE CAPS		preferred drug(s) exists.
			MC/DEL	1	PLETAL TABS		and a single) contact
			MC	1	TRENTAL TBCR		
			MC	1	YOSPRALA		
				4			
		HEMATOLOGICALS					
						*	

MONOCLONAL ANTIBODY			MC	EMPAVELI	Use PA Form# 20420	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
1	1	1	MC/DEL	ENSPRYNG	,	vaccine at least 2 weeks prior to the start of therapy.
1	1	1	MC	FABHALTA	1	Gamifant is recommended for the treatment of adult and pediatric (newborn and older) patients with primary hemophagocytic lymphohisticocytosis (HLH) with refractory, recurrent, or
1	1	1	MC/DEL	GAMIFANT	1	progressive disease or intolerance with conventional HLH therapy.
<b> </b>	1	1	MC/DEL	GAMIFAN I PIASKY	1	progressive diseases of interioring man controllation. The controllation is a state of the controllation of the co
<b> </b>	1	1			1	· 1
<b> </b>	1	1		SOLIRIS	1	
<b> </b>	1	1	MC/DEL	ULTOMIRIS	1	Fabhalta and Ultomiris are recommended for the treatment of adults with paroxysmal nocturnal hemoglobinuria (PNH).
/ [	1	1	MC	UPLIZNA	1	· 1
/ L	4	1	MC	VOYDEYA		
IMMUNE GLOBULIN	MC	BIVIGAM¹	MC	ALYGLO		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
<b> </b>	MC/DEL	CUTAQUIG <sup>1</sup>	MC	ASCENIV <sup>2</sup>		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
l I ,	MC	GAMUNEX-C	MC/DEL	CUVITRU	2. For the treatment of	preferred drug(s) exists.
	MC	GAMMAGARD S-D <sup>1</sup>	MC	GAMMAPLEX INJ	patients between 12 to 17	· I
<b> </b>	MC/DEL	HIZENTRA <sup>1</sup>		HYQVIA	' -	Alyglo is indicated for treatment of primary humoral immunodeficiency in adults ages 17 or older.
l I	MC/DEL	PANZYGA <sup>1</sup>	MC MC	OCTAGAM INJ <sup>1</sup>	ſ	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
l I	MC/DEL	PRIVIGEN <sup>1</sup>		XEMBIFY	1	· I
<b> </b>	MC	PRIVIGEN	MC/DEL	XEMBIFY	1	O to 1 1 1 1-district and a second the second for primary hymeral immunately in a polyton
	1	1	1 1	•	1	Cutaquig is indicated as replacement therapy for primary humoral immunodeficiency (PI) in adults.
<b> </b>	<i>i</i>	1	1 1	•	1	
	1	1	1 1	•	1	Xembify is indicated for treatment of primary humoral immunodeficiency (PI) in patients 2 years of age and older.
/1	1	1	1 1	•	1	· 1
/ 1	<i>i</i>	1	1 1	•	1	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
/1 /	<i>i</i>	1	1 1	•		Asceniv indicated for the treatment of primary numbral immunodeficiency (PI) in adults and adolescents (12 to 17 years of age). PI includes but is not limited to the numbral immune defect in congenital agammaglobulinemia, common variable immunodeficiency (CVID), X-linked agammaglobulinemia, Wiskott-Aldrich syndrome, and severe combined
/1 /	1	1	1 1	•	1	immunodeficiencies (SCID).
HEREDITARY ANGIOEDEMA		PROPHYLAXIS		PROPHYHLAXIS	1. Clinical PA is required to	
/I ,	мс	CINRYZE <sup>1</sup>	<del></del>	+	establish diagnosis and	Haegarda is indicated for routine prophylaxis to prevent Hereditary Angioedema (HAE) attacks in adolescent and adult patients
/I ,	MC	HAEGARDA <sup>1</sup>	1 1	•	medical necessity.	Thought to managed for routing property
/I ,	MC	ORLADEYO <sup>1,2</sup>	1 1	•	2. For the treatment of	· 1
/I ,		TAKHZYRO <sup>1</sup>	1 1	•	patients ≥ 12 years of age.	· I
/I ,	MC/DEL	TAKHZYKU	1 1	•	pationic = 1= ,=	
/I ,	<i>i</i>	1	1 1	•	1	· I
/I ,	1		<u> </u>			
/I ,	1	TREATMENT		TREATMENT	1	
/I ,	MC/DEL	BERINERT KIT <sup>1</sup>	MC/DEL	KALBITOR VIAL	1	· I
/I ,	MC	FIRAZYR <sup>1</sup>	1 1	•	1	· 1
/I ,	MC/DEL	RUCONEST VIAL <sup>1</sup>	1 1	•	1	· [
/I ,	<i>i</i>	1	1 1	•	1	· [
/I ,	<i>i</i>	1	1 1	•	Use PA Form# 20420	· [
HEMATOLOGICAL AGENTS-	<del></del>	+	1	<b>+</b>	USE FA I OIIIIII 20-120	+
THROMBOPOIETIN RECEPTOR	·	PROMACTA <sup>1</sup>	1	117417	U DA Farm# 20420	· [
AGONISTS	MC MC		MC	ALVAIZ	Use PA Form# 20420	·]
/	MC	NPLATE <sup>1</sup>		DOPTELET	<ol> <li>Clinical PA required. Must see prior trial with insufficient</li> </ol>	ont in the second secon
/I ,	1	1	MC/DEL	MULPLETA	response to corticosteroids	Dobleiet and Mulbella. For the treatment of thrombocytopenia in adults with chronic liver disease who are scheduled to undergo a procedure.
/I ,	1	1	1 1	•	and immunoglobulins.	
/I ,	1	1	1 1	•	,	
/I ,	1	1	1 1	•	1	
/I ,	1	1	1 1	•	1	
HEMATOLOGICAL AGENTS-IgAN	$\overline{}$	<del></del>	MC/DEL	FILSPARI <sup>1</sup>	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 (in step-order) will be approved, unless an acceptable clinical
1	1	1	MC	TARPEYO	1. PA required to confirm	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
/I ,	1	1	imo	TARLETO	FDA approved indication.	another drug and the preferred drug(s) exists
/I ,	1	1	1 1	•	, sept	
<u>                                     </u>		<u> </u>	<del></del>	·		
ANEMIA- BETA THALASSEMIA	<i>ī</i>	· [	MC	REBLOZYL		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
/I ,	1	1	MC	ZYNTEGLO	1	RBC transfusions in patients who require immediate correction of anemia.
/I ,	<i>i</i>	1	1 1	•	1	
/	1	1	1 1	•	1	Zynteglo is indicated for the treatment of adult and pediatric patients with β-thalassemia who require regular red blood cell (RBC) transfusions.
·		_	_		- 1	
11	'	· I	<b>.</b>	. <b>I</b>		
HEMATOLOGIC DISORDER TREATMENT	<u> </u>	<u> </u>	MC/DEL	CABLIVI	Use PA Form# 20420	Tavalisse is recommended for patients at risk of bleeding when one line of therapy (steroids, IVIG, splenectomy) has failed.

AGENIS			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	
					OSE FA FOITH# 20420	
WHIM SYNDROME AGENTS			MC	XOLREMDI		
WHIM STNDROME AGENTS			IVIC	XOLKEWIDI		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.
					<u>Use PA Form#20420</u>	
	•	HEMOSTATIC				
HEMOSTATIC	MC/DEL	AMICAR	MC	FIBRYGA	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
	MC	AMINOCAPROIC ACID	MC	RIASTAP		hypofibrinogenemia. Fibryga® is not indicated for dysfibrinogenemia.
		ACUTE HEPATIC PORPHYRIA (A	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 A	GENTS			
PYRUVATE KINASE DEFICIENCY		1	MC	PYRUKYND¹	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
AGENTS						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
					<ol> <li>PA required to confirm FDA approved indication.</li> </ol>	mintament day afal
					FDA approved indication.	
OP ANTIBIOTICS	MC	AK-SPORE OINT	NC I	AK-POLY-BAC OINT	L. D. E. W. 00400	Designed drawn with bright and failed due to look of officers with a first before any professed drawn will be approved unless an apparatulation in affected any
OP ANTIBIOTICS	IVIC		MC		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
	MC	BACITRACIN/NEOMYCIN/POLYM	MC	AK-SULF OINT		preferred drug(s) exists.
	MC/DEL	BACITRACIN/POLYMYXIN B OINT	MC	AK-TOB SOLN		product diagraphical
	MC	CHLOROPTIC SOLN	MC	AZASITE		
	MC/DEL	ERYTHROMYCIN OINT	MC	BACITRACIN OINT		
	MC	NEOSPORIN SOLN	MC	BLEPH-10 SOLN		
	MC	POLYSPORIN	MC/DEL	GATIFLOXACIN DROPS		
	MC/DEL	TRIMETHOPRIM SULFATE/POLY	MC/DEL	GENTAMICIN SULFATE		
	MC/DEL	TOBRAMYCIN SULFATE SOLN	MC	GENTAK		
			MC	ILOTYCIN OINT		
			MC/DEL	LEVOFLOXACIN DROPS		
			MC/DEL	NEOMYCIN/BACI/POLYM OINT		
			MC/DEL	NEOMYCIN/POLYMYXIN/GRAMIC		
			мс	NEOSPORIN OINT		
			MC	OCUSULF-10 SOLN		
			MC	OCUTRICIN SOLN		
			MC/DEL	POLYTRIM DROPS		
			MC/DEL	SULFACETAMIDE SODIUM DROPS		
			MC/DEL	SULFACETAMIDE SODIUM OINT		
			MC	TERAK OINT		<u> </u>
OPANTI-PARASITIC			MC	XDEMVY <sup>1</sup>	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
					1. For the treatment of	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
					Demodex blepharitis.	aratarrad drunie) aviete
						<u> </u>
OP RHO KINASE INHIBITORS		DUODDESS	+			
UP KHU KINASE INHIBITUKS	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)
					<u>Use PA Form# 20420</u>	
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

<u> </u>	MC/DEL	CIPROFLOXACIN SOL 0.3%	MC/DEL	CILOXAN SOLN	Ī	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
	MC/DEL	OFLOXACIN	MC	OCUFLOX SOLN		preferred drug(s) exists.
	MC/DEL	QUIXIN SOLN	1 1	000. 201. 002		
	1	,	1			
OPQUINOLONES-4TH GENERATION	MC/DEL	MOXIFLOXACIN 0.5% SOLN (Generic Vigamox)	MC	ZYMAXID	Use PA Form# 20420	
OP ARTIFICIAL TEARS AND	MC/DEL	ARTIFICIAL TEARS OINT	MC/DEL	ARTIFICIAL TEARS SOLN OP	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
LUBRICANTS	MC/DEL	ARTIFICIAL TEARS SOLN	MC	BION TEARS SOLN	1. Dosing limits apply,	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
	MC	CELLUVISC SOLN	MC	DRY EYES OINT	please see dose	preferred drug(s) exists.
	MC	EYE LUBRICANT OINT	MC	DURATEARS OINT	consolidation list.	
!	MC/DEL	GENTEAL	MC/DEL	HYPO TEARS		
	MC	LIQUITEARS SOLN	MC/DEL	ISOPTO TEARS SOLN		
	MC	MAJOR TEARS SOLN	MC	LACRI-LUBE		
!	MC	PURALUBE OINT	MC	LUBRIFRESH P.M. OINT		
	MC	PURALUBE TEARS SOLN	MC	MURINE SOLN		
<u> </u>	MC	REFRESH SOLN OP	MC/DEL	MUROCEL SOLN		
!	MC	REFRESH PLUS SOLN <sup>1</sup>	MC/DEL	NATURE'S TEARS SOLN		
<b>!</b>	MC	REFRESH PM OINT	MC	REFRESH SOLN		
	<b>1</b>	'	MC	REFRESH TEARS SOLN <sup>1</sup>		
	<b>1</b>	'	MC	TEARGEN SOLN		
<b>!</b>	<b>1</b>	'	MC	TEARISOL SOLN		
<b>!</b>	<b>1</b>	'	MC/DEL	TEARS NATURALE		
1	<b>1</b>	·	MC/DEL	TEARS PURE SOLN		
!	<b>1</b>	'	MC	TEARS RENEWED OINT		
	<b>1</b>	'	MC/DEL	THERATEARS SOLN		
	<b>1</b>	'	MC	V-R ARTIFICIAL TEARS SOLN		
	1	'	1 -			
	l <u>L</u>		1		<u> </u>	
OP BETA - BLOCKERS	MC/DEL	BETOPTIC-S SUSP	MC	BETAGAN 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
	MC/DEL	CARTEOLOL HCL SOLN	MC/DEL	BETAXOLOL HCL SOLN		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
1	MC/DEL	LEVOBUNOLOL HCL SOLN	MC	ISTALOL		preferred drug(s) exists.
	MC/DEL	METIPRANOLOL SOLN	MC/DEL	OCUPRESS SOLN		
	<b>1</b>	· • • • • • • • • • • • • • • • • • • •	MC		_	
	1		IVIC	OPTIPRANOLOL SOLN	l	
		<b>.</b> [	MC/DEL	OPTIPRANOLOL SOLN TIMOPTIC SOLN		
			MC/DEL	TIMOPTIC SOLN		
			MC/DEL MC	TIMOPTIC SOLN TIMOLOL DROP		
			MC/DEL MC MC/DEL	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL		
OP ANTI-INFLAMMATORY / STEROIDS	MC	AK-SPORE HC OINT	MC/DEL MC MC/DEL	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-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
OP ANTI-INFLAMMATORY / STEROIDS OPHTH.	MC MC/DEL	AK-SPORE HC OINT ALREX SUSP	MC/DEL MC MC/DEL MC/DEL	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL TIMOPTIC-XE SOLG	Use PA Form# 20420	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
			MC/DEL MC MC/DEL MC/DEL	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL TIMOPTIC-XE SOLG  AK-TROL SUSP	<u>Use PA Form# 20420</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.
	MC/DEL	ALREX SUSP	MC/DEL MC MC/DEL MC/DEL MC/DEL MC MC	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL TIMOPTIC-XE SOLG  AK-TROL SUSP BAC/POLY/NEOMY/HC OINT	<u>Use PA Form# 20420</u>	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
	MC/DEL MC/DEL	ALREX SUSP DEXAMETH SOD PHOS SOLN	MC/DEL MC MC/DEL MC/DEL MC/DEL MC MC MC	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL TIMOPTIC-XE SOLG  AK-TROL SUSP BAC/POLY/NEOMY/HC OINT BLEPHAMIDE S.O.P. OINT	Use PA Form# 20420	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
	MC/DEL MC/DEL MC/DEL	ALREX SUSP DEXAMETH SOD PHOS SOLN FLUOROMETHOLONE SUSP	MC/DEL MC MC/DEL MC/DEL MC/DEL MC MC MC MC MC	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL TIMOPTIC-XE SOLG  AK-TROL SUSP BAC/POLY/NEOMY/HC OINT BLEPHAMIDE S.O.P. OINT BLEPHAMIDE SUSP	Use PA Form# 20420	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
	MC/DEL MC/DEL MC/DEL MC	ALREX SUSP DEXAMETH SOD PHOS SOLN FLUOROMETHOLONE SUSP FML DROPS SUSP 1%	MC/DEL MC MC/DEL MC/DEL MC/DEL MC MC MC MC MC MC MC	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL TIMOPTIC-XE SOLG  AK-TROL SUSP BAC/POLY/NEOMY/HC OINT BLEPHAMIDE S.O.P. OINT BLEPHAMIDE SUSP BROMDAY	Use PA Form# 20420	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
	MC/DEL MC/DEL MC/DEL MC MC	ALREX SUSP DEXAMETH SOD PHOS SOLN FLUOROMETHOLONE SUSP FML DROPS SUSP 1% FML FORTE SUSP	MC/DEL MC MC/DEL MC/DEL MC MC MC MC MC MC MC MC MC MC	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL TIMOPTIC-XE SOLG  AK-TROL SUSP BAC/POLY/NEOMY/HC OINT BLEPHAMIDE S.O.P. OINT BLEPHAMIDE SUSP BROMDAY EFLONE SUSP	Use PA Form# 20420	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
	MC/DEL MC/DEL MC MC MC MC	ALREX SUSP DEXAMETH SOD PHOS SOLN FLUOROMETHOLONE SUSP FML DROPS SUSP 1% FML FORTE SUSP FML S.O.P. OINT	MC/DEL MC MC/DEL MC/DEL MC MC MC MC MC MC MC MC MC MC MC MC MC	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL TIMOPTIC-XE SOLG  AK-TROL SUSP BAC/POLY/NEOMY/HC OINT BLEPHAMIDE S.O.P. OINT BLEPHAMIDE SUSP BROMDAY EFLONE SUSP FLAREX SUSP	Use PA Form# 20420	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
	MC/DEL MC/DEL MC MC MC MC MC MC MC	ALREX SUSP DEXAMETH SOD PHOS SOLN FLUOROMETHOLONE SUSP FML DROPS SUSP 1% FML FORTE SUSP FML S.O.P. OINT LOTEMAX OINT	MC/DEL MC MC/DEL MC/DEL MC MC MC MC MC MC MC MC MC MC MC MC MC	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL TIMOPTIC-XE SOLG  AK-TROL SUSP BAC/POLY/NEOMY/HC OINT BLEPHAMIDE S.O.P. OINT BLEPHAMIDE SUSP BROMDAY EFLONE SUSP FLAREX SUSP FLUOR-OP SUSP	Use PA Form# 20420	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
	MC/DEL MC/DEL MC MC MC MC MC MC MC MC/DEL	ALREX SUSP DEXAMETH SOD PHOS SOLN FLUOROMETHOLONE SUSP FML DROPS SUSP 1% FML FORTE SUSP FML S.O.P. OINT LOTEMAX OINT LOTEMAX GEL	MC/DEL MC MC/DEL MC/DEL MC MC MC MC MC MC MC MC MC MC MC MC MC	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL TIMOPTIC-XE SOLG  AK-TROL SUSP BAC/POLY/NEOMY/HC OINT BLEPHAMIDE S.O.P. OINT BLEPHAMIDE SUSP BROMDAY EFLONE SUSP FLAREX SUSP FLUOR-OP SUSP ILUVIEN IMPLANT	Use PA Form# 20420	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
	MC/DEL MC/DEL MC MC MC MC MC MC MC/DEL MC/DEL MC/DEL MC/DEL	ALREX SUSP DEXAMETH SOD PHOS SOLN FLUOROMETHOLONE SUSP FML DROPS SUSP 1% FML FORTE SUSP FML S.O.P. OINT LOTEMAX OINT LOTEMAX GEL LOTEMAX SUSP	MC/DEL MC MC/DEL MC/DEL MC MC MC MC MC MC MC MC MC MC MC MC MC	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL TIMOPTIC-XE SOLG  AK-TROL SUSP BAC/POLY/NEOMY/HC OINT BLEPHAMIDE S.O.P. OINT BLEPHAMIDE SUSP BROMDAY EFLONE SUSP FLAREX SUSP FLUOR-OP SUSP ILUVIEN IMPLANT INVELTYS	Use PA Form# 20420	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
	MC/DEL MC/DEL MC MC MC MC MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	ALREX SUSP DEXAMETH SOD PHOS SOLN FLUOROMETHOLONE SUSP FML DROPS SUSP 1% FML FORTE SUSP FML S.O.P. OINT LOTEMAX OINT LOTEMAX GEL LOTEMAX SUSP NEO/POLY/DEXAMETH OINT	MC/DEL MC MC/DEL MC/DEL MC MC MC MC MC MC MC MC MC MC MC MC MC	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL TIMOPTIC-XE SOLG  AK-TROL SUSP BAC/POLY/NEOMY/HC OINT BLEPHAMIDE S.O.P. OINT BLEPHAMIDE SUSP BROMDAY EFLONE SUSP FLAREX SUSP FLUOR-OP SUSP ILUVIEN IMPLANT INVELTYS LOTEMAX SM DROPS GEL 0.38%	Use PA Form# 20420	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
	MC/DEL MC/DEL MC MC MC MC MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	ALREX SUSP DEXAMETH SOD PHOS SOLN FLUOROMETHOLONE SUSP FML DROPS SUSP 1% FML FORTE SUSP FML S.O.P. OINT LOTEMAX OINT LOTEMAX GEL LOTEMAX SUSP NEO/POLY/DEXAMETH OINT NEO/POLY/DEXAMETH SUSP	MC/DEL MC MC/DEL MC/DEL MC MC MC MC MC MC MC MC MC MC MC MC MC	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL TIMOPTIC-XE SOLG  AK-TROL SUSP BAC/POLY/NEOMY/HC OINT BLEPHAMIDE S.O.P. OINT BLEPHAMIDE SUSP BROMDAY EFLONE SUSP FLAREX SUSP FLUOR-OP SUSP ILUVIEN IMPLANT INVELTYS LOTEMAX SM DROPS GEL 0.38% MAXITROL OPTH OINT 0.1%	Use PA Form# 20420	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
	MC/DEL MC/DEL MC MC MC MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC	ALREX SUSP DEXAMETH SOD PHOS SOLN FLUOROMETHOLONE SUSP FML DROPS SUSP 1% FML FORTE SUSP FML S.O.P. OINT LOTEMAX OINT LOTEMAX GEL LOTEMAX SUSP NEO/POLY/DEXAMETH OINT NEO/POLY/DEXAMETH SUSP PRED-G SUSP	MC/DEL MC MC/DEL MC MC MC MC MC MC MC MC MC MC MC MC MC	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL TIMOPTIC-XE SOLG  AK-TROL SUSP BAC/POLY/NEOMY/HC OINT BLEPHAMIDE S.O.P. OINT BLEPHAMIDE SUSP BROMDAY EFLONE SUSP FLAREX SUSP FLUOR-OP SUSP ILUVIEN IMPLANT INVELTYS LOTEMAX SM DROPS GEL 0.38% MAXITROL OPTH OINT 0.1% NEO/POLY/BAC/HC OINT NEOM/POLY/DEX OPTH OINT 0.1%	Use PA Form# 20420	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
	MC/DEL MC/DEL MC MC MC MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC MC	ALREX SUSP DEXAMETH SOD PHOS SOLN FLUOROMETHOLONE SUSP FML DROPS SUSP 1% FML FORTE SUSP FML S.O.P. OINT LOTEMAX OINT LOTEMAX GEL LOTEMAX SUSP NEO/POLY/DEXAMETH OINT NEO/POLY/DEXAMETH SUSP PRED FORTE SUSP 1%	MC/DEL MC MC/DEL MC MC MC MC MC MC MC MC MC MC MC MC MC	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL TIMOPTIC-XE SOLG  AK-TROL SUSP BAC/POLY/NEOMY/HC OINT BLEPHAMIDE S.O.P. OINT BLEPHAMIDE SUSP BROMDAY EFLONE SUSP FLAREX SUSP FLUOR-OP SUSP ILUVIEN IMPLANT INVELTYS LOTEMAX SM DROPS GEL 0.38% MAXITROL OPTH OINT 0.1% NEO/POLY/BAC/HC OINT	Use PA Form# 20420	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
	MC/DEL MC/DEL MC MC MC MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC MC MC MC	ALREX SUSP DEXAMETH SOD PHOS SOLN FLUOROMETHOLONE SUSP FML DROPS SUSP 1% FML FORTE SUSP FML S.O.P. OINT LOTEMAX OINT LOTEMAX GEL LOTEMAX SUSP NEO/POLY/DEXAMETH OINT NEO/POLY/DEXAMETH SUSP PRED FORTE SUSP 1% PRED MILD SUSP	MC/DEL MC MC/DEL MC MC MC MC MC MC MC MC MC MC MC MC MC	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL TIMOPTIC-XE SOLG  AK-TROL SUSP BAC/POLY/NEOMY/HC OINT BLEPHAMIDE S.O.P. OINT BLEPHAMIDE SUSP BROMDAY EFLONE SUSP FLAREX SUSP FLUOR-OP SUSP ILUVIEN IMPLANT INVELTYS LOTEMAX SM DROPS GEL 0.38% MAXITROL OPTH OINT 0.1% NEO/POLY/BAC/HC OINT NEOM/POLY/BAC/HC OINT NEOM/POLY/DEX OPTH OINT 0.1% OMNIPRED DROPS SUSP	Use PA Form# 20420	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
	MC/DEL MC/DEL MC MC MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC MC MC MC MC MC MC MC MC MC	ALREX SUSP DEXAMETH SOD PHOS SOLN FLUOROMETHOLONE SUSP FML DROPS SUSP 1% FML FORTE SUSP FML S.O.P. OINT LOTEMAX OINT LOTEMAX GEL LOTEMAX SUSP NEO/POLY/DEXAMETH OINT NEO/POLY/DEXAMETH SUSP PRED FORTE SUSP 1% PRED MILD SUSP PREDNISOLONE	MC/DEL MC MC/DEL MC MC MC MC MC MC MC MC MC MC MC MC MC	TIMOPTIC SOLN TIMOLOL DROP TIMOLOL SOL-GEL TIMOPTIC-XE SOLG  AK-TROL SUSP BAC/POLY/NEOMY/HC OINT BLEPHAMIDE S.O.P. OINT BLEPHAMIDE SUSP BROMDAY EFLONE SUSP FLAREX SUSP FLUOR-OP SUSP ILUVIEN IMPLANT INVELTYS LOTEMAX SM DROPS GEL 0.38% MAXITROL OPTH OINT 0.1% NEOM/POLY/BAC/HC OINT NEOM/POLY/BAC/HC OINT NEOM/POLY/DEX OPTH OINT 0.1% OMNIPRED DROPS SUSP OZURDEX	Use PA Form# 20420	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

1							,
1	MC	SULFACETAMIDE/PREDNISOLONE	MC/DEL	,	RETISERT IMPLANT	•	
<u>'</u>	MC/DEL	ZYLET SUSP	MC/DEL	,	SULFACET SOD/PRED SOLN	'	
!	1 .	•	MC/DEL	,	TRIESENCE VIAL	'	·
<u>'</u>	1		MC/DEL	,	TOBRADEX ST	'	
	1 1	1	MC/DEL	,	TOBRAMYCIN SUSP DEXAMETHASONE	'	
	1	1	MC	,	VASOCIDIN SOLN	'	·
ı <b>l</b>	1	1	MC/DEL	,	VEXOL SUSP	'	
/ <b> </b>	1	1	MC	,	XIPERE	'	
<b>/                                    </b>	1	1	l live	,	AIFERE	'	·   •
<b> </b>	1	1		,	1	'	
/ <b> </b>	1	1		,	1	'	
OP PROSTAGLANDINS	MC/DEL	LATANOPROST SOL 0.005%	MC/DEL	7	ZIOPTAN	1 All preferred must be trie	d. 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,
OF FROSTAGEARDING	MC/BEL	LUMIGAN SOLN					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
/ <b> </b>			MC/DEL	8	BIMATOPROST 0.03% DROPS		drug interaction between another drug and the preferred drug(s) exists.
<b>/                                    </b>	MC/DEL	ROCKLATAN TRAVATAN 7	MC	8	DURYSTA	'	
<b> </b>	MC/DEL	TRAVATAN-Z	MC	8	IYUZEH		
/ <b> </b>	1	1	MC	8	RESCULA <sup>1,2,3</sup>	2. Dosing limits apply,	
/ <b> </b>	1	1	- 1 - 1	,	1	please see dosing consolidation list.	
/ <b> </b>	1	<b> </b>		4 .	1	COllsonidation hat.	
<b>/                                    </b>	1	<b> </b>	MC/DEL		TRAVATAN SOLN	3. Clinical PA is required to	,
/ <b> </b>	1	<b> </b>	MC/DEL	8	TRAVOPROST	establish diagnosis and	
<b>/                                    </b>	1	1	MC/DEL	8	VYZULTA	medical necessity.	
<b>/                                    </b>	1	1	MC/DEL	8	XALATAN SOLN <sup>1</sup>	Use PA Form# 20420	
<b>/                                    </b>	1	1	MC/DEL	8	XELPROS	'	
OP CYCLOPLEGICS	MC	AK-PENTOLATE SOLN	MC/DEL	. —	CYCLOGYL 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
/ <b> </b>	MC/DEL	ATROPINE SULFATE	MC	,	ISOPTO ATROPINE SOLN		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
/ <b> </b>	MC/DEL	CYCLOPENTOLATE HCL SOLN	MC/DEL	,	ISOPTO HOMATROPINE SOLN	,	preferred drug(s) exists.
/	MC/DEL	ISOPTO HYOSCINE SOLN	MC	,	MUROCOLL-2 SOLN	· [	· I
OP MIOTICS - DIRECT ACTING	MC/DEL	ISOPTO CARBACHOL SOLN	+	,	<del></del>	Use PA Form# 20420	<del>                                     </del>
1	MC	ISOPTO CARPINE SOLN	l j	,	1	030 17(1 01.1., 25 .25	· I
/ <b> </b>	MC	PILOCAR SOLN	l j	,	1	,	· I
/ <b> </b>	MC/DEL	PILOCAR SOLIN PILOCARPINE HCL SOLIN	l j	,	1	,	·
/	MC/DEL	PILOPINE HS GEL	l j	,	1	,	1
OP SELECTIVE ALPHA ADRENERGIC	MC	ALPHAGAN SOLN	MC/DEL	,'	BRIMONIDINE TARTRATE DROPS 0.15 %	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
AGONISTS	MC	ALPHAGAN SOLIN ALPHAGAN P 0.1% SOLN	MC/DEL	,	IOPIDINE SOLN		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
1   1		ALPHAGAN P 0.1% SOLN ALPHAGAN P 0.15% SOLN	WIC/DLL	,	IOPIDINE SOLIN		preferred drug(s) exists.
/	MC MC/DEL		l j	,	1	,	
/	MC/DEL	BRIMONIDINE DROPS 0.2 %	l j	,	1	,	·
l	MC/DEL	SIMBRINZA					
OP ANTI-ALLERGICS	MC/DEL	AZELASTINE HCL DROPS	MC	8	ALOCRIL SOLN		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 as the Price Authorization form such as the presence of a condition that proved unless an acceptable clinical exception is offered
/ <b> </b>	MC	BEPREVE	MC/DEL	8	ALOMIDE SOLN	,	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.
<b>/                                    </b>	MC/DEL	CROMOLYN SODIUM DROPS	MC/DEL	8	EMADINE SOLN	'	preferred drug(s) exists.
<b>/                                    </b>	MC/DEL	KETOTIFEN FUMARATE DROPS	MC	8	OPTICROM SOLN	'	!
<b>/                                    </b>	1	LASTACAFT	MC/DEL	8	PATANOL SOLN	'	· [
<b>/                                    </b>	MC	1	- 1 - 1	,	1	'	
<b>/                                    </b>	MC/DEL	OLOPATADINE HCL 0.1%	MC	8	ZERVIATE	'	
<b>/                                    </b>	MC/DEL	OLOPATADINE HCL 0.2%	MC/DEL		EPINASTINE	'	·
<b>/                                    </b>	MC/DEL	ZADITOR SOLN	- 1 - 1	,	1	'	·
<b>/                                    </b>	1	1		,	1	'	
OP. ANTI-ALLERGICS- MASTCELL	<del></del>	+	MC/DEL	,'	ALAMAST SOLN	Use PA Form# 20420	+
STABILIZER CLASS	1			, 4			
OP CARBONIC ANHYDRASE	MC/DEL	AZOPT SUSP	MC/DEL	, ——	COSOPT SOLN PF	Use PA Form# 20420	
INHIBITORS/COMBO	MC	COMBIGAN	- 1 - 1	,	1		
<b>                                     </b>	MC/DEL	DORZOLAMIDE	- 1 - 1	,	1	'	
<b>/                                    </b>	MC/DEL	DORZOLAMIDE/TIMOLOL	1 )	,	1	,	
OP NSAID'S	MC	ACULAR SOLN <sup>1</sup>	MC	8	ACULAR LS <sup>1</sup>	Must fail all 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
4   O							i rotation a arage made by area and area and area and area and area area area area area area area are
1	MC/DEL	DUREZOL	MC		BROMSITE <sup>1</sup>		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

	MC/DEL	KETOROLAC OPTH 0.4%	MC/DEL	8	DEXAMETHASONE DROPS	prererrea.	preferred drug(s) exists.
	MC/DEL	KETOROLAC OPTH 0.5%	MC/DEL	8	DICLOFENAC OPTH 0.1%		
	MC/DEL	MAXIDEX SUSP	MC	8	FLURBIPROFEN SODIUM SOLN		
	MC/DEL	NEVANAC		•			
			MC/DEL	8	ILEVRO		
	MC/DEL	PREDNISOLONE DROPS	MC/DEL	8	LOTEMAX SM DROPS GEL 0.38%		
			MC/DEL	8	PROLENSA		
			MC	8	OCUFEN SOLN <sup>1</sup>		
			MC	8	XIBROM <sup>1</sup>		
			MC	8	VOLTAREN SOLN <sup>1</sup>		
			MC	8	ACUVAIL <sup>1</sup>		
			MC/DEL	9	BROMFENAC		
			IIIO/DEE	Ü	Errom Ervio	Use PA Form# 20420	
OP OF INTEREST		2001 222 222 222			2000007		Next fell adequate trials of sulfi assats from adjustic tops and lubricant selector.
OP OF INTEREST	MC/DEL	CYCLOSPORINE OPTH 0.05%	MC		BYOOVIZ	PA required to confirm appropriate diagnosis and	Must fail adequate trials of multi agents from artificial tears and lubricant category.
	MC	EYSUVIS <sup>2</sup>	MC		BEOVU	clinical parameters for use.	Beovu is non-preferred and indicated for the treatment of Neovascular (wet) Age-Related Macular Degeneration (AMD)
	MC	LUCENTIS	MC		BOTOX SOLR	cimical parameters for use.	
	MC	RESTASIS DROPPERETTE	MC/DEL		CEQUA		
	MC	XIIDRA	MC		CIMERLI		
			MC		CYCLOSPORINE DROPERETTE		
			MC		CYSTADROPS <sup>1</sup>	2. For the short-term (up to	
					CYSTARAN <sup>1</sup>	two weeks) treatment of the	
			MC			signs and symptoms of dry	
			MC		EYLEA	eye disease.	Eylea is non-preferred and indicated for the treatment of: Neovascular (wet) Age-Related Macular Degeneration (AMD), Macular Edema following Retinal Vein Occlusion (RVO), Diabetic
			MC		EYLEA HD <sup>1</sup>		Macular Edema (DME), Diabetic Retinopathy (DR)
			MC		IZERVAY <sup>1</sup>		Luxturna will be considered for the treatment of patients with confirmed biallelic RPE65 mutation-associated retinal dystrophy. Patients must have viable retinal cells as determined by the
			MC		LUCENTIS		treating physician(s).
			МС		LUXTURNA		
			MC/DEL		MIEBO		Miebo is non-preferred and is indicated for the treatment of the signs and symptoms of dry eye disease (DED).
			MC/DEL		OXERVATE		Oxervate is non-preferred and is indicated for the treatment of neurotrophic keratitis.
			MC		PAVBLU		Pavblu: Clinical rationale for why Eylea cannot be used
			MC/DEL		RESTASIS MULTIDOSE DROPS		
			MC		SUSVIMO		
			MC		SYFOVRE		Syfovre is non-preferred and is indicated for the treatment of geographic atrophy (GA) secondary to age-related macular degeneration (AMD).
			MC		TYRVAYA		
			MC		VABYSMO		
			MC		VERKAZIA		
							Vivvo Must fell adaptivate trials of multi-graphs from additional techniques and a professed audiopassing alternative
			MC		VEVYE	H DA F# 00400	Vevye - Must fail adequate trials of multi agents from artificial tears and lubricant category and a preferred cyclosporine alternative.
						Use PA Form# 20420	
		DERMATOLOGICAL				_	
ISOTRETINION, ACNE	MC	AMNESTEEM <sup>1</sup>	MC		ABSORICA	<ol> <li>Users 24 or under, PA will</li> </ol>	
	MC	CLARAVIS <sup>1</sup>	MC		ABSORICA LD	not be required.	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
	MC	MYORISAN <sup>1</sup>					preferred drug(s) exists.
	MC	ZENATANE'				Use PA Form# 20420	
TOPICAL - ACNE PREPARATIONS	MC	ERYDERM SOLN	MC/DEL		ADAPALENE 0.3% GEL	1. Users 24 or under, PA will	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/DEL	ERYTHROMYCIN GEL	MC/DEL		AKLIEF <sup>6</sup>	not be required	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
	MC/DEL	ERYTHROMYCIN SOLN	MC		ALTINAC CREA	Dosing limits allowing one	preferred drug(s) exists.
	MC/DEL	EVOCLIN	MC/DEL		ALTRENO	package per month. Please	
	MC	ISOTRETINOIN			AMZEEQ <sup>6</sup>	refer to Dose Consolidation	
			MC		ARAZLO LOTION <sup>6</sup>	List.	
	MC	METRONIDAZOLE CREA <sup>2</sup>	MC				
	MC	METRONIDAZOLE GEL <sup>2</sup>	MC		AVITA CREA	Only available if	
	MC	METRONIDAZOLE LOTN <sup>2</sup>	MC		BENZAC	component ingredients are	
	MC/DEL	TRETINOIN .025%, .05%, .01% GEL1	MC/DEL		BENZACLIN GEL <sup>3</sup>	unavailable.	
	MC	TRETINOIN CREA <sup>1,2</sup>	MC/DEL		BENZAGEL-10 GEL	4. Dosing limits apply,	
	l		MC/DEL		BENZAMYCIN GEL	please see dosing	
			MC/DEL		BENZAMYCINPAK PACK	consolidation list.	
						5 11 16 1	
			MC		BENZEFOAM	5. Not approved for use in	
	l l	l	MC		BENZOYL PEROXIDE	children <12 years of age	I and the second

				MC MC MC/DEL MC MC MC MC/DEL MC MC MC MC MC MC MC MC MC MC MC MC MC		6. For the treatment of patients ≥ 9 years of age.  Use PA Form# 10220 for Brand Name requests  Use PA Form# 20420 for all other requests	
				MC MC MC/DEL MC MC MC MC MC MC MC MC MC	EPSOLAY ERYCETTE PADS FINEVIN CREA KLARON LOTN METROCREAM CREA <sup>2</sup> METROGEL GEL <sup>2</sup> METROLOTION LOTN <sup>2</sup> NEOBENZ MICRO NORITATE CREA ONEXTON <sup>5</sup>		
				MC/DEL MC MC MC MC MC MC/DEL MC MC/DEL MC MC MC MC	PLIXDA RETIN-A GEL <sup>2</sup> RETIN-A CREA <sup>2</sup> RETIN-A MICRO GEL RHOFADE SODIUM SULFACET/SULF LOTN SOOLANTRA <sup>4</sup> TRIAZ TWYNEO VELTIN WINLEVI <sup>5</sup>		
TOPICAL- ATOPIC DERMATITIS	MC/DEL	1	ELIDEL CREA	MC MC MC/DEL MC	ZENCIA WASH ZETACET ZIANA ZILXI CIBINQO		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
	MC/DEL MC/DEL MC MC/DEL MC MC/DEL MC	1 1 1 2 2 2 2	PIMECROLIMUS CRE (AUTH GENERIC LABELER 68682 Oceanside Pharmaceuticals) PROTOPIC OINT TACROLIMUS OINT ADBRY <sup>2,4</sup> DUPIXENT <sup>1,2,4</sup> EUCRISA <sup>2,4</sup> OPZELURA <sup>2,3,4</sup>	MC MC	EBGLYSS <sup>2,3</sup> NEMLUVIO	1.Avoid live vaccines if treated with Dupixent 2. Clinical PA required. 3. For the treatment of patients ≥ 12 years of age. 4. Preferred after a trial and failure of TCI.  Use PA Form# 20420	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.
	MC MC/DEL MC/DEL		BACIT/NEOMYCIN/POLYM OINT BACITRACIN OINT GENTAMICIN SULFATE	MC/DEL MC/DEL MC/DEL		please see dosing	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	MUPIROCIN OINT <sup>1</sup>	МС		XEPI	<u>Use PA Form# 20420</u>	
TOPICAL - ANTIFUNGALS	MC/DEL MC/DEL MC MC MC/DEL MC MC/DEL	BETAMETHASONE CLOTRIMAZOLE CREA BETAMETHASONE CLOTRIMAZOLE LOT CICLOPIROX 0.77 CREA CICLOPIROX 0.77 SUSP CLOTRIMAZOLE ECONAZOLE NITRATE CREA KETOCONAZOLE CREA KETOCONAZOLE SHAM LOPROX 1.0 CREA LOPROX 1.0 LOTN LOPROX GEL LOPROX TS LOTN MICONAZOLE NITRATE CREA MYCO-TRIACET II CREA NYSTATIN NYSTATIN/TRIAMCINOLONE CREA NYSTOP POWD TRI-STATIN II CREA	MC/DEL MC MC MC MC MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC MC/DEL MC MC/DEL MC MC/DEL MC MC/DEL MC MC/DEL MC MC/DEL MC MC MC/DEL MC MC MC/DEL MC MC MC/DEL MC MC MC MC MC MC MC MC MC MC MC MC MC	8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	CICLOPIROX SOLN EXELDERM FUNGIZONE CREA HYDROCORT/IODOQ CREA JUBLIA KERYDIN¹ LOPROX 0.77 CREA LOPROX 0.77 CREA LOPROX 0.77 SUSP LOPROX SHAMPOO SHAM LOTRISONE LOT LOTRISONE CREA LUZU MENTAX CREA MYCOGEN II CREA NAFTIN NIZORAL SHAM NYSTATIN/TRIAMCINOLONE OINT NYSTAT-RX POWD OXISTAT PENLAC NAIL LACQUER SOLN	Use PA Form# 10120  1. Diagnosis required	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: 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, Onglyza or Omeprazole.  Kerydin- Verify prior trials and failures or intolerance of preferred treatments, including both topical and oral agents
TOPICAL - ANTIPRURITICS	MC	ZONALON CREA	MC MC		Korsuva Prudoxin 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 - ANTIPSORIATICS	MC/DEL	CALCIP/BETAMETHASONE SUS	MC/DEL MC/DEL MC MC MC MC MC MC MC/DEL MC MC MC	7 8 8 8 8 8 8	TACLONEX <sup>1</sup> DUOBRII ENSTILAR OXSORALEN ULTRA CAPS <sup>1</sup> PSORIATEC CREA <sup>1</sup> SORIATANE CK KIT <sup>1</sup> VECTICAL <sup>1</sup> VTAMA ZORYVE	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.
TOPICAL - ANTISEBORRHEICS	MC/DEL	SELENIUM SULFIDE SHAM	MC MC		CARMOL SCALP TREATMENT KIT ZNP BAR ZORYVE 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.  Zoryve Foam: For the treatment of seborrheic dermatitis in adult and pediatric patients 9 years of age and older.
TOPICAL - ANTIVIRALS			MC/DEL MC/DEL MC		ACYCLOVIR OINT DENAVIR CREA <sup>1,3</sup> YCANTH	Must fail oral treatment with Acyclovir or Valacyclovir.	

efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered on condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the
efficacy or intolerable side effects before non-preferred drugs will be approved, unless an acceptable clinical exception is offered of a condition that prevents usage of the preferred drug or a significant potential drug interaction between another drug and the
ust be tried and failed due to lack of efficacy or intolerable side effects before non-preferred drugs will be approved, unless an
thorization form, such as the presence of a condition that prevents usage of the preferred drug or a significant potential drug
g(s) exists.
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		VERY HIGH POTENCY	MC		BETAMETHASONE DIPROPIONATE		
		VERY HIGH POTENCY	MC/DEL		DESOXIMETASONE 0.25% CREA/OINT		
	MC/DEL	AUGMENTED BETA DIP			VERY HIGH POTENCY		
<b> </b>	MC/DEL	BETAMETHASONE VALERATE	MC/DEL		BRYHALI LOTN		
<b>                                     </b>	MC	DIFLORASONE DIACETATE	MC/DEL		CLOBETASOL PROPINATE LOTN		
<b> </b>	MC	HALOBETASOL	MC/DEL		CLOBETASOL PROPINATE SHAMPOO 0.05%		
<b>                                     </b>			MC/DEL		CORMAX		
<b> </b>			MC/DEL		DIPROLENE		
<b>[ ]</b>			MC/DEL		IMPEKLO <sup>4</sup>		
<b>                                     </b>		MISCELLANEOUS	MC/DEL		LEXETTE		
<b>                                     </b>	MC	PROCTO-KIT CREA 1%	MC/DEL		OLUX FOAM		
<b>                                     </b>			MC/DEL		PSORCON		
<b>                                     </b>							
<b>                                     </b>			MC/DEL		PSORCON E		
<b>                                     </b>			WIC/DEL		PSURCON E		
<b>                                     </b>					_		
<b>                                     </b>			MC		SERNIVO SPRAY <sup>2</sup>		
<i>1</i>			MC/DEL		TEMOVATE		
<i>1</i>			MC		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
TOPICAL - STEROID COMBINATIONS	MC	DERMA-SMOOTHE-FS SCALP	MC		CARMOL-HC CREA		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
<b>/</b>					<u> </u>		essfersed drug(e) eviate
TOPICAL - EMOLLIENTS	MC/DEL	AMMONIUM LACTATE CREA <sup>1</sup>	MC		LAC-HYDRIN CREA <sup>1</sup>	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
<b>/</b> 1	MC	AMMONIUM LACTATE LOTN 12% <sup>1</sup>	MC		LAC-HYDRIN LOTN 12%		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.
l I	MC	VITAMIN A & D MEDICATED OINT	MC		MEDERMA GEL	Dosing limits still apply.	preterred drug(s) exists.
<b>[ ]</b>			MC		MIMYX	Please see dose	
<b>[ ]</b>			MC		RENOVA CREA	consolidation list.	
<b>[ ]</b>					1		
TOPICAL - ENZYMES / KERATOLYTICS /	<del>                                     </del>	<del></del>	MC		CARMOL 40 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
UREA			MC		SALEX CREA	00017110111 20120	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
'			MC		SALEX LOTN		preferred drug(s) exists.
<b>[ ]</b>					OALEA LOTTE		
<b>[ ]</b>					1		Ziox, Panafil and Papain products have been removed from the PDL due to FDA safety concerns regarding drugs containing Papain.
<b>[ ]</b>							ZIOX, Panaili and Papain products have been removed from the PDL due to PDA salety concerns regarding drugs containing rapain.
TOTION OF WITH WARTO			<del></del>		POR OF HOVE ON HE		<u> </u>
TOPICAL - GENITAL WARTS	MC/DEL	IMIQUIMOD 5% <sup>2</sup>	MC/DEL	5	PODOFILOX SOLN	Use PA Form# 20420	
'			MC/DEL	8	CONDYLOX <sup>1</sup>	Non-preferred products	
<b>[ ]</b>			MC/DEL	8	ALDARA <sup>1</sup>	must be used in specified order.	
<b>[ ]</b>			MC	8	PICATO	Oluei.	
<b>[ ]</b>			MC	8	VEREGEN <sup>1</sup>	<ol><li>Dosing limits still apply.</li></ol>	
<b>[ ]</b>			MC	8	ZYCLARA <sup>1</sup>	Please see dose	
<b>4</b> I					1	consolidation list.	·
TOPICAL - LOCAL ANESTHETICS	MC	AF CAPSICUM OLEORESIN CREA	MC/DEL		EMLA PADS	1. Lidocaine/Prilocaine	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
<b>4</b> 1	MC/DEL	CAPSAICIN CREA	MC/DEL		EMLA CREA	cream and Ela-Max products	ts 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
<b>[ ]</b>	MC/DEL	CAPSAICIN PATCH	MC		LIDA MANTLE CREA	require PA for users over 18	
<b>[ ]</b>	MC/DEL	DIBUCAINE OINT	MC		PONTOCAINE SOLN	years of age.	
<b>[ ]</b>	MC	ELA-MAX <sup>1</sup>	MC		SYNERA		
<b>     </b>	MC/DEL		MC		ZOSTRIX		
<b>     </b>	MC/DEL	LIDOCAINE/PRILOCAINE CREA <sup>1</sup> LIDOCAINE CREAM			ZTLIDO <sup>2</sup>	Dosing limits still apply.	
<b>[ ]</b>			MC/DEL		ZILIDO	Please see dose	
<b>[ ]</b>	MC/DEL	LIDOCAINE GEL			1	consolidation list.	
<b>[ ]</b>	MC/DEL	LIDOCAINE PTCH 5%			1		
						Use PA Form# 20420	
TOPICAL - DEPIGMENTING AGENTS			MC	8	ALUSTRA 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.
<b>4  </b>			MC	8	EPIQUIN MICRO		
41			MC	8	GLYQUIN CREA		
<b>∐</b> •			•		1	•	

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 <sup>1</sup>	MC/DEL MC MC MC MC MC MC MC MC MC MC MC MC MC/DEL MC MC MC MC MC/DEL	8 8 8 8 9	HYDROQUINONE CREA HYDROQUINONE/SUNSCREENS SOLAQUIN FORTE CREA TRI-LUMA CREA ELDOQUIN ELIMITE CREA EURAX LINDANE MALATHION OVIDE LOTN SPINOSAD SUSP	Use PA Form# 20420  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		FILSUVEZ 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 02 >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		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		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	AK-DILATE SOLN	MC		LENS PLUS REWETTING DROPS	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	EYE WASH SOLN	MC/DEL	i	MURO 128	0301711 5	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
	MC	NAPHAZOLINE HCL SOLN	MC	1	NEO-SYNEPHRINE SOLN		preferred drug(s) exists.
	MC	PHENYLEPHRINE HCL SOLN	1 " )	i	NEO-OTHERTHANE GGE.		
/ <b> </b>	MC	PONTOCAINE SOLN	1 )	i			
/ <b> </b>	MC/DEL	SODIUM CHLORIDE	1 )	i			
	MODEL	MISCELLANEOUS EAR					
EAR	MC/DEL	A/B OTIC SOLN	MC		ANTIBIOTIC EAR 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
EAR	MC MC	ACETASOL SOLN	MC	4	ANTIBIOTIC EAR SOLN ANTIBIOTIC EAR SUSP	Use PA FORM# ZU4ZU	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
. 1		ACETASOL SOLN ACETASOL HC SOLN		4			preferred drug(s) exists.
. 1	MC/DEL MC/DEL	ACETIC ACID	MC/DEL	4	CIPRODEX		
. 1		ACETIC ACID  ACETIC ACID/HYDROCORTISON	MC/DEL MC/DEL	4	CIPROFLOXACIN HCL DEBROX SOLN		
1	MC/DEL			4			
1	MC/DEL	ALLERGEN SOLN	MC	4	DERMOTIC		
1	MC	CARBAMIDE PEROXIDE 6.5% OTIC SOLN.	MC	4	FLOXIN		
1	MC/DEL	CIPRO HC SUSP	MC	4	OTIPRIO		
. 1	MC/DEL	CORTISPORIN-TC SUSP	MC	4	OTOVEL		
i <b>I</b>	MC/DEL	CORTOMYCIN	1 )	4			
. 1	MC	COLY-MYCIN-S SUSP	1 )	4			
i	MC	EAR DROPS SOLN	1 )	4			
11 !	MC	EAR DROPS RX SOLN	1 )	4			
/ <b> </b>	MC/DEL	EAR WAX REMOVAL DROPS	l ,	1			
•	]	1	1 1		•	•	

	MC	FLUOCINOLONE ACETONIDE OIL DROPS 0.01%	1	Ī	I	
	MC/DEL	NEOMYCIN/POLYMYXIN/HC	1 1			1
	MC/DEL	OFLOXACIN 0.3% OTIC	1			1
		MOUTH ANTISEPTICS	_			
MOUTH ANTI-INFECTIVES	MC	NILSTAT SUSP	MC	MYCELEX TROC	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/DEL	NYSTATIN SUSP	MC	ORAVIG	USE FA FUITH ZUYZU	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
	MICIDEE	INTOTATIN SOSI		ORAVIG		preferred drug(s) exists.
MOUTH ANTISEPTICS	MC/DEL	CHLORHEXIDINE GLUCONATE	MC	APHTHASOL PSTE <sup>1</sup>	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/DEL	LIDOCAINE VISCOUS SOLN	MC	PERIOGARD SOLN <sup>1</sup>	Must fail all preferred	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
	MC	TRIAMCINOLONE IN ORABASE PSTE	MC	TRIAMCINOLONE ACETONIDE PSTE <sup>1</sup>	products before non-	preferred drug(s) exists.
	MC	TRIAMCINOLONE ORADENT PSTE	1 1	MANONOLONE AGETORISE TOTE	preferred.	
		DENTAL PRODUCTS				
DENTAL PRODUCTS	MC/DEL	ETHEDENT CREA	MCOMC	APF 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
DEMINE: NOSCO.	MC/DEL	GEL-KAM CONC	MC/DEL	DENTAGEL GEL	USE FA FUITH ZUYZU	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
	MC/DEL	GEL-KAM GEL 0.4%	MC/DEL	PHOS-FLUR GEL		preferred drug(s) exists.
	MC/DEL	PHOS FLUR SOLN	MC MC	THERA-FLUR-N GEL		
	MC/DEL	SF 5000 PLUS CREA		IIILIVA LOIVIN GLE		
		SF GEL	1 1			
	MC/DEL	STANNOUS FLUORIDE ORAL RI CONC	1 1			
	MC	STANNOUS FLUORIDE ORAL RI CONC	1 1			
		<b> </b>	1			
TOTAL CALL DE CALLED		ARTIFICIAL SALIVA/STIMULANTS			-	
ARTIFICIAL SALIVA/STIMULANTS	MC	SALIVA SUBSTITUTE SOLN	MC	EVOXAC 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
ı <b>İ</b>		<b>'</b>	MC	RADIACARE SOLR		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.
	<u> </u>		MC	SALAGEN TABS		prototou drug(o) onioto.
		MISCELLANEOUS ANORECTAL				
ANORECTAL - MISC.	MC	CORTENEMA ENEM	MC/DEL	ANUSOL-HC CREA	Use PA Form# 20420	
	MC	ELA-MAX 5 CREA	MC/DEL	CORTIFOAM FOAM		
ı.	MC/DEL	HYDROCORTISONE ENEM	MC/DEL	PROCTOFOAM HC FOAM		
.I	MC/DEL	PROCTOSOL HC CREA	MC/DEL	PROCTO-KIT CREA 2.5%		
	MC/DEL	PROCTOZONE-HC CREA	MC	RECTIV OINT		
.I		<b> </b>	1			
	<u> </u>		4			
		T-CELL ACTIVATION INHIBITOR				
PSORIASIS BIOLOGICALS		ADALIMUMAB-FKJP	MC	AMJEVITA	Dosing limits apply,  places refer to decade.	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	ENBREL <sup>1,5</sup>	MC/DEL	BIMZELX <sup>3</sup>	please refer to dosage consolidation list.	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
1	MC	ENBREL SURECLICK <sup>1</sup>	MC	COSENTYX <sup>4</sup>		preferred drug(s) exists.
ı.	MC	HUMIRA <sup>1,5</sup>	MC/DEL	CYLTEZO	2.Clinical PA required and	
1	MC	OTEZLA	MC	HADLIMA	will be preferred for the indication of plaque	Cosentyx approvals for 300mg dose(s) must use "300DOSE" package (containing 2 x 150mg pens or syringes).
1		SIMLANDI	MC/DEL	HULIO	psoriasis, psoriatic arthritis	
1	MC/DEL	SKYRIZI <sup>6</sup>	MC/DEL	HYRIMOZ	and ankylosing spondylitis.	It is recommended to assess for TB infection prior to starting treatment with Taltz®.
ı <b>İ</b>	MC	TALTZ <sup>2</sup>	MC	IDACIO		
1		<b>'</b>	MC/DEL	ILUMYA <sup>3</sup>		Stelara will require using preferred trial of Skyrizi if unable please provide clinical rational as why inappropriate.
ı.		<b>,</b>	MC	SOTYKTU	3. For the treatment of adults	s.
ı.		<b>,</b>	MC/DEL	SPEVIGO	with moderate-to-severe	
ı.		<b>,</b>	MC	SILIQ	plaque psoriasis who are candidates for systemic	
ı.		<b>,</b>	MC	STELARA	therapy or phototherapy.	
1		<b>,</b>	MC	TREMFYA	(10.0p) 0. p	
1		<b>,</b>	MC	YUFLYMA		
1		<b>,</b>	MC	YUSIMRY	Please see criteria section	n n
1		<b>'</b>	IVIC	TOSINICI	7.1 10000 000 0.1.0.1.0 000	1
1		<b>'</b>	1		5. Will not require a PA if at	
ı <b>İ</b>		<b>,</b>	1 1		least one systemic drug such	
/ <b>[</b>		<b>'</b>	1		as methotrexate,	
<b>∤ I</b>	I I	<b>.</b>	1 1	I	cyclosporine, methoxsalen	1

					or acitretin is in members	
					drug profile.	
				<b>!</b>	6. Clinical PA required and	
				<b>!</b>	will be preferred for the indication of plaque	
					psoriasis, psoriatic arthritis,	
					Crohn's disease and	
					ulcerative colitis.	
					Use PA Form# 20910	
		ALTERNATIVE MEDICIN	ES			
ALTERNATIVE MEDICINES	MC	DIMETHYL SULFOXIDE SOLN	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.
	MC	MELATONIN				
		CHELATING AGENTS				
CHELATING AGENTS	MC/DEL	CUPRIMINE CAPS	MC	CLOVIQUE	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 (in step order), unless an acceptable clinical
<b>1                                    </b>			MC	DEPEN TITRATABS TABS	1. FDA indication of	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
			MC/DEL	EXJADE <sup>1</sup>	treatment of chronic iron	another drug and the preferred drug(s) exists.
			MC	SYPRINE	overload due to blood transfusions in members 2	
			MC/DEL	TRIENTINE CAPS	Transfusions in members z	Clovique® should be used when continued treatment with penicillamine is no longer possible because of intolerable or life endangering side effects.
				<b>!</b>		
				<b>!</b>		
		ANTILEPROTIC				
ANTILEPROTIC			MC	THALOMID CAPS <sup>1</sup>		Approved for indications of leprosy, treatment-resistant multiple myeloma and AIDS.
					150mg dosing will require	
			I I		use of Thalomid 100mg and	
					■hima cansilles	
					50mg capsules.	
					Use PA Form# 20420	
		ANTINEOPLASTIC AGEN				
ANTINEOPLASTIC AGENTS -	MC/DEL	ANTINEOPLASTIC AGEN	ITS MC/DEL	CASODEX	<u>Use PA Form# 20420</u>	
ANTIADNDROGENS		BICALUTAMIDE	MC/DEL		Use PA Form# 20420 Use PA Form# 20420	
ANTIADNDROGENS ANTINEOPLASTIC AGENTS- LHRH	MC/DEL MC/DEL			CASODEX  LUPRON DEPOT SYRINGEKIT	Use PA Form# 20420  Use PA Form# 20420  1. Dosing limits apply,	
ANTIADNDROGENS	MC/DEL	BICALUTAMIDE  LUPRON DEPOTSYRINGEKIT <sup>1</sup>	MC/DEL	LUPRON DEPOT SYRINGEKIT	Use PA Form# 20420  Use PA Form# 20420  1. Dosing limits apply, please refer to dosage	
ANTIADNDROGENS ANTINEOPLASTIC AGENTS- LHRH	MC/DEL	BICALUTAMIDE  LUPRON DEPOTSYRINGEKIT <sup>1</sup> LUPRON DEPOT- PED KIT <sup>1</sup> (1-month)	MC/DEL MC/DEL		Use PA Form# 20420  Use PA Form# 20420  1. Dosing limits apply, please refer to dosage consolidation list.	
ANTIADNDROGENS ANTINEOPLASTIC AGENTS- LHRH	MC/DEL	BICALUTAMIDE  LUPRON DEPOTSYRINGEKIT <sup>1</sup>	MC/DEL  MC/DEL  MC/DEL	LUPRON DEPOT SYRINGEKIT FIRMAGON <sup>2</sup>	Use PA Form# 20420  Use PA Form# 20420  1. Dosing limits apply, please refer to dosage consolidation list.  2. PA required to confirm	
ANTIADNDROGENS ANTINEOPLASTIC AGENTS- LHRH	MC/DEL MC/DEL MC/DEL	BICALUTAMIDE  LUPRON DEPOTSYRINGEKIT <sup>1</sup> LUPRON DEPOT- PED KIT <sup>1</sup> (1-month)  LUPRON DEPOT-PED SYRINGEKIT (3-month	MC/DEL  MC/DEL  MC/DEL  th)	LUPRON DEPOT SYRINGEKIT  FIRMAGON <sup>2</sup> SUPPRELIN LA (IMPLANT) KIT	Use PA Form# 20420  Use PA Form# 20420  1. Dosing limits apply, please refer to dosage consolidation list.	
ANTIADNDROGENS ANTINEOPLASTIC AGENTS- LHRH	MC/DEL	BICALUTAMIDE  LUPRON DEPOTSYRINGEKIT <sup>1</sup> LUPRON DEPOT- PED KIT <sup>1</sup> (1-month)	MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL	LUPRON DEPOT SYRINGEKIT  FIRMAGON <sup>2</sup> SUPPRELIN LA (IMPLANT) KIT  TRELSTAR	Use PA Form# 20420  Use PA Form# 20420  1. Dosing limits apply, please refer to dosage consolidation list.  2. PA required to confirm	
ANTIADNDROGENS ANTINEOPLASTIC AGENTS- LHRH	MC/DEL MC/DEL MC/DEL	BICALUTAMIDE  LUPRON DEPOTSYRINGEKIT <sup>1</sup> LUPRON DEPOT- PED KIT <sup>1</sup> (1-month)  LUPRON DEPOT-PED SYRINGEKIT (3-month	MC/DEL  MC/DEL  MC/DEL  th)	LUPRON DEPOT SYRINGEKIT  FIRMAGON <sup>2</sup> SUPPRELIN LA (IMPLANT) KIT	Use PA Form# 20420  Use PA Form# 20420  1. Dosing limits apply, please refer to dosage consolidation list.  2. PA required to confirm	
ANTIADNDROGENS ANTINEOPLASTIC AGENTS- LHRH	MC/DEL MC/DEL MC/DEL	BICALUTAMIDE  LUPRON DEPOTSYRINGEKIT <sup>1</sup> LUPRON DEPOT- PED KIT <sup>1</sup> (1-month)  LUPRON DEPOT-PED SYRINGEKIT (3-month	MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL	LUPRON DEPOT SYRINGEKIT  FIRMAGON <sup>2</sup> SUPPRELIN LA (IMPLANT) KIT  TRELSTAR	Use PA Form# 20420  Use PA Form# 20420  1. Dosing limits apply, please refer to dosage consolidation list.  2. PA required to confirm	
ANTIADNDROGENS ANTINEOPLASTIC AGENTS- LHRH	MC/DEL MC/DEL MC/DEL	BICALUTAMIDE  LUPRON DEPOTSYRINGEKIT <sup>1</sup> LUPRON DEPOT- PED KIT <sup>1</sup> (1-month)  LUPRON DEPOT-PED SYRINGEKIT (3-month	MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL	LUPRON DEPOT SYRINGEKIT  FIRMAGON <sup>2</sup> SUPPRELIN LA (IMPLANT) KIT  TRELSTAR	Use PA Form# 20420  1. Dosing limits apply, please refer to dosage consolidation list.  2. PA required to confirm FDA approved indication.	
ANTIADNDROGENS ANTINEOPLASTIC AGENTS- LHRH ANALOGS	MC/DEL MC/DEL MC/DEL	BICALUTAMIDE  LUPRON DEPOTSYRINGEKIT <sup>1</sup> LUPRON DEPOT- PED KIT <sup>1</sup> (1-month)  LUPRON DEPOT-PED SYRINGEKIT (3-month	MC/DEL  MC/DEL  MC/DEL  th)  MC/DEL  MC/DEL  MC/DEL  MC/DEL	LUPRON DEPOT SYRINGEKIT  FIRMAGON <sup>2</sup> SUPPRELIN LA (IMPLANT) KIT  TRELSTAR  VANTAS <sup>2</sup>	Use PA Form# 20420  Use PA Form# 20420  1. Dosing limits apply, please refer to dosage consolidation list.  2. PA required to confirm	
ANTINEOPLASTIC AGENTS - LHRH ANALOGS  ANTINEOPLASTIC AGENTS - TYROSINE	MC/DEL MC/DEL MC/DEL	BICALUTAMIDE  LUPRON DEPOTSYRINGEKIT <sup>1</sup> LUPRON DEPOT- PED KIT <sup>1</sup> (1-month)  LUPRON DEPOT-PED SYRINGEKIT (3-month	MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC	LUPRON DEPOT SYRINGEKIT  FIRMAGON <sup>2</sup> SUPPRELIN LA (IMPLANT) KIT  TRELSTAR  VANTAS <sup>2</sup> SPRYCEL <sup>1</sup>	Use PA Form# 20420  1. Dosing limits apply, please refer to dosage consolidation list. 2. PA required to confirm FDA approved indication.  Use PA Form# 20420  Use PA Form# 20420	
ANTIADNDROGENS ANTINEOPLASTIC AGENTS- LHRH ANALOGS	MC/DEL MC/DEL MC/DEL	BICALUTAMIDE  LUPRON DEPOTSYRINGEKIT <sup>1</sup> LUPRON DEPOT- PED KIT <sup>1</sup> (1-month)  LUPRON DEPOT-PED SYRINGEKIT (3-month	MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC  MC	LUPRON DEPOT SYRINGEKIT  FIRMAGON <sup>2</sup> SUPPRELIN LA (IMPLANT) KIT  TRELSTAR  VANTAS <sup>2</sup> SPRYCEL <sup>1</sup> TYKERB <sup>2</sup>	Use PA Form# 20420  1. Dosing limits apply, please refer to dosage consolidation list.  2. PA required to confirm FDA approved indication.  Use PA Form# 20420  Use PA Form# 20420  1. Verification of diagnosis	
ANTINEOPLASTIC AGENTS - LHRH ANALOGS  ANTINEOPLASTIC AGENTS - TYROSINE	MC/DEL MC/DEL MC/DEL	BICALUTAMIDE  LUPRON DEPOTSYRINGEKIT <sup>1</sup> LUPRON DEPOT- PED KIT <sup>1</sup> (1-month)  LUPRON DEPOT-PED SYRINGEKIT (3-month	MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC	LUPRON DEPOT SYRINGEKIT  FIRMAGON <sup>2</sup> SUPPRELIN LA (IMPLANT) KIT  TRELSTAR  VANTAS <sup>2</sup> SPRYCEL <sup>1</sup>	Use PA Form# 20420  1. Dosing limits apply, please refer to dosage consolidation list.  2. PA required to confirm FDA approved indication.  Use PA Form# 20420  Use PA Form# 20420  1. Verification of diagnosis is required.	
ANTINEOPLASTIC AGENTS - LHRH ANALOGS  ANTINEOPLASTIC AGENTS - TYROSINE	MC/DEL MC/DEL MC/DEL	BICALUTAMIDE  LUPRON DEPOTSYRINGEKIT <sup>1</sup> LUPRON DEPOT- PED KIT <sup>1</sup> (1-month)  LUPRON DEPOT-PED SYRINGEKIT (3-month	MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC  MC	LUPRON DEPOT SYRINGEKIT  FIRMAGON <sup>2</sup> SUPPRELIN LA (IMPLANT) KIT  TRELSTAR  VANTAS <sup>2</sup> SPRYCEL <sup>1</sup> TYKERB <sup>2</sup>	Use PA Form# 20420  1. Dosing limits apply, please refer to dosage consolidation list.  2. PA required to confirm FDA approved indication.  Use PA Form# 20420  Use PA Form# 20420  1. Verification of diagnosis is required.  2. PA required to confirm	
ANTINEOPLASTIC AGENTS - LHRH ANALOGS  ANTINEOPLASTIC AGENTS - TYROSINE	MC/DEL MC/DEL MC/DEL	BICALUTAMIDE  LUPRON DEPOTSYRINGEKIT <sup>1</sup> LUPRON DEPOT- PED KIT <sup>1</sup> (1-month)  LUPRON DEPOT-PED SYRINGEKIT (3-month	MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC  MC	LUPRON DEPOT SYRINGEKIT  FIRMAGON <sup>2</sup> SUPPRELIN LA (IMPLANT) KIT  TRELSTAR  VANTAS <sup>2</sup> SPRYCEL <sup>1</sup> TYKERB <sup>2</sup>	Use PA Form# 20420  1. Dosing limits apply, please refer to dosage consolidation list.  2. PA required to confirm FDA approved indication.  Use PA Form# 20420  Use PA Form# 20420  1. Verification of diagnosis is required.  2. PA required to confirm FDA approved indication and	
ANTINEOPLASTIC AGENTS - LHRH ANALOGS  ANTINEOPLASTIC AGENTS - TYROSINE	MC/DEL MC/DEL MC/DEL	BICALUTAMIDE  LUPRON DEPOTSYRINGEKIT <sup>1</sup> LUPRON DEPOT- PED KIT <sup>1</sup> (1-month)  LUPRON DEPOT-PED SYRINGEKIT (3-month	MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC  MC	LUPRON DEPOT SYRINGEKIT  FIRMAGON <sup>2</sup> SUPPRELIN LA (IMPLANT) KIT  TRELSTAR  VANTAS <sup>2</sup> SPRYCEL <sup>1</sup> TYKERB <sup>2</sup>	Use PA Form# 20420  1. Dosing limits apply, please refer to dosage consolidation list.  2. PA required to confirm FDA approved indication.  Use PA Form# 20420  Use PA Form# 20420  1. Verification of diagnosis is required.  2. PA required to confirm FDA approved indication and to monitor for potential drug-	
ANTINEOPLASTIC AGENTS - LHRH ANALOGS  ANTINEOPLASTIC AGENTS - TYROSINE	MC/DEL MC/DEL MC/DEL	BICALUTAMIDE  LUPRON DEPOTSYRINGEKIT <sup>1</sup> LUPRON DEPOT- PED KIT <sup>1</sup> (1-month)  LUPRON DEPOT-PED SYRINGEKIT (3-month	MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC  MC	LUPRON DEPOT SYRINGEKIT  FIRMAGON <sup>2</sup> SUPPRELIN LA (IMPLANT) KIT  TRELSTAR  VANTAS <sup>2</sup> SPRYCEL <sup>1</sup> TYKERB <sup>2</sup>	Use PA Form# 20420  1. Dosing limits apply, please refer to dosage consolidation list.  2. PA required to confirm FDA approved indication.  Use PA Form# 20420  Use PA Form# 20420  1. Verification of diagnosis is required.  2. PA required to confirm FDA approved indication and	
ANTINEOPLASTIC AGENTS - LHRH ANALOGS  ANTINEOPLASTIC AGENTS - TYROSINE	MC/DEL MC/DEL MC/DEL	BICALUTAMIDE  LUPRON DEPOTSYRINGEKIT <sup>1</sup> LUPRON DEPOT- PED KIT <sup>1</sup> (1-month)  LUPRON DEPOT-PED SYRINGEKIT (3-month	MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC/DEL  MC  MC	LUPRON DEPOT SYRINGEKIT  FIRMAGON <sup>2</sup> SUPPRELIN LA (IMPLANT) KIT  TRELSTAR  VANTAS <sup>2</sup> SPRYCEL <sup>1</sup> TYKERB <sup>2</sup>	Use PA Form# 20420  1. Dosing limits apply, please refer to dosage consolidation list.  2. PA required to confirm FDA approved indication.  Use PA Form# 20420  Use PA Form# 20420  1. Verification of diagnosis is required.  2. PA required to confirm FDA approved indication and to monitor for potential drug-	

	MC/DEL	MERCAPTOPURINE	MC/DEL	ELOXATIN	l	
	MC/DEL	OXALIPLATIN	MC/DEL	ETHYOL		
			MC	LEUPROLIDE		
			MC/DEL	PURINETHOL		
			MC/DEL	ZOLINZA		
			O,DEE	LOLINAN.		
ANTINEOPLASTICS- MONOCLONAL	MC/DEL	TRAZIMERA	+ +			
ANTIBODIES			MC/DEL	ENHERTU		
•			MC/DEL	HERCEPTIN		
			MC	HERCESSI		•
			MC.DEL	HERZUMA		
I			MC	KANJINTI		
			MC	OGIVRI		
			MC/DEL	ONTRUZANT	Use PA Form# 20420	
		CANCER	WO/DEE	ONTROZANT	03e1 A 1 0111# 20420	
CANCER	MC	ALIMTA	MC	ABECMA	PA required to confirm	
	MC/DEL	ANASTROZOLE TABS	MC	AKEEGA	appropriate diagnosis and	
	MC	ERBITUX	MC	ALECENSA	testing.	All non-preferred: A clinical PA is required to confirm appropriate clinical indication for the individual drug request. Specific to each drug all age, clinical testing requirements, previous step
				ALIQOPA <sup>3</sup>		therapies, adjunctive drug therapy requirements, and response without disease progression will be also be evaluated for clinical appropriateness. The standard for the appropriate
	MC	IMATINIB MESYLATE	MC/DEL			indication will include the FDA label as well as current NCCN guidelines
	MC/DEL	LETROZOLE	MC	ALUNBRIG <sup>1</sup>	<ol><li>Avoid CYP3A drug interaction.</li></ol>	
	MC	RUXIENCE	MC	ALYMSYS		
	MC/DEL	VIDAZA	MC/DEL	ARIMIDEX		Scemblix is for the treatment of adult patients with: Philadelphia chromosome-positive chronic myeloid leukemia (Ph+ CML) in chronic phase (CP), previously treated with two or more tyrosine kinase inhibitors (TKIs).
	MC	ZIRABEV	MC	AUGTYRO	<ol><li>Clinical PA required for</li></ol>	
			MC	AYVAKIT	appropriate diagnosis	
			MC/DEL	AVASTIN	4. Re-approval will require	
			MC/DEL	BALVERSA	documentation of response	
			MC	BAVENCIO <sup>1,8</sup>	without disease progression	
			MC/DEL	BENDEKA <sup>3</sup>	and tolerance to treatment	
			MC/DEL	BESPONSA <sup>3</sup>	5. Dosing limits apply,	
				BESREMI <sup>1</sup>	please see dosage	
			MC		consolidation list.	
			MC	BIZENGRI		
			MC	BLENREP BOSULIF	C M d-ib. df 200	
			MC/DEL		<ol><li>Max daily dose of 300mg.</li></ol>	
			MC/DEL	BRAFTOVI <sup>1</sup>	L	
			MC	BREYANZI	7. Monitor liver enzymes	
			MC	BRUKINSA	periodically and stop treatment upon Grade 3 or	
			MC	CABOMETYX <sup>3</sup>	higher elevation of liver	
			MC	CAMCEVI	enzymes approved	
			MC/DEL	CALQUENCE <sup>3</sup>	indication	
			MC	COMETRIQ <sup>3,4,5</sup>	8. For patients ≥ 12 years of	
			MC	COTELLIC	age	
			MC/DEL	COPIKTRA	9. For the treatment of	
			MC	DANZITEN	patients up to 25 years of	
			MC	DARZALEX <sup>3</sup>	age with B-cell acute	
			MC/DEL	DAURISMO	lymphoblastic leukemia (ALL) that is refractory or in	
			MC/DEL	ELREXFIO	second or later relapse.	
			MC/DEL	EMPLICITI(IV) <sup>8</sup>		
			MC	EPKINLY		
			MC/DEL	ERLEADA		
			MC/DEL	ERIVEDGE		
			MC	EXKIVITY		
			MC	FARYDAK		
			MC/DEL	FEMARA	Use PA Form# 20420	
			MC	FOLOTYN		
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		Inner
	MC	FOTIVDA
	MC	FRUZAQLA
	MC	GAVRETO
	MC/DEL	GILOTRIF <sup>4,5</sup>
	MC/DEL	IBRANCE
	MC	ICLUSIG <sup>3</sup>
	MC/DEL	IDHIFA <sup>3</sup>
	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 <sup>1,2</sup>
	MC	JEMPERLI
	MC/DEL	JEMPERLI KEYTRUDA <sup>1</sup>
	MC	
		KIMMTRAK
	MC	KISQALI <sup>1</sup>
	MC/DEL	KOSELUGO
	MC	KRAZATI <sup>3</sup>
	MC	KYMRIAH <sup>3,9</sup>
	MC	KYPROLIS <sup>1</sup>
	MC	LARTRUVO <sup>1</sup>
	MC	LAZCLUZE
	MC	LENVIMA
	MC/DEL	LIBTAYO <sup>1</sup>
	MC	LONSURF
	MC/DEL	LORBRENA
	MC	LOQTORZI
	MC	LUMAKRAS
	MC/DEL	LUMOXITI <sup>1</sup>
	МС	LUNSUMIO <sup>1</sup>
	MC	LYNPARZA <sup>1</sup>
	MC	LYTGOBI
	MC	NEXAVAR <sup>1</sup>
	MC	NERLYNX <sup>3</sup>
	MC	NINLARO(PO)
	MC/DEL	NUBEQA
	MC MC	MARGENZA
	MC/DEL	MARGENZA MEKINIST <sup>3,4</sup>
	MC/DEL	MEKTOVI <sup>1</sup>
	MC	MONJUVI MYLOTARG <sup>3</sup>
	MC/DEL	
	MC/DEL	MVASI
	MC	ODOMZO <sup>1,2,5</sup>
	MC	OGSIVEO
	MC	OJEMDA
	MC	OJJAARA
	MC	OMISIRGE
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IMMUNOSUPPRESSANTS	MC/DEL MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL	IMMUNOSUPPRESSANTS  CYCLOSPORINE MODIFIED GENGRAF CAPS MYCOPHENOLATE MYFORTIC NEORAL SOL RAPAMUNE SANDIMMUNE TACROLIMUS CAPS	MC MC MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC/DEL MC MC/DEL MC MC MC MC MC MC MC MC MC MC MC MC MC	VONJO VORANIGO VYLOY WELIREG XALKORI XPOVIO XOSPATA XTANDI YERVOY YESCARTA³ ZALTRAP ZEJULA¹ ZELBORAF ZEPZELCA ZIIHERA ZYDELIG ZYKADIA ZYNLONTA ZYNYZ¹ ZYTIGA  CELLCEPT CYCLOSPORINE CAPS CYCLOSPORINE SOL. MODIFIED ENVARSUS XR MYHIBBIN² NEORAL CAP PROGRAF CAPS REZUROCK¹ ZORTRESS	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  2. Clinical PA is required.	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: 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 ovestatin (doses greater than 20mg).  DDI: Cyclosporine will require prior authorization when used with Livalo.  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.
IMMUNOSUPPRESSANTS- Misc.			MC	HYFTOR <sup>12</sup>		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.
		PURINE ANALOG		THE PARTY TARE		
PURINE ANALOG	MC/DEL	AZASAN TABS AZATHIOPRINE TABS	MC/DEL	IMURAN 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
		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	Use PA Form# 20420	

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New drugs are initially non-preferred until reviewed by the DUR Committee and the State. According to State policy, any drug requiring specific diagnosis still requires the specific diagnosis unless otherwise noted within this document.