



The following tables provide a summary of the final Preferred Drug List (PDL) selections made by the Commissioner for the Department for Medicaid Services (DMS) based on the Drug Review and Options for Consideration document prepared for the Pharmacy and Therapeutics (P&T) Advisory Committee’s review on **January 28, 2025**, and the resulting official recommendations.

## NEW PRODUCTS TO MARKET

### Cobenfy™

#### Central Nervous System – Antipsychotics, Second Generation (Atypical) and Injectable: Non-Preferred

##### Approval Duration: 1 year

- *Xanomeline is a M1 and M4 muscarinic acetylcholine receptor agonist in the central nervous system, influencing dopaminergic activity. Trospium is a muscarinic antagonist in the peripheral tissues, mitigating xanomeline’s side effects.*

##### Initial Approval Criteria:

- Diagnosis of schizophrenia; **AND**
- Trial and failure, allergy, contraindication (including potential drug-drug interactions with other medications) or intolerance to one preferred agent; **AND**
- Prescriber attests that liver enzymes and bilirubin were measured prior to initiation; **AND**
- Patient meets the minimum age recommended by the package insert for the provided indication.

##### Renewal Criteria:

- Prescriber attestation of clinically significant improvement or stabilization in clinical signs and symptoms.

**Age Limit:** 18 years of age or older

**Quantity Limit:** 2 capsules per day

Preferred Agents	Non-Preferred Agents
aripiprazole tablet <sup>CC, QL</sup>	Abilify MyCite starter kit <sup>CC, QL</sup>
asenapine tablet <sup>CC, QL</sup>	Abilify MyCite maintenance kit <sup>CC, QL</sup>
clozapine tablet <sup>CC, QL</sup>	Abilify tablet <sup>QL</sup>
lurasidone tablet <sup>CC, QL</sup>	aripiprazole ODT
olanzapine ODT <sup>CC, QL</sup>	aripiprazole solution
olanzapine tablet <sup>CC, QL</sup>	Caplyta capsule <sup>CC, QL</sup>
quetiapine tablet <sup>CC, QL</sup>	clozapine ODT <sup>QL</sup>
quetiapine ER tablet <sup>CC, QL</sup>	Clozaril tablet <sup>QL</sup>
risperidone ODT <sup>CC, QL</sup>	<b>Cobenfy capsule <sup>AE, CC, QL</sup></b>
risperidone solution <sup>CC, QL</sup>	Fanapt tablet dose pack <sup>QL</sup>
risperidone tablet <sup>CC, QL</sup>	Fanapt tablet <sup>QL</sup>
Vraylar capsule dose pack <sup>AE, CC, QL</sup>	Geodon capsule <sup>QL</sup>



Preferred Agents	Non-Preferred Agents
Vraylar capsule <sup>AE, CC, QL</sup>	Invega ER tablet <sup>QL</sup>
ziprasidone capsule <sup>CC, QL</sup>	Latuda tablet <sup>QL</sup>
	Lybalvi tablet <sup>AE, CC, QL</sup>
	Nuplazid capsule <sup>CC, QL</sup>
	Nuplazid tablet <sup>CC, QL</sup>
	olanzapine/fluoxetine capsule <sup>CC, QL</sup>
	Opipza film <sup>CC, QL</sup>
	paliperidone ER tablet <sup>QL</sup>
	Rexulti tablet <sup>QL</sup>
	Risperdal solution <sup>QL</sup>
	Risperdal tablet <sup>QL</sup>
	Saphris SL tablet <sup>QL</sup>
	Secuado patch <sup>QL</sup>
	Seroquel tablet <sup>QL</sup>
	Seroquel XR tablet <sup>QL</sup>
	Symbyax capsule <sup>CC, QL</sup>
	Versacloz suspension <sup>QL</sup>
	Zyprexa tablet <sup>QL</sup>
	Zyprexa Zydis ODT <sup>QL</sup>

**Livdelzi®**

**Gastrointestinal, Bile Salts: Non-Preferred**

**Approval Duration: 1 year**

- *Seladelpar decreases bile acid synthesis by activating peroxisome proliferator-activated receptor (PPAR)-delta. This causes downregulation of CYP7A1, an enzyme used to synthesize bile acids from cholesterol, through Fibroblast Growth Factor 21 (FGF21).*

**Initial Approval Criteria:**

- Diagnosis of primary biliary cholangitis (PBC); **AND**
- Prescribed by, or in consultation with, a gastroenterologist, hepatologist, or other disease state specialist; **AND**
- Patient meets one of the following:
  - Patient has had a 12-month trial and failure of ursodiol, and will take Livdelzi in addition to current therapy; **OR**
  - Patient has a contraindication or intolerance to ursodiol and will take Livdelzi as monotherapy; **AND**
- Patient has an alkaline phosphatase (ALP) level greater than 200 IU/L; **AND**
- Patient does not have decompensated cirrhosis; **AND**
- Patient meets the minimum age recommended by the package insert for the provided indication.



**Renewal Criteria:**

- Documentation (e.g., progress notes, labs) of improvement or stabilization in alkaline phosphatase (ALP); **AND**
- Patient meets one of the following:
  - Patient has had a 12-month trial and failure of ursodiol and will take Livdelzi in addition to current therapy; **OR**
  - Patient has a contraindication or intolerance to ursodiol and will take Livdelzi as monotherapy.

**Age Limit:** 18 years of age or older

**Quantity Limit:** 1 capsule per day

Preferred Agents	Non-Preferred Agents
ursodiol capsule	Bylvay capsule <sup>CC, QL</sup>
ursodiol tablet	Bylvay pellet <sup>CC, QL</sup>
	Chenodal tablet
	Cholbam capsule
	Iqirvo tablet <sup>CC, QL</sup>
	<b>Livdelzi<sup>AE, CC, QL</sup></b>
	Livmarli solution <sup>CC, QL</sup>
	Ocaliva tablet <sup>CC, QL, AE</sup>
	Reltone capsule
	Urso Forte tablet
	Urso tablet

**Vyalev™**

**Central Nervous System – Parkinson’s Disease (Antiparkinson’s Agents): Non-Preferred**

**Approval Duration: 1 year**

- *Foscarbidopa and foslevodopa are prodrugs that are converted to carbidopa and levodopa in vivo. Carbidopa inhibits peripheral levodopa from decarboxylation, allowing more levodopa to be delivered to the brain, where it is converted to dopamine.*

**Initial Approval Criteria:**

- Diagnosis of Parkinson’s disease (PD); **AND**
- Receiving PD therapy with carbidopa/levodopa; **AND**
- Experiencing “off” episodes with carbidopa/levodopa for at least 2 hours per day; **AND**
  - Trial and failure of at least 2 adjunctive therapies, such as: Dopamine agonists (e.g., pramipexole, ropinirole)
  - Monoamine oxidase-B inhibitors (e.g., selegiline)
  - Catechol-O-methyltransferase inhibitors (e.g., entacapone); **AND**
- Patient will not take within two weeks of a non-selective monoamine oxidase (MAO) inhibitor (e.g., phenelzine, isocarboxazid, tranylcypromine); **AND**



- Patient meets the minimum age recommended by the package insert for the provided indication.

**Renewal Criteria:**

- Patient has clinically meaningful response of treatment (e.g., patient shows a reduction in time of “off” episodes).

**Age Limit:** 18 years of age or older

**Quantity Limit:** 2 vials (20 mL) per day

Preferred Agents	Non-Preferred Agents
amantadine capsule	Azilect tablet
amantadine solution	carbidopa tablet
amantadine tablet	Comtan tablet
benztropine tablet	Crexont IR ER capsule
carbidopa/levodopa ER tablet	Dhivy tablet
carbidopa/levodopa ODT	Duopa suspension
carbidopa/levodopa tablet	Gocovri capsule
carbidopa/levodopa/entacapone tablet	Inbrija inhalation
entacapone tablet	Kynmobi film <sup>CC, QL</sup>
selegiline capsule	Lodosyn tablet
selegiline tablet	Nourianz tablet <sup>CC QL</sup>
trihexyphenidyl solution	Ongentys capsule <sup>CC, QL</sup>
trihexyphenidyl tablet	Osmolex ER tablet
	rasagiline tablet
	Rytary ER capsule
	Sinemet tablet
	Stalevo tablet
	Tasmar tablet
	<b>Vyalev vial <sup>AE, CC, QL</sup></b>
	Xadago <sup>CC, QL</sup>
	Zelapar ODT

**Ebglyss™**

**Immunomodulators – Atopic Dermatitis: Non-Preferred**

**Approval Duration: 4 months initial, 1 year renewal**

- *Lebrikizumab-lbkz is an IgG4 monoclonal antibody that binds to and allows interleukin (IL)-13 to bind to IL-13R-α1, but inhibits human IL-13 signaling. IL-13 is a cytokine involved in type 2 inflammation seen in atopic dermatitis.*

**Initial Approval Criteria:**

- Diagnosis of moderate-to-severe atopic dermatitis (AD) with ≥ 1 of the following:
  - Involvement of at least 10% of body surface area (BSA); **OR**



- Scoring Atopic Dermatitis (SCORAD) score of 25 or more; **OR** Investigator’s Global Assessment (IGA) with a score  $\geq$  3; **OR**
- Eczema Area and Severity Index (EASI) score of  $\geq$  16; **OR**
- Incapacitation due to AD lesion location (e.g., head and neck, palms, soles, or genitalia); **AND**
- Prescribed by, or in consultation with, a dermatologist, allergist/immunologist, or other specialist in the treatment of atopic dermatitis; **AND**
- Trial and failure, contraindication, or intolerance to  $\geq$  1 agent in 2 or more of the following categories (total prior agent use of  $\geq$  90 days):
  - Topical corticosteroid of medium to high potency (e.g., mometasone, fluocinolone) unless inappropriate for the location (e.g., face, groin); **AND**
  - Topical calcineurin inhibitor (i.e., tacrolimus or pimecrolimus); **OR**
  - Immunosuppressive systemic agent (e.g., cyclosporine, azathioprine, methotrexate, mycophenolate mofetil); **AND**
- Trial and failure, contraindication, or intolerance to at least one preferred injectable agent (Adbry or Dupixent); **AND**
- Patient must meet the minimum age and weight recommended by the package insert for the provided indication.

**Renewal Criteria:**

- Patient must continue to meet initial approval criteria; **AND**
- Patient must have disease improvement and/or stabilization based on an objective measure.

**Age Limit:** 12 years of age or older

**Quantity Limit:** 1 pen/syringe (2 mL) per 28 days

Preferred Agents	Non-Preferred Agents
Adbry autoinjector <sup>AE, CC, QL</sup>	<b>Ebglyss</b> <sup>AE, CC, QL</sup>
Adbry syringe <sup>AE, CC, QL</sup>	Elidel
Dupixent pen <sup>CC, QL</sup>	<b>Nemluvio</b> <sup>AE, CC, QL</sup>
Dupixent syringe <sup>CC, QL</sup>	
Eucrisa <sup>CC, QL</sup>	
Opzelura cream <sup>CC, AE</sup>	
pimecrolimus cream	
tacrolimus ointment	



### Xdemvy™

#### Non-PDL

#### Approval Duration: 3 months initial, 1 year renewal

- *Lotilaner is an inhibitor of the gamma-aminobutyric acid (GABA)-gated chloride channel. This drug is selective for channels present in mites, causing a paralytic action leading to death.*

#### Initial Approval Criteria:

- Diagnosis of Demodex Blepharitis; **AND**
- Prescribed by, or in consultation with, an ophthalmologist or other specialist for the requested condition; **AND**
- Prescriber attests that the patient currently has active disease.

#### Renewal Criteria:

- Patient has a diagnosis of Demodex Blepharitis [H01.00]; **AND**
- Prescribed by, or in consultation with, an ophthalmologist or other specialist for the requested condition; **AND**
- Prescriber attests patient has experienced a response to previous therapy.

**Age Limit:** 18 years of age or older

**Quantity Limit:** 1 bottle (10 mL) per month

### Yorvipath™

#### Non-PDL

#### Approval Duration: 6 months initial, 1 year renewal

- *Palopegteriparatide mimics endogenous parathyroid hormone to increase serum calcium and decrease serum phosphate. This drives homeostasis and a number of downstream effects, such as maintaining appropriate laboratory levels and stimulating bone turnover.*

#### Initial Approval Criteria:

- Diagnosis of hypoparathyroidism; **AND**
- Prescriber attests that this medication is NOT being prescribed for acute hypoparathyroidism post-surgery; **AND**
- Patient has not received therapy with parathyroid hormone analogs (e.g. abaloparatide, teriparatide) for 24 months or more (lifetime cumulative); **AND**
- Documentation that the following labs are within normal limits:
  - **Corrected Serum Calcium:** 7.8-10.2 mg/dL; **AND**
  - **Serum Phosphate:** 2.5-4.5 mg/dL; **AND**



- Prescriber attestation that the patient is not well-controlled despite appropriate utilization (trial and failure of 3 months) of calcium and active forms of vitamin D; **AND**
- Prescribed by, or in consultation with, an endocrinologist or other specialist for the requested condition.

**Renewal Criteria:**

- Patient continues to have the above listed diagnosis; **AND**
- Prescribed by, or in consultation with, an endocrinologist or other specialist for the requested condition; **AND**
- Documentation (e.g., progress note) of response to therapy.

**Age Limit:** 18 years of age or older

**Quantity Limit:** 2 pens per month

**Duvyza<sup>TM</sup>**

**Muscular Dystrophy Agents: Nonpreferred**

**Approval Duration: 6 months initial, 1 year renewal**

- *Givinostat is a histone deacetylase inhibitor. The precise mechanism by which givinostat exerts its effect in patients with DMD is unknown.*

**Initial Approval Criteria:**

- Diagnosis of Duchenne muscular dystrophy (DMD) [G71.01]; **AND**
- Platelet count within the last 30 days equals to or is greater than  $150 \times 10^9/L$ ; **AND**
- Prescribed by, or in consultation with, a neuromuscular specialist with expertise in the treatment of DMD; **AND**
- Patient is ambulatory (e.g., ability to walk with or without assistive devices, not wheelchair dependent); **AND**
- Patient's baseline ambulatory function has been or will be assessed prior to therapy initiation; **AND**  
Patient has been on a stable systemic corticosteroid therapy for at least 6 months and will continue to be on the systemic corticosteroid therapy unless contraindicated or clinically significant adverse effects are experienced; **AND**
- Prescriber provides a patient weight obtained within the past 3 months; **AND**
- The requested dose meets the FDA-approved dosing recommendation.

**Renewal Criteria:**

- Documentation (e.g., progress note) of stabilized or improved ambulatory function from baseline; **AND**
- Patient will continue systemic corticosteroid therapy unless contraindicated or clinically significant adverse effects are experienced; **AND**
- Prescriber provides a patient weight obtained within the past 3 months; **AND**
- The requested dose meets the FDA-approved dosing recommendation.



**Age Limit:** 6 years of age or older  
**Quantity Limit:** 12 mL per day

Preferred Agents	Non-Preferred Agents
Emflaza suspension <sup>AE, CC</sup>	Agamree suspension <sup>AE, CC, QL</sup>
Emflaza tablet <sup>AE, CC, QL</sup>	deflazacort suspension <sup>AE, CC</sup>
	deflazacort tablet <sup>AE, CC, QL</sup>
	<b>Duvyzat <sup>AE, CC, QL</sup></b>

**Nemluvio®**

**Immunomodulators, Atopic Dermatitis Class: Nonpreferred**

**Approval Duration: 4 months initial, 1 year renewal**

- *Nemolizumab-ilot is a humanized IgG2 monoclonal antibody that inhibits IL-31 signaling by binding selectively to IL-31 RA. IL-31 is a naturally occurring cytokine that is involved in pruritus, inflammation, epidermal dysregulation, and fibrosis. Nemolizumab-ilot inhibited IL-31-induced responses including the release of proinflammatory cytokines and chemokines.*

**Initial Approval Criteria:**

**Atopic Dermatitis:**

- Diagnosis of moderate-to-severe atopic dermatitis (AD) with ≥ 1 of the following:
  - Involvement of at least 10% of body surface area (BSA); **OR**
  - Investigator’s Global Assessment (IGA) with a score ≥ 3; **OR**
  - Eczema Area and Severity Index (EASI) score of ≥ 16; **OR** Peak Pruritus Numeric Rating Scale (PP-NRS) score ≥ 4; **OR**
  - Incapacitation due to AD lesion location (e.g., head and neck, palms, soles, or genitalia); **AND**
- Prescribed by, or in consultation with, a dermatologist, allergist/immunologist, or other specialist in the treatment of atopic dermatitis; **AND**
- Trial and failure, contraindication, or intolerance to ≥ 1 agent in 2 or more of the following categories (total prior agent use of ≥ 90 days):
  - Topical corticosteroid of medium to high potency (e.g., mometasone, fluocinolone) unless inappropriate for the location (e.g., face, groin); **AND**
  - Topical calcineurin inhibitor (i.e., tacrolimus or pimecrolimus); **OR**
  - Immunosuppressive systemic agents (e.g., cyclosporine, azathioprine, methotrexate, mycophenolate mofetil, etc.); **AND**
- Trial and failure, allergy, contraindication (including potential drug-drug interactions with other medications) or intolerance of 1 preferred injectable (Adbry, Dupixent) agent; **AND**
- Nemluvio will be taken with topical corticosteroids and/or calcineurin inhibitors (e.g., pimecrolimus, tacrolimus); **AND**
- Patient must meet the minimum age recommended by the package insert for this FDA approved indication.





**Prurigo Nodularis:**

- Diagnosis of prurigo nodularis; **AND**
- At least 20 nodular lesions; **AND**
- Other causes of pruritis have been ruled out; **AND**
- Trial and failure, contraindication, or intolerance to one of the following:
  - Moderate to super potent topical corticosteroids [e.g., betamethasone dipropionate, (augmented), fluocinonide 0.1%, flurandrenolide, betamethasone dipropionate 0.05%, clobetasol propionate 0.025%, or desoximetasone 0.05%] for a minimum of 2 weeks; **OR**
  - Narrowband ultraviolet B (NBUVB) phototherapy or psoralen plus ultraviolet A (PUVA) phototherapy; **AND**
- Trial and failure, contraindication, or intolerance to Dupixent; **AND**
- Patient must meet the minimum age recommended by the package insert for this FDA-approved indication.

**Renewal Criteria:**

- Patient must continue to meet initial approval criteria; **AND**
- Patient must have disease improvement and/or stabilization based on an objective measure

**Quantity Limit:** 2 pens (60 mg) per 28 days

Preferred Agents	Non-Preferred Agents
Adbry autoinjector <sup>AE, CC, QL</sup>	<b>Ebglyss</b> <sup>AE, CC, QL</sup>
Adbry syringe <sup>AE, CC, QL</sup>	Elidel
Dupixent pen <sup>CC, QL</sup>	<b>Nemluvio</b> <sup>AE, CC, QL</sup>
Dupixent syringe <sup>CC, QL</sup>	
Eucrisa <sup>CC, QL</sup>	
Opzelura cream <sup>CC, AE</sup>	
pimecrolimus cream	
tacrolimus ointment	



**Nefly®**

**Self-injectable Epinephrine: Nonpreferred**

**Approval Duration: 6 months initial, 1 year renewal**

- *Epinephrine acts on both alpha- and beta-adrenergic receptors and is a potent vasoconstrictor via its effects on the alpha- receptors.*

**Approval Criteria:**

- Patient has had a trial and failure, allergy, contraindication (including potential drug-drug interactions with other medications), or intolerance of 1 preferred agent.

**Quantity Limit: 2 bottles per fill**

Preferred Agents	Non-Preferred Agents
<b>epinephrine 0.3 mg autoinjector (all manufacturers) <sup>QL</sup></b>	Auvi-Q autoinjector <sup>QL</sup>
<b>epinephrine 0.15 mg autoinjector (all manufacturers) <sup>QL</sup></b>	<b>Nefly <sup>QL</sup></b>
EpiPen <sup>QL</sup> EpiPen Jr. <sup>QL</sup>	Symjepi <sup>QL</sup>

**Miplyffa™**

**Non-PDL**

**Approval Duration: 6 months initial, 1 year renewal**

- *Miplyffa is a first-in-class oral heat shock protein (HSP) amplifier that can stabilize lysosomal membranes and improve lysosomal function so that cells can clear away waste.*

**Initial Approval Criteria:**

- Diagnosis of Niemann-Pick Disease Type C (NPC) [ICD-10 code E75.242]; **AND**
- Confirmed diagnosis of NPC by ≥ 1 of the following:
  - Positive genetic test for mutations on both alleles of NPC1 or NPC2; **OR**
  - Positive genetic test for mutations on one allele NPC1 or NPC2; **AND**
    - Elevated biomarker; **OR**
    - Positive filipin staining; **AND**
- Prescribed by, or in consultation with, a neurologist or geneticist or other specialist in the treatment of Niemann-Pick Disease Type C; **AND**
- Prescriber attests patient presents with at least one neurological symptom of the disease (e.g., hearing loss, ataxia, dystonia, seizures, speech delay); **AND**
- Prescriber attests medication will be used in combination with miglustat; **AND**
- Patient must meet the minimum age recommended by the package insert.



**Renewal Criteria:**

- Prescriber provides documentation (i.e., NPC Neurologic Severity Scale, cognitive function tests, motor function assessment, etc.) that patient has experienced disease improvement or stabilization or a reduction in disease progression

**Age Limit:** 2 years of age or older

**Quantity Limit:** 3 capsules per day

**Aqneursa™**

**Non-PDL**

**Approval Duration: 3 months initial, 1 year renewal**

- *Aqneursa is modified amino acid (acetylleucine) kilograms. Once ingested, ubiquitous monocarboxylate transporters deliver Aqneursa to all tissues and is thought to serve as a neuroprotectant that reduces neuroinflammation and stabilizes the nerve cells responsible for balance and the coordination of movement.*

**Initial Approval Criteria:**

- Diagnosis of Niemann-Pick Disease Type C (NPC) [ICD-10 code E75.242]; **AND**
- Confirmed diagnosis of NPC by ≥ 1 of the following:
  - Positive genetic test for mutations on both alleles of NPC1 or NPC2; **OR**
  - Positive genetic test for mutations on one allele NPC1 or NPC2; **AND**
    - Elevated biomarker; **OR**
    - Positive filipin staining; **AND**
- Prescribed by, or in consultation with, a neurologist or geneticist or other specialist in the treatment of Niemann-Pick Disease Type C; **AND**
- Prescriber attests patient presents with at least one neurological symptom of the disease (e.g., hearing loss, ataxia, dystonia, seizures, speech delay); **AND**
- Patient must meet the minimum age and weight recommended by the package insert for the provided indication
  - ≥ 4 years of age
  - ≥ 15 kg

**Renewal Criteria:**

- Prescriber provides documentation (i.e., NPC Neurologic Severity Scale, cognitive function tests, motor function assessment, etc.) that patient has experienced disease improvement or stabilization or a reduction in disease progression

**Age Limit:** 4 years of age or older

**Quantity Limit:** 4 packets (4 grams) per day



## FULL CLASS REVIEWS

### Antibiotics: Gastrointestinal

#### Class Selection & Guidelines

- DMS to select preferred agent(s) based on economic evaluation; however, at least 2 unique chemical entities should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the Antibiotics, Gastrointestinal class, require PA until reviewed by the P&T Committee.

Preferred Agents	Non-Preferred Agents
metronidazole 250 mg, 500 mg tablet	Aemcolo
neomycin	Dificid suspension, tablet <sup>CC, QL</sup>
tinidazole	<b>Firvanq</b> <sup>CC</sup>
<b>vancomycin</b> capsule, <b>solution</b> <sup>CC</sup>	Flagyl
Xifaxan <sup>CC, QL</sup>	Likmez
	metronidazole capsule
	metronidazole 125 mg tablet
	nitazoxanide
	paromomycin
	Solosec <sup>AE, CC, QL</sup>
	Vancocin
	Vowst <sup>AE, CC, QL</sup>

### Antibiotics: Vaginal

#### Class Selection & Guidelines

- DMS to select preferred agent(s) based on economic evaluation; however, at least 2 unique chemical entities should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the Antibiotics, Vaginal class, require PA until reviewed by the P&T Committee.

Preferred Agents	Non-Preferred Agents
Cleocin Ovule	Cleocin cream
<b>clindamycin vaginal 2% cream</b>	<b>Clindesse vaginal cream</b>
metronidazole vaginal 0.75% gel	metronidazole vaginal 1.30% gel
Nuversa gel	Vandazole gel
	Xaciato gel



## Antibiotics: Penicillins

### Class Selection & Guidelines

- DMS to select preferred agent(s) based on economic evaluation; however, at least 2 unique chemical entities should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the Antibiotics, Penicillins class, require PA until reviewed by the P&T Committee.

Preferred Agents	Non-Preferred Agents
amoxicillin	amoxicillin/clavulanate ER
<b>amoxicillin/clavulanate chewable tablet</b> , tablet,	Augmentin
suspension	
Ampicillin capsule	Augmentin XR
Dicloxacillin capsule	
penicillin V potassium tablet, suspension	

## Antibiotics: Sulfonamides, Folate Antagonists

### Class Selection & Guidelines

- DMS to select preferred agent(s) based on economic evaluation; however, at least 2 unique chemical entities should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the Antibiotics: Sulfonamides, Folate Antagonists class, require PA until reviewed by the P&T Committee.

Preferred Agents	Non-Preferred Agents
sulfamethoxazole/trimethoprim	Bactrim
<b>Sulfatrim suspension</b>	Bactrim DS
trimethoprim	sulfadiazine

## Antifungal, Oral

### Class Selection & Guidelines

- DMS to select preferred agent(s) based on economic evaluation; however, at least 2 unique chemical entities should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the Antifungal, Oral class, require PA until reviewed by the P&T Committee.

Preferred Agents	Non-Preferred Agents
clotrimazole troche	Ancobon
fluconazole suspension, tablet	Brexafemme CC, QL



Preferred Agents	Non-Preferred Agents
griseofulvin suspension	Cresemba
itraconazole capsule <sup>CC, QL</sup>	Diflucan
<b>ketoconazole</b>	flucytosine
nystatin suspension, tablets	griseofulvin microsize tablet, ultramicrosize tablet
terbinafine	itraconazole solution
	Noxafil
	Oravig
	posaconazole
	Sporanox <sup>QL</sup>
	Tolsura
	Vfend
	Vivjoa <sup>CC, QL</sup>
	voriconazole

## Hepatitis C Agents: Interferons and Ribavirins

### Class Selection & Guidelines

- DMS to select preferred agent(s) based on economic evaluation; however, at least 2 unique chemical entities should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the Immunological and Genetic Hepatitis C Agents: Interferons and Ribavirins class, require PA until reviewed by the P&T Committee.

Preferred Agents	Non-Preferred Agents
<b>PEGASYS</b> syringe, <b>via</b> <sup>CC, QL</sup> ribavirin capsule, tablet <sup>CC</sup>	

## Chronic Obstructive Pulmonary Disease (COPD) Agents

### Class Selection & Guidelines

- DMS to select preferred agent(s) based on economic evaluation; however, at least 2 unique chemical entities should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the Chronic Obstructive Pulmonary Disease (COPD) Agents class, require PA until reviewed by the P&T Committee.

Preferred Agents	Non-Preferred Agents
albuterol-ipratropium inhalation solution <sup>QL</sup>	Bevespi Aerosphere <sup>QL</sup>
Anoro Ellipta <sup>QL</sup>	Daliresp tablet <sup>CC, QL</sup>
Atrovent HFA <sup>QL</sup>	Duaklir Pressair
<b>Breztri Aerosphere</b> <sup>CC, QL</sup>	Incruse Ellipta <sup>QL</sup>
Combivent Respimat <sup>QL</sup>	Ohtuvayre <sup>AE, CC, QL</sup>
ipratropium inhalation solution <sup>QL</sup>	Spiriva Respimat <sup>QL</sup>
<b>roflumilast tablet</b> <sup>CC, QL</sup>	Tiotropium <sup>QL</sup>
Spiriva Handihaler <sup>QL</sup>	



Preferred Agents	Non-Preferred Agents
Stiolto Respimat <sup>QL</sup>	Trelegy Ellipta <sup>CC, QL</sup>
	Tudorza Pressair <sup>QL</sup>
	Yupelri solution <sup>CC, QL</sup>

## Epinephrine, Self-Injectable

### Class Selection & Guidelines

- DMS to select preferred agent(s) based on economic evaluation; however, at least 2 unique chemical entities should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the Epinephrine, Self-Injectable class, require PA until reviewed by the P&T Committee.

Preferred Agents	Non-Preferred Agents
<b>epinephrine 0.3 mg autoinjector (all manufacturers)<sup>QL</sup></b>	Auvi-Q autoinjector <sup>QL</sup>
<b>epinephrine 0.15 mg autoinjector (all manufacturers)<sup>QL</sup></b>	<b>Nefly<sup>QL</sup></b>
EpiPen <sup>QL</sup>	Symjepi <sup>QL</sup>
EpiPen Jr. <sup>QL</sup>	

## Glucocorticoids, Inhaled

### Class Selection & Guidelines

- DMS to select preferred agent(s) based on economic evaluation; however, at least 2 unique chemical entities should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the Glucocorticoids, Inhaled class, require PA until reviewed by the P&T Committee.

Preferred Agents	Non-Preferred Agents
Asmanex Twisthaler <sup>QL</sup>	Alvesco <sup>QL</sup>
budesonide inhalation suspension <sup>AE, QL</sup>	ArmonAir Digihaler <sup>QL</sup>
Flovent HFA <sup>QL</sup>	Arnuity Ellipta <sup>QL</sup>
fluticasone propionate HFA <sup>QL</sup>	Asmanex HFA <sup>QL</sup>
<b>Pulmicort Flexhaler<sup>QL</sup></b>	Flovent Diskus <sup>QL</sup>
	Pulmicort Respules <sup>QL</sup>
	Qvar Redihaler



## CONSENT AGENDA REVIEWS

For the following therapeutic classes, there were no changes in PDL status:

### Therapeutic Classes

- **Antibiotics, Cephalosporins 1<sup>st</sup> Generation**
- **Antibiotics, Cephalosporins 2<sup>nd</sup> Generation**
- **Antibiotics, Cephalosporins 3<sup>rd</sup> Generation**
- **Antibiotics, Inhaled**
- **Antibiotics, Macrolides**
- **Antibiotics, Oxazolidinones**
- **Antibiotics, Quinolones**
- **Antibiotics, Tetracyclines**
- **Antihistamines, Minimally Sedating**
- **Antiretrovirals, HIV/AIDS**
- **Antivirals, Oral**
- **Bronchodilators, Beta Agonist**
- **Hepatitis B Agents**
- **Hepatitis C Agents: Direct-Acting Antivirals**
- **Intranasal Rhinitis Agents**
- **Leukotriene Modifiers**