# NYRx, the New York Medicaid Pharmacy Program

#### **OVERVIEW OF CONTENTS**

Beginning Summer 2025, NYRx will accept electronic PA requests via CoverMyMeds®. For more info, reach out to NYRxEO@primetherapeutics.com.

#### Preferred Drug Program (PDP) (Pages 4–56)

The PDP promotes the use of less expensive, equally effective drugs when medically appropriate through a Preferred Drug List (PDL). All drugs currently covered by NYRx, the Medicaid Pharmacy Program, remain available under the PDP and the determination of preferred and non-preferred drugs does not prohibit a prescriber from obtaining any of the medications covered under Medicaid.

- Non-preferred drugs in these classes require prior authorization (PA), unless indicated otherwise.
- Preferred drugs that require prior authorization are indicated by footnote.
- Specific Clinical, Frequency/Quantity/Duration, Step Therapy criteria are listed in column at the right.

**Note:** Not all drugs covered by NYRx are subject to programs included in this document. For a complete list of drugs covered by NYRx see the <u>Medicaid Pharmacy List of Reimbursable Drugs</u>

#### **Clinical Drug Review Program (CDRP)** (Page 57)

The CDRP is aimed at ensuring specific drugs are utilized in a medically appropriate manner. Under the CDRP, certain drugs require prior authorization because there may be specific safety issues, public health concerns, the potential for fraud and abuse, or the potential for significant overuse and misuse.

#### Drug Utilization Review (DUR) Program (Pages 58-69)

The DUR helps to ensure that prescriptions for outpatient drugs are appropriate, medically necessary, and not likely to result in adverse medical consequences. This program uses professional medical protocols and computer technology and claims processing to assist in the management of data regarding the prescribing and dispensing of prescriptions. Frequency/Quantity/Duration (F/Q/D) Program and Step Therapy parameters are implemented to ensure clinically appropriate and cost-effective use of these drugs and drug classes.

#### Medication Assisted Treatment (MAT) Formulary (Page 70)

Prior authorization will not be required for medications used for the treatment of substance use disorder prescribed according to generally accepted national professional guidelines for the treatment of a substance use disorder.

#### Brand Less Than Generic (BLTG) Program (Pages 71–72)

The Brand Less Than Generic Program is a cost containment initiative which promotes the use of certain multi-source brand name drugs when the cost of the brand name drug is less expensive than the generic equivalent. This program is in conformance with State Education Law, which intends that patients receive the lower cost alternative.

For more information on NYRx, the Medicaid Pharmacy Program: <a href="http://www.health.ny.gov/health\_care/medicaid/program/pharmacy.htm">http://www.health.ny.gov/health\_care/medicaid/program/pharmacy.htm</a> To contact the NYRx Clinical Call Center please call 1-877-309-9493 To download a copy of the Prior Authorization fax form go to <a href="https://newyork.fhsc.com/providers/PA">https://newyork.fhsc.com/providers/PA</a> forms.asp

Disclaimer: Branded generics are included with the single generic name listing; they are not listed as separate agents.

#### Mandatory Generic Drug Program (Page 73)

State law excludes Medicaid coverage of brand name drugs that have a Federal Food and Drug Administration (FDA) approved A-rated generic equivalent unless a prior authorization is obtained. Drugs subject to the Preferred Drug Program (PDP), Clinical Drug Review Program (CDRP), and/or the Brand Less Than Generic (BLTG) Program are not subject to the Mandatory Generic Program.

#### **Dose Optimization Program** (Pages 74–78)

Dose optimization can reduce prescription costs by reducing the number of pills a patient needs to take each day. The Department has identified drugs to be included in this program, the majority of which have FDA approval for once-a-day dosing, have multiple strengths available in correlating increments at similar costs and are currently being utilized above the recommended dosing frequency.

# NYRx, the Medicaid Pharmacy Program Preferred Drug List PREFERRED DRUG LIST – TABLE OF CONTENTS

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Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	I. Anal	gesics
	Non-Steroidal Anti-Inflar	nmatory Drugs (NSAIDS)
Celebrex <sup>®</sup> celecoxib diclofenac 1% topical gel diclofenac sodium oral ibuprofen Rx tablet, suspension indomethacin capsule ketorolac meloxicam tablet nabumetone naproxen tablet piroxicam sulindac	Arthrotec <sup>®</sup> Daypro <sup>®</sup> diclofenac epolamine patch diclofenac capsule diclofenac potassium diclofenac potassium (gen Cambia <sup>®</sup> ) diclofenac sodium ER diclofenac topical soln diflunisal Dolobid Elyxyb <sup>™</sup> F/Q/D etodolac etodolac ER Feldene <sup>®</sup> fenoprofen Fenopron <sup>™</sup> flurbiprofen ibuprofen/famotidine (gen Duexis <sup>®</sup> ) indomethacin ER indomethacin suspension ketoprofen ketoprofen ER ketorolac nasal spray (gen Sprix <sup>®</sup> ) Kiprofen <sup>™</sup> meclofenamate mefenamic acid meloxicam capsule (gen Vivlodex <sup>®</sup> ) Nalfon <sup>®</sup> Naprelan <sup>®</sup> naproxen Susp naproxen CR naproxen EC	FREQUENCY/QUANTITY/DURATION (F/Q/D) Elyxyb <sup>™</sup> (celecoxib) – 4.8 mL bottle (120 mg) maximum quantity: 9 bottles / 30 days

Standard PA fax form:

1 = Preferred as of 2/6/2025

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	I. Anal	gesics
		ong-Acting <sup>CC</sup>
buprenorphine patch fentanyl patch (12 mcg, 25 mcg, 50 mcg, 75 mcg, 100 mcg) morphine sulfate ER tablet	Belbuca <sup>®</sup> Butrans <sup>®</sup> ConZip <sup>® ST</sup> fentanyl patch (37.5 mcg, 62.5 mcg, 87.5 mcg) hydrocodone ER hydrocodone ER (gen Hysingla ER) hydromorphone ER Hysingla <sup>®</sup> ER morphine ER capsule (gen Avinza) morphine ER capsule (gen Kadian) MS Contin <sup>®</sup> oxycodone ER Oxycontin <sup>®</sup> oxymorphone ER tramadol ER <sup>ST</sup>	<ul> <li>CLINICAL CRITERIA (CC) *</li> <li>Limited to a total of 4 opioid prescriptions every 30 days; Exemption for diagnosis of cancer, hospice or palliative care, or sickle cell disease</li> <li>PA required for initiation of opioid therapy for patients on established opioid dependence therapy</li> <li>PA required for use if ≥ 90 MME (MME = morphine milligram equivalents) of opioid per day for management of non-acute pain (pain lasting &gt; 7 days)</li> <li>PA required for any additional long-acting opioid therapy in opioid-naïve patients.</li> <li>PA required for any additional long-acting opioid prescription for patients currently on long-acting opioid therapy.</li> <li>PA required for initiation of opioid therapy in patients currently on benzodiazepine therapy</li> <li>PA required for any codeine- or tramadol-containing products in pts &lt; 12 years</li> <li>STEP THERAPY (ST)</li> <li>Tramadol ER (tramadol naïve patients): Attempt treatment with IR formulations before the following ER formulations: ConZip<sup>®</sup>, tramadol ER</li> <li>*Exemption from requirements for diagnosis of cancer, sickle cell disease, or hospice or palliative care.</li> </ul>

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	I. Ana	Igesics
	Opioids – Sl	hort-Acting <sup>CC</sup>
butalbital/APAP/caffeine/codeine codeine codeine/APAP hydrocodone/APAP hydrocodone/ibuprofen hydromorphone tablets morphine IR oxycodone IR tablets, solution oxycodone/APAP tramadol tablet	butalbital compound/codeine butorphanol nasal spray dihydrocodeine/APAP/caffeine Dilaudid <sup>®</sup> hydromorphone solution levorphanol meperidine Nalocet <sup>®</sup> oxycodone IR capsules, concentrate oxycodone/APAP (Prolate) solution, tablets oxymorphone pentazocine/naloxone Percocet <sup>®</sup> RoxyBond Roxicodone <sup>®</sup> Seglentis <sup>®</sup> tramadol solution tramadol 25mg, 75mg tablet tramadol/APAP	<ul> <li>CLINICAL CRITERIA (CC) * <ul> <li>Limited to a total of 4 opioid prescriptions every 30 days.</li> <li>Initial prescription for opioid-naïve patients limited to a 7-day supply.</li> <li>PA required for initiation of opioid therapy for patients on established opioid dependence therapy.</li> <li>PA required for use if ≥ 90 MME of opioid per day for management of non-acute pain (&gt; 7 days) <ul> <li>Exception for diagnosis of cancer or sickle cell disease, or hospice or palliative care programs</li> </ul> </li> <li>PA is required for opioid-naïve patients for prescription requests ≥ 50 MME per day.</li> <li>PA required for continuation of opioid therapy beyond an initial 7-day supply in patients established on gabapentin or pregabalin</li> <li>PA required for any codeine- or tramadol-containing products in pts &lt; 12 years</li> </ul> </li> <li>PA required for continuation of opioid therapy for &gt;7days for patients on established CNS stimulant therapy</li> <li>STEP THERAPY (ST)</li> <li>For Non-opioid Pain management alternatives please visit: https://health.ny.gov/health_care/medicaid/program/opioid_management.pdf</li> </ul>

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	II. Anti-li	nfectives
	Antibiotics –	nhaled <sup>CC, F/Q/D</sup>
Bethkis <sup>® <u>BLTG</u> Cayston<sup>®</sup> Kitabis<sup>®</sup> Pak <sup>BLTG</sup> TOBI Podhaler™ tobramycin (gen TOBI<sup>®</sup>) solution</sup>	TOBI <sup>®</sup> solution tobramycin (gen Bethkis <sup>®</sup> , Kitabis <sup>®</sup> ) solution	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>Confirm diagnosis of FDA-approved or compendia-supported indication</li> <li>FREQUENCY/QUANTITY/DURATION (F/Q/D)</li> <li>Aztreonam (Cayston) <ul> <li>3 ampules (3 mL) per day</li> <li>84 ampules (84 mL) per 56-day regimen (28 days on, 28 days off)</li> </ul> </li> <li>Tobramycin inhalation solution (Bethkis, TOBI, Kitabis Pak) <ul> <li>2 ampules (8 mL Bethkis, 10 mL TOBI, Kitabis Pak) per day</li> <li>56 ampules (224 mL Bethkis, 280 mL TOBI, Kitabis Pak) per 56-day regimen (28 days on-28 days off)</li> </ul> </li> <li>Tobramycin capsules with inhalation powder (TOBI Podhaler) <ul> <li>8 capsules per day 224 capsules per 56-day regimen (28 days on-28 days off)</li> </ul> </li> </ul>
	Anti-Fungals – Oral	for Onychomycosis
griseofulvin suspension, ultramicronized terbinafine tablet	griseofulvin tablet itraconazole itraconazole solution (gen Sporanox) Sporanox <sup>®</sup>	
		als – Oral
acyclovir valacyclovir	famciclovir Valtrex®	
Cephalosporins – Third Generation		
cefdinir	cefixime cefpodoxime	
Cipro® augpopoion BLTG	Fluoroquino Baxdela®	olones – Oral
Cipro <sup>®</sup> suspension <sup>BLTG</sup> ciprofloxacin tablet	Cipro <sup>®</sup> tablet	
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2 = Non-Preferred as of 2/6/2025

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Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	II. Anti-Ir	nfectives
levofloxacin tablet	ciprofloxacin suspension levofloxacin solution moxifloxacin ofloxacin tablet	
	Hepatitis	B Agents
adefovir dipivoxil Baraclude <sup>®</sup> solution entecavir Iamivudine HBV	Baraclude <sup>®</sup> tablet Vemlidy <sup>®</sup>	
	Hepatitis C Agents – D	Direct Acting Antivirals
Mavyret <sup>™</sup> ribavirin sofosbuvir/velpatasvir (gen Epclusa <sup>®</sup> ) Vosevi <sup>®</sup>	Epclusa <sup>®</sup> Harvoni <sup>®</sup> ledipasvir/sofosbuvir (gen Harvoni <sup>®</sup> ) Sovaldi <sup>®</sup> Zepatier <sup>®</sup>	
	Tetrac	yclines
demeclocycline doxycycline hyclate minocycline capsule tetracycline capsule	Doryx <sup>® ST</sup> Doryx MPC <sup>® ST</sup> doxycycline hyclate DR <sup>ST</sup> doxycycline monohydrate minocycline tablet minocycline ER tablet Nuzyra <sup>™</sup> Solodyn <sup>®</sup> tetracycline tablet	<ul> <li>STEP THERAPY (ST)</li> <li>Trial of doxycycline IR before progressing to doxycycline DR</li> </ul>

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	III. Cardio	ovascular
	Angiotensin Converting E	Enzyme Inhibitors (ACEIs)
benazepril	Accupril®	
enalapril	Altace®	
lisinopril	captopril	
ramipril	enalapril (gen Epaned®)	
lampin	Epaned®	
	fosinopril	
	Lotensin <sup>®</sup>	
	moexipril	
	perindopril	
	Qbrelis™	
	quinapril	
	trandolapril	
	Vasotec®	
	Zestril <sup>®</sup>	
		Combinations
benazepril/amlodipine	Accuretic®	
benazepril/HCTZ	fosinopril/HCTZ	
captopril/HCTZ	Lotensin HCT <sup>®</sup>	
enalapril/HCTZ	quinapril/HCTZ	
lisinopril/HCTZ	Vaseretic <sup>®</sup>	
Lotrel®	Zestoretic <sup>®</sup>	
trandolapril/verapamil ER		
	Angiotensin Recep	tor Blockers (ARBs)
irbesartan	Atacand®	DOSE OPTIMIZATION (DO)
losartan	Avapro®	<ul> <li>See Dose Optimization Chart for affected drugs and strengths</li> </ul>
olmesartan	Benicar <sup>® <u>DO</u></sup>	
telmisartan	candesartan	
valsartan tablet	Cozaar®	
	Diovan <sup>® <u>DO</u></sup>	
	Edarbi <sup>®</sup>	
	eprosartan	
	Micardis <sup>® <u>DO</u></sup>	

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Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	III. Cardio	ovascular
	valsartan solution	
	Antianginals and	d Anti-Ischemics
ranolazine	Aspruzyo Sprinkle™	
	ARBs Con	nbinations
Entresto <sup>®</sup> Exforge HCT <sup>®</sup> irbesartan/HCTZ losartan/HCTZ olmesartan/amlodipine olmesartan/HCTZ telmisartan/HCTZ valsartan/amlodipine valsartan/amlodipine/HCTZ valsartan/HCTZ	Atacand HCT <sup>®</sup> Avalide <sup>®</sup> Azor <sup>®</sup> Benicar HCT <sup>® DO</sup> candesartan/HCTZ Diovan HCT <sup>® DO</sup> Edarbyclor <sup>® DO</sup> Entresto <sup>®</sup> Sprinkle Exforge <sup>® DO</sup> Hyzaar <sup>®</sup> Micardis HCT <sup>® DO</sup> olmesartan/amlodipine/HCTZ telmisartan/amlodipine Tribenzor <sup>®</sup>	<ul> <li>DOSE OPTIMIZATION (DO)</li> <li>See Dose Optimization Chart for affected drugs and strengths</li> </ul>
	Beta B	lockers
atenolol carvedilol labetalol metoprolol succ. XL metoprolol tartrate propranolol tablet propranolol ER	acebutolol betaxolol bisoprolol Bystolic <sup>®</sup> <sup>DO</sup> carvedilol ER Inderal LA <sup>®</sup> Inderal XL <sup>®</sup> InnoPran XL <sup>®</sup> Kapspargo <sup>™</sup> Sprinkle Lopressor <sup>®</sup> nadolol <sup>DO</sup> nebivolol (gen Bystolic <sup>®</sup> ) pindolol propranolol solution	<ul> <li>DOSE OPTIMIZATION (DO)</li> <li>See Dose Optimization Chart for affected drugs and strengths</li> </ul>
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Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	III. Cardiova	ascular
	Tenormin <sup>®</sup> timolol Toprol XL <sup>® <u>DO</u></sup>	
	Beta Blockers	/ Diuretics
atenolol/chlorthalidone bisoprolol/HCTZ propranolol/HCTZ	metoprolol tartrate/ HCTZ Tenoretic <sup>®</sup>	
	Calcium Channel Blocker	rs (Dihydropyridine)
amlodipine felodipine ER isradipine nicardipine HCI nifedipine nifedipine ER/SA	Katerzia <sup>™</sup> levamlodipine nisoldipine Norliqva <sup>®</sup> Norvasc <sup>®</sup> Procardia XL <sup>®</sup> Sular <sup>®</sup>	
	Cholesterol Absorp	otion Inhibitors
cholestyramine cholestyramine light Colestid <sup>®</sup> tablet colestipol tablet ezetimibe	colesevelam Colestid granules, packet colestipol granules, packet Questran <sup>®</sup> Questran Light <sup>®</sup> Welchol <sup>®</sup> Zetia <sup>®</sup>	
	HMG-CoA Reductase I	nhibitors (Statins)
atorvastatin lovastatin pravastatin rosuvastatin simvastatin	Altoprev <sup>®</sup> Atorvaliq <sup>®</sup> atorvastatin/amlodipine Caduet <sup>®</sup> Ezallor™ Sprinkle ezetimibe/simvastatin FloLipid™ fluvastatin fluvastatin ER	
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Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	III. Cardio	ovascular
	Lescol XL <sup>®</sup> Lipitor <sup>®</sup> Livalo <sup>®</sup> pitavastatin (gen Livalo <sup>®</sup> ) Vytorin <sup>®</sup> Zocor <sup>®</sup> Zypitamag™	
	Phosphodiesterase Type-5 (	PDE-5) Inhibitors for PAH <sup>CC</sup>
sildenafil tadalafil	Adcirca <sup>®</sup> Opsynvi <sup>®</sup> Revatio <sup>®</sup> Tadliq <sup>®</sup>	<ul> <li>CLINICAL CRITERIA</li> <li>Prior authorization is required for all drugs in this class</li> <li>Prescribers or their authorized agents are required to respond to a series of questions that identify prescriber, patient, and reason for prescribing drug</li> <li>Please be prepared to fax clinical documentation upon request</li> <li>Prescriptions can be written for a 30-day supply with up to 11 refills</li> </ul>
	Pulmonary Arterial Hypertensi	ion (PAH) Agents, Other – Oral
ambrisentan (gen Letairis) bosentan tablets (gen Tracleer <sup>®</sup> )	Adempas <sup>®</sup> Letairis <sup>®</sup> Opsumit <sup>®</sup> Orenitram <sup>®</sup> ER tablet, dosepack Tracleer <sup>®</sup> tablet for suspension, tablet Uptravi <sup>®</sup>	
	Triglyceride Lo	owering Agents
fenofibrate tablet (gen Tricor <sup>®</sup> ) fenofibrate capsule (gen Lofibra <sup>®</sup> ) fenofibric acid capsule (gen Trilipix <sup>®</sup> ) gemfibrozil icosapent <sup>F/Q/D</sup> omega-3-acid ethyl esters (gen Lovaza <sup>®</sup> ) <sup>F/Q/D</sup>	fenofibrate caps (gen Lipofen <sup>®</sup> ) fenofibrate micronized capsule fenofibrate tablet (gen Fenoglide <sup>®</sup> ) fenofibric acid tablet (gen Fibricor <sup>®</sup> ) Fenoglide <sup>®</sup> Fibricor <sup>®</sup> Lipofen <sup>®</sup> Lopid <sup>®</sup> Tricor <sup>®</sup>	<ul> <li>FREQUENCY/QUANTITY/DURATION (F/Q/D)</li> <li>omega-3-acid ethyl esters (gen Lovaza<sup>®</sup>) and icosapent ethyl – Required dosage equal to 4 grams per day</li> </ul>

2 = Non-Preferred as of 2/6/2025

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
III. Cardiovascular		
	Trilipix®	

	Non-Preferred Drugs	Prior Authorization/Coverage Parameters	
IV. Central Nervous System			
	Alzheime	r's Agents	
donepezil 5 mg, 10 mg, ODT galantamine galantamine ER memantine Namenda <sup>®</sup> rivastigmine	Adlarity <sup>®</sup> Aricept <sup>®</sup> donepezil 23 mg Exelon <sup>®</sup> memantine ER memantine-donepezil ER (gen Namzaric <sup>®</sup> ) Namenda XR <sup>®</sup> Namzaric <sup>®</sup> Zunveyl <sup>®</sup> DR		
		oamazepine Derivatives	
carbamazepine chewable Carbatrol <sup>®</sup> <sup>BLTG</sup> Equetro <sup>®</sup> oxcarbazepine tablet Oxtellar XR <sup>®</sup> <u>DO</u> , <u>BLTG</u> Tegretol <sup>®</sup> suspension, tablet <u>BLTG</u> Tegretol XR <sup>®</sup> <u>BLTG</u> Trileptal <sup>®</sup> suspension <u>BLTG</u>	Aptiom <sup>® CC, DO</sup> carbamazepine suspension carbamazepine tablet carbamazepine ER capsule carbamazepine XR tablet oxcarbazepine suspension oxcarbazepine ER (gen Oxtellar XR <sup>®</sup> ) Trileptal <sup>®</sup> tablet <sup>CC</sup>	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>Clinical editing will allow patients currently stabilized on a non-preferred agent to continue to receive that agent without PA</li> <li>DOSE OPTIMIZATION (DO)</li> <li>See Dose Optimization Chart for affected drugs and strengths</li> </ul>	
		ants - Nasal	
Nayzilam <sup>®</sup> Valtoco <sup>®</sup>			
	Anticonvuls	ants – Other	
clobazam tablet <sup>ST, CC</sup> gabapentin capsule, solution, tablet F/Q/D, CC acosamide tablet, solution amotrigine tablet, chew evetiracetam evetiracetam ER	Banzel <sup>®</sup> Briviact <sup>®</sup> clobazam suspension <sup>ST</sup> Diacomit <sup>® CC</sup> Elepsia <sup>®</sup> XR Epidiolex <sup>® CC</sup> Eprontia <sup>™ CC, F/Q/D</sup>	<ul> <li>DOSE OPTIMIZATION (DO)</li> <li>See Dose Optimization Chart for affected drugs and strengths CLINICAL CRITERIA (CC)</li> <li>Clinical editing will allow patients currently stabilized on a non- preferred agent to continue to receive that agent without PA</li> </ul>	

	, ,				
Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters			
IV. Central Nervous System					
Lyrica® capsule <u>PQ</u> , F/Q/D, CC pregabalin capsule F/Q/D, CC tiagabine topiramate <sup>CC</sup> zonisamide	felbamate Felbatol® Fintepla® Fycompa® DO Keppra® Keppra XR® Lamictal® tablet, chew, dosepak Lamictal® ODT tablet, dosepak Lamictal® XR DO tablet, dosepak Lamotrigine dosepak lamotrigine ODT dosepak levetiracetam 250mg tablet for suspension (gen Spritam®) Lyrica® solution F/Q/D, CC Lyrica® CR F/Q/D, CC Motpoly XR Neurontin® F/Q/D, CC Onfi® ST, CC pregabalin solution F/Q/D, CC pregabalin ER (gen Lyrica® CR) F/Q/D, CC Qudexy® XR CC, DO rufinamide (gen Banzel®) Sabril® Sympazan® film ST, CC Topamax® CC topiramate 50mg Sprinkle CC topiramate ER CC, DO (gen Qudexy® XR) topiramate ER CC, DO (gen Trokendi XR®)	<ul> <li>Cannabidiol extract (Epidiolex<sup>®</sup>) – Confirm diagnosis of FDA-approved or compendia-supported indication, or; Institutional Review Board (IRB) approval with signed consent form</li> <li>Lyrica<sup>®</sup>/Lyrica<sup>®</sup> CR (pregabalin/pregabalin ER) – PA required for the initiation of pregabalin at &gt; 150 mg per day in patients currently on an opioid at &gt; 50 MME per day</li> <li>Neurontin<sup>®</sup> (gabapentin) – PA required for initiation of gabapentin at &gt; 900 mg per day in patients currently on an opioid at &gt; 50 MME per day</li> <li>Stiripentol (Diacomit<sup>®</sup>) – Require diagnosis of FDA-approved or compendia-supported indication, or; Institutional Review Board (IRB) approval with signed consent form</li> <li>Topiramate IR/ER (Eprontia™, Qudexy<sup>®</sup> XR, Topamax<sup>®</sup>, Trokendi XR™) – Require confirmation of FDA-approved, compendia-supported, or Medicaid covered diagnosis</li> <li>Onfi<sup>®</sup>/Sympazan<sup>®</sup> (clobazam):         <ul> <li>Require confirmation of clobazam therapy in patients currently on opioid or oral buprenorphine therapy</li> <li>PA required for initiation of clobazam therapy in patients currently on benzodiazepine therapy</li> <li>PA required for any clobazam prescription in patients currently on benzodiazepine therapy</li> <li>PA required for R (pregabalin/pregabalin ER) – Maximum daily dose of IR: 600 mg per day, and ER: 660 mg per day</li> </ul> </li> <li>Neurontin<sup>®</sup> (gabapentin) – Maximum daily dose of 3,600 mg per day</li> <li>Neurontin<sup>®</sup> (clobazam) – Requires a trial with an SSRI or SNRI for treatment of anxiety</li> </ul>			
= Preferred as of 2/6/2025	Trokendi XR <sup>® CC, DO</sup> Standard PA fax form:				

vigabatrin Vigafyde™ Vimpat <sup>®</sup> Xcopri <sup>®</sup> Zonisade™ Ztalmy <sup>®</sup>	Agents, Other <sup>F/Q/D</sup> CLINICAL CRITERIA (CC)	
vigabatrin Vigafyde™ Vimpat <sup>®</sup> Xcopri <sup>®</sup> Zonisade™ Ztalmy <sup>®</sup> <b>Antimigraine</b> Qulipta™ Reyvow™ ST	Agents, Other <sup>F/Q/D</sup> CLINICAL CRITERIA (CC) • Confirm diagnosis of FDA-app indication STEP THERAPY (ST) Acute treatment of migraine	
Vigafyde™ Vimpat <sup>®</sup> Xcopri <sup>®</sup> Zonisade™ Ztalmy <sup>®</sup> <b>Antimigraine</b> Qulipta™ Reyvow™ ST	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>Confirm diagnosis of FDA-app indication</li> <li>STEP THERAPY (ST) Acute treatment of migraine</li> </ul>	
Emgality <sup>®</sup> 100mg syringe Qulipta™ Reyvow™ <sup>ST</sup>	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>Confirm diagnosis of FDA-app indication</li> <li>STEP THERAPY (ST) Acute treatment of migraine</li> </ul>	
Qulipta™ Reyvow™ <sup>ST</sup>	<ul> <li>Confirm diagnosis of FDA-app indication</li> <li>STEP THERAPY (ST) Acute treatment of migraine</li> </ul>	
		The Antimigranie Agents- mptan Cla
	Agent	F/Q/D
	Aimovig	1 syringe/30 days
	Emgality 120 mg	2 syringes/30 days
	Emgality 100 mg	3 syringes/30 days
	Ajovy	3 syringes/90 days
	Reyvow	8 units/30 days
	Ubrelvy	16 units/30 days
	Nurtec™ ODT	24 units/40 days
	Qulipta	30 units/30 days
	Zavzpret <sup>®</sup>	8 units/30 days
	Agents – Triptans	
	FREQUENCY/QUANTITY/DURA	TION (F/Q/D)
eletriptan <sup>F/Q/D</sup>	Agent	F/Q/D
frovatriptan <sup>F/Q/D</sup> Imitrex <sup>® F/Q/D</sup>	almotriptan eletriptan (Relpax <sup>®</sup> ) frovatriptan (Frova <sup>®</sup> )	18 units/30 days
	almotriptan <sup>F/Q/D</sup> eletriptan <sup>F/Q/D</sup> Frova <sup>® F/Q/D</sup> frovatriptan <sup>F/Q/D</sup> Imitrex <sup>® F/Q/D</sup> Maxalt <sup>® F/Q/D</sup> Standard PA fax form:	Agent         Aimovig         Emgality 120 mg         Emgality 120 mg         Emgality 100 mg         Ajovy         Reyvow         Ubrelvy         Nurtec™ ODT         Qulipta         Zavzpret®         Almotriptan F <sup>/Q/D</sup> Frova® F <sup>/Q/D</sup> Frova® F <sup>/Q/D</sup> Imitrex® F <sup>/Q/D</sup> Maxalt® F <sup>/Q/D</sup> Maxalt® F <sup>/Q/D</sup>

2 = Non-Preferred as of 2/6/2025

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Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters		
IV. Central Nervous System				
	Maxalt <sup>®</sup> MLT <sup>F/Q/D</sup> naratriptan <sup>F/Q/D</sup> Relpax <sup>® F/Q/D</sup> sumatriptan-naproxen <sup>F/Q/D</sup> Tosymra <sup>™ F/Q/D</sup> Zembrace <sup>™</sup> SymTouch <sup>™</sup> zolmitriptan <sup>F/Q/D</sup> Zomig <sup>® F/Q/D</sup>	naratriptanrizatriptan (Maxalt®)rizatriptan (Maxalt® MLT)sumatriptan nasal spray (Imitrex®)sumatriptan (Imitrex®)sumatriptan-naproxenTosymra™ nasal sprayzolmitriptan (Zomig®)zolmitriptan nasal spray (Zomig®)		
	Antipsychotic	s – Injectable		
Abilify Asimtufii <sup>®</sup> Abilify Maintena <sup>®</sup> Aristada <sup>®</sup> Aristada Initio <sup>®</sup> fluphenazine decanoate Haldol <sup>®</sup> decanoate haloperidol decanoate Invega Hafyera <sup>™</sup> Invega Sustenna <sup>®</sup> Invega Trinza <sup>®</sup> Perseris <sup>™</sup> Risperdal Consta <sup>® <u>BLTG</u> Uzedy<sup>™</sup> Zyprexa Relprevv<sup>®</sup></sup>	Erzofri <sup>®</sup> risperidone injection (gen Risperdal Consta <sup>®</sup> ) Rykindo <sup>®</sup>			
	Antipsychotics – Sec	ond Generation <sup>CC, ST</sup>		
aripiprazole tablet <sup>DO</sup> asenapine (gen Saphris <sup>®</sup> ) clozapine lurasidone (gen Latuda <sup>®</sup> ) olanzapine tablet <sup>DO</sup> paliperidone ER <sup>DO</sup> quetiapine <sup>F/Q/D</sup> quetiapine ER <sup>F/Q/D</sup> , <u>DO</u>	Abilify <sup>®</sup> tablet <sup>DO</sup> Abilify MyCite <sup>®</sup> aripiprazole solution aripiprazole ODT Caplyta <sup>™</sup> clozapine ODT Clozaril <sup>®</sup> Cobenfy <sup>™</sup> capsules, starter pack	<ul> <li>DOSE OPTIMIZATION (DO)</li> <li>See Dose Optimization Chart for affected drugs and strengths</li> <li>CLINICAL CRITERIA (CC)</li> </ul>		

Standard PA fax form:

1 = Preferred as of 2/6/2025

Preferred Drugs	Non-Preferred Drugs		Prior Authorization/Coverage Pa	arameters
	IV. Central Ne	ous System		
risperidone capsule	Fanapt <sup>®</sup> Geodon <sup>®</sup> Invega <sup>®</sup> D <sup>Q</sup> Latuda <sup>®</sup> D <sup>Q</sup> Lybalvi <sup>®</sup> Nuplazid <sup>®</sup> olanzapine ODT D <sup>Q</sup> olanzapine / fluoxetine Opipza <sup>™</sup> Rexulti <sup>®</sup> D <sup>Q</sup> Risperdal <sup>®</sup> Saphris <sup>®</sup> Secuado <sup>®</sup> Seroquel XR <sup>®</sup> D <sup>Q</sup> , F/Q/D Versacloz <sup>®</sup> Vraylar <sup>®</sup> D <sup>Q</sup> Zyprexa <sup>®</sup> D <sup>Q</sup> Zyprexa <sup>®</sup> Zydis	•	Confirm diagnosis of FDA-approved or comperindication Clinical editing will allow patients currently state preferred agent to continue to receive that age Prior authorization is required when an oral SC highest MDD according to FDA labeling. Prior authorization is required for patients less when there is concurrent use of 2 or more different antipsychotics for greater than 90 days. Prior authorization is required for patients 21 y when 3 or more different oral second-generation used for more than 180 days. PA is required for initial prescription for beneficd drug-specific minimum age as indicated below aripiprazole (Abilify®, Opipza™) aripiprazole (Abilify®, Opipza™) asenapine (Saphris®) asenapine (Secuado®) brexpiprazole (Rexulti®) cariprazine (Vraylar®) clozapine (Clozaril®, Versacloz®) iloperidone (Fanapt®) lumateperone (Caplyta™) lurasidone HCI (Latuda®) olanzapine / fluoxetine (Symbyax®) olanzapine / samidorphan (Lybalvi®) paliperidone ER (Invega®) pimavanserin (Nuplazid®) quetiapine fum. (Seroquel®, Seroquel XR®) risperidone (Risperdal®) xanomeline-trospium (Cobenfy™) ziprasidone HCI (Geodon®)	pilized on a non- ent without PA GA is utilized above the than 21 years of age erent oral rears of age or older on antipsychotics are ciaries younger than the

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Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters			
	IV. Central Nervous System				
		<ul> <li>Require confirmation of diagnosis that supports the concurrent use of a Second Generation Antipsychotic and a CNS Stimulant for patients &lt; 18 years of age</li> </ul>			
		<ul> <li>STEP THERAPY (ST)</li> <li>For all Second Generation Antipsychotics used in the treatment of Major Depressive Disorder in the absence of other psychiatric comorbidities, trial with at least two different antidepressant agents is required</li> <li>olanzapine / fluoxetine: When prescribing for the treatment of major depressive disorder (MDD) in the absence of other psychiatric comorbidities, trial with at least one different antidepressant agent is required</li> <li>FREQUENCY/QUANTITY/DURATION (F/Q/D)</li> <li>quetiapine/quetiapine ER (Seroquel®/Seroquel XR®): Minimum 50 mg/day</li> <li>quetiapine (Seroquel®): Maximum 3 units per day, 90 units per 30 days</li> <li>quetiapine ER (Seroquel XR®): 50mg, maximum 2 units/day, 60 units/30 days</li> </ul>			
	Central Nervous System	n (CNS) Stimulants <sup>CC, F/Q/D</sup>			
amphetamine salt combo IR (gen Adderall <sup>®</sup> ) amphetamine salt combo ER (gen Adderall XR <sup>®</sup> ) <sup>DO</sup> Daytrana <sup>® BLTG</sup> dexmethylphenidate (gen Focalin <sup>®</sup> ) dexmethylphenidate ER <sup>DO</sup> (gen Focalin XR <sup>®</sup> ) dextroamphetamine tablet lisdexamfetamine chewable tablet (gen Vyvanse® chew tablet) methylphenidate solution (gen Methylin <sup>®</sup> )	Adderall XR <sup>®</sup> DO Adzenys XR-ODT <sup>®</sup> amphetamine (gen Adzenys ER <sup>®</sup> ) amphetamine (gen Evekeo <sup>®</sup> ) Aptensio XR <sup>®</sup> armodafinil (gen Nuvigil <sup>®</sup> ) Azstarys <sup>™</sup> Concerta <sup>®</sup> DO Cotempla <sup>®</sup> XR-ODT <sup>™</sup> Dexedrine <sup>®</sup> dextroamphetamine / amphetamine (gen Mydayis <sup>™</sup> )	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>Confirm diagnosis of FDA-approved, compendia-supported and Medicaid covered indication</li> <li>Prior authorization is required for initial prescriptions for stimulant therapy for beneficiaries less than 3 years of age</li> <li>Confirm diagnoses that support concurrent use of CNS Stimulant and Second Generation Antipsychotic agent for beneficiaries less than 18 years of age</li> <li>Patient-specific considerations for drug selection include treatment of narcolepsy, excessive daytime sleepiness, sleepiness associated with shift work sleep disorder, or sleepiness associated with obstructive sleep apnea.</li> </ul>			
1 = Preferred as of $2/6/2025$ 2 = Non-Preferred as of $2/6/2025$	Standard PA fax form: https://newyork.fhsc.com/down	loads/providers/NYRx_PDP_PA_Fax_Standardized.pdf 19			

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters			
	IV. Central Nervous System				
methylphenidate tablet (gen Ritalin <sup>®</sup> ) methylphenidate CD <sup>DQ</sup> methylphenidate ER (gen Aptensio <sup>®</sup> XR) methylphenidate ER (gen Metadate CD) methylphenidate ER (gen Ritalin LA <sup>®</sup> ) Ritalin LA <sup>®</sup> <sup>DQ</sup> Vyvanse <sup>®</sup> capsule <sup>DQ, BLTG</sup>	dextroamphetamine ER (gen Dexedrine®) dextroamphetamine solution (gen ProCentra®) dextroamphetamine tablet (gen Zenzedi®) Dyanavel XR® Evekeo® Evekeo® ODT Focalin® Focalin XR® № Jornay PM <sup>™</sup> lisdexamfetamine capsule (gen Vyvanse®) methamphetamine (gen Desoxyn®) Methylin® methylphenidate (gen Daytrana®) methylphenidate chewable tablet (gen Methylin®) methylphenidate ER 45 mg, 63 mg, 72 mg tablet modafinil (gen Provigil®) № Mydayis™ Nuvigil® ProCentra® Provigil® № QuilliChew ER <sup>™</sup> № QuilliChew ER <sup>™</sup> № Relexxii® Ritalin® Sunosi™ Vyvanse® chewable tablet Wakix® Xelstrym™ Zenzedi®	<ul> <li>PA required for initiation of CNS Stimulant for patients currently on an opioid</li> <li>PA required for initiation of CNS Stimulant for patients currently on a benzodiazepine</li> <li>DOSE OPTIMIZATION (DO)</li> <li>See Dose Optimization Chart for affected drugs and strengths</li> <li>FREQUENCY/QUANTITY/DURATION (F/Q/D)</li> <li>Quantity limits based on daily dosage as determined by FDA labeling</li> </ul>			

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters		
	IV. Central Nervous System			
	Movement Disc	order Agents <sup>CC</sup>		
Austedo <sup>®</sup> Austedo <sup>®</sup> XR Austedo <sup>®</sup> XR titration pack Ingrezza <sup>®</sup> Ingrezza <sup>®</sup> Sprinkle Ingrezza <sup>®</sup> titration pack tetrabenazine	Xenazine®	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>Confirm diagnosis for an FDA-approved or compendia-supported indication</li> </ul>		
	Multiple Scle	rosis Agents		
Avonex <sup>®</sup> Copaxone <sup>®</sup> 20 mg/mL <sup>BLTG</sup> dimethyl fumarate DR fingolimod (gen Gilenya <sup>®</sup> ) Kesimpta <sup>®</sup> teriflunomide (gen Aubagio <sup>®</sup> )	Aubagio <sup>®</sup> Bafiertam <sup>™</sup> Betaseron <sup>®</sup> Copaxone <sup>®</sup> 40 mg/mL Gilenya <sup>®</sup> glatiramer Mavenclad <sup>®</sup> Mayzent <sup>®</sup> Plegridy <sup>®</sup> Ponvory <sup>™</sup> F/Q/D Rebif <sup>®</sup> Rebif <sup>®</sup> Rebidose <sup>®</sup> Tascenso ODT <sup>™</sup> Tecfidera <sup>®</sup> Vumerity <sup>®</sup> Zeposia <sup>®</sup> <sup>CC, ST</sup>	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>Zeposia<sup>®</sup> (ozanimod): Confirm diagnosis for FDA- or compendia- supported use</li> <li>STEP THERAPY (ST)</li> <li>Zeposia<sup>®</sup> (ozanimod): For an indication of Ulcerative Colitis <ul> <li>Trial of a non-specific anti-inflammatory drug such as an aminosalicylate or immunosuppressant, or a disease-modifying anti-rheumatic drug (DMARD), and;</li> <li>Trial of a preferred systemic immunomodulator</li> </ul> </li> <li>FREQUENCY/QUANTITY/DURATION (F/Q/D)</li> <li>Ponvory™ (ponesimod) starter pack; maximum quantity is 14, no refills</li> <li>Ponvory™ (ponesimod); maintenance limited to a 30-day supply</li> </ul>		
	Non-Ergot Dopamine	e Receptor Agonists		
pramipexole ropinirole	Neupro <sup>®</sup> pramipexole ER ropinirole ER			

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Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	IV. Central Ne	ervous System
C	ther Agents for Attention Deficit	t Hyperactivity Disorder (ADHD) <sup>CC</sup>
atomoxetine <sup>DO</sup> clonidine ER guanfacine ER <sup>DO</sup>	Intuniv <sup>®</sup> <u>PO</u> Onyda <sup>™</sup> XR Qelbree <sup>™</sup> Strattera <sup>®</sup> <u>PO</u>	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>Confirm diagnosis for an FDA-approved or compendia-supported indication for beneficiaries &lt; 18 years of age.</li> <li>Prior authorization is required for initial prescriptions for non-stimulant therapy for beneficiaries less than 6 years of age</li> <li>DOSE OPTIMIZATION (DO)</li> <li>See Dose Optimization Chart for affected strengths</li> </ul>
		s/Sleep Agents <sup>F/Q/D</sup>
estazolam <sup>CC</sup> eszopiclone ramelteon (gen Rozerem <sup>®</sup> ) temazepam 15 mg, 30 mg <sup>CC</sup> zolpidem tablet <sup>CC</sup> zolpidem ER <sup>CC</sup>	Ambien <sup>® CC</sup> Ambien CR <sup>® CC</sup> Belsomra <sup>®</sup> Dayvigo <sup>™</sup> Doral <sup>® CC</sup> doxepin Edluar <sup>® CC</sup> flurazepam <sup>CC</sup> Halcion <sup>® CC</sup> Lunesta <sup>® DO</sup> quazepam <sup>CC</sup> (gen Doral <sup>®</sup> ) Quviviq <sup>™</sup> Restoril <sup>® CC</sup> Rozerem <sup>®</sup> temazepam 7.5 mg, 22.5 mg <sup>CC</sup> triazolam <sup>CC</sup> zaleplon zolpidem sublingual, capsule <sup>CC</sup>	<ul> <li>DOSE OPTIMIZATION (DO)</li> <li>See Dose Optimization Chart for affected strengths</li> <li>CLINICAL CRITERIA (CC)</li> <li>Zolpidem products: Confirm dosage is consistent with FDA labeling for initial prescriptions</li> <li>Benzodiazepine Agents (estazolam, Doral®, flurazepam, Halcion®, quazepam, Restoril®, temazepam, triazolam): <ul> <li>Confirm diagnosis of FDA-approved or compendia-supported indication</li> <li>PA required for initiation of benzodiazepine therapy in patients currently on opioid or oral buprenorphine therapy</li> <li>PA required for any additional benzodiazepine prescription in patients currently on benzodiazepine therapy</li> <li>PA required when greater than a 14-day supply of a benzodiazepine is prescribed for someone on a CNS stimulant</li> </ul> </li> <li>FREQUENCY/QUANTITY/DURATION (F/Q/D)</li> <li>Frequency and duration limits for the following products: <ul> <li>30 dosage units per fill/1 dosage unit per day/30 days</li> <li>For zaleplon-containing products: <ul> <li>60 dosage units per fill/2 dosage units per day/30 days</li> </ul> </li> </ul></li></ul>

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters		
IV. Central Nervous System				
	Selective Serotonin Re	<ul> <li>o 180 days for immediate-release zolpidem (Ambien<sup>®</sup>, Edluar<sup>®</sup>) products</li> <li>o 180 days for eszopiclone and ramelteon (Rozerem<sup>®</sup>) products</li> <li>o 180 days for lemborexant (Dayvigo<sup>™</sup>)</li> <li>o 168 days for zolpidem ER (Ambien CR<sup>®</sup>) products</li> <li>o 90 days for daridorexant (Quviviq<sup>™</sup>)</li> <li>o 90 days for suvorexant (Belsomra<sup>®</sup>)</li> <li>o 90 days for doxepin</li> <li>o 30 days for benzodiazepine agents (Doral<sup>®</sup>, estazolam, flurazepam, Halcion<sup>®</sup>, quazepam, Restoril<sup>®</sup>, temazepam, triazolam) for the treatment of insomnia</li> <li>Additional/Alternate parameters:</li> <li>For patients naïve to non-benzodiazepine sedative hypnotics (NBSH): First-fill duration and quantity limit of 10 dosage units as a 10-day supply, except for zaleplon-containing products which the quantity limit is 20 dosage units as a 10-day supply</li> </ul>		
citalopram tablet, solution	Celexa <sup>®</sup>	DOSE OPTIMIZATION (DO)		
escitalopram tablet	citalopram capsule	See Dose Optimization Chart for affected strengths		
fluoxetine capsule, solution	escitalopram solution			
paroxetine tablet sertraline tablet, concentrate vilazodone (gen Viibryd <sup>®</sup> )	fluoxetine tablet fluoxetine DR weekly fluvoxamine CC fluvoxamine ER CC Lexapro <sup>® DO</sup> paroxetine capsule paroxetine CR paroxetine suspension Paxil <sup>®</sup> Paxil CR <sup>®</sup> Prozac <sup>®</sup> sertraline capsule	<ul> <li>Clinical editing will allow patients currently stabilized on fluvoxamine or fluvoxamine ER to continue to receive that agent without PA</li> <li>Clinical editing to allow patients with a diagnosis of Obsessive-Compulsive Disorder (OCD) to receive fluvoxamine and fluvoxamine ER without prior authorization</li> </ul>		

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Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters	
	IV. Central Net	rvous System	
	Trintellix <sup>®</sup> DO Viibryd <sup>®</sup> DO Zoloft <sup>®</sup>		
	Serotonin-Norepinephrine Reuptake Inhibitors (SNRIs)		
duloxetine 20 mg, 30 mg, 60 mg (gen Cymbalta <sup>®</sup> ) venlafaxine venlafaxine ER capsule	Cymbalta <sup>®</sup> desvenlafaxine ER desvenlafaxine succinate ER <sup>DO</sup> Drizalma Sprinkle <sup>™</sup> duloxetine 40 mg Effexor XR <sup>® DO</sup> Fetzima <sup>®</sup> Pristiq <sup>® DO</sup> Savella <sup>®</sup> venlafaxine ER tablet	<ul> <li>DOSE OPTIMIZATION (DO)</li> <li>See Dose Optimization Chart for affected strengths</li> </ul>	

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters		
	V. Dermatologic Agents			
	Acne Ager	nts, Topical		
adapalene/benzoyl peroxide (gen Epiduo®) adapalene cream adapalene OTC gel Retin-A® cream <sup>CC</sup> , <u>BLTG</u> tazarotene cream <sup>CC</sup> tretinoin gel (Retin-A) <sup>CC</sup>	adapalene Rx gel, gel pump adapalene/benzoyl peroxide (gen Epiduo® Forte) Aklief® Altreno® <sup>CC</sup> Arazlo <sup>™</sup> <sup>CC</sup> Cabtreo <sup>™</sup> clindamycin/tretinoin <sup>CC</sup> dapsone Differin® cream, gel pump, lotion, OTC gel Epiduo® Forte gel pump Fabior® <sup>CC</sup> Klaron® Ovace® Plus Retin-A <sup>®</sup> gel <sup>CC</sup> Retin-A Micro® <sup>CC</sup> SSS® cream, foam sulfacetamide sulfacetam	CLINICAL CRITERIA  • Confirm diagnosis of FDA-approved, compendia-supported, and Medicaid-covered indication		

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	V. Dermatol	ogic Agents
	Actinic Kera	tosis Agents
diclofenac 3% gel fluorouracil solution fluorouracil 0.5% cream (gen Carac) fluorouracil 5% cream (gen Efudex cream) imiquimod (gen Aldara) mupirocin ointment	Carac <sup>®</sup> Efudex <sup>®</sup> imiquimod (gen Zyclara) Zyclara <sup>®</sup> Antibiotics Centany <sup>®</sup>	s – Topical
	mupirocin cream Xepi™	
	Anti-Funga	ls – Topical
ciclopirox cream, suspension, shampoo ciclopirox 8% solution clotrimazole OTC clotrimazole Rx clotrimazole/betamethasone cream ketoconazole cream ketoconazole 2% shampoo miconazole OTC nystatin cream, ointment, powder nystatin/triamcinolone terbinafine OTC tolnaftate OTC	butenafine Ciclodan <sup>®</sup> cream ciclopirox gel clotrimazole/betamethasone lotion econazole Ertaczo <sup>®</sup> Extina <sup>®</sup> Jublia <sup>®</sup> ketoconazole foam Loprox <sup>®</sup> cream, suspension luliconazole Luzu <sup>®</sup> miconazole/zinc/petrolatum (gen Vusion <sup>®</sup> ) <sup>F/Q/D</sup> naftifine Naftin <sup>®</sup> oxiconazole Oxistat <sup>®</sup> tavaborole Vusion <sup>®</sup> <sup>F/Q/D</sup>	<ul> <li>FREQUENCY/QUANTITY/DURATION (F/Q/D)</li> <li>Vusion<sup>®</sup> 50 gm ointment –Maximum 100 grams in a 90-day time period</li> </ul>

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	V. Dermatol	ogic Agents
Anti-Infectives – Topical		
clindamycin solution, gel, lotion, swab clindamycin/benzoyl peroxide (gen Duac <sup>®</sup> ) erythromycin solution, gel	Acanya <sup>®</sup> Benzamycin <sup>®</sup> Cleocin T <sup>®</sup> Clindagel <sup>®</sup> clindamycin phos gel (gen Clindagel <sup>®</sup> ) clindamycin/benzoyl peroxide (gen BenzaClin <sup>®</sup> ) clindamycin/benzoyl peroxide (gen Onexton <sup>®</sup> ) clindamycin/benzoyl peroxide (gen Acanya <sup>®</sup> ) erythromycin swab erythromycin swab erythromycin/benzoyl peroxide Evoclin <sup>®</sup> Neuac <sup>®</sup> Onexton <sup>®</sup>	
	Anti-Virals	s – Topical
acyclovir cream docosanol (gen Abreva)	acyclovir ointment Denavir <sup>®</sup> penciclovir (gen Denavir <sup>®</sup> ) Xerese <sup>®</sup> Zovirax <sup>®</sup> cream, ointment	
	Immunomodulators & Re	lated Agents – Topical <sup>CC</sup>
Eucrisa <sup>®</sup> pimecrolimus tacrolimus	Elidel <sup>®</sup> Opzelura <sup>®</sup> Vtama <sup>®</sup> Zoryve <sup>®</sup> 0.3% foam, 0.15% cream	<ul> <li>CLINICAL CRITERIA</li> <li>Confirm diagnosis of FDA-approved, compendia-supported, and Medicaid-covered indication</li> <li>Plaque psoriasis – Trial of a Preferred agent from the Psoriasis Agents, Topical class</li> </ul>

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters		
	V. Dermatol	ogic Agents		
	Psoriasis Agents – Topical			
calcipotriene cream, ointment, scalp solution	calcipotriene foam (gen Sorilux <sup>®</sup> ) calcipotriene/betamethasone dipropionate (gen Taclonex <sup>®</sup> ) calcitriol ointment (gen Vectical <sup>®</sup> ) Duobrii™ Enstilar <sup>®</sup> Sorilux <sup>®</sup> Taclonex <sup>®</sup> Vectical <sup>®</sup> Zoryve <sup>®</sup> 0.3% cream			
	Rosacea Ager	nts, Topical <sup>cc</sup>		
azelaic acid metronidazole cream, gel	Epsolay <sup>®</sup> Finacea <sup>®</sup> ivermectin Metrocream <sup>®</sup> Metrogel <sup>®</sup> metronidazole gel pump, lotion Noritate <sup>®</sup> Rosadan <sup>®</sup> Soolantra <sup>®</sup>	<ul> <li>CLINICAL CRITERIA</li> <li>Confirm diagnosis of FDA-approved, compendia-supported, and Medicaid-covered indication</li> </ul>		
	Steroids, Topica	I – Low Potency		
hydrocortisone acetate OTC hydrocortisone acetate Rx	alclometasone Capex <sup>®</sup> shampoo Derma-Smoothe/FS <sup>®</sup> desonide fluocinolone oil hydrocortisone 2.5% soln (gen Texacort <sup>®</sup> ) Texacort <sup>®</sup>			

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Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters	
	V. Dermatolo	ogic Agents	
	Steroids, Topical – Medium Potency		
fluocinolone acetonide solution fluticasone propionate cream, ointment hydrocortisone valerate cream mometasone furoate	Beser lotion betamethasone valerate foam clocortolone fluocinolone acetonide cream, ointment flurandrenolide fluticasone propionate lotion hydrocortisone butyrate cream, lotion, ointment, solution hydrocortisone valerate ointment Locoid <sup>®</sup> Locoid Lipocream <sup>®</sup> Pandel <sup>®</sup> prednicarbate Synalar <sup>®</sup>		
	Steroids, Topical	I – High Potency	
betamethasone dipropionate lotion, cream, ointment betamethasone dipropionate augmented cream betamethasone valerate cream, ointment fluocinonide cream, ointment, solution triamcinolone acetonide	amcinonide cream ApexiCon-E <sup>®</sup> betamethasone dipropionate augmented gel, ointment, lotion betamethasone valerate lotion desoximetasone diflorasone Diprolene <sup>®</sup> fluocinonide gel, emollient halcinonide cream, solution (gen Halog <sup>®</sup> ) Halog <sup>®</sup> cream, solution, ointment Kenalog <sup>®</sup> Topicort <sup>®</sup> triamcinolone spray Vanos <sup>®</sup>		

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters	
	V. Dermatolo	ogic Agents	
	Steroids, Topical – Very High Potency		
clobetasol cream, emollient, gel, ointment, solution halobetasol cream, ointment	Bryhali™ clobetasol foam, lotion, spray, shampoo clobetasol 0.025% cream Clobex <sup>®</sup> halobetasol foam Olux <sup>®</sup> Ultravate <sup>®</sup>		

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	VI. Endocrine and	I Metabolic Agents
	Anabolic Steroids	– Topical <u>CDRP</u> , F/Q/D
testosterone gel packets (gen Vogelxo <sup>®</sup> ) testosterone gel pump (gen Androgel)	AndroGel <sup>®</sup> pump Natesto <sup>®</sup> Testim <sup>®</sup> testosterone gel packets (gen AndroGel <sup>®</sup> ) testosterone pump Vogelxo <sup>®</sup>	<ul> <li>CLINICAL DRUG REVIEW PROGRAM (CDRP)</li> <li>For diagnosis of hypogonadotropic or primary hypogonadism:         <ul> <li>Requires documented low testosterone concentration with two tests prior to initiation of therapy.</li> <li>Require documented testosterone therapeutic concentration to confirm response after initiation of therapy.</li> </ul> </li> <li>For diagnosis of delayed puberty:         <ul> <li>Requires documentation that growth hormone deficiency has been ruled out prior to initiation of therapy.</li> </ul> </li> <li>For diagnosis of fact form can be found at:         <ul> <li>https://newyork.fhsc.com/downloads/providers/NYRx_CDRP_PA_W orksheet_Prescribers_Anabolic_Steroids.pdf</li> <li>For diagnosis of gender dysphoria, see Hormone Replacement Therapy for Treatment of Gender Dysphoria coverage in the <u>DUR section</u> of this document</li> </ul> </li> <li>FREQUENCY/QUANTITY/DURATION (F/Q/D)</li> <li>Limitations for anabolic steroid products based on approved FDA labeled daily dosing and documented diagnosis:             <ul> <li>Duration limit of 6 months for delayed puberty</li> </ul> </li> </ul>

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Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters	
	VI. Endocrine and	Metabolic Agents	
	Bigua	nides	
glipizide/metformin glyburide/metformin Glumetza <sup>® <u>BLTG</u> metformin HCI metformin ER (gen Glucophage XR<sup>®</sup>)</sup>	metformin solution (gen Riomet <sup>®</sup> ) metformin 625mg, 750mg tablets metformin ER <sup>DO</sup> (gen Fortamet <sup>®</sup> , Glumetza <sup>®</sup> ) Riomet <sup>®</sup>	<ul> <li>DOSE OPTIMIZATION (DO)</li> <li>See Dose Optimization Chart for affected strengths</li> </ul>	
	Bisphosphonates – Oral		
alendronate	Actonel <sup>®</sup> Atelvia <sup>®</sup> Binosto <sup>®</sup> Fosamax <sup>®</sup> Fosamax <sup>®</sup> Plus D ibandronate risedronate		
	Dipeptidyl Peptidase	e-4 (DPP-4) Inhibitors	
alogliptin alogliptin/metformin Glyxambi <sup>®</sup> Janumet <sup>®</sup> Janumet <sup>®</sup> XR Januvia® <sup>DO</sup> Jentadueto <sup>®</sup> Jentadueto <sup>®</sup> Kazano <sup>®</sup> Nesina <sup>®</sup> Tradjenta <sup>®</sup>	alogliptin/pioglitazone (gen Oseni <sup>®</sup> ) Qtern <sup>®</sup> saxagliptin (gen Onglyza <sup>®</sup> ) saxagliptin/metformin sitagliptin (gen Zituvio <sup>™</sup> ) sitagliptin/metformin (gen Zituvimet) Steglujan <sup>®</sup> Zituvimet Zituvimet XR Zituvio <sup>™</sup>	<ul> <li>DOSE OPTIMIZATION (DO)</li> <li>See Dose Optimization Chart for affected strengths</li> </ul>	

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	VI. Endocrine and	Metabolic Agents
	Glucago	n Agents
Baqsimi <sup>®</sup> glucagon vial glucagon HCI emergency kit (Fresenius) Gvoke <sup>®</sup> pen, syringe, vial Zegalogue <sup>®</sup> pen, syringe	glucagon emergency kit (Eli Lilly, Amphastar)	
	Glucagon-like Peptide	-1 (GLP-1) Agonists <sup>CC</sup>
Byetta <sup>®</sup> Ozempic <sup>®</sup> Trulicity <sup>®</sup> Victoza <sup>®</sup> <sup>BLTG</sup>	Bydureon <sup>®</sup> BCise <sup>™</sup> liraglutide (gen Victoza <sup>®</sup> ) Mounjaro <sup>®</sup> Rybelsus <sup>®</sup> Soliqua <sup>®</sup> Xultophy <sup>®</sup>	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>Confirm diagnosis of FDA-approved, compendia-supported, and Medicaid-covered indication</li> </ul>

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters	
	VI. Endocrine and Metabolic Agents		
	Glucocortic	oids – Oral	
budesonide EC, DR dexamethasone tablet hydrocortisone methylprednisolone dose-pack prednisolone solution prednisone dose-pack, tablet	Agamree® Alkindi® Sprinkle budesonide ER Cortef® cortisone deflazacort (gen Emflaza®) dexamethasone elixir, solution dexamethasone intensol Emflaza® Eohilia™ Hemady™ Medrol® dose-pack, tablet methylprednisolone 4 mg, 8 mg, 16 mg, 32 mg Millipred® Millipred® Millipred® DP prednisolone tablet (gen Millipred®) prednisone intensol, solution Rayos® TaperDex™ Uceris®		

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Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	VI. Endocrine and	d Metabolic Agents
	Growth He	ormones <sup>cc</sup>
Genotropin <sup>®</sup> Norditropin <sup>®</sup>	Humatrope <sup>®</sup> Ngenla™ Nutropin AQ <sup>®</sup> NuSpin Omnitrope <sup>®</sup> Skytrofa <sup>®</sup> Sogroya <sup>®</sup> Zomacton <sup>®</sup>	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>Confirm diagnosis of FDA-approved or compendia-supported indication</li> <li>For Diagnosis of Growth Hormone Deficiency (GHD) or Short for Gestational Age (SGA):         <ul> <li>Prior to initiating growth hormone treatment, documentation of a recommended GHD diagnostic and / or laboratory test (e.g., provocative test and / or IGF-1 test)</li> </ul> </li> <li>Continuation of GH treatment, documentation of a recommended GHD laboratory test annually (e.g., IGF-1 test) and documentation of positive treatment response</li> </ul>
	Insulin – L	Long-Acting
insulin glargine solostar, vial (gen Lantus <sup>®</sup> Solostar <sup>®</sup> , vial) insulin glargine-YFGN <sup>1</sup> Lantus <sup>®</sup> Solostar <sup>®</sup> , vial	Basaglar <sup>®</sup> Basaglar <sup>®</sup> Tempo <sup>™</sup> insulin degludec vial, pen (gen Tresiba) insulin glargine max solostar (gen Toujeo <sup>®</sup> Max Solostar <sup>®</sup> ) insulin glargine solostar (gen Toujeo <sup>®</sup> Solostar <sup>®</sup> ) Levemir <sup>® 2</sup> Rezvoglar <sup>™</sup> Semglee <sup>®</sup> -YFGN: vial, pen Toujeo <sup>®</sup> Solostar <sup>®</sup> Toujeo <sup>®</sup> Max Solostar <sup>®</sup> Tresiba <sup>®</sup>	
	Insulin	– Mixes
Humalog <sup>®</sup> 50/50 Mix: pen insulin lispro 75/25 mix: pen (gen Humalog <sup>®</sup> Mix) insulin aspart prot/insulin aspart: vial, pen (gen Novolog)	Humalog <sup>®</sup> 75/25 mix: pen, vial Novolog <sup>®</sup> Mix: vial, pen	
1 = Preferred as of $2/6/2025$ 2 = Non-Preferred as of $2/6/2025$	Standard PA fax form: https://newyork.fhsc.com/down	loads/providers/NYRx_PDP_PA_Fax_Standardized.pdf 34

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
VI. Endocrine and Metabolic Agents		
	Insulin – Ra	apid-Acting
Apidra <sup>®</sup> insulin aspart (gen Novolog <sup>®</sup> ) cartridge, vial, pen insulin lispro (gen Humalog <sup>®</sup> U100) vial, pen insulin lispro junior (gen Humalog <sup>®</sup> Jr.)	Admelog <sup>®</sup> Afrezza <sup>®</sup> Fiasp <sup>®</sup> Penfill, FlexTouch, Pumpcart, vial Humalog <sup>®</sup> Jr. 100 U/mL Kwikpen Humalog <sup>®</sup> 100 U/mL vial, pen, cartridge, Tempo™ Humalog <sup>®</sup> 200 U/mL Lyumjev <sup>®</sup> Lyumjev <sup>®</sup> Tempo™ Novolog <sup>®</sup> cartridge, vial, FlexPen	
	Pancreatio	Enzymes
Creon <sup>®</sup> Zenpep <sup>®</sup>	Pertzye <sup>®</sup> Viokace <sup>®</sup>	
	Sodium Glucose Co-Trans	porter 2 (SGLT2) Inhibitors
Farxiga <sup>® <u>BLTG</u> Jardiance<sup>®</sup> Synjardy<sup>®</sup> Synjardy<sup>®</sup> XR Trijardy<sup>®</sup> XR Xigduo<sup>®</sup> XR <sup><u>BLTG</u></sup></sup>	dapagliflozin (gen Farxiga <sup>®</sup> ) dapagliflozin/metformin (gen Xigduo <sup>®</sup> XR) Inpefa <sup>™</sup> Invokamet <sup>® 2</sup> Invokamet <sup>® 2</sup> Segluromet <sup>®</sup> Steglatro <sup>®</sup>	
Thiazolidinediones (TZDs)		
pioglitazone	ACTOplus Met <sup>®</sup> Actos <sup>® <u>DO</u> Duetact<sup>®</sup> pioglitazone/glimepiride pioglitazone/metformin</sup>	<ul> <li>DOSE OPTIMIZATION (DO)</li> <li>See Dose Optimization Chart for affected strengths</li> </ul>

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Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters	
	VII. Gastrointestinal		
	Anti-E	metics	
aprepitant pack Diclegis <sup>® CC</sup> doxylamine succ/pyridoxine (gen Diclegis <sup>®</sup> ) <sup>CC</sup> ondansetron ODT, solution, tablet	Akynzeo <sup>®</sup> Anzemet <sup>®</sup> aprepitant capsule Bonjesta <sup>® CC</sup> Emend <sup>®</sup> capsule, powder packet, TriPack granisetron tablet Sancuso <sup>®</sup>	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>doxylamine succ/pyridoxine (Diclegis<sup>®</sup>, Bonjesta<sup>®</sup>): Confirm diagnosis of FDA-approved or compendia-supported indication</li> </ul>	
	Gastrointestir	nal Antibiotics	
metronidazole tablet neomycin vancomycin capsule, solution	Dificid <sup>®</sup> Firvanq <sup>®</sup> Flagyl <sup>®</sup> Likmez <sup>™</sup> metronidazole capsule metronidazole 125 mg tablet nitazoxanide paromomycin tinidazole Vancocin <sup>®</sup> Xifaxan <sup>® CC, ST, F/Q/D</sup>	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>Xifaxan<sup>®</sup>: Confirm diagnosis of FDA-approved or compendia- supported indication</li> <li>STEP THERAPY (ST)</li> <li>Xifaxan<sup>®</sup>: Requires trial of a fluoroquinolone antibiotic or azithromycin before Xifaxan<sup>®</sup> for treatment of Traveler's Diarrhea</li> <li>QUANTITY LIMITS: Xifaxan<sup>®</sup>:</li> <li>Irritable bowel syndrome with diarrhea (550 mg tablets) – 42 tablets per 30 days (Dose = 550 mg three times a day for 14 days)</li> <li>Maximum of 42 days' supply (126 units) per 365 days (3 rounds of therapy).</li> <li>Small Intestine Bacterial Overgrowth (550mg tablets) - 42 tablets per 30 days (Dose = 550mg three times a day for 10-14 days);</li> <li>Maximum of 28 days' supply (84 units) per 365 days (2 rounds of therapy).</li> </ul>	
Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters	
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	VII. Gastro	intestinal	
	Helicobacter p	oylori Agents	
Pylera <sup>®</sup> <sup>BLTG</sup>	bismuth/metronidazole/tetracycline (gen Pylera <sup>®</sup> ) lansoprazole/amoxicillin/clarithromycin Omeclamox-Pak <sup>®</sup> Talicia <sup>®</sup> Voquezna <sup>®</sup> Dual Pak Voquezna <sup>®</sup> Triple Pak <b>Proton Pump (PPI)/Acid S</b>		
esomeprazole magnesium Rx capsule lansoprazole capsule (Rx, OTC) lansoprazole OTC solutab omeprazole Rx pantoprazole tablet Protonix suspension <sup>BLTG</sup> rabeprazole	dexlansoprazole (gen Dexilant <sup>®</sup> ) Dexilant <sup>®</sup> <sup>DO</sup> esomeprazole magnesium tablet OTC esomeprazole capsules OTC esomeprazole suspension esomeprazole DR packets Konvomep <sup>™</sup> lansoprazole Rx solutab Nexium <sup>®</sup> Rx <sup>DO</sup> omeprazole OTC omeprazole/sodium bicarbonate Rx pantoprazole suspension Prevacid <sup>®</sup> OTC Prevacid <sup>®</sup> Rx Protonix <sup>®</sup> tablet Voquezna <sup>®</sup>	<ul> <li>DOSE OPTIMIZATION (DO)</li> <li>See Dose Optimization Chart for affected strengths</li> <li>FREQUENCY/QUANTITY/DURATION (F/Q/D)</li> <li>Quantity limits: <ul> <li>Once daily dosing for:</li> <li>GERD</li> <li>erosive esophagitis</li> <li>healing and maintenance of duodenal/gastric ulcers (including NSAID-induced)</li> <li>prevention of NSAID-induced ulcers</li> <li>Twice daily dosing for:</li> <li>hypersecretory conditions</li> <li>Barrett's esophagitis</li> <li>H. pylori</li> <li>refractory GERD</li> </ul> </li> <li>Duration limits: <ul> <li>90 days for:</li> <li>GERD</li> <li>365 days for:</li> <li>Maintenance treatment of duodenal ulcers, or erosive</li> </ul> </li> </ul>	
		esophagitis – 14 days for: o H. pylori	

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Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	VII. Gastro	intestinal
	Sulfasalazine	Derivatives
Apriso® <sup>BLTG</sup> mesalamine DR (gen Lialda®) Pentasa® <sup>BLTG</sup> sulfasalazine DR sulfasalazine IR	Azulfidine® Azulfidine Entab® balsalazide Colazal® Delzicol® Dipentum® Lialda® mesalamine DR (gen Delzicol®) mesalamine ER (gen Apriso®) mesalamine ER (gen Pentasa®) mesalamine DR	

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	VIII. Hemato	logical Agents
		s – Injectable <sup>F/Q/D</sup>
enoxaparin sodium Fragmin <sup>®</sup> vial	Arixtra <sup>® CC</sup> fondaparinux <sup>CC</sup> Fragmin <sup>®</sup> syringe Lovenox <sup>®</sup>	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>For patients requiring &gt; 30 days of therapy: Require confirmation of FDA-approved or compendia-supported indication</li> <li>Arixtra<sup>®</sup> (fondaparinux) Clinical editing to allow patients with a diagnosis of Heparin Induced Thrombocytopenia (HIT) to receive therapy without prior authorization.</li> <li>FREQUENCY/QUANTITY/DURATION (F/Q/D)</li> <li>Duration Limit: No more than 30 days for members initiating therapy</li> </ul>
	Anticoag	ulants – Oral
Eliquis <sup>®</sup> Pradaxa <sup>®</sup> capsule <sup>BLTG</sup> warfarin Xarelto <sup>®</sup> tablet <sup>DO</sup> Xarelto <sup>®</sup> 2.5 mg tablet <sup>BLTG</sup>	dabigatran (gen Pradaxa <sup>®</sup> ) Pradaxa <sup>®</sup> pellet pack rivaroxaban (gen Xarelto <sup>®</sup> ) Savaysa <sup>®</sup> Xarelto <sup>®</sup> dose pack, suspension	<ul> <li>DOSE OPTIMIZATION (DO)</li> <li>See Dose Optimization Chart for affected strengths</li> </ul>
	Colony Stim	ulating Factors
Neupogen <sup>®</sup> Nyvepria™	Fylnetra <sup>®</sup> Fulphila <sup>™</sup> Granix <sup>®</sup> Leukine <sup>®</sup> Neulasta <sup>®</sup> Nivestym <sup>™</sup> Releuko <sup>™</sup> Rolvedon <sup>®</sup> Stimufend <sup>®</sup> Udenyca <sup>®</sup> Zarxio <sup>®</sup> Ziextenzo <sup>®</sup>	

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters		
	VIII. Hematological Agents			
	Erythropoiesis Stimul	ating Agents (ESAs) <sup>cc</sup>		
Aranesp <sup>®</sup>	Mircera®	CLINICAL CRITERIA (CC)		
Epogen <sup>®</sup>	Procrit <sup>®</sup>	Confirm diagnosis for FDA- or compendia-supported uses		
Retacrit <sup>®</sup>				
	Hemophilia Age	ents – Factor VIII		
Advate <sup>®</sup>				
Adynovate®				
Afstyla®				
Altuviiio™				
Eloctate <sup>®</sup>				
Esperoct®				
Hemofil <sup>®</sup> M				
Humate-P <sup>®</sup>				
Jivi <sup>®</sup>				
Koate®				
Kogenate <sup>®</sup> FS				
Kovaltry®				
Novoeight <sup>®</sup>				
Nuwiq <sup>®</sup>				
Obizur®				
Recombinate™				
Xyntha <sup>®</sup>				
Xyntha <sup>®</sup> Solofuse				

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters		
	VIII. Hematological Agents			
	Hemophilia Ag	ents – Factor IX		
AlphaNine <sup>®</sup> SD				
Alprolix <sup>®</sup>				
BeneFIX®				
Idelvion®				
lxinity <sup>®</sup>				
Profilnine®				
Rebinyn <sup>®</sup>				
Rixubis®				
	Hemophilia A	gents – Other		
Alphanate <sup>®</sup> (von Willebrand	Alhemo®			
factor/Factor VIII)	Hympavzi™			
Coagadex <sup>®</sup> (Factor X)				
Corifact <sup>®</sup> (Factor XIII)				
Feiba <sup>®</sup> NF (activated prothrombin				
complex)				
Hemlibra <sup>®</sup> (emicizumab-kxwh)				
Novoseven <sup>®</sup> RT (Factor VIIa)				
Sevenfact <sup>®</sup> (Factor VIIa-jncw)				
Tretten <sup>®</sup> (Factor XIII) Vonvendi <sup>®</sup> (von Willebrand factor)				
Wilate <sup>®</sup> (von Willebrand				
factor/Factor VIII)				
Platelet Inhibitors				
Brilinta <sup>®</sup>	Effient®			
clopidogrel	Plavix®			
dipyridamole	prasugrel			
dipyridamole/aspirin				

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	IX. Immunol	ogic Agents
	Immunomodulato	rs – Systemic <sup>CC, st</sup>
Cosentyx® Dupixent® Enbrel® Fasenra® Humira® Nucala® Kolair®	Abrilada <sup>™</sup> (adalimumab-AFZB)         Actemra <sup>®</sup> subcutaneous         adalimumab-AACF (gen Idacio <sup>®</sup> )         adalimumab-AATY (gen Yuflyma <sup>®</sup> )         adalimumab-ADAZ (gen Hyrimoz <sup>®</sup> )         adalimumab-ADBM (gen Cyltezo <sup>®</sup> )         adalimumab-FKJP (gen Hulio <sup>®</sup> )         adalimumab-RYVK (gen Simlandi <sup>®</sup> )         adalimumab-RYVK (gen Simlandi <sup>®</sup> )         adalimumab-RYVK         Adbry <sup>™</sup> Amjevita <sup>™</sup> Bimzelx <sup>®</sup> Cibinqo <sup>™</sup> Cimzia <sup>®</sup> Cyltezo <sup>®</sup> (adalimumab-ADBM)         Ebglyss <sup>™</sup> Entyvio <sup>®</sup> SQ         Hadlima <sup>™</sup> Hulio <sup>®</sup> (adalimumab-FKJP)         Hyrimoz <sup>®</sup> (adalimumab-AACF)         Ilumya <sup>®</sup> Kevzara <sup>®</sup> Kineret <sup>®</sup> Nemluvio <sup>®</sup> Olumiant <sup>®</sup> Orrencia <sup>®</sup> SQ         Ortezla <sup>®</sup> Otulfi <sup>™</sup> Pyzchiva <sup>®</sup> Rinvoq <sup>™</sup> ER	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>Confirm diagnosis for FDA- or compendia-supported uses</li> <li>STEP THERAPY (ST)</li> <li>For indications not specified below</li> <li>Trial of a non-specific anti-inflammatory drug such as an aminosalicylate or immunosuppressant, or a disease-modifying anti-rheumatic drug (DMARD)</li> <li>Trial of a TNF inhibitor prior to treatment with a JAK inhibitor</li> <li>INDICATION-SPECIFIC REQUIREMENTS: <ul> <li>Asthma:</li> <li>history and concurrent use of a corticosteroid</li> </ul> </li> <li>Nasal polyps: <ul> <li>Trial of dermatitis:</li> <li>Trial with a topical prescription product for a duration of at leas 3 months.</li> <li>For JAK inhibitors: Trial of topical prescription product and systemic product for a combined duration of at least 6 months.</li> </ul> </li> <li>COPD: <ul> <li>History and concurrent use of a long acting beta agonist (LABA) + long acting muscarinic agonist (LAMA) + inhaled corticosteroid (ICS)</li> </ul> </li> </ul>

2 = Non-Preferred as of 2/6/2025

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Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	IX. Immunol	ogic Agents
	Rinvoq <sup>®</sup> LQ Selarsdi <sup>™</sup> Siliq <sup>™</sup> Simlandi <sup>®</sup> (adalimumab-RYVK) Simponi <sup>®</sup> Skyrizi <sup>®</sup> Skyrizi <sup>®</sup> On-Body Sotyktu <sup>™</sup> Spevigo <sup>®</sup> Stelara <sup>®</sup> Steqeyma <sup>®</sup> Taltz <sup>®</sup> Tezspire <sup>®</sup> pen Tremfya <sup>®</sup> Tyenne <sup>®</sup> Velsipity <sup>™</sup> Xeljanz <sup>®</sup> Xeljanz <sup>®</sup> XR Yesintek <sup>™</sup> Yuflyma <sup>®</sup> (adalimumab-AATY) Yusimry <sup>™</sup> Zymfentra <sup>™</sup>	
	Immunosupp	ressives, Oral
azathioprine CellCept <sup>®</sup> suspension <sup>BLTG</sup> cyclosporine softgel, capsule cyclosporine modified capsule, solution mycophenolic acid mycophenolate mofetil capsule, tablet Rapamune <sup>®</sup> solution Rapamune <sup>®</sup> tablet	Astagraf XL <sup>®</sup> Azasan <sup>®</sup> CellCept <sup>®</sup> capsule, tablet Envarsus XR <sup>®</sup> everolimus (gen Zortress <sup>®</sup> ) Imuran <sup>®</sup> Lupkynis <sup>™</sup> <sup>CC, F/Q/D</sup> mycophenolate mofetil suspension Myfortic <sup>®</sup> Myhibbin <sup>™</sup>	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>Lupkynis<sup>™</sup> (voclosporin):         <ul> <li>Confirm diagnosis for FDA- or compendia-supported uses</li> <li>Confirm concurrent therapy with mycophenolate</li> </ul> </li> <li>FREQUENCY/QUANTITY/DURATION (F/Q/D)</li> <li>Lupkynis<sup>™</sup> limited to 30-day supply</li> </ul>

2 = Non-Preferred as of 2/6/2025

https://newyork.fhsc.com/downloads/providers/NYRx\_PDP\_PA\_Fax\_Standardized.pdf

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
IX. Immunologic Agents		
tacrolimus	Neoral <sup>®</sup> Prograf <sup>®</sup> Sandimmune <sup>®</sup> capsule, solution Zortress <sup>®</sup>	

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	X. Miscellane	eous Agents
Progestins (for Cachexia)		
megestrol acetate suspension	megestrol 625 mg/5 mL suspension	
Epinephrine – Self- administered		
EpiPen <sup>® <u>BLTG</u> EpiPen Jr.® <u>BLTG</u></sup>	Auvi-Q <sup>®</sup> epinephrine (gen Adrenaclick <sup>®</sup> )	
	epinephrine (gen EpiPen®) epinephrine (gen EpiPen Jr.®) Neffy <sup>®</sup>	

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters		
	XI. Musculoskeletal Agents			
	Skeletal Muse	cle Relaxants		
baclofen tablet chlorzoxazone 500 mg cyclobenzaprine 5 mg, 10 mg tablet dantrolene methocarbamol orphenadrine ER tizanidine tablet	Amrix <sup>®</sup> baclofen 15mg tablet baclofen solution baclofen suspension (gen Fleqsuvy <sup>™</sup> ) carisoprodol <sup>ST, F/Q/D</sup> carisoprodol compound <sup>ST, F/Q/D</sup> carisoprodol compound/codeine <sup>CC,</sup> <sub>ST, F/Q/D</sub> chlorzoxazone (gen Lorzone) 375 mg, 750 mg chlorzoxazone 250 mg tablet cyclobenzaprine 7.5 mg cyclobenzaprine ER capsule (gen Amrix) Dantrium <sup>®</sup> Fexmid <sup>®</sup> Fleqsuvy <sup>™</sup> Lorzone <sup>®</sup> Lyvispah <sup>™</sup> metaxalone metaxalone 640 mg tablet orphenadrine-aspirin-caffeine Soma <sup>® ST, F/Q/D</sup> Soma <sup>®</sup> 250 <sup>ST, F/Q/D</sup> Tanlor <sup>®</sup> tizanidine capsule Zanaflex <sup>®</sup>	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>For carisoprodol/codeine products:</li> <li>Limited to a total of 4 opioid prescriptions every 30 days; exemption for diagnosis of cancer or sickle cell disease</li> <li>Medical necessity rationale for opioid therapy is required for patients on established opioid dependence therapy</li> <li>PA required for initiation of opioid therapy in patients currently on benzodiazepine therapy</li> <li>STEP THERAPY (ST)</li> <li>Trial with 1 analgesic and 2 skeletal muscle relaxants prior to use of carisoprodol containing products</li> <li>FREQUENCY/QUANTITY/DURATION (F/Q/D)</li> <li>Carisoprodol – Maximum 4 units per day, 21-day supply</li> <li>Carisoprodol combinations – Maximum 8 units per day, 21-day supply (not to exceed the 84 cumulative units per year limit)</li> </ul>		

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	XII. Opht	halmics
	Alpha-2 Adrenergic Agonists	(for Glaucoma) – Ophthalmic
Alphagan P <sup>®</sup> 0.1% <sup>BLTG</sup> Alphagan P <sup>®</sup> 0.15% <sup>BLTG</sup> brimonidine 0.2%	apraclonidine brimonidine 0.1% (gen Alphagan P <sup>®</sup> )	
Simbrinza®	brimonidine 0.15% (gen Alphagan P <sup>®</sup> ) Iopidine <sup>®</sup>	
	Antibiotics –	Ophthalmic
bacitracin/polymyxin B erythromycin gentamicin Natacyn <sup>®</sup> neomycin/gramicidin/polymyxin polymyxin/trimethoprim sulfacetamide solution tobramycin	Azasite <sup>®</sup> bacitracin neomycin/bacitracin/polymyxin sulfacetamide ointment Tobrex <sup>®</sup>	
	Antibiotics/Steroid Com	binations – Ophthalmic
neomycin/polymyxin/dexamethasone sulfacetamide/prednisolone TobraDex <sup>®</sup> ointment tobramycin/dexamethasone suspension	Maxitrol <sup>®</sup> neomycin / bacitracin/polymyxin /HC neomycin/polymyxin/HC TobraDex <sup>®</sup> ST Zylet <sup>®</sup>	
Antihistamines – Ophthalmic		
azelastine ketotifen OTC olopatadine OTC	bepotastine (gen Bepreve <sup>®</sup> ) Bepreve <sup>®</sup> epinastine Lastacaft <sup>®</sup> olopatadine Rx Pataday <sup>®</sup> Zaditor <sup>®</sup> OTC Zerviate <sup>™</sup>	

Standard PA fax form:

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters		
	XII. Ophthalmics			
	Anti-inflammatories/Immuno	omodulators – Ophthalmic <sup>CC</sup>		
Eysuvis <sup>®</sup> Restasis <sup>® <u>BLTG</u> Restasis MultiDose<sup>®</sup> Xiidra<sup>®</sup></sup>	Cequa <sup>®</sup> cyclosporine (gen Restasis <sup>®</sup> ) Miebo™ Tyrvaya™ Verkazia <sup>®</sup> Vevye <sup>®</sup>	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>Diagnosis documentation required to justify utilization as a first line agent or attempt treatment with an artificial tear, gel, or ointment.</li> </ul>		
	Beta Blockers	– Ophthalmic		
betaxolol Betoptic S <sup>®</sup> carteolol Combigan <sup>® <u>BLTG</u> Istalol<sup>® <u>BLTG</u> levobunolol timolol maleate gel</sup></sup>	Betimol <sup>®</sup> brimonidine/timolol (gen Combigan <sup>®</sup> ) timolol 0.5% (gen Betimol <sup>®</sup> ) timolol maleate (gen Timoptic <sup>®</sup> Ocudose <sup>®</sup> ) timolol maleate solution (gen Istalol <sup>®</sup> ) Timoptic <sup>®</sup> Ocudose <sup>®</sup>			
	Fluoroquinolone	es – Ophthalmic		
ciprofloxacin moxifloxacin (gen Vigamox®) ofloxacin	Besivance <sup>®</sup> Ciloxan <sup>®</sup> gatifloxacin moxifloxacin (gen Moxeza <sup>®</sup> ) Ocuflox <sup>®</sup> Vigamox <sup>®</sup>			
1	Non-Steroidal Anti-Inflammatory Drugs (NSAIDS) – Ophthalmic			
Acular LS <sup>® <u>BLTG</u> diclofenac flurbiprofen ketorolac</sup>	Acular <sup>®</sup> Acuvail <sup>®</sup> bromfenac BromSite <sup>®</sup> Ilevro <sup>®</sup> ketorolac LS Nevanac <sup>®</sup> Prolensa <sup>®</sup>			

Standard PA fax form:

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters				
	XII. Ophthalmics					
	Prostaglandin Agor	nists – Ophthalmic				
latanoprost	bimatoprost Iyuzeh™ Lumigan® Rocklatan® tafluprost (gen Zioptan®) Travatan Z® travoprost (gen Travatan Z®) Xalatan® Xelpros® Vyzulta™ Zioptan®					

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters				
	XIII. Otics					
	Fluoroquinolones – Otic					
Cipro HC®ciprofloxacinciprofloxacin/dexamethasone (gen Ciprodex®)ciprofloxacin/fluocinolone (gen Otovel™)						

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters		
XIV. Renal and Genitourinary				
	Alpha Reductase I	nhibitors for BPH		
finasteride	nasteride dutasteride dutasteride/tamsulosin Proscar®			
	Antihyper	uricemics		
allopurinol 100 mg, 300 mg colchicine tablet febuxostat probenecid probenecid/colchicine	allopurinol 200 mg colchicine capsule Colcrys Gloperba <sup>®</sup> Mitigare <sup>®</sup> Uloric <sup>®</sup> Zyloprim <sup>®</sup>			
	Cystine Deplet	ing Agents <sup>cc</sup>		
Cystagon®	Procysbi®	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>Confirm diagnosis of FDA-approved or compendia-supported indication</li> </ul>		
	Electrolyte	Depleters		
Lokelma <sup>®</sup> sodium polystyrene Veltassa <sup>®</sup>				
	Phosphate Bind	lers/Regulators		
calcium acetate sevelamer carbonate powder, tablet (gen Renvela)	Auryxia <sup>™</sup> ferric citrate 210 mg tablet (gen Auryxia <sup>™</sup> ) Fosrenol <sup>®</sup> lanthanum carbonate Renvela <sup>®</sup> tablet, powder pack sevelamer HCI (gen Renagel) Velphoro <sup>®</sup> Xphozah <sup>®</sup>			

1 = Preferred as of 2/6/2025 Standard PA fax form:

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	XIV. Renal and	l Genitourinary
	Selective Alpha A	drenergic Blockers
alfuzosin tamsulosin	Flomax <sup>®</sup> Rapaflo <sup>®</sup> silodosin	
	Urinary Tract A	Antispasmodics
fesoterodine ER (gen Toviaz <sup>®</sup> ) Myrbetriq <sup>® DO, BLTG</sup> oxybutynin oxybutynin ER <sup>DO</sup> solifenacin Toviaz <sup>® DO</sup>	darifenacin Detrol <sup>®</sup> Detrol LA <sup>® DO</sup> flavoxate Gemtesa <sup>®</sup> mirabegron (gen Myrbetriq®) Myrbetriq <sup>®</sup> solution <sup>F/Q/D</sup> Oxytrol <sup>®</sup> tolterodine tolterodine ER trospium trospium ER Vesicare <sup>® DO</sup> Vesicare <sup>® DO</sup>	<ul> <li>DOSE OPTIMIZATION (DO)</li> <li>See Dose Optimization Chart for affected strengths</li> <li>FREQUENCY/QUANTITY/DURATION (F/Q/D)</li> <li>Myrbetriq<sup>®</sup> solution; limited to a 30-day supply</li> </ul>
	Urea Cycle	e Disorders
Buphenyl <sup>®</sup> powder, tablet Carbaglu <sup>®</sup> <sup>BLTG</sup> Olpruva <sup>™</sup> Pheburane <sup>®</sup> Ravicti <sup>®</sup> sodium phenylbutyrate powder, tablet (gen Buphenyl <sup>®</sup> )	carglumic acid	

Preferred Drugs	Non-Preferred Drugs Prior Authorization/Coverage Parameters					
	XIV. Renal and Genitourinary					
	Uterine Disorder Treatments <sup>F/Q/D</sup>					
Myfembree <sup>®</sup> Oriahnn <sup>®</sup> Orilissa <sup>®</sup>		<ul> <li>LIFETIME QUANTITY LIMIT:</li> <li>Myfembree<sup>®</sup>, Oriahnn<sup>®</sup>, Orilissa<sup>®</sup> 150 mg: maximum of 24 months cumulative use</li> <li>Orilissa<sup>®</sup> 200 mg: maximum of 6 months cumulative use</li> </ul>				

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters				
XV. Respiratory						
	COPD Agents					
Anoro Ellipta <sup>®</sup> Atrovent HFA <sup>®</sup> Bevespi <sup>®</sup> Aerosphere <sup>®</sup> Combivent Respimat <sup>®</sup> Incruse Ellipta <sup>®</sup> ipratropium ipratropium / albuterol roflumilast (gen Daliresp <sup>®</sup> ) Spiriva <sup>®</sup> HandiHaler <sup>®</sup> <u>BLTG</u> Spiriva Respimat <sup>®</sup> Stiolto Respimat <sup>®</sup> Trelegy Ellipta <sup>®</sup> Tudorza Pressair <sup>®</sup>	Breztri <sup>™</sup> Aerosphere Daliresp <sup>®</sup> Duaklir <sup>®</sup> Pressair Ohtuvayre <sup>™</sup> tiotropium (gen Spiriva <sup>®</sup> Handihaler <sup>®</sup> ) Yupelri <sup>®</sup>					
	Antihistamines	s – Intranasal				
azelastine olopatadine						
	Antihistamines – S	econd Generation				
cetirizine OTC tablet cetirizine OTC syrup/solution 1 mg/ 1 mL fexofenadine OTC tablet levocetirizine tablet loratadine OTC	cetirizine OTC chewable cetirizine-D OTC Clarinex <sup>®</sup> Clarinex-D <sup>®</sup> desloratadine (gen Clarinex <sup>®</sup> ) levocetirizine solution loratadine-D OTC					
Beta2 Adrenergic Agents – Inhaled Long-Acting <sup>CC, F/Q/D</sup>						
Brovana <sup>® <u>BLTG</u> formoterol (gen Perforomist<sup>®</sup>) Serevent Diskus<sup>®</sup></sup>	arformoterol (gen Brovana <sup>®</sup> ) Perforomist <sup>®</sup> Striverdi Respimat <sup>®</sup>	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>PA is required for all new long-acting beta agonist prescriptions for beneficiaries under FDA- or compendia-supported age as indicated:</li> </ul>				

Non-Preferred Drugs	Prior Authorization/Coverage Parameters				
XV. Respiratory					
		Brovana <sup>®</sup> / arformoterol		≥ 18 years	
		Perforomist <sup>®</sup> / formoterol		≥ 18 years	
		Serevent Diskus®		≥ 4 years	
		Striverdi Respimat®		≥ 18 years	
	F	REQUENCY/QUANTITY/	DURATION (F/Q/D)		
			. ,		
		Brovana <sup>®</sup> / arformoterol	60 units (1 carton of 6	0 vials or 120 mL)	
		Perforomist <sup>®</sup> / formoterol	60 units (1 carton of 6	0 vials or 120 mL)	
		Serevent Diskus®	1 diskus (60 blisters)		
		Striverdi Respimat®	1 unit (one cartridge a inhaler)	ind one Respimat	
Beta2 Adrenergic Agents	s –	- Inhaled Short-Acting			
Airsupra™ albuterol HFA (gen Ventolin HFA®)					
levalbuterol HFA					
Corticosteroi	ids	s – Inhaled			
ArmonAir <sup>®</sup> Digihaler <sup>®</sup>					
fluticasone DISKUS					
QVAR RediHaler <sup>®</sup>					
	XV. Res Beta2 Adrenergic Agent Airsupra™ albuterol HFA (gen Ventolin HFA®) levalbuterol solution levalbuterol HFA Corticostero ArmonAir® Digihaler® Asmanex® HFA	XV. Respir         KV. Respir         F         M         Beta2 Adrenergic Agents -         Airsupra™         albuterol HFA (gen Ventolin HFA®)         levalbuterol solution         levalbuterol HFA         Corticosteroids         ArmonAir® Digihaler®         Asmanex® HFA         fluticasone DISKUS	XV. Respiratory         Brovana® / arformoterol         Perforomist® / formoterol         Serevent Diskus®         Striverdi Respimat®         FREQUENCY/QUANTITY/         Maximum units per 30 days         Brovana® / arformoterol         Perforomist® / formoterol         Serevent Diskus®         Brovana® / arformoterol         Perforomist® / formoterol         Perforomist® / formoterol         Serevent Diskus®         Striverdi Respimat®         Beta2 Adrenergic Agents – Inhaled Short-Acting         Airsupra™         albuterol HFA (gen Ventolin HFA®)         levalbuterol solution         levalbuterol HFA         Gorticosteroids – Inhaled         ArmonAir® Digihaler®         Asmanex® HFA         fluticasone DISKUS	XV. Respiratory         Brovana® / arformoterol         Perforomist® / formoterol         Serevent Diskus®         Striverdi Respimat®         FREQUENCY/QUANTITY/DURATION (F/Q/D)         Maximum units per 30 days         Brovana® / arformoterol         60 units (1 carton of 6         Perforomist® / formoterol         60 units (1 carton of 6         Perforomist® / formoterol         60 units (1 carton of 6         Serevent Diskus®         Striverdi Respimat®         1 diskus (60 blisters)         Striverdi Respimat®         1 unit (one cartridge a inhaler)         Beta2 Adrenergic Agents – Inhaled Short-Acting         Airsupra™         albuterol HFA (gen Ventolin HFA®)         levalbuterol solution         levalbuterol HFA         fuevalbuterol HFA         ArmonAir® Digihaler®         Asmanex® HFA         fluticasone DISKUS	

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters	
	XV. Resp	biratory	
Corti	costeroid/Beta2 Adrenergic Agent (Lo	ng-Acting) Combinations – Inhaled <sup>CC, F/Q/D</sup>	
Advair Diskus <sup>® <u>BLTG</u> Advair HFA<sup>® <u>BLTG</u> Dulera<sup>®</sup> Symbicort<sup>® <u>BLTG</u></sup></sup></sup>	AirDuo <sup>®</sup> Digihaler <sup>®</sup> AirDuo <sup>™</sup> RespiClick <sup>®</sup> Breo Ellipta <sup>®</sup> budesonide/formoterol (gen Symbicort) fluticasone-salmeterol (gen AirDuo <sup>™</sup> RespiClick <sup>®</sup> ) fluticasone-salmeterol (gen Advair Diskus <sup>®</sup> ) fluticasone-salmeterol (gen Advair HFA <sup>™</sup> ) fluticasone-vilanterol (gen Breo Ellipta <sup>®</sup> )	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>PA is required for all new long-acting beta age beneficiaries under FDA-or compendia-support Advair Diskus<sup>®</sup></li> <li>Advair HFA<sup>®</sup></li> <li>AirDuo<sup>™</sup> RespiClick<sup>®</sup></li> <li>Dulera<sup>®</sup> 100 mcg and 200 mcg</li> <li>Dulera<sup>®</sup> 50 mcg</li> <li>fluticasone-salmeterol</li> <li>budesonide-formoterol (Symbicort<sup>®</sup>) 80/4.5 mcg</li> </ul>	rted age as indicated: ≥ 4 years ≥ 12 years > 12 years ≥ 12 years ≥ 4 years ≥ 4 years ≥ 4 years ≥ 4 years
		budesonide-formoterol (Symbicort <sup>®</sup> ) 160/4.5 mcc fluticasone/vilanterol (Breo Ellipta <sup>®</sup> ) FREQUENCY/QUANTITY/DURATION (F/Q/D) Advair Diskus <sup>®</sup> Advair HFA <sup>®</sup>	g ≥ 12 years ≥ 18 years One inhaler/diskus
		AirDuo <sup>™</sup> RespiClick <sup>®</sup> fluticasone-salmeterol fluticasone/vilanterol (Breo Ellipta <sup>®</sup> ) Budesonide/formoterol (Symbicort <sup>®</sup> ) Dulera <sup>®</sup>	every 30 days Up to 8 inhalers every 180 days
	Corticosteroid	s – Intranasal	
budesonide OTC Dymista <sup>® <u>BLTG</u> fluticasone fluticasone OTC Nasonex<sup>®</sup> OTC Omnaris<sup>®</sup> triamcinolone OTC Zetonna<sup>®</sup></sup>	azelastine-fluticasone (gen Dymista <sup>®</sup> ) flunisolide mometasone Rx, OTC QNASL <sup>® CC</sup> Ryaltris <sup>®</sup> Xhance™	<ul> <li>CLINICAL CRITERIA (CC)</li> <li>Clinical consideration in regard to drug interac patients with HIV/AIDs diagnosis or antiretrovi</li> </ul>	

1 = Preferred as of 2/6/20252 = Non-Preferred as of 2/6/2025

Preferred Drugs	Non-Preferred Drugs	n-Preferred Drugs Prior Authorization/Coverage Parameters				
	XV. Respiratory					
	Leukotriene Modifiers					
montelukast tablet, chew tab	Accolate <sup>®</sup> montelukast granules Singulair <sup>®</sup> zafirlukast					

# NYS Medicaid NYRx Clinical Drug Review Program (CDRP)

The Clinical Drug Review Program (CDRP) is aimed at ensuring specific drugs are utilized in a medically appropriate manner. Under the CDRP, certain drugs require prior authorization because there may be specific safety issues, public health concerns, the potential for fraud and abuse or the potential for significant overuse and misuse.

#### **Prior Authorization**

Prior authorization for some drugs subject to the CDRP must be obtained through a representative at the clinical call center. For some drugs subject to the CDRP, only prescribers, not their authorized agents, can initiate the prior authorization process.

Please be prepared to respond to a series of questions that identify prescriber, patient, and reason for prescribing drug, and to fax clinical documentation upon request. Clinical guidelines for the CDRP as well as prior authorization worksheets are available online at <a href="https://newyork.fhsc.com/providers/CDRP\_about.asp">https://newyork.fhsc.com/providers/CDRP\_about.asp</a>.

The following drugs are subject to the Clinical Drug Review Program:

- <u>fentanyl mucosal agents</u>: <u>https://newyork.fhsc.com/providers/CDRP\_fentanyl\_mucosal\_agents.asp</u>
- palivizumab (Synagis<sup>®</sup>): <u>https://newyork.fhsc.com/providers/CDRP\_synagis.asp</u>
- <u>sodium oxybate products (Xyrem<sup>®</sup>, Xywav<sup>™</sup>)</u>: <u>https://newyork.fhsc.com/providers/CDRP\_xyrem.asp</u>
- <u>somatropin (Serostim®)</u>: <u>https://newyork.fhsc.com/providers/CDRP\_serostim.asp</u>

The following drug classes are subject to the Clinical Drug Review Program and are also included on the Preferred Drug List:

<u>Anabolic Steroids</u>: <u>https://newyork.fhsc.com/providers/CDRP\_anabolic\_steroids.asp</u>

#### NYS Medicaid NYRx Drug Utilization Review (DUR) Program

Frequency/Quantity/Duration (F/Q/D) Program and Step Therapy parameters are implemented to ensure clinically appropriate and cost-effective use of these drugs and drug classes.

For additional Step Therapy and Frequency/Quantity/Duration parameters for drugs and drug classes that are also included on the Preferred Drug List (PDL), please see pages 4 through 56.

Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Quantity / Duration (F/Q/D) Parameters	Additional / Alternate Parameter(s)
Corticotropin (Acthar <sup>®</sup> , Cortrophin <sup>®</sup> )	all FDA-approved indications, other than infantile spasms. <b>Note</b> : It is first line therapy for infantile spasms in children less than 2 years of age – step therapy not required.	<ul> <li>QUANTITY LIMITS:</li> <li>Infantile spasms – 30 mL (six 5 mL vials)</li> <li>Multiple sclerosis – 35 mL (seven 5 mL vials)</li> <li>Multiple sclerosis - 40 u/0.5mL and 80 u/mL Selfject = 21 syringes</li> <li>DURATION LIMITS:</li> <li>Infantile spasms – 4 weeks; indicated for &lt; 2 years of age</li> <li>Multiple sclerosis – 5 weeks</li> <li>Rheumatic disorders – 5 weeks</li> <li>Dermatologic conditions – 5 weeks</li> <li>Allergic states (serum sickness) – 5 weeks</li> </ul>	<ul> <li>Confirm diagnosis of FDA- approved or compendia- supported indication</li> <li>Not covered for diagnostic purposes</li> </ul>

Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Quantity / Duration (F/Q/D) Parameters	Additional / Alternate Parameter(s)
Corticotropin (Acthar® Cortrophin®) (	continued)	FDA Indication	First Line Therapy
		<ul> <li>Multiple Sclerosis (MS) exacerbations</li> <li>Polymyositis/ dermatomyositis</li> <li>Idiopathic nephrotic syndrome Systemic lupus erythematosus (SLE)</li> <li>Nephrotic syndrome due to SLE</li> <li>Rheumatic disorders (specifically: psoriatic arthritis, rheumatoid arthritis, juvenile rheumatoid arthritis, ankylosing spondylitis)</li> <li>Dermatologic diseases (specifically Stevens-Johnson syndrome and erythema multiforme)</li> <li>Allergic states (specifically serum sickness)</li> <li>Ophthalmic diseases (keratitis, iritis, iridocyclitis, diffuse posterior uveitis/choroiditis, optic neuritis, chorioretinitis, anterior segment inflammation)</li> <li>Respiratory diseases (systemic sarcoidosis)</li> </ul>	<ul> <li>Corticosteroid or plasmapheresis</li> <li>Corticosteroid</li> <li>ACE Inhibitor, diuretic, corticosteroid (and for refractory patients: an immunosuppressive)</li> <li>Corticosteroid, antimalarial, or cytotoxic/immunosuppressive agent</li> <li>Immunosuppressive, corticosteroid, or ACE Inhibitor</li> <li>Corticosteroid, topical retinoid, biologic disease-modifying antirheumatic drugs (DMARD), non-biologic DMARD, or a non- steroidal anti-inflammatory drug (NSAID)</li> <li>Corticosteroid or analgesic</li> <li>Topical or oral corticosteroid, antihistamine, or NSAID</li> <li>Analgesic, anti-infective agent, and agents to reduce inflammation, such as NSAIDs and steroids</li> <li>Oral corticosteroid or an immunosuppressive.</li> </ul>

Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Quantity / Duration (F/Q/D) Parameters	Additional / Alternate Parameter(s)
<ul> <li>Anabolic Steroids – Injectable</li> <li>testosterone cypionate (Depo- Testosterone<sup>®</sup>, Azmiro<sup>™</sup>)</li> <li>testosterone enanthate (Xyosted<sup>®</sup>)*</li> <li>Anabolic Steroids – Oral</li> <li>testosterone undecanoate (Jatenzo<sup>®</sup>, Tlando<sup>®</sup>, Undecatrex)</li> <li>methyltestosterone (Methitest<sup>®</sup>)</li> </ul>		<ul> <li>Limitations for anabolic steroid products is based on approved FDA labeled daily dosing and documented diagnosis not to exceed a 90-day supply</li> <li>Xyosted<sup>®</sup> is limited to no more than 3 boxes for 90 days (1 box per 30 days)</li> <li>Initial duration limit of 3 months requiring documented follow-up monitoring for response and/or adverse effects before continuing treatment</li> <li>Duration limit of 6 months for delayed puberty</li> </ul>	*for additional parameters, see Hormone Replacement Therapy for Treatment of Gender Dysphoria section below.
<ul> <li>Anti-Diarrheal Agents</li> <li>alosetron (Lotronex<sup>®</sup>)</li> <li>crofelemer (Mytesi<sup>®</sup>)</li> <li>eluxadoline (Viberzi<sup>®</sup>)</li> </ul>	<ul> <li>Irritable Bowel Syndrome w/Diarrhea         <ul> <li>Trial of eluxadoline and rifaximin prior to alosetron.</li> </ul> </li> <li>Symptomatic relief of non- infectious diarrhea in patients with HIV/AIDS on anti-retroviral therapy         <ul> <li>Trial with an alternative anti- diarrheal agent.</li> </ul> </li> </ul>		<ul> <li>Confirmation of FDA-approved or compendia-supported indication.</li> </ul>

Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Quantity / Duration (F/Q/D) Parameters	Additional / Alternate Parameter(s)
Anti-Retroviral (ARV) Interventions		<ul> <li>QUANTITY LIMITS:</li> <li>Limit ARV active ingredient duplication</li> <li>Limit ARV utilization to a maximum of five products concurrently - excluding boosting with ritonavir (dose limit 600 mg or less) or cobicistat</li> <li>Limit Protease Inhibitor utilization to a maximum of two products concurrently</li> <li>Limit Integrase inhibitor utilization to a maximum of one product concurrently</li> <li>Limit non-nucleoside reverse transcriptase inhibitor utilization to a maximum of 1 product concurrently</li> <li>Limit ARV booster utilization to 1 product concurrently</li> <li>Limit co-formulated and copackaged complete ARV regimens listed in Appendix A to a maximum of 1 product concurrently with no additional ARVs.</li> </ul>	<ul> <li>Require confirmation of FDA- approved or compendia- supported use</li> <li>Point-of-service edit for antiretroviral / antiretroviral combinations to be avoided: https://newyork.fhsc.com/downl oads/providers/NYRx_PDP_ref erence_Antiretroviral_Antiretrov iral_Drug2Drug_Interactions.pdf</li> </ul>
belimumab (Benlysta®)	Trial of a disease-modifying anti-rheumatic drug (DMARD) prior to treatment with an immunomodulator		<ul> <li>Confirm diagnosis of FDA- approved or compendia- supported indication</li> </ul>
biotin			Confirm diagnosis of FDA- approved or compendia- supported indication

Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Quantity / Duration (F/Q/D) Parameters	Additional / Alternate Parameter(s)
<ul> <li>Benzodiazepine agents – oral</li> <li>alprazolam (Xanax<sup>®</sup>, Xanax<sup>®</sup> XR)</li> <li>chlordiazepoxide</li> <li>chlordiazepoxide/amitriptyline</li> <li>clonazepam (Klonopin<sup>®</sup>)</li> <li>clorazepate</li> <li>diazepam (Valium<sup>®</sup>)</li> <li>lorazepam (Ativan<sup>®</sup>, Lorazepam Intensol<sup>®</sup>, Loreev XR<sup>™</sup>)</li> <li>oxazepam</li> </ul>	<ul> <li>Generalized Anxiety Disorder (GAD) or Social Anxiety Disorder (SAD)</li> <li>Require trial with a Selective- Serotonin Reuptake Inhibitor (SSRI) or a Serotonin- Norepinephrine Reuptake Inhibitor (SNRI) prior to initial benzodiazepine prescription</li> <li>Panic Disorder requires concurrent therapy with an antidepressant (SSRI, SNRI, or Tricyclic antidepressant [TCA]).</li> <li>Skeletal muscle spasms</li> <li>Require trial with a skeletal muscle relaxant prior to a benzodiazepine</li> </ul>	<ul> <li><b>DURATION LIMIT:</b></li> <li>For Insomnia: 30 consecutive days</li> <li>For Panic Disorder: 30 consecutive days</li> </ul>	<ul> <li>Require confirmation of FDA- approved or compendia- supported use</li> <li>PA required for initiation of benzodiazepine therapy in patients currently on opioid or oral buprenorphine therapy</li> <li>PA required for any additional oral benzodiazepine prescription in patients currently on benzodiazepine therapy</li> <li>PA required when greater than a 14-day supply of a benzodiazepine is prescribed for someone on a CNS stimulant</li> </ul>
<ul> <li>Constipation Agents</li> <li>linaclotide (Linzess<sup>®</sup>)</li> <li>lubiprostone (Amitiza<sup>®</sup>)</li> <li>methylnaltrexone (Relistor<sup>®</sup>)</li> <li>naldemedine (Symproic<sup>®</sup>)</li> <li>naloxegol (Movantik<sup>®</sup>)</li> <li>plecanatide (Trulance<sup>®</sup>)</li> <li>prucalopride (Motegrity<sup>™</sup>)</li> <li>tenapanor (Ibsrela<sup>®</sup>)</li> </ul>	<ul> <li>Opioid Induced Constipation (OIC) and Chronic Idiopathic Constipation (CIC)</li> <li>Trial with an osmotic laxative, a stimulant laxative, and a stool softener prior to use.</li> <li>Irritable Bowel Syndrome w/ Constipation (IBS-C)</li> <li>Trial with a bulking agent and an osmotic laxative within 89 days of use.</li> </ul>	<ul> <li>QUANTITY LIMIT:</li> <li>linaclotide, naldemedine, naloxegol, plecanatide: 1 tablet/day</li> <li>lubiprostone: 2 capsules/day</li> <li>methylnaltrexone: 1 vial or syringe/day, 4 kits/28 days</li> <li>prucalopride: 2 mg/day max; 1 tablet per day</li> <li>tenapanor 2 tablets/day</li> </ul>	Confirmation of FDA-approved or compendia-supported indication.

Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Quantity / Duration (F/Q/D) Parameters	Additional / Alternate Parameter(s)
<ul> <li>Hormone Replacement Therapy for Treatment of Gender Dysphoria</li> <li>conjugated estrogens</li> <li>estradiol</li> <li>testosterone cypionate (Azmiro<sup>™</sup>)</li> <li>testosterone enanthate (Xyosted<sup>™</sup>)</li> <li>testosterone gel 1.62% (AndroGel<sup>®</sup>)*</li> <li>testosterone patch*</li> </ul>			<ul> <li>Confirm diagnosis of FDA- approved or compendia- supported indication</li> <li>For diagnosis of gender dysphoria please refer to October 2023 edition of the Medicaid Update:</li> <li><u>https://www.health.ny.gov/health</u> <u>care/medicaid/program/update/</u> <u>2023/no15_2023-</u> <u>10.htm#hormones</u></li> </ul>
			*Subject to Anabolic Steroids – Topical PDL class criteria
dextromethorphan / quinidine (Nuedexta®)		<ul> <li>QUANTITY LIMIT:</li> <li>2 capsules per day; 60 units per 30 days</li> <li>DURATION LIMIT:</li> <li>90 days of therapy</li> </ul>	<ul> <li>For patients ≥ 18 years of age:</li> <li>Confirm diagnosis of FDA- approved or compendia- supported indication</li> </ul>
Diabetic Test Strips		Preferred diabetic supply program <u>https://newyork.fhsc.com/provid</u> ers/diabeticsupplies.asp	
dronabinol (Marinol®)	<ul> <li>Step therapy for beneficiaries with HIV/AIDS, or cancer, AND eating disorder:</li> <li>Trial with megestrol acetate suspension prior to dronabinol</li> <li>Step therapy for beneficiaries with diagnosis of cancer and nausea/vomiting:</li> <li>Trial with a NYS Medicaid- preferred 5-HT3 receptor antagonist prior to dronabinol</li> </ul>		Confirm diagnosis of FDA- approved or compendia- supported indication

Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Quantity / Duration (F/Q/D) Parameters	Additional / Alternate Parameter(s)
risdiplam (Evrysdi <sup>®</sup> ) Fentanyl Transmucosal Agents • fentanyl (lozenge) • fentanyl (Fentora <sup>®</sup> ) (buccal tablet)		<ul> <li>QUANTITY LIMIT: fentanyl lozenge, Fentora®:</li> <li>4 units per day, 120 units per 30 days</li> <li>DURATION LIMIT:</li> <li>90 days</li> <li>Exemption for diagnosis of cancer, sickle cell disease, or hospice care</li> </ul>	<ul> <li>Confirm diagnosis of FDA-approved indication</li> <li>Confirm absence of advanced disease</li> <li>Confirm diagnosis</li> <li>Limited to a total of 4 opioid prescriptions every 30 days;</li> <li>For opioid-naïve patients: limited to a 7 days' supply for all initial opioid prescriptions,</li> <li>PA required for use if &gt; 90 MME (MME = morphine milligram equivalents) of opioid per day for management of non-acute pain (pain lasting &gt; 7 days).</li> <li>PA required for initiation of opioid therapy for patients on established opioid dependence therapy</li> <li>PA is required for initiation of opioid therapy in patients currently on benzodiazepine therapy</li> <li>Exemption for diagnosis of cancer, sickle cell, or hospice care</li> </ul>
<ul> <li>HIV PrEP (Pre-Exposure Prophylaxis Agents):</li> <li>cabotegravir (Apretude)</li> <li>emtricitabine/tenofovir disoproxil fumarate (Truvada<sup>®</sup>)</li> <li>emtricitabine/tenofovir alafenamide (Descovy<sup>®</sup>)</li> </ul>			Confirmation of negative HIV test every 3 months

Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Quantity / Duration (F/Q/D) Parameters	Additional / Alternate Parameter(s)
Imcivree™ (setmelanotide)			<ul> <li>Confirm diagnosis of FDA approved or compendia supported indication.</li> <li>Please be prepared to respond to a series of questions that identify the prescriber, the patient, and the reason for prescribing this drug.</li> <li>Please be prepared to fax clinical documentation upon request.</li> <li>The Imcivree fax form can be found at: <u>https://newyork.fhsc.com/downloads</u> /providers/NYRx_PDP_PA_Worksh <u>eet_Prescribers_Imcivree.pdf</u></li> </ul>
ivermectin (oral)			<ul> <li>Confirm diagnosis of FDA- approved or compendia- supported indication</li> </ul>
Lidocaine patches ● lidocaine (Lidoderm <sup>®</sup> , ZTLido™)			<ul> <li>Prescribers, or their authorized agents, are required to respond to a series of questions that identify the prescriber, the patient, and the reason for prescribing this drug.</li> <li>Prescriptions can be written for a 30-day supply with up to 2 refills</li> </ul>

Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Quantity / Duration (F/Q/D) Parameters	Additional / Alternate Parameter(s)
methadone	Requires a trial of a long-acting opioid prior to initiation for the management of chronic non- cancer pain	<ul> <li>QUANTITY LIMIT:</li> <li>12 units per day, 360 units per 30 days</li> <li>Exemption for diagnosis of cancer, hospice care, or sickle cell disease</li> </ul>	<ul> <li>Confirm diagnosis of chronic non-cancer pain</li> <li>Limited to a total of 4 opioid prescriptions every 30 days;</li> <li>PA required for initiation of methadone for patients on established opioid dependence therapy</li> <li>PA required for methadone prescriptions for patients currently on long-acting opioid therapy.</li> <li>PA required for initiation of long-acting opioid therapy in opioid-naïve patients.</li> <li>PA required for use if &gt; 90 MME (MME = morphine milligram equivalents) of opioid per day for management of non-acute pain (pain lasting &gt; 7 days).</li> <li>PA required for initiation of methadone therapy in patients currently on benzodiazepine therapy</li> <li>Exemption for diagnosis of cancer, sickle cell, or hospice care</li> </ul>
metoclopramide nasal spray (Gimoti™)			Metoclopramide nasal spray confirm diagnosis of diabetes

Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Quantity / Duration (F/Q/D) Parameters	Additional / Alternate Parameter(s)
<ul> <li>Ovulation Enhancing Drugs</li> <li>bromocriptine</li> <li>clomiphene</li> <li>letrozole</li> <li>tamoxifen</li> </ul>			<ul> <li>Confirm diagnosis of FDA- approved or compendia- supported indication and Medicaid covered indication</li> <li>Refer to <u>https://www.health.ny.gov/healt</u> <u>h_care/medicaid/program/updat</u> e/2019/2019-06.htm#ovulation</li> </ul>
Oxazolidinone Antibiotics <ul> <li>linezolid (Zyvox<sup>®</sup>)</li> <li>tedizolid (Sivextro<sup>®</sup>)</li> </ul>			<ul> <li>Please be prepared to respond to a series of questions that identify the prescriber, the patient, and the reason for prescribing this drug.</li> <li>Please be prepared to fax clinical documentation upon request.</li> <li>The Oxazolidinone Antibiotics fax form can be found at: <u>https://newyork.fhsc.com/download</u> <u>s/providers/NYRx_PDP_PA_Works</u> <u>heet_Prescribers_oxazolidinone_a</u> <u>ntibiotic.pdf</u></li> </ul>
<ul> <li>Pubertal Suppressants</li> <li>leuprolide acetate (Lupron Depot-PED<sup>®</sup>, Eligard<sup>®</sup>, Fensolvi<sup>®</sup>, Lupron Depot<sup>®</sup>)</li> <li>nafarelin acetate (Synarel<sup>®</sup>)</li> <li>triptorelin (Triptodur<sup>®</sup>)</li> </ul>			<ul> <li>Confirm diagnosis of FDA- approved or compendia- supported indication</li> <li>Refer to <u>https://www.health.ny.gov/healt</u> <u>h_care/medicaid/program/updat</u> <u>e/2017/2017-</u> <u>01.htm#transgender</u> for Transgender Related Care and Services Update</li> </ul>

Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Quantity / Duration (F/Q/D) Parameters	Additional / Alternate Parameter(s)
esketamine (Spravato®)	Treatment Resistant     Depression: trial of at least two     oral antidepressants		<ul> <li>Confirm diagnosis of FDA approved indication for patients ≥18 years of age</li> <li>Before initiating esketamine nasal spray (Spravato), prescribers must attest that they have obtained a baseline score using a validated clinical assessment tool for depression (e.g., HAMD17, QIDS-C16C, MADRS).</li> </ul>
			<ul> <li>After the initiation of esketamine nasal spray (Spravato) therapy, every six months prescribers must attest that esketamine nasal spray (Spravato) has resulted in an improvement of depressive symptoms (from baseline) using the same baseline clinical assessment tool for depression (e.g., HAMD17, QIDS-C16C, MADRS).The esketamine worksheet can be accessed at: https://newyork.fhsc.com/downlo ads/providers/NYRx_PDP_PA_ Worksheet_Prescribers_Spravat o.pdf</li> </ul>
tasimelteon (Hetlioz <sup>®</sup> ,Hetlioz <sup>®</sup> LQ)		<ul> <li>QUANTITY LIMIT:</li> <li>One unit per day; 30 units per 30 days</li> </ul>	Confirm diagnosis of FDA- approved or compendia- supported indication

Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Quantity / Duration (F/Q/D) Parameters	Additional / Alternate Parameter(s)
<ul> <li>Parathyroid Hormone Analogs</li> <li>teriparatide (Forteo<sup>®</sup>)</li> <li>abaloparatide (Tymlos<sup>®</sup>)</li> </ul>	Requires a trial with a preferred oral bisphosphonate	<ul> <li>QUANTITY LIMIT:</li> <li>One unit per 28-day period</li> <li>LIFETIME QUANTITY LIMIT:</li> <li>25 months' cumulative use of a PTH analog</li> </ul>	
Topical Compounded Prescriptions			<ul> <li>Confirm diagnosis of FDA- approved or compendia- supported indication</li> <li>For non-opioid pain management alternatives please visit: <u>https://health.ny.gov/health_car</u> <u>e/medicaid/program/opioid_ma</u> <u>nagement/docs/non_opioid_alte</u> <u>rnatives_to_pain_management.</u> <u>pdf</u></li> </ul>

For more information on DUR Program, please refer to <a href="https://www.health.ny.gov/health\_care/medicaid/program/dur/index.htm">https://www.health.ny.gov/health\_care/medicaid/program/dur/index.htm</a>.

#### **Medication Assisted Treatment (MAT) Formulary**

<b>Medication Assisted Treatment (MAT) Formulary</b> **Prior authorization will not be required for medications used for the treatment of substance use disorder when prescribed according to generally accepted national professional guidelines for the treatment of a substance use disorder. **		
Drugs	Coverage Parameters	
	Opioid Antagonists	
Kloxxado <sup>™</sup> naloxone (syringe, vial, nasal spray) naloxone (nasal spray) OTC naltrexone Narcan <sup>®</sup> (nasal spray) Narcan <sup>®</sup> OTC Opvee <sup>®</sup> Rextovy® Zimhi <sup>™</sup> *		
	Opioid Dependence Agents – Injectable	
Brixadi™ Sublocade™ Vivitrol®		
	Opioid Dependence Agents – Oral/Transmucosal <sup>F/Q/D</sup>	
buprenorphine (tablet) buprenorphine/naloxone (tablet) buprenorphine/naloxone (film) Suboxone <sup>®</sup> (film) Zubsolv <sup>®</sup>	<ul> <li>QUANTITY LIMIT:</li> <li>buprenorphine sublingual (SL): Eight tablets dispensed as a 2-day supply; not to exceed 32 mg per day</li> <li>buprenorphine / naloxone tablet and film (Suboxone<sup>®</sup> 2mg/0.5mg, Zubsolv<sup>®</sup> 1.4mg/0.36mg, 0.7mg/0.18mg strength; Up to 12 sublingual tablets or films per day.</li> <li>buprenorphine/naloxone tablet and film (Suboxone<sup>®</sup> up to 4mg/1mg and 8mg/2mg strength, Zubsolv<sup>®</sup> 2.9mg/0.71mg and 5.7mg/1.4mg strength; Four sublingual tablets or films per day; maximum of 120 tablets or films dispensed as a 30-day supply, not to exceed 32 mg-8 mg of Suboxone<sup>®</sup>, or its equivalent per day</li> <li>buprenorphine/naloxone tablet: Suboxone<sup>®</sup> 12mg/3mg, Zubsolv<sup>®</sup> 8.6 mg/2.1 mg and Zubsolv<sup>®</sup> 11.4 mg/2.9 mg strength: Maximum of 60 tablets dispensed as a 30-day supply</li> <li>RELATED CLINICAL CRITERIA (CC)</li> <li>PA required for initiation of opioid therapy for patients established on opioid dependence therapy **</li> </ul>	

#### NYS Medicaid NYRx Brand Less Than Generic (BLTG) Program

On April 26, 2010, NYRx, the Medicaid Pharmacy Program, implemented a new cost-containment initiative, which promotes the use of certain multisource brand name drugs when the cost of the brand name drug is less expensive than the generic equivalent.

In conformance with State Education Law, which intends that patients receive the lower cost alternative, brand name drugs included in this program:

- Do not require "Dispense as Written" (DAW) or "Brand Medically Necessary" on the prescription
- Have a generic copayment
- Are paid at the Brand Name Drug reimbursement rate or usual and customary price, whichever is lower (SMAC/FUL are not applied)
- Do not require a new prescription if the drug is removed from this program

#### Effective May 01, 2025:

- Acular LS<sup>®</sup>, Brovana<sup>®</sup>, Carbatrol<sup>®</sup>, Cipro<sup>®</sup> oral suspension, Oxtellar XR<sup>®</sup>, Targretin<sup>®</sup> gel, Tegretol<sup>®</sup> tablet, Xarelto<sup>®</sup> 2.5 mg tablet will be **added** to the program
- Oseni<sup>®</sup> and Onglyza<sup>®</sup> Rx will be **removed** from the program

List of	Brand Name Drugs included in this pro	gram**
Acular LS <sup>®</sup>	EpiPen, Jr	Sprycel®
Advair Diskus <sup>®</sup>	Farxiga <sup>®</sup>	Symbicort®
Advair HFA®	Forteo®	Targretin <sup>®</sup> gel
Alphagan P <sup>®</sup> 0.15%	Glumetza®	Tegretol <sup>®</sup> suspension
Alphagan P <sup>®</sup> 0.1%	Istalol®	Tegretol <sup>®</sup> tablet
Apriso <sup>®</sup>	Kitabis <sup>®</sup> Pak	Tegretol <sup>®</sup> XR
Azopt™	Myrbetriq <sup>®</sup>	Trileptal <sup>®</sup> suspension
Bethkis <sup>®</sup>	Nexavar®	Ventolin <sup>®</sup> HFA
Brovana®	NuvaRing®	Victoza®
Carbaglu®	Oxtellar XR <sup>®</sup>	Votrient®
<b>Carbatrol</b> <sup>®</sup>	Pentasa®	Vyvanse <sup>®</sup> capsules
CellCept <sup>®</sup> suspension	Pradaxa®	Xarelto <sup>®</sup> 2.5 mg tablet
Cipro <sup>®</sup> oral suspension	Protonix <sup>®</sup> suspension	Xigduo <sup>®</sup> XR
Combigan <sup>®</sup>	Pylera®	Xopenex HFA <sup>®</sup>
Copaxone <sup>®</sup> 20 mg SQ	Restasis®	Zavesca®
Daytrana®	Retin-A <sup>®</sup> cream	
Depakote <sup>®</sup> Sprinkle	Risperdal Consta <sup>®</sup>	
Dymista <sup>®</sup>	Sandostatin LAR®	
EpiPen	Spiriva <sup>®</sup> Handihaler <sup>®</sup>	

\*\*List is subject to change

Please keep in mind that drugs in this program may be subject to prior authorization requirements of other pharmacy programs, promoting the use of the most cost-effective product.

#### **Important Billing Information**

- Pursuant to this program, prescription claims submitted to the Medicaid program do not require the submission of Dispense as Written/Product Selection Code of '1'; Pharmacies should submit DAW code 9 (Substitution Allowed by Prescriber but Plan Requests Brand). Pharmacies will receive an NCPDP reject response of "22" which means missing/invalid DAW code if other DAW codes are submitted. The only exception to this is DAW code 1 and "Brand Medically Necessary" on the prescription.
- For more information on the Brand Less Than Generic (BLTG) Program, please refer to https://newyork.fhsc.com/providers/bltgp\_about.asp

#### NYS Medicaid NYRx Mandatory Generic Drug Program

State law excludes Medicaid coverage of brand name drugs that have a Federal Food and Drug Administration (FDA) approved A-rated generic equivalent unless a prior authorization is obtained.

Coverage parameters under the Preferred Drug Program (PDP), Clinical Drug Review Program (CDRP), and/or the Brand Less Than Generic (BLTG) Program are applicable for certain products subject to the Mandatory Generic Drug Program (MGDP), including exemptions (as listed below).

#### **Prior Authorization Process**

- Prescribers, or an agent of the prescriber, must call the prior authorization line at 1-877-309-9493 and respond to a series of questions that identify the prescriber, the patient, and the reason for prescribing this drug.
- The prescriber must write "DAW and Brand Medically Necessary" on the face of the prescription.
- The call line 1-877-309-9493 is in operation 24 hours a day, seven days a week.

#### **Exempt Drugs**

• Based on specific characteristics of the drug and/or disease state generally treated, the following brand name drugs are exempt from the program and do **NOT** require PA:

Exempt Drugs	
Clozaril®	Neoral®
Dilantin®	Sandimmune®
Gengraf <sup>®</sup>	Tegretol®
Lanoxin®	Zarontin <sup>®</sup>
Levothyroxine Sodium (Unithroid <sup>®</sup> , Synthroid <sup>®</sup> , Levoxyl <sup>®</sup> )	

For more information on the Mandatory Generic Program, please refer to <u>https://newyork.fhsc.com/providers/MGDP\_about.asp</u>.

#### NYS Medicaid NYRx Dose Optimization Program

On November 14, 2013, the Medicaid NYRx program instituted a Dose Optimization initiative. Dose optimization can reduce prescription costs by reducing the number of pills a patient needs to take each day. The Department has identified drugs to be included in this program, the majority of which have FDA approval for once-a-day dosing, have multiple strengths available in correlating increments at similar costs and are currently being utilized above the recommended dosing frequency. Prior authorization will be required to obtain the following medication beyond the following limits:

#### Dose Optimization Chart

Brand Name			Dose Optimization Limitations	
CARDIOVASCULAR				
	Angioter	nsin Receptor	Blockers (ARBs)	
Benicar <sup>®</sup> 20 mg	1 daily	Tablet		
Micardis <sup>®</sup> 20 mg, 40 mg	1 daily	Tablet		
Diovan <sup>®</sup> 40 mg, 80 mg, 160 mg	1 daily	Tablet		
		Antiarrhyth	mics	
Amiodarone 100 mg	1 daily	Tablet	In case of dose titration for these medications, the department will allow for multiday dosing (up to 2 doses daily) for loading dose for 30 days	
ARBs Combinations				
Exforge <sup>®</sup> 5–160mg	1 daily	Tablet		
		ARBs/Diure	etics	
Benicar <sup>®</sup> HCT 20–12.5 mg	1 daily	Tablet		
Diovan <sup>®</sup> HCT 80–12.5 mg, 160–12.5 mg	1 daily	Tablet		
Edarbyclor <sup>®</sup> 40–12.5 mg	1 daily	Tablet		
Micardis <sup>®</sup> HCT 40–12.5 mg, 80–12.5 mg	1 daily	Tablet		
Beta Blockers				
Bystolic <sup>®</sup> 2.5 mg, 5 mg, 10 mg	1 daily	Tablet		
nadolol 40 mg	1 daily	Tablet		
Toprol <sup>®</sup> XL 25 mg, 50 mg, 100 mg	1 daily	Tablet		

Brand Name			Dose Optimization Limitations
	CEN	ITRAL NERVOU	IS SYSTEM
		Anticonvuls	ants
Aptiom <sup>®</sup> 200 mg, 400 mg	1 daily	Tablet	
Fycompa <sup>®</sup> 4 mg, 6 mg	1 daily	Tablet	
topiramate ER 100 mg (Qudexy <sup>®</sup> XR, Trokendi XR <sup>®</sup> )	1 daily	Capsule	
Lamictal XR <sup>®</sup> 50 mg	1 daily	Tablet	In case of dose titration for these medications, the department will allow for multiday dosing (up to 2 doses daily) for titration purposes for 90 days
Oxtellar XR <sup>®</sup> 300 mg	1 daily	Tablet	In case of dose titration for these medications, the department will allow for multiday dosing (up to 2 doses daily) for titration purposes for 90 days
Lyrica <sup>®</sup> 25 mg, 50 mg, 75 mg, 100 mg, 150 mg, 200 mg	3 daily	Capsule	Electronic bypass for diagnosis of seizure disorder identified in medical claims data. In case of dose titration for these
Qudexy <sup>®</sup> XR 100 mg	1 daily	Capsule	medications, the department will allow for multiday dosing (up
Lyrica <sup>®</sup> 225 mg and 300 mg	2 daily	Capsule	to 2 doses daily) for titration purposes for 3 months
Trokendi XR <sup>®</sup> 100 mg	1 daily	Capsule	
		Antiparkinson A	Agents
Azilect <sup>®</sup> 0.5 mg	1 daily	Tablet	
	Antipsy	chotics – Seco	nd Generation
Abilify <sup>®</sup> 2 mg	4 daily	Tablet	
Abilify <sup>®</sup> 5 mg, 10 mg, 15 mg	1 daily	Tablet	In case of dose titration for these medications, the Department
aripiprazole 5 mg, 10 mg, 15 mg	1 daily	Tablet	will allow for multiday dosing (up to 2 doses/daily) for titration
Invega <sup>®</sup> 1.5 mg, 3 mg	1 daily	Tablet	purposes for three months
Latuda <sup>®</sup> 20 mg, 40 mg, 60 mg	1 daily	Tablet	
olanzapine 5 mg, 10 mg	1 daily	Tablet	
olanzapine ODT 5 mg, 10 mg	1 daily	Tablet	
paliperidone er 1.5 mg, 3 mg	1 daily	Tablet	
quetiapine fumarate er 200 mg, 150 mg	1 daily	Tablet	
Rexulti <sup>®</sup> 0.25 mg, 0.5 mg, 1 mg, 2 mg	1 daily	Tablet	
Seroquel <sup>®</sup> XR 150 mg, 200 mg	1 daily	Tablet	
Vraylar <sup>®</sup> 1.5 mg, 3 mg	1 daily	Capsule	
Zyprexa <sup>®</sup> Zydis 5 mg, 10 mg	1 daily	Tablet	

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Brand Name		D	ose Optimization Limitations		
CENTRAL NERVOUS SYSTEM					
CNS Stimulants					
Adderall <sup>®</sup> XR 5 mg, 10 mg, 15 mg	1 daily	Capsule			
amphetamine salt combo ER 5 mg, 10 mg, 15 mg	1 daily	Capsule			
Concerta <sup>®</sup> ER 18 mg, 27 mg	1 daily	Tablet			
dexmethylphenidate ER 10 mg, 20 mg (Focalin XR generic)	1 daily	Capsule			
Focalin <sup>®</sup> XR 5 mg, 10 mg, 15 mg, 20 mg	1 daily	Capsule			
methylphenidate CD 10 mg, 20 mg	1 daily	Capsule			
methylphenidate er 18 mg (Concerta® generic)	1 daily	Tablet			
methylphenidate la 20 mg (Ritalin <sup>®</sup> LA generic)	1 daily	Capsule			
modafinil 100 mg	1 daily	Tablet			
Provigil <sup>®</sup> 100 mg	1 daily	Tablet			
QuilliChew <sup>®</sup> ER 20 mg	1 daily	Tablet			
Ritalin <sup>®</sup> LA 10 mg, 20 mg	1 daily	Capsule			
Vyvanse <sup>®</sup> 10 mg, 20 mg, 30 mg, 40 mg	1 daily	Capsule			
Other Agen	ts for Attenti	ion Deficit Hyper	activity Disorder (ADHD)		
guanfacine ER 1 mg, 2 mg	1 daily	Tablet			
atomoxetine 40 mg	1 daily	Capsule			
Intuniv <sup>®</sup> 1 mg, 2 mg	1 daily	Tablet			
Strattera <sup>®</sup> 40 mg	1 daily	Capsule			
	S	edative Hypnoti	cs		
Lunesta <sup>®</sup> 1 mg	1 daily	Tablet			
Seroto	Serotonin-Norepinephrine Reuptake Inhibitors (SNRIs)				
Effexor <sup>®</sup> XR 37.5 mg, 75 mg	1 daily	Capsule	In the case of dose titration for these medications, the		
desvenlafaxine succinate ER (Pristiq <sup>®</sup> ER 50 mg)	1 daily	Tablet	Department will allow for multiday dosing (up to 2 doses/daily) for titration purposes for three months.		
Selective Serotonin Reuptake Inhibitors (SSRIs)					
Lexapro <sup>®</sup> 5 mg, 10 mg	1 daily	Tablet			
Trintellix <sup>®</sup> 5 mg, 10 mg	1 daily	Tablet			

Brand Name	Dose Optimization Limitations			
CENTRAL NERVOUS SYSTEM				
Viibryd <sup>®</sup> 10 mg, 20 mg	1 dailyTabletIn the case of dose titration for these once daily medicatio the Department will allow for multiday dosing (up to 2 doses/daily) for titration purposes for three months.			
Miscellaneous Antidepressants				
bupropion xl 150 mg	1 daily	Tablet	In case of dose titration for these medications, the Department	
mirtazapine 7.5 mg	1 daily	Tablet	will allow for multiday dosing (up to 2 doses/daily) for titration purposes for three months	

Brand Name	Dose Optimization Limitations			
ENDOCRINE AND METABOLIC				
Biguanides				
metformin ER 500 mg (Glumetza ER, Fortamet ER generic)	1 daily	Tablet		
Dipeptidyl Peptidase-4 (DPP-4) Inhibitors				
Januvia <sup>®</sup> 25 mg, 50 mg	1 daily	Tablet		
Thiazolidinediones (TZDs)				
Actos <sup>®</sup> 15 mg	1 daily	Tablet		

Brand Name	Dose Optimization Limitations		
GASTROINTESTINAL			
Proton Pump / Acid Secretion Inhibitors			
Dexilant <sup>®</sup> 30 mg	1 daily	Capsule	
Nexium <sup>®</sup> 5 mg, 10 mg, 20 mg	1 daily	Packet	
Nexium <sup>®</sup> 20 mg	1 daily	Capsule	
Prevacid <sup>®</sup> DR 15 mg	1 daily	Capsule	

Brand Name		Dose Optimization Limitations		
HEMATOLOGICAL				
Anticoagulants - Oral				
Xarelto <sup>®</sup> 10 mg	1 daily	Tablet		

Brand Name	Dose Optimization Limitations		
RENAL AND GENITOURINARY			
	Urinary	y Tract Antispasmodics	
Detrol <sup>®</sup> LA 2 mg	1 daily	Capsule	
Myrbetriq <sup>®</sup> 25 mg	1 daily	Tablet	
oxybutynin chloride ER 5 mg	1 daily	Tablet	
Toviaz <sup>®</sup> ER 4 mg	1 daily	Tablet	
VESIcare <sup>®</sup> 5 mg	1 daily	Tablet	

PA requirements are not dependent on the date a prescription is written. New prescriptions and refills on existing prescriptions require PA even if the prescription was written before the date the drug was determined to require PA.

To obtain a prior authorization (PA), please call the prior authorization Clinical Call Center at 1-877-309-9493. The Clinical Call Center is available 24 hours per day, 7 days per week with pharmacy technicians and pharmacists who will work with you, or your agent, to quickly obtain PA.

Medicaid enrolled prescribers with an active e-PACES account can initiate PA requests through the web-based application PAXpress<sup>®</sup>. The website for PAXpress is <u>https://paxpress.nypa.hidinc.com</u>.

When, in the judgment of the prescriber or the pharmacist, an emergency condition exists, the prescriber or pharmacist can call the Clinical Call center and obtain authorization for a 72-hour emergency supply of the drug prescribed to allow time for the prior authorization to be obtained.