Clinical Criteria, Step Therapy, and Quantity Limits for TennCare Preferred Drug List (PDL)

June 1, 2025

Note: All agents must be prescribed by a provider with a Tennessee Medicaid Provider ID.

	ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.						
Medication	PDL		Qty. Limits	PA Form			
		Agents for Dependency					
Lucemyra®	Р	 Initial Criteria: Must be ≥ 18 years of age; AND Patient is not pregnant or breast feeding; AND Attestation that if patient is at risk for QT interval prolongation (congestive heart failure, bradyarrhythmia, hepatic impairment, renal impairment, or taking other medicinal products that lead to QT prolongation), baseline electrocardiogram (ECG) has been performed; AND Patient has tried and failed, had a contraindication to, or experienced an adverse reaction/intolerance to clonidine; AND Prescriber to provide verbal attestation of a comprehensive treatment plan between provider and patient; AND In the case of opioid use disorder (OUD), provide verbal attestation that patient:	16/day	General PA Form			
Vivitrol® injection	Р		1 vial per 28 days				
lofexidine	NP	See Lucemyra® prior authorization criteria; AND • Trial and failure, contraindication, or intolerance to preferred Lucemyra®	16/day				



		ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicat	ed.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Buprenorphine and Buprenorphine/Naloxone		
		Buprenorphine Enhanced Supportive Medication-Assisted Recovery and Treatment (BESMART) Network Provider	onl <u>y</u> :	
buprenorphine/ naloxone tablets	Р		8/2 mg: 3/day; 2/0.5 mg: 3/day ^	
buprenorphine/ naloxone film	Р		12/3 mg: 2/day; 8/2 mg: 3/day; 4/1 mg: 2/day; 2/0.5 mg: 3/day ^	
buprenorphine	NP	 Criteria: Diagnosis of opiate addiction; AND Prescriber is enrolled and in good standing in the BESMART program; AND Buprenorphine will not be approved for treatment of depression or pain; AND ONE of the following: Patients is actively pregnant (must provide estimated due date) Patient is actively breastfeeding (must provide delivery date) Request is for a two-day induction for patients transitioning off of Methadone Patient is unable to take naloxone containing products due to a contraindication, drug to drug interaction, or history of toxic side effects that caused immediate or long-term damage (DOCUMENTATION REQUIRED). Note: Mild rash, itching, and GI intolerance are not accepted as intolerance to naloxone. PA Approval durations: Pregnancy: 3 months past due date; Breastfeeding: 6 months (maximum 4 approvals); Contraindication to Naloxone: Initial Authorization 6-months, Reauthorization 12 months Note: The PA for buprenorphine monotherapy sublingual tablets in pregnant or breastfeeding patients diagnosed with opioid use disorder can be bypassed using Professional Pharmacy Service (PPS) Codes. 	8 mg: 3/day; 2 mg: 3/day ^	Buprenorphine Products PA Form
Suboxone® film	NP	Criteria: (6-month duration for initial request; 12-month duration for reauthorization) Diagnosis of opiate addiction; AND Prescriber is enrolled and in good standing in the BESMART program; AND Buprenorphine will not be approved for treatment of depression or pain; AND Patient has a documented allergy to inactive ingredient in preferred product that is not in requested product (DOCUMENTATION REQUIRED)	12/3 mg: 2/day; 8/2 mg: 3/day; 4/1 mg: 2/day; 2/0.5 mg: 3/day^	
Zubsolv®	NP	See Suboxone film prior authorization criteria	11.4/2.9 mg: 1/day 8.6/2.1 mg: 2/day; 5.7/1.4 mg: 3/day; 2.9/0.71 mg: 2/day; 1.4/0.36 mg: 3/day; 0.7/0.18 mg: 3/day	



ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
^ Requests for 4/	day w	uantities may be approved as medically necessary. ill only be approved if dose is being titrated or patient's condition is too unstable to attempt to change to a higher strength. er (NP) , Physician Assistant (PA) and other mid-level prescribers in the BESMART network are limited to MDD 16mg/day per TN s	state law.		
		All other TennCare Providers:			
buprenorphine/ naloxone tablets	Р	 Criteria: (6-month duration for initial request; 12-month duration for reauthorization) Diagnosis of opiate addiction; AND Prescriber is NOT a nurse practitioner or physician assistant; AND Physician attests they have reviewed the Tennessee Controlled Substances Database for this patient on the date of the prior authorization request to ensure that concomitant narcotic or benzodiazepine use is not occurring. Additional Information: Buprenorphine will not be approved for treatment of depression or pain. Buprenorphine will not be approved for recipients whose medication history indicates use of concomitant narcotics or benzodiazepines without a clinically valid reason and drug tapering plan. Quantity limit is as a single daily dose. Twice daily dosing may be approved as clinically necessary. Physicians will be asked to provide an anticipated treatment plan for the patient (including anticipated dosing for induction & maintenance phases, anticipated frequency of office visits, & anticipated plan for psychosocial counseling). The "Here to Help" program as an exclusive provider of counseling will not be accepted. Prior Authorizations will be assigned to the prescribing physician. Requests for buprenorphine from a different physician will require a new prior authorization request and documentation that the previous prescribing physician has communicated transfer of care. 	8/2 mg: 2/day for 6 months then 1/day*; 2/0.5 mg: 3/day* ^		
buprenorphine	NP	Criteria: See buprenorphine/naloxone tab prior authorization criteria; AND ONE of the following: Patients is actively pregnant (must provide estimated due date) Patient is actively breastfeeding (must provide delivery date) Request is for a two-day induction for patients transitioning off of Methadone Patient is unable to take naloxone containing products due to a contraindication, drug to drug interaction, or history of toxic side effects that caused immediate or long-term damage (DOCUMENTATION REQUIRED). Note: Mild rash, itching, and GI intolerance are not accepted as intolerance to naloxone. PA Approval Durations: Pregnancy: 3 months past due date; Breastfeeding: 6 months (maximum 4 approvals); Contraindication to Naloxone: Initial Authorization 6-months, Reauthorization 12 months	8 mg: 2/day for 6 months then 1/day*; 2 mg: 3/day* ^	Buprenorphin Products PA Form	
buprenorphine/ naloxone film	NP	See buprenorphine/naloxone tab prior authorization criteria; AND • Patient has a documented allergy to inactive ingredient in preferred product that is not in requested product (DOCUMENTATION REQUIRED)	12/3mg & 8/2 mg: 2/day for 6 months		
Suboxone® film	NP	See buprenorphine/naloxone tab prior authorization criteria; AND • Patient has a documented allergy to inactive ingredient in preferred product that is not in requested product (DOCUMENTATION REQUIRED)	then 1/day*; 4/1 mg: 2/day 2/0.5 mg: 3/day* ^		



		ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indica	ited.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Zubsolv®	NP	See buprenorphine/naloxone tab prior authorization criteria • Patient has a documented allergy to inactive ingredient in preferred product that is not in requested product (DOCUMENTATION REQUIRED)	11.4/2.9 & 8.6/2.1 mg: 1.day; 5.7/1.4mg :2/day for 6 months then 1/day*; 2.9/0.71 mg: 2/day; 1.4/0.36 mg: 3/day; 0.7/0.18 mg:3/day*	
		nantities may be approved as medically necessary. Il only be approved if dose is being titrated or patient's condition is too unstable to attempt to change to a higher strength		
		Transmucosal Fentanyl Products		
fentanyl lozenge	NP	 Medication is ordered for the treatment of breakthrough cancer pain Recipient must be receiving around-the-clock scheduled long-acting opioids Recipient must be tolerant to opioids, defined as one of the following: ≥ 60 mg oral morphine per day for at least one week without adequate pain relief ≥ 25 mcg/hr transdermal fentanyl for at least one week without adequate pain relief ≥ 30 mg oral oxycodone/day for at least one week without adequate pain relief ≥ 8 mg oral hydromorphone/day for at least one week without adequate pain relief ≥ 25 mg oral oxymorphone/day for at least one week without adequate pain relief Equianalgesic dose of another opioid for at least one week without adequate pain relief Trial and failure, contraindication, intolerance, or drug-to-drug interaction with at least 2 immediate release opioid products Note: Prescription should be written by or in consultation with an oncologist or pain management specialist unless patient is enrolled in or eligible for hospice care. 	4/day	General PA Form
Fentora®	NP	See fentanyl lozenge prior authorization criteria	4/day	
Subsys®	NP	See fentanyl lozenge prior authorization criteria	4/day	
		Naloxone Products	,	
Kloxxado®	Р		2 sprayers/30 days	
naloxone injection	Р		2 injections/30 days	General PA
naloxone nasal spray (Rx & OTC)	Р		2 sprayers/30 days	Form
Narcan®	Р		2 sprayers/30 days	
Opvee®	Р		2 sprayers/30 days	



ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
		Narcotic Agonist/Antagonists			
nalbuphine	Р	 Trial and failure of at least 2 short acting narcotics; OR Documented contraindication, or intolerance to short acting narcotics; AND Unable to swallow, OR Unable to absorb medications through the GI tract. 	10 mg/mL: 4 mL/day 20 mg/mL: 8 mL/day		
butorphanol nasal spray	NP	 Documented inability to swallow or absorb PO narcotics, OR For the treatment of migraines; AND Recipient MUST be receiving prophylactic therapy for migraines, AND Trial and failure, intolerance, or contraindication to at least ONE agent in EACH of the following categories:	2.5 mL/30 days	General PA Form	
pentazocine/ naloxone	NP	 Contraindication, or intolerance to ALL short acting narcotics Prescriber has checked the Tennessee Controlled Substance Database for this patient within the last 30 days 	12/day		

ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
	Narcotics, Long-Acting Approval of non-preferred agents in the Long-Acting Narcotics class requires contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage (NOTE: this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated. *** Edits on agents in the Short-Acting and Long-Acting Narcotics classes of the PDL impact all first-time (acute) and non-chronic opioid users. For details, visit: Acute Use Opioid Criteria				
fentanyl patch 12, 25, 50, 75, & 100 mcg	Р	See morphine ER tablets prior authorization criteria	10 patches/30 days; *^Max Total: Non-Chronic: 60 <u>MME/day;</u> Chronic: 200 <u>MME/day</u>		



		ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise ind	licated.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Narcotics, Long-Acting		
		preferred agents in the Long-Acting Narcotics class requires contraindication, drug to drug interaction, or history of toxic side effects that (NOTE: this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated. gents in the Short-Acting and Long-Acting Narcotics classes of the PDL impact all first-time (acute) and non-chronic opioid users. For details		
norphine ER ablets	Р	 Management of severe pain with need for around-the-clock analgesia for an extended period; AND Prescriber must have checked the Tennessee Controlled Substance Database for this patient within the last 7 days; AND Pain agreement required. Please refer to the Opioid and Controlled Substance Agreement; AND Concomitant use of benzodiazepines and opioids will only be approved under the care of, or referral to, a mental health provider; AND Requests for strengths ≥ 90mg: (Please refer to the TennCare MME Conversion Chart) Recipient must be opioid tolerant (as demonstrated by at least a week or longer history of morphine ≥ 60 mg/day, oral oxycodone ≥ 30 mg/day, oral hydromorphone ≥8 mg/day, or an equianalgesic dose of another opioid); AND If request is for a female of child-bearing age (14-44 years), patient is not pregnant; AND Using contraception; OR Has an intrauterine device (IUD) or implant; OR Has history of hysterectomy, tubal ligation, or endometrial ablation; AND The provider attests to investigating ALL of the following before submitting a PA: History of substance abuse Frequent requests for early refills Reported frequent instances of lost tablets Requests for odd quantities which requires fractional dosing Requests for short-term or prn usage Medication history indicates concurrent use of other extended-release opioids Note: Use of opioid analgesics during pregnancy has been associated with Neonatal Opioid Withdrawal Syndrome. Providers MUST counsel women of childbearing age on the risks of becoming pregnant while receiving opioids, including the risk of Neonatal Opioid Withdrawal		Acute Opic PA Form Chronic Opioid PA Form Exception Opioid PA Form
ramadol ER	Р	See morphine ER prior authorization criteria	1/day; *^Max Total: Non-Chronic: 60 MME/day; Chronic: 200 MME/day	



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Narcotics, Long-Acting preferred agents in the Long-Acting Narcotics class requires contraindication, drug to drug interaction, or history of toxic side effects that (NOTE: this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated. gents in the Short-Acting and Long-Acting Narcotics classes of the PDL impact all first-time (acute) and non-chronic opioid users. For detail.	•	, i
Belbuca®	NP	 Management of severe pain with need for around-the-clock analgesia for an extended period; AND Prescriber has checked the Tennessee Controlled Substance Database for this patient within the last 7 days; AND Patients who have not been titrated down to no more than 30 mg morphine (or morphine equivalents) per day will NOT be approved; AND Pain agreement required. Please refer to the Opioid and Controlled Substance Agreement; AND Concomitant use of benzodiazepines and opioids will only be approved under the care of, or referral to, a mental health provider; AND If request is for a female of child-bearing age (14-44 years), patient is not pregnant; AND Using contraception; OR Has an intrauterine device (IUD) or implant; OR Has history of hysterectomy, tubal ligation, or endometrial ablation; AND The prescriber attests to investigating all of following before submitting a PA: History of substance abuse Frequent requests for early refills Reported frequent instances of lost tablets Requests for short-term or prn usage Medication history indicates concurrent use of other extended-release opioids; AND Contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage (this does not include Gl intolerance) with ALL preferred agents, unless otherwise indicated Note: Use of opioid analgesics during pregnancy has been associated with Neonatal Opioid Withdrawal Syndrome. Providers MUST counsel women of childbearing age on the risks of becoming pregnant while receiving opioids, including risk of Neonatal Opioid Withdrawal Syndrome. 	2/day; *^Max Total: Non-Chronic: 60 <u>MME/day;</u> Chronic: 200 <u>MME/day</u>	Acute Opioid PA Form Chronic Opioid PA Form Exceptions Opioid PA Form
buprenorphine patch	NP	See Belbuca® prior authorization criteria Additionally, Butrans® 7.5, 10, 15, and 20 mcg/hr will be approved for opioid-experienced patients only.	4 patches/28 days; *^Max Total: Non-Chronic: 60 <u>MME/day;</u> Chronic: 200 <u>MME/day</u>	
Butrans®	NP	See Belbuca® prior authorization criteria Additionally, Butrans® 7.5, 10, 15, and 20 mcg/hr will be approved for opioid-experienced patients only.	4 patches/28 days; *^Max Total: Non-Chronic: 60 <u>MME/day;</u> Chronic: 200 <u>MME/day</u>	



ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. PDL Medication **Prior Authorization Criteria Qty. Limits PA Form** Narcotics, Long-Acting Approval of non-preferred agents in the Long-Acting Narcotics class requires contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage (NOTE: this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated. *** Edits on agents in the Short-Acting and Long-Acting Narcotics classes of the PDL impact all first-time (acute) and non-chronic opioid users. For details, visit: Acute Use Opioid Criteria *** Management of severe pain with need for around-the-clock analgesia for an extended period; AND Prescriber must have checked the Tennessee Controlled Substance Database for this patient within the last 7 days; AND Pain agreement required. Please refer to the Opioid and Controlled Substance Agreement; AND • Concomitant use of benzodiazepines and opioids will only be approved under the care of, or referral to, a mental health provider; AND • If request is for a female of child-bearing age (14-44 years), patient is not pregnant; AND Using contraception; OR o Has an intrauterine device (IUD) or implant; OR o Has history of hysterectomy, tubal ligation, or endometrial ablation; AND • The prescriber attests to investigating ALL of the following before submitting a PA: History of substance abuse Frequent requests for early refills Reported frequent instances of lost tablets o Requests for odd quantities which requires fractional dosing **Acute Opioid** o Requests for short-term or prn usage PA Form o Medication history indicates concurrent use of other extended-release opioids; AND • If patient is 12 to 18 years of age: (For patients less than 12 years of age, approval will not be granted) Patient does not have any of the following: Obesity (BMI ≥ 30) 1/day; Chronic Obstructive Sleep Apnea Opioid PA NP ConZip® Severe Lung Disease (acute or severe asthma, COPD, Cystic Fibrosis, hypoxemia, hypercapnia, pneumonia, *^Max Total: Form Non-Chronic: 60 MME/day; pulmonary hypertension, etc.) Chronic: 200 MME/day Recent adenectomy/tonsillectomy; AND o Trial and failure or contraindication to acetaminophen; AND o Trial and failure or contraindication to ALL NSAIDs; AND **Exceptions** Contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage Opioid PA (this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated Form **Note**: Use of opioid analgesics during pregnancy has been associated with Neonatal Opioid Withdrawal Syndrome. Providers MUST counsel women of childbearing age on the risks of becoming pregnant while receiving opioids, including risk of Neonatal Opioid Withdrawal Syndrome. Providers should offer access to contraceptive services when necessary.

ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.							
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
Narcotics, Long-Acting Approval of non-preferred agents in the Long-Acting Narcotics class requires contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage (NOTE: this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated. *** Edits on agents in the Short-Acting and Long-Acting Narcotics classes of the PDL impact all first-time (acute) and non-chronic opioid users. For details, visit: Acute Use Opioid Criteria ***							
fentanyl patch 37.5, 62.5, & 37.5 mcg	NP	See hydromorphone ER prior authorization criteria	10 patches/30 days; *^Max Total: Non-Chronic: 60 <u>MME/day;</u> Chronic: 200 <u>MME/day</u>				
hydrocodone ER	NP	 The prescriber has checked the Tennessee Controlled Substance Database for this patient within the last 7 days; AND Pain agreement required. Please refer to the Opioid and Controlled Substance Agreement; AND Concomitant use of benzodiazepines and opioids will only be approved under the care of, or referral to, a mental health provider If request is for a female of child-bearing age (14-44 years), patient is not pregnant; AND Using contraception; OR Has an intrauterine device (IUD) or implant; OR Has history of hysterectomy, tubal ligation, or endometrial ablation; AND Approval of non-preferred agents requires: Contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage (NOTE: this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated. The following should be investigated before a PA is granted: History of substance abuse Frequent requests for early refills Reported frequent instances of lost tablets Requests for short-term or prn usage Medication history indicates concurrent use of other extended-release opioids Requests for strengths ≥ 90mg: (Please refer to the TennCare MME Conversion Chart) Recipient must be opioid tolerant (as demonstrated by at least a week or longer history of morphine ≥ 60 mg/day, oral oxycodone > 30 mg/day, oral hydromorphone > 8 mg/day, or an equianalgesic dose of another opioid) 	Tabs: 1/day; Caps: 2/day; *^Max Total: Non-Chronic: 60 <u>MME/day;</u> Chronic: 200 <u>MME/day</u>	Acute Opioid PA Form Chronic Opioid PA Form Exceptions Opioid PA Form			



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
		Narcotics, Long-Acting preferred agents in the Long-Acting Narcotics class requires contraindication, drug to drug interaction, or history of toxic side effects that of (NOTE: this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated. gents in the Short-Acting and Long-Acting Narcotics classes of the PDL impact all first-time (acute) and non-chronic opioid users. For details Management of severe pain with need for around-the-clock analgesia for an extended period; AND Prescriber must have checked the Tennessee Controlled Substance Database for this patient within the last 7 days; AND	•	, i	
hydromorphone ER	NP	 Pain agreement required. Please refer to the Opioid and Controlled Substance Agreement; AND Concomitant use of benzodiazepines and opioids will only be approved under the care of, or referral to, a mental health provider; AND Requests for strengths ≥ 90mg: (Please refer to the TennCare MME Conversion Chart) Recipient must be opioid tolerant (as demonstrated by at least a week or longer history of morphine ≥ 60 mg/day, oral oxycodone ≥ 30 mg/day, oral hydromorphone ≥ 8 mg/day, or an equianalgesic dose of another opioid); AND If request is for a female of child-bearing age (14-44 years), patient is not pregnant; AND Using contraception; OR Has an intrauterine device (IUD) or implant; OR Has history of hysterectomy, tubal ligation, or endometrial ablation; AND The provider attests to investigating ALL of the following before submitting a PA: History of substance abuse Frequent requests for early refills Reported frequent instances of lost tablets Requests for odd quantities which requires fractional dosing Requests for short-term or prn usage Medication history indicates concurrent use of other extended-release opioids; AND Contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage (this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated Note: Use of opioid analgesics during pregnancy has been associated with Neonatal Opioid Withdrawal Syndrome. Providers MUST counsel women of childbearing age on the risks of becoming pregnant while receiving opioids, including the risk of Neonatal Opioid Withdrawal Syndrome. 	Tablet: 1/day; *^Max Total: Non-Chronic: 60 MME/day; Chronic: 200 MME/day	Acute Opio PA Form Chronic Opioid PA Form Exception: Opioid PA Form	
Hysingla® ER	NP	See hydromorphone ER prior authorization criteria	1/day; *^Max Total: Non-Chronic: 60 <u>MME/day;</u> Chronic: 200 <u>MME/day</u>		



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Narcotics, Long-Acting preferred agents in the Long-Acting Narcotics class requires contraindication, drug to drug interaction, or history of toxic side effects that contraindication (NOTE: this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated. gents in the Short-Acting and Long-Acting Narcotics classes of the PDL impact all first-time (acute) and non-chronic opioid users. For details One of the following:	•	_
methadone	NP	 Diagnosis of Metastatic Neoplasia Infants up to 1 year of age who are discharged from hospital on a methadone taper will be approved for up to 30 days Management of severe pain with need for around-the-clock analgesia for an extended period AND patient has contraindication to all other long-acting opioids; AND Prescriber must have checked the Tennessee Controlled Substance Database for this patient within the last 7 days; AND Pain agreement required. Please refer to the Opioid and Controlled Substance Agreement; AND Concomitant use of benzodiazepines & opioids will only be approved under the care of, or referral to, a mental health provider; AND Requests for strengths ≥ 90mg: (Please refer to the TennCare MME Conversion Chart) Recipient must be opioid tolerant (as demonstrated by at least a week or longer history of morphine ≥ 60 mg/day, oral oxycodone ≥ 30 mg/day, oral hydromorphone ≥ 8 mg/day, or an equianalgesic dose of another opioid); AND If request is for a female of child-bearing age (14-44 years), patient is not pregnant; AND Using contraception; OR Has nistory of hysterectomy, tubal ligation, or endometrial ablation; AND The following should be investigated before a PA is granted: History of substance abuse Frequent requests for early refills Reported frequent instances of lost tablets Requests for odd quantities which requires fractional dosing Requests for short-term or prn usage Medication history indicates concurrent use of other extended-release opioids; AND Note: TennCare does not cover any form of methadone for the treatment of opioid addiction. Note: Use of opioid analgesics during pregnancy has been associated with Neonatal Opioid Withdrawal Syndrome. Providers MUST counsel women of childbearing age on the risks of becoming	5 mg: 8/day; 10 mg: 4/day; 5 mg/5 mL: 40mL/day; 10 mg/5 mL: 20 mL/day; 10 mg/mL: 4 mL/day; *^Max Total: Non-Chronic: 60 MME/day; Chronic: 200 MME/day	Acute Opioic PA Form Chronic Opioid PA Form Exceptions Opioid PA Form
Methadose®	NP	See methadone prior authorization criteria	See methadone	
morphine ER capsules	NP	See hydromorphone ER prior authorization criteria	Beads Caps: 1/day; Caps: 2/day *^Max Total: Non-Chronic: 60 <u>MME/day</u> Chronic: 200 <u>MME/day</u>	



ANALGESICS

Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.

Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
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Narcotics, Long-Acting

Approval of non-preferred agents in the Long-Acting Narcotics class requires contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage (NOTE: this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated.

*** Edits on agents in the Short-Acting and Long-Acting Narcotics classes of the PDL impact all first-time (acute) and non-chronic opioid users. For details, visit: Acute Use Opioid Criteria ***

MS Contin®	NP	See hydromorphone ER prior authorization criteria	15, 30, 60 mg: 3/day; 100 mg: 2/day; 200 mg: 1/day; *^Max Total:	Acute Opioid PA Form
			Non-Chronic: 60 MME/day; Chronic: 200 MME/day	<u>Chronic</u> <u>Opioid PA</u>
oxycodone ER	NP	See hydromorphone ER prior authorization criteria	-4.	<u>Form</u>
Oxycontin®	NP	See hydromorphone ER prior authorization criteria	2/day; *^Max Total:	Exceptions
Oxymorphone ER	NP	See hydromorphone ER prior authorization criteria Note : Due to cross-reactivity with morphine, oxymorphone SR will not be approved for patients with immune-mediated morphine allergy.	Non-Chronic: 60 MME/day; Chronic: 200 MME/day	Opioid PA Form

*^Morphine Milligram Equivalent (MME) Criteria:

- Indication or diagnosis is Cancer pain or Hospice
 - Prescriber must have checked the Tennessee Controlled Substance Database for this patient within the last 7 days (document date); AND
 - Document prescriber's specialty; AND
 - Patient has a written treatment plan with established objectives; AND
 - Patient has a signed Pain Management Agreement; AND
 - If request is for a female of child-bearing age (14-44 years), patient is not pregnant; AND
 - Using contraception (e.g., barrier, oral contraceptive, rhythm method); OR
 - Has an intrauterine device (IUD) or implant; **OR**
 - Has history of hysterectomy, tubal ligation, or endometrial ablation



ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. **PDL** Medication **Prior Authorization Criteria** Qty. Limits **PA Form Narcotics, Short-Acting** Approval of non-preferred agents in the Short-Acting Narcotics class requires contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage (NOTE: this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated. *** Edits on agents in the Short-Acting and Long-Acting Narcotics classes of the PDL impact all first-time (acute) and non-chronic opioid users. For details, visit: Acute Use Opioid Criteria *** • Patient is > 12 years of age and < 18 years of age; AND Trial and failure of acetaminophen; AND Contraindication to ALL NSAIDs; AND 12/day: *^Max Total: • Patient does not have any of the following: codeine/APAP Obesity Non-Chronic: 60 MME/day Chronic: 200 MME/day Obstructive Sleep Apnea Severe Lung Disease (acute or severe asthma, COPD, Cystic Fibrosis, hypoxemia, hypercapnia, pneumonia, etc.) Recent adenectomy/tonsillectomy 2.5/325 mg tab: 12/day; All other tabs: 8/day; Endocet® *^Max Total: Non-Chronic: 60 MME/day Chronic: 200 MME/day **Acute Opioid** 5/325 mg tab: 12/day; **PA Form** 7.5/325 & 10/325 mg tabs: 8/day; hydrocodone/ Ρ soln: 120 mL/day; APAP 325 mg **Chronic Opioid** *^Max Total: PA Form Non-Chronic: 60 MME/day Chronic: 200 MME/day 5/200 mg tab: 12/day; 7.5/200 mg tab: 8/day; Exceptions hydrocodone/ 10/200 mg tab: 6/day; **Opioid PA Form** ibuprofen *^Max Total: Non-Chronic:60 MME/day; Chronic: 200 MME/day 2 mg: 7/day; 4 mg: 3/day; hydromorphone 8 mg: 1/day; tabs *^Max Total: Non-Chronic:60 MME/day Chronic: 200 MME/day 6/day; *^Max Total: morphine IR tabs Non-Chronic: 60 MME/day Chronic: 200 MME/day



		ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise in	ndicated.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Narcotics, Short-Acting		
Approval o	of non-p	referred agents in the Short-Acting Narcotics class requires contraindication, drug to drug interaction, or history of toxic side effects the (NOTE: this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated.	at cause immediate or long-term	damage
*** Edit	s on ag	ents in the Short-Acting and Long-Acting Narcotics classes of the PDL impact <u>all first-time (acute) and non-chronic opioid users</u> . For deta	nils, visit: Acute Use Opioid Criteri	<u>a</u> ***
morphine solution	P	 Prescriber has checked the Tennessee Controlled Substance Database for this patient within the last 7 days (document date); OR request is for a hospice patient, HIV/AIDS patient, active cancer patient, OR long-term care facility resident (document name of facility); AND Concomitant use of benzodiazepines and opioids will only be approved under the care of, or referral to, a mental health provider; AND Pain agreement required. Please refer to the Opioid and Controlled Substance Agreement; AND If patient is females and of child-bearing age (14-44 years), patient is not pregnant; AND One of the following: Using contraception Has an intrauterine device (IUD) or implant Has history of hysterectomy, tubal ligation, or endometrial ablation; AND Recipient must be opioid tolerant (as demonstrated by ≥1 week history of morphine ≥ 60 mg/day, oral oxycodone ≥ 30 mg/day, oral hydromorphone ≥8 mg/day, or an equianalgesic dose of another opioid) 	*^Max Total: Non-Chronic: 60 <u>MME/day</u> Chronic: 200 <u>MME/day</u>	Acute Opioic PA Form
oxycodone/ APAP 325mg	Р		2.5/325 mg tab: 12/day; All other tabs: 8/day; soln: 40 mL/day *^Max Total: Non-Chronic: 60 MME/day Chronic: 200 MME/day	Chronic Opioid PA Form
oxycodone concentrate	Р	See morphine solution prior authorization criteria	*^Max Total: Non-Chronic: 60 MME/day Chronic: 200 MME/day	Exceptions Opioid PA Form
oxycodone tabs	Р		5 & 10 mg: 8/day; 15, 20, & 30 mg: 4/day; *^Max Total: Non-Chronic: 60 <u>MME/day</u> Chronic: 200 <u>MME/day</u>	
oxycodone soln	Р		*^Max Total: Non-Chronic: 60 MME/day Chronic: 200 MME/day	



ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. **PDL** Medication **Prior Authorization Criteria Qty. Limits PA Form** Narcotics, Short-Acting Approval of non-preferred agents in the Short-Acting Narcotics class requires contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage (NOTE: this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated. *** Edits on agents in the Short-Acting and Long-Acting Narcotics classes of the PDL impact all first-time (acute) and non-chronic opioid users. For details, visit: Acute Use Opioid Criteria *** Patient is > 12 years of age and < 18 years of age; AND • Patient does not have any of the following: o Obesity (BMI ≥ 30) 8 tabs/day; Obstructive Sleep Apnea 80 mL/day tramadol Severe Lung Disease (acute or severe asthma, COPD, Cystic Fibrosis, hypoxemia, hypercapnia, pneumonia, etc.) *^Max Total: Recent adenectomy/tonsillectomy; AND Non-Chronic: 60 MME/day • Trial and failure or contraindication to acetaminophen; AND Chronic: 200 MME/day Trial and failure or contraindication to ALL NSAIDs Note: Patients 18 years and older will only be subject to the quantity limit and opioid criteria 12/day; *^Max Total: tramadol/APAP See tramadol prior authorization criteria Acute Opioid Non-Chronic: 60 MME/day **PA Form** Chronic: 200 MME/day 6.12/325 mg tab: 8/day; 8.16/325 mg tab: 6/day; Apadaz® NΡ 4.08/325 mg tab: 12/day Chronic Max: 4 g APAP/day Opioid PA benzhydrocodone/ Form See Apadaz® APAP Contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage (NOTE: this does not include GI intolerance) with ALL preferred short-acting narcotic agents; AND **Exceptions** • One of the following: **Opioid PA** Patients ≥ 18 years of age Form Patient is > 12 years of age and < 18 years of age; AND - Trial and failure of acetaminophen; AND **Butalbital-containing** butalbital/APAP/ - Contraindication to ALL NSAIDs; AND products: 20/30 days** caffeine/codeine - Patient does not have any of the following: Max: 4 g APAP/day Obesity Obstructive Sleep Apnea Severe Lung Disease (acute or severe asthma, COPD, Cystic Fibrosis, hypoxemia, hypercapnia, pneumonia, etc.) Recent adenectomy/tonsillectomy butalbital/ASA/ **Butalbital-containing** See butalbital/APAP/caffeine/codeine prior authorization criteria caffeine/codeine products: 20/30 days**



ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. **PDL** Medication **Prior Authorization Criteria Qty. Limits PA Form** Narcotics, Short-Acting Approval of non-preferred agents in the Short-Acting Narcotics class requires contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage (NOTE: this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated. *** Edits on agents in the Short-Acting and Long-Acting Narcotics classes of the PDL impact all first-time (acute) and non-chronic opioid users. For details, visit: Acute Use Opioid Criteria *** 15 mg & 30 mg: 12/day; 60 mg: 6/day; codeine See butalbital/APAP/caffeine/codeine prior authorization criteria *^Max Total: Non-Chronic: 60 MME/day Chronic: 200 MME/day dihydrocodeine/ 8 tabs/day; See butalbital/APAP/caffeine/codeine prior authorization criteria APAP/caffeine Max: 4 g APAP/day Contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage (NOTE: this does not include GI intolerance) with ALL preferred agents; AND Prescriber must have checked the Tennessee Controlled Substance Database for this patient within the last 7 days; AND Pain agreement required. Please refer to the Opioid and Controlled Substance Agreement; AND 2 mg: 7/day; • Concomitant use of benzodiazepines and opioids will only be approved under the care of, or referral to, a mental health Acute Opioid 4 mg: 3/day; provider. **PA Form** 8 mg: 1/day; If request is for a female of child-bearing age (14-44 years), patient is not pregnant; AND Dilaudid® *^Max Total: Using contraception; OR Non-Chronic: 60 MME/day o Has an intrauterine device (IUD) or implant; OR Chronic Chronic: 200 MME/day Has history of hysterectomy, tubal ligation, or endometrial ablation; AND Opioid PA • Has history of hysterectomy, tubal ligation, or endometrial ablation Form Note: Use of opioids during pregnancy has been associated with Neonatal Opioid Withdrawal Syndrome. Providers MUST counsel women of childbearing age regarding the risks of becoming pregnant while receiving opioids, including the risk of Neonatal Opioid Withdrawal Syndrome. Providers should offer access to contraceptive services when necessary. Exceptions **Butalbital-containing** Fioricet® with Opioid PA See butalbital/APAP/caffeine/codeine prior authorization criteria products: 20/30 days** codeine Form Max: 4 g APAP/day 5/300 mg tab: 12/day; 10/300 mg tab: 6/day; hydrocodone/ Soln: 89 mL/day; See Dilaudid® prior authorization criteria APAP 300 mg *^Max Total: Non-Chronic: 60 MME/day Chronic: 200 MME/day 15 mL/day; *^Max Total: hydromorphone Non-Chronic: 60 MME/day See Dilaudid® prior authorization criteria liquid Chronic: 200 MME/day 5/day; *^Max Total: hydromorphone See Dilaudid® prior authorization criteria Non-Chronic: 60 MME/day suppositories Chronic: 200 MME/day



ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. **PDL** Medication **Prior Authorization Criteria Qty. Limits PA Form** Narcotics, Short-Acting Approval of non-preferred agents in the Short-Acting Narcotics class requires contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage (NOTE: this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated. *** Edits on agents in the Short-Acting and Long-Acting Narcotics classes of the PDL impact all first-time (acute) and non-chronic opioid users. For details, visit: Acute Use Opioid Criteria *** 6/day; *^Max Total: levorphanol See Dilaudid® prior authorization criteria Non-Chronic: 60 MME/day Chronic: 200 MME/day 5/325 mg tabs: 8/day; All other tabs: 8/day; soln: 89 mL/day; Lortab® See Dilaudid® prior authorization criteria *^Max Total: Non-Chronic: 60 MME/day Chronic: 200 MME/day tabs: 12/day; **Acute Opioid** soln: 60 mL/day; **PA Form** See Dilaudid® prior authorization criteria *^Max Total: meperidine Non-Chronic: 60 MME/day Chronic: 200 MME/day 5 mg: 12/day; Chronic All others: 6/day; Opioid PA morphine See Dilaudid® prior authorization criteria *^Max Total: Form suppositories Non-Chronic: 60 MME/day Chronic: 200 MME/day 12/day; *^Max Total: **Exceptions** Nalocet® See Dilaudid® prior authorization criteria Non-Chronic: 60 MME/day **Opioid PA** Chronic: 200 MME/day Form 8/day; *^Max Total: Oxaydo® See Dilaudid® prior authorization criteria Non-Chronic: 60 MME/day Chronic: 200 MME/day 2.5/325 mg tab: 12/day; All other tabs: 8/day; soln: 40 mL/day oxycodone/ See Dilaudid® prior authorization criteria



APAP 300 mg

*^Max Total: Non-Chronic: 60 MME/day Chronic: 200 MME/day

ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. PDL Medication **Prior Authorization Criteria Qty. Limits PA Form** Narcotics, Short-Acting Approval of non-preferred agents in the Short-Acting Narcotics class requires contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage (NOTE: this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated. *** Edits on agents in the Short-Acting and Long-Acting Narcotics classes of the PDL impact all first-time (acute) and non-chronic opioid users. For details, visit: Acute Use Opioid Criteria *** 8/day; *^Max Total: oxycodone caps NP | See Dilaudid® prior authorization criteria Non-Chronic: 60 MME/day Chronic: 200 MME/day 4/day; *^Max Total: oxymorphone See Dilaudid® prior authorization criteria Non-Chronic: 60 MME/day Chronic: 200 MME/day **Acute Opioid** 2.5/325 mg: 12/day; **PA Form** All others: 8/day; Percocet® *^Max Total: See Dilaudid® prior authorization criteria Non-Chronic: 60 MME/day Chronic: 200 MME/day Chronic tabs: 8/day; Opioid PA soln: 40 mL/day; Form Prolate® *^Max Total: See Dilaudid® prior authorization criteria Non-Chronic: 60 MME/day Chronic: 200 **Exceptions** *^Max Total: Qdolo® NP Non-Chronic: 60 MME/day Opioid PA Chronic: 200 Form 4/day; *^Max Total: Roxicodone® See Dilaudid® prior authorization criteria Non-Chronic: 60 MME/day Chronic: 200 MME/day 4/day; *^Max Total: Roxybond® See Dilaudid® prior authorization criteria Non-Chronic: 60 MME/day Chronic: 200 MME/day



		ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise inc	dicated.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Narcotics, Short-Acting		
Approval	of non-p	referred agents in the Short-Acting Narcotics class requires contraindication, drug to drug interaction, or history of toxic side effects that (NOTE: this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated.	cause immediate or long-term	damage
*** Edi	s on ag	ents in the Short-Acting and Long-Acting Narcotics classes of the PDL impact <u>all first-time (acute) and non-chronic opioid users</u> . For detail	ls, visit: Acute Use Opioid Criteri	<u>a</u> ***
Seglentis®	NP	 Patient is > 12 years of age and < 18 years of age; AND Patient does not have any of the following: Obesity (BMI ≥ 30) Obstructive Sleep Apnea Severe Lung Disease (acute or severe asthma, COPD, Cystic Fibrosis, hypoxemia, hypercapnia, pneumonia, pulmonary hypertension, etc.) Recent adenectomy/tonsillectomy; AND Trial and failure or contraindication to acetaminophen; AND Trial and failure or contraindication to ALL NSAIDs; AND Contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage (NOTE: this does not include GI intolerance) with ALL preferred agents; AND Patient is ≥ 18 years of age: If request is for a female of child-bearing age (14-44 years), patient is not pregnant; AND Using contraception; OR Has an intrauterine device (IUD) or implant; OR Has history of hysterectomy, tubal ligation, or endometrial ablation; AND Contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage (NOTE: this does not include GI intolerance) with ALL preferred agents; AND Note: Use of opioids during pregnancy has been associated with Neonatal Opioid Withdrawal Syndrome. Providers MUST counsel women of childbearing age regarding the risks of becoming pregnant while receiving opioids, including the risk of Neonatal Opioid Withdrawal Syndrome. Providers should offer access to contraceptive services when necessary. 	12/day; *^Max Total: Non-Chronic: 60 <u>MME/day</u> Chronic: 200 <u>MME/day</u>	Acute Opioid PA Form Chronic Opioid PA Form Exceptions Opioid PA Form
Ultracet®	NP	See Seglentis® prior authorization criteria	12/day; *^Max Total: Non-Chronic: 60 <u>MME/day</u> Chronic: 200 <u>MME/day</u>	



ANALGESICS

Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.

Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
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Narcotics, Short-Acting

Approval of non-preferred agents in the Short-Acting Narcotics class requires contraindication, drug to drug interaction, or history of toxic side effects that cause immediate or long-term damage (NOTE: this does not include GI intolerance) with ALL preferred agents, unless otherwise indicated.

*** Edits on agents in the Short-Acting and Long-Acting Narcotics classes of the PDL impact all first-time (acute) and non-chronic opioid users. For details, visit: Acute Use Opioid Criteria ***

**Quantity Limit Override Criteria for Butalbital-Containing Products:

Requests for butalbital-containing products for quantities greater than 20 per 30 days will be approved for patients meeting the following criteria:

• Trial and failure of at least 2 prophylactic headache treatments: a tricyclic antidepressant (unless contraindicated) PLUS at least one of the following: divalproex sodium, sodium valproate, topiramate, frovatriptan or beta-blocker

*^Morphine Milligram Equivalent (MME) Criteria:

- Indication or diagnosis is Cancer pain or Hospice
 - Prescriber must have checked the Tennessee Controlled Substance Database for this patient within the last 7 days (document date); AND
 - Document prescriber's specialty; AND
 - Patient has a written treatment plan with established objectives; AND
 - Patient has a signed Pain Management Agreement; AND
 - Female of child-bearing age (14-44 years):
 - Is not pregnant; AND
 - Using contraception; OR
 - Has an intrauterine device (IUD) or implant; OR
 - Has history of hysterectomy, tubal ligation or endometrial ablation

	ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
		NSAIDs				
celecoxib	Р		2/day			
diclofenac 1% gel	Р		10 g/day			
ketorolac tabs	Р		20/60 days			
Pennsaid	Р	Diagnosis of osteoarthritis pain of the knee				
Voltaren® gel	Р		10 g/day			
Celebrex®	NP		2/day	General PA		
diclofenac caps, packet, and solution	NP	Clinically valid reason why the preferred NSAIDs cannot be used		<u>Form</u>		
diclofenac patch	NP	Clinically valid reason why the preferred NSAIDs cannot be used	2 patches/day			
diclofenac potassium 25 mg tabs		Clinically valid reason why the preferred diclofenac products cannot be used				



	ANALGESICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.			
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Elyxb®	NP	 Diagnosis of migraine; AND Patient is unable to swallow solid dosage forms 	120 mg/day	
Lofena®	NP	Clinically valid reason why the preferred diclofenac products cannot be used		
ketorolac spray	NP	Trial and failure, contraindication, or intolerance of oral ketorolac; OR Patient is unable to swallow solid dosage forms	5 bottles/60 days	
Flector®	NP	Clinically valid reason why the preferred NSAIDs cannot be used	2 patches/day	
meloxicam caps	NP	Clinically valid reason why the preferred meloxicam tablets cannot be used	1/day	
Sprix®	NP	 Trial and failure, contraindication, or intolerance of oral ketorolac; OR Patient is unable to swallow solid dosage forms 	5 bottles/60 days	General PA
Toradol®	NP		20/60 days	Form
Zorvolex®	NP	Clinically valid reason why the preferred NSAIDs cannot be used		
		NSAID/Anti-Ulcer Agents	<u>.</u>	
diclofenac/ misoprostol	Р	 Patient is ≥ 60 years old; OR Patients < 60 years old and is at high risk for GI side effects as indicated by ANY of the following: History of peptic ulcer disease/GI bleed/NSAID gastropathy GERD (gastroesophageal reflux disease) due to conventional NSAIDS Patient on anticoagulants Patient on chronic corticosteroids History of platelet dysfunction or coagulopathy, including use of clopidogrel or aspirin Patient on methotrexate 	50 mg/200 mcg: 4/day; 75 mg/200 mcg: 2/day	
Ibuprofen/ famotidine	Р	 Patient is at high risk for GI side effects as indicated by ANY of the following: History of peptic ulcer disease/GI bleed/NSAID gastropathy GERD (gastroesophageal reflux disease) due to conventional NSAIDS Patient on anticoagulants Patient on chronic corticosteroids History of platelet dysfunction or coagulopathy, including use of clopidogrel or aspirin Patient on methotrexate 	3/day	General PA Form
Vimovo®	Р	See Duexis® prior authorization criteria	2/day	
Arthrotec®	NP		50 mg/200 mcg: 4/day; 75 mg/200 mcg: 2/day	
Duexis®	NP		3/day	
naproxen/ esomeprazole	NP		2/day	
		Salicylates		
salsalate	Р		500 mg: 6/day; 750 mg: 4/day	General PA
diflunisal	NP		3/day	<u>Form</u>



		ANTI-INFECTIVES		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Antibiotics: Agents for Diarrhea		
vancomycin soln	Р	 Patient is unable to swallow sold dosage forms; OR Patient is < 12 years of age 	2,000 mg/day	
Aemcolo®	NP	 Patient is being treated for traveler's diarrhea; AND Trial and failure, contraindication, intolerance, drug-drug interaction or resistance to a fluoroquinolone or azithromycin 	12 tabs/Rx; max 24 tabs/year	
Firvanq®	NP	Trial and failure, contraindication, or intolerance to generic vancomycin solution	2,000 mg/day	
Vancocin®	NP	Trial and failure, contraindication, or intolerance to vancomycin capsules		
		Antibiotics: Aminoglycosides, Oral		
Arikayce®	NP	Initial Criteria: Patient is ≥ 18 years of age; AND Diagnosis of Mycobacterium avium complex (MAC) lung disease as determined by the following: Chest radiography or high-resolution computed tomography (HRCT) scan; AND At least two positive sputum cultures; AND Other conditions such as tuberculosis and lung malignancy have been ruled out; AND Patient has failed a multi-drug regimen with a macrolide (clarithromycin or azithromycin), rifampin, and ethambutol. (Failure is defined as continual positive sputum cultures for MAC while adhering to a multi-drug treatment regimen for a minimum duration of 6-months); AND Prescribed in conjunction with a multi-drug antimycobacterial regimen Renewal Criteria: Patient has demonstrated response to therapy defined as having three consecutive monthly negative sputum cultures by month six of treatment; AND Patient has not experienced toxicity to amikacin treatment (e.g., ototoxicity, renal toxicity, neuromuscular blockade) Antibiotics: Anti-Tuberculosis, Oral	8.4 mL/day	General PA Form
Sirturo®	NP	 Criteria: (9-month approval duration) Patient is ≥ 5 years of age and weighs ≥ 15 kg; AND Patient has a diagnosis of pulmonary multi drug-resistant tuberculosis (MDR-TB); AND Sirturo is prescribed as part of a combination regimen with at least 3 other drugs to which the patient's MDR-TB isolate has been shown to be susceptible; AND Sirturo is prescribed by, or in consultation with, an infectious disease specialist 		
		Antibiotics: Cephalosporins Third Generation		
cefpodoxime suspension	NP	 Patient less than 12 years of age and treatment is for genitourinary infection; OR Patient is unable to swallow solid dosage forms 		General PA Form
		Antibiotics: Lincosamides, Oral	,	
clindamycin pediatric solution	Р	 Patient less than 12 years of age; OR Patient is unable to swallow solid dosage forms 		General PA
Cleocin® Pediatric granules	NP	Patient is unable to swallow solid dosage forms		<u>Form</u>



		ANTI-INFECTIVES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Antibiotics: Macrolides		
azithromycin packet	Р		2 g/Rx	
azithromycin suspension	Р			
azithromycin tablets	Р		250, 500 mg: 12/Rx 600 mg: 8/month	
clarithromycin ER/XL	NP		2/day	General PA
Dificid® tablets & suspension	NP	• Diagnosis of Clostridium difficile (C. diff) associated diarrhea Note: Individuals started on Dificid® therapy in the hospital will be approved for this agent following hospital discharge to allow for completion of the course of therapy.	Tabs: 2/day Susp: 1 bottle/Rx	General PA Form
Zithromax® packet	NP		2 g/Rx	
Zithromax® susp	NP			
Zithromax® tablet	NP		250, 500 mg: 12/Rx 600 mg: 8/month	
		Antibiotics: Nitrofurans, Oral		
nitrofurantoin	Р	Patient is unable to swallow solid dosage forms Note: PA not required for patients less than 12 years of age.		General PA Form
suspension		Antibiotics: Oxazolidinones		FOITH
linezolid tablets	Р	 Treatment is for ONE of the following: Vancomycin Resistant Enterococcus faecalis infections Healthcare-associated Methicillin-Resistant Staph Aureus (MRSA) infections or community-acquired MRSA with polyresistance Community-acquired pneumonia (CAP) caused by S. pneumoniae or S. aureus (MSSA) Nosocomial pneumonia caused by S. pneumoniae or S. aureus (including MSSA and MRSA) Complicated skin and skin structure infections (SSSI) caused by S. aureus (MSSA and MRSA), S. pyogenes, or S. agalactiae. Uncomplicated SSTI caused by S. aureus (MSSA only) or S. pyogenes Patient is continuing therapy from an inpatient hospital stay (to facilitate completion of therapy) 	2/day	General PA
linezolid suspension	Р	 One of the following: Patient is less than 12 years of age Patient is unable to swallow oral dosage forms Patient is continuing therapy from an inpatient hospital stay (to facilitate completion of therapy) 		Form
Sivextro®	NP	 Diagnosis of acute bacterial skin and skin structure infection; AND Patient must be resistant to or have a contraindication, or intolerance, to all other treatment options; OR Patient is continuing therapy from an inpatient hospital stay (to facilitate completion of therapy) 	1/day	
Zyvox® suspension	NP		60 mL/day	
Zyvox® tablets	NP		2/day	



ANTI-INFECTIVES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.				
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Antibiotics: Quinolones, Oral		
Baxdela®	NP	 Patient age ≥ 18 years of age; AND ONE of the following: Diagnosis of acute bacterial skin and skin structure infection (ABSSSI); AND Trial and failure to, contraindication, or resistance to ONE preferred standard of care agents for ABSSSI (e.g., linezolid, clindamycin, doxycycline, SMX-TMP, vancomycin, cephalosporin, a preferred fluoroquinolone) Diagnosis of community-acquired bacterial pneumonia (CABP); AND Trial and failure to, contraindication, or resistance to TWO preferred standard of care agents for CABP (e.g., macrolide, doxycycline, a preferred fluoroquinolone, beta-lactam, linezolid) Patient is continuing therapy from an inpatient hospital stay (to facilitate completion of therapy) 	2/day; Max 14-day supply	General PA Form
Cipro® suspension	NP	Patient is unable to swallow solid dosage forms		
ciprofloxacin suspension	NP	Patient is unable to swallow solid dosage forms		
Levofloxacin solution	NP	Patient is unable to swallow solid dosage forms		
moxifloxacin	NP	 Trial and failure, contraindication, or intolerance to 2 preferred agents; OR Patient is continuing therapy from an inpatient hospital stay (to facilitate completion of therapy) 		
		Antibiotics: Tetracyclines		
doxycycline hyclate caps	Р		50 mg: 3/day; All others: 2/day	
doxycycline hyclate tabs 50, 100 mg			50 mg: 3/day; All others: 2/day	
doxycycline monohydrate caps 50, 100 mg	Р		50 mg: 3/day; All others: 2/day	
demeclocycline	NP	 Trial and failure of 2 preferred agents; OR Treatment is for syndrome of inappropriate antidiuretic hormone secretion (SAIDH) 		
Doryx®	NP		50 mg: 3/day; All others: 2/day	General PA Form
doxycycline DR	NP		50 mg: 3/day; All others: 2/day	
doxycycline hyclate tabs 20, 75, 150 mg	NP	Agent is used as an adjunct to scaling and root planting to promote attachment level gain and to reduce pocket depth for adult periodontitis	2/day	
doxycycline monohydrate caps 75, 150 mg	NP		2/day	
doxycycline suspension	NP	Patient is unable to swallow solid dosage forms		



		ANTI-INFECTIVES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
minocycline ER	NP	 Patient is ≤ 21 years old; AND Diagnosis of non-nodular moderate to severe acne vulgaris with inflammatory lesions; AND Patient requires long-term therapy with an oral tetracycline; AND Trial and failure, contraindication, or intolerance of TWO of the following topical agents: Metronidazole (Metrogel®) Azelaic acid (Azelex®, Finacea®) Erythromycin (A/T/S® solution, gel) Clindamycin (Cleocin T®) Topical keratolytic agents (such as benzoyl peroxide, salicylic acid preparations); AND Clinically valid reason why the preferred minocycline capsules cannot be used 	1/day				
Minolira® ER	NP	See minocycline ER prior authorization criteria	1/day				
Nuzyra®	NP	 Criteria: (approval duration: 14 days) Patient is ≥ 18 years of age; AND One of the following: Community-acquired bacterial pneumonia (CABP); AND Trial and failure to, contraindication, or resistance to TWO preferred standard of care agents for CABP (e.g., macrolide, doxycycline, a preferred fluoroquinolone, beta-lactam, linezolid) Diagnosis of acute bacterial skin and skin structure infections (ABSSSI); AND Trial and failure to, contraindication, or resistance to ONE preferred standard of care agents for ABSSSI (e.g., linezolid, clindamycin, doxycycline, SMX-TMP, vancomycin, cephalosporin, a preferred fluoroquinolone) Patient is continuing therapy from an inpatient hospital stay (to facilitate completion of therapy) Diagnosis of inflammatory lesions (papules and pustules) of rosacea; AND 	3/day; Max 14-day supply	General PA Form			
Oracea®	NP	 Patient is < 21 years of age; AND Patient requires long-term therapy (greater than 3 months) with an oral antibiotic; AND Trial and failure, contraindication, or intolerance to ONE of the following topical agents: Metronidazole (e.g., MetroGel®, MetroCream®) Azelaic Acid (e.g., Azelex®, Finacea®) Erythromycin gel or solution 	2/day				
Solodyn®	NP	See minocycline ER prior authorization criteria	1/day				
Targadox®	NP		3/day				
Vibramycin®	NP		50 mg: 3/day; All others: 2/day	General PA Form			
Ximino®	NP	See minocycline ER prior authorization criteria	1/day				
	Antibiotics: UTI Agents, Miscellaneous						
fosfomycin	NP	 Trial and failure, contraindication, intolerance, or resistance to at least 2 of the following agents: Sulfamethoxazole/trimethoprim Quinolones Nitrofurantoin 	1 packet (3 g) per course of therapy	General PA Form			



		ANTI-INFECTIVES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Antibiotics, Vaginal		
Cleocin® cream	Р		40 g/Rx	
metronidazole 0.75% vaginal gel	Р		70 g/Rx	
Nuvessa®	Р		5 g/Rx	General PA
Vandazole®	Р		70 g/Rx	Form Form
clindamycin phos 2% cream	NP		40 g/Rx	
Clindesse® vaginal cream	NP		5 g/Rx	
		Antifungals, Oral		
fluconazole suspension	Р	Patient is unable to swallow solid dosage forms; OR Patients ≤ 20 years of age		
fluconazole tablets	Р		150 mg: 4/28 days	
Sporanox® capsules	Р		4/day	
Sporanox® solution	Р	Patient is unable to swallow sold dosage forms	40 mL/day	
terbinafine tablets	Р		84/year	
Ancobon®	NP	 Diagnosis of systemic candidiasis or cryptococcosis; OR Patient is continuing therapy from an inpatient hospital stay (to facilitate completion of therapy) 		
Brexafemme®	NP	 Diagnosis of vulvovaginal candidiasis; AND One of the following: ○ Patient is ≥ 18 years of age ○ Patient is a post-menarchal female; AND Patient is not pregnant; AND Trial and failure, contraindication, or intolerance to 1 preferred oral agent (fluconazole tablets) OR 1 preferred topical agent (miconazole-3 kit or terconazole) 	4 tabs/Rx	
Cresemba® oral	NP	 Patient is ≥ 6 years of age; AND Diagnosis of one of the following: Invasive aspergillosis; AND Trial and failure, contraindication, or intolerance to voriconazole OR posaconazole Invasive mucormycosis; AND A fungal culture and relevant laboratory study (including histopathology) has been obtained to isolate and identify the causative organism(s); OR Patient is continuing therapy from an inpatient hospital stay (to facilitate completion of therapy) 		
Diflucan® susp	NP	Patient is unable to swallow solid dosage forms		
Diflucan® tablets	NP		150 mg: 4/28 days	



ANTI-INFECTIVES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. PDL Medication **Prior Authorization Criteria Qty. Limits PA Form** • Diagnosis of systemic candidiasis or cryptococcosis; **OR** NP flucytosine Patient is continuing therapy from an inpatient hospital stay (to facilitate completion of therapy) itraconazole caps NP • Trial and failure of preferred Sporanox® capsules 4/day Patient is unable to swallow solid dosage forms; AND NP itraconazole soln 40 mL/day • Trial and failure of preferred Sporanox® solution Trial and failure, contraindication, or intolerance to TWO preferred agents; OR ketoconazole NP Patient is continuing therapy from an inpatient hospital stay (to facilitate completion of therapy) ONE of the following: o As indicated for the prophylaxis of invasive aspergillus and/or candida in patients who are at high risk of developing these infections due to being severely immunocompromised, such as hematopoietic stem cell transplant (HSCT) recipients with Graft versus Host Disease (GVHD), recipients with hematologic malignancies (leukemia, lymphoma, myelodysplastic syndromes) with prolonged neutropenia from chemotherapy, or recipients with AIDS. Noxafil® NP o Treatment of Fusariosis disease Treatment of Zygomycetes disease o Treatment of other fungal infections or molds that are refractory or resistant to, or in patient who have a contraindication, or intolerance to itraconazole or voriconazole Patient is continuing therapy from an inpatient hospital stay (to facilitate completion of therapy) · Patient is 18 years of age or older; AND Oravig® Patient has a diagnosis of oropharyngeal candidiasis; AND 1/day Patient has a contraindication, allergic reaction, or drug-drug interaction to clotrimazole troche and nystatin See Noxafil® prior authorization criteria posaconazole **General PA** · Diagnosed of ONE of the following: Aspergillosis (pulmonary and extrapulmonary) Form Tolsura® NP 4/day Blastomycosis (pulmonary and extrapulmonary) Histoplasmosis (including chronic cavitary pulmonary disease, disseminated, or nonmeningeal); AND Clinically valid reason why the patient cannot use the other itraconazole capsules or solution • Treatment is for ONE of the following: Candidemia (in non-neutropenic patients) o Esophageal candidiasis Invasive aspergillosis NP 18/84 days Vfend® o Serious fungal infections caused by S. apiospermum and Fusarium species including F. solani o Part of standard anti-fungal regimen in febrile neutropenic patients o Other fungal infections that are refractory or resistant to other oral triazole agents (i.e., fluconazole, ketoconazole); OR Patient is continuing therapy from an inpatient hospital stay (to facilitate completion of therapy) Diagnosis of recurrent vulvovaginal candidiasis (RVCC); AND Provider attests patient is NOT of reproductive potential; AND NP Vivjoa® The member has experienced ≥ 3 episodes of VVC in less than one year; AND • Failure of a maintenance course of oral fluconazole defined as 100-mg, 150-mg, or 200-mg taken weekly for 6-months voriconazole NP | See Vfend prior authorization criteria



		ANTI-INFECTIVES		
Medication	PDL	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated and Prior Authorization Criteria	Qty. Limits	PA Form
Wedledion		Antifungals, Vaginal	Qty. Ellinis	TATOM
Gynazole-1	Р		5 gm/day	
miconazole-3 kit	Р		1 box/Rx	
miconazole-3 vaginal supp	Р		1 box/Rx	
terconazole	Р		1 box/Rx	
		Anti-Infectives: Anthelmintics	·	
albendazole	Р	 Treatment of neurocysticercosis caused by <i>Taenia solium</i>; AND Prescribed by, or in consultation with, an Infectious Disease specialist; OR Treatment of cystic hydatid disease caused by <i>Echinococcus granulosus</i>; OR Treatment of hookworm 		
ivermectin tablets	Р		20/90 days	
Emverm®	NP	 Treatment of Enterobius vermicularis (pinworm) in single or mixed infections; AND Recipient has tried and failed, has an intolerance, OR contraindication to pyrantel pamoate; OR Treatment of Ancylostoma duodenale (common hookworm) or Necator americanus (American hookworm); AND Recipient has tried and failed, has an intolerance, OR contraindication to albendazole; OR Treatment of Trichuris trichiura (whipworm) or Ascaris lumbricoides (common roundworm); AND Recipient has tried and failed, has an intolerance, OR contraindication to ivermectin Length of authorization: Will be based on FDA indication 		General PA Form
Stromectol®	NP		20/90 days	
	1	Anti-Infectives: Antiprotozoals, Oral	-	•
atovaquone	Р	 Treatment is for Pneumocystis pneumonia (PCP) prevention or treatment; AND Trial and failure, contraindication, intolerance to sulfamethoxazole/trimethoprim; OR Diagnosis of Toxoplasmosis gondii encephalitis; AND Trial and failure, contraindication, intolerance to sulfamethoxazole/trimethoprim; OR Diagnosis of Babesiosis 		
benznidazole	NP	Diagnosis of American trypanosomiasis (Chagas disease) caused by Trypanosoma cruzi	12.5 mg: 6/day 100 mg: 4/day	
Lampit®	NP	Diagnosis of American trypanosomiasis (Chagas disease) caused by Trypanosoma cruzi		
Likmez [®]		 Patient is unable to swallow solid dosage forms; OR Patients less than 12 years of age 		General PA
Mepron®	NP	See atovaquone prior authorization criteria: AND • Trial and failure, contraindication, intolerance, or drug-drug interaction to sulfamethoxazole/trimethoprim		<u>Form</u>



		ANTI-INFECTIVES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
nitazoxanide tablets	NP	 Patient is > 12 years of age or older; AND One of the following: Treatment of diarrhea caused by Cryptosporidium parvum (Note: Will not be approved for the treatment of diarrhea caused by C. parvum in HIV-infected or immunodeficient patients); OR Treatment of diarrhea caused by Giardia lamblia; AND Patient has failed and failed, or has a contraindication, intolerance, or adverse drug reaction to tinidazole and metronidazole 	6/day	General PA Form
pyrimethamine	NP	Treatment of toxoplasmosis when used in combination with a sulfonamide		
Solosec®	NP	 Patient is 12 years of age or older; AND One of the following: Diagnosis of bacterial vaginosis; AND Trial and failure, contraindication, or intolerance to one of the following: Cleocin® vaginal cream Cleocin® vaginal suppository clindamycin capsules metronidazole tablets metronidazole vaginal gel Diagnosis of trichomoniasis caused by <i>Trichomonas vaginalis</i> (<i>T. vaginalis</i>); AND Trial and failure, contraindication, or intolerance to preferred metronidazole tablets 	1 pack/month	General PA Form
sulfadiazine	NP	 Treatment of <i>Toxoplasma gondii</i> encephalitis in combination with pyrimethamine; OR Rheumatic fever prophylaxis in patients who have a contraindication or intolerance to penicillin 		
		Antivirals: COVID Treatment		
Lagevrio®	Р	Patient is ≥ 18 years of age and older	40/5 days	General PA
Paxlovid [®]	Р	Patient is > 12 years of age and older	30/5 days	<u>Form</u>
		Antivirals: Cytomegalovirus Agents		
Livtencity®	NP	 Patient is ≥ 12 years of age and weighs ≥ 35kg; AND Diagnosis of post-transplant cytomegalovirus (CMV) infection; AND Infection is refractory to prior treatment with at least one of the following: Ganciclovir, valganciclovir, cidofovir or foscarnet 	4/day	General PA Form



	ANTI-INFECTIVES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
Prevymis [®]	NP	 Patient is > 18 years of age and older; AND One of the following: Patient is scheduled or has received an allogeneic hematopoietic stem cell transplant (HSCT) and ONE of the following:	1/day	General PA Form		
		Antivirals: Hepatitis B				
entecavir	Р		1/day	General PA		
lamivudine-HBV	Р		1/day	<u>Form</u>		
tenofovir	Р		1/day			
adefovir	NP		1/day			
Baraclude® solution	NP	 Diagnosis of chronic hepatitis B virus infection with evidence of active viral replication and either evidence of persistent elevations in serum aminotransferases (ALT or AST) or histologically active disease; AND Patient is unable to swallow tablets; AND Prescriber will monitor hepatic function closely for at least several months in patients who discontinue therapy Note: Prior authorization is not required for patients 2 through 11 years of age 	20 ml/day			
Baraclude® tablets	NP		1/day			
Vemlidy®	NP	 Patient is 6 years of age and older; AND Diagnosis of Chronic Hepatitis B virus (HBV) infection in adults with compensated liver disease; AND Inadequate treatment response (detectable HBV DNA level after 24 weeks of therapy), virologic breakthrough, resistance, intolerance, or contraindication to entecavir; AND Patient has ONE of the following: History of osteoporosis or osteopenia Renal impairment defined by CrCL <50 mL/min Clinically valid reason as to why the preferred tenofovir disoproxil fumarate (TDF) cannot be used; AND Patient is not using Vemlidy® as monotherapy if (HIV)-1 positive (must have additional antiviral therapy if HIV-1 positive for coverage of both disease states); AND Prescriber will monitor hepatic function closely at repeated intervals for at least several months in patients who discontinue therapy 	1/day			
Viread® powder	NP	 Patient has had a trial and failure, contraindication, or intolerance to 2 preferred agents; OR Patient is 6 years of age or younger and being treated for post-exposure prophylaxis (PEP) 				
Viread® tablets	NP		1/day			



	ANTI-INFECTIVES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL		Qty. Limits	PA Form		
		Antivirals: Hepatitis C Antivirals				
Epclusa® tablet	P	One of the following: Diagnosis of Chronic Hepatitis C, Genotype 1, 2, 4, 5, and 6 Treatment naïve patients with OR without compensated cirrhosis (Child-Pugh A) (Total duration – 12 weeks); OR Diagnosis of Chronic Hepatitis C, Genotype 3 Treatment naïve patient without cirrhosis (Total Duration-12 weeks) Treatment naïve patient with compensated cirrhosis (Child-Pugh A) without baseline NS5A RAS Y93H (Total duration – 12 weeks); Treatment naïve patient with compensated cirrhosis (Child-Pugh A) with baseline NS5A RAS Y93H AND given in combination with ribavirin (Total duration – 12 weeks); Treatment naïve patient with compensated cirrhosis (Child-Pugh A) with baseline NS5A RAS Y93H AND given in combination with ribavirin (Total duration – 12 weeks); Diagnosis of Chronic Hepatitis C, Genotype 1, 2, 3, 4, 5, and 6 Patients with decompensated cirrhosis (Child-Pugh B or C) AND given in combination with ribavirin (Total duration – 12 weeks); OR Patients with decompensated cirrhosis (Child-Pugh B or C) who are ribavirin ineligible (Total duration – 24 weeks); AND If patient has a history of HBV co-infection, prior history with direct acting hepatitis C antivirals, or decompensated cirrhosis, authorization is being requested by or in consultation with a physician specialist with experience in the treatment of hepatitis C infection (e.g., Hepatology, Infectious Disease, or Gastroenterology); AND Patients requiring retreatment of HCV or 2 nd course of therapy with Hepatitis C Direct Acting Antiviral, requires escalation and documentation of ALL the following: Requested HCV treatment regimen is recommended by the AASLD/IDSA guidelines for treatment-experienced patients (HCV Guidance - Treatment Experienced) Current quantitative HCV RNA levels Quantitative HCV RNA levels Quantitative HCV RNA level measured 12 weeks after completion of previous treatment Previous treatment history Genotype testing from current and previous infections; AND Patient has been screened for Hepatitis B prior to treatment with any direct-acting a	1/day	Epclusa P. Form		



ANTI-INFECTIVES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
Harvoni® tablet	P	 One of the following: 	1/day	Harvoni PA Form	
ledipasvir/sofosbuvir	Р	See Harvoni® tablet prior authorization criteria	1/day	Harvoni PA Form	



		ANTI-INFECTIVES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL		Qty. Limits	PA Form
Mavyret [®]	P	Diagnosis of Chronic Hepatitis C, all genotypes Patients with or without cirrhosis: Treatment naïve patients (Total authorization 8 weeks); OR Liver or kidney transplant recipients (Total duration – 12 weeks); OR If patient has a history of HBV co-infection, prior history with direct acting hepatitis C antivirals, or decompensated cirrhosis, authorization is being requested by or in consultation with a physician specialist with experience in the treatment of hepatitis C infection (e.g., Hepatology, Infectious Disease or Gastroenterology); AND Patients requiring retreatment of HCV or 2 nd course of therapy with Hepatitis C Direct Acting Antiviral, requires escalation and documentation of ALL the following: Requested HCV treatment regimen is recommended by the AASLD/IDSA guidelines for treatment-experienced patients (HCV Guidance - Treatment Experienced) Current quantitative HCV RNA levels Quantitative HCV RNA level measured 12 weeks after completion of previous treatment Previous treatment history Genotype testing from current and previous infections; AND Patient has been screened for Hepatitis B prior to treatment with any direct-acting antiviral agent for Chronic Hepatitis C Note: Patients previously treated with one the following are considered treatment-naïve: sofosbuvir+ daclatasvir, peginterferon alfa + ribavirin, paritaprevir/ritonavir/ombitasvir/dasabuvir, and telaprevir or boceprevir + pegylated interferon, ribavirin	3/day	Mavyret PA Form
Mavyret® pellet	Р	See Mavyret® prior authorization criteria; AND Patient is unable to swallow tablets	5/day	
sofosbuvir/ velpatasvir	Р	See Epclusa® tablet prior authorization criteria	1/day	Epclusa PA
Epclusa® pellet	NP	See Epclusa® tablet prior authorization criteria; AND Patient is unable to swallow tablets	150 mg: 1/day 200 mg: 2/day	<u>Form</u>
Harvoni® pellet	NP	See Harvoni® tablet prior authorization criteria; AND Patient is unable to swallow tablets	1 pak/28 days	Harvoni PA Form



	ANTI-INFECTIVES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL		Qty. Limits	PA Form		
Sovaldi® tablets	NP	 One of the following: Diagnosis of Chronic Hepatitis C, genotype 1 or 4 (Total duration −12 weeks) Used in combination with ribavirin and peginterferon affa; OR Patient must have a contraindication or drug-drug interaction with two preferred agents; OR Patients must be treatment naïve to all HCV therapy (including therapies with pegylated interferon or ribavirin); OR If patient has a documented contraindication to interferon; may use in combination with ribavirin alone (Total duration − 24 weeks); AND Diagnosis of Chronic Hepatitis C, genotype 2 (Total duration − 12 weeks): Treatment-naïve and treatment-experienced with or without cirrhosis (Child-Pugh A); AND Requires contraindication or drug-drug interaction with two preferred agents; AND Used in combination with ribavirin Diagnosis of Chronic Hepatitis C, genotype 3 (Total duration − 24 weeks): Treatment-naïve and treatment-experienced without cirrhosis or with compensated cirrhosis (Child-Pugh A); AND Requires contraindication or drug-drug interaction with Mavyret and Epclusa; AND Used in combination with ribavirin If patient has a history of HBV co-infection, prior history with direct acting hepatitis C antivirals, or decompensated cirrhosis, authorization is being requested by or in consultation with a physician specialist with experience in the treatment of hepatitis C infection (e.g., Hepatology, Infectious Disease, or Gastroenterology); AND Patients requiring retreatment of HCV or 2nd course of therapy with Hepatitis C Direct Acting Antiviral, requires escalation and documentation of ALL the following:	1/day	Sovaldi PA Form		
Sovaldi® pellet	NP	See Sovaldi® tablet prior authorization criteria; AND • Patient is unable to swallow tablets	1 pack/28 days			



ANTI-INFECTIVES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
Vosevi®	NP	 Diagnosis of chronic Hepatitis C, genotype 1–6 Sofosbuvir- based treatment failures, with or without compensated cirrhosis (Total duration – 12 weeks); OR Glecaprevir/Pibrentasvir treatment failure with or without compensated cirrhosis (Total duration – 12 weeks); OR Multiple Direct-Acting Antiviral (DAA) treatment failures in combination with weight-based ribavirin (Total duration- 24 weeks); AND If patient has a history of HBV co-infection, prior history with direct acting hepatitis C antivirals, or decompensated cirrhosis, authorization is being requested by or in consultation with a physician specialist with experience in the treatment of hepatitis C infection (e.g., Hepatology, Infectious Disease or Gastroenterology); AND Patients requiring retreatment of HCV or 2nd course of therapy with Hepatitis C Direct Acting Antiviral, requires escalation and documentation of all the following: Requested HCV treatment regimen is recommended by the AASLD/IDSA guidelines for treatment-experienced patients (HCV Guidance - Treatment Experienced) Current quantitative HCV RNA levels Quantitative HCV RNA level measured 12 weeks after completion of previous treatment Previous treatment history Genotype testing from current and previous infections; AND Patient does not have, nor has ever had, decompensated cirrhosis [Child-Pugh score greater than 6 (class B or C)]; AND Patient has been screened for Hepatitis B prior to treatment with any direct-acting antiviral agent for Chronic Hepatitis C 	1/day	Vosevi PA Form	
Zepatier®	NP	 One of the following: Diagnosis of Chronic Hepatitis C, genotype 1a without NS5A polymorphism, genotype 1b, genotype 4 (Total duration – 12 weeks); Patient must have a contraindication or drug-drug interaction with two preferred agents Diagnosis of Chronic Hepatitis C, genotype 1a WITH NS5A polymorphism (Total duration – 16 weeks); Patient must have a contraindication or drug-drug interaction with two preferred agents; OR Diagnosis of Chronic Hepatitis C, genotype 4 (Total duration – 16 weeks) Patient failed prior treatment with peginterferon alfa + ribavirin; AND Patient must have a contraindication or drug-drug interaction with two preferred agents; AND If patient has a history of HBV co-infection, prior history with direct acting hepatitis C antivirals, or decompensated cirrhosis, authorization is being requested by or in consultation with a physician specialist with experience in the treatment of hepatitis C infection (e.g., Hepatology, Infectious Disease or Gastroenterology); AND Patients requiring retreatment of HCV or 2nd course of therapy with Hepatitis C Direct Acting Antiviral, approval requires: Requested HCV treatment regimen is recommended by the AASLD/IDSA guidelines for treatment-experienced patients (HCV Guidance - Treatment Experienced) Current quantitative HCV RNA levels Quantitative HCV RNA level measured 12 weeks after completion of previous treatment Previous treatment history Genotype testing from current and future infections; AND Patient does not have decompensated cirrhosis (defined as a Child-Pugh score > 6 [class B or C]); AND Patient has been screened for Hepatitis B prior to treatment	1/day	Zepatier PA Form	



		ANTI-INFECTIVES				
	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
		Antivirals: Hepatitis C Pegylated Interferons				
Pegasys® syringes	Р	 Chronic Hepatitis C and one of the following: Adult Patients: In combination therapy with other hepatitis C virus drugs for adults with compensated liver disease. Pegasys monotherapy is indicated only if patient has contraindication or significant intolerance to other Hepatitis C drugs Pediatric Patients: In combination with ribavirin for pediatric patients 5 years of age and older with compensated liver disease Chronic Hepatitis B and one of the following: Adult Patients: Treatment of adults with HBeAg-positive and HBeAg-negative chronic hepatitis B (CHB) infection who have compensated liver disease and evidence of viral replication and liver inflammation; OR Pediatric Patients: Treatment of non-cirrhotic pediatric patients 3 years of age and older with HBeAg-positive CHB and evidence of viral replication and elevations in serum alanine aminotransferase (ALT) Note: Prior authorization will be required after 24 weeks of therapy 	4/24 days	General PA Form		
Pegasys® vials	Р	See prior authorization criteria for Pegasys® syringes	4/24 days			
		Antivirals: Herpes Agents, Oral				
famciclovir	Р		125 mg: 20/30 days; 250 mg: 60/30 days; 500 mg: 3/day & 21/Rx			
valacyclovir	Р		500 mg: 60/30 days 1000 mg: 30/Rx	Form		
Sitavig® buccal tabs	NP		2/Rx			
Valtrex®	NP		See valacyclovir			
		Antivirals: HIV Attachment Inhibitors				
Rukobia®	NP	 Initial Criteria: Diagnosis of treatment-experienced multidrug-resistant HIV-1 infection; AND HIV-1 RNA levels ≥ 200 copies/mL; AND Prescriber attests that the patient lacks sufficient treatment options due to resistance, intolerability, contraindication, or other safety concerns to construct a fully suppressive antiretroviral regimen; AND Will not be used with strong cytochrome P450 (CYP)3A inducers Prescribed by, or in consultation with or by an infectious disease specialist Renewal Criteria: Patient demonstrates documented efficacy (e.g., reduced viral load/improved CD4, remain virologically suppressed) 	2/day			



		ANTI-INFECTIVES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Antivirals: HIV Capsid Inhibitors		•
Sunlenca®	Р	 Diagnosis of treatment-experienced multidrug-resistant HIV-1 infection; AND HIV-1 RNA levels ≥ 200 copies/mL; AND Prescriber attests that the patient lacks sufficient treatment options due to resistance, intolerability, contraindication, or other safety concerns to construct a fully suppressive antiretroviral regimen; AND Agent will be used in combination with an optimized antiretroviral regimen; AND Prescriber attests the patient has received or will receive the subcutaneous dose; AND Prescribed by, or in consultation with or by an infectious disease specialist 	1 pack/year	General P Form
		Antivirals: HIV CCR5 Antagonists		
maraviroc tablets	Р	 Diagnosis of CCR5-tropic HIV-1 via a co-receptor tropism; AND Verification that agent will be administered in combination with other antiretroviral agents 	150 mg: 2/day; 300 mg: 4/day	
Selzentry® tablets	NP	 Diagnosis of CCR5-tropic HIV-1 via a co-receptor tropism; AND Verification that agent will be administered in combination with other antiretroviral agents; AND Trial and failure, contraindication, or intolerance to maraviroc tablets 	150 mg: 2/day; 300 mg: 4/day	
Selzentry® solution	NP	 Diagnosis of CCR5-tropic HIV-1 via a co-receptor tropism; AND Verification that agent will be administered in combination with other antiretroviral agents; AND Patient is 11 years of age or younger OR patient is unable to swallow tablets 		
		Antivirals: HIV Fusion Inhibitors	•	
Fuzeon®	Р	 Initial Criteria: Diagnosis of treatment-experienced multidrug-resistant HIV-1 infection; AND HIV-1 RNA levels > 200 copies/mL; AND Prescriber attests that the patient lacks sufficient treatment options due to resistance, intolerability, contraindication, or other safety concerns to construct a fully suppressive antiretroviral regimen; AND Agent will be used in combination with an optimized antiretroviral regimen therapy (ART); AND Prescribed by, or in consultation with or by an infectious disease specialist Renewal Criteria: Patient demonstrates documented efficacy (e.g., reduced viral load/improved CD4, remain virologically suppressed)	1 kit/30 days (2 vials/day)	General PA Form



	ANTI-INFECTIVES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
		Antivirals: HIV Integrase Inhibitors	•	•		
Isentress®	Р		tabs: 2/day; chews: 6/day; granules: 2 packs/day			
Tivicay®	P		2/day			
Tivicay PD®	Р	 Patient is ≤ 6 years of age; OR Patient is unable to swallow solid dosage forms; OR Clinically valid reason why the patient cannot use Tivicay tablets 	3 bottles/30 days	General PA Form		
Isentress® HD	NP	 Verification that agent will be administered in combination with other antiretroviral agents; AND Clinically valid reason why the patient cannot use the preferred agents 	2/day			
Juluca®	NP	 Patient has a diagnosis of HIV; AND Patient does not have any prior history of treatment failure to other HIV agents OR known resistance to the individual components (dolutegravir/rilpivirine); AND Patient is virologically suppressed (HIV-1 RNA < 50 copies/mL) on a current ART regimen for ≥ 6-months 	1/day			
		Antivirals: HIV NNRTIs				
efavirenz	Р		50 mg: 7/day; 200 mg: 2/day; 600 mg: 1/day	General PA Form		
Intelence®	Р	 Patient is treatment-experienced; AND Patient will concomitantly take at least two additional antiretroviral agents; AND Patient has documented non-nucleoside reverse transcriptase inhibitor (NNRTI) resistance 	2/day			
nevirapine	Р		200 mg 2/day; Susp: 40 mL/day			
Pifeltro®	Р		1/day			
etravirine	NP	See Intelence prior authorization criteria	2/day			
nevirapine ER	NP		1/day			
		Antivirals: HIV NRTIs	_ _			
abacavir	Р		tabs: 2/day soln: 30mL/day			
emtricitabine	Р		1/day	=		
lamivudine	P		100 & 300 mg: 1/day; 150 mg: 2/day; soln: 30 mL/day	General PA Form		
stavudine	Р		caps: 2/day; soln: 80 mL/day			



ANTI-INFECTIVES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.				
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
zidovudine	Р		100 mg: 6/day; 300 mg: 2/day; syrup: 60 mL/day	
Emtriva®	NP		caps: 1/day; soln: 24 mL/day	
Epivir®	NP		150 mg: 2/day; 300 mg: 1/day; soln: 30 mL/day	
Retrovir®	NP		100 mg: 6/day; syrup: 60 mL/day	
Ziagen®	NP		tabs: 2/day; soln: 30 mL/day	
		Antivirals: HIV NRTI Combos		
abacavir/ lamivudine	Р		1/day	
Biktarvy®	Р		1/day	
Complera®	Р		1/day	
Delstrigo®	Р		1/day	
Descovy®	Р		1/day	
Dovato®	Р		1/day	
emtricitabine/ tenofovir	Р		1/day	
efavirenz/emtricita- bine/tenofovir	Р		1/day	
Genvoya®	Р		1/day	
lamivudine/ zidovudine	Р		2/day	
Odefsey®	Р		1/day	
Stribild®	Р		1/day]
Symtuza®	Р	 Initial Criteria: Patient has a diagnosis of HIV-1; AND Patient has no known substitutions associated with resistance to darunavir or tenofovir; AND One of the following: Patient is ARV treatment-naïve; OR Patient is ARV treatment-experienced and meets the following requirements: 	1/day	General PA Form



ANTI-INFECTIVES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.				
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Triumeq®	Р		1/day	
Trizivir®	Р		2/day	
Cimduo®	NP		1/day	
efavirenz/lamivudin e/tenofovir	NP		1/day	
Epzicom®	NP		1/day	
Symfi®	NP		1/day	
Symfi® Lo®	NP		1/day	
Triumeq PD®	NP		6/day	
Truvada®	NP		1/day	-
		Antivirals: HIV Pharmacokinetic Enhancers	,	I.
ritonavir tablet	Р			
Norvir® pack	NP	 Patient has a diagnosis of HIV-1; AND The requested will be used in combination with other antiretroviral agents; AND One of the following: Patient is ≤ 18 years of age; OR Clinically valid reason why the preferred ritonavir tablets cannot be used Note: Norvir oral powder is dosed in increments of 100 mg; prescription should not be written for <100 mg increments 	12/day	
Norvir® tablet	NP		12/day	1
Tybost®	NP	 Verification that agent will be administered in combination with Prezista® (darunavir) OR atazanavir; AND Patient has a contraindication or intolerance to ritonavir; AND Patient is not pregnant; AND Patient does not have renal impairment 	1/day	
		Antivirals: HIV Protease Inhibitors		l
atazanavir caps	Р		See Reyataz®	
darunavir	NP		800 mg: 1/day; All other strengths: 2/day; susp: 12 mL/day	
Evotaz®	Р		1/day	
fosamprenavir	Р		4/day	1
lopinavir/ritonavir	Р		soln: 6 mL/day tabs: 1/day	1
Prezcobix®	Р		1/day	-
Prezista® suspension			12 mL/day	1
Reyataz® powder	Р		5/day	General PA
Viracept®	Р		tabs: 4/day	Form



	ANTI-INFECTIVES					
Medication	PDL	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Prior Authorization Criteria	Qty. Limits	PA Form		
Aptivus®	Р	Confirmation that patient has had previous exposure to at least one PI indicated for first line therapy.	caps: 4/day; soln: 10 mL/day			
Kaletra®	NP		soln: 15 mL/day tabs: 6/day			
Prezista® tabs	NP		800 mg: 1/day; All other strengths: 2/day			
Reyataz® caps	NP		300 mg: 1/day; 150, 200 mg: 2/day			
		Antivirals: Influenza				
oseltamivir capsules and suspension	Р		caps: 20/180 days; susp: 300 mL/180 days			
Relenza®	Р		40/180 days			
Tamiflu® capsules and suspension	NP		See oseltamivir			
Xofluza®	NP	 Agent is being used for treatment of influenza OR post-exposure prophylaxis of influenza; AND Treatment is being used for ONE of the following: Acute uncomplicated influenza in patients ≥ 5 years of age who have been symptomatic for no more than 48 hours and who are otherwise healthy Acute uncomplicated influenza in patients ≥ 5 years of age who are at high risk of developing influenza-related complications Post-exposure prophylaxis of influenza in patients > 5 years of age; AND One of the following: Contraindication to both Relenza® and Tamiflu® that is not associated with requested agent Area surveillance data that indicates an oseltamivir resistant strain Recurrent documented influenza in the same flu season that was previously treated with a preferred agent 	2/Rx	Influenza Antiviral PA Form		



		CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwis	e indicated.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
	•	ACE Inhibitors (AEIs)		
ramipril	Р		2/day	
Altace®	NP		2/day	
captopril	NP	Trial and failure, contraindication, or intolerance of TWO preferred agents Note: PA is not required for members 18 years of age and younger		
Epaned®	NP	Patient is unable to swallow solid dosage forms Note: PA is not required for members 8 years of age and younger		
enalapril suspension	NP	See Epaned® prior authorization criteria Note: PA is not required for members 8 years of age and younger		General PA Form
moexipril	NP		7.5 mg: 1/day; 15 mg: 2/day	
perindopril	NP		2 mg, 4 mg: 1/day; 8 mg: 2/day	
Qbrelis® solution	NP	 Patient is unable to swallow solid dosage forms Note: PA is not required for members 7 years of age and younger 		
trandolapril	NP		1/day	
	,	ACEIs + Calcium Channel Blockers		-
benazepril/ amlodipine	Р		5/40 mg: 2/day; All others: 1/day	
Lotrel®	NP	Patient is unable to take the two components separately	5/40 mg: 2/day; All others: 1/day	General PA
Prestalia®	NP	Patient is unable to take the two components separately	1/day	<u>Form</u>
trandolapril/ verapamil	NP	Patient is unable to take the two components separately	1/day	
		ACEIs + Diuretics		
benazepril/HCTZ	NP	Patient is unable to take the two components separately		General PA Form
		Alpha-Beta Blockers	·	
carvedilol	Р		2/day	
carvedilol ER	NP		1/day	General PA
Coreg®	NP		2/day	<u>Form</u>
Coreg CR®	NP		1/day	



CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication **PDL Prior Authorization Criteria Qty. Limits PA Form** Angiotensin II Receptor Antagonists (ARBs) irbesartan Ρ 1/day 25 mg, 100 mg: Ρ 1/day; losartan 50 mg: 2/day Р olmesartan 1/day Ρ valsartan 1/day Atacand® NΡ 1/day NΡ 1/day Avapro® Benicar® NΡ 1/day General PA Form 4 & 32 mg: 1/day; NΡ candesartan 8 mg & 16 mg: 2/day 25 mg, 100 mg: Cozaar® NP 1/day; 50 mg: 2/day NΡ 1/day Diovan® Edarbi™ NP 1/day Micardis® NP 1/day NΡ 1/day telmisartan 80 mL/day valsartan solution NP • Patient is unable to swallow solid dosage forms **ARB + Calcium Channel Blocker** valsartan/ amlodipine Ρ 1/day valsartan/ • Patient is unable to take the components separately 1/day amlodipine/HCTZ Azor® NP 1/day Exforge® NP 1/day **General PA** Exforge HCT® NΡ • Patient is unable to take the components separately 1/day **Form** olmesartan/ NP 1/day amlodipine 20/5/12.5 mg: 2/day; olmesartan/ NP • Patient is unable to take the components separately All others: 1/day amlodipine/HCTZ telmisartan/ NP 1/day amlodipine



		CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherw.	ise indicated.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Tribenzor®	NP	Patient is unable to take the components separately	20/5/12.5 mg: 2/day; All others: 1/day	
		ARB + Diuretic		
irbesartan/HCTZ	Р		1/day	
losartan/HCTZ	Р		1/day	
olmesartan/HCTZ	Р		1/day	
valsartan/HCTZ	Р		1/day	
Atacand HCT®	NP		1/day	
Avalide®	NP		1/day	
Benicar HCT®	NP		1/day	General PA Form
candesartan/HCTZ	NP		1/day	101111
Diovan HCT®	NP		1/day	
Edarbyclor®	NP		1/day	
Hyzaar®	NP		1/day	
Micardis HCT®	NP		1/day	
telmisartan/HCTZ	NP		1/day	
		ARB + Neprilysin Inhibitor		
Entresto® tablets	Р	Diagnosis of chronic heart failure (NYHA Class II-IV)	2/day	General PA
Entresto® sprinkles	NP	 Diagnosis of chronic heart failure (NYHA Class II-IV); AND Clinically valid reason why Entresto tablets cannot be used 	8/day	Form
		Antianginals: Nitrates		
Rectiv®	Р	 Diagnosis of history of anal fissure; AND Patient is a candidate for surgery 		General PA Form
GoNitro® powder	NP	 Clinically valid reason why the preferred agents cannot be used; OR Patient is unable to swallow solid dosage forms or sublingual formulations (e.g., spray, tablet) 		General PA
nitroglycerin spray	NP	 Trial and failure, contraindication, or intolerance of TWO preferred agents; OR Clinically valid reason why the preferred agent cannot be used 		<u>Form</u>



		CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicate	ed.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Antiarrhythmics, Oral		
dofetilide	Р		2/day	
Multaq®	NP	 Not on concurrent Class I or III anti-arrhythmic agent; AND Not hospitalized for exacerbation of heart failure in past 30 days; AND Patient does not have NYHA class IIIb or IV heart failure; AND Trial and failure, contraindication, or intolerance of TWO of the following preferred antiarrhythmic agents: (Note: Requirement is waived if patient has structural heart disease) amiodarone flecainide propafenone sotalol 		General P. Form
Sotylize®	NP	Patient is unable to swallow tablets and capsules Note: PA is not required for patients 8 years of age and younger		
Tikosyn®	NP		2/day	
	•	Anticoagulants, Injectable	•	
enoxaparin	Р		2 injections/day	
fondaparinux	Р		1 injection/day	General PA
Arixtra®	NP		1 injection/day	<u>Form</u>
Lovenox®	NP		2 syringes/day	1
		Anticoagulants, Oral	•	
Eliquis®	Р		2/day	
Pradaxa® caps	Р		2/day	General PA
Xarelto®	Р		2.5 & 15 mg: 2/day 10 & 20 mg: 1/day;	Form Form
dabigatran	NP	Clinically valid reason why the preferred Pradaxa cannot be used	2/day	
Pradaxa® packs	NP	 Patient is unable to swallow sold dosage forms; OR Clinically valid reason why the patient cannot use Pradaxa oral pellets 	2/day	
Savaysa®	NP	 One of the following: Diagnosis of non-valvular atrial fibrillation; AND Documentation that CrCl NOT ≥ 95 mL/min as calculated by Cockcroft-Gault equation Diagnosis of deep vein thrombosis or pulmonary embolism; AND Trial and failure, intolerance, or contraindication to Xarelto® and Pradaxa® 	1/day	General Pa
Xarelto® suspension	NP	Patient is unable to swallow solid dosage forms		



	CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.						
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
	•	Antihypertensives, Miscellaneous					
clonidine weekly patch	Р		0.1, 0.2 mg: 4/28 days; 0.3 mg: pt ≤21: 4/28 days pt >21: 8/28 days				
clonidine 24hr ER	NP		1/day				
minoxidil	NP	 Diagnosis of severe hypertension (symptomatic or associated with target organ damage only); AND Trial and failure, contraindication, or intolerance to TWO of the following: ACEI or ARBs Beta-blocker Calcium channel blockers Methyldopa Clonidine; AND Patient is concomitantly taking a diuretic (e.g., hydrochlorothiazide, chlorthalidone, furosemide, etc.); AND Patient does not have diagnosis of pheochromocytoma (minoxidil may stimulate secretions of catecholamines from the tumor) Note: Minoxidil will not be approved for alopecia 		General PA Form			
Vecamyl®	NP	 Diagnosis of Essential Hypertension or Malignant Hypertension, AND Trial and failure, contraindication, or intolerance to ALL the following: ACE inhibitor-or-ARB Beta blocker Calcium Channel Blocker Clonidine Hydralazine; AND Patient is concomitantly taking a diuretic (e.g., hydrochlorothiazide, chlorthalidone, furosemide) 	10/day				



	CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
	•	Beta Blockers				
metoprolol succinate ER	Р		1/day			
Hemangeol®	NP	 Diagnosis of Infantile Hemangioma; AND Clinically valid reason why the preferred propranolol solution cannot be used 				
InnoPran XL®	NP		80 mg: 2/day; 120 mg: 1/day			
Kapspargo Sprinkle®	NP	 Diagnosis of ONE of the following: Heart Failure or LVEF ≤ 40% Hypertension Angina Pectoris; AND Patient is unable to swallow tablets and capsules 	1/day	General PA Form		
Toprol XL®	NP	 Diagnosis of one of the following: ○ Heart Failure or LVEF ≤ 40% ○ Paroxysmal Atrial Fibrillation 	1/day			
	•	Calcium Channel Blockers (DHP)				
amlodipine	Р		2.5 & 5 mg (1.5/day); 10 mg (1/day)			
nifedipine ER/SA/XL	Р		1/day			
Norliqva®	Р	 Diagnosis of one of the following: Hypertension Chronic stable angina or treatment Vasospastic Angina (Prinzmetal's or Variant Angina) Confirmed or suspected vasospastic angina Angiographically documented Coronary Artery Disease in patients without heart failure and an ejection fraction ≥ 40%; AND One of the following: Patient is unable to swallow solid dosage forms; OR Clinically valid reason why nimodipine capsules cannot be used 	10 mL/day	General PA Form		
isradipine	NP		2.5 mg (2/day); 5 mg (4/day)			
Katerzia®	NP	See Norliqva prior authorization criteria; AND • Trial and failure, contraindication, or intolerance to Norliqva®	10 mL/day			
nimodipine	NP	Diagnosis of subarachnoid hemorrhage (SAH)		General PA		
nisoldipine	NP		1/day	<u>Form</u>		
Norvasc®	NP		See amlodipine			



		CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Nymalize®	NP	Diagnosis of Subarachnoid Hemorrhage; AND One of the following: Patient is unable to swallow solid dosage forms Clinically valid reason why nimodipine capsules cannot be used	120 mL/day	
Procardia® XL	NP		1/day	
Sular®	NP		1/day	
	<u> </u>	Calcium Channel Blockers (Non-DHP)	1	
verapamil ER/SR	Р		1/day	
Cardizem LA®	NP		1/day	General Pa
diltiazem ER caps	NP		1/day	<u> </u>
	·	Cardiac Agents, Miscellaneous		·
ranolazine ER	Р		2/day	
ivabradine	P	 Diagnosis of Congestive Heart Failure (NYHA class II to IV) and documentation of the following: Left ventricular ejection fraction ≤ 35%; AND In sinus rhythm with resting heart rate ≥ 70 beats per minute; AND One of the following: Currently taking a maximum tolerated dose of a beta-blocker and still experiencing heart failure symptoms; OR Patient has a contraindication, adverse reaction, or drug-drug interaction to a beta-blocker; OR Diagnosis of Congestive Heart Failure (NYHA class II to IV) due to dilated cardiomyopathy (DCM); AND Left ventricular ejection fraction ≤ 45%; AND 	2/day	General P Form
Aspruzyo Sprinkle®	NP	 Diagnosis of chronic angina; AND Failure to achieve an adequate response, or intolerance to, at least ONE agent from TWO of the following classes: Beta-blocker (e.g., atenolol, bisoprolol, metoprolol, propranolol) Long-acting nitrate (e.g., nitroglycerin, isosorbide dinitrate, isosorbide mononitrate) Non-DHP calcium channel blocker (e.g., diltiazem, verapamil); AND Patient is unable to swallow solid dosage form 	2/day	



CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Prior Authorization Criteria Qty. Limits PA Form Initial Criteria:** Diagnosis of obstructive hypertrophic cardiomyopathy (HCM); AND · Left ventricular hypertrophy (LVH) confirmed by cardiac imaging (i.e., echocardiography, cardiac MRI); AND Heart failure is classified New York Heart Association (NYHA) class II or III Patient has New York Heart Association (NYHA) Class II or III symptoms (e.g., shortness of breath, chest pain); AND Patient has left ventricular outflow tract (LVOT) peak gradient > 50 mmHg at rest or with provocation; AND • Patient has a left ventricular ejection fraction > 55% (for initiation of therapy); AND Prescribed by or in consultation with a cardiologist; AND **General PA** Camzyos® NP • Trial and failure, contraindication, or intolerance to TWO of the following at a maximally tolerated dose: 1/day Form o Non-vasodilating beta blocker (e.g., bisoprolol, propranolol) Calcium channel blocker (e.g., verapamil, diltiazem) Disopyramide **Renewal Criteria:** • Documentation of positive clinical response to therapy (e.g., NHYA class remains stable or improves improved symptom relief, improvement of LVOT gradient); AND • Patient has a left ventricular ejection fraction > 50%; AND • Prescribed by, or in consultation with, a cardiologist See ivabradine prior authorization criteria; AND **General PA** Corlanor® NΡ 2/day · Clinically valid reason why ivabradine cannot be used Form · Diagnosis of chronic angina; AND • Failure to achieve an adequate response, or intolerance to, at least ONE agent from TWO of the following classes: o Beta-blocker (e.g., atenolol, bisoprolol, metoprolol, propranolol) General PA Ranexa® NΡ 2/day o Long-acting nitrate (e.g., nitroglycerin, isosorbide dinitrate, isosorbide mononitrate) Form Non-DHP calcium channel blocker (e.g., diltiazem, verapamil); AND • Clinically valid reason why the patient cannot use generic ranolazine ER



CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.						
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
Tryvio®	NP	Initial Criteria: (6-month duration) Diagnosis of resistant hypertension; AND Patient is 18 years of age or older; AND Submission of medical records (e.g. chart notes) documenting patient has not achieved target blood pressure (e.g. systolic <130/80) following concurrent maximumly tolerated treatment with at least THREE of the following antihypertensive classes unless contraindicated or intolerance: ACE inhibitors or Angiotensin II receptor blockers (ARBs) Beta blockers Calcium channel blockers Mineralocorticoid Receptor Antagonists Triyvio will be used in combination with at least one other hypertensive drug Renewal Criteria Submission of medical records (e.g. chart notes) documenting a positive clinical response to therapy (e.g. decrease of systolic blood pressure from baseline); AND Patient is receiving concomitant therapy with other hypertensive drugs (documented by claims or medical records)	1/day	General PA Form		
Verquvo®	NP	 Diagnosis of symptomatic chronic heart failure (NYHA class II-IV) with reduced ejection fraction (≤45%); AND Prescribed by, or in consultation with, a cardiologist (initial approval only); AND Patient has had a heart failure hospitalization in the last 6-months OR has received outpatient IV diuretics for heart failure in the last 3 months; AND Patient is 18 years of age or older; AND Patient is currently being treated with an ACEI, ARB, or Entresto; AND Patient is currently being treated with a beta blocker; AND Patient is not pregnant or breastfeeding; AND Female patients of reproductive potential will be counseled to use effective contraception during treatment with therapy and for at least one month after the last dose; AND Patient does not meet any of the following: Concomitant use with another soluble guanylate cyclase (sGC) stimulator (e.g., Adempas) Concomitant use with a PDE-5 inhibitor (e.g., tadalafil, sildenafil) 	1/day	General PA Form		
		Direct Renin Inhibitors				
aliskiren	NP	Patient has a diagnosis of hypertension; AND Trial and failure, contraindication, or intolerance to an agent from at least TWO of the following drug classes: ACEI/ARB Calcium channel blocker Thiazide diuretic	1/day	General PA Form		
Tekturna®	NP	See aliskiren prior authorization criteria	1/day			



		CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicates.	ed.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Tekturna HCT®	NP	 Trial and failure, contraindication, or intolerance to an agent from at least TWO of the following drug classes: ACEI/ARB Calcium channel blocker Thiazide diuretic Patient is unable to take the individual components 	1/day	
	•	Diuretics: Carbonic Anhydrase	<u>, </u>	
dichlorphenamide	NP	See Keveyis criteria; AND • Trial and failure of Keveyis®	2/day	
Keveyis®	NP	Initial Criteria: (2-month duration) Diagnosis of Primary Hypokalemic/Hyperkalemic Periodic Paralysis, and related variants; AND Patient does not have any of the following: Hepatic insufficiency Severe pulmonary disease Hypersensitivity to dichlorphenamide or other sulfonamides Avoid concomitant use with high dose aspirin Renewal Criteria: Clinical documentation that patient has exhibited a reduction in symptoms or attacks; AND Patient's serum potassium and bicarbonate levels are being monitored	2/day	General PA Form
		Diuretics: Loop		
Furoscix®	NP	 Diagnosis of chronic heart failure; AND Patient has signs and symptoms of congestive heart failure due to fluid overload; AND The patient is currently receiving maximal oral diuretic therapy; AND Prescriber attests that additional oral diuretic therapy would be ineffective; AND Prescribed by, or in verbal consultation with, a cardiologist; AND Prescriber has demonstrated appropriate administration use of the On-Body Infusor® 	4 devices/month	
		Diuretics: Potassium Sparing		
CaroSpir®	NP	One of the following: Diagnosis of hypertension Diagnosis of heart failure Diagnosis of edema associated with hepatic cirrhosis; AND Patient is unable to swallow solid dosage forms Note: PA not required for patients < 6 years of age	15 mL/day	General PA Form
eplerenone	NP	One of the following: Patient has a diagnosis of hypertension Patient has a diagnosis of congestive heart failure Patient has a diagnosis of Duchenne muscular dystrophy (DMD); AND Trial and failure, contraindication, or intolerance of spironolactone		General PA Form
Inspra®	NP	See eplerenone prior authorization criteria		



		CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Kerendia®	NP	 Patient is ≥ 18 years of age; AND Diagnosis of chronic kidney disease (CKD) associated with type 2 diabetes (T2D); AND Currently taking the maximum tolerated dose of an ACE inhibitor or ARB, unless contraindicated or intolerant; AND Currently taking an antidiabetic agent (e.g., insulin, metformin, GLP-1 receptor agonist, SGLT2 inhibitor) 	1/day	General PA Form
spironolactone susp	NP	One of the following: Diagnosis of hypertension Diagnosis of heart failure Diagnosis of edema associated with hepatic cirrhosis; AND Patient is unable to swallow solid dosage forms; AND Trial and failure of Carospir®		
		Diuretics: Thiazide and Related		
Diuril®	NP	Patient is unable to swallow solid dosage forms		General PA Form
	•	Hemostatics, Oral		<u> </u>
tranexamic acid	Р	 Diagnosis of acute uterine or cyclic heavy menstrual bleeding; AND Trial and failure, contraindication, or intolerance to ALL the following:		



		CARDIOVASCULAR				
Medication	PDL	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Prior Authorization Criteria	Qty. Limits	PA Form		
	Lipotropics: Antihyperlipidemic Agents					
Praluent®	P	Initial Criteria (6-month duration):	2 pens /28 days	PCSK9 Inhibitors PA Form		



CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.							
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
Repatha®	Р	See Praluent® prior authorization criteria	Repatha: 2/28 days Repatha Pushtronex: 1/28 days	PCSK9 Inhibitors PA Form			
Juxtapid®	NP	 Initial Criteria (6-month duration): Diagnosis of homozygous familial hypercholesterolemia (HoFH) confirmed by one of the following:	5 mg, 10mg: 1/day 20mg: 3/day	General PA Form			



		CARDIOVASCULAR		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Nexletol®	NP	Primary Prevention of Cardiovascular Disease Initial Criteria (6-month duration): Age ≥ 18 years; AND Agent is being use for primary prevention of cardiovascular disease; AND Documented current LDL-C value (within 3 months); AND Patient specific target LDL-C value is provided; AND Patient prevention of concurrent therapy with BOTH the following, unless contraindicated or intolerance: High-intensity statin (atorvastatin/rosuvastatin) Ezetimibe	1/day	General PA Form
Nexlizet®	NP	See Nexeltol® prior authorization criteria	1/day	General PA Form
		Lipotropics: Bile Acid Sequestrant		
colesevelam packets	NP	Patient is unable to swallow solid dosage forms		General PA
Welchol® packets	NP	Patient is unable to swallow solid dosage forms		<u>Form</u>



		CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
	_	Lipotropics: Cholesterol Absorption Inhibitors		
Zetia®	NP	 One of the following: Patient is currently taking a high-intensity statin and has experienced less than anticipated therapeutic response Patient is unable to tolerate lower doses of high-intensity therapy Use in combination with a bile acid sequestrant, fibrate, or niacin will be approved. For requests as monotherapy, recipients must have been intolerant to, or have a contraindication to, a statin 	1/day	General PA Form
		Lipotropics: Combination Agents		
ezetimibe/ simvastatin	NP	 For patients that require ≤45% LDL reduction: 4-week trial and failure of both atorvastatin and simvastatin; OR For patients that require >45% LDL reduction: 4-week trial and failure of atorvastatin 	1/day	
Roszet®	NP	 One of the following: For patients that require ≤45% LDL reduction: 4-week trial and failure of both atorvastatin and rosuvastatin For patients that require >45% LDL reduction: 4-week trial and failure of atorvastatin; AND Clinically valid reason as to why the patient is unable to take components individually 	1/day	General PA Form
Vytorin®	NP	See ezetimibe/simvastatin prior authorization criteria	1/day	
	,	Lipotropics: Fibric Acid Derivatives	•	<u> </u>
fenofibrate caps	NP	 Patient will take fenofibrate concomitantly with a sulfonylurea, thiazolidinedione, repaglinide, or a statin; OR Clinically valid reason why a preferred agent cannot be used (e.g., gemfibrozil, fenofibrate tabs 48, 145, & 160 mg) 		
fenofibrate tabs 40, 54, & 120 mg	NP	See fenofibrate caps prior authorization criteria		
fenofibric acid	NP	See fenofibrate caps prior authorization criteria		
Fenoglide®	NP	See fenofibrate caps prior authorization criteria		
Fibricor®	NP	See fenofibrate caps prior authorization criteria		
Lipofen®	NP	See fenofibrate caps prior authorization criteria		
Lofibra®	NP	See fenofibrate caps prior authorization criteria		
TriCor®	NP	See fenofibrate caps prior authorization criteria		



		CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
	1	Lipotropics: Niacin Derivatives		
niacin ER	Р	One of the following: • Triglycerides > 500 mg/dL; AND ○ Trial and failure. contraindication, or intolerance to BOTH gemfibrozil and fenofibrate; OR • Diagnosis of hyperlipidemia; AND ○ Use in combination with a statin will be approved if the dose of the statin tried is considered sufficient to achieve ≥35% LDL reduction; OR ○ For requests as monotherapy, recipients must have been intolerant to, or have a contraindication to a statin		General PA Form
Niacor®	NP	See niacin ER prior authorization criteria		
		Lipotropics: Omega-3 Fatty Acids		
omega-3 acid ethyl esters	Р	Initial Criteria: Diagnosis of Severe Hypertriglyceridemia (TG level is above 500 mg/dl) Renewal Criteria: Documentation of positive clinical response (e.g., reduction in TG from baseline)	4/day	
icosapent ethyl	NP	Initial Criteria: Diagnosis of Severe Hypertriglyceridemia (TG level is above 500 mg/dl); OR Patient is on maximally tolerated statin AND has triglyceride levels ≥ 135 Renewal Criteria: Documentation of positive clinical response (e.g., reduction in TG from baseline)	0.5 g: 2/day 1 g: 4/day	General PA Form
Lovaza®	NP	Initial Criteria: Diagnosis of Severe Hypertriglyceridemia (TG level is above 500 mg/dl); AND Trial and failure, contraindication, or intolerance to preferred omega-3 acid ethyl esters Renewal Criteria: Documentation of positive clinical response (e.g., reduction in TG from baseline)	4/day	
		Lipotropics: Low and Moderate Intensity Statins		
atorvastatin	Р		1/day	
lovastatin	Р		1/day	General PA
pravastatin	Р		1/day	<u>Form</u>
simvastatin 5, 10, 20, & 40 mg	Р		1/day	
Altoprev®	NP		1/day	
Atorvaliq®	NP	Patient is unable to swallow solid dosage forms	80 mg/day	Gonoral BA
Ezallor Sprinkles®	NP	Patient is unable to swallow solid dosage forms	1/day	General PA Form
Flolipid®	NP	 Patient is 10 to 17 years of age; AND Patient is unable to swallow solid dosage forms 	40 mg/day	<u> </u>
fluvastatin	NP		1/day	
fluvastatin ER	NP		1/day	



		CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Lescol XL®	NP		1/day	
Livalo®	NP		1/day	
pitavastatin	NP		1/day	
Zocor®	NP		1/day	
Zypitamag®	NP		1/day	
		Lipotropics: High Intensity Statins		
atorvastatin	Р		1/day	
rosuvastatin	Р		1/day	<u>High</u>
simvastatin 80 mg	Р	Patient has previously received simvastatin 80 mg for 12 months or longer with no evidence of myopathy	1/day	Potency
Crestor®	NP		1/day	Statin PA
Ezallor Sprinkles®	NP	Patient is unable to swallow solid dosage forms	1/day	<u>Form</u>
Lipitor®	NP		1/day	
		Lipotropics: Statin + Calcium Channel Blocker		
amlodipine/ atorvastatin	NP	Patient is unable to take the 2 components separately	1/day	General PA
Caduet®	NP	Patient is unable to take the 2 components separately	1/day	<u>Form</u>
		Pheochromocytoma Agents		
Demser®	NP	 Documentation of pheochromocytoma diagnosis; AND Trial and failure of an alpha and beta blocker 		
dibenzyline	NP	Diagnosis of pheochromocytoma diagnosis	4/day	General PA
metyrosine	NP	See Demser prior authorization criteria		<u>Form</u>
phenoxybenzamine	NP	See dibenzyline prior authorization criteria	4/day	
		Platelet Inhibitors		
Brilinta®	Р	 History of Myocardial Infarction (MI); OR ACS initial event (USA, NSTEMI or STEMI) or recurrence within previous 12 months; OR Patient has diagnosis of coronary artery disease (CAD) and is at high risk for myocardial infarction (MI) or stroke, OR Acute ischemic stroke or transient ischemic attack (TIA) risk reduction Note: Will NOT be approved if patient is receiving aspirin doses > 100mg/day (includes Rx & OTC aspirin containing products) 		General PA
prasugrel	Р	 Patients has unstable angina, NSTEMI, or STEMI; AND PCI will be performed, or PCI is planned; AND Age < 75 years; AND Weight ≥ 60 kg; AND No history of stroke or TIA 		<u>Form</u>



CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Qty. Limits PA Form Prior Authorization Criteria** Criteria: (2-month duration) Diagnosis of acquired thrombotic thrombocytopenic purpura (aTTP); AND • Used in combination with both of the following: o Plasma exchange until at least 2 days after normalization of the platelet count Cablivi® NP o Immunosuppressive therapy (e.g., corticosteroids); AND • Date Cablivi IV was initiated/administered by a healthcare provider; AND Total treatment duration will be limited to 58 days beyond the last therapeutic plasma exchange; AND • The patient has not experienced more than two recurrences of aTTP while on Cablivi Note: If started as an inpatient hospital regimen and this is continuation of therapy, Cablivi® will be approved • Trial and failure, contraindication, or intolerance to 2 preferred platelet inhibitors with the same indication; AND Durlaza® NΡ 1/day Clinically valid reason why OTC aspirin cannot be used • Patients has unstable angina, NSTEMI, or STEMI; AND PCI has been performed, or PCI is planned; AND • Age < 75 years; **AND** Effient® NP • Weight ≥ 60 kg; AND • No history of stroke or TIA; AND Trial and failure of prasugrel Diagnosis of one of the following: o Ischemic stroke, Transient ischemia of the brain, Previous myocardial infarction, Unstable angina pectoris, Chronic stable angina pectoris; OR Patient has had ONE of the following: Yosprala® NP 1/dav Coronary Artery Bypass Graft (CABG) o Percutaneous Transluminal Coronary Angioplasty (PTCA); AND · Patient meets ALL the following: o Patient is considered a high-risk candidate for aspirin-associated gastric ulcers due to **ONE** of the following: Age ≥ 55, OR Documented history of gastric ulcers; AND O Patient had an inadequate treatment response, or intolerance to use of aspirin and omeprazole separately Pulmonary Arterial Hypertension (PAH) Agents • Diagnosis of Pulmonary arterial hypertension (PAH)/ elevated pulmonary vascular resistance or primary pulmonary Alyq® Ρ hypertension (PPH); OR 2/day • Diagnosis of Congenital heart disease with elevated pulmonary vascular resistance General PA ambrisentan See Alyq® prior authorization criteria 1/day Form bosentan See Alyg® prior authorization criteria 2/day sildenafil See Alyg® prior authorization criteria 3/day



CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Qty. Limits PA Form Prior Authorization Criteria** tadalafil Ρ See Alyg® prior authorization criteria 2/day Diagnosis of pulmonary arterial hypertension (PAH)/ elevated pulmonary vascular resistance or primary pulmonary hypertension (PPH); OR Ρ • Diagnosis of pulmonary hypertension associated with interstitial lung disease (PH-ILD; WHO Group 3) to improve exercise 2.9 mL/day Tyvaso® • Diagnosis of congenital heart disease with elevated pulmonary vascular resistance Ventavis® See Alyg® prior authorization criteria 3 mL/day • Diagnosis of one of the following: o Pulmonary arterial hypertension (PAH)/elevated pulmonary vascular resistance or primary pulmonary hypertension General PA NP Adcirca® 2/day o Congenital heart disease with elevated pulmonary vascular resistance; AND Form Clinically valid reason why the preferred generic cannot be used • One of the following: Diagnosis of pulmonary arterial hypertension (PAH)/ elevated pulmonary vascular resistance or primary pulmonary hypertension (PPH); AND - Trial of ONE preferred agent with persistent signs or symptoms **General PA** Diagnosis of congenital heart disease with elevated pulmonary vascular resistance; AND Adempas® NΡ 3/day - Trial of ONE preferred agent with persistent signs or symptoms Form o Diagnosis of chronic thromboembolic pulmonary hypertension (CTEPH) with one of the following: - Patient has disease that is inoperable; **OR** - Patient has residual post-pulmonary endarterectomy hypertension Note: Use of Adempas[®] is contraindicated in patients also taking PDE-5 inhibitors General PA Letairis® NP | See Adcirca® prior authorization criteria 1/day **Form** Diagnosis of one of the following: o Pulmonary arterial hypertension (PAH)/ elevated pulmonary vascular resistance or primary pulmonary hypertension o Congenital heart disease with elevated pulmonary vascular resistance; AND **General PA** • One of the following: Ligrev® 240mg/day Form Patient is unable to swallow tablets Patient is < 6 years of age Clinically valid reason why a preferred tablet formulation cannot be used • Diagnosis of one of the following: o Pulmonary arterial hypertension (PAH)/ elevated pulmonary vascular resistance or primary pulmonary hypertension NP Opsumit® 1/day o Congenital heart disease with elevated pulmonary vascular resistance; AND • Trial of one preferred agent with persistent signs or symptoms **General PA** • Diagnosis of Pulmonary arterial hypertension (PAH)/ elevated pulmonary vascular resistance or primary pulmonary Form Opsynvi® NP hypertension: AND 1/day Clinically valid reason as to why the patient is unable to take components of Opsynvi individually Orenitram® NP See Opsumit® prior authorization criteria 3/day Revatio® tab See Adcirca® prior authorization criteria 3/dav

June 1, 2025



CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Qty. Limits PA Form Prior Authorization Criteria** 6 ml/day; Revatio® suspension NP | See Ligrev® prior authorization criteria Max day supply=60 6 ml/day; sildenafil suspension NP | See Ligrev® prior authorization criteria Max day supply=60 Tadlig® NP | See Ligrev® prior authorization criteria 10mL/dav Diagnosis of one of the following: o Pulmonary arterial hypertension (PAH)/ elevated pulmonary vascular resistance or primary pulmonary hypertension Tracleer® soluble tabs 2.9 mL/day o Diagnosis of congenital heart disease with elevated pulmonary vascular resistance; AND • Patient is unable to swallow solid dosage forms Tracleer® tabs NP | See Adcirca® prior authorization criteria 2/day • Diagnosis of one of the following: Single cartridges: Pulmonary arterial hypertension (PAH)/elevated pulmonary vascular resistance or primary pulmonary hypertension 4/day; Combo NP Tyvaso DPI® o Pulmonary hypertension associated with interstitial lung disease; AND cartridges: 8/day; Clinically valid reason why the preferred Tyvaso inhalation solution cannot be used Kits: 2/year Tabs: 2 /day; Uptravi® NP | See Opsumit® prior authorization criteria Pack: 1 /Rx • Patient is 18 years of age or older; AND • Diagnosis of Pulmonary arterial hypertension (PAH)/ elevated pulmonary vascular resistance or primary pulmonary hypertension; AND Patient is Functional Class II or III; AND Winrevair® NP • Trial and failure of 1 agent with persistent signs and symptoms from TWO different classes for PAH such as: 1 kit/21 days o Endothelin receptor antagonist (e.g. ambrisentan, bosentan) Phosphodiesterade-5 inhibitors (e.g. sildenafil, tadalafil) o Prostacyclin analogue or receptor agonist (e.g., treprostinil, epoprostenol, Uptravi, Ventavis); AND • Winrevair will be used in combination with other PAH therapies **Pulmonary Fibrosis Agents** • Diagnosis of one of the following: Idiopathic pulmonary fibrosis Interstitial Lung Disease Associated with Systemic Sclerosis- associated interstitial lung disease (SSc-ILD) Ofev® 2/day o Chronic Fibrosing Interstitial Lunch Diseases (ILDs) with a progressive phenotype (at least 10% of the lungs show presence of fibrotic ILD); AND **General PA** Prescribed by, or in consultation with, a pulmonologist (initial approval only) Form • Patient has a diagnosis of idiopathic pulmonary fibrosis; AND 534, 801 mg: 3/day; pirfenidone tablets • Prescribed by, or in consultation with, a pulmonologist (initial approval only) 267 mg: 9/day • Patient has a diagnosis of idiopathic pulmonary fibrosis; AND 3/day: 801 mg: 3/day Esbriet® NP Prescribed by, or in consultation with, a pulmonologist (initial approval only); AND 9/day: 267 mg



• Clinically valid reason as to why the preferred pirfenidone cannot be used

		CARDIOVASCULAR		
Medication	PDL	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Prior Authorization Criteria	Qty. Limits	PA Form
pirfenidone capsules	NP	See Esbriet prior authorization criteria	9/day: 267 mg	
		Thrombopoietin Agonists, Orals		<u> </u>
Promacta® tabs	NP	 Diagnosis of persistent or chronic thrombocytopenia purpura (ITP) in patients ≥1 year of age; AND Documentation of failure or insufficient response to adequate treatment with corticosteroids AND immunoglobulins, OR ITP related splenectomy; AND Documentation that patient's thrombocytopenia and clinical condition puts the patient at increased risk of bleeding; OR Diagnosis of thrombocytopenia in patient with chronic hepatitis C; AND Patient receiving (or planning to initiate) interferon-based anti-viral therapy; OR Diagnosis of severe aplastic anemia in patients 2 years of age or older; AND Patient will use in combination with standard immunosuppressive therapy for first-line treatment; OR Diagnosis of severe aplastic anemia; AND Patient has tried and failed or has intolerance to immunosuppressive therapy 	1/day	
Doptelet®	NP	 Patient is ≥ 18 years old; AND Patient must have a diagnosis of thrombocytopenia and meet one of the following: Chronic liver disease AND scheduled to undergo a medical procedure; AND Patient is scheduled to take the requested agent 10 to 13 days prior to the procedure, with the procedure occurring 5 to 8 days following the last dose of Doptelet®; OR Prescribed dose is according to baseline platelet count (10 tabs per 5 days ≥ 40 x 10⁹/L or 15 tabs per 5 days for platelets < 40 x 10⁹/L) PA Duration: single course of treatment per scheduled procedure, QL=15 per treatment Chronic Immune Thrombocytopenia (ITP); AND Patient has had an insufficient response to a previous treatment; AND Patient has a platelet count of < 50 x 109/L PA Duration: 1 year, QL= 2/day 	See criteria	
Mulpleta®	NP	 Criteria: (PA duration – single course of treatment per scheduled procedure): Patient is ≥ 18 years old; AND Patient has a diagnosis of Chronic Liver Disease (CLD); AND Patient does NOT have Child-Pugh class C liver disease, absence of hepatopetal blood flow, a prothrombotic condition other than CLD nor a history of splenectomy, partial splenic embolization, or thrombosis; AND Patient has a platelet count of < 50 x 10⁹/L; AND Patient has an upcoming invasive procedure scheduled; AND Patient is scheduled to take the requested agent 8 to 14 days prior to the procedure, with the procedure occurring 2 to 8 days following the last dose of Mulpleta®; AND Patient is NOT scheduled for a thoracotomy, laparotomy, open-heart surgery, craniotomy, or organ resection. 	7 tabs/Rx	<u>General I</u> <u>Form</u>
Promacta® suspension	NP	See Promacta® prior authorization criteria • Patient is unable to swallow solid dosage forms	4 packets/day	1



		CARDIOVASCULAR Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Tavalisse®	NP	Initial Criteria: Patient has a diagnosis of chronic immune thrombocytopenia; AND Trial and failure (platelet count ≥ 50 x 10°/L not achieved) of ONE of the following: Corticosteroids Thrombopoietin receptor antagonists (e.g., Promacta) Splenectomy Azathioprine (Azasan, Imuran), cyclosporine (Neoral, Sandimmune), cyclophosphamide (Cytoxan), mycophenolate mofetil (CellCept), danazol, or rituximab (Rituxan); AND Patient is not on concomitant therapy with a strong CYP3A4 inducer; AND Patient has received a baseline and will receive ongoing routine monitoring that includes: Neutropenia (measure ANC monthly) Hepatotoxicity (measure LFTs monthly) Hypertension (measure blood pressure every 2 weeks until stable dose established, then monthly) Renewal Criteria: Patient has laboratory values documenting platelet response to therapy (platelet count ≥ 50 x 10°/L; AND Patient has not experienced severe adverse effect as a result of fostamatinib therapy	2/day	General PA Form
droxidopa	NP	 Diagnosis of symptomatic neurogenic orthostatic hypotension secondary to primary autonomic failure, dopamine beta-hydroxylase deficiency, or non-diabetic autonomic neuropathy; AND Trial and failure, contraindication, or intolerance to midodrine OR fludrocortisone 	100 & 200 mg: 3/day 300 mg: 6/day	General PA
Northera®	NP	See droxidopa prior authorization criteria	100 & 200 mg: 3/day 300 mg: 6/day	Form Form



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication **PDL Prior Authorization Criteria Qty. Limits PA Form** Agents for Neuropathic Pain and Fibromyalgia Note: The maximum daily dose limit for gabapentin, including all formulations and Brand products, is 3,600 mg. duloxetine 20,30, & **SNRI PA** Р 2/day 60 mg Form 100 mg: 6/day; gabapentin capsules 300 mg: 12/day; 400 mg: 9/day 1/dav · Diagnosis of post-herpetic neuralgia; OR Horizant® Ρ Max daily gabapentin • Diagnosis of Restless Leg Syndrome General PA dose: 3600 mg Form lidocaine 5% patch · Diagnosis of post-herpetic neuralgia 2/day • Diagnosis of neuropathic pain; OR • Diagnosis of postherpetic neuralgia; OR pregabalin capsules • Diagnosis of fibromyalgia; OR · Diagnosis of seizure disorder See pregabalin capsules prior authorization criteria: AND • Patient is less than 12 years of age; OR pregabalin solution • Patient is unable to swallow solid oral dosage forms **SNRI PA** Cymbalta® NP 2/day Form duloxetine 40 mg NP Clinically valid reason as to why the preferred duloxetine strengths (20 mg, 30 mg, 60 mg) cannot be used 2/day • One of the following: gabapentin solution NP o Patient is less than 12 years of age; OR 72 mL/day Inability to swallow solid oral dosage forms 600 mg: 6/day; gabapentin tablets NP Documented allergy or contraindication to an inactive ingredient in the capsules that is NOT present in the tablets 800 mg: 4.5/day Gralise® NP • Clinically valid reason why the preferred gabapentin agents cannot be used 3/day See pregabalin capsules prior authorization criteria; AND Lyrica® capsules • Trial and failure of preferred pregabalin capsules **General PA** See pregabalin solution prior authorization criteria; AND Form Lyrica® solution Trial and failure of preferred pregabalin solution • Diagnosis of postherpetic neuralgia OR neuropathic pain associated with-diabetic peripheral neuropathy; AND 82.5 mg & 165 mg: Trial and failure, contraindication, or intolerance to a tricyclic antidepressant OR gabapentin; AND Lyrica® CR NP 1/day Provider must provide peer-reviewed medical literature documenting why the requested drug for the requested indication 330 mg: 2/day is the only appropriate choice versus immediate-release pregabalin 100 mg: 6/day; Neurontin[®] capsules NP 300 mg: 12/day; 400 mg: 9/day



		CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless other	herwise indicated.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Neurontin® solution	NP	See gabapentin solution prior authorization criteria	72 mL/day	
Neurontin® tablets	NP		600 mg: 6/day; 800 mg: 4.5/day	
pregabalin CR	NP	See Lyrica® CR prior authorization criteria	82.5 mg & 165 mg: 1/day 330 mg: 2/day	General P
Savella®	NP	 Patient has a diagnosis of fibromyalgia accompanied by fatigue; AND Patient is 18 years of age or older; AND Patient MUST have tried and failed, or have contraindication, or intolerance to duloxetine 	2/day	<u>Form</u>
		Agents for Restless Leg Syndrome (RLS)	<u>,</u>	
pramipexole	Р		3/day	
Horizant®	Р	 Diagnosis of Restless Leg Syndrome; OR Diagnosis of post-herpetic neuralgia 	1/day Max daily gabapentin dose: 3600 mg	General PA
Neupro®	NP	 Diagnosis of Parkinson's Disease or Restless Leg Syndrome, AND Trial and failure, contraindication, or intolerance to Horizant, pramipexole, and ropinirole, OR Inability to swallow 		<u>Form</u>
	,	Alzheimer's: Cholinesterase Inhibitors	·	
donepezil (excluding 23 mg)	Р		1/day	
donepezil ODT	Р	 Patient is unable to swallow; OR Unable to absorb medications through the GI tract 	1/day	
Exelon®	Р		1/day	
Adlarity®	NP		4 patch/month	General PA
Aricept®	NP		1/day	Form
Aricept® 23 mg	NP	Patient has been established (at least 3 months) on therapy with Aricept 10mg daily	1/day	101111
Aricept® ODT	NP	 Patient is unable to swallow; OR Unable to absorb medications through the GI tract 	1/day	
donepezil 23 mg	NP	Patient has been established (at least 3 months) on therapy with donepezil 10mg daily	1/day	
galantamine ER	NP		1/day	
rivastigmine patch	NP		1/day	

June 1, 2025



		CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated		
Medication	PDL		Qty. Limits	PA Form
	1	Alzheimer's: NMDA Receptor Agent		
memantine tablets	Р		5, 10 mg: 2/day; Titration Pack: 1/Rx	
memantine ER	NP	Diagnosis of moderate to severe Alzheimer's disease	1/day	
memantine solution	NP	Diagnosis of moderate to severe Alzheimer's disease	10mL/day	General PA
Namzaric®	NP	 Diagnosis of moderate to severe dementia associated with Alzheimer's disease; AND Concomitantly taking donepezil and memantine (immediate release or extended release) [≥10mg/day on both agents]; AND Clinical reason why recipient is unable to take the components individually 	1/day	- <u>Form</u>
Zunvey®	NP	,, ,,	2/day	
,		Analeptics	. ,	
caffeine citrate soln	NP	 Criteria (2-month duration) Diagnosis of apnea in premature infants (born between 28 and <33 weeks gestational age); AND Patient is continuing therapy from an inpatient hospital stay (to facilitate transition to outpatient for completion of therapy); AND Infant does not have renal impairment, hepatic impairment, or cardiovascular disease; AND Prescriber must attest that they are aware of the risks of fatal necrotizing enterocolitis in premature infants and will monitor patient for efficacy and to avoid serious toxicity; AND Prescribed by, or in consultation with a board-certified neonatologist 		General PA Form
	1	Antiparkinson Agents: Adenosine Antagonists	T	1
Nourianz [®]	NP	Initial Criteria: (6-month duration) Diagnosis of Parkinson's disease; AND Patient is experiencing "off" episode; AND Patient is 18 years of age or older; AND Patient is currently being treated with a stable dosage of levodopa/carbidopa; AND Prescriber advises women of childbearing potential to use contraception during treatment; AND Prescriber agrees to monitor the following: Patients with moderate hepatic impairment (Child-Pugh B) for adverse reactions Exacerbation of pre-existing dyskinesia Presence of hallucinations/psychotic behavior Presence of impulse control/compulsive behaviors; AND Trial and failure, intolerance, or contraindication to ONE agent in TWO different antiparkinson classes (e.g., Dopamine Agents, Decarboxylase Inhibitors, COMT Inhibitors, MAO-B inhibitors, NMDA Antagonists) Renewal Criteria: Patient is currently being treated with levodopa/carbidopa; AND Patient has a positive clinical response to therapy (e.g., reduction in number or total daily hours of "off" episodes, increase "on" time without troublesome dyskinesia)	1/day	General PA Form



		CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL		Qty. Limits	PA Form
		Antiparkinson Agents: COMT Inhibitors		
Ongentys®	NP	 Initial Criteria: (6-month duration) Diagnosis of Parkinson's disease; AND Patient is experiencing "off" episodes; AND Patient is currently being treated with a stable dose of carbidopa/levodopa; AND Trial and failure, intolerance, or contraindication to ONE agent in TWO different antiparkinson classes (e.g., dopamine agents, decarboxylase inhibitors, COMT inhibitors, MAO-B inhibitors, NMDA antagonists); AND Will not be taken concomitantly with a non-selective monoamine oxidase inhibitor (MAOI); AND Patient does not have a history of pheochromocytoma, paraganglioma, or other catecholamine secreting neoplasms Renewal Criteria: Patient is currently being treated with levodopa/carbidopa; AND Patient has a positive clinical response to therapy (e.g., reduction in number or total daily hours of "off" episodes, increase "on" time without troublesome dyskinesia) 	1/day	General P Form
	•	Antiparkinson Agents: Dopamine Agents		
pramipexole	Р		3/day	
Apokyn®	NP	 Patient has a diagnosis of Parkinson's disease; AND Patient is experiencing acute, intermittent treatment of "off" episodes; AND Must be 18 years of age or older; AND Patient is currently being treated with a carbidopa/levodopa agent; AND Patient has had a trial and failure, contraindication, or intolerance of TWO of the following preferred adjunct drugs prescribed in combination with levodopa/carbidopa, each from different classes: MAO-B inhibitor: selegiline COMT inhibitor: entacapone, carbidopa/levodopa/entacapone, Stalevo Dopamine agonist: pramipexole, ropinirole; AND Patient must not meet any of the following: Patient is on concomitant 5HT3 antagonist Patient has a sensitivity to sulfites 		General P Form
apomorphine injection	NP	See prior authorization criteria for Apokyn®		
Neupro®	NP	 Diagnosis of Parkinson's Disease OR Restless Leg Syndrome, AND Trial and failure, contraindication, or intolerance to BOTH pramipexole AND ropinirole, OR Inability to swallow 		
pramipexole ER	NP		1/day	
	•	Antiparkinson Agents: Levodopa Combinations		
Dhivy®	NP	Clinically valid reason as to why all the preferred carbidopa/levodopa agents cannot be used		
Inbrija®		Initial Criteria: (6-month duration)	60 blisters/month	General PA



		CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
	NP	 Diagnosis of Parkinson's disease; AND Experiencing "off" episodes; AND Patient is currently being treated with a stable dose of carbidopa/levodopa; AND Trial and failure, intolerance, or contraindication to ONE agent in TWO different antiparkinson classes (e.g., dopamine agents, decarboxylase inhibitors, COMT inhibitors, MAO-B inhibitors, NMDA antagonists); AND Will not be taken concomitantly with a non-selective monoamine oxidase inhibitor (MAOI); AND Patient does not have asthma, COPD, or other chronic lung disease Renewal Criteria: Patient is currently being treated with levodopa/carbidopa; AND Patient has a positive clinical response to therapy (e.g., reduction in number or total daily hours of "off" episodes, increase "on" time without troublesome dyskinesia) 		Form
	•	Antiparkinson Agents: MAOI-Bs		
Xadago®	NP		1/day	General PA
Zelapar®	NP	 Inability to swallow solid dosage forms; OR Clinically valid reason why the preferred selegiline formulation cannot be used 		Form Form
	•	Antiparkinson Agents: NMDA Antagonists		
Gocovri®	NP	 Initial Criteria: One of the following:	68.5 mg: 1/day; 137 mg: 2/day	
Osmolex® ER tabs	NP	Initial Criteria: One of the following: Diagnosis of Parkinson's disease Treatment of drug-induced extrapyramidal reactions; AND Patient does not have end-stage renal disease (creatinine clearance below 15 mL/min/1.73 m2); AND Patient has had an adequate trial of or is intolerant to amantadine IR (capsules) Renewal Criteria: Documentation of decreased Parkinson's disease symptoms OR decreased extrapyramidal effects	193 mg & 258 mg: 1/day; 129 mg: 2/day	



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL Prior Authorization Criteria Qty. Limits PA Form

CLASS PRIOR AUTHORIZATION CRITERIA FOR PATIENT'S WITH INTELLECTUAL AND DEVELOPMENTAL DISABILITIES (I/DD):

Anti-anxiety agents prescribed for disorders related to Intellectual and Developmental Disabilities will be approved if ONE of the following is met:

- Prescribed bay a Gold Card prescriber; OR
- Mental health assessment applicable to behavioral symptoms for which the medication is being prescribed; AND
 - Underlying physical condition such as pain, discomfort, and environmental issues have been evaluated, treated, and behavioral symptoms persist; AND
 - Non-pharmacological interventions have been tried and behavioral symptoms persist (e.g., behavior plan, crisis intervention, and stabilization strategies or training) and training and support have been provided to family or other caregivers
- Short-term therapy (less than 90 days) has been prescribed; AND
 - o Behavioral symptoms are significant enough to place the person at potential risk or needing higher level of care or loss of community placement; OR
 - Continuation of existing therapy to address serious and ongoing behavioral symptoms while other prior authorization criteria are met; AND
 - o Efficacy and potential side effects to be monitored; AND
 - o Need for requested medication will be evaluated once other non-pharmacological interventions have been tried

Note the following:

- Drug specific step therapy, NP criteria and Brand Medically Necessary Criteria will still apply to this patient population.
- The I/DD Worksheet can be found at: I/DD Prior Authorization Form

Anti-Anxiety and Anti-Panic Agents				
alprazolam tabs	Р	 Diagnosis of one of the following: Anxiety disorder Panic disorder with or without agoraphobia; AND Documented trial (at least 3 weeks) of non-pharmacological therapies (e.g., cognitive behavioral therapy, worry exposure, applied relaxation, muscle relaxation, short-term psychodynamic psychotherapy, mindfulness-based therapy); AND Trial and failure, contraindication, or intolerance to therapy with TWO of the following: SSRI (minimum trial duration of 4 weeks) SNRI (minimum trial duration of 4 weeks) Buspirone; AND Due to increased risk of toxicity, patient should not be pregnant OR concurrently taking CNS stimulants, opiates, carisoprodol, meprobamate, or barbiturates; AND Due to increased risk of dependency, patient does not have a history of alcohol or drug dependence/abuse; AND Prescriber has checked the Tennessee Controlled Substance Monitoring Database (CSMD) on the date of the request for concomitant controlled substance use 	3/day	Anti-anxiety PA Form
alprazolam ER tabs	Р	See alprazolam tablets prior authorization criteria	2/day	
buspirone	Р		30 mg: 2/day; All other strengths: 3/day	General PA Form



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Qty. Limits Prior Authorization Criteria PA Form** • Diagnosis of acute alcohol withdrawal syndrome; OR · Diagnosis of anxiety disorder; AND o Documented trial (at least 3 weeks) of non-pharmacological therapies (e.g., cognitive behavioral therapy, worry exposure, applied relaxation, muscle relaxation, mindfulness-based therapy); AND o Trial and failure, contraindication, or intolerance to therapy with TWO of the following: SSRI (minimum trial duration of 4 weeks) chlordiazepoxide Ρ 4/day SNRI (minimum trial duration of 4 weeks) - Buspirone; AND Due to increased risk of dependency, patient does not have a history of alcohol or drug dependence/abuse; AND Due to increased risk of toxicity, patient should not be pregnant OR concurrently taking CNS stimulants, opiates, carisoprodol, meprobamate, or barbiturates; AND Prescriber has checked the Tennessee Controlled Substance Monitoring Database (CSMD) on the date of the request for concomitant controlled substance use Diagnosis of seizure disorder; OR Diagnosis of panic disorder; AND Trial and failure, contraindication, or intolerance to therapy with TWO of the following: - SSRI (minimum trial duration of 4 weeks) - SNRI (minimum trial duration of 4 weeks) clonazepam Ρ - Buspirone; AND 3/day • Due to increased risk of toxicity, patient should not be pregnant OR concurrently taking CNS stimulants, opiates, Anti-anxiety carisoprodol, meprobamate, or barbiturates; AND **PA Form** · Due to increased risk of dependency, patient does not have a history of alcohol or drug dependence/abuse Prescriber has checked the Tennessee Controlled Substance Monitoring Database (CSMD) on the date of the request for concomitant controlled substance use Diagnosis of acute alcohol withdrawal syndrome; OR · Diagnosis of seizure disorder; AND o Must be used in conjunction with another anticonvulsant; **OR** · Diagnosis of anxiety disorder; AND o Documented trial (at least 3 weeks) of non-pharmacological therapies (e.g., cognitive behavioral therapy, worry exposure, applied relaxation, muscle relaxation, mindfulness-based therapy); AND o Trial and failure, contraindication, or intolerance to therapy with TWO of the following: SSRI (minimum trial duration of 4 weeks) Ρ clorazepate 3/day SNRI (minimum trial duration of 4 weeks) Buspirone; AND Due to increased risk of toxicity, patient should not be pregnant OR concurrently taking CNS stimulants, opiates, carisoprodol/meprobamate, or barbiturates; AND Due to increased risk of dependency, patient does not have a history of alcohol or drug dependence/abuse (does not apply to diagnosis of acute alcohol withdrawal syndrome); AND Prescriber has checked the Tennessee Controlled Substance Monitoring Database (CSMD) on the date of the request for



concomitant controlled substance use

		CENTRAL NERVOUS SYSTEM		
B.d. adioaking	DDI	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated		DA Forms
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
	ı	Anti-anxiety Agents (continued)	1	
diazepam tablets, solution, concentrate	Р	 Diagnosis of acute alcohol withdrawal syndrome; OR Diagnosis of seizure disorder; AND Must be used in conjunction with another anticonvulsant; OR Diagnosis of muscle spasms; AND Patient has tried and failed at least TWO preferred skeletal muscle relaxants; OR Diagnosis of anxiety disorder; AND Documented trial (at least 3 weeks) of non-pharmacological therapies (e.g., cognitive behavioral therapy, worry exposure, applied relaxation, muscle relaxation, mindfulness-based therapy); AND Trial and failure, contraindication, or intolerance to therapy with TWO of the following: SSRI (minimum trial duration of 4 weeks) SNRI (minimum trial duration of 4 weeks) Buspirone; AND Due to increased risk of toxicity, patient should not be pregnant OR concurrently taking CNS stimulants, opiates, carisoprodol, meprobamate, or barbiturates; AND Due to increased risk of dependency, patient does not have a history of alcohol or drug dependence/abuse (does not apply to diagnosis of acute alcohol withdrawal syndrome); AND Prescriber has checked the Tennessee Controlled Substance Monitoring Database (CSMD) on the date of the request for concomitant controlled substance use 	tabs: 4/day soln: 10 mL/day concentrate: 2 mL/day	Anti-anxiety
lorazepam tablets and concentrate	Р	 Patient is < 1 year of age and completing taper following inpatient hospital use for Neonatal Withdrawal symptoms; OR Diagnosis of anxiety disorder; AND Documented trial (at least 3 weeks) of non-pharmacological therapies (e.g., cognitive behavioral therapy, worry exposure, applied relaxation, muscle relaxation, mindfulness-based therapy); AND Trial and failure, contraindication, or intolerance to therapy with TWO of the following:	tabs: 3/day concentrate: 3 mL/day	PA Form
Xanax®	Р	See alprazolam tablets prior authorization criteria	3/day	
Xanax® XR	Р	See alprazolam tablets prior authorization criteria	2/day	
alprazolam ODT	NP	 See alprazolam prior authorization criteria; AND Patient is unable to swallow solid dosage forms or unable to absorb medications through the GI tract; AND Trial and failure, contraindication, or intolerance to the BOTH preferred concentrate solutions 	3/day	
alprazolam concentrate	NP	See alprazolam prior authorization criteria; AND • Patient is unable to swallow solid dosage forms or unable to absorb medications through the GI tract; AND • Patient must have a trial and failure, contraindication, or intolerance to the BOTH preferred concentrate solutions	6 mL/day	Anti-anxiety PA Form



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.				
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Ativan®	NP	See lorazepam prior authorization criteria; AND • Clinically valid reason as to why the preferred lorazepam tablets or concentrate cannot be used	3/day	
Loreev XR®	NP	See lorazepam prior authorization criteria; AND • Clinically valid reason as to why the preferred lorazepam tablets or concentrate cannot be used	1/day	
meprobamate	NP	See alprazolam prior authorization criteria; AND • Trial and failure, contraindication, or intolerance of TWO preferred agents		
oxazepam	NP	See chlordiazepoxide prior authorization criteria; AND Trial and failure, contraindication, or intolerance of TWO preferred agents	4/day	
Valium®	NP	 Diagnosis of acute alcohol withdrawal syndrome; OR Diagnosis of seizure disorder; AND Must be used in conjunction with another anticonvulsant; AND Trial and failure of the following preferred agents:	3/day	
Aptiom [®]	Р	Use as monotherapy for partial onset seizures and trial and failure with ONE preferred anticonvulsant with the same indication; OR		General PA
clobazam tablets	Р	 Use as adjunctive therapy for partial onset seizures when used in combination with at least ONE other anticonvulsant. Diagnosis of Lennox-Gastaut Syndrome; AND Used as adjunct therapy with at least one other anticonvulsant 		Form



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Prior Authorization Criteria Qty. Limits PA Form** · Diagnosis of seizure disorder; OR Diagnosis of panic disorder; AND o Trial and failure, contraindication, or intolerance to therapy with TWO of the following: - SSRI (minimum trial duration of 4 weeks) SNRI (minimum trial duration of 4 weeks) Anti-anxiety Buspirone: AND 3/day clonazepam PA Form • Due to increased risk of toxicity, patient should not be pregnant OR concurrently taking CNS stimulants, opiates, carisoprodol, meprobamate, or barbiturates; AND Due to increased risk of dependency, patient does not have a history of alcohol or drug dependence/abuse; AND • Prescriber has checked the Tennessee Controlled Substance Database on the date of the request for concomitant controlled substance use • Prior Authorization will not be required for patients less than 21 years of age. diazepam rectal gel 2 packs/30 days • Will be approved for patients 21 years of age and older with a Diagnosis of Seizure Disorder or Epilepsy. Initial Criteria: • Diagnosis of one of the following: Dravet Syndrome (DS) Lennox-Gastaut Syndrome (LGS) Tuberous sclerosis complex (TSC) Epidiolex® Ρ o Treatment-Refractory Epilepsy; AND Trial of 2 anticonvulsants within the past 12 months (documented by claims); AND Epidiolex will be used as adjunct therapy with > 1 anticonvulsant (documented by claims) Renewal Criteria Epidiolex will be used as adjunct therapy with ≥ 1 anticonvulsant (documented by claims) 100 mg: 6/day 300 mg: 12/day gabapentin capsules 400 mg: 9/day Max daily gabapentin dose: 3600 mg • Inability to swallow solid oral dosage forms, AND 72 mL/day gabapentin solution Ρ o Patient and caregiver are unable to open capsule and empty contents in food or drink; OR Max daily gabapentin dose: 3600 mg • Patient is < 12 years of age • Use as monotherapy for partial onset seizures requires trial and failure with at least ONE other preferred anticonvulsant for the same indication; OR General PA Use as adjunctive therapy for partial onset seizures when used in combination with at least ONE other anticonvulsant: OR lacosamide tablets **Form** • Used as adjunctive therapy in the treatment of primary generalized tonic-clonic (PGTC) seizures in patients 4 years of age and older



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Prior Authorization Criteria Qty. Limits PA Form** Initial Criteria (6-month duration): Diagnosis of intermittent, stereotypic episodes of frequent seizure activity (e.g., seizure clusters, acute repetitive seizures) that are distinct from a patient's usual seizure pattern; AND • Patient is 2 to 5 years of age; AND Prescribed by, or in consultation with, a neurologist; AND • Patient is on a stable antiepileptic regimen; AND • Prescriber has counseled patient on the following: Risks if combined with opioids or benzodiazepines o Identification of a seizure cluster **General PA** o Proper administration Libervant® 10 doses/30 days o When to seek emergency medical treatment; AND **Form** Patient is not using moderate or strong CYP2C19 and CYP 3A4 inhibitors or, if unavoidable, prescriber will monitor toxicity risk during concomitant use; AND Patient does not have acute narrow-angle glaucoma Renewal Criteria: · Patient continues to meet initial criteria; AND Patient does not have treatment-limiting adverse effects (e.g., treatment-limiting central nervous system depression or cognitive impairment, worsened glaucoma, respiratory depression, suicidal ideation, clinically significant changes in blood pressure or heart rate); AND Prescriber to provide verbal attestation of diazepam effectiveness (e.g., decreased typical length of repetitive seizures) Initial Criteria (6-month duration): • Patient has diagnosis of intermittent, stereotypic episodes of frequent seizure activity (e.g., seizure clusters, acute repetitive seizures) that are distinct from a patient's usual seizure pattern; AND Patient is 12 years of age or older; AND Prescribed by, or in consultation with, a neurologist; AND Patient is on a stable antiepileptic regimen; AND • Prescriber has counseled patient on the following: Risks if combined with opioids o Identification of a seizure cluster Proper administration o When to seek emergency medical treatment; AND 10 doses/ **General PA** Nayzilam® Patient is not using moderate or strong CYP 3A4 inhibitors or, if unavoidable, prescriber will monitor toxicity risk during 30 days Form concomitant use; AND Due to increased risk of toxicity, patient should not be pregnant OR concurrently taking CNS stimulants, CNS depressants, carisoprodol, meprobamate, or barbiturates; AND • Patient does not have acute narrow-angle glaucoma **Renewal Criteria:** Patient continues to meet initial criteria; AND Patient does not have treatment-limiting adverse effects (e.g., treatment-limiting central nervous system depression or cognitive impairment, worsened glaucoma, respiratory depression, suicidal ideation, clinically significant changes in blood pressure or heart rate); AND Prescriber to provide verbal attestation of midazolam effectiveness (e.g., decreased typical length of repetitive seizures)



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Prior Authorization Criteria Qty. Limits PA Form** • Diagnosis of neuropathic pain; OR • Diagnosis of postherpetic neuralgia; OR **General PA** Ρ pregabalin capsules • Diagnosis of fibromyalgia; OR **Form** · Diagnosis of seizure disorder • One of the following: o Diagnosis of neuropathic pain o Diagnosis of postherpetic neuralgia **General PA** o Diagnosis of fibromyalgia pregabalin solution Ρ Form o Diagnosis of seizure disorder; AND • Patient is less than 12 years of age; OR Patient is unable to swallow solid oral dosage forms phenobarbital Ρ • Will be approved for use ONLY in patients with diagnosis of seizure disorders. • Will be approved for use ONLY in patients with diagnosis of seizure disorders. phenobarbital elixir **Note**: PA is not required for patients less than 2 years of age **General PA** • Diagnosis of Lennox-Gastaut Syndrome; AND Ρ rufinamide tablets Form • Used as adjunct therapy with at least one other anticonvulsant Diagnosis of Lennox-Gastaut Syndrome; AND rufinamide susp • Used as adjunct therapy with at least one other anticonvulsant; AND • Unable to swallow solid dosage forms • Adjunctive therapy for patients with partial-onset seizures or primary generalized tonic-clonic seizures; OR seizures associated with Lennox-Gastaut syndrome; AND o Will be used approved in combination with at least one other anticonvulsant; AND 25, 50, & 100 mg: General PA Trokendi XR o Trial and failure of preferred immediate release product and one additional preferred agent; OR 1/day; Form • Initial monotherapy in patients with partial-onset or primary generalized tonic-clonic seizures; AND 200 mg: 2/day o Trial and failure of preferred immediate release product and one additional preferred agent; OR



• Migraine Prophylaxis in patients ≥ 12 years of age

CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Prior Authorization Criteria Qty. Limits PA Form** Initial Criteria (6-month duration): • Patient has diagnosis of intermittent, stereotypic episodes of frequent seizure activity (e.g., seizure clusters, acute repetitive seizures) that are distinct from a patient's usual seizure pattern; AND • Patient is 2 years of age or older; AND Prescribed by, or in consultation with, a neurologist; AND · Patient is on a stable antiepileptic regimen; AND • Prescriber has counseled patient on the following: Risks if combined with opioids Identification of a seizure cluster Proper administration When to seek emergency medical treatment; AND 5 boxes/30 days Valtoco® • Patient is not using CYP 2C19 and CYP 3A4 inhibitors or, if unavoidable, prescriber will monitor toxicity risk during concomitant use; AND Due to increased risk of toxicity, patient should not be pregnant OR concurrently taking CNS stimulants, CNS depressants, carisoprodol, meprobamate, or barbiturates; AND • Patient does not have acute narrow-angle glaucoma Renewal Criteria (1 year duration): • Patient continues to meet initial criteria; AND Patient does not have treatment-limiting adverse effects (e.g., treatment-limiting CNS depression or cognitive impairment, worsened glaucoma, respiratory depression, suicidal ideation, clinically significant changes in blood pressure); AND Prescriber to provide verbal attestation of diazepam effectiveness (e.g., decreased typical length of repetitive seizures) 25 mg (4/day); Р 50 mg (2/day); zonisamide 100 mg (6/day) Initial Criteria: Patient is 2 years of age and older; AND Diagnosis of seizure disorder associated with cyclin-dependent kinase-like 5 deficiency disorder; AND • Prescriber has confirmed that patient is not pregnant (if applicable) and counseled patient on risks of pregnancy while taking Ztalmy; AND • Prescriber has confirmed member does not have hepatic disease and will monitor hepatic function (dose reductions may 36 mL/day Ztalmy® be required in impaired hepatic function) Renewal Criteria: General PA Prescriber has confirmed that patient is not pregnant (if applicable); AND Form Prescriber has confirmed member does not have hepatic disease and will monitor hepatic function (dose reductions may be required in impaired hepatic function) Diagnosis of Lennox-Gastaut Syndrome; AND Banzel® tablets NP Used as adjunct therapy with at least one other anticonvulsant



Banzel® suspension

NP

Diagnosis of Lennox-Gastaut Syndrome; AND

Unable to swallow solid dosage forms

• Used as adjunct therapy with at least one other anticonvulsant; AND

CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Prior Authorization Criteria Qty. Limits PA Form** See Briviact® tablets prior authorization criteria Briviact® solution NP 20 mL/day Additionally, patient must be unable to swallow tablets Patient is ≥ 1 month of age; AND · Have diagnosis of partial-onset seizures; AND Briviact® tablets NP Have tried and failed at least 1 other medication indicated for partial-onset seizures 2/dav NOTE: A dose reduction is required for all stages of hepatic impairment (Child-Pugh A, B, and C) and use is not recommended in end- stage renal disease patients. • Must meet clobazam tablets prior authorization criteria; AND clobazam suspension NP Patient must be unable to swallow tablets Must meet clonazepam prior authorization criteria; AND 3/day clonazepam ODT NP · Patient must be unable to swallow, OR unable to absorb medications through the GI tract. Initial Criteria: Patient is ≥ 6 months of age: AND Patient must also be taking clobazam concomitantly; AND · Diagnosis of Dravet syndrome (DS); AND Prescribed by neurologist or epileptologist; AND Prescriber attests that baseline serum hematologic testing has been completed; AND • Prescriber attests the patient has refractory epilepsy (failed to become seizure free after trials of 2 antiepileptic drugs); 250 mg (1/day); General PA Diacomit® NΡ 500 mg (6/day) Form If the oral powder for suspension is prescribed, the patient does not have phenylketonuria (PKU). Renewal Criteria: · Patient continues to meet initial criteria; AND • Patient has no treatment-limiting adverse effects (e.g., thrombocytopenia, neutropenia, new onset or worsened depression; suicidal thoughts, worsened seizure control); AND Prescriber to provide verbal attestation of Diacomit effectiveness (e.g., reduced seizure frequency and/or duration) Patient has a diagnosis or history of partial-onset seizures, juvenile myoclonic epilepsy, or primary generalized tonic-clonic seizure: AND 1000 mg: 3/day; Elepsia® XR • Will be used as adjunctive therapy with another anticonvulsant; AND 1500 mg: 2/day Patient must be 12 years of age or older; AND Clinically valid reason why the preferred levetiracetam ER cannot be used **General PA** • One of the following: Form o Will be used as initial monotherapy for the treatment of partial-onset or primary generalized tonic-clonic seizures in patients 2 years of age and older Eprontia® solution NP o Will be used as adjunctive therapy for the treatment of partial-onset seizures, primary generalized tonic-clonic 16 ml/day seizures, and seizures associated with Lennox-Gastaut syndrome in patients 2 years of age and older



Will be used as preventive treatment of migraine in patients 12 years and older; AND

Patient is unable to swallow tablets

CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Prior Authorization Criteria Qty. Limits PA Form** Initial Criteria: Used as adjunctive therapy for the treatment of partial and generalized seizures associated with Lennox-Gastaut Syndrome in children 2-14 years of age with a contraindication to, or trial and failure of, TWO of the following: Valproic acid/divalproex sodium Lamotrigine o Topiramate • Used as monotherapy and adjunctive therapy for the treatment of partial seizures with or without generalization in adults > 14 years of age with a contraindication to, or trial and failure of, **THREE** of the following: Felbatol® and NP Carbamazepine felbamate Oxcarbazepine o Phenytoin Gabapentin o Lamotrigine Topiramate Valproic acid/divalproex sodium **Note:** Will not be approved if there is a history of blood dyscrasia or liver disease unless the prescriber can make a compelling clinical case demonstrating that the benefits of the drug outweigh the risks. Initial Criteria: Patient must be ≥ 2 years of age; AND • Diagnosis of Dravet syndrome (DS) or Lennox-Gastaut syndrome (LGS); AND Prescribed by a neurologist or epileptologist; AND Patient has not received MAOI therapy within 14 days and will not receive during Fintepla therapy; AND • Prescriber attestation that baseline echocardiogram has been completed and will be monitored throughout treatment and Fintepla® NP 3 - 6 months after the final dose; AND 1 bottle/30 days • Inadequate response to trials of 2 preferred anticonvulsant agents Renewal Criteria: **General PA** · Patient continues to meet initial criteria; AND Form Patient has no treatment-limiting adverse effects (e.g., serotonin syndrome, abnormal AST/ALT, CrCl, abnormal echocardiogram); AND Patient is responding to therapy (e.g., reduced seizure frequency and/or duration) • Diagnosis of partial onset seizures with or without secondarily generalized seizures; AND o Patient is ≥ 4 years of age; AND 2, 4, 8, 10, & 12 mg: o Trial and failure, contraindication, or intolerance to 2 preferred agents, one of which must be lacosamide **OR** Fycompa® 1/day; • Will be used as adjunctive therapy for the treatment of primary generalized tonic-clonic (PGTC) seizures; AND 6 mg: 2/day Patient is ≥ 12 years of age; AND



o Trial and failure, contraindication, or intolerance to TWO preferred agents

CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication **PDL Prior Authorization Criteria Qty. Limits PA Form** 100 & 600 mg: 6/day; 800 mg: 4.5/day; All other strengths: gabapentin tablets Clinically valid reason why the preferred gabapentin capsules cannot be used 3/day Max daily gabapentin dose: 3600 mg Anti-anxiety See clonazepam prior authorization criteria; AND Klonopin® NP 3/day • Trial and failure of clonazepam PA Form Lamictal® ODT NP Unable to swallow solid dosage forms **General PA** • Trial and failure of a regular-release lamotrigine product and 1 other preferred agent Lamictal® XR NP **Form** NΡ lamotrigine ER • Trial and failure of a regular-release lamotrigine product and 1 other preferred agent lamotrigine ODT NΡ Unable to swallow solid dosage forms See pregabalin capsules prior authorization criteria; AND Lyrica® capsules NP • Trial and failure of preferred pregabalin capsules See pregabalin solution prior authorization criteria; AND Lyrica[®] solution NΡ • Trial and failure of preferred pregabalin solution Diagnosis of postherpetic neuralgia OR neuropathic pain associated with-diabetic peripheral neuropathy; AND 82.5 mg & 165 mg: • Trial and failure, contraindication, or intolerance to a tricyclic antidepressant OR gabapentin; AND 1/day NP Lyrica® CR • Provider must provide peer-reviewed medical literature documenting why the requested drug for the requested indication 330 mg: 2/day **General PA** is the only appropriate choice versus immediate-release pregabalin Form • One of the following: Initial monotherapy for partial onset seizures Motpoly ®XR NP o Adjunctive therapy for partial onset seizures and will be used in combination with at least one other anticonvulsant; Trial and failure of preferred immediate release product and one additional preferred agent 72 mL/day Max total daily See gabapentin solution prior authorization criteria. Neurontin® solution Note: Prior authorization criteria is waived for recipients 12 years of age and under gabapentin dose: 3600mg Anti-anxiety Onfi® NP | See clobazam tablets prior authorization criteria PA Form



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Qty. Limits PA Form Prior Authorization Criteria** • Will be used as monotherapy or adjunctive therapy in patients with focal (partial) onset or primary generalized tonic-clonic seizures; OR 200 mg: 2/day **General PA** Qudexy® XR NP • Will be used as adjunctive therapy in patients with seizures associated with Lennox-Gastaut syndrome; OR All other strengths: Form • Migraine Prophylaxis in patients ≥ 12 years of age; AND 1/day o Trial and failure of an Trokendi XR and 1 other preferred agent Diagnosis of Lennox-Gastaut Syndrome; AND **General PA** NP rufinamide tablet • Used as adjunct therapy with at least one other anticonvulsant Form Adjunctive therapy for patients with refractory complex partial seizures who have responded inadequately to several alternative treatments; AND General PA NP Patient has tried and failed 2 preferred anticonvulsants; OR Sabril® Form • Monotherapy for patients with infantile spasms Note: This drug is subject to REMS requirements to ensure the benefits of treatment outweigh the risks of vision loss 250, 500, & 1000 mg: • Patient is unable to swallow solid oral dosage form; AND General PA NP Spritam® 2/dav: • Provider must have a clinically valid reason as to why the generic levetiracetam solution cannot be used Form 750 mg: 4/day • Patient has a diagnosis of Lennox-Gastaut syndrome (LGS); AND Requested drug will be used as adjunctive therapy in combination with at least one other anticonvulsant; AND **General PA** Sympazan® NP 2/dav • Provider must have a clinically valid reason as to why both clobazam tablets and suspension cannot be used. (NOTE: Patient **Form** convenience is NOT an approvable reason) • Will be used as monotherapy or adjunctive therapy in patients with focal (partial) onset or primary generalized tonic-clonic seizures: OR 200 mg: 2/day **General PA** NP • Will be used as adjunctive therapy in patients with seizures associated with Lennox-Gastaut syndrome; OR topiramate ER All other strengths: Form • Migraine Prophylaxis in patients ≥ 12 years of age; AND 1/day o Trial and failure of an Trokendi XR and 1 other preferred agent vigabatrin NP | See Sabril® prior authorization criteria General PA • Treatment is for monotherapy for patients with infantile spasms; AND Form Clinically valid reason why vigabatrin 50 mg/mL powder oral solution cannot be used Vigafyde® Note: This drug is subject to REMS requirements to ensure the benefits of treatment outweigh the risks of vision loss Vigadrone® NP See Sabril® prior authorization criteria **General PA** See lacosamide prior authorization criteria; AND Form NP Vimpat[®] • Trial and failure, contraindication, or intolerance to lacosamide



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication **PDL Prior Authorization Criteria Qty. Limits** PA Form Initial criteria: • Diagnosis of partial-onset seizures; AND • Prescribed by, or in consultation with, a neurologist; AND • Must be 18 years of age and older; AND • Trial and failure, contraindication, or intolerance to 2 preferred anticonvulsants indicated for partial-onset seizures; AND **General PA** NP Xcopri® 2/day • Patient does not have Familial Short QT syndrome Form Renewal criteria: • Patient must demonstrate disease improvement and stabilization as a result of the medication; AND • Patient is absent of unacceptable toxicity from the drug; AND • Patient's QT interval is being monitored • Diagnosis of partial-onset seizures; AND **General PA** Zonisade® • Zonisade will be used as adjunctive therapy; AND 30 mL/day NP <u>Form</u> • Patient must be unable to swallow solid dosage forms



		CENTRAL NERVOUS SYSTEM		
Medication	PDL	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Prior Authorization Criteria	Qty. Limits	PA Form
		Movement Disorders		
Austedo®	P	 Diagnosis of tardive dyskinesia: Patient age ≥ 18 years; AND Documentation that AIMS test has been completed (e.g., score or copy of AIMS assessment); AND Prescribed by, or in consultation with, a neurologist or psychiatrist (or other mental health provider), provided patient has reasonable access; AND Documentation or claims history of current or former chronic patient use of a dopamine antagonist (e.g., antipsychotic, metoclopramide, prochlorperazine, droperidol, promethazine) Diagnosis of chorea related to Huntington's Disease: Physician is experienced in the treatment of Huntington's Disease or is in a Center of Excellence for Huntington's Disease; AND Patient does not have a history of untreated or inadequately treated depression or suicidal ideation due to a boxed warning that it increases the risk of depression and suicidal thoughts and behavior Patients meeting any of the following criteria will NOT be approved: Concurrent therapy with tetrabenazine, reserpine, or MAOIs Hepatic impairment Hypersensitivity to the active ingredient Pregnancy 	4/day	
Austedo XR®	Р	See Austedo prior authorization criteria	1/day	
Ingrezza [®]	P	 Diagnosis of tardive dyskinesia: Patient age ≥ 18 years; AND Documentation that AIMS test has been completed (e.g., score or copy of AIMS assessment); AND Prescribed by, or in consultation with, a neurologist or psychiatrist (or other mental health provider), provided patient has reasonable access; AND Documentation or claims history of current or former chronic patient use of a dopamine antagonist (e.g., antipsychotic, metoclopramide, prochlorperazine, droperidol, promethazine) Diagnosis of chorea related to Huntington's Disease: Physician is experienced in the treatment of Huntington's Disease or is in a Center of Excellence for Huntington's Disease; AND Patient does not have a history of untreated or inadequately treated depression or suicidal ideation due to a boxed warning that it increases the risk of depression and suicidal thoughts and behavior Patients meeting any of the following criteria will NOT be approved: Concurrent use of MAOIs or strong CYP3A4 inducers Hypersensitivity to the active ingredient Pregnancy 	40 mg: 2/day 60, 80 mg: 1/day	General PA Form
tetrabenazine	Р	Will only be approved for the treatment of chorea associated with Huntington's disease.		
Xenazine®	Р	 Diagnosis of chorea associated with Huntington's disease; AND Clinically valid reason why preferred generic tetrabenazine cannot be used 		



CENTRAL NERVOUS SYSTEM

Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.

Medication	PDL Prior Authorization Criteria	Qty. Limits	PA Form
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Antidepressants: MAOIs

CLASS PRIOR AUTHORIZATION CRITERIA FOR PATIENT'S WITH INTELLECTUAL AND DEVELOPMENTAL DISABILITIES (I/DD):

Antidepressants prescribed for disorders related to Intellectual and Developmental Disabilities will be approved if ONE of the following is met:

- Prescribed by a Gold Card prescriber; OR
- Mental health assessment applicable to behavioral symptoms for which the medication is being prescribed; AND
 - Underlying physical condition such as pain, discomfort, and environmental issues have been evaluated, treated, and behavioral symptoms persist; AND
 - Non-pharmacological interventions have been tried and behavioral symptoms persist (e.g., behavior plan, crisis intervention, and stabilization strategies or training) and training and support have been provided to family or other caregivers; OR
- Short-term therapy (less than 90 days) has been prescribed; AND
 - o Behavioral symptoms are significant enough to place the person at potential risk or needing higher level of care or loss of community placement; **OR**
 - Continuation of existing therapy to address serious and ongoing behavioral symptoms while other prior authorization criteria are met; AND
 - o Efficacy and potential side effects to be monitored; AND
 - o Need for requested medication will be evaluated once other non-pharmacological interventions have been tried

Note the following:

- Drug specific step therapy, NP criteria and Brand Medically Necessary Criteria will still apply to this patient population.
- The I/DD Worksheet can be found at: I/DD Prior Authorization Form

phenelzine	P	 Diagnosis of major depression; AND Trial and failure of THREE antidepressant agents from TWO different following drug classes: SSRIs SNRIs New generation antidepressants 	6 tabs/day	
Emsam®	NP	See Marplan® prior authorization criteria; AND • Patient must be 13 years of age or older	1/day	
Marplan®	NP	Diagnosis of major depression; AND Trial and failure of THREE antidepressant agents from TWO different following drug classes: SSRIS SNRIS New generation antidepressants; AND Trial and failure, contraindication, or intolerance to preferred phenelzine	6 tabs/day	General PA Form
Nardil®	NP	See Marplan® prior authorization criteria	6 tabs/day	
Parnate®	NP	See Marplan® prior authorization criteria	6 tabs/day	
tranylcypromine	NP	See Marplan® prior authorization criteria	6 tabs/day	



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.

Prior Authorization Criteria Qty. Limits PA Form

Antidepressants: New Generation

CLASS PRIOR AUTHORIZATION CRITERIA FOR PATIENT'S WITH INTELLECTUAL AND DEVELOPMENTAL DISABILITIES (I/DD):

Antidepressants prescribed for disorders related to Intellectual and Developmental Disabilities will be approved if ONE of the following is met:

Prescribed by a Gold Card prescriber; OR

PDL

- Mental health assessment applicable to behavioral symptoms for which the medication is being prescribed; AND
 - Underlying physical condition such as pain, discomfort, and environmental issues have been evaluated, treated, and behavioral symptoms persist; AND
 - o Non-pharmacological interventions have been tried and behavioral symptoms persist (e.g., behavior plan, crisis intervention, and stabilization strategies or training) and training and support have been provided to family or other caregivers
- Short-term therapy (less than 90 days) has been prescribed; AND
 - Behavioral symptoms are significant enough to place the person at potential risk or needing higher level of care or loss of community placement; OR
 - Continuation of existing therapy to address serious and ongoing behavioral symptoms while other prior authorization criteria are met; AND
 - Efficacy and potential side effects to be monitored; AND
 - Need for requested medication will be evaluated once other non-pharmacological interventions have been tried

Note the following:

Medication

- Drug specific step therapy, NP criteria and Brand Medically Necessary Criteria will still apply to this patient population.
- The I/DD Worksheet can be found at: I/DD Prior Authorization Form

Aplenzin®	Р			
bupropion IR/SR	Р			
bupropion XL	Р		1/day	
mirtazapine	Р			
mirtazapine ODT	Р	Patient is unable to swallow solid dosage forms		
trazodone (excluding 300mg)	Р			
Auvelity®	NP	 Diagnosis of Major Depressive Disorder (MDD); AND Patient is 18 years of age or older; AND Trial and failure, or contraindication, intolerance to 2 preferred antidepressants; AND Patient does not have ANY of the following: Seizure disorder Current or prior diagnosis of bulimia or anorexia nervosa Undergoing abrupt discontinuation of alcohol, benzodiazepines, barbiturates, or antiepileptic drugs; AND Prescriber attests patient has not received MAOI therapy within 14 days and will not receive during therapy 		General PA Form
Forfivo XL®	NP	 Trial and failure, contraindication, or intolerance of 2 preferred agents; AND Patient must currently be on a bupropion product titrated to a dose of 300 mg per day 		
nefazodone	NP	 Diagnosis of major depression; AND Trial and failure, contraindication, or intolerance of 2 preferred agents; AND Patient does not have hepatic impairment 		
Remeron®	NP			Conoral DA
Remeron SolTab®	NP	Patient is unable to swallow solid dosage forms		General PA

June 1, 2025



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.						
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
trazodone 300mg	NP	 Trial and failure, contraindication, or intolerance of 2 preferred agents; AND Clinically valid reason why the preferred lower strength tablets cannot be used (i.e., trazodone 50mg, 100mg, 150mg) 		<u>Form</u>		
Wellbutrin® IR & SR	NP					
Wellbutrin XL®	NP		1/day			
Zurzuvae®	NP	 Criteria: (3 month-duration) Patient is 18 years of age or older; AND Diagnosis of postpartum depression (PPD); AND Patient's symptoms began in the third trimester or within 4 weeks of delivery; AND Prescriber attests that the PPD requires rapid improvement and resolution of symptoms; AND Prescribed by, or in consultation with, a psychiatrist, psychologist, or an obstetrician-gynecologist; AND Prescriber attests to ALL of the following: Patient has been advised not to drive or operate machinery until at least 12 hours after administration due central nervous system (CNS) depressant effects such as somnolence and confusion Females of reproductive potential have been advised to use effective contraception during treatment and for 1 week after the final dose due to potential risk to fetus and to notify healthcare provider if they become pregnant during treatment Lactating women have been counseled on risk versus benefits of breastfeeding while on treatment 	1 treatment course/year	General PA Form		

Antidepressants: SNRIs

CLASS PRIOR AUTHORIZATION CRITERIA FOR PATIENT'S WITH INTELLECTUAL AND DEVELOPMENTAL DISABILITIES (I/DD):

Antidepressants prescribed for disorders related to Intellectual and Developmental Disabilities will be approved if ONE of the following is met:

- Prescribed by a Gold Card prescriber; OR
- Mental health assessment applicable to behavioral symptoms for which the medication is being prescribed; AND
 - Underlying physical condition such as pain, discomfort, and environmental issues have been evaluated, treated, and behavioral symptoms persist; AND
 - Non-pharmacological interventions have been tried and behavioral symptoms persist (e.g., behavior plan, crisis intervention, and stabilization strategies or training) and training and support have been provided to family or other caregivers; OR
- Short-term therapy (less than 90 days) has been prescribed; AND
 - Behavioral symptoms are significant enough to place the person at potential risk or needing higher level of care or loss of community placement; OR
 - Continuation of existing therapy to address serious and ongoing behavioral symptoms while other prior authorization criteria are met; AND
 - o Efficacy and potential side effects to be monitored; AND
 - Need for requested medication will be evaluated once other non-pharmacological interventions have been tried

Note the following:

- Drug specific step therapy, NP criteria and Brand Medically Necessary Criteria will still apply to this patient population.
- The I/DD Worksheet can be found at: I/DD Prior Authorization Form

desvenlafaxine ER	Р	1/day	
duloxetine 20, 30, & 60 mg	Р	2/day	SNRI PA
Effexor XR®	Р	1/day	Form
venlafaxine IR tabs	Р	2/day	



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication **PDL Prior Authorization Criteria PA Form** Qty. Limits 37.5, 75 mg: 1/day 150 mg: 2/day venlafaxine ER caps Ρ Note: for 225 & 375 mg doses: use 150 mg & 75 mg caps Cymbalta® NP 2/day duloxetine 40 mg NΡ • Clinically valid reason why the preferred duloxetine capsules (20, 30, or 60 mg) cannot be used 2/day Titration Pack: 1/day Fetzima® NP (56 tabs/lifetime)

Antidepressants: SSRI

Clinically valid reason why preferred venlafaxine agents cannot be used (Effexor XR, venlafaxine ER caps, venlafaxine IR

CLASS PRIOR AUTHORIZATION CRITERIA FOR PATIENTS WITH INTELLECTUAL AND DEVELOPMENTAL DISABILITIES (I/DD):

Antidepressants prescribed for disorders related to Intellectual and Developmental Disabilities will be approved if ONE of the following is met:

• Prescribed by a Gold Card prescriber; OR

NP

NP

NP

- Mental health assessment applicable to behavioral symptoms for which the medication is being prescribed; AND
 - Underlying physical condition such as pain, discomfort, and environmental issues have been evaluated, treated, and behavioral symptoms persist; AND
 - Non-pharmacological interventions have been tried and behavioral symptoms persist (e.g., behavior plan, crisis intervention, and stabilization strategies or training) and training and support have been provided to family or other caregivers; OR
- Short-term therapy (less than 90 days) has been prescribed; AND

tabs)

- Behavioral symptoms are significant enough to place the person at potential risk or needing higher level of care or loss of community placement; OR
 - Continuation of existing therapy to address serious and ongoing behavioral symptoms while other prior authorization criteria are met; AND
- o Efficacy and potential side effects to be monitored; AND
- Need for requested medication will be evaluated once other non-pharmacological interventions have been tried

Note the following:

Pristiq®

ER tabs

venlafaxine besylate

venlafaxine ER tabs

- Drug specific step therapy, NP criteria and Brand Medically Necessary Criteria will still apply to this patient population.
- The I/DD Worksheet can be found at: I/DD Prior Authorization Form

citalopram	Р	10, 20 mg: 1.5/day 40 mg: 1/day	
escitalopram	Р	1.5/day	
escitalopram solution	Р		General PA
fluoxetine capsules	Р	3/day	<u>Form</u>
fluoxetine solution	Р		
fluvoxamine	Р	3/day	



1/day

1/day

1/day

SNRI PA

Form

CENTRAL NERVOUS SYSTEM

Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Wiedleation	100	Thor Authorization Criteria	-	TATOM
paroxetine tablets	Р		10, 20 mg: 1/day;	
-	+		30, 40 mg: 2/day	_
sertraline	Р		25, 50 mg: 1.5/day; 100 mg: 2/day	
vilazodone	P		1/day	-
VIIdZOGOTIC	+ '		10, 20 mg: 1.5/day	-
Celexa®	NP		40 mg: 1/day	
		Stabilized at a dose of 20 mg/day of fluoxetine for > one month; AND		
fluoxetine DR caps	NP	Documented reason why the patient is unable to continue fluoxetine 20 mg daily	4/28 days	
fluoxetine tablets	NP		20 mg: 3/day;	
nuoxetine tablets	INP		60 mg: 1/day	
fluvoxamine ER	NP		100 mg: 3/day;	
navoxanime En	14.		150 mg: 2/day	
Lexapro®	NP		1.5/day	
		Diagnosis of hot flashes associated with menopause; AND		
paroxetine 7.5 mg	NP	Trial and failure, contraindication, or intolerance to estrogen therapy; AND		
	-	An allergy or intolerance to an inactive ingredient in paroxetine		_
paroxetine CR	NP		12.5, 25 mg: 1/day;	
			37.5 mg: 2/day	General PA
Paxil® tablets	NP		10, 20 mg: 1/day; 30, 40 mg: 2/day	<u>Form</u>
Paxil® CR	NP		See paroxetine CR	_
Paxil® solution	NP		See paroxetine CK	
Prozac®	NP		3/day	1
				_
sertraline capsules	NP	Di i fati D i Di Land	1/day	1
Trintellix®	NP	 Diagnosis of Major Depression Disorder; AND Adequate trial and failure of TWO agents at an appropriate dose (3 weeks at the maximum tolerated dose within the 	1/454	
Trintellix -	INP	recommended therapeutic range) within the following drug classes: SSRI, SNRI, or New Generation Antidepressants	1/day	
Viibryd	NP	and the second s	1/day	1
- 1 (. 0	1		25, 50 mg: 1.5/day;	1
Zoloft®	NP		100 mg: 2/day	

June 1, 2025



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL Prior Authorization Criteria Qty. Limits PA Form Antidepressants: Tricyclics

CLASS PRIOR AUTHORIZATION CRITERIA FOR PATIENTS WITH INTELLECTUAL AND DEVELOPMENTAL DISABILITIES (I/DD):

Antidepressants prescribed for disorders related to Intellectual and Developmental Disabilities will be approved if ONE of the following is met:

- Prescribed by a Gold Card prescriber; OR
- Mental health assessment applicable to behavioral symptoms for which the medication is being prescribed; AND
 - Underlying physical condition such as pain, discomfort, and environmental issues have been evaluated, treated, and behavioral symptoms persist; AND
 - Non-pharmacological interventions have been tried and behavioral symptoms persist (e.g., behavior plan, crisis intervention, and stabilization strategies or training) and training and support have been provided to family or other caregivers; OR
- Short-term therapy (less than 90 days) has been prescribed; AND
 - Behavioral symptoms are significant enough to place the person at potential risk or needing higher level of care or loss of community placement; OR
 - Continuation of existing therapy to address serious and ongoing behavioral symptoms while other prior authorization criteria are met; AND
 - o Efficacy and potential side effects to be monitored; AND
 - o Need for requested medication will be evaluated once other non-pharmacological interventions have been tried

Note the following:

- Duration of short-term therapy is 90 days for antidepressants
- Drug specific step therapy, NP criteria and Brand Medically Necessary Criteria will still apply to this patient population.
- The I/DD Worksheet can be found at: I/DD Prior Authorization Form

amitriptyline	Р		
doxepin caps	Р		
imipramine tabs	Р		
nortriptyline	Р		
amoxapine	NP		
Anafranil®	NP	See prior authorization criteria for clomipramine	
clomipramine	NP	 Diagnosis of obsessive-compulsive disorder; AND Trial and failure of at least 2 unique SSRIs 	General PA Form
desipramine	NP		
imipramine caps	NP		
Norpramin®	NP		
nortriptyline solution	NP	Patient is unable to swallow nortriptyline capsules	
Pamelor®	NP		
protriptyline	NP		



	CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
		Antihyperkinesis: Stimulants		•		
Adderall® XR	Р	See amphetamine salt ER combination prior authorization criteria	5, 10, 15 mg: 1/day 25 & 30mg: 2/day 20mg: 3/day Max total amphetamine dose (Age ≥ 21): 60mg/day	Anti- hyperkinesis: Stimulants PA Form		
amphetamine salt ER combination	Р	 Agent must not be prescribed by a pain clinic Patient will not concurrently take a benzodiazepine, barbiturate, sedative hypnotic, opioid (including buprenorphine), MAOI (monoamine oxidase inhibitor) agent, or meprobamate/carisoprodol; AND Patient has NOT had active alcohol or substance abuse for last 3 years, if patient ≥ 21 years of age; AND One of the following: Diagnosis of Attention Deficit Disorder and/or Hyperactivity Disorder (ADD/ADHD); AND Documentation that the symptoms affect the patient's ability to function in daily life tasks or creates significant difficulties in at least 2 major settings (school, work, social settings, and/or home) Diagnosis of Narcolepsy supported with documentation of polysomnography or multiple sleep latency test (MSLT) Diagnosis of neurocognitive disorder (also known as organic brain disorder) Diagnosis of treatment resistant Major Depressive Disorder; AND Adequate trial and failure of 3 agents at an appropriate dose (defined as: 3 weeks at the maximum tolerated dose within the recommended therapeutic range) from at least 3 distinct drug classes:	5, 10, 15 mg: 1/day 25 & 30 mg: 2/day 20 mg: 3/day Max total amphetamine dose (Age ≥ 21): 60 mg/day	Anti- hyperkinesis: Stimulants PA Form		
amphetamine salt IR combo	Р	See amphetamine salt ER combination prior authorization criteria	5, 7.5, 10, & 12.5 mg: 4/day 15 & 30 mg: 2/day 20 mg: 3/day Max total amphetamine dose (Age ≥ 21): 60 mg/day	Anti- hyperkinesis: Stimulants PA Form		
amphetamine sulfate	Р	See amphetamine salt ER combination prior authorization criteria	See Evekeo®	<u> </u>		
Aptensio XR®	Р	See amphetamine salt ER combination prior authorization criteria	1/day	Anti- hyperkinesis: Stimulants PA Form		



CENTRAL NERVOUS SYSTEM

		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated	<i>t.</i>	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Concerta®	Р	See amphetamine salt ER combination prior authorization criteria	18, 27, 54 mg: 1/day; 36 mg: 2/day	Anti-
Daytrana®	Р	See amphetamine salt ER combination prior authorization criteria	1/day	hyperkinesis:
dexmethylphenidate	Р	See amphetamine salt ER combination prior authorization criteria	1/day	Stimulants PA
dexmethylphenidate XR	Р	See amphetamine salt ER combination prior authorization criteria	1/day	<u>Form</u>
dextroamphetamine tablets	Р	See amphetamine salt ER combination prior authorization criteria	20 mg: 3/day 30 mg: 2/day All others: 4/day Max total amphetamine dose (Age ≥ 21): 60mg/day	Anti- hyperkinesis: Stimulants PA
Focalin XR®	Р	See amphetamine salt ER combination prior authorization criteria	1/day	<u>Form</u>
methylphenidate (generic for Ritalin®)	Р	See amphetamine salt ER combination prior authorization criteria	1/day	
methylphenidate solution	Р	See amphetamine salt ER combination prior authorization criteria		
methylphenidate ER tablets	Р	See amphetamine salt ER combination prior authorization criteria	See Metadate ER®	
ProCentra®	Р	See amphetamine salt ER combination prior authorization criteria	20 mL/day Max (Age ≥ 21): 60mg/day	Anti- hyperkinesis: Stimulants PA Form
Vyvanse® capsules and chewables	Р	See amphetamine salt ER combination prior authorization criteria	1/day; Max total amphetamine dose (Age ≥ 21): 60mg/day	



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Prior Authorization Criteria Qty. Limits PA Form** • Agent must not be prescribed by a pain clinic Patient will not concurrently take a benzodiazepine, barbiturate, sedative hypnotic, opioid (including buprenorphine), MAOI (monoamine oxidase inhibitor) agent, or meprobamate/carisoprodol; AND Patient has NOT had active alcohol or substance abuse for last 3 years, if patient ≥ 21 years of age; AND • One of the following: Diagnosis of Attention Deficit Disorder and/or Hyperactivity Disorder (ADD/ADHD); AND - Documentation that the symptoms affect the patient's ability to function in daily life tasks or creates significant difficulties in at least 2 major settings (school, work, social settings, and/or home) Diagnosis of Narcolepsy supported with documentation of polysomnography or multiple sleep latency test (MSLT) See amphetamine salt NP Adderall® o Diagnosis of neurocognitive disorder (also known as organic brain disorder) IR combo o Diagnosis of treatment resistant Major Depressive Disorder; AND Adequate trial and failure of 3 agents at an appropriate dose (defined as: 3 weeks at the maximum tolerated dose within the recommended therapeutic range) from at least 3 distinct drug classes: Anti- SSRI hyperkinesis: SNRI Stimulants PA New Generation Antidepressants Form TCAs; AND Trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated 5, 10, 15 mg: 1/day 25 & 30mg: 2/day 20mg: 3/day Adderall® XR NP | See Adderall® prior authorization criteria Max total amphetamine dose (Age ≥ 21): 60mg/day Adhansia XR® NP | See Adderall® prior authorization criteria 1/day See Adderall® prior authorization criteria Adzenys ER® solution NP 10mL/day • Patient must have clinical reason as to why the preferred generic methylphenidate solution cannot be used. Adzenys XR® ODT NP See Adderall® prior authorization criteria 1/day amphetamine ER See Adderall® prior authorization criteria Anti-NP 10mL/day suspension Patient must have clinical reason as to why the preferred generic methylphenidate solution cannot be used. hyperkinesis: Stimulants PA Azstarys® NP | See Adderall® prior authorization criteria 1/day **Form** Cotempla XR® ODT NP | See Adderall® prior authorization criteria 1/day Anti-4/day hyperkinesis: Max total Stimulants PA NP | See Adderall® prior authorization criteria Desoxyn® amphetamine dose **Form** (Age \geq 21): 60 mg/day



CENTRAL NERVOUS SYSTEM

	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.			
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
dextroamphetamine solution	NP	See Adderall® prior authorization criteria	20 mL/day Max total amphetamine dose (Age ≥ 21): 60 mg/day	Anti- hyperkinesis:
Dexedrine®	NP	See Adderall® prior authorization criteria	4/day Max total amphetamine dose (Age ≥ 21): 60 mg/day	Stimulants PA Form
Dyanavel XR®	NP	See Adderall® prior authorization criteria	8 mL/day Max total amphetamine dose (Age ≥ 21): 60 mg/day	
Evekeo® tab & ODT	NP	See Adderall® prior authorization criteria	5 mg tab & ODT: 3/day 10 mg tab & ODT: 6/day 15 mg ODT: 4/day 20 mg ODT: 6/day Max total amphetamine dose (Age ≥ 21): 60 mg/day	Anti- hyperkinesis: Stimulants PA Form
Focalin®	NP	See Adderall® prior authorization criteria		
Jornay PM®	NP	See Adderall® prior authorization criteria	1/day	Anti-
lisdexamfetamine caps and chewables	NP	See Adderall® prior authorization criteria	1/day; Max total amphetamine dose (Age ≥ 21): 60mg/day	hyperkinesis: Stimulants PA Form
methamphetamine	NP	See Adderall® prior authorization criteria	4/day Max total amphetamine dose (Age ≥ 21): 60 mg/day	Anti- hyperkinesis: Stimulants PA
Methylin® solution	NP	See Adderall® prior authorization criteria		Form
methylphenidate chewables	NP	See Adderall® prior authorization criteria		<u>FOIIII</u>
methylphenidate patch	NP	See Adderall® prior authorization criteria	1/day	<u>Anti-</u>
methylphenidate ER 24hr capsules (generic for Aptensio XR, Ritalin LA)	NP	See Adderall® prior authorization criteria	1/day	hyperkinesis: Stimulants PA Form

June 1, 2025



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Prior Authorization Criteria Qty. Limits PA Form** methylphenidate ER OSM tablets (generic NP | See Adderall® prior authorization criteria See Concerta® Antifor Concerta® & hyperkinesis: Relexxii®) Stimulants PA methylphenidate XR **Form** ODT (generic for NP | See Adderall® prior authorization criteria 1/day Cotempla® XR ODT) Mydavis ER® NP See Adderall® prior authorization criteria 1/day Quillichew ER® NP See Adderall® prior authorization criteria 1/day Anti-Quillivant XR® NP See Adderall® prior authorization criteria 12 mL/day hyperkinesis: Relexxii® ER NP Stimulants PA See Adderall® prior authorization criteria 1/day **Form** Ritalin® NP See Adderall® prior authorization criteria 1/day Ritalin® LA NP See Adderall® prior authorization criteria 1/day 20 mg: 3/day 30 mg: 2/day Anti-All others: 4/day hyperkinesis: Zenzedi® NP | See Adderall® prior authorization criteria Stimulants PA Max total **Form** amphetamine dose (Age ≥ 21): 60mg/day **Antihyperkinesis: Non-Stimulants** 60 mg, 80 mg, 100 mg: 1/day Ρ atomoxetine All other strengths: **General PA** 2/day **Form** Ρ clonidine 12hr ER 4/day guanfacine ER Ρ 1/day • Diagnosis of Attention Deficit Hyperactivity Disorder (ADHD); AND · Patient is 6 years of age or older; AND 100 mg: 2/day Patient's blood pressure and heart rate will be assessed prior to therapy and monitored throughout therapy; AND Oelbree® Patient will not concomitantly use monoamine oxidase inhibitors (MAOIs); AND 150 mg: 2/day General PA Patient will not concomitantly use CYP1A2 substrates or CYP1A2 substrates with a narrow therapeutic range; AND 200 mg: 3/day Form · Patient is not pregnant; AND • Trial and failure, contraindication, or intolerance to 2 preferred antihyperkinesis stimulant and/or non-stimulant agents • Clinically valid reason why preferred guanfacine ER cannot be used Intuniv® 1/dav

June 1, 2025



		CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated	<i>t</i> .	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Onyda XR®	NP	 Diagnosis of attention deficit hyperactivity disorder (ADHD); AND Patient is 6 years of age or older; AND One of the following: Trial &failure, contraindication, or intolerance of 2 preferred non-stimulant antihyperkinesis agents; OR Patient is unable to swallow solid dosage forms 	4mL/day	
Strattera®	NP		60, 80, 100 mg: 1/day All others: 2/day	
	•	Agents for Narcolepsy		
modafinil	Р	 Diagnosis of ADD/ADHD; AND Contraindication, adverse reaction, or drug-drug interaction to ALL preferred antihyperkinesis agents; OR Daytime sleepiness/hypersomnolence occurring for at least 3 months; AND Diagnosis is associated with ONE of the following: 	2/day	Narcolepsy Agents PA Form
Provigil®	Р	See modafinil prior authorization criteria	2/day	1
Xyrem®	Р	 Enrolled in the Xyrem Program (1-866-997-3688); AND One of the following: Diagnosis of cataplexy associated with narcolepsy Diagnosis of excessive daytime sleepiness/hypersomnolence associated with narcolepsy occurring ≥ 3 months; AND Trial and failure, intolerance, or contraindication to modafinil; AND Hypersomnolence secondary to another sleep disorder, neurologic disorder, medical condition, or by medicine or substance use has been ruled out 	9 grams/day	
armodafinil	NP	 Daytime sleepiness/hypersomnolence occurring for at least 3 months; AND Diagnosis is associated with ONE of the following: Diagnosis of Narcolepsy Obstructive sleep apnea/hypopnea syndrome supported by a documented sleep study, AND Trial and failure (minimum duration 3 months with documented compliance) of Continuous Positive Airway Pressure (CPAP) or BiPAP device, unless contraindications Diagnosis of Shift Work Sleep Disorder; AND Statement of patient's work schedule showing a minimum of 6 hours work between 10 pm and 8 am; AND Hypersomnolence secondary to another sleep disorder, neurologic disorder, medical condition, or by medicine or substance use has been ruled out; AND Trial and failure, contraindication, or intolerance to modafinil 	50mg: 2/day 150mg, 200mg, 250mg: 1/day	Narcolepsy Agents PA Form



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Prior Authorization Criteria Qty. Limits PA Form** 50mg: 2/day 150mg, 200mg, Nuvigil® NP | See armodafinil prior authorization criteria 250mg: 1/day See Xyrem® prior authorization criteria; AND NΡ sodium oxybate 9 grams/day Trial and failure of Xyrem® • Daytime sleepiness/hypersomnolence occurring for at least 3 months; AND • Diagnosis is associated with ONE of the following: Diagnosis of Narcolepsy Obstructive sleep apnea/hypopnea syndrome supported by a documented sleep study, AND Sunosi® NP Trial and failure (minimum duration 3 months with documented compliance) of Continuous Positive Airway 1/dav Pressure (CPAP) or BiPAP device, unless contraindications; AND Hypersomnolence secondary to another sleep disorder, neurologic disorder, medical condition, or by medicine or substance use has been ruled out; AND • Trial and failure, contraindication, or intolerance to modafinil • Daytime sleepiness/hypersomnolence occurring for at least 3 months; AND • ONE of the following: o Diagnosis of cataplexy associated with narcolepsy; AND Trial and failure, contraindication, or intolerance to Xyrem Wakix[®] NP 2/day Diagnosis of excessive daytime sleepiness (EDS) associated with Narcolepsy; AND - Trial and failure, contraindication, or intolerance to modafinil; AND Hypersomnolence secondary to another sleep disorder, neurologic disorder, medical condition, or by medicine or substance use has been ruled out Enrolled in the Xywav Program (1-866-997-3688); AND · One of the following: Diagnosis of cataplexy associated with narcolepsy; AND - Clinically valid reason is given why the patient requires Xyway over Xyrem Diagnosis of excessive daytime sleepiness/hypersomnolence associated with narcolepsy occurring > 3 months; AND **Narcolepsy** Agents PA Xywav[®] NP - Trial and failure, intolerance, or contraindication to modafinil; AND 18 mL per day - Clinically valid reason is given why the patient requires Xywav over Xyrem Form



Diagnosis of idiopathic hypersomnia (IH) in patients ≥ 18 years of age; AND
 Trial and failure, intolerance, or contraindication to modafinil; AND

substance use has been ruled out

Hypersomnolence secondary to another sleep disorder, neurologic disorder, medical condition, or by medicine or

CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.						
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
		Antimigraine Preparations: CGRP Antagonists				
Aimovig®	Р	 Initial Criteria: Patient has a diagnosis of migraine with or without aura; AND Patient has ≥ 4 migraine days per month; AND Patient is utilizing prophylactic interventions (e.g., behavioral therapy, physical therapy, life-style modifications); AND Trial (duration ≥ 8weeks) and failure of TWO of the following oral medication classes, unless contraindicated: Antidepressants (i.e., amitriptyline, venlafaxine) Beta blockers (i.e., propranolol, metoprolol, timolol, atenolol) Antiepileptics (i.e., valproate, topiramate) Renewal Criteria: Patient has experienced positive response to therapy (e.g., decrease in the number, frequency, and/or intensity of headaches, improved function, decreased reliance on acute treatments for migraine headaches); AND Patient has absence of unacceptable toxicity (e.g., intolerable injection site pain or constipation) 	1 syringe/30 days	General PA Form		
Emgality® syringe & pen	P	 Initial Criteria: Diagnosis of episodic cluster headache; OR Diagnosis of migraine with or without aura; AND	1 syringe/month (120 mg for migraine and 300 mg for cluster headache)	General PA Form		



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Prior Authorization Criteria Qty. Limits PA Form** Initial Criteria: Diagnosis of migraine with or without aura; AND · One of one of the following: o Acute treatment of migraine, AND Medication will not be used in combination with another acute CGRP inhibitor; AND - Trial and failure or intolerance to TWO triptans (e.g., eletriptan, rizatriptan, sumatriptan) OR contraindication to Acute treatment: all triptans 1 dose pack (8 o Preventative treatment of migraine; AND tablets)/30 days Patient has ≥ 4 migraine days per month; AND Nurtec ODT® Ρ Patient is utilizing prophylactic interventions (e.g., behavioral therapy, physical therapy, life-style modifications); Prophylaxis: 2 dose packs (16 - Trial (duration ≥ 8weeks) and failure of TWO of the following oral medication classes, unless contraindicated: tablets)/30 days • Antidepressants (i.e., amitriptyline, venlafaxine) • Beta blockers (i.e., propranolol, metoprolol, timolol, atenolol) Antiepileptics (i.e., valproate, topiramate); AND Renewal Criteria: Patient has experienced positive response to therapy (e.g., decrease in the number, frequency, and/or intensity of headaches, improved function, decreased reliance on acute treatments for migraine headaches) Initial Criteria: **General PA** · Patient has a diagnosis of migraine with or without aura; AND Form • Patient has ≥ 4 migraine days per month; AND · Patient is utilizing prophylactic interventions (e.g., behavioral therapy, physical therapy, lifestyle modifications); AND • Trial (duration > 8weeks) and failure of TWO of the following oral medication classes, unless contraindicated: Qulipta® Antidepressants (i.e., amitriptyline, venlafaxine) 1/day o Beta blockers (i.e., propranolol, metoprolol, timolol, atenolol) Antiepileptics (i.e., valproate, topiramate); AND Renewal Criteria: Patient has experienced positive response to therapy (e.g., decrease in the number, frequency, and/or intensity of headaches, improved function, decreased reliance on acute treatments for migraine headaches) Initial Criteria: • Diagnosis of migraine with or without aura and will be used for the acute treatment of migraine, AND • Trial and failure or intolerance to TWO triptans (e.g., eletriptan, rizatriptan, sumatriptan) OR contraindication to all triptan; 1 box (10 tablets) / Ubrelvy® 30 days Medication will not be used in combination with another acute CGRP inhibitor Renewal Criteria: Patient has experienced a positive response to therapy (e.g., reduction in pain, photophobia, phonophobia, nausea) Ajovy® autoinjector See Aimovig prior authorization criteria; AND 3 injections/90 days and prefilled syringe Trial and failure of Aimovig and Emgality



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Prior Authorization Criteria Qty. Limits PA Form** Initial Criteria: Diagnosis of migraine with or without aura and will be used for the acute treatment of migraine, AND 60 mg/30 days **General PA** • Trial and failure or intolerance to Nurtec ODT and Ubrelvy; AND (6 devices) Zavzpret® Medication will not be used in combination with another acute CGRP inhibitor Form Renewal Criteria: Patient has experienced a positive response to therapy (e.g., reduction in pain, photophobia, phonophobia, nausea) **Antimigraine: Ergotamine Derivatives** Р Migranal® 8 mL/30 days dihydroergotamine Ρ 8 mL/30 days nasal spray General PA dihydroergotamine NP 8 mL/30 days Form injection NΡ Migergot® 15/30 days Trudhesa® NP See dihydroergotamine injection prior authorization criteria 1 package/30 days **Antimigraine: Barbiturate Combination Agents** **Quantity Limit Override Criteria for Butalbital-Containing Products: Butalbital-containing products have a quantity limit of 20 caps per 30 days. Requests for quantities greater than 20/30 will be approved if the following criteria is met: • Trial and failure of a tricyclic antidepressant (unless contraindicated); AND Trial and failure of divalproex sodium, sodium valproate, topiramate, frovatriptan, or a beta-blocker 20/30 days** Ρ butalbital/APAP APAP: 4 g/day 20/30 days** butalbital/APAP/ Ρ caffeine APAP: 4 g/day 20/30 days** Allzital® NP APAP: 4 g/day **General PA** butalbital/ASA/ Form · Allergy or intolerance to APAP 20/30 days** NP caffeine 20/30 days** Fioricet® NΡ APAP: 4 g/day 20/30 davs** Esgic® NP APAP: 4 g/day **Antimigraine: Selective 5-HT1 Agonists** Ρ eletriptan 6/30 days Ρ rizatriptan 12/30 days **General PA** Ρ rizatriptan ODT 12/30 days Form Ρ 9/30 days sumatriptan tabs sumatriptan vials Ρ 8 vials/30 days



CENTRAL NERVOUS SYSTEM

	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
zolmitriptan nasal spray	Р		6/30 days			
Frova®	NP		9/30 days			
frovatriptan	NP		9/30 days			
Imitrex Injectable®	NP		8 vials/30 days			
Imitrex Kit®	NP	• Clinically valid reason why the injectable vials cannot be used (NOTE: Patient convenience is NOT an approvable reason)	4/30 days			
Imitrex Nasal®	NP		6/30 days			
Imitrex® tablets	NP		9/30 days			
Maxalt®	NP		12/30 days			
Maxalt MLT®	NP		12/30 days			
naratriptan	NP		9/30 days			
Onzetra Xsail®	NP	 Patient has an allergy to an inactive ingredient found in the preferred sumatriptan containing agents; AND Patient has a contraindication, allergic reaction, or drug-drug interaction to preferred rizatriptan containing agents; AND Clinically valid reason why the patient requires a nasal powder (NOTE: Patient convenience is NOT an approval reason) 	16/30 days			
Relpax®	NP		6/30 days			
Reyvow [®]	NP	 Initial Criteria (3-month duration): Agent is being used for acute treatment of migraine with or without aura; AND Patient is 18 years of age or older; AND Trial and failure, contraindication, or intolerance to TWO triptans (e.g., eletriptan, rizatriptan, sumatriptan); AND Renewal Criteria: Patient has experienced a positive response to therapy (e.g., reduction in pain, photophobia, phonophobia, nausea) 	4/30 days	General PA Form		
sumatriptan autoinjector	NP	Clinically valid reason as to why the patient cannot use the injectable vials. (Note: Patient convenience is NOT an approvable reason)	4/30 days			
sumatriptan cartridge		Clinically valid reason as to why the patient cannot use the injectable vials. (Note: Patient convenience is NOT an approvable reason)				
sumatriptan nasal	NP		6/30 days	1		
sumatriptan/ naproxen	NP		9/30 days			
Tosymra®	NP		12/30 days			
Treximet®	NP		9/30 days			
zolmitriptan nasal spray and tablets	NP		6/30 days			
Zembrace Symtouch®	NP	 Patient has an allergy to an inactive ingredient found in the preferred sumatriptan containing agents; AND Patient has a contraindication, allergic reaction, or drug-drug interaction to preferred rizatriptan containing agents; AND Clinically valid reason why the patient requires an autoinjector device (NOTE: Patient convenience is NOT an approval reason) 	2 mL/30 days	General PA Form		
Zomig® nasal spray	NP	•	6/30 days			
Zomig® tablets	NP		6/30 days			

June 1, 2025



	CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
		Atypical Antipsychotic/SSRI Combos				

CLASS PRIOR AUTHORIZATION CRITERIA FOR PATIENTS WITH INTELLECTUAL AND DEVELOPMENTAL DISABILITIES (I/DD):

Antipsychotics prescribed for disorders related to Intellectual and Developmental Disabilities will be approved if ONE of the following is met:

- Prescribed by a Gold Card prescriber; OR
- Mental health assessment applicable to behavioral symptoms for which the medication is being prescribed; AND
 - Underlying physical condition such as pain, discomfort, and environmental issues have been evaluated, treated, and behavioral symptoms persist; AND
 - Non-pharmacological interventions have been tried and behavioral symptoms persist (e.g., behavior plan, crisis intervention, and stabilization strategies or training) and training and support have been provided to family or other caregivers; OR
- Short-term therapy (less than 90 days) has been prescribed; AND
 - o Behavioral symptoms are significant enough to place the person at potential risk or needing higher level of care or loss of community placement; OR
 - Continuation of existing therapy to address serious and ongoing behavioral symptoms while other prior authorization criteria are met; AND
 - o Efficacy and potential side effects to be monitored; AND
 - Need for requested medication will be evaluated once other non-pharmacological interventions have been tried

Note the following:

- Duration of short-term therapy is 90 days for antipsychotics
- Drug specific step therapy, NP criteria and Brand Medically Necessary Criteria will still apply to this patient population.
- The I/DD Worksheet can be found at: I/DD Prior Authorization Form

fluoxetine/ olanzapine	NP	 For diagnosis of depressive episodes associated with bipolar disorder; AND Refractory to treatment with components taken separately For diagnosis of major depressive disorder: Must have undergone an adequate trial of at least ONE agent in THREE of the following classes of antidepressants (unless contraindicated or intolerant to):	1/day	Atypical Antipsychotic PA form
Symbyax [®]	NP	See fluoxetine/olanzapine prior authorization criteria	1/day	



CENTRAL NERVOUS SYSTEM

Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.

Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
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Atypical Antipsychotics

CLASS PRIOR AUTHORIZATION CRITERIA FOR PATIENTS WITH INTELLECTUAL AND DEVELOPMENTAL DISABILITIES (I/DD):

Antipsychotics prescribed for disorders related to Intellectual and Developmental Disabilities will be approved if ONE of the following is met:

- Prescribed by a Gold Card prescriber; OR
- Mental health assessment applicable to behavioral symptoms for which the medication is being prescribed; AND
 - Underlying physical condition such as pain, discomfort, and environmental issues have been evaluated, treated, and behavioral symptoms persist; AND
 - Non-pharmacological interventions have been tried and behavioral symptoms persist (e.g., behavior plan, crisis intervention, and stabilization strategies or training) and training and support have been provided to family or other caregivers
- Short-term therapy (less than 90 days) has been prescribed; AND
 - o Behavioral symptoms are significant enough to place the person at potential risk or needing higher level of care or loss of community placement; **OR**
 - Continuation of existing therapy to address serious and ongoing behavioral symptoms while other prior authorization criteria are met; AND
 - o Efficacy and potential side effects to be monitored; AND
 - o Need for requested medication will be evaluated once other non-pharmacological interventions have been tried

Note the following:

- Duration of short-term therapy is 90 days for antipsychotics
- Drug specific step therapy, NP criteria and Brand Medically Necessary Criteria will still apply to this patient population.
- The I/DD Worksheet can be found at: I/DD Prior Authorization Form

Note: A list of ICD-10 to allow PA bypass for preferred atypical antipsychotics that require PA can be found at Appropriate Diagnosis for PA Bypass List

Abilify Asimtufii®	Р	 Patient is > 18 years of age; AND Patient has documented tolerance to the oral active ingredient 	1 injection/60 days	
Abilify Maintena®	Р	 Patient is ≥ 18 years of age; AND Patient has documented tolerance to the oral active ingredient 	1/30 days	
aripiprazole ODT	Р		1/day	
aripiprazole solution	Р		10 mL/day	
aripiprazole tablets	Р		1/day	<u>Atypical</u>
Aristada®	Р	 Patient is ≥ 18 years of age; AND Patient has documented tolerance to the oral active ingredient 	1064 mg: 1/60 days; All other strengths: 1/30 days	Antipsychotic PA form
Aristada® Initio	Р	 Patient is ≥ 18 years of age; AND Patient has documented tolerance to the oral active ingredient 	2.4 mL/60 days	
asenapine	NP	See lurasidone prior authorization criteria	2/day	1
clozapine	Р		1/day	1
Erzofri®	Р	 Patient is ≥ 18 years of age; AND Patient has documented tolerance to the oral active ingredient 	1 injection/28 days	



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Prior Authorization Criteria Qty. Limits PA Form** Patient is > 18 years of age; AND TennCare prescription claims history must indicate patient has been on Invega Sustenna® for 4 months OR Invega Trinza Invega Hafyera® 1 syringe/168 days for at least one three-month cycle **Atypical** Antipsychotic Patient is > 18 years of age; AND Invega Sustenna® Ρ 1 syringe/28 days PA form · Patient has documented tolerance to the oral active ingredient Patient is > 18 years of age; AND Р Invega Trinza® 1 syringe/76 days • TennCare prescription claims history must indicate patient has been on Invega Sustenna® for 4 months • Diagnosis of ONE of the following: o Agitation in dementia o Bipolar and manic disorders o Bipolar depression, bipolar maintenance, bipolar mania-acute, bipolar mixed states Brief psychotic disorder o Delusional disorder Depression with psychotic symptoms Drug-induced psychotic disorder with hallucinations o Impulse control disorders, including Oppositional Defiant Disorder and Intermittent Explosive Disorder Organic psychotic condition o Psychosis secondary to a medical condition, psychotic depression, psychotic disorders Schizoaffective disorder, schizoid/schizotypal personality disorder, schizophrenia, schizophrenic disorders **Atypical lurasidone** Ρ Substance-induced psychotic disorder, substance-induced withdrawal psychotic disorder 1/day Antipsychotic Severe refractory OCD or PTSD PA form o Tourette's/Severe tic disorder; OR • Diagnosis of major depressive disorder (MDD); AND o Atypical agents will be approved only as adjunctive treatment for MDD; AND o Adequate trial(4 - 6 weeks) of ONE agent from any of the following classes (unless contraindication or intolerance): - SSRIs - SNRIs TCAs New generation antidepressants (including bupropion, mirtazapine, etc.); OR • For patients without one of the above diagnoses: o May be approved if the physician can provide documented clinical evidence supporting the use of the requested medication for the requested indication Ρ olanzapine tablets 1/day Atypical olanzapine IM Antipsychotic See lurasidone prior authorization criteria 1/day injection PA form See lurasidone prior authorization criteria; AND olanzapine ODT • Patient is unable to swallow solid dosage forms or absorb medications through the GI tract; OR 1/day • Non-response due to noncompliance



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication **PDL Prior Authorization Criteria Qty. Limits PA Form** 6 mg: 2/day; All other paliperidone ER Ρ strengths: 1/day • Patient is ≥ 18 years of age; AND Р 1 injection/month Perseris® • Patient has documented tolerance to oral risperidone **Atypical** Ρ quetiapine 4/day **Antipsychotic** PA form Р See lurasidone prior authorization criteria 2/day quetiapine ER risperidone ODT Ρ See olanzapine ODT prior authorization criteria 2/day risperidone solution Ρ See lurasidone prior authorization criteria Ρ risperidone tabs 2/day 50, 75, 100, & 125 mg: 1 injection/30 days • Patient is ≥ 18 years of age; AND Atypical Uzedy • Documented tolerance to the oral active ingredient 150, 200, & 250 mg: Antipsychotic 1 injection/60 days PA form Vraylar® See lurasidone prior authorization criteria 1/day ziprasidone injection See lurasidone prior authorization criteria 2/day Ρ 2/day ziprasidone caps



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Qty. Limits Prior Authorization Criteria PA Form** · Approval of non-preferred atypical antipsychotics requires trial and failure of ONE preferred agent; AND • Diagnosis of ONE of the following: o Agitation in dementia o Bipolar and manic disorders o Bipolar depression, bipolar maintenance, bipolar mania-acute, bipolar mixed states Brief psychotic disorder o Delusional disorder Depression with psychotic symptoms Drug-induced psychotic disorder with hallucinations o Impulse control disorders, including Oppositional Defiant Disorder and Intermittent Explosive Disorder o Organic psychotic condition o Psychosis secondary to a medical condition, psychotic depression, psychotic disorders Schizoaffective disorder, schizoid/schizotypal personality disorder, schizophrenia, schizophrenic disorders Abilify® tablets NP 1/day Substance-induced psychotic disorder, substance-induced withdrawal psychotic disorder Severe refractory OCD or PTSD o Tourette's/Severe tic disorder; OR **Atypical** Diagnosis of major depressive disorder (MDD); AND Antipsychotic Atypical agents will be approved only as adjunctive treatment for MDD; AND PA form o Adequate trial(4 - 6 weeks) of ONE agent from any of the following classes (unless contraindication or intolerance): - SSRIs SNRIs TCAs New generation antidepressants (including bupropion, mirtazapine, etc.); OR • For patients without one of the above diagnoses: o May be approved if the physician can provide documented clinical evidence supporting the use of the requested medication for the requested indication See lurasidone prior authorization criteria; AND Abilify MyCite® 1/day Clinically valid reason why none of the other forms of aripiprazole cannot be used Caplyta® NP | See Abilify® tablets prior authorization criteria 1/day See Abilify® tablets prior authorization criteria; AND 12.5 & 25 mg: 2/day; clozapine ODT NP Patient is unable to swallow solid dosage forms or absorb medications through the GI tract; OR 100mg: 9/day; 150mg: 6/day; 200mg: 4/day Non-response due to noncompliance Clozaril® NP | See Abilify® tablets prior authorization criteria 1/day Patient is 18 years of age or older; AND Diagnosis of schizophrenia; AND Approval of non-preferred atypical antipsychotics requires trial and failure of ONE preferred agent; AND • Prescriber attests patient does NOT have any of the following: Cobenfy® 2/day Urinary retention o Moderate (Child-Pugh Class B) or severe (Child-Pugh Class C) hepatic impairment o Gastric retention



Untreated narrow-angle glaucoma

CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Qty. Limits PA Form Prior Authorization Criteria** Fanapt® NP See Abilify® tablets prior authorization criteria 2/day Geodon® NP | See Abilify® tablets prior authorization criteria 2/day Atypical Antipsychotic 6 mg: 2/day; All Invega® NP | See Abilify® tablets prior authorization criteria PA form others: 1/day Latuda® NP | See Abilify tablets prior authorization criteria Patient is ≥18 years of age; AND • One of the following: o Diagnosis of schizophrenia o Diagnosis of Bipolar I disorder and will be used for the acute treatment of manic or mixed episodes o Diagnosis of Bipolar I disorder and will be used as maintenance monotherapy treatment • Prescriber must attest that patient does not meet any of the following: o Patient is using opioids or has used a short-acting opioid in the last 7 days or a long-acting opioid in the last 14 days o Patient is undergoing acute opioid withdrawal; AND Lybalvi® NP 1/day Approval of non-preferred atypical antipsychotics requires trial and failure of ONE preferred agent; AND • Submission of medical records (e.g. chart notes) documenting ONE of the following: Atypical Patient has a BMI of 30 kg/m2 or greater; OR Antipsychotic o Patient has a BMI of 27 kg/m2 or greater with a weight-related comorbidity (e.g., dyslipidemia, hypertension, type 2 PA form diabetes, sleep apnea); OR o Patient has a documented history of weight gain of greater than or equal to 10% of their baseline weight after trial and failure of a preferred atypical antipsychotic; **OR** o Patient is stable on Lybalvi (minimum trial duration 4 weeks) and the request is for continuation of therapy Hallucinations and/or delusions associated with Parkinson's disease psychosis; AND Must be ≥18 years of age; AND · Trial of dose adjustment or withdrawal of anti-Parkinson medications (anticholinergics, amantadine, dopamine agonists, Nuplazid® NP 2/day COMT inhibitors, selegiline) prior to treatment with Nuplazid® • Trial and failure of ONE preferred agent **Note:** Coverage will not be approved for psychosis not related to Parkinson's disease 2mg, 5mg: 1/day; See lurasidone prior authorization criteria; AND NΡ Opipza® Clinically valid reason why none of the other forms of aripiprazole (e.g. aripiprazole ODT) cannot be used 10mg: 3/day Atypical See Abilify® tablets prior authorization criteria Antipsychotic Rexulti® NP 1/day Note: Rexulti used for the diagnosis of agitation in dementia does NOT require trial and failure of ONE preferred agent PA form Risperdal® NP See Abilify® tablets prior authorization criteria 2/day Patient is > 18 years of age; ANDa • Documented tolerance to the oral active ingredient; AND **Atypical** Risperdal Consta® One of the following: 2 vials/28 days NP o Diagnosis of Bipolar Disorder Antipsychotic o Clinically valid reason why the patient cannot use the preferred long-acting injectables PA form risperidone ER



injection

2 vials/28 days

See Risperdal Consta® prior authorization criteria

CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Prior Authorization Criteria Qty. Limits PA Form** Rvkindo® NP | See Risperdal Consta® prior authorization criteria 2 injections/28 days Saphris® NP | See Abilify tablets prior authorization criteria 2/day See Abilify® tablets prior authorization criteria; AND Secuado® NP Patient is unable to swallow solid dosage forms or absorb medications through the GI tract; OR 1/day • Non-response due to noncompliance **Atypical** Seroquel® NP See Abilify® tablets prior authorization criteria 4/day Antipsychotic Seroquel® XR NP See Abilify® tablets prior authorization criteria 2/day PA form See Abilify tablets prior authorization criteria; AND Versacloz® • Allergy or intolerance to inactive ingredient in clozapine ODT tab (i.e., dye, filler, excipient, etc); OR • Dose not achievable with ODT tab • Patient is > 18 years of age; AND Zyprexa® IM • Patient has documented tolerance to the oral active ingredient; AND 1/day injection • Trial and failure of ONE preferred atypical antipsychotic Zyprexa® tablets See Abilify® tablets prior authorization criteria NP 1/day **Atypical** 210mg, 300mg: Patient is > 18 years of age; AND **Antipsychotic** 1 injection/2 weeks; Zyprexa Relprevv® NP • Documented tolerance to the oral active ingredient; AND 450mg: PA form Clinically valid reason why the patient cannot use the preferred long-acting injectables 1 injection/month See Abilify® tablets prior authorization criteria; AND Zvprexa Zvdis® NP Patient is unable to swallow solid dosage forms or absorb medications through the GI tract; OR 1/day Non-response due to noncompliance Miscellaneous CNS Agents Diagnosis of Pseudobulbar Affect (PBA); AND • The following patient circumstances have been excluded: o Heart failure or high grade (second/third degree) atrioventricular block (AV) without an implanted pacemaker **General PA** Nuedexta® NP 2/day o Patient receiving drugs that prolong QT interval and are metabolized by CYP2D6 system Form o Prolonged QT interval (including congenital long QT syndrome) or a history of torsades de pointes o Concomitantly taking monoamine oxidase inhibitors (MAOIs) or have used a MAOI in the past 14 days Mood Stabilizers Unable to swallow; OR **General PA** Lamictal® ODT NP Unable to absorb medications through the GI tract Form



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL Prior Authorization Criteria Qty. Limits PA Form

Sedative Hypnotics

CLASS PRIOR AUTHORIZATION CRITERIA FOR PATIENTS WITH INTELLECTUAL AND DEVELOPMENTAL DISABILITIES (I/DD):

Sedative hypnotics prescribed for disorders related to Intellectual and Developmental Disabilities will be approved if ONE of the following is met:

- Prescribed by a Gold Card prescriber; OR
- Mental health assessment applicable to behavioral symptoms for which the medication is being prescribed; AND
 - Underlying physical condition such as pain, discomfort, and environmental issues have been evaluated, treated, and behavioral symptoms persist; AND
 - Non-pharmacological interventions have been tried and behavioral symptoms persist (e.g., behavior plan, crisis intervention, and stabilization strategies or training) and training and support have been provided to family or other caregivers
- Short-term therapy (less than 90 days) has been prescribed; AND
 - Behavioral symptoms are significant enough to place the person at potential risk or needing higher level of care or loss of community placement; OR
 - Continuation of existing therapy to address serious and ongoing behavioral symptoms while other prior authorization criteria are met; AND
 - o Efficacy and potential side effects to be monitored; AND
 - o Need for requested medication will be evaluated once other non-pharmacological interventions have been tried

Note the following:

- Drug specific step therapy, NP criteria and Brand Medically Necessary Criteria will still apply to this patient population.
- The I/DD Worksheet can be found at: I/DD Prior Authorization Form

· ·		<u> </u>		
doxepin concentrate 10mg/mL	Р			
eszopiclone	Р		14/30 days*	
ramelteon	Р		14/30 days*	
zaleplon	Р		14/30 days*	
zolpidem	Р		14/30 days*	
Ambien®	NP		14/30 days*	
Ambien CR®	NP		14/30 days*	
Belsomra®	NP		14/30 days*	General PA
Dayvigo®	NP	 Diagnosis of insomnia characterized by difficulties with sleep onset and/or sleep maintenance; AND Patient is 18 years of age or older; AND Narcolepsy and other insomnia related disorders have been ruled out (e.g., movement, breathing, psychiatric disorders, medications); AND Trial (at least 3 weeks) of non-pharmacological therapies (e.g., stimulus control, sleep restriction, sleep hygiene measures and relaxation therapy); AND Trial and failure, contraindication, or intolerance of 2 preferred agents; AND Patients who are pregnant should be registered in the Dayvigo® pregnancy registry 	14/30 days*	<u>Form</u>
Doral®	NP	See Halcion® prior authorization criteria	14/30 days*	
doxepin soln	NP	Documented trial/failure (defined as ≥ 1 week) at an appropriate dose of the doxepin 10mg/mL concentrated solution	14/30 days*	
Edluar®	NP	Patient is unable to swallow solid dosage forms	14/30 days*	
estazolam	NP	See flurazepam prior authorization criteria	14/30 days*	Anti-anxiety



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Prior Authorization Criteria Qty. Limits PA Form** · Diagnosis of Insomnia; AND Form • Medical documentation that rules out other insomnia related disorders (e.g., movement, breathing, psychiatric disorders and medication); AND Documented trial (at least 3 weeks) of non-pharmacological therapies (e.g., stimulus control, sleep restriction, sleep flurazepam NP hygiene measures and relaxation therapy); AND 14/30 days* Use of 2 preferred agents, unless patient has a contraindication or allergy; AND • Due to increased risk of toxicity, patient should not be pregnant; AND Will not be taken concurrently with CNS stimulants, opiates, carisoprodol, meprobamate or barbiturates; AND • Due to increased risk of dependency, patient does not have a history of alcohol or drug dependence/abuse Diagnosis of Insomnia; AND Medical documentation that rules out other insomnia related disorders (e.g., movement, breathing, psychiatric disorders & medication/substance use); AND Documented trial (at least 3 weeks) of non-pharmacological therapies (e.g., stimulus control, sleep restriction, sleep hygiene measures & relaxation therapy); AND Halcion® 14/30 days* • Use of 2 preferred agents, unless patient has a contraindication or allergy; AND Clinical reason as to why patient cannot use generic equivalent; AND • Due to increased risk of toxicity, patient should not be pregnant; AND • Will not be taken concurrently with CNS stimulants, opiates, carisoprodol, meprobamate or barbiturates; AND • Due to increased risk of dependency, patient does not have a history of alcohol OR drug dependence/abuse • Treatment of non-24-hour sleep wake disorder (non-24 or N24) in members who are unable to distinguish between light and darkness in both eyes; OR Treatment of nighttime sleep disturbances in Smith-Magenis Syndrome (SMS) in patients 16 years of age and older; AND Hetlioz® capsules NP • Trial and failure or contraindication to melatonin; AND 30/60 days* • Patient will not take any of the following: Strong CYP1A2 inhibitors (e.g., fluvoxamine) Strong CYP3A4 inducers (e.g., rifampin) • Treatment of nighttime sleep disturbances in Smith-Magenis Syndrome (SMS); AND Patient is at least 3 years of age but not greater than 15 years of age; AND **General PA** Trial and failure or contraindication to melatonin: AND Form 5 mL per day Hetlioz® suspension NP Patient is unable to swallow/absorb medications through the GI tract; AND 158 mL/60 days* • Patient will not take any of the following: Strong CYP1A2 inhibitors (e.g., fluvoxamine) Strong CYP3A4 inducers (e.g., rifampin) Intermezzo® NΡ 14/30 days* NP 14/30 days* Lunesta® NP 14/30 days* Rozerem® NP 14/30 davs* quazepam See flurazepam prior authorization criteria



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
Quviviq®	NP	 Patient must 18 years of age or older; AND Diagnosis of insomnia characterized by difficulties with sleep onset and/or sleep maintenance; AND Narcolepsy and other insomnia related disorders have been ruled out (e.g., movement, breathing, psychiatric disorders, medications); AND Trial (at least 3 weeks) of non-pharmacological therapies (e.g., stimulus control, sleep restriction, sleep hygiene measures and relaxation therapy); AND Trial and failure, contraindication, or intolerance of 2 preferred agents; AND Patient should not have any of the following diagnoses: Narcolepsy, COPD, or moderate to severe OSA; AND Concurrently not taking CNS stimulants, opiates, carisoprodol, meprobamate or barbiturates; AND Due to increased risk of dependency, patient does not have a history of alcohol or drug dependence/abuse; AND Patients who are pregnant should be registered in the Quviviq® pregnancy registry 	14/30 days*		
Restoril®	NP	See Halcion® prior authorization criteria	14/30 days*	Anti-anxiety Form	
Rozerem®	NP		14/30 days*		
tasimelteon capsules		See Hetlioz capsules prior authorization criteria; AND Clinically valid reason why Hetlioz® cannot be used	30/60 days*		
tasimelteon suspension	NP	See Hetlioz suspension prior authorization criteria; AND • Clinically valid reason why Hetlioz® cannot be used	5 mL per day 158 mL/60 days*		
temazepam	NP	See flurazepam prior authorization criteria	14/30 days*	Anti-anxiety	
triazolam	NP	See flurazepam prior authorization criteria	14/30 days*	<u>Form</u>	
zolpidem ER	NP		14/30 days*	General PA	
zolpidem tartrate SL	NP		14/30 days*	<u>Form</u>	
* For children, larger	quan	tities may be approved as medically necessary.			
		Skeletal Muscle Relaxants			
Amrix ®	NP	 Diagnosis of an FDA-approved indication; AND Provider must provide peer-reviewed medical literature documenting why the requested drug for the requested indication is the only appropriate choice versus the preferred cyclobenzaprine 	1/day		
baclofen solution	NP	 Diagnosis of spasticity with flexor spasms and concomitant pain, clonus, and/or muscular rigidity (e.g., multiple sclerosis, spinal cord injury, other spinal cord disease); AND Documented inability to swallow baclofen tablets 	16 mL/day		
baclofen suspension	NP	 Diagnosis of spasticity with flexor spasms and concomitant pain, clonus, and/or muscular rigidity (e.g., multiple sclerosis, spinal cord injury, other spinal cord disease); AND Documented inability to swallow baclofen tablets; AND Trial and failure of baclofen solution 	16 mL/day	General PA Form	
carisoprodol	NP	 Patient is 16 years of age or older; AND Contraindication, drug to drug interaction, or history of toxic side effects that will cause immediate or long-term damage with ALL preferred skeletal muscle relaxants; AND 	4/day		

June 1, 2025



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication **PDL Prior Authorization Criteria Qty. Limits PA Form** • Patient does not have a history of, or received treatment for, drug dependency or drug abuse; AND Prescriber must have checked the Tennessee Controlled Substance Database for this patient within the last 30 days; AND · Patient is not concurrently utilizing any other opioid therapy cyclobenzaprine ER NP | See Amrix® prior authorization criteria 1/day Fleqsuvy® NΡ See baclofen suspension prior authorization criteria 16 mL/day Lyvispah® NP See baclofen suspension prior authorization criteria 4 packets/day • Diagnosis of an FDA-approved indication; AND Norgesic Forte® NP · Provider must provide peer-reviewed medical literature documenting why the requested drug for the requested indication is the only appropriate choice versus the preferred agents Soma® NP | See carisoprodol prior authorization criteria 4/day



CENTRAL NERVOUS SYSTEM Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL Prior Authorization Criteria Qty. Limits PA Form Typical Antipsychotics

CLASS PRIOR AUTHORIZATION CRITERIA FOR PATIENTS WITH INTELLECTUAL AND DEVELOPMENTAL DISABILITIES (I/DD):

Antipsychotics prescribed for disorders related to Intellectual and Developmental Disabilities will be approved if ONE of the following is met:

- Prescribed by a Gold Card prescriber; OR
- There has been a mental health assessment applicable to behavioral symptoms for which the medication is being prescribed; AND
 - Underlying physical condition such as pain, discomfort, and environmental issues have been evaluated, treated, and behavioral symptoms persist; AND
 - Non-pharmacological interventions have been tried and behavioral symptoms persist (e.g., behavior plan, crisis intervention, and stabilization strategies or training) and training and support have been provided to family or other caregivers; OR
- Short-term therapy (less than 90 days) has been prescribed; AND
 - Behavioral symptoms are significant enough to place the person at potential risk or needing higher level of care or loss of community placement; OR
 - Continuation of existing therapy to address serious and ongoing behavioral symptoms while other prior authorization criteria are met; AND
 - o Efficacy and potential side effects to be monitored; AND
 - Need for requested medication will be evaluated once other non-pharmacological interventions have been tried

Note the following:

- Drug specific step therapy, NP criteria and Brand Medically Necessary Criteria will still apply to this patient population.
- The I/DD Worksheet can be found at: I/DD Prior Authorization Form

,		·	
chlorpromazine	Р		
fluphenazine	Р		
haloperidol	Р		
loxapine	Р		
perphenazine	Р		
pimozide	Р		General PA Form
thioridazine	Р		<u>101111</u>
thiothixene	Р		
trifluoperazine	Р		
molindone	NP		
Orap®	NP		



DERMATOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
		Acne Agents, Topical (Covered for recipients < 21 years old only)		•	
benzoyl peroxide 2.5%, 5%, 10% (excluding cleanser, gel, microspheres, and towelettes)	Р		1 package/Rx		
clindamycin phosphate (excluding foam, lotion, & 75 mL bottle of gel)	Р		1 package/Rx		
clindamycin/benzoyl peroxide gel	Р		1 package/Rx		
erythromycin (excluding swab & gels)	Р		1 package/Rx		
sodium sulfacetamide/ sulfur	Р		1 package/Rx		
Aczone®	NP	 Patient is at least 12 years of age and less than 21 years of age; AND Patient has a diagnosis of acne vulgaris; AND Clinically valid reason why generic dapsone gel cannot be used 	1 package/Rx	General PA	
Amzeeq [®]	NP	 Diagnosis of non-nodular moderate to severe acne vulgaris; AND Patient is at least 9 years of age and less than 21 years of age; AND Trial and failure, contraindication, or intolerance to ALL the following: 2 preferred agents minocycline capsules; AND Prescriber must provide peer-reviewed medical literature documenting why the requested drug for the requested indication is the only appropriate choice versus the preferred agents 	1 package/28 days	Form	
benzoyl peroxide (excluding preferred products)	NP		1 package/Rx		
Cabtreo®	NP	 Patient is at least 12 years of age and less than 21 years of age; AND Patient has a diagnosis of acne vulgaris; AND Provider must provide peer-reviewed medical literature documenting why the requested drug for the requested indication is the only appropriate choice versus the preferred agents 	1 package/Rx		
clindamycin (excluding preferred products)	NP		1 package/Rx		
dapsone gel	NP	 Patient is at least 12 years of age and less than 21 years of age; AND Patient has a diagnosis of acne vulgaris; AND Clinically valid reason why the preferred agents cannot be used 	1 package/Rx		



		DERMATOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
dermatological kits	NP	 Trial and failure of 3 preferred agents; AND Trial and failure of the individual components of the kit 	1 package/Rx	
erythromycin/benzol peroxide	NP		1 package/Rx	
erythromycin swab & gel	NP		1 package/Rx	
sulfacetamide suspension	NP		1 package/Rx	
Winlevi®	NP	 Diagnosis of acne vulgaris; AND Patient is at least 12 years of age and less than 21 years of age; AND Trial and failure, contraindication, or intolerance of 2 preferred agents; AND Prescriber provides peer-reviewed medical literature documenting why the requested drug for the requested indication is the only appropriate choice versus the preferred agents 	1 tube/30 days	General PA Form
All branded single agent and combination products of benzoyl peroxide, clindamycin, erythromycin, and sodium sulfacetamide	NP		1 package/Rx	
		Agents for Burns, Topical		
silver sulfadiazine	Р		1 package/Rx	
SSD®	Р		1 package/Rx]
mafenide	NP		1 package/Rx	General PA Form
Silvadene®	NP		1 package/Rx	
Sulfamylon®	NP		1 package/Rx	7
		Agents for Rosacea, Topical (Covered for recipients < 21 years old only)		-
Finacea® foam	Р		50 g/Rx	
metronidazole cream, lotion, and gel	Р		60 g/Rx	General PA
brimonidine gel	NP		30 g/Rx	Form
ivermectin cream	NP		45 g/Rx	
Mirvaso®	NP		30 g/Rx	



		DERMATOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Noritate® cream	NP		60 g/Rx	
Rhofade®	NP	 Patient age < 21 years of age; AND Patient has a diagnosis of persistent facial erythema associated with rosacea; AND Trial and failure, or contraindication, of 2 of the following: brimonidine, ivermectin, tetracycline, minocycline, doxycycline, erythromycin, clindamycin, benzoyl peroxide; AND Trial and failure of 2 preferred topical agents for rosacea 	30 g/30 days	
		Anesthetics, Topical		
lidocaine (excluding lotion, solution, kits)	Р		1 tube/Rx	
lidocaine patch 5%	Р	Diagnosis of post-herpetic neuralgia	3/day	
lidocaine/prilocaine	Р		30 g/Rx	
ZTLido®	Р	Diagnosis of Postherpetic neuralgia	3/day	
lidocaine kits	NP	 Diagnosis of FDA-approved indication; AND Clinically valid reason why the preferred topical anesthetics cannot be used; AND For combination kits, trial and failure of individual agents 		General PA Form
lidocaine/ hydrocortisone	NP	 Diagnosis of FDA-approved indication; AND Clinically valid reason why the preferred topical anesthetics cannot be used 	1 package/Rx	
Pliaglis®	NP		1 package/Rx	
Pramosone® 2.5-1% lotion	NP		1 package/Rx	
		Antibiotics, Topical		
mupirocin ointment	Р		44 g/Rx	General PA
mupirocin cream	NP		30 g/Rx	<u>Form</u>
		Antifungal Agents, Topical		•
ciclopirox cream	Р		1 package/Rx	
ciclopirox solution 8%	Р		1 -0-/	7
clotrimazole 1% cream & soln (OTC)	Р		1 package/Rx	
clotrimazole 1% cream (Rx)	Р		1 package/Rx	General PA
clotrimazole/ betamethasone	Р		1 package/Rx	Form
ketoconazole (cream and shampoo)	Р		1 package/Rx	
nystatin/ triamcinolone	Р		1 package/Rx	



DERMATOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. PDL Medication **Prior Authorization Criteria Qty. Limits PA Form** Ρ 120 g/Rx nystatin powder Vusion® Р 1 package/Rx ciclopirox gel and NP 1 package/Rx suspension ciclopirox nail kit NP • Clinically valid reason for why the preferred topical ciclopirox 8% solution cannot be used clotrimazole 1% NP 1 package/Rx solution (Rx) NP econazole 1 package/Rx NP Ertaczo® 1 package/Rx • Diagnosis of mild to moderate onychomycosis of the toenails; AND Jublia® NP 1 package/Rx • Trial and failure, contraindication, or intolerance to the preferred topical ciclopirox 8% solution Klayesta® NP 1 package/Rx luliconazole NP 1 package/Rx General PA Luzu® NP 1 package/Rx Form miconazole/zinc/ NP • Clinically valid reason for why the preferred Vusion cannot be used 1 package/Rx petrolatum naftifine gel NP 1 package/Rx Nyamyc® NP 1 package/Rx NP oxiconazole 1 package/Rx Oxistat® NP 1 package/Rx tavaborole soln See Jublia® prior authorization criteria 1 package/Rx Antineoplastics, Topical diclofenac 3% gel Ρ • Diagnosis of actinic keratosis 1 package/Rx fluorouracil 5% Р 1 package/Rx cream Р 1 package/Rx imiquimod Targretin® Р 1 package/Rx bexarotene NP 1 package/Rx Efudex® NP 1 package/Rx fluorouracil 0.5% NP 1 package/Rx cream



		DERMATOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Hyftor®	NP	Initial Criteria (4-month duration): Diagnosis of facial angiofibroma associated with tuberous sclerosis complex; AND Patient is 6 years of age or older; AND Prescribed by or in consultation with a dermatologist or neurologist; AND Patient is not a candidate for laser therapy or surgical treatments Renewal Criteria: Documentation of positive clinical response to therapy (e.g., improvement in size or redness of facial angiofibroma)	30 g/month	General PA Form
Valchlor®	NP	Diagnosis of stage IA or IB mycosis fungoides-type cutaneous T-cell lymphoma; AND Patient has received skin directed therapy	1 package/Rx	
Zyclara®	NP	 Diagnosis of actinic keratosis; OR Diagnosis of basal cell carcinoma 	1 package/Rx	
		Antipruritics/Antihistamines, Topical		
doxepin cream	NP	 Patient has moderate pruritus due to various forms of eczematous dermatitis, including atopic dermatitis and lichen simplex chronicus; AND Inadequate response, intolerance, or contraindication to BOTH of the following: A topical corticosteroid An oral antihistamine (first or second generation) or a topical antihistaminic agent 	45g/90 days	General PA Form
Prudoxin®	NP	See doxepin cream prior authorization criteria	45g/90 days	1
Zonalon®	NP	See doxepin cream prior authorization criteria	45g/90 days	
		Antipsoriatics, Oral		
acitretin	NP	 Patient has a diagnosis of severe psoriasis; AND Minimum duration of a 4-week trial and failure, contraindication, or intolerance to at least two of the following: Corticosteroids (e.g., betamethasone, clobetasol) Vitamin D analogs (e.g., calcitriol, calcipotriene) Tazarotene Calcineurin inhibitors (e.g., tacrolimus, pimecrolimus) Prescriber attests to each of the following: Patient does-NOT have impaired liver or kidney function, or abnormally elevated lipid levels Patient will NOT be receiving concomitant methotrexate (due to risk of hepatitis) or tetracyclines (due to risk of increased intracranial pressure) If applicable, appropriate laboratory assessments and counseling have been conducted regarding risks associated with pregnancy Note: Will not be covered for the diagnosis of acne or rosacea for recipients ≥ 21 years of age. 	10 mg (3/day); 17.5, 22.5, & 25 mg (2/day)	General PA Form
methoxsalen	NP	 Diagnosis of severe, recalcitrant, disabling psoriasis supported by biopsy; AND Minimum duration of a 4-week trial and failure, contraindication, or intolerance to at least two of the following: Corticosteroids (e.g., betamethasone, clobetasol) Vitamin D analogs (e.g., calcitriol, calcipotriene) Tazarotene Calcineurin inhibitors (e.g., tacrolimus, pimecrolimus) 		



		DERMATOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Antipsoriatics, Topical		
calcipotriene cream	Р	Trial and failure, contraindication, or intolerance to ≥ 1 topical steroid	1 package/Rx	
calcipotriene foam		Trial and failure, contraindication, or intolerance to ≥ 1 topical steroid	1 package/Rx	
calcipotriene scalp soln	Р	Trial and failure, contraindication, or intolerance to ≥ 1 topical steroid		
Sorilux®	Р	Trial and failure, contraindication, or intolerance to ≥ 1 topical steroid	1 package/Rx	
Taclonex®	Р	Trial and failure, contraindication, or intolerance to ≥ 1 topical steroid		_
tazarotene 0.1% cream	Р	 Diagnosis of psoriasis; OR Diagnosis of acne in patients less than 21 years of age 	1 package/Rx	
calcipotriene ointment	NP	 Trial and failure, contraindication, or intolerance to ≥ 1 topical steroid; AND Trial and failure, contraindication, or intolerance to a preferred topical antipsoriatic agent 	1 package/Rx	
calcitriol ointment	NP	See calcipotriene foam prior authorization criteria	1 package/Rx	
calcipotriene/ betamethasone	NP	See calcipotriene ointment prior authorization criteria	1 package/Rx	
Calcitrene® ointment		See calcipotriene ointment prior authorization criteria	1 package/Rx	
Duobrii®	NP	 Patient has a diagnosis of plaque psoriasis; AND Trial and failure, contraindication, or intolerance to at least one topical steroid; AND Clinically valid reason why the preferred individual components cannot be taken separately 	200 mg/30 days	
Enstilar®	NP	See calcipotriene ointment prior authorization criteria	1 package/Rx	
Sorilux®	NP	See calcipotriene ointment prior authorization criteria	1 package/Rx	General PA Form
tazarotene 1% gel	NP	 Diagnosis of psoriasis AND Both of the following: Trial and failure, contraindication, or intolerance to at least one topical steroid; Trial and failure, contraindication, or intolerance to a preferred topical antipsoriatic agent; OR Diagnosis of acne in patients less than 21 years of age; AND Trial and failure, contraindication, or intolerance to TWO preferred topical retinoids 	1 package/Rx	- <u>FOIIII</u>
Vtama®	NP	Initial Criteria: Diagnosis of plaque psoriasis; AND Minimum duration of a 4-week trial and failure, contraindication, or intolerance to at least two of the following: Corticosteroids (e.g., betamethasone, clobetasol) Vitamin D analogs (e.g., calcitriol, calcipotriene) Tazarotene Calcineurin inhibitors (e.g., tacrolimus, pimecrolimus; OR Diagnosis of atopic dermatitis and Both of the following: Trial and failure, contraindication, or intolerance to a topical corticosteroid AND a topical calcineurin inhibitor; AND Prescribed by, or in consultation with, a dermatologist Renewal Criteria: Documentation of positive clinical response to therapy as evidenced by one of the following: Reduction in the body surface area (BSA) involvement from baseline Improvement in symptoms (e.g., pruritus, inflammation) from baseline	60 grams/28 days	



	DERMATOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
Zoryve® 0.3% cream	NP	 Initial Criteria: Diagnosis of mild to plaque psoriasis; AND Patient is 6 years of age or older; AND Patient does not have moderate to severe liver impairment (Child-Pugh B or C); AND Request is for Zoryve 0.3% cream; AND Trial and failure, contraindication, or intolerance to TWO preferred topical antipsoriatic agents Renewal Criteria: Patient continues to be monitored for liver impairment; AND Documented clinical improvement in response to treatment (e.g. reduction in itch, rash, inflammation) 	60 grams/28 days	General PA Form		
		Antiseborrheic Agents				
selenium sulfide 2.5% lotion	Р		1 package/Rx			
Zoryve® 0.3% topical foam	NP	Initial Criteria: (3-month duration) Diagnosis of seborrheic dermatitis; AND Patient is 9 years of age or older; AND Patient does not have moderate to severe liver impairment (Child-Pugh B or C); AND Trial and failure, contraindication, or intolerance to BOTH of the following agents: Topical antifungals (ketoconazole, ciclopirox, miconazole, clotrimazole) Topical corticosteroids Renewal Criteria: Patient continues to be monitored for liver impairment; AND Documented clinical improvement in response to treatment (e.g., decreased erythema, scaling, inflammation); AND Patient does not have any treatment limiting adverse effects	1 can (60 gr)/30 days	General PA Form		
	•	Antivirals, Topical				
acyclovir 5% ointment	Р		1 tube/Rx	General PA		
penciclovir cream	Р		1 tube/Rx	<u>Form</u>		
acyclovir cream	NP		1 tube/Rx			
Denavir® cream	NP		1 tube/Rx	_		
Xerese®	NP	 Patient must be 6 years of age and older; AND Diagnosis of recurrent herpes labialis; AND Trial and failure of the individual components of the kit 	1 tube/Rx	General PA Form		
Zovirax® cream	NP	·	1 tube/Rx			
Zovirax® ointment	NP		1 tube/Rx			
		Atopic Dermatitis, Topical				
Elidel®	Р		1 package/Rx			
tacrolimus ointment	Р		1 package/Rx			



DERMATOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
Eucrisa®	NP	 Diagnosis of atopic dermatitis; AND One of the following: Patient is ≥ 2 years and meets BOTH of the following; AND Trial and failure, contraindication, or intolerance to BOTH of the following:	1 tube/month	General PA Form	
Opzelura®	NP	Initial Criteria: One of the following: Diagnosis of mild to moderate atopic dermatitis (3-month approval duration) and BOTH of the following: Patient is not immunocompromised; AND Opzelura will only be used for short-term and/or non-continuous chronic treatment; OR Diagnosis of Nonsegmental Vitiligo (12-month approval duration); AND Patient is 12 years of age or older; AND Patient is not breastfeeding; AND Trial and failure, contraindication, or intolerance to a topical corticosteroid; AND Trial and failure, contraindication, or intolerance to a topical calcineurin inhibitor; AND Prescriber attests to each of the following: Patient is not concurrently taking biologic agents (i.e., adalimumab, anakinra, etanercept, rituximab, tocilizumab, infliximab, abatacept) OR potent immunosuppressants (i.e., azathioprine, cyclosporine); AND Benefits of using this agent outweigh the heart-related or cardiovascular risk factors; AND Risk of malignancy has been considered and it has been determined that Jak Kinase inhibitor therapy is appropriate Renewal Criteria: Positive response to therapy [e.g., reduction in symptoms (itch, rash, etc.), re-pigmentation, etc.]	240 g/month	Topical Immuno- modulators PA Form	
pimecrolimus	NP	 Patient must have a diagnosis of atopic dermatitis; AND Trial and failure of 1 preferred agent (e.g., Elidel® or tacrolimus ointment) 	1 package/Rx		
Zoryve® 0.15% cream	NP	Initial Criteria: Diagnosis of mild to moderate atopic dermatitis; AND Patient is 6 years of age or older; AND Patient does not have moderate to severe liver impairment (Child-Pugh B or C); AND Request is for Zoryve 0.15% cream; AND Trial and failure, contraindication, or intolerance to ONE topical corticosteroid; AND Trial and failure, contraindication, or intolerance to ONE topical calcineurin inhibitor Renewal Criteria: Patient continues to be monitored for liver impairment; AND Documented clinical improvement in response to treatment (e.g. reduction in itch, rash, inflammation)	60 gram/28 days		



		DERMATOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Emollients		
ammonium lactate (Rx & OTC)	Р		1 package/Rx	General PA Form
		Genital Warts		
imiquimod	Р		1 package/Rx	
Condylox®	Р		1 package/Rx	General PA
Imiquimod pump	NP		1 package/Rx	<u>Form</u>
Veregen [®]	NP		1 package/Rx	
Zyclara®	NP		1 package/Rx	
		Keratolytic Agents		
generic urea products	Р		1 package/Rx	
generic salicylic acid products	Р		1 package/Rx	General PA
brand urea products	NP		1 package/Rx	<u>Form</u>
brand salicylic acid products	NP		1 package/Rx	
		Pediculocides/Scabicides		
Natroba®	Р		2 bottles/Rx	
permethrin	Р		2 tubes/Rx	
VanaLice®	Р		1 bottle/Rx	
Crotan®	NP	 Patient is being treated for scabies or pruritis; AND Patient has tried/failed permethrin (unless patient has a contraindication) 	1 bottle/Rx	General PA
ivermectin lotion	NP		1 tube/Rx	<u>Form</u>
malathion	NP		2 bottles/Rx	
Ovide®	NP		2 bottles/Rx	
Sklice®	NP		1 tube/Rx	
spinosad	NP		2 bottles/Rx	
		Retinoids, Oral		
Absorica® & Absorica LD®	NP	 Diagnosis of chronic myelogenous leukemia, head or neck cancer, ichthyosis, keratosis follicularis, neuroblastoma, or pityriasis rubra pilaris will be reviewed on a case-by-case basis; OR Diagnosis of severe recalcitrant nodular acne AND Patient is < 21 years of age (will not be covered for acne or rosacea for recipients ≥ 21 years of age) Note: Active registration and compliance with the iPLEDGE program is required by prescriber, patient, and pharmacy. 		General PA
Accutane®	NP	See Absorica® prior authorization criteria		Form
Amnesteem®	NP	See Absorica® prior authorization criteria		
Claravis®	NP	See Absorica® prior authorization criteria		
isotretinoin	NP	See Absorica® prior authorization criteria		
Zenatane®	NP	See Absorica® prior authorization criteria		



		DERMATOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indic	ated.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Retinoids, Topical		
adapalene	Р	See tretinoin prior authorization criteria	1 package/Rx	
tazarotene 0.1% cream	Р	See Tazorac® prior authorization criteria (Topical Antipsoriatics section)	1 package/Rx	
tretinoin cream	Р	 Patient is < 21 years old; AND Diagnosis of acne, keratosis follicularis, verruca plana, or actinic keratosis; OR Patient is ≥ 21 years old: AND Diagnosis of keratosis follicularis (1 year approval duration); OR Diagnosis of verruca plana (2-month approval duration); OR Diagnosis of actinic keratosis for the prevention of future lesions (1 year approval duration) Note: Will not be covered for patients > 21 years old with a diagnosis of acne 	1 package/Rx	General PA Form
adapalene/benzoyl peroxide	NP	See tretinoin prior authorization criteria In addition, non-preferred criteria and trial and failure of individual components is required.	1 package/Rx	
Altreno®	NP	See Aklief® prior authorization criteria	1 package/Rx	
Atralin®	NP	See tretinoin prior authorization criteria	1 package/Rx	
Arazio®	NP	 Patient is 9 years of age or older and less than 21 years of age; AND Diagnosis of acne; AND Patient is not pregnant; AND Trial and failure, contraindication, or intolerance to 2 preferred agents; AND Clinically valid reason why the requested drug is the only appropriate choice versus the preferred agents 	1 package/28 days	General PA Form
clindamycin/tretinoin	NP	See tretinoin prior authorization criteria	1 package/Rx	
Fabior®	NP	See Tazorac® prior authorization criteria (Topical Antipsoriatics section)	1 package/Rx	
Retin A®	NP	See tretinoin prior authorization criteria	1 package/Rx	
Retin A Micro®	NP	See tretinoin prior authorization criteria	1 package/Rx	
tretinoin gel	NP	See tretinoin prior authorization criteria	1 package/Rx	
Ziana®	NP	See tretinoin prior authorization criteria		
		Topical Steroids: Least Potent		
hydrocortisone 0.5% cream and ointment (Rx & OTC)	Р		1 package/Rx	
hydrocortisone 1% cream, lotion, gel, and ointment (Rx & OTC)	Р		1 package/Rx	General PA Form
hydrocortisone 2.5% cream, lotion, and ointment	Р		1 package/Rx	



		DERMATOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Topical Steroids: Mild		1
desonide 0.05% cream& ointment	Р		1 package/Rx	
fluocinolone 0.01% cream, oil, solution	Р		1 package/Rx	
Synalar® 0.01% solution	NP		1 package/Rx	
	_	Topical Steroids: Lower Mid-Strength		_
betamethasone dipropionate 0.05% lotion	Р		1 package/Rx	
betamethasone valerate 0.1% cream	Р		1 package/Rx	
Locoid Lipocream	Р		1 package/Rx	
clocortolone 0.1% cream and pump	NP		1 package/Rx	
desonide 0.05% lotion	NP		1 package/Rx	General PA Form
hydrocortisone butyrate 0.1% cream, lotion, ointment, solution	NP		1 package/Rx	
hydrocortisone valerate 0.2% cream	NP		1 package/Rx	
Locoid® lotion	NP		1 package/Rx	
Pandel® 0.1% cream	NP		1 package/Rx	
		Topical Steroids: Mid-Strength		,
triamcinolone acetonide 0.1% cream	Р		1 package/Rx	General PA
hydrocortisone valerate 0.2% ointment	NP		1 package/Rx	<u>Form</u>
		Topical Steroids: Upper Mid-Strength		•
betamethasone valerate 0.1% ointment	Р		1 package/Rx	General PA
fluticasone propionate 0.005% ointment	Р		1 package/Rx	<u>Form</u>



	DERMATOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.				
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
triamcinolone acetonide 0.025% cream, lotion and ointment	Р		1 package/Rx		
triamcinolone acetonide 0.05% ointment	Р		1 package/Rx		
triamcinolone acetonide 0.1% lotion and ointment	Р		1 package/Rx		
triamcinolone acetonide 0.5% cream and ointment	Р		1 package/Rx		
amcinonide 0.1% cream and lotion	NP		1 package/Rx		
betamethasone dipropionate 0.05% cream	NP		1 package/Rx		
betamethasone valerate 0.12% foam	NP		1 package/Rx		
desoximetasone 0.05% cream	NP		1 package/Rx		
fluocinonide 0.05% emulsified base cream	NP		1 package/Rx		
		Topical Steroids: Potent			
betamethasone dipropionate, augmented 0.05% cream	Р		1 package/Rx		
ApexiCon E® 0.05% cream	NP		1 package/Rx		
betamethasone dipropionate, augmented 0.05% lotion	NP		1 package/Rx		
betamethasone dipropionate 0.05% ointment	NP		1 package/Rx		
desoximetasone 0.05% gel and	NP		1 package/Rx	General PA	



DERMATOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
ointment				<u>Form</u>	
desoximetasone 0.25% cream, ointment, spray	NP		1 package/Rx		
diflorasone diacetate 0.05% cream and ointment	NP		1 package/Rx		
fluocinonide 0.05% cream, gel, and ointment	NP		1 package/Rx		
Halog® 0.1% ointment and cream	NP		1 package/Rx		
Halog [®] solution	NP		120 mL per 30 days		
		Topical Steroids: Super Potent			
clobetasol propionate 0.05% cream, gel, ointment, lotion, and solution	Р		1 package/Rx		
clobetasol propionate emollient base 0.05% cream	Р		1 package/Rx		
fluocinonide 0.1% cream	Р		1 package/Rx		
Bryhali [®] lotion	NP	 Diagnosis of an FDA-approved indication; AND Clinically valid reason why the preferred individual components cannot be taken concomitantly 	200 g/28 days		
betamethasone dipropionate, augmented 0.05% gel and ointment	NP		1 package/Rx	General PA Form	
clobetasol 0.025% cream	NP		1 package/Rx		
clobetasol propionate 0.05% foam, shampoo, and spray	NP		1 package/Rx		
clobetasol propionate emollient base 0.05% foam	NP		1 package/Rx		
halobetasol propionate 0.05%	NP		1 package/Rx		



		DERMATOLOGICS		
Medication	PDL	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Prior Authorization Criteria	Qty. Limits	PA Form
cream, foam, and ointment			-	
Lexette®	NP	See Bryhali [®] prior authorization criteria	100 g/Rx	1
Ultravate® 0.05% lotion	NP		1 package/Rx	
		Wound Care, Topical		•
Filsuvez®	NP	Initial Criteria: (6-month duration) One of the following: Diagnosis of Dystrophic Epidermolysis bullosa (EB); Diagnosis of Junctional Epidermolysis bullosa (EB); Wound is clean in appearance and does not appear to be infected Wound is 10 cm² to 50 cm²; AND Patient will continue standard treatments for EB such as appropriate wound management and avoiding skin trauma; AND Prescribed by or in consultation with a dermatologist or wound management Renewal Criteria: Patient has a clinical response to therapy (e.g., decreased wound size, decreased frequency of wound dressing changes, reduction in pain) Note: New wounds untreated with Filsuvez or recurrent reopened wounds are subject to initial criteria. Unhealed wounds > 6 months should rule out squamous cell and basal cell carcinoma.	15 tubes/per 30 days	General P Form
		DIABETIC SUPPLIES		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Blood Glucose Meters and Test Strips (OTC)		
		Abbott Products		
FreeStyle Meters: Lite, Freedom Lite, InsuLinx	Р		Meters: 1/730 days	<u>Diabetic</u>
Freestyle Test Strips: Lite, InsuLinx	Р		Test Strips: Age ≤ 5: 306/30 days	Supply PA Form
All other Abbott diabetic supplies	Р		Age > 6: 204/30 days	
		AgaMatrix Products		



	DIABETIC SUPPLIES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.				
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
Various	NP	See prior authorization criteria for Breeze-2 Meter (Bayer Products)	Meters: 1/365 days Test Strips: Age ≤ 5: 306/30 days Age > 6: 204/30 day	<u>Diabetic</u> <u>Supply PA</u> <u>Form</u>	
		Bayer Products			
Bayer Meters: Breeze-2 & Contour	NP	 Non-preferred meters will be approved for patients meeting ONE of the following criteria: Patient is using an insulin pump that does not adequately communicate with a preferred meter. Patient requires a special meter due to visual impairment 	Meters: 1/365 days;	<u>Diabetic</u> Supply PA	
Bayer Test Strips All other Bayer diabetic supplies	NP NP	Will be approved for individuals who meet prior authorization criteria and receive a prior authorization for a Bayer diabetes meter.	Test Strips: Age ≤ 5: 306/30 days Age > 6: 204/30 days	Form Form	
		Home Diagnostics Products			
Various	NP	See prior authorization criteria for Breeze-2 Meter (Bayer Products)	See Bayer Products	<u>Diabetic</u> <u>Supply PA</u> <u>Form</u>	
		Johnson and Johnson Products	1		
OneTouch Meters: UltraMini, Ping, Ultra-2, UltraLink, UltraSmart	NP	See prior authorization criteria for Breeze-2 Meter (Bayer Products)	Meters: 1/365 days;	<u>Diabetic</u>	
Johnson & Johnson Test Strips All other OneTouch diabetic supplies	NP NP	Will be approved for individuals who meet prior authorization criteria and receive a prior authorization for a OneTouch diabetes meter.	Test Strips: Age ≤ 5: 306/30 days Age > 6; 204/30 days	Supply PA Form	
,,	1	LifeScan Products	1		
Various	NP	See prior authorization criteria for Breeze-2 Meter (Bayer Products)	Meters: 1/365 days; Test Strips: Age ≤ 5: 306/36 days Age > 6: 204/30 days	<u>Diabetic</u> <u>Supply PA</u> <u>Form</u>	
		Roche Products			
Accu-Chek Meters: Aviva & Compact Plus	NP	See prior authorization criteria for Breeze-2 Meter (Bayer Products)	Meters: 1/365 days;	<u>Diabetic</u>	
Roche Test Strips All other Roche diabetic supplies	NP NP	Will be approved for individuals who meet prior authorization criteria and receive a prior authorization for an Accu-Chek diabetes meter.	Test Strips: Age ≤ 5: 306/36 days Age > 6: 204/30 days	Supply PA Form	



		DIABETIC SUPPLIES		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		All Manufacturers	1	ı
Ketone Testing Strips			50 /30 days	General PA Form
		Continuous Glucose Monitors and Supplies		
		Dexcom		
Dexcom G6; Dexcom G7	Р	 One of the following: Diagnosis of Gestational Diabetes Mellitus with suboptimal glycemic control that is likely to cause risk or harm to the mother/fetus; OR Patient has Diagnosis of Type 1 Diabetes Mellitus OR Diagnosis of Type 2 Diabetes Mellitus and meets ONE of the following:	G6 Sensor: 3/30 days Transmitter: 1/90 days Receiver: 1/year G7 Sensor: 3/month Receiver: 1/year	<u>Diabetic</u> Supply PA Form
		See Dexcom prior authorization criteria; AND	Sensor: 1/90 days	
Eversense Mis	NP	Prescriber must submit clinical documentation (e.g., medical records) why the patient cannot use CGM Dexcom	Transmitter: 1/90 days	Diabetic
Eversense E3	NP	 See Dexcom prior authorization criteria; AND Prescriber must submit clinical documentation (e.g., medical records) why the patient cannot use CGM Dexcom 	Sensor: 2/year Transmitter: 1/year	Supply PA Form
Eversense E365	NP	 See Dexcom prior authorization criteria; AND Prescriber must submit clinical documentation (e.g., medical records) why the patient cannot use CGM Dexcom 	Sensor: 1/year Transmitter: 1/year	



	DIABETIC SUPPLIES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
		Abbot				
Freestyle; Freestyle Libre 2 Freestyle Libre 3	NP	 See Dexcom prior authorization criteria; AND Prescriber must submit clinical documentation (e.g., medical records) why the patient cannot use CGM Dexcom 	Readers: 1/year Sensors:2/28 days	<u>Diabetic</u> <u>Supply PA</u> Form		
Freestyle Libre Plus (2 and 3)		 See Dexcom prior authorization criteria; AND Prescriber must submit clinical documentation (e.g., medical records) why the patient cannot use CGM Dexcom 	2 kits/30 days	<u> </u>		
		Medtronic				
Guardian 3; Guardian 4	NP	 One of the following: Patient is a currently using MiniMed insulin pump; OR See Dexcom prior authorization criteria; AND Prescriber must submit clinical documentation (e.g., medical records) why the patient cannot use CGM Dexcom 	Transmitter: 1/year Sensors: 5/30 days			
Guardian Connect	NP	 See Dexcom prior authorization criteria; AND Prescriber must submit clinical documentation (e.g., medical records) why the patient cannot use CGM Dexcom 	1 transmitter/year			
Guardian CGM supplies	NP	 See Dexcom prior authorization criteria; AND Prescriber must submit clinical documentation (e.g., medical records) why the patient cannot use CGM Dexcom 	Charger: 1/year Test plug: 1/year	<u>Diabetic</u> <u>Supply PA</u> <u>Form</u>		



DIABETIC SUPPLIES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.							
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
		Insulin Management Systems		<u> </u>			
Omnipod 5® Omnipod 5 G7® Omnipod Dash®	P	Initial Criteria: (6-month duration) One of the following: Diagnosis of Type 1 Diabetes Mellitus; OR Diagnosis of Type 2 Diabetes Mellitus has had ONE of the following in the last 9-months: Patient has an HgA1c of greater than 7% with 2 consecutive HbA1c; OR Patient has not met individual goal for A1c or time in range (if on a CGMS) with 2 consecutive HbA1c; AND Prescribed by, or in consultation with, an endocrinologist or diabetologist; AND Patient or caregiver has completed a physician-directed comprehensive diabetes management program which included a visit with a dietician; AND Patient has met one of the following insulin administration methods within the last 6-months: If patient has used insulin pump within the last 6-months, clinically valid reason why current insulin pump is no longer appropriate; OR Administration of at least three daily insulin injections with frequent self- adjustments of insulin dose and exhibits one or more of the following criteria while on a regimen of multiple daily injections of insulin: Glycosylated hemoglobin level (HbA1c) > 7% History of reoccurring hypoglycemia Wide fluctuations in blood glucose before mealtime Dawn phenomenon with fasting blood glucose frequently exceeding 200 mg/dL History of severe glycemic excursions; AND Documented monitored blood glucose self-testing ≥ 4 times a day or regular use of calibrated CGMS during 2 months prior to initiation of insulin pump; AND Provider must submit documentation (such as office chart notes, lab results or other clinical information) supporting the member's insulin administration methods and blood glucose monitoring methods. Renewal Criteria: Documentation of a positive clinical response (e.g. decrease HbA1C from baseline, decrease hypoglycemia episodes, decrease fasting and mealtime blood glucose levels)	Pods: 10/30 days; Device: 1/year	General PA Form			



		DIABETIC SUPPLIES		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Omnipod Go®	P	Initial Criteria (6-month duration): Patient is ≥ 18 years of age; AND Diagnosis of Type 2 Diabetes Mellitus has had ONE of the following in the last 9-months: Patient has an HgA1c of greater than 7% with 2 consecutive HbA1c; OR Patient has not met individual goal for A1c or time in range (if on a CGMS) with 2 consecutive HbA1c; AND Is not using more than 40 units of basal insulin per day; AND Prescriber by or in consultation with an endocrinologist or diabetologist; AND Prescriber must provide a clinically valid reason as to why the Omnipod GO insulin management system is needed for the patient versus standard insulin injections; AND Patient or caregiver has completed a physician-directed comprehensive diabetes management program Renewal Criteria: Patient is ≥ 18 years of age; AND Patient has Diagnosis of Type 2 diabetes; AND Is not using more than 40 units of basal insulin per day; AND Documentation of a positive clinical response (e.g. decrease HbA1C from baseline)	Pods: 10/30 days; Device: 1/year	General PA Form
Cequr Simplicity®	P	Initial Criteria: (6-month duration) One of the following: ○ Diagnosis of Type 1 Diabetes Mellitus; OR ○ Diagnosis of Type 2 Diabetes Mellitus has had ONE of the following in the last 9-months: Patient has an HgA1c of greater than 7% with 2 consecutive HbA1c; OR Patient has not met individual goal for A1c or time in range (if on a CGMS) with 2 consecutive HbA1c; AND Patient is ≥ 21 years old; AND Prescribed by, or in consultation with, an endocrinologist or diabetologist; AND Patient or caregiver has completed a physician-directed comprehensive diabetes management program which included a visit with a dietician; AND Patient has met ONE of the following insulin administration methods within the last 6-months: ○ If patient has used an insulin pump within the last 6-months, clinically valid reason why current insulin pump is no longer appropriate; OR ○ Administration of at least three daily insulin injections with frequent self- adjustments of insulin dose and exhibits one or more of the following criteria while on a regimen of multiple daily injections of insulin: Glycosylated hemoglobin level (HbA1c) > 7% History of reoccurring hypoglycemia Wide fluctuations in blood glucose before mealtime Dawn phenomenon with fasting blood glucose frequently exceeding 200 mg/dL History of severe glycemic excursions; AND Documented monitored blood glucose self-testing ≥ 4 times a day or regular use of calibrated CGMS during 2 months prior to initiation of insulin pump; AND Provider must submit documentation (such as office chart notes, lab results or other clinical information) supporting the member's insulin administration methods and blood glucose monitoring methods. Renewal Criteria: Documentation of a positive clinical response (e.g. decrease HbA1C from baseline, decrease hypoglycemia episodes, decrease fasting and mealtime blood glucose levels)	3-day patch: 10 /30 days 4-day patch: 8 /32 days	General PA Form



	DIABETIC SUPPLIES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
InPen®	NP	Prescriber must submit clinical documentation (e.g., medical records) why the patient cannot use the preferred products		General PA		
V-Go® products	NP	Prescriber must submit clinical documentation (e.g., medical records) why the patient cannot use the preferred products	30 patches/30 days	<u>Form</u>		
	Insulin Syringes and Pen Needles (OTC)					
BD/Embecta products	Р	Refer to OTC List for covered NDCs		General PA Form		



		ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Adrenocorticotropins		
Acthar® gel	NP	 Appropriate FDA-approved diagnosis (e.g., diuresis in nephrotic syndrome, treatment of SLE or polymyositis, or acute MS exacerbation) for use AND has a contraindication, or intolerance to oral and injectable glucocorticoids; OR Diagnosis of infantile spasms 	1/day	General PA
Cortrophin® gel	NP	See Acthar® gel prior authorization criteria; AND • Clinically valid reason why Acthar® gel cannot be used	1/day	<u>Form</u>
		Agents for Dyspareunia		
Intrarosa [®]	NP	 Female younger than 21 years of age; AND Cessation of menses due to menopause; AND Painful intercourse Note: This product is excluded from coverage in patients 21 years of age and older. Not a Covered Benefit. 		
Osphena [®]	NP	See Intrarosa® prior authorization criteria Note: This product is excluded from coverage in patients 21 years of age and older. Not a Covered Benefit.		
		Agents for Gout		
colchicine tablet	Р	 Diagnosis of Familial Mediterranean Fever; OR Diagnosis of acute pericarditis, AND must be taken concurrently with NSAID (unless contraindicated); OR For initiation of colchicine for acute gout attack; OR For continuation of colchicine prophylaxis for gout: Current history of urate lowering therapy with compliance in the past three months; AND One of the following: 		General PA Form
colchicine capsules	NP	See colchicine tablet prior authorization criteria; AND • Trial and failure of the preferred colchicine product		



ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Qty. Limits PA Form Prior Authorization Criteria** Initial Criteria (3 months): · Diagnosis or history of gout flares; AND · Patient is 18 years of age or older; AND • Patient has had a trial and failure of colchicine tablets; OR o Patient is unable to swallow or has difficulty swallowing colchicine tablets/capsules; AND • Females of reproductive potential and males with female partners of reproductive potential should be advised to use effective contraception; AND • Patient does not meet the following: o Presence of an active gout flare Gloperba® NP o Renal or hepatic impairment 300 ml/28 days o In combination with CYP3A4 and P-gp inhibitors; AND • Prescriber attests that the following will be monitored: o CBC, ALTs, ASTs, Scr o Serum uric acid levels o Neuromuscular toxicity (creatine phosphokinase (CPK), SGOT, SGPT, and LDH) Renewal Criteria (3 months): · Patient continues to meet the initial criteria; AND Patient has not experienced any treatment-restricting adverse effects (e.g., colchicine toxicity, neuromuscular toxicity, blood dyscrasias, liver and renal toxicity) See colchicine tablet prior authorization criteria; AND Mitigare® NΡ • Trial and failure of the preferred colchicine product • Trial and failure, contraindication, or intolerance to allopurinol; AND NP Uloric® • Clinically valid reason as to why the preferred febuxostat cannot be used



	ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.						
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
	,	Androgens					
AndroGel® pump	P	Initial Criteria: Patient age 21 years of age or less AND Diagnoses of Micropenis, Congenital or Acquired Anorchia, Kallmann Syndrome, Klinefelter Syndrome OR Hypogonadotrophic hypogonadism/central hypogonadism due to one of the following etiologies: ○ Congenital midline brain defects: Septo-optic Dysplasia, Holoprosencephaly, Hypopituitarism ○ CNS tumors and treatment including irradiation, surgery, and chemotherapy ○ Significantly delayed puberty ○ Approval requires: - Baseline Luteinizing Hormone - Baseline Luteinizing Hormone - Baseline stosterone level [faxed labs required] • Patient age 21 years of age or less: diagnosis not specified above: ○ Diagnosis of hypogonadism as confirmed by 2 baseline fasting testosterone levels drawn in the AM on separate dates demonstrating low testosterone [faxed labs required] and requires: - Baseline hematocrit ≤ 50% - Baseline Luteinizing Hormone • Patient age 22 years of age and older: ○ Diagnosis of hypogonadism as confirmed by 2 baseline fasting testosterone levels drawn in the AM on separate dates demonstrating low testosterone [faxed labs required] and requires: - Baseline hematocrit ≤ 50% - Baseline hematocrit ≤ 50% - Baseline Luteinizing Hormone - PSA level < 3 ng/mL • Requests for diagnosis of gender dysphoria will be reviewed on a case-by-case basis for determination Renewal Requests: • Documentation of low or normal fasting testosterone level from previous 12 months [faxed labs required] • Hematocrit ≤ 50% • PSA level < 3 ng/mL [not required for <21]	1 package/Rx	General PA Form			
testosterone gel (excluding 2%)	Р	See AndroGel® pump prior authorization criteria	1 package/Rx				
testosterone cypionate	Р	See AndroGel® pump prior authorization criteria	4 mL/30 days				



	ENDOCRINE/METABOLIC AGENTS							
Medication	PDL	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Prior Authorization Criteria	Qty. Limits	PA Form				
AndroGel® 1% and 1.62% packets	NP	Initial Criteria: Patient age 21 years of age or less AND Diagnoses of Micropenis, Congenital or Acquired Anorchia, Kallmann Syndrome, Klinefelter Syndrome OR Hypogonadotrophic hypogonadism/central hypogonadism due to one of the following etiologies: Congenital midline brain defects: Septo-optic Dysplasia, Holoprosencephaly, Hypopituitarism CNS tumors and treatment including irradiation, surgery, and chemotherapy Significantly delayed puberty Approval requires: Baseline Luteinizing Hormone Baseline testosterone level [faxed labs required] Intolerance or contraindication to an inactive ingredient in ALL preferred topical testosterone that is not in the requested product Patient age 21 years of age or less: diagnosis not specified above: Diagnosis of hypogonadism as confirmed by 2 baseline fasting testosterone levels drawn in the AM on separate dates demonstrating low testosterone [faxed labs required] and requires: Baseline hematocrit ≤ 50% Baseline Luteinizing Hormone Intolerance or contraindication to an inactive ingredient in ALL preferred topical testosterone that is not in the requested product Patient age 22 years of age and older: Diagnosis of hypogonadism as confirmed by 2 baseline fasting testosterone levels drawn in the AM on separate dates demonstrating low testosterone [faxed labs required] and requires: Baseline Luteinizing Hormone PSA level < 3 ng/mL Intolerance or contraindication to an inactive ingredient in ALL preferred topical testosterone that is not in the requested product Requests for diagnosis of gender dysphoria will be reviewed on a case-by-case basis for determination Renewal Requests: Documentation of low or normal fasting testosterone level from previous 12 months [faxed labs required] Hematocrit ≤ 50% PSA level < 3 ng/mL [not required for <21]	1 package/Rx	General PA Form				
Depo-Testosterone®	NP	See AndroGel® 1% and 1.62% packets prior authorization criteria	4 mL/30 days					
Jatenzo®	NP	See AndroGel® 1% and 1.62% packets prior authorization criteria	2/day					



ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Qty. Limits Prior Authorization Criteria PA Form Initial Criteria:** Patient age 21 years of age or less AND Diagnoses of Micropenis, Congenital or Acquired Anorchia, Kallmann Syndrome, Klinefelter Syndrome OR Hypogonadotrophic hypogonadism/central hypogonadism due to one of the following etiologies: o Congenital midline brain defects: Septo-optic Dysplasia, Holoprosencephaly, Hypopituitarism o CNS tumors and treatment including irradiation, surgery and chemotherapy Significantly delayed puberty Approval requires: - Baseline Luteinizing Hormone Baseline testosterone level [faxed labs required] Intolerance or contraindication to at least ONE preferred testosterone product • Patient age 21 years of age or less: diagnosis not specified above: o Diagnosis of hypogonadism as confirmed by 2 baseline fasting testosterone levels drawn in the AM on separate dates demonstrating low testosterone [faxed labs required] and requires: - Baseline hematocrit ≤ 50% Methitest® NP - Baseline Luteinizing Hormone Intolerance or contraindication to at least ONE preferred testosterone product • Patient age 22 years of age and older: o Diagnosis of hypogonadism as confirmed by 2 baseline fasting testosterone levels drawn in the AM on separate dates demonstrating low testosterone [faxed labs required] and requires: Baseline hematocrit ≤ 50% - Baseline Luteinizing Hormone PSA level < 3 ng/mL o Intolerance or contraindication to at least ONE preferred testosterone product Requests for diagnosis of gender dysphoria will be reviewed on a case-by-case basis for determination **Renewal Requests:** • Documentation of low or normal fasting testosterone level from previous 12 months [faxed labs required] Hematocrit ≤ 50% • PSA level < 3 ng/mL [not required for <21] methyltestosterone NP See Methitest[®] prior authorization criteria Natesto® nasal gel NP See AndroGel® 1% and 1.62% packets prior authorization criteria Testim® NP See AndroGel® 1% and 1.62% packets prior authorization criteria 1 package/Rx See AndroGel® 1% and 1.62% packets prior authorization criteria; OR testosterone NP Palliative treatment of androgen-responsive, advanced, inoperable, metastatic breast cancer in women who are 1-5 years 4 mL/30 days enanthate injection postmenopausal and in premenopausal women who have benefited from oophorectomy See AndroGel® 1% and 1.62% packets prior authorization criteria Testosterone 2% gel 1 package/Rx Tlando® NP | See AndroGel® 1% and 1.62% packets prior authorization criteria 2/day Undecatrex® See AndroGel® 1% and 1.62% packets prior authorization criteria 4/day



		ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Vogelxo®	NP	See AndroGel® 1% and 1.62% packets prior authorization criteria		
Xyosted®	NP	See testosterone enanthate injection prior authorization criteria	2 mL/30 days	
		Bone: Bisphosphonate		
alendronate	Р		10, 40 mg: 1/day , 70 mg: 4/28 days	
alendronate solution	Р		10 mL/day	
Atelvia®	Р		4/28 days	
ibandronate	Р		1/28 days	
Actonel®	NP	3	5, 30 mg: 1/day 35 mg: 4/28 days 50 mg: 1/28 days	General PA Form
Binosto®	NP		4/28 days	
Fosamax®	NP		see alendronate	
Fosamax Plus D®	NP		4/28 days	
risedronate	NP	3	5, 30 mg: 1/day 35 mg: 4/28 days 50 mg: 1/28 days	



		ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Bone: Calcitonin		
calcitonin nasal spray	Р	 Diagnosis of osteoporosis in postmenopausal women greater than five years post menopause, AND Trial and failure, contraindication, or intolerance to BOTH bisphosphonates AND raloxifene. 	3.7 mL/30 days	
calcitonin injection	NP	 Diagnosis of Paget's disease of the bone; AND Trial and failure, contraindication, or intolerance to bisphosphonates; OR Treatment of hypercalcemia; OR Diagnosis of osteoporosis in postmenopausal women greater than five years post-menopause; AND Trial and failure, contraindication, or intolerance to BOTH bisphosphonates AND raloxifene; AND Trial and failure, contraindication, or intolerance to the preferred agent 	1 mL/day	General PA Form
Miacalcin® injection	NP	See calcitonin injection prior authorization criteria	1 mL/day	
		Bone: Parathyroid Hormone		
Forteo®	Р	 Diagnosis of ONE of the following: Postmenopausal osteoporosis and patient is female; OR Osteoporosis and patient is male; OR Glucocorticoid-induced osteoporosis; AND	1 pen/28 days	General PA Form
teriparatide	NP	See Forteo prior authorization criteria; AND Clinically valid reason why preferred Forteo® cannot be used	1 pen/28 days	General PA



		ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Tymlos®	NP	 Initial Criteria: Diagnosis of ONE of the following:	1/28 days	General PA Form
		Bone: SERMs		
raloxifene	Р		1/day	General PA
Evista®	NP		1/day	<u>Form</u>
		Contraceptives, Non-Oral		
Depo IM Provera ®	Р		1 vial/ 90 days	
Depo SubQ Provera®	Р		1 vial/ 90 days	
medroxyprogesteron e acetate injection	Р		1 vial/ 90 days	
Nuvaring®	Р		1/28 days	General PA Form
Xulane®	Р		3/28 days	
Annovera®	NP	Patient has tried and failed or had a contraindication to two preferred non-oral contraceptives; AND Clinically valid reason as to why preferred Nuvaring cannot be used	1/year	
Eluryng®	NP		1/28 days	
Etonogestrel-ethinyl estradiol vaginal ring	NP		1/28 days	General PA
Haloette®	NP		1/28 days	<u>Form</u>



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Phexxi®	NP	Patient has tried and failed or had a contraindication to two preferred non-oral contraceptives; AND Provider attests the patient will be monitored for cystitis and pyelonephritis	12/month	
Twirla ®	NP	Trial and failure, or contraindication/intolerance of two preferred non-oral contraceptives AND Avoid concomitant use of hepatitis C drug combinations containing ombitasvir/paritaprevir/ritonavir, with or without dasabuvir	3/28 days	
Zafemy®	NP		3/28 days	
		Contraceptives, Oral		
Various	Р		1/day	
Emergency contraceptives	Р		1/21 days	General PA Form
Various	NP		1/day	
		Corticosteroids, Oral		
Alkindi Sprinkles®	NP	 Diagnosis of adrenocortical insufficiency; AND Patient is 18 years of age or younger; AND Patient does not have ANY of the following: Hypersensitivity to hydrocortisone Untreated fungal and bacterial infections; AND Clinically valid reason as to why the preferred prednisolone solution cannot be used 	0.5 mg: 3/day 1 mg: 3/day 2 mg: 3/day 5 mg: 4/day	General PA Form
Eohilia®	NP	 Criteria: (3-month duration) Patient is 11 years of age or older; AND Diagnosis of Eosinophilic esophagitis (EoE); AND Prescribed by, or in consultation with, a gastroenterologist, allergist, or immunologist; AND Prescriber attest patient meets both of the following: Esophageal biopsy consists of ≥ 15 intraepithelial eosinophils per high-power field (eos/hpf) following treatment course of a proton pump inhibitor; AND Patient has symptoms of esophageal dysfunction (e.g., feeding difficulties, vomiting, pain, dysphagia); AND Trial and failure, or contraindication, to swallowed inhaled corticosteroids such as budesonide or fluticasone 	180 packs / 365 days	General PA Form



	ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.						
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
Hemady®	NP	 Patient must be 18 years of age or older; AND Patient must have a diagnosis of Multiple Myeloma; AND Must be used in combination with other anti-myeloma agents; AND Patient must NOT have any of the following: Systemic fungal or bacterial infection Glaucoma Herpes Simplex Keratitis Ocular infection Tympanic membrane perforation Prior hypersensitivity with dexamethasone Strong CYP3A4 inhibitors or inducers Pregnant or breastfeeding; AND Female patients should use effective contraception during treatment and for at least 1 week after treatment; AND Trial and failure, contraindication, or intolerance to two preferred dexamethasone products; AND Clinically valid reason why the preferred agents cannot be used 	2/day	General PA Form			
prednisolone ODT	NP	 Unable to swallow, OR Unable to absorb medications through the GI tract 		General PA			
Rayos®	NP	 Trial and failure, contraindication, or intolerance to TWO preferred products (trial must include prednisone); AND Clinically valid reason why the preferred agents cannot be used 	1 mg: 3/day 2 mg: 2/day 5 mg: 12/day	Form			
		Diabetes: Alpha-Glucosidase Inhibitors					
acarbose	Р	Trial and failure, contraindication, or intolerance to metformin monotherapy		General PA			
miglitol	NP	 Trial and failure, contraindication, or intolerance to metformin monotherapy; AND Trial and failure, contraindication, or intolerance of TWO preferred agents 		<u>Form</u>			
		Diabetes: Amylin Analogs					
Symlin Pen®	NP	 Diagnosis of Type 1 or 2 diabetes; AND On insulin therapy; AND Failure to achieve adequate glycemic control (HbA1c ≤ 6.5%); AND Patients meeting any of the following will NOT be approved: Recurrent, severe hypoglycemia requiring assistance during the past 6-months Confirmed diagnosis of gastroparesis Requiring the use of drugs that stimulate gastrointestinal motility 		General PA Form			
Diabetes: Biguanides							
metformin	netformin P 500 mg: 4/day 850 & 1000 mg: 2/day						



	ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
metformin ER	Р		500 mg: 1/day 1000 mg: 2/day			
Glumetza®	NP		500 mg: 1/day 1000 mg: 2/day	General PA		
metformin ER osmotic	NP		500 mg: 3/day 1000 mg: 2/day	<u>Form</u>		
metformin solution	NP	See Riomet prior authorization criteria	20 mL/day			
	•	Diabetes: DPP-4 Inhibitors and Combinations				
Janumet®	Р		2/day			
Janumet XR®	Р		50/500 mg, 100/1000 mg: 1/day; 50/1000 mg: 2/day	DPP-4 PA Form		
Januvia®	Р		1/day			
Jentadueto®	Р		2/day			
Jentadueto® XR	Р		2.5/1000 mg: 2/day; 5/1000 mg: 1/day			
saxagliptin	Р		1/day	DPP-4		
Tradjenta®	Р		1/day	PA Form		
alogliptin	NP	 Diagnosis of type 2 diabetes; AND Patient's HbA1c level is greater than 6.5 (for initial approval); AND Trial and failure, contraindication, or intolerance to TWO preferred single entity DPP-4 inhibitors (Januvia, Tradjenta) 	1/day			
alogliptin/metformin	NP	 Diagnosis of type 2 diabetes; AND Patient's HbA1c level is greater than 6.5 (for initial approval); AND Trial and failure, contraindication, or intolerance to TWO preferred DPP-4/metformin combination products (Janumet, Janumet XR, Jentadueto, Jentadueto XR) 	2/day	DPP-4		
alogliptin/ pioglitazone	NP	See alogliptin/metformin prior authorization criteria	1/day	PA Form		
saxagliptin/ metformin	NP	See alogliptin/metformin prior authorization criteria	2/day			
Zituvimet®	NP	Clinically valid reason why Janumet or Janumet XR cannot be used	50/500 mg, 100/1000 mg: 1/day; 50/1000 mg: 2/day	DPP-4		
Zituvimet XR®	NP	Clinically valid reason why Janumet or Janumet XR cannot be used	2/day	PA Form		
Zituvio®	NP	Clinically valid reason why Januvia® cannot be used	1/day			



ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
		Diabetes: Rapid-Acting Insulins			
Apidra SoloStar®	Р	 Prior authorization not required for patients < 21 years of age; OR Patient is 21 years of age or older; AND Recipient or caregiver has problems with manual dexterity which may result in dosing errors (e.g., Parkinson's Disease, rheumatoid arthritis in the finger/hand joints, multiple sclerosis, lupus, osteoarthritis, stroke); OR Recipient or caregiver has poor eyesight such that dosing errors may occur 		General PA Form	
Humalog KwikPen®	Р	See Apidra® Solostar® prior authorization criteria			
Humalog Jr Kwik Pen®	Р	 Prior authorization not required for patients < 21 years of age; OR Patient is 21 years of age or older; AND Patient requires half unit (0.5) dosing or adjustments that cannot be achieved with Humalog® Kwik Pen® 		General PA Form	
insulin lispro KwikPen	Р	See Apidra® Solostar® prior authorization criteria		General PA	
insulin lispro Jr Kwikpen	Р	See Humalog® Jr KwikPen prior authorization criteria		<u>Form</u>	
Admelog SoloStar®	NP	 Patient < 21 years of age; AND Trial and failure or intolerance of TWO preferred rapid acting insulin agents; OR Patients ≥ 21 years old; AND Trial and failure or intolerance of 2 preferred rapid acting insulin agents; AND Recipient or caregiver has problems with manual dexterity which may result in dosing errors (e.g., Parkinson's Disease, rheumatoid arthritis in the finger/hand joints, multiple sclerosis, lupus, osteoarthritis, stroke); OR Recipient or caregiver has poor eyesight such that dosing errors may occur Recipient or caregiver has poor eyesight such that dosing errors may occur 		General PA Form	
Afrezza®	NP	 Patient is not a current smoker and does not have a history of smoking in the past 6-months; AND Prescriber attests that baseline spirometry has been performed prior to therapy and will be performed after 6-months of therapy, and every year thereafter; AND Patient does not have a history of chronic lung disease (e.g., asthma, COPD); AND Patient has ONE of the following diagnoses: Type 2 Diabetes Type 1 Diabetes while concurrently taking a long-acting insulin; AND Recipient or caregiver has problems with manual dexterity which may result in dosing errors (i.e., Parkinson's Disease, rheumatoid arthritis in the finger/hand joints, multiple sclerosis, lupus, osteoarthritis, stroke); OR Recipient or caregiver has poor eyesight such that dosing errors may occur 	Cartridges: 4-unit: 3/day 8-unit: 6/day 12-unit:6/day Combo package: 1 box/month	General PA Form	
Fiasp FlexTouch®	NP	See Admelog [®] SoloStar [®] prior authorization criteria			
Fiasp PenFill®	NP	See Admelog [®] SoloStar [®] prior authorization criteria		General PA	
Humalog U-200 KwikPen®	NP	See Admelog® SoloStar® prior authorization criteria; AND • Provider must provide peer-reviewed medical literature documenting why the requested drug for the requested indication is the only appropriate choice versus the preferred agents		<u>Form</u>	
Humalog TempoPen®		See Admelog [®] SoloStar [®] prior authorization criteria			



		ENDOCRINE/METABOLIC AGENTS		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Lyumjev® vial	NP	 Trial and failure or intolerance of 2 preferred, rapid-acting insulin agents; AND Provider must provide peer-reviewed medical literature documenting why the requested drug for the requested indication is the only appropriate choice versus the preferred agents 		
Lyumjev KwikPen®	NP	See Admelog® SoloStar® prior authorization criteria; AND • Provider must provide peer-reviewed medical literature documenting why the requested drug for the requested indication is the only appropriate choice versus the preferred agents		General PA Form
Lyumjev TempoPen®	NP	See Admelog® SoloStar® prior authorization criteria		
Novolog FlexPen®	NP	See Admelog® SoloStar® prior authorization criteria		
		Diabetes: Intermediate-Acting Insulins		
Humulin N KwikPen®	Р	Prescriber must provide valid clinical rationale as to why patient is unable to utilize preferred Novolin® N FlexPen®		General PA Form
	,	Diabetes: Mixed Insulins		•
Humalog Mix 50/50 KwikPen®	Р	 Prior authorization not required for patients < 21 years of age; OR Patient is 21 years of age or older; AND Recipient or caregiver has problems with manual dexterity which may result in dosing errors (e.g., Parkinson's Disease, rheumatoid arthritis in the finger/hand joints, multiple sclerosis, lupus, osteoarthritis, stroke); OR Recipient or caregiver has poor eyesight such that dosing errors may occur 		General PA Form
Humalog Mix 75/25 KwikPen®	Р	See Humalog® Mix 50/50® KwikPen prior authorization criteria		
Humulin 70/30 KwikPen®	Р	See Humalog® Mix 50/50® KwikPen prior authorization criteria		General PA Form
insulin aspart mix 70/30 FlexPen	Р	See Humalog® Mix 50/50® KwikPen prior authorization criteria		
insulin lispro mix 75/25 KwikPen®	NP	 Patient < 21 years of age; AND Trial and failure or intolerance of TWO preferred rapid acting insulin agents; OR Patients ≥ 21 years old; AND Trial and failure or intolerance of 2 preferred rapid acting insulin agents; AND Recipient or caregiver has problems with manual dexterity which may result in dosing errors (e.g., Parkinson's Disease, rheumatoid arthritis in the finger/hand joints, multiple sclerosis, lupus, osteoarthritis, stroke); OR Recipient or caregiver has poor eyesight such that dosing errors may occur Recipient or caregiver has poor eyesight such that dosing errors may occur 		General PA Form
Novolog Mix 70/30 FlexPen®	NP	See insulin lispro mix 75/25 KwikPen® prior authorization criteria		



	ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.						
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
	•	Diabetes: Long-Acting Insulins					
Basaglar KwikPen®	NP	 Patients < 21 years of age approval requires a contraindication to a preferred insulin glargine pen that is not observed with the requested agent; OR For patients ≥ 21 years old approval requires a contraindication to a preferred insulin glargine pen that is not observed with the requested agent; AND Recipient or caregiver has problems with manual dexterity which may result in dosing errors (e.g., Parkinson's Disease, rheumatoid arthritis in the finger/hand joints, multiple sclerosis, lupus, osteoarthritis, stroke); OR Recipient or caregiver has poor eyesight such that dosing errors may occur 		General PA Form			
Basaglar Tempo Pen®	NP	See prior authorization criteria for Basaglar KwikPen®					
insulin degludec FlexTouch	NP	 For patients < 21 years of age, trial and failure, contraindication, or intolerance of 2 preferred agents; OR For patients ≥ 21 years of age, trial and failure, contraindication, or intolerance of 2 agents; AND Recipient or caregiver has problems with manual dexterity which may result in dosing errors (e.g., Parkinson's disease, rheumatoid arthritis in the finger/hand joints, multiple sclerosis, lupus, osteoarthritis, stroke, etc.); OR Recipient or caregiver has poor eyesight such that dosing errors may occur 		General PA			
Rezvoglar®	NP	See prior authorization criteria for Basaglar KwikPen®		<u>Form</u>			
Semglee®	NP	See prior authorization criteria for Basaglar KwikPen®					
Tresiba FlexTouch®	NP	See prior authorization criteria for insulin degludec FlexTouch					
		Diabetes: Meglitinides					
nateglinide	Р	Trial and failure, contraindication, or intolerance of metformin monotherapy	3/day	General PA			
repaglinide	Р	Trial and failure, contraindication, or intolerance of metformin monotherapy	0.5mg, 1mg: 4/day 2 mg: 8 day	<u>Form</u>			
		Diabetes: SGLT2 Inhibitors and Combinations					
Farxiga®	Р		1/day				
Glyxambi [®]	Р		1/day				
Invokana®	Р		1/day				
Invokamet®	Р		2/day	General PA Form			
Jardiance®	Р		1/day				
Synjardy®	Р		2/day				
Xigduo® XR	Р		1/day				



	ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.						
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
dapagliflozin	NP	Clinically valid reason why the preferred Farxiga® cannot be used	1/day				
dapagliflozin/ metformin ER	NP	Clinically valid reason why the preferred Xigduo XR ® cannot be used	1/day				
Inpefa®	NP	 Requested medication is being used to reduce the risk of cardiovascular death, hospitalization for heart failure, and urgent heat failure visit in adults with one of the following: Heart Failure Type 2 diabetes mellitus, chronic kidney disease, and other cardiovascular risk factors; AND Trial and failure, or intolerance, to TWO preferred agents 	1/day				
Invokamet XR®	NP	 Diagnosis of Type 2 Diabetes; AND Trial and failure or intolerance to TWO preferred single-entity SGLT2 agents (Farxiga, Invokana, Jardiance); AND Clinically valid reason as to why patient cannot use Invokamet® 	2/day				
Qtern®	NP	Trial and failure or intolerance to separate components (Farxiga and Onglyza)	1/day				
Steglatro®	NP	 Diagnosis of Type 2 Diabetes; AND Trial and failure or intolerance to TWO preferred single-entity SGLT2 agents (Farxiga, Invokana, Jardiance) 	2/day (5 mg); 1/day (15 mg)				
Segluromet®	NP	 Diagnosis of Type 2 Diabetes; AND Trial and failure or intolerance to TWO preferred single-entity SGLT2 agents (Farxiga, Invokana, Jardiance); AND Clinically valid reason as to why the patient cannot use a preferred single-entity SGLT2 agent and metformin as separate agents; AND Patient does not have metabolic acidosis 	2/day	General PA			
Steglujan®	NP	 Diagnosis of Type 2 Diabetes; AND Trial and failure or intolerance to TWO preferred single-entity SGLT2 agents (Farxiga, Invokana, Jardiance); AND Patient does not have metabolic acidosis 	1/day	<u>Form</u>			
Synjardy XR®	NP	 Diagnosis of Type 2 Diabetes; AND Trial and failure or intolerance to TWO preferred single-entity SGLT2 agents (Farxiga, Invokana, Jardiance); AND Clinically valid reason as to why patient cannot use Synjardy 	1/day (25/1000 mg); 2/day (all other strengths)				
Trijardy XR®	NP	 Diagnosis of Type 2 Diabetes; AND Trial and failure or intolerance to TWO preferred single-entity SGLT2 agents (Farxiga, Invokana, Jardiance); AND Clinically valid reason as to why patient cannot use the patient cannot use Glyxambi and metformin ER as separate agents 	10/5/1000 mg, 2.5/5/1000 mg: 1/day; 5/2.5/1000 mg, 12.5/2.5/1000 mg: 2/day				
		Diabetes: Sulfonylureas and Combinations					
glimepiride	Р		2/day	General PA			
Glucotrol XL®	NP	 Trial and failure, contraindication, or intolerance to, metformin monotherapy; AND Trial and failure, contraindication, or intolerance of TWO preferred agents 		Form			



ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Medication PDL **Prior Authorization Criteria Qty. Limits PA Form Diabetes: TZDs and Combinations** pioglitazone • Trial and failure, contraindication, or intolerance to metformin or a metformin containing product 1/day pioglitazone/ Ρ • Trial and failure, contraindication, or intolerance to metformin or a metformin containing product 2/day metformin • Trial and failure, contraindication, or intolerance to metformin or a metformin containing product; AND Actos® NP 1/day · Patient must have an allergy or intolerance to an inactive ingredient in the generic equivalent TZD and Combos ACTOplus Met® NΡ See Actos® prior authorization criteria 2/day PA Form • Trial and failure, contraindication, or intolerance to metformin or a metformin containing product; AND **Duetact**® NP • Trial and failure, contraindication, or intolerance to pioglitazone; AND 1/day • Clinically valid reason why the patient cannot use pioglitazone and glimepiride as separate agents pioglitazone/ NP | See Duetact® prior authorization criteria 1/day glimepiride



		ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
	· ·	Diabetes: Glucagon Agents		•
Baqsimi®	Р		2/Rx	
Gvoke	Р		2/Rx	General PA Form
Zegalogue®	NP		2/Rx	101111
	•	GLP-1 Agonists		
Ozempic® Victoza®	P	Initial Criteria: Diagnosis of type 2 diabetes; AND Submission of lab test for one of the following: ○ HbA1C level* ○ Oral glucose tolerance test ○ Random plasma glucose ≥ 200 mg/dL with classic symptoms of hyperglycemia or hyperglycemic crisis; AND One of the following: ○ Patient has or is at high-risk of atherosclerotic cardiovascular disease (ASCVD), chronic kidney disease (CKD), or heart failure (HF); OR ○ Trial and failure, contraindication, or intolerance TWO of the following; ○ Metformin or metformin containing product ○ SGLT2 or combination product ○ TZD ○ Sulfonylurea ○ Insulin; AND GLP-1 Receptor Agonists will NOT be covered for the following: ○ Diagnosis of Type I diabetes; OR ○ Treatment of diabetic ketoacidosis; OR ○ Use for weight loss; OR ○ Personal or immediate family history of medullary thyroid carcinoma or multiple endocrine neoplasia type 2 (MEN2) Renewal Criteria: Submission of recent medical records (e.g., chart notes and/or labs) documenting one of the following: ○ Reduction of HbA1c from baseline ○ Achievement or maintenance of therapeutic HbA1c goal ○ Improvement in fasting blood glucose levels ○ Patient is at increased risk of ASCVD, CKD, or HF Note*: HbA1c level can be from early stages in patient treatment. If original HbA1c is unknown, or current HbA1c is controlled due to another current diabetic regimen, please include current regimen and current HbA1c.	5 mcg: 1.2 mL/ 30 days 10 mcg: 2.4 mL/30 days	GLP-1 Agonist PA Form
Rybelsus®	NP	See Ozempic® prior authorization criteria; AND Trial and failure, contraindication, or intolerance to 2 preferred drugs (e.g., Ozempic, Victoza)	1/day	GLP-1



ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. PDL **PA Form** Medication **Prior Authorization Criteria Qty. Limits** See Ozempic® prior authorization criteria; AND <u>Form</u> Soliqua® NP • Trial and failure, contraindication, or intolerance to 2 preferred drugs (e.g., Ozempic, Victoza); AND 5 pens/30 days · Patient is currently taking, but inadequately controlled on, a long-acting insulin documented per TennCare paid claims See Ozempic® prior authorization criteria; AND NΡ Trulicity® 2 mL/28 days • Trial and failure, contraindication, or intolerance to 2 preferred drugs (e.g., Ozempic, Victoza) See Ozempic® prior authorization criteria; AND Mounjaro® NΡ 2 mL/28 days • Trial and failure, contraindication, or intolerance to 2 preferred drugs (e.g., Ozempic, Victoza)



ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.							
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
Wegovy®	NP	 Initial Criteria Treatment is being requested to reduce the risk of major adverse cardiovascular events; AND Patient is 21 years of age or older; AND Submitted medical documentation (e.g. chart notes) of initial body mass index (BMI) of ≥ 27 kg/m2; AND Submitted medical documentation (e.g. chart notes) of ONE of the following: Prior myocardial infarction Prior stroke (ischemic and hemorrhagic stroke) Symptomatic peripheral arterial disease as evidenced by intermittent claudication with ankle–brachial index < 0.85, peripheral arterial revascularization procedure, or amputation due to atherosclerotic disease; AND Submitted documentation HbA1C ≤ 6.5%; AND Patient is on optimized guideline-directed therapy including beta-blockers, RAS inhibitors, and lipid lowering agents to ensure reduced cardiovascular risk unless contraindicated (supported by medical documentation or claims history); AND Prescriber attests patient is participating in a supervised comprehensive weight management program that encourages behavioral modification, reduced calorice diet, and increased physical activity; AND Patient does not have any of the following: Diagnosis of type 1 or type 2 diabetes New York Heart Association class is Iv heart failure; Personal or family history of medullary thyroid carcinoma (MTC) OR MEN 2 History or presence of chronic pancreatitis End-stage renal disease or currently receiving dialysis; AND For female patients of reproductive potential, the following has been addressed: Patient is not pregnant or breastfeeding Patient is participating in a supervised comprehensive weight management program that encourage	4 injectors/28 days	GLP-1 Agonist PA Form			



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
Xultophy®	NP	See Ozempic® prior authorization criteria; AND • Trial and failure, contraindication, or intolerance to 2 preferred drugs (e.g., Ozempic, Victoza); AND • Patient is currently taking, but inadequately controlled on, a long-acting insulin documented per TennCare paid claims	5 pens/30 days				
	•	GnRH Agonist/Antagonist & LNRH Analogs					
Myfembree®	Р	Initial Criteria: Patient age is ≥ 18 years; AND Diagnosis of one of the following: Heavy menstrual bleeding associated with uterine leiomyomas/fibroids Moderate to severe pain associated with endometriosis; AND Patient must be premenopausal; AND Patient has tried and failed 2 medications in the following drug classes: Hormonal contraceptives (including oral or transdermal formulations, vaginal ring, or intrauterine device) NSAIDs Hemostatics (e.g., tranexamic acid) Oral progesterone; AND Prescribed by, or in consultation with, an obstetrics/gynecology or reproductive specialist; AND Patient will use effective non-hormonal contraception during treatment and 1 week after stopping therapy; AND Total treatment duration should not exceed 24 months due to risk of continued bone loss Renewal Criteria (only for 150 mg strength): Patient has positive response to therapy (e.g., reduction in pain and discomfort from baseline, sustained reduction in menstrual blood loss per cycle); AND Patient will use effective non-hormonal contraception during treatment and 1 week after stopping therapy; AND Total treatment duration should not exceed 24 months	1/day	General PA Form			
Oriahnn®	Р	See Myfembree® prior authorization criteria	1 box/28 days	1			



ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.						
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
Orilissa®	Р	Initial Criteria: Patient age is ≥ 18 years; AND Patient has confirmed diagnosis of endometriosis; AND Patient has tried and failed 2 medications in the following drug classes: Hormonal contraceptives (including oral or transdermal formulations, vaginal ring, or intrauterine device) NSAIDS Hemostatics (e.g., tranexamic acid) Oral progesterone; AND Prescribed by, or in consultation with, an obstetrics/gynecology or reproductive specialist; AND Pregnancy is excluded prior to initiating treatment; AND Total treatment duration should not exceed 24 months due to risk of continued bone loss Renewal Criteria (only for 150 mg strength): Patient continues to meet the initial criteria; AND Patient is considered to have clinically meaningful response to treatment	1/day: 150 mg; 2/day: 200 mg	General PA Form		
		GnRH Agonists / LHRH Analogs				
leuprolide	Р	Diagnosis of prostate cancer in male patient; OR Diagnosis of central precocious puberty in children (onset of secondary sexual development before 8 [girls] or 9 years of age [boys])				
Fensolvi®	NP	See leuprolide prior authorization criteria				
Lupron Ped-Depot®	NP	Diagnosis of central precocious puberty in children (onset of secondary sexual development before 8 years of age [girls] or 9 years of age [boys])				



ENDOCRINE/METABOLIC AGENTS						
Medication	PDL	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Prior Authorization Criteria	Qty. Limits	PA Form		
	Growth Hormone Agents PA					
Genotropin®	P	NOTE: Growth hormone therapy will NOT be approved for idiopathic short stature (ISS) Criteria: Agent is prescribed by, or in consultation with, an endocrinologist; AND Daily dose within approved dosage range for somatotropin for requested indication per clinical compendium; AND Daily dose based on weight of the enrollee, supported by submitted growth charts; AND Approval will be based on dosage form resulting in least wastage of product For patients < 21 years old, will be approved if ANY of the following criteria are met: Patient has failed two GH stimulation tests (defined as peak GH level < 10 ng/mL) OR has failed one GH stimulation test (adfined as peak GH level < 10 ng/mL) OR has failed one GH stimulation test (adfined as peak GH level < 10 ng/mL) OR has failed one GH stimulation test and has a documented low IGF-1 level based on age normal values Ocontinuation of therapy will be approved only if height velocity is within normal range for patient's age or bone age Therapy will not be approved once epiphyseal closure occurs Patient has evidence of hypothalamic-pituitary disease or structural lesions/trauma to the pituitary, including pituitary tumor, pituitary surgical damage, head trauma, or cranial irradiation and meets any of the following: Failed a GH stimulation test (peak GH level < 10ng/mL) Documented low IGF-1 level (below normal for patient's age) Has deficiencies in 3 or more pituitary axes Patient is a newborn infant and has evidence of hypoglycemia AND either a low GH level (<20 ng/mL) or a low for age IGF-1/GEBP-3 level Diagnosis of short stature associated with Short stature homeobox (SHOX) gene deficiency Diagnosis of Prader-Willi Syndrome Patient has chronic renal insufficiency (CrCl < 30 mL/min/1.73 m²) Diagnosis of Prader-Willi Syndrome Patient has chronic renal insufficiency (CrCl < 30 mL/min/1.73 m²) Diagnosis of Small for Gestational Age (SGA) or Intrauterine Growth Retardation (IGR) and patient is > 2 years old with a height at least 2 standard deviations below the		Growth Hormone PA Form		



ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.						
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
Egrifta SV®	NP	 Recipient must be at least 18 years of age, but less than 21 years old; AND Diagnosis of Acquired Immunodeficiency Syndrome (AIDs) or Human Immunodeficiency Virus (HIV); AND Prescribed by, or in consultation with, an endocrinologist or provider with expertise in HIV; AND Waist circumference greater than or equal to 95 cm for males, or greater than or equal to 94 cm for females; AND Waist to hip ratio greater than or equal to 0.94 for males, or greater than or equal to 0.88 for females Note: For recipients > 21 years of age, these agents are a non-covered benefit 		Growth Hormone PA Form		
Humatrope®	NP	See Genotropin® prior authorization criteria		Growth		
Norditropin®	NP	See Genotropin® prior authorization criteria		Hormone		
Nutropin AQ®	NP	See Genotropin® prior authorization criteria		PA Form		
Ngenla®	NP	Initial Criteria: Patient is at least 3 years of age and less than 18 years of age; AND Patient weighs at least 11.5kg; AND Diagnosis of growth failure due to inadequate secretion of endogenous growth hormone (GH); AND Agent is prescribed by, or in consultation with, an endocrinologist; AND Documentation that diagnosis of growth hormone deficiency has been confirmed by two evidence-based diagnostics (e.g., imaging, measurement of insulin-like growth factor 1 (IGF-1) levels, growth hormone stimulation test); AND Prescriber attests that a baseline fundoscopic eye examination to exclude preexisting papilledema; AND Patient provides a clinically valid reason why preferred Genotropin injection cannot be used Renewal Criteria: Patient continues to meet initial criteria; AND Patient has open epiphyses; AND Prescriber attests that patient has an annualized height velocity of > 2.5 cm/year		Growth Hormone PA Form		
Omnitrope®	NP	See Genotropin® prior authorization criteria				



ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.						
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
Serostim®	NP	Initial Criteria: Diagnosis of HIV-associated wasting syndrome or cachexia; AND One of the following: Unintentional weight loss of >10% over the last 12 months Unintentional weight loss of >7.5% over the last 6-months Body mass index (BMI) < 20 kg/m2; AND Body cell mass (BCM) below 40% total body weight in males or 35% total body weight in females; AND Nutritional evaluation since onset of wasting first occurred; AND Patient has not had weight loss due to other underlying treatable conditions (e.g., depression, mycobacterium avium complex, chronic infectious diarrhea, malignancy); AND Anti-retroviral therapy has been optimized to decrease the viral load and will be continued throughout the course of treatment; AND Trial and failure of megestrol Renewal Criteria: Evidence of positive response to therapy (i.e., > 2% increase in body weight and/or BCM); AND A target goal has not been achieved (i.e., weight, BCM, BMI)		Growth Hormone PA Form		
Skytrofa®	NP	Initial Criteria: Patient is at least 1 year of age and less than 18 years of age; AND Patient weighs at least 11.5kg; AND Diagnosis of growth failure due to inadequate secretion of endogenous growth hormone (GH); AND Agent is prescribed by, or in consultation with, an endocrinologist; AND Documentation that diagnosis of growth hormone deficiency has been confirmed by two evidence-based diagnostics (e.g., imaging, measurement of insulin-like growth factor 1 (IGF-1) levels, growth hormone stimulation test); AND Prescriber attests that a baseline fundoscopic eye examination to exclude preexisting papilledema; AND Patient provides a clinically valid reason why preferred Genotropin injection cannot be used Renewal Criteria: Patient continues to meet initial criteria; AND Patient has open epiphyses; AND Prescriber attests that patient has an annualized height velocity of > 2.5 cm/year		Growth Hormone PA Form		



		ENDOCRINE/METABOLIC AGENTS		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Sogroya®	NP	 Netial Criteria: Agent is prescribed by, or in consultation with, an endocrinologist; AND Daily dose based on weight of the enrollee, supported by submitted growth charts; AND Clinically valid reason as to why the patient cannot take the preferred product Genotropin; AND For patients < 21 years old, will be approved if ANY of the following criteria are met: Patient has evidence of hypothalamic-pituitary disease or structural lesions/trauma to the pituitary, including pituitary tumor, pituitary surgical damage, trauma, or cranial irradiation and meets any of the following:		Growth Hormone PA Form



ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.							
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
Voxzogo®	NP	 Initial Criteria: Diagnosis of achondroplasia; AND Prescribed by, or in consultation with, an endocrinologist; AND Patient has open epiphyses; AND Patient will not have limb-lengthening surgery during treatment with Voxzogo®; AND Provider attests that patient/caregiver has been properly trained on preparation and administration of Voxzogo Renewal Criteria: Patient continues to meet initial criteria; AND Provider attests that patient has an annualized growth velocity ≥ 1.5 cm/year 		General PA Form			
Zomacton®	NP	See Genotropin® prior authorization criteria					
	Hematopoietic Agents						
Retacrit®	Р	See Epogen® prior authorization criteria					
Aranesp®	NP	See Epogen® prior authorization criteria					



ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
Epogen®	NP	 Lab values obtained within 30 days of the date of administration; AND Adequate iron stores demonstrated by serum ferritin ≥ 100 ng/mL (mcg/L) and transferrin saturation (TSAT) ≥ 20%; AND Hemoglobin (Hb) < 10 g/dL and/or hematocrit (Hct) < 30% (unless otherwise specified); AND One of the following: One of the following: Patient is at least 5 years of age and receiving concurrent myelosuppressive chemotherapy; AND — Patient is at least 5 years of age and receiving concurrent myelosuppressive chemotherapy; AND — Patient's chemotherapy is not intended to cure their disease (i.e., paliative treatment) O Anemia secondary to idovudine treated, HIV-infected patient; AND — Zidovudine dose is ≤ 4,200 mg/week; AND Endogenous serum erythropoietin (EPO) levels ≤ 500 mUnits/mL; OR O Anemia secondary to hepatitis C virus (HCV) treatment in patients receiving ribavirin and interferon-alfa therapy; OR O Anemia secondary to myelodysplastic syndrome (MDS); AND — Treatment of lower risk disease associated with symptomatic anemia; AND — Endogenous serum erythropoietin (EPO) level ≤ 500 mUnits/mL; OR O Anemia secondary to myeloproliferative neoplasms (MPN) – Myelofibrosis; AND — Endogenous serum EPO ≤ 500 mUnits/mL; OR O Anemia secondary to multiple myeloma; OR O Anemia secondary to thematoid arthritis; OR O Anemia secondary to chronic kidney disease end (KD) and hemoglobin (Hb) is ≤ 12.9 g/dL; OR Patient is NOT willing to donate autologous blood		General PA Form	



ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
Mircera®	NP	 Lab values obtained within 30 days of the date of administration; AND Adequate iron stores demonstrated by serum ferritin ≥ 100 ng/mL (mcg/L) and transferrin saturation (TSAT) ≥ 20%; AND One of the following: Diagnosis of anemia secondary to chronic kidney disease (CKD) in adult patients and BOTH of the following:			
Procrit®	NP	See Epogen® prior authorization criteria			



		ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise	e indicated.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Vafseo®	NP	Initial Criteria: (6-month duration) Diagnosis of anemia due to CKD; AND Patient has been receiving dialysis for ≥ 4 months; AND Recent documentation (within 30 days or request) of ALL the following: Hemoglobin level <10 g/dL Serum ferritin ≥ 100 ng/mL (mcg/L) Trial and failure, contraindication, or intolerance to erythropoiesis-stimulating agents (ESAs); AND Prescriber attests to ALL of the following: Will not use in combination with ESAs Will not use in combination with strong CYP2C8 inhibitor such as gemfibrozil Patient does not have uncontrolled hypertension Renewal Criteria: Patient is receiving dialysis for anemia due to CKD; AND Submitted documentation demonstrating an increase hemoglobin from baseline; AND Recent documentation (within 30 days or request) of ALL the following: Serum ferritin ≥ 100 ng/mL (mcg/L) Transferrin saturation (TSAT) ≥ 20%; AND Prescriber attests to ALL of the following: Will not use in combination with ESAs Will not use in combination with strong CYP2C8 inhibitor such as gemfibrozil Patient does not have uncontrolled hypertension	100mg, 450mg: 1/day 300mg: 2/day	
		Estrogen/Progestin, Oral		
Premphase®	Р		1/day	General PA
Prempro®	Р		1/day	<u>Form</u>
		Estrogen / Progestin, Transdermal		
CombiPatch®	Р		8/28 days	General PA
Climara Pro®	NP		4/28 days	<u>Form</u>



		ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Estrogens, Transdermal		
estradiol biweekly patch	Р		8/28 days	
estradiol weekly patch	Р		4/28 days	
Dotti®	Р		4/28 days	
Lyllana®	Р		4/28 days	
Alora®	NP		8/28 days	
Climara®	NP		4/28 days	General PA Form
Divigel®	NP		1/day	101111
Elestrin®	NP		1/28 days	
estradiol gel	NP		1/day	
Menostar®	NP		4/28 days	
Minivelle®	NP		8/28 days	
Vivelle-Dot®	NP		8/28 days	
	•	Estrogens, Vaginal		
Premarin® cream	Р		2 grams/day	
Estrace®	NP		42.5 g/Rx	General PA Form
estradiol cream	NP		42.5 g/Rx	<u>101111</u>
		Estrogen/SERM Combinations		
Duavee®	NP	 Patient has an intact uterus with a diagnosis of moderate to severe vasomotor symptoms associated with menopause; OR Patient has an intact uterus with a diagnosis of post-menopausal osteoporosis 	1/day	General PA Form
		Progestins, Oral		
megestrol suspension 40 mg/mL	Р		20 mL/day	
norethindrone acetate	Р	Diagnosis of endometriosis		General PA
Aygestin®	NP	Diagnosis of endometriosis		<u>Form</u>
megestrol suspension 625 mg/5 mL	NP	Inability to swallow the 10 mL (400 mg) or 20 mL (800 mg) dose of the regular-strength suspension	5 mL/day	



ENDOCRINE/METABOLIC AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
	•	Hyperparathyroid Agents	•		
cinacalcet	Р	 Secondary Hyperparathyroidism due to Chronic Kidney Disease (CKD), AND patient must be on dialysis; OR Parathyroid Carcinoma resulting in hypercalcemia; OR Severe Hypercalcemia in patients with primary HPT who are unable to undergo parathyroidectomy 			
doxercalciferol capsules	NP	 Recipients experiencing (or with a history of) hypercalcemia and/or hyperphosphatemia with calcitriol use; AND Trial and failure, contraindication, or intolerance to cinacalcet 	0.5, 2.5 mcg: 1/day; 1 mcg: 3/day		
paricalcitol capsules	NP	See doxercalciferol capsules prior authorization criteria	1/day	General PA	
Rayaldee®	NP	 Secondary Hyperparathyroidism due to Stage 3 or Stage 4 Chronic Kidney Disease (CKD); AND Serum total 25-hydroxyvitamin D levels less than 30 ng/mL; AND Trial and failure, contraindication, or intolerance of cinacalcet 	2/day	<u>Form</u>	
Sensipar®	NP	See cinacalcet prior authorization criteria; AND • Clinically valid reason why the preferred cinacalcet agent cannot be used			
Zemplar® capsules	NP	See doxercalciferol capsules prior authorization criteria	1/day		
	•	NK3 Antagonists			
Veozah®	NP	 Diagnosis of moderate to severe vasomotor symptoms due to menopause; AND Trial and failure, contraindication, or intolerance to TWO of the following: Gabapentin Menopausal hormone therapy (e.g., estrogen monotherapy or estrogen + progesterone) Oxybutynin SSRI (e.g., paroxetine, escitalopram, citalopram) SNRI (e.g., venlafaxine and desvenlafaxine) 	1/day	General PA Form	



	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.						
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
		Vasopressor Antagonists					
lynarque®	NP	Initial Criteria (6-month duration): Patient has a diagnosis of autosomal dominant polycystic kidney disease (ADPKD); AND Patient is 18 years of age or older; AND Prescribed by, or in consultation with, a nephrologist; AND Prescriber and patient are enrolled in the Jynarque REMS program; AND Patient does not have a known hypersensitivity to tolvaptan; AND Patient does not have any of the following: History of symptoms of significant liver impairment or injury (not including uncomplicated polycystic liver disease) Uncorrected abnormal blood sodium concentration Inability to sense or respond to thirst Hypovolemia Uncorrected urinary outflow obstruction Anuria; AND Patient does not concurrently use a strong CYP 3A inhibitors; AND A baseline alanine aminotransferase (ALT), aspartate aminotransferase (AST), and bilirubin have been performed and are within normal range (results must be within 3 months of request). Labs must also be repeated 2 weeks and 4 weeks after initiation and then continued monthly for the first 18 months and every 3 months thereafter. Renewal Criteria (6-month duration): Patient's most recent ALT, AST, and bilirubin are within normal range (results must be within 3 months of request)		General P. Form			
ynarque Pak®	NP	See Jynarque® prior authorization criteria					
amsca®	NP	 Diagnosis of hyponatremia; AND Medication was initiated in a hospital setting 					
olvaptan	NP	See Samsca® prior authorization criteria					



		GASTROINTESTINAL Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated	ı.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		5-ASA Derivatives, Oral		
Apriso®	Р		4/day	
Delzicol®	Р		6/day]
sulfasalazine	Р		8/day]
sulfasalazine EC	Р		8/day]
Azulfidine®	NP		8/day	
Azulfidine® EN	NP		8/day	
balsalazide	NP		9/day	1
Colazal®	NP		9/day	1
Dipentum®	NP		4/day	General PA
Lialda®	NP		4/day	<u>Form</u>
mesalamine DR caps	NP		6/day	1
mesalamine DR tabs	NP		800 mg: 6/day 1.2 gm: 4/day	
mesalamine ER 24 Hour caps	NP		4/day	
mesalamine ER caps			500 mg: 8/day	1
Pentasa®	NP		250 mg: 16/day; 500 mg: 8/day	
		Agents for Chronic Constipation		
Linzess®	Р		1/day	General PA
lubiprostone	Р		2/day	<u>Form</u>
Movantik®	P	 Age ≥ 18 years; AND One of the following: Diagnosis of opioid-induced constipation with chronic non-cancer pain Diagnosis of opioid-induced constipation with chronic pain related to prior cancer or its treatment who do not require frequent (e.g., weekly) opioid dosage escalation; AND Submission of medical records (e.g., chart notes, control substance monitoring data base) or confirmed pharmacy claims documenting at ≥1 of opioid therapy within the past 90 days; AND Prescriber attests that Movantik® will be discontinued when opioid treatment is discontinued 	1/day	General PA Form
Amitiza®	NP		2/day	



GASTROINTESTINAL Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. PDL Medication **Prior Authorization Criteria** Qty. Limits **PA Form** Age ≥ 18 years; AND · Patient has diagnosis of chronic idiopathic constipation (CIC); AND **General PA** Motegrity® Patient does not have intestinal perforation or obstruction due to structural or functional disorder of the gut wall, 1/day Form obstructive ileus, or severe inflammatory conditions of the intestinal tract (e.g., Crohn's disease, ulcerative colitis); AND Trial and failure of, or contraindication, or intolerance to lubiprostone AND Linzess® See Motegrity® prior authorization criteria; AND NP prucalopride · Clinically valid why Motegrity® cannot be used • Age ≥ 18 years; AND • One of the following: o Diagnosis of opioid-induced constipation with chronic non-cancer pain o Diagnosis of opioid-induced constipation with chronic pain related to prior cancer or its treatment who do not require frequent (e.g., weekly) opioid dosage escalation General PA Relistor® injectable NP o Diagnosis of opioid-induced constipation with advanced illness or pain caused by active cancer who requires opioid Form dosage escalation for palliative care; AND · Submission of medical records (e.g., chart notes, control substance monitoring data base) or confirmed pharmacy claims documenting at >1 of opioid therapy within the past 90 days; AND Prescriber attests that Relistor® will be discontinued when opioid treatment is discontinued Age ≥ 18 years; AND • One of the following: o Diagnosis of opioid-induced constipation with chronic non-cancer pain o Diagnosis of opioid-induced constipation with chronic pain related to prior cancer or its treatment who do not **General PA** Relistor® tablets NP require frequent (e.g., weekly) opioid dosage escalation; AND 3/day Form Submission of medical records (e.g., chart notes, control substance monitoring data base) or confirmed pharmacy claims documenting at >1 of opioid therapy within the past 90 days; AND Prescriber attests that the requested drug will be discontinued when opioid treatment is discontinued • Trial and failure of, or contraindication, or intolerance to Movantik® See Relistor® tablets prior authorization criteria; AND Symproic® NP 1/day Patient does not have known or suspected gastrointestinal obstruction Age ≥ 18 years; AND • Diagnosis of one of the following: o Chronic idiopathic constipation (CIC) **General PA** NP Trulance® 1/day o Irritable bowel syndrome with constipation (IBS-C); AND Form Patient does not have a known or suspected mechanical gastrointestinal obstruction; AND Trial and failure of, or contraindication, or intolerance to, lubiprostone OR Linzess®



	GASTROINTESTINAL Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.						
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
		Agents for Irritable Bowel Syndrome (IBS)		_			
alosetron	P	Initial Criteria: Patient is female and ≥ 18 years of age; AND Diagnosis of severe, diarrhea-predominant, irritable bowel syndrome (IBS); AND Chronic IBS symptoms lasting 6-months or more; AND Provider has ruled out anatomic or biochemical abnormalities of the GI tract; AND Patient is not concomitantly using fluvoxamine; AND Patient does not have a history of the following conditions: Chronic or severe constipation or sequalae from constipation Intestinal obstruction, stricture, toxic megacolon, gastrointestinal perforation, and/or adhesions Ischemic colitis, impaired intestinal circulation, thrombophlebitis, or hypercoagulable state Crohn's disease or ulcerative colitis Diverticulitis Severe hepatic impairment Renewal Criteria: Patient continues to meet the initial criteria; AND Patient has not experienced any treatment-restricting adverse effects (e.g., severe constipation); AND Positive response to therapy (e.g., decrease stool frequency, frequent bowel urgency, and abdominal pain)	2/day	General PA Form			
Linzess®	Р		1/day				
lubiprostone	Р		2/day				
Amitiza®	NP		2/day				
lbsrela®	NP	Initial Criteria: Patient is ≥ 18 years of age; AND Diagnosis of irritable bowel syndrome with constipation (IBS-C); AND Patient does not have known or suspected mechanical gastrointestinal obstruction; AND Trial and failure, contraindication, or intolerance to lubiprostone AND Linzess® Renewal Criteria: Patient continues to meet the initial criteria; AND Patient has not experienced any treatment-restricting adverse effects (e.g., severe diarrhea); AND Positive response to therapy (e.g., decrease stool frequency, frequent bowel urgency, and abdominal pain)	2/day				
Lotronex®	NP	Clinically valid reason why the preferred generic alosetron cannot be used	2/day				



		GASTROINTESTINAL Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated	<i>1</i> .	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Viberzi®	NP	Initial Criteria: Patient is ≥ 18 years of age; AND Diagnosis of severe, diarrhea-predominant, irritable bowel syndrome (IBS); AND Patient does not have history of the following: alcohol abuse/addiction or drink more than 3 alcoholic drinks per day pancreatitis or structural diseases of the pancreas severe hepatic impairment (Child Pugh Class-C) severe constipation absence of gallbladder biliary duct (gallbladder) obstruction or Sphincter of Oddi disease/dysfunction Renewal Criteria: Patient continues to meet the initial criteria; AND Patient has not experienced any treatment-restricting adverse effects (e.g., severe diarrhea); AND Positive response to therapy (e.g., decrease stool frequency, frequent bowel urgency, and abdominal pain)	2/day	
Xifaxan®	NP	One of the following: Treatment of uncomplicated traveler's diarrhea (1-month approval duration); AND Request is for Rifaximin 200 mg tablets; AND Trial and failure, contraindication, intolerance, or resistance to a fluoroquinolone or azithromycin Treatment of diarrhea-predominant IBS (3-month approval duration) Documented use for reduction in risk of overt hepatic encephalopathy (12-month approval duration)	3/day	
		Antidiarrheals		
Mytesi®	NP	 Patient has non-infectious diarrhea of at least one month duration; AND Patient has a diagnosis of HIV or AIDS; AND Patiently is currently receiving anti-retroviral therapy 		
		Antiemetics: 5-HT3 Receptor Antagonists		
ondansetron tablets and ODT	Р	Note: Prior authorization is not required for quantities up to 30 tablets per 90 days. For requests that exceed the quantity limit, one of the following must be met: Receiving highly or moderately emetogenic chemotherapy Receiving radiation therapy Treatment is for post-operative nausea and vomiting (PONV) Nausea or vomiting associated with pregnancy and trial and failure of TWO conventional antiemetics (i.e., metoclopramide, prochlorperazine, dexamethasone, Diclegis)	30/90 days	General PA
Anzemet®		 ONE of the following: Receiving highly or moderately emetogenic chemotherapy Receiving radiation therapy Treated for post-operative nausea and vomiting (PONV); AND Trial and failure, contraindication, or intolerance to a preferred 5HT3 antagonist 	2/30	<u>Form</u>
granisetron	NP	See Anzemet® prior authorization criteria	Tabs: 60/30 days Inj: 2 mL/30 days	



		GASTROINTESTINAL		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated	l.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
ondansetron solution	NP	 Patient < 6 years of age; OR The requested dose is not achievable with ondansetron ODT; OR Allergy or intolerance to inactive ingredient in ODT tab (e.g., dye, filler, excipient) 		
Sancuso®	NP	See Anzemet® prior authorization criteria	1/30 days	
		Antiemetics: Anticholinergics		
promethazine	Р	 Patients < 2 years of age; AND Prescriber documents medical necessity; AND Prescriber is aware of contraindication and agrees to accept risk Note: Prior authorization is not required for patients 2 years of age or older 		Promethazine PA Form
Transderm-Scop®	Р	One of the following:	10 patches/30 days	General PA Form
Phenergan®	NP	One of the following: ○ Patient is ≥ 2 years of age, AND - Clinical reason as to why patient cannot use generic equivalent ○ Patients < 2 years of age; AND - Prescriber documents medical necessity; AND - Prescriber is aware of contraindication and agrees to accept risk; AND - Clinical reason as to why patient cannot use generic equivalent		Promethazine PA Form
promethazine suppositories	NP	See promethazine prior authorization criteria Note: Prior authorization is not required for patients 2 years of age or older		
scopolamine patches	NP	See Transderm-Scop® prior authorization criteria; AND • Clinically valid reason as to why preferred Transderm-Scop® cannot be used	10 patches/30 days	General PA Form
		Antiemetics: Delta-9-THC Derivatives		
dronabinol	NP	 Request is for the treatment of severe nausea/vomiting associated with cancer chemotherapy for patients actively being treated for cancer; AND Trial and failure, intolerance, intolerance, medical reason, or contraindication that prohibits taking Emend + 5HT3 receptor antagonist + corticosteroid; OR Request is for the treatment of AIDS-related wasting; AND Trial and failure, intolerance, or contraindication to megestrol acetate oral suspension 		
Marinol®	NP	See dronabinol prior authorization criteria]
Syndros®	NP	See dronabinol prior authorization criteria; AND • Unable to swallow solid dosage forms		



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
	1	Antiemetics: NK-1 Antagonists				
aprepitant	Р	 Receiving a highly emetogenic chemotherapy regimen; OR Receiving a moderately emetogenic chemotherapy regimen and has failed two other antiemetic regimens; OR Treatment for PONV with trial and failure or contraindication to a 5HT3-receptor antagonist; OR Refractory nausea that would require hospitalization 	40 mg: 1/30 days 80 mg: 4/30 days 125 mg: 2/30 days			
Akynzeo®	NP	 ONE of the following: Receiving a highly emetogenic chemotherapy regimen Receiving a moderately emetogenic chemotherapy regimen and has failed other previous antiemetic regimens; AND Trial and failure, contraindication, or intolerance to aprepitant 	2/30 days	General PA Form		
Emend®	NP	See aprepitant prior authorization criteria; AND • Clinically valid reason preferred aprepitant cannot be used	80 mg: 4/30 days Tri-Pack: 2 packs/30 days			
	•	Antiemetics: Miscellaneous				
Diclegis®	Р		4/day			
Bonjesta®	NP	 Patient has a diagnosis of pregnancy-induced nausea or vomiting; AND Patient has failed documented conservative measures (e.g., dietary changes, trigger avoidance); AND Clinically valid reason as to why preferred Diclegis® cannot be used 	2/day	General PA Form		
doxylamine/ pyridoxine	NP	Clinically valid reason as to why preferred Diclegis® cannot be used	4/day			
		Antispasmodics/Anticholinergics				
glycopyrrolate solution	Р	 Patients unable to swallow tablets; OR Patient is < 8 years of age 		General PA		
Cuvposa®	NP	 Patients unable to swallow tablets; OR Patient is < 8 years of age 		<u>Form</u>		
		Combination Products for H. pylori				
Pylera®	Р	Documentation of recent positive <i>H. pylori</i> test	1 box/Rx; 2 courses of			
Talicia®	Р	Documentation of recent positive <i>H. pylori</i> test	therapy/year)			
bismuth subcitrate/ metronidazole/ tetracycline	NP	 Documentation of recent positive <i>H. pylori</i> test; AND Trial and failure, contraindication, or intolerance to a preferred combination agent 		General PA		
lansoprazole/amox/ clarithromycin	NP	 Documentation of recent positive <i>H. pylori</i> test; AND Trial and failure, contraindication, or intolerance to a preferred combination agent 	1 box/Rx; 2 courses of therapy/year)	<u>Form</u>		
Omeclamox-Pak®	NP	 Documentation of recent positive <i>H. pylori</i> test; AND Trial and failure, contraindication, or intolerance to a preferred combination agent 				



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
Voquezna Dual Pak®	NP	 Documentation of recent positive H. pylori test; AND Trial and failure, contraindication, or intolerance to a preferred combination agent 	1 box/Rx; 2 courses of therapy/year)	General PA Form			
Voquezna Triple Pak®	NP	 Documentation of recent positive H. pylori test; AND Trial and failure, contraindication, or intolerance to a preferred combination agent 	1 box/Rx; 2 courses of therapy/year)	General PA Form			
		Fecal Microbiota		-			
Vowst®	NP	 Criteria: (2-month duration) Patient is ≥ 18 years old; AND Treatment is to prevent the recurrence of Clostridioides difficile infection (CDI); AND Patient has had three or more episodes of CDI within the past year; AND Submission of medical records (e.g. chart notes, lab test) of a positive C. difficile stool test with toxin A/B results within the previous 30 days; AND Patient has completed a full treatment course with ONE of the following antibiotic therapies 2 to 4 days prior to initiating Vowst: Fidaxomicin Vancomycin; AND Prescriber by or in consultation with an infectious disease specialist or gastroenterologist; AND The agent will not to be used in combination with other products for prevention of CDI, such as Zinplava or Rebyota 	12 caps/year	General PA Form			
		Gallstone Solubilizing Agents					
ursodiol	Р		200, 250, 300, & 400 mg: 3/day; 500 mg: 2/day:				
Cholbam [®]	NP	 Diagnosis of Bile Acid Synthesis Disorders due to Single Enzyme Defects (SED); OR Agent will be used as adjunctive treatment for manifestations of Peroxisomal Disorders (PDs); AND Prescribed by a hepatologist or gastroenterologist 					
lqirvo® Livdelzi®	NP NP	 Patient has a diagnosis of primary biliary cholangitis (PBC) AND Prescribed by a hepatologist or gastroenterologist AND ONE of the following: Both of the following: Will be taken in combination with ursodiol; AND Submitted lab documentation indicates the patient had an inadequate response (no reduction in ALP or total bilirubin after 1-year trial) to ursodiol; OR Patient has a contraindication, or intolerance to ursodiol See Iqirvo® prior authorization criteria 	1/day	General PA Form			



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
Ocaliva®	NP	See Iqirvo® prior authorization criteria	1/day			
Reltone®	NP		3/day			
Urso Forte®	NP		2/day			
		Hepatotrophics				
Rezdiffra®	NP	 Initial Criteria Patient is 18 years of age or older; AND Diagnosis of noncirrhotic nonalcoholic steatohepatitis (NASH) or metabolic dysfunction associated steatohepatitis (MASH); AND Submission of medical records (e.g. chart notes) confirming disease is fibrosis stage F2 or F3 as confirmed by ONE of the following: FibroScan Fibrosis-4 index (FIB-4) Magnetic Resonance Elastography (MRE) Liver Biopsy; AND Prescriber attests patient is participating in a supervised comprehensive weight management program that encourages behavioral modification, reduced calorie diet, and increased physical activity; AND Patient does not have decompensated cirrhosis (Child-Pugh Class B or C); AND Prescribed by or in consultation with a gastroenterologist or hepatologist Renewal Criteria Prescriber attest patient is participating in a supervised comprehensive weight management program that encourages behavioral modification, reduced calorie diet, and increased physical activity; AND Submission of medical records (e.g., chart notes) documenting a positive clinical response to therapy (e.g. NASH resolution, fibrosis stage improvements) 	1/day	General PA Form		
	ı	Inflammatory Bowel Disease, Miscellaneous Agents				
budesonide foam	Р		66.8 g/day			
Uceris® tablet	Р		1/day			
hydrocortisone AC suppository	Р		12/30 days	General PA Form		
budesonide ER tabs	NP	Trial and failure of preferred Uceris tablets	1/day			
Uceris® foam	NP		66.8 g/day			
		Laxatives				
Sutab [®]	NP		24 tabs per colonoscopy	General PA Form		



GASTROINTESTINAL Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
	•	Motility Agents		•	
metoclopramide	Р		12-week duration limit		
metoclopramide solution	Р		12-week duration limit		
Gimoti®	NP	 Patient must have acute and recurrent diabetic gastroparesis; AND Patient is ≥ 18 years of age; AND Patient does not have a history of tardive dyskinesia (TD) or dystonic reaction to metoclopramide; AND Clinically valid reason why metoclopramide tablets or solution cannot be used 	1 bottle per Rx	General PA Form	
metoclopramide ODT	NP	 Unable to swallow, OR Unable to absorb medications through the GI tract 	12-week duration limit		
Reglan®	NP		12-week duration limit		
		Mucosal Protectants			
Carafate® suspension	NP	 Patient is < 13 years of age; OR Trial and failure, or intolerance to, sucralfate tablets, OR Has documented difficulty swallowing/dysphagia 		General PA Form	
sucralfate suspension	NP	See Carafate suspension prior authorization criteria			



GASTROINTESTINAL

Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.

Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form

Proton Pump Inhibitors

The quantity limit for proton pump inhibitors is 1 per day. If request is for twice daily dosing, one of the following must be met:

- Treatment of H. Pylori (1-month duration)
- Treatment of GI Bleed/Hemorrhagic Gastritis (12-motnh duration)
- Patient has a diagnosis of Barrett's Esophagus with documentation of uncontrolled reflux symptoms or esophagitis (following a trial of once daily PPI therapy)
- Uncontrolled symptoms following a 30-day trial of once daily PPI therapy (1-month duration); renewals will require member to attempt step down to once daily PPI therapy. If patient fails step down to once daily dosing, they will not be asked to step down again

Dexilant®	Р		1/day	
esomeprazole	Р		1/day	C
lansoprazole	Р		1/day	General PA Form
Nexium® pack	Р	Unable to swallow solid dosage forms	1/day	101111
omeprazole	Р		1/day	1
omeprazole ODT	Р		1/day	
omeprazole/sodium bicarbonate	Р		1/day	General PA
pantoprazole	Р		1/day	<u>Form</u>
Protonix® packs	Р		1/day	
dexlansoprazole	NP		1/day	
esomeprazole packs	NP	 Unable to sallow solid dosage forms; AND Trial, failure, contraindication, or intolerance to Protonix[®] suspension and Nexium granules 	1/day	General PA Form
First-Lansoprazole®	NP	 Unable to sallow solid dosage forms; AND Trial, failure, contraindication, or intolerance to Protonix suspension packets; OR Patient is < 6 years of age 	1/day	
Konvomep®	NP	See First-Lansoprazole® prior authorization criteria	1/day	
lansoprazole ODT	NP		1/day	General PA
Nexium®	NP		1/day	<u>Form</u>
pantoprazole pack	NP	Clinically valid reason why the preferred Protonix® suspension cannot be used	1/day	<u> </u>
Prevacid®	NP		1/day	General PA
Prevacid SoluTab®	NP	 Unable to swallow solid oral dosage forms; AND Trial, failure, contraindication, or intolerance to Protonix[®] suspension 	1/day	Form Form



GASTROINTESTINAL Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.				
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Voquezna®	NP	 Patient is 18 years of age or older; AND One of the following diagnoses: Active treatment of erosive esophagitis (2-month approval duration); OR Maintenance treatment of healed erosive esophagitis (6-month approval duration); AND Request is for Voquezna 10 mg tablets; OR Non-erosive gastroesophageal reflux disease (GERD) (1-month approval duration); AND Request is for Voquezna 10 mg tablets; AND Trial and failure, contraindication, or intolerance to TWO preferred proton pump inhibitors 	1/day; AND 10 mg: 180 days/ year 20 mg: 60 days/ year	
Prilosec®	NP		1/day	
Protonix® tablets	NP		1/day	
rabeprazole	NP		1/day	General PA
Zegerid®	NP		1/day	<u>Form</u>



		IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDI	Prior Authorization Criteria	Qty. Limits	PA Form
		Allergen Specific Immunotherapy		•
Grastek®	NP	 Diagnosis of grass pollen-induced allergic rhinitis with or without conjunctivitis; AND Documentation initial dose was administered in the physician office or medical facility; AND Must be prescribed by an allergy/immunology specialist; AND Patient's diagnosis is confirmed with documentation of ONE of the following: A positive skin test to ONE of the pollen extracts contained in the requested agent Pollen specific IgE antibodies to ONE of the pollen extracts contained in the requested_agent; AND Trial and failure, contraindication, or intolerance to ONE agent from TWO of the following classes: Oral antihistamine Intranasal antihistamine Intranasal corticosteroid Leukotriene receptor antagonist; AND Documented trial/failure or drug-drug interaction of subcutaneous allergen immunotherapy (SCIT, or allergy shots) [Note: Failure defined as lack of efficacy, allergic reaction, documented intolerable side effects; agent will not be approved for needle phobia]; AND Patient has been prescribed and trained to administer epinephrine in case of severe allergic reaction; AND Oral Anti-allergens will NOT be approved if patient meets ANY of the following:	1/day	General PA Form
Odactra®	NP	 Diagnosis of house dust mite (HDM) induced allergic rhinitis with or without conjunctivitis; AND Patient's diagnosis confirmed with documentation of ONE of the following: Confirmed in vitro IgE antibodies to Dermatophagoides farinae or Dermatophagoides pteronyssinus HDMs Confirmed skin testing to licensed HDM allergen extracts; AND Prescribed by or in consultation with an allergy/immunology specialist; AND Documentation initial dose was administered in the physician office or medical facility; AND Trial and failure, contraindication, or intolerance to ONE agent from TWO of the following classes: Oral antihistamine Intranasal antihistamine Intranasal corticosteroid Leukotriene receptor antagonist; AND Patient has been prescribed and trained to administer epinephrine in case of severe allergic reaction; AND Oral Anti-allergens will NOT be approved if patient meets ANY of the following: Patient experienced a severe reaction post initial dose administered in the physician's office Patient has a history of severe, unstable, or uncontrolled asthma Patient has a history of eosinophilic esophagitis 	1/day	General PA Form



		IMMUNOLOGICS		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Oralair®	NP	 Diagnosis of grass pollen-induced allergic rhinitis with or without conjunctivitis; AND Documentation initial dose was administered in the physician office or medical facility; AND Must be prescribed by an allergy/immunology specialist; AND Patient's diagnosis is confirmed with documentation of ONE of the following: A positive skin test to ONE of the pollen extracts contained in the requested agent Pollen specific IgE antibodies to ONE of the pollen extracts contained in the requested agent; AND Trial and failure, contraindication, or intolerance to ONE agent from TWO of the following classes: Oral antihistamine Intranasal antihistamine Intranasal corticosteroid Leukotriene receptor antagonist; AND Documented trial/failure or drug-drug interaction of subcutaneous allergen immunotherapy (SCIT, or allergy shots) [Note: Failure defined as lack of efficacy, allergic reaction, documented intolerable side effects; agent will not be approved for needle phobia]; AND Patient has been prescribed and trained to administer epinephrine in case of severe allergic reaction; AND Oral Anti-allergens will NOT be approved if patient meets ANY of the following: Patient experienced a severe reaction post initial dose administered in the physician's office Patient has a history of severe, unstable, or uncontrolled asthma Patient has a history of severe, unstable, or uncontrolled asthma Patient has a history of severe, unstable, or uncontrolled asthma Patient has a history of severe, unstable, or uncontrolled asthma Patient has a history of severe, unstable, or uncontrolled asthma Patient has	tabs: 1/day; Dose Pak: total max limit 100 mg IR/300 mg IR	General PA Form
Palforzia®	NP	Initial Criteria: Diagnosis of peanut allergy confirmed by one of the following: ○ Serum peanut-specific immunoglobulin E (IgE) of greater than or equal to 0.35 kUA/L ○ Mean wheal diameter greater than or equal to 3 mm compared to control on skin prick testing for peanut; AND Initial doses for each up-dose will be administered and monitored at the prescriber's office and distributed by the specialty pharmacy; AND Prescribed by, or in consultation with, an allergist or immunologist that is enrolled in Palforzia REMS Program; AND Provider must prescribe injectable epinephrine, instruct, and train patients on its appropriate use; AND Must be used in conjunction with a peanut-avoidant diet; AND Patient must not have ANY of the following: ○ Severe, persistent, or uncontrolled Asthma ○ History of eosinophilic esophagitis or other eosinophilic gastrointestinal disease ○ History of severe or life-threatening episode(s) of anaphylaxis or anaphylactic shock within the past 2 months Renewal Criteria: Documentation (medical records, chart notes, etc.) of tolerance to therapy during the initial dose escalation and up-dosing phases; AND Documentation of positive clinical response to Palforzia therapy; AND Patient continues to use in conjunction with a peanut-avoidant diet; AND Prescribed by, or in consultation with, an allergist or immunologist that is enrolled in the Palforzia REMS Program		General PA Form



		IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Ragwitek®	NP	 Diagnosis of grass pollen-induced allergic rhinitis with or without conjunctivitis; AND Documentation initial dose was administered in the physician office or medical facility; AND Must be prescribed by an allergy/immunology specialist; AND Patient's diagnosis is confirmed with documentation of ONE of the following: A positive skin test to ONE of the pollen extracts contained in the requested agent Pollen specific IgE antibodies to ONE of the pollen extracts contained in the requested agent; AND Trial and failure, contraindication, or intolerance to ONE agent from TWO of the following classes: Oral antihistamine Intranasal antihistamine Intranasal corticosteroid Leukotriene receptor antagonist; AND Documented trial/failure or drug-drug interaction of subcutaneous allergen immunotherapy (SCIT, or allergy shots) [Note: Failure defined as lack of efficacy, allergic reaction, documented intolerable side effects; agent will not be approved for needle phobia]; AND Patient has been prescribed and trained to administer epinephrine in case of severe allergic reaction; AND Oral Anti-allergens will NOT be approved if patient meets ANY of the following: Patient has concomitant allergen immunotherapy Patient has a history of severe, unstable, or uncontrolled asthma Patient has a history of eosinophilic esophagitis; AND Treatment is requested within 12 wks prior to season of allergen being treated (Ragweed season: August-December) Note: Prior authorizations may be processed for Ragwitek® between May 1st thru July 31st; with PA requests being accepted 2 weeks prior to this period. Requests received after July 31st will	1/day	General PA Form



		IMMUNOLOGICS		
Medication	PDL	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Prior Authorization Criteria	Qty. Limits	PA Form
		Anti-Rheumatic: Kinase Inhibitors	<u> </u>	
Rinvoq [®]	P	Prescriber attests to each of the following:	1/day	General PA Form



IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.						
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
Rinvoq LQ®	Р	Initial Criteria (6-month duration): Prescriber attests to each of the following: Patient is not concurrently taking biologic agents (i.e., adalimumab, anakinra, etanercept, rituximab, tocilizumab, infliximab) OR potent immunosuppressants (i.e., azathioprine, cyclosporine); AND Benefits of using this agent outweigh heart-related events or cardiovascular risk factors; AND Risk of malignancy has been considered, and it has been determined that Jak inhibitor therapy is appropriate; AND One of the following: Diagnosis of active Polyarticular Juvenile Idiopathic Arthritis (pJIA); AND Trial and failure, contraindication, or intolerance to leflunomide, methotrexate, OR sulfasalazine; OR Diagnosis of active psoriatic arthritis (PSA); AND Trial and failure, contraindication, or intolerance to a TNF-inhibitor (e.g. Humira, Enbrel); AND One of the following: Patient weighs <30 kg Patient is unable to swallow solid dosage forms Renewal criteria: One of the following: Patient weighs <30 kg Patient weighs <30 kg Patient is unable to swallow oral dosage forms; AND Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of swollen joint counts)	30 mL/day			
Xeljanz® tablet	Р	Initial Criteria (6-month duration): Prescriber attests to each of the following: Patient is not concurrently taking biologic agents (i.e., adalimumab, anakinra, etanercept, rituximab, tocilizumab, infliximab) OR potent immunosuppressants (i.e., azathioprine, cyclosporine); AND Benefits of using this agent outweigh heart-related events or cardiovascular risk factors; AND Risk of malignancy has been considered and it has been determined that Jak inhibitor therapy is appropriate; AND One of the following: Diagnosis of moderately to severely active Rheumatoid Arthritis (RA), or active Polyarticular Course Juvenile Idiopathic Arthritis (pcJIA); AND Trial and failure, contraindication, or intolerance to leflunomide, methotrexate, OR sulfasalazine; AND Trial and failure or intolerance to a TNF-inhibitor (e.g. Humira, Enbrel) Diagnosis of active Psoriatic Arthritis (PsA); AND Trial and failure, contraindication, or intolerance to a TNF-inhibitor (e.g. Humira, Enbrel); OR Diagnosis of moderately to severely active Ulcerative Colitis (UC); AND Trial and failure, contraindication, or intolerance to Humira Diagnosis of Ankylosing spondylitis; AND Trial and failure, contraindication, or intolerance to a TNF-inhibitor (e.g. Humira, Enbrel) Renewal Criteria: Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of tender and swollen joint counts, lower UC disease activity index)	2/day			



		IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Olumiant [®]	NP	Initial Criteria (6-month duration): Prescriber attests to each of the following: Prescriber attests to each of the following: Prescriber is not concurrently taking biologic agents (i.e., adalimumab, anakinra, etanercept, rituximab, tocilizumab, infliximab, abatacept) OR potent immunosuppressants (i.e., azathioprine, cyclosporine); AND Benefits of using this agent outweigh heart-related events or cardiovascular risk factors Risk of malignancy has been considered and it has been determined that Jak inhibitor therapy is appropriate; AND One of the following: Diagnosis of moderately to severely active Rheumatoid Arthritis; AND Trial and failure, contraindication, or intolerance to leflunomide, methotrexate, OR sulfasalazine; AND Trial and failure, contraindication, or intolerance a preferred TNF-inhibitors (e.g., Enbrel, Humira); AND Trial and failure, contraindication, or intolerance to ONE preferred agent; OR Diagnosis of Severe alopecia areata; AND Patient is at ≥ 18 years old but less than 21 years old (indication is not a covered benefit in patients ≥ 21 years old); AND Recipient has ≥ 50% scalp hair loss; AND Prescriber attest patient does not have other underlying causes of hair loss (e.g. male pattern hair loss (androgenic alopecia), female pattern hair loss, telogen effluvium, traction alopecia, and tinea capitis); AND Recipient must be evaluated every 4 months by a physician and submit chart documentation indicating patient has had improved hair growth/decreased hair loss Renewal Criteria: Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of tender and swollen joint counts, lower UC disease activity index, Alopecia areata: improvement in extent and density of scalp hair loss) Note: Will not be covered for COVID-19 treatment in post hospitalized patients	1/day	General PA Form
Xeljanz® solution	NP	Initial Criteria: Prescriber attests to each of the following: Patient is not concurrently taking biologic agents (i.e., adalimumab, anakinra, etanercept, rituximab, tocilizumab, infliximab, abatacept) OR potent immunosuppressants (i.e., azathioprine, cyclosporine); AND Benefits of using this agent outweigh heart-related events or cardiovascular risk factors; AND Risk of malignancy has been considered and it has been determined that Jak inhibitor therapy is appropriate; AND Diagnosis of active Polyarticular Juvenile Idiopathic Arthritis (pcJIA); AND Trial and failure, contraindication, or intolerance to leflunomide, methotrexate, OR sulfasalazine; AND Trial and failure or intolerance to a TNF-inhibitor (e.g. Humira, Enbrel); AND One of the following: Patient weighs <40 kg Patient is unable to swallow oral dosage forms Renewal Criteria One of the following: Patient weighs <40 kg Patient usuable to swallow oral dosage forms Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of tender and swollen joint counts, lower UC disease activity index)	10 mL/day	General PA Form



		IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Xeljanz® XR 11 mg	NP	See Xeljanz® tablet prior authorization criteria; AND Clinically valid reason why the preferred Xeljanz immediate release product cannot be used	1/day	
Xeljanz® XR 22 mg	NP	Initial Criteria (6-month duration): Prescriber attests to each of the following: Patient is not concurrently taking biologic agents (i.e., adalimumab, anakinra, etanercept, rituximab, tocilizumab, infliximab, abatacept) OR potent immunosuppressants (i.e., azathioprine, cyclosporine); AND Benefits of using this agent outweigh heart-related events or cardiovascular risk factors; AND Risk of malignancy has been considered and it has been determined that Jak inhibitor therapy is appropriate; AND Diagnosis of moderately to severely active Ulcerative Colitis (UC); AND Trial and failure, contraindication, or intolerance a preferred adalimumab product; AND Clinically valid reason why the preferred Xeljanz immediate release product cannot be used Renewal Criteria: Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of tender and swollen joint counts, lower UC disease activity index)	1/day	General PA Form
		Disease Modifying Anti-Rheumatic Drugs (DMARDs)		
sulfasalazine	Р		8/day	General PA
sulfasalazine EC	Р		8/day	<u>Form</u>
Azulfidine®	NP		8/day	
Azulfidine EN®	NP			General PA
Jylamvo®	NP	 Dosing that will not allow the use of preferred methotrexate tablets; OR Patient unable to swallow methotrexate tablets 	4 syringes/28 days	<u>Form</u>
Otrexup®	NP	 Diagnosis of Rheumatoid Arthritis (RA) or polyarticular Juvenile Idiopathic Arthritis (pJIA); AND Trial/failure of TWO preferred DMARD agents; AND Must have an allergy or contraindication to benzoyl alcohol or other preservative contained in injectable methotrexate that is not in requested agent; OR Patient is experiencing dexterity issues without assistance to a caregiver who can administer the requested agent; OR Diagnosis of psoriasis: Trial and failure of TWO topical antipsoriatic agents; AND Clinically valid reason why oral methotrexate cannot be used; AND One of the following:	4 syringes/28 days	General PA Form
Rasuvo®	NP	See Otrexup® prior authorization criteria	4 syringes/28 days	
Xatmep®	NP	 Age ≤ 12 years; AND One of the following: Dosing that will not allow the use of preferred methotrexate tablets Patient unable to swallow methotrexate tablet 		General PA Form



		IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL		Qty. Limits	PA Form
		Immunomodulators: TNF Inhibitors		•
Enbrel®; Enbrel Mini Cartridge®; Enbrel Sureclick®	P	Initial Criteria (6-month duration): ■ Diagnosis of Ankylosing Spondylitis; OR ■ Diagnosis of chronic, moderate to severe Plaque Psoriasis; AND □ Trial and failure of ONE topical treatment of a corticosteroid, calcipotriene, OR tazarotene; AND □ Trial and failure, or contraindication, of ONE oral treatment with acitretin, methotrexate, OR cyclosporine; OR ■ Diagnosis of active Psoriatic Arthritis (PsA); OR ■ Diagnosis of active Juvenile Psoriatic arthritis (JPsA); OR ■ Diagnosis of Rheumatoid Arthritis (RA) or Polyarticular Juvenile Idiopathic Arthritis (pJIA); AND □ Trial and failure, contraindication, or intolerance to methotrexate, leflunomide, OR sulfasalazine Renewal Criteria: ■ Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of tender and swollen joint counts, a 50% reduction of total Psoriasis Area Severity Index (PASI) score)	25 mg dose: 8 syringes/28 days 50 mg dose: 4 syringes/28 days	General F Form
Humira®; Hadlima®; Simlandi®	P	Initial Criteria (6-month duration):	2 syringes/28 days Starter Packs: 1 kit/28 days Hidradenitis Suppurativa (HS) diagnosis only: 4 syringes/28 days	General P. Form



IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. PDL Medication **Prior Authorization Criteria Qty. Limits PA Form** Initial Criteria (6-month duration): • Diagnosis of one of the following: Ankylosing Spondylitis (AS) - Active Psoriatic Arthritis (PsA) Rheumatoid Arthritis (RA) Polyarticular Juvenile Idiopathic Arthritis (pJIA) Plaque Psoriasis (PsO) 2 injectors/28 days **General PA** Abrilada® Moderately to severely active Crohn's Disease (CD) NP Form - Hidradenitis Suppurativa (HS) Moderately to severely active Ulcerative Colitis (UC) - Non-infectious intermediate, posterior or panuveitis (UV); AND • Clinically valid reason why ALL the preferred adalimumab products cannot be used Renewal Criteria: Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of swollen joint counts, a 50% reduction of total PASI score, endoscopic remission) adalimumab NP | See Abrilada® prior authorization criteria 2 injectors/28 days **General PA** Form Amjevita® NP | See Abrilada® prior authorization criteria 2 injectors/28 days Initial Criteria (6-month duration): • One of the following: o Diagnosis of one of the following: Ankylosing spondylitis (AS) Axial spondyloarthritis, nonradiographic (nr-axSpA) Active psoriatic arthritis (PsA) Rheumatoid arthritis (RA) 2 kits/28 days **General PA** Cimzia[®] Plaque psoriasis (PsO); (4 syringes) Form - Polyarticular Juvenile Idiopathic Arthritis (pJIA); AND o Trial and failure, contraindication, or intolerance to TWO preferred immunomodulators with same indication; OR • Diagnosis of moderately to severely active Crohn's disease; AND o Trial and failure, contraindication, or intolerance to a preferred adalimumab product Entyvio, or infliximab Renewal Criteria: • Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of swollen joint counts, a 50% reduction of total PASI score, endoscopic remission) Cyltezo® See Abrilada® prior authorization criteria 2 injectors/28 days Hulio® See Abrilada® prior authorization criteria 2 injectors/28 days **General PA Form** Hyrimoz® See Abrilada® prior authorization criteria 2 injectors/28 days Idacio® NP | See Abrilada® prior authorization criteria 2 injectors/28 days



		IMMUNOLOGICS		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Simponi®	NP	Initial Criteria (6-month duration): Diagnosis of Ankylosing Spondylitis, active Psoriatic Arthritis (PsA), or Rheumatoid Arthritis (RA): Trial and failure, contraindication, or intolerance to 2 preferred immunomodulators with same indication; OR Diagnosis of moderately to severely active Ulcerative Colitis (UC): Trial and failure to two of the following (or have an intolerance or contraindication to all agents): A preferred adalimumab product Entyvio Infliximab A preferred JAK inhibitor (e.g. Xeljanz and Rinvoq) Renewal Criteria: Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of swollen joint counts, a 50% reduction of total PASI score, endoscopic remission)	1 syringe /28 days	General PA Form
Yuflyma®	NP	See Abrilada® prior authorization criteria	2 injectors/28 days	General PA
Yusimry®	NP	See Abrilada® prior authorization criteria	2 injectors/28 days	<u>Form</u>
Zymfentra®	NP	 Initial Criteria Patient is 18 years of age or older; AND Diagnosis of ONE of the following: Moderately to severely active Crohn's disease Moderately to severely active Ulcerative Colitis; AND Prescriber attests that patient has received three IV doses of infliximab prior to transitioning to subcutaneous therapy; AND Submission of medical records demonstrating a positive clinical response following a treatment minimum of 10 weeks of infliximab IV Renewal Criteria Diagnosis of ONE of the following: Moderately to severely active Crohn's disease Moderately to severely active Ulcerative Colitis; AND Disease response to therapy and tolerability compared to baseline (e.g., decreased UC disease activity index, endoscopic remission, decreased number of soft stools, decreased abdominal pain) 	2/28 days	General PA Form
		Immunomodulators: Interleukin Inhibitors		
Kineret®	Р	Initial Criteria (6-month duration): Diagnosis of Rheumatoid Arthritis; AND Trial and failure, contraindication, or intolerance to methotrexate, leflunomide, OR sulfasalazine; OR Diagnosis of Neonatal-Onset Multisystem Inflammatory Disease (NOMID) Diagnosis of Deficiency of Interleukin-1 Receptor Antagonist (DIRA) Renewal Criteria: Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of swollen joint counts)	1 syringe/day	General PA Form



IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.						
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
Taltz®	Р	Initial Criteria (6-month duration): Diagnosis of chronic, moderate to severe Plaque Psoriasis; AND Trial and failure to ONE topical treatment of a corticosteroid, calcipotriene, OR tazarotene; AND Trial and failure, or contraindication, to ONE oral treatment with acitretin, methotrexate, OR cyclosporine; OR Diagnosis of active Psoriatic Arthritis (PsA); OR Diagnosis of Axial spondyloarthritis (axSpA), Active Ankylosing Spondylitis (AS), or Active non-radiographic axial spondyloarthritis (nr-axSpA) Renewal Criteria: Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of swollen joint counts, a 50% reduction of total PASI score)	1 syringe/28 days	General PA Form		
Tyenne®	Р	Initial Criteria: (6-month duration) ■ Diagnosis of Rheumatoid Arthritis; AND □ Trial and failure, contraindication, or intolerance to methotrexate, leflunomide, OR sulfasalazine; OR ■ Diagnosis of active Systemic Juvenile Idiopathic Arthritis (SJIA); OR ■ Diagnosis of Giant Cell Arteritis (GCA) and ONE of the following: □ Trial and failure of > 90 days of therapy with systemic glucocorticoids, azathioprine, or methotrexate unless contraindicated or intolerance; OR □ Occurrence of GCA relapse while patient on prednisone doses greater than 20 mg/day, OR □ Patient will be utilizing systemic glucocorticoids with tocilizumab Renewal Criteria: ■ Disease response to therapy and tolerability compared to baseline (e.g., decrease number of tender/swollen joint course)	4 injections (3.6mL)/ 28 days	General PA Form		
Actemra®, Actemra ACTPen®	NP	Initial Criteria (6-month duration): Diagnosis of Rheumatoid Arthritis or active Polyarticular Juvenile Idiopathic Arthritis (pJIA); AND Trial and failure, contraindication, or intolerance to methotrexate, leflunomide, OR sulfasalazine; AND Clinically valid reason why the preferred product Tyenne cannot be used; OR Diagnosis of active Systemic Juvenile Idiopathic Arthritis; AND Clinically valid reason why the preferred product Tyenne cannot be used; OR Diagnosis of Giant Cell Arteritis (CGA); AND One of the following: Trial and failure of > 90 days of drug therapy with systemic glucocorticoids, azathioprine, or methotrexate Occurrence of GCA relapse while patient on prednisone doses greater than 20 mg/day Patient will be utilizing systemic glucocorticoid with tocilizumab; AND Clinically valid reason why the preferred product Tyenne cannot be used Diagnosis of systemic sclerosis-associated interstitial lung disease (SSc-ILD); AND Patient is 18 years of age or older; AND Patient's onset of disease was 5 years ago or less; AND Patient has active disease with elevated inflammatory markers or platelets Renewal Criteria: Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of swollen joint counts)	3.6 mL/28 days	General PA Form		



	IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
Arcalyst®	NP	 Diagnosis of Cryopyrin-Associated Periodic Syndromes (CAPS), including Familial Cold Auto-inflammatory Syndrome (FCAS), and Muckle-Wells Syndrome (MWS); OR Diagnosis of Deficiency of Interleukin-1 Receptor Antagonist (DIRA); AND Patient has tried and failed or have contraindication or intolerance to preferred agent Kineret; OR Diagnosis of recurrent pericarditis (RP) and meets all of the following; AND Trial and failure, contraindication, or intolerance to ONE of the following: Colchicine Corticosteroids NSAIDS 	8 vials/month	General PA Form		
Bimzelx®	NP	 Initial Criteria: Diagnosis of moderate to severe Plaque Psoriasis (PsO), Active Psoriatic Arthritis (PsA), or Ankylosing Spondylitis (AS); AND Trial and failure, contraindication, or intolerance to TWO preferred immunomodulators with the same indication; OR Diagnosis of Non-Radiographic Axial Spondyloarthritis (nr-axSpA); AND Trial and failure, contraindication, or intolerance to Taltz; OR Diagnosis of moderate to severe Hidradenitis Suppurativa (HS); AND Trial and failure, contraindication, or intolerance of a preferred adalimumab product Renewal Criteria: Disease response to therapy and tolerability compared to baseline (e.g., reduction of total PASI score, reduction in inflammatory bumps/abscesses) 	1 injections/28 days	General PA Form		
Cosentyx®	NP	Initial Criteria (6-month duration): • Diagnosis of chronic, moderate to severe Plaque Psoriasis(PsO), Ankylosing Spondylitis (AS), active Psoriatic Arthritis (PsA); AND • Trial and failure, contraindication, or intolerance to TWO preferred immunomodulators with same indication; OR • Diagnosis of active non-radiographic axial spondyloarthritis (nr-axSpA) with objective signs of inflammation; AND • Trial and failure, contraindication, or intolerance of Taltz; OR • Diagnosis of Active Enthesitis-related arthritis (ERA); AND • Failed an adequate trial of ONE NSAID (unless contraindicated); AND • Diagnosis of moderate to severe Hidradenitis Suppurativa (HS); AND • Trial and failure, contraindication, or intolerance of a preferred adalimumab product Renewal Criteria: • Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of tender and swollen joint counts, a 50% reduction of total PASI score, reduction in inflammatory bumps/abscesses, decreases in flares)	300 mg dose: 2 pens/28 days; 150 mg dose: 1 pen /28 days Hidradenitis Suppurativa (HS) diagnosis only- 300 mg dose: 4 syringes/28 days	General PA Form		
Kevzara®	NP	 Initial Criteria (6-month duration): Diagnosis of Rheumatoid Arthritis (RA) or active Polyarticular Juvenile Idiopathic Arthritis (pJIA); AND ○ Trial and failure, contraindication, or intolerance to 2 preferred immunomodulators with same indication: OR Diagnosis of Polymyalgia Rheumatic (PR); AND ○ Trial and failure, contraindication, or intolerance to systemic corticosteroids; AND Renewal Criteria (6-month duration): Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of swollen joint counts) 	2 pens or syringes /30 days	General PA Form		



		IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated		
Medication	PDL		Qty. Limits	PA Form
Omvoh® Auto- injector	NP	Initial Criteria: (6-month duration) ■ Diagnosis of Ulcerative Colitis; AND □ Trial and failure to two of the following (or have an intolerance or contraindication to all agents): □ A preferred adalimumab product □ Entyvio □ Infliximab □ A preferred JAK inhibitor (e.g. Xeljanz or Rinvoq); OR ■ Diagnosis of moderately to severely active Crohn's disease (CD); AND □ Trial and failure, contraindication, or intolerance to a preferred adalimumab product, Entyvio, or infliximab; AND □ Prescriber attests that patient has received three IV doses of Omvoh prior to transitioning to subcutaneous therapy Renewal Criteria: ■ Disease response to therapy and tolerability compared to baseline (e.g., endoscopic remission)	2 auto-injectors/28 days	General PA Form
Otulfi®	NP	See prior authorization criteria for Stelara®	See Stelara®	
Pyzchiva®	NP	See prior authorization criteria for Stelara®	See Stelara®	General PA Form
Selarsdi®	NP	See prior authorization criteria for Stelara®	See Stelara®	
Siliq®	NP	 Initial Criteria (6-month duration): Diagnosis of moderate to severe plaque psoriasis (PsO); AND Patient has a contraindication, drug-drug interaction, or adverse reaction to TWO preferred immunomodulator agents with same indication; AND Patient does not have a history of Crohn's disease; AND Prescriber and patient have met the requirements of the Siliq REMS program Renewal Criteria: Disease response to therapy and tolerability compared to baseline (e.g., a 50% reduction of total PASI score) 	2 syringes/28 days	General PA Form
Skyrizi® Steqeyma®	NP	Initial Criteria (6-month duration): Diagnosis of Plaque Psoriasis (PsO) or active psoriatic arthritis (PsA); AND Trial and failure to ALL preferred immunomodulator agents with the same indication; OR Diagnosis of moderately to severely active Crohn's disease (CD); AND Trial and failure, contraindication, or intolerance to a preferred adalimumab product, Entyvio, or infliximab; OR Diagnosis of moderately to severely active Ulcerative Colitis (UC); AND Trial and failure to two of the following (or have an intolerance or contraindication to all agents): A preferred adalimumab product Entyvio Infliximab A preferred JAK inhibitor (e.g. Xeljanz or Rinvoq) Renewal Criteria: Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of swollen joint counts, a 50% reduction of total PASI score)	Cartridge: 1 per 8 weeks Auto-injector, pre- filled syringe, and pre- filled syringe kit: 2 per 84 days	General PA Form



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Medication	PDL	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated Prior Authorization Criteria	Qty. Limits	PA Form
Spevigo®	NP	Initial Criteria: Diagnosis of Generalized pustular psoriasis (GPP) confirmed by ONE of the following: Presence of sterile, macroscopically visible pustules on non-acral skin and pustulation is NOT restricted to psoriatic plaques (i.e. occurs outside of psoriatic plaques) Skin biopsy confirming presence of Kogoj's spongiform pustules; Genetic confirmation of IL36RN, CARD14, or AP1S3 mutation; AND Patient is 12 years of age and older and weights at least 40 kg; AND Prescriber attest to ALL of the following: Treatment is NOT for an active GPP flare Patient has been evaluated and screened for the presence of latent TB infection prior to initiating treatment Patient will not receive live vaccines during therapy and 16 weeks after treatment; AND Trial and failure to BOTH of the following (or have an intolerance or contraindication to all agents): ATNF inhibitor (e.g. adalimumab, infliximab, and etanercept) Taltz; AND Prescribed by, or in consultation with, a dermatologist Renewal Criteria: Submission of medical records (e.g. chart notes) documenting disease response to therapy and tolerability compared to baseline (e.g., decreased number of GPP flares) Note: The Spevigo subcutaneous formulation is not FDA approved for the treatment of GPP flare and will not be approved for that diagnosis. A SQ loading dose is not required following treatment of a GPP flare with IV Spevigo	2/28 days	General PA Form
Stelara® prefilled syringe and 45 mg/0.5 mL vial	NP	Initial Criteria (6-month duration): • Diagnosis of Plaque Psoriasis (PsO) or active Psoriatic Arthritis (PsA); AND • Trial and failure, contraindication, or intolerance to 2 preferred immunomodulators with same indication; OR • Diagnosis of moderately to severely active Crohn's disease (CD); AND • Trial and failure, contraindication, or intolerance to a preferred adalimumab product, Entyvio, or infliximab; OR • Diagnosis of moderately to severely active Ulcerative Colitis (UC); AND • Trial and failure to two of the following (or have an intolerance or contraindication to all agents): - A preferred adalimumab product - Entyvio - Infliximab - A preferred JAK inhibitor (e.g. Xeljanz or Rinvoq) Renewal Criteria: • Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of swollen joint counts, a 50% reduction of total PASI score)	Plaque Psoriasis, Psoriatic Arthritis: 1 injection/84 days Crohn's Disease and Ulcerative Colitis: 1 injection/56 days	General PA Form



		IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Tremfya®	NP	Initial Criteria (6-month duration): • Diagnosis of Plaque Psoriasis (PsO) or active psoriatic arthritis (PsA); AND ○ Trial and failure to ALL preferred immunomodulator agents with the same indication; OR • Diagnosis of moderately to severely active Ulcerative Colitis (UC); AND ○ Trial and failure to two of the following (or have an intolerance or contraindication to all agents): - A preferred adalimumab product - Entyvio - Infliximab - A preferred JAK inhibitor (e.g. Xeljanz or Rinvoq) Renewal Criteria: • Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of swollen joint counts, a 50% reduction of total PASI score)	1 syringe (1 mL) / 56 days 1 autoinjector/ 56 days	General PA Form
ustekinumab	NP	See prior authorization criteria for Stelara®	See Stelara®	
Yesintek®	NP	See prior authorization criteria for Stelara®	See Stelara®	
		Immunomodulators: Miscellaneous		•
Orencia®	Р	 Initial Criteria (6-month duration): Diagnosis of Rheumatoid Arthritis (RA) or Polyarticular Juvenile Idiopathic Arthritis (pJIA); AND ○ Trial and failure, contraindication, or intolerance to leflunomide, methotrexate, or sulfasalazine; OR Diagnosis of active Psoriatic Arthritis PsA) Renewal Criteria: Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of swollen joint counts, a 50% reduction of total PASI score) 	4 mL/28 days	General PA Form
Otezla®	P	Initial Criteria (6-month duration): Diagnosis of Plaque Psoriasis (PsO); AND Trial and failure to a topical treatment of a corticosteroid, calcipotriene, OR tazarotene; AND Trial and failure, or contraindication, to oral treatment with Soriatane®, methotrexate, cyclosporine; OR Diagnosis of active severe Psoriatic Arthritis (PsA); OR Diagnosis of oral lesions associated with Behçet's Disease; AND Patient has active oral ulcers; AND Trial and failure, contraindication, or intolerance to colchicine Renewal Criteria: Disease response to therapy and tolerability compared to baseline (e.g., decrease in number of swollen joint counts, a 50% reduction of total PASI score)	30 mg: 2/day Starter Pack: 1/Rx	General PA Form



		IMMUNOLOGICS		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Entyvio®	NP	 Initial Criteria: (4-month duration) One of the following: Diagnosis of moderate to severe Crohn's disease Diagnosis of moderate to severe ulcerative colitis (UC); AND Trial and failure, contraindication, or intolerance of a TNF- inhibitor (e.g., Humira, Infliximab) supported by paid claims or chart notes; AND Prescriber attests that patient has or will receive ≥ 2 intravenous doses of Entyvio prior to transitioning to subcutaneous therapy Renewal Criteria: Patient is established on Entyvio therapy for ≥ 14 weeks (supported by paid claims or chart notes); AND Documentation of positive disease response to therapy and tolerability compared to baseline (e.g., decreased UC disease activity index, endoscopic remission, decreased stool frequency) 		General PA Form
Sotyktu®	NP	 Initial Criteria (6-month duration): Diagnosis of moderate to severe Plaque Psoriasis (PsO); AND Trial and failure, contraindication, or intolerance to 2 preferred immunomodulators with same indication Renewal Criteria: Disease response to therapy and tolerability compared to baseline (e.g., a 50% reduction of total PASI score) 	1/day	General PA Form
Velsipity [®]	NP	 Initial Criteria (3-month duration) Diagnosis of moderately to severely active ulcerative colitis (UC); AND Patient is ≥ 18 years old; AND Trial and failure to two of the following (or have an intolerance or contraindication to all agents): A preferred adalimumab product Entyvio Infliximab A preferred JAK inhibitor (e.g. Xeljanz or Rinvoq); AND Patient does not have a recent (within the previous 6 months) myocardial infarction, unstable angina, stroke, transient ischemic attack (TIA), decompensated heart failure with hospitalization, or Class III/IV heart failure; AND Patient does not have a history or presence of Mobitz Type II second-degree, or third-degree atrioventricular block, sick sinus syndrome, or sino-atrial block (unless treated with a functioning pacemaker) Renewal Criteria Disease response to therapy and tolerability compared to baseline (e.g., endoscopic remission, decreased stool frequency, decreased rectal bleeding) 	1/day	General PA Form



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B.d. diantian	DDI	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		DA Forms
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
	1	Immunosuppressants		I
Astagraf XL®	NP	Trial and failure, contraindication, or intolerance to ONE preferred agent Note: The PA requirement may be overridden at POS via an ICD-10 code override.		General PA Form
Azasan®	NP	 Diagnosis of rheumatoid arthritis; AND Trial and failure of ONE preferred agent with the same indication (e.g., azathioprine); OR All transplant recipients will be allowed a prior authorization for any drug Note: The PA requirement may be overridden at POS via an ICD-10 code override. 		General PA Form
CellCept® tablets	NP	All transplant recipients will be allowed a prior authorization for any drug. Note: The PA requirement may be overridden at POS via an ICD-10 code override.		General PA Form
CellCept® capsules	NP	See CellCept® tablets prior authorization criteria		Comoral DA
Envarsus® XR	NP	Trial and failure, contraindication, or intolerance to ONE preferred agent Note: The PA requirement may be overridden at POS via an ICD-10 code override.		General PA Form
Imuran®	NP	See Azasan® prior authorization criteria		General PA
Myfortic [®]	NP	See CellCept® tablets prior authorization criteria		<u>Form</u>
Neoral*	NP	 Diagnosis of rheumatoid arthritis or plaque psoriasis; AND Trial and failure of ONE preferred agent with the same indication (e.g., cyclosporine, GENGRAF); OR All transplant recipients will be allowed a prior authorization for any drug Note: The PA requirement may be overridden at POS via an ICD-10 code override. 		General PA Form
Prograf® capsules	NP	See CellCept® tablets prior authorization criteria		General PA
Prograf [®] packets	NP	Trial and failure, contraindication, or intolerance to ONE preferred agent Note: The PA requirement may be overridden at POS via an ICD-10 code override.		Form
Sandimmune [®] caps	NP	See CellCept® tablets prior authorization criteria		General PA
Zortress®	NP	See CellCept® tablets prior authorization criteria		<u>Form</u>
		Lupus and Lupus Nephritis Agents		
Benlysta®	NP	Initial Criteria: (6-month duration) Patient is ≥ 5 years of age; AND Diagnosis of one of the following: Active systemic lupus erythematosus (SLE); OR Active lupus nephritis; AND Diagnosis has been confirmed by biopsy or biopsy is contraindicated in the patient; AND Prescribed by, or consultation with, a rheumatologist or nephrologist; AND Must be used in combination with standard treatment regimens (e.g., corticosteroids, mycophenolate, azathioprine); AND Patient does not have severe active central nervous system lupus Renewal Criteria: Patient meets the initial criteria; AND Positive clinical response to therapy (e.g., reduction in corticosteroid dose, reduction in flares, improvement in organ dysfunction)	4 syringes/28 days	General PA Form



		IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated		
Medication	PDL		Qty. Limits	PA Form
Lupkynis®	NP	Initial Criteria: Patient is ≥ 18 years of age; AND Diagnosis of active lupus nephritis; AND Diagnosis has been confirmed by biopsy or biopsy is contraindicated in the patient; AND Must take in combination with mycophenolate mofetil and corticosteroids; AND Will NOT take in combination with cyclophosphamide or Benlysta; AND Prescribed by, or in consultation with, a rheumatologist or nephrologist; AND Will not be used concomitantly with strong CYP3A4 inhibitors (e.g., ketoconazole, itraconazole, clarithromycin); AND Patient is not pregnant Renewal Criteria: Must take in combination with mycophenolate mofetil and corticosteroids; AND Will NOT take in combination with cyclophosphamide or Benlysta; AND Prescribed by, or in consultation with, a rheumatologist or nephrologist; AND Patient has experienced a positive response to therapy (evidence of long-term preservation of kidney function, prevention of disease flares, prevention of organ damage); AND Patient has not experienced treatment-limiting adverse effects (eGFR decline, blood pressure increase, hypertensive crisis)	6/day	General PA Form
		Multiple Sclerosis Agents, Injectable		
Avonex®	Р		4/28 days	
Avonex Pack®	Р		4/28 days	General PA
Copaxone®	Р		20mg: 1 mL/day 40mg: 12 mL/30 days	Form
Betaseron®	NP		14/28 days	
glatiramer	NP		20mg: 1 mL/day 40mg: 12 mL/30 days	General PA
Glatopa®	NP		1/day	<u>Form</u>



		IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Kesimpta®	NP	Initial Criteria: Patient is ≥ 18 years of age; AND Prescribed by, or in consultation with, a neurologist; AND Patient has a relapsing form of multiple sclerosis (i.e. clinically Isolated syndrome, relapsing-remitting Multiple Sclerosis [RRMS], active secondary progressive disease [SPMS]); AND Prescriber attests that initial dose was administered under the guidance of a healthcare professional; AND Patient meets ONE of the following: Patient has active secondary progressive disease (SPMS); OR Trial and failure, contraindication, or intolerance to one disease modifying therapy for MS; OR Submission of medical records (e.g., chart notes) documenting clinical features of highly active MS such as high radiological burden of disease, high relapse frequency, severe relapse(s) requiring corticosteroids and/or hospitalization, severe physical or cognitive impairment; AND Patient has been evaluated and screened for the presence of hepatitis B virus (HBV) prior to initiating treatment; AND Patient has been counseled to use effective contraception during treatment and for 6-months after the last dose; AND Actating women will be counseled to discontinue breast feeding during treatment; AND Provider has confirmed (via pregnancy test) that the patient is not pregnant prior to receiving treatment entermore. Renewal Criteria: Documentation of positive clinical response to therapy (e.g., improvement or stabilization in radiologic disease activity, clinical relapses, disease progression)	Initiation: 3 pens the 1st month Maintenance: 1 pen/month	General PA Form
Plegridy®	NP	 Patient is ≥ 18 years old; AND Prescribed by, or in consultation with, a neurologist; AND Diagnosis of relapsing forms of Multiple Sclerosis which include clinically isolated syndrome, relapsing-remitting disease, and active secondary progressive disease; AND Trial and failure, contraindication, or intolerance of 2 preferred injectable MS agents 	2 pens/28 days	General PA Form
Rebif®	NP		6 mL /28 days	
		Multiple Sclerosis (MS) Agents, Oral		
dalfampridine ER	Р		2/day	
dimethyl fumarate	Р	 Initial Criteria: Patient must be the labeled age minimum; AND Prescribed by, or in consultation with, a neurologist; AND Patient has a relapsing form of multiple sclerosis (i.e., clinically isolated syndrome, relapsing-remitting disease, active secondary progressive disease) Renewal Criteria: Documentation of positive clinical response to therapy (e.g., improvement or stabilization in radiologic disease activity, clinical relapses, disease progression) 	2/day	General PA Form
fingolimod	Р	See dimethyl fumarate prior authorization criteria	1/day	1
teriflunomide	Р	See dimethyl fumarate prior authorization criteria		1



		IMMUNOLOGICS (2)		
Medication	PDL	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated Prior Authorization Criteria	. Qty. Limits	PA Form
			-	PA FORM
Ampyra®	NP	Clinically valid reason why preferred dalfampridine cannot be used	2/day	General PA
Aubagio®	NP	See dimethyl fumarate prior authorization criteria; AND Clinically valid reason why preferred dalfampridine cannot be used		<u>Form</u>
Bafiertam®	NP	 Initial Criteria: Patient is ≥ 18 years old; AND Prescribed by, or in consultation with, a neurologist; AND Patient has a relapsing form of multiple sclerosis (i.e. clinically Isolated syndrome, relapsing-remitting disease, active secondary progressive disease); AND Trial and failure, contraindication, or intolerance of teriflunomide or fingolimod; AND Trial and failure, contraindication, or intolerance of dimethyl fumarate; AND Renewal Criteria: Documentation of positive clinical response to therapy (e.g., improvement or stabilization in radiologic disease activity, clinical relapses, disease progression) 	4/day	General PA Form
Gilenya®	NP	 Initial Criteria: Patient is ≥ 10 years of age; AND Prescribed by, or in consultation with, a neurologist; AND Diagnosis of a relapsing form of multiple sclerosis (i.e., clinically isolated syndrome, relapsing-remitting disease, active secondary progressive disease); AND Clinically valid reason why preferred fingolimod cannot be used Renewal Criteria: Documentation of positive clinical response to therapy (e.g., improvement or stabilization in radiologic disease activity, clinical relapses, disease progression) 	1/day	General PA Form
Mavenclad®	NP	Initial Criteria: Patient is ≥ 18 years of age; AND Prescribed by, or in consultation with, a neurologist; AND ONE of the following: Diagnosis of relapsing-remitting multiple sclerosis [RRMS]; AND Trial and failure, contraindication, or intolerance to teriflunomide, dimethyl fumarate, or fingolimod; OR Diagnosis of active secondary progressive disease [SPMS]; AND For patients of reproductive potential: Provider has counseled patient to use contraception during treatment and for 6-months after the last dose Lactating women will be counseled to discontinue breast feeding during treatment Provider has confirmed (via pregnancy test) that the patient is not pregnant prior to receiving treatment; AND Prescriber attests to ALL of the following: Patient does not have a currently malignancy Patient does not have serious active chronic infections such as HIV, tuberculosis, and active hepatitis Patient has been screened for the presence of tuberculosis Renewal Criteria: At least 43 weeks (approx. 10 months) has/will have elapsed since the end of the first treatment course; AND Documentation of positive clinical response to therapy (e.g., improvement or stabilization in radiologic disease activity, clinical relapses, disease progression)	40 tabs/2 years	General PA Form



IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. PDL Medication **Prior Authorization Criteria Qty. Limits PA Form Initial Criteria:** Patient is ≥ 18 years of age; AND Prescribed by, or in consultation with, a neurologist; AND Patient has a relapsing form of multiple sclerosis (i.e. clinically Isolated syndrome, relapsing-remitting disease, active secondary progressive disease); AND • Patient does NOT have any of the following: o Recent (within 6-months) myocardial infarction, unstable angina, stroke, transient ischemic attack (TIA), decompensated heart failure with hospitalization, or Class III/IV heart failure; OR Starter pack: o Prolonged QTc interval at baseline (> 500 msec); OR 1 pack/Rx; o History of Mobitz Type II second- or third-degree atrioventricular block or sick sinus syndrome (unless treated with a **General PA** NP 0.25 mg: Mayzent® functioning pacemaker); OR Form 4 tabs/day; CYP2C9*3/*3 genotype; OR 2 mg: 1 tab/day Active infection (including clinically important localized infections); AND Patient will not be initiating therapy after previous treatment with alemtuzumab (Lemtrada); AND • For female patients of reproductive potential, the following has been addressed: Provider has counseled patient to use effective contraception during treatment with therapy o Lactating patient has been counseled on the risks versus benefits of breastfeeding while on treatment Renewal Criteria: Documentation of positive clinical response to therapy (e.g., improvement or stabilization in radiologic disease activity, clinical relapses, disease progression) Initial Criteria: Patient ≥ 18 years old; AND Prescribed by, or in consultation with, a neurologist; AND ONE of the following: Diagnosis of relapsing-remitting multiple sclerosis [RRMS]; AND - Trial and failure, contraindication, or intolerance to teriflunomide, dimethyl fumarate, or fingolimod; **OR** Diagnosis of active secondary progressive disease [SPMS]; AND • Patient must NOT meet any of the following: o Recent (within 6-months) myocardial infarction, unstable angina, stroke, transient ischemic attack (TIA), **General PA** Ponvory® NP 1/day decompensated heart failure with hospitalization, or Class III/IV heart failure; OR Form o Presence of Mobitz type II second-degree, third degree atrioventricular (AV) block, sick sinus syndrome unless the patient has a functioning pacemaker; AND • For female patients of reproductive potential, all the following has been addressed: Provider has counseled patient to use effective contraception during treatment Lactating patients have been counseled on the risks versus benefits of breastfeeding while on treatment Renewal Criteria: Documentation of positive clinical response to therapy (e.g., improvement or stabilization in radiologic disease activity, clinical relapses, disease progression)



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Tascenso ODT®	NP	 Initial Criteria: Patient is ≥ 10 years of age; AND Prescribed by, or in consultation with, a neurologist; AND Patient has a relapsing form of multiple sclerosis (i.e. clinically Isolated syndrome, relapsing-remitting disease, active secondary progressive disease); AND Patient is unable to swallow sold dosage forms Renewal Criteria: Patient is unable to swallow sold dosage forms; AND Documentation of positive clinical response to therapy (e.g., improvement or stabilization in radiologic disease activity, clinical relapses, disease progression) 	1/day	General PA Form
Tecfidera®	NP	 Initial Criteria: Patient is ≥ 18 years of age; AND Prescribed by, or in consultation with, a neurologist; AND Patient has a relapsing form of multiple sclerosis (i.e. clinically Isolated syndrome, relapsing-remitting disease, active secondary progressive disease); AND Clinically valid reason why preferred dimethyl fumarate cannot be used Renewal Criteria: Documentation of positive clinical response to therapy (e.g., improvement or stabilization in radiologic disease activity, clinical relapses, disease progression) 	2/day	General PA Form
Vumerity®	NP	 Initial Criteria: Patient is ≥ 18 years of age; AND Prescribed by, or in consultation with, a neurologist; AND Patient has a relapsing form of multiple sclerosis (i.e. clinically Isolated syndrome, relapsing-remitting disease, active secondary progressive disease); AND Trial and failure, contraindication, or intolerance of teriflunomide or fingolimod; AND Trial and failure, contraindication, or intolerance of dimethyl fumarate Renewal Criteria: Documentation of positive clinical response to therapy (e.g., improvement or stabilization in radiologic disease activity, clinical relapses, disease progression) 	4/day	General PA Form



		IMMUNOLOGICS		
Medication	PDL	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Prior Authorization Criteria	Qty. Limits	PA Form
Zeposia®	NP	Initial Criteria: Patient is ≥ 18 years of age; AND ONE of the following: Diagnosis of relapsing forms of multiple sclerosis, including clinical isolated syndrome, relapsing-remitting disease, and active secondary progressive disease: AND Prescribed by, or in consultation with, a neurologist; AND Trial and failure, contraindication, or intolerance to teriflunomide, dimethyl fumarate, OR fingolimod (not required for SPMS); OR Diagnosis of moderately to severely active ulcerative colitis (UC) in adults; AND Trial and failure to two of the following (or have an intolerance or contraindication to all agents): A preferred adalimumab product Entyvio Infliximab A preferred JAK inhibitor (e.g. Xeljanz and Rinvoq); AND Patient does NOT have any of the following: Recent (within the previous 6-months) myocardial infarction, unstable angina, stroke, transient ischemic attack (TIA), decompensated heart failure with hospitalization, or Class III/IV heart failure Severe untreated sleep apnea History or presence of Mobitz Type II second-degree, or third-degree atrioventricular block, sick sinus syndrome, or sino-atrial block (unless treated with a functioning pacemaker) Concomitantly taking a monoamine oxidase (MAO) inhibitor; AND For female patients of reproductive potential, all the following has been addressed: Patient has been counseled to use effective contraception during treatment Lactating patients have been counseled on the risks versus benefits of breastfeeding while on treatment Renewal Criteria: Documentation of positive clinical response to therapy (e.g., improvement or stabilization in radiologic disease activity, clinical relapses, disease progression, endoscopic remission)	1/day	General PA Form



		IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicate	ed.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Respiratory and Allergy Biologics		
Adbry®	P	 Initial Criteria (6-month duration): Patient is ≥ 12 years of age; AND Diagnosis of moderate to severe atopic dermatitis with ≥ 1 of the following: Involvement of at least 10% of body surface area (BSA) Scoring Atopic Dermatitis (SCORAD) score of 20 or more Investigator's Global Assessment (IGA) with a score ≥ 3 Eczema Area and Severity Index (EASI) score of ≥ 16 Incapacitation due to AD lesion location (e.g., head and neck, palms, soles, or genitalia); AND Trial and failure (documented by claims) or contraindication to both of the following: A topical corticosteroid of medium to high potency (e.g., mometasone, fluocinolone) A topical calcineurin inhibitor; AND Prescribed by, or in consultation with, a dermatologist, allergist, or immunologist Renewal Criteria: Documented positive response to therapy (e.g., improved pruritus, BSA involvement, EASI, IGA, or SCORAD) 	Initial month: 6 syringes Maintenance: 4 syringes/28 days	General PA Form
Ebglyss®	P	See Adbry prior authorization criteria	Initial 6 months: 4 pens Maintenance: 1 pens/28 days	General PA Form



		IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Dupixent®	P	Initial Criteria (6-month duration): • Diagnosis of moderate to severe asthma; AND • Patient is ≥ 6 years old; AND • One of the following: • Asthma is an eosinophilic phenotype as defined by a baseline (pre- treatment) peripheral blood eosinophil level ≥ 150 cells/µL or peripheral blood eosinophil levels > 300 cells/mc; OR • Patient is currently dependent on oral corticosteroids for the treatment of asthma; AND • Asthma is inadequately controlled as shown by one of the following: • Two more exacerbations requiring systemic corticosteroids within the past 12 months; OR • Asthma-related emergency treatment (e.g., hospitalization, intensive care unit stay, or mechanical ventilation in the past 12 months; OR • Patient is currently dependent on oral corticosteroids for the treatment of asthma; AND • Patient is currently being treated with ONE of the following, unless there is a contraindication: • Combination therapy including BOTH of the following: • One medium or high dose inhaled corticosteroid (ICS); AND • One maximally dosed combination inhaled ICS/ LABA or ICS/LAMA/LABA product; AND • Will be used as adjunct therapy along with above asthma treatment; AND • Prescribed by, or in consultation with, a pulmonologist, allergist, or immunologist Renewal Criteria: • Documentation of a positive clinical response (e.g., reduction in exacerbations, improvement of FEV1, decreased use of rescue medications); AND • Patient is being treated with ONE of the following, unless there is a contraindication: • Combination therapy including both a high-dose ICS and an additional asthma controller medication • One maximally dosed combination inhaled ICS/ LABA or ICS/LAMA/LABA product Atopic Dermatitis Diagnosis Initial Criteria (6-month duration): • Patient is ≥ 6-months of age; AND • Patient is ≥ 6-months of age; AND • Patient of the following: • Involvement of at least 10% of body surface area (BSA) • Scoring Atopic Dermatitis (SCORAD) score of 20 or more • Investigator's Global Assessment (IGA) with a	2 syringes/28 days	General PA Form



		IMMUNOLOGICS		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicate	ed.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Dupixent® (continued)	P	Chronic Idiopathic Urticaria (CIU) Initial Criteria (6-month duration): Patient is ≥ 12 years of age; AND Diagnosis of chronic spontaneous idiopathic urticaria (CSU) or chronic idiopathic urticaria (CIU); AND Patient remains symptomatic despite a 2-week trial to BOTH the following taken in combination: ○ A second generation H1-antihistamine (e.g., cetirizine, fexofenadine, levocetirizine, loratadine); AND ○ One of the following: □ Different second generation H1-antihistamine (e.g., cetirizine, fexofenadine, levocetirizine) □ First generation H1-antihistamine (e.g., diphenhydramine, chlorpheniramine, hydroxyzine) □ H2-receptor antihistamine (e.g., fiphenhydramine, chlorpheniramine, hydroxyzine) □ Leukotriene modifier (e.g., montelukast); AND □ Prescribed by, or in consultation with, an allergist, dermatologist, or immunologist Renewal Criteria: □ Documentation of positive clinical response to therapy (e.g., reduction in exacerbations, itch severity, hives) Chronic rhinosinusitis with nasal polyposis (CRSwNP) Initial Criteria (6-month duration): □ Diagnosis of chronic rhinosinusitis with nasal polyposis (CRSwNP) confirmed by ONE of the following: □ Presence of bilateral nasal polyps □ Patient has previously required surgical removal of bilateral nasal polyps; AND □ Patient has required prior sinus surgery; OR □ Patient has required prior sinus surgery; OR □ Patient has required systemic corticosteroids for CRSwNP □ Symptoms persist after trial of TWO of the following classes of agents: □ Nasal saline irrigations □ Intranasal corticosteroids □ Antileukotriene agents; AND ○ Must be used in combination with intranasal corticosteroid, unless contraindication or intolerance; AND Prescribed by, or in consultation with intranasal corticosteroids ○ Documentation of positive clinical response to therapy; AND	2 syringes/28 days	General PA Form



	IMMUNOLOG	SICS	
	Approval of NP agents requires trial and failure, contraindication, or int		
Medication	DL Prior Authorization Crite	ria Qty. Limits	PA Form
Dupixent® (continued)	Chronic Obstructive Pulmonary Disease (COPD) Initial Criteria (G-month duration): Diagnosis of COPD; AND Patient is ≥ 18 years of age; AND COPD is inadequately controlled as shown by one of the following: OTWO or more exacerbations requiring systemic corticosteroids and, COPD-related emergency treatment (e.g., hospitalization in the pa Patient is currently receiving standard of care COPD treatment, unless of Post-bronchodilator FEV1/FVC ratio <0.7 and FEV1 ≤ 79%; AND Medication will be used as maintenance add-on therapy Renewal Criteria Positive clinical response to treatment (e.g., improved FEV1 from baseli Patient is currently receiving standard of care COPD treatment, unless of Medication will be used as maintenance add-on therapy Renewal Criteria Patient is currently receiving standard of care COPD treatment, unless of Medication will be used as maintenance add-on therapy Eosinophilic Esophagitis (EoE) Initial Criteria (G-month duration): Diagnosis of eosinophilic esophagitis (EOE) as confirmed by an esophage eosinophilis per high power field (documentation required); AND Patient is ≥ 1 years of age and weighs at least 15 kg; AND Patient is experiencing symptoms related to esophageal dysfunction (e. not respond to antacids, gastroesophageal reflux disease-like symptom. Symptoms are inadequately controlled after a trial of ONE of the follow of Proton pump inhibitor (e.g., pantoprazole, omeprazole); OR Topical esophageal corticosteroids (e.g., budesonide, fluticasone); Prescribed by, or in consultation with, a gastroenterologist, allergist, or Renewal Criteria: Diagnosis of Prurigo Nodularis (PN) Initial Criteria: Diagnosis of Prurigo Nodularis (PN); AND Patient has 20 or more nodular lesions (IGA PN-S ≥ 3); AND Inadequate response, intolerance, or contraindication to a topical stero Prescribed by, or in consultation with, a dermatologist, allergist, or imm Renewal Criteria: Documentation of positive clinical response (e.g., improved symptoms of the process of the process of the process of the process of	for antibiotics within the past 12 months; OR st 12 months, mechanical ventilation); AND contraindicated (i.e., ICS/LAMA/LABA); AND me, reduction in COPD exacerbations); AND contraindicated (i.e., ICS/LAMA/LABA); AND eal biopsy demonstrating ≥ 15 intraepithelial 2 syringes/28 day g., dysphagia, food impaction, chest pain that may high and immunologist of eosinophil count, improvement of symptoms) id; AND unologist	General PA Form



		IMMUNOLOGICS		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Fasenra®	P	Asthma Initial Criteria (6-month duration): Diagnosis of moderate to severe asthma; AND Patient is ≥ 6 years old; AND One of the following: Asthma is an eosinophilic phenotype as defined by a baseline (pre-treatment) peripheral blood eosinophil level ≥ 150 cells/µL or peripheral blood eosinophil level > 300 cells/mcl; OR Patient is currently dependent on oral corticosteroids for the treatment of asthma; AND Asthma is inadequately controlled as shown by one of the following: Two more exacerbations requiring systemic corticosteroids within the past 12 months; OR Asthma-related emergency treatment (e.g., hospitalization, intensive care unit stay, or mechanical ventilation); OR Patient is currently being treated with ONE of the following; Patient is currently being treated with ONE of the following, unless there is a contraindication: Combination therapy including both of the following: One high dose inhaled corticosteroid (ICS) One maximally dosed combination ICS/ LABA or ICS/LAMA/LABA product; AND Will be used as adjunct therapy along with above asthma treatment; AND Prescribed by, or in consultation with, a pulmonologist, allergist, or immunologist Renewal Criteria: Documentation of a positive clinical response (e.g., reduction in exacerbations, improvement of FEV1, decreased use of rescue medications); AND Patient is being treated with ONE of the following, unless there is a contraindication: Combination therapy including both a high-dose ICS and an additional asthma controller medication One maximally dosed combination ICS/ LABA or ICS/LAMA/LABA product Essinophilic granulomatosis with polyangilitis (EGPA) Diagnosis of eosinophilitis (granulomatosis with polyangilitis (EGPA); AND Patient is ≥ 18 years of age; AND Patient is ≥ 18 years of age; AND Patient is 2 18 years of age; AND Patient is 2 18 years of age; AND Patient has relapsing disease (defined as a relapse requiring additional or dose escalation of corticosteroids or immunosuppressant, or hospitalization); OR Renewal Criteria: Documentation of posi	Initial (first 3 doses): 1/30 days Maintenance: 1/56 days	General PA Form



		IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicate	d.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Nemluvio [®]	Р	Atopic Dermatitis Diagnosis Initial Criteria (6-month duration): Patient is ≥ 12 years of age; AND Diagnosis of moderate to severe atopic dermatitis with ≥ 1 of the following: Involvement of at least 10% of body surface area (BSA) Scoring Atopic Dermatitis (SCORAD) score of 20 or more Investigator's Global Assessment (IGA) with a score ≥ 3 Eczema Area and Severity Index (EASI) score of ≥ 16 Incapacitation due to AD lesion location (e.g., head and neck, palms, soles, or genitalia); AND Trial and failure (documented by claims) or contraindication to both of the following: A topical corticosteroid of medium to high potency (e.g., mometasone, fluocinolone) A topical calcineurin inhibitor; AND Prescribed by, or in consultation with, a dermatologist, allergist, or immunologist Renewal Criteria: Documented positive response to therapy (e.g., improved pruritus, BSA involvement, EASI, IGA, or SCORAD) Prurigo Nodularis (PN) Initial Criteria: Diagnosis of Prurigo Nodularis (PN); AND Patient age ≥ 18 years; AND Patient age ≥ 18 years; AND Patient has 20 or more nodular lesions (IGA PN-S ≥ 3); AND Inadequate response, intolerance, or contraindication to a topical steroid; AND Prescribed by, or in consultation with, a dermatologist, allergist, or immunologist; AND Trial and failure, contraindication, or intolerance to Dupixent Renewal Criteria: Documentation of positive clinical response (e.g., improved symptoms or IGA PN-S)	2 injections/28 days	General P/ Form



		IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Nucala®	P	Initial Criteria (6-month duration): Diagnosis of moderate to severe asthma; AND Patient is ≥ 6 years old; AND One of the following: Asthma is an eosinophilic phenotype as defined by a baseline (pre-treatment) peripheral blood eosinophil level ≥ 150 cells/µL or peripheral blood eosinophil levels > 300 cells/mcl; OR Patient is currently dependent on oral corticosteroids for the treatment of asthma; AND Asthma is inadequately controlled as shown by one of the following: Two more exacerbations requiring systemic corticosteroids within the past 12 months Asthma-related emergency treatment (e.g., hospitalization, intensive care unit stay, or mechanical ventilation) Patient is currently begendent on oral corticosteroids for the treatment of asthma; AND Patient is currently being treated with ONE of the following; One high dose inhaled corticosteroid (ICS) One high dose inhaled corticosteroid (ICS) One additional asthma controller medication [e.g., LABA, leukotriene receptor antagonist, theophylline]; OR Will be used as adjunct therapy along with above asthma treatment; AND Prescribed by, or in consultation with, a pulmonologist, allergist, or immunologist Renewal Criteria: Documentation of a positive clinical response (e.g., reduction in exacerbations, improvement of FEV1, decreased use of rescue medications); AND Patient is being treated with ONE of the following, unless there is a contraindication: Combination therapy including both a high-dose ICS and an additional asthma controller medication One maximally dosed combination ICS/ LABA or ICS/LAMA/LABA product Eosinophilic granulomatosis with polyangitis (EGPA) Initial Criteria (6-month duration): Diagnosis of eosinophilic granulomatosis with polyangitis (EGPA); AND Patient is 2 18 years of age; AND Patient is 2 18 years of age; AND Patient is acreently taking standard therapy (e.g., cyclophosphamide, rituximab)]; AND One of the following: Patient's disease as defined as a relapse requiring additional or dose escalation of corticosteroi	3 pens or syringes / 28 days	General PA Form



		IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated	ı.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Nucala® (continued)	P	Hypereosinophilic syndrome (HES) Initial Criteria (6-month duration): Patient is ≥ 12 years of age; AND Patient has had HES for > 6-months without an identifiable non-hematologic secondary cause (e.g., drug hypersensitivity, parasitic helminth infection, HIV infection, non-hematologic malignancy, etc.); AND Patient does not have FiP1L1-PDGFRα kinase-positive HES; AND Baseline (pre-Nucala treatment) blood eosinophil level ≥1000 cells/μL (documentation required); AND Patient is currently receiving a stable dose of background HES therapy (e.g., oral corticosteroid, immunosuppressor, or cytotoxic therapy); AND Prescribed by, or in consultation with a pulmonologist, rheumatologist, allergist, or immunologist; AND Renewal Criteria: Documentation of positive clinical response to therapy Chronic rhinosinusitis with nasal polyps (CRSwNP) Diagnosis Initial Criteria (6-month duration): Diagnosis of chronic rhinosinusitis with nasal polyposis (CRSwNP) confirmed by ONE of the following: Presence of bilateral nasal polyps Patient has previously required surgical removal of bilateral nasal polyps; AND Patient has required systemic corticosteroids for CRSwNP; OR Patient has required systemic corticosteroids for CRSwNP; OR Symptoms persist after trial of TWO of the following classes of agents: Nasal saline irrigations Intranasal corticosteroids Antileukotriene agents; AND Must be used in combination with intranasal corticosteroid, unless contraindication or intolerance; AND Prescribed by, or in consultation with, an allergist, immunologist, otolaryngologist, or pulmonologist Renewal Criteria: Documentation of positive clinical response to therapy; AND	3 pens or syringes /28 days	General PA Form



		IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated	<i>t.</i>	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Tezspire®	P	Initial Criteria (6-month duration): Diagnosis of severe asthma; AND Patient is ≥ 12 years old; AND Patient has inadequately controlled asthma as shown by one of the following: Two more exacerbations requiring systemic corticosteroids within the past 12 months; OR Asthma-related emergency treatment (e.g., hospitalization, intensive care unit stay, or mechanical ventilation); OR Patient is currently dependent on oral corticosteroids for the treatment of asthma; AND Patient is currently being treated with ONE of the following, unless there is a contraindication: Combination therapy including both of the following: One medium or high dose inhaled corticosteroid (ICS); AND One additional asthma controller medication [e.g., LABA, leukotriene antagonist, theophylline]; OR One maximally dosed combination inhaled ICS/ LABA or ICS/LAMA/LABA product; AND Prescribed by, or in consultation with, a pulmonologist, allergist, or immunologist Renewal Criteria: Documentation of a positive clinical response (e.g., reduction in exacerbations, improvement of FEV1, decreased use of rescue medications); AND Patient is being treated with ONE of the following, unless there is a contraindication: Combination therapy including both a high-dose ICS and an additional asthma controller medication One maximally dosed combination ICS/ LABA or ICS/LAMA/LABA product	4 pens or syringes /28 days	General PA Form



		IMMUNOLOGICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Xolair®	P	Initial Criteria (6-month duration): Diagnosis of moderate to severe persistent; AND Patient is ≥ 6 years old; AND Dose requested is consistent with corresponding weight and IgE level per manufacturer's dosing chart; AND Baseline (pre-treatment) serum total IgE level greater than or equal to 30 IU/mL and less than or equal to 1300 IU/mL; AND Positive skin test or in vitro reactivity to a perennial aeroallergen; AND Positive skin test or in vitro reactivity to a perennial aeroallergen; AND Positive skin test or in vitro reactivity to a perennial aeroallergen; AND Patient has inadequately controlled asthma as shown by one of the following: Two more exacerbations requiring systemic corticosteroids within the past 12 months; OR Asthma-related emergency treatment (e.g., hospitalization, intensive care unit stay, or mechanical ventilation; OR Patient is currently bependent on oral corticosteroids for the treatment of asthma; AND Patient is currently bependent on oral corticosteroids for the treatment of asthma; AND Patient is currently being treated with ONE of the following, unless there is a contraindication: One medium or high dose inhaled corticosteroid (ICS); AND One maximally dosed combination inhaled ICS/ LABA or ICS/LAMA/LABA product; AND Will be used as adjunct therapy along with above asthma treatment; AND Prescribed by, or in consultation with, a pulmonologist, allergist, or immunologist Renewal Criteria: Documentation of a positive clinical response (e.g., reduction in exacerbations, improvement of FEV1, decreased use of rescue medications); AND Patient is being treated with ONE of the following, unless there is a contraindication: Combination therapy including both a high-dose ICS and an additional asthma controller medication One maximally dosed combination ICS/ LABA or ICS/LAMA/LABA product Chronic Idiopathic Urticaria (CIU) Initial Criteria (6-month duration): Patient is ≥ 12 years of age; AND Diagnosis of chronic spontaneous idiopathic urticaria (CSU) or chronic idiopathic		General PA Form



		IMMUNOLOGICS IMMUNOLOGICS		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Xolair [®] (continued)		IgE-mediated food allergy		General PA Form
Cibinqo®	NP	 Initial criteria (6-month duration): Patient is ≥ 12 years of age; AND Diagnosis of moderate to severe atopic dermatitis with ≥ 1 of the following: Involvement of at least 10% of body surface area (BSA) Scoring Atopic Dermatitis (SCORAD) score of 20 or more Investigator's Global Assessment (IGA) with a score ≥ 3 Eczema Area and Severity Index (EASI) score of ≥ 16 Incapacitation due to AD lesion location (e.g., head and neck, palms, soles, or genitalia); AND Trial and failure (documented by claims) or contraindication to both of the following: A topical corticosteroid of medium to high potency (e.g., mometasone, fluocinolone) A topical calcineurin inhibitor; AND Prescribed by, or in consultation with, a dermatologist, allergist, or immunologist Trial and failure, contraindication, or intolerance of a preferred agent indicated for atopic dermatitis (e.g., Adbry, Dupixent, Ebglyss, Nemluvio) Renewal Criteria: Documented positive response to therapy (e.g., improved pruritus, BSA involvement, EASI, IGA, or SCORAD) 	1/day	General PA Form



		MISCELLANEOUS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise in	ndicated	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Oral Iron Chelators	-	
deferiprone	NP	 Patient has a diagnosis of ONE of the following: Transfusional iron overload due to thalassemia syndromes regardless of prior chelation exposure Transfusional iron overload in patients with sickle cell disease or other anemias; AND Patient is 8 years of age and up (tablets); OR 3 years of age and up (solution); AND ONE of the following: Serum ferritin > 1,000 mcg/L Liver iron concentration is > 3.2 Fe/g dw L; AND Clinically valid reason as to why patient cannot use Exjade® 		General PA Form
deferasirox	NP	See Exjade® prior authorization criteria; AND • Clinically valid reason as to why patient cannot use Exjade®		General PA Form
Exjade®	NP	Patient has a diagnosis of ONE of the following: Chronic iron overload due to blood transfusions in patients 2 years of age and older Non-transfusion-dependent thalassemia (NTDT) in patients aged 10 and older; AND ONE of the following: Serum ferritin > 1,000 mcg/L; OR Liver iron concentration is > 3.2 Fe/g dw L If platelet count is less than 50x109/L., creatinine clearance is greater than 40 mL/min		General PA Form
Ferriprox®	NP	See deferiprone prior authorization criteria		
Ferriprox Twice-A-Day®	NP	See deferiprone prior authorization criteria		General PA Form
Jadenu®	NP	See Exjade® prior authorization criteria; AND • Clinically valid reason as to why patient cannot use Exjade®		General PA Form
		Oral Iron Supplements		
Accrufer®	NP	 Patient has iron deficiency; AND Patient is 18 years of age or older; AND Patient must NOT meet any of the following: Hemochromatosis and other iron overload syndromes Receiving repeated blood transfusions or intravenous iron supplementation Irritable bowel disease (IBD) flare Concomitant use of dimercaprol 	2/day	General PA Form
		Saliva Stimulating Agents		•
pilocarpine	Р		3/day	
cevimeline	NP	Trial and failure, contraindication, or intolerance of pilocarpine	3/day	General PA Form
Evoxac®	NP	Trial and failure, contraindication, or intolerance of pilocarpine	3/day	rom



	ONCOLOGY AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL		Qty. Limits	PA Form		
		Acute Myeloid Leukemia (AML) Agents		•		
Venclexta®	Р		Ramp-Up Phase Dosing: Dispense 7- day supply of 10mg tabs (for 20mg dose); followed by 7-day supply of 50mg tabs	General PA Form		
Daurismo®	NP	Initial Approval Criteria (6-month duration): ONE of the following: Patient ≥ 75 years of age Patient has comorbidities that preclude the use of intensive induction chemotherapy (i.e., Severe Cardiac Disease, Baseline serum creatinine > 1.3 mg/dL, or Baseline Eastern Cooperative Oncology Group (ECOG) performance status of 2); AND Women of child-bearing potential must have a negative pregnancy test; AND Female patients of reproductive potential and males undergoing treatment with female partners of reproductive should use effective contraception during treatment and for at least 30 days after treatment; AND Daurismo® will be used in conjunction with low-dose subcutaneous cytarabine; AND Renewal Criteria (6-month duration): Patient continues to meet initial criteria; AND Patient does not show evidence of progressive disease or unacceptable toxicity (e.g., QTc- interval prolongation)	25 mg: 84/28 days; 100 mg: 28/28 days	General PA Form		
Onureg [®]	NP	 Initial Criteria (6-month duration): Diagnosis of acute myeloid leukemia; AND Patient has achieved first complete remission (CR) or complete remission with incomplete blood count recovery (CRi) following intensive induction chemotherapy and are not able to complete intensive curative therapy; AND Female patients of child-bearing potential have a negative pregnancy test and have been advised that: Female patients should use effective contraception during treatment and for at least 6-months after treatment Males undergoing treatment with female partners of reproductive age should use effective contraception during treatment and for at least 3 months after treatment due to male mediated teratogenicity; AND Renewal Criteria: Patient must continue to meet the initial criteria; AND Patient does not show evidence of progressive disease or unacceptable toxicity (e.g., myelosuppression, renal impairment, hepatic impairment) 	1/day	General PA Form		



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
Vanflyta®	NP	 Initial Criteria: Patient has newly diagnosed acute myeloid leukemia (AML); AND AML is FLT3 internal tandem duplication (ITD)-positive as detected by an FDA-approved test; AND Vanflyta will be used in combination with cytarabine and anthracycline induction and high dose cytarabine consolidation therapy followed by maintenance monotherapy therapy; AND Vanflyta will not be used as maintenance monotherapy following allogeneic hematopoietic stem cell transplantation (HSCT); AND Patient and prescriber are enrolled in the Vanflyta REMS program Renewal Criteria: Patient continues to meet initial criteria; AND Patient does not show evidence of progressive disease or unacceptable toxicity (e.g., hypokalemia, hypomagnesemia, long QT syndrome) 	2/day	General PA Form		
Xospata®	NP	Initial Criteria: Patient has a diagnosis of acute myeloid leukemia (AML); AND AML is positive for FLT3 mutation as detected by an FDA-approved; AND Females of child-bearing potential had a negative pregnancy test within 7 days before starting Xospata®; AND Female and male patients of reproductive potential have been advised to use effective contraception during treatment and for at least 6 and 4 months, respectively, after the last dose Renewal Criteria: Patient continues to meet initial criteria; AND Patient does not show evidence of progressive disease or unacceptable toxicity (e.g., QT prolongation)	3/day	General PA Form		



		ONCOLOGY AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL		Qty. Limits	PA Form
	•	Antimetabolites		
nqovi®	NP	Initial Criteria: (3-month duration) Diagnosis of myelodysplastic syndromes (MDS), patients previously treated and untreated, de novo and secondary MDS with the following French American-British subtypes: Refractory anemia Refractory anemia with ringed sideroblasts Refractory anemia with excess blasts Chronic myelomonocytic leukemia [CMML]) Intermediate-1, intermediate-2, and high-risk international prognostic IPSS groups; AND Patient has tried and failed or is not a candidate for Allogenic stem cell transplantation; AND Prescriber will obtain baseline CBC, creatinine clearance (CrCl), and liver enzymes prior to therapy and prior to each cycle; AND Prescriber must not be pregnant or breastfeeding; AND Patient must not be pregnant or breastfeeding; AND Males undergoing treatment with female partners of reproductive age should use effective contraception during treatment and 3 months after treatment due to male mediated teratogenicity; AND Mill not be used concomitantly with drugs metabolized by cytidine deaminase enzyme (i.e., gemcitabine, capecitabine, cytarabine, azacytidine) Renewal Criteria: (3-month duration) Continues to meet initial criteria; AND Patient has positive disease response, defined as disease stabilization; AND Prescriber attests to delay next cycle and reduce dose if patient experiences elevated liver enzymes or renal impairment OR if patient's absolute neutrophil count (ANC) is less than 1,000 cells/microL and platelet count is less than 50,000 cell/microL	5 per 28-day cycle	General Form
urixan®	NP	 Diagnosis of acute lymphocytic leukemia (ALL); AND ONE of the following: For patients ≤ 11 years of age, no prior authorization required For patients > 11 years of age, Purixan will be approved for patients unable to swallow tablets 		General P Form



	ONCOLOGY AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL		Qty. Limits	PA Form		
		Colorectal Cancer Agents, Miscellaneous				
Lonsurf®	Р		8/day			
Fruzaqla®	NP	Initial Criteria: • Diagnosis of metastatic colorectal cancer; AND • Patient has tried and failed, contraindication, or intolerance to ALL of the following chemotherapy-based regimens: ○ Fluoropyrimidine, ○ Oxaliplatin ○ Irinotecan ○ Anti- vascular endothelial growth factor (VEGF) therapy (e.g., bevacizumab); AND • If RAS wild-type, patient has tried and failed, contraindication, or intolerance to anti-epidermal growth factor receptor (EGFR) therapy (e.g., cetuximab, panitumumab); AND • Prescribed by or in consultation with an oncologist Renewal Criteria: Patient continues to meet initial criteria; AND Patient has positive disease response defined as disease stabilization or decrease in size of tumor or tumor spread; AND Patient does not have unacceptable toxicity (e.g., hypertension, hemorrhagic events)	5 mg: 21/28 days 1 mg: 84/28 days	General P Form		
		EGFR Inhibitors	1			
Lazcluze®	NP	 Initial Criteria Diagnosis of locally advanced or metastatic non-small cell lung cancer (NSCLC); AND Disease is positive for epidermal growth factor receptor (EGFR) exon 19 deletions or exon 21 L858R substitution mutations as detected by an FDA-approved test; AND Lazcluze will be used in combination with Rybrevant; AND Prescribed by, or in consultation with, an oncologist Renewal Criteria Patient continues to meet the initial criteria; AND Patient does not have unacceptable toxicity (e.g., interstitial lung disease, keratitis, venous thromboembolic events) 	80mg: 2/day 240mg: /day	General PA Form		
Vizimpro®	NP	 Patient has a diagnosis of metastatic non-small cell lung cancer (NSCLC); AND Disease is positive for EGFR mutations as confirmed by FDA approved Test (e.g. cobas® EGFR Mutation Test v2); AND Requested agent will be prescribed by, or in consultation with, an oncologist 	1/day	General P		
		Enzyme Inhibitors: ALK				
Lorbrena®	Р		3/day: 25 mg; 1/day: 100 mg	General P		
(alkori sprinkles®	NP	Patient is unable to swallow oral dosage forms				



		ONCOLOGY AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Enzyme Inhibitors: BCR-ABL Kinase		<u>'</u>
dasatinib	NP	Clinically valid reason why preferred Sprycel cannot be used		
Imkeldi®	NP	 Patient is <8 years old; OR Patient is unable to swallow oral dosage forms 	10mL/day	
Scemblix®	NP	Patient has ONE of the following: Newly diagnosed Philadelphia chromosome-positive chronic myeloid leukemia (Ph+ CML) in chronic phase; OR Philadelphia chromosome-positive CML in chronic phase (Ph+ CML-CP) previously treated with two or more tyrosine kinase inhibitors (TKIs); OR Ph+ CML-CP with the T315I mutation; AND Prescribed by, or in consultation with, an oncologist; AND Females of reproductive potential will use effective contraception during treatment and for 1 week after receiving the last dose of Scemblix; AND Patient will not breastfeed during treatment with Scemblix and for 1 week after the last dose		General PA Form
		Enzyme Inhibitors: BRAF Kinase & MEK		
Braftovi®	Р	Initial Criteria: Prescribed by, or in consultation with, an oncologist; AND One of the following: Diagnosis of unresectable or metastatic melanoma; AND Patient is positive for BRAF V600E or V600K mutation as confirmed by an FDA-approved test; AND Prescribed in combination with Mektovi® Diagnosis of metastatic colorectal cancer (CRC); AND Cancer is positive for BRAF V600E mutation as confirmed by an FDA-approved test after prior therapy Diagnosis of metastatic non-small cell lung cancer (NSCLC); Cancer is positive for BRAF V600E mutation, as detected by an FDA-approved test; AND Prescribed in combination with Mektovi® Renewal Criteria: Patient continues to meet initial criteria; AND No unacceptable disease progression or unacceptable toxicity	6/day	General PA Form



ONCOLOGY AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. **PDL** Medication **Prior Authorization Criteria Qty. Limits PA Form Initial Criteria:** Prescribed by, or in consultation with, an oncologist; AND • Prescribed in combination with Braftovi®; AND • One of the following: o Diagnosis of unresectable or metastatic melanoma; AND **General PA** Mektovi® Patient is positive for BRAF V600E or V600K mutation as confirmed by an FDA-approved test; AND 6/day Form Diagnosis of metastatic non-small cell lung cancer (NSCLC); Cancer is positive for BRAF V600E mutation as detected by an FDA-approved test; AND Renewal Criteria: Patient continues to meet initial criteria; AND No unacceptable disease progression or unacceptable toxicity **Initial Criteria:** Diagnosis of neurofibromatosis type 1 (NF1); AND • Patient has plexiform neurofibromas that are BOTH of the following: Inoperable o Causing significant morbidity (e.g., disfigurement, motor dysfunction, pain, airway dysfunction, visual impairment); 1mg tab: 6/day 2 mg: tab 4/day Gomekli® NP Prescribed by, or in consultation with, an oncologist or neurologist; AND 1mg susp tab: 8/day • If the request is for the tablets for oral suspension, patient is unable to swallow solid dosage forms **Renewal Criteria:** · Patient continues to meet initial criteria; AND Prescriber attests that patient has experienced improvement in disease severity and/or symptoms; AND • Patient does not have unacceptable toxicity (e.g., retinal pigment epithelial detachment, ventricular dysfunction, rash) Initial Criteria: Diagnosis of neurofibromatosis type 1 (NF1); AND • Patient has symptomatic, inoperable plexiform neurofibromas (PN); AND Prescribed by, or in consultation with, an oncologist or neurologist 10 mg: 10/day **General PA** Renewal Criteria: Koselugo® 25 mg: 4/day **Form** Patient continues to meet initial criteria; AND Prescriber attests that patient has experienced improvement in disease severity and/or symptoms; AND Patient does not have unacceptable toxicity (e.g., retinal pigment epithelial detachment, severe diarrhea, rash, increased bleeding, myalgia) • Patient is <8 years old; OR General PA NP Mekinist® solution · Patient is unable to swallow solid dosage forms Form



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Ojemda®	NP	Initial Criteria: Diagnosis of relapsed or refractory pediatric low-grade glioma (PLGG); AND Patient has BRAF fusion or rearrangement or BRAF V600 mutation; AND Prescribed by, or in consultation with, an oncologist Renewal Criteria: Patient demonstrates disease stabilization or clinical response to therapy (e.g., stabilized or decrease tumor size, decreased pain, improved vision, increased quality of life)	24/28 days	General PA Form
Tafinlar® solution	NP	 Patient is <8 years old; OR Patient is unable to swallow solid dosage forms 		
		Enzyme Inhibitors: BTK		
Brukinsa®	Р		4/day	
Calquence®	Р		2/day	General PA
Imbruvica® suspension	NP	Patient is unable to swallow capsules		<u>Form</u>
Jaypirca®	NP	Initial Criteria: Diagnosis of mantle cell lymphoma (MCL); AND Patient has received TWO prior therapies including a BTK inhibitor (e.g., Ibrutinib, acalabrutinib, zanubrutinib); AND Jaypirca® will be used as monotherapy; OR Diagnosis of chronic lymphocytic leukemia (CLL) or small lymphocytic lymphoma (SLL); AND Patient has received TWO prior therapies including a BTK inhibitor (e.g., Ibrutinib, acalabrutinib, zanubrutinib) and a BCL-2 inhibitor (e.g., Venclexta); AND Jaypirca® will be used as monotherapy Renewal Criteria: Patient continues to meet the initial criteria; AND Absence of unacceptable toxicity from Jaypirca (e.g., hemorrhage, severe infections, myelosuppression (neutropenia, thrombocytopenia, anemia), atrial fibrillation/flutter, second primary malignancies); AND Patient has positive disease response defined as disease stabilization or decrease in size of tumor or tumor spread	50 mg: 1/day 100 mg: 2/day	General PA Form
	•	Enzyme Inhibitors: CDK		•
Kisqali®	Р		63 tabs/28 days	
Kisqali/Femara®	Р		200mg pack: 49 tabs/28 days; 400 mg pack: 70 tabs/28 days; 600 mg pack: 91 tabs/28 days	General PA Form



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
		Enzyme Inhibitors: FGFR					
Balversa®	NP	 Initial Criteria: Patient has a diagnosis of locally advanced or metastatic urothelial carcinoma; AND Patient has a susceptible FGFR3 or FGFR2 genetic alteration as confirmed by an FDA-approved diagnostic; AND Patient has progressed during or following ≥ 1 prior line of platinum-containing chemotherapy, including within 12 months of neoadjuvant or adjuvant platinum-containing chemotherapy; AND Prescribed by, or in consultation with, an oncologist; AND Provider attests to ALL the following:	3 mg (3/day); 4 mg (2/day); 5 mg (1/day)	General PA Form			
Lytgobi®	NP	Initial Criteria (6-month duration): Patient has diagnosis of unresectable, locally advanced, or metastatic intrahepatic cholangiocarcinoma; AND Patient has fibroblast growth factor receptor 2 (FGFR2) gene fusions or other rearrangements as detected by FDA approved test; AND The patient has progressed on at least one systemic therapy; AND The prescriber attest to ALL of the following: Patient will have an ophthalmological examination including optical coherence tomography (OCT) performed prior to initiation of therapy, every 2 months for the first 6-months of treatment and every 3 months thereafter, and urgently at any time for visual symptoms Prescriber will obtain baseline phosphate levels and monitor for hyperphosphatemia throughout treatment Patient is not pregnant Female patients of reproductive potential and males with female partners of reproductive age have been advised to use effective contraception during treatment and for at least 1 week after the last dose Patient is not concomitantly taking strong dual P-gp and CYP3A Inducers (e.g. rifampin) Renewal Criteria: Positive disease response defined as disease stabilization or decrease in size of tumor or tumor spread; AND Patient does not have unacceptable toxicity (e.g., retinal pigment epithelial detachment, severe hyperphosphatemia)	12 mg: 84/month 16 mg: 112/month 20 mg: 140/month	General PA Form			



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Medication	PDL		Qty. Limits	PA Form
Pemazyre®	NP	Initial Criteria: One of the following: Diagnosis of previously treated unresectable locally advanced or metastatic cholangiocarcinoma with a fibroblast growth factor receptor 2 (FGFR2) fusion or other rearrangement as detected by an FDA-approved test Diagnosis of relapsed or refractory myeloid/lymphoid neoplasms (MLNs) with FGFR1 rearrangement; AND Prescriber attests to ALL the following: Patient will have an ophthalmological examination including optical coherence tomography (OCT) performed prior to initiation of therapy, every 2 months for the first 6-months of treatment and every 3 months thereafter, and urgently at any time for visual symptoms Prescriber will obtain baseline phosphate levels and monitoring for hyperphosphatemia Females and males with female partners will be advised to use effective contraception during treatment and for 1 week after the final dose due to embryo-fetal toxicity Patient is not concomitantly taking strong and moderate CYP3A Inducers Renewal Criteria: Patient continues to meet initial criteria; AND Patient has positive disease response defined as disease stabilization or decrease in size of tumor or tumor spread; AND Patient does not have unacceptable toxicity (e.g., retinal pigment epithelial detachment, severe hyperphosphatemia)	14 tablets/ 21 days	General PA Form
	•	Enzyme Inhibitors: HER2 Targeted Therapies		
Tukysa®	NP	Initial Criteria: ONE of the following: Diagnosis of advanced unresectable or metastatic HER2-positive breast cancer and both of the following: Patient has received at least one or more prior anti-HER2 based regimen; Must be used in combination with trastuzumab and capecitabine; OR Diagnosis of RAS wild-type, HER2-positive unresectable or metastatic colorectal cancer; AND Must be used in combination with trastuzumab; AND Prescribed by, or in consultation with, an oncologist; AND Prescriber attests to ALL of the following: Patient has baseline ALT, AST, and bilirubin measured and within normal limits Patient continues to receive ALT/AST and bilirubin monitoring every 3 weeks during treatment Renewal Criteria: Patient continues to meet initial criteria; AND Patient has positive disease response defined as disease stabilization or decrease in size of tumor or tumor spread; AND Patient does not have unacceptable toxicity (e.g., diarrhea, hepatotoxicity)	50 mg: 10/day 150 mg: 4/day	General PA Form



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
	Enzyme Inhibitors: Isocitrate Dehydrogenase (IDH)						
Tibsovo®	Р		2/day	General PA Form			
Rezlidhia®	NP	Initial Criteria (6-month duration): Patient has diagnosis of relapsed or refractory acute myeloid leukemia (AML); AND Patient has an isocitrate dehydrogenase-1 (IDH1) mutation, as detected by an FDA-approved test Renewal Criteria: Patient continues to meet initial criteria; AND Patient does not show evidence of progressive disease or unacceptable toxicity (e.g., hepatoxicity, differentiation syndrome)	2/day	General PA Form			
Rezlidhia [®]	NP	Initial Criteria: Diagnosis of grade 2 astrocytoma or oligodendroglioma; AND Tumor has susceptible isocitrate dehydrogenase-1 (IDH1) or isocitrate dehydrogenase-2 (IDH2) mutation; AND Patient has had prior surgery including biopsy, sub-total resection, or gross total resection; AND Prescribed by, or in consultation with, an oncologist Renewal Criteria: Patient continues to meet initial criteria; AND Patient does not show evidence of progressive disease or unacceptable toxicity (e.g., hepatoxicity)	40mg: 1/day 10mg: 2/day	General PA Form			



Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Enzyme Inhibitors: KRAS		
Krazati®	NP	ONE of the following: Diagnosis of KRAS G12C-mutated locally advanced or metastatic non-small cell lung cancer (NSCLC) as confirmed by an FDA-approved test; AND Patient has received at least one prior systemic therapy (e.g., immune checkpoint inhibitors [anti-PD-1, PD-L1 immunotherapy], platinum-based chemotherapy); AND Patient has at least one measurable lesion as defined by Response Evaluation Criteria in Solid Tumors (RECIST v1.1); OR Diagnosis of KRAS G12C-mutated locally advanced or metastatic colorectal cancer (CRC) as confirmed by an FDA-approved test; AND Patient has tried and failed, contraindication, or intolerance to ALL the following chemotherapy-based regimens: Fluoropyrimidine Oxaliplatin Irinotecan; AND Rrazati will be used in combination with cetuximab; AND Prescribed by, or in consultation with, an oncologist; AND Prescriber attests that patient is not pregnant or breastfeeding during treatment and for 1 week after the final dose; AND Prescriber attests that patient will be monitored for the following: Hepatotoxicity: Monitor liver function tests ((ALT, AST, and total bilirubin) prior to the start of Krazati, every 3 weeks for the first 3 months of treatment then once monthly as clinically indicated Interstitial Lung Disease (ILD)/Pneumonitis: Monitor for new or worsening pulmonary symptoms; AND Prescriber attests that Patient will not take Krazati with: Acid-reducing agents (e.g., proton pump inhibitors, H2 receptor antagonists, antacids, etc.)	6/day	General PA Form



		ONCOLOGY AGENTS		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Lumakras®	NP	ONE of the following: Diagnosis of KRAS G12C-mutated locally advanced or metastatic non-small cell lung cancer (NSCLC) as confirmed by an FDA-approved test for detection of KRAS G12C; AND Patient has at least one measurable lesion as defined by Response Evaluation Criteria in Solid Tumors (RECIST v1.1); AND Diagnosis of KRAS G12C-mutated locally advanced or metastatic colorectal cancer (CRC) as confirmed by an FDA-approved test; AND Trial and failure, contraindication, or intolerance to ALL the following chemotherapy-based regimens: Fluoropyrimidine Oxaliplatin Irinotecan; AND Agent will be used in combination with panitumumab; AND Prescribed by, or in consultation with, an oncologist; AND Patient has received at least one prior systemic therapy (e.g., immune checkpoint inhibitors [anti-PD-1, PD-L1 immunotherapy], platinum-based chemotherapy); AND Prescriber attests that patient is not pregnant or breastfeeding during treatment with Lumakras and for 1 week after the final dose; AND Prescriber attests that Patient will be monitored for the following: Hepatotoxicity: Liver function tests ((ALT, AST, and total bilirubin) prior to the start of Lumakras, every 3 weeks for the first 3 months of treatment then once monthly as clinically indicated Interstitial Lung Disease (ILD)/Pneumonitis: New or worsening pulmonary symptoms; AND Prescriber attests that the patient will not take Lumakras with: Acid-reducing agents (e.g., proton pump inhibitors, H ₂ receptor antagonists, antacids.) Strong CYP3A4 inducers (e.g., rifampin, carbamazepine)		General PA Form
		Enzyme Inhibitors: MET	T	
Tabrecta®	P	Initial Criteria: Patient must have metastatic non-small cell lung cancer (NSCLC); AND Prescribed by, or in consultation with, an oncologist; AND Patient must have a mutation that leads to mesenchymal-epithelial transition (MET) exon 14 skipping in tumor specimens as confirmed by an FDA-approved test; AND Renewal Criteria: Patient continues to meet the initial criteria; AND Patient has positive disease response defined as disease stabilization or decrease in size of tumor or tumor spread; AND Patient does not have unacceptable toxicity (e.g., interstitial lung disease, liver enzymes outside of normal limits)	4/day	General PA Form



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Tepmetko®	NP	Initial Criteria: Diagnosis of metastatic non-small cell lung cancer (NSCLC); AND Disease is harboring mesenchymal-epithelial transition (MET) exon 14 skipping alterations; AND Prescribed by, or in consultation with, an oncologist Renewal Criteria: Patient continues to meet the initial criteria; AND Patient has positive disease response defined as disease stabilization or decrease in size of tumor or tumor spread; AND Patient does not have unacceptable toxicity (e.g. interstitial lung disease, liver enzymes outside of normal limits)	2/day	General PA Form
		Enzyme Inhibitors: MTOR		
Afinitor Disperz®	NP	Patient is unable to swallow solid dosage forms		General PA
everolimus soluble tabs	NP	Patient is unable to swallow solid dosage forms		Form Form
		Enzyme Inhibitors: PARP Inhibitors		
Lynparza®	Р		4/day	General PA
Rubraca®	Р		4/day	<u>Form</u>
Talzenna®	Р	Initial Criteria (6-month duration): One of the following: Diagnosis of HER2-negative locally advanced or metastatic breast cancer; AND Patient has a BRCA-positive mutated germline confirmed by an FDA-approved test (e.g., BRACAnalysis CDx); AND Patient must have received treatment with an anthracycline and/or a taxane (unless contraindicated) as neoadjuvant, adjuvant, and/or metastatic treatment; AND If patient received prior platinum-based chemotherapy, disease progression nor relapse were experienced within 6-months of receiving neoadjuvant or adjuvant platinum therapy; OR Diagnosis of metastatic castration-resistant prostate cancer; AND Patient has homologous recombination repair (HRR) gene mutation; AND Patient must use in combination with Xtandi; AND Patient has had a bilateral orchiectomy OR will receive a gonadotropin-releasing hormone (GnRH)-analog (e.g., leuprolide, goserelin, triptorelin); AND Provider will monitor complete blood counts at baseline and monthly thereafter; AND Patient does not have untreated CNS metastases (patient has completed definitive local therapy and may have stable CNS lesions on repeat brain imaging); AND Patient will not use requested agent in combination with any other PARP inhibitors; AND Patient has not received prior therapy with a PARP-inhibitor (e.g., Lynparza) Renewal Criteria (6-month duration): Patient continues to meet initial criteria; AND Tumor response has been demonstrated with either stabilization of disease or decrease in size of tumor or tumor spread; AND Absence of unacceptable toxicity from; AND Patient has not developed myelodysplastic syndrome (MDS)/acute myeloid leukemia (AML)	1/day	General PA Form



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
Zejula®	Р		3/day	General PA Form	
	•	Enzyme Inhibitors: RET		•	
Retevmo®	P	Initial Criteria: Patient must have ONE of the following diagnoses: Locally advanced or metastatic RET fusion-positive non-small cell lung cancer (NSCLC) Advanced or metastatic RET-mutant medullary thyroid cancer (MTC) who require systemic therapy Advanced or metastatic RET fusion-positive thyroid cancer who require systemic therapy and who are radioactive iodine-refractory Locally advanced or metastatic solid tumors with a RET gene fusion that have progressed on or following prior systemic treatment or who have no satisfactory alternative treatment options; AND Prescribed by, or in consultation with, an oncologist Renewal Criteria: Patient continues to meet the initial criteria; AND Patient has positive disease response defined as disease stabilization or decrease in size of tumor or tumor spread; AND Patient does not have unacceptable toxicity (e.g., severe or life-threatening hemorrhaging, uncontrolled blood pressure, interstitial lung disease, elevated liver enzymes, QT interval prolongation)	80mg: 4/day 40mg: 6/day	General PA Form	
Gavreto®	NP	Initial Criteria: Diagnosis of ONE of the following: Metastatic RET fusion-positive non-small cell lung cancer (NSCLC) that is detected by an FDA approved test Advanced or metastatic RET fusion-positive thyroid cancer who require systemic therapy and who are radioactive iodine-refractory (if radioactive iodine is appropriate); AND Requested agent will be prescribed by, or in consultation with, an oncologist Renewal Criteria: Patient continues to meet the initial criteria; AND Patient has positive disease response defined as disease stabilization or decrease in size of tumor or tumor spread; AND Patient does not have unacceptable toxicity (e.g., interstitial lung disease, elevated liver enzymes, severe or life-threatening hemorrhaging, uncontrolled blood pressure)	4/day	General PA Form	



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
	•	Enzyme Inhibitors: Tropomyosin Receptor Kinase (TRK)				
Augtyro®	NP	Initial Criteria: (6-month duration) Patient has diagnosis of ONE of the following: Diagnosis of locally advanced or metastatic Non-Small Cell Lung Cancer (NSCLC); AND Tumor is ROS1 rearrangement positive; OR AND NTRK Gene Fusion-Positive Solid Tumor and BOTH of the following: Disease is locally advanced or metastatic or where surgical resection is likely to result in severe morbidity Disease has progressed following treatment or there is no satisfactory alternative treatment; OR Diagnosis of secretory breast cancer or mammary analogue secretory cancer; AND Tumor is ROS1 rearrangement positive; AND Prescribed by, or in consultation with, an oncologist; AND For patients with reproductive potential, prescriber attest to all of the following: Patient is not pregnant prior to initiation of therapy Female patients have been advised to use effective contraception during treatment and for 2 months after the final dose Female patients have been advised to not breastfeed during treatment and 10 days after the final dose Male patients with female partners of reproductive potential have been advised to use effective contraception during treatment and for 4 months after the final dose Renewal Criteria: Patient must continue to meet the initial criteria; AND Patient has positive disease response defined as disease stabilization or decrease in size of tumor or tumor spread; AND Patient does not have unacceptable toxicity (e.g., hepatotoxicity, central nervous system effects, hyperuricemia, skeletal fractures, creatine phosphokinase elevation, interstitial lung disease/pneumonitis)	8/day	General PA Form		



ONCOLOGY AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. PDL Medication **Prior Authorization Criteria Qty. Limits PA Form** Initial Criteria (6-month duration): Patient meets ONE of the following disease specific criteria: Diagnosis of recurrent, advanced, or metastatic Non-Small Cell Lung Cancer (NSCLC); AND Disease is ROS1 positive as detected by an FDA-approved test NTRK Gene Fusion-Positive Solid Tumor; AND Presence of a neurotrophic tyrosine receptor kinase (NTRK) gene fusion as detected by an FDA-approved test, without a known acquired resistance mutation; AND Disease is metastatic or where surgical resection is likely to result in severe morbidity; AND Disease has progressed following treatment or there is no satisfactory alternative treatment; AND Prescribed by, or in consultation with, an oncologist; AND Patient does not have a history of prolonged QTc interval (e.g., QTc interval > 450 milliseconds); AND Patient will not use therapy in combination with drugs which prolong QT-interval; AND Patient will not use therapy with other NTRK-inhibitor therapy or ROS1-directed therapy; AND Patient does not have signs and symptoms of hyperuricemia as evidenced by a baseline serum; AND Patient will avoid concomitant use with moderate or strong CYP3A inducers or inhibitors; AND Provider attests to perform ALL the following: General PA 100 mg: 5/day; Rozlytrek® capsules Assess left ventricular ejection fraction (LVEF) prior to initiation of Rozlytrek in patients with symptoms or known risk factors 200 mg: 3/day Form for CHF Monitor liver tests, including ALT and AST, every 2 weeks during the first month of the patient's treatment, then monthly thereafter, and as clinically indicated Assess serum uric acid levels prior to initiation and periodically during treatment with Rozlytrek Assess QT interval and electrolytes at baseline and periodically during treatment patients who have or who are at risk for QTc interval prolongation Advise females of reproductive potential of the potential risk to a fetus and use of effective contraception during treatment and for 5 weeks following the final dose Advise males with female partners of reproductive potential to use effective contraception during treatment and for 3 months after the final dose Renewal Criteria (6-month duration): Patient must continue to meet the initial criteria; AND Documented disease response with treatment, as defined by stabilization of disease or decrease in size of tumor or tumor spread; AND There is absence of unacceptable toxicity from the drug (e.g., congestive heart failure, hepatotoxicity, central nervous system effects, hyperuricemia, QT-interval prolongation, visual disturbances) See Rozlytrek capsules prior authorization criteria; AND **General PA** Rozlytrek® pack 600mg/day Clinically valid reason why Rozlytrek capsules cannot be used Form



		ONCOLOGY AGENTS		
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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Vitrakvi®	NP	Initial Criteria: Patient has a solid tumor (e.g., soft tissue sarcoma, salivary gland, infantile fibrosarcoma, thyroid, lung, or gastrointestinal stromal tumors); AND Prescribed by, or in consultation with, an oncologist; AND Patient meets ALL the following: Presence of a neurotrophic tyrosine receptor kinase (NTRK) gene fusion without a known acquired resistance mutation Disease is metastatic or surgical resection is likely to result in severe morbidity Disease has progressed following treatment or there is no satisfactory alternative treatment; AND Provider attests to ALL the following: Monitor liver tests including ALT and AST every 2 weeks during the first month of treatment, then monthly thereafter and as clinically indicated Advise females with reproductive potential and males with female partners of reproductive potential to use effective contraception during treatment and for 1 week after the final dose Renewal Criteria: Patient continues to meet initial criteria; AND Patient has tumor response with stabilization of disease or decrease in size of tumor or tumor spread; AND Patient does not have unacceptable toxicity such as severe neurotoxicity, hepatotoxicity; (adverse effects resolve following dose recommendations/no permanent discontinuation required)	25 mg: 3/day; 100 mg: 2/day; 20 mg/mL: 10 mL/day	General PA Form
		Hormonal Agents: Aromatase Inhibitors		·
anastrozole	Р	For male patients, diagnosis of breast cancer For female patients, no PA required		General PA Form
		Hormonal Agents: Anti-Androgens Second Generation		
Akeega®	NP	 Initial Criteria (6-month duration) Diagnosis of metastatic castration-resistant prostate cancer (mCRPC); AND Patient has a deleterious or suspected deleterious BRCA-mutated (BRCAm) germline confirmed by an FDA approved test; AND Will be taken in combination with prednisone; AND ONE of the following: Medication will used in combination with androgen deprivation therapy (ADT) (e.g., leuprolide, goserelin, triptorelin, degarelix, relugolix) Patient has had a bilateral orchiectomy Renewal Criteria Patient continues to meet the initial criteria; AND Tumor response with stabilization of disease or decrease in size of tumor or tumor spread; AND Absence of unacceptable toxicity from the drug (e.g., hepatotoxicity, fractures, hypertension) 	2/day	General PA Form



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Erleada®	NP	Initial Criteria (6-month duration): Patient has diagnosis of ONE of the following: Non-metastatic castration-resistant disease prostate cancer (nmCRPC) Metastatic castration-sensitive disease prostate cancer (mCSPC); AND ONE of the following: Medication will used in combination with androgen deprivation therapy (ADT) (e.g., leuprolide, goserelin, triptorelin, degarelix, relugolix) Patient has had a bilateral orchiectomy Renewal Criteria: Patient continues to meet the initial criteria; AND Tumor response with stabilization of disease or decrease in size of tumor or tumor spread; AND Absence of unacceptable toxicity from the drug. Examples of unacceptable toxicity include seizures, excessive falls and/or fractures and any other Grade 3 or above side effects that are intolerable to patient, etc.	4/day	General PA Form
Nubeqa®	NP	Initial Criteria (6-month duration): Diagnosis of ONE of the following: Non-metastatic castration-resistant prostate cancer (nmCRPC); AND Metastatic hormone-sensitive prostate cancer (mHSPC); AND ONE of the following: Medication will used in combination with androgen deprivation therapy (ADT) (e.g., leuprolide, goserelin, triptorelin, degarelix, relugolix); OR Patient has had a bilateral orchiectomy Renewal Criteria: Patient continues to meet the initial criteria; AND Tumor response with stabilization of disease or decrease in size of tumor or tumor spread; AND Absence of unacceptable toxicity from the drug. Examples of unacceptable toxicity include elevated hepatic enzymes, hyperbilirubinemia, neutropenia, or any other Grade 3 or above side effects that are intolerable to patient, etc.	4/day	General PA Form
Xtandi® tablets	NP	Documented allergy or contraindication to an inactive ingredient in the capsules that is NOT in the tablets		General PA Form
Yonsa®	NP	Initial Criteria (6-month duration): Patient has metastatic castration-resistant prostate cancer (mCRPC); AND Will be taken in combination with methylprednisolone; AND ONE of the following: Medication will used in combination with androgen deprivation therapy (ADT) (e.g., leuprolide, goserelin, triptorelin, degarelix, relugolix) Patient has a bilateral orchiectomy; AND Renewal Criteria: Patient continues to meet the initial criteria; AND Tumor response with stabilization of disease or decrease in size of tumor or tumor spread Absence of unacceptable toxicity from the drug (e.g., elevated hepatic enzymes, hypokalemia, fluid retention, hypertension)		General PA Form



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Hormonal Agents: GnRH Agonists & LNRH Analogs		
Eligard®	Р	Diagnosis of prostate cancer in male patient		General PA Form
leuprolide	Р	 Leuprolide will be approved for patients meeting ONE of the following criteria: Diagnosis of prostate cancer in male patient Diagnosis of central precocious puberty in children (onset of secondary sexual development before 8 [girls] or 9 years of age [boys]) 		General PA Form
Lupron Depot®	NP	 Will be approved for self-administering patients with ONE of the following: Diagnosis of prostate cancer in male patient Diagnosis of endometriosis in female patient Diagnosis of uterine leiomyomas in female patient Diagnosis of recurrent ovarian carcinoma 		General PA Form
Orgovyx®	NP	 Diagnosis of advanced prostate cancer in male patient; AND Male patients with female partners of reproductive potential have been advised to use effective contraception during treatment and for two weeks after the last dose; AND Patient will not take requested medication with ANY of the following: P-GP Inhibitors Strong CYP3A Inducers cisapride pimozide thioridazine 	30/month (32 tablets for initial month of therapy)	General PA Form
		Hormonal Agents: SERM/SERD		
Orserdu®	NP	Initial Criteria (6-month duration): Patient has hormone receptor-positive, HER2-negative advanced breast; AND Patient has received at least one endocrine based regimen; AND Patient has ESR1 mutation detected by FDA-approved test; AND If female, patient is postmenopausal; AND Orserdu will be used as monotherapy; AND Prescribed by, or in consultation with, an oncologist; AND Patient must not be pregnant or breastfeeding; AND Females of reproductive potential and males undergoing treatment with female partners of reproductive age should be advised to use effective contraception during treatment and for 1 week after the final dose Renewal Criteria: Patient continues to meet initial criteria; AND Patient has positive disease response defined as disease stabilization or decrease in size of tumor or tumor spread; AND Patient does not have unacceptable toxicity (e.g., dyslipidemia, musculoskeletal pain)	345 mg: 1/day 86 mg: 3/day	General PA Form



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
	•	Multikinase Inhibitors: Renal and Thyroid Cancers		
Fotivda®	NP	Initial Criteria (6-month duration): Patient has diagnosis of relapsed or refractory advanced renal cell carcinoma (RCC); AND Patient has had two or more prior systemic therapies [two kinase inhibitors (KIs), a KI plus an immune checkpoint inhibitor, or a KI plus other systemic agents]; AND Prescriber attests to ALL the following: Patient will be closely monitored due to increased risk of Arterial and venous Thromboembolic Events, Hemorrhagic Events, Proteinuria, and Thyroid Dysfunction Fotivda will be withheld for at least 24 days before elective surgery and will not administer for at least 2 weeks following major surgery and adequate wound healing Patient's baseline liver function tests will be assessed Female and male patients of reproductive potential have been advised to use effective contraception during treatment and for one month after the last dose Agent will not be co-administered with strong CYP3A inducers Patient does not have a history of allergic reactions to tartrazine (only applies to requests for Fotivda 0.89 mg) Female patients are not pregnant or breastfeeding; AND Will not use in patients with any of the following: Strong CYP3A inducers History of allergic reactions to tartrazine Renewal Criteria: Patient continues to meet initial criteria; AND Prescriber attests to positive response to therapy indicated by tumor response with stabilization of disease OR decrease in size of tumor or tumor spread; AND Patient has absence of unacceptable toxicity from the drug (e.g., uncontrolled hypertension, onset of cardiac failure, arterial and venous Thromboembolic Events, hemorrhagic events, proteinuria, thyroid dysfunction, onset of Reversible Posterior Leukoencephalopathy Syndrome (RPLS), or increased LFT's)	21/28 days	General PA Form
lenalidomide	NP	Clinically valid reason why Revlimid cannot be used		General PA



	ONCOLOGY AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
Xpovio®	NP	Initial Criteria (6-month duration): One of the following: Diagnosis of multiple myeloma; AND Patient has received at least one prior therapy Diagnosis of diffuse large B-cell lymphoma (DLBCL), not otherwise specified, including DLBCL arising from follicular lymphoma; AND Patient has relapsed or refractory disease; AND Patient has received at least 2 lines of systemic therapy; AND Prescribed by, or in consultation with, an oncologist Renewal Criteria: Patient continues to meet initial criteria; AND Prescriber attests that the patient has experienced lack of disease progression, and/or improvement in symptoms; AND Patient has absence of unacceptable toxicity from the drug (e.g., thrombocytopenia, neutropenia, gastrointestinal toxicity, hyponatremia, neurological toxicity)	4 packs/month	General PA Form		
	•	Myelofibrosis		•		
Jakafi®	Р		2/day	General PA Form		
Besremi®	NP	 Diagnosis of polycythemia vera; AND Prescribed by, or in consultation with, an oncologist or hematologist; AND Patient does not have ANY of the following: Severe, acute, or unstable cardiovascular disease Existence of, or history of severe psychiatric disorders, particularly severe depression, suicidal ideation, or suicide attempt Hypersensitivity to interferon or to any component of BESREMI Hepatic impairment (Child-Pugh B or C) History or presence of active serious or untreated autoimmune disease; AND Patient is not an immunosuppressed transplant recipient; AND For women of childbearing age, provider has confirmed that the patient is not pregnant prior to receiving treatment; AND Patients of reproductive potential will be counseled to use effective contraception during treatment and for at least 8 weeks after the final dose 		General PA Form		
Inrebic®	NP	 Initial Criteria: Patient has a diagnosis of primary or secondary (post-polycythemia vera or post-essential thrombocythemia) myelofibrosis; AND Patient is considered intermediate-2 risk or high-risk; AND Patient's platelet count ≥ 50 x 109/L; AND Provider attests patient is not thiamine deficient (vitamin B1) and will monitor thiamine level during treatment Renewal Criteria: Patient has positive clinical response to therapy (e.g., reduction in symptoms, decreased spleen size); AND	4/day	General PA Form		



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Ojjaara®	NP	 Initial Criteria: Patient has a diagnosis of primary or secondary (post-polycythemia vera or post-essential thrombocythemia) myelofibrosis; AND Patient is considered intermediate-1, intermediate-2, or high-risk; AND Patient is anemic (e.g., hemoglobin (Hb) < 10 g/dL and/or hematocrit (Hct) < 30%) Renewal Criteria: Patient has positive clinical response to therapy (e.g., reduction in symptoms, decreased spleen size, decreased number of transfusion); AND Absence of unacceptable toxicity (e.g., thrombocytopenia, neutropenia, hepatotoxicity, major adverse cardiovascular events, thrombosis, and malignancies) 	1/day	General PA Form
Vonjo®	NP	 Initial Criteria Patient has a diagnosis of primary or secondary (post-polycythemia vera or post-essential thrombocythemia) myelofibrosis; AND Patient is considered intermediate risk or high-risk; AND Platelet count is below 50 x 10⁹/L Renewal Criteria: Patient has positive clinical response to therapy (e.g., reduction in symptoms, decreased spleen size, decreased number of transfusions); AND Absence of unacceptable toxicity (e.g., thrombocytopenia, major adverse cardiovascular events, thrombosis, malignancies) 	4/day	General PA Form
	u .	PI3K Inhibitors		I
Itovebi®	NP	Initial Criteria Patient has hormone receptor-positive, HER2-negative locally advanced or metastatic breast cancer; AND Patient has a PIK3CA mutation as detected by an FDA-approved test; AND Patient has experienced disease recurrence on or after completing adjuvant endocrine therapy; AND Agent is being given in combination with Ibrance and fulvestrant; AND Agent is prescribed by, or in consultation with, an oncologist Renewal criteria Patient continues to meet initial criteria; AND Patient has clinical response defined as disease stabilization or decrease in size or spread of tumor; AND Patient does not have unacceptable toxicity (e.g., hyperglycemia, diarrhea, stomatitis)	3mg: 2/day 9mg: 1/day	General PA Form



		ONCOLOGY AGENTS		
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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Piqray®	NP	Initial Criteria: Patient has hormone receptor-positive, HER2-negative advanced breast cancer; AND Agent is prescribed by, or in consultation with, an oncologist; AND Patient has experienced disease progression on after an endocrine based regimen for advanced disease OR has relapsed disease within 12 months after completion of adjuvant endocrine therapy; AND Patient has a PIK3CA-mutation as detected by an FDA-approved test; AND Piqray® will be given in combination with fulvestrant; AND Renewal Criteria: Patient continues to meet initial criteria; AND Patient has tumor response with stabilization of disease or decrease in the size of tumor or tumor spread; AND Patient does not have unacceptable toxicity such as severe cutaneous reaction or pneumonitis		General PA Form
Truqap®	NP	 Initial Criteria Patient has hormone receptor-positive, HER2-negative locally advanced or metastatic breast cancer; AND Patient has one or more PIK3CA/AKT1/PTEN-alterations as detected by an FDA-approved test; AND Patient has experienced disease progression on after an endocrine based regimen for advanced disease OR has relapsed disease within 12 months after completion of adjuvant endocrine therapy; AND Agent is being given in combination with fulvestrant; AND Agent is prescribed by, or in consultation with, an oncologist Renewal criteria Patient continues to meet initial criteria; AND Patient has clinical response defined as disease stabilization or decrease in size of tumor or tumor spread; AND Patient does not have unacceptable toxicity (e.g., hyperglycemia, diarrhea, cutaneous adverse reactions) 	64/28 days	General PA Form
		Rare/Miscellaneous Oncology Conditions		
Ayvakit®	NP	Initial Criteria: Diagnosis of ONE of the following: Unresectable or metastatic gastrointestinal stromal tumors (GIST) with platelet-derived growth factor-alpha (PDGFRA) exon 18 mutation, including PDGFRA D842V mutations Indolent systemic mastocytosis (ISM) Advanced systemic mastocytosis (AdvSM) Note: Includes aggressive systemic mastocytosis (ASM), systemic mastocytosis with an associated hematological neoplasm (SM-AHN), and mast cell leukemia (MCL); AND Prescribed by, or in consultation with, an oncologist; AND Renewal Criteria: Patient continues to meet initial criteria; AND Patient does not show evidence of progressive disease or unacceptable toxicity (e.g., intracranial hemorrhage, cognitive dysfunction)	1/day	General PA Form



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
lwilfin®	NP	Initial Criteria: Diagnosis of high-risk neuroblastoma (HRNB); AND Patient had a partial response to prior multiagent, multimodality therapy; AND Patient has received anti-GD2 immunotherapy (e.g., dinutuximab); AND Prescribed by, or in consultation with, an oncologist Renewal Criteria: Patient continues to meet initial criteria; AND Patient does not show evidence of progressive disease or unacceptable toxicity (e.g., hepatotoxicity, hearing loss)		General PA Form			
Ogsiveo®	NP	 Initial Criteria: Diagnosis of progressing desmoid tumor (also known as aggressive fibromatosis); AND Prescriber has reviewed and evaluated appropriate treatment options and attests that the patient requires systemic therapy; AND Prescribed by, or in consultation with, an oncology, hematology, or gastroenterology specialist Renewal Criteria: Patient demonstrates disease stabilization or clinical response to therapy (e.g., decrease tumor size, decreased pain, improved physical function, increased quality of life) 	6/day	General PA Form			
Qinlock®	NP	 Initial Criteria: Diagnosis of unresectable, locally advanced, or metastatic gastrointestinal stromal tumor (GIST); AND Prescribed by, or in consultation with, an oncologist; AND Patient has been previously treated with at least THREE kinase systemic therapies (e.g., imatinib, avapritinib, sunitinib, regorafenib) Renewal Criteria: Patient continues to meet initial criteria; AND Patient does not show evidence of progressive disease and unacceptable toxicity (e.g., uncontrolled hypertension, cardiac dysfunction) 	3/day	General PA Form			
Rezurock®	NP	 Initial Criteria (6-month duration): Patient has diagnosis of Chronic Graft-Versus-Host Disease; AND Patient is 12 years of age or older; AND Patient has a history of allogenic hematopoietic cell transplant (HCT); AND Agent is prescribed by, or in consultation with, an oncologist, hematologist, or bone marrow transplant specialist; AND Patient has had a previous failure of at least one systemic corticosteroid therapy (i.e., methylprednisolone, prednisone, etc.); AND Patient has had a previous failure of at least one non-steroidal systemic immunosuppressant therapy (e.g., abatacept, alemtuzumab, calcineurin inhibitor, etanercept, hydroxychloroquine, ibrutinib, imatinib, interleukin-2, low-dose methotrexate, mTOR inhibitor, mycophenolate mofetil, pentostatin, rituximab, ruxolitinib, etc.); AND Prescriber attests, if applicable, that patient will be advised that effective contraception should be used during treatment and for at least one week after last dose Renewal Criteria: Patient continues to meet the initial criteria; AND Patient is responding positively to treatment 	1/day	General PA Form			



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form			
Romvimza®	NP	 Initial Criteria Diagnosis of symptomatic tenosynovial giant cell tumor (TGCT); AND Patient has severe morbidity or functional limitations (e.g., worst stiffness numeric rating scale [NRS] of > 4); AND Patient is not a candidate for surgical resection associated with potential worsening, functional limitation, or severe morbidity; AND Prescribed by, or in consultation with, a hematologist or oncologist Renewal Criteria Patient continues to meet initial criteria; AND Patient does not show evidence of progressive disease and unacceptable toxicity (e.g., hepatotoxicity) 					
Tazverik [®]	NP	Initial Criteria: Diagnosis of ONE of the following: Metastatic or locally advanced epithelioid sarcoma; AND Patient not eligible for complete resection Relapsed or refractory follicular lymphoma; AND Tumor is positive for an EZH2 mutation as detected by an FDA approved test; AND Patient has received ≥ 2 prior systemic therapies OR has not had satisfactory alternative treatment option; AND Prescribed by, or in consultation with, an oncologist; AND Renewal Criteria: Patient continues to meet initial criteria; AND Patient does not show evidence of progressive disease and unacceptable toxicity (e.g. secondary malignancy)	8/day	General PA Form			
Turalio®	NP	 Initial Criteria: Diagnosis of symptomatic tenosynovial giant cell tumor (TGCT); AND Patient has severe morbidity or functional limitations (e.g., worst stiffness numeric rating scale [NRS] of ≥ 4); AND Patient is not a candidate for surgical resection associated with potential worsening, functional limitation, or severe morbidity; AND Prescribed by, or in consultation with, a hematologist or oncologist; AND Prescriber is enrolled in the Turalio REMS Program Renewal Criteria: Patient continues to meet initial criteria; AND Patient does not show evidence of progressive disease and unacceptable toxicity (e.g., hepatotoxicity) 	4/day	General PA Form			



Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Forn
Welireg®	NP	Initial Criteria: ONE of the following: Diagnosis of Von Hippel-Lindau (VHL) disease and require therapy for ONE of the following VHL-associated cancers, not requiring immediate surgery: Renal cell carcinoma (RCC) Central nervous system (CNS) hemangioblastomas Pancreatic neuroendocrine tumors (pNET); AND Diagnosis of advanced renal cell carcinoma (RCC) and patient has tried and failed, contraindication, or intolerance to ALL of the following: Programmed death receptor-1 (PD-1) or programmed death-ligand 1 (PD-L1) inhibitor (e.g., nivolumab, avelumab, pembrolizumab) Vascular endothelial growth factor tyrosine kinase inhibitor (VEGF-TKI) (e.g., Cabometyx, Inlyta, Lenvima, Nexavar, Sutent); AND Prescribed by, or in consultation with, a hematologist or oncologist; AND Patient is not pregnant or breastfeeding; AND Females of reproductive potential and males with female partners of reproductive potential should be advised to use effective non-hormonal contraception during treatment and for 1 week after the last dose Renewal Criteria: Patient continues to meet initial criteria; AND Patient does not show evidence of progressive disease and unacceptable toxicity (e.g., anemia, hypoxia)	3/day	General I Form



	OPHTHALMICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.				
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
		Dry Eye Disease Agents	,	•	
Lacrisert	Р		60 inserts/30 days		
Restasis®	Р	 Treatment of vernal keratoconjunctivitis (VKC) (i.e., severe atopic keratoconjunctivitis); OR Diagnosis of dry eye disease [i.e., dry eye syndrome, keratoconjunctivitis sicca (KCS)] 	60 vials/30 days	General PA Form	
Xiidra®	Р	 Diagnosis of dry eye disease [i.e., dry eye syndrome, keratoconjunctivitis sicca (KCS)]; AND Trial and failure or contraindication to Restasis® (trial duration ≥ 12 weeks confirmed by paid claims) 	2 vials/day	FOIIII	
Cequa®	NP	 Diagnosis of dry eye disease [i.e., dry eye syndrome, keratoconjunctivitis sicca (KCS)]; AND Trial and failure, or contraindication, to both the following: Restasis® (trial duration > 12 weeks confirmed by paid claims) Xiidra® (trial duration > 12 weeks confirmed by paid claims) 	2 vials/day	General PA Form	
cyclosporine emulsion 0.05%	NP	 One of the following: Treatment of vernal keratoconjunctivitis (VKC) (i.e., severe atopic keratoconjunctivitis) Diagnosis of dry eye disease [i.e., dry eye syndrome, keratoconjunctivitis sicca (KCS)]; AND Clinically valid reason why the preferred Restasis® cannot be used 	60 vials/30 days	General PA Form	
Meibo®	NP	See Cequa® prior authorization criteria	3 bottles/30 days	General PA	
Restasis Multidose®		See cyclosporine emulsion 0.05% prior authorization criteria	1 bottle/30 days	<u>Form</u>	
Tyrvaya®	NP	See Cequa® prior authorization criteria		General PA	
Vevye®	NP	See Cequa® prior authorization criteria	3 bottles/30 days	<u>Form</u>	
		Ophthalmic Alpha-2 Agonists			
apraclonidine	Р		1 package/Rx		
brimonidine 0.2%	Р		1 package/Rx		
Alphagan P®	Р		1 package/Rx	General PA	
brimonidine 0.1%, 0.15%	NP		1 package/Rx	<u>Form</u>	
Iopidine®	NP		1 package/Rx		
		Ophthalmic Antibiotics	·		
ciprofloxacin	Р		10 mL/Rx		
erythromycin	Р		1 package/Rx	General PA	
moxifloxacin	Р		1 package/Rx	<u>Form</u>	
neomycin/bac/poly B	Р		1 package/Rx		
neomycin/poly B/gramicidin	Р		1 package/Rx		
polymyxin B/TMP	Р		1 package/Rx		
sulfacetamide soln	Р		1 package/Rx		
tobramycin	Р		1 package/Rx		
AzaSite®	NP		1 package/Rx		



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
Besivance®	NP		1 package/Rx		
Ciloxan®	NP		10 mL/Rx	General PA	
gentamicin	NP		15 mL/Rx	<u>Form</u>	
gatifloxacin 0.5% soln	NP		1 package/Rx	Cananal DA	
levofloxacin 0.5% soln	NP		1 package/Rx	General PA Form	
moxifloxacin (3X Day)	NP		1 package/Rx	<u>FOITH</u>	
sulfacetamide oint	NP		1 package/Rx	General PA	
Tobrex®	NP		1 package/Rx	<u>Form</u>	
	I	Ophthalmic Antibiotic/Steroid Combos	1		
neomycin/BAC/poly B/HC	Р		1 package/Rx	General PA	
sulfacetamide/ prednisolone	Р		1 package/Rx	<u>Form</u>	
tobramycin/ dexamethasone	Р		1 package/Rx	General PA	
Maxitrol®	NP		1 package/Rx	Form	
neomycin/poly B/HC	NP		1 package/Rx		
TobraDex®	NP		1 package/Rx		
TobraDex ST®	NP		1 package/Rx	General PA	
Zylet®	NP	 Trial and failure, contraindication, or intolerance of TWO preferred agents; OR There is concern over a potential increase in intra-ocular pressure (IOP) with other steroids (i.e., glaucoma, recipient is pre- or post-cataract surgery and a known steroid-responder) 	1 package/Rx	Form Form	
		Ophthalmic Antifungals			
Natacyn®	NP	Diagnosis of ophthalmic fungal infection	1 package/Rx	General PA Form	
		Ophthalmic Antivirals			
trifluridine	Р		1 package/Rx	General PA	
Zirgan®	Р		1 package/Rx	<u>Form</u>	
		Ophthalmic Anti-Allergics			
azelastine	Р		6 mL/Rx		
Bepreve®	Р		10 mL/Rx	1	
cromolyn sodium	Р		1 package/Rx	General PA	
ketotifen	Р		10 mL/Rx	<u>Form</u>	
olopatadine	Р		5 mL/Rx]	
Alocril®	NP		1 package/Rx	General PA	



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Alomide®	NP			<u>Form</u>
epinastine	NP		5 mL/Rx	
Lastacaft®	NP		3 mL/Rx	
Pataday®	NP		5 mL/Rx	
Verkazia [®]	NP	Initial Criteria (6-month duration): Diagnosis of moderate to severe vernal keratoconjunctivitis; AND Trial and failure, contraindication, or intolerance of one agent in ALL the following categories: Ophthalmic antihistamines (e.g., azelastine, olopatadine) Ophthalmic mast cell stabilizers (e.g., cromolyn sodium) Ophthalmic corticosteroids (e.g., dexamethasone, prednisolone, fluorometholone) Renewal Criteria: Patient demonstrates positive clinical response to therapy as evidenced by an improvement in clinical signs and symptoms (e.g., itching, photophobia, papillary hypertrophy, mucus discharge, conjunctival hyperaemia)	120/30 days	General PA Form
Zerviate®	NP	Clinically valid reason as to why patient cannot use a preferred ophthalmic antihistamine product	30 vials/Rx	
		Ophthalmic Beta Blockers		
carteolol	Р		1 package/Rx	
timolol maleate	Р		1 package/Rx	General PA
Betaxolol	NP		1 package/Rx	<u>Form</u>
Betoptic-S®	NP		1 package/Rx	1
Istalol®	NP		1 package/Rx	
levobunolol	NP		1 package/Rx	General PA
timolol gel solution	NP		1 package/Rx	<u>Form</u>
Timoptic Ocudose®	NP		1 package/Rx	
		Ophthalmic Carbonic Anhydrase Inhibitors		
Azopt®	Р		15 mL/30 days	General PA
dorzolamide	Р		10 mL/30 days	<u>Form</u>
dorzolamide/timolol	Р		10 mL/30 days	General PA
brinzolamide	NP		15 mL/30 days	Form
Cosopt®	NP		10 mL/30 days	General PA
Cosopt PF®	NP		2 vials/day	<u>Form</u>
		Ophthalmic Kinase Inhibitors		
Rhopressa®	Р	Patient has a diagnosis of ocular hypertension or open-angle glaucoma; AND • Patient has tried/failed or is intolerant to BOTH a prostaglandin inhibitor AND beta-adrenergic antagonist	5 ml/30 days	General PA
Rocklatan®	Р	See Rhopressa® prior authorization criteria	5 ml/Rx	<u>Form</u>



		OPHTHALMICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indic	ated.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Glaucoma Combinations	<u> </u>	
Combigan®	Р	 Patient is on simultaneous therapy with brimonidine and timolol for at least 60 days; AND Patient demonstrates non-compliance with 2 products individually. 	1 package/Rx	
Simbrinza®	Р	Patient is on simultaneous therapy with brimonidine and Azopt® for at least 60 days	1 package/Rx	General PA
brimonidine/timolol	NP	 Patient is on simultaneous therapy with brimonidine and timolol for at least 60 days; AND Trial and failure, contraindication, or intolerance of Combigan. 	1 package/Rx	<u>Form</u>
	•	Miotics		
phospholine iodide	NP		1 package/Rx	
Vuity®	NP	 Diagnosis of presbyopia; AND Patient is 18 years of age or older; AND Patient is not a candidate for surgery or surgery was non-curative; AND Clinically valid reason as to why the preferred pilocarpine cannot be used 	2.5 mL/30 days	General PA Form
	1	Miscellaneous Ophthalmics		•
Cystaran®	NP	Diagnosis of cystinosis	1 package/Rx	
Cystadrops®	NP	Patient is being treated for Corneal cystine crystal deposits with cystinosis; AND • Prescriber must provide a clinically valid reason as to why Cystaran cannot be used	1 package/Rx	General PA Form
Oxervate®	NP	 Patient must be ≥ 2 years of age; AND Patient must have a diagnosis of moderate to severe (stage 2 or stage 3) neurotrophic keratitis (NK); AND Prescribed by, or in consultation with, an ophthalmologist; AND Prescriber attests that patient or caregiver has been counseled on proper administration technique 	2 ml/day (lifetime therapy QL=112 ml for 8 weeks of therapy)	General PA Form
Xdemvy®	NP	Criteria: (2-month duration) Diagnosis of Demodex blepharitis; AND Patient has collarettes, cylindrical deposits at the base of eyelashes, confirmed by slit lamp examination; AND Prescribed by or in consultation with an ophthalmologist or optometrist	1 bottle (10 ml)/ 50 days	General PA Form
		Ophthalmic NSAIDs Approval of NP agents requires trial and failure, contraindication, or intolerance of ONE preferred agent		
diclofenac	Р		1 package/Rx	
flurbiprofen	Р		1 package/Rx	Ophthalmic
ketorolac	Р		1 package/Rx	NSAIDs PA
Acular LS®	NP		1 package/Rx	<u>Form</u>
Acuvail®	NP		1 package/Rx]
BromSite®	NP		1 package/Rx	
bromfenac	NP		1 package/Rx	Ophthalmic
Ilevro®	NP		1 package/Rx	NSAIDs PA
Nevanac®	NP		1 package/Rx	<u>Form</u>
Prolensa®	NP		1 package/Rx	1



		OPHTHALMICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indica	ated.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Ophthalmic Prostaglandin Agonists		
latanoprost	Р		5 mL/Rx	
Lumigan®	Р		5 mL/Rx	General PA Form
Travatan Z®	Р		5 mL/Rx	FOITH
Zioptan®	Р		1 container/day	
bimatoprost	NP		5 mL/ Rx	General PA
tafluprost	NP		1 container/day	<u>Form</u>
travoprost	NP	Clinically valid reason why preferred Travatan Z® cannot be used	5 mL/ Rx	1
lyuzeh®	NP	Clinically valid reason why preferred Travatan Z® cannot be used	1 container/day	
Vyzulta®	NP		5 mL/ Rx	General PA
Xalatan®	NP		5 mL/ Rx	<u>Form</u>
Xelpros®	NP		5 mL/ Rx	1
		Ophthalmic Steroids	·	•
Alrex®	Р		1 package/Rx	
difluprednate	Р		1 package/Rx]
fluorometholone	Р		1 package/Rx	General PA Form
Lotemax® suspension	Р		1 package/Rx	<u>FOITH</u>
Pred Mild®	Р		1 package/Rx	1
prednisolone acetate	Р		1 package/Rx	
dexamethasone	NP		1 package/Rx	1
Durezol®	NP		1 package/Rx	General PA
Eysuvis®	NP	Patient is being treated for symptoms of Dry Eye disease; AND Patient has had a trial and failure of Restasis; AND Patient has had a trial and failure of a preferred loteprednol product (e.g., Alrex, Lotemax suspension)	1 package/Rx	<u>Form</u>
Flarex®	NP		1 package/Rx	
FML Forte®	NP		1 package/Rx	
FML Liquifilm®	NP		1 package/Rx	General PA
Lotemax SM® gel	NP		1 package/Rx	<u>Form</u>
Lotemax ointment	NP		1 package/Rx	1
loteprednol gel	NP		1 package/Rx	
loteprednol suspension	NP		15 ml/Rx	General PA
Maxidex®	NP		1 package/Rx	<u>Form</u>
prednisolone sodium phosphate	NP		1 package/Rx	



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
Pred Forte®	NP		1 package/Rx		
	Ophthalmic Vasoconstrictors				
phenylephrine	Р			General PA	
p,				<u>Form</u>	

		OTICS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.			
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
		Otic Quinolones			
ciprofloxacin otic	Р		14 mL/Rx	General PA	
ofloxacin otic	Р		10 mL/Rx	<u>Form</u>	
		Otic Steroid/Antibiotic Combinations			
HC/neomycin/ polymyxin B	Р		1 package/Rx		
ciprofloxacin- dexamethasone	Р		7.5 mL/Rx	General PA Form	
Cipro® HC	NP		10 mL/Rx		
	Miscellaneous Otics				
acetic acid/HC	Р		10 mL/Rx	General PA	
DermOtic®	Р		20 mL/Rx	<u>Form</u>	



		RARE CONDITIONS		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Activated PI3K Delta Syndrome (APDS)		
Joenja®	NP	 Initial Criteria (6-month duration): Patient is ≥ 12 years of age and weighs ≥ 45kg; AND Diagnosis of activated phosphoinositide 3-kinase delta syndrome (APDS); AND Diagnosis has been confirmed by the presence of an APDS-associated genetic variant in either PIK3CD or PIK3R1; AND Documentation of clinical findings and manifestations consistent with APDS (e.g., recurrent respiratory tract infections and viral infections, lymphadenopathy, hepatosplenomegaly, autoimmune cytopenia); AND Prescribed by, or in consultation with, hematologist, allergist, or immunologist; AND For patients with reproductive potential, the prescriber attests to all of the following: Patient is not pregnant prior to initiation of therapy Patient has been counseled on potential risk during pregnancy Patient has been advised to use effective contraception during treatment and for 1 week after the last dose Patient has been advised to not breastfeed during treatment and for 1 week after the last dose Renewal Criteria: Disease response to therapy and tolerability compared to baseline (e.g., decreased lymph node size, increased functional B cell counts, decreased infections/hospitalizations, and decreased utilization of immunoglobulin replacement therapy) 	2/day	General PA Form
		Amyotrophic Lateral Sclerosis (ALS)		
Radicava ORS®	NP	 Initial Criteria (6-month duration): Submission of medical records (e.g., chart notes, diagnostic tests, nerve conduction studies, lab values) to support a diagnosis of "definite" or "probable" ALS per the revised EL Escorial diagnostic criteria; AND Prescribed by, or in consultation with, a neurologist; AND Patient has scores ≥ 2 in all items of the ALS Functional Rating Scale-Revised (ALSFRS-R) criteria at the start of treatment; AND Patient has a forced vital capacity (%FVC) greater than or equal to 80% at the start of treatment Patient must not be pregnant Renewal Criteria (6-month duration): Prescribed by, or in consultation with, a neurologist; AND Documentation of positive clinical response to therapy (e.g., slowing in the decline of functional abilities); AND Patient is not dependent on invasive ventilation or tracheostomy 		General PA Form
Teglutik®	NP	 Diagnosis of Amyotrophic Lateral Sclerosis (ALS); AND Patient is unable to swallow tablets; AND 	20 mL/day	



		RARE CONDITIONS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Antineutrophil Cytoplasmic Autoantibody (ANCA)		
Tavneos®	NP	Initial criteria (6-month duration): Diagnosis of severe active anti-neutrophil cytoplasmic autoantibody ANCA-associated vasculitis (granulomatosis with polyangiitis [GPA] or microscopic polyangiitis [MPA]) confirmed by ONE of the following: ANCA test positive for proteinase 3 (PR3) antigen ANCA test positive for myeloperoxidase (MPO) antigen Tissue biopsy; AND Prescribed by, or in consultation with, a rheumatologist, nephrologist, pulmonologist, or a provider with expertise in vascular medicine; AND Will be used as adjunctive therapy with standard therapy (e.g., cyclophosphamide, azathioprine, mycophenolate, rituximab); AND Patient is concurrently on glucocorticoids or has an intolerance or contraindication to glucocorticoids Renewal Criteria: Patient continues to meet initial approval criteria; AND Disease response to therapy and tolerability compared to baseline	6 caps/day	General P/ Form
		CHAPLE Disease		
Veopoz®	NP	Initial Criteria: Diagnosis of CD55-deficient protein-losing enteropathy (CHAPLE disease); AND Patient has documentation of genetic testing confirming biallelic CD55 loss-of-function mutation; AND Prescriber attests to ALL of the following: Patient has received or will receive Veopoz IV loading dose; Patient has completed or updated meningococcal vaccination at least 2 weeks prior to administering the first dose of Veopoz unless the risk of delaying therapy outweighs the risk; AND Prescribed by or in consultation with a clinical specialist knowledgeable in appropriate disease management (e.g., geneticist, gastroenterologist, hematologist) Renewal Criteria: Patient has positive clinical response to therapy (e.g., normalization of serum albumin, decreased abdominal pain, diarrhea, facial edema, and peripheral edema)	8 vials/28 days	General PA Form



		RARE CONDITIONS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL		Qty. Limits	PA Form
	•	Duchenne Muscular Dystrophy (DMD)		•
Duvyzat®	NP	 Initial Criteria: Documentation of a confirmed diagnosis of Duchenne muscular dystrophy (DMD); AND Age ≥ 6 years; AND Patient retains meaningful voluntary motor function (e.g., patient can speak, manipulate objects using upper extremities, ambulate); AND Will be used in combination with a stable dose of corticosteroids (e.g. prednisone, Emflaza, Agamree); AND Prescribed by, or in consultation, with a neuromuscular specialist or medical geneticist Renewal Criteria: Patient retains meaningful voluntary motor function (e.g., patient can speak, manipulate objects using upper extremities, ambulate); AND Will be used in combination with a stable dose of corticosteroids (e.g. prednisone, Emflaza, Agamree); AND Patient has received benefit from therapy [e.g., stability or slowing in the decline of symptoms (motor function, respiratory function, musculature strength), quality of life] 	12 mL/day	General PA Form
Emflaza®	Р	 Initial Criteria: Documentation of a confirmed diagnosis of Duchenne muscular dystrophy (DMD); AND Age ≥ 2 years; AND Patient has experienced ≥ ONE of the following adverse reactions directly attributable to therapy with prednisone:		General PA Form
Agamree®	NP	See Emflaza prior authorization criteria; AND Trial and failure, contraindication, or intolerance to Emflaza	3 bottles (300mL)/ month	
deflazacort	NP	See Emflaza prior authorization criteria; AND Clinically valid reason why preferred Emflaza cannot be used		
		Fatty Acid Oxidation Disorder (FAOD)		
Dojolvi®	NP	Initial Criteria: Diagnosis of long-chain fatty acid oxidation disorders (LC-FAOD) as confirmed by two of the following: Acylcarnitine profile Molecular/genetic test Fibroblast test; AND Patient does not have pancreatic insufficiency; AND Prescribed by, or in consultation with, a clinical specialist knowledgeable in appropriate disease-related dietary management (e.g., geneticist, cardiologist, gastroenterologist, etc.); AND For patients receiving another medium-chain triglyceride product, discontinue prior to the first dose of Dojolvi® Renewal Criteria: Evidence of positive clinical response from baseline (e.g., reduction in signs/symptoms such as hypoglycemia, hepatopathy, skeletal myopathy, rhabdomyolysis, cardiomyopathy, etc.)		General PA Form



		RARE CONDITIONS		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Fibrodysplasia ossificans progressive (FOP)		
Sohonos®	NP	 Diagnosis of fibrodysplasia ossificans progressive (FOP); AND One of the following: Female aged ≥ 8 years of age Male aged ≥ 10 years of age; AND Diagnosis of FOP confirmed by one of the following: Mutation in the ALK2/ACVR1 gene Classic FOP clinical features such as malformation of big toe and progressive heterotopic endochondral ossification in ribbons, sheets, and plates Radiographic bone scans detecting heterotopic ossification (HO); AND Prescriber attests to all of the following: Patient is not pregnant Female patients of reproductive potential will be counseled to use effective contraception during treatment with therapy and for at least 1 month after last dose For pediatric patients, premature epiphyseal closure has not occurred 		General PA Form
		Friedreich's Ataxia		
Skyclarys®	NP	Initial Criteria • Patient is ≥ 16 years old; AND • Patient has diagnosis of Friedreich's ataxia (FA); AND • Patient has documentation of genetic testing confirming frataxin (FXN) gene mutation; AND • Prescribed by, or in consultation with, a neurologist, geneticist, or cardiologist Renewal Criteria • Patient has disease stabilization or clinical response to therapy	3/day	General PA Form
		Glucagon-Like Peptide-2 (GLP-2) Analog		
Gattex®	NP	 Initial Criteria: Diagnosis of short bowel syndrome, AND Patient is dependent on parenteral nutrition and/or fluids/electrolytes; AND Submission of medical records (e.g. chart notes) documenting that patient has been unable to significantly reduce PN/IV support Renewal Criteria: Submission of medical records (e.g. chart notes) demonstrating a positive response to therapy (e.g. decreased frequency or volume of parenteral nutrition and/or fluids/electrolytes from baseline) 		General PA Form



		RARE CONDITIONS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Hereditary Angioedema (HAE) Agents		•
Sajazir®	Р	 Prescribed by, or in consultation with, a specialist in allergy, immunology, hematology, pulmonology, or genetics; AND Patient is ≥ 18 years of age; AND Submission of medical records (e.g. chart notes and labs) documenting a diagnosis consistent with 1 of the following HAE subtypes: Type I:	6 injections/28 days	General PA Form
Takhzyro®	P	Initial Criteria: Prescribed by, or in consultation with, a specialist in allergy, immunology, hematology, pulmonology, or genetics; AND Patient must be ≥ the labeled age minimum (Haegarda ≥6 years; Orladeyo ≥12 years; Takhzyro ≥2 years); AND Submission of medical records (e.g. chart notes and labs) documenting a diagnosis consistent with 1 of the following HAE subtypes: Type I: Low C1 inhibitor (C1-INH) antigenic level (below the limit of normal defined by the lab performing the test); AND Low C1-INH functional level (below the limit of normal defined by the lab performing the test); OR Type II: Normal to elevated C1-INH antigenic level; AND Low C1-INH functional level (below the limit of normal defined by the lab performing the test); AND Type III: Normal C1-INH antigenic level; AND One of the following: Confirmed presence of a FXII, angiopoietin-1, plasminogen, KNG1, MYOF, or HS3ST6 gene mutation; OR Recurring angioedema attacks that are refractory to high-dose antihistamines with confirmed family history of angioedema; AND Will not be used in combination with other routine prophylaxis HAE agents (e.g., Haegarda, Takhzyro, Orladeyo); AND Prescriber attests patient is avoiding all possible triggers for HAE attacks Renewal Criteria: Improvement in severity and duration of attacks have been achieved and sustained; AND Absence of unacceptable toxicity from the drug (e.g., severe hypersensitivity reactions, thromboembolic events)	2 injections /28 days	General PA Form



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Firazyr®	NP	See Sajazir® prior authorization criteria; AND • Trial and failure, contraindication, or intolerance to 2 preferred agents	6 injections/28 days	
Haegarda®	NP	See Takhzyro® prior authorization criteria	2 injections/28 days	General PA
icatibant	NP	See Sajazir® prior authorization criteria; AND • Trial and failure, contraindication, or intolerance to 2 preferred agents	6 injections/28 days	<u>Form</u>
Orladeyo®	NP	See Takhzyro® prior authorization criteria	1/day	
	1	Homocystinuria Agents	•	,
Cystadane®	Р	 Diagnosis of moderate to severe hyperhomocysteinemia Genetic test confirming ONE of the following: cystathionine beta-synthase (CBS) deficiency 5,10-methylenetetrahydrofolate reductase (MTHRF) deficiency cobalamin cofactor metabolism (cbl) defect; AND; AND Prescribed by, or in consultation with, a physician who specializes in the treatment of inherited metabolic disorders; AND Patient had an inadequate response or is unable to be managed by diet and vitamin supplementation with folic acid, vitamin B12, and vitamin B6 	6 g/day	General PA Form
betaine anhydrous powder	NP	See Cystadane® prior authorization criteria; AND • Clinically valid reason why preferred Cystadane® cannot be used	6 g/day	
		Hutchinson-Gilford Progeria Syndrome		
Zokinvy [®]	NP	Initial Criteria (6-month duration): Patient has a diagnosis of Hutchinson-Gilford Progeria Syndrome; OR Patient has processing deficient Progeroid Laminopathies with either: Heterozygous LMNA mutation with progerin-like protein accumulation Homozygous or compound heterozygous ZMPSTE24 mutations; AND Patient must be 12 months of age or older; AND Patient must have a body surface area (BSA) of 0.39 m2 and above; AND Females must use effective contraception due to embryo-fetal toxicity; AND Patient must not meet any of the following: Other Progeroid Syndromes or processing proficient Progeroid Laminopathies Concomitant use of strong or moderate CYP3A inhibitors or inducers Concomitant use of midazolam Concomitant use of lovastatin, simvastatin, and atorvastatin Patient is pregnant Renewal Criteria: Patient continues to meet initial criteria; AND Patient has experienced a positive response to therapy, as documented by provider; AND Patient has not experienced treatment-limiting adverse effects (e.g., laboratory Abnormalities: changes in electrolytes, complete blood counts, and liver enzymes, decrease in renal function, retinal toxicity)		General PA Form



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Medication	PDL		Qty. Limits	PA Form
		Hyperoxaluria Agents		<u>'</u>
Rivfloza®	NP	 Initial Criteria: (6-month duration) Patient is 9 years of age or older; AND Diagnosis of primary hyperoxaluria type 1 (PH1) confirmed by ONE of the following: Genetic testing demonstrating mutation in the alanine-glyoxylate aminotransferase (AGXT) gene Liver biopsy demonstrating absent or reduced alanine-glyoxylate aminotransferase (AGT) activity; AND Patient has ONE of the following: Elevated urinary oxalate excretion Elevated plasma oxalate levels Urinary oxalate creatinine ratio above the age-specific upper limit of normal Patient has relatively preserved kidney function (e.g., eGFR ≥ 30 mL/min/1.73 m²); AND Prescribed by, or in consultation with, a hematologist, nephrologist, urologist, or geneticist Renewal Criteria: Patient has positive clinical response to therapy (e.g., decreased urinary oxalate excretion or plasma concentration, decreased number or size of kidney stones, improved kidney function) 	1/28 days	General PA Form
		Hypophosphatasia (HPP) Agents		
Strensiq®	NP	 Initial Criteria (6-month duration): Diagnosis of perinatal, infantile, or juvenile-onset hypophosphatasia (HPP); AND Onset of clinical signs and symptoms of HPP prior to age 19 years (e.g., rickets, skeletal deformities, fractures, respiratory compromise, vitamin B6 dependent seizure, craniosynostosis, dental abnormalities, severe osteopenia); AND Clinical diagnosis of HPP evidenced by one of the following: Serum alkaline phosphatase (ALP) below age-adjusted normal range Genetic confirmation of ALPL mutation; Elevated plasma pyridoxal 5'-phosphate (PLP) levels; AND Prescribed by, or in consultation with, a physician who specializes in the treatment of inherited metabolic disorders Note: 80 mg/0.8 mL vial will not be approved for pediatric patients weighing < 40 kg Renewal Criteria: Documentation of positive clinical response to therapy (e.g., healing of the skeletal manifestations, improved respiratory, motor function, and linear growth); AND Prescribed by, or in consultation with, a physician who specializes in the treatment of inherited metabolic disorders 		General PA Form



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		IBAT (Ileal Bile Acid Transporter) Inhibitors		•
Bylvay®	NP	 One of the following: Diagnosis of progressive familial intrahepatic cholestasis (PFIC); AND Patient does not have ABCB11 variant resulting in non-functional or complete absence of bile salt export pump protein (BSEP-3) Diagnosis of Alagille syndrome (ALGS) confirmed by presence of the JAG1 or Notch2 gene mutation; AND Prescribed by, or in consultation with, hepatologist or gastroenterologist; AND Patient is experiencing moderate to severe pruritus confirmed by ONE of the following: Total serum bile acid > 3x the upper limit of normal Conjugated bilirubin > 1 mg/dL. Fat soluble vitamin deficiency otherwise unexplainable. GGT > 3x the upper limit of normal Intractable pruritus explainable only by liver disease; AND Trial and failure to at TWO other conventional treatments for the symptomatic relief of pruritus (e.g., bile acid-binding agents, naltrexone, phenobarbital, rifampin, ursodiol); AND Provider attests to monitor the following: Liver-function tests at baseline and during treatment Fat-soluble vitamin (FSV) levels at baseline and during treatment 		General P Form
Livmarli®	NP	See Bylvay® prior authorization criteria		
		IgA Nephropathy (IgAN)		
Fabhalta®	NP	Initial Criteria (6-month duration) One of the following: Diagnosis of primary IgA immunoglobulin A nephropathy (IgAN) confirmed by kidney biopsy; AND Patient is at risk of rapid disease progression (e.g., proteinuria > 0.5 g/day or UPCR > 0.8 g/g); AND Patient has an eGFR > 20 mL/min/1.73 m2; OR Diagnosis of paroxysmal nocturnal hemoglobinuria (PNH) confirmed by flow cytometry; AND Patient has symptoms attributed to PNH (e.g., anemia, extreme fatigue, difficulty swallowing, recurrent abdominal pain, smooth muscle dystonia, thrombosis, CKD, organ damage secondary to chronic hemolysis); AND Prescribed by, or in consultation with, a hematologist, nephrologist, or oncologist; AND Prescriber is enrolled in the Fabhalta REMS Program; AND One of the following: Will not be used concurrently with another complement inhibitor; OR Patient is currently receiving another complement inhibitor (e.g., Empaveli, Soliris, Ultomiris) which will be discontinued and Fabhalta will be initiated in accordance with the FDA approved labeling Renewal Criteria: Submission of medical records (e.g., chart notes) documenting a positive clinical response to therapy (e.g., increased or stabilization of hemoglobin levels, reduction in transfusions, improvement in hemolysis, decrease in LDH); AND Prescribed by, or in consultation with, a hematologist or oncologist; AND Patient is not receiving Empaveli in combination with another complement inhibitor	2/day	



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Filspari®	NP	 Initial Criteria: (6-month duration) Patient is 18 years of age or older; AND Diagnosis of primary IgA immunoglobulin A nephropathy (IgAN) confirmed by kidney biopsy; AND Patient is at risk of rapid disease progression (e.g., proteinuria > 0.5 g/day or UPCR > 0.8 g/g); AND Patient has an eGFR > 30 mL/min/1.73 m2; AND Inadequate response to a stable max tolerated doses of an ARB or an ACE inhibitor; AND Use of renin-angiotensin-aldosterone system (RAAS) inhibitors (e.g., ACE inhibitors, ARBs), endothelin receptor antagonists (e.g., Letairis, Opsumit, Tracleer), and aliskiren will be discontinued prior to initiating treatment; AND Prescribed by, or in consultation with, a nephrologist; AND Prescriber and patient have met the requirements of Filspari REMS Program Renewal Criteria: Patient has positive clinical response to Filspari therapy (e.g., reduction of proteinuria from baseline, decreased UPCR) 	1/day	General PA Form
Tarpeyo®	NP	 Patient is 18 years of age or older; AND Diagnosis of primary IgA immunoglobulin A nephropathy (IgAN) confirmed by kidney biopsy; AND Patient is at risk of rapid disease progression (e.g., proteinuria > 0.5 g/day or UPCR > 0.8 g/g); AND Patient has an eGFR > 30 mL/min/1.73 m2; AND Inadequate response to a stable max tolerated doses of an ARB or an ACE inhibitor, unless contraindicated; AND Prescribed by, or in consultation with, a nephrologist 	4/day	General PA Form
Vanrafia®		See Filspari® prior authorization criteria	1/day	



		RARE CONDITIONS		
Medication	PDL	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Prior Authorization Criteria	Qty. Limits	PA Form
		IGF-1 Deficiency		
Increlex®	P	 Initial Criteria: Patient is < 21 years old; AND Epiphyses is open (therapy will not be approved once epiphyseal fusion occurs); AND One of the following: Diagnosis of growth failure due to severe primary IGF-1 deficiency defined by the following (documentation required): Height standard deviation score ≤ -3 Basal IGF-1 standard deviation score ≤ -3 Normal or elevated growth hormone Diagnosis of growth hormone (GH) gene deletion in a patient who has developed neutralizing antibodies to GH; AND Secondary causes of IGF-1 deficiency have been ruled out (e.g., hypothyroidism, malnutrition, hepatic disease, GHD, chronic corticosteroid treatment); AND Secondary causes of IGF-1 deficiency have been ruled out (e.g., hypothyroidism, malnutrition, hepatic disease, GHD, chronic corticosteroid treatment); AND Patient will not be treated with concurrent growth hormone therapy Note: Will not be approved for patients with active or secondary neoplasms, secondary forms of IGF-1 deficiency, weight loss management, nor as a substitute for growth hormone. Renewal Criteria: Patient is < 21 years old; AND Prescriber attests patient has had a height increase of ≥ 2 cm/year over the previous year of treatment; AND		General PA Form
		Lambert-Eaton Myasthenic Syndrome (LEMS)		
Firdapse®	NP	 Initial Criteria: Diagnosis of Lambert-Eaton Myasthenic Syndrome (LEMS) confirmed by a positive anti-P/Q type voltage-gated calcium ch annel antibody test; AND Patient is ≥ 6 years old; AND Patient does not have a history of seizures; AND Patient does not have a hypersensitivity to amifampridine or another aminopyridine (such as dalfampridine [Ampyra®]) Renewal Criteria: Patient has not experienced any treatment-restricting adverse effects; AND Patient must demonstrate disease improvement, stabilization, and/or slowing in the rate of decline due to the medication 	10/day	General PA Form



		RARE CONDITIONS		
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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Leptin Deficiency		
Myalept®	NP	Initial Criteria: Diagnosis of congenital or acquired lipodystrophy; AND Leptin deficiency confirmed by laboratory testing; AND Patient has one of the following complications of lipodystrophy: Diabetes mellitus Hypertriglyceridemia Hepatic steatosis Polycystic ovarian syndrome Acanthosis nigricans; AND Requested agent will be used as adjunct to dietary management of lipodystrophy; AND Documented baseline HbA1C, fasting glucose, triglycerides, and liver enzymes provided; AND Patient does NOT have HIV-related or partial lipodystrophy or metabolic disease without concurrent evidence of generalized lipodystrophy; AND Prescriber is enrolled in the Myalept REMS program Renewal Criteria: Documented positive clinical response to therapy (e.g., improved glycemic control, decrease in triglycerides)		General PA Form
		Lysosomal Storage Disease		
Aqneursa®	NP	 Initial Criteria Patient weighs at least 15 kg; AND Diagnosis of Niemann-Pick disease type C confirmed by one of the following: Genetically confirmed mutations in both alleles of NPC1 or NPC2; OR Mutation in only one allele of NPC1 or NPC2 plus either positive filipin staining or elevated cholestane triol/oxysterols (>2 x upper limit of normal); AND Patient has at least one neurological symptom of the disease (e.g., hearing loss, vertical supranuclear gaze palsy, ataxia, dementia, dystonia, seizures, dysarthria, or dysphagia); AND Prescribed by, or in consultation with, a neurologist, a medical geneticist, a metabolic disorder sub-specialist, or a physician who specializes in the treatment of lysosomal storage disorders Renewal Criteria Submission of medical records (e.g., chart notes) documenting a positive clinical response to therapy (e.g., improvement or stabilization in neurological symptoms of disease) 		General PA Form
Cerdelga®	NP		2/day	



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Galafold®	NP	 Initial Criteria (6-month duration): Patient is ≥ 18 years old; AND Documented diagnosis of Fabry disease with biochemical/genetic confirmation by 1 of the following:	14/28 days	General PA Form
Miplyffa®	NP	Initial Criteria: Patient is 2 years of age or older; AND Diagnosis of Niemann-Pick disease type C confirmed by ONE of the following: Genetically confirmed mutations in both alleles of NPC1 or NPC2; OR Mutation in only one allele of NPC1 or NPC2 plus either positive filipin staining or elevated cholestane triol/oxysterols (>2 x upper limit of normal); AND Patient has at least one neurological symptom of the disease (e.g., hearing loss, vertical supranuclear gaze palsy, ataxia, dementia, dystonia, seizures, dysarthria, dysphagia); AND Miplyffa will be used in combination with miglustat; AND Prescribed by, or in consultation with, a neurologist, a medical geneticist, a metabolic disorder sub-specialist, or a physician who specializes in the treatment of lysosomal storage disorders Renewal Criteria: Patient continued to meet initial criteria; AND Submission of medical records (e.g., chart notes) documenting a positive clinical response to therapy (e.g., improvement or stabilization in neurological symptoms of disease)		General PA Form
Opfolda®	NP	 Patient is ≥ 18 years old and weighs at least 40 kg; AND Diagnosis of late-onset Pompe disease confirmed by ONE of the following: Documentation demonstrating deficiency of acid alpha-glucosidase (GAA) enzyme activity Molecular genetic test demonstrating pathogenic variants in GAA; AND Prescriber attest patient did not have clinical improvement on enzyme replacement therapy (e.g., Lumizyme, Nexviazyme, Elfabrio); AND Must be used in combination with Pombiliti (cipaglucosidase alfa); AND Prescribed by, or in consultation with, a neurologist, a medical geneticist, a metabolic disorder sub-specialist, or a physician who specializes in the treatment of lysosomal storage disorders 	8/28 days	General PA Form



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Procysbi®	NP	Initial Criteria (6-month duration): Diagnosis of nephropathic cystinosis; AND Patient is ≥ 1 year old; AND Trial and failure, contraindication, or intolerance to Cystagon®; AND WBC cystine levels or plasma cysteamine concentration will be monitored Renewal Criteria: Documentation of positive clinical response to therapy; AND WBC cystine levels or plasma cysteamine concentration will be monitored		General PA Form
		Neuromyelitis Optica Spectrum Disorder (NMOSD)		
Enspryng®	NP	Initial Criteria (6-month duration): • Diagnosis of neuromyelitis optica spectrum disorder (NMOSD); AND • Patient ≥ 12 years old of age; AND • Patient is anti-aquaporin-4 (AQP4) antibody positive; AND • Patient has been screened, and does not have any of the following: • Active Hepatitis B infection • Active or untreated latent tuberculosis • Active infection; AND • Prescribed by or in consultation with a neurologist or ophthalmologist Renewal criteria: • Patient has demonstrated positive response to therapy	Loading Dose: 3 syringes/28 days Maintenance: 1/28 days	General PA Form
		Myasthenia Gravis		
Zilbrysq®	NP	Initial Criteria: (6- month duration) Diagnosis of generalized myasthenia gravis (gMG); AND Documented positive serology for acetylcholine receptor (AChR) autoantibodies; AND Patient has a baseline MG-Activities of Daily Living (MG-ADL) total score of ≥6; AND Patient has tried and failed, or has contraindication, or intolerance to TWO of the following: Corticosteroids Azathioprine Cyclosporine mycophenolate mofetil methotrexate tacrolimus; AND Prescribed by, or in consultation with, a neurologist or neuromuscular specialist; AND Prescriber is enrolled in the Zilbrysq REMS Program; AND Patient has not failed a previous course of Zilbrysq, Ultomiris, or Soliris therapy; AND Patient is not receiving Zilbrysq in combination with another complement inhibitor (e.g., Soliris, Ultomiris) Renewal Criteria: Submission of medical records (e.g., chart notes) documenting a positive clinical response to therapy (e.g., reduction in MG-ADL score or improvement in talking, chewing, swallowing, breathing, double vision, eyelid drop, movement)	1/day	General PA Form



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form	
		Paroxysmal Nocturnal Hemoglobinuria (PNH)			
Empaveli®	NP	 Initial Criteria: Diagnosis of paroxysmal nocturnal hemoglobinuria (PNH) confirmed by flow cytometry; AND Prescribed by, or in consultation with, a hematologist or oncologist; AND Patient has symptoms attributed to PNH (e.g., anemia, extreme fatigue, difficulty swallowing, recurrent abdominal pain, smooth muscle dystonia, thrombosis, CKD, organ damage secondary to chronic hemolysis); AND One of the following: Will not be used concurrently combination with another complement inhibitor (e.g., Soliris, Ultomiris) Patient is currently receiving another complement inhibitor (e.g., Empaveli, Soliris, Ultomiris) which will be discontinued and Fabhalta will be initiated in accordance with the FDA approved labeling; AND One of the following: The requested quantity does not exceed 1,080 mg twice weekly The requested quantity is for 1,080 mg every 3 days and lactate dehydrogenase (LDH) is >2 the upper limit of the normal range (LDH level documentation is required) Renewal Criteria: Documentation of positive clinical response to therapy (e.g., increased or stabilization of hemoglobin levels, reduction in transfusions, improvement in hemolysis, decrease in LDH); AND Prescribed by, or in consultation with, a hematologist or oncologist; AND Patient is not receiving Empaveli in combination with another complement inhibitor; AND One of the following: The requested quantity does not exceed 1,080 mg twice weekly The requested quantity is for 1,080 mg every 3 days and lactate dehydrogenase (LDH) is >2 the upper limit of the normal range (LDH level documentation is required) 	200 mL/30 days	General PA Form	



	RARE CONDITIONS					
Medication	PDL	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. Prior Authorization Criteria	Qty. Limits	PA Form		
Fabhalta®	NP	Initial Criteria: (6-month duration) One of the following: Diagnosis of primary IgA immunoglobulin A nephropathy (IgAN) confirmed by kidney biopsy; AND Patient is at risk of rapid disease progression (e.g., proteinuria > 0.5 g/day or UPCR > 0.8 g/g); AND Patient has ar isk of rapid disease progression (e.g., proteinuria > 0.5 g/day or UPCR > 0.8 g/g); AND Patient has an eGFR > 20 mL/min/1.73 m²; OR Diagnosis of paroxysmal nocturnal hemoglobinuria (PNH) confirmed by flow cytometry; AND Patient has symptoms attributed to PNH (e.g., anemia, extreme fatigue, difficulty swallowing, recurrent abdominal pain, smooth muscle dystonia, thrombosis, CKD, organ damage secondary to chronic hemolysis); AND Prescribed by, or in consultation with, a hematologist, nephrologist, or oncologist; AND Prescriber is enrolled in the Fabhalta REMS Program; AND One of the following: Will not be used concurrently with another complement inhibitor; OR Patient is currently receiving another complement inhibitor (e.g., Empaveli, Soliris, Ultomiris) which will be discontinued and Fabhalta will be initiated in accordance with the FDA approved labeling Renewal Criteria: Submission of medical records (e.g., chart notes) documenting a positive clinical response to therapy (e.g., increased or stabilization of hemoglobin levels, reduction in transfusions, improvement in hemolysis, decrease in LDH); AND Prescribed by, or in consultation with, a hematologist or oncologist; AND Patient is not receiving Empaveli in combination with another complement inhibitor	2/day	General PA Form		
Voydeya®	NP	 Initial Criteria: (6-month duration) Diagnosis of paroxysmal nocturnal hemoglobinuria (PNH) confirmed by flow cytometry; AND Patient has symptoms attributed to PNH (e.g., anemia, extreme fatigue, difficulty swallowing, recurrent abdominal pain, smooth muscle dystonia, thrombosis, CKD, organ damage secondary to chronic hemolysis); AND Patient is experiencing extravascular hemolysis (EVH) while on complement C5 inhibitor Ultomiris or Soliris; AND Prescriber attests Voydeya will be used in combination with Ultomiris or Soliris; AND Prescriber is enrolled in the Voydeya REMS Program; AND Prescribed by, or in consultation with, a hematologist or oncologist Renewal Criteria: Submission of medical records (e.g., chart notes) documenting a positive clinical response to therapy (e.g. hemoglobin stabilization, decreased number of blood transfusions, improvement in signs and symptoms of the disease); AND Prescribed by, or in consultation with, a hematologist or oncologist; AND Patient will continue to use Voydeya in combination with Ultomiris or Soliris 	6/day	General PA Form		



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Medication	PDL		Qty. Limits	PA Form
		Phenylketonuria (PKU)		'
Palynziq®	Р	 Diagnosis of Phenylketonuria (PKU); AND Prescribed by, or in consultation with, a physician who specializes in the treatment of inherited metabolic disorders; AND Patient will receive first dose of Palynziq® in prescribing MD's office; AND Patient meets ONE of the following: Patient has blood phenylalanine (Phe) concentrations > 600 µmol/L; OR Prescriber attests patient cannot maintain a healthy diet with Phe restriction; OR Patient has neurocognitive deficits; OR Trial and failure, contraindication, or intolerance of sapropterin 		General PA Form
sapropterin	Р	 Patient has diagnosis of Phenylketonuria (PKU); AND Prescribed by, or in consultation with, a metabolic specialist; AND Prescriber attests that Phenylalanine (Phe) levels cannot be maintained within recommended range (120-360 umol/L) with dietary intervention alone; AND Medication will be used in conjunction with a phenylalanine restricted diet 		General PA Form
Javygtor®	NP	See sapropterin prior authorization criteria; AND Clinically valid reason why the preferred sapropterin agents cannot be used		General PA
Kuvan®	NP	See sapropterin prior authorization criteria; AND Clinically valid reason why the preferred sapropterin agents cannot be used		<u>Form</u>
	l	PIK3CA-Related Overgrowth Spectrum (PROS)		
Vijoice®	NP	Initial Criteria (6-month duration): Diagnosis of PIK3CA-Related Overgrowth Spectrum (PROS); AND Patient has a mutation of the PIK3CA gene; AND Patient has a mutation of the PIK3CA gene; AND Patient has severe manifestations of PROS and requires systemic therapy; AND Vijoice will NOT be used for an oncology diagnosis; AND Prescriber attests to monitor, and potentially discontinue Vijoice treatment, if patient shows any of the following: Signs or symptoms of severe cutaneous adverse reactions (SCARs) New or worsening respiratory symptoms or is suspected to have developed pneumonitis Severe diarrhea Severe hyperglycemia Severe hypersensitivity; AND Female and male patients of reproductive potential have been advised to use effective contraception during treatment and for one week after the last dose Renewal Criteria: Patient continues to meet initial criteria; AND Prescriber attests patient has had ≥ 20% reduction from baseline in the measurable target lesion volume confirmed by at least one subsequent imaging assessment		General PA Form



		RARE CONDITIONS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Pyruvate Kinase (PK) Deficiency		
Pyrukynd®	NP	Initial Criteria (6-month duration): • Patient has diagnosis of hemolytic anemia with pyruvate kinase (PK) deficiency; AND • Patient has at least 2 variant alleles in the PK liver and red blood cell gene of which at least 1 was a missense variant; AND • Hemoglobin is <10 g/dL; AND • One of the following: • Patient has symptomatic anemia • Patient is transfusion dependent; AND • Prescribed by or in consultation with a hematologist Renewal Criteria: • Documentation of positive clinical response to therapy as evidenced by one of the following: • Hemoglobin increase ≥ 1.5 g/dL from baseline • Reduction in the number of red blood cell units transfused from baseline	2 tabs/day	General PA Form
	•	Rett Syndrome		
Daybue®	NP	Initial Criteria: Patient is > 2 years old; AND Diagnosis of Rett Syndrome; AND Prescribed by, or in consultation with, a neurologist, clinical geneticist, or developmental pediatrician Renewal Criteria: Documentation of positive clinical response to Daybue® (e.g. improvement or stabilization in purposeful hand skills, spoken language, repetitive hand movements, and gait abnormalities)	120 mL/day	General PA Form
		Sickle Cell Disease		
Endari®	NP	Initial Criteria: Diagnosis of sickle cell disease; AND Patient meets ONE of the following: Endari will be used in combination with hydroxyurea; OR Trial and failure, contraindications, or intolerance to hydroxyurea; AND Dosed according to weight-based dosing found in package insert: < 30 kg, up to 2 packets per day 30-65 kg, up to 4 packets per day > 65 kg, up to 6 packets per day Renewal Criteria: Documentation of positive clinical response to therapy(e.g., decrease in number of days in crisis, number of days in hospital, occurrence of Acute Chest Syndrome)	6 packs/day	General PA Form
L-glutamine pack	NP	See Endari prior authorization criteria; AND • Trial and failure, contraindications, or intolerance to Endari®	6 packs/day	General PA



		RARE CONDITIONS		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indica	ted.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Siklos®	NP	 Patient has a diagnosis of sickle cell anemia with recurrent moderate to severe painful crisis; AND ONE of the following: Documentation of need for dosing that will not allow the use of a preferred hydroxyurea agent Patient unable to swallow hydroxyurea capsules 		General PA Form
Xromi®	NP	 Diagnosis of sickle cell anemia with recurrent moderate to severe painful crises; AND One of the following: Documentation of need for dosing that will not allow the use of a preferred hydroxyurea agent; OR Patient is unable to swallow solid oral dosage forms of hydroxyurea 		General PA Form
	ı	Somatostatins and Related Agents	1	
Korlym®	Р	 Diagnosis of Cushing's Syndrome; AND Patient has type 2 diabetes mellitus or glucose intolerance; AND Patient has failed surgical treatment OR is not candidate for surgery; AND Will NOT be approved for use during pregnancy 		General PA Form
octreotide	Р	 Diagnosis of acromegaly; OR Treatment is for severe diarrhea/flushing episodes associated with metastatic carcinoid tumors; OR Treatment is for profuse watery diarrhea associated with VIP-secreting tumors 		<u> 101111</u>
Isturisa®	NP	Initial Criteria (6-month duration): Diagnosis of Cushing's disease; AND Patient has failed surgical treatment OR is not candidate for surgery; AND Trial and failure (trial duration ≥ 90 days) or intolerance to oral ketoconazole; AND Patient is 18 years of age or older; AND Prescribed by, or in consultation with, an endocrinologist Renewal Criteria: Documentation of positive clinical response to therapy (e.g., normalization or reduction of urinary free cortisol, improvement in signs or symptoms of the disease)	1 mg: 4/day 5 mg: 2/day 10 mg: 6/day	General PA Form
Mifepristone 300 mg tablet	NP	See Korlym prior authorization criteria; AND Clinically valid reason why the preferred Korlym® cannot be used		
Mycapssa®	NP	 Diagnosis of acromegaly; AND Patient has previously taken, responded to, and tolerated treatment with octreotide or lanreotide; AND Clinically valid reason why the patient is unable to be maintained on current octreotide or lanreotide therapy 	4/day	General PA Form



RARE CONDITIONS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. PDL Medication **Prior Authorization Criteria Qty. Limits** PA Form **Initial Criteria:** Diagnosis of Cushing's Syndrome; AND • Patient is being treated for endogenous hypercortisolemia (e.g., pituitary adenoma, ectopic tumor, adrenal · adenoma); AND Surgery is not an option or has not been curative; AND • Trial and failure (trial duration > 90 days) or intolerance to oral ketoconazole; AND Patient is 18 years of age or older; AND **General PA** Recorlev® NP 8/day • Prescribed by or in consultation with an endocrinologist; AND Form • Patient has had baseline liver enzymes and an electrocardiogram prior to initiating therapy, and prescriber attests to monitor regularly thereafter; AND · Patient does not have hypokalemia and hypomagnesemia, or has been corrected prior to therapy Renewal Criteria: • Documentation of positive clinical response to therapy (e.g., normalization or reduction of urinary free cortisol, improvement in signs or symptoms of the disease) See prior authorization criteria for octreotide; AND Sandostatin® NP • Clinically valid reason why preferred octreotide cannot be used General PA • Diagnosis of Cushing's Disease; AND Form Signifor® NP • Surgery is not an option or has not been curative; AND 2 injections/day • Prescribed by, or in consultation with, an endocrinologist Diagnosis of acromegaly; AND NP Somavert® • Tral and failure, intolerance, or contraindication to octreotide Initial Criteria: Patient has a carcinoid/neuroendocrine tumor and has been diagnosed with carcinoid syndrome; AND • Patient has had an inadequate treatment response to at least a 3-month trial of SSA (somatostatin analog) therapy at the highest tolerated dose; AND **General PA** Patient will continue to receive somatostatin analog therapy; AND Xermelo® NP 3/day Form Patient has tried and received an inadequate response to antidiarrheals (e.g., loperamide); AND • Patient has at least 4 bowel movements per day Renewal Criteria: • Documentation of positive clinical response to therapy (e.g., decrease in number of bowel movements per day)



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Spinal Muscular Atrophy (SMA)		
Evrysdi®	NP	Initial Criteria: Diagnosis of Spinal Muscular Atrophy (SMA); AND Submission of medical records (e.g., chart notes, laboratory values) confirming the mutation or deletion of genes in chromosome 5q confirming in one of the following: Homozygous gene deletion or mutation of SMN1 gene; OR Compound heterozygous mutation of SMN1 gene; AND Prescribed by, or in consultation with, a neurologist with expertise in the diagnosis and treatment of SMA; AND Patient will not receive concomitant survival motor neuron (SMN) modifying therapy (e.g., Spinraza); AND One of the following: Patient has not previously received gene replacement therapy for the treatment of SMA (e.g., Zolgensma); OR Both of the following: Patient has previously received gene therapy for the treatment of SMA (e.g., Zolgensma); AND Provider attests that there has been an inadequate response to gene therapy (e.g., sustained decrease in ≥ 1 motor test score over a period of 6-months); AND Advise female patients of reproductive potential to use effective contraception during treatment and for at least 1 month after the last dose Renewal criteria Patient continues to meet initial criteria; AND Patient has clinically significant improvement in SMA associated signs and symptoms (progression, stabilization, or decreased decline in motor function)	Tabs: 1/day Soln:3 bottles/ 28 days	General PA Form
		Transthyretin Amyloidosis Agents		_
Attruby®	NP	 Initial Criteria Patient is 18 years of age or older; AND Prescribed by or in consultation with a cardiologist; AND Diagnosis of wild type or hereditary transthyretin-mediated amyloidosis (ATTR-CM) with cardiomyopathy; AND Patient has New York Heart Association Class I, II or III heart failure; AND Patient does not meet have any of the following: Impaired renal function (glomerular filtration rate < 15 mL/min/1.73 m2) History of heart transplantation New York Heart Association Class IV; AND Patient will not use in combination with another medication indicated for the management cardiomyopathy or neuropathy of transthyretin-mediated amyloidosis (e.g., Amvuttra, Onpattro, Vyndaqel, Vyndamax) Renewal Criteria Patient meets initial criteria; AND Patient has demonstrated a positive benefit from therapy (e.g., improved clinical symptoms of heart failure) 	4/day	General PA Form



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Vyndamax®	NP	Initial Criteria: Patient is 18 years of age or older; AND Must be prescribed in consultation with a cardiologist; AND Diagnosis of wild type or hereditary transthyretin-mediated amyloidosis (ATTR-CM) with cardiomyopathy; AND Patient has New York Heart Association Class I, II or III heart failure; AND Patient does not have any of the following: Impaired renal function (glomerular filtration rate < 25 mL/min/1.73 m2) History of liver or heart transplantation Implanted left ventricular assist device (LVAD) [pacemaker or cardiac defibrillator allowed] Patient is pregnant or breastfeeding New York Heart Association Class IV; AND Patient will not use in combination with another medication indicated for the management cardiomyopathy or neuropathy of transthyretin-mediated amyloidosis (e.g., Amvuttra, Onpattro, Attruby) Renewal Criteria: Patient continues to meet initial criteria; AND Patient has demonstrated a positive benefit from therapy (e.g., improved clinical symptoms of heart failure)	1/day	General PA Form
Vyndaqel®	NP	See prior authorization criteria for Vyndamax	4/day	
Wainua®	NP	Initial Criteria: Patient is 18 years of age or older; AND Diagnosis of hereditary transthyretin-mediated amyloidosis (ATTRv amyloidosis) with polyneuropathy; AND Documentation that patient has a transthyretin (TTR) mutation (e.g., V30M); AND Prescribed by, or in consultation with, a neurologist, cardiologist, or specialist with knowledge of ATTRv; AND Documentation of ONE of the following: Patient has a baseline polyneuropathy disability (PND) score ≤ IIIb Patient has a baseline familial amyloidotic polyneuropathy (FAP) stage of 1 or 2 Patient has not had a liver transplant; AND Patient has not had a liver transplant; AND Will not be used in combination with another medication indicated for the management cardiomyopathy or neuropathy of transthyretin-mediated amyloidosis (e.g., Amvuttra, Onpattro, Attruby, Vyndaqel, Vyndamax) Renewal Criteria: Patient has demonstrated a benefit from therapy (e.g., improved neurologic impairment, motor function, slowing of disease progression, quality of life assessment); AND Will not be used in combination with another medication indicated for the management cardiomyopathy or neuropathy of transthyretin-mediated amyloidosis (e.g., Amvuttra, Onpattro, Attruby, Vyndaqel, Vyndamax)	1 injector/28 days	General PA Form
		Tyrosinemia Type 1		
Orfadin® suspension	NP NP	 Diagnosis of hereditary tyrosinemia type 1; AND Agent is prescribed by a physician specializing in the condition being treated; AND Patient has a clinically valid reason as to why the Orfadin® capsules cannot be utilized See Orfadin® suspension prior authorization criteria		General PA Form



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Nityr® tablet	NP	See Orfadin® suspension prior authorization criteria		
		Urea Cycle Disorders		
Carbaglu®	Р	Diagnosis of urea cycle disorders		
Pheburane®	Р	Diagnosis of urea cycle disorders		
carglumic acid	NP	 Diagnosis of urea cycle disorders; AND Trial and failure, contraindication, or intolerance of Carbaglu® 		
Olpruva®	NP	 Diagnosis of urea cycle disorders; AND Trial and failure, contraindication, or intolerance of Pheburane® 		General PA Form
Ravicti®	NP	See Olpruva® prior authorization criteria		
sodium phenylbutyrate	NP	 Diagnosis of urea cycle disorders; AND Trial and failure, contraindication, or intolerance of Buphenyl® 		
		Wilson Disease		
Galzin®	NP	 Diagnosis of Wilson's disease; AND Intolerance to zinc sulfate 		
Syprine®	NP		8/day	General PA Form
trientine	NP		250mg: 8/day 500mg: 4/day	<u> </u>
		WHIM Syndrome		
Xolremdi [®]	NP		4/day	General PA Form

	RENAL AND GENITOURINARY					
	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
		Alpha Blockers for BPH				
alfuzosin	Р		1/day			
tamsulosin	Р		2/day	<u>General PA</u>		
Cardura XL	NP		1/day	<u>Form</u>		
Flomax®	NP		2/day			



		RENAL AND GENITOURINARY		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indication.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Androgen Hormone Inhibitors		
dutasteride	Р		1/day	
finasteride	Р		1/day	General PA
Avodart®	NP		1/day	<u>Form</u>
Proscar®	NP		1/day	
Tezruly®	NP	Patient is unable to swallow solid dosage forms	20mL/day	
		Agents for BPH		
Cialis®	NP	 Diagnosis of Benign Prostatic Hypertrophy; AND Patient is NOT concurrently receiving nitrates or guanylate cyclase stimulators; AND Trial and failure, contraindication, or intolerance to at least ONE agent from each of the following classes: Alpha blockers for BPH Androgen Hormone Inhibitors 		
dutasteride/ tamsulosin	NP	Patient has a diagnosis of benign prostatic hyperplasia (BPH) with an enlarged prostate; AND Patient has a contraindication or adverse event to finasteride; AND Patient is unable to use the individual components	1/day	General PA
Entadfi [®]	NP	Criteria (6-month duration): Diagnosis of Benign Prostatic Hyperplasia (BPH) with an enlarged prostate; AND Total length of therapy has not exceeded 26 weeks; AND Trial and failure, contraindication, or intolerance to combination therapy with alpha blocker and androgen hormone inhibitor; AND Clinically valid reason why the individual components of Entadfi® cannot be used (finasteride and tadalafil); AND Patient is NOT concurrently receiving nitrates or guanylate cyclase stimulators	1/day; 182/year	<u>Form</u>
Jalyn®	NP	See dutasteride/tamsulosin prior authorization criteria	1/day	
		Phosphorus Depletors		
sevelamer carbonate tablets	Р		9/day	
Renvela® packs	Р	Patient is unable to swallow solid dosage forms	0.8 g packets: 6/day 2.4 g packets: 5/day	
Auryxia®	NP	 Diagnosis of hyperphosphatemia in chronic kidney disease on dialysis; AND Trial and failure, contraindication, or intolerance to TWO preferred agents; OR Diagnosis of iron deficiency anemia in chronic kidney disease NOT on dialysis; AND Trial and failure, contraindication, or intolerance to TWO oral iron products (e.g., ferrous sulfate, ferrous gluconate) 		General PA Form
Fosrenol® packs	NP	 Trial and failure, contraindication, or intolerance of TWO preferred phosphorus depletors; AND Contraindication to sevelamer powder for suspension; AND Patient is unable to swallow solid dosage forms 		General PA Form
Renvela® tablets	NP		9/day	
sevelamer carbonate packs	NP	Patient is unable to swallow solid dosage forms	0.8 g packets: 6/day 2.4 g packets: 5/day	
Xphozah®	NP	Patient is 18 years of age or older; AND	2/day	General PA



		RENAL AND GENITOURINARY Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicate	ed	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		 Diagnosis of chronic kidney disease (CKD); AND Patient is currently on dialysis; AND Trial and failure, contraindication, or intolerance of TWO preferred agents; AND Agent will be used as adjunctive therapy to reduce serum phosphorus; AND Patient does not have known or suspected mechanical gastrointestinal obstruction 		<u>Form</u>
		Kidney Stone Agents		
Thiola EC®	NP	 Patient has tried/failed an adequate trial of or is intolerant to two preferred agents; AND Clinically valid reason why preferred Thiola cannot be used 		General PA Form
		Urinary Tract Antispasmodics		
fesoterodine	Р		1/day	General PA
Myrbetriq® tabs	Р		1/day	<u>Form</u>
oxybutynin ER tabs	Р		5 mg: 1/day; 10, 15 mg: 2/day	
Oxytrol®	Р		8 patches/28 days	General PA Form
solifenacin	Р		1/day	FOITH
tolterodine ER caps	Р		1/day	
tolterodine tabs	Р		2/day	
darifenacin	NP		1/day	
Detrol®	NP		2/day	General PA
Detrol LA®	NP		1/day	<u>Form</u>
flavoxate	NP		2 fills/ 60 days	
Gelnique®	NP		1 pack (1 gr)/day	
Gemtesa®	NP	Patient is 18 years of age or older: AND Diagnosis of overactive bladder (OAB); AND Trial and failure of one preferred anticholinergic agent (e.g., fesoterodine, oxybutynin, solifenacin, tolterodine); AND Trial and failure, or contraindication, or intolerance to Myrbetriq	1/day	General PA Form
mirabegron tabs	NP	Clinically valid reason why preferred Myrbetriq® cannot be used	1/day	
Myrbetriq® susp	NP	 Clinically valid reason why Myrbetriq tablets cannot be used; OR Diagnosis of neurogenic detrusor overactivity (NDO); AND Trial and failure, contraindication, or intolerance to oxybutynin solution 		General PA Form
Toviaz®	NP		1/day	
trospium	NP		2/day	General PA
trospium XR	NP		1/day	<u>Form</u>
VESIcare® susp	NP	 Diagnosis of neurogenic detrusor overactivity (NDO); AND Trial and failure, contraindication, or intolerance to oxybutynin solution 	10 mL/day	General PA
VESIcare® tabs	NP		1/day	<u>Form</u>



		RESPIRATORY Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indica	ted.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Anaphylaxis Therapy Agents	'	
epinephrine auto injector	Р		2/Rx	
Auvi-Q	NP		2/Rx	General PA
EpiPen®	NP		2/Rx	<u>Form</u>
EpiPen-Jr®	NP		2/Rx	
Neffy®	NP	Clinically valid reason why the preferred epinephrine auto-injector cannot be used	2/Rx	
		Anticholinergics, Nasal		
ipratropium 0.3%	Р		2 boxes/30days	General PA
ipratropium 0.6%	Р		3 boxes/30days	<u>Form</u>
		Antihistamines, Nasal	1	_
Azelastine	Р		2 bottles/30 days	
Dymista®	Р		1 bottle/30 days	General PA
olopatadine	Р		1 bottle/30 days	Form
azelastine/ fluticasone	NP	Trial and failure of preferred Dymista®	1 bottle/30 days	<u>101111</u>
Ryaltris®	NP	 Diagnosis of Seasonal Allergic Rhinitis; AND Patient is 12 years of age or older; AND Trial and failure, contraindication, or intolerance to Dymista; AND Clinically valid reason as to why the patient is unable to take components of Ryaltris individually (Note: Patient convenience is not an approvable reason) 	1 bottle/30 days	General PA Form
		Antihistamines: Non-Sedating, Oral (Covered for recipients < 21 years old only)		
cetirizine	Р		1/day	
cetirizine chewable	Р	Clinically valid reason why the liquid formulation cannot be used	1/day	
cetirizine/PSE	Р		2/day	
levocetirizine tablets	Р		1/day	
loratadine tablets	Р		1/day	
loratadine syrup	Р		10 mL/day	
loratadine chewable	Р		1/day	
Ioratadine RDT	Р	Patient is unable to swallow solid dosage forms	1/day	General PA
loratadine/PSE	Р		12 Hour: 2/day;	Form
,			24 Hour (1/day)	4
Allegra®	NP		60mg: 2/day); 180mg (1/day)	
			12 Hour: 2/day;	╡ !
Allegra D®	NP		24 Hour: 1/day	
Allegra® ODT	NP	Patient is unable to swallow solid dosage forms	2/day	1
Clarinex D®	NP	-	12 Hour (2/day);	7
Ciai iiiex D	INP		24 Hour (1/day)	



		RESPIRATORY Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated	t.	
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Clarinex RediTabs®	NP	Patient is unable to swallow solid dosage forms	1/day	
Clarinex® tabs	NP		1/day	
Clarinex® syrup	NP		10mg/day	
Claritin D®	NP		12 Hour: 2/day; 24 Hour: 1/day	
Claritin® chewable	NP	Clinically valid reason why the liquid formulation cannot be used	1/day	
Claritin® tabs	NP		1/day	
Claritin RediTabs®	NP	Patient is unable to swallow solid dosage forms	1/day	
desloratadine	NP		1/day	General PA
desloratadine ODT	NP	Patient is unable to swallow solid dosage forms	1/day	Form
fexofenadine	NP		60 mg: 2/day); 180 mg (1/day)	101111
fexofenadine/PSE	NP		12 Hour: 2/day; 24 Hour: 1/day	
levocetirizine solution	NP		10 mL/day	
Semprex®-D	NP		4/day	
Xyzal®	NP		5 mg/day	
Zyrtec® chewable	NP	Clinically valid reason why the liquid formulation cannot be used	1/day	General PA
Zyrtec® tabs	NP		1/day	<u>Form</u>
Zyrtec® ODT	NP	Patient is unable to swallow solid dosage forms	1/day	
Zyrtec D®	NP		1/day	
		Antitussives, Non-Narcotic		
benzonatate	Р	 Patient is ≥ 10 years of age; OR Patient is < 10 years of age and prescriber is aware that, if chewed, benzonatate may cause numbness of the mouth, tongue, throat, and esophagus, increasing the risk of choking 	3/day	General PA Form
		Cystic Fibrosis Agents, Inhaled/Injectable		
Bethkis®	Р	Diagnosis of Cystic Fibrosis or <i>Pseudomonas</i> infection	224 mL/56 days	
Kitabis Pak®	Р	Diagnosis of Cystic Fibrosis or <i>Pseudomonas</i> infection	280 mL/56 days	
Pulmozyme®	Р	Diagnosis of Cystic Fibrosis or <i>Pseudomonas</i> infection	5 mL/day	
tobramycin solution 300 mg/5 mL	Р	Diagnosis of Cystic Fibrosis or <i>Pseudomonas</i> infection	280 mL/56 days	Con and DA
tobramycin vial (excluding 1.2 g vials)	Р	Claims exceeding \$200 will only be approved for diagnoses of Cystic Fibrosis or <i>Pseudomonas</i> infection		General PA Form
Bronchitol	NP	 Diagnosis of Cystic Fibrosis; AND Patient must not have an episode of hemoptysis (>60 mL) in the last 3 months; AND Must be 18 years of age or older; AND Patient must have baseline FEV1 >40% to <90%; AND 	20/day	



		RESPIRATORY		
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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		 Patient has passed the Bronchitol Tolerance Test; AND Must be used concomitantly with a short-acting bronchodilator; AND Prescriber attests that the patient has been instructed to administer the agent 5-15 minutes after a short-acting bronchodilator 		
Cayston®	NP	 Diagnosis of Cystic Fibrosis or Pseudomonas Infection; AND Trial and failure, contraindication, intolerance, or resistance to preferred inhaled tobramycin product 	84 mL/56 days	
tobramycin solution 300 mg/4 mL (generic for Bethkis)	NP	 Diagnosis of Cystic Fibrosis or <i>Pseudomonas</i> infection; AND Clinically valid reason why preferred Bethkis® cannot be used 	224 mL/56 days	General P
TOBI® Podhaler and inhalation	NP	 Diagnosis of Cystic Fibrosis or Pseudomonas Infection; AND Provider must provide peer-reviewed medical literature documenting why the requested drug for the requested indication is the only appropriate choice versus the preferred agents 	Podhaler: 224 caps/56 days; Solution: 280 mL/56 days	- <u>Form</u>
		Cystic Fibrosis Agents, Oral		-
Alyftrek®	NP	Initial Criteria (6-month duration): • Diagnosis of cystic fibrosis (CF); AND • Must be prescribed by, or in consultation with, a provider at a CF Center of Excellence or pulmonologist; AND • Age ≥ 6 years old; AND • Lab documentation confirming ONE of the following: ○ Patient has at least one copy of the F508del mutation in the CFTR gene ○ Patient has a mutation in the CFTR gene that is responsive based on in vitro data; AND • For patients 6- 12 years of age, prescriber attests to obtain ophthalmic examination before and during treatment Renewal Criteria: • Disease response to therapy and tolerability compared to baseline (e.g., decreased pulmonary exacerbations, improvement, or stabilization of lung function)	4-20-50 mg: 3/day 10-50-125 mg: 2/day	General P Form
Kalydeco®	NP	 Initial Criteria (6-month duration): Diagnosis of cystic fibrosis (CF); AND Must be prescribed by, or in consultation with, a provider at a CF Center of Excellence or pulmonologist; AND Lab documentation confirming patient has one mutation in the CFTR gene that is responsive to Kalydeco®; AND For patients 2- 12 years of age, prescriber attests to obtain ophthalmic examination before and during treatment Renewal Criteria: Disease response to therapy and tolerability compared to baseline (e.g., decreased pulmonary exacerbations, improvement, or stabilization of lung function) Note: will NOT be approved for homozygous F508del mutation in the CFTR gene 	2/day	General P. Form



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Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Orkambi®	NP	 Initial Criteria (6-month duration): Diagnosis of cystic fibrosis (CF); AND Must be prescribed by, or in consultation with, a provider at a CF Center of Excellence or pulmonologist; AND Age ≥ 1 years old; AND Lab documentation confirming patient has homozygous F508del mutation in the CFTR gene For patients 2- 12 years of age, prescriber attests to obtain ophthalmic examination before and during treatment Renewal Criteria: Disease response to therapy and tolerability compared to baseline (e.g., decreased pulmonary exacerbations, improvement, or stabilization of lung function) 	Tablets: 4/day Granules: 2/day	General PA Form
Symdeko®	NP	 Initial Criteria (6-month duration): Diagnosis of cystic fibrosis (CF); AND Must be prescribed by, or in consultation with, a provider at a CF Center of Excellence or pulmonologist; AND Age ≥ 6 years old; AND Lab documentation confirming ONE of the following:	2/day	General PA Form
Trikafta®	NP	 Initial Criteria (6-month duration): Diagnosis of cystic fibrosis (CF); AND Must be prescribed by, or in consultation with, a provider at a CF Center of Excellence or pulmonologist; AND Patient is ≥ 2 years of age; AND Lab documentation confirming ONE of the following:	3/day	General PA Form



		RESPIRATORY		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
		Inhaled: Anticholinergics and Anticholinergic Combinations		
Anoro Ellipta®	Р		2 blisters/day	
albuterol/ ipratropium	Р		18 mL/day	
Atrovent HFA®	Р		2 inhalers/month	
ipratropium solution	Р		10 mL/day	
Spiriva HandiHaler®	Р		1 capsule/day	General PA
Spiriva Respimat®	Р	 Diagnosis of Asthma; AND Patient age ≥ 6 years; AND Diagnosis of step 4 or higher asthma; AND Optimal doses of inhaled steroids and long-acting beta-agonists are being used and breakthrough symptoms require frequent use of inhaled short-acting bronchodilators; OR Diagnosis of Chronic Obstructive Pulmonary Disorder (COPD); AND Must be used as maintenance therapy only; AND Trial and failure, contraindication, or intolerance to Spiriva HandiHaler® 	1 inhaler/month	Form
Trelegy Ellipta®	Р	 Initial Criteria: Diagnosis of chronic obstructive pulmonary disease (COPD); AND Trial and failure (as defined by continued symptoms, including exacerbations) of adequate treatment with a long-acting beta-agonist + long-acting antimuscarinic; AND Must be used as maintenance therapy only; OR A diagnosis of asthma in patients 12 years of age or older; AND Trial and failure (as defined by continued symptoms, including exacerbations) of adequate treatment with 2 dual combination inhaled corticosteroid + long-acting beta-agonist therapies; AND Must be used as maintenance therapy only; AND Patient does not have known hypersensitivity to milk proteins Renewal Criteria: Documentation of continued efficacy via prescriber's medical opinion on patient evaluation; AND Patient has not experienced any intolerable adverse effects (e.g., hypersensitivity, bronchospasm, worsening of intraocular pressure, increased severe infections) 	2 blisters/day	General PA Form
Bevespi Aerosphere®	NP	Diagnosis of Chronic Obstructive Pulmonary Disorder (COPD); AND Must be used as maintenance therapy only; AND Trial and failure, contraindication, or intolerance to TWO preferred inhaled anticholinergic/anticholinergic combination agents	1 inhaler/ month	



	RESPIRATORY Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
Breztri Aerosphere®	NP	 Initial Criteria: Diagnosis of chronic obstructive pulmonary disease (COPD); AND Must be used as maintenance therapy only; AND Trial and failure (as defined by continued symptoms, including exacerbations) of adequate treatment with a long-acting beta-agonist + long-acting antimuscarinic; AND Trial and failure, contraindication, or intolerance to the preferred product Trelegy Ellipta Renewal Criteria: Documentation of continued efficacy via prescriber's medical opinion on patient evaluation; AND Patient has not experienced any intolerable adverse effects (e.g., hypersensitivity, bronchospasm, worsening of intraocular pressure, increased severe infections) 	1 inhaler/month			
Combivent Respimat®	NP	Diagnosis of Chronic Obstructive Pulmonary Disorder (COPD); AND Trial and failure, contraindication, or intolerance to TWO preferred inhaled anticholinergic/anticholinergic combination agents	2 inhalers/month			
Duaklir Pressair®	NP	Diagnosis of Chronic Obstructive Pulmonary Disorder (COPD); AND Must be used as maintenance therapy only; AND Trial and failure, contraindication, or intolerance to TWO preferred inhaled anticholinergic/anticholinergic combination agents	1 inhaler/month	General PA		
Incruse Ellipta®	NP	Diagnosis of Chronic Obstructive Pulmonary Disorder (COPD); AND Must be used as maintenance therapy only; AND Trial and failure, contraindication, or intolerance to TWO preferred inhaled anticholinergic/anticholinergic combination agents Patient must not have severe hypersensitivity to milk proteins	1 blister/day	<u>Form</u>		
Stiolto Respimat®	NP	See Duaklir Pressair prior authorization criteria	1 inhaler/month			
tiotropium inhalation capsules	NP	Clinically valid reason why the patient cannot use the preferred brand Spiriva HandiHaler	1 capsule/day			
Tudorza®	NP	See Incruse Ellipta® prior authorization criteria	1 inhaler/month			
Yupelri®	NP	Initial Criteria: Patient must be ≥ 18 years of age; AND Diagnosis of Chronic Obstructive Pulmonary Disorder (COPD); AND Trial and failure, contraindication, or intolerance to TWO preferred inhaled anticholinergic/anticholinergic combination agents; AND Must be used as maintenance therapy only; AND Patient is unable to master proper inhaler technique, as attested by prescriber; AND Patient is not prescribed other inhaled long-acting anticholinergic agents. Renewal Criteria: Patient continues to meet initial criteria; AND Patient symptoms are clinically improving, as documented by provider; AND Patient demonstrates continued compliance, based on fill history (not using PRN); AND Prescriber documents that nebulized therapy continues to be required.	3 mL/day	General PA Form		



		RESPIRATORY				
Medication	PDL	Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicat		DA Forms		
Medication	PUL	Prior Authorization Criteria	Qty. Limits	PA Form		
		Inhaled: Beta Agonists-Corticosteroid Combination Products				
Advair HFA®	Р		1 inhaler/month			
Advair Diskus®	Р		2 blisters/day			
Dulera®	Р		2 inhalers/month			
fluticasone/ salmeterol aerosol	Р		1 inhaler/month			
Symbicort®	Р		2 inhalers/month			
AirDuo RespiClick®	NP	 Agent will be used for the treatment of asthma in patients 12 years of age or older; AND Trial and failure, contraindication, or intolerance of TWO preferred agents; AND Patient must not have severe hypersensitivity to milk proteins 	1 inhaler/month			
Airsupra®	NP	 Agent will be used for the treatment of asthma in patients 18 years of age and older; AND Trial and failure, contraindication, or intolerance to preferred agents Symbicort and Dulera 	2 inhalers/month			
Breo Ellipta®	NP	 Agent will be used for the treatment of asthma in patients 18 years of age or older; OR Agent will be used for the treatment of COPD where optimal doses of a long-acting beta agonist and/or long-acting muscarinic antagonists are being used and symptoms are still uncontrolled (100/25 mcg strength only); AND Trial and failure, contraindication, or intolerance of TWO preferred agents; AND Patient must not have severe hypersensitivity to milk proteins 	2/day	Beta Agonist Combos		
Breyna®	NP	 Trial and failure, contraindication, or intolerance of TWO preferred agents; AND Clinically valid reason why the patient cannot use the preferred brand Symbicort® 	2 inhalers/month			
budesonide/ formoterol	NP	 Trial and failure, contraindication, or intolerance of TWO preferred agents; AND Clinically valid reason why the patient cannot use the preferred brand Symbicort® 	2 inhalers/month			
fluticasone/ salmeterol powder	NP	 Trial and failure, contraindication, or intolerance of TWO preferred agents; AND Clinically valid reason why the patient cannot use the preferred Advair HFA® or Advair Diskus® cannot be used. 	55, 113, 232, -14mcg: 1 inhaler/month 100, 250, 500 -50mcg: 2 blisters/day			
fluticasone/ vilanterol	NP	See Breo Ellipta® prior authorization criteria; AND • Clinically valid reason why the patient cannot use the brand Breo Ellipta®	2/day			
Wixela®	NP	 Trial and failure, contraindication, or intolerance of TWO preferred agents; AND Clinically valid reason why the patient cannot use the preferred Advair HFA® or Advair Diskus® 	2 blisters/day			
		Inhaled: Beta Agonists, Long Acting				
Serevent Diskus®	Р		2 blisters/day	Compand DA		
Striverdi Respimat®	NP	 Diagnosis of COPD; AND Trial and failure, contraindication, or intolerance of the preferred agent (Serevent Diskus) 	1/day	General PA Form		
	Inhaled: Beta Agonists, Short Acting					
albuterol HFA	Р		2 inhalers/month			
Proventil® HFA	Р		2 inhalers/month			



		RESPIRATORY Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Ventolin® HFA	Р		2 inhalers/month	
Xopenex® HFA	Р	Patients has experienced intolerable side effects to albuterol (e.g., tachycardia, etc.)	2 canisters/month	
levalbuterol HFA	NP	 Patients has experienced intolerable side effects to albuterol (e.g., tachycardia, etc.); AND Clinically valid rationale for why patient cannot use brand Xopenex HFA® 	2 canisters /month	
ProAir RespiClick®	NP		2 inhalers/month	
		Inhaled: Nebulizers, Beta Agonists		
albuterol nebulizer solution	Р		125 nebs/month (3 bottles/month	
arformoterol	Р		60 nebs/month	
Brovana®	NP	 Diagnosis of COPD; AND Difficulty using a dry powder inhaler (DPI); AND Trial and failure, contraindication, or intolerance of the preferred agent (arformoterol nebulizer) 	60 nebs/month (120 mL/month)	General PA Form
formoterol	NP	See Brovana® prior authorization criteria	60 nebs/month	
levalbuterol	NP	Patients has experienced intolerable side effects to albuterol (e.g., tachycardia)	96 nebs/month	
Perforomist®	NP	See Brovana® prior authorization criteria	60 nebs/month	
	I	Inhaled: Nebulizers, Mast Cell Stabilizers	1	
cromolyn solution	Р	Diagnosis of asthma	120 vials/month	General PA Form
		Inhaled: Steroids	•	
Alvesco®	Р	 Diagnosis of asthma; AND Patient is 12 years of age or older 	2/30 days	General PA Form
Arnuity Ellipta®	Р		1 blister/day	
Asmanex HFA®	Р		1/30 days	
Asmanex Twisthaler®	Р		1/30 days	
budesonide suspension	Р	 ONE of the following: Diagnosis of asthma; AND Patient is < 8 years old; OR Diagnosis of Eosinophilic esophagitis (EoE); AND Prescribed by, or in consultation with, a gastroenterologist, allergist, or immunologist; AND Prescriber attest to both of the following:	0.25, 0.5 mg: 2 vials/day; 1 mg: 1 vial/day	General PA Form



		RESPIRATORY Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
Flovent Diskus®	Р		50 mcg: 2/day; 100 mcg: 4/day; 250 mcg: 8/day	
Flovent HFA®	Р		2/30 days	
fluticasone HFA	Р		2/30 days	
Pulmicort Flexhaler®	Р	 Diagnosis of asthma; AND Patient is 6 years of age or older 	2/30 days	
Pulmicort Respules®	Р	 Diagnosis of asthma; AND Patient is ≤ 8 years old 	0.25, 0.5 mg: 2 vials/day; 1 mg: 1 vial/day	
QVAR RediHaler®	Р		2/30 days	
		Intranasal: Steroids		
budesonide nasal (OTC)	Р		2/30 days	
fluticasone propionate	Р		1/30 days	General PA Form
Nasacort® (<u>OTC</u>)	Р		2/30 days	
budesonide nasal (Rx only)	Р		2/30 days	
flunisolide	NP		2/30 days	
mometasone furoate	NP		1/30 days	
Nasonex®	NP		1/30 days	General PA
Omnaris®	NP		1/30 days	<u>Form</u>
Qnasl®	NP		1/30 days	
triamcinolone acetonide	NP		1/30 days	
Xhance®	NP	 Patient has a trial/failure, contraindication, or intolerance to at least 2 preferred nasal corticosteroid agents; AND Patient has a clinically valid reason as to why preferred fluticasone propionate products cannot be used 	2/30 days	
		Leukotriene Modifiers		
montelukast tabs and chewables	Р		1/day	General PA
Accolate®	NP	 Trial and failure, contraindication, or intolerance of a preferred agent (montelukast tablets or chewables); AND Patient is 5 years of age or older and has a diagnosis of asthma 	2/day	<u>Form</u>



		RESPIRATORY		
		Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.		
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form
montelukast granules	NP	 One of the following: Diagnosis of asthma in patients 12 months of age or older; OR Diagnosis of exercise-induced bronchoconstriction (EIB) documented with concomitant use of at least one other asthma medication in patients 6 years of age or older; OR For treatment of seasonal allergic rhinitis in patients 2 years of age or older OR perennial allergic rhinitis in patients 6-months of age or older, patient must have failed trial of an intranasal corticosteroid OR a non-sedating antihistamine; AND Will be approved ONLY for patients who have clinically valid reason not to use chewable tablets Note: For patients less than 3 years of age, no prior authorization is required 	1/day	
Singulair® tabs and chewables	NP	 One of the following: Diagnosis of asthma in patients 12 months of age or older; OR Diagnosis of exercise-induced bronchoconstriction (EIB) documented with concomitant use of at least one other asthma medication in patients 6 years of age or older; OR For treatment of seasonal allergic rhinitis in patients 2 years of age or older OR perennial allergic rhinitis in patients 6-months of age or older, patient must have failed trial of an intranasal corticosteroid OR a non-sedating antihistamine; AND Trial and failure, contraindication, or intolerance of a preferred agent (montelukast tablets or chewables) 	1/day	
Singulair® granules	NP	See montelukast granules prior authorization criteria; AND Trial and failure, contraindication, or intolerance of a preferred agent (montelukast tablets or chewables)	1/day	
zafirlukast	NP	See Accolate® prior authorization criteria	2/day	
zileuton CR	NP	 Trial and failure, contraindication, or intolerance of a preferred agent (montelukast tablets or chewables); AND Patient is 12 years of age or older and has a diagnosis of asthma 	4/day	
Zyflo®	NP	See zileuton CR prior authorization criteria	4/day	
		Miscellaneous: OTC Products		
Peak Flow Meters			4 per 365 days	General PA
Spacers			4 per 365 days	<u>Form</u>
		Phosphodiesterase 4 (PDE-4) Inhibitors		
roflumilast	Р	 Initial Criteria (6-month duration): Diagnosis of COPD associated with chronic bronchitis, AND Patient has forced expiratory volume in 1 second [FEV1] < 50%; AND Patient is currently receiving standard of care COPD treatments, unless contraindicated (short acting β agonists OR short acting anticholinergics PLUS long acting β agonists OR long-acting anticholinergics), AND Patient has a history of continued COPD exacerbations on their current COPD treatment regimen Renewal Criteria Positive clinical response to treatment (e.g., improvement in FEV1 from baseline, reduction in COPD exacerbations); AND Patient is currently receiving standard of care COPD treatments, unless contraindicated (short acting β agonists OR short acting anticholinergics PLUS long acting β agonists OR long-acting anticholinergics) 	250 mcg: 28/year 500 mcg: 1/day	



	RESPIRATORY Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.					
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form		
Daliresp®	NP	See roflumilast prior authorization criteria; AND • Clinically valid reason why the patient cannot use the preferred generic roflumilast	250 mcg: 28/year 500 mcg: 1/day			
Ohtuvayre®	NP	 Patient is ≥ 18 years of age; AND Diagnosis of Chronic Obstructive Pulmonary Disorder (COPD); AND Submission of medical records (e.g. chart notes) that patient meets ALL the following: Bronchodilator FEV1/FVC ratio of <0.7 FEV1 % predicted of ≤ 79% Modified medical research council (mMRC) dyspnea scale score of ≥ 2; AND Trial and failure (as defined by continued symptoms, including exacerbations) of adequate treatment concomitantly with ONE of the following: A long-acting beta-agonist (LABA) + long-acting antimuscarinic (LAMA) + inhaled corticosteroid; OR A long-acting beta-agonist (LABA) and long-acting antimuscarinic (LAMA); AND Medication must be used as maintenance therapy only 	2 ampules/day			

SMOKING CESSATION AGENTS Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated. **Qty. Limits** Medication **PDL Prior Authorization Criteria PA Form Smoking Cessation Agents** 2/day; apo-varenicline Ρ 24 weeks/yr* 2/day; bupropion sustained Ρ release 24 weeks/yr* 2/day; Chantix® Р 24 weeks/yr* nicotine polacrilex Р 24 weeks/yr* gum nicotine polacrilex Ρ 24 weeks/yr* lozenge General PA nicotine transdermal Form Ρ 24 weeks/yr* patch 2/day; Varenicline Р 24 weeks/yr* 24 weeks/yr* Nicotrol® inhaler NP Nicotrol® nasal spray 24 weeks/yr* NP 2/day; Zyban® NP 24 weeks/yr*

VITAMINS/ELECTROLYTES Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.								
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form				
Folic Acid Preparations								
L-methylfolate	NP	Patient has documented methylenetetrahydrofolate reductase (MTHFR) mutation/deficiency						
Potassium Depletors								
Lokelma®	NP	 Initial Criteria: Patient must be ≥ 18 years of age; AND Patient has a diagnosis of chronic hyperkalemia; AND One of the following: Trial and failure, contraindication, or intolerance to a loop or thiazide diuretic; OR Trial and failure, contraindication, or intolerance to a preferred potassium deplete agent Renewal Criteria: Patient has a positive clinical response to therapy [e.g., decreased serum potassium levels, levels within normal limits) 	1/day	General PA Form				



* For children, larger quantities may be approved as medically necessary.

VITAMINS/ELECTROLYTES								
Approval of NP agents requires trial and failure, contraindication, or intolerance of 2 preferred agents, unless otherwise indicated.								
Medication	PDL	Prior Authorization Criteria	Qty. Limits	PA Form				
Veltassa®	NP	 Initial Criteria: Patient must be ≥ 12 years of age; AND Patient has a diagnosis of chronic hyperkalemia; AND One of the following: Trial and failure, contraindication, or intolerance to a loop or thiazide diuretic; OR Trial and failure, contraindication, or intolerance to a preferred potassium deplete agent Renewal Criteria: Patient has a positive clinical response to therapy [e.g., decreased serum potassium levels, levels within normal limits) 	1 packet/day					
		Vitamin B Products		•				
cyanocobalamin injection	Р	 Diagnosis of Pernicious Anemia; AND Product is being administered by the patient, patient's caregiver, or in a long-term care facility NOTE: If the medication is being administered in the prescriber's office OR by a Home Health Nurse, coverage must be obtained through the patient's MCO. 						
cyanocobalamin nasal spray	Р	Diagnosis of one of the following: Pernicious Anemia B12 deficiency; AND Provider must submit lab documentation confirming deficiency		General PA Form				
hydroxocobalamin injection	Р	See cyanocobalamin injection prior authorization criteria						
cyanocobalamin, <u>OTC</u>	Р	 Will be approved for patients who meet the following criteria: Diagnosis of Pernicious Anemia Patient must be UNDER 21 years old (not a covered benefit for adults) Diagnosis of B12 deficiency Patient must be UNDER 21 years old (not a covered benefit for adults) Provider must submit lab documentation confirming deficiency 						
Nascobal [®] nasal spray	NP							
Vitamin K Products								
phytonadione	Р		5/Rx					

