

			STANDARD FORMULARY EXCEPTION CRITERIA		
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
ACE-Inhibitor/CCB Combination Product	Lotrel	amlodipine/benazepril	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
					MSB Exclusion
Acne Vulgaris Agents (Topical)	Clindagel 1% gel	clindamycin 1% gel	<b>NOTE:</b> A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product. <b>Criteria:</b> Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	*This criteria applies only to the NPF
Acne Vulgaris Agents (Topical)	Winlevi	clascoterone cream 1%	Acne Vulgaris in a patient ≥ 12 years of age.  Approve if the patient meets both of the following (A and B):  A. Patient has tried at least one prescription topical retinoid for the treatment of acne vulgaris [documentation required]; AND  Note: Examples of prescription topical retinoids include adapalene (Differin, generic), Aklief (trifarotene 0.005% cream), Arazlo (tazarotene 0.045% lotion), Fabior (tazarotene 0.1% foam), tazarotene 0.1% cream and gel (Tazorac, generic), tretinoin.  B. Patient has tried at least three other prescription non-retinoid topical therapies for the treatment of acne vulgaris [documentation required].  Note: Topical retinoids do not count. Examples of other prescription non-retinoid topical therapies for acne include: benzoyl peroxide, dapsone gel (Aczone, generic), Azelex (azelaic acid 20% cream), topical clindamycin, topical erythromycin, and topical minocycline (Amzeeq [minocycline 4% foam]). For combination products, each active chemical entity counts as one trial. Example: If one topical prescription product has 2 non-retinoids, this would fulfill a trial of 2 non-retinoid topical therapies.	1 year	Yes
Acne Vulgaris Agents (Topical) – Combination Products	Acanya Gel	benzoyl peroxide 2.5% and clindamycin phosphate 1.2% gel	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Acne Vulgaris Agents (Topical) – Combination Products	Veltin	clindamycin phosphate and tretinoin gel	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Acne Vulgaris Agents (Topical) – Combination Products	. Cahtreo	clindamycin phosphate, adapalene and benzoyl peroxide topical gel	Acne vulgaris in a patient ≥ 12 years of age.  Approve if the patient meets BOTH of the following (A and B):  A. Patient has concomitantly tried ALL three of the following products [documentation required]: 1) a topical benzoyl peroxide product, 2) a topical tretinoin-containing or adapalene-containing product, and 3) a topical clindamycin-containing product; AND  B. According to the prescriber, there is a significant clinical concern such that the patient is unable to continue to use the products in criterion A.	1 year	Yes
Acne Vulgaris Agents (Topical) – Retinoid		top.ou. go.	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the	. , ,	MSB Exclusion *This criteria applies only to
Products	Atralin	tretinoin gel (0.05%)	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Acne Vulgaris Agents (Topical) – Retinoid Products	Retin-A Micro 0.1% & 0.04% gel	tretinoin 0.1% & 0.04% gel	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Acne Vulgaris Agents (Topical) – Retinoid	Fabior and		Other diagnoses (e.g., acne vulgaris). Approve if the patient meets the following (A and B):  A. Patient has tried and cannot take one of tazarotene cream (Tazorac cream, generics) or tazarotene gel (Tazorac gel, generics), if one is formulary. If none are formulary, approve; AND  B. Patient has tried and, according to the prescriber, has experienced inadequate efficacy or significant intolerance with one topical tretinoin-containing product.  Note: Examples of topical retinoid products include tretinoin cream (Retin-A cream, generics), tretinoin gel (Retin-A gel, generics).		
Products		tazarotene 0.1% foam		1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Actinic Keratosis	Drana Name	tirbanibulin ointment	Approve if the patient has tried two of the following products: diclofenac 3% gel, a fluorouracil-containing product (e.g., fluorouracil cream, Carac, fluorouracil topical solution), or an imiquimod-containing product (e.g., imiquimod 5%	Duration	Wicarcarton
Agents (Topical)	Klisyri	1%	cream, Zyclara).	1 year	Yes
	Carac and				
Actinic Keratosis Agents (Topical)	authorized generic 0.5%	fluorouracil 0.5% cream	Approve if the patient has tried one of the following products, if formulary: Tolak, Fluoroplex, fluorouracil 2% solution, fluorouracil 5% solution, or fluorouracil 5% cream (Efudex, generics). If none are formulary, approve.	1 vear	Yes
Actinic Keratosis	Zyclara 2.5% and	imiguimod 2.5% and	Approve if the patient has thed one of the following products, in formularly. Totak, Pridorophex, Indorodracil 2% solution, or indorodracil 3% cream (Eludex, generics). If none are formularly, approve.	i year	res
Agents (Topical)	3.75%	3.75% cream	Approve if the patient has tried imiquimod 5% cream (Aldara, generics), if formulary. If imiquimod 5% cream (Aldara, generics) is non-formulary, approve.	1 year	Yes
9 (р)			For allogeneic hematopojetic stem cell transplantation in patients with acute myeloid leukemia or myelodysplastic syndrome.	, , ,	1
		treosulfan intravenous	1. Approve if the patient has tried OR, according to the prescriber, is unable to use one other conditioning regimen (e.g., a regimen containing busulfan, cyclophosphamide, fludarabine, or etoposide).		
Alkylating Agent	Grafapex	infusion	2. Approve if the patient has already been started on therapy with Grafapex.	30 days	Yes
		peanut [Arachis			
A.II		hypogaea] allergen			
Allergen Immunotherapy	Palforzia	powder-dnfp for oral administration	See standard Allergen Immunotherapy - Palforzia Prior Authorization Policy criteria.	See PA duration	Yes
пппипопетару	FallUIZIA	aummistration	рее stantuarti Aniergen inimianotrierapy – Paniorzia Prior Autriorization Policy Citteria.	duration	MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Alpha and beta-			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
blocker	Coreg	carvedilol tablet	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Alpha-1 Adrenoceptor			1. Approve if the patient tried and cannot take terazosin capsules. If terazosin capsules are non-formulary, approve.		
Antagonist	Tezruly		2. Approve if the patient is unable to swallow capsules or has difficulty swallowing capsules.	1 year	Yes
Alabad Badalaa		alpha1-proteinase	A LAA MARINA DE CARANTA DE LA CARANTA DE LA CARANTA DE LA CARANTA DE CARANTA		
Alpha1 Proteinase Inhibitors	Zemaira	inhibitor [human] lyophilized powder	Alpha1-Antitrypsin Deficiency with Emphysema (or Chronic Obstructive Pulmonary Disease); Alpha1-Antitrypsin Deficiency-Associated Panniculitis:  Approve if the patient has tried two formulary alternatives from the following list, if formulary (or one if one is formulary): Aralast NP, Glassia, or Prolastin-C (powder or liquid). If none are formulary, approve.	1 vear	Yes
ITITIDIOIS	Zemana	Tyoprillized powder	paper over it the patient has their two formulary alternatives from the following list, informularly (or one if one is formularly). Araiast five, Glassia, or Profastine (powder or liquid). If note are formularly, approve.	i yeai	MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Alpha-2 Agonists	Lucemyra	lofexidine tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
		clonidine ER tablet			
Alpha-adrenergic	Nexiclon XR and	and authorized			
Agonist	authorized generic	generic	Approve if the patient tried and is unable to use both clonidine immediate-release tablets AND clonidine transdermal patches.	1 year	Yes
Aluminum Chloride		aluminum chloride	Hyperhidrosis in the axillae, palms, or soles.		
Agents	Drysol	20% topical solution	Approve if the patient has tried, for at least 4 weeks, and experienced inadequate efficacy with one over-the-counter aluminum-containing product (such as Certain Dri, Bromi-lotion) [documentation required].	1 year	Yes
Alzheimer's Agent -	Diyooi	2070 topical colution	Approve if the parent has thed, for acreast 4 weeks, and experienced madequate cineary with the over-the-counter and minimum-containing product (such as octain bit, broth-routin) (accumentation required).	1 your	100
Amyloid beta-directed		donanemab-azbt	No exceptions are recommended. Due to safety concerns and the lack of clinically significant efficacy data, an exception is not recommended for Kisunla. (NOTE: It is not appropriate to use standard global criteria for this medication;		
antibody	Kisunla	intravenous infusion	Denial reason is: No exceptions are recommended. There are safety concerns and a lack of clinically significant efficacy data with use of Kisunla.)	N/A	Yes
Alzheimer's Agent -					
Amyloid beta-directed		lecanemab-irmb	No exceptions are recommended. Due to safety concerns and the lack of clinically significant efficacy data, an exception is not recommended for Legembi. (NOTE: It is not appropriate to use standard global criteria for this medication;		
antibody	Leqembi	intravenous infusion	Denial reason is: No exceptions are recommended. There are safety concerns and a lack of clinically significant efficacy data with use of Leqembi.)	N/A	Yes
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		MSB Exclusion *This criteria
Alzheimer's Disease		memantine extended-	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Agents	Namenda XR	release capsule	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
			1. Approve if the patient has tried two of the following (if formulary): 1) denegation (accept, generics), 2) galantamine immediate-release tablets (Razadyne, generics) or extended-release capsules (generics), or 3) rivastigmine		
		benzgalantamine	capsules (generic) or patches (Exelon, generics) [documentation required]. If none are formulary, approve.		
Alzheimer's Disease		delayed-release	Note: All products with the same chemical count as one alternative. For example, all galantamine products count as one alternative.		
Agents	Zunveyl	tablets	2. Approve if the patient has already been started on therapy with Zunveyl.	1 year	Yes
Amyloidoisis-					
		noticiron for	Polyneuropathy of Hereditary Transthyretin-Mediated Amyloidosis (hATTR) in a patient ≥18 years of age.  Approve if noticet has tried and according to the prescriber has experienced indexwate effects OR a circuit continuous transfer or Maintain in the prescriber has experienced indexwate effects OR a circuit continuous transfer or Maintain in the prescriber has experienced indexwate effects OR a circuit continuous transfer or Maintain in the prescriber has experienced indexwate effects or the prescriber has experienced in the prescriber has exp		
associated		patisiran for	1. Approve if patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with one of Amvuttra or Wainua, if formulary. If neither are formulary, approve.		.,
Polyneuropathy	Onnattro	intravenous use	2 Approve if the Patient has already been started on therapy with Oppattro	1 vear	
Polyneuropathy Agents	Onpattro	intravenous use	2. Approve if the Patient has already been started on therapy with Onpattro.  Polyneuropathy of Hereditary Transthyretin-Mediated Amyloidosis (hATTR) in a patient >18 years of age	1 year	Yes
Polyneuropathy Agents Amyloidoisis-	Onpattro		Polyneuropathy of Hereditary Transthyretin-Mediated Amyloidosis (hATTR) in a patient ≥18 years of age.	1 year	Yes
Polyneuropathy Agents	Onpattro	intravenous use eplontersen subcutaneous		1 year	Yes

					2025 NPF
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	Excluded Medicaiton
Analgesics - Butalbital-Containing Products	Bupap tablet	butalbital 50 mg, acetaminophen 300 mg tablet	Approve if the patient has tried one other formulary butalbital-containing product, if one is formulary (e.g., butalbital/acetaminophen capsule or tablet, butalbital/acetaminophen/caffeine capsule, butalbital/aspirin/caffeine capsule or tablet, butalbital/aspirin/caffeine/codeine capsule). If none are formulary, approve.	1 year	Yes
Products	Bupap tablet	mg tablet	acetaminophen/carierne/coderne capsule, butaibitai/aspirin/carierne capsule or tablet, butaibitai/aspirin/carierne/coderne capsule). If none are formularly, approve.	i year	res
Analgesics – Combination Products	Combogesic	acetaminophen 325 mg and ibuprofen 97.5 mg tablets	<ol> <li>Patient is directed to acetaminophen AND ibuprofen, used concurrently.</li> <li>Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use acetaminophen and ibuprofen, used concurrently.</li> </ol>	1 year	Yes
Angiotensin Converting Enzyme (ACE) Inhibitors	Epaned	enalapril maleate powder for oral solution, enalapril maleate oral solution	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Angiotensin Converting Enzyme (ACE) Inhibitors	Qbrelis	lisinopril oral solution	<ol> <li>Approve if the patient has tried lisinopril tablets (Prinivl, Zestril, generics), if formulary. If lisinopril tablets (Prinivil, Zestril, generics) are non-formulary, approve.</li> <li>Approve if the patient cannot swallow or has difficulty swallowing tablets.</li> </ol>	1 year	Yes
Angiotensin Receptor Blockers	Valsartan oral solution (previously Prexxartan)	valsartan oral solution	1. Direct the patient to valsartan tablets. 2. Approve if the patient is unable to or has difficulty swallowing oral tablets.	1 year	Yes
Angiotensin Receptor Blockers (ARBs) and Combination Products	Atacand	candesartan cilexetil tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Angiotensin Receptor Blockers (ARBs) and Combination Products	Atacand HCT	candesartan/hydrochl orothiazide tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Angiotensin Receptor Blockers (ARBs) and Combination Products	Avalide	irbesartan/hydrochlor othiazide tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Angiotensin Receptor Blockers (ARBs) and Combination Products	Avapro	irbesartan tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Angiotensin Receptor Blockers (ARBs) and Combination Products	AZOR	amlodipine besylate/olmesartan medoxomil tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Angiotensin Receptor Blockers (ARBs) and Combination Products	Benicar	olmesartan medoxomil tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Angiotensin Receptor Blockers (ARBs) and Combination Products	Benicar HCT	olmesartan/hydrochlo rothiazide tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
Angiotensin Receptor Blockers (ARBs) and Combination Products	s Cozaar	losartan tablet	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Angiotensin Receptor Blockers (ARBs) and Combination Products	s Diovan	valsartan tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Angiotensin Receptor Blockers (ARBs) and Combination Products	s Diovan HCT	valsartan/hydrochlorot hiazide tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Angiotensin Receptor Blockers (ARBs) and Combination Products	s Exforge	valsartan/amlodipine tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Angiotensin Receptor Blockers (ARBs) and Combination Products	Exforge HCT	valsartan/amlodipine/ hydrochlorothiazide tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Angiotensin Receptor Blockers (ARBs) and Combination Products	s Hyzaar	losartan/hydrochloroth	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Angiotensin Receptor Blockers (ARBs) and Combination Products	s Micardis	telmisartan tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Angiotensin Receptor Blockers (ARBs) and Combination Products	Micardis HCT	telmisartan/hydrochlor othiazide tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Angiotensin Receptor Blockers (ARBs) and Combination Products	s Tribenzor		NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Angiotensin Receptor Blockers (ARBs) and Combination Products	s Edarbyclor	azilsartan and chlorthalidone tablets	<ol> <li>Approve if the patient has tried five of the following formulary angiotensin receptor blocker/diuretic combination products, if five are formulary, or four if four are formulary, or three if three are formulary, or two are formulary, or one if only one is formulary): candesartan-hydrochlorothiazide (Atacand HCT, generics), irbesartan-hydrochlorothiazide (Avalide, generics), losartan-hydrochlorothiazide (Diovan HCT, generics), elmisartan-hydrochlorothiazide (Diovan HCT, generics).</li> <li>Approve if the patient has tried chlorthalidone AND Edarbi, if Edarbi is formulary. If Edarbi is non-formulary, approve if the patient has tried five of the following formulary angiotensin receptor blockers (ARBs), if five are formulary or thore if three are formulary, or two if only two are formulary; or one if only one is formulary): candesartan (Atacand, generics), irbesartan (Avapro, generics), olmesartan (Benicar, generics), losartan (Cozaar, generics), valsartan (Diovan, generics), telmisartan (Micardis, generics), or eprosartan. If none are formulary, approve.</li> </ol>	1 year	Yes
Angiotensin Receptor Blockers (ARBs) and Combination Products	s Edarbi	azilsartan	Approve if the patient has tried five of the following, if five are formulary (or four if four are formulary or three if three are formulary or two if two are formulary; or one if only one is formulary): candesartan (Atacand, generics), irbesartan (Avapro, generics), olmesartan (Benicar, generics), losartan (Cozaar, generics), valsartan (Diovan, generics), telmisartan (Micardis, generics), or eprosartan. If none are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.	1 year	Yes

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
Anti-arrhythmic agents	Norpace and disopyramide capsules	disopyramide phosphate capsules	<ol> <li>Approve if the patient has tried two other anti-arrhythmic agents (e.g., amiodarone, quinidine, sotalol).</li> <li>Approve if the patient has already been started on therapy with disopyramide (Norpace, generics) or Norpace CR.</li> </ol>	1 year	Yes
agenis	capsules	disopyramide	2. Approve if the patient has already been started on therapy with disopyramide (Norpace, generics) or Norpace CR.	i yeai	162
Anti-arrhythmic agents	Norpace CR	extended-release	1. Approve if the patient has tried two other anti-arrhythmic agents (e.g., amiodarone, quinidine, sotalol). 2. Approve if the patient has already been started on therapy with disopyramide (Norpace, generics) or Norpace CR.	1 year	Yes
<u> </u>	'		, and the state of		MSB Exclusion
Antibiotics (Inhaled)	ТОВІ	tobramycin solution for inhalation	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	*This criteria applies only to the NPF
randouse (milaisa)	1.02.	To minatagon	producer, wear research a significant tribungy or senses across research required.	. you.	MSB Exclusion
Antibiotics (Oral)	Doryx 50 mg, 200	doxycycline hyclate delayed-release tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	*This criteria applies only to the NPF
Artibiotics (Oral)	Firvang and	lablets	prescriber, would result in a significant anergy or serious adverse reaction tuocumentation required.	i yeai	ule NFF
	authorized generic	vancomycin oral	1. Approve if the patient has tried vancomycin capsules (Vancocin oral capsule, generics) or vancomycin oral solution (Vancocin oral solution, generics), if formulary. If neither are formulary, approve.		
Antibiotics (Oral)	solution	solution	2. If the patient is unable to swallow or has difficulty swallowing capsules, approve if the patient has tried vancomycin oral solution (Vancocin oral solution, generics), if formulary. If vancomycin oral solution is non-formulary, approve.	1 year	Yes
Antibiotics (Oral)	Minolira and authorized generic	minocycline FR tablet	Approve if the patient has tried minocycline extended-release tablets (Solodyn, generics), if formulary. If none are formulary, approve.	1 year	Yes - Authorized generic
ranabioaco (Orai)	Ximino and	minocycyline ER	reprove it the patient rate that immedyante extended to least ying generally, in formularly, approve.	1 your	gonono
	authorized generic	capsule	Approve if the patient has tried minocycline extended-release tablets (Solodyn, generics), if formulary. If none are formulary, approve	1 year	Yes
Antibiotics (Oral)	Nitrofurantoin 50 mg/5 ml suspension (brand)	ml suspension	<ol> <li>Direct to nitrofurantoin 25 mg/5 ml oral suspension.</li> <li>Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use the nitrofurantoin 25 mg/5 ml oral suspension.</li> </ol>	1 year	Yes
	Doryx DR 80 mg	doxycycline hyclate			
Antibiotics (Oral)	and authorized generic	delayed-release tablets	<ol> <li>Direct patient to other doxycycline products.</li> <li>Approve if, per the prescriber, the 80 mg tablet is required to meet the prescribed dosing requirement.</li> </ol>	1 year	Yes
Antibiotics (Oral)	generio	doxycycline hyclate	2. Approve it, per tile presented, tile do riig tablet is required to meet tile presented dosing requirement.	i you	103
		tablet, delayed-	1. Direct patient to other doxycycline products.		
Antibiotics (Oral)	Doryx MPC	release	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use a generic doxycycline product.	1 year	Yes
A 411- 1 - 41 (O1)	1.31	metronidazole oral	1. Direct the patient to metronidazole tablets.	4	
Antibiotics (Oral)	Likmez	suspension	<ol> <li>Approve if the patient is unable to swallow tablets or has difficulty swallowing tablets.</li> <li>Approve if the patient has tried linezolid tablets or oral suspension (Zyvox, generics), if formulary. If none are formulary, approve.</li> </ol>	1 year	Yes
			2. Approve if the patient is currently taking a medication that interacts with linezolid (Zyvox, generics) [e.g., monoamine oxidase inhibitors {MAOIs} or selective serotonin reuptake inhibitors {SSRIs}].		
		tedizolid phosphate	3. Approve if the patient is being treated for an organism that is resistant to linezolid (Zyvox, generics), but sensitive to Sivextro.		
Antibiotics (Oral)	Sivextro	tablets	4. Approve if the patient has been started on a course of therapy with Sivextro (to allow for completion of a course of therapy).	1 year	Yes
			1. Approve if the patient has tried one of metronidazole 250 mg tablets or metronidazole 500 mg tablets. If neither are formulary, approve.		
	metronidazole 125	metronidazole 125	<ol> <li>Approve if the prescribed dose cannot be obtained with whole tablets of the higher metronidazole strengths (i.e., 250 mg, 500 mg).</li> <li>Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use metronidazole 250mg or 500mg tablets.</li> </ol>		
Antibiotics (Oral)	mg tablets	ma tablets	4. Approve if, according to the prescriber, there is a significant crimical concern such that the patient is unable to use metronidazole 250mg of 500mg tablets.	1 year	Yes
7 1112701100 (0141)	ing tableto	mg tablete	1. Approve if the patient has tried griseofulvin ultra 125 mg or 250 mg tablets, if formulary. If neither are formulary, approve.	. you.	
		griseofulvin	2. Approve if the prescribed dose cannot be obtained with whole tablets of the griseofulvin ultramicrosize 125 mg or 250 mg strengths.		
			3. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use griseofulvin ultra 125 mg or 250 mg tablets.		
Antibiotics (Oral)	Fulvicin P/G	tablets	4. Approve if the patient has been started on a course of therapy with Fulvicin P/G (to allow for completion of a course of therapy).	1 year	Yes
			<ol> <li>Approve if the patient has tried one of dabigatran capsules, Eliquis, Savaysa, or Xarelto, if one is formulary [documentation required]. If none are formulary, approve.</li> <li>Patient is less than (&lt;) 18 years of age: approve if the patient has tried Xarelto (tablets or oral suspension) [documentation required], if formulary. If neither are formulary, approve.</li> </ol>		
		dabigatran etexilate	3. Patients currently receiving Pradaxa for treatment of thrombosis (e.g., deep vein thrombosis [DVT] or pulmonary embolism (PEI), approve.		
Anticoagulants (Oral)	Pradaxa	mesylate capsules	4. Patients currently receiving Pradaxa for prophylaxis of DVT or PE after orthopedic surgery (e.g., hip or knee replacement surgery), approve.	1 year	Yes
,			1. Regardless of the patient's age, approve if the patient is currently receiving Pradaxa (oral pellets or tablets) for treatment of thrombosis (e.g., deep vein thrombosis [DVT] or pulmonary embolism [PE]).		
			2. Patient is ≥ 8 years of age and < 12 years of age, approve if the patient meets one of the following (A or B):		
		dahinahan	A. Patient has tried dabigatran capsules (Pradaxa, generics) [documentation required], if formulary. If dabigatran capsules (Pradaxa, generics) are non-formulary, approve; OR		
Anticoagulants (Oral)	Pradaxa oral pellets	dabigatran oral	B. Patient is not able to swallow capsules, approve if the patient has tried Xarelto (tablets or oral suspension) [documentation required], if formulary. If neither are formulary, approve.  3. Patient is < 8 years of age, approve if the patient has tried Xarelto (tablets or oral suspension) [documentation required], if formulary, approve.	1 vear	Yes
	. radana orai policio	I P S / O C C	to a substitutive of your or ago, approve it the patient has the Aurente (aurene for ordinary), provided in the international formulary, approve.	. your	

					2025 NPF
				Approval	Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
			<ol> <li>Approve if the patient has tried one of the following, if one is formulary: dabigatran (Pradaxa, generics), Xarelto, or Eliquis [documentation required]. If none are formulary, approve.</li> <li>Patients currently receiving Savaysa for treatment of thrombosis (e.g., deep vein thrombosis [DVT] or pulmonary embolism (PEI), approve.</li> </ol>		
			3. Patients using Savaysa for treatment of DVT or PE associated with cancer: approve if the patient has tried Eliquis [documentation required], if formulary. If Eliquis is non-formulary, approve.		
Anticoagulants (Oral)	Savavsa	edoxaban tablets	4. Patients currently receiving Savaysa for prophylaxis of DVT or PE after orthopedic surgery (e.g., hip replacement surgery), approve.	1 vear	Yes
				. ,	MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Antidepressants -			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Other	Wellbutrin SR	bupropion HCl tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
			<u></u>		MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Antidepressants -	\\\ -	h	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the	1	applies only to
Other	Wellbutrin XL	bupropion XL tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
		bupropion hydrobromide			
Antidepressants -		extended-release	Approve if the patient has tried one product from the following list: bupropion hydrochloride extended-release tablets (Wellbutrin XL, generics), if formulary. If bupropion hydrochloride extended-release tablets (Wellbutrin XL, generics)		
Other	Aplenzin	tablets	Approve in the platent has the one product not the following list. Duplopion hydrocilloride extended release dates (Wellouthi Ac, generos), informaliary, approve.	1 vear	Yes
0 11.01	7 (510112111	bupropion	and the terminal of approximation of the terminal of the termi	. ,	1.00
		hydrochloride			
Antidepressants -	Forfivo XL and	extended-release	1. Patient is directed to bupropion hydrochloride XL 150 mg and/or 300 mg tablets (Wellbutrin XL, generics).		
Other	authorized generic	tablets	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use the buproprion hydrochloride XL 150 mg and/or 300 mg tablets (Wellbutrin XL, generics).	1 year	Yes
Antiemetic Agents -					
Substance					
P/Neurokinin-1 (NK1)			1. Approve if the patient has tried ONE of fosaprepitant dimeglumine injection (IV) [Emend IV, generics] or Focinvez IV, if formulary. If neither are formulary, approve.		
receptor antagonists	Cimumati IV	aprepitant injectable emulsion	2. In patients with hypersensitivities to polysorbate 80, approve if the patient has tried Focinvez IV, if formulary. If Focinvez IV is non-formulary, approve.	1	V
(Injectable)	Cinvanti IV	emuision	3. Approve if the patient has already started Cinvanti IV to complete all cycles in the current course of chemotherapy.	1 year	Yes
Antiomotic Agents					
Antiemetic Agents - Substance			1. Approve if the patient has tried ONE of generic fosaprepitant dimeglumine injection (IV) [Emend IV, generics] or Cinvanti IV, if formulary. If neither are formulary, approve.		
P/Neurokinin-1 (NK1)			2. Patients < 18 years of age, approve if the patient has tried fosaprepitant dimeglumine IV (Emend IV, generics), if formulary. If fosaprepitant dimeglumine IV (Emend IV, generics) are non-formulary, approve.		
receptor antagonists			3. In patients with hypersensitivities to polysorbate 80, approve if the patient has tried Cinvanti IV, if formulary. If Cinvanti IV is non-formulary, approve.		
(Injectable)	Focinvez IV		4. Approve if the patient has already started Focinvez IV to complete all cycles in the current course of chemotherapy.	1 year	Yes
Antiemetic Agents -					
Substance					MSB Exclusion
P/Neurokinin-1 (NK1)			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
receptor antagonists			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
(Injectable)	Emend IV	dimeglumine injection	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Antiemetic Agents -			Approve if the patient meets ONE of the following (1 or 2):		
Combination			1. Patient meets BOTH of the following (A and B):		
Substance P/NK1 receptor antagonist			A. Patient has tried two formulary oral or transdermal serotonin 5-HT3 receptor antagonists from the following list (if two are formulary or one if one is formulary): ondansetron oral (generics), granisetron oral (generics), or Sancuso;		
and serotonin (5-HT3)			B. Patient has tried one oral formulary Substance P/NK1 antagonists from the following list: aprepitant capsules (Emend, generics) or Varubi tablets, if one is formulary; OR		
receptor antagonist.		netunitant/nalonsetron	Note: If there are no formulary 5-HT3 receptor antagonists, approve. If there are no Substance P/NK1 antagonists, approve.		
(Oral)	Akynzeo capsules	capsules	2. Approve if the patient has already started Akynzeo to complete all cycles in the current course of chemotherapy.	1 year	Yes
Antiemetics -	,				
Serotonin (5-HT3)					
Receptor Antagonists	ondansetron ODT	ondansetron ODT 16			
(Oral)	16 mg (brand)	mg	Approve if the patient has tried ondansetron ODT 4 mg or ondansetron ODT 8 mg AND is unable to continue to use these products. If both ondansetron ODT 4 mg and ondansetron ODT 8 mg are non-formulary, approve.	1 year	Yes
			1. Approve if the patient has tried, and according to the prescriber, experienced inadequate efficacy OR significant intolerance with BOTH of the following: 1) granisetron tablets (generics) and 2) ondansetron tablets (generics), if		
Antiemetics -			formulary (or only one if one is formulary). If neither are formulary, approve.		
Serotonin Receptor			2. Patient < 18 years of age, approve if the patient tried, and according to the prescriber, experienced inadequate efficacy OR significant intolerance with ondansetron tablets (generics), if formulary. If ondansetron tablets (generics) are		
Antagonists (Oral and		4.1	non-formulary, approve.	4	V
Inejctable)	Anzemet tablets	dolasetron tablets	3. Approve if the patient has already started Anzemet to complete all cycles in the current course of chemotherapy.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Antiemetics and Antivertigo Agents	Emend capsules and Emend Trifold Pack	aprepitant oral capsules	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antiemetics and Antivertigo Agents	Emend oral suspension	aprepitant oral suspension	<ol> <li>Approve if the patient has tried one formulary alternative from the following list: aprepitant capsules (Emend, generics) or Varubi tablets. If none are formulary, approve.</li> <li>Patients ≥ 12 and &lt;18 years of age: approve if the patient has tried aprepitant capsules (Emend, generics), if formulary. If aprepitant capsules (Emend, generics) are non-formulary, approve.</li> <li>Patients &lt; 12 years of age: approve.</li> <li>Patients who cannot swallow or have difficulty swallowing capsules, approve.</li> <li>Approve if the patient has already started Emend oral suspension to complete all cycles in the current course of chemotherapy.</li> </ol>	1 year	Yes
Antiemetics and Antivertigo Agents		doxylamine succinate and pyridoxine hydrochloride extended-release tablets	Approve if the patient has tried, and according to the prescriber, has experienced inadequate efficacy OR significant intolerance with doxylamine-pyridoxine (Diclegis, generics), if formulary. If doxylamine-pyridoxine (Diclegis, generics) are non-formulary, approve if the patient has tried doxylamine AND pyridoxine (Vitamin B6).	1 year	Yes
Antifungals (Oral)		posaconazole delayed-release tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antifungals (Oral)	Tolsura	itraconazole capsules	<ol> <li>Approve if the patient has tried one of itraconazole capsules (Sporanox, generics) or itraconazole oral solution (Sporanox liquid, generics).</li> <li>NOTE: A trial of either the conventional intraconazole capsules or intraconazole solution would count toward meeting criteria regardless of the formulary status of the product.</li> <li>Patient has been started on a current course of therapy with Tolsura (for a non-oncychomycosis diagnosis): approve to complete the current course.</li> <li>Deny: If the patient is requesting Tolsura for a diagnosis of onychomycosis, the request should be denied regardless of what the patient has tried for the current condition or if the patient has already been started on the product.</li> </ol>	1 year	Yes
Antifungals (Topical)		oxiconazole nitrate cream	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antifungals (Topical)	Ertaczo	sertaconazole nitrate 2% cream	<ol> <li>Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with four topical antifungals.</li> <li>If the patient has an occluded fungal infection, such as intertrigo or erosio interdigitalis blastomycetica, approve if the patient has tried and, according to the prescriber has experienced inadequate efficacy OR a significant intolerance with two topical antifungals.</li> <li>Note: Examples of topical antifungals include: naftifine 1% or 2% cream or 1% gel (Naftin, generics), Naftin 2% gel, Lotrimin Ultra (over-the-counter [OTC]), clotrimazole 1% cream (OTC formulation), econazole 1% cream, ketoconazole 2% cream or foam (Extina, generics), Oxistat 1% lotion, oxiconazole 1% cream (Oxistat, generics), Ecoza foam, Exelderm 1% cream or solution, ciclopirox 0.77% cream or gel (generics), Mentax 1% cream, Xolegel 2% gel.</li> </ol>	1 year	Yes
Antifungals (Topical)		oxiconazole nitrate	1. Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with four topical antifungals.  2. If the patient has an occluded fungal infection, such as intertrigo or erosio interdigitalis blastomycetica, approve if the patient has tried and, according to the prescriber has experienced inadequate efficacy OR a significant intolerance with two topical antifungals.  Note: Examples of topical antifungals include: naftifine 1% or 2% cream or 1% gel (Naftin, generics), Naftin 2% gel, Lotrimin Ultra (over-the-counter [OTC]), clotrimazole 1% cream (OTC formulation), econazole 1% cream, ketoconazole 2% cream or foam (Extina, generics), oxiconazole 1% cream (Oxistat, generics), Ertaczo 2% cream, Exelderm 1% cream or solution, ciclopirox 0.77% cream or gel (generics), Mentax 1% cream, Xolegel 2% gel.	1 year	Yes
Antifungals (Topical)	Ecoza foam	econazole nitrate topical foam	1. Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with four topical antifungals.  2. If the patient has an occluded fungal infection, such as intertrigo or erosio interdigitalis blastomycetica, approve if the patient has tried and, according to the prescriber has experienced inadequate efficacy OR a significant intolerance with two topical antifungals.  Note: Examples of topical antifungals include: naftifine 1% or 2% cream or 1% gel (Naftin, generics), Naftin 2% gel, Lotrimin Ultra (over-the-counter [OTC]), clotrimazole 1% cream (OTC formulation), econazole 1% cream, ketoconazole 2% cream or foam (Extina, generics), Oxistat 1% lotion, oxiconazole 1% cream (Oxistat, generics), Ertaczo 2% cream, Exelderm 1% cream or solution, ciclopirox 0.77% cream or gel (generics), Mentax 1% cream, Xolegel 2% gel.	1 year	Yes
Antifungals (Topical)	(sulconazole nitrate	sulconazole nitrate 1% (cream and solution)	1. Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with four topical antifungals. 2. If the patient has an occluded fungal infection, such as intertrigo or erosio interdigitalis blastomycetica, approve if the patient has tried and, according to the prescriber has experienced inadequate efficacy OR a significant intolerance with two topical antifungals.  Note: Example of topical antifungals include: naftifine 1% or 2% cream or 1% gel (Naftin, generics), Naftin 2% gel, Lotrimin Ultra (over-the-counter [OTC]), clotrimazole 1% cream (OTC formulation), econazole 1% cream, Ecoza 1% foam, ketoconazole 2% cream or foam (Extina, generics), Oxistat 1% lotion, oxiconazole 1% cream (Oxistat, generics), Ertaczo 2% cream, ciclopirox 0.77% cream or gel (generics), Luzu 1% cream, Mentax 1% cream, Xolegel 2% gel.	1 year	Yes - Authorized generic only

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
	Luzu and authorized generic (Iuliconazole	Iuliconazole 1%	1. Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with four topical antifungals.  2. If the patient has an occluded fungal infection, such as intertrigo or erosio interdigitalis blastomycetica, approve if the patient has tried and, according to the prescriber has experienced inadequate efficacy OR a significant intolerance with two topical antifungals.  Note: Examples of topical antifungals include: naftifine 1% or 2% cream or 1% gel (Naftin, generics), Naftin 2% gel, Lotrimin Ultra (over-the-counter [OTC]), clotrimazole 1% cream (OTC formulation), econazole 1% cream, Ecoza 1% foam, ketoconazole 2% cream or foam (Extina, generics), Oxistat 1% lotion, oxiconazole 1% cream (Oxistat, generics), Ertaczo 2% cream, Exelderm 1% cream or solution, ciclopirox 0.77% cream or gel (generics), Mentax 1% cream,		
Antifungals (Topical)	1% cream)	cream	Xolegel 2% gel.	1 year	Yes
Antifungals (Topical)	Xolegel	ketoconazole 2% gel	1. Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with four topical antifungals.  2. If the patient has an occluded fungal infection, such as intertrigo or erosio interdigitalis blastomycetica, approve if the patient has tried and, according to the prescriber has experienced inadequate efficacy OR a significant intolerance with two topical antifungals.  Note: naftifine 1% or 2% cream or 1% gel (Naftin, generics), Naftin 2% gel, Lotrimin Ultra (over-the-counter [OTC]), clotrimazole 1% cream (OTC formulation), econazole 1% cream, Ecoza 1% foam, ketoconazole 2% cream or foam (Extina, generics), Oxistat 1% lotion, oxiconazole 1% cream (Oxistat, generics), Etaczo 2% cream, Exelderm 1% cream or solution, ciclopirox 0.77% cream or gel (generics), Luzu 1% cream, Mentax 1% cream.	1 year	Yes
3 ( 1 /	3	J			
Antihistamines (Oral)	clemastine syrup	clemastine syrup	<ol> <li>Approve if the patient has tried five oral antihistamines.</li> <li>Note: Examples of oral antihistamine include, but are not limited to: diphenhydramine, chlorpheniramine, carbinoxamine, hydroxyzine, cetirizine, levocetirizine, desloratadine.</li> <li>If the patient is unable to swallow or has difficulty swallowing tablets, approve if the patient has tried at least two of the following: carbinoxamine syrup, diphenhydramine solution, or hydroxyzine solution or syrup.</li> </ol>	1 year	Yes
	clemastine oral				
Antihistamines (Oral)	tablet, Clemasz tablet	clemastine oral tablet	Approve if the patient has tried five oral antihistamines.  Note: Examples of oral antihistamine include, but are not limited to: diphenhydramine, chlorpheniramine, carbinoxamine, hydroxyzine, cetirizine, levocetirizine, desloratadine.	1 year	Yes
Antihistamines (oral) - First-generation	carbinoxamine maleate 4 mg/5 ml oral suspension (brand) [authorized generic of Karbinal ER] and Karbinal ER	carbinoxamine maleate 4 mg/5 ml	1. Approve if the patient has tried five oral antihistamines (e.g., clemastine, diphenhydramine, chlorpheniramine, carbinoxamine [generic], hydroxyzine, cetirizine, loratadine).  Note: OTC products would count toward meeting the requirement.  2. If the patient is unable to swallow or has difficulty swallowing solid dosage forms, approve if the patient has tried at least two oral liquid antihistamines (e.g., carbinoxamine solution [generic], diphenhydramine solution, hydroxyzine solution or syrup, clemastine syrup, cetirizine solution, or loratadine solution or syrup).  Note: OTC products would count toward meeting the requirement.	1 year	Yes
i iist-generation	Lity and Naibinal Lit	orar suspension	Note: OTO products would count toward meeting the requirement.	i yeai	MSB Exclusion
Antimuscarinic Agents	Transderm-Scop	scopolamine patches	<b>NOTE:</b> A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product. <b>Criteria:</b> Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	*This criteria applies only to the NPF
Antiparkinson Drugs	Gocovri ER	amantadine extended- release capsules	Approve if the patient has tried one product from the following list: amantadine capsules, amantadine tablets, or amantadine oral solution AND meets one of the following (A or B):  A. Patient derived benefit from immediate-release amantadine, but had intolerable adverse events, as determined by the prescriber [documentation required]; OR  B. Patient could not achieve a high enough dosage to gain adequate benefit, as determined by the prescriber [documentation required].	1 year	Yes
7 mapanancen Brage	Good III Z. K	release superiles	Approve if the patient has tried one product from the following list: amantadine capsules, amantadine tablets, or amantadine oral solution AND meets one of the following (A or B):	. you.	
Antiparkinson Drugs	O-malay ED	amantadine extended- release tablets	A. Patient derived benefit from immediate-release amantadine, but had intolerable adverse events, as determined by the prescriber [documentation required]; OR  B. Patient could not achieve a high enough dosage to gain adequate benefit, as determined by the prescriber [documentation required].	1 vear	Yes
Antiparkinson brugs	Ositiolex ER	release tablets	B. Patient could not achieve a riigh enough dosage to gain adequate benefit, as determined by the prescriber [documentation required].	i yeai	res
Antiparkinson Drugs - Carbidopa and/or Levodopa Agents	Dhivy	carbidopa and levodopa immediate- release tablets	Approve if dose prescribed cannot be obtained withcarbidopa-levodopa tablets (Sinemet, generics) or half-tablets.  Note: Dhivy can be split into a ¼ of a tablet (i.e., 6.25 mg of carbidopa and 25 mg of levodopa).	N/A	Yes
Antiparkinson Drugs – Carbidopa and/or	July	foslevodopa- foscarbidopa	1. Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with Onapgo, if formulary. If Onapgo is non formulary, approve if the patient meets ONE of the following (A or B):  A. Patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with one of the following: Crexont capsules, or carbidopa-levodopa extended-release tablets, if formulary. If none are formulary, approve; OR  B. Patient is unable to swallow oral dosage forms or has difficulty swallowing oral dosage forms.		.55
Levodopa Agents	Vyalev	subcutaneous infusion	2. Approve if the patient has already been started on therapy with Vyalev.	1 year	Yes
Antiparkinson Drugs - Inhibitor of Monoamine Oxidase	V. J.		1. Approve if the patient has tried two products from the following list, if formulary (or one if one is formulary): selegiline tablets/capsules, rasagiline tablets (Aziliect, generics), or Zelapar. If none are formulary, approve.		V
Type B Inhibitors	Xadago	safinamide tablets	2. Patients already started on Xadago, approve.	1 year	Yes

				Annuaral	2025 NPF
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	Excluded Medicaiton
Antiparkinson Drugs - Inhibitor of Monoamine Oxidase Type B Inhibitors	Zelapar	selegiline orally	<ol> <li>Approve if the patient has tried one product from the following list, if formulary (or one if one is formulary): selegiline tablets/capsules, rasagiline tablets (Azilect, generics), or Xadago. If none are formulary, approve.</li> <li>Approve if the patient cannot swallow or has difficulty swallowing selegiline tablets.</li> </ol>	1 year	Yes
Antiparkinson Drugs – Apomorphine products	Apokyn	apomorphine injection	See standard <i>Parkinsons Disease Apomorphine Subcutaneous Prior Authorization Policy</i> criteria	See PA duration	Yes
Antiparkinson Drugs – Apomorphine products	Onapgo	apomorphine subcutaneous infusion	<ol> <li>Approve if the patient has tried, and according to the prescriber, has experienced inadequate efficacy OR significant intolerance with Vyalev, if formulary. If Vyalev is non-formulary, approve if the patient meets ONE of the following (A or B):         <ul> <li>A. Patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with one of the following: Crexont capsules, Rytary capsules, or carbidopa-levodopa extended-release tablets, if formulary. If none are formulary, approve; OR</li> <li>B. Patient is unable to swallow oral dosage forms or has difficulty swallowing oral dosage forms.</li> </ul> </li> <li>Approve if the patient has already been started on therapy with Onapgo.</li> </ol>	1 year	Yes
Antiplatelet Agents	Plavix	clopidogrel bisulfaste tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antiprotozoals (Oral)	Alinia tablets	nitazoxanide tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antipsychotics (Long- Acting Injectables) – Risperidone or Paliperidone Based	Invega Hafyera		<ol> <li>Approve if the patient has been established on therapy with Invega Sustenna for ≥ 4 months OR Invega Trinza for ≥ one 3-month cycle AND the prescriber attests the patient requires an extended dosing interval due to a demonstrated significant concern for non-adherence with a 4-week or 3-month dosing interval.</li> <li>NOTE: Invega Sustenna/Invega Trinza Formulary Exception Criteria will apply.</li> <li>Approve if the patient has already been started on therapy with Invega Hafyera.</li> </ol>	1 year	Yes
Antipsychotics (Oral)	Abilify	aripiprazole tablets and oral solution	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antipsychotics (Oral)	Saphris	asenapine sublingual tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antipsychotics (Oral)	Seroquel	quetiapine fumarate tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antipsychotics (Oral)	Seroquel XR	quetiapine fumarate extended-release tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antipsychotics (Oral)	Latuda	lurasidone tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
		· ·	<ol> <li>Approve if the patient has tried two oral antipsychotics (e.g., risperidone tablets/orally disintegrating tablets [ODT]{Risperdal, generics}, olanzapine tablets/ODT [Zyprexa/Zydis, generics], quetiapine tablets [Seroquel, generics], quetiapine extended-release tablets [Seroquel XR, generics], aripiprazole tablets [Abilify, generics], paliperidone ER tablets [Invega, generics], ziprasidone capsules [Geodon, generics], Latuda tablets, Rexulti tablets, Vraylar capsules, asenapine sublingual tablets [Saphris, generics], Caplyta).</li> <li>Approve if the patient is currently taking Fanapt.</li> </ol>		
Antipsychotics (Oral)	Fanapt	and titration pack	3. Approve if the patient has taken Fanapt at any time in the past.	1 year	Yes
Antipsychotics (Oral)	Opipza	aripiprazole oral film	Approve if the patient has tried and cannot take ONE of aripiprazole ODT or aripiprazole oral solution. If neither are formulary, approve.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Antipsychotics (Oral)	Quetiapine 150 mg tablets	quetiapine 150 mg tablet	1. Direct to quetiapine 50 mg and/or quetiapine 100 mg tablets. 2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use the quetiapine 50 mg and/or 100 mg tablet.	1 year	Yes
Antipsychotics (Oral) [Muscarinic Agonist and Muscarinic Antagonist]	Cobenfy	xanomeline and	1. Approve if the patient has tried two other novel (atypical) antipsychotics.  Note: Examples of novel (atypical) antipsychotics include risperidone tablets/orally disintegrating tablets (ODT) [Risperdal, generics], Fanapt tablets, olanzapine tablets/ODT (Zyprexa/Zydis, generics), quetiapine tablets (Seroquel, generics), quetiapine extended-release tablets (Seroquel XR, generics), aripiprazole tablets (Abilify, generics), paliperidone ER tablets (Invega, generics), ziprasidone capsules (Geodon, generics), Latuda tablets, Rexulti tablets, asenapine sublingual tablets (Saphris, generics), Caplyta.  2. Approve if the patient has already started therapy with Cobenfy.  3. Approve if the patient has taken Cobenfy at any time in the past.	1 year	Yes
Antiseizure Medications	Banzel	rufinamide tablets and oral suspension	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antiseizure Medications	Керрга	levetiracetam tablets and solution	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antiseizure Medications	Keppra XR	levetiracetam exteended-release tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antiseizure Medications	Lamictal	lamotrigine tablets and chewable tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antiseizure Medications	Lamictal ODT	lamotrigine oral disintegrating tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antiseizure Medications	Lamictal XR	lamotrigine extended- release tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antiseizure Medications	Onfi	clobazam tablets and suspension	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antiseizure Medications	Sabril	vigabatrin tablets and powder packet	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antiseizure Medications	Торатах	topiramate tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antiseizure Medications	Trileptal	oxcarbazepine tablets and suspension	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antiseizure Medications	Vimpat	lacosamide tablets and oral solution and vials	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antiseizure Medications	Zonegran	zonisamide capsule	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
	Primidone 125 mg		Approve if the patient's prescribed dose cannot be obtained with primidone 50 mg or 250 mg tablets.	4	V
Medications	(brand)		Note: The patient is NOT required to split the 250 mg tablets in half.	1 year	Yes
Antiseizure Medications	Vigafyde	vigabatrin oral solution	Approve if the patient tried and cannot take vigabatrin granules for oral solution (Sabril powder for solution, generics), if formulary. If vigabatrin granules for oral solution (Sabril powder for solution, generics) is non-formulary, approve.	1 year	Yes
Antiseizure Medications	Motpoly XR		Approve if the patient is unable to use lacosamide immediate-release tablets (Vimpat tablets, generics), if formulary. If lacosamide immediate-release tablets (Vimpat tablets, generics) are non-formulary, approve.	1 year	Yes
Antiseizure	Zonisade oral	zonisamide oral			
Medications	suspension	suspension	Approve if the patient is unable to swallow or has difficulty swallowing zonisamide capsules. If zonisamide capsules are non-formulary, approve.	1 year	Yes
Antiseizure	topiramate 50 mg	topiramate 50 mg	1. Approve if the patient has tried topiramate 25 mg sprinkle capsules. If topiramate 25 mg sprinkle capsules are non-formulary, approve.		
Medications	sprinkle capsule	sprinkle capsule	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use topiramate 25mg sprinkle capsules.	1 year	Yes
Antiseizure		fenfluramine oral		See PA	
Medications	Fintepla	solution	See standard Antiepileptics - Fintepla Prior Authorization Policy criteria.	duration	Yes
Antiseizure Medications	Eprontia	topiramate oral solution	Approve if the patient has tried and cannot take topiramate sprinkle capsules (Topamax Sprinkle capsules, generics). If topiramate sprinkle capsules (Topamax Sprinkle capsules, generics) are non-formulary, approve.	1 year	Yes
Antiseizure Medications - Buccal	Libervant		1. Approve if the patient has tried diazepam rectal gel (Diastat, generics), if formulary. If diazepam rectal gel (Diastat, generics) is non-formulary, approve.  Note: If the patient has tried a benzodiazepine nasal spray (e.g., Valtoco or Nayzilam), this would satisfy the requirement for approval.  2. If the patient's caregiver is unable to administer diazepam rectal gel (Diastat, generics), approve.	1 year	Yes
Antivirals (Oral)	Valtrex	valacyclovir HCl	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antivirals (Topical)	Zovirax ointment		NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Antivirals (Topical)	Xerese		Approve if the patient has tried two of the following: acyclovir capsules/tablets, famciclovir tablets (generics), valacyclovir tablets (Valtrex, generics), acyclovir 5% cream (Zovirax 5% cream, generics), penciclovir 1% cream (Denavir, generics), Sitavig tablets, or over-the-counter (OTC) Abreva 10% cream.	1 year	Yes

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  OR		
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  1. For brand Arimidex requests, approve one of the following (A or B):  A) The patient meets both of the following (i and ii):  i. The requested brand non-formulary drug is being prescribed for the primary prevention of breast cancer for a post-menopausal patient aged 35 years or greater who is at increased risk of breast cancer and at low risk for adverse medication effects and who does NOT have a current or previous diagnosis of breast cancer or ductal carcinoma in situ (DCIS); AND		
			ii. The patient meets one of the following (a <u>or</u> b):  a. According to the prescriber, the brand product is being requested because the preferred bioequivalent generic product would not be as medically appropriate for the patient as the requested brand non-formulary drug; OR  b. According to the prescriber, other formulary alternatives would not be as medically appropriate for the patient as the requested non-formulary drug.*  B) The patient meets both of the following (i <u>and</u> ii):  i. The requested brand non-formulary drug is being prescribed for a use OTHER THAN the primary prevention of breast cancer for a post-menopausal patient aged 35 years or greater who is at increased risk of breast cancer and at low risk for adverse medication effects and who does NOT have a current or previous diagnosis of breast cancer or ductal carcinoma in situ (DCIS); AND		
			ii. The Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].		
			2. For generic anastrozole requests,** approve if the requested non-formulary drug is being prescribed for the primary prevention of breast cancer for a post-menopausal patient aged 35 years or greater who is at increased risk of breast cancer and at low risk for adverse medication effects and who does NOT have a current or previous diagnosis of breast cancer or ductal carcinoma in situ (DCIS) AND, according to the prescriber, other formulary alternatives would not be as medically appropriate for the patient as the requested non-formulary drug.  *Applicable for clients who are not using Multi-Source Brand criteria.		MSB Exclusion *This criteria applies only to
Aromatase inhibitor	Arimidex	anastrozole tablets	**Note: When compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required, these products would be reviewed under the Standard Commercial Default Criteria.	1 year	the NPF
Benign Prostatic		finasteride 5 mg and tadalafil 5 mg	Benign Prostatic Hyperplasia (BPH).		
Hyperplasia – Combination Agents	Entadfi	capsules	Approve if, according to the prescriber, the patient has a clinical reason they cannot take finasteride 5 mg and tadalafil 5 mg as separate agents.	1 year	Yes
Benign Prostatic Hyperplasia (Alpha			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.	,	MSB Exclusion *This criteria
Blockers and 5-Alpha Reductase Inhibitors)	Avodart	dutasteride capsules	<b>Criteria:</b> Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	applies only to the NPF
Benign Prostatic Hyperplasia (Alpha	rttodart	dutational suppures	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.	r your	MSB Exclusion *This criteria
Blockers and 5-Alpha			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Reductase Inhibitors)	Rapaflo	silosodin capsules	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Benign Prostatic Hyperplasia (Alpha			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		MSB Exclusion *This criteria
Blockers and 5-Alpha			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Reductase Inhibitors)	Uroxatral	alfuzosin tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		MSB Exclusion *This criteria applies only to
Benzodiazepines	Klonopin	clonazepam tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		MSB Exclusion *This criteria applies only to
Benzodiazepines	Valium	diazepam tablets	prescriber, would result in a significant allergy or serious adverse reaction (documentation required).	1 year	the NPF

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
Benzodiazepines	Xanax	alprazolam tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Benzodiazepines	Xanax XR	alprazolam entended- release tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Benzodiazepines	Loreev XR	lorazepam extended- release capsules	<ol> <li>Direct the patient to use lorazepam tablets.</li> <li>Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use lorazepam immediate-release tablets.</li> </ol>	1 year	Yes
Benzodiazepines	Doral and authorized generic	guazepam tablets	Approve if the patient has tried estazolam or lorazepam, if formulary. If neither are formulary, approve.	1 year	Yes
Benzodiazepines and Combination Products		chlordiazepoxide/clidi nium bromide capsules			Yes *This criteria applies only to the NPF
Beta-Blocker Products	Bystolic	nebivolol tablets	Approve if the patient has tried clidinium-chlordiazepoxide capsules. If clidinium-chlordiazepoxide capsules are non-formulary, approve.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Beta-Blocker	labetalol 400 mg	labetalol 400 mg	1. Direct the patient to labetalol 100 mg, 200 mg, or 300 mg tablets.		
Products  Beta-Blocker and Beta-Blocker Combination Products	tablets	propranolol HCl	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use the labetalol 100 mg, 200 mg, or 300 mg tablets.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	Yes MSB Exclusion *This criteria applies only to the NPF
Beta-Blocker and Beta-Blocker Combination Products		metoprolol succinate extended-release tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Beta-Blocker and Beta-Blocker Combination Products	Kapspargo Sprinkle	metoprolol succinate extended-release capsules	<ol> <li>Approve if the patient has tried metoprolol succinate extended-release tablets, if formulary. If non-formulary, approve.</li> <li>If the patient requires a dosage form which can be opened and sprinkled for alternative administration (e.g., for patients unable to swallow capsules, for nasogastric tube administration), approve.</li> </ol>	1 year	Yes
Beta-Blocker and Beta-Blocker Combination Products	Inderal XL	propranolol hydrochloride capsule, extended release	<ol> <li>Direct the patient to propranolol extended-release capsules.</li> <li>Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use generic propranolol extended-release capsules.</li> </ol>	1 year	Yes
Beta-Blocker and Beta-Blocker		propranolol hydrochloride capsule, extended	1. Direct the patient to propranolol extended-release capsules.		
Combination Products  Beta-Blockers	bisoprolol fumarate 2.5 mg tablets	bisoprolol fumarate 2.5 mg tablets	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use generic propranolol extended-release capsules.  1. Approve if the prescribed dose cannot be obtained with whole bisoprolol 5 mg or 10 mg tablets.  Note: The patient is NOT required to split the 5 mg or 10 mg tablets in half.  2. Patient with bronchospastic disease, approve.	1 year	Yes
Bone Modifiers - Other	Evenity	romosozumab-aqqg	<ol> <li>Approve if patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with one of the following products: an oral bisphosphonate (e.g., alendronate [Fosamax, Fosamax Plus D, generics], ibandronate tablets [Boniva, generics], alendronate oral solution, Binosto, risedronate [Actonel, Atelvia, generics], a teriparatide product (i.e., Forteo, teriparatide), Tymlos, or Prolia.</li> <li>Patient has already tried ibandronate injection (Boniva IV) or zoledronic acid injection (Reclast): approve.</li> <li>According to the prescriber, patient has severe renal impairment or chronic kidney disease: approve.</li> <li>Note: An example of severe renal impairment is a creatinine clearance &lt; 35 mL/min).</li> <li>Patients who have had an osteoporotic fracture or a fragility fracture: approve.</li> <li>Patients who cannot swallow or has difficulty swallowing tablets, cannot remain in an upright position (post oral bisphosphonate administration), or has a pre-existing gastrointestinal medical condition: approve.</li> <li>Note: Examples of pre-existing gastrointestinal medical conditions include esophageal lesions, esophageal ulcers, or abnormalities of the esophagus that delay esophageal emptying [stricture, achalasia].</li> </ol>	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Bone Modifiers - Other	Forteo		Approve if the patient has tried generic teriparatide (generic Forteo), if formulary.  If generic teriparatide (generic Forteo) is non-formulary or if generic teriparatide (generic Forteo) is being requested, approve if the patient meets one of the following (1, 2, or 3):  1. Approve if the patient has tried brand teriparatide, if formulary. If brand teriparatide is non-formulary, approve if patient has tried Tymlos, if formulary. If Tymlos is non-formulary, approve.  2. Approve if the patient has tried brand teriparatide, if formulary. If brand teriparatide is non-formulary, patients with glucocorticoid-induced osteoporosis (GIO): approve.  Note: For approvals above under criteria (1 and 2): Use of teriparatide (Forteo [generics] or teriparatide) exceeding 2 years during a patient's lifetime, approve if the patient is at high risk for fracture as determined by the prescriber.  3. Approve if the patient has tried brand teriparatide, if formulary. If brand teriparatide is non-formulary, approve if the patient has a diagnosis of chronic hypoparathyroidism	1 year	Yes - Forteo brand
Bone Modifiers - Other	Prolia	denosumab injection	If either Jubbonti or Stobodo are formulary, approve if the patient meets BOTH of the following (1 and 2):  1. Patient has tried BOTH Jubbonti and Stobodo, if formulary (or one if one is formulary); AND  2. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant alterity or serious adverse reaction.  If neither Jobbonti nor Stobodo are formulary, approve if the patient meets ONE of the following (1, 2, 3, 4, 5, 6, 7, 8, or 9):  1. Approve if patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with one of the following products: an oral bisphosphonate (e.g., alendronate [Fosamax, Fosamax Plus D, generics], binardronate tablets [Boniva, generics], alendronate rail solution, Binosot, risedronate [Actone], Altelya, generics, a teriparatide product (i.e., Forteo, teriparatide), Tymlos, or Evenity.  2. Patient has already tried ibandronate injection (Boniva IV) or zoledronic acid injection (Reclast): approve.  3. According to the prescriber, the patient has severe renal impairment or chronic kidney disease: approve.  Note: An example of severe renal impairment is a creatinine clearance < 35 mL/minute.  4. Patients who cannot swallow or has difficulty swallowing tablets, cannot remain in an upright position post oral bisphosphonate administration, or has a pre-existing gastrointestinal medical conditions include esophageal lesions; esophageal ulcers, or abnormalities of the esophagus that cleay esophageal emptying (stricture, achalasia).  5. Treatment of bone loss (to increase bone mass) in patients with nonmetastatic prostate cancer at high risk for fracture who are receiving androgen deprivation therapy are. Lupron Depot [leurprolide or depot suspension], Eligard [leurprolide acetate for injectable suspension], Treistar (triptorelin pamoate	1 уеаг	Yes
Botulinum Toxin Products	Daxxify		<ol> <li>Patient has a sensitivity or allergy to cow's milk protein, approve if the patient has tried one of Botox or Xeomin, if formulary. If neither are formulary, approve.</li> <li>Approve if the patient has already been started on therapy with Daxxify.</li> </ol>	1 year	Yes

Approval	2025 NPF Excluded
Duration	Medicaiton
1 1	
1	
1	
1	
1	
1 1	
1	
1	
1	
1	
1 1	
1	
1	
1	
1 1	
1 1	
1 1	
1 1	
1 1 1000	Yes
1 ye	ear

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
morapy crace		Constitution Number	Hyperhidrosis, Primary Axillary, in a patient ≥ 18 years of age. Approve if the patient has tried at least one prescription topical agent for axillary hyperhidrosis.  Note: Examples of prescription topical agents for the treatment of axillary hyperhidrosis include Drysol (aluminum chloride 20% topical solution), Qbrexza (glycopyrronium cloth 2.4% for topical use), Sofdra (glycopyrronium 12.45% topical gel).  Hyperhidrosis, Primary Palmar/Plantar and Facial, in a patient ≥ 18 years of age.  Approve if the patient has tried at least one topical agent for the treatment of hyperhidrosis (e.g., aluminum chloride).  Migraine Headache Prevention in a patient ≥ 18 years of age.  1. Approve if the patient has tried one of Aimovig, Ajovy, Emgality, Vyepti, or Qulipta [documentation required], if formulary. If none are formulary, approve.  2. Approve if the patient has already been started on therapy with Botox.  Blepharospasm in a patient ≥ 12 years of age.  Note: This includes blepharospasm associated with dystonia, benign essential blepharospasm, seventh (VII) nerve disorders.  1. Approve if the patient has tried Xeomin, if formulary. If Xeomin is non-formulary, approve.  2. If the patient is < 18 years of age, approve.		
			3. Approve if the patient has already been started on therapy with Botox.		
			Strabismus in a patient ≥ 12 years of age: Approve.  Note: Common types of strabismus include esotropia, exotropia, hypotropia.		
			Cervical Dystonia in a patient ≥ 18 years of age.  Note: Cervical dystonia is also referred to as spasmodic torticollis.		
			1. Approve if the patient has tried one of Dysport, Xeomin, or Daxxify, if formulary. If none are formulary, approve.		
			2. Patient has a sensitivity or allergy to cow's milk protein, approve if the patient has tried one of Xeomin or Daxxify, if formulary. If neither are formulary, approve.		
			3. Approve if the patient has already been started on therapy with Botox.		
			Spasticity, Limb(s), in a patient ≥ 2 years of age.  1. Approve if the patient has tried one of Dysport or Xeomin, if formulary. If neither are formulary, approve.  2. Patients with lower limb spasticity, approve if the patient has tried Dysport. If Dysport is non-formulary, approve.  a. Patient has a sensitivity or allergy to cow's milk protein, approve.		
Botulinum Toxin	Botox (NOT	onabotulinumtoxinA	3. Patient has a sensitivity or allergy to cow's milk protein, approve.		
	cosmetic) (1 of 2)	for injection	4. Approve if the patient has already been started on therapy with Botox.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Botulinum Toxin Products - Botox	Botox (NOT cosmetic) [2 of 2]		Sialorrhea_Chronic, in a patient ≥ 18 years of age.  1. Approve if the patient has tried one of Dysport, Xeomin, or Myobloc, if formulary. If none are formulary, approve.  2. Patient has a sensitivity or allergy to cow's milk protein, approve if the patient has tried one of Xeomin or Myobloc, if formulary. If neither are formulary, approve.  3. Approve if the patient has already been started on therapy with Botox.  Anal Fissure, Chronic, in a patient ≥ 18 years of age.  1. Approve if the patient has tried Dysport, if formulary. If Dysport is non-formulary, approve.  2. Patient has a sensitivity or allergy to cow's milk protein, approve.  3. Approve if the patient has already been started on therapy with Botox.  Hemifacial Spasm in a patient ≥ 18 years of age.  1. Approve if the patient has already been started on therapy with Botox.  Commandibular Dystonia in a patient ≥ 18 years of age.  1. Approve if the patient has already been started on therapy with Botox.  Commandibular Dystonia in a patient ≥ 18 years of age.  1. Approve if the patient has already been started on therapy with Botox.  Commandibular Dystonia in a patient ≥ 18 years of age.  1. Approve if the patient has already been started on therapy with Botox.  Commandibular Dystonia in a patient ≥ 18 years of age.  1. Approve if the patient has already been started on therapy with Botox.  Neurogenic Detrusor Overactivity in patient ≥ 18 years of age; Curinary Incontinence Due to Detrusor Overactivity Neuropenic Detrusor Overactivity in patient ≥ 18 years of age; Curinary Incontinence Due to Detrusor Overactivity    2. 18 years of age; Dystonia, Focal Upper Limb in a patient ≥ 18 years of age; Laryngeal Dystonia (Spasmodic Dysphonia) in a patient ≥ 18 years of age; Approve.  Botos: not covered in the following situations: Cosmetic Uses.  Note: Examples of cosmetic uses include facial rhytides, from lines, glabellar wrinkling, horizontal neck rhytides, mid and lower face and neck rejuvenation, platsymal bands, or rejuvenation of the periorbital region.	1 year (continued)	Yes
Bowel Evacuants – Low Volume – Polyethylene Glycol (PEG)-based Preparations	Moviprep	PEG-3350, sodium sulfate, sodium chloride, potassium chloride, sodium ascorbate, ascorbic acid	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve one of the following (A or B):  A. The patient meets both of the following (i and ii):  i. The requested non-formulary drug is being prescribed for bowel preparation as part of a colorectal cancer screening procedure in a patient between the ages of 45 years and 75 years old; AND  ii. According to the prescriber, the brand product is being requested because the preferred bioequivalent generic product would not be as medically appropriate for the patient as the requested brand non-formulary drug; OR  B. The patient meets both of the following (i and ii):  i. The requested non-formulary drug is being prescribed for a use OTHER THAN bowel preparation as part of a colorectal cancer screening procedure in a patient between the ages of 45 years and 75 years old; AND  ii. The patient meets both of the following (i and ii):  ii. The patient meets being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 уеаг	MSB Exclusion *This criteria applies only to the NPF

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  Approve if the patient meets one of the following criteria (i or ii):  i. Patient has tried PEG3350/Ascorbic Acid powder packet (Moviprep, generics). If PEG3350/Ascorbic Acid powder packet (Moviprep, generics) are non-formulary, approve; OR  ii. Patients with phenylketonuria.  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.		
Bowel Evacuants – Low Volume – Polyethylene Glycol (PEG)-based		polyethylene glycol;	Approve if the patient meets one of the following criteria (i, ii, or iii): i. Patient has tried PEG3350/Ascorbic Acid powder packet (Moviprep, generics) are non-formulary, approve; OR ii. Patients with phenylketonuria; OR iii. Patient meets both of the following (a and b): a. The requested non-formulary drug is being prescribed for bowel preparation as part of a colorectal cancer screening procedure in a patient between the ages of 45 years and 75 years old; AND		
Preparations	Plenvu	solution	b. Other formulary alternative(s) would not be as medically appropriate for the patient as the requested non-formulary drug.	1 month	Yes
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  1. Approve if the patient meets one of the following (a or b):  a. Patient has tried one of the following, if formulary: 1) PEG3350/Ascorbic Acid powder pack (Moviprep, generic) OR 2) Sod Sul-Potass Sul-Magnesium sol prep kit (Suprep, generic). If neither are formulary, approve; OR b. Patient is < 12 years of age.  2. If only PEG3350/Ascorbic Acid powder (Moviprep, generic) is formulary, approve if the patient meets one of the following criteria (a, b, c, or d):  a. Patient has tried PEG3350/Ascorbic Acid powder packet (Moviprep, generics). If PEG3350/Ascorbic Acid powder packet (Moviprep, generics) are non-formulary, approve; OR  b. The patient is less than 18 years of age; OR  c. Patients with phenylketonuria; OR  d. Patients with glucose-6-phosphate dehydrogenase deficiency.		
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  1. Approve if the patient meets one of the following (a or b):  a. Patient has tried one of the following, if formulary:  1) PEG3350/Ascorbic Acid powder pack (Moviprep, generic) OR 2) Sod Sul-Potass Sul-Magnesium sol prep kit (Suprep, generic). If neither are formulary, approve; OR  b. Patient is < 12 years of age.  2. If only PEG3350/Ascorbic Acid powder (Moviprep, generic) is formulary, approve if the patient meets one of the following criteria (a, b, c, or d):  a. Patient has tried PEG3350/Ascorbic Acid powder packet (Moviprep, generics). If PEG3350/Ascorbic Acid powder packet (Moviprep, generics) are non-formulary, approve; OR  b. The patient is less than 18 years of age; OR  c. Patients with phenylketonuria; OR		
Bowel Evacuants –		sodium picosulfate;	d. Patients with glucose-6-phosphate dehydrogenase deficiency.		
Low Volume – Sodium Picosulfate-		magnesium oxide; anhydrous citric acid	<ol> <li>Patient meets both of the following (a and b):</li> <li>The requested non-formulary drug is being prescribed for bowel preparation as part of a colorectal cancer screening procedure in a patient between the ages of 45 years and 75 years old; AND</li> </ol>		
	Clenpiq	solution	b. Other formulary alternative(s) would not be as medically appropriate for the patient as the requested non-formulary drug.	1 month	Yes

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
Therapy olass	Diana Name	Ceneric Hame	Sommercian E Street	Duration	Wedication
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  OR		
Bowel Evacuants – Low Volume – Sodium Sulfate-		magnesium sulfate; potassium sulfate; sodium sulfate	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve one of the following (A or B):  A. The patient meets both of the following (i and ii):  i. The requested non-formulary drug is being prescribed for bowel preparation as part of a colorectal cancer screening procedure in a patient between the ages of 45 years and 75 years old; AND  ii. According to the prescriber, the brand product is being requested because the preferred bioequivalent generic product would not be as medically appropriate for the patient as the requested brand non-formulary drug; OR  B. The patient meets both of the following (i and ii):  i. The requested non-formulary drug is being prescribed for a use OTHER THAN bowel preparation as part of a colorectal cancer screening procedure in a patient between the ages of 45 years and 75 years old; AND  ii. The Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber,		MSB Exclusion *This criteria applies only to
	Suprep	solution	would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  1. Approve if the patient has tried one of the following, if formulary: 1) PEG3350/Ascorbic Acid powder pack (Moviprep, generic) OR 2) Sod Sul-Potass Sul-Magnesium sol prep kit (Suprep, generic). If neither are formulary, approve.  2. If only PEG3350/Ascorbic Acid powder (Moviprep, generic) is formulary, approve if the patient meets one of the following criteria (a, b, or c):  a. Patient has tried PEG3350/Ascorbic Acid powder packet (Moviprep, generics). If PEG3350/Ascorbic Acid powder packet (Moviprep, generics) are non-formulary, approve; OR  b. Patients with phenylketonuria; OR c. Patients with glucose-6-phosphate dehydrogenase deficiency.  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  1. Approve if the patient has tried one of the following, if formulary: 1) PEG3350/Ascorbic Acid powder pack (Moviprep, generic) OR 2) Sod Sul-Potass Sul-Magnesium sol prep kit (Suprep, generic). If neither are formulary, approve.		
		polyethylene glycol	2. If only PEG3350/Ascorbic Acid powder (Moviprep, generics) is formularly, approve if the patient meets one of the following criteria (a, b, <u>or</u> c):  a. Patient has tried PEG3350/Ascorbic Acid powder packet (Moviprep, generics). If PEG3350/Ascorbic Acid powder packet (Moviprep, generics) are non-formularly, approve:		
Bowel Evacuants –		3350, sodium sulfate,			
Low Volume –		potassium chloride,	c. Patients with glucose-6-phosphate dehydrogenase deficiency.		
Polyethylene Glycol		magnesium sulfate,	3. Patient meets both of the following (a and b):		
(PEG)-based		and sodium chloride	a. The requested non-formulary drug is being prescribed for bowel preparation as part of a colorectal cancer screening procedure in a patient between the ages of 45 years and 75 years old; AND		
Preparations	Suflave	for oral solution	b. Other formulary alternative(s) would not be as medically appropriate for the patient as the requested non-formulary drug.	1 month	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  1. Approve if the patient has tried one of the following, if formulary: 1) PEG3350/Ascorbic Acid powder pack (Moviprep, generic) OR 2) Sod Sul-Potass Sul-Magnesium sol prep kit (Suprep, generic). If neither are formulary, approve.  2. If only PEG3350/Ascorbic Acid powder (Moviprep, generic) is formulary, approve if the patient meets one of the following criteria (a, b, or c):  a. Patient has tried PEG3350/Ascorbic Acid powder packet (Moviprep, generics). If PEG3350/Ascorbic Acid powder packet (Moviprep, generics) are non-formulary, approve; OR  b. Patients with phenylketonuria; OR  c. Patients with glucose-6-phosphate dehydrogenase deficiency.		
Bowel Evacuants – Low Volume –		sodium sulfate, magnesium sulfate,	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  1. Approve if the patient has tried one of the following, if formulary: 1) PEG3350/Ascorbic Acid powder pack (Moviprep, generic) OR 2) Sod Sul-Potass Sul-Magnesium sol prep kit (Suprep, generic). If neither are formulary, approve.  2. If only PEG3350/Ascorbic Acid powder (Moviprep, generic) is formulary, approve if the patient meets one of the following criteria (a, b, or c):  a. Patient has tried PEG3350/Ascorbic Acid powder packet (Moviprep, generics). If PEG3350/Ascorbic Acid powder packet (Moviprep, generics) are non-formulary, approve; OR  b. Patients with phenylketonuria; OR  c. Patients with glucose-6-phosphate dehydrogenase deficiency.  3. Patient meets both of the following (a and b):		
Sodium Sulfate-based Preparations	Sutab	and potassium chloride tablets	a. The requested non-formulary drug is being prescribed for bowel preparation as part of a colorectal cancer screening procedure in a patient between the ages of 45 years and 75 years old; AND b. Other formulary alternative(s) would not be as medically appropriate for the patient as the requested non-formulary drug.	1 month	Yes
Calcium Channel Blockers (CCBs)	Norvasc		NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Calcium Channel	Norvasc	amlodipine oral	1. Direct the patient to amlodipine tablets.	i yeai	uie ivi i
Blockers (CCBs)	Katerzia	suspension	2. If the patient is unable to swallow or has difficulty swallowing amlodipine tablets, approve if the patient has tried Norliqva oral solution, if formulary. If Norliqva oral solution is non-formulary, approve	1 year	Yes
Calcium Channel		· ·	1. Direct the patient to amlodipine tablets.		
Blockers (CCBs)	Norliqva		2. If the patient is unable to swallow or has difficulty swallowing amlodipine tablets, approve if the patient has tried Katerzia oral suspension, if formulary. If Katerzia oral suspension is non-formulary, approve.	1 year	Yes
Calcium Channel Blockers (CCBs)	Conjupri and levamlodipine		<ol> <li>Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with four formulary products from the following list: amlodipine, felodipine, nifedipine LA, nisoldipine (i four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary).</li> <li>If the patient is &lt; 18 years of age, approve if the patient has tried amlodipine, if formulary. If amlodipine is non-formulary, approve.</li> </ol>	1 year	Yes
Cancer (Injectable) – Bortezomib Agents	Boruzu	bortezomib injection	Approve if the patient has tried bortezomib injection (Velcade, generics) if formulary. If bortezomib injection (Velcade, generics) are non-formulary, approve.	1 year	Yes
Cancer (Injectable) -		pemetrexed	1. Approve if the patient has tried one other pemetrexed injectable product.		
	Axtle	l'	2. If generic pemetrexed injectable is not available or formulary, approve.	1 year	Yes
J			FLT3-ITD Mutation-positive Acute Myeloid Leukemia.  1. Approve if the patient has tried Rydapt. If Rydapt is non-formulary, approve.  2. If, according to the prescriber, the patient has or is at risk for pulmonary toxicity, approve.	. , , ,	
Cancer (Oral) – FMS-			3. Approve if the patient has already been started on Vanflyta therapy.		
Like Tyrosine Kinase 3 Inhibitors for AML	Vanflyta	guizartinib tablets	Myeloid/Lymphoid Neoplasms: Approve.	1 vear	Yes
Cancer (Oral) –	vannyta	'	Acute myeloid leuemia with isocitrate dehydrogenase-1 (IDH1) mutation positive disease in a patient ≥ 18 years of age.  1. Approve if the patient has tried Tibsovo. If Tibsovo is non-formulary, approve.	i you	100
Isocitrate			2. Approve if the patient has QTc prolongation OR is or will be taking medications that can prolong the QTc interval.		
Dehydrogenase-1	5		3. Patients with Guillain-Barre, approve.		.,
Inhibitors	Rezlidhia	olutasidenib capsules	4. Approve if the patient has already been started on Rezlidhia therapy.	1 year	Yes

Ministry   Commercial PE Criteria   Part						2025 NPF
A Protect file or foliated report (Per position travels CDE of the Scholary In Age (Per position of the Scholary In Age (P	Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	Excluded Medicaiton
Intravenous) Bispecific Human Bispecif	Cancer Agent – Multiple Myeloma Nuclear Export	Xpovio	selinexor tablets	Approve if the patient meets ONE of the following (A or B):  A. Patient meets ONE of the following (i, ii, ii); iii):  i. Patient has tried at least FOUR prior regimens for multiple myeloma; OR  ii. Patient has tried at least FOUR prior regimens for multiple myeloma; AND  b) The medication will be taken in combination with Pomalyst (pomalidomide capsules); OR  iii. Patient meets both of the following (a and b):  a) Patient has tried at least FOUR prior regimens for multiple myeloma; AND  b) The medication will be taken in combination with Pomalyst (pomalidomide capsules); OR  iii. Patient meets both of the following (a and b):  a) Patient has tried at least ONE prior regimen for multiple myeloma; AND  b) The medication will be taken in combination with bortezomib, Darzalex (daratumumb infusion), Darzalex Faspro (daratumumab and hyaluronidase-fihj injection), or Kyprolis (carfilzomib intravenous infusion); OR  Note: Examples of prior regimens include Darzalex (daratumumab intravenous infusion)/bortezomib/lenalidomide/dexamethasone, Kyprolis (carfilzomib intravenous infusion)/lenalidomide/dexamethasone, Darzalex (daratumumab intravenous infusion)/bortezomib or Kyprolis/dexamethasone, bortezomib/lenalidomide/dexamethasone, Kyprolis (carfilzomib intravenous infusion)/lenalidomide/dexamethasone, Darzalex (daratumumab intravenous infusion)/bortezomib or Kyprolis/dexamethasone, or other regimens containing a proteasome inhibitor, immunomodulatory drug, and/or anti-CD38 monoclonal antibody.  B. Patient has already been started on therapy with Xpovio.  Diffuse Large B-Cell Lymphoma; High-grade B-Cell Lymphoma; HiV-related B-Cell Lymphoma; Post-transplant lymphoproliferative disorders.  Note: Diffuse Large B-Cell Lymphoma includes patients with histologic transformation of indolent lymphomas to diffuse large B-cell lymphoma.  Approve if the patient meets ONE of the following (A or B):  A. Patient has tried at least TVO prior therapies; OR	1 уеаг	Yes
NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Cancer Agents – (Injectable) - Docetaxe intravenous infusion Docityx infusion Docityx infusion Docityx infusion Docityx infusion Docityx infusion Docetaxe intravenous infusion Intusion Docetaxe intravenous infusion Docetaxe intravenous infusion Docetaxe intravenous infusion Intusion Docetaxe intravenous infusion Docetaxe intravenous infusion Intusion Docetaxe intr	(Intravenous) - Bispecific Human Epidermal Growth Factor Receptor 2 (HER2)-Directed	77		Biliary Tract Cancer in which the tumor is human epidermal growth factor receptor 2 (HER2) positive with immunohistochemistry score of 3+ (IHC3+) as determined by an approved test in a patient ≥ 18 years of age.  1. Approve if the patient has tried one of the following regimens or, according to the prescriber, all the regimens are contraindicated (A, B, or C):  A. Enhertu; OR  B. Trastuzumab plus Perjeta; OR  C. Trastuzumab plus Tukysa.  Note: If none of these products are formulary approve: Enhertu, Perjeta, Tukysa.		
Cancer Agents – (Injectable) - Docetaxel intravenous infusion  Docityx  Infusion  Docityx  Infusion  Docityx  Infusion  Approve if the patient has tried generic docetaxel. If generic decetaxel is non-formulary, approve.  Acute Myeloid Leukemia; Approve if the patient meets ONE of the following (1 OR 2):  1. Patient is 2 18 years of age and using the decidation for post-remission maintenance; OR  2. The patient has been started on therapy with Onureg.  Leukemia (AML) Agents  Onureg  azacitadine tablets  Peripheral T-Cell Lymphoma: Approve.  Approve if the patient has rived and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with one of bendamustine vial (Treanda, generics), Bendeka, bendamustine hydrochoride injection, or Belrapzo. If none are formulary, approve.  MOTE: A trial of the requested agent would NOT count toward this requirement.  Approve if the patient meets BOTH of the following (A and B):  A. The patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with one of bendamustine vial (Treanda, generics), Bendeka, bendamustine hydrochoride injection, or Belrapzo. If none are formulary, approve.  NOTE: A trial of the requested agent would NOT count toward this requirement.  Approve if the patient meets BOTH of the following (A and B):  A. The patient has tried three of the following: A vastin, Mvasi, Vegzelma, or Zirabev, if three are formulary or one if one is formulary). If none are formulary, approve; AND  Approve if the patient meets BOTH of the following (A and B):  A. The patient has tried three of the following: Avastin, Mvasi, or Zirabev, if three are formulary or noe if one is formulary). If none are formulary, approve; AND  Approve if the patient meets BOTH of the following (A and B):  A. The patient has tried three of the following: Avastin, Mvasi, or Zirabev, if three are formulary or noe if one is formulary). If none are formulary, approve; AND  Approve if the patient meets BOTH of th	,			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		MSB Exclusion *This criteria applies only to the NPF
infusion Docivyx infusion Approve if the patient has tried generic docetaxel. If generic decetaxel is non-formulary, approve.  Cancer Agents - Acute Myeloid Leukemia: Approve if the patient meets ONE of the following (1 OR 2):  1. Patient is 2: 18 years of age and using the medication for post-remission maintenance; OR  Acute Myeloid Leukemia: Approve if the patient has been started on therapy with Onureg.  2. The patient has been started on therapy with Onureg.  Peripheral T-Cell Lymphoma: Approve.  Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with one of bendamustine vial (Treanda, generics), Bendeka, bendamustine hydrochloride injection, or Belrapzo. If none are formulary, approve.  NOTE: A trial of the requested agent would NOT count toward this requirement.  Approve if the patient meets BOTH of the following: (A and B):  A The patient has tried three of the following: (A and B):  A The patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with one of bendamustine vial (Treanda, generics), Bendeka, bendamustine hydrochloride injection, or Belrapzo. If none are formulary, approve.  NOTE: A trial of the requested agent would NOT count toward this requirement.  Approve if the patient meets BOTH of the following: (A and B):  A The patient has tried three of the following: (A and B):  A The patient has tried and, according to the prescriber, would result in a significant allergy or serious adverse reaction.  Approve if the patient meets BOTH of the following: (A and B):  A The patient has tried three of the following: (A and B):  A The patient has tried and, according to the prescriber, would result in a significant allergy or serious adverse reaction.  Approve if the patient meets BOTH of the following: (A and B):  A The patient has tried three of the following: (A and B):  A The patient has tried and, according to the prescriber, would result in a significant all	Cancer Agents – (Injectable) -		·		. you.	
Agents Onureg azacitadine tablets Peripheral T-Cell Lymphoma: Approve.  Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with one of bendamustine vial (Treanda, generics), Bendeka, bendamustine hydrochloride injection, or hydrochloride intravenous infusion  NOTE: A trial of the requested agent would NOT count toward this requirement.  Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with one of bendamustine vial (Treanda, generics), Bendeka, bendamustine hydrochloride injection, or Belrapzo. If none are formulary, approve.  NOTE: A trial of the requested agent would NOT count toward this requirement.  Approve if the patient meets BOTH of the following: Avastin, Mvasi, Vegzelma, or Zirabev, if three are formulary or one if one is formulary). If none are formulary, approve; AND  B. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.  Yes  Cancer Agents -  Cancer Agents -  Approve if the patient meets BOTH of the following (A and B):  1 year Yes  Approve if the patient meets BOTH of the following: Alymsys, Avastin, Mvasi, or Zirabev, if three are formulary or one if one is formulary). If none are formulary, approve; AND	Cancer Agents - Acute myeloid	Docivyx	infusion	Acute Myeloid Leukemia: Approve if the patient meets ONE of the following (1 <u>OR</u> 2):  1. Patient is ≥ 18 years of age and using the medication for post-remission maintenance; OR	1 year	Yes
Bendamustine Agents - Bendamustine Agents - Vivimusta bevacizumab-maly containing Agents - Alymsys Agents - Alymsys - Alymsys - Alprove if the patient meets BOTH of the following: A lymsys, Avastin, Mvasi, or Zirabev, if three are formulary (or two if two are formulary). If none are formulary, approve; AND - Alymsys - Alprove if the patient meets BOTH of the following: A vastin, Mvasi, or Zirabev, if three are formulary or one if one is formulary). If none are formulary, approve; AND - Alymsys - Alprove if the patient has tried three of the following: Alymsys, Avastin, Mvasi, or Zirabev, if three are formulary (or two if two are formulary). If none are formulary, approve; AND - Alymsys - Alprove if the patient meets BOTH of the following: Alymsys, Avastin, Mvasi, or Zirabev, if three are formulary or one if one is formulary). If none are formulary, approve; AND - Alprove if the patient meets BOTH of the following: Alymsys, Avastin, Mvasi, or Zirabev, if three are formulary or one if one is formulary). If none are formulary, approve; AND - Alprove if the patient meets BOTH of the following: Alymsys, Avastin, Mvasi, or Zirabev, if three are formulary or one if one is formulary). If none are formulary, approve; AND	` ,	Onureg	azacitadine tablets	Peripheral T-Cell Lymphoma: Approve.	1 year	Yes
Cancer Agents - Bevacizumab- containing Agents - Alymsys  Cancer Agents -  Cancer Agents -  Bevacizumab- containing Agents -  Cancer Agents -		Vivimusta	hydrochloride	Belrapzo. If none are formulary, approve.  NOTE: A trial of the requested agent would NOT count toward this requirement.	1 year	Yes
Cancer Agents - A. The patient has tried three of the following: Alymsys, Avastin, Mvasi, or Zirabev, if three are formulary (or two if two are formulary or one if one is formulary, approve; AND	Bevacizumab-	Alymsys	injection for	A. The patient has tried three of the following: Avastin, Mvasi, Vegzelma, or Zirabev, if three are formulary (or two if two are formulary or one if one is formulary). If none are formulary, approve; AND B. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the	1 year	Yes
Bevacizumab- bevacizumab-adcd bevacizumab-adcd containing Agents Vegzelma bevacizumab-adcd intravenous infusion prescriber, would result in a significant allergy or serious adverse reaction.  B. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the intravenous infusion prescriber, would result in a significant allergy or serious adverse reaction.	Bevacizumab-	Vegrelme		A. The patient has tried three of the following: Alymays, Avastin, Mvasi, or Zirabev, if three are formulary (or two if two are formulary or one if one is formulary). If none are formulary, approve; AND B. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the	1,400-	Voc

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Cancer Agents - Bevacizumab- containing Agents	Avastin	bevacizumab injection for intravenous use	Approve if the patient meets BOTH of the following (A and B):  A. The patient has tried three of the following: Alymsys, Mvasi, Vegzelma, or Zirabev, if three are formulary (or two if two are formulary or one if one is formulary). If none are formulary, approve; AND  B. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.	1 year	Yes
Cancer Agents – Bispecific Antibodies for B-Cell Lymphomas	Columvi	glofitamab intravenous infusion	Diffuse Large B-Cell Lymphoma. Note: Examples of diffuse large B-cell lymphoma (DLBCL) include DLBCL not otherwise specified, high-grade B-cell lymphoma, and DLBCL arising from indolent lymphoma. Approve if the patient meets ONE of the following (1 or 2):  1. Patient has tried Epkinly. If Epkinly is non-formulary, approve; OR 2. Patient has already been started on therapy with Columvi.  Human Immunodeficiency Virus (HIV)-Related B-Cell Lymphomas Note: HIV-related B-cell lymphomas includes HIV-related diffuse large B-cell lymphoma (DLBCL), primary effusion lymphoma, and human herpes virus-8 (HHV8) positive DLBCL. Post-Transplant Lymphoproliferative Disorders: Approve.	1 year	Yes
Cancer Agents – Bispecific Antibodies for B-Cell			Diffuse Large B-Cell Lymphoma. Note: Examples of diffuse large B-cell lymphoma (DLBCL) include DLBCL not otherwise specified, DLBCL arising from indolent lymphoma, and high-grade B-cell lymphoma. Approve if the patient meets ONE of the following: (1, 2, 3, or 4):  1. Patient has tried Columvi. If Columvi is non-formulary, approve; OR  2. Patient is unable to obtain and/or maintain intravenous access; OR  4. Patient has already been started on therapy with Epkinly.  Follicular Lymphoma. Approve if the patient meets ONE of the following: (1, 2, or 3):  1. Patient has tried Lunsumio. If Lunsumio is non-formulary, approve; OR  2. Patient is unable to obtain and/or maintain intravenous access; OR  3. Patient has tried Lunsumio. If Lunsumio is non-formulary, approve; OR  2. Patient is unable to obtain and/or maintain intravenous access; OR  3. Patient has already been started on therapy with Epkinly.  Human Immunodeficiency Virus (HIV)-Related B-Cell Lymphomas  Note: HIV-related B-cell lymphomas includes HIV-related diffuse large B-cell lymphoma (DLBCL), primary effusion lymphoma, and human herpes virus-8 (HHV8) positive DLBCL.		
Lymphomas  Cancer Agents - Bruton Tyrosine	Epkinly	injection	Post-Transplant Lymphoproliferative Disorders: Approve.	1 year	Yes
Kinase Inhibitors  Cancer Agents -	Jaypirca	pirtobrutinib tablets	Richter's Transformation to Diffuse Large B-Cell Lymphoma; Marginal Zone Lymphoma; Waldenstrom Macroglobulinemia/Lymphoplasmacytic Lymphoma: Approve.	1 year	Yes
Kinase inhibitor (phosphatidylinositol	lah:	in a continuit de la	Con Canadamy. Has sale Drive Authorization Delices with size	See PA	V
3-kinase [PI3K])	Itovebi	inavolisib tablets	See Oncology – Itovebi Prior Authorization Policy criteria.	duration	Yes

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
			Non-Small Cell Lung Cancer - KRAS G12C-mutated.  1. Approve if the patient has tried Lumakras. If Lumakras is non-formulary, approve.  2. Patient with brain metastases, approve.  3. Approve if the patient has already been started on therapy with Krazati.  Colon or Rectal Cancer - KRAS G12C-mutated.		
			1. Approve if the patient has tried Lumakras. If Lumakras is non-formulary, approve.		
Cancer Agents -			2. Approve if the patient has already been started on therapy with Krazati OR has already been started on therapy with Erbitux.		
Kirsten rat sarcoma					
` /	Krazati	adagrasib tablets	Ampullary Adenocarcinoma - KRAS G12C-mutated; Biliary Tract Cancer - KRAS G12C-mutated; Pancreatic Adenocarcinoma - KRAS G12C-mutated; Small Bowel Adenocarcinoma - KRAS G12C-mutated: Approve.	1 year	Yes
Cancer Agents - NSCLC (Oral) -MET			Non-Small Cell Lung Cancer (NSCLC) with mesenchymal-epithelial transition (MET) exon 14 skipping alterations or high-level MET amplification:		
receptor tyrosine			1. Approve if the patient has tried Tabrecta. If Tabrecta is non-formulary, approve.		
kinase inhibitor	Tepmetko	tepotinib tablets	2. Approve if the patient has already been started on Tepmetko.	1 year	Yes
Cancer Agents - PARP inhibitor/Prostate Cancer Agent	Akeega	niraparib and abiraterone acetate tablets	BRCA-mutated Prostate Cancer.  1. Approve if the patient has tried ONE of the following: 1) Lynparza +/- abiraterone or 2) Talzenna plus Xtandi.  Note: If either medication in the regimens above are non-formulary, then that regimen does not need to be tried.  Note: If Lynparza is non-formulary, approve.  2. Approve if the patient has already been started on therapy with Akeega.	1 year	Yes
Cancer Agent	Akeeya	labiels	1. Ovarian, Fallopian Tube, or Primary Peritoneal Cancer in the Maintenance setting (after complete or partial response to chemotherapy): Approve if the patient meets one of the following criteria (A or B):	i yeai	165
Cancer Agents - PARP Inhibitors (oral)	Rubraca	rucaparib tablets	A) Patient has tried one of Zejula or Lynparza. If neither are formulary, approve; OR B) Patient has already started on Rubraca.  2. Prostate cancer: Approve if the patient meets one of the following criteria (A or B): A) Patient has tried Lynparza. If Lynparza is non-formulary, approve; OR B) Patient has already started on Rubraca.  3. <u>Uterine Leiomyosarcoma</u> : Approve if the patient meets one of the following (A or B): A) Patient has tried one of Zejula or Lynparza. If neither is formulary, approve; OR B) Patient has already started on Rubraca.  4. <u>Pancreatic Adenocarcinoma</u> : approve.	1 year	Yes- 7/1
Cancer Agents -		niraparib capsules	1. Ovarian, Fallopian Tube, or Primary Peritoneal Cancer in the Maintenance setting (after complete or partial response to chemotherapy): Approve if the patient meets one of the following (i or ii):  i. Patient has tried Lynparza [documentation required]. If Lynparza is non-formulary, approve; OR  ii. Patient has already been started on therapy with Zejula; OR  B) Patient meets both of the following (i and ii):  i. Patient has had a complete or partial response to first-line platinum-based chemotherapy; AND  ii. Patient does not have a BRCA mutation [documentation required].  2. Uterine Leiomyosarcoma: Approve if the patient meets one of the following (A or B):  A) Patient has tried one of Rubraca or Lynparza [documentation required]. If neither is formulary, approve; OR		
PARP Inhibitors (oral)	Zejula	and tablets	B) Patient has already started on Zejula.	1 year	Yes - 7/1
Cancer Agents - Prostate Cancer (Oral)	Zytiga		NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Cancer Agents - Renal Cell Carcinoma (Oral)	Afinitor Disperz	everolimus tablets for oral suspension	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Cancer Agents - Renal Cell Carcinoma (Oral)	Afinitor tablet		NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Cancer Agents - Renal Cell Carcinoma (Oral)	ı Fotivda	tivozanib capsules	Renal Cell Carcinoma. Approve if the patient meets one of the following (1, 2, or 3):  1. Patient has tried one of Inlyta, Lenvima, or Cabometyx. If none are formulary, approve; OR  2. If there are toxicity concerns with a trial of Lenvima (and other concomitantly given medications), according to the prescriber, approve if the patient has tried Inlyta or Cabometyx. If neither are formulary, approve; OR  3. Patient has already been started on therapy with Fotivda.	1 year	Yes
Cancer Agents - Trastuzumab- containing Agents	Hercessi	trastuzumab-strf for intravenous injection	Approve if the patient meets BOTH of the following (A and B):  A. Patient has tried four products from the following list (if four are formulary or three if three are formulary or two if two are formulary or one if one is formulary): Herceptin intravenous, Kanjinti, Trazimera, Ogivri, Ontruzant, or Herzuma; AND  Note: If none are formulary, approve.  B. Patient cannot continue to use each of the alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.	1 year	Yes
Cancer Agents - Trastuzumab- containing Agents	Herceptin	trastuzumab for intravenous injection	Approve if the patient meets BOTH of the following (A and B):  A. Patient has tried four products from the following list (if four are formulary or three if three are formulary or two if two are formulary or one if one is formulary): Hercessi, Kanjinti, Trazimera, Ogivri, Ontruzant, or Herzuma; AND Note: If none are formulary, approve.  B. Patient cannot continue to use each of the alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.	1 year	Yes
Cancer Agents - Trastuzumab- containing Agents	Herceptin Hylecta	trastuzumab and hyaluronidase-oysk for subcutaneous use	<ol> <li>Approve if the patient has tried one product from the following list (if one is formulary): Herceptin intravenous, Hercessi, Kanjinti, Trazimera, Ogivri, Ontruzant, or Herzuma. If none are formulary, approve.</li> <li>Approve if the patient is unable to obtain and/or maintain intravenous access.</li> <li>If the patient has already been started on therapy with Herceptin Hylecta, approve.</li> </ol>	1 year	Yes
Cancer Agents - Trastuzumab- containing Agents	Herzuma	trastuzumab-pkrb for intravenous injection	Approve if patient meets BOTH of the following (A and B):  A. Patient has tried four products from the following list (if four are formulary or three if three are formulary or two if two are formulary or one if one is formulary): Herceptin intravenous, Hercessi, Kanjinti, Ogivri, Ontruzant, or Trazimera; AND  Note: If none are formulary, approve.  B. Patient cannot continue to use each of the alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.	1 year	Yes
Cancer Agents - Trastuzumab- containing Agents	Ogivri	trastuzumab- dkst	Approve if the patient meets BOTH of the following (A and B):  A. Patient has tried four products from the following list (if four are formulary or three if three are formulary or two if two are formulary or one if one is formulary): Hercessi, Herceptin intravenous, Trazimera, Kanjinti, Ontruzant, or Herzuma; AND  Note: If none are formulary, approve.  B. Patient cannot continue to use each of the alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.	1 year	Yes
Cancer Agents - Trastuzumab-		trastuzumab-dttb for	Approve if the patient meets BOTH of the following (A and B):  A. Patient has tried four products from the following list (if four are formulary or three if three are formulary or two if two are formulary or one if one is formulary): Hercessi, Kanjinti, Trazimera, Ogivri, Herzuma, or Herceptin intravenous; AND  Note: If none are formulary, approve.  B. Patient cannot continue to use each of the alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would		W
containing Agents  Cancer Agents - Tyrosine Kinase Inhibitors	Ontruzant  Gleevec	intravenous injection	result in a significant allergy or serious adverse reaction.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year 1 year	Yes  MSB Exclusion *This criteria applies only to the NPF
Cancer Agents - Tyrosine Kinase Inhibitors	Sprycel	dasatinib tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Cancer Agents - Tyrosine Kinase Inhibitors	Tykerb	lapatinib tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF

					2025 NPF
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	Excluded Medicaiton
Cancer Agents - Tyrosine Kinase Inhibitors	Qinlock	ripretinib tablets	Gastrointestinal stromal tumor.  1. Approve if the patient has been previously treated with at least two other kinase inhibitors.  Note: Examples of kinase inhibitors are an imatinib product (Gleevec, generics, Imkeldi), sunitinib (Sutent), Stivarga, sorafenib (Nexavar), pazopanib (Votrient), Tasigna, dasatinib (Sprycel), Ayvakit.  2. Approve if the patient has already been started on therapy with Qinlock.  Melanoma. Cutaneous.  1. Approve if the patient meets all of the following (A, B and C):  A. Patient has metastatic or unresectable disease; AND  B. Patient has an activating KIT mutation; AND  C. Patient has tried at least one systemic regimen.  Note: Examples of a systemic regimen include: Opdivo (nivolumab intravenous infusion) + Yervoy (ipilimumab intravenous infusion), Opdualag (nivolumab/relatlimab-rmbw intravenous infusion), Keytruda (pembrolizumab intravenous infusion), Opdivo, Tafinlar (dabrafenib capsules) + Mekinist (trametinib tablets), Zelboraf (vemurafenib tablets) + Cotellic (cobimetinib tablets), Braftovi (encorafenib capsules) + Mektovi (binimetinib tablets).  2. Approve if the patient has already been started on therapy with Qinlock.	1 year	Yes
Cancer Agents (Injectable) –					
melphalan	Ivra	melphalan injection	Approve if the patient has tried and cannot use melphalan injection (Alkeran, generics). If melphalan injection (Alkeran, generics) is non-formulary, approve.	1 year	Yes
Carbonic Anhydrase Inhibitors	Keveyis and generics (including dichlorphenamide tablets, Ormalvi)	dichlorphenamide tablets	Approve if the patient has tried one of dichlorphenamide tablets or Ormalvi, if formulary.  If BOTH dichlorphenamide tablets and Ormalvi are non-formulary, or generic dichlorphenamide or Ormalvi is being requested, approve if the patient meets one of the following (1 or 2):  1. For the treatment of primary hyperkalemic periodic paralysis (HyperPP), primary hypokalemic periodic paralysis (HypoPP), and related variants: approve if the patient has tried one of acetazolamide tablets (generics) or acetazolamide ER capsules, if one is formulary. If neither are formulary, approve.  2. For the treatment of primary hyperkalemic periodic paralysis (HyperPP), primary hypokalemic periodic paralysis (HypoPP), and related variants: approve if the patient has been started on therapy with Keveyis, Ormalvi or dichlorphenamide.	1 year	Yes - brand only MSB Exclusion
Cardiovascular Medications - Other	BiDil	isosorbide dinitrate and hydralazine tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	*This criteria applies only to the NPF
Cardiovascular Medications - Other	Tikosyn	dofetilide capsules	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Cardiovascular Medications - Other	Corlanor	ivabradine tablets and solution	If requesting brand Corlanor tablets or Corlanor solution, approve if the patient meets ONE of the following (1 or 2):  1. Patient has tried and cannot take generic ivabradine tablets; OR  2. Patient cannot swallow or has difficulty swallowing tablets, approve Corlanor solution.  If requesting generic ivabradine tablets or generic ivabradine tablets are non-formulary, approve if the patient meets ONE of the following (1, 2, or 3):  1. Patient has tried, or is currently receiving a beta-blocker (e.g., bisoprolol, carvedilol, metoprolol) OR the patient has a contraindication to beta-blockers; OR  2. Heart failure due to dilated cardiomyopathy, approve if the patient is < 18 years of age; OR  3. Patient has already been started on Corlanor or ivabradine.	1 year	Yes
Cardiovascular Medications - Other	Aspruzyo Sprinkle	ranolazine extended-release granules	<ol> <li>Approve if the patient meets one of the following (A or B):         <ul> <li>A. Patient is unable to or has difficulty swallowing ranolazine extended-release tablets (Ranexa, generics); OR</li> <li>B. Patient requires administration by nasogastric or gastrostomy/gastric tube.</li> </ul> </li> <li>If ranolazine extended-release tablets (Ranexa, generics) are non-formulary, approve if the patient meets one of the following (A, B, or C):</li></ol>	1 year	Yes
Control Nonvous			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		MSB Exclusion *This criteria
Central Nervous System Non- Stimulants	Intuniv	guanfacine HCl tablets	Criteria: Approve if the Brand product is being requested up to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	applies only to the NPF
Central Nervous System Non- Stimulants	Strattera	atomoxetine HCI capsules	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF

					2025 NPF
				Approval	Excluded
Therapy Class	Brand Name	Generic Name clonidine	Commercial FE Criteria	Duration	Medicaiton
Central Nervous		hydrochloride			
System Non-			1. Approve if the patient has tried clonidine ER tablets (generic of Kapvay), if formulary. If clonidine ER tablets (generic of Kapvay) is non-formulary, approve.		
Stimulants	Onyda XR	suspension	2. Approve if the patient is unable to swallow tablets or has difficulty swallowing tablets.	1 year	Yes
Central Nervous			<u></u>		MSB Exclusion
System Stimulants –		d	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Amphetamine Products	Adderall	mphetamine tablets	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	applies only to the NPF
Central Nervous	Adderail	Inprietamine tablets	prescriber, would result in a significant allergy or serious adverse reaction (documentation required).	i yeai	MSB Exclusion
System Stimulants –		dextroamphetamine/a	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Amphetamine		mphetamine extended	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Products	Adderall XR	release capsules	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Central Nervous			<u></u>		MSB Exclusion
System Stimulants –			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Amphetamine Products	Evekeo	amphetamine sulfate tablet	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	applies only to the NPF
Central Nervous	LVCKCO	tabict	1. Approve if the patient has tried Dyanavel XR oral suspension, if formulary.	i yeai	uic ivi i
System Stimulants –		amphetamine	2. If Dyanavel XR oral suspension is non-formulary, approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with three products (or two if two are formulary or one if		
Amphetamine		extended-release	one is formulary) from the following list: 1) amphetamine mixed extended-release (Er) capsules (Adderall XR, generics), or 2) Adzenys XR ODT tablets, or 3) lisdexamfetamine capsules (Vyvanse capsule, generics) or lisdexamfetamine		
Products	Dyanavel XR tablets	tablets	chewable tablets (Vyvanse chewable tablet, generics). If none are formulary, approve.	1 year	Yes
0 1 N			1. Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with three products (or two if two are formulary or one if one is formulary) from the following list: 1)		
Central Nervous System Stimulants –		amphetamine	amphetamine mixed extended-release (Er) capsules (Adderall XR, generics), or 2) Adzenys XR ODT tablets, or 3) lisdexamfetamine capsules (Vyvanse capsule, generics) or lisdexamfetamine chewable tablets (Vyvanse chewable tablet, generics). If none are formulary, approve.		
Amphetamine	Dyanavel XR		Lastics, generous). In notice are forminging, approve.  2. If the patient cannot swallow solid oral dosage forms or has difficulty swallowing solid oral dosage forms AND the patient is unable to ingest the prescribed dosage when using a product that can be opened and sprinkled on food,		
Products	suspension	suspension	approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with Adzenys XR ODT tablets, if formulary. If Adzenys XR ODT tablets are non-formulary, approve.	1 year	Yes
Central Nervous	,	'			MSB Exclusion
System Stimulants –		methylphenidate	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Methylphenidate			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Products	Aptensio XR	capsule	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Central Nervous System Stimulants –		methylphenidate hcl	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		MSB Exclusion *This criteria
Methylphenidate		extended-release	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Products	Concerta	tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Central Nervous					MSB Exclusion
System Stimulants –		dexmethylphenidate	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Methylphenidate	Focalin and Focalin	tablets and extended-	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Products	XR	release capsules	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Central Nervous System Stimulants –			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		MSB Exclusion *This criteria
Methylphenidate		methylphenidate	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Products	Ritalin	tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Central Nervous					MSB Exclusion
System Stimulants –			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Methylphenidate	Ditalia I A		Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the	1 11000	applies only to the NPF
Products	Ritalin LA	acting capsules	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	ule NPF
Central Nervous			Approve if the patient has tried Quillivant XR suspension, if formulary.		
System Stimulants –		methylphenidate HCI	If Quillivant XR suspension is non-formulary, approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with four products (or three if three are formulary, or two if two		
Methylphenidate		extended-release	are formulary, or one if one is formulary) from the following list: 1) dexmethylphenidate extended-release capsules (Focalin XR, generics), 2) methylphenidate extended-release capsules (Aptensio XR, generics), 3) Jornay PM, 4)		
Products	QuilliChew ER	chewable tablets	Azstarys.	1 year	Yes
Central Nervous			Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with five products (or four if four are formulary, or three if three are formulary, or two if two are formulary, or		
System Stimulants –	D. L		one if one is formulary) from the following list: 1) dexmethylphenidate extended-release capsules (Focalin XR, generics), 2) methylphenidate extended-release capsules (Aptensio XR, generics), 3) Jornay PM, or 4) Azstarys or 5)		
Methylphenidate Products	Relexxii and authorized generic	methylphenidate ER tablet	QuilliChew ER tablets or Quillivant XR suspension.  Note: QuilliChew ER tablets and Quillivant XR suspension count as one alternative.	1 year	Yes
Toddels	Tautionzeu genend	เฉมเซเ	Table. Quinto new List table to and Quinty and Art out-perioded to the attendance.	ı yeai	103

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Central Nervous System Stimulants – Methylphenidate		methylphenidate hydrochloride for extended-release oral	1. Approve if the patient has tried QuilliChew ER tablets, if formulary. If QuilliChew ER tablets are non-formulary, approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with four products (or three if three are formulary, or two if two are formulary, or one if one is formulary) from the following list: 1) dexmethylphenidate extended-release capsules (Focalin XR, generics), 2) methylphenidate extended-release capsules (Aptensio XR, generics), 3) Jornay PM, 4) Azstarys.  2. If the patient cannot swallow solid oral dosage forms or has difficulty swallowing solid oral dosage forms AND the patient is unable to ingest the prescribed dosage when using a product that can be opened and sprinkled on food,		
Products	Quillivant XR	suspension	approve.	1 year	Yes
Central Nervous System Stimulants –Amphetamine Products	Xelstrym	dextroamphetamine transdermal system	Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with three products, if formulary (or two if two are formulary or one if one is formulary) from the following list:  1) amphetamine mixed extended-release (Er) capsules (Adderall XR, generics), 2) Adzenys XR-ODT tablets, 3) lisdexamfetamine capsules (Vyvanse capsule, generics) or lisdexamfetamine chewable tablets (Vyvanse chewable tablet, generics) 4) Dyanavel XR oral suspension, or 5) dextroamphetamine extended-release capsules. If none are formulary, approve.	1 year	Yes
Central Nervous	Sodium oxybate oral		Cataplexy Treatment in Patients with Narcolepsy:  Direct the patient to one of 1) Xyrem (brand) OR 2) sodium oxybate oral solution (by Hikma), if formulary. If neither are formulary, approve if the patient meets (1 or 2):  1. Patients ≥ 18 years of age: approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with one of Lumryz, Xywav, or Wakix, if formulary. If none are formulary, approve.  2. Patients ≥ 7 years of age and < 18 years of age: approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with one of Lumryz or Xywav, if formulary. If neither are formulary, approve.  Excessive Daytime Sleepiness in Patients with Narcolepsy:		
System/Autonomic Drugs	solution (AG to Xyrem) by AMNEAL	sodium oxybate oral solution	Direct the patient to one of 1) Xyrem (brand) OR 2) sodium oxybate oral solution (by Hikma), if formulary. If neither are formulary, approve if the patient (≥ 7 years of age) has tried and, according to the prescriber, has experienced linadequate efficacy OR a significant intolerance with one of Lumryz, Xywav, or Wakix, if formulary. If none are formulary, approve.	1 year	Yes
			Cataplexy Treatment in Patients with Narcolepsy:  Direct the patient to one of 1) sodium oxybate oral solution (authorized generic of Xyrem) [by Hikma] OR 2) sodium oxybate oral solution (authorized generic of Xyrem) [by Amneal], if formulary. If neither are formulary, approve if the patient meets (1 or 2):  1. Patients ≥ 18 years of age: approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with one of Lumryz, Xywav, or Wakix, if formulary. If neither are formulary, approve.  2. Patients ≥ 7 years of age and < 18 years of age: approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with one of Lumryz or Xywav, if formulary. If neither are formulary, approve.		
Central Nervous System/Autonomic Drugs	Xyrem (brand)	sodium oxybate oral solution	Excessive Daytime Sleepiness in Patients with Narcolepsy:  Direct the patient to one of 1) sodium oxybate oral solution (authorized generic of Xyrem) [by Hikma] OR 2) sodium oxybate oral solution (authorized generic of Xyrem) [by Amneal], if formulary. If neither are formulary, approve if the patient (≥ 7 years of age) has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with one of Lumryz, Xywav, or Wakix, if formulary. If none are formulary, approve.	1 year	Yes
Central Nervous System/Autonomic Drugs	Northera and generic droxidopa capsules	droxydopa capsules	Neurogenic Orthostatic Hypotension.  Approve if the patient has tried two of the following products: 1) midodrine tablets, 2) fludrocortisone tablets, 3) dihydroergotamine injection/nasal spray, 4) indomethacin capsules/injection, 5) pyridostigmine tablets, or 6) atomoxetine.	1 year	Yes
Chagas Disease	1		1. Approve if the patient has tried benznidazole, if formulary. If benznidazole is non-formulary, approve. 2. Approve if the patient is less than 2 years of age.		V
Agents  Chelating Agents	Lampit	nifurtimox tablets  deferasirox tablets for oral suspension	3. Approve if the patient has already started on therapy with Lampit.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Chelating Agents	Jadenu	deferasirox tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Chelating Agents	Jadenu Sprinkles	deferasirox oral granules	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Chelating Agents - Wilson's Disease	Cuprimine	penicillamine capsules	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
			Approve if the patient has tried generic trientine 250 mg capsules, if formulary.		
			If generic trientine 250 mg capsules are non-formulary, approve if the patient meets one of the following:		
			1. Approve if the patient has tried one penicillamine product: penicillamine (Cuprimine, generics) or penicillamine (Depen, generics), if one is formulary. If neither are formulary, approve.		
			2. Approve if per the prescriber, the patient is intolerant to penicillamine or the patient has clinical features indicating the potential for intolerance to penicillamine (i.e., history of any renal disease, congestive splenomegaly causing severe thrombocytopenia, autoimmune tendency).		
			3. Approve if, per the prescriber, the patient has a contraindication to penicillamine.		
			4. Approve if the patient has neurological manifestations of Wilson's Disease.		
Chelating Agents -	trientine 500 mg	trientine 500 mg	5. Approve if the patient is pregnant.		
Wilson's Disease	capsules	capsules	6. Approve if the patient has been started on therapy with a trientine product.	1 year	Yes
771100110 2100000	- Capoulos	- Capoaioo	Approve if the patient has tried trientine capsules (Syprine, generics), if formulary.	. ,	
			(-),,,,,,		
			If trientine capsules (Syprine, generics) are non-formulary, approve if the patient meets one of the following:		
i			1. Approve if the patient has tried one penicillamine product: penicillamine (Cuprimine, generics) or penicillamine (Depen, generics), if one is formulary. If neither are formulary, approve.		
			2. Approve if per the prescriber, the patient is intolerant to penicillamine or the patient has clinical features indicating the potential for intolerance to penicillamine (i.e., history of any renal disease, congestive splenomegaly causing severe		
			thrombocytopenia, autoimmune tendency).		
			3. Approve if, per the prescriber, the patient has a contraindication to penicillamine.		
0		trientine	4. Approve if the patient has neurological manifestations of Wilson's Disease.		
Chelating Agets -	C	tetrahydrochloride	5. Approve if the patient is pregnant.	1	V
Wilson's Disease	Cuvrior	300 mg tablets	6. Approve if the patient has been started on therapy with a trientine product or Cuvrior.	1 year	Yes
			Atherosclerotic Disease. Approve if the patient meets ALL of the following (1, 2, 3, and 4):		
			1. Patient is ≥ 18 years of age; AND		
			2. Lodoco is being added onto a background regimen(s) of other atherosclerotic disease medication(s) [documentation required]: AND		
			Note: Examples of medications recommended in guideline-directed therapy for patients with atherosclerotic disease can include aspirin, antiplatelet agents (e.g., clopidogrel, Brilinta [ticagrelor tablets]), anticoagulants, lipid-lowering		
			agents (e.g., statins such as atorvastatin and rosuvastatin), beta blockers, angiotensin-converting enzyme inhibitors, and/or angiotensin receptor blockers.		
		colchicine 0.5 mg	3. Patient has a creatinine clearance ≥ 50 mL/min; AND		
Colchicine Agents	Lodoco	tablets	4. Patient has tried colchicine 0.6 mg tablets or capsules [documentation required].	1 year	Yes
			Chronic Graft-Versus-Host Disease.		
			1. Approve if the patient has tried at least TWO systemic medications for chronic graft-versus-host disease.		
Colony Stimulating			Note: Examples of systemic therapy may include Jakafi (ruxolitinib tablets), Rezurock (belumosudil tablets), Impruvica (ibrutinib tablets, capsules, and oral suspension), imatinib, hydroxychloroquine, methotrexate, rituximab, pentostatin,		
Factor-1 Receptor- Blocking Antibody	Niktimvo	axatilimab-csfr intravenous infusion	interleukin-2 (e.g., Proleukin [aldesleukin intravenous infusion]), methylprednisolone, cyclosporine, tacrolimus, sirolimus, etanercept products, and mycophenolate mofetil.	1 vear	Yes
blocking Antibody	INIKUITIVO	intravenous iniusion	2. Approve if the patient has already been started on therapy with Niktimvo.  Cancer in a Patient ≥ 18 Years of Age Receiving Myelosuppressive Chemotherapy.	i yeai	res
			Approve if the patient meets the following (A and B):		
			A. Patient has tried Ryzneuta, if formulary; AND		
		eflapegrastim-xnst	B. Patient has tried one pegfilgrastim product [documentation required].		
Colony Stimulating		subcutaneous	Note: Pegfilgrastim products are Neulasta, Fulphila, Fylnetra, Nyvepria, Udenyca, Stimufend, and Ziextenzo.		
Factors	Rolvedon	injection	Note: If neither Ryzneuta nor any pegfilgrastim products are formulary, approve.	1 year	Yes
			1. Approve if the patient meets BOTH of the following (A and B):		
			a. Patient has tried four of the following products, if formulary (or three if three are formulary, or two if two are formulary, or one if one is formulary): Neupogen, Releuko, Zarxio, Nivestym, or Granix [documentation required]; AND		
			Note: If none are formulary, approve.		
			B. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.		
		filgrastim-txid	2. Patients who require administration by intravenous infusion: approve if the patient meets the following (A and B):		
		subcutaneous or	A. Patient has tried one of Neupogen, Releuko, Zarxio, or Nivestym [documentation required], if formulary; AND		
0 1 0" 1 "		intravenous injection	Note: If none are formulary, approve.		
Colony Stimulating	Nymozi	(biosimilar to	B. Patient cannot continue to use the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would require a property of the prescriber alternative agent, and/or surfactant which according to the prescriber, would be a surfactant agent, and/or surfact	1	Voc
Factors - Filgrastim	Nypozi	Neupogen)	result in a significant allergy or serious adverse reaction.	1 year	Yes

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
			1. Approve if the patient meets BOTH of the following (A and B):  A. Patient has tried four of the following products, if formulary (or three if three are formulary, or two if two are formulary, or one if one is formulary): Nypozi, Releuko, Neupogen, Nivestym, or Zarxio [documentation required]; AND Note: If none are formulary, approve.  B. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.		
Colony Stimulating Factors - Filgrastim	Granix	tbo-filgrastim subcutaneous injection	<ul> <li>2. Patients requiring a dose &lt; 180 mcg: approve if the patient meets the following (A and B):         <ul> <li>A. Patient has tried one of Neupogen or Nivestym [documentation required], if formulary; AND</li> </ul> </li> <li>Note: If neither are formulary, approve.</li> <li>B. Patient cannot continue to use the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.</li> </ul>	1 year	Yes
			1. Approve if the patient meets BOTH of the following (A and B):  A. Patient has tried four of the following products, if formulary (or three if three are formulary, or two if two are formulary, or one if one is formulary):  Note: If none are formulary, approve.  B. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.		
			2. Patients who require administration by intravenous infusion: approve if the patient meets the following (A and B):  A. Patient has tried one of Nypozi, Releuko, Zarxio, or Nivestym [documentation required], if formulary; AND  Note: If none are formulary, approve.  B. Patient cannot continue to use the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.  Note: If Nivestym is non-formulary and the patient requires a dose of < 180 mcg, approve.		
Colony Stimulating Factors - Filgrastim	Neupogen	filgrastim intravenous or subcutaneous injection	3. Patients requiring a dose < 180 mcg: approve if the patient meets the following (A and B):  A. Patient has tried one of Nivestym or Granix [documentation required]; AND  Note: If neither are formulary, approve.  B. Patient cannot continue to use the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.  Note: If the only formulary alternative is Granix and the patient requires intravenous administration, approve.	1 vear	Yes
,	. 0	filgrastim-ayow subcutaneous or intravenous injection	<ol> <li>Approve if the patient meets BOTH of the following (A and B):         <ul> <li>A. Patient has tried four of the following products, if formulary (or three if three are formulary, or two if two are formulary, or one if one is formulary): Nypozi, Nivestym, Neupogen, Granix, or Zarxio [documentation required]; AND Note: If none are formulary, approve.</li> <li>B. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.</li> </ul> </li> <li>Patients who require administration by intravenous infusion: approve if the patient meets BOTH of the following (A and B):         <ul> <li>A. Patient has tried one of Nypozi, Nivestym, Neupogen, or Zarxio [documentation required], if formulary; AND Note: If none are formulary, approve.</li> </ul> </li> </ol>		
Colony Stimulating Factors - Filgrastim	Releuko	(biosimilar to Neupogen)	B. Patient cannot continue to use the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.	1 year	Yes
			1. Approve if the patient meets BOTH of the following (A and B):  A. The patient has tried four of the following products, if formulary (or three if three are formulary, or two if two are formulary, or one if one is formulary): Nypozi, Releuko, Neupogen, Nivestym, or Granix [documentation required];  AND  Note: If none are formulary, approve.  B. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.		
Colony Stimulating		filgrastim-sndz subcutaneous or intravenous injection (biosimilar to	2. Patients who require administration by intravenous infusion: approve if the patient has meets the following (A and B):  A. Patient has tried one of Nypozi, Releuko, Neupogen, or Nivestym [documentation required], if formulary; AND  Note: If none are formulary, approve.  B. Patient cannot continue to use the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber,		
,	Zarxio	Neupogen)	would result in a significant allergy or serious adverse reaction.	1 year	Yes

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
			Approve if the patient meets BOTH of the following (a and b):  a. The patient has tried five of the following, if five are formulary (or four if four are formulary or three if three are formulary or two if two are formulary or one if one is formulary): Neulasta, Nyvepria, Fulphila, Udenyca, Ziextenzo, or		
			a. The patient has thed live of the following, if live are formularly of flour if four are formularly of two if two are formularly of one if one is formularly. Neulasta, Nyvepha, Pulphila, odenyca, ziexterizo, of Fvinetra [documentation required]; AND		
Colony Stimulating			Note: If none are formulary, approve.		
Factors -			b. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the		
Pegfilgrastim	Stimufend	pegfilgrastim-fpgk	prescriber, would result in a significant allergy or serious adverse reaction.	1 year	Yes
- 0 0		1 0 0 10	Approve if the patient meets BOTH of the following (a and b):	ĺ	
			a. The patient has tried five of the following, if five are formulary (or four if there are formulary or two if two are formulary or one if one is formulary): Neulasta, Fulphila, Udenyca, Ziextenzo, Nyvepria, or		
			Stimufend [documentation required]; AND		
Colony Stimulating		pegfilgrastim-pbbk	Note: If none are formulary, approve.		
Factors -		subcutaneous	b. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the		
Pegfilgrastim	Fylnetra	injection	prescriber, would result in a significant allergy or serious adverse reaction.	1 year	Yes
			Approve if the patient meets BOTH of the following (a and b):		
			a. The patient has tried five of the following, if five are formulary (or four if four are formulary or three if three are formulary or two if two are formulary or one if one is formulary): Fulphila, Udenyca, Ziextenzo, Nyvepria, Fylnetra, or		
			Stimufend [documentation required]; AND		
Colony Stimulating		pegfilgrastim	Note: If none are formulary, approve.		
Factors - Pegfilgrastim	Neulasta	subcutaneous injection	b. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.	1 vear	Yes
Pegiligrasiiiii	iveulasta	Injection		i yeai	res
			Approve if the patient meets BOTH of the following (a and b):  a. The patient has tried five of the following, if five are formulary (or four if four are formulary or three if three are formulary or two if two are formulary or one if one is formulary): Neulasta, Fulphila, Udenyca, Ziextenzo, Fylnetra, or		
			Stimufend [documentation required]: AND		
Colony Stimulating		pegfilgrastim-apgf	Note: If none are formulary, approve.		
Factors -		subcutaneous	b. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the		
Pegfilgrastim	Nyvepria	injection	prescriber, would result in a significant allergy or serious adverse reaction.	1 year	Yes
			Approve if the patient meets BOTH of the following (a and b):		
			a. The patient has tried five of the following, if five are formulary (or four if four are formulary or three if three are formulary or two if two are formulary or one if one is formulary): Neulasta, Fulphila, Ziextenzo, Nyvepria, Fylnetra, or		
			Stimufend [documentation required]; AND		
Colony Stimulating		pegfilgrastim-cbqv	Note: If none are formulary, approve.		
Factors -		subcutaneous	b. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the		
Pegfilgrastim	Udenyca	injection	prescriber, would result in a significant allergy or serious adverse reaction.	1 year	Yes
			Paroxysmal nocturnal hemoglobinuria in a patient ≥ 13 years of age.		
			1. Approve if the patient has tried ONE of 1) an eculizumab product (Soliris, Bkemv, Epysqli) or 2) Ultomiris, if formulary. If neither are formulary, approve.		
		crovalimab-akkz	Note: All of the eculizumab products would count as one alternative (Soliris, Bkemv, Epysqli).  2. Patient < 18 years of age, approve if the patient has tried Ultomiris, if formulary. If Ultomiris is non-formulary, approve.		
Complement		intravenous infusion and subcutaneous	3. Patient is unable to maintain intravenous access, approve.		
Inhibitors	PiaSky	iniection	4. Patient has already been started on therapy with PiaSky, approve.	1 vear	Yes
THI IDIO 3	Паску	Injection		i year	103
			Anti-acetylcholine receptor antibody positive generalized myasthenia gravis in a patient ≥ 18 years of age.		
			Approve if the patient meets one of the following (1 or 2):  1. Patient meets BOTH of the following (A and B):		
			A. Patient meets one of the following (i or ii):		
			i. Patient has tried, and according to the prescriber, experienced inadequate efficacy OR significant intolerance with ONE of 1) an eculizumab product (Soliris, Bkemv, Epysqli) or 2) Ultomiris, if formulary; OR		
			I. Faterinas tred, and according to the prescriber, experience indeequate entacty on significant molerance with one or 1) an equipment ground (solins, brenty, epysqri) or 2) originally, on Note: All of the equipment products would count as one alternative (Solins, Bkenry, Epysqri).		
			ii. Patient is unable to obtain intravenous access; AND		
			Note: If neither are formulary, would still need to meet criterion B.		
Complement			B. Patient has tried, and according to the prescriber, experienced inadequate efficacy OR significant intolerance with one of Imaavy, Vyvgart intravenous, Vyvgart Hytrulo, or Rystiggo, if formulary; OR		
Inhibitors -		zilucoplan	Note: If none are formulary, would still need to meet criterion A.		
Complement C5		subcutaneous	Note: If there are no formulary alternatives from criterion A or B, approve.		
inhibitor	Zilbrysq	injection	2. Approve if the patient has already been started on therapy with Zilbrysq.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
			If either Soliris or Epysqli are formulary:  Approve if the patient meets BOTH of the following (A and B):  A. Patient has tried both of Soliris and Epysqli, if formulary; AND  B. Patient cannot continue to use all the formulary eculizumab products due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.  If neither Soliris nor Epysqli are formulary:  Approve if the patient meets ONE of the following (1, 2, 3, or 4):  1. Atypical hemolytic uremic syndrome: Approve.  2. Anti-acetylcholine receptor antibody positive generalized myasthenia gravis in a patient ≥ 6 years of age.  Approve if the patient meets one of the following (A, B, or C):  A. Patient meets ONE of the following (i or ii):		
			i. Patient is < 18 years of age, approve if the patient has tried, and according to the prescriber, experienced inadequate efficacy or significant intolerance with Imaavy, if formulary. If Imaavy is non-formulary, approve; OR ii. Patient is < 12 years of age; OR  B. Patient meets BOTH of the following (i and ii): i. Patient has tried, and according to the prescriber, experienced inadequate efficacy or significant intolerance with one of Ultomiris or Zilbrysq, if formulary; AND  Note: If neither are formulary, would still need to meet criterion ii. ii. Patient has tried, and according to the prescriber, experienced inadequate efficacy or significant intolerance with one of Imaavy, Vyvgart Intravenous, Vyvgart Hytrulo, or Rystiggo, if formulary; OR  Note: If none are formulary, would still need to meet criterion i. Note: If none are formulary alternatives from criterion i or ii, approve.  C. Patient has already been started on therapy with an eculizumab product (Soliris, Bkemy, Epysgli).		
Complement			<ul> <li>3. Paroxysmal nocturnal hemoglobinuria in a patient ≥ 18 years of age: Approve if the patient meets one of the following (A or B):         <ul> <li>A. Patient has tried, and according to the prescriber, experienced inadequate efficacy or significant intolerance with one of Ultomiris IV or PiaSky, if formulary. If neither are formulary, approve; OR</li> <li>B. Patient has already been started on therapy with an eculizumab product (Soliris, Bkemv, Epysqli).</li> </ul> </li> <li>4. Anti-aquaporin (AQ4P) antibody-positive Neuromyelitis optica spectrum disorder in a patient ≥ 18 years of age. Approve if the patient meets one of the following (A or B):         <ul> <li>A. The patient has tried, and according to the prescriber, experienced inadequate efficacy or significant intolerance to BOTH Uplizna and Enspryng, if formulary, would need to try one or if neither are formulary,</li> </ul> </li> </ul>		
Inhibitors – Eculizumab Products	Bkemy	eculizumab-aeeb intravenous infusion	approve; OR  B. The patient has already been started on therapy with an eculizumab product (Soliris, Bkemv, Epysqli).	1 vear	Yes
Constipation Agents	Relistor	methylnaltrexone bromide tablets	Approve if the patient has tried two products from the following list [documentation required]: Movantik, Symproic, or Amitiza (lubiprostone), if two are formulary or one if one is formulary. If none are formulary, approve if the patient has tried two laxative agents (e.g., bisacodyl-containing products, senna-containing products, milk of magnesia, lactulose).	1 year	Yes
Constipation Agents – Chronic Idiopathic Constipation Agents	Motegrity	prucalopride tablets	Patient ≥ 18 years of age.  Approve if the patient has tried, and according to the prescriber, experienced inadequate efficacy OR significant intolerance with BOTH products from the following list: Linzess AND Trulance [documentation required], if two are formulary or one if one is formulary. If neither are formulary, approve.	1 year	Yes
Constipation Agents – Chronic Idiopathic Constipation Agents/Irritable Bowel Syndrome	I Amitiza	lubiprostone capsules	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Constipation Agents – Irritable Bowel Syndrome	Ibsrela	tenapanor tablets	Patient ≥ 18 years of age.  Approve if the patient has tried, and according to the prescriber, experienced inadequate efficacy OR significant intolerance with BOTH products from the following list: Linzess AND Trulance, if two are formulary or one if one is formulary. If neither are formulary, approve.	1 year	Yes

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
Contraceptives	NuvaRing	etonogestrel/ethinyl	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve if the patient meets one of the following criteria (i or ii):  i. The requested brand non-formulary drug is being prescribed primarily for the prevention of pregnancy AND, according to the prescriber, the brand product is being requested because the preferred bioequivalent generic product would not be as medically appropriate for the patient as the requested brand non-formulary drug is being prescribed for a use OTHER THAN primarily for the prevention of pregnancy AND the Brand product is being requested due to a formulation difference in the inactive ingredient(s) and the prescriber and th	1 year	MSB Exclusion *This criteria applies only to the NPF
Опивориче	- Curaning	L-lactic acid, citric	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required. Approve if the patient has tried or is unable to tolerate THREE other barrier methods of contraception, such as diaphragms, condoms, spermicides (over-the-counter), or sponges.  Note: Examples include, but may not be limited to, Caya contoured diaphragm, condom, FC2 Female Condom, FemCap, Gynol II contraceptive gel, VCF contraceptive gel.  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve if the patient meets one of the following (i or ii):  i. Patient has tried or is unable to tolerate THREE other barrier methods of contraception, such as diaphragms, condoms, spermicides (over-the-counter), or sponges; OR	ryou	uio ini i
Contraceptives	Phexx	acid, and potassium bitartrate vaginal gel	Note: Examples include, but may not be limited to, Caya contoured diaphragm, condom, FC2 Female Condom, FemCap, Gynol II contraceptive gel, VCF contraceptive gel.  ii. The requested non-formulary drug is being prescribed primarily for the prevention of pregnancy and other barrier methods of contraception would not be as medically appropriate for the patient as the requested non-formulary drug.	1 vear	Yes
Contraceptives	Twirla	levonorgestrel and ethinyl estradiol transdermal system	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  Approve if the patient has tried five other contraceptive agents (e.g., oral contraceptive tablets, Xulane [contraceptive patch], Annovera [contraceptive ring]), NuvaRing or generics [contraceptive ring]).  Note: Examples include, but may not be limited to, Blisovi Fe, Eluryng, etonogestrel-ethinyl estradiol vaginal ring, Hailey Fe, Junel Fe, Larin Fe, Xulane.  Note: A trial of five different oral contraceptive agents would meet the requirement.  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve if the patient meets one of the following (i or ii):  i. The patient has tried five other contraceptive agents (e.g., oral contraceptive tablets, Xulane [contraceptive patch], Annovera [contraceptive ring]), NuvaRing or generics [contraceptive ring]); OR  Note: Examples include, but may not be limited to, Blisovi Fe, Eluryng, etonogestrel-ethinyl estradiol vaginal ring, Hailey Fe, Junel Fe, Larin Fe, Xulane.  Note: A trial of five different oral contraceptive agents would meet the requirement.	1 year	Yes
Contraceptives – Oral		drospirenone tablet	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required. Approve if the patient has tried one progesterone-only contraceptives containing norethindrone include Camila, Deblitane, Emzahh, Errin, Nora-BE, norethindrone, Heather, Jencycla, Lyza, Sharobel, Tulana, Lyleq, Incassia.  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve if the patient meets one of the following criteria (i or ii):  i. Patient has tried one progesterone-only contraceptive containing norethindrone; OR  Note: Examples of progesterone-only contraceptives containing norethindrone include Camila, Deblitane, Emzahh, Errin, Nora-BE, norethindrone, Heather, Jencycla, Lyza, Sharobel, Tulana, Lyleq, Incassia.  ii. The requested non-formulary drug is being prescribed primarily for the prevention of pregnancy and other progesterone-only contraceptives containing norethindrone would not be as medically appropriate for the patient as the requested non-formulary drug.	1 year	Yes

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
Contraceptives – Oral	Balcoltra	ethinyl estradiol 0.02 mg; levonorgestrel 0.1 mg; ferrous bisglycinate tablet	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve if the patient meets one of the following criteria (i or ii):  i. The requested brand non-formulary drug is being prescribed primarily for the prevention of pregnancy AND, according to the prescriber, the brand product is being requested because the preferred bioequivalent generic product would not be as medically appropriate for the patient as the requested brand non-formulary drug is being prescribed for a use OTHER THAN primarily for the prevention of pregnancy AND the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Contraceptives – Oral	Loestrin and Loestrin FE	ethinyl estradiol/norethindron e and ferrous fumarate tablets	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve if the patient meets one of the following criteria (i or ii):  i. The requested brand non-formulary drug is being prescribed primarily for the prevention of pregnancy AND, according to the prescriber, the brand product is being requested because the preferred bioequivalent generic product would not be as medically appropriate for the patient as the requested brand non-formulary drug is being prescribed for a use OTHER THAN primarily for the prevention of pregnancy AND the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Contraceptives – Oral	Minastrin 24 FE	norethindrone - ethinyl estradiol - iron chewable tablets	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve if the patient meets one of the following criteria (i or ii):  i. The requested brand non-formulary drug is being prescribed primarily for the prevention of pregnancy AND, according to the prescriber, the brand product is being requested because the preferred bioequivalent generic product would not be as medically appropriate for the patient as the requested brand non-formulary drug is being prescribed for a use OTHER THAN primarily for the prevention of pregnancy AND the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Contraceptives – Oral	Quartette	levonorgestrel-ethinyl estradiol and ethinyl estradiol tablets	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve if the patient meets one of the following criteria (i or ii):  i. The requested brand non-formulary drug is being prescribed primarily for the prevention of pregnancy AND, according to the prescriber, the brand product is being requested because the preferred bioequivalent generic product would not be as medically appropriate for the patient as the requested brand non-formulary drug is being prescribed for a use OTHER THAN primarily for the prevention of pregnancy AND the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
		drospirenone/ethinyl estradiol-levomefolate	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve if the patient meets one of the following criteria (i or ii):  i. The requested brand non-formulary drug is being prescribed primarily for the prevention of pregnancy AND, according to the prescriber, the brand product is being requested because the preferred bioequivalent generic product would not be as medically appropriate for the patient as the requested brand non-formulary drug is being prescribed for a use OTHER THAN primarily for the prevention of pregnancy AND the Brand product is being requested due to a formulation difference in the inactive ingredient(s)		MSB Exclusion *This criteria applies only to
Contraceptives – Oral	Safyral	tablets	[e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve if the patient meets one of the following criteria (i or ii):  i. The requested brand non-formulary drug is being prescribed primarily for the prevention of pregnancy AND, according to the prescriber, the brand product is being requested because the preferred bioequivalent generic product	1 year	MSB Exclusion
Contraceptives – Oral	Seasonique	levonorgestrel-ethinyl estradiol and ethinyl estradiol tablets	would not be as medically appropriate for the patient as the requested brand non-formulary drug; OR  ii. The requested brand non-formulary drug is being prescribed for a use OTHER THAN primarily for the prevention of pregnancy AND the Brand product is being requested due to a formulation difference in the inactive ingredient(s)  [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  OR	1 year	*This criteria applies only to the NPF
Contraceptives – Oral	Taytulla	norethindrone and ethinyl estradiol and ferrous fumarate capsules	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve if the patient meets one of the following criteria (i or ii):  i. The requested brand non-formulary drug is being prescribed primarily for the prevention of pregnancy AND, according to the prescriber, the brand product is being requested because the preferred bioequivalent generic product would not be as medically appropriate for the patient as the requested brand non-formulary drug; OR  ii. The requested brand non-formulary drug is being prescribed for a use OTHER THAN primarily for the prevention of pregnancy AND the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
		ethinyl estradial/	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve if the patient meets one of the following criteria (i or ii):  i. The requested brand non-formulary drug is being prescribed primarily for the prevention of pregnancy AND, according to the prescriber, the brand product is being requested because the preferred bioequivalent generic product would not be as medically appropriate for the patient as the requested brand non-formulary drug is being prescribed for a use OTHER THAN primarily for the prevention of pregnancy AND the Brand product is being requested due to a formulation difference in the inactive ingredient(s).		MSB Exclusion *This criteria
Contraceptives – Oral	Yasmin	ethinyl estradiol/ drospirenone tablets	ii. The requested brand non-formulary drug is being prescribed for a use OTHER THAN primarily for the prevention of pregnancy AND the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	applies only to the NPF

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
			Compliance with the Affordable Care Act. HRSA Guidelines, and PHS Act section 2713 is NOT required.		
			Approve if the patient has tried two other oral contraceptive agents.		
			Note: Examples include, but may not be limited to, Hailey Fe, Junel Fe, Larin Fe, Mibelas 24 Fe, Microgestin Fe, norethindrone-ethinyl estradiol-iron.		
			OR		
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.		
		ethinyl estradiol 0.01	Approve if the patient meets one of the following criteria (i or ii):		
		mg; norethindrone acetate 1 mg; ferrous	i. Patient has tried two other oral contraceptive agents; OR  Note: Examples include, but may not be limited to, Hailey Fe, Junel Fe, Larin Fe, Mibelas 24 Fe, Microgestin Fe, norethindrone-ethinyl estradiol-iron.		
Contraceptives - Oral	Lo Loestrin FE	fumarate tablet	ii. The requested non-formulary drug is being prescribed primarily for the prevention of pregnancy and other oral contraceptive agents would not be as medically appropriate for the patient as the requested non-formulary drug.	1 year	Yes
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  Approve if the patient has tried four other oral contraceptive agents.		
			Note: Examples include, but may not be limited to, Blisovi Fe, drospirenone-ethinyl estradiol, Estarylla, Junel Fe, Tri-Sprintec, Sprintec.		
			OR .		
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve if the patient meets one of the following criteria (i or ii):		
			i. Patient has tried four other oral contraceptive agents; OR		
Contraceptives – Oral	Natazia	dienogest; estradiol valerate tablet	Note: Examples include, but may not be limited to, Blisovi Fe, drospirenone-ethinyl estradiol, Estarylla, Junel Fe, Tri-Sprintec, Sprintec.  ii. The requested non-formulary drug is being prescribed primarily for the prevention of pregnancy and other oral contraceptive agents would not be as medically appropriate for the patient as the requested non-formulary drug.	1 year	Yes
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  Approve if the patient has tried four other oral contraceptive agents.		
			Note: Examples include, but may not be limited to, Aurovela Fe, Blisovi Fe, drospirenone-ethinyl estradiol, Estarylla, Junel Fe, Tri-Sprintec, Sprintec.		
			OR		
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve if the patient meets one of the following criteria (i or ii):		
			i. Patient has tried four other oral contraceptive agents; OR		
Contraceptives – Oral	Nextstellis	estetrol and drospirenone tablets	Note: Examples include, but may not be limited to, Aurovela Fe, Blisovi Fe, drospirenone-ethinyl estradiol, Estarylla, Junel Fe, Tri-Sprintec, Sprintec.  ii. The requested non-formulary drug is being prescribed primarily for the prevention of pregnancy and other oral contraceptive agents would not be as medically appropriate for the patient as the requested non-formulary drug.	1 year	Yes
	. Noxuotomo	ar oopii onono tabioto	in the requestion for the protein of	. you.	
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.		
			Approve if the patient has tried four other oral contraceptive agents.  Note: Examples include, but may not be limited to, Altavera, Aviane, Falmina, Lessina, levonorgestrel-ethinyl estradiol, Portia, Vienva.		
			OR CONTRACTOR OF THE PROPERTY		
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.		
		levonorgestrel 0.1 mg	Approve if the patient meets one of the following criteria (i or ii):  i. Patient has tried four other oral contraceptive agents; OR		
		and ethinyl estradiol	Note: Examples include, but may not be limited to, Altavera, Aviane, Falmina, Lessina, levonorgestrel-ethinyl estradiol, Portia, Vienva.		
Contraceptives – Oral	Tyblume	0.02 mg tablets	ii. The requested non-formulary drug is being prescribed primarily for the prevention of pregnancy and other oral contraceptive agents would not be as medically appropriate for the patient as the requested non-formulary drug.	1 year	Yes

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  1. Approve if the patient has tried four other oral contraceptive agents.		
			Note: Examples include, but may not be limited to, Charlotte 24 Fe, Finzala, Kaitlib Fe, Layolis Fe, Mibelas 24 Fe, norethindrone-ethinyl estradiol, Wymzya Fe.		
			2. If the patient is unable to swallow tablets or has difficulty swallowing tablets, approve if the patient has tried one oral chewable birth control product (e.g., Finzala, Mibelas, Charlotte, Wymzya, Kaitlib, Layolis).		
			OR .		
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.		
		norethindrone acetate and ethinyl estradiol	1. Approve if the patient has tried four other oral contraceptive agents.  Note: Examples include, but may not be limited to, Charlotte 24 Fe, Finzala, Kaitlib Fe, Layolis Fe, Mibelas 24 Fe, norethindrone-ethinyl estradiol, Wymzya Fe.		
		orally disintegrating	2. If the patient is unable to swallow tablets or has difficulty swallowing tablets, approve if the patient has tried one oral chewable birth control product (e.g., Finzala, Mibelas, Charlotte, Wymzya, Kaitlib, Layolis).		
Contraceptives – Oral	Femlyv	tablets	3. The requested non-formulary drug is being prescribed primarily for the prevention of pregnancy and other oral contraceptive agents would not be as medically appropriate for the patient as the requested non-formulary drug.	1 year	Yes
			Patient meets Gastroenterology – Eohilia Prior Authorization Policy AND		
Corticosteroid (oral) -			Patient meets ONE of the following (1 or 2):		
Eosinophilic		budesonide oral	1. Approve if the patient has tried budesonide inhalation suspension (Pulmicort Respules, generic) that was made into a slurry or suspension and swallowed (not inhaled).		
Esophagitis Agent	Eohilia	suspension	2. Approve if the patient has already been started on a 12-week course of therapy with Eohilia (to allow for completion of up to a 12-week course of therapy).	12 weeks	Yes
		hydrocortisone oral	<ol> <li>Approve if the patient has tried and cannot take hydrocortisone tablets.</li> <li>Approve if the patient cannot swallow or has difficulty swallowing hydrocortisone tablets.</li> </ol>		
Corticosteroids (Oral)	Alkindi Sprinkle	granules	3. Approve if the prescribed dose cannot be obtained using whole hydrocortisone tablets.	1 year	Yes
0 11 1 11 10 10		dexamethasone 20			.,
Corticosteroids (Oral)	Hemady	mg tablets	Approve if the patient has tried generic dexamethasone tablets, if formulary. If dexamethasone tablets are non-formulary, approve.	1 year	Yes Yes
					*This criteria
Corticosteroids	Anusol-HC	hydrocortisone			applies only to
(Rectal Formulations)	suppository	acetate suppository	Approve if the patient has tried hydrocortisone acetate suppositories. If hydrocortisone acetate suppositories are non-formulary, approve.	1 year	the NPF
	Hydrocortisone-	hydrocortisone-			
Corticosteroids	pramoxine	pramoxine			
(Rectal Formulations)	suppository	suppository 25-18 mg	Approve if the patient has tried one of 1) hydrocortisone acetate suppositories or 2) a rectal topical product containing hydrocortisone and pramoxine (e.g., topical foam, topical cream).	1 year	Yes
		pramoxine			
		hydrochloride			
Corticosteroids	Description of LIC	hydrocortisone	Annual if the national beautiful and according to the annual bank and according to the second independent of the second in	4	V
(Rectal Formulations)	Proctoloam-HC	acetate aerosoi, ioam	Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with pramoxine-hydrocortisone cream.	1 year	Yes
			1. Approve if the patient has tried budesonide foam (Uceris foam, generics), if formulary. If budesonide foam (Uceris foam, generics) are non-formulary, approve if the patient has tried one corticosteroid enema from the following list (if		
Corticosteroids	0	hydrocortisone	one is formulary): Cortenema or hydrocortisone enema. If neither are formulary, approve.	4	
(Rectal Formulations)	Cortifoam	acetate aerosol toam	2. Patients who are unable to retain a corticosteroid enema: approve if the patient has tried budesonide foam (Uceris foam, generics), if formulary. If budesonide foam (Uceris foam, generics) are non-formulary, approve.	1 year	Yes MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Corticosteroids		hydrocortisone	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
(Topical)	Anusol-HC cream	acetate cream hydrocortisone	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF MSB Exclusion
		butyrate cream,	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Corticosteroids		lotion, ointment,	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
(Topical)	Locoid	solution	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Corticosteroids		hydrocortisone	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
(Topical)	Locoid Lipocream	butyrate 0.1% cream	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		MSB Exclusion *This criteria
Corticosteroids		desoximetasone	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
(Topical)	Topicort spray	sprav	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF

					2025 NPF
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	Excluded Medicaiton
Corticosteroids (Topical)	Vanos	fluocinonide 0.1% cream	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Corticosteroids (Topical)	Impoyz (and authorized generic)	clobetasol propionate cream, 0.025%	Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with five unique, generic prescription-strength topical corticosteroid products. <u>Note</u> : Examples of topical steroid products include: desoximetasone, triamcinolone, betamethasone, clobetasol, fluocinonide, mometasone, halcinonide, diflorasone. <u>NOTE</u> : The products must be chemically unique.	1 year	Yes
Corticosteroids (Topical)	Sernivo spray	dipropionate spray	Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with five unique, generic prescription-strength topical corticosteroid products.  Note: Examples of topical steroid products include: desoximetasone, triamcinolone, betamethasone, clobetasol, fluocinonide, mometasone, halcinonide.  NOTE: The five products must be chemically unique.	1 year	Yes
Corticosteroids (Topical)	Verdeso		Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with five unique, generic prescription-strength topical corticosteroid products.  Note: Examples of topical steroid products include: desonide, alclometasone dipropionate, betamethasone valerate, fluocinolone acetonide, triamcinolone, flurandrenolide, hydrocortisone butyrate.  NOTE: The five products must be chemically unique (i.e., a trial of desoximetasone 0.05% and 0.25% would NOT fulfill the requirement).	1 year	Yes
Corticosteroids (Topical)	clocortolone 0.1% cream pump; clocortolone pivalate 0.1% cream	clocortolone pivalate 0.1% cream	Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with three unique, generic prescription-strength topical corticosteroid products.  Note: Examples of topical steroid products include: betamethasone, fluocinolone acetonide, hydrocortisone valerate, mometasone, triamcinolone acetonide.	1 year	Yes
Corticosteroids (Topical)	diflorasone 0.05% ointment and 0.05% cream; Apexicon E 0.05% cream	diflorasone 0.05% ointment and 0.05% cream	Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with three unique, generic prescription-strength topical corticosteroid products.  Note: Examples of topical steroid products include: betamethasone, triamcinolone, fluocinolone, clobetasol, fluocinonide, amcinonide, mometasone furoate, fluticasone.	1 year	Yes
Corticosteroids (Topical)	flurandrenolide 0.05% cream; flurandrenolide 0.05% ointment; flurandrenolide 0.05% lotion	flurandrenolide 0.05% cream; flurandrenolide 0.05% ointment; flurandrenolide 0.05% lotion	Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with three unique, generic prescription-strength topical corticosteroid products.  Note: Examples of topical steroid products include: betamethasone, desoximetasone, triamcinolone, fluocinolone, mometasone, fluticasone.	1 year	Yes
Corticosteroids (Topical)	halcinonide 0.1% cream; halcinonide 0.1% solution	halcinonide 0.1% cream; halcinonide 0.1% solution	Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with three unique, generic prescription-strength topical corticosteroid products.  Note: Examples of topical steroid products include: betamethasone, clobetasol, fluocinonide, amcinonide, desoximetasone, mometasone furoate.	1 year	Yes
			Cushing's Disease in a patient ≥ 18 years of age.  Approve if the patient meets one of the following (A or B):  A. Patient has tried, or is currently taking, one of Signifor or Signifor LAR. If neither are formulary, approve; OR  B. Patient has already been started on Isturisa.  Endogenous Cushing's Syndrome in a patient ≥ 18 years of age.  Note: This includes patients awaiting surgery and patients awaiting therapeutic response after pituitary radiotherapy.  Approve if the patient meets one of the following (A or B):  A. Patient has tried, or is currently taking, ONE of ketoconazole, Metopirone (metyrapone capsules), or Recorlev. If ketoconazole is non-formulary, approve; OR  B. Patient has already been started on Isturisa.		
Cushing's - Cortisol Synthesis Inhibitor	Isturisa	osilodrostat tablets	Note: A trial of any of the products above would count toward meeting criteria regardless of the formulary status of the product.  Endogenous Cushing's Syndrome in a patient ≥ 18 years of age.	1 year	Yes
Cushing's - Cortisol Synthesis Inhibitor	Recorlev	levoketoconazole tablets	Note: If Isturisa is non-formulary, would still need to try ketoconazole.  In the patient has tried would still need to try ketoconazole.	1 year	Yes
Cushing's -Cortisol Receptor Blocker	Korlym	mifepristone 300 mg tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Cystinuria Agents	Thiola	tiopronin tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Diabetes - Oral Sulfonylurea	Glimepiride 3 mg (brand)	glimepiride 3 mg	Approve if the patient cannot use one of the following: generic glimepiride 1mg, 2mg, or 4 mg. If generic glimepiride is non-formulary, approve.	1 year	Yes
Culloriylarda	(brand)	giiiiopiiido o nig	Approve if the patient has tried Janumet XR, if formulary.	1 your	100
Diabetes Agents - Dipeptidyl Peptidase- 4 (DPP-4) Inhibitor Combination Products	Zituvimet XR and	sitagliptin and metformin hydrochloride extended-release tablets	If Janumet XR is non-formulary, approve if the patient meets ONE of the following (1 or 2):  1. Approve if the patient has tried three metformin-DPP-4 inhibitor combination products from the following list (if three are formulary, or two if two are formulary): Janumet (NOT XR), alogliptin and metformin tablets, Jentadueto, Jentadueto, Jentadueto XR, Kazano, or saxagliptin plus metformin extended-release tablets (Kombiglyze XR, generics). If none are formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics). AND two of the following, if two are formulary (or one if only one is formulary): saxagliptin tablets (Onglyza, generics), Januvia, Tradjenta, or alogliptin tablets (Nesina, authorized generics). If none are formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics).  Note: Jentadueto and Jentadueto XR would count as one alternative. Alogliptin/metformin tablets and Kazano would count as one alternative.  2. Patients with a history of heart failure (HF) or renal impairment: approve if the patient has tried Janumet (NOT XR). If Janumet (NOT XR) is non-formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics).  Note: A brand product and its generic or authorized generic would count as one alternative.	1 year	Yes
			Approve if the patient has tried Janumet, if formulary.	,	
Diabetes Agents - Dipeptidyl Peptidase- 4 (DPP-4) Inhibitor Combination Products	hydrochloride tablets	sitagliptin and metformin hydrochloride tablets 50-1000; 50-500	If Janumet is non-formulary, approve if the patient meets ONE of the following (1 or 2):  1. Approve if the patient has tried three metformin-DPP-4 inhibitor combination products from the following list (if three are formulary, or two if two or formulary or one if one is formulary): alogliptin and metformin tablets, Janumet XR, Jentadueto, Jentadueto XR, Kazano, or saxagliptin plus metformin extended-release tablets (Kombiglyze XR, generics). If none are formulary, approve if the patient has tried metformin (Glucophage, Glucophage, Glucopha		Yes
Diabetes Agents - Dipeptidyl Peptidase- 4 (DPP-4) Inhibitor Combination Products	Oseni and	alogliptin and pioglitazone tablets	Approve if the patient has tried pioglitazone (Actos, generics) AND two of the following, if two are formulary (or one if only one is formulary): saxagliptin (Onglyza, generics), alogliptin tablets (Nesina, authorized generics), Tradjenta, or Januvia. If none are formulary, approve if the patient has tried pioglitazone (Actos, generics).  Note: A brand product and its generic or authorized generic would count as one alternative.  NOTE: A trial of Oseni or is authorized generic would not count toward this requirement.	1 year	Yes - Authorized generic only
Diabetes Agents - Dipeptidyl Peptidase- 4 (DPP-4) Inhibitor Combination Products		linagliptin and metformin tablets	1. Approve if the patient has tried three metformin-DPP-4 inhibitor combination products from the following list (if three are formulary, or two if two are formulary, or one if one is formulary): Jentadueto XR, alogliptin and metformin tablets, Janumet XR, Kazano, or saxagliptin plus metformin extended-release tablets (Kombiglyze XR, generics). If none are formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics) AND two of the following, if two are formulary (or one if only one is formulary): alogliptin tablets (Nesina, authorized generics), Tradjenta, saxagliptin tablets (Onglyza, generics), or Januvia. If none are formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics).  Note: Janumet and Janumet XR would count as one alternative. Alogliptin/metformin tablets and Kazano would count as one alternative.  Patients with a history of heart failure (HF) or renal impairment: approve if the patient has tried ONE of Jentadueto XR, Janumet or Janumet XR, if one is formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics) AND one of the following, if one is formulary: Januvia or Tradjenta. If neither are formulary, approve if the patient has tried metformin (Glucophage, Glucophage, Gluco	1 year	Yes
Diabetes Agents - Dipeptidyl Peptidase- 4 (DPP-4) Inhibitor Combination Products		linagliptin and metformin extended- release tablets	1. Approve if the patient has tried three metformin-DPP-4 inhibitor combination products from the following list: (if three are formulary, or two if two are formulary, or one if one is formulary): Jentadueto (NOT XR), alogliptin and metformin tablets, Janumet XR, Kazano, saxagliptin plus metformin extended-release tablets (Kombiglyze XR, generics). If none are formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics), or Januvia. If none are formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics).  Note: Janumet and Janumet XR would count as one alternative. Alogliptin/metformin tablets and Kazano would count as one alternative.  Patients with a history of heart failure or renal impairment: approve if the patient has tried one of Jentadueto (NOT XR), Janumet or Janumet XR, if formulary. If none are formulary, approve if the patient has tried metformin (Glucophage, Glucophage, Glucoph	1 year	Yes

					2025 NPF
				Approval	Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
Diabetes Agents - Dipeptidyl Peptidase- 4 (DPP-4) Inhibitor Combination Products	Kazano and authorized generic	alogliptin and metformin tablets	Approve if the patient has tried three metformin-DPP-4 inhibitor combination products from the following list (if three are formulary, or two if two are formulary, or one if one is formulary): Jentadueto, Jentadueto XR, Janumet, Janumet XR, or saxagliptin plus metformin extended-release tablets (Kombiglyze XR, generics). If none are formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics) AND two of the following, if two are formulary (or one if only one is formulary): saxagliptin tablets (Onglyza, generics), alogliptin tablets (Nesina, authorized generics), Tradjenta, or Januvia. If none are formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics).  Note: Jentadeuto and Jentadueto XR would count as one alternative. Janumet and Janumet XR would count as one alternative.	1 year	Yes
Diabetes Agents - Dipeptidyl Peptidase- 4 (DPP-4) Inhibitor Combination Products	Kombiglyze XR	saxagliptin plus metformin extended- release tablets	If requesting brand Kombiglyze XR: Approve if the patient has tried generic Kombiglyze XR tablets (saxagliptin plus metformin ER tablets), if formulary.  If requesting brand Kombiglyze XR and generic Kombiglyze XR tablets (saxagliptin plus metformin ER tablets) are non-formulary (or if requesting generic Kombiglyze), approve if the patient has tried three metformin-DPP-4 inhibitor combination products from the following list (if three are formulary, or two if two are formulary, or one if one is formulary): alogliptin and metformin tablets, Jentadueto XR, Kazano, Janumet, or Janumet XR. If none are formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics) AND two of the following, if two are formulary (or one if only one is formulary): alogliptin tablets (Nesina, authorized generics), saxagliptin tablets (Onglyza, generic), Tradjenta, or Januvia. If none are formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics).  Note: Jentadueto and Jentadueto XR would count as one alternative. A brand product and its generic or alternative.  A brand product and its generic or alternative.	1 year	Yes
Diabetes Agents - Dipeptidyl Peptidase- 4 (DPP-4) Inhibitor/ Sodium Glucose Co- Transporter-2 (SGLT- 2) Inhibitors	Steglujan	ertugliflozin/ sitagliptin tablets	<ol> <li>Approve if the patient has tried, and according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with both Qtern and Glyxambi, if formulary [documentation required]. If one is formulary, try one, if neither are formulary, approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with three formulary SGLT-2 inhibitors (or two if two are formulary or one if one is formulary) [documentation required].</li> <li>Patient with a history of heart failure or renal impairment: Approve if the patient has tried, and according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with Glyxambi, if formulary [documentation required]. If Glyxambi is not formulary, approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with one of Farxiga or Jardiance, if formulary [documentation required], AND one of Tradjenta or Januvia, if formulary [documentation required]. If Farxiga and Jardiance are both non-formulary, approve. If Tradjenta and Januvia are both non-formulary, approve.</li> <li>Note: SGLT-2 inhibitors: Brenzavvy, Farxiga, Invokana, Jardiance, Steglatro.</li> <li>DPP-4 inhibitors: Januvia, Nesina (alogliptin), Onglyza (saxagliptin), Tradjenta.</li> <li>Note: If the patient has tried a combination product containing the DPP-4 inhibitor or the SGLT-2 inhibitor, this would count as a trial of the respective product. A trial of the request agent would NOT count toward this requirement.</li> </ol>	1 year	Yes
Diabetes Agents - Dipeptidyl Peptidase- 4 (DPP-4) Inhibitor/ Sodium Glucose Co- Transporter-2 (SGLT- 2) Inhibitors	Qtern	dapagliflozin/ saxaqliptin tablets	Approve if the patient has tried, and according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with both Glyxambi and Steglujan, if formulary. If one is formulary, try one, if neither are formulary, approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with three formulary SGLT-2 inhibitors (or two if two are formulary or one if one is formulary).  SGLT-2 inhibitors: Brenzavvy, Farxiga, Invokana, Jardiance, Steglatro.  DPP-4 inhibitors: Januvia, Nesina (alogliptin), Onglyza (saxagliptin), Tradjenta.  Note: If the patient has tried a combination product containing a DPP-4 inhibitor or an SGLT-2 inhibitor, this would count as a trial of the respective product. A trial of the request agent would NOT count toward this requirement.	1 vear	Yes
Diabetes Agents - Dipeptidyl Peptidase- 4 (DPP-4) Inhibitors	Zituvio and authorized generic sitagliptin		Approve if the patient has tried Januvia, if formulary.  If Januvia is non-formulary, approve if the patient meets one of the following (1 or 2):  1. Approve if the patient has tried three products from the following list (if three are formulary or two if two are formulary, or one if only one is formulary): saxagliptin (Onglyza, generics), Tradjenta, or alogliptin tablets (Nesina, authorized generics). If none are formulary, approve.  Note: Saxagliptin and Onglyza count as one alternative. Alogliptin and Nesina count as one alternative.  2. Patients with a history of heart failure or a history of renal impairment: approve if the patient has tried Tradjenta, if formulary. If Tradjenta is non-formulary, approve.	1 year	Yes
Diabetes Agents - Dipeptidyl Peptidase- 4 (DPP-4) Inhibitors	Nesina and authorized generic	alogliptin tablets	Approve if the patient has tried three products from the following list (if three are formulary or two if two are formulary, or one if only one is formulary): saxagliptin (Onglyza, generics), Tradjenta, or a sitagliptin product (Januvia or Zituvio). If none are formulary, approve.  Note: Saxagliptin and Onglyza count as one alternative. Januvia and Zituvio would count as one alternative.	1 year	Yes
Diabetes Agents - Dipeptidyl Peptidase- 4 (DPP-4) Inhibitors	Onglyza	saxagliptin tablets	If requesting brand Onglyza: Approve if the patient has tried saxagliptin tablets (generic for Onglyza), if formulary.  If requesting brand Onglyza and generic saxagliptin tablets are non-formulary (or if requesting generic Onglyza), approve if the patient has tried three products from the following list (if three are formulary or two if two are formulary, or one if only one is formulary): alogliptin tablets (Nesina, authorized generics), Tradjenta, or a sitagliptin product (Januvia or Zituvio). If none are formulary, approve.  Note: Alogliptin and Nesina count as one alternative. Januvia and Zituvio count as one alternative.	1 year	Yes
Diabetes Agents - Dipeptidyl Peptidase- 4 (DPP-4) Inhibitors	Tradjenta	linagliptin tablets	<ol> <li>Approve if the patient has tried three products from the following list (if three are formulary or two if two are formulary, or one if only one is formulary): alogliptin tablets (Nesina, authorized generics), saxagliptin (Onglyza, generics), or a sitagliptin product (Januvia or Zituvio). If none are formulary, approve.</li> <li>Note: Alogliptin and Nesina count as one alternative. Saxagliptin and Onglyza count as one alternative.</li> <li>Patients with a history of heart failure or a history of renal impairment: Approve if the patient has tried a stiagliptin product (Januvia or Zituvio), if formulary. If neither Januvia nor Zituvio is formulary, approve.</li> </ol>	1 year	Yes

Dabetes Agents Chazgors I. Policy of Characteristic Special Section Medical Section of Section Medical Section of Section Medical Section of Section Medical Section Medical Section Section Medical Section M					Approval	2025 NPF Excluded
Cuckey-is Agents - Image of the common to th	Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
Lantus and Insuling dispring (by bringhouse)  Lantus and Insuling dispring (by Winthrop, AS, 20 decided to use Semglee (YFGN) or Insulin glargine (YFGN) plaufhorized generic of Semglee (YFGN)); in formulary. If formulary. If neither are formulary, approve.  1. Pallers is discided to use Semglee (YFGN) or Insulin glargine (YFGN) and a not construct the factor of the formulary status of these products. 1 year 1 ye	Glucagon-Like			If requesting brand Victoza:  Approve if the patient has tried and cannot take generic liraglutide due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product, if the generic is formulary.  If requesting generic liraglutide or generic liraglutide is non-formulary:		
Dabetes Agents - Insulin (glasgne Ly Commission (glasgne) (PCR) (and social color) (and s	Agonists	Victoza and generic	origin) injection	2. If the patient is less than 18 years of age, approve if the patient has tried Trulicity [documentation required], if formulary. If Trulicity is non-formulary, approve.	1 year	Yes - brand only
Diabetes Agents - Insulin (Basa)  Provided the patient has tried and cannot use Semigle (FCN) or insulin glargine (FCN) use in a flarging to a formulation difference in the inactive ingredient(s) [6.g., differences in stabilizing agent, buffering agent, and/or surfactant) which, listed the patient has been described by the patient has been desc		glargine (by Winthrop, A-S	vial and SoloStar	2. Approve if the patient has tried and cannot use Semglee (YFGN) or Insulin glargine (YFGN) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	Yes
are formulary, see, 1, 2, and 3 below.  Type 2 Diabetes (Initial user and a patient Currently Receiving Basaglar) [and all otheres]  1. If all the following products: Rezvoglar, Lantus, Semglee (YFGN), Insulin Glargine (YFGN) are non-formulary, approve if the patient meets the following (a and b): a. Patient has tried one of Tosibo or Insulin glargine U300, If formulary, AND b. Patient has tried one of Tosibo or Insulin glargine U300, If formulary slatus, that criterion would be satisfied.  Note: If the patient has tried one of Tosibo or Insulin glargine U300, If formulary, AND b. Patient has tried one of Tosibo or Insulin glargine U300, If formulary, AND b. Patient has tried one of Tosibo or Insulin glargine U300, If formulary, AND b. Patient has tried one of Tosibo or Insulin Degludce, If formulary, Note: If the patient has tried or Insulin Degludce, If formulary, Note: If the patient has tried or Insulin glargine U300, If formulary, and the patient has Type 1 diabetes, and the patient is currently taking Basaglar, approve if the patient has tried one of Tosibo  Insulin glargine U-100 or Insulin glargine U300, If formulary, If neither are formulary spardess of formulary status, the criterion would be satisfied.  Very Poliabetes, Continuation of Therapy with Basaglar, Insulin (Basal)  Basaglar  KWkPen Note: If the patient has tried or formulary status, the criterion would be satisfied.  1 year Yes  Approve if the patient has tried one of Tosipo or Insulin glargine U-100 or Insulin glargine U300, If formulary, If neither are formulary, spardess of formulary status, the criterion would be satisfied.  1 year Yes  Approve if the patient has tried dien or is using Basaglar Tempo Pen with the Tempo Smart Button, AND 2. Patient will use or is using Basaglar Tempo Pen with the Tempo Smart Button, AND 3. Patient will use or is using Basaglar Tempo Pen with the Tempo Smart Button, AND 4. Patient will use or is using Basaglar Tempo Pen with the Tempo Smart Button, AND 5. Patient will use or is using Basaglar Tempo Pen wi		Rezvoglar	insulin glargine-aglr	2. Approve if the patient has tried and cannot use Semglee (YFGN) or Insulin glargine (YFGN) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	Yes
Diabetes Agents - Insulin glargine U-100 or Insulin glargine U300, if formulary. If neither are formulary, approve.    Insulin (Basal)   Basaglar   KwikPen   Note: If the patient has tried either product above, regardless of formulary status, the criterion would be satisfied.    Approve if the patient meets the following (1, 2, and 3):   Approve if the patient mee				are formulary, see, 1, 2, and 3 below.  Type 2 Diabetes (Initial user and a patient Currently Receiving Basaglar) [and all others].  1. If all the following products: Rezvoglar, Lantus, Semglee (YFGN), Insulin Glargine (YFGN) are non-formulary, approve if the patient meets the following (a and b):  a. Patient has tried one of Toujeo or Insulin plargine U300, if formulary; AND  b. Patient has tried any product from a. or b. regardless of formulary status, that criterion would be satisfied.  Note: If the patient has tried any products in a or b, approve.  Type 1 Diabetes (initial user).  2. If the patient has Type 1 diabetes and all the following products: Rezvoglar, Lantus, Semglee (YFGN), Insulin Glargine (YFGN) are non-formulary, approve if the patient meets (a and b):  a. Patient has tried one of Toujeo or Insulin plargine U300, if formulary; AND  b. Patient has tried one of Tresiba or Insulin Degludec, if formulary.  Note: If the patient has tried any product from a. or b. regardless of formulary status, that criterion would be satisfied.  Note: If the patient has tried any product from a. or b. regardless of formulary status, that criterion would be satisfied.  Note: If there are no formulary products in a or b, approve.  Type 1 Diabetes, Continuation of Therapy with Basaglar.		
Approve if the patient meets the following (1, 2, and 3):  1. Patient will use or is using Basaglar Tempo Pen with the Tempo Smart Button; AND  2. Patient has tried a basal insulin pen; AND  Diabetes Agents - Insulin glargine U-100  3. Patient was unable to adhere to a regimen using a standard basal insulin pen, according to the prescriber [documentation required].	•		Insulin glargine U-100	or Insulin glargine U300, if formulary. If neither are formulary, approve.		
		Basaglar		Approve if the patient meets the following (1, 2, and 3):  1. Patient will use or is using Basaglar Tempo Pen with the Tempo Smart Button; AND  2. Patient has tried a basal insulin pen; AND	1 year	Yes
INSUITED DASARE TRANSPORT FEMORE PROTECTION FED. IN THE PROTECTION FED. IN MONTHS TYPES	Insulin (Basal)	Basaglar Tempo Per		Note: Document the specific issue(s) with adherence that would be solved by the use of a Tempo Pen.	6 months	Yes

				A	2025 NPF
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	Excluded Medicaiton
			Type 2 Diabetes (Initial user and a patient Currently Receiving Levemir); AND Type 1 Diabetes (Initial user) [and all others].		
			1. Approve if the patient meets the following (a and b):		
			a. Patient has tried one of Tresiba or Insulin Degludec, if formulary; AND		
			Note: If the patient has tried any product from a. regardless of formulary status, criterion a. would be satisfied.  b. Patient has tried one of Rezvoglar, Toujeo, Basaglar, Lantus, Insulin Glargine (YFGN), or Semglee (YFGN), if formulary.		
			Note: If the patient has tried any product from b. regardless of formulary status, criterion b. would be satisfied.		
			Note: If there are no formulary products in a or b, approve.		
			2. Patients < 6 years of age: approve if the patient has tried one of Tresiba or Insulin Degludec, if formulary. If neither are formulary, approve.		
			Note: If the patient has tried either product listed in 2. regardless of formulary status, criterion 2. would be satisfied.		
			3. Pregnant patients: approve.		
		insulin detemir U-100			
Diabetes Agents -	1	vial and FlexTouch	Type 1 Diabetes, Continuation of Therapy with Levemir.	4	
Insulin (Basal)	Levemir	pen	4. If the patient has Type 1 diabetes and is currently taking Levemir, approve.  1. Patient is directed to use Semglee (YFGN) [brand] or Insulin glargine-YFGN, if formulary.	1 year	Yes
Diabetes Agents -	Semglee (non	insulin glargine H-100	2. If neither are formulary, approve if the patient has tried one of Rezvoglar, Lantus, or Basaglar, if formulary. If Rezvoglar, Lantus, and Basaglar are non-formulary, approve.		
Insulin (Basal)	YFGN)	vial and pen	Note: If the patient has tried any product from 2. regardless of formulary status, criterion 2 would be satisfied.	1 year	Yes
			If Toujeo (brand) is non-formulary,  If Toujeo (brand) is non-formulary, approve if the patient meets (1, 2, 3 or 4):  Type 2 Diabetes, (initial user) OR taking Toujeo/Insulin glargine U300 < 100 Units/injection (all others taking < 100 units/injection).  1. Approve if the patient meets the following (a and b):  a. Patient has tried one of Tresiba or Insulin Degludec, if formulary; AND  Note: If the patient has tried one of Rezvogiar, Basagiar, Lantus, Semglee (YFGN), or Insulin glargine (YFGN), if formulary.  Note: If the patient has tried one of Rezvogiar, Basagiar, Lantus, Semglee (YFGN), or Insulin glargine (YFGN), if formulary.  Note: If the patient has tried one of Rezvogiar, Basagiar, Lantus, Semglee (YFGN), or Insulin glargine (YFGN), if formulary.  Note: If there are no formulary products in a or b, approve.  Type 2 Diabetes, Continuation of Therapy with Toujeo or Insulin glargine U300 ≥ 100 units per injection (and all others taking ≥ 100 units/injection).  2. Patients currently taking Toujeo or Insulin glargine U300 dose of ≥ 100 units per injection, approve if the patient has tried one of Tresiba U-200 or Insulin Degludec U-200, if formulary, approve.  Note: If the patient has tried either product listed in 2. regardless of formulary status, criterion 2 would be satisfied.  Note: A patient who has previously tried Tresiba U-100 is not required to try Tresiba U-200.  Type 1 Diabetes (initial user).  3. Patients with Type 1 diabetes-approve if the patient meets the following (a and b):  a. Patient has tried one of Tresiba or Insulin Degludec, if formulary; AND  Note: If the patient has tried one of Rezvogiar, Basagiar, Lantus, Semglee (YFGN), or Insulin Glargine (YFGN), if formulary.  Note: If the patient has tried one of Rezvogiar, Basagiar, Semglee (YFGN), or Insulin Glargine (YFGN), or Lantus, if formulary. If none are formulary, approve; OR		
Diabetes Agents -	Insulin glargine		Note: If the patient has tried any product from a. regardless of formulary status, criterion a would be satisfied.		
Insulin (Basal)	U300	SoloStar pen	b. Patient is currently receiving a Toujeo or Insulin glargine U300 dose of ≥ 100 units per injection.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
		insulin degludec vial	Patient is directed to use Tresiba (brand), if formulary.  If Tresiba (brand) is non-formulary, approve if the patient meets 1, 2, 3, or 4 below:  All patients < 6 years (Type 1, Type 2, all others).  1. Patients   Separate   Separ		
Diabetes Agents – Insulin (Basal)	Insulin Degludec	and FlexTouch pen U- 100 and U-200	- Type 1 Diabetes, Continuation of therapy with Insulin Degludec or Tresiba.  4. If the patient has Type 1 diabetes and is currently taking Tresiba or Insulin Degludec, approve.	1 vear	Yes
Diabetes Agents – Insulin (Basal) and Glucagon-Like Peptide-1 (GLP-1) Agonist Combination	Xultophy	insulin degludec/liraglutide injection	Approve if the patient has tried Soliqua, if formulary. If Soliqua is non-formulary, approve if the patient has tried two formulary basal insulins (if two are formulary or one if one is formulary): a glargine product (Basaglar, Lantus, Insulin Glargine [YFGN], Semglee [YFGN], Toujeo, Insulin glargine U300), or a degludec product (Tresiba or Insulin Degludec) AND three formulary glucagon-like peptide-1 (GLP-1) agonists (if three are formulary, or two if two are formulary or one if one is formulary): 1) exenatide injection, 2) Ozempic, 3) Trulicity, or 4) liraglutide (Victoza, generics). If none of the basal insulin products or none of the GLP-1 agonists are formulary, approve.  Note: Lantus, Insulin Glargine (YFGN), Semglee (YFGN), Basalgar, Toujeo, and Insulin glargine U300 would count as one alternative.  Note: Victoza and its generic would count as one alternative.  Note: A trial of Rybelsus or Mounjaro would also count as a trial of a GLP-1 agonist.  Note: If exenatide injection is formulary, a previous trial of Byetta or Bydureon BCise would satisfy a trial of exenatide injection.	1 year	Yes
Diabetes Agents - Insulin (Human)	Novolin 70/30 Flexpen and Relion Novolin 70/30 Flexpen	insulin, 70/30 pen	<ol> <li>Approve if the patient has tried Humulin 70/30 Kwikpens or Humulin 70/30 vials, if formulary. If both Humulin 70/30 Kwikpens and Humulin 70/30 vials are non-formulary, approve.</li> <li>If only Humulin 70/30 vials are formulary, approve in patients who are visually impaired, disabled (unable to draw up dose because of arthritis or otherwise physically disabled), or have coordination issues.</li> </ol>	1 year	Yes
Diabetes Agents - Insulin (Human)	Novolin 70/30 vials and Relion Novolin 70/30 vials	insulin, 70/30 vials	Approve if the patient has tried Humulin 70/30 vials or Humulin 70/30 Kwikpens, if formulary. If both Humulin 70/30 vials and Humulin 70/30 Kwikpens are non-formulary, approve.	1 year	Yes
Diabetes Agents - Insulin (Human)	Novolin N Flexpen and Relion Novolin N Flexpen Novolin N vials and	insulin, NPH pen	<ol> <li>Approve if the patient has tried Humulin N Kwikpens or Humulin N vials, if formulary. If both Humulin N Kwikpens and Humulin N vials are non-formulary, approve.</li> <li>If only Humulin N vials are formulary, approve in patients who are visually impaired, disabled (unable to draw up dose because of arthritis or otherwise physically disabled), or have coordination issues.</li> </ol>	1 year	Yes
Diabetes Agents - Insulin (Human)	Relion Novolin N	insulin, NPH vials	Approve if the patient has tried Humulin N vials or Humulin N Kwikpens, if formulary. If both Humulin N vials and Humulin N Kwikpens are non-formulary, approve.	1 year	Yes
Diabetes Agents - Insulin (Human)	Novolin R Flexpen and Relion Novolin R U-100 Flexpen	insulin, regular pen	<ol> <li>Approve if the patient has tried Humulin R U-100 vials, if formulary. If Humulin R U-100 vials are non-formulary, approve.</li> <li>Approve in patients who are visually impaired, disabled (unable to draw up dose because of arthritis or otherwise physically disable), or have coordination issues.</li> </ol>	1 year	Yes
Diabetes Agents - Insulin (Human)	Novolin R R U-100 vials and Relion Novolin R vials	insulin, regular vials	Approve if the patient has tried Humulin R U-100 vials, if formulary. If Humulin R U-100 vials are non-formulary, approve.	1 year	Yes

					2025 NPF
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	Excluded Medicaiton
			Direct the patient to Humalog JR (brand). If Humalog JR (brand) is non-formulary, approve if the patient meets the following (A or B):  A. Patient meets the following (i, ii, and iii):  i. Patient has tried Apidra, if formulary; AND  ii. Patient has tried one of the following, if formulary: Novolog, Insulin Aspart (authorized generic of Novolog), Fiasp, or Merilog; AND  Note: If the patient has tried any product from ii. regardless of formulary status, criterion ii would be satisfied.		
Diabetes Agents - Insulin (Rapid-Acting			iii. Patient has tried one of Admelog or Lyumjev, if formulary; OR  Note: If the patient has tried any product from iii. regardless of formulary status, criterion iii would be satisfied.  B. Patient requires ½ unit dosing.		
and Other)	Insulin Lispro JR	Insulin lispro JR	Note: If no products in A i, ii, or iii are formulary, approve.	1 year	Yes
Diabetes Agents - Insulin (Rapid-Acting		insulin aspart injection vial, pen, cartridge,	Approve if the patient meets one of the following (1 or 2):  1. Patient meets all of the following (A, B, and C):  A. Patient has tried Apidra, if formulary; AND  B. Patient has tried one of the following, if formulary: Insulin Lispro (authorized generic of Humalog), Humalog, Lyumjev, or Admelog; AND  Note: If the patient has tried any product from B. regardless of formulary status, criterion B would be satisfied.  C. Patient has tried one of the following, if formulary: NovoLog, Insulin Aspart (authorized generic of NovoLog), or Merilog; OR  Note: If the patient has tried any product from C. regardless of formulary status, criterion C would be satisfied.  Note: If no products in A, B, or C are formulary, approve.		
and Other)	Fiasp	PumpCart	2. Patient is using an insulin pump that is not compatible with the formulary alternative(s), approve.	1 year	Yes
Diabetes Agents - Insulin (Rapid-Acting and Other)	Afrezza	insulin human [rDNA origin] inhalation powder	Approve if the patient meets the following (A, B, and C):  A. Patient has tried Apidra, if formulary; AND  B. Patient has tried one of the following, if formulary: NovoLog, Insulin Aspart (authorized generic), Fiasp, or Merilog; AND  Note: If the patient has tried any product from B. regardless of formulary status, criterion B would be satisfied.  C. Patient has tried one of the following, if formulary: Insulin Lispro (authorized generic), Humalog, Lyumjev, or Admelog.  Note: If the patient has tried any product from C. regardless of formulary status, criterion C would be satisfied.  Note: If no products in A, B, or C are formulary, approve.	1 year	Yes
` '	Insulin Lispro Mix 75/25	75% Insulin lispro protamine/25% insulin lispro Kwikpen	Direct the patient is Humalog 75/25 (brand), if formulary. If Humalog 75/25 (brand) is non-formulary, approve if the patient has tried one of Novolog 70/30 or Insulin Aspart Protamine-Insulin Aspart Mix, if formulary. If neither are formulary, approve.	1 year	Voc
Diabetes Agents - Insulin (Rapid-Acting	NovoLog 70/30 and authorized generic (insulin aspart protamine-insulin aspart) and Relion Novolog 70/30	insulin aspart protamine/insulin aspart, Flexpen	Approve if the patient has tried Humalog 75/25, if formulary. If Humalog 75/25 is non-formulary, approve.	1 year	Yes
Diabetes Agents - Insulin (Rapid-Acting and Other)	,	insulin lispro vial, SoloStar (prefilled pen)	Approve if the patient meets one of the following (1 or 2):  1. Patient meets all of the following (A, B, and C):  A. Patient has tried Apidra, if formulary; AND  B. Patient has tried one of the following, if formulary: Insulin Lispro (authorized generic of Humalog), Humalog, or Lyumjev; AND  Note: If the patient has tried any product from B. regardless of formulary status, criterion B would be satisfied.  C. Patient has tried one of the following, if formulary: NovoLog, or Insulin Aspart (authorized generic of NovoLog), Fiasp, or Merilog; OR  Note: If the patient has tried any product from C. regardless of formulary status, criterion C would be satisfied.  Note: If no products in A, B, or C are formulary, approve.  2. Patient is using an insulin pump that is not compatible with the formulary alternative(s), approve.	1 year	Yes
Diabetes Agents - Insulin (Rapid-Acting	Apidra	insulin glulisine	Approve if the patient meets one of the following (1 or 2):  1. Patient meets both of the following (A and B):  A. Patient has tried one of the following, if formulary: Insulin Lispro (authorized generic of Humalog), Humalog, Lyumjev, or Admelog; AND  Note: If the patient has tried any product from A. regardless of formulary status, criterion A would be satisfied.  B. Patient has tried one of the following, if formulary: NovoLog or Insulin Aspart (authorized generic of NovoLog), Fiasp, or Merilog; OR  Note: If the patient has tried any product from B. regardless of formulary status, criterion B would be satisfied.  Note: If no products in A or B are formulary, approve.  2. Patient is using an insulin pump that is not compatible with the formulary alternative(s), approve.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
			If Merilog is formulary: Approve if the patient meets one of the following (1 or 2):  1. Approve if the patient meets BOTH of the following (A and B):  A. Patient has tried Merilog; AND  B. Patient cannot continue to use Merilog due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.  2. Patient is using an insulin pump.		
Diabetes Agents - Insulin (Rapid-Acting	NovoLog and authorized generic (insulin aspart) and	insulin aspart syringe, cartridge/Flexpen	Note: If no products in A, B, or C are formulary, approve.		
and Other)	Relion Novolog	(prefilled syringe)/vial	2. Patient is using an insulin pump that is not compatible with the formulary alternative(s), approve.	1 year	Yes
Diabetes Agents - Insulin (Rapid-Acting and Other)	Humalog	insulin lispro syringe, cartridge/Kwikpen/vial 100 units/mL, and	1. If the patient is requesting Humalog vial 100 units/mL or Humalog Kwikpen 100 units/mL, direct the patient to Insulin Lispro (authorized generic of Humalog), if formulary. If Insulin Lispro (authorized generic of Humalog) is non-formulary, then approve if the patient meets one of the following (i, ii, and iii):  i. Patient meets all of the following (i, ii, and iii): i. Patient has tried Apidra, if formulary: AND ii. Patient has tried one of the following, if formulary: NovoLog, Insulin Aspart (authorized generic of NovoLog), Fiasp, or Merilog; AND Note: If the patient has tried any product from ii. regardless of formulary status, criterion ii would be satisfied. iii. Patient has tried one of Admelog or Lyumjev, if formulary; OR Note: If the patient has tried any product from iii. regardless of formulary status, criterion iii would be satisfied. Note: If no products in i, ii, or iii are formulary, approve.  B. Patient is using an insulin pump that is not compatible with the formulary alternative(s), approve.  If the patient has tried any product from iii. regardless of formulary alternative(s), approve.  If the patient is requesting Humalog cartridge, or Humalog KwikPen U-200, approve if the patient has tried Insulin Lispro (authorized generic of Humalog), if formulary. If Insulin Lispro (authorized generic of Humalog) is non-formulary, then approve if the patient meets all of the following (A, B, and C):  A. Patient has tried Apidra, if formulary; NND  B. Patient has tried any product from B. regardless of formulary status, criterion B would be satisfied.  C. Patient has tried any product from C. regardless of formulary status, criterion C would be satisfied.  Note: If the patient has tried any product from C. regardless of formulary.	1 year	Yes - vial only
,		·			,
Diabetes Agents -	metformin immediate release	metformin immediate- release tablet 625 mg			
Other	625 mg and 750 mg	and 750 mg	Approve if the patient had inadequate efficacy OR significant intolerance with metformin 500 mg, 850 mg, or 1000 mg immediate-release tablets.	1 year	Yes
B: 1			Approve if the patient has tried BOTH one metformin immediate-release tablet product AND two other formulary metformin extended-release products (if two are formulary or one if one is formulary): metformin extended-release tablets		
Diabetes Agents - Other	Glumetza	metformin extended- release tablets	or Fortamet (brand or generic).  NOTE: A trial of Glumetza would NOT count toward this requirement.	1 year	Yes
Diabetes Agents - Sodium Glucose Co- Transporter-2 (SGLT- 2) Inhibitor			Approve if the patient has tried two formulary alternatives from the following list (or one if only one is formulary): Synjardy, Synjardy XR, or Xigduo XR. If none are formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics) AND three formulary alternatives from the following list, if formulary (or two if two are formulary or one if one is formulary: Farxiga, Steglatro, or Jardiance.		Voc
Combination Products	segiuromet	metformin tablets	Note: Synjardy and Synjardy XR would count as one alternative.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Diabetes Agents - Sodium Glucose Co- Transporter-2 (SGLT- 2) Inhibitor Combination Products	metformin ER	1 3	Direct to Xigduo XR (brand), if formulary.  If Xigduo XR (brand) is non-formulary, approve if the patient meets one of the following (1 or 2):  1. Approve if the patient has tried two formulary alternatives from the following list, if formulary (or one if one is formulary): Synjardy XR, or Segluromet. If none are formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics) AND three formulary alternatives from the following list, if formulary (or three if three are formulary or two if two are formulary or one if one is formulary): Farxiga, Jardiance, or Steglatro.  Note: Synjardy and Synjardy XR would count as one alternative.  2. Patients ≥ 10 years of age to < 18 years of age with type 2 diabetes mellitus: Approve if the patient has tried Synjardy, if formulary. If Synjardy is non-formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics) AND BOTH of 1) Farxiga or dapadiflozin tablet and 2) Jardiance, if formulary. If neither are formulary, approve.	1 year	Yes
Diabetes Agents - Sodium Glucose Co- Transporter-2 (SGLT- 2) Inhibitor Combination Products			1. Approve if the patient has tried four formulary alternatives from the following list, if formulary (or three if three are formulary or two if two are formulary or one if one is formulary): Invokamet XR, Synjardy XR, Segluromet, or Xigduo XR. If none are formulary, approve if the patient has tried metformin (Glucophage ER, generics) AND four formulary alternatives from the following list, if formulary (or three if three are formulary or two if two are formulary or one if only one is formulary): Farxiga, Invokana, Jardiance, or Steglatro.  Note: Synjardy and Synjardy XR would count as one alternative.  2. Patients ≥ 10 years of age to < 18 years of age with type 2 diabetes mellitus: Approve if the patient has tried BOTH of 1) Synjardy and 2) one of dapagliflozin-metformin ER tablets (authorized generic of Xigduo XR), or Xigduo XR (brand), if formulary. If none are formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics) AND BOTH of 1) Farxiga or dapagliflozin tablet and 2) Jardiance, if formulary. If neither are formulary, approve.	1 year	Yes
Diabetes Agents - Sodium Glucose Co- Transporter-2 (SGLT- 2) Inhibitor Combination Products	Invokamet XR	canagliflozin and metformin extended- release tablets	1. Approve if the patient has tried four formulary alternatives from the following list, if formulary (or three if three are formulary or two if two are formulary or one if one is formulary): Invokamet (not XR), Synjardy XR, Xigduo XR, or Segluromet. If none are formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics) AND four formulary alternatives from the following list, if formulary (or three if three are formulary or two if two are formulary or one if one is formulary): Farxiga, Invokana, Jardiance, or Steglatro.  Note: Synjardy and Synjardy XR would count as one alternative.  2. Patients ≥ 10 years of age to < 18 years of age with type 2 diabetes mellitus: Approve if the patient has tried BOTH of 1) Synjardy and 2) one of dapagliflozin-metformin ER tablets (authorized generic of Xigduo XR), or Xigduo XR (brand), if formulary. If none are formulary, approve if the patient has tried metformin (Glucophage, Glucophage ER, generics) AND BOTH of 1) Farxiga or dapagliflozin tablet and 2) Jardiance, if formulary. If neither are formulary, approve.	1 year	Yes
Diabetes Agents - Sodium Glucose Co- Transporter-2 (SGLT- 2) Inhibitors	dapagliflozin tablets (authorized generic of Farxiga)	dapagliflozin tablets	Direct to Farxiga (brand), if formulary.  If Farxiga (brand) is non-formulary:  Approve if the patient has tried, according to the prescriber, and experienced inadequate efficacy OR significant intolerance with Jardiance, if formulary. If Jardiance is non-formulary, approve.	1 year	Yes
Diabetes Agents - Sodium Glucose Co- Transporter-2 (SGLT- 2) Inhibitors	Inpefa	sotagliflozin tablets	Patients with one of the following: 1) Heart Failure OR 2) Type 2 diabetes, Chronic Kidney Disease (CKD), and Other cardiovascular (CV) risk factors.  Approve if the patient has tried, and according to the prescriber, experienced inadequate efficacy OR significant intolerance with BOTH Farxiga and Jardiance, if formulary (or one if one is formulary). If neither are formulary, approve.	1 year	Yes
Diabetes Agents - Sodium Glucose Co- Transporter-2 (SGLT- 2) Inhibitors	Steglatro	ertugliflozin tablets	Approve if the patient has tried, and according to the prescriber, experienced inadequate efficacy OR significant intolerance with BOTH products from the following list (or one if only one is formulary): Farxiga and Jardiance. If neither are formulary, approve.	1 year	Yes
Diabetes Agents - Sodium Glucose Co- Transporter-2 (SGLT- 2) Inhibitors	Brenzavvy	bexagliflozin tablets	1. Approve if the patient has tried BOTH Farxiga AND Jardiance, if both are formulary (or one if one is formulary). If neither are formulary, approve.  2. If the patient's estimated glomerular filtration rate is less than 45 mL/minute, approve if the patient has tried Jardiance, if formulary. If Jardiance is non-formulary, approve.	1 year	Yes
Diabetes Agents - Sodium Glucose Co- Transporter-2 (SGLT- 2) Inhibitors	Invokana	canagliflozin tablets	<ol> <li>Approve if the patient has tried, and according to the prescriber, experienced inadequate efficacy OR significant intolerance with BOTH products from the following list (or one if only one is formulary): Farxiga and Jardiance. If neither are formulary, approve.</li> <li>If Invokana is being used for glycemic control and the patient's estimated glomerular filtration rate is less than 45 mL/minute, approve if the patient has tried and, according to the prescriber, experienced inadequate efficacy OR significant intolerance with Jardiance, if formulary. If Jardiance is non-formulary, approve.</li> </ol>	1 year	Yes
Diabetes Agents – Sulfonylurea	glipizide 2.5 mg	glipizide 2.5 mg	Approve if the patient's prescribed dose cannot be obtained with glipizide 5 mg.  Note: The patient is NOT required to split the 5 mg tablets in half.	1 year	Yes

					2025 NPF
Th	B d N	O N	Ourself FE Office	Approval	Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
	Pen needles by Arkray, Home Aide	Pen needles by Arkray, Home Aide			
	Diagnostics, HTL-	Diagnostics, HTL-			
	Strefa, Nipro	Strefa, Nipro			
	Diagnostics, Novo	Diagnostics, Novo			
	Nordisk, Owen	Nordisk, Owen			
	Mumford, Simple	Mumford, Simple			
	Diagnostics,	Diagnostics, Ultimed,			
	Ultimed, all other	all other diabetic pen	1. Approve if the patient has tried one formulary pen needle. If none are formulary, approve.		
Diabetic Pen Needles		needles that are not BD	2. Approve if the prescriber states the patient requires a needle of the requested length and/or gauge which is not available as a formulary product.  Note: NPF prefers BD products.	1 11005	Voc
Diabetic Pen Needles	s that are not BD	BD	1. Approve if the patient has tried one formulary meter/test strip/control solution. If none are formulary, approve.	1 year	Yes
			2. Patients using an insulin pump/meter system that is not compatible with one of the available formulary alternatives: approve.		
			3. If the request is for Freestyle Precision Neo strips for use in a Freestyle Libre reader, approve.		
		Blood glucose	4. Patients who are blind or significantly visually impaired who are requesting a meter with audio capabilities. If there are no formulary meters with audio capabilities.		
		meters/test	capabilities, approve.		Yes - certain
		strips/control	Note: Meters with audio capabilities include Advocate (Redi-Code plus speaking meter), Arkray (Glucocard Expression, Glucocard Shine Express), Foracare (Fora D40D, Fora D40D, F		diabetic
Diabetic Supplies	Diabetic Supplies	solutions	Advance Voice, Fora Tn'G Voice, Fora V30), Oak Tree Health (EasyMax V, Fortiscare V3), Omnis Health (Embrace Talk), Prodigy (Prodigy Autocode, Prodigy Voice), Relion Premier Voice.	1 year	supplies
Diabetic Supplies – Continuous Glucose Monitoring Systems	[That are NOT Dexcom G6, Dexcom G7, Freestyle Libre 2, Freestyle Libre 2 Plus, Freestyle Libre	Other continuous glucose monitoring systems (receiver/reader, transmitter, sensor) [That are NOT Dexcom G6, Dexcom G7, or Freestyle Libre 2, Freestyle Libre 2 Plus, or Freestyle Libre 3 Freestyle Libre 3 Plus]	Note: If only a Freestyle Libre product is formulary and the patient has tried a different Freestyle Libre product (e.g., Freestyle Libre 10- or 14 day- product), approve.  Note: If only a Dexcom product is formulary and the patient has tried a different Dexcom product (e.g., Dexcom G4 or G5), approve.  If the patient is using an insulin pump system that is not compatible with one of the formulary alternatives: approve.  If requesting the Guardian Connect and the patient is pregnant: approve if the patient has tried BOTH of the following systems, if formulary: 1) Freestyle Libre 2, Freestyle Libre 2 Plus, Freestyle Libre 3, or Freestyle Libre 3 Plus AND 2) Dexcom G7 system. If none are formulary, approve.	1 year	Yes - Bigfoot Unity Program Kit, Eversense 365
			Approve if the patient meets the following (1, 2, and 3):		
			1. Patient will use or is using this product concomitantly with a Tempo Insulin Pen; AND		
District Committee		Tempo Lancets,	2. Patient has tried standard insulin products; AND		
Diabetic Supplies – Other	Tempo Refill Kit	strips, and Pen needles	3. Patient was unable to adhere to a regimen using standard insulin products, according to the prescriber [documentation required].  Note: Document the specific issue(s) with adherence that would be solved by the use of a Tempo product.	6 months	Vos
Outer	rempo Reilli Kit	needles	Approve if the patient meets the following (1, 2, and 3):	6 months	Yes
			1. Patient will use or is using this product concomitantly with a Tempo Insulin Pen; AND		
			2. Patient has tried standard insulin products; AND		
Diabetic Supplies -			3. Patient was unable to adhere to a regimen using standard insulin products, according to the prescriber [documentation required].		
Other	Tempo Smart Button	Tempo Smart Button	Note: Document the specific issue(s) with adherence that would be solved by the use of a Tempo product.	6 months	Yes
		Tempo Smart Button;			
		Tempo Blood	Approve if the patient meets the following (1, 2, and 3):		
		Glucose Monitoring	1. Patient will use or is using this product concomitantly with a Tempo Insulin Pen; AND		
		System, Lancets,	2. Patient has tried standard insulin products; AND		
		Strips, and Pen	3. Patient was unable to adhere to a regimen using standard insulin products, according to the prescriber [documentation required].		
Diabetic Supplies – Other	Tempo Welcome Kit		Note: Document the specific issue(s) with adherence that would be solved by the use of a Tempo product.	6 months	Yes

					2025 NPF
Th	Barried Marrie	Our and a Name	Our months FT O Work	Approval	Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
	Syringes by Arkray, Home Aide	Syringes by Arkray,			
	Diagnostics, HTL-	Home Aide			
	Strefa, Nipro	Diagnostics, HTL-			
	Diagnostics, Novo	Strefa, Nipro			
	Nordisk, Owen	Diagnostics, Novo			
	Mumford, Simple Diagnostics.	Nordisk, Owen Mumford, Simple			
	Ultimed, all other	Diagnostics, Ultimed,	1. Approve if the patient has tried one formulary syringe. If none are formulary, approve.		
	syringes that are not		2. Approve if the prescriber states the patient requires a needle of the requested length and/or gauge which is not available as a formulary product.		
	BD	are not BD	Note: NPF prefers BD products	1 year	Yes
, 0					MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Direct Renin Inhibitors	Tekturna	aliskiren tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Duchenne Muscular		delandistrogene	A CONTRACTOR OF THE CONTRACTOR		
Dystrophy (DMD) Agents	Elevidys	moxeparvovec-rokl intravenous infusion	No exceptions are recommended. The effectiveness of Elevidys has not been established at this time. (NOTE: It is not appropriate to use standard global criteria for this medication; Denial reason is: No exceptions are recommended. The effectiveness of Elevidys has not been established at this time.)	N/A	Yes
Duchenne Muscular	Elevidys	ilitiavellous illiusion	The enectiveness of Elevity's has not been established at this time.)	IN/A	165
Dystrophy (DMD)		casimersen	No exceptions are recommended. The effectiveness of Amondys 45 has not been established at this time. (NOTE: It is not appropriate to use standard global criteria for this medication; Denial reason is: No exceptions are		
Agents	Amondvs 45	intravenous	recommended. The effectiveness of Amondys 45 has not been established at this time.)	N/A	Yes
Duchenne Muscular	,				
Dystrophy (DMD)		eteplirsen injection for	No exceptions are recommended. The effectiveness of Exondys 51 has not been established at this time. (NOTE: It is not appropriate to use standard global criteria for this medication; Denial reason is: No exceptions are		
•	Exondys 51	intravenous use	recommended. The effectiveness of Exondys 51 has not been established at this time.)	N/A	Yes
Duchenne Muscular		viltolarsen injection			
Dystrophy (DMD)	\ m.	for intravenous	No exceptions are recommended. The effectiveness of Viltepso has not been established at this time. (NOTE: It is not appropriate to use standard global criteria for this medication; Denial reason is: No exceptions are recommended.		V
Agents	Viltepso	infusion	The effectiveness of Viltepso has not been established at this time.)	N/A	Yes
Duchenne Muscular Dystrophy (DMD)		golodirsen injection	No exceptions are recommended. The effectiveness of Vyondys 53 has not been established at this time. (NOTE: It is not appropriate to use standard global criteria for this medication; Denial reason is: No exceptions are		
	Vyondys 53	for intravenous use	recommended. The effectiveness of Vyondys 53 has not been established at this time. (NOTE: It is not appropriate to use standard global criteria for this inedication, benian leason is. No exceptions are	N/A	Yes
Duchenne Muscular	r you ayo oo	Tot militaronous dos			
Dystrophy (DMD)		vamorolone oral			
Agents	Agamree	suspension	See standard Muscular Dystrophy – Agamree Prior Authorization Policy criteria.	1 year	Yes
Duchenne Muscular	Brand Emflaza				
Dystrophy (DMD)	(tablets and oral	deflazacort tablets		See PSM	
Agents	suspension)	and oral suspension	See standard Muscular Dystrophy – Deflazacort Preferred Specialty Management Policy criteria.	duration	Yes
Duchenne Muscular					
Ducnenne Muscular Dystrophy (DMD)					
Agents - Histone		givinostat oral		See PA	
Deacetylase Inhibitor	Duvvzat	suspension	See standard Muscular Dystrophy – Duvyzat Prior Authorization Policy criteria	duration	Yes
,	,	repository			
		corticotropin			
Endocrine Drugs –		subcutaneous or			
	Cortrophin Gel	intramuscular	No exceptions are recommended. There is a lack of updated clinical efficacy data and potential safety concerns with long-term use. (NOTE: It is not appropriate to use standard global criteria for this medication; Denial reason is: No		
Corticotropin	(Purified)	injection	exceptions are recommended. There is a lack of updated clinical efficacy data and insufficient information to determine clinically meaningful benefits.)	N/A	Yes
			NOTE: A multipayers Drond product in being requested. The notices should use the professed biological state of the state o		MSB Exclusion
Endocrino Drugo			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		*This criteria applies only to
Endocrine Drugs - Miscellaneous	Samsca	tolvaptan tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
i i i i i i i i i i i i i i i i i i i	Camboa	torrapian tablets	production, model result in a significant anergy or solitons adverse reduction function required.	i your	MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Endocrine Drugs -			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Endothelin Receptor Antagonist	Tryvio	aprocitentan tablets	Approve if the patient has tried, or is currently receiving, at least three other antihypertensive agents for the treatment of hypertension from at least three of the following pharmacological classes [documentation required] (i, ii, iii, iv, v, vi, vii, vii, vi, x).  Note: A combination product from two or more different classes would count as an alternative from each class.  i. Anglotensin converting enzyme (ACE) inhibitor or angiotensin receptor blocker (ARB);  Note: Examples of ACE inhibitors include benazepril, captopril, enalapril, fosinopril, lisinopril, moexipril, perindopril, ramipril, and trandolapril. Examples of ARBs include azilsartan, candesartan, eprosartan, irbesartan, losartan, olimisartan, and valsartan.  ii. Non-dihydropyridine calcium channel blocker;  Note: Examples include dilitizarem and verapamil.  iii. Dihydropyridine calcium channel blocker;  Note: Examples include dilitizarem and verapamil.  iii. Dihydropyridine calcium channel blocker;  Note: Examples of thiazide diuretics include chlorthalidone, chlorothiazide, hydrochlorothiazide, indapamide, and metolazone. Examples of potassium-sparing diuretics are amilioride and triamterene.  v. Mineralocorticoid receptor antagonist;  Note: Examples of thiazide diuretics include acebutolol, atenolol, betaxolol, bispoprolol, carvedilol, metoprolol, nadolol, nebivolol, pindolol, propranolol, and timolol.  vii. Alpha-adrenergic blocker;  Note: Examples of elablockers include acebutolol, atenolol, betaxolol, bispoprolol, carvedilol, metoprolol, nadolol, nebivolol, pindolol, propranolol, and timolol.  viii. Central alpha-adrenergic agonists are clonidine, guanfacine, and methyldopa.  ix. Direct vasodilator;  Note: Examples of direct rein inhibitor is alliskiren.	1 year	Yes
Epinephrine Self- Administered Injectables	epinephrine auto-	epinephrine 0.15 mg, 0.3 mg auto-injector authorized generic (Amneal Pharmace, Avkare, A-S Medication)	Approve if the patient has tried one product from the following list, if one is formulary: epinephrine auto-injector (EpiPen/EpiPen Jr., generics). If none are formulary, approve.	1 vear	Yes
Erectile Dysfunction Agents - Phosphodiesterase Type 5 (PDE-5)	Cialis	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].		MSB Exclusion *This criteria applies only to the NPF
Erectile Dysfunction Agents - Phosphodiesterase Type 5 (PDE-5) Inhibitors	Viagra		NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Erythropoiesis- Stimulating Agents (ESAs)	Aranesp	darbepoetin alfa	Approve if the patient has tried one product from the following list: Epogen, Procrit or Retacrit [documentation required], if one is formulary. If none are formulary, approve.	1 year	Yes

					2025 NPF
The server Olever	Down d Massa	Oursella Name		Approval	Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
			Approve if the patient meets the following criteria (A <u>and</u> B):     A. Patient meets the following criteria (i and ii):		
			i. Patient has tried both products from the following list, if formulary (or one if only one is formulary): Procrit and Retacrit [documentation required]; AND		
			Note: If neither are formulary, would still need to meet criteria B, if Aranesp is formulary.		
			ii. Patient cannot continue to use each formulary alternative due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction; AND		
			B. Patient has tried Aranesp, if formulary [documentation required].		
			Note: If none of the following products are formulary: Aranesp, Procrit, and Retacrit, approve.		
			2. Pediatric patients with anemia due to cancer chemotherapy; Patients undergoing surgery requesting agent for the reduction of allogeneic red blood cell transfusion; Patients with anemia and human immunodeficiency virus (HIV)		
			infection who are receiving zidovudine:		
			Patient meets the following criteria (i and ii):    Patient has tried both products from the following list if formulars (or one if only one is formulars). Progrit and Retarit fide automated in AND.		
Erythropoiesis-			i. Patient has tried both products from the following list, if formulary (or one if only one is formulary): Procrit and Retacrit [documentation required]; AND Note: If neither are formulary, approve.		
Stimulating Agents			ii. Patient cannot continue to use each formulary alternative due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would		
(ESAs)	Epogen	epoetin alfa	result in a significant allergy or serious adverse reaction.	1 year	Yes
			Approve if the patient meets the following criteria (1 AND 2):		
			1. Patient has tried one epoetin alpha product from the following list, if formulary: Epogen, Procrit, or Retacrit [documentation required]; AND  Note: If none are formulary, would still need to meet criteria 2, if Aranesp is formulary.		
			2. Patient has tried Aranesp, if formulary [documentation required].		
Erythropoiesis-		methoxy polyethylene			
Stimulating Agents	M4:	glycol-epoetin beta	Note: The requirements are that one epoetin alpha product and Aranesp have been tried, if both are formulary. If only epoetin alpha product(s) is/are formulary and the patient has tried an epoetin alpha product, then the request should	4	V
(ESAs) Estrogen and	Mircera	solution for injection	be approved.	1 year	Yes MSB Exclusion
Estrogen			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Combination Products			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
(Topical)	Divigel	estradiol gel 0.1%	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Estrogen and Estrogen			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		MSB Exclusion *This criteria
Combination Products		estradiol transdermal	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
(Topical)	Minivelle	patch	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].		the NPF
Estrogen and			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		MSB Exclusion
Estrogen Combination Products		estradiol transdermal	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		*This criteria applies only to
(Topical)	Vivelle-Dot	patch	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Estrogen and					MSB Exclusion
Estrogen			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Combination Products (Topical)	Estrogel	estradiol gel 0.06%	<b>Criteria:</b> Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	applies only to the NPF
Estrogen and		garana	processes, weath read significant analysis of control for the control of the cont	, , ,	
Estrogen					
Combination Products (Topical)	Climara Pro	estradiol/ levonorgestrel patch	Approve if the patient has tried CombiPatch, if formulary. If CombiPatch is non-formulary, approve if the patient has tried one oral estrogen/progestin combination product (e.g., estradiol/norethindrone [Activella, generics], Prempro, Premphase, ethinyl estradiol/norethindrone acetate [Femhrt, generics], Prefest, Angelig).	1 year	Yes
(Topical)	Cililiara i 10	levollorgestier pateri	i remprase, enimy estracionio enimo die acetate (i enimi, generios), i relest, Angenq).	i yeai	163
			Approve if the patient meets BOTH of the following (A and B):		
Estrogen and			A. Patient has tried at least one formulary non-patch topical estradiol product: estradiol 0.06% gel (transdermal) [Estrogel, generics], estradiol 0.1% gel (transdermal) [Divigel, generics], Evamist, if one is formulary; AND		
Estrogen Combination Products			B. Patient has tried at least one estradiol patch.  Note: Examples of estradiol patch products include: estradiol patch [Climara, Vivelle Dot, generics], Minivelle [generics).		
(Topical)	Elestrin	estradiol gel 0.06%	Note: If no transdermal gels or sprays are formulary, the patient would still need to try an estradiol patch.	1 year	Yes
			Approve if the patient meets the following (A, B and C):		
			A. Patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with one product from the following list, if formulary: Activella (including generics of Activella: Mimvey and estradiol-		
			norethindrone); AND  B. Patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with one product from the following list: Femhrt (including generics of Femhrt: Jinteli, Fyavolv, norethindrone-ethiny)		
Estrogen		estradiol and	B. Patient has thed and, according to the prescriber, has experienced inadequate enlicacy OR significant intolerance with one product from the following list: Feminit (including generics of Feminit: Jintell, Fyavolv, norethindrone-ethiny) lestradiol); AND		
Combination Products		progesterone	C. Patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with one of Premphase or Prempro, if formulary.		
(Oral)	Bijuva	capsules	Note: If none are formulary in A, B and C, approve. If none are formulary in A, B, or C, the other(s) would still need to be tried.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Estrogen Combination Products (Oral)	Premphase	conjugated estrogens/medroxypr ogesterone tablets	Approve if the patient meets the following (A, B, and C):  A. Patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with one product from the following list, if formulary: Femhrt (including generics of Femhrt: Jinteli, Fyavolv, norethindrone-ethinyl estradiol); AND  B. Patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with one product from the following list, if formulary: Activella (including generics of Activella: Mimvey and estradiol-norethindrone); AND  C. Patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with Bijuva.  Note: If none are formulary in A, B, and C, approve. If none are formulary in A, B, or C, the other(s) would still need to be tried.	1 year	Yes
Estrogen Combination Products	·	conjugated estrogens, medroxyprogesterone tablets	Approve if the patient meets the following (A, B, and C):  A. Patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with one product from the following list, if formulary: Femhrt (including generics of Femhrt: Jinteli, Fyavolv, norethindrone-ethinyl estradiol); AND  B. Patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with one product from the following list, if formulary: Activella (including generics of Activella: Mimvey and estradiol-norethindrone); AND  C. Patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with Bijuva.		
(Oral) Estrogen Products	Prempro	esterified estrogens	Note: If none are formulary in A, B, and C, approve. If none are formulary in A, B, or C, the other(s) would still need to be tried.  Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with two products (or one if one is formulary) from the following list: estradiol tablets (Estrace, generics)	1 year	Yes
(Oral)	Menest	tablets	and Premarin tablets. If neither are formulary, approve.	1 year	Yes
Estrogen Products (Oral)	Premarin	conjugated estrogens tablets	Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with two products (or one if one is formulary) from the following list: estradiol tablets (Estrace, generics) and Menest tablets. If neither are formulary, approve.	1 year	Yes
Estrogen Products (Vaginal)	Estrace Cream	estradiol cream	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Estrogen Products (Vaginal)	Vagifem	estradiol vaginal tablet	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Estrogen Products (Vaginal)	Femring	estradiol vaginal ring (0.05 mg and 0.10 mg)	Approve if the patient has tried two formulary alternatives from the following list (or one if only one is formulary): Invexxy vaginal insert, Premarin cream, estradiol 0.01% cream (Estrace Cream, generics), Estring vaginal ring, estradiol vaginal tablet (e.g., Yuvafem, Vagifem, generics), estradiol patch (Climara, generics), estradiol patch (Vivelle Dot, generics), Menostar patch, estradiol tablets (Estrace, generics), Menest tablets, or Premarin tablets. If none are formulary, approve.	1 year	Yes
Estrogen Products (Vaginal)	Estring	estradiol 2 mg vaginal	1. Approve if the patient has tried two formulary alternatives from the following list (or one if only one is formulary): Imvexxy vaginal insert, Femring vaginal ring, Premarin Cream, estradiol 0.01% cream (Estrace Cream, generics), or estradiol vaginal tablet (e.g., Yuvafem, Vagifem, generics). If none are formulary, approve.  2. If according to the prescriber, the patient requires a low-dose vaginal product, approve if the patient has tried one of Imvexxy vaginal insert or estradiol vaginal tablets (e.g., Yuvagen, Vagifem, generics), if formulary. If neither are formulary, approve.	1 year	Yes
Estrogen Products (Vaginal)	Imvexxy	estradiol vaginal insert	<ol> <li>Approve if the patient has tried two formulary alternatives from the following list (or one if only one is formulary): Premarin vaginal cream, Femring vaginal ring, estradiol 0.01% cream (Estrace Cream, generics), Estring vaginal ring, or estradiol vaginal tablet (e.g., Yuvafem, Vagifem, generics). If none are formulary, approve.</li> <li>If according to the prescriber, the patient requires a low-dose vaginal product, approve if the patient has tried one of Estring or estradiol vaginal tablets (e.g., Yuvagen, Vagifem, generics), if formulary. If neither are formulary, approve.</li> </ol>	1 year	Yes
Factor Deficiency Agents and Related Products	NovoSeven RT	Factor VIIa (recombinant) powder for injection	<ol> <li>Hemophilia A with Inhibitors; Hemophilia B with Inhibitors: Approve if the patient meets the following criteria (a, b, c, or d):         <ul> <li>The patient has tried Sevenfact, if formulary. If Sevenfact is non-formulary, approve; OR</li> <li>The patient is less than 12 years of age; OR</li> <li>The patient has an allergy to rabbits or rabbit-derived products; OR</li> <li>The patient is currently receiving NovoSeven RT or has received NovoSeven RT in the past.</li> </ul> </li> <li>Congenital Factor VII Deficiency, approve.</li> <li>Glanzmann's Thrombasthenia, approve.</li> <li>Hemophilia, Aquired, approve.</li> <li>Generalized Myasthenia Gravis, anti-acetylcholine receptor antibody positive in a patient ≥18 years of age.</li> </ol>	1 year	Yes
		rozanolixizumab-noli	<ol> <li>Approve if the patient has tried one of 1) Imvaavy, or 2) an efgartigimod alfa product (Vyvgart intravenous or Vyvgart Hytrulo), if formulary. If none are formulary, approve.</li> <li>If the patient is unable to obtain and/or maintain intravenous access, approve if the patient has tried Vyvgart Hytrulo, if formulary. If Vyvgart Hytrulo is non-formulary, approve.</li> <li>Approve if the patient has already been started on therapy with Rystiggo.</li> <li>Generalized Myasthenia Gravis, anti-muscle-specific tyrosine kinase antibody-positive in a patient ≥ 18 years of age.</li> <li>Approve if the patient has tried Imaavy, if formulary. If Imaavy is non-formulary, approve.</li> <li>If the patient is unable to obtain and/or maintain intravenous access, approve.</li> </ol>		
Fc receptor blocker	Rystiggo	subcutaneous infusion	3. Approve if the patient has already been started on therapy with Rystiggo.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Fenofibrates	Tricor	fenofibrate tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Fenofibrates	Antara, Lipofen and authorized generics	fenofibrate capsules or tablets	Approve if the patient has tried three other formulary fenofibrate products (e.g., TriCor or generic, Lipofen, Fenoglide or generic, Trilipix or generic, generic fenofibrate capsule/ tablets, Fibricor or generic, generic fenofibric acid tablets) or two if only two are formulary, or one if only one is formulary. If none are formulary approve the requested agent.	1 year	Yes
Fentanyl Transmucosal Products	Fentora and authorized generic	fentanyl buccal tablet	See Opioids Transmucosal – Fentora FE	1 year	Yes
Fertility Agents – Follitropin Ovulatory Stimulants	Follistim AQ	follitropin beta	<ol> <li>Approve if the patient has tried one product from the following list: Gonal-F/Gonal-F RFF, if formulary. If Gonal-F/Gonal-F RFF is non-formulary, approve.</li> <li>Patient has been started on a current cycle of therapy with Follistim AQ: approve to complete the current cycle.</li> </ol>	1 year	Yes
Fertility Agents – Gonadotropin- Releasing Hormone (GnRH) Antagonists	ganirelix injection	ganirelix acetate injection	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Gabapentin and Gabapentin-Like Medications	Lyrica	pregabalin capsules	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Gabapentin and Gabapentin-Like Medications	Lyrica CR	pregabalin controlled- release capsules	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Gabapentin and Gabapentin-Like Medications	Neurontin	gabapentin tablet,	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Gabapentin and Gabapentin-Like Medications	Gabarone	gabapentin tablets	Approve if the patient tried and CANNOT TAKE gabapentin CAPSULES.	1 year	Yes
Gastrointestinal Drugs - Miscellaneous	Carafate	sulcralfate tablets and oral suspension	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Gastrointestinal Drugs - Miscellaneous		glycopyrrolate oral solution	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Gastrointestinal Drugs - Miscellaneous	Dartisla ODT	glycopyrrolate orally disintegrating tablets	<ol> <li>Direct to glycopyrrolate tablets.</li> <li>Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use glycopyrrolate tablets.</li> </ol>	1 year	Yes
Gastrointestinal Drugs - Miscellaneous	Mytesi	crofelemer delayed- release tablets	For the symptomatic relief of non-infectious diarrhea in adult patients with Human immunodeficiency virus (HIV) or Acquired immunodeficiency syndrome (AIDS): Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR a significant intolerance with both diphenoxylate-atropine tablets AND loperamide.	1 year	Yes
Gastroparesis Agents	Gimoti	metoclopramide nasal spray	No exceptions are recommended. Due to insufficient clinical efficacy data, approval is not recommended for Gimoti. ( <b>NOTE</b> : It is not appropriate to use standard global criteria for this medication; Denial reason is: No exceptions are recommended. Due to insufficient clinical efficacy data, approval is not recommended.)  1. Patients with Gaucher Disease Type 1, approve if the patient has tried one product from the following list: Cerezyme or Vpriv, if formulary. If neither are formulary, approve.	N/A	Yes
Gaucher Disease		taliglucerase alfa for	Note: Type 1 Gaucher Disease Type 3, approve if the patient has tried one product from the following list: Cerezyme or Vpriv, if formulary. If neither are formulary, approve.  Note: Type 3 Gaucher disease is also known as non-neuronopathic Gaucher disease.		
Medications	Elelyso	injection	3. Patients with Gaucher Disease Type I or Type 3 currently established on treatment with Elelyso: approve.	1 year	Yes

					2025 NPF
				Approval	Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
			1. Patients with Gaucher Disease Type 1, approve if the patient has tried one product from the following list: Cerezyme or Elelyso, if formulary. If neither are formulary, approve.  Note: Type 1 Gaucher disease is also known as non-neuronopathic Gaucher disease.		
			2. Patients with Gaucher Disease Type 3, approve if the patient has tried one product from the following list: Cerezyme or Elelyso, if formulary. If neither are formulary, approve.		
Gaucher Disease			Note: Type 3 Gaucher disease is also known as chronic neuronopathic Gaucher disease.		
Medications	Vpriv	injection	3. Patients with Gaucher Disease Type 1 or Type 3 currently established on treatment with Vpriv: approve.	1 year	Yes
					MSB Exclusion
					*This criteria
Gaucher Disease	_		NOTE: A multisource Brand product is being requested.		applies only to
Medications	Zavesca	miglustat capsules	See standard Gaucher Disease – Substrate Reduction Therapy Preferred Specialty Management Policy criteria	1 year	the NPF
		aluggan human	<ol> <li>Approve if the patient has tried TWO products from the following list: Baqsimi intranasal, Gvoke, or Zegalogue, if formulary (or only one if one is formulary). If none are formulary, approve.</li> <li>Patient is ≥ 2 years of age but &lt; 6 years of age, approve if the patient has tried one of Baqsimi or Gvoke, if formulary. If neither are formulary, approve.</li> </ol>		
Glucose-Elevating	GlucaGen/GlucaGen		3. Patient is ≤ 2 years of age and ≥ 1 year of age, approve if the patient has tried ble of Bagsimi, if formulary. If Bagsimi is non-formulary, approve.		
Drugs	HypoKit		4. If the patient is < 1 year of age, approve.	1 year	Yes
5.490	1.1500.111	joodo	1. Approve if the patient has tried TWO products from the following list: Baqsimi intranasal, Gvoke, or Zegalogue, if formulary (or only one if one is formulary). If none are formulary, approve.	. you.	1.00
			2. If the patient is ≥ 2 years of age but < 6 years of age, approve if the patient has tried one of Bagsimi or Gvoke, if formulary, If neither are formulary, approve.		
Glucose-Elevating	Glucagon/Glucagon	glucagon/glucagon	3. If the patient is < 2 years of age and ≥ 1 year of age, approve if the patient has tried Baqsimi, if formulary. If Baqsimi is non-formulary, approve.		
Drugs	Emergency Kit	Emergency Kit	4. If the patient is < 1 year of age, approve.	1 year	Yes
		dasiglucagon			
Glucose-Elevating		subcutaneous			
Drugs	Zegalogue	injection	Approve if the patient has tried BOTH of Gvoke and Baqsimi, if formulary (or one if one is formulary). If neither are formulary, approve.	1 year	Yes
Gonadotropin-			Central Precocious Puberty; Gender-Dysphoric/Gender-Incongruent Persons; Persons Undergoing Gender Reassignment (Female-To-Male or Male-To-Female).		
Releasing Hormone (GnRH) Analogs -		leuprolide acetate for	1. Approve if the patient has tried both Triptodur and Fensolvi, if formulary (or one if one is formulary) [documentation required]. If neither are formulary, approve.		
CPP	Lupron Depot-Ped	depot suspension	1. Approve it < 2 years of age, approve.	1 vear	Yes
0	Zapron Bopot i ou	a opor o a oporioron	and the district of the distri	. you.	1.00
Gonadotropin-			Prostate Cancer:		
Releasing Hormone	Leuprolide Depot	leuprolide acetate	Approve if patient has tried Lupron Depot 22.5 mg, if formulary. If Lupron Depot, 22.5 mg is non-formulary, approve if the patient meets (1 or 2):		
(GnRH) Analogs -		22.5 mg for depot	1. Approve if the patient has tried one of Camcevi, Eligard, Firmagon, Trelstar, or Orgovyx, if formulary. If none are formulary, approve.		
Prostate Cancer	Depot 22.5 mg	suspension	2. Patient who has already been started on therapy with Leuprolide Depot, approve if the patient has tried one of Camcevi or Eligard, if formulary. If neither are formulary, approve.	1 year	Yes
			Prostate Cancer.		
Canadatranin			1. Approve if the patient has tried one of the following: leuprolide depot 22.5 mg (formerly Lutrate), Lupron Depot (7.5 mg, 22.5 mg, 30 mg, or 45 mg), Eligard, Firmagon, Trelstar or Orgovyx. If none are formulary, approve.  2. Patients currently receiving therapy with Camcevi, approve if the patient has tried one of Lupron Depot or Eligard. If neither are formulary, approve.		
Gonadotropin- Releasing Hormone		leuprolide injectable	2. Fallettis Currently freceiving therapy with Carticevi, approve it the patient has thed one of Eupron Depot of Engand. If heldrer are formularly, approve.		
(GnRH) Analogs -		emulsion for	Head and Neck Cancer – Salivary Gland Tumors.		
Prostate Cancer	Camcevi	subcutaneous use	Approve if the patient has tried one of Lupron Depot or Eligard. If neither are formulary, approve.	1 year	Yes
				Í	
			Prostate Cancer:		
			1. Approve if the patient has tried one of the following: leuprolide depot 22.5 mg (formerly Lutrate), Camcevi, Lupron Depot (7.5 mg, 22.5 mg, 30 mg, or 45 mg), Eligard, Firmagon, or Orgovyx. If none are formulary, approve.		
Gonadotropin-			2. If only leuprolide depot 22.5 mg (formerly Lutrate) is formulary and the prescriber prefers monthly dosing, approve the patient has tried one of Lupron Depot 7.5 mg, Eligard, or Firmagon. If none are formulary, approve.		
Releasing Hormone			3. Patients currently receiving therapy with Trelstar: approve.		
(GnRH) Analogs - Prostate Cancer	Trelstar	triptorelin pamoate for injectable suspension	Hand and Nack Concer. Salivery Cland Tumore: approve	1 voor	Yes
1 TOSIALE CATICEI	TICISIAI	injectable suspension	Head and Neck Cancer – Salivary Gland Tumors: approve.	1 year	MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Gout Medications	Colcrys	colchicine tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
					MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Gout Medications	Uloric	febuxostat tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Therapy Class	Dranu Name	Generic Name	Growth hormone deficiency (GHD) in children, Turner syndrome, GHD in adults, Prader Willi syndrome, Idiopathic short stature in children, Children with chronic kidney disease, Short Stature Homeobox-Containing Gene deficiency,	Duration	Wedicalton
			Children Born Small for Gestational Age or with Intrauterine Growth Restriction, Noonan syndrome, Short bowel syndrome:		
			Approve if the patient meets BOTH of the following (1 and 2):		
			1. Patient has tried five of the following (if five are formulary, or four if four are formulary, or three if three are formulary, or two if only two are formulary, or one if only one is formulary): Genotropin, Norditropin Flexpro, Nutropin AQ,		
			Omnitrope, Saizen, or Zomacton (documentation required): AND		
			Note: If none are formulary, approve.		
Growth Hormone			2. Patient cannot constitue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber,		
Products	Humatrope	somatropin injection	would result in a significant allergy or serious adverse reaction [documentation required].	1 year	Yes
			Growth hormone deficiency (GHD) in children, Turner syndrome, GHD in adults, Prader Willi syndrome, Idiopathic short stature in children, Children with chronic kidney disease. Short Stature Homeobox-Containing Gene deficiency,	. <b>,</b>	
			Children Born Small for Gestational Age or with Intrauterine Growth Restriction, Noonan syndrome, Short bowel syndrome:		
			Approve if the patient meets BOTH of the following (1 and 2):		
			1. Patient has tried five of the following (if five are formulary, or four if four are formulary, or three if three are formulary, or two if only two are formulary, or one if only one is formulary): Genotropin, Humatrope, Nutropin AQ, Omnitrope,		
			Saizen, or Zomacton [documentation required]; AND		
			Note: If none are formulary, approve.		
Growth Hormone			2. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber,		
Products	Norditropin Flexpro	somatropin injection	would result in a significant allergy or serious adverse reaction [documentation required].	1 year	Yes
			Growth hormone deficiency (GHD) in children, Turner syndrome, GHD in adults, Prader Willi syndrome, Idiopathic short stature in children, Children with chronic kidney disease, Short Stature Homeobox-Containing Gene deficiency,		
			Children Born Small for Gestational Age or with Intrauterine Growth Restriction, Noonan syndrome, Short bowel syndrome:		
			Approve if the patient meets BOTH of the following (1 and 2):		
			1. Patient has tried five of the following (if five are formulary, or four if four are formulary, or three if three are formulary, or two if only two are formulary, or one if only one is formulary). Genotropin, Humatrope, Norditropin Flexpro,		
			Omnitrope, Saizen, or Zomacton [documentation required]; AND		
			Note: If none are formulary, approve.		
Growth Hormone			2. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber,		
Products	Nutropin AQ Nuspin	somatropin injection	would result in a significant allergy or serious adverse reaction [documentation required].	1 year	Yes
			Growth hormone deficiency (GHD) in children, Turner syndrome, GHD in adults, Prader Willi syndrome, Idiopathic short stature in children, Children with chronic kidney disease, Short Stature Homeobox-Containing Gene deficiency		
			Children Born Small for Gestational Age or with Intrauterine Growth Restriction, Noonan syndrome, Short bowel syndrome:		
			Approve if the patient meets BOTH of the following (1 and 2):		
			1. Patient has tried five of the following (if five are formulary, or four if four are formulary, or three if three are formulary, or two if only two are formulary, or one if only one is formulary). Genotropin, Humatrope, Norditropin Flexpro,		
			Nutropin AQ, Omnitrope, or Zomacton [documentation required]; AND		
			Note: If none are formulary, approve.		
Growth Hormone			2. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber,		
Products	Saizen/SaizenPrep	somatropin injection	would result in a significant allergy or serious adverse reaction [documentation required].	1 year	Yes
			Growth hormone deficiency (GHD) in children, Turner syndrome, GHD in adults, Prader Willi syndrome, Idiopathic short stature in children, Children with chronic kidney disease, Short Stature Homeobox-Containing Gene deficiency,		
			Children Born Small for Gestational Age or with Intrauterine Growth Restriction, Noonan syndrome, Short bowel syndrome:		
			Approve if the patient meets BOTH of the following (1 and 2):		
			1. Patient has tried five of the following (if five are formulary, or four if four are formulary, or three if three are formulary, or two if only two are formulary, or one if only one is formulary): Genotropin, Humatrope, Norditropin Flexpro,		
			Nutropin AQ, Omnitrope, or Saizen [documentation required];AND		
			Note: If none are formulary, approve.		
Growth Hormone	Zomacton (formerly		2. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber,		
Products	Tev-Tropin)	somatropin injection	would result in a significant allergy or serious adverse reaction [documentation required].	1 year	Yes
			1. Growth hormone deficiency in a patient ≥ 2.5 years of age to < 3 years of age.		
			patient meets criteria #3.		
			2. Growth hormone deficiency in patients ≥ 3 years of age to < 18 years of age, approve if the patient has tried one of Sogroya or Ngenla for 6 months OR experienced an intolerance with the respective agent, if one is formulary.		
			3. If neither Sogroya nor Ngenla are formulary (in patients ≥ 2.5 years of age to < 18 years of age) OR the patient is ≥ 1 year of age and < 2.5 years of age, approve if the patient meets ONE of the following (A or B):		
			A. Patient has been able to adhere to somatropin product(s) administered daily AND has experienced inadequate efficacy (i.e., patient has tried for 12 months and has a growth rate of less than 2 cm per year) [documentation		
			required] with ONE product from the following list: Genotropin, Humatrope, Norditropin Flexpro, Nutropin AQ, Omnitrope, Saizen, or Zomacton; OR		
			B. Patient meets BOTH of the following (i and ii):		
			i. Patient has tried TWO of the following products: Genotropin, Humatrope, Norditropin Flexpro, Nutropin AQ, Omnitrope, Saizen, or Zomacton [documentation required]; AND		
			ii. Patient cannot continue to use each of the TWO products due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber,		
Growth Hormone		Ionapegsomatropin-	would result in a significant allergy or serious adverse reaction [documentation required].		
Products – Weekly		tcgd subcutaneous	Note: Meeting the criteria above with a trial of any daily growth hormone product(s) would count toward meeting the requirements regardless of formulary status.		
Dosed	Skytrofa	injection	Note: If there is only ONE growth hormone product on a formulary, then the patient would NOT be required to try TWO products- only ONE.	1 vear	Yes

					2025 NPF
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	Excluded Medicaiton
			1. Growth hormone deficiency in patients ≥ 2.5 years of age to < 3 years of age, approve if the patient has tried Skytrofa for 6 months OR experienced an intolerance with Skytrofa, if formulary. If Skytrofa is non-formulary, approve if the patient meets criteria #3.		
			2. Growth hormone deficiency in patients ≥ 3 years of age to < 18 years of age, approve if the patient has tried one of Skytrofa or Ngenla for 6 months OR experienced an intolerance with the respective agent, if one is formulary.		
			3. If neither Skytrofa nor Ngenla are formulary (in patients ≥ 2.5 years of age to < 18 years of age), approve if the patient meets ONE of the following (A or B):  A. Patient has been able to adhere to somatropin product(s) administered daily AND has experienced inadequate efficacy (i.e., patient has tried for 12 months and has a growth rate of less than 2 cm per year) [documentation required] with ONE product from the following list: Genotropin, Humatrope, Norditropin Flexpro, Nutropin AQ, Omnitrope, Saizen, or Zomacton; OR  B. Patient meets BOTH of the following (i and ii):  i. Patient has tried TWO of the following products: Genotropin, Humatrope, Norditropin Flexpro, Nutropin AQ, Omnitrope, Saizen, or Zomacton [documentation required]; AND		
			ii. Patient cannot continue to use each of the TWO products due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  Note: Meeting the criteria above with a trial of any daily growth hormone product(s) would count toward meeting the requirements regardless of formulary status.  Note: If there is only ONE growth hormone product on a formulary, then the patient would NOT be required to try TWO products- only ONE.		
Growth Hormone			<ul> <li>4. Adults with growth hormone deficiency (patients ≥ 18 years of age).</li> <li>Approve if the patient meets BOTH of the following (A and B):         <ul> <li>A. Patient has tried TWO of the following products: Genotropin, Humatrope, Norditropin Flexpro, Nutropin AQ, Omnitrope, Saizen, or Zomacton [documentation required]; AND</li> <li>B. Patient cannot continue to use each of the TWO products due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].</li> </ul> </li> </ul>		
Products – Weekly Dosed	Sogrovo	somapacitan-beco subcutaneous injection	Note: If there is only ONE growth hormone product on a formulary, then the patient would NOT be required to try TWO products- only ONE.	1 year	Yes
Dosed	Sogroya			1 year	MSB Exclusion
Head Lice Treatments (Topical)	Natroba		<b>NOTE:</b> A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product. <b>Criteria:</b> Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 vear	*This criteria applies only to the NPF
(*			If requesting brand Pylera: Approve if the patient has tried generic Pylera (bismuth-metronidazole-tetracycline 140-125-125), if formulary.	, ,	
Helicobacter Pylori Agents	Pylera	bismuth subcitrate potassium, metronidazole plus tetracycline capsules	If requesting brand Pylera and generic Pylera (bismuth-metronidazole-tetracycline 140-125-125), is non-formulary (or if requesting generic Pylera), approve if the patient meets ONE of the following (A or B):  A. The patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with TWO different regimens of single-entity products (e.g., clarithromycin + amoxicillin + proton pump inhibitor [e.g., omeprazole, lansoprazole]; bismuth-containing product + tetracycline + metronidazole + proton pump inhibitor [e.g., omeprazole, lansoprazole]; by Voquezna + amoxicillin +/- clarithromycin); OR  B. The patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with any TWO pre-packaged products (e.g., amoxicillin/clarithromycin/lansoprazole [Prevpac, generics], Omeclamox-Pak, Voquezna Pak, or Talicia).	1 month	Yes
- ·g-····	. ,	'	Myelodysplastic Syndromes with Transfusion-Dependent Anemia who are relapsed, refractory or ineligible for erythropoiesis-stimulating agents.		
			1. Approve if the patient tried Reblozyl, if formulary. If Reblozyl is non-formulary, approve. 2. Approve if the patient meets ALL of the following (A, B, and C):  A. Patient does NOT have a deletion 5q [del(5q)]; AND  B. Patient has ring sideroblasts < 15% (or ring sideroblasts < 5% with an SF3B1 pathogenic variant); AND		
Hematology Agents - Miscellaneous	Rytelo	imetelstat intravenous injection	C. Patient has tried or has a poor probability to respond to immunosuppressive therapy.  3. Approve if the patient has already been started on therapy with Rytelo.	1 year	Yes
		motixafortide	Peripheral blood stem cell mobilization for collection and subsequent autologous transplantation in patients with Multiple myeloma.		
Hematopoietic/Throm bopoietic Agents	Aphexda		1. Approve if the patient has tried plerixafor injection (Mozobil, generics), if formulary. If plerixafor injection (Mozobil, generics) are non-formulary, approve.  2. Approve if the patient has already started therapy with Aphexda.	1 year	Yes
Hemophilia - Factor IX Products (recombinant		coagulation Factor IX [recombinant],			
extended half-life products)	Rebinyn	glycoPEGylated for IV injection	1. Approve if the patient has tried one product from the following list (if one is formulary): Alprolix or Idelvion. If neither are formulary, approve.  2. Approve if the patient is currently receiving Rebinyn or has received Rebinyn in the past.	1 year	Yes
Hemophilia - Factor IX Products (recombinant					
standard half-life			1. Approve if the patient has tried one of BeneFIX or Ixinity, if formulary. If neither are formulary, approve.		
products)	Rixubis	[recombinant]	2. Approve if the patient is currently receiving Rixubis or has received Rixubis in the past.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Hemophilia - Factor					
IX Products		coagulation factor IX			
(recombinant standard half-life		[recombinant] solution for intravenous	1. Approve if the patient has tried one product from the following list (if one is formulary): Rixubis or BeneFIX. If neither are formulary, approve.		
products)	Ixinity	injection	2. Approve if the patient is currently receiving Ixinity or has received Ixinity in the past.	1 vear	Yes
Hemophilia - Factor	IXIIIIty	Injection	2. Approve if the patient is currently receiving tainity or has received rainity in the past.	i yeai	163
VIII Products		antihemophilic factor	1. Patient has tried two formulary recombinant Factor VIII products from the following list (if two are formulary, or one if one is formulary): Advate, Recombinate, Kogenate FS, Xyntha, Novoeight, Koyaltry, Afstyla. If none are formulary,		
(recombinant		[recombinant] for	approve.		
standard half-life)	Nuwig		2. Patient is currently receiving Nuwig or has received Nuwig in the past: approve	1 year	Yes
Hemophilia - Factor	'	,		Í	
VIII Products		antihemophilic factor	1. Patient has tried two formulary recombinant Factor VIII products from the following list (if two are formulary or one if one is formulary): Advate, Kogenate FS, Xyntha, Novoeight, Nuwiq, Kovaltry, Afstyla. If none are formulary,		
(recombinant		[recombinant]	approve.		
standard half-life)	Recombinate	injection	2. Patient is currently receiving Recombinate or has received Recombinate in the past: approve.	1 year	Yes
			For Hemophilia A with inhibitors.		
			1. Approve if the patient has tried Hemlibra, if formulary. If Hemlibra is non-formulary, approve.		
			2. Approve if, according to the prescriber, there is concern for a drug-drug interaction (e.g., drug-drug interaction with Hemlibra and Feiba).		
			3. Approve if the patient has already been started on therapy with Alhemo.		
Hemophilia – Non-		concizumab-mtci	For Hemophilia B with inhibitors.		
Factor Routine			1. Approve if the patient has tried Qfittiia, if formulary. If Qfittia is non-formulary, approve.		
Prophylaxis Products			2. Approve if the patient has a lieu dinitially. In dinitially in dinitial short-inditionally, approve.  2. Approve if the patient has already been started on therapy with Alhemo.	1 year	Yes
1 Tophylaxis I Toddots	Allicino	Injection	For Hemophilia A without inhibitors.	i year	103
			1. Approve if the patient has tried Hemlibra, if formulary. If Hemlibra is non-formulary, approve.		
			2. Approve if the patient has already been started on therapy with Hympavzi.		
			, , , , , , , , , , , , , , , , , , , ,		
Hemophilia – Non-		marstacimab-hncq	For Hemophilia B without inhibitors.		
Factor Routine		subcutaneous	1. Approve if the patient has tried Qfittia, if formulary. If Qfittia is non-formulary, approve.		
Prophylaxis Products	Hympavzi	injection	2. Approve if the patient has already been started on therapy with Hympavzi.	1 year	Yes
			Patient meets the following standard Hemophilia – Gene Therapy – Beqvez Prior Authorization Policy criteria AND		
			Patient meets ONE of the following (1 or 2):		
			1. If Hemgenix is non-formulary, approve; OR		
			2. Hemgenix is not available at the treatment facility or treatment center in which the patient is enrolled to receive the gene therapy, approve.		
		<b>6</b> 1			
Homonbilio Con-		fidanacogene elaparvovec-dzkt	If the patient does not meet 1 or 2 above, direct the patient to Hemgenix.	Coo DA	
Hemophilia Gene Therapy	Begvez	intravenous infusion	All reviews (approvals and denials) will be forwarded to the Medical Director for evaluation.	See PA Duration	Yes
Погару	DOGVOZ	maavonous musion	A minorate (approvide and defined of the medical principle for evaluation).	Daration	MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Hepatitis B Agents	Baraclude tablets	entecavir tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
	ledipasvir/sofosbuvir				
	tablets 90 mg/400				
Hepatitis C - Oral	mg (Authorized	ledipasvir/sofosbuvir			
Agents	generic for Harvoni)	tablets 90 mg/400 mg	Patient is directed to use Harvoni 90 mg/400 mg. If Harvoni 90 mg/400 mg is non-formulary, approve.	24 weeks	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
			If Epclusa (brand) is formulary:  1. Genotype 2 or 3 Chronic Hepatitis C Virus, Pediatric Patient (≥ 3 Years of Age and < 18 Years of Age) – New Start: Sovaldi is not approved. Offer to review for Epclusa (brand only) using the standard Hepatitis C – Epclusa PA Policy criteria.  2. Patient Continuing Therapy with Sovaldi: Refer to the standard Hepatitis C – Sovaldi PA Policy criteria.		
			If Epclusa (brand) is non-formulary and sofosbuvir/velpatasvir is formulary:  1. Genotype 2 or 3 Chronic Hepatitis C Virus, Pediatric Patient (≥ 3 Years of Age and < 18 Years of Age) – New Start. Approve for the duration specified in the standard Hepatitis C – Sovaldi PA Policy criteria if the patient has met the standard Hepatitis C-Sovaldi PA Policy criteria.		
Hepatitis C - Oral	Sovaldi 200 mg tablets and oral	sofosbuvir tablets and	2. Patient Continuing Therapy with Sovaldi. Refer to the standard Hepatitis C – Sovaldi PA Policy criteria.		
Agents	pellets	oral pellets	If neither Epclusa (brand) nor sofosbuvir/velpatasvir are formulary, approve.	Varies	Yes
7 igorito	ponote	ordi politico	## Epclusa (brand) is formulary:  1. Genotype 2 or 3 Chronic Hepatitis C Virus, Pediatric Patient (≥ 3 Years of Age and < 18 Years of Age) – New Start: Sovaldi is not approved. Offer to review for Epclusa (brand only) using the standard Hepatitis C – Epclusa PA Policy criteria.  2. Patient Continuing Therapy with Sovaldi: Refer to the standard Hepatitis C – Sovaldi PA Policy criteria.	vanos	
Hepatitis C - Oral	Sovaldi 400 mg		If Epclusa (brand) is non-formulary and sofosbuvir/velpatasvir is formulary:  1. Genotype 2 or 3 Chronic Hepatitis C Virus, Pediatric Patient (≥ 3 Years of Age and < 18 Years of Age) – New Start. Sovaldi is not approved. Offer to review for sofosbuvir/velpatasvir 400 mg/100 mg tablets (generic only) using the standard Hepatitis C – Epclusa PA Policy criteria.  2. Patient Continuing Therapy with Sovaldi. Refer to the standard Hepatitis C – Sovaldi PA Policy criteria.		
Agents	tablets	sofosbuvir tablets	If neither Epclusa (brand) nor sofosbuvir/velpatasvir are formulary, approve.	Varies	Yes
· ·9		glecaprevir/			
Hepatitis C - Oral		pibrentasvir tablets		See PSM	
Agents	Mavyret	and oral pellets	See Hepatitis C Virus Direct Acting Antivirals Preferred Specialty Management (PSM) for National Preferred Formulary and Basic Formulary (Mavyret Criteria)	duration	Yes
•	sofosbuvir/velpatasv ir (Authorized generic for Epclusa) 400 mg/100 mg tablets	sofosbuvir/velpatasvir tablets 400 mg/100 mg tablets	Patient is directed to use Epclusa. If Epclusa is non-formulary, approve.	24 weeks	Vos
Agents Hereditary	lablets	ing tablets	Pratient is directed to use Epiciusa. Il Epiciusa is non-iornidiary, approve.	24 weeks	Yes
Angioedema – Acute		icatibant injection for			
Treatment	Firazyr	subcutaneous use	See standard Hereditary Angioedema – Icatibant Preferred Specialty Management Policy criteria.	1 year	Yes
Hereditary Angioedema Products - IV C1 Esterase		C1 esterase inhibitor [human] powder for			
Products	Berinert	intravenous injection	See Hereditary Angioedema Medications - Berinert FE	1 year	Yes
HMG-CoA Reductase Inhibitors and Combination Products	Vytorin	ezetimibe/simvastatin tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  1. Approve if the patient has tried, and according to the prescriber, has experienced inadequate efficacy OR significant intolerance with three statins from the following list (if three are formulary or two if only two are formulary, or one if only is formulary): lovastatin, rosuvastatin (Crestor, generics), atorvastatin (Lipitor, generics), fluvastatin (Lescol/XL, generics), a pitavastatin product (Livalo, Zypitamag), pravastatin (Pravachol), or simvastatin (Zocor, generics). If none are formulary, approve.  2. Patients who cannot swallow or have difficulty swallowing tablets or capsules, approve.  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.		
HMG-CoA Reductase Inhibitors and Combination Products			1. Approve if the patient has tried, and according to the prescriber, has experienced inadequate efficacy OR significant intolerance with three statins from the following list (if three are formulary or two if only two are formulary, or one if only is formulary): lovastatin, rosuvastatin (Crestor, generics), atorvastatin (Lipitor, generics), fluvastatin (Lescol/XL, generics), a pitavastatin product (Livalo, Zypitamag), pravastatin (Pravachol), or simvastatin (Zocor, generics). If none are formulary, approve.  2. Patients who cannot swallow or have difficulty swallowing tablets or capsules, approve.  3. The patient meets both of the following (i and ii):  i. The requested non-formulary drug is being prescribed for the primary prevention of cardiovascular disease (CVD) for an adult aged 40 to 75 years who has one or more CVD risk factors (i.e., dyslipidemia, diabetes, hypertension, or smoking) and an estimated 10-year CVD event risk of 10% or greater and who does NOT have a history of (or signs or symptoms of) CVD (for example: symptomatic coronary artery disease, ischemic stroke); AND  ii. Other formulary alternative(s) would not be as medically appropriate for the patient as the requested non-formulary drug.	1 year	Yes
		·	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  OR	, , , ,	
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve one of the following (A or B):  A. The patient meets both of the following (i and ii):  i. The requested brand non-formulary drug is being prescribed for the primary prevention of cardiovascular disease (CVD) for an adult aged 40 to 75 years who has one or more CVD risk factors (i.e., dyslipidemia, diabetes, hypertension, or smoking) and an estimated 10-year CVD event risk of 10% or greater and who does NOT have a history of (or signs or symptoms of) CVD (for example: symptomatic coronary artery disease, ischemic stroke); AND ii. According to the prescriber, the brand product is being requested because the preferred bioequivalent generic product would not be as medically appropriate for the patient as the requested brand non-formulary drug; OR B. The patient meets both of the following (i and ii):  i. The requested brand non-formulary drug is being prescribed for a use OTHER THAN the primary prevention of cardiovascular disease (CVD) for an adult aged 40 to 75 years who has one or more CVD risk factors (i.e., dyslipidemia, diabetes, hypertension, or smoking) and an estimated 10-year CVD event risk of 10% or greater and who does NOT have a history of (or signs or symptoms of) CVD (for example: symptomatic coronary artery disease,		MSB Exclusion
HMG-CoA Reductase Inhibitors and Combination Products	Crestor		ischemic stroke); AND  ii. The Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	*This criteria applies only to the NPF

Therapy Class	Prand Name	Gaparia Nama	Commercial EE Critoria	Approval	2025 NPF Excluded
HMG-CoA Reductase Inhibitors and Combination Products		Generic Name	Compliance with the Affordable Care Act. HRSA Guidelines, and PHS Act section 2713 is NOT required.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve one of the following (A or B):  A. The patient meets both of the following (i and ii):  i. The requested brand non-formulary drug is being prescribed for the primary prevention of cardiovascular disease (CVD) for an adult aged 40 to 75 years who has one or more CVD risk factors (i.e., dyslipidemia, diabetes, hypertension, or smoking) and an estimated 10-year CVD event risk of 10% or greater and who does NOT have a history of (or signs or symptoms of) CVD (for example: symptomatic coronary artery disease, ischemic stroke); AND ii. According to the prescriber, the brand product is being requested because the preferred bioequivalent generic product would not be as medically appropriate for the patient as the requested brand non-formulary drug; is being prescribed for a use OTHER THAN the primary prevention of cardiovascular disease (CVD) for an adult aged 40 to 75 years who has one or more CVD risk factors (i.e., dyslipidemia, diabetes, hypertension, or smoking) and an estimated 10-year CVD event risk of 10% or greater and who does NOT have a history of (or signs or symptoms of) CVD (for example: symptomatic coronary artery disease, ischemic stroke); AND ii. The requested brand non-formulary drug is being prescribed for a use OTHER THAN the primary prevention of cardiovascular disease (CVD) for an adult aged 40 to 75 years who has one or more CVD	Duration  1 year	MSB Exclusion *This criteria applies only to the NPF
HMG-CoA Reductase Inhibitors and Combination Products		simvastatin tablets	Compliance with the Affordable Care Act. HRSA Guidelines. and PHS Act section 2713 is NOT required.  A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  Approve one of the following (A or B):  i. The patient meets both of the following (i and ii):  i. The requested brand non-formulary drug is being prescribed for the primary prevention of cardiovascular disease (CVD) for an adult aged 40 to 75 years who has one or more CVD risk factors (i.e., dyslipidemia, diabetes, hypertension, or smoking) and an estimated 10-year CVD event risk of 10% or greater and who does NOT have a history of (or signs or symptoms of) CVD (for example: symptomatic coronary artery disease, ischemic stroke); AND ii. According to the prescriber, the brand product is being requested because the preferred bioequivalent generic product would not be as medically appropriate for the patient as the requested brand non-formulary drug; or a substitution of the following (i and ii):  i. The requested brand non-formulary drug is being prescribed for a use OTHER THAN the primary prevention of cardiovascular disease (CVD) for an adult aged 40 to 75 years who has one or more CVD risk factors (i.e., dyslipidemia, diabetes, hypertension, or smoking) and an estimated 10-year CVD event risk of 10% or greater and who does NOT have a history of (or signs or symptoms of) CVD (for example: symptomatic coronary artery disease, ischemic stroke); AND iii. According to the prescriber, or specially appropriate for the patient as the requested brand non-formulary	1 year	MSB Exclusion *This criteria applies only to the NPF

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
			Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  Approve if the patient has tried, and according to the prescriber, has experienced inadequate efficacy OR significant intolerance with five statins from the following list (if five are formulary or four if four are formulary or three if three are formulary or two if only two are formulary, or one if only one is formulary): lovastatin, atorvastatin (Lipitor, generics), rosuvastatin (Crestor, generics), fluvastatin (Lescol/XL, generics), a pitavastatin product (Livalo, Zypitamag), pravastatin (Pravachol), or simvastatin (Zocor, generics). If none are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  1. 1. Approve if the patient has tried, and according to the prescriber, has experienced inadequate efficacy OR significant intolerance with five statins from the following list (if five are formulary or four if four are formulary or three if three are formulary or two if only two are formulary, or one if only one is formulary): lovastatin, atorvastatin (Lipitor, generics), rosuvastatin (Crestor, generics), fluvastatin (Lescol/XL, generics), a pitavastatin product (Livalo, Zypitamag), pravastatin (Pravachol), or simvastatin (Zocor, generics). If none are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.		
HMG-CoA Reductase Inhibitors and		lovastatin extended-	i. The requested non-formulary drug is being prescribed for the primary prevention of cardiovascular disease (CVD) for an adult aged 40 to 75 years who has one or more CVD risk factors (i.e., dyslipidemia, diabetes, hypertension, or smoking) and an estimated 10-year CVD event risk of 10% or greater and who does NOT have a history of (or signs or symptoms of) CVD (for example: symptomatic coronary artery disease, ischemic stroke); AND		
Combination Products	Altoprev	release tablets	ii. Other formulary alternative(s) would not be as medically appropriate for the patient as the requested non-formulary drug.	1 year	Yes
HMG-CoA Reductase Inhibitors and Combination Products	Ezallor Sprinkle	rosuvastatin capsules	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  1. Approve if the patient has tried, and according to the prescriber, has experienced inadequate efficacy OR significant intolerance with three statins from the following list (if three are formulary or two if only two are formulary, or one if only is formulary): lovastatin, (rosuvastatin (Crestor, generics), atorvastatin (Lipitor, generics), fluvastatin (Lescol/XL, generics), a pitavastatin product (Livalo, Zypitamag), pravastatin (Pravachol), or simvastatin (Zocor, generics). If none are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.  2. Patients who cannot swallow or have difficulty swallowing tablets or capsules, approve.  OR  Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is required.  1. Approve if the patient has tried, and according to the prescriber, has experienced inadequate efficacy OR significant intolerance with three statins from the following list (if three are formulary or two if only two are formulary, or one if only is formulary): lovastatin, rosuvastatin (Crestor, generics), atorvastatin (Lescol/XL, generics), a pitavastatin product (Livalo, Zypitamag), pravastatin (Pravachol), or simvastatin (Zocor, generics). If none are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.  2. Patients who cannot swallow or have difficulty swallowing tablets or capsules, approve.  3. The patient meets both of the following (i and ii):  1. The requested non-formulary drug is being prescribed for the primary prevention of cardiovascular disease (CVD) for an adult aged 40 to 75 years who has one or more CVD risk factors (i.e., dyslipidemia, diabetes, hypertension, or smoking) and an estimated 10-year CVD event risk of 10% or greater and who does NOT have a history of (or sig	1 year	Yes
HMG-CoA Reductase		pitavastatin calcium	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  Approve if the patient has tried, and according to the prescriber, has experienced inadequate efficacy OR significant intolerance with three statins from the following list (if three are formulary or two if only two are formulary, or one if only one is formulary): lovastatin, simvastatin (Zocor, generics), pravastatin (Pravachol, generics), atorvastatin (Lipitor, generics), rosuvastatin (Crestor, generics), or fluvastatin (Lescol/XL, generics), or Altoprev. If none are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.  OR  Compliance with the Affordable Care Act. HRSA Guidelines, and PHS Act section 2713 is required.  1. 1. Approve if the patient has tried, and according to the prescriber, has experienced inadequate efficacy OR significant intolerance with three statins from the following list (if three are formulary or two if only two are formulary, or one if only one is formulary): lovastatin, simvastatin (Zocor, generics), pravastatin (Pravachol, generics), atorvastatin (Lipitor, generics), rosuvastatin (Crestor, generics), or fluvastatin (Lescol/XL, generics), or Altoprev. If none are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.  2. The patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.  2. The patient meets both of the following (i and ii):  i. The requested non-formulary drug is being prescribed for the primary prevention of cardiovascular disease (CVD) for an adult aged 40 to 75 years who has one or more CVD risk factors (i.e., dyslipidemia, diabetes, hypertension, or smoking) and an estimated 10-year CVD event risk of 10% or greater and who does NOT have a history of (or signs or symptoms of) CVD (for example: symptomatic coronary ar		
Combination Products	121.	tablets	ii. Other formulary alternative(s) would not be as medically appropriate for the patient as the requested non-formulary drug.	1 vear	Yes

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
HMG-CoA Reductase Inhibitors and Combination Products	Roszet and authorized generic	rosuvastatin and ezetimibe	Approve if the patient meets the following criteria (A <u>and</u> B):  A. Patient has tried ezetimibe; AND  B. Patient has tried, and according to the prescriber, has experienced inadequate efficacy OR significant intolerance with atorvastatin (Lipitor, generics) or rosuvastatin (Crestor, generics). If neither of atorvastatin (Lipitor, generics) nor rosuvastatin (Crestor, generics) are formulary, approve.	1 year	Yes - Authorized generic only
	chorionic gonadotropin		<ol> <li>Approve if the patient has tried one product from the following list (if one is formulary): Pregnyl, Novarel or Ovidrel. If none are formulary, approve.</li> <li>For a diagnosis of cryptorchidism or hypogonadism, approve if the patient has tried Pregnyl or Novarel, if formulary. If neither are formulary, approve.</li> <li>For a diagnosis related to infertility or induction of ovulation, approve a one-time fill if the patient may be at risk of missing the optimal administration timeframe window of the product (in order to avoid disruption of the current fertility medication cycle).</li> </ol>	1 year	Yes
Human Chorionic Gonadotropin, HCG Agents	Novarel	intramuscular	<ol> <li>Approve if the patient has tried one product from the following list (if one is formulary): chorionic gonadotropin, Pregnyl or Ovidrel. If none are formulary, approve.</li> <li>For a diagnosis of cryptorchidism or hypogonadism, approve if the patient has tried chorionic gonadotropin or Pregnyl, if formulary. If neither are formulary, approve.</li> <li>Patients with a latex allergy: approve if the patient has tried Pregnyl, if formulary. If Pregnyl is non-formulary, approve.</li> <li>For a diagnosis related to infertility or induction of ovulation, approve a one-time fill if the patient may be at risk of missing the optimal administration timeframe window of the product (in order to avoid disruption of the current fertility medication cycle).</li> </ol>	1 year	Yes
Human Immunideficiency Virus (HIV-1) - Non- Nucleoside Reverse Transcriptase	D. L.		1. Approve if the patient has tried one non-nucleoside reverse transcriptase inhibitor (NNRTI) or a NNRTI-containing product (e.g., Sustiva, Edurant, Delstrigo, Complera, Odefsey, Atripla, Symfi, Smyfi Lo).		W.
Human Immunideficiency Virus (HIV-1) – Protease Inhibitor (PI)	Pifeltro		<ol> <li>Patients already started on therapy with Pifeltro, approve.</li> <li>Approve if the patient has tried one protease inhibitor (PI) or a PI-containing product (e.g., Aptivus, atazanavir [Reyataz, generics], Viracept, ritonavir [Norvir, generics], fosamprenavir, Prezista, Evotaz, lopinavir-ritonavir [Kaletra, generics]).</li> <li>Approve if, according to the prescriber, the patient meets BOTH of the following (A and B):         <ul> <li>A. Patient has a history of Apretude (cabotegravir extended-release injectable suspension) for pre-exposure prophylaxis (PrEP); AND</li> <li>B. Patient meets ONE of the following (i or ii):</li></ul></li></ol>	1 year	Yes
, ,	Prezcobix	cobicistat tablets	4. Approve if the patient has already been started on therapy with Prezcobix.  Human Immunodeficiency Virus (HIV) infection, multi-drug treatment-resistant.  1. Approve if the patient has tried Sunlenca or is concomitantly receiving Sunlenca, if formulary. If Sunlenca is non-formulary, approve.  2. Approve if the patient has exhausted at least FOUR of the following antiretroviral classes defined as elimination of all antiretrovirals within a given class due to demonstrated or projected resistance to the agent(s) in that class OR due to significant intolerance (FOUR of a, b, c, d, e, or f):  a) Nucleoside reverse transcriptase inhibitor; OR  Note: Examples of nucleoside reverse transcriptase inhibitor; OR  Note: Examples of nucleoside reverse transcriptase inhibitor; include delaviridine, efavirenz, etravirine, nevirapine, nevirapine XR, rilpivirine.  c) Protease inhibitor; OR  Note: Examples of protease inhibitors include atazanavir, darunavir, fosamprenavir, indinavir, nelfinavir, ritonavir, saquinavir, tipranavir.  d) Fusion inhibitor; OR  Note: Examples of fusion inhibitors include Fuzeon (enfuviritide for injection).  e) Integrase strand transfer inhibitor; include raltegravir, dolutegravir, elvitegravir.	1 year	Yes
Human	Rukobia	fostemsavir extended- release tablets	f) CCR5 antagonist.  Note: Examples of CCR5 antagonists include Selzentry (maraviroc tablets).  3. Approve if the patient has already been started on Rukobia therapy.	1 year	Yes
Immunodeficiency Virus (HIV-1) - integrase strand transfer inhibitor (INSTI) Combination Products	Stribild	elvitegravir/ cobicistat/ emtricitabine/ tenofovir tablets	<ol> <li>Approve if the patient has tried Biktarvy, if formulary. If Biktarvy is non-formulary, approve.</li> <li>Approve if the patient has tried one integrase strand transfer inhibitor (INSTI) or an INSTI-containing product (e.g., Genvoya, Tivicay, Triumeq, Juluca, Isentress or Intress-HD).</li> <li>Patients already started on therapy with Stribild: approve.</li> </ol>	1 year	Yes

				Ammroval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	Medicaiton
Human Immunodeficiency Virus (HIV-1) - Non- Nucleoside Reverse Transcriptase Inhibitors (NNRTI)- Based Combination		efavirenz 600 mg, emtricitabine 200 mg, tenofovir disoproxil fumarate 300 mg	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		MSB Exclusion *This criteria applies only to
-	Atripla	tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Human Immunodeficiency Virus (HIV-1) - Non- Nucleoside Reverse Transcriptase Inhibitors (NNRTI)- Based Combination Products	Complera	emtricitabline/rilpivirin e/tenofovir disoproxil fumarate (TDF) tablets	<ol> <li>Approve if the patient has tried Odefsey, if formulary. If Odefsey is non-formulary, approve if the patient has tried one of the following products: Biktarvy, Genvoya, Stribild, Triumeq, Symtuza, efavirenz-emtricitabine-tenofovir disoproxil fumarate (Atripla, generics), efavirenz-lamivudine-tenofovir (Symfi, Symfi Lo, generics), if formulary. If none are formulary, approve.</li> <li>Approve if the patient is currently taking single-entity or combination products containing emtricitabine, rilpivirine, and tenofovir disoproxil fumarate and is requesting Complera for a single-table regimen.</li> <li>Patients already started on therapy with Complera: approve.</li> </ol>	1 year	Yes
Human Immunodeficiency Virus (HIV-1) - Non- Nucleoside Reverse Transcriptase Inhibitors (NNRTI)- Based Combination Products	Delstrigo	doravirine/lamivudine/ tenofovir disoproxil fumarate tablets	<ol> <li>Approve if the patient has tried one of the following products: Biktarvy, Genvoya, Odefsey, Stribild, Complera, Triumeq, Symtuza, efavirenz-lamivudine-tenofovir (Symfi, Symfi Lo, generics), if formulary. If none are formulary, approve.</li> <li>Patient &lt; 18 years of age AND weighing ≥ 35 kg (77 pounds), approve if the patient has tried one of Biktarvy, Genvoya, Odefsey, Stribild, Complera, or efavirenz-lamivudine-tenofovir (Symfi Lo, generics), if formulary. If none are formulary, approve.</li> <li>Approve if the patient is currently taking single-entity or combination products containing doravirine, lamivudine, and tenofovir disoproxil fumarate and is requesting Delstrigo for a single tablet regimen.</li> <li>Patients already started on therapy with Delstrigo, approve.</li> </ol>	1 year	Yes
Human Immunodeficiency Virus (HIV-1) – NRTI Based Combination	Truvada	emtricitabine/ tenofovir tablets	Compliance with the Affordable Care Act, HRSA Guidelines, and PHS Act section 2713 is NOT required.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  Compliance with the Affordable Care Act. HRSA Guidelines, and PHS Act section 2713 is required.  Approve if the patient meets one of the following criteria (i or ii):  i. The requested brand non-formulary drug is being prescribed for HIV Pre-Exposure Prophylaxis (PrEP) AND, according to the prescriber, the brand product is being requested because the preferred bioequivalent generic product would not be as medically appropriate for the patient as the requested brand non-formulary drug is being prescribed for a use OTHER THAN HIV Pre-Exposure Prophylaxis (PrEP) AND the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].		MSB Exclusion *This criteria applies only to the NPF
Hyaluronic Acid Derivatives	Synojoynt	sodium hyaluronate injection	1. Approve if the patient has tried five formulary intra-articular hyaluronic acid product from the following list ((if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Durolane, Euflexxa, Gel-One, Gel-Syn, GenVisc 850, Hyalgan, Monovisc, Orthovisc, Supartz FX, Synvisc-One, Hymovis, Trivisc, Triluron, or Visco-3 [documentation required]. If none are formulary, approve.  2. Patient has a known allergy to avian or avian-derived products (e.g., eggs, feathers): approve if the patient has tried five formulary products from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Durolane, Euflexxa, Orthovisc, Monovisc, Hymovis, Gel-Syn, Trivisc, or GenVisc 850 [documentation required]. If none are formulary, approve.  3. Patients who have already been started on an injection series with Synojoynt: approve to complete the series.  Note: If the patient has received all the injections in a completed series (for each targeted joint), refer to criteria above.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Hyaluronic Acid Derivatives	Durolane	hyaluronic acid intraarticular injection	<ol> <li>Approve if the patient has tried five formulary intra-articular hyaluronic acid products from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Euflexxa, Gel-One, Gel-Syn, GenVisc 850, Hyalgan, Monovisc, Orthovisc, Supartz FX, Synvisc, Synvisc-One, Hymovis, Visco-3, Synojoynt, Triluron, or Trivisc [documentation required]. If none are formulary, approve Durolane.</li> <li>Patient has a known allergy to avian or avian-derived products (e.g., eggs, feathers): approve if the patient has tried five formulary products from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Euflexxa, Orthovisc, Monovisc, Hymovis, Gel-Syn, GenVisc 850, Synojoynt, or Trivisc [documentation required]. If none are formulary, approve Durolane.</li> </ol>	1 year	Yes
Hyaluronic Acid Derivatives	Euflexxa	sodium hyaluronate	<ol> <li>Approve if the patient has tried five formulary intra-articular hyaluronic acid products from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Durolane, Gel-One, Gel-Syn, GenVisc 850, Hyalgan, Monovisc, Orthovisc, Supartz FX, Synvisc-One, Hymovis, Visco-3, Synojoynt, Triluron, or Trivisc [documentation required]. If none are formulary, approve Euflexxa.</li> <li>Patient has a known allergy to avian or avian-derived products (e.g., eggs, feathers): approve if the patient has tried five formulary products from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Orthovisc, Monovisc, Hymovis, Durolane, Gel-Syn, GenVisc 850, Synojoynt, or Trivisc [documentation required]. If none are formulary, approve Euflexxa.</li> <li>Patients who have already been started on an injection series with Euflexxa: approve to complete the series.</li> <li>Note: If the patient has received all the injections in a completed series (for each targeted joint), refer to criteria above.</li> </ol>	1 year	Yes
Hyaluronic Acid Derivatives	Gel-One	hyaluronate gel	Approve if the patient has tried five formulary intra-articular hyaluronic acid products from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Durolane, Euflexxa, Gel-Syn, GenVisc 850, Hyalgan, Monovisc, Orthovisc, Supartz FX, Synvisc, Synvisc-One, Hymovis, Visco-3, Synojoynt, Triluron, or Trivisc [documentation required]. If none are formulary, approve Gel-One.	1 year	Yes
Hyaluronic Acid Derivatives	Gel-Syn-3	sodium hyaluronate	1. Approve if the patient has tried five formulary intra-articular hyaluronic acid products from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Durolane, Euflexxa, Gel-One, GenVisc 850, Hyalgan, Hymovisc, Monovisc, Orthovisc, Supartz FX, Synvisc, Synvisc-One, Visco-3, Synojoynt, Triluron, or Trivisc [documentation required]. If none are formulary, approve Gel-Syn.  2. Patient has a known allergy to avian or avian-derived products (e.g., eggs, feathers): approve if the patient has tried five formulary products from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Orthovisc, Monovisc, Durolane, Euflexxa, Hymovis, GenVisc 850, Synojoynt, or Trivisc [documentation required]. If none are formulary, approve Gel-Syn.  3. Patients who have already been started on an injection series with Gel-Syn: approve to complete the series.  Note: If the patient has received all the injections in a completed series (for each targeted joint), refer to criteria above.	1 year	Yes
Hyaluronic Acid Derivatives	GenVisc 850	sodium hyaluronate	1. Approve if the patient has tried five formulary intra-articular hyaluronic acid products from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Durolane, Euflexxa, Gel-One, Gel-Syn, Hyalgan, Monovisc, Orthovisc, Supartz FX, Synvisc, Synvisc-One, Hymovis, Visco-3, Synojoynt, Triluron, or Trivisc [documentation required]. If none are formulary, approve GenVisc 850.  2. Patient has a known allergy to avian or avian-derived products (e.g., eggs, feathers): approve if the patient has tried five formulary product from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Orthovisc, Monovisc, Durolane, Gel-Syn, Hymovis, Euflexxa, Synojoynt, or Trivisc [documentation required]. If none are formulary, approve GenVisc 850.  3. Patients who have already been started on an injection series with Genvisc 850: approve to complete the series.  Note: If the patient has received all the injections in a completed series (for each targeted joint), refer to criteria above.	1 year	Yes
Hyaluronic Acid Derivatives	Hyalgan	sodium hyaluronate	1. Approve if the patient has tried five formulary intra-articular hyaluronic acid products from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Durolane, Euflexxa, Gel-One, Gel-Syn, GenVisc 850, Monovisc, Orthovisc, Supartz FX, Synvisc, Synvisc-One, Hymovis, Visco-3, Synojoynt, Triluron, or Trivisc [documentation required]. If none are formulary, approve Hyalgan.  2. Patients who have already been started on an injection series with Hyalgan: approve to complete the series.  Note: If the patient has received all the injections in a completed series (for each targeted joint), refer to criteria above.	1 year	Yes
Hyaluronic Acid Derivatives	Hymovis	hyaluronic acid	<ol> <li>Approve if the patient has tried five formulary intra-articular hyaluronic acid products from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Durolane, Euflexxa, Gel-One, Gel-Syn, GenVisc 850, Hyalgan, Monovisc, Orthovisc, Supartz FX, Synvisc-One, Visco-3, Synojoynt, Triluron, or Trivisc [documentation required] If none are formulary, approve Hymovis.</li> <li>Patient has a known allergy to avian or avian-derived products (e.g., eggs, feathers): approve if the patient has tried five formulary products from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Orthovisc, Monovisc, Durolane, Euflexxa, Gel-Syn, GenVisc 850, Synojoynt, or Trivisc [documentation required]. If none are formulary, approve Hymovis.</li> <li>Patients who have already been started on an injection series with Hymovis: approve to complete the series.</li> <li>Note: If the patient has received all the injections in a completed series (for each targeted joint), refer to criteria above.</li> </ol>	1 year	Yes
Hyaluronic Acid Derivatives	Supartz FX	sodium hyaluronate	1. Approve if the patient has tried five formulary intra-articular hyaluronic acid products from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Durolane, Euflexxa, Gel-One, Gel-Syn, GenVisc 850, Hyalgan, Monovisc, Orthovisc, Synvisc-One, Hymovis Visco-3, Synojoynt, Triluron, or Trivisc [documentation required]. If none are formulary, approve Supartz FX.  2. Patients who have already been started on an injection series with Supartz FX: approve to complete the series.  Note: If the patient has received all the injections in a completed series (for each targeted joint), refer to criteria above.		
Derivatives	Supariz FX	injection	Note: If the patient has received all the injections in a completed series (for each targeted joint), refer to criteria above.  1. Approve if the patient has tried five other formulary intra-articular hyaluronic acid products from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Durolane, Euflexxa, Gel-One, Gel-Syn, GenVisc 850, Hyalgan, Monovisc, Orthovisc, Supartz FX, Synvisc-One, Hymovis Visco-3, Synojoynt, Triluron, or Trivisc [documentation required]. If none are formulary, approve Synvisc.	1 year	Yes
Hyaluronic Acid Derivatives	Synvisc	sodium hyaluronate injection	2. Patients who have already been started on an injection series with Synvisc: approve to complete the series.  Note: If the patient has received all the injections in a completed series (for each targeted joint), refer to criteria above.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Hyaluronic Acid Derivatives	Synvisc-One	sodium hyaluronate injection	Approve if the patient has tried five formulary intra-articular hyaluronic acid products from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Durolane, Euflexxa, Gel-One, Gel-Syn, GenVisc 850, Hyalgan, Monovisc, Orthovisc, Supartz FX, Synvisc, Hymovis Visco-3, Synojoynt, Triluron, or Trivisc [documentation required]. If none are formulary, approve Synvisc-One.	1 year	Yes
Hyaluronic Acid Derivatives	Triluron	sodium hyaluronate 1% injection	<ol> <li>Approve if the patient has tried five formulary alternatives from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Durolane, Euflexxa, Gel-One, Gel-Syn, GenVisc 850, Hyalgan, Monovisc, Orthovisc, Supartz FX, Synvisc-One, Hymovis, Synojoynt, Visco-3, or Trivisc [documentation required]. If none are formulary, approve.</li> <li>Patients who have already been started on an injection series with Triluron: approve to complete the series.</li> <li>Note: If the patient has received all the injections in a completed series (for each targeted joint), refer to criteria above.</li> </ol>	1 year	Yes
Hyaluronic Acid Derivatives	Trivisc	sodium hyaluronate injection	<ol> <li>Approve if the patient has tried five formulary intra-articular hyaluronic acid product from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Durolane, Euflexxa, Gel-One, Gel-Syn, GenVisc 850, Hyalgan, Monovisc, Orthovisc, Supartz FX, Synvisc, Synvisc-One, Hymovis, Synojoynt, Triluron, or Visco-3 [documentation required]. If none are formulary, approve Trivisc.</li> <li>Patient has a known allergy to avian or avian-derived products (e.g., eggs, feathers): approve if the patient has tried five formulary products from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Durolane, Euflexxa, Orthovisc, Monovisc, Hymovis, Gel-Syn, Synojoynt, or GenVisc 850 [documentation required]. If none are formulary, approve Trivisc.</li> <li>Patients who have already been started on an injection series with Trivisc: approve to complete the series.</li> <li>Note: If the patient has received all the injections in a completed series (for each targeted joint), refer to criteria above.</li> </ol>	1 year	Yes
Hyaluronic Acid Derivatives	Visco-3	sodium hyaluronate injection	1. Approve if the patient has tried five formulary alternatives from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if one is formulary): Durolane, Euflexxa, Gel-One, Gel-Syn, GenVisc 850, Hyalgan, Monovisc, Orthovisc, Supartz FX, Synvisc-One, Hymovis, Synojoynt, Triluron, or Trivisc [documentation required]. If none are formulary, approve Visco-3.  2. Patients who have already been started on an injection series with Visco-3: approve to complete the series.  Note: If the patient has received all the injections in a completed series (for each targeted joint), refer to criteria above.	1 year	Yes
Hyperlipidemia - Proprotein Convertase Subtilisin Kexin Type 9 (PCSK9) Inhibitors and Related Agents	Leqvio	inclisiran subcutaneous injection	Established Cardiovascular Disease; Heterozygous Familial Hypercholesterolemia; Primary Hyperlipidemia (all diagnoses in a patient ≥ 18 years of age).  Approve if the patient has tried Repatha or Praluent, if formulary. If neither are formulary, approve.	1 year	Yes
Hyperlipidemia - Proprotein Convertase Subtilisin Kexin Type 9 (PCSK9) Inhibitors and Related Agents	Praluent	alirocumab injection for subcutaneous use	See Proprotein Convertase Subtilisin Kexin Type 9 Related Products Care Value Policy criteria  **For Praluent only**	1 year	Yes
Hypolipoproteinemics	Welchol packets and tablets	colesevelam packets and tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Hypolipoproteinemics	Zetia	ezetimibe tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Hypoxia-Inducible Factor Prolyl Hydroxylase Inhibitor	Vafseo	vadadustat tablets	Treatment of anemia due to chronic kidney disease in a patient ≥ 18 years of age.  Approve if the patient meets the following (1 and 2):  1. Patient has been receiving dialysis for at least 3 consecutive months; AND  2. Patient has tried, and according to the prescriber, experienced inadequate efficacy OR significant intolerance with one of the following: an epoetin alfa product or Aranesp or Mircera.  Note: Examples of epoetin alfa products are Procrit, Epogen, and Retacrit.	1 year	Yes
Idiopathic Pulmonary Fibrosis Agents	Esbriet	pirfenidone tablets and capsules	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Idiopathic Pulmonary Fibrosis Agents		pirfenidone 534 mg tablet	Idiopathic pulmonary fibrosis. Patient meets both of the following (i and ii):  i. Patient has tried generic pirfenidone tablets; AND  Note: True generic tablets are available in 267 mg tablets.  ii. Patient cannot continue to use generic pirfenidone tablets due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent product which, per the prescriber, would result in a significant allergy or serious adverse reaction.	1 year	Yes
Immune Globulins - Intravenous (IVIG) and Subcutaneous (SCIG)	Gammaked	immune globulin injection (human), 10%	<ol> <li>If using via the subcutaneous (SC) route: approve if the patient has tried three products from the following list, if formulary (or two if two are formulary or one if one is formulary): Cuvitru, Hizentra, Xembify, Cutaquig, Gamunex-C or Gammagard Liquid. If none are formulary, approve.</li> <li>If using via the intravenous (IV) route: approve if the patient has tried three formulary IVIG products from the following list, if formulary (or two if two are formulary or one if only one is formulary): Alyglo, Asceniv, Bivigam, Flebogamma DIF, Gammagard Liquid, Gammagard S/D, Gammaplex, Gamunex-C, Octagam, Privigen or Panzyga. If none are formulary, approve.</li> </ol>	1 year	Yes
Immune Globulins - Subcutaneous (SCIG)	Cutaquig		Primary Immunodeficiencies:  Note: Examples of primary immunodeficiences include, but are not limited to, congenital agammaglobulinemia, X-linked agammaglobulinemia, severe combined immunodeficiency, common variable immunodeficiency.  Approve if the patient has tried three products from the following list, if formulary (or two if two are formulary or one if one is formulary): Cuvitru, Hizentra, Xembify, Gamunex-C, Gammagard Liquid, or Gammaked. If none are formulary, approve.	1 year	Yes
Immunological Agents	Cingair	reslizumab for intravenous injection	Asthma with an eosinophilic phenotype.  Approve if the patient meets one of the following (A or B):  A. Initial therapy in a patient ≥ 18 years of age: Patient has tried one formulary alternative from the following list: Nucala or Fasenra. If neither is formulary, approve if the patient has tried Dupixent. If Dupixent is non-formulary, approve;  OR  B. Patient has already been started on therapy with Cingair.	1 year	Yes
Immunosuppressant Agents	Envarsus XR	tacrolimus extended- release tablets	<ol> <li>Approve if the patient has tried and cannot take tacrolimus immediate-release capsules (Prograf, generics), if formulary. If tacrolimus immediate-release capsules (Prograf, generics) are non-formulary, approve.</li> <li>Approve if the patient has the CYP3A5*1 allele.</li> <li>Note: The CYP3A5*1 allele is a gene variant determined by testing that may confer faster metabolism of certain medications.</li> <li>If the patient has already started on therapy with Envarsus XR, approve.</li> </ol>	1 year	Yes
Immunosuppressant Agents – Methotrexate Injections	Otrexup	methotrexate injection for subcutaneous use; 10mg, 12.5 mg, 15 mg, 17.5 mg, 20 mg, 22.5 mg, 25 mg	Approve if the patient has tried Rasuvo, if formulary. If Rasuvo is non-formulary, approve if, according to the prescriber, the patient and caregiver are unable to administer methotrexate injection (NOT including Otrexup or Rasuvo).	1 year	Yes
Immunosuppressant Agents – Oral Methotrexate Agents	Xatmep	methotrexate 2.5 mg/mL oral solution	Approve if the patient has tried Jylamvo, if formulary. If Jylamvo is non-formulary, approve if the patient meets one of the following (1 or 2):  1. Patient cannot swallow or has difficulty swallowing oral methotrexate tablets; OR  2. The dose prescribed cannot be obtained using whole methotrexate tablets.	1 year	Yes
Immunosuppressant Agents – Oral Methotrexate Agents	Jylamvo	methotrexate 2 mg/mL oral solution	Approve if the patient has tried Xatmep, if formulary. If Xatmep is non-formulary, approve if the patient meets one of the following (1 or 2):  1. Patient cannot swallow or has difficulty swallowing oral methotrexate tablets; OR  2. The dose prescribed cannot be obtained using whole methotrexate tablets.	1 year	Yes
Inflammatory Bowel Agents	Canasa	mesalamine rectal suppository	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Inflammatory Bowel Agents	Delzicol	mesalamine delayed- release capsule	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Inflammatory Bowel Agents	Lialda	mesalamine delayed- release tablet	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Inflammatory Bowel Agents	Dipentum	olsalazine capsule	Approve if the patient has tried two products from the following list (if two are formulary, or one if one is formulary): mesalamine delayed-release tablets (Asacol HD, generics), sulfasalazine (generics), mesalamine delayed-release tablets (Lialda, generics), mesalamine delayed-release capsules (Apriso, generics) or Pentasa. If none are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Inflammatory Conditions	Plaquenil	sulfate tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Inflammatory Conditions	Sovuna	hydroxychloroquine sulfate 200 mg, 300 mg	<ol> <li>Direct to generic hydroxychloroquine tablets.</li> <li>Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use generic hydroxychloroquine tablets.</li> </ol>	1 year	Yes
Inflammatory Conditions – Infused Non-TNF Biologics	Orencia IV	abatacept injection for intravenous use	Juvenile Idiopathic Arthritis; Psoriatic Arthritis; Rheumatoid Arthritis.  1. Patient has tried at least one biologic: Approve.  Examples: a tocilizumab product (e.g., Actemra intravenous [IV] or subcutaneous), a sarilumab product (Kevzara), an etanercept product (e.g., Enbrel, biosimilars), an adalimumab product (e.g., Humira, biosimilars), a certolizumab pegol product (e.g., Cimzia), a golimumab product (e.g., Simponi Aria or subcutaneous), an infliximab IV product (e.g., Remicade, biosimilars), a rituximab product (e.g., Rituxan intravenous, biosimilars), a secukinumab product (e.g., Cosentyx IV or SC), an ixekizumab product (e.g., Taltz), a guselkumab product (e.g., Tremfya), or a ustekizumab product (e.g., Stelara SC). If none are formulary, approve.  2. According to the prescriber, the patient previously experienced a serious infection: Approve.  3. Patient is currently taking Orencia intravenous or subcutaneous: Approve if the patient has been established on Orencia intravenous or subcutaneous for ≤ 3 months.  4. Patient has been started on Orencia intravenous or subcutaneous for < 3 months: Refer to the appropriate criteria above.  Caraft-Versus-Host Disease – Prevention: Approve.	1 year	Yes
Inflammatory Conditions – Infused Non-TNF Biologics	Entyvio SC	vedolizumab for subcutabeous injection	See standard Inflammatory Conditions (Entyvio SC) Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, Alternate, 2) Legacy OR FLEX Formulary policies.	See PSM duration	Yes
			If Actemra IV AND Tyenne IV are both formulary (or one is formulary) [For all indications]: Approve if the patient meets ONE of the following (1 or 2):  1. Patient meets BOTH of the following (A and B):  A. Patient has tried BOTH Actrema IV and Tyenne IV (if both are formulary or one if one is formulary); AND  B. Patient cannot continue to use each of the formulary alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.  2. Patient has already been started on therapy with Tofidence.  If both Actemra IV AND Tyenne IV are non-formulary: Polyarticular Juvenile Idiopathic Arthritis; Rheumatoid Arthritis.		
Inflammatory Conditions – Infused Non-TNF Biologics –			1. Patient has tried at least one biologic: Approve.  Examples: an abatacept product (e.g., Orencia intravenous [IV] or subcutaneous), a sarilumab product (e.g., Kevzara), an etanercept product (e.g., Enbrel, biosimilars), an adalimumab product (e.g., Humira, biosimilars), a certolizumab pegol product (e.g., Cimzia), a golimumab product (e.g., Simponi Aria or subcutaneous), an infliximab IV product (e.g., Remicade, biosimilars), a rituximab product (e.g., Rituxan intravenous, biosimilars). If none are formulary, approve.  2. Patient is currently taking a tocilizumab product (e.g., Actemra IV or SC or Tyenne IV or Tofidence IV): Approve if the patient has been established on a tocilizumab product (e.g., Actemra IV or SC or Tyenne IV or Tofidence IV) for ≥ 90 days.  If both Actemra IV and Tyenne IV are non-formulary:  Giant Cell Arteritis; Polymyalqia Rhematica; Systemic Juvenile Idiopathic Arthritis, Castleman's Disease, Chimeric Antigen Receptor (CAR) T Cell-Induced Severe or Life-Threatening Cytokine Release Syndrome, or Inflormatical Associated with Characters Interior Polymyalqia Rhematica, Astociated with Characters Interior Polymyalqia Rhematica, Systemic Juvenile Idiopathic Appropriate Interior Polymyalqia Rhematica Interior Poly		
Tocilizumab Intravenous Agents	Tofidence IV	tocilizumab-bavi intravenous infusion	Inflammatory Arthritis Associated with Checkpoint Inhibitor Therapy: Approve.  Note: Examples of checkpoint inhibitors are Keytruda (pembrolizumab intravenous infusion), Opdivo (nivolumab intravenous infusion), Yervoy (ipilimumab intravenous infusion), Tecentriq (atezolizumab intravenous infusion), Bavencio (avelumab intravenous infusion), Imfinzi (durvalumab intravenous infusion), and Libtayo (cemiplimab-rwlc intravenous infusion).	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
			If any of the following ustekinumab intravenous (IV) products are formulary, Selarsdi IV, Stelara IV or ustekinumab unbranded IV, Steqeyma IV, Wezlana IV, Pyzchiva IV or ustekinumab-ttwe IV, or Yesintek IV, approve if the patient meets BOTH of the following (A and B):  A. Patient has tried ALL of the following: 1) Stelara IV or ustekinumab unbranded IV, 2) Steqeyma IV, 3) Wezlana IV, 4) Pyzchiva IV or ustekinumab-ttwe IV, 5) Yesintek IV, and 6) Selarsdi IV, if formulary; AND  Note: Pyzchiva IV and ustekinumab-ttwe IV count as one alternative. Stelara IV and ustekinumab unbranded IV count as one alternative.  Patient cannot continue to use all the formulary ustekinumab products due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.		
Inflammatory			If none of the ustekinumab intravenous (IV) products are formulary, Selarsdi IV, Stelara IV or ustekinumab unbranded IV, Steqeyma IV, Wezlana IV, Pyzchiva IV or ustekinumab-ttwe IV, Yesintek IV, approve if the patient meets ONE of the following (1 or 2):  1. <u>Ulcerative Colitis, for an induction regimen in a patient ≥ 18 years of age.</u> Approve if the patient has tried two of 1) Entyvio intravenous (IV)/subcutaneous (SC), 2) Skyrizi IV/SC, or 3) Tremfya IV/SC, if two are formulary (or one if one is formulary). If none are formulary, approve.		
Conditions – Infused Non-TNF Biologics – ustekinumab Agents	Otulfi IV	ustekinumab- aauz for IV infusion	2. Crohn's Disease, for an induction regimen in a patient ≥ 18 years of age.  Approve if the patient has tried one of 1) Entyvio IV/SC, 2) Skyrizi IV/On Body, or 3) Tremfya IV/SC, if formulary. If none are formulary, approve.	1 dose	Yes
Inflammatory Conditions – Infused			Direct to ustekinumab-ttwe IV, if formulary.  If ustekinumab-ttwe IV is non-formulary, If any of the following ustekinumab intravenous (IV) products are formulary, Selarsdi IV, Stelara IV or ustekinumab unbranded IV, Steqeyma IV, Wezlana IV, Yesintek IV, Otulfi IV, approve if the patient meets BOTH of the following (A and B):  A. Patient has tried ALL of the following: 1) Stelara IV or ustekinumab unbranded IV, 2) Steqeyma IV, 3) Wezlana IV, 4) Yesintek IV, 5) Otulfi IV, and 6) Selarsdi IV, if formulary; AND  Note: Stelara IV and ustekinumab unbranded IV count as one alternative.  B. Patient cannot continue to use all the formulary ustekinumab products due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.  If none of the ustekinumab intravenous (IV) products are formulary, Selarsdi IV, Stelara IV or ustekinumab unbranded IV, Steqeyma IV, Wezlana IV, Yesintek IV, Otulfi IV, approve if the patient meets ONE of the following (1 or 2):  1. Ulcerative Colitis, for an induction regimen in a patient ≥ 18 years of age.  Approve if the patient has tried two of 1) Entyvio intravenous (IV)/subcutaneous (SC), 2) Skyrizi IV/SC, or 3) Tremfya IV/SC, if two are formulary (or one if one is formulary). If none are formulary, approve.		
Non-TNF Biologics – ustekinumab Agents	Pyzchiva IV	ustekinumab- ttwe for IV infusion	2. Crohn's Disease, for an induction regimen in a patient ≥ 18 years of age.  Approve if the patient has tried one of 1) Entyvio IV/SC, 2) Skyrizi IV/On Body, or 3) Tremfya IV/SC, if formulary. If none are formulary, approve.	1 dose	Yes
			If any of the following ustekinumab intravenous (IV) products are formulary, Selarsdi IV, Stelara IV or ustekinumab unbranded IV, Wezlana IV, Yesintek IV, Pyzchiva IV or ustekinumab-ttwe IV, or Otulfi IV, approve if the patient meets BOTH of the following (A and B):  A. Patient has tried ALL of the following: 1) Stelara IV or ustekinumab unbranded IV, 2) Wezlana IV, 3) Yesintek IV, 4) Pyzchiva IV or ustekinumab-ttwe IV, 5) Otulfi IV, and 6) Selarsdi IV, if formulary; AND Note: Pyzchiva IV and ustekinumab-ttwe IV count as one alternative. Stelara IV and ustekinumab unbranded IV count as one alternative.  B. Patient cannot continue to use all the formulary ustekinumab products due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.		
Inflammatory			If none of the ustekinumab intravenous (IV) products are formulary, Selarsdi IV, Stelara IV or ustekinumab unbranded IV, Wezlana IV, Yesintek IV, Pyzchiva IV or ustekinumab-ttwe IV, Otulfi IV, approve if the patient meets ONE of the following (1 or 2):  1. <u>Ulcerative Colitis, for an induction regimen in a patient ≥ 18 years of age.</u> Approve if the patient has tried two of 1) Entyvio intravenous (IV)/subcutaneous (SC), 2) Skyrizi IV/SC, or 3) Tremfya IV/SC, if two are formulary (or one if one is formulary). If none are formulary, approve.		
Conditions – Infused Non-TNF Biologics – ustekinumab Agents	Steqeyma IV	ustekinumab-stba for IV infusion	2. Crohn's Disease, for an induction regimen in a patient ≥ 18 years of age.  Approve if the patient has tried one of 1) Entyvio IV/SC, 2) Skyrizi IV/On Body, or 3) Tremfya IV/SC, if formulary. If none are formulary, approve.	1 dose	Yes

					2025 NPF
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	Excluded Medicaiton
Inflammatory Conditions – Infused Non-TNF Biologics – ustekinumab Agents	ustekinumab IV (unbranded version of Stelara IV)	ustekinumab IV	Direct to Stelara intravenous (IV) [brand], if formulary.  If Stelara IV (brand) is non-formulary:  If any of the following ustekinumab IV products are formulary, Selarsdi IV, Steqeyma IV, Wezlana IV, Yesintek IV, Pyzchiva IV, or ustekinumab-ttwe IV, or Otulfi IV, approve if the patient meets BOTH of the following (A and B):  A. Patient has tried ALL of the following: 1) Steqeyma IV, 2) Wezlana IV, 3) Yesintek IV, 4) Pyzchiva IV or ustekinumab-ttwe IV, 5) Otulfi IV, and 6) Selarsdi IV, if formulary; AND  Note: Pyzchiva IV and ustekinumab-ttwe IV count as one alternative.  B. Patient cannot continue to use all the formulary ustekinumab products due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.  If none of the ustekinumab intravenous (IV) products are formulary, Selarsdi IV, Steqeyma IV, Wezlana IV, Yesintek IV, Pyzchiva IV, or ustekinumab-ttwe IV, or Otulfi IV, approve if the patient meets ONE of the following (1 or 2):  1. Ulcerative Colitis, for an induction regimen in a patient ≥ 18 years of age.  Approve if the patient has tried two of 1) Entyvio intravenous (IV)/subcutaneous (SC), 2) Skyrizi IV/SC, or 3) Tremfya IV/SC, if formulary. If none are formulary, approve.	1 dose	Yes
Inflammatory Conditions – Infused Non-TNF Biologics –	,		If any of the following ustekinumab intravenous (IV) products are formulary, Selarsdi IV, Stelara IV or ustekinumab unbranded IV, Steqeyma IV, Yesintek IV, Pyzchiva IV or ustekinumab-ttwe IV, or Otulfi IV, approve if the patient meets BOTH of the following (A and B):  A. Patient has tried ALL of the following: 1) Stelara IV or ustekinumab unbranded IV, 2) Steqeyma IV, 3) Yesintek IV, 4) Pyzchiva IV or ustekinumab-ttwe IV, 5) Otulfi IV, and 6) Selarsdi IV, if formulary; AND  Note: Pyzchiva IV and ustekinumab-ttwe IV count as one alternative. Stelara IV and ustekinumab unbranded IV count as one alternative.  B. Patient cannot continue to use all the formulary ustekinumab products due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.  If none of the ustekinumab intravenous (IV) products are formulary, Selarsdi IV, Stelara IV or ustekinumab unbranded IV, Steqeyma IV, Yesintek IV, Pyzchiva IV, or ustekinumab-ttwe IV, or Otulfi IV, approve if the patient meets ONE of the following (1 or 2):  1. Ulcerative Colitis, for an induction regimen in a patient ≥ 18 years of age.  Approve if the patient has tried two of 1) Entyvio intravenous (IV)/subcutaneous (SC), 2) Skyrizi IV/SC, or 3) Tremfya IV/SC, if formulary, approve.  2. Crohn's Disease, for an induction regimen in a patient ≥ 18 years of age.  Approve if the patient has tried one of 1) Entyvio IV/SC, 2) Skyrizi IV/On Body, or 3) Tremfya IV/SC, if formulary. If none are formulary, approve.	1 dose	Yes
Inflammatory Conditions – Infused Non-TNF Biologics –	ustekinumab-aekn SC (unbranded version of Selarsdi SC)	ustekinumab-aekn SC (unbranded version of Selarsdi SC)	See standard Inflammatory Conditions – ustekinumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, 2) Alternate, 3) Legacy OR FLEX Formulary policies.	See PSM duration	Yes
Inflammatory Conditions – Infused Non-TNF Biologics – ustekinumab Agents  Inflammatory Conditions – Infused Non-TNF Biologics –	Otulfi SC  Pyzchiva SC	ustekinumab- ttwe SC	See standard Inflammatory Conditions – ustekinumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, 2) Alternate, 3) Legacy OR FLEX Formulary policies.  See standard Inflammatory Conditions – ustekinumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, 2) Alternate, 3) Legacy OR FLEX Formulary policies.	See PSM duration	Yes
ustekinumab Agents	Steqeyma SC	ustekinumab-stba SC	See standard Inflammatory Conditions – ustekinumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, 2) Alternate, 3) Legacy OR FLEX Formulary policies.	duration	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
			Patient meets standard Inflammatory Conditions – Infliximab Intravenous Products Prior Authorization Policy criteria AND		
Inflammatory Conditions - Infused		Infliximab- axxq for	Approve if the patient meets BOTH of the following (A and B):  A. Patient has tried Inflectra; AND  B. Patient cannot continue to use Inflectra due to a formulation difference in the inactive ingredients(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.	See PA	
TNF antagonists	Avsola	intravenous use	Note: An approval will be entered for Inflectra if the Infliximab Intravenous Products Prior Authorization criteria are met, but the remaining criteria are not met.	duration	Yes
Inflammatory Conditions - Infused TNF antagonists	Remicade and authorized generic infliximab	infliximab injection for intravenous use	Patient meets standard Inflammatory Conditions – Infliximab Intravenous Products Prior Authorization Policy criteria AND  Approve if the patient meets BOTH of the following (A and B):  A. Patient has tried Inflectra; AND  B. Patient cannot continue to use Inflectra due to a formulation difference in the inactive ingredients(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.  Note: An approval will be entered for Inflectra if the Infliximab Intravenous Products Prior Authorization criteria are met, but the remaining criteria are not met.	See PA duration	Yes
			Patient meets standard Inflammatory Conditions – Infliximab Intravenous Products Prior Authorization Policy criteria AND		
Inflammatory Conditions - Infused TNF antagonists	Renflexis	Infliximab-abda for intravenous use	Approve if the patient meets BOTH of the following (A and B):  A. Patient has tried Inflectra; AND  B. Patient cannot continue to use Inflectra due to a formulation difference in the inactive ingredients(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.  Note: An approval will be entered for Inflectra if the Infliximab Intravenous Products Prior Authorization criteria are met, but the remaining criteria are not met.	See PA duration	Yes
Inflammatory Conditions – Janus				See PSM	
Kinase Inhibitors	Olumiant	baricitinib tablets	See standard Inflammatory Conditions (Olumiant) Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, Alternate, 2) Legacy OR FLEX Formulary policies.	duration	Yes
Inflammatory		coulding make for	Approve if the patient has tried Cosentyx subcutaneous, if formulary, AND the patient is unable to continue to use a subcutaneous dosage form.  If Cosentyx subcutaneous is non-formulary:  Ankylosing Spondylitis: Psoriatic Arthritis. Approve if the patient meets ONE of the following (1 or 2):  1. Patient meets BOTH of the following (A and B):  A. Patient has tried Taltz, if formulary; AND  Note: If Taltz is non-formulary, would still need to meet criterion B.  B. Patient has tried at least one other biologic.  Examples of other biologics: an adalimumab product (e.g., Humira, biosimilars), a certolizumab pegol product (e.g., Cimzia), an etanercept product (e.g., Enbrel, biosimilars), an infliximab product (e.g., Remicade, biosimilars), a golimumab product (e.g., Simponi Aria or subcutaneous), or an abatacept product (e.g., Orencia intravenous or subcutaneous). If none are formulary, approve.  2. Patient is currently receiving Cosentyx (IV or SC) for ≤ 90 days, approve.  Note: If the patient has been on Cosentyx (IV or SC) for < 90 days, refer to criterion #1.  Non-Radiographic Spondyloarthritis.  Non-Radiographic Spondyloarthritis.  Approve if the patient has tried Taltz, if formulary, OR  Note: If Taltz is non-formulary, approve.  2. Patient is currently receiving Cosentyx (IV or school of the following (1 or 2):  1. Approve if the patient has tried Taltz, if formulary, OR  Note: If Taltz is non-formulary, approve.		
Conditions – SC Non- TNF Biologics	Cosentvx IV	secukinumab for intravenous injection	2. Patient is currently receiving Cosentyx (intravenous (IV) or subcutaneous (SC): Patient has been established on Cosentyx (IV or SC) for ≥ 90 days, approve.  Note: If the patient has been on Cosentyx (IV or SC) for < 90 days, refer to criterion #1.	1 vear	Yes
Inflammatory Conditions – SC Non- TNF Biologics	Kevzara	sarilumab subcutaneous injection	See standard Inflammatory Conditions (Kevzara) Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, Alternate, 2) Legacy OR FLEX Formulary policies.	See PSM duration	Yes
Inflammatory Conditions – SC Non-TNF Biologics	Kineret	anakinra SC injection	See standard Inflammatory Conditions (Kineret) Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, Alternate, 2) Legacy OR FLEX Formulary policies.	See PSM duration	Yes
Inflammatory Conditions – SC Non- TNF Biologics	Ilumya	tildrakizumab SC injection	See standard Inflammatory Conditions (Ilumya) Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, Alternate, 2) Legacy OR FLEX Formulary policies.	See PSM duration	Yes

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
Inflammatory Conditions – SC Non-		abatasant injection for		See PSM	
TNF Biologics	Orencia for SC use	abatacept injection for subcutaneous use	See standard Inflammatory Conditions (Orencia SC) Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, Alternate, 2) Legacy OR FLEX Formulary policies.	duration	Yes
Inflammatory	Cronola for CO acc	bimekizumab-bkzx	dee standard illiminatory conditions ( Grencia GO) interior depending management in only for industrial referred, right chomitance, and basic informations in ordinary policies.	duration	100
Conditions – SC Non-		subcutaneous		See PSM	
TNF Biologics	Bimzelx	injection	See standard Inflammatory Conditions (Bimzelx) Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, Alternate, 2) Legacy OR FLEX Formulary policies.	duration	Yes
Inflammatory		brodalumab for			
Conditions – SC Non-		subcutaneous		See PSM	
TNF Biologics	Siliq	injection	See standard Inflammatory Conditions (Siliq) Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, Alternate, 2) Legacy OR FLEX Formulary policies.	duration	Yes
Inflammatory					
Conditions – SC Non-	ustekinumab SC	ustekinumab SC		C DCM	
TNF Biologics – ustekinumab Agents	(unbranded version of Stelara SC)	(unbranded version of Stelara SC)	See standard Inflammatory Conditions – ustekinumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, 2) Alternate, 3) Legacy OR FLEX Formulary policies.	See PSM duration	Yes
ustekillulliab Agelits	oi Stelala SC)	Stelala SC)	See standard illinaminatory Conditions – ustexinumab Froducts Freienred Specialty Management Policy for National Freienred, Flight Performance, and basic Politicians 1) Choice, 2) Attendate, 3) Legacy OK PLEA Politicians policies.	uurauon	162
Inflammatory					
Conditions – SC Non-					
TNF Biologics –		ustekinumab for SC		See PSM	
ustekinumab Agents	Wezlana SC	injection	See standard Inflammatory Conditions – usteknumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, 2) Alternate, 3) Legacy OR FLEX Formulary policies.	duration	Yes
Inflammatory		golimumab			
Conditions – SC TNF		subcutaneous		See PSM	
Antagonists	Simponi SC	injection	See standard Inflammatory Conditions (Simponi SC) Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, Alternate, 2) Legacy OR FLEX Formulary policies.	duration	Yes
Inflammatory					
Conditions – SC TNF		certolizumab powder		See PSM	
Antagonists	Cimzia	for injection	See standard Inflammatory Conditions (Cimzia) Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, Alternate, 2) Legacy OR FLEX Formulary policies.	duration	Yes
Inflammatory					
Conditions – SC TNF Antagonists -	Yuflyma and	adalimumab-aaty subcutaneous		See PSM	
Adalimumab Agents	adalimumab-aaty	injection	See standard Inflammatory Conditions – Adalimumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, 2) Alternate, 3) Legacy OR FLEX Formulary policies.	duration	Yes
/ taaiiiiaiiiab / tgonto	addilliand daty	Injudation	Constitution of the state of th	duration	100
Inflammatory					
Conditions – SC TNF		adalimumab-aqvh			
Antagonists -		subcutaneous		See PSM	
Adalimumab Agents	Yusimry	injection	See standard Inflammatory Conditions - Adalimumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, 2) Alternate, 3) Legacy OR FLEX Formulary policies.	duration	Yes
Inflammatory					
Conditions – SC TNF		adalimumab-afzb			
Antagonists -		subcutaneous		See PSM	v
Adalimumab Agents	Abrilada	injection	See standard Inflammatory Conditions – Adalimumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, 2) Alternate, 3) Legacy OR FLEX Formulary policies.	duration	Yes
1					
Inflammatory Conditions – SC TNF		adalimumab-fkjp subcutaneous			
Antagonists -		injection (unbranded		See PSM	
Adalimumab Agents	Adalimumab-fkip	version of Hulio)	See standard Inflammatory Conditions – Adalimumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, 2) Alternate, 3) Legacy OR FLEX Formulary policies.	duration	Yes
			The state of the s	30.0001	
Inflammatory					
Conditions – SC TNF		adalimumab-atto			
Antagonists -		subcutaneous		See PSM	
Adalimumab Agents	Amjevita	injection	See standard Inflammatory Conditions - Adalimumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, 2) Alternate, 3) Legacy OR FLEX Formulary policies.	duration	Yes
		_			
Inflammatory					
Conditions – SC TNF		adalimumab-bwwd			
Antagonists -		subcutaneous		See PSM	v
Adalimumab Agents	Hadlima	injection	See standard Inflammatory Conditions – Adalimumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, 2) Alternate, 3) Legacy OR FLEX Formulary policies.	duration	Yes

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
Inflammatory					
Conditions – SC TNF		adalimumab-fkjp			
Antagonists -		subcutaneous		See PSM	
Adalimumab Agents	Hulio	injection	See standard Inflammatory Conditions - Adalimumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, 2) Alternate, 3) Legacy OR FLEX Formulary policies.	duration	Yes
Inflammatory Conditions – SC TNF					
Antagonists -				See PSM	
Adalimumab Agents	Humira	adalimumab injection	See standard Inflammatory Conditions – Adalimumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, 2) Alternate, 3) Legacy OR FLEX Formulary policies.	duration	Yes
_					
Inflammatory					
Conditions – SC TNF		adalimumab-adaz		See PSM	
Antagonists - Adalimumab Agents	Hyrimoz	subcutaneous injection	  See standard Inflammatory Conditions – Adalimumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, 2) Alternate, 3) Legacy OR FLEX Formulary policies.	duration	Yes
/ tadimamab / tgonto	riyiiiioz	Injection	dee standard illiminatory conditions—Administration for the formation of t	duration	100
Inflammatory					
Conditions – SC TNF		adalimumab-aacf			
Antagonists -	Idacio and	subcutaneous		See PSM	V
Adalimumab Agents	adalimumab-aacf	injection	See standard Inflammatory Conditions – Adalimumab Products Preferred Specialty Management Policy for National Preferred, High Performance, and Basic Formularies 1) Choice, 2) Alternate, 3) Legacy OR FLEX Formulary policies.	duration	Yes
		ropeginterferon alfa-			
		2b-njft subcutaneous			
Interferons	Besremi	injection	See Oncology (Injectable) – Besremi Prior Authorization Policy criteria.	1 year	Yes
					MSB Exclusion
Iron Donlagoment			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		*This criteria
Iron Replacement (Injectable)	Feraheme	ferumoxytol injection	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	applies only to the NPF
(mjestažie)	- Granding	i or annoxytor injection	processor, was read to the read to the significant and the read to	. , ,	
		ferric derisomaltose	1. Approve if the patient has tried three products from the following list (if three are formulary, or two if two are formulary, or one if one is formulary): Venofer, sodium ferric gluconate complex (Ferrlecit, generics), or Injectafer. If none are		
Iron Replacement		injection for	formulary, approve.		
(Injectable)	Monoferric	intravenous use	2. Patient does NOT have chronic kidney disease, approve if the patient has tried Injectafer. If Injectafer is non-formulary, approve.	1 year	Yes
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		MSB Exclusion *This criteria
Irritable Bowel			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Syndrome Agents	Lotronex	alosetron tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].		the NPF
		isotretinoin capsules	Approve if the patient has tried three of the following: isotretinoin capsules (Absorica [not LD]), Accutane, Amnesteem, Claravis, or Zenatane, if formulary (or two if two are formulary or one if one is formulary). If none are formulary,		
Isotretinoin Products	Absorica LD	low dose	approve.	1 year	Yes
		montelukast sodium	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		MSB Exclusion *This criteria
Leukotriene Pathway		tablets, chewable	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Inhibitors	Singulair tablets	tablets, granules	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Leukotriene Pathway	zileuton extended-	zileuton extended-	1. Approve if the patient has tried one of the following, if one is formulary: montelukast (Singulair, generics) or zafirlukast (Accolate, generics). If none are formulary, approve.		
Inhibitors	release tablets	release tablets	2. Approve if the patient has already been started on therapy with a zileuton-containing product (e.g., zileuton ER tablets, Zyflo).	1 year	Yes
Leukotriene Pathway Inhibitors	Zuflo	zileuton tablets	1. Approve if the patient has tried one of the following, if one is formulary: montelukast (Singulair, generics) or zafirlukast (Accolate, generics). If neither are formulary, approve.	1 year	Voc
HIMBROIS	Zyflo	Zileutori tablets	2. Approve if the patient has already been started on therapy with a zileuton-containing product (e.g., zileuton ER tablets, Zyflo).  1. Approve if the patient has tried Striverdi Respimat, if formulary. If Striverdi is non-formulary, approve.	1 year	Yes
			2. Patient who is unable to coordinate breath and actuation with a metered-dose inhaler (MDI): approve.		
			3. Patient with asthma: Approve if the patient is using Serevent Diskus concomitantly with an inhaled corticosteroid or an inhaled corticosteroid-containing product.		
Long-Acting Beta-		salmeterol xinafoate	4. Patient with exercise induced bronchospasm without asthma: approve.		
Agonists (Inhalers)	Serevent Diskus	inhalation powder	Note: A patient with exercise-induced bronchospasm and asthma should be referred to criterion #3.	1 year	Yes
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		MSB Exclusion
Long-Acting Beta-		formoterol fumarate	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		*This criteria applies only to
Agonists (nebulized)	Perforomist	inhalation solution	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
,		•			

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Therapy Class	Diana Name	Generic Name	Commercian E Criteria	Duration	Wedication
Long-Acting Muscarinic Antagonist (LAMA)/Long-Acting Beta-Agonist (LABA) Combination Inhalers	Bevespi Aerosphere	glycopyrrolate and formoterol fumarate inhalation aerosol	<ol> <li>Approve if the patient has tried three of Anoro Ellipta, Duaklir Pressair, or Stiolto Respimat, if three are formulary, or two if two are formulary, or one if one is formulary. If none are formulary, approve.</li> <li>If the patient has a low inspiratory flow rate and is unable to use a dry powder inhaler (DPI), approve if the patient has tried Stiolto Respimat, if formulary. If Stiolto Respimat is non-formulary, approve.</li> </ol>	1 year	Yes
Long-Acting Muscarinic Antagonist (LAMA)/Long-Acting Beta-Agonist (LABA) Combination Inhalers	Duaklir Pressair	aclidinium bromide and formoterol fumarate inhalation powder	<ol> <li>Approve if the patient has tried three of Anoro Ellipta, Bevespi Aerosphere, or Stiolto Respimat, if three are formulary, or two if two are formulary or one if one is formulary. If none are formulary, approve.</li> <li>If the patient is unable to coordinate breath and actuation with a metered-dose inhaler (MDI), approve if the patient has tried Anoro Ellipta, if formulary. If Anoro Ellipta is non-formulary, approve.</li> </ol>	1 year	Yes
, , ,	umeclidinium and vilanterol inhalation powder	umeclidinium and vilanterol inhalation powder	Direct to Anoro Ellipta (brand), if formulary. If Anoro Ellipta (brand) is non-formulary:  1. Approve if the patient has tried three of Bevespi Aerosphere, Duaklir Pressair, Stiolto Respimat, if three are formulary, or two if two are formulary, or one if one is formulary. If none are formulary, approve.  2. If the patient is unable to coordinate breath and actuation with a metered-dose inhaler (MDI), approve if the patient has tried Duaklir Pressair, if formulary. If Duaklir Pressair is non-formulary, approve.	1 year	Yes
Long-Acting Opioids (Oral)	Nucynta ER	tapentadol extended- release tablets	<ol> <li>Approve if the patient has tried three other oral long-acting opioid products.</li> <li>For example: morphine sulfate controlled-release/extended-release capsules or tablets [MS Contin, Kadian, generics], OxyContin, oxycodone ER tablets [generics], Xtampza ER, hydromorphone extended-release tablets [Exalgo, generics], oxymorphone extended-release tablets, or hydrocodone ER (Zohydro ER, Hysingla ER, generics).</li> <li>Patient is intolerant or allergic to morphine: approve if the patient has tried one product from the following list (if one is formulary): hydrocodone ER (Zohydro ER, Hysingla ER, generics), OxyContin, Xtampza ER, or oxycodone ER tablets (generics).</li> <li>Patient has renal insufficiency: approve if the patient has tried one product from the following list (if one is formulary): hydrocodone ER (Zohydro ER, Hysingla ER, generics), OxyContin, Xtampza ER, or oxycodone ER tablets (generics). If none are formulary, approve.</li> </ol>	1 year	Yes
Long-Acting Opioids		oxycodone extended-	1. Approve if the patient has tried three other oral long-acting opioid products.  For example: morphine sulfate controlled-release/extended-release capsules or tablets [MS Contin, Kadian, generics], hydromorphone extended-release tablets [Exalgo, generics], oxymorphone extended-release, Nucynta ER, hydrocodone ER (Zohydro ER, Hysingla ER, generics), OxyContin, or Xtampza ER.  2. Patient is intolerant or allergic to morphine: approve if the patient has tried one product from the following list (if one is formulary): Nucynta ER, hydrocodone ER (Zohydro ER, Hysingla ER, generics), Xtampza ER, or OxyContin. If none are formulary, approve.  3. Patient has renal insufficiency: approve if the patient has tried one product from the following list (if one is formulary): Nucynta ER, hydrocodone ER (Zohydro ER, Hysingla ER, generics), Xtampza ER, or OxyContin. If none are formulary, approve.		W.
Long-Acting Opioids	oxycodone ER  Xtampza ER	oxycodone extended- release capsules (with DETERX)	<ol> <li>4. Patients ≥ 11 years and &lt; 18 years of age: approve if the patient has tried OxyContin, if formulary. If Oxycontin is non-formulary, approve.</li> <li>1. Approve if the patient has tried three other oral long-acting opioid products.</li> <li>For example: morphine sulfate controlled-release/extended-release capsules or tablets [MS Contin, Kadian, generics], hydromorphone extended-release tablets [Exalgo, generics], oxymorphone extended-release, Nucynta ER, hydrocodone ER (Zohydro ER, Hysingla ER, generics), OxyContin, or oxycodone ER tablets [generics].</li> <li>2. Patient is intolerant or allergic to morphine: approve if the patient has tried one product from the following list (if one is formulary): Nucynta ER, hydrocodone ER (Zohydro ER, Hysingla ER, generics), oxycodone ER tablets (generics), or OxyContin. If none are formulary, approve.</li> <li>3. Patient has renal insufficiency: approve if the patient has tried one product from the following list (if one is formulary): Nucynta ER, hydrocodone ER (Zohydro ER, Hysingla ER, generics), oxycodone ER tablets (generics), or OxyContin. If none are formulary, approve.</li> </ol>	1 year	Yes
Long-Acting Opioids (Transdermal)	Butrans	buprenorphine transdermal system	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  Approve if the patient has tried three products from the following list, if formulary (or two if two are formulary or one if one is formulary): torsemide tablets, burnetanide (Bumex, generics), furosemide (Lasix, generics). If none are	1 year	MSB Exclusion *This criteria applies only to the NPF
Loop diuretics	Soaanz	furosemide subcutaneous injection by on-body	formulary, approve.  For the treatment of edema in a patient ≥ 18 years of age with chronic heart failure or chronic kidney disease, including the nephrotic syndrome.  Approve if the patient has tried at least one loop diuretic [documentation required] or the patient is currently taking a loop diuretic.	1 year	Yes
oop diuretics	Furoscix	infusor	Note: Examples of loop diuretics include furosemide, burnetanide, torsemide.	30 days	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Thorapy Glaco	Diana Hamo	Contrib Name	COMMODULY E CHOILE	Burution	MSB Exclusion
Low Molecular			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Weight Heparins and		enoxaparin sodium	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Related Agents	Lovenox	injection (syringe/vial)	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
		betaine	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		MSB Exclusion *This criteria
		trimethylglycine	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Metabolic Agents	Cystadane	powder for solution	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
			Patient meets the following: Metabolic Disorders - Phenylbutyrate Products Prior Authorization Policy criteria AND		
			Patient meets one of the following (1, 2, 3, or 4):		
			1. Approve if the patient has tried two of the following, if two are formulary (or one if only one is formulary): Olpruva and Pheburane [documentation required]. If neither are formulary, approve; OR		
			2. Patient has a feeding tube: Approve if the patient has tried sodium phenylbutyrate powder for oral administration (Buphenyl powder, generic) [documentation required], if formulary. If sodium phenylbutyrate powder for oral administration (Buphenyl powder, generic) is non-formulary, approve; OR		
			3. Patient is < 20 kg: approve if the patient meets one of the following (a or b):		
Metabolic Agents -		glycerol	a. Patient has tried Pheburane [documentation required], if formulary, if Pheburane is non-formulary, approve; OR		
Phenylbutyrate		phenylbutyrate oral	b. Patient is NOT eating solid food AND does NOT have a feeding tube (e.g., young infant): Approve; OR	See PA	
Agents	Ravicti	liquid	4. Patient is on a sodium-restricted diet OR, according to the prescriber, a high sodium diet is contraindicated [documentation required]: Approve.	duration	Yes
		nedosiran	Primary Hyperoxaluria Type 1 in a patient ≥ 2 years of age.		
Metabolic Disorder		subcutaneous	1. Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with Oxlumo, if formulary. If Oxlumo is non-formulary, approve.		
Agent	Rivfloza	injection	2. Approve if the patient has already been started on therapy with Rivfloza.	1 year	Yes
Metabolic Disorders – Cysteamine Ophthalmic Products	Cystadrops	cysteamine ophthalmic solution	Cystinosis with Corneal Cysteine Crystal Deposits: Approve, if the patient has tried Cystaran, if formulary. If Cystaran is non-formulary, approve.	1 year	Yes
			Approve if the patient has tried dihydroergotamine nasal spray (Migranal, generics), if formulary. If dihydroergotamine nasal spray (Migranal, generics) are non-formulary, approve if the patient meets one of the following (A or B):  A. Patient meets one of the following (i or ii):		
			i. Patient has tried one of sumatriptan nasal spray (Imitrex Nasal Spray, generics), Tosymra, or Onzetra Xsail, if formulary; OR ii. Patient has tried Zomig Nasal Spray or zolmitriptan nasal spray, if formulary: OR		
Migraine –		dihydroergotamine	ii. Fauterii rias ureu Zoring Nasai Spray of Zoriniurpian riasai spray, ii formulary. ON Note: If no products from i. or ii. are formulary, approve.		
Ergotamine Agents	Trudhesa	mesylate nasal spray	B. Patient has already experienced inadequate efficacy or a contraindication with a triptan product.	1 year	Yes
			Approve if the patient meets the following (A <u>and B</u> ):  A. Patient meets one of the following (i <u>or</u> ii):  i. Patient has tried both Nurtec ODT AND Ubrelvy, if both are formulary (or only one if one is formulary); OR  ii. If the patient is unable to swallow or has difficulty swallowing tablets, the patient has tried Nurtec ODT, if formulary. If Nurtec ODT is non-formulary, criteria A is met; AND  Note: The patient would still need to meet criteria B even if criteria A is met.  B. Patient meets one of the following (i or ii):		
Migraine Agent –			i. Patient meets one of the following (1 or 2):		
Treatment			1. Patient has tried two triptan products (for example, almotriptan [Axert, generics], eletriptan [Relpax, generics], frovatriptan [Frova, generics], naratriptan [Amerge, generics], rizatriptan [Maxalt, generics], sumatriptan [Imitrex,		
Medications -			generics], zolmitriptan [Zomig, generics]); OR		
Calcitonin gene-			2. Patient has tried one tripitan/non-steroidal anti-inflammatory drug (NSAID) combination product (e.g., Treximet or Symbravo) OR the patient has tried a triptan taken concomitantly with an NSAID; OR		
related peptide		zavogopont posol	ii. Patient meets one of the following (1 or 2):		
(CGRP) receptor antagonist	Zavzpret	zavegepant nasal spray	1. Per the prescriber, the patient has a contraindication to triptans; OR     2. Per the prescriber, the patient has had a significant intolerance to one or more triptans.	1 vear	Yes
Migraine Agents -	Zavzpict	оргау	2. For the processing, the person had a dignificant interestine to the or more aparts.	, your	1.03
Calcitonin Gene-		eptinezumab-jjmr			
Related Peptide		injection for			
(CGRP) Inhibitors	Vyepti	intravenous use	Approve if the patient has tried four of the following products, if formulary (or three if three are formulary or two if two are formulary or one if one is formulary): Aimovig, Emgality, Ajovy, and Qulipta. If none are formulary, approve.	1 year	Yes
Address in a America	Treximet and		Approve if the patient meets BOTH of the following (A and B):  A. Patient has used concurrently any oral non-steroidal anti-inflammatory drug (NSAID) with an oral triptan AND, according to the prescriber experienced inadequate efficacy or significant intolerance; AND		
Migraine Agents – Triptan Combination	generic sumatriptan- naproxen sodium	sumatriptan/ naproxen sodium	B. Patient has used concurrently any oral NSAID with another oral triptan (NOT the same triptan that was used in criterion A) AND, according to the prescriber experienced inadequate efficacy or significant intolerance.		
Products	tablets	tablets	Note: Examples of oral NSAIDs include etodolac, flurbiprofen, ibuprofen, ketoprofen, meloxicam, nabumetone, naproxen, oxaprozin.  Note: Examples of oral triptans include rizatriptan, almotriptan, eletriptan, frovatriptan, naratriptan, sumatriptan, zolmitriptan.	1 vear	Yes
			1	. , 500.	

Thomas Class	Daniel Name	Consulta Nama	Communical FE Oritoria	Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
Migraine Agents -		sumatriptan succinate solution for injection (injectable	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		MSB Exclusion *This criteria applies only to
Triptans	Imitrex injection	pen/cartridges)	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Migraine Agents - Triptans	Imitrex nasal spray		NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Migraine Agents - Triptans	Imitrex tablets	sumatriptan succinate tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Migraine Agents - Triptans	Maxalt	rizatriptan tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Migraine Agents - Triptans	Maxalt MLT	rizatriptan orally disintegrating tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Migraine Agents - Triptans	Relpax	eletriptan tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Migraine Agents - Triptans	Zomig tablets	zolmitriptan tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Migraine Agents -		sumatriptan nasal	Approve if the patient meets both of the following (a <u>and</u> b): a. Patient has tried one of sumatriptan nasal spray (Imitrex Nasal Spray, generics) or Tosymra, if formulary; AND b. Patient has tried Zomig Nasal Spray or zolmitriptan nasal spray, if formulary.		
Triptans  Miscellaneous	Onzetra Xsail	powder	Note: If no products from a. or b. are formulary, approve.  Hyperhidrosis, Primary Axillary in a patient ≥ 9 years of age.  Note: Softra is not intended for application to areas other than the axillae.  Approve if the patient meets BOTH of the following (1 and 2):  1. Approve if the patient meets ONE of the following (A or B):  A. Patient has tried, for at least 4 weeks, and experienced inadequate efficacy with one of Drysol, Xerac AC, or Bromi-lotion [documentation required]; OR  B. According to the prescriber, the patient has experienced a significant intolerance with one of these products [documentation required]; AND  2. Patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with Qbrexza, if formulary.	1 year	Yes
anticholinergic	Sofdra	gel, 12.45%  pyridostigmine tablet,	Note: If Qbrexa is non-formulary, criterion 2 is met.  NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.	1 year	Yes MSB Exclusion *This criteria
Miscellaneous anticholinergic	Mestinon	solution, exteneded- release tablet	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  Hyperhidrosis, Primary Axillary in a patient ≥ 9 years of age.	1 year	applies only to the NPF
			Note: Obrexza is not intended for application to areas other than the axillae.  Approve if the patient meets BOTH of the following (1 and 2):  1. Patient meets ONE of the following (A or B):  A. Patient has tried, for at least 4 weeks, and experienced inadequate efficacy with one of Drysol, Xerac AC, or Bromi-lotion [documentation required]; OR  B. According to the prescriber, the patient has experienced a significant intolerance with one of these products [documentation required].		
Miscellaneous	Ohrovza		2. Patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with Sofdra, if formulary.	1 1100	Vaa
anticholinergic	Qbrexza	2.4%, for topical use	Note: If Sofdra is non-formulary, criterion 2 is met.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Miscellaneous Urologicals	Urimar-T	methenamine 120 mg, sodium phosphate monobasic 40.8 mg, phenyl salicylate 36.2 mg, methylene blue 10.8 mg, hyoscyamine sulfate 0.12 mg capsule	Apporve if the patient has tried, and according to the prescriber, has experienced inadequate efficacy OR significant intolerance with BOTH of the following, if formulary: Uro-MP capsules AND Uro-SP capsules. If neither are formulary, approve.	1 year	Yes
Miscellaneous Urologicals	Urneva	methenamine 120 mg, sodium phosphate monobasic 40.8 mg, phenyl salicylate 36.2 mg, methylene blue 10.8 mg, hyoscyamine sulfate 0.12 mg capsule	Apporve if the patient has tried, and according to the prescriber, has experienced inadequate efficacy OR significant intolerance with BOTH of the following, if formulary: Uro-MP capsules AND Uro-SP capsules. If neither are formulary, approve.	1 year	Yes
Multiple Sclerosis Drugs -Injectable glatiramer	Copaxone	glatiramer acetate injection	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Multiple Sclerosis Drugs (Injectable) - CD20-directed cytolytic antibodies	Briumvi	ublituximab-xiiy	Relapsing forms of multiple sclerosis.  Note: Examples of relapsing forms of multiple sclerosis (MS) include clinically isolated syndrome, relapsing remitting disease, and active secondary progressive disease.  1. Approve if the patient has tried and, according to the prescriber, has had inadequate efficacy or significant intolerance with ONE of 1) Ocrevus intravenous or Ocrevus Zunovo or 2) Kesimpta, if formulary. If none are formulary, approve.  2. Approve if the patient has already been started on Briumvi therapy.	1 year	Yes
Multiple Sclerosis Drugs (Oral)	Ampyra	dalfampridine extended-release tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Multiple Sclerosis Drugs (Oral)	Gilenya 0.5 mg	fingolimod capsule	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  Patient meets all of the following (A, B, C and D):  A. Patient with relapsing form of multiple sclerosis; AND	1 year	MSB Exclusion *This criteria applies only to the NPF
Multiple Sclerosis Drugs (Oral)	Gilenya 0.25 mg	fingolimod capsule	Note: Examples of relapsing forms of multiple sclerosis (MS) include clinically isolated syndrome, relapsing remitting disease, and active secondary progressive disease.  B. Patients ≥ 10 years of age; AND  C. Patient weighs less than or equal to 40 kg [documentation required]; AND  D. Patient has tried Tascenso 0.25 mg orally disintegrating tablets (ODT), if formulary. If Tascenso 0.25 ODT are non-formulary, approve.	1 year	Yes

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
			Relapsing forms of multiple sclerosis.  Note: Examples of relapsing forms of multiple sclerosis (MS) include clinically isolated syndrome, relapsing remitting disease, and active secondary progressive disease.  Approve if the patient has tried teriflunomide tablets, if formulary.		
			If teriflunomide tablets are non-formulary or generic teriflunomide is being requested, approve if the patient meets one of the following (1, 2, or 3):		
			1. Patient meets the following (A and B): A. Patient has tried and, according to the prescriber, has had inadequate efficacy OR significant intolerance with one fumarate-based product, if formulary: Bafiertam, dimethyl fumarate (Tecfidera, generics), or Vumerity. If none are		
			formulary, approve; AND  B. Patient has tried and, according to the prescriber, has had inadequate efficacy OR significant intolerance with one of the following: fingolimod (Gilenya, generics), Zeposia, Mayzent, or Ponvory, if formulary. If none are formulary, would still need to try a fumarate-based product, if one is formulary.		
			2. For patients with an underlying cardiovascular condition (e.g., heart failure, myocardial infarction, stroke, transient ischemic attack, unstable angina, atrioventricular [AV] block, cardiac arrhythmias, bradyarrhythmias), patient has tried and, according to the prescriber, has had inadequate efficacy OR significant intolerance with one other oral disease-modifying therapy (e.g., dimethyl fumarate, Vumerity).		
Multiple Sclerosis Drugs (Oral)	Aubagio	teriflunomide tablets		1 year	Yes- brand only
			Patient with relapsing form of multiple sclerosis.  Note: Examples of relapsing forms of multiple sclerosis (MS) include clinically isolated syndrome, relapsing remitting disease, and active secondary progressive disease.		
Multiple Sclerosis Drugs (Oral)	Tascenso ODT 0.5 mg	fingolimod orally disintegrating tablets	<ol> <li>Approve if the patient is unable to swallow or has difficulty swallowing fingolimod 0.5 mg capsules or Gilenya 0.5 mg capsules [documentation required].</li> <li>Approve if neither fingolimod 0.5 mg capsule nor Gilenya 0.5 mg capsules are formulary.</li> </ol>	1 year	Yes
			Relapsing forms of multiple sclerosis.  Note: Examples of relapsing forms of multiple sclerosis include clinically isolated syndrome, relapsing remitting disease, and active secondary progressive disease.		
			Approve if the patient meets one of the following (1 or 2):  1. Patient meets the following (A and B):		
			A. Patient has tried and, according to the prescriber, has had inadequate efficacy OR significant intolerance with one fumarate-based product, if formulary: Bafiertam, dimethyl fumarate (Tecfidera, generics), or Vumerity. If none are formulary, approve; AND		
Multiple Sclerosis			B. Patient has tried and, according to the prescriber, has had inadequate efficacy ORor significant intolerance with one of the following: fingolimod (Gilenya, generics), Mayzent, or Zeposia, if formulary. If none are formulary, would still need to try a fumarate-based product, if one is formulary.		
Drugs (Oral)	Ponvory	ponesimod capsules		1 year	Yes
			Approve if the patient meets the following 1 AND 2:  1. Patient meets all of the following (A, B, and C):		
			A. Patient with relapsing form of multiple sclerosis; AND  Note: Examples of relapsing forms of multiple sclerosis (MS) include clinically isolated syndrome, relapsing remitting disease, and active secondary progressive disease.		
			B. Patients ≥ 10 years of age; AND C. Patient weighs less than or equal to 40 kg [documentation required]; AND		
Multiple Colonsois	T ODT 0.05	fin maline and a malle.	2. Patients meets one of the following (A, B, <u>OR</u> C):  A. Patient is unable to swallow or has difficulty swallowing Gilenya [documentation required]; OR  D. D		
Multiple Sclerosis Drugs (Oral)	Tascenso ODT 0.25 mg	disintegrating tablets	B. Patient is unable to obtain Gilenya 0.25 mg capsules from the manufacturer; OR C. Gilenya 0.25 mg is non-formulary.	1 year	Yes
Multiple Sclerosis Drugs (Oral) –		dimethyl fumarate			
Fumarate-based		delayed-release			
Agents	Tecfidera Methocarbamol	capsules	See standard Multiple Sclerosis (Tecfidera) Preferred Specialty Management Policy criteria.	1 year	Yes
	1,000 mg tablets		1. Direct the patient to methocarbamol 500 mg tablets.		
Muscle Relaxants	(brand)	mg tablets cyclobenzaprine	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use the methocarbamol 500 mg tablets.	1 year	Yes
		extended-release 15			
Muscle Relaxants	Amrix and generic	mg and 30 mg capsule	Approve if the patient has tried and cannot take cyclobenzaprine 5 mg or 10 mg tablets (generics), if formulary. If cyclobenzaprine 5 mg or 10 mg tablets (generics) are non-formulary, approve.	1 year	Yes
Muscle Relaxants	metaxalone 640 mg tablet	metaxalone 640 mg tablet	<ol> <li>Approve if the patient has tried and cannot take one of metaxolone 400 mg or 800 mg tablets.</li> <li>According to the prescriber, there is a significant clinical concern such that the patient is unable to use metaxolone 400 mg or 800 mg tablets.</li> </ol>	1 year	Yes
		baclofen oral			
Muscle Relaxants –		suspension, concentrated	1. Direct to oral baclofen tablets. 2. Patient is unable to or has difficulty swallowing oral tablets, approve if the patient has tried baclofen 25 mg/5ml oral suspension (generic of Fleqsuvy), if formulary. If baclofen 25 mg/5ml oral suspension (generic of Fleqsuvy) is non-		
Baclofen Agents	Fleqsuvy	formulation	formulary, approve if the patient has tried one of 1) Ozobax solution or 2) Lyvispah oral gransules, if formulary. If neither are formulary, approve.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
morupy oraco	Brana Hamo	Concrio Humo	1. Direct the patient to oral baclofen tablets.	Burution	incurcuitori
Muscle Relaxants – Baclofen Agents	Lyvispah	baclofen oral granules	2. If Lvyispah will be administered via a feeding tube, approve. 3. Patient is unable to or has difficulty swallowing oral tablets, approve if the patient has tried one of 1) Ozobax solution or 2) baclofen 25mg/5ml oral suspension (Fleqsuvy suspension, generics), if formulary. If neither are formulary, approve.	1 year	Yes
	- <b>y</b> p	g		. ,	
	Ozobax, Ozobax DS, and authorized generics	baclofen oral solution	1. Direct to oral baclofen tablets. 2. Patient is unable to or has difficulty swallowing oral tablets, approve if the patient has tried one of 1) baclofen 25 mg/5ml oral suspension (Fleqsuvy suspension, generics) or 2) Lyvispah oral granules, if formulary. If neither are formulary, approve.	1 year	Yes
			Chronic Myelomonocytic Leukemia; Myelodysplastic Syndrome with Myeloproliferative Neoplasm Overlap Syndrome; Myelodysplastic Syndromes (Note: Examples of myelodysplastic syndromes include: refractory anemia, refractory		
Myelodysplastic		decitabine and	<ul> <li>anemia with ringed sideroblasts, and refractory anemia with excess blasts.).</li> <li>Approve if the patient has tried decitabine injection (Dacogen, generics), if formulary. If decitabine injection (Dacogen, generics) is non-formulary, approve.</li> <li>Approve if the patient is unable to obtain and/or maintain intravenous access.</li> </ul>		
syndrome Agent	Inqovi	cedazuridine tablets	3. Approve if the patient has already started therapy with Inqovi.	1 year	Yes
			Myelofibrosis. Note: Myelofibrosis includes primary myelofibrosis, post-polycythemia vera myelofibrosis, and post-essential thrombocythemia myelofibrosis.  1. Approve if the patient has tried Jakafi. If Jakafi is non-formulary, approve.  2. Approve if the patient has already been started on Inrebic.		
Myelofibrosis Agents	Inrebic	febratinib capsules	Accelerated or Blast Phase Myeloproliferative Neoplasm: approve.  Myeloid/Lymphoid Neoplasms: approve.	1 year	Yes
Myelulibiosis Agents	Illiebic		Myelofibrosis.  Myelofibrosis, Dote: This includes primary myelofibrosis, post-polycythemia vera myelofibrosis, and post-essential thrombocythemia myelofibrosis.  1. Approve if the patient has tried Jakafi. If Jakafi is non-formulary, approve.  Note: If the patient has tried Vonjo, this would satisfy requirement for approval.  2. If the patient has myelofibrosis anemia, approve.  3. Approve if the patient has already started on therapy with Ojjaara.	i yeai	165
Myelofibrosis Agents  – JAK Inhibitors	Oijaara	momelotinib tablets	Accelerated or Blast Phase Myeloproliferative Neoplasm: approve.	1 year	Yes
Naloxone Products for	Zimhi	naloxone hydrochloride intramuscular or subcutaneous injection 5 mg/0.5 ml	1. Approve if the patient has tried naloxone syringes, if formulary. If naloxone syringes are non-formulary, approve.  2. Approve, if according to the prescriber, a higher-strength naloxone product is needed.	1 year	Yes
органа в напада		,gg	т. фр. т.	. ,	MSB Exclusion
Nasal Antihistamines and Combination Products	Duminto		NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the	4	*This criteria applies only to the NPF
Products	Dymista	nasal spray beclomethasone	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  Approve if the patient has tried five nasal steroids from the following list, if formulary (or four if four are formulary or three if three are formulary or two if two are formulary or one if one is formulary): fluticasone nasal spray, mometasone nasal spray (Nasonex, generics), flunisolide nasal spray (generics), Omnaris, Qnasl, or Zetonna.	1 year	THE NPP
Nasal Steroids	Beconase AQ	nasal spray	Note: Over-the-counter (OTC) nasal steroids would count as a trial of an alternative.	1 year	Yes
Nasal Steroids	Omnaris	ciclesonide nasal spary	Approve if the patient has tried five nasal steroids from the following list, if formulary (or four if four are formulary or three if three are formulary or two if two are formulary or one if one is formulary): fluticasone propionate spray, mometasone nasal spray (Nasonex, generics), flunisolide nasal spray (generics), Qnasl, or Zetonna.  Note: Over-the-counter (OTC) nasal steroids would count as a trial of an alternative.	1 year	Yes
		beclomethasone	Approve if the patient has tried five nasal steroids from the following list, if formulary (or four if four are formulary or three if three are formulary or two if two are formulary or one if one is formulary): fluticasone propionate spray,		
Nasal Steroids	Qnasl	dipropionate nasal aersol	mometasone nasal spray (Nasonex, generics), flunisolide nasal spray (generics), Omnaris, or Zetonna.  Note: Over-the-counter (OTC) nasal steroids would count as a trial of an alternative.	1 year	Yes
		ciclesonide nasal	Approve if the patient has tried five nasal steroids from the following list, if formulary (or four if four are formulary or three if three are formulary or two if two are formulary or one if one is formulary): fluticasone propionate spray, mometasone nasal spray (Nasonex, generics), flunisolide nasal spray (generics), Omnaris, or Qnasl.	j	
Nasal Steroids	Zetonna	aerosol	Note: Over-the-counter (OTC) nasal steroids would count as a trial of an alternative.	1 year	Yes
Nephropathic		cysteamine bitartrate dealyed-release			
Cystinosis			Nephropathic cystinosis.		
Medications	Procysbi	packets	Approve if the patient has tried Cystagon [documentation required], if formulary. If Cystagon is non-formulary, approve.	1 year	Yes

				A	2025 NPF
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	Excluded Medicaiton
Neurology -					
Amyotrophic Lateral					
Sclerosis (ALS) Agents	Qalsody	tofersen intrathecal injection	See standard Neurology – Qalsody Prior Authorization Policy criteria.  Note: No conditions of approval are recommended in the prior authorization policy.	N/A	Yes
Agents	Qaisody	injection	Anti-aquaporin (AQ4P) antibody-positive Neuromyelitis Optica Spectrum Disorder in a patient ≥ 18 years of age.	IN/A	res
			1. Approve if the patient has tried and, according to the prescriber, has inadequate efficacy or significant intolerance to Enspryng, if formulary. If Enspryng is non-formulary, approve.		
Neuromyelitis optica		inebilizumab-cdon	2. Approve if the patient has already started on therapy with Uplinza.		
spectrum disorder		injection for			.,
(NMOSD) Agents	Uplizna	intravenous infusion	Immunoglobulin G4-Related Disease in a patient ≥ 18 years of age: Approve.	1 year	Yes
			Patient meets the following Niemann-Pick Disease Type C – Miplyffa Prior Authorization Policy AND		
			Patient meets ONE of the following (1, 2, or 3):		
Niemann-Pick			1. Patient has tried Agneursa, if formulary. If Agneursa is non-formulary, criterion A is met; OR		
disease type C			2. Patient is < 4 years of age.		
Agents	Miplyffa	arimoclomol capsules	3. Patient has already been started on therapy with Miplyffa.	1 year	Yes
			1. For the diagnosis of Treatment-Resistant Depression: approve if the patient meets the following criteria (A, B, C, D, and E):		
			A. Patient is ≥ 18 years of age; AND		
			B. Patient meets both of the following (i and ii):  i. Patient has demonstrated nonresponse (≤ 25% improvement in depression symptoms or scores) to at least two different antidepressants, each from a different pharmacologic class, according to the prescriber; AND		
			Note: Different pharmacologic classes of antidepressants include selective serotonin reuptake inhibitors, serotonin-norepinephrine reuptake inhibitors, tricyclic antidepressants, bupropion, mirtazapine, etc.		
			ii. Each antidepressant was used at therapeutic dosages for at least 6 weeks in the current episode of depression, according to the prescriber; AND		
			C. Patient has one of the following (i or ii):		
			i. No history of psychosis; OR		
			ii. History of psychosis and the prescriber believes that the benefits of Spravato outweigh the risks; AND  D. The patient's history of controlled substance prescriptions has been checked using the state prescription drug monitoring program, according to the prescriber; AND		
			E. The medication is prescribed by a psychiatrist.		
			2. Major Depressive Disorder with Acute Suicidal Ideation or Behavior: approve if the patient meets the following criteria (A, B, C, D, and E):		
			A. Patient is ≥ 18 years of age; AND		
			B. Patient has major depressive disorder that is considered to be severe, according to the prescriber; AND C. Patient is concomitantly receiving at least one oral antidepressant; AND		
			Note: Antidepressants may include, but are not limited to, selective serotonin reuptake inhibitors, serotonin-norepinephrine reuptake inhibitors, tricyclic antidepressants, mirtazapine, and bupropion.		
			D. Patient has one of the following (i or ii):		
			i. No history of psychosis; OR		
N-methyl D-aspartate (NMDA) receptor		esketamine nasal	ii. History of psychosis and the prescriber believes that the benefits of Spravato outweigh the risks; AND  E. The medication is prescribed by a psychiatrist.		
antagonists	Spravato	spray	3. For the diagnosis of Treatment-Resistant Depression OR Major Depressive Disorder with acute suicidal ideation or behavior: approve if the patient has already started therapy with Sprayato.	1 year	Yes
<u> </u>			יייייייייייייייייייייייייייייייייייייי		
NSAID and Acid	Duexis and generic				
Reducing Agent	ibuprofen-famotidine		Annual if the matient has brind and assert that if the second of the sec	4	Yes- generic
Combination Products	ableis	famotidine tablets	Approve if the patient has tried and cannot take famotidine AND has concurrently tried ibuprofen (brand or generic).	1 year	only
	Vimovo and generic				
	naproxen-	naproxen and			
NSAID and Acid	esomeprazole	esomeprazole			
Reducing Agent	magnesium delayed-	magnesium delayed-	A STATE OF THE STA	4	V
Combination Products	release tablets	release tablets	Approve if the patient has tried and cannot take esomeprazole AND has concurrently tried naproxen (e.g., Naprosyn, Naprelan, generics).	1 year	Yes MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
NSAIDS (Cox2)	Celebrex	celecoxib capsules	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
			Acute treatment of migraine.		
NSAIDS (Cox2)	Elvado	celecoxib oral solution	1. Direct the patient to celecoxib capsules. If celecoxib capsules (Celebrex, generics) are non-formulary, approve.	1 voor	Vos
NOAIDO (COXZ)	Elyxyb	SUIULIUII	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use celecoxib capsules.	1 year	Yes

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
			NOTE: A multipayers Provider requested. The national about the preferred big squivelent generic product		MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		*This criteria applies only to
NSAIDs (Oral)	Nalfon	fenoprofen capsules	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
NOAID3 (Olai)	INGIIOII	ichoprotett capsules	prosince, would result in a significant energy or scripts adverse reaction required.	i you	MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
		diclofenac potassium	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
NSAIDs (Oral)	Zipsor	capsule	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
					MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
NCAID- (OI)	Indesia Communica	indomethacin oral	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the	4	applies only to
NSAIDs (Oral)	Indocin Suspension	suspension	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  Approve if the patient has tried five prescription-strength, oral NSAIDs.	1 year	the NPF
			Note: For example: fenoprofen (tablets/generics), etodolac (generics), flurbiprofen (Ansaid, generics), ibuprofen (e.g., Motrin, generics), ketoprofen (generics), meloxicam (Mobic, generics), nabumetone (generics), naproxen (e.g.,		
	Fenoprofen		Naprosyn, Naprelan, generics), oxaprozin (Daypro, generics), idiclofenac (Voltaren XR, generics), inconcern (generics), inconcern (g		
	capsules [brand] and		Note: Over-the-counter (OTC) NSAIDs would count if prescription-strength doses were used.		
NSAIDs (Oral)	Fenopron	fenoprofen capsules	Note: Five unique NSAIDs should be tried.	1 year	Yes
			Approve if the patient has tried five prescription-strength oral NSAIDs.		
			Note: For example: nabumetone (generics), etodolac (generics), flurbiprofen (Ansaid, generics), ibuprofen (e.g., Motrin, generics), ketoprofen (generics), meloxicam (Mobic, generics), naproxen (e.g., Naprosyn, Naprelan, generics),		
			oxaprozin (Daypro, generics), diclofenac (Voltaren XR, generics), piroxicam (Feldene, generics), indomethacin (generics).		
NOAID (0 I)	D 1 ( DO	nabumetone 1,000	Note: Over-the-counter (OTC) NSAIDs would count if prescription-strength doses were used.		
NSAIDs (Oral)	Relafen DS	mg tablets	Note: Five unique NSAIDs should be tried.	1 year	Yes
			1. Approve if the patient has tried five prescription-strength, oral NSAIDs.  Note: Examples include: etodolac (generics), flurbiprofen (Ansaid, generics), ibuprofen (e.g., Motrin, generics), ketoprofen (generics), meloxicam (Mobic, generics), nabumetone (generics), naproxen (e.g., Naprosyn, Naprelan,		
			generics), oxaprozin (Daypro, generics), idiolofenac (Voltaren XR, generics), piroxicam (Feldene, generics), indomethacin (generics).		
		ketorolac	Note: Over-the-counter (OTC) NSAIDs would count if prescription-strength doses were used.		
	Sprix and authorized	tromethamine nasal	Note: Five unique NSAIDs should be tried.		Yes - Authorized
NSAIDs (Oral)	generic	spray	2. Approve for patients with difficulty swallowing or for patients who cannot swallow.	1 year	generic only
			Approve if the patient has tried five prescription-strength, oral NSAIDS.		
			Note: Examples include: diclofenac (Voltaren XR, generics), etodolac (generics), flurbiprofen (Ansaid, generics), ibuprofen (e.g., Motrin, generics), ketoprofen (generics), meloxicam (Mobic, generics), nabumetone (generics), naproxen		
			(e.g., Naprosyn, Naprelan, generics), oxaprozin (Daypro, generics), piroxicam (Feldene, generics), indomethacin (generics).		
NICAID= (O==1)	Zorvolex and	diala6	Note: Over-the-counter (OTC) NSAIDs would count if prescription-strength doses were used.	4	V
NSAIDs (Oral)	authorized generic Meloxicam	diclofenac capsules meloxicam	Note: Five unique NSAIDs should be tried.  Approve if the patient has tried one of ibuprofen suspension (e.g., Motrin, generics) or naproxen suspension (e.g., Naprosyn, generics), if formulary. If neither are formulary, approve.	1 year	Yes
NSAIDs (Oral)	suspension	suspension	Note: Over-the-counter ibuprofen suspension would count as an alternative, regardless of formulary status.	1 year	Yes
NOAIDS (Olai)	заэрспэюн	зазрензіон	Approve if the patient has tried five prescription-strength, oral NSAIDs.	i you	103
			Note: Examples include: meloxicam (Mobic, generics), etodolac (generics), flurbiprofen (Ansaid, generics), ibuprofen (e.g., Motrin, generics), ketoprofen (generics), nabumetone (generics), naproxen (e.g., Naproxyn, Naprelan,		
			generics), oxaprozin (Daypro, generics), diclofenac (Voltaren XR, generics), piroxicam (Feldene, generics), indomethacin (generics).		
			Note: Over-the-counter (OTC) NSAIDs would count if prescription-strength doses were used.		
NSAIDs (Oral)	Vivlodex	meloxicam capsules	Note: Five unique NSAIDs should be tried.	1 year	Yes
			Approve if the patient has tried five prescription-strength oral NSAIDs.		
			Note: For example: nabumetone (generics), etodolac (generics), flurbiprofen (Ansaid, generics), ibuprofen (e.g., Motrin, generics), ketoprofen (generics), meloxicam (Mobic, generics), naproxen (e.g., Naprosyn, Naprelan, generics),		
			oxaprozin (Daypro, generics), diclofenac (Voltaren XR, generics), piroxicam (Feldene, generics), indomethacin (generics).  Note: Over-the-counter (OTC) NSAIDs would count if prescription-strength doses were used.		
NSAIDs (Oral)	Dolobid	diflunisal tablets	Note: Five unique NSAIDs should be tried.	1 year	Yes
NOMIDS (Olal)	Dolobia	amanisar labicis	Approve if the patient has tried five prescription-strength, oral NSAIDs.	i you	1.03
			Note: Examples include: oxaprozin (Daypro, generics), etodolac (generics), flurbiprofen (Ansaid, generics), ibuprofen (e.g., Motrin, generics), ketoprofen (generics), meloxicam (Mobic, generics), nabumetone (generics), naproxen (e.g.,		
	Coxanto and		Naprosyn, Naprelan, generics), diclofenac (Voltaren XR, generics), indomethacin (generics).		
	oxaprozin 300 mg		Note: Over-the-counter (OTC) NSAIDs would count if prescription-strength doses were used.		
NSAIDs (Oral)	(brand)	oxaprozin capsule	Note: Five unique NSAIDs should be tried.	1 year	Yes
	Indocin	indomethacin	No exceptions are recommended. There are multiple therapeutic alternatives available. (NOTE: It is not appropriate to use standard global criteria for this medication; Denial reason is: No exceptions are recommended. There are		
NSAIDs (Suppository)	Suppositories	suppositories	multiple therapeutic alternatives available.)	N/A	Yes- brand only
		P. 1. 6	NOTE: A multipayers Provider to being requested. The national should use the professed big equivalent generic ac-direct		MSB Exclusion
		diclofenac sodium topical solution 2.0%	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		*This criteria
NSAIDs (Topical)	Pennsaid	pump	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	applies only to the NPF
(Topical)	. Jilloulu	L 2.11b	processor, was read to the significant allergy or solices acressed to desired to quite allerge.	. 3001	

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
NSAIDs (Topical)	diclofenac epolamine 1.3% topical patch (authorized generic of Flector Patch)	diclofenac epolamine 1.3% topical patch	Direct the patient to use Flector patch (brand), if formulary. If Flector patch (brand) is non-formulary, approve if the patient has tried, and according to the prescriber, has experienced inadequate efficacy OR significant intolerance with four products from the following list (if four are formulary or three if three are formulary or two if two are formulary or one if one is formulary): Licart 1.3% topical system, diclofenac 2% solution pump (Pennsaid 2.0%, generics), diclofenac sodium 1.5% topical solution (generics), or prescription diclofenac sodium topical 1% gel (Voltaren 1% gel, generics), if one is formulary. If none are formulary, approve if the patient has tried over-the-counter Voltaren 1% gel.	; 1 year	Yes
	,				
Nuclear Factor (erythroid-derived 2)- like 2 (Nrf2) Activator	Skyclarys	omaveloxolone capsules	See standard Neurology – Skyclarys Prior Authorization Policy criteria.	See PA duration	Yes
Omega-3 Fatty Acid Products	Lovaza	omega-3 acid ethyl esters capsules	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Ophthalmic – Antibiotic/Corticostero id Combination	,	tobramycin 0.3%/dexamethasone 0.05% ophthalmic	1. Approve if the patient has tried tobramycin-dexamethasone ophthalmic suspension (Tobradex, generics), if formulary. If tobramycin-dexamthasone ophthalmic suspension (Tobradex, generics) are non-formulary, approve.		
Products	TobraDex ST	suspension	2. For the treatment of currently active eye infections: approve in patients already receiving TobraDex ST to complete the course of therapy.	1 year	Yes
Ophthalmic – Antibiotic/Corticostero id Combination Products	Zylet	tobramycin 0.3%/loteprednol etabonate 0.5% ophthalmic suspension	<ol> <li>Approve if the patient has tried one of tobramycin-dexamethasone ophthalmic suspension (Tobradex, generics) or TobraDex ST, if formulary. If neither are non-formulary, approve.</li> <li>Patients &lt; 2 years of age, approve.</li> <li>For the treatment of currently active eye infections: approve in patients already receiving Zylet to complete the course of therapy.</li> </ol>	1 year	Yes
Ophthalmic - Calcineurin Inhibitor Immunosuppressant	Verkazia	cyclosporine 0.1% ophthalmic emulsion	Moderate to Severe Vernal Keratoconjunctivitis.  1. Approve if the patient meets one of the following (A or B):  A. Patient has tried two single-action ophthalmic medications (i.e., ophthalmic mast-cell stabilizers or ophthalmic antihistamines) for the maintenance treatment of vernal keratoconjunctivitis; OR  Note: Examples of single-action ophthalmic medications for the maintenance treatment of vernal keratoconjunctivitis include ophthalmic mast-cell stabilizers (e.g., cromolyn ophthalmic solution, Alomide ophthalmic solution) and ophthalmic antihistamines (e.g., Zerviate [cetirizine solution]).  B. Patient has tried one dual-action ophthalmic mast-cell stabilizer/antihistamine product for the maintenance treatment of vernal keratoconjunctivitis.  Note: Examples of dual-action ophthalmic mast-cell stabilizer/antihistamine products include azelastine ophthalmic solution, beoptastine ophthalmic solution, epinastine ophthalmic solution, ketotifen ophthalmic solution, lopatadine ophthalmic solution.  Note: An exception to the requirement for a trial of two single-action ophthalmic medications (i.e., ophthalmic mast-cell stabilizers or ophthalmic antihistamines) or one dual-action ophthalmic mast-cell stabilizer/antihistamine product for the maintenance treatment of vernal keratoconjunctivitis can be made if the patient has already tried at least one ophthalmic cyclosporine product.	1 year	Yes
Ophthalmic Agent – Mydriatics/ Cycloplegics	Atropine sulfate 1% ophthalmic solution (preservative free) [brand]	atropine sulfate 1%	1. Direct the patient to generic atropine sulfate 1% ophthalmic solution. 2. Patients with a known sensitivity to a preservative (e.g., benzalkonium chloride [BAK]), approve.	1 year	Yes
Ophthalmic Agents -					
Complement Protein C5 Inhibitor	Izervay	avacincaptad pegol intravitreal injection	See standard Ophthalmology – Izervay Prior Authorization Policy criteria.	See PA duration	Yes
Ophthalmic Agents - VEGF Inhibitors	Susvimo	ranibizumab intravitreal injection via ocular implant and implant/insert tool		N/A	Yes
Ophthalmic Agents -		aflibercept intravitreal	Approve if the patient meets BOTH of the following (1 and 2):  1. Patient has tried ONE of Eylea (not HD) or Pavblu [documentation required], if one is formulary; AND  2. Patient has experienced a significant intolerance with ONE of Eylea (not HD) or Pavblu [documentation required].		
VEGF Inhibitors	Eylea HD	injection	Note: If BOTH Eylea (not HD) and Pavblu are non-formulary, approve.	1 year	Yes

					2025 NPF
				Approval	Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
			Pavblu is formulary:		
			Approve if the patient meets BOTH of the following (A <u>and</u> B):		
			A. Patient has tried Pavblu; AND		
			B. Patient cannot continue to use Pavblu due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a		
			significant allergy or serious adverse reaction.		
			Payblu is non-formulary:		
			Approve if the patient meets ONE of the following (A, B, C, or D):		
			A. Patient has tried Eylea HD, if formulary. If Eylea HD is non-formulary, approve; OR		
			B. For the diagnosis macular edema following retinal vein occlusion; OR		
			C. For the diagnosis retinopathy of prematurity; OR		
Ophthalmic Agents -		aflibercept intravitreal	D. For the diagnosis of other neovascular diseases of the eye.		
VEGF Inhibitors	Eylea	injection	Note: Examples of other neovascular diseases of the eye include neovascular glaucoma, sickle cell neovascularization, and choroidal neovascular conditions.	1 year	Yes
			1. If Byooviz and Cimerli are both formulary or only one is formulary: Approve if the patient meets both of the following (A and B):		
			A. Patient has tried both Byooviz and Cimerli (or one if one is formulary); AND		
			B. Patient cannot continue to use the alternative(s) due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a		
			significant allergy or serious adverse reaction.		
			2. If both Byooviz and Cimerli are non-formulary, approve if the patient meets one of the following (A, B, <u>or</u> C):		
			A. Patient has tried one of Eylea or Pavblu, if formulary. If neither are formulary, approve; OR		
Ophthalmic Agents -		ranibizumab	B. Patient with myopic choroidal neovascularization (mCNV); OR		
VEGF Inhibitors	Lucentis	intravitreal injection	C. Patient is currently receiving therapy with Lucentis.	1 year	Yes
			Neovascular (Wet) Age-Related Macular Degeneration; Diabetic Macular Edema.		
			1. Approve if the patient has tried one of 1) Eylea (not HD) or Pavblu OR Eylea HD, if formulary. If none are formulary, approve.		
			2. Patient is currently receiving therapy with Vabysmo: approve.		
			Macular Edema following Retinal Vein Occlusion.		
Ophthalmic Agents -		faricimab-svoa	1. Approve if the patient has tried one of Eylea (not HD) or Pavblu, if formulary. If neither are formulary, approve.		
VEGF Inhibitors	Vabysmo	intravitreal injection	2. Patient is currently receiving therapy with Vabysmo: approve.	1 year	Yes
	,	oxymetazoline	and the second s	,	
Ophthalmic alpha		hydrochloride 0.1%	No exceptions are recommended. Due to insufficient clinical efficacy data, approval is not recommended for Upneeq. (NOTE: It is not appropriate to use standard global criteria for this medication; Denial reason is: No exceptions are		
adrenoceptor agonist	Upneeq	ophthalmic solution	recommended. Due to insufficient clinical efficacy data, approval is not recommended.)	N/A	Yes
					MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Ophthalmic Anti-		bepotastine besilate	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Allergics	Bepreve	ophthalmic solution	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
		lodoxamide	1. Approve if the patient has tried three products from the following list (if three are formulary; or two if two are formulary; or one if one is formulary): Alocril, cromolyn sodium 4% solution (generics), azelastine 0.05% solution (generics),		
Ophthalmic Anti-		tromethamine 0.1%	bepotastine solution (Bepreve, generics), epinastine 0.05% solution (generics), Lastacaft, olopatadine solution (generics), or Zerviate. If none are formulary, approve.		
Allergics	Alomide	ophthalmic solution	2. For a diagnosis of vernal keratoconjunctivitis, vernal conjunctivitis, and vernal keratitis, approve if the patient has tried cromolyn sodium 4% solution (generics). If cromolyn sodium 4% solution (generics) is non-formulary, approve.	1 year	Yes
			1. Approve if the patient has riged three products from the following list (if three are formulary, or two if only two are formulary, or one if only one is formulary): bepotastine ophthalmic drops (Bepreve, generics), cromolyn ophthalmic		
			drops (generics), epinastine 0.05% solution (generics), Lastacaft, azelastine 0.05% solution (generics), olopatadine ophthalmic solution (generics), Zerviate. If none are formulary, approve.		
Ophthalmic Anti-	A1==	0.2% ophthalmic	2. Patients who require concurrent use of Alrex with an H1 antagonist or an H1 antagonist/mast cell stabilizer (e.g. azelastine [generics], bepotastine, epinastine solution [generics], Lastacaft, olopatadine ophthalmic solution [generics],	4	V
Allergics	Alrex	suspension	Zerviate): approve.	1 year	Yes
Ophthalmic Anti- Allergics	Zerviate	cetirizine 0.24% ophthalmic solution	Approve if the patient has tried three products from the following list (if three are formulary; or two if two are formulary; or one if one is formulary): Alocril, Alomide, cromolyn sodium 4% solution (generics), azelastine 0.05% solution (generics), bepotastine solution (generics), epinastine 0.05% solution (generics), Lastacaft, or olopatadine solution (generics).	1 year	Yes
Allergics	Zerviale	ophinalinic solution		ı year	165
Onbthalmic		hosiflovacia	1. Approve if the patient has tried two products from the following list, (if two are formulary, or one if one is formulary): 1) gatifloxacin ophthalmic solution (generics), 2) moxifloxacin ophthalmic solution (Vigamox, Moxeza, generics), or 3) levofloxacin ophthalmic solution (generics). If none are formulary, approve.		
Ophthalmic Antibiotics -		besifloxacin ophthalmic	2. Approve if there is laboratory data that the patient has an eye infection due to pathogens resistant to ciprofloxacin and one other ophthalmic quinolone.		
Quinolones	Besivance	suspension 0.6%	3. For the treatment of currently active eye infections: approve in patients already receiving Besivance therapy to complete the course of therapy.	1 year	Yes
		2.2.00.0.0.0.00		. , ,	
			1. Approve if the patient has tried four products from the following list, if formulary (or three if three are formulary, or two if two are formulary, or one if one is formulary): ciprofloxacin ophthalmic solution (Ciloxan, generics), gatifloxacin		
		ciprofloxacin	ophthalmic solution (generics), moxifloxacin ophthalmic solution (Vigamox, Moxeza, generics), levofloxacin ophthalmic solution (generics), or ofloxacin 0.3% ophthalmic solution (Ocuffox, generics). If none are formulary, approve.		
Ophthalmic		CIPIUIUXACIII			
Ophthalmic Antibiotics -			2. If the patient is allergic to benzalkonium chloride, approve if the patient has tried moxifloxacin (Vigamox, Moxeza, generics), if formulary. If moxifloxacin (Vigamox, Moxeza, generics) are non-formulary, approve.		

					2025 NPF
				Approval	Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton  MSB Exclusion
Ophthalmic Anti-			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Inflammatory Agents -			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
NSAIDs	BromSite	ophthalmic solution	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Ophthalmic Anti- Inflammatory Agents - NSAIDs	Acuvail	ketorolac tromethamine 0.45%	<ol> <li>Approve if the patient has tried two products from the following list (if two are formulary, or one of the following if one is formulary): diclofenac ophthalmic solution (generics), a bromfenac product (0.09% ophthalmic solution [generics], bromfenac ophthalmic solution 0.07% [Prolensa, generics], or bromfenac 0.075% [BromSite, generics]), Nevanac, Ilevro, or ketorolac ophthalmic solution (Acular, Acular LS, generics). If none are formulary, approve.</li> <li>Patients with a sulfite allergy: approve if the patient has tried two of the following, if two are formulary (or one if only one is formulary): bromfenac 0.075% (BromSite, generics), diclofenac ophthalmic solution (generics), Nevanac, Ilevro, or ketorolac ophthalmic solution (Acular, Acular LS, generics). If none are formulary, approve.</li> <li>Patients with a known sensitivity to a preservative (e.g., benzalkonium chloride [BAK]): approve if the patient has tried diclofenac ophthalmic solution (generics), if formulary. If diclofenac ophthalmic solution is non-formulary, approve.</li> <li>Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.</li> </ol>	1 year	Yes
Ophthalmic Anti- Inflammatory Agents - NSAIDs	Nevanac		1. Approve if the patient has tried two products from the following list (if two are formulary, or one of the following if one is formulary): diclofenac ophthalmic solution (generics), ketorolac ophthalmic solution (Acular, Acular LS, generics), Acuvail, llevro, or a bromfenac product (0.09% ophthalmic solution [generics], bromfenac ophthalmic solution 0.07% [Prolensa, generics], or bromfenac 0.075% [BromSite, generics]). If none are formulary, approve.  2. Patients with a sulfite allergy: approve if the patient has tried two of the following, if two are formulary (or one if only one is formulary): bromfenac 0.075% (BromSite, generics), diclofenac ophthalmic solution (generics), llevro, ketorolac ophthalmic solution (Acular, Acular LS, generics), or Acuvail. If none are formulary, approve.  3. Patients < 18 years of age: approve if the patient has tried ketorolac ophthalmic solution (Acular, Acular LS, generics) or llevro, if one is formulary. If neither are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.	1 year	Yes
					MSB Exclusion
Ophthalmic Corticosteroids	Durezol	difluprednate 0.05% ophthalmic emulsion	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	*This criteria applies only to the NPF 7/1/2022
Ophthalmic Corticosteroids	Clobetasol propionate 0.05% ophthalmic suspension	clobetasol propionate	1. Approve if patient has tried three formulary ophthalmic corticosteroids from the following list: 1) a dexamethasone product (generics or Maxidex), 2) a fluorometholone product (FML Liquifilm, generics; FML Forte, Flarex), 3) difluprednate (Durezol, generics), 4) a loteprednol product (Lotemax, generics; Lotemax SM; Inveltys), or 5) a prednisolone product (Pred Forte, Omnipred, generics; Pred Mild), if three are formulary or two if two are formulary or one if one is formulary. If none are formulary, approve.  2. If the patient has elevated intraocular pressure (IOP) or glaucoma after a trial of another corticosteroid (i.e., topical, inhaled, oral, ophthalmic): approve if the patient has tried one of the following, if one is formulary: 1) a fluorometholone product (FML Liquifilm, generics; FML Forte, Flarex), 2) a loteprednol product (Lotemax, generics; Lotemax SM; Inveltys), or 3) difluprednate (Durezol, generics). If none are formulary, approve.	1 year	Yes
Ophthalmic Corticosteroids	Flarex	fluorometholone	1. Approve if patient has tried three formulary ophthalmic corticosteroids from the following list: 1) a dexamethasone product (generics or Maxidex), 2) a fluorometholone product (FML Liquifilm, generics; FML Forte), 3) difluprednate (Durezol, generics), 4) a loteprednol product (Lotemax, generics; Lotemax SM; Inveltys), 5) a prednisolone product (Pred Forte, Omnipred, generics; Pred Mild), or 6) clobetasol propionate ophthalmic suspension, if three are formulary or two if two are formulary or one if one is formulary. If none are formulary, approve.  2. If the patient has elevated intraocular pressure (IOP) or glaucoma after a trial of another corticosteroid (i.e., topical, inhaled, oral, ophthalmic): approve if the patient has tried one of the following, if one is formulary: 1) a fluorometholone product (FML Liquifilm, generics; FML Forte), 2) a loteprednol product (Lotemax, generics; Lotemax SM; Inveltys), or 3) difluprednate (Durezol, generics). If none are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.	1 year	Yes
Ophthalmic Corticosteroids	FML Forte	fluorometholone	1. Approve if patient has tried three formulary ophthalmic corticosteroids from the following list: 1) a dexamethasone product (generics or Maxidex), 2) a fluorometholone product (FML Liquifilm, generics; Flarex), 3) difluprednate (Durezol, generics), 4) a loteprednol product (Lotemax, generics; Lotemax SM; Inveltys), 5) prednisolone (Pred Forte, Omnipred, generics; Pred Mild), or 6) clobetasol propionate ophthalmic suspension, if three are formulary or two if two are formulary or one if one is formulary. If none are formulary, approve.  2. If the patient has elevated intraocular pressure (IOP) or glaucoma after a trial of another corticosteroid (i.e., topical, inhaled, oral, ophthalmic): approve if the patient has tried one of the following, if one is formulary: 1) a loteprednol product (Lotemax, generics; Lotemax SM; Inveltys), 2) a fluorometholone product (FML Liquifilm, generics; Flarex), or 3) difluprednate (Durezol, generics). If none are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.	1 year	Yes
Ophthalmic Corticosteroids	Maxidex		1. Approve if the patient has tried three formulary ophthalmic corticosteroids from the following list (if three are formulary or two if two are formulary or one if one is formulary): 1) dexamethasone (generics), 2) difluprednate (Durezol, generics), 3) a loteprednol product (Lotemax, generics; Lotemax SM; Inveltys), or 4) a fluorometholone product (FML Liquifilm, generics; FML Forte; Flarex), 5) a prednisolone product (Pred Forte, Omnipred, generics; Pred Mild) or 6) clobetasol propionate ophthalmic suspension. If none are formulary, approve.  2. If the patient has elevated intraocular pressure (IOP) or glaucoma after a trial of another corticosteroid (i.e., topical, inhaled, oral, ophthalmic): approve if the patient has tried one product from the following list (if one is formulary): 1) a fluorometholone product (FML Liquifilm, generics; FML Forte; Flarex), 2) a loteprednol product (Lotemax, generics; Lotemax SM; Inveltys), or 3) difluprednate (Durezol, generics). If none are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.	1 year	Yes

					2025 NPF
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	Excluded Medicaiton
Ophthalmic Corticosteroids	Pred Mild	prednisolone acetate 0.12% ophthalmic suspension	1. Approve if the patient has tried three formulary ophthalmic corticosteroids from the following list (if three are formulary or two if two are formulary; or one if one is formulary): 1) a dexamethasone (generics or Maxidex), 2) diffuprednate (Durezol, generics), 3) a loteprednol product (Lotemax, generics; Lotemax SM; Inveltys), 4) a fluorometholone product (FML Liquifilm, generics; FML Forte; Flarex), 5) a prednisolone product (Pred Forte, Omnipred, generics), or 6) clobetasol propionate ophthalmic suspension. If none are formulary, approve.  2. If the patient has elevated intraocular pressure (IOP) or glaucoma after a trial of another corticosteroid (i.e., topical, inhaled, oral, ophthalmic): approve if the patient has tried one product from the following list (if one is formulary): 1) a fluorometholone product (FML Liquifilm, generics; Flarex; FML Forte), 2) a loteprednol product (Lotemax, generics; Lotemax SM; Inveltys), or 3) diffuprednate (Durezol, generics). If none are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.	1 year	Yes
Ophthalmic Drugs for Glaucoma - Beta- Adrenergic Blocker	Istalol	timolol maleate 0.5% ophthalmic solution	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Ophthalmic Drugs for Glaucoma - Beta- Adrenergic Blocker		timolol maleate 0.25% and 0.5%	1. Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with four products from the following list: 1) a timolol product (Istalol, Timoptic/XE, generics), 2) levobunolol ophthalmic solution (generics), 3) betaxolol ophthalmic solution (generics), if four are formulary (or three if three are formulary or two if two are formulary or one if one is formulary). If none are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.  2. Approve if the patient has a known sensitivity to a preservative or when use of a preservative-free topical medication is advisable.	1 year	Yes
Ophthalmic Drugs for Glaucoma - Beta- Adrenergic Blocker	·	timolol hemihydrates 0.25% and 0.5% ophthalmic solution	Approve if the patient has tried four of the following, if formulary (or three if three are formulary or two if two are formulary or one if one is formulary): 1) levobunolol ophthalmic solution, 2) a timolol product (Istalol, Timoptic/XE, generics), 3) a betaxolol ophthalmic solution (generics or Betoptic S), or 4) carteolol ophthalmic solution (generics). If none are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.	1 year	Yes
Ophthalmic Drugs for Glaucoma - Carbonic Anhydrase Inhibitor	Azopt	brinzolamide 1% ophthalmic suspension	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Ophthalmic Drugs for Glaucoma - Carbonic Anhydrase Inhibitor/Beta- Adrenergic Blocker		dorzolamide 2%/timolol 0.5% ophthalmic solution	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Ophthalmic Drugs for Glaucoma - Prostaglandins	Travatan Z	travoprost 0.004% ophthalmic solution (benzalkonium chloride-free)	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Ophthalmic Drugs for Glaucoma - Prostaglandins	Xalatan	latanoprost 0.005% ophthalmic solution	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Ophthalmic Drugs for Glaucoma - Prostaglandins	Zioptan	tafluprost 0.0015% ophthalmic solution	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Ophthalmic Drugs for Glaucoma - Prostaglandins	Lumigan	bimatoprost 0.01% ophthalmic solution	Approve if the patient has tried four formulary alternatives from the following list (or three if three are formulary or two if two are formulary or one if only one is formulary): latanoprost ophthalmic solution (Xalatan, generics), bimatoprost 0.03% ophthalmic solution (generics), travoprost ophthalmic solution (Travatan Z, generics), Vyzulta, or tafluprost ophthalmic solution (Zioptan, generics). If none are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.	1 year	Yes
Ophthalmic Drugs for Glaucoma - Prostaglandins	Vyzulta	latanoprostene bunod ophthalmic solution 0.024%	Approve if the patient has tried four formulary alternatives from the following list (or three if three are formulary or two if two are formulary or one if only one is formulary): latanoprost ophthalmic solution (Xalatan, generics), travoprost ophthalmic solution (Travatan Z, generics), bimatoprost 0.03% ophthalmic solution (generics), Lumigan, or tafluprost ophthalmic solution (Zioptan, generics). If none are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.  1. Approve if the patient has tried, and according to the prescriber, experienced inadequate efficacy OR significant intolerance with one of 1) latanoprost ophthalmic solution (Xalatan, generics) or 2) Xelpros, if formulary. If none are	1 year	Yes
Ophthalmic Drugs for Glaucoma -		0.005%; preservative-	formulary, approve.  2. If according to the prescriber, the patient has a significant benzalkonium chloride allergy/sensitivity: approve if the patient has tried, and according to the prescriber, experienced inadequate efficacy OR significant intolerance with Xelpros, if formulary. If Xelpros is non-formulary, approve.		
Prostaglandins	lyuzeh	free	3. If, according to the prescriber, the patient has a significant allergy/sensitivity to other preservatives (OTHER than benzalkonium chloride), approve.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
			1. Approve if the patient has tried, and according to the prescriber, experienced inadequate efficacy OR significant intolerance with one of 1) latanoprost ophthalmic solution (Xalatan, generics) or 2) lyuzeh, if formulary. If none are		
Ophthalmic Drugs for Glaucoma -		latanoprost 0.005%	formulary, approve.  2. If, according to the prescriber, the patient has a significant benzalkonium chloride allergy/sensitivity: approve if the patient has tried, and according to the prescriber, experienced inadequate efficacy OR significant intolerance with		
Prostaglandins	Xelpros	ophthalmic emulsion	lyuzeh, if formulary. If lyuzeh is non-formulary, approve.	1 year	Yes
Ophthalmic Drugs for Glaucoma – Prostaglandins – Implants	Durysta	bimatoprost implant	Approve if the patient meets the following (A, B and C):  A. The patient has tried and, according to the prescriber, experienced inadequate efficacy OR significant intolerance with at least two ophthalmic prostaglandins (either as monotherapy or as concomitant therapy); AND  Note: Examples of ophthalmic prostaglandins include bimatoprost 0.03% ophthalmic solution, latanoprost 0.005% ophthalmic solution, travoprost 0.004% ophthalmic solution; Lumigan® (bimatoprost 0.01% ophthalmic solution), Vyzulta® (latanoprostene bunod 0.024% ophthalmic solution), Xelpros™ (latanoprost 0.005% ophthalmic solution, Iyuzeh (latanoprost 0.005% ophthalmic solution), and Omlonti (omidenepag isopropyl 0.002% ophthalmic solution).  B. The patient has tried and, according to the prescriber, experienced inadequate efficacy OR significant intolerance with at least two other ophthalmic products from two different pharmacological classes (either as monotherapy or as concomitant therapy); AND  Note: Examples of pharmacological classes of ophthalmic products for the treatment of open-angle glaucoma or ocular hypertension include beta-blockers, alpha-agonist (brimonidine), carbonic anhydrase inhibitors, and rho kinase inhibitor (netarsudil).  C. The product is NOT being used for re-treatment of an eye previously treated with Durysta.  Note: Durysta is approved for a one-time use in each eye. Repeat administration in previously treated eye(s) is not approvable.	30 days	Yes
Ophthalmic Drugs for Glaucoma – Prostaglandins – Implants	iDose TR	travoprost intracameral implant	Approve if the patient meets the following (A, B and C):  A. The patient has tried and, according to the prescriber, experienced inadequate efficacy OR significant intolerance with at least two ophthalmic prostaglandins (either as monotherapy or as concomitant therapy); AND Note: Examples of ophthalmic prostaglandins include bimatoprost 0.03% ophthalmic solution, latanoprost 0.005% ophthalmic solution, travoprost 0.004% ophthalmic solution; Lumigan® (bimatoprost 0.01% ophthalmic solution), Vyzulta® (latanoprost bunod 0.024% ophthalmic solution), Xelpros™ (latanoprost 0.005% ophthalmic solution, latanoprost 0.0015% ophthalmic solution, lyuzeh (latanoprost 0.005% ophthalmic solution), and Omlonti (omidenepage isopropyl 0.002% ophthalmic solution).  B. The patient has tried and, according to the prescriber, experienced inadequate efficacy OR significant intolerance with at least two other ophthalmic products from two different pharmacological classes (either as monotherapy or as concomitant therapy); AND Note: Examples of pharmacological classes of ophthalmic products for the treatment of open-angle glaucoma or ocular hypertension include beta-blockers, alpha-agonist (brimonidine), carbonic anhydrase inhibitors, and rho kinase inhibitor (netarsudil).  C. The product is NOT being used for re-treatment of an eye previously treated with iDose TR.  Note: iDose TR is approved for a one-time use in each eye. Repeat administration in previously treated eye(s) is not approvable.	30 days	Yes
mpianto	IDOGC TIT	muddamorai impiant	15000 Trito approved for a one while add in each eye. He pear administration in providing about a operation.	oo aayo	MSB Exclusion
Opiate			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		*This criteria applies only to
Agonists/Antagonists Opioids (Oral) - Other	Suboxone	ne sublingual film oxycodone/acetamino phen tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year 1 year	the NPF MSB Exclusion *This criteria applies only to the NPF
Opioids (Oral) - Other	Prolate solution	oxycodone and acetaminophen 10- 300 mg/5 oral solution	1. Approve if the patient has tried and cannot take oxycodone-acetaminophen 10-325 mg tablets. 2. Approve if the patient is unable to swallow or has difficulty swallowing tablets.	1 year	Yes
Opioids (Oral) - Other	Nucynta	tapentadol immediate- release tablets	<ol> <li>Approve if the patient has tried three other oral immediate-release (NOT long-acting) centrally acting/opioid analgesics. Examples of oral immediate-release (NOT long-acting) centrally acting/opioid analgesics include, but are not limited to: hydromorphone (Dilaudid, generics), oxycodone hydrochloride tablets (Roxicodone, generics), oxymorphone (generics), morphine (generics), hydrocodone/acetaminophen (Vicodin, Vicodin ES, Norco, Lortab, Lorcet, multiple generics), oxycodone/acetaminophen (Percocet, Endocet, Roxicet, multiple generics), tramadol (Ultram, generics), tramadol/acetaminophen (Ultracet, generics).</li> <li>NOTE: A trial of the requested product does not count toward this requirement.</li> <li>Patients ≥ 6 years of age to &lt; 18 years of age, approve if the patient meets ONE of the following (A, B, or C):         <ul> <li>A. Patient has tried one of morphine sulfate immediate-release tablets or morphine sulfate immediate-release oral solution. If neither are formulary, approve; OR</li> <li>B. Patient has renal insufficiency; OR</li> <li>C. Patient is intolerant or allergic to morphine.</li> </ul> </li> </ol>	1 year	Yes
	Qdolo and	tramadol hydrochloride oral			
Opioids (Oral) - Other	authorized generic	solution	Approve if the patient is unable to swallow or has difficulty swallowing tramadol tablets.	1 year	Yes
Opioids (Oral) - Other	Oxaydo	oxycodone hydrochloride tablets	Approve if the patient has tried and cannot take one of the following formulary products: oxycodone hydrochloride tablets (Roxicodone, generics). If oxycodone hydrochloride tablets (Roxicodone, generics) are non-formulary, approve.	1 year	Yes

					2025 NPF
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	Excluded Medicaiton
Thorapy Glado	Diana itamo	oxycodone	Other Control of the	Duration	Modicalion
	Roxybond and	hydrochloride tablet,			
Opioids (Oral) - Other		coated	Approve if the patient has tried and cannot take one of the following formulary products: oxycodone hydrochloride tablets (Roxicodone, generics). If oxycodone hydrochloride tablets (Roxicodone, generics) are non-formulary, approve.	1 year	Yes
Opioids (Oral) - Other	tramadol 25 mg	tramadol 25 mg tablets	Approve if the prescribed dose cannot be obtained with tramadol 50 mg.  Note: The patient is NOT required to split the 50 mg tablets in half.	1 vear	Yes
Opiolus (Orai) - Otilei	tablets (blaild)	tablets	NOTE: The patient is NOT required to spirit the 50 mg tablets in rian.	i yeai	163
	oxycodone-				
	acetaminophen 10-	oxycodone-			
Opioids (Oral) - Other		acetaminophen 10- 300 mg tablets	1. Direct to oxycodone-acetaminophen 10-325 mg tablets.	1 vear	Yes - Primlev only
Opioids (Orai) - Other	Primiev, Prolate)	300 mg tablets	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use oxycodone-acetaminophen 10-325 mg tablets.	i year	only
	oxycodone-				
	acetaminophen 5-	oxycodone-			
	300 tablets (includes		1. Direct to oxycodone-acetaminophen 5-325 mg tablets.		Yes - Primlev
Opioids (Oral) - Other	Primlev, Prolate)	mg tablets	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use oxycodone-acetaminophen 5-325 mg tablets.	1 year	only
	oxvcodone-				
	,	oxycocodne-			
	300 tablets (includes	•	1. Direct to oxycodone-acetaminophen 7.5-325 mg tablets.		Yes - Primlev
Opioids (Oral) - Other	Primlev and Prolate)	300 mg tablets	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use oxycodone-acetaminophen 7.5-325 mg tablets.	1 year	only
Opioids (Oral) - Other	Conzip and tramadol	tramadol ER capsule	Approve, if per the prescriber, the patient is unable to use generic tramadol ER tablets.	1 year	Yes
Opiolus (Oral) - Otilei	tramadol 100 mg	tramadol 100 mg	Approve, if per title prescriber, the patient is unable to use generic trainautor EN tablets.	i yeai	165
Opioids (Oral) - Other	tablets (brand)	tablets	Approve, if per the prescriber, the patient is unable to use generic tramadol 50 mg tablets.	1 year	Yes
	levorphanol 2mg	levorphanol 2mg and	Approve if the patient has tried three medications (each from a different group) of the following: a morphine-containing product, a hydrocodone-containing product, a hydrocodone-containing product, an oxycodone-containing product,		
Opioids (Oral) - Other	and 3mg	3mg tablets	an oxymorphone-containing product, a fentanyl-containing product, a methadone-containing product, or a tapentadol-containing product.	1 year	Yes
		celecoxib and			
Opioids (Oral) –		tramadol	1. Direct the patient to tramadol tablets and celecoxib capsules as separate agents. If celecoxib capsules (Celebrex, generics) are non-formulary, approve.		
- 1 ( - /	Seglentis	hydrochloride tablets	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use tramadol and celecoxib as separate agents.	1 year	Yes
		ciprofloxacin 0.2%			
Otic Antibiotics	Cetraxal	otic solution	Approve if the patient has tried one of the following, if one is formulary: ofloxacin otic solution (generics) or ciprofloxacin 0.2% otic solution (generic). If none are formulary, approve.	1 year	Yes
			A Association to the stind both and the following link of the stind both and the following link of the stind both and the stind		
Otic Antibiotics and	Cipro HC Otic	ciprofloxacin/ hydrocortisone otic	1. Approve if the patient has tried both products from the following list: 1) ciprofloxacin-dexamethasone otic suspension (Ciprodex otic suspension, generics) and 2) ciprofloxacin-fluocinolone otic (authorized generic of Otovel) or Otovel otic solution, if formulary. If none are formulary, approve.		
Combination Products		,	2. Patient has a benzalkonium chloride sensitivity: approve if the patient has tried one of ciprofloxacin-fluocinolone otic (authorized generic of Otovel) or Otovel, if formulary, If neither are formulary, approve.	1 year	Yes
			1. Direct the patient to Otovel (brand), if formulary.		
		•	2. If Otovel (brand) is non-formulary, approve if the patient has tried both 1) ciprofloxacin-dexamethasone otic suspension (Ciprodex otic suspension, generics) and 2) Cipro HC otic suspension (or one if one is formulary). If neither are		
	ciprofloxacin/ fluocinolone otic	fluocinolone acetonide otic	formulary, approve.  3. If Otovel (brand) is non-formulary, patients treating acute otitis media through tympanostomy tubes (AOMT), patients with a perforated ear drum (tympanic membrane), or patients < 1 year of age: approve if the patient has tried		
	solution (authorized		ciprofloxacin- dexamethasone otic suspension (Ciprodex otic suspension, generics), if formulary. If ciprofloxacin- dexamethasone otic suspension, generics) are non-formulary, approve.		
Combination Products		0.3%/0.025%	4. If Otovel (brand) is non-formulary, patient has a known hypersensitity to a preservative (e.g., benzalkonium chloride [BAK], benzyl alcohol), approve.	1 year	Yes
Overactive Bladder			Approve if the patient has tried oxybutynin oral solution or syrup, if formulary. If neither oxybutynin oral solution nor syrup is formulary approve if the patient meets one of the following (A or B):		
	Oxybutynin 2.5 mg tablet (brand)	oxybutynin 2.5 mg tablet	A. Patient has tried other strengths of oxybutynin tablets; OR	1 year	Voc
Topical)	tablet (brand)	lablet	B. Patient's dose requires a 2.5 mg increment.	1 year	Yes MSB Exclusion
Overactive Bladder			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Agents (Oral and			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Topical)	Detrol	tolterodine tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
0 " -:			NOTE. A multi-come Daniel and the being a multi-chair and the multi-chair and big multi-chair and big multi-chair and the second big multi-chair and the se		MSB Exclusion
Overactive Bladder Agents (Oral and		tolterodine ovtonded	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		*This criteria applies only to
•	Detrol LA	tolterodine, extended- release capsules	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
· opioui)	DOMOI LIV	. c. saco oapsaics	research result in a significant directly of sortion forest internation required.	. you	

					2025 NDF
				Approval	2025 NPF Excluded
Class Brai	rand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
Bladder and	:	solifenacin succinate	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		MSB Exclusion *This criteria applies only to
Vesicare	are	tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
sladder ) Toviaz			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].  1. Approve if the patient has tried oxybutynin solution/syrup OR Myrbetrig Granules, if formulary. If neither are formulary, approve.	1 year	MSB Exclusion *This criteria applies only to the NPF
sladder ) Vesicare			2. Patient is < 5 years of age: approve if the patient has tried Myrbetriq Granules, if formulary. If Myrbetriq Granules are non-formulary, approve.  3. Patients < 3 years of age, approve.  Note: If the patient has tried any oxybutynin-containing product (e.g., immediate-release or extended-release tablets), this would meet the requirement for a trial of an oxybutynin product.  Note: If the patient has tried Mybetriq tablets, this would meet the requirement for a trial of Myrbetriq granules.	1 year	Yes
nzymes Pertzye		pancrelipase delayed- release capsules	Approve if the patient has tried three products from the following list (if three are formulary, or two if two are formulary, or one if one is formulary): Creon, Pancreaze, or Zenpep. If none are formulary, approve.	1 year	Yes
uria Kuvan	:	·	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	i yeai	MSB Exclusion *This criteria applies only to the NPF
Fosreno inders tablets			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
inders Renage			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Auryxia inders authoriz		ferric citrate tablets	1. Approve if the patient has tried two formulary alternative from the following list (if two are formulary, or one if one is formulary): sevelamer hydrochloride tablets, sevelamer carbonate tablets/powder for oral suspension (Renvela, generics), Velphoro chewable tablets, lanthanum carbonate chewable tablets (Fosrenol, generics), or Fosrenol oral powder. If none are formulary, approve.  2. Patients with iron deficient anemia and chronic kidney disease who are NOT on dialysis: approve.	1 year	Yes - only authorized generic
Fosreno inders powder		lanthanum carbonate oral powder	1. Approve if the patient has tried two formulary alternatives from the following list (if two are formulary or one if one is formulary): sevelamer hydrochloride tablets, lanthanum carbonate chewable tablets (Fosrenol, generics), Velphoro chewable tablets, Auryxia tablets, or sevelamer carbonate tablets/powder for oral suspension (Renvela, generics). If none are formulary, approve.  Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.  2. Patients who are unable to chew and swallow tablets: approve if the patient has tried sevelamer carbonate powder for oral suspension (Renvela powder, generics), if formulary. If sevelamer carbonate powder for oral suspension (Renvela powder, generics) is non-formulary, approve.	1 year	Yes
paring Carospi		spironolactone oral suspension	<ol> <li>Approve if the patient has tried and cannot take spironolactone tablets (Aldactone, generics), if formulary. If spironolactone tablets (Aldactone, generics) are non-formulary, approve.</li> <li>Approve if the patient cannot swallow spironolactone tablets.</li> </ol>	1 year	Yes
		potassium chloride			
Pokonza		powder, for solution ascorbic acid, cholecalciferol, thiamine hydrochloride, riboflavin,pyridoxal phosphate,levomefola te glucosamine, folic acid, methylcobalamin, calcium carbonate, ferrous gluconate, potassium iodide	Approve if the patient has tried one other oral potassium chloride product (e.g., potassium chloride powder for oral solution, potassium chloride oral solution).  1. Direct to generic prenatal vitamins.	1 year	Yes
amins Natal Pi			2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use generic prenatal vitamins.	1 year	

				A	2025 NPF
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	Excluded Medicaiton
	Citranatal prenatal				
	vitamins (examples				
	include Citranatal RX tablets,				
	Citranatal Harmony		1. Direct to generic prenatal vitamins.		
Prenatals vitamins	capsules)	various	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use generic prenatal vitamins.	1 year	Yes
		beta carotene,			
		ascorbic acid, cholecalciferol,			
		.alphatocopherol			
		acetate, pyridoxine			
		hydrochloride, biotin,			
		folic acid, levomefolate calcium,			
		cyanocobalamin,			
		calcium carbonate,			
		magnesium oxide,			
		ferrous bisglycinate,	1. Direct to generic prenatal vitamins.		
Prenatals vitamins	Pregenna	tablet	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use generic prenatal vitamins.	1 year	Yes
		ascorbic acid,			
		cholecalciferol,			
		thiamine			
		hydrochloride, riboflavin, pyridoxal			
		phosphate anhydrous,			
		folic acid,			
		methylcobalamin,			
		calcium carbonate, ferrous gluconate,			
			1. Direct to generic prenatal vitamins.		
Prenatals vitamins	Trinaz		2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use generic prenatal vitamins.	1 year	Yes
		pilocarpine 1.25%	No exception is recommended.		,
Presbyopia Agents	Vuity	ophthalmic solution	(NOTE: It is not appropriate to use standard global criteria for this medication; Denial reason is: Formulary coverage is not provided for this medication.)	N/A	Yes
		pilocarpine hydrochloride			
		ophthalmic solution,	No exception is recommended.		
Presbyopia Agents	Qlosi	0.4%	(NOTE: It is not appropriate to use standard global criteria for this medication; Denial reason is: Formulary coverage is not provided for this medication.)	N/A	Yes
Primary				Coo DA	
Immunoglobulin A Nephropathy Agents	Filspari	sparsentan tablets	See standard Nephrology – Filspari Prior Authorization Policy criteria.	See PA duration	Yes
pinopaary / gorito	opan		1. Approve if the patient has tried Crinone 8% gel [documentation required], if formulary. If Crinone 8% gel is non-formulary, approve.	24.4.011	
Progestin – Vaginal		progesterone vaginal	2. A patient already started on a course of therapy with Endometrin for progesterone supplementation/replacement to achieve or maintain pregnancy: approve to complete the current course of therapy.		
Agents	Endometrin	insert	Note: Approve for 9 months or to complete the course of therapy.	1 year	Yes
Progestin – Vaginal	Crinono 40/ Csl	progestorone gel 40/	Approve if the patient has tried one product from the following list (if one is formulary): medroxyprogesterone [Provera, generics], megestrol acetate, norethindrone tablets, or progesterone capsules (Prometrium, generics). If none are	1 1000	Vac
Agents	Crinone 4% Gel	progesterone gel 4%	portitularly, approve.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Proton Pump Inhibitor Combination	Yosprala and authorized generic	aspirin and omeprazole delayed-release tablets	Approve if the patient has tried aspirin AND at least five proton pump inhibitors (PPIs).  Note: Examples of PPIs include a rabeprazole delayed-release (DR) product (Aciphex, generics; Aciphex Sprinkle, authorized generics), dexlansoprazole DR capsules (Dexilant, generics), an esomeprazole DR product (Nexium, generics; Nexium granules for oral suspension, generics), a pantoprazole DR product (Protonix, generics), an omeprazole DR product (Priosec, generics; Priosec granules for oral suspension), an omeprazole DR product (Priosec, generics; Konvomep).  Note: The requested agent would NOT count as a trial of an alternative.	1 year	Yes - brand only
Proton Pump Inhibitors (PPIs)	Aciphex	rabeprazole sodium tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Proton Pump Inhibitors (PPIs)	Nexium capsules	esomeprazole delayed-release capsules	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Proton Pump Inhibitors (PPIs)	Nexium packet (granules for oral suspension) 10 mg, 20 mg, 40 mg packet	esomeprazole delayed-release granules for oral suspension (packet)	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Proton Pump Inhibitors (PPIs)	Prevacid	lansoprazole delayed- release (DR) capsules	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Proton Pump Inhibitors (PPIs)	Protonix	pantoprazole sodium delayed-release (DR) tablets and intravenous (IV) injection	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Proton Pump Inhibitors (PPIs)	Protonix oral suspension	release oral	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Proton Pump Inhibitors (PPIs)	Prevacid SoluTab	lansoprazole orally disintegrating tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Proton Pump Inhibitors (PPIs)	Konvomep	omeprazole and sodium bicarbonate oral suspension	1. Approve if the patient has tried at least five proton pump inhibitors (PPIs).  Note: Examples of PPIs include a rabeprazole delayed-release (DR) product (Aciphex, generics; Aciphex Sprinkle, authorized generics), dexlansoprazole DR capsules (Dexilant, generics), an esomeprazole DR product (Nexium, generics; Nexium granules for oral suspension, generics), a lansoprazole DR product (Prevacid, generics; Prevacid SoluTab, generics), an omeprazole DR product (Priosec, generics; Prilosec granules for oral suspension), an omeprazole DR/sodium bicarbonate product (Zegerid, generics; Zegerid powder for oral suspension, generics).  Note: The requested agent would NOT count as a trial of an alternative.  2. Patients who have difficulty swallowing tablets/capsules: Approve if the patient has tried at least four proton pump inhibitors (PPIs) from the following list, if formulary (or three if three are formulary, or one if one is formulary): 1) rabeprazole sprinkle; 2) an esomeprazole product (esomeprazole DR capsules [Nexium, generics], esomeprazole packet [Nexium granules for oral suspension, generic]); 3) pantoprazole suspension (granules) [Protonix suspension, generic]; 4) a lansoprazole product (Innsoprazole DR capsules [Prevacid, generics], lansoprazole oral disintegrating tablets [Prevacid SoluTab, generics]); 5) an omeprazole product (omeprazole DR capsules [Prilosec, generics], Prilosec DR suspension). If none are formulary, approve.	1 year	Yes

					2025 NPF
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	Excluded Medicaiton
Proton Pump Inhibitors (PPIs)	Dexilant and generic dexlansoprazole delayed-release capsules	dexlansoprazole delayed-release capsules	1. Approve if the patient has tried at least five proton pump inhibitors (PPIs).  Note: Examples of PPIs include a rabeprazole delayed-release (DR) product (Aciphex, generics; Aciphex Sprinkle, authorized generics), dexlansoprazole DR capsules (Dexilant, generics), an esomeprazole DR product (Nexium, generics; Nexium granules for oral suspension, generics), a pantoprazole DR product (Protonix, generics; Protonix granules for oral suspension, generics; Product (Prevacid, generics; Prevacid SoluTab, generics), an omeprazole DR product (Prilosec, generics; Prilosec granules for oral suspension), an omeprazole DR/sodium bicarbonate product (Zegerid, generics; Zegerid powder for oral suspension, generics; Konvomep).  Note: The requested agent would NOT count as a trial of an alternative.  2. Patients < 18 years of age OR patients who have difficulty swallowing tablets/capsules: Approve if the patient has tried at least four proton pump inhibitors (PPIs) from the following list, if formulary (or three if three are formulary, or two if two are formulary), or one if one is formulary: 1) rabeprazole sprinkle; 2) an esomeprazole DR capsules [Nexium, generics], esomeprazole packet [Nexium granules for oral suspension, generics]); 3) pantoprazole suspension (granules) [Protonix suspension, generics]; 4) a lansoprazole product (lansoprazole DR capsules [Prevacid, generics], lansoprazole oral disintegrating tablets [Prevacid Solutab, generics]); 5) an omeprazole product (omeprazole DR capsules [Prilosec, generics], Prilosec DR suspension). If none are formulary, approve.	1 year	Yes
Proton Pump Inhibitors (PPIs)	Aciphex Sprinkle and authorized generic	rabeprazole sodium delayed-release capsules	1. Approve if the patient has tried at least five proton pump inhibitors (PPIs).  Note: Examples of PPIs include dexlansoprazole DR capsules (Dexilant, generics), an esomeprazole DR product (Nexium, generics; Nexium granules for oral suspension, generics), a pantoprazole DR product (Protonix, generics; Protonix granules for oral suspension, generics), a lansoprazole DR product (Prevacid, generics; Prevacid SoluTab, generics), an omeprazole DR product (Prilosec, generics; Prilosec granules for oral suspension), an omeprazole DR/sodium bicarbonate product (Zegerid, generics; Zegerid powder for oral suspension, generics; Konvomep). Note: The requested agent would NOT count as a trial of an alternative.  2. Patients < 18 years of age OR patients who have difficulty swallowing tablets/capsules: Approve if the patient has tried at least two proton pump inhibitors (PPIs).	1 year	Yes Authorized generic only
Proton Pump Inhibitors (PPIs)	Nexium packet (granules for oral suspension) 5 mg and 2.5 mg packets and esomeprazole delayed-release 5 mg and 2.5 mg granules for oral suspension (packets)	esomeprazole delayed-release granules for oral suspension (packet)	If requesting brand Nexium oral suspension (packet):  Approve if the patient has tried and cannot take generic esomeprazole delayed-release granules for oral suspension (packet) due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product, if the generic is formulary.  If requesting generic esomeprazole delayed-release granules for oral suspension or generic esomeprazole delayed-release granules for oral suspension are non-formulary:  1. Approve if the patient has tried at least five proton pump inhibitors (PPIs).  Note: Examples of PPIs include a rabeprazole delayed-release (DR) product (Aciphex, generics; Aciphex Sprinkle, authorized generics), dexlansoprazole DR capsules (Dexilant, generics), an esomeprazole DR product (Nexium, generics), a pantoprazole DR product (Protonix, generics; Protonix granules for oral suspension, generics; Prevacid, generics; Prevacid SoluTab, generics), an omeprazole DR product (Prilosec, generics; Prilosec granules for oral suspension), an omeprazole DR/sodium bicarbonate product (Zegerid, generics; Zegerid powder for oral suspension, generics; Konvomep).  Note: The requested agent would NOT count as a trial of an alternative.  2. Patients < 18 years of age OR patients who have difficulty swallowing tablets/capsules: Approve if the patient has tried at least two proton pump inhibitors (PPIs).  3. Patients < 19 years of age and require administration via a feeding tube, approve if the patient has tried Prilosec DR suspension, if formulary. If Prilosec DR suspension is non-formulary, approve.	1 year	Yes
Proton Pump Inhibitors (PPIs)	Prilosec oral suspension	omeprazole delayed- release oral suspension	1. Approve if the patient has tried at least five proton pump inhibitors (PPIs).  Note: Examples of PPIs include a rabeprazole delayed-release (DR) product (Aciphex, generics; Aciphex Sprinkle, authorized generics), dexlansoprazole DR capsules (Dexilant, generics), an esomeprazole DR product (Nexium, generics; Nexium granules for oral suspension, generics), a pantoprazole DR product (Protonix, generics; Protonix granules for oral suspension, generics), a lansoprazole DR product (Prevacid, generics; Prevacid SoluTab, generics), an omeprazole DR product (Prilosec, generics), an omeprazole DR product (Prilosec, generics), an omeprazole DR product (Protonix, generics; Prevacid SoluTab, generics), an omeprazole generics (Prilosec, generics), an omeprazole DR product (Protonix, generics), an omeprazole generics (Prilosec, generics), an omeprazole DR product (Prilosec, generics), an omeprazole DR product (Protonix, generics), an omeprazole generics), an omeprazole generics, a lansoprazole DR product (Prevacid, generics; Prevacid SoluTab, generics), an omeprazole DR product (Protonix, generics), a lansoprazole DR product (Protonix, generics), a lansoprazole DR product (Protonix, generics), a lansoprazole DR product (Protonix, generics)	1 year	Yes
Proton Pump Inhibitors (PPIs)	Zegerid capsules	omeprazole/ sodium bicarbonate capsules	Approve if the patient has tried at least five proton pump inhibitors (PPIs).  Note: Examples of PPIs include a rabeprazole delayed-release (DR) product (Aciphex, generics; Aciphex Sprinkle, authorized generics), dexlansoprazole DR capsules (Dexilant, generics), an esomeprazole DR product (Nexium, generics; Nexium granules for oral suspension, generics), a pantoprazole DR product (Protonix, generics; Protonix granules for oral suspension, generics; Prevacid SoluTab, generics), an omeprazole DR product (Prilosec, generics; Prilosec granules for oral suspension), Konvomep.  Note: The requested agent would NOT count as a trial of an alternative.	1 year	Yes
Proton Pump Inhibitors (PPIs)	Zegerid packets	omeprazole/ sodium bicarbonate powder for oral suspension (packets)	Approve if the patient has tried at least five proton pump inhibitors (PPIs).  Note: Examples of PPIs include a rabeprazole delayed-release (DR) product (Aciphex, generics; Aciphex Sprinkle, authorized generics), dexlansoprazole DR capsules (Dexilant, generics), an esomeprazole DR product (Nexium, generics; Nexium granules for oral suspension, generics), a pantoprazole DR product (Protonix, generics; Protonix granules for oral suspension, generics; Prevacid SoluTab, generics), an omeprazole DR product (Prilosec, generics; Prilosec granules for oral suspension), Konvomep.  Note: The requested agent would NOT count as a trial of an alternative.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Pulmonary Arterial Hypertension (PAH) - Endothelin Receptor Antagonists	Letairis		NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Pulmonary Arterial Hypertension (PAH) - Phosphodiesterase 5 Inhibitors	Adcirca		NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Pulmonary Arterial Hypertension (PAH) - Phosphodiesterase 5 Inhibitors	Ligrev		Pulmonary arterial hypertension World Health Organization Group 1.  1. Direct the patient to sildenafil powder for oral suspension 10 mg/mL (Revatio oral suspension, generics), if formulary.  2. Approve if, according to the prescriber, there is a significant clinical concern (e.g., a significant allergy or serious adverse reaction due to inactive ingredients) such that the patient is unable to use sildenafil powder for oral suspension 10 mg/mL (Revatio oral suspension, generics).  3. If sildenafil powder for oral suspension (10 mg/mL) is non-formulary, approve if the patient meets one of the following (A or B):  A. Patient has tried Tadliq, if formulary. If Tadliq is non-formulary, approve; OR  Note: This criterion would also be satisfied if the patient tried any other tadalafil product.  B. Patient has already been started on a sildenafil product (e.g., sildenafil tablets or suspension, Revatio, or Liqrev).	1 year	Yes
Pulmonary Arterial Hypertension (PAH) - Phosphodiesterase 5 Inhibitors	Tadliq		Pulmonary arterial hypertension World Health Organization Group 1.  1. Approve if the patient is unable to swallow or has difficulty swallowing tadalafil tablets (Adcirca tablets, Alyq tablets, generics), if formulary.  2. If tadalafil tablets (Adcirca tablets, Alyq tablets, generics) are non-formulary, approve if the patient meets one of the following (A or B):  A. Patent has tried sildenafil powder for oral suspension (Revatio oral suspension, generics), if formulary. If sildenafil powder for oral suspension, generics) is non-formulary approve; OR  Note: This criterion would also be satisfied if the patient tried any other sildenafil product.  B. Patient has already been started on a tadalafil product (e.g., tadalafil tablets, Adcirca tablets, Alyq, Tadliq).	1 year	Yes
Respiratory - Corticosteroid			1. Approve if the patient has tried four formulary alternatives from the following list (if four are formulary, or three if three are formulary, or two if two are formulary, or one if only one is formulary): a fluticasone inhaler (Arnuity Ellipta, fluticasone propionate diskus [authorized generic of Flovent Diskus], fluticasone propionate HFA [authorized generic of Flovent HFA]), a mometasone inhaler (Asmanex HFA, Asmanex Twisthaler), Pulmicort Flexhaler, or Qvar RediHaler. If none are formulary, approve.  2. If the patient has a low inspiratory flow rate and is unable to use a dry powder inhaler, approve if the patient has tried three formulary alternatives from the following list (if three are formulary, or two if two are formulary, or one if only one is formulary): Asmanex HFA, fluticasone propionate HFA (authorized generic of Flovent HFA), or Qvar RediHaler. If none are formulary, approve.  Note: If the patient tried the brand version of an authorized generic equivalent product, then this trial would count towards the requirement.  Note: Arnuity Ellipta, fluticasone propionate diskus [authorized generic of Flovent Diskus], and fluticasone propionate HFA (authorized generic of Flovent HFA) would count as one alternative. Asmanex HFA and Asmanex Twisthaler would count as one alternative.		
Inhalers	Alvesco	aerosol	3. Approve if the patient is pregnant AND currently receiving the requested medication for asthma.	1 year	Yes

					2025 NPF
Therany Class	Brand Name	Generic Name	Commercial FF Criteria	Approval	Excluded Medicaiton
Respiratory - Corticosteroid	Brand Name		1. Approve if the patient has tried five formulary alternatives from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if only one is formulary): Alvesco, a fluticasone inhaler (Amulty Ellipta, fluticasone propionate diskus [authorized generic of Flovent Diskus], fluticasone propionate HFA [authorized generic of Flovent HFA]), a mometasone inhaler (Asmanex HFA, Asmanex Twisthaler), Pulmicort Flexhaler, or Ovar Redithaler. If none are formulary, approve.  a. If the patient is < 12 years of age, approve if the patient has tried four formulary alternatives from the following list (if four are formulary, or two if two are formulary, or one if only one is formulary): a fluticasone inhaler (Armulty Ellipta, fluticasone propionate diskus [authorized generic of Flovent Diskus], fluticasone propionate HFA [authorized generic of Flovent HFA]), a mometasone inhaler (Asmanex HFA, Asmanex Twisthaler), Pulmicort Flexhaler, or Ovar Redithaler. If none are formulary, approve.  1. If the patient is < 12 years of age and is unable to coordinate breath and actuation with a conventional metered-dose inhaler (Armulty Ellipta, fluticasone propionate diskus [authorized generic of Flovent Diskus]), Pulmicort Flexhaler, or Ovar Redithaler. If none are formulary, or one if only one is formulary): a fluticasone inhaler (Armulty Ellipta, fluticasone propionate diskus [authorized generic of Flovent Diskus]), Pulmicort Flexhaler, or two if two are formulary, or one if only one is formulary): a fluticasone inhaler (Armulty Ellipta, fluticasone propionate diskus [authorized generic of Flovent Diskus]), Pulmicort Flexhaler, or two if two are formulary, or one if only one is formulary): Asmanex Twisthaler, a fluticasone inhaler (Armulty Ellipta, fluticasone propionate diskus [authorized generic of Flovent Diskus]), or one if only one is formulary): Asmanex Twisthaler, a fluticasone inhaler (Armulty Ellipta, fluticasone propionate diskus [authorized gene	Duration	Medicaiton
Respiratory - Corticosteroid	ArmonAir Digihaler  Flovent Diskus (brand and	powder, metered	1. Approve if the patient is pregnant AND currently receiving the requested medication for asthma.  1. Approve if the patient is pregnant and the patient has tried five formulary alternatives from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if only one is formulary); Alvesco, a fluticasone inhaler (Armuity Ellipta, fluticasone propionate HFA [authorized generic of Flovent HFA]), a mometasone inhaler (Asmanex Twisthaler), Pulmicort Flexhaler, or Ovar RediHaler. If none are formulary, approve.  a. If the patient is < 12 years of age, approve if the patient has tried four formulary alternatives from the following list (if four are formulary, or three if three are formulary, or two if two are formulary, or one if only one is formulary); a fluticasone inhaler (Armuity Ellipta, Fluticasone propionate HFA [authorized generic of Flovent HFA]), a mometasone inhaler (Asmanex Twisthaler, Asmanex HFA), Pulmicort Flexhaler, or Ovar RediHaler. If none are formulary, or three if three are formulary, or two if two are formulary, or one if only one is formulary): Asmanex Twisthaler, a fluticasone inhaler (Armuity Ellipta), Pulmicort Flexhaler, or Ovar RediHaler. If none are formulary or three if three are formulary, or two if two are formulary, or one if only one is formulary): Asmanex Twisthaler, a fluticasone inhaler (Armuity Ellipta), Pulmicort Flexhaler, or Ovar RediHaler. If none are formulary, approve.  b. If the patient is < 6 years of age, approve if the patient has tried three formulary alternatives from the following list (if three are formulary, or two if two are formulary, or one if only one is formulary): Asmanex Twisthaler, a fluticasone inhaler (Armuity Ellipta), Pulmicort Flexhaler, or Ovar RediHaler. If none are formulary alternatives from the following list (if three are formulary or one if only one is formulary): Asmanex Twisthaler, a fluticasone inhaler (Armuity Ellipta), or Ovar RediHaler. If none are formular		Yes
nhalers	authorized generic)	powder	3. Approve if the patient is pregnant AND currently receiving the requested medication for asthma.	1 year	Yes

Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
		Direct the patient to fluticasone propionate HFA, if formulary. If fluticasone propionate HFA is non-formulary:  1. Approve if the patient has tried five formulary alternatives from the following list (if five are formulary, or three if three are formulary, or two if two are formulary, or one if only one is formulary: approve.  a. If the patient is < 12 years of age, approve if the patient has tried four formulary alternatives from the following list (if four are formulary, or three if three are formulary, or two if two are formulary, or one if only one is formulary: approve.  a. If the patient is < 12 years of age, approve if the patient has tried four formulary alternatives from the following list (if four are formulary, or three if three are formulary, or two if two are formulary, or one if only one is formulary: approve.  i. If the patient is < 12 years of age and has a low inspiratory flow rate and is unable to use a dry powder inhaler (DPI), approve if the patient has tried both formulary alternatives from the following list (if both are formulary, approve.  ii. If the patient is < 12 years of age, approve if usuable to use BOTH a DPI AND a breath-actuated metered-dose inhaler (MDI) [i.e., Qvar Redihaler], approve if the patient has tried Asmanex HFA, if formulary. If Asmanex HFA is non-formulary, approve.  b. If the patient is < 6 years of age, approve if the patient has tried three formulary: alternatives from the following list (if three are formulary, or two if two are formulary, or one if only one is formulary): a fluticasone propionate diskus [authorized generic of Flovent Diskus]), a mometasone inhaler (Asmanex HFA), or Ovar Redihaler, if none are formulary, approve.  ii. If the patient is < 6 years of age, approve if the patient has tried three formulary alternatives from the following list (if three are formulary, or two if two are formulary, or one if only one is formulary): a fluticasone inhaler (Asmanex HFA) in the patient is see of a ge, approve if the patient has tried three formulary, appro		
		4. Approve if the patient is pregnant AND currently receiving the requested medication for asthma.	1 year	Yes
			Direct the patient to fluticasone propionate HFA, if formulary. If fluticasone propionate HFA is non-formulary:  1. Approve if the patient has tried five formulary alternatives from the following list (if five are formulary, or five if there are formulary, or two if two are formulary, or one if only one is formulary): Alvesco, a studioscope inhaler (Annuty Ellipia, fluticasone propionate diskus jeuthorized generic of Flovent Diskus), a mometasone inhaler (Asmanex HFA, Asmanex Twisthaler), Pulmicort Flexhaler, or Ovar Redilhaler. If none are formulary, a fluticasone inhaler (Annuty Ellipia, fluticasone propionate diskus jeuthorized generic of Flovent Diskus)), a mometasone inhaler (Asmanex HFA, Asmanex Twisthaler, Asmanex Twisthaler, Asmanex HFA), Pulmicort Flexhaler, or Ovar Redilhaler. If none are formulary, approve.  a. If the patient is < 12 years of age, approve if the patient is described in the patient is set in the patient is < 12 years of age and has a low inspiratory flow rate and is unable to use a dry powder inhaler (DPI), approve if the patient has tried both formulary alternatives from the following list (if both are formulary, approve.  b. If the patient is < 6 years of age, approve if the patient has tried three formulary, approve.  b. If the patient is < 6 years of age, approve if the patient has tried three formulary alternatives from the following list (if the are formulary, approve).  b. If the patient is < 6 years of age, approve if the patient has tried three formulary alternatives from the following list (if the are formulary, or one if only one is formulary): a fluticasone proplemate diskus jeuthorized generic of Flovent Diskus), a mometasone inhaler (Asmanex HFA), or Ovar Redilhaler, if none are formulary, approve.  b. If the patient is < 12 years of age, approve if the patient has tried three formulary alternatives from the following list (if the are formulary, approve).  i. If the patient is < 6 years of age and has a low inspiratory flow rate and be alternative and the patient has tri	Direct the patient to fluicasone propionate HFA, if formulary, if fluicasone propionate HFA is non-formulary;  1. Approve if the patient to fluicasone propionate HFA, if formulary, if fluicasone propionate diskus [authorized generic of Flovent Diskus], a mometasone inhaler (Amane, HFA, Asmanex Twisthaler, Asmanex Twisthaler, Asmanex Twisthaler, Asmanex Twisthaler, Asmanex HFA, Pulmicort Flexhaler, or Ovar RediHaler. If none are formulary; approve.  a. If the patient is < 12 years of age, approve if the patient has tried four formulary alternatives from the following list (if four are formulary, or three if three are formulary, or two if two are formulary, or one if only one is formulary; approve.  i. If the patient is < 12 years of age, approve if the patient has tried four formulary, approve.  ii. If the patient is < 12 years of age, approve if the patient and is unable to use a dry powder inhaler (API), approve if the patient has tried both formulary alternatives from the following list (if both are formulary, or one if only one is formulary). If Asmanex HFA is non-formulary, approve.  iii. If the patient is < 12 years of age, approve if the patient has tried both formulary alternatives from the following list (if both are formulary) or one if only one is formulary. If Asmanex HFA is non-formulary, approve.  iii. If the patient is < 6 years of age, approve if the patient has tried both formulary alternatives from the following list (if three are formulary, or two if two are formulary, or one if only one is formulary): a fluidicasone propionate diskus [authorized generic of Flovent Diskus]), a mometasone inhaler (Asmanex Twishaler, Asmanex HFA), or Ovar RediHaler, if normulary, approve.  b. If the patient is < 6 years of age, approve if the patient has tried both formulary alternatives from the following list (if three are formulary, or one if only one is formulary). Care RediHaler, approve.  c. If the patient is < 6 years of age, and has a low inspiratory flow rate and is unable to use a dry powder inhaler, a

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
	Fluticasone propionate HFA (authorized generic of Flovent HFA)	fluticasone propionate HFA	1. Approve if the patient has tried five formulary alternatives from the following list (if five are formulary, or four if four are formulary, or three if three are formulary, or two if two are formulary, or one if only one is formulary): Alvesco, a fluticasone inhaler (Amuity Ellipta, fluticasone propionate diskus [authorized generic of Flovent Diskus]), a mometasone inhaler (Asmanex HFA, Asmanex Twisthaler, or Ovar Redithaler. If none are formulary, or three if three are formulary, or two if two are formulary, or own if only one is formulary; a fluticasone inhaler (Armuity Ellipta, fluticasone propionate diskus [authorized generic of Flovent Diskus]), a mometasone inhaler (Asmanex HFA), Pulmicont Flexhaler, or Ovar Redithaler. If none are formulary, approve.  i. If the patient is < 12 years of age and has a low inspiratory flow rate and is unable to use a dry powder inhaler (DPI), approve if the patient has tried both formulary alternatives from the following list (if both are formulary, approve.  ii. If the patient is < 12 years of age and is unable to use BOTH a DPI AND a breath-actuated metered-dose inhaler (MDI) [i.e., Ovar Redithaler], approve if the patient has tried Asmanex HFA, if formulary. If Asmanex HFA is non-formulary, approve.  b. If the patient is < 12 years of age, approve if the patient has tried three formulary alternatives from the following list (if three are formulary, or two if two are formulary, or one if only one is formulary): a fluticasone inhaler (Asmanex HFA, indicasone propionate diskus [authorized generic of Flovent Diskus]), a mometasone inhaler (Asmanex HFA, or Ovar Redithaler), approve if the patient has tried both formulary alternatives from the following list (if three are formulary, or two if two are formulary, approve.  i. If the patient is < 4 years of age, approve if the patient Diskus is a low inspiratory flow or at each is unable to use BOTH a DPI AND a breath-actuated MDI (i.e., Ovar Redithaler), approve if the patient has tried both formulary, approve if only one is	1 year	Yes
Respiratory - Corticosteroid Inhalers	Pulmicort Flexhaler		1. Approve if the patient has tried four formulary alternatives from the following list (if four are formulary, or three if three are formulary, or two if two are formulary, or one if only one is formulary): Alvesco, a fluticasone inhaler (Arnuity Ellipta, fluticasone propionate diskus [authorized generic of Flovent Diskus], fluticasone propionate HFA [authorized generic of Flovent HFA]), a mometasone inhaler (Asmanex HFA, Asmanex Twisthaler), or Qvar RediHaler. If none are formulary, approve.  a. If the patient is < 12 years of age, approve if the patient has tried three formulary alternatives from the following list (if three are formulary, or two if two are formulary, or one if only one is formulary): a fluticasone inhaler (Arnuity Ellipta, fluticasone propionate diskus [authorized generic of Flovent HFA]), a mometasone inhaler (Asmanex Twisthaler, Asmanex HFA), or Qvar RediHaler. If none are formulary, approve.  i. If the patient is < 12 years of age and is unable to coordinate breath and actuation with a conventional metered-dose inhaler, approve if two are formulary or one if only one is formulary): Asmanex Twisthaler, a fluticasone inhaler (Arnuity Ellipta, fluticasone propionate diskus [authorized generic of Flovent Diskus]), or Qvar RediHaler. If none are formulary approve.  2. If the patient is unable to coordinate breath and actuation with a conventional metered-dose inhaler, approve if the patient has tried three formulary alternatives from the following list (if three are formulary or one if only one is formulary): Asmanex Twisthaler, a fluticasone inhaler (Arnuity Ellipta, fluticasone propionate diskus [authorized generic of Flovent Diskus]), or Qvar RediHaler. If none are formulary or one if only one is formulary): Asmanex Twisthaler, a fluticasone propionate diskus [authorized generic of Flovent Diskus]), or Qvar RediHaler. If none are formulary or one if only one is formulary): Asmanex Twisthaler, a fluticasone propionate diskus [authorized generic of Flovent Diskus]), or Qvar RediHaler. If non	1 year	Yes MSB Exclusion
Respiratory - Corticosteroid Nebulized Solutions	Pulmicort	budesonide respules	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	*This criteria applies only to the NPF
Respiratory - Corticosteroid/Long- Acting Beta-Agonist Combination Inhalers	Advair Diskus	fluticasone propionate/salmeterol inhalation powder	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF

				Anneous	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	Medicaiton
Respiratory - Corticosteroid/Long- Acting Beta-Agonist	Dianu reante	fluticasone	1. Approve if the patient has tried four of the following (if four are formulary, or three if three are formulary, or two if two are formulary, or one if only one is formulary): fluticasone-salmeterol HFA (Advair HFA, authorized generic), fluticasone propionate/salmeterol inhalation powder (Advair Diskus, Wixela, generics), fluticasone-vilanterol powder inhalation (Breo Ellipta, authorized generic), Dulera, fluticasone propionate/salmeterol multidose dry powder inhalation (Breo Ellipta, authorized generic), Dulera, fluticasone propionate/salmeterol multidose dry powder inhalation powder (Advair Diskus, Wixela, generics), fluticasone propionate/salmeterol inhalation powder (Advair Diskus, Wixela, generics), fluticasone propionate/salmeterol inhalation powder (Advair Diskus, Wixela, generics), fluticasone propionate/salmeterol multidose DPI (authorized generic) or one if only one is formulary); fluticasone-vilanterol powder inhalation (Breo Ellipta, authorized generic), budesonide-formoterol aerosol (Symbicort, Breyna, generics), fluticasone-salmeterol HFA (Advair HFA, authorized generic), budesonide-formoterol aerosol (Symbicort, Breyna, generics), fluticasone-salmeterol multidose DPI (authorized generic) inhalation powder (Advair Diskus, Wixela, generics), fluticasone propionate/salmeterol multidose DPI (authorized generic) inhalation powder (Advair Diskus, Wixela, generics), fluticasone propionate/salmeterol multidose DPI (authorized generic of AirDuo RespiClick), or Dulera. If none are formulary, approve.  4. Patients < 18 years of age who are unable to coordinate breath and actuation with a metered-dose inhaler (MDI): approve if the patient has tried one of fluticasone-vilanterol powder inhalation (Breo Ellipta, authorized generic), fluticasone propionate/salmeterol inhalation (Advair Diskus, Wixela, generics) or fluticasone propionate/salmeterol multidose DPI (authorized generic of AirDuo RespiClick), if one is formulary. If none are formulary, approve.  5. Approve if the patient is pregnant AND currently r	Durauon	wedication
0 0	AirDuo RespiClick	inhalation powder	or generic count as one alternative.	1 year	Yes
Respiratory - Corticosteroid/Long- Acting Beta-Agonist Combination Inhalers	fluticasone propionate/salmeter ol HFA	fluticasone propionate/salmeterol HFA	Direct to Advair HFA (brand), if formulary. If Advair HFA (brand) is non-formulary:  1. Approve if the patient has tried four of the following, if (four are formulary, or three if three are formulary, or two if two are formulary, or one if only one is formulary): budesonide-formoterol aerosol (Symbicort, Breyna, generics), Dulera, fluticasone-vilanterol powder inhalation (Breo Ellipta, authorized generic), fluticasone propionate/salmeterol multidose dry powder inhaler (AirDuo RespiClick, authorized generics), or fluticasone propionate/salmeterol inhalation powder (Advair Diskus, Wixela, generics). If none are formulary, approve.  2. Patients < 18 years of age: approve if the patient has tried three of the following, if three are formulary (if three are formulary, or two if two are formulary, or one if only one is formulary): fluticasone-vilanterol powder inhalation (Breo Ellipta, authorized generic), budesonide-formoterol aerosol (Symbicort, Breyna, generics), Dulera, fluticasone propionate/salmeterol multidose dry powder inhaler (AirDuo RespiClick, authorized generics), or fluticasone propionate/salmeterol inhalation powder (Advair Diskus, Wixela, generics). If none are formulary, approve.  3. Patients with a low inspiratory flow rate who are unable to use a dry powder inhaler (DPI): approve if the patient has tried both 1) budesonide-formoterol (Symbicort, Breyna, generics) and 2) Dulera (if both are formulary or one if only one is formulary). If neither are formulary, approve.  4. Approve if the patient is pregnant AND currently receiving the requested medication for asthma.  8. Note: Fluticasone proprionate-salmetrol inhalation powder, Wixela, and Advair Diskus count as one alternative. Each product and its authorized generic or generic count as one alternative.  8. Budesonide-formoterol aerosol, Breyna, and Symbicort count as one alternative. Each product and its authorized generic or generic count as one alternative.	1 year	Yes
	fluticasone propionate/salmeter ol multidose dry powder inhaler	inhalation powder	1. Approve if the patient has tried four of the following (if four are formulary, or three if three are formulary, or two if two are formulary, or one if only one is formulary): fluticasone-salmeterol HFA (Advair HFA, authorized generic), fluticasone propionate/salmeterol inhalation powder (Advair Diskus, Wixela, generics), AirDuo RespiClick, fluticasone-vilanterol powder inhalation (Breo Ellipta, authorized generic), Dulera or budesonide-formoterol (Symbicort, Breyna, generics). If none are formulary, approve.  2. Patients who are unable to coordinate breath and actuation with a metered-dose inhaler (MDI): approve if the patient has tried two of the following (if two are formulary or one if only one is formulary): fluticasone propionate/salmeterol inhalation powder (Advair Diskus, Wixela, generics), AirDuo RespiClick, or fluticasone-vilanterol powder inhalation (Breo Ellipta, authorized generic), budesonide-formoterol aerosol, (Symbicort, Breyna, generics), fluticasone-salmeterol HFA (Advair HFA, authorized generic), fluticasone propionate/salmeterol inhalation powder (Advair Diskus, Wixela, generics), AirDuo RespiClick, or Dulera. If none are formulary, approve.  4. Patients < 18 years of age who are unable to coordinate breath and actuation with a metered-dose inhaler (MDI): approve if the patient has tried one of fluticasone-vilanterol powder inhalation (Breo Ellipta, authorized generic), fluticasone propionate/salmeterol inhalation powder (Advair Diskus, Wixela, generics), or AirDuo RespiClick, if formulary. If neither are formulary, approve.  5. Approve if the patient is pregnant AND currently receiving the requested medication for asthma.  Note: Fluticasone proprionate-salmetrol inhalation powder, Wixela, and Advair Diskus count as one alternative. Each product and its authorized generic or generic count as one alternative.	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Respiratory - Corticosteroid/Long-	Eluticasana	fluticasone furoate	Direct the patient to Breo Ellipta (brand), if formulary. If Breo Ellipta (brand) is non-formulary:  1. Approve if the patient has tried three of the following (if three are formulary, or two if two are formulary, or one if only one is formulary): fluticasone propionate/salmeterol inhalation powder (Advair Diskus, Wixela, generics), or Dulera. If none are formulary, approve.  2. Patients < 12 years of age: Approve if the patient has tried one of the following (if formulary): fluticasone propionate/salmeterol inhalation powder (Advair Diskus, Wixela, generics), Dulera, or budesonide-formoterol aerosol (Symbicort, Breyna, generics). If none are formulary, approve.  3. Patients ≤ 5 years of age: Approve if the patient has tried one of the following (if formulary): fluticasone propionate/salmeterol inhalation powder (Advair Diskus, Wixela, generics) or Dulera. If neither are formulary, approve.  4. Patients who are unable to coordinate breath and actuation with a metered-dose inhaler (MDI): approve if the patient has tried one of fluticasone propionate/salmeterol multidose dry powder inhaler (AirDuo RespiClick, authorized generics), if one is formulary, approve.  a. Patient < 12 years of age who are unable to coordinate breath and actuation with a metered-dose inhaler (MDI): Approve if the patient has tried fluticasone propionate/salmeterol inhalation powder (Advair Diskus, Wixela, generics). If fluticasone propionate/salmeterol inhalation powder (Advair Diskus, Wixela, generics) if the patient has tried fluticasone propionate/salmeterol inhalation powder (Advair Diskus, Wixela, generics) if obtained formulary or one if only one is formulary, approve.  5. Patients with COPD: Approve if the patient has tried both 1) fluticasone propionate/salmeterol inhalation powder (Advair Diskus, Wixela, generics) if formulary. If none are formulary, approve.  6. Patients with COPD: Approve if the patient has tried both 1) fluticasone propionate/salmeterol inhalation powder (Advair Diskus, Wixela, generics) if formulary. If flut		
Acting Beta-Agonist Combination Inhalers	Fluticasone- vilanterol	and vilanterol inhalation powder	Note: Fluticasone proprionate-salmetrol inhalation powder, Wixela, and Advair Diskus count as one alternative. Fluticasone propionate-salmeterol multidose dry powder inhaler and AirDuo RespiClick count as one alternative. Budesonide-formoterol aerosol, Breyna, and Symbicort count as one alternative. Each product and its authorized generic or generic count as one alternative.	1 year	Yes
Respiratory - Inhaled Phosphodiesterase (PDE)-3 and PDE-4 Inhibitor	Ohtuvayre	ensifentrine inhalation suspension	Chronic obstructive pulmonary disease (COPD) in a patient ≥ 18 years of age.  Approve if the patient has tried, and according to the prescriber, experienced inadequate efficacy OR significant intolerance with BOTH of the following products used concurrently: 1) a Long-Acting Muscarinic Antagonist (LAMA) product AND a Long-Acting Beta-Agonist (LABA) product.  LAMA/LABA Inhalers: Anoro Ellipta, Bevespi Aerosphere, Duaklir Pressair, Stiolto Respimat.  LAMA Inhalers: Incruse Ellipta, tiotropium inhaler (Spiriva HandiHaler, generics), Spiriva Respimat, Tudorza Pressair.  LABA Inhalers/Nebulized: Serevent Diskus, Striverdi Respimat, formoterol fumarate inhalation solution (Perforomist, generics).  ICS/LABA Inhalers: fluticasone-salmeterol HFA (Advair HFA, authorized generic), fluticasone-salmeterol respiclock (AirDuo RespiClick, authorized generic), AirDuo Digiplaler, or budesonide-formoterol (Symbicort, generics).	1 year	Yes
Respiratory - Long- Acting Muscarinic Antagonist (LAMA)	,	aclidinium bromide	Approve if the patient has tried, and according to the prescriber, has experienced inadequate efficacy OR significant intolerance with BOTH products from the following list, if formulary (or one if one is formulary): 1) Incruse Ellipta, and 2)		
Inhalers	Tudorza Pressair	inhalation powder	a tiotropium inhaler (tiotropium cap-inhaler [Spiriva HandiHaler, generics], or Spiriva Respimat). If neither are formulary, approve.	1 year	Yes
Respiratory Drugs - Other	Daliresp	roflumilast tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Rett Syndrome	D t	trofinetide oral		See PA	V
Agents  Rituximab-containing	Daybue	injection for	See standard Neurology – Daybue Prior Authorization Policy criteria. Note: No conditions of approval are recommended in the prior authorization policy.  1. Approve if the patient has tried one the following: Rituxan, Truxima, Ruxience, Riabni, but cannot continue to use the product.  2. Approve if, according to the prescriber, cannot use rituximab intravenous due to an inability to obtain IV access.	duration	Yes
Agents	Rituxan Hycela	subcutaneous use	3. If the patient has already been started on or has previously received therapy with Rituxan Hycela, approve.  Approve if the patient meets BOTH of the following (A and B):  A. The patient has tried three of the following, if three are formulary (or one or two of the following, if one or two is formulary): Truxima, Rituxan intravenous, Ruxience; AND  Note: If none are formulary, approve.	1 year	Yes
Rituximab-containing Agents	Riabni	rituximab-arrx intravenous injection	B. Patient cannot continue to use each formulary alternative due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.  Approve if the patient meets BOTH of the following (A and B):  A. The patient has tried three of the following, if three are formulary (or one or two of the following, if one or two is formulary): Truxima, Riabni, Ruxience; AND	1 year	Yes
Rituximab-containing Agents	Rituxan	rituximab intravenous injection	Note: If none are formulary, approve.  B. Patient cannot continue to use each formulary alternative due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would result in a significant allergy or serious adverse reaction.	1 year	Yes

					2025 NPF
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	Excluded Medicaiton
Therapy Class	Brand Name	Generic Name	Approve if the patient meets BOTH of the following (A and B):	Duration	Wedication
			A. The patient has tried three of the following, if three are formulary (or one or two of the following, if one or two is formulary): Rituxan intravenous, Riabni, Ruxience; AND		
			Note: If none are formulary, approve.		
Rituximab-containing	T	rituximab-abbs	B. Patient cannot continue to use each formulary alternative due to a formulation difference in the inactive ingredient(s) [e.g., differences in stabilizing agent, buffering agent, and/or surfactant] which, according to the prescriber, would	4	V
Agents	Truxima	intravenous injection	result in a significant allergy or serious adverse reaction.	1 year	Yes
			Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with one formulary product from three of the four groups below, if there is a formulary product in the group:		
			Group 1: An topical azelaic acid product (azelaic acid 15% gel [Finacea 15% gel, generics], Finacea 15% foam, Azelex 20% cream);		A
D		4 F0/	Group 2: A topical sodium sulfacetamide 10%/sulfur 5% product. (any generic sodium sulfacetamide10%/sulfur 5% product, Rosula);		
Rosacea Agents (Topical)	Zilxi	minocycline 1.5% topical foam	Group 3: A topical metronidazole product (metronidazole 0.75% or 1% [MetroGel, generics; MetroCream, generics; MetroLotion, generics, Noritate]); Group 4: a topical ivermectin product (generic ivermectin cream or Soolantra).	1 vear	Yes
(10000)		topiour rouni	State : a topical nominous product (general nominous accusing).	. , ,	1.00
			1. Direct the patient to a topical metronidazole product.		
Rosacea Agents		metronidazole cream	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use a generic topical metronidazole agent.		
(Topical)	Noritate	1%	Note: Examples of topical metronidazole products include metronidazole 0.75% cream (MetroCream, generics), metronidazole 0.75% or 1% gel (Metrogel, generics), metronidazole 0.75% lotion (MetroLotion, generics).	1 year	Yes
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		MSB Exclusion *This criteria
Sedative-Hypnotics			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
and Related Agents	Ambien	zolpidem tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
					MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Sedative-Hypnotics and Related Agents	Ambien CR	zolipidem extended- release tablets	<b>Criteria:</b> Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	applies only to the NPF
and Related Agents	Ambien on	release tablets	prescriber, would result in a significant anergy or serious adverse reaction to commentation required.	i yeai	MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Sedative-Hypnotics			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
and Related Agents	Lunesta	eszopiclone tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		MSB Exclusion
Sedative-Hypnotics			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		*This criteria applies only to
and Related Agents	Rozerem	ramelteon tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Sedative-Hypnotics	zolpidem 7.5 mg	zolpidem 7.5 mg	Approve if the patient has tried three of the following agents, if three are formulary (or two if two are formulary, or one if only one is formulary): zolpidem tablets (other strengths) [Ambien, Ambiren CR, generics], eszopiclone tablets		
and Related Agents	capsules (brand)	capsules	(Lunesta, generics), or zaleplon. If none are formulary, approve.	1 year	Yes
Selective Estrogen Receptor Modifiers			Approve if the patient has tried and varied state and residual state a		
and Antiestrogens	Osphena	ospemifene tablets	Approve if the patient has tried one vaginal estrogen product from the following list (if one is formulary): estradiol cream (Estrace cream, generics), Femring vaginal ring, Premarin vaginal cream, Estring vaginal ring, estradiol vaginal tablet (e.g., Yuvafem, Vagifem, generics), or Imvexxy If none are formulary, approve.	1 year	Yes
and / massa sgens	Сортона	oopenmene tablete	take (2-9) - take (1-9) - take	. , ,	1.00
Selective Serotonin					
Reuptake Inhibitor					
(SSRI) and a		trazodone	A Approve if the national host triangles to be triangles to be the national host transplant to be triangles to be triangles.		
Serotonin (5HT)2 Receptor Antagonist	Raldesy	hydrochloride oral solution	<ol> <li>Approve if the patient has tried trazodone tablets, if formulary. If trazodone tablets are non-formulary, approve.</li> <li>Approve if the patient has difficulty swallowing tablets or the patient cannot swallow tablets.</li> </ol>	1 year	Yes
Selective Serotonin	. tarassy	ooiuori	e. 7 pp. 010 in the patient rad difficulty distribution of the patient definet distribution.	. , ,	1.00
Reuptake Inhibitors	Viibryd 10/20 mg				
(SSRIs)	starter pack	vilazodone tablets	Approve if the patient is unable to use vilazodone tablets (which are not packaged in a starter pack).	1 year	Yes
Selective Serotonin Reuptake Inhibitors	oitalanram 20 mg		1. Direct to citalopram 10 mg or 20 mg tablets.		
(SSRIs)	citalopram 30 mg capsules	citalopram capsules	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use the citalopram 10 mg and/or 20 mg tablets.	1 vear	Yes
Selective Serotonin	Zercapli and		2.7 Pp. 5.7 II, according to the precession, the color of a dignimization of the first the patient to another of the officer to the officer t	. ,	
Reuptake Inhibitors	sertraline 150 mg,	sertraline 150 mg,	1. Direct the patient to sertraline 50 mg and/or 100 mg tablets.		
(SSRIs)	200 mg capsules	200 mg capsules	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use the sertraline 50 mg and/or 100 mg tablet.	1 year	Yes
Coloctive Ct			NOTE: A multisource Brand product is being requested. The nations should use the preferred bioequivalent generic product		MSB Exclusion
Selective Serotonin Reuptake Inhibitors			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		*This criteria applies only to
(SSRIs)	Celexa	citalopram tablets	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
			1	/	

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
Selective Serotonin Reuptake Inhibitors (SSRIs)	Lexapro	escitalopram oxalate tablets and oral solution	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Selective Serotonin Reuptake Inhibitors (SSRIs)	Prozac	fluoxetine HCI pulvules	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Reuptake Inhibitors	Viibryd (non- starter pack) 10 mg, 20 mg, 40 mg	vilazodone tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Selective Serotonin Reuptake Inhibitors (SSRIs)	Zoloft	sertraline HCl tablets and oral solution	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Serotonin and Norepinephrine Reuptake Inhibitors (SNRIs)	Cymbalta	duloxetine HCl capsules	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Serotonin and Norepinephrine Reuptake Inhibitors (SNRIs)	Effexor XR	venlafaxine HCl extended-release capsules	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Serotonin and Norepinephrine Reuptake Inhibitors (SNRIs)	Pristiq	dexvenlafaxine succinate tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Serotonin and Norepinephrine Reuptake Inhibitors (SNRIs)	Drizalma Sprinkle	duloxetine delayed- release capsules	<ol> <li>Approve if the patient has tried one product from the following list (if one is formulary): duloxetine capsules (Cymbalta, generics), Fetzima, desvenlafaxine succinate extended-release (ER) [Pristiq, generics], venlafaxine ER capsules (Effexor XR, generics), or venlafaxine extended-release tablets. If none are formularly, approve.</li> <li>NOTE: If patient has tried venlafaxine immediate-release tablets. If none are formularly, approve.</li> <li>Approve if the patient is unable to swallow, has difficulty swallowing, or requires administration via a nasogastric tube.</li> </ol>	1 year	Yes
Norepinephrine	Venlafaxine besylate ER 112.5 mg (formerly Venbysi	venlafaxine extended-	1. Approve if the patient has tried two products from the following list (if two are formulary; or one if one is formulary): desvenlafaxine succinate ER (Pristiq, generics), Fetzima, Drizalma Sprinkle, venlafaxine ER capsules (Effexor XR, generics), duloxetine capsules (Cymbalta, generics), or venlafaxine ER tablets. If none are formulary, approve.  NOTE: If patient has tried venlafaxine immediate-release, a trial of venlafaxine extended-release is not required.  2. Approve if the patient is currently taking or has taken venlafaxine besylate ER at any time in the past.	,	
	XR)	tablets	3. Suicidal ideation: approve.  1. Approve if the patient has tried one other single-entity albuterol inhaler.	1 year	Yes
Short-Acting Beta- Agonists (Inhaled)	ProAir Digihaler	albuterol sulfate inhalation powder	For example, albuterol HFA (ProAir HFA or generics) or albuterol HFA (Proventil HFA or generics).  Note: If there are no single-entity albuterol-containing formulary alternatives, approve.  2. Patients who are unable to coordinate breath and actuation with a metered-dose inhaler (MDI): approve if the patient has tried ProAir Respiclick, if formulary. If ProAir Respiclick is non-formulary, approve.	1 year	Yes
Short-Acting Beta- Agonists (Inhaled)	ProAir Respiclick	albuterol sulfate inhalation powder	<ol> <li>Approve if the patient has tried one other single-entity albuterol inhaler.</li> <li>For example, albuterol HFA (ProAir HFA or generics) or albuterol HFA (Proventil HFA or generics).</li> <li>Note: If there are no single-entity albuterol-containing formulary alternatives, approve.</li> <li>Patients who are unable to coordinate breath and actuation with a metered-dose inhaler (MDI): approve if the patient has tried ProAir Digihaler, if formulary. If ProAir Digihaler is non-formulary, approve.</li> </ol>	1 year	Yes
Short-Acting Beta- Agonists (Inhaled)	Ventolin HFA and authorized generic	albuterol sulfate inhalation aerosol	Approve if the patient has tried one other single-entity albuterol inhaler.  For example, albuterol HFA (ProAir HFA or generics) or albuterol HFA (Proventil HFA or generics).  Note: If there are no single-entity albuterol-containing formulary alternatives, approve.  Approve if the patient has tried one single-entity albuterol inhaler.	1 year	Yes
	Xopenex HFA and levalbuterol HFA	levalbuterol inhalation aerosol	Approve it the patient has tried one single-entity abouterol inhaler.  For example, albuterol HFA (ProAir HFA or generics) or albuterol HFA (Proventil HFA or generics).  Note: If there are no single-entity albuterol-containing formulary alternatives, approve.  1. Approve is the patient has tried Droxia, if formulary. If Droxia is non-formulary, approve.	1 year	Yes
Agents – Hydroxyurea	Siklos	hydroxyurea tablets	<ol> <li>Approve is the patient has the Droxia, in formulary. In Droxia is non-formulary, approve.</li> <li>If the patient requires Siklos 100 mg or 1,000 mg tablets to achieve a dosage that cannot be achieved with the available strengths of Droxia, approve.</li> <li>If the patient cannot swallow Droxia capsules or has difficulty swallowing Droxia capsules, approve.</li> </ol>	1 year	Yes

					0005 ND5
				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
Sickle Cell Disease			1. Approve if the patient has tried one of Droxia or Siklos, if formulary. If neither are formulary, approve.		
Agents – Hydroxyurea		hydroxyurea oral	2. Patient < 2 years of age, approve.	4	V
Agents	Xromi	solution	3. If the patient is unable to swallow tablets or capsules OR has difficulty swallowing tablets or capsules, approve.	1 year	Yes
			Approve if the patient meets one of the following (1 or 2):		
			1. Patient meets BOTH of the following (A and B):  A. Patient has tried at least two phosphate binders; AND		
			Note: Examples of phosphate binders include: sevelamer, lanthanum, ferric citrate, and sucroferric oxyhydroxide, calcium carbonate, and calcium acetate.		
			B. Patient had an inadequate response and/or intolerance to at least two phosphate binders; OR		
			2. Patient meets one of the following (A or B):		
			A. Patient has a contraindication to at least two phosphate binders; OR		
			Note: Contraindication to phosphate binders includes bowel obstruction, iron overload, or hypercalcemia.		
			B. Patient meets BOTH of the following (i and ii):  i. Patient has inadequate response and/or intolerance to at least one phosphate binder; AND		
Sodium Hydrogen			ii. Patient has a contraindications to at least one phosphate binder.		
Exchanger 3 Inhibitor	Xphozah	tenapanor tablets	Note: Contraindication to phosphate binders include bowel obstruction, iron overload, or hypercalcemia.	1 year	Yes
J	1		1. Acromegaly: Approve if the patient has tried one of Sandostatin LAR Depot, Somatuline Depot, or lanreotide subcutaneous injection, if one is formulary. If none are formulary, approve.		
			2. Cushing's Disease. Approve if the patient has tried Signifor (not LAR). If Signifor (not LAR) is non-formulary, approve.		
		pasireotide IM	3. Endogenous Cushing's Syndrome. Note: This includes patients awaiting surgery and patients awaiting therapeutic response after pituitary radiotherapy. Approve if patient has tried Signifor (not LAR), if formulary. If Signifor (not LAR)	)	
Somatostatin Analogs	Signifor LAR	injection	is non-formulary, approve.	1 year	Yes
			1. <u>Acromegally: neuroendocrine tumors: pheochromoctoma/paraganglioma</u> . Approve if the patient has tried Somatuline Depot, if formulary.		
			If Somatuline Depot is non-formulary, approve if the patient meets (A or B):		
			A. <u>Acromegaly: pheochromoctoma/paraganglioma</u> : Approve if the patient has tried Sandostatin LAR Depot, if formulary. If Sandostatin LAR Depot is non-formulary, approve.  B. Patients with neuroendocrine tumors: Approve if the patient meets the following (i or ii):		
			Note: This includes (but is not limited to) carcinoid tumors, vasoactive intestinal peptide tumors (VIPomas), glucagonomas, gastrinomas, insulinomas.		
	lanreotide	lanreotide	i. Patient has tried Sandostatin LAR Depot, if formulary. If Sandostatin LAR Depot is non-formulary, approve; OR		
	subcutaneous	subcutaneous	ii. Patient has already been started on therapy with lanreotide subcutaneous injection.		
Somatostatin Analogs	injection [Cipla]	injection	2. Carcinoid syndrome: Approve if the patient has tried Somatuline Depot, if formulary. If Somatuline Depot is non-formulary, approve.	1 year	Yes
			1. Acromegaly: Approve if the patient has tried one of Somatuline Depot or lanreotide subcutaneous injection, if formulary. If neither are formulary, approve.		
			2. Patient with neuroendocrine tumors: approve if the patient meets the following (A or B):		
			Note: This includes (but is not limited to) carcinoid tumors, vasoactive intestinal peptide tumors (VIPomas), glucagonomas, gastrinomas, insulinomas.  A. Patient has tried one of Somatuline Depot or lanreotide subcutaneous injection, if formulary, approve; OR		
			B. Patient has already been started on therapy with Sandostatin LAR.		
			3. Patient with pheochromocytoma/paraganglioma: approve if the patient meets the following (A or B):		
			A. Patient has tried Somatuline Depot, if formulary. If Somatuline Depot is non-formulary, approve; OR		
	Sandostatin LAR	octreotide injectable	B. Patient has already been started on therapy with Sandostatin LAR.		
Somatostatin Analogs	Depot	suspension	4. Patient with diarrhea associated with chemotherapy; enterocutaneous fistula; meningioma; Merkel cell carcinoma; pancreatic fistula; thymoma/thymic carcinoma: approve.	1 year	Yes
04			1. Approve if the patient has tried one formulary alternative from the following list: Imvexxy, Femring vaginal ring, Premarin Cream, Estring vaginal ring, estradiol 0.01% cream (Estrace cream, generics), or estradiol vaginal tablet (e.g.,		
Steroid Products (Vaginal)	Intrarosa	prasterone vaginal inserts	Yuvafem, Vagifem, generics). If none are formulary, approve.  2. Approve if, according to the prescriber, the patient is at an increased risk of endometrial cancer, stroke, or deep vein thrombosis (DVT).	1 year	Yes
(vagiliai)	IIIIIaiUSa	testosterone	2. Approve it, according to the prescriber, the patient is at an increased risk of endomental cancer, stroke, or deep vein thrombosis (DVT).	i yeai	165
		cypionate for	Approve if the patient has tried one of the following injectable testosterone products, if one is formulary: testosterone cypionate injection (Depo-Testosterone, generics) or testosterone enanthate injection (generics). If none are		
Testosterone		intramuscular	formulary, approve.		
Products (Injectable)	Azmiro	injection	Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.	1 year	Yes
		testosterone	Approve if the patient has tried one of the following injectable testosterone products, if one is formulary: testosterone enanthate injection [generics], testosterone cypionate injection [Depo-Testosterone, generics], Azmiro, or Xyosted. If		
Testosterone	A	undecanoate for	none are formulary, approve.	4	
Products (Injectable)	Aveed	intramuscular use testosterone	Note: If the patient tried the brand version of a generic equivalent product, then this trial would count towards the requirement.	1 year	Yes
Testosterone	Kyzatrex and	undecanoate			
Products (Oral)	Undecatrex	capsules	Approve if the patient has tried both of Jatenzo and Tlando capsules, if formulary (or one if one is formulary). If neither are non-formulary, approve if the patient has tried two forms of topical testosterone (e.g., gel, solution, patches).	1 year	Yes
(=:=:)			1. Approve if the patient meets BOTH of the following, if formulary (or one if one is formulary) [a and b]:		
			a. Patient has tried Jatenzo, if formulary; AND		
		testosterone	b. Patient has tried one of Kyzatrex or Undecatrex, if formulary.		
Testosterone	T11.	undecanoate oral	Note: Kyzatrex and Undecatrex count as one alternative.		V
Products (Oral)	Tlando	capsules	2. If neither are formulary, approve if the patient has tried two forms of topical testosterone (e.g., gel, solution, patches).	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
					MSB Exclusion
		testosterone 1% gel	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Testosterone		packets and pump,	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Products (Topical)	Androgel	1.62% (2021)	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
					MSB Exclusion
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Testosterone	T		Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the	4	applies only to
\	Testim	testosterone gel	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Testosterone	Matanta		According to the standard stan	4	V
Products (Topical)	Natesto	testosterone nasal gel	Approve if the patient has tried three other topical testosterone products (e.g., Androgel 1% or generics, Axiron [generics only], Androgel 1.62% or generics, Fortesta or generics, Testim or generics, Vogelxo or generics.)	1 year	Yes
Tetracycline- Derivatives - Oral	Oracea and doxycycline 40 mg capsules (authorized generic of Oracea)	doxycycline 40 mg capsules	Inflammatory Rosacea.  Approve if the patient meets both of the following (A and B):  A. Patient has tried two of the following: 1) a topical metronidazole-containing product, 2) a topical azelaic acid-containing product or 3) topical ivermectin; AND  B. Patient meets one of the following (i or ii):  i. Patient has tried, and according to the prescriber, has experienced inadequate efficacy with one other generic, oral doxycycline product after a 4 week duration with the product; OR  ii. Patient has tried, and according to the prescriber, has experienced a significant intolerance with one other generic, oral doxycycline product.	9 months	Yes - brand only
Tetracycline- Derivatives -Oral	<u> </u>	minocycline extended-	Inflammatory Rosacea.  Approve if the patient meets both of the following (A and B):  A. Patient has tried two of the following: 1) a topical metronidazole-containing product, 2) a topical azelaic acid-containing product or 3) topical ivermectin; AND  B. Patient meets one of the following (i or ii):		
Agents for Rosacea	Emrosi	release capsules	ii. Patient has tried, and according to the prescriber, has experienced a significant intolerance with one other generic, oral minocycline product.	1 year	Yes
Thiazide Diuretics		hydrochlorothiazide oral suspension chlorthalidone 15 mg	<ol> <li>Approve if the patient tried and cannot take hydrochlorothiazide tablets or capsules. If hydrochlorothiazide tablets and capsules are non-formulary, approve.</li> <li>Approve if the patient is unable to swallow tablets or capsules or has difficulty swallowing tablets or capsules.</li> <li>Approve if the prescribed dose cannot be obtained with whole tablets or capsules of any hydrochlorothiazide strengths.</li> <li>Direct the patient to generic chlorthalidone tablets. Generics available as 25 mg, 50 mg.</li> </ol>	1 year	Yes
Thiazide-like Diuretics	Thalitone 15 mg	tablets	2. Approve if the prescribed dose cannot be obtained with the 25 mg and/or 50 mg strength tablets.	1 year	Yes
			Mulpleta is being used pre-procedure and the patient has thrombocytopenia and chronic liver disease.	1	
Thrombocytopenia			1. Approve if the patient has tried Doptelet, if formulary. If Doptelet is non-formulary, approve.		
	Mulpleta	lusutrombopag tablets		1 month	Yes
Thrombocytopenia		eltrombopag choline	Immune Thrombocytopenia.  1. Approve if the patient has tried one of Promacta or Nplate, if formulary. If neither are formulary, approve.  2. Approve if the patient has already been started on therapy with Alvaiz.  Aplastic Anemia; Thrombocytopenia in a Patient with Chronic Hepatitis C; Thrombocytopenia in a Patient with Myelodysplastic Syndrome; Thrombocytopenia in a Patient Post-Allogeneic Transplantation; Thrombocytopenia due to Immune Checkpoint Inhibitor Therapy.  1. Approve if the patient has tried Promacta, if formulary. If Promacta is non-formulary, approve.  2. Approve if the patient has already been started on therapy with Alvaiz.  Note: Examples of checkpoint inhibitors are Keytruda (pembrolizumab intravenous infusion), Opdivo (nivolumab intravenous infusion), Yervoy (ipilimumab intravenous infusion), Tecentriq (atezolizumab intravenous infusion), Bavencio		
	Alvaiz	tablets	(avelumab intravenous infusion), Imfinzi (durvalumab intravenous infusion), and Libtavo (cemiplimab-rwlc intravenous infusion), Televoy (priminimab intravenous infusion), Imfinzi (durvalumab intravenous infusion), and Libtavo (cemiplimab-rwlc intravenous infusion).	1 year	Yes
Thyroid Supplements		liothyronine sodium tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Thyroid Supplements	,	levothyroxine tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Th	Tirosint and	levothyroxine	Approve if the patient has tried five formulary levothyroxine products from the following list (if five are formulary or four are formulary or three if three are formulary or two if two are formulary or one if one is formulary): levothyroxine		v
Thyroid Supplements	authorized generic	capsules	[Synthroid, generics), Levoxyl (generics), Unithroid (generics), Evoxyl (generics), Unithroid (generics), or Tirosint oral solution [documentation required]. If none are formulary, approve.	1 year	Yes
			1. Approve if the patient has tried five formulary levothyroxine products from the following list (if five are formulary or four if four are formulary or three if three are formulary or two if two are formulary or one if one is formulary):		
		lavada mario e e elle	levothyroxine (Synthroid, generics), Levoxyl (generics), Unithroid (generics), Euthyrox (generics), or Tirosint capsules [documentation required]. If none are formularly, approve.		
Thyroid Supplements	Thyquidity	levothyroxine sodium oral solution	2. If the patient cannot swallow or has difficulty swallowing tablets or capsules [documentation required], approve if the patient has tried both Tirosint oral solution and Ermeza oral solution, if formulary (or one if one is formulary). If neither are formulary, approve.	1 vear	Yes
myroid Supplements	rriyquiuity	orar Solution	niciuiei aic iorinidaty, approve.	i yeai	162

					2025 NPF
				Approval	Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria  1. Approve if the patient has tried five formulary levothyroxine products from the following list (if five are formulary or four if four are formulary or three if three are formulary or two if two are formulary or one if one is formulary):	Duration	Medicaiton
			levothyroxine (Synthroid, generics), Levoxyl (generics), Euthyrox (generics), or Tirosint capsules [documentation required]. If none are formulary, approve.		A
		levothyroxine oral	2. If the patient cannot swallow or has difficulty swallowing tablets or capsules [documentation required], approve if the patient has tried both Thyquidity oral solution and Ermeza oral solution, if formulary (or one if one is formulary). If		A .
Thyroid Supplements	Tirosint-SOL	solution	neither are formulary, approve.	1 year	Yes
Thyroid Supplements			1. Approve if the patient has tried one levothyroxine product (e.g., levothyroxine, Synthroid, Levoxyl) AND one other desiccated thyroid product (e.g., Armour Thyroid, NP thyroid).		
Desiccated Thyroid			2. Patient currently receiving Adthyza: Approve if the patient has tried one other desiccated thyroid product (e.g., Armour Thyroid, NP thyroid).		A .
Supplements	Adthyza	thyroid tablets	Note: Some desiccated thyroid products are currently not available, such as Nature thyroid, WP thyroid, Westhroid, and Thyroid tablet, but a previous trial of these would count as a trial of a desiccated thyroid product.	1 year	Yes
			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		MSB Exclusion *This criteria
Topical Agents for			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
Atopic Dermatitis	Elidel	pimecrolimus cream	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
			NOTE A sufficient Development of The office development of the original development of		MSB Exclusion
Topical agents for Condyloma	Condylox 0.5%		NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		*This criteria applies only to
acuminatum	topical gel	podofilox 0.5% gel	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
	·				
Topical Corticosteroid	l <del>-</del>	halobetasol	Approve if the patient has tried, and according to the prescriber has experienced inadequate efficacy OR significant intolerance with four generic prescription-strength topical corticosteroid products.		A
containing Agents – Halobetasol Agents	Ultravate Lotion	propionate lotion 0.05%	Note: Examples of prescription-strength topical corticosteroids products include: halobetasol propionate, betamethasone dipropinate, clobetasol propionate, diflorasone diacetate.  NOTE: The products must be chemically unique.	1 year	Yes
riaiozotaco: / tgonto	Citarate Letteri	0.00%	The state of the s	. you.	MSB Exclusion
Topical			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Dermatological Drugs	1.1.1	11.1	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the	4	applies only to
- Miscellaneous	Lidoderm	lidocaine 5% patch	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF MSB Exclusion
Topical			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		*This criteria
Dermatological Drugs		tazarotene 0.1%	Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
- Miscellaneous	Tazorac 0.1% cream	cream	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Topical					A
Dermatological Drugs		sinecatechins	1. Approve if the patient has tried both 1) podofilox topical solution or gel (e.g., Condylox, generics) AND 2) imiquimod 5% cream (Aldara, generics), if formulary. If none are formulary, approve.		A
- Miscellaneous	Veregen	ointment 15%	2. For perianal warts, approve if the patient has tried both 1) podofilox topical gel (e.g., Condylox gel, generics) AND 2) imiquimod 5% cream (Aldara, generics), if formulary. If neither are formulary approve.	1 year	Yes
Taminal		sodium sulfacetamide			A
Topical Dermatological Drugs	Clenia Plus and	9%- sulfur 4.25%	1. Direct the patient to a topical product containing sodium sulfacetamide-sulfur (e.g., generic sodium sulfacetamide-sulfur 9.8%-4.8% topical cleanser, generic sodium sulfacetamide-sulfur 8%-4% topical suspension).		A
- Miscellaneous	authorized generic	suspension	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use a generic topical product containing sodium sulfacetamide/sulfur.	1 year	Yes
Topical Dermatological Drugs	sulfacetamide-sulfur	culfocatomido culfur 9	3-1. Direct the patient to a topical product containing sodium sulfacetamide-sulfur (e.g., generic sodium sulfacetamide-sulfur 8%-4% topical suspension).		4
- Miscellaneous	8-4% cleanser	4% cleanser	2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use a generic topical product containing sodium sulfacetamide-sulfur.	1 year	Yes
				,	
Topical		sodium sulfacetamide			A
Dermatological Drugs - Miscellaneous	Zma Clear	9% and sulfur 4.5% suspension	1. Direct the patient to a topical product containing sodium sulfacetamide-sulfur (e.g., generic sodium sulfacetamide-sulfur 9.8%-4.8% topical cleanser, generic sodium sulfacetamide-sulfur 8%-4% topical suspension).  2. Approve if, according to the prescriber, there is a significant clinical concern such that the patient is unable to use a generic topical product containing sodium sulfacetamide-sulfur.	1 year	Yes
- Miscellarieous	Pliaglis and	Suspension	2. Approve it, according to the prescriber, there is a significant clinical concern such that the patient is unable to use a generic topical product containing sodium sunacetamide-sund.	i yeai	163
Topical	lidocaine 7% and				
Dermatological Drugs		lidocaine 7% and	Approve if the patient has tried and cannot use two of the following, if two are formulary (or one if only one is formulary): lidocaine and prilocaine cream (generics), lidocaine cream (generics, multiple strengths), Livixil Pak, DermacinRx		
- Miscellaneous	(brand)	tetracaine 7% cream	Prizopak. If none are formulary, approve.	1 year	Yes
			Approve if the patient has tried and, according to the prescriber, has experienced inadequate efficacy OR significant intolerance with five single-entity corticosteroid topical agents AND one prescription topical anti-infective agent.		
Topical		hydrocortisone 2%/	Note: Examples of topical corticosteroids include: hydrocortisone cream/lotion/ointment [multiple brand and generic products], betamethasone cream/ointment/lotion [Diprolene, generics], clobetasol cream/gel/lotion [Temovate, Clobex,		
Dermatological Drugs		iodoquinol 1%/ aloe	generics], fluocinolone ointment/cream [Synalar, generics], fluocinonide cream/ointment/gel [generics], mometasone cream/lotion/ointment [Elocon, generics], triamcinolone cream/ointment/lotion [generics].	1 year	Voc
- Miscellaneous	Alcortin A	1% gel	Note: Examples of prescription topical anti-infectives include: mupirocin 2% cream [Bactroban, generics], mupirocin 2% ointment [Bactroban, generics], Centany ointment, Centany AT ointment, Altabax ointment).	1 year	Yes

Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Approval Duration	2025 NPF Excluded Medicaiton
Topical Dermatological Drugs - Tazarotene	Tazorac gel	tazarotene gel 0.05% and 0.1%	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
- razaroterie	Tazorac 0.05%	and 0.176	prescriber, would result find significant differgy or serious deverse reaction (documentation required).	i yeai	ule INFF
Topical	cream and				
Dermatological Drugs		tazarotene cream	If requesting brand Tazorac 0.05% cream: Approve if the patient has tried generic tazarotene 0.05% cream, if formulary. If generic tazarotene 0.05% cream is non-formulary or generic tazarotene is being requested, approve if the		
- Tazarotene	0.05% cream	0.05%	patient has tried one of 1) tazarotene 0.1% cream (Tazorac 0.1% cream, generics) or 2) tazarotene gel (Tazorac gel, generics), if one is formulary. If neither are formulary, approve.	1 year	Yes
	Vusion and				
Topical Diaper	miconazole-zinc	miconazole-zinc oxide	Approve if the patient has tried one topical antifungal agent.		
Dermatitis Agents	oxide-petroleum	petroleum ointment	Note: Examples include: miconazole, clotrimazole, ketoconazole, nystatin.	1 year	Yes
	doxepin, prudoxin		Approve if the patient has tried TWO topical corticosteroids.		
Topical Doxepin	5% cream	doxepin 5% cream	Note: Examples of topical corticosteroids include alclometasone cream or ointment, desonide cream or ointment, fluocinolone cream or ointment, hydrocortisone cream or ointment.	1 year	Yes
Topical Products -		fluocinolone acetonide 0.01%/hydroquinone 4%/tretinoin 0.05%	Direct the patient to the separate entities:		
Miscellaneous	Tri-luma cream	cream	fluocinolone 0.01% cream- hydroquinone 4% cream- tretinoin 0.05% cream.	N/A	Yes
Urinary Tract Analgesic	Pyridium	tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
			1. Approve if the patient has tried generic ursodiol capsules or tablets.		
Ursodiol Products	Reltone	mg, 400 mg	2. Approve, if according to the prescriber, the patient is unable to achieve the appropriate dosage requirement with ursodiol capsules.	1 year	Yes
Vertigo Agents	Antivert 50 mg tablet and authorized generic meclizine 50 mg	meclizine 50 mg tablet	Patient meets both of the following (i and ii): i. Patient has tried generic 25 mg tablets; AND ii. Patient cannot take generic 25 mg tablets due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction.	1 year	Yes
Vesicular Monoamine			NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.		MSB Exclusion *This criteria
Transporter Type 2			Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		applies only to
(VMAT2) Inhibitors	Xenazine	tetrabenazine tablets	prescriber, would result in a significant allergy or serious adverse reaction (documentation required).	1 year	the NPF
(******=) *********			1. Approve if the patient has tried calcipotriene solution, if formulary. If calcipotriene solution is non-formulary, approve.	, , =	
Vitamin D Analogs	Sorilux and		2. Approve if the patient has tried calcipotriene cream or ointment.		
(Topical)	authorized generic	calcipotriene foam	3. If the patient is using the requested medication for plaque psoriasis and is between the ages ≥ 4 and < 18 years of age, approve.	1 year	Yes
Vitamin D Analogs		calcipotriene 0.005%/betamethaso ne dipropionate 0.064% topical	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the		MSB Exclusion *This criteria applies only to
(Topical)	Taclonex	suspension	prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	the NPF
Wakefulness Agents	Nuvigil	armodafinil tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF
Wakefulness Agents	Provigil	modafinil tablets	NOTE: A multisource Brand product is being requested. The patient should use the preferred bioequivalent generic product.  Criteria: Approve if the Brand product is being requested due to a formulation difference in the inactive ingredient(s) [e.g., difference in dyes, fillers, preservatives] between the Brand and the bioequivalent generic product which, per the prescriber, would result in a significant allergy or serious adverse reaction [documentation required].	1 year	MSB Exclusion *This criteria applies only to the NPF

				Approval	2025 NPF Excluded
Therapy Class	Brand Name	Generic Name	Commercial FE Criteria	Duration	Medicaiton
			Weight loss in a patient ≥ 18 years of age.  Approve if the patient meets the following (A and B):  A. At baseline, the patient has or had a body mass index (BMI) ≥ 30 kg/m2; or a BMI ≥ 27 kg/m2 and at least one of the following weight-related comorbidities: hypertension, type 2 diabetes, dyslipidemia, obstructive sleep apnea, cardiovascular disease, knee osteoarthritis, asthma, chronic obstructive pulmonary disease, metabolic-dysfunction associated steatotic liver disease/non-alcoholic fatty liver disease, polycystic ovarian syndrome, or coronary artery disease; AND  Note: This refers to baseline prior to any glucagon-like peptide-1 (GLP-1) agonist (e.g., Saxenda, Wegovy) or GLP-1/glucose-dependent insulinotropic polypeptide (GIP) receptor agonist (e.g., Zepbound).  B. Patient has tried one of Wegovy or Zepbound, if formulary. If neither are formulary, approve.		
			Weight loss in a patient is ≥ 12 years of age and < 18 years of age.  Approve if the patient meets the following (A <u>and</u> B):  A. At baseline, the patient has or had a BMI ≥ 95th percentile for age and sex; AND		
ight Loss – GLP-1 eptor agonists		liraglutide [rDNA] injeciton	Note: This refers to baseline prior to any glucagon-like peptide-1 (GLP-1) agonist (e.g., Saxenda, Wegovy) or GLP-1/glucose-dependent insulinotropic polypeptide (GIP) receptor agonist (e.g., Zepbound).  B. Patient has tried Wegovy, if formulary. If Wegovy is non-formulary, approve.	1 year	Yes