



Commissioner for the Department for Medicaid Services Selections for Preferred Products

This is a summary of the final Preferred Drug List (PDL) selections made by the Commissioner of 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 **November 15, 2018**, and the recommendations delivered by the P&T Committee members in attendance.

New Products to Market

Lucemyra[™] – Non-prefer in the PDL class: Opiate Dependence Treatments Length of Authorization: 5 days

• Lucemyra[™] (lofexidine) is a central alpha-2 adrenergic agonist indicated for mitigation of opioid withdrawal symptoms to facilitate abrupt opioid discontinuation in adults.

Criteria for Approval:

- Medication is being used to mitigate opioid withdrawal symptoms and facilitate abrupt discontinuation of opioids; AND
- Patient is NOT pregnant or breastfeeding; AND
- Patient does NOT have a prolonged QT interval (> 450 msec for males, > 470 msec for females);
 AND
- If patient is currently taking methadone, prescriber attestation that a baseline electrocardiogram (ECG) has been performed; AND
- Patient has tried and failed, had a contraindication to, or experienced an adverse reaction/intolerance to buprenorphine OR methadone; AND
- Patient has tried and failed, had a contraindication to, or experienced an adverse reaction/intolerance to clonidine; AND
- Prescriber to provide verbal attestation of a comprehensive treatment plan between provider and patient; AND
- Prescriber to provide verbal attestation that the patient is capable of and instructed how to self-monitor for hypotension, orthostasis, bradycardia, and associated symptoms; AND
- Prescriber to provide verbal attestation that patient is NOT receiving prescribed concurrent opioid medication based on current medication list/orders, medical records, patient history and verified by KASPER query; AND
- Prescriber to provide verbal attestation that the patient has been provided with a tapering schedule and instructions on when to contact their healthcare provider for further guidance.

Age Limit: > 18 years

Quantity Limit: 48 tablets with 1 refill (96 tabs per treatment course; 1 course per year)





Drug Class	Preferred Agents	Non-Preferred Agents
Opiate Dependence	Suboxone® film ^{CC, QL}	Bunavail® ^{QL}
Treatments		buprenorphine ^{CC, QL}
		buprenorphine/naloxone ^{QL}
		Lucemyra™ ^{CC, QL}
		Probuphine® CC, QL
		Sublocade™ ^{CC, QL}
		Zubsolv ^{® QL}

Tibsovo® – Prefer with clinical criteria in the PDL class: Oncology, Oral – Hematologic Cancer (Oral Oncology, Hematologic Cancer)

Length of Authorization: 1 year

• Tibsovo (ivosidenib) is an isocitrate dehydrogenase-1 (IDH1) indicated for the treatment of adult patients with relapsed or refractory acute myeloid leukemia (AML) with a susceptible IDH1 mutation as detected by the Abbott RealTime™ IDH1 FDA-approved companion diagnostic.

Criteria for Approval:

- Diagnosis acute myeloid leukemia; AND
- Documentation showing susceptible isocitrate dehydrogenase-1 (IDH1) mutation, as detected by an FDA-approved test; AND
- Must be used as single agent; AND
- Patient has relapsed or refractory disease; OR
- Patient is not a candidate for intensive remission induction therapy; OR
- Patient declines intensive therapy.

Renewal Criteria

- Patient continues to meet the above conditions; AND
- Evidence of tumor response or lack of disease progression.

Age Limit: > 18 years

Quantity Limit: 2 tablets per day

Drug Class	Preferred Agents	Non-Preferred Agents
Oral Oncology,	Alkeran®	Bosulif® QL
Hematologic Cancer	Gleevec® QL	Calquence® CC, QL
	hydroxyurea	Farydak® QL
	Imbruvica® CC, QL	Hydrea®
	Jakafi ^{® CC, QL}	Iclusig® QL
	Leukeran®	Idhifa® ^{CC, QL}
	mercaptopurine	imatinib ^{QL}
	Purixan®	melphalan
	Revlimid [®]	Ninlaro®
	Rydapt® CC, QL	Pomalyst®
	Sprycel® QL	Tasigna® ^{QL}
	Thalomid [®]	Venclexta™ ^{QL}





Drug Class	Preferred Agents	Non-Preferred Agents
	Tibsovo® ^{cc, QL}	
	Zolinza ^{® QL}	
	Zydelig ^{® CC, QL}	

Braftovi[™] – Prefer with clinical criteria in the PDL class: *Oncology, Oral – Skin (Oral Oncology, Skin Cancer)*

Length of Authorization: 1 year

 Braftovi[™] (encorafenib) is a kinase inhibitor indicated, in combination with binimetinib, for the treatment of patients with unresectable or metastatic melanoma with a BRAF V600E or V600K mutation, as detected by an FDA-approved test.

Criteria for Approval:

- Diagnosis of unresectable or metastatic melanoma with a BRAF V600E or V600K mutation, as detected by an FDA-approved test; AND
- Used in combination with binimetinib.

Renewal Criteria:

- Meet initial approval criteria; AND
- Evidence of tumor response or lack of disease progression.

Age Limit: ≥ 18 years

Quantity Limit: 75 mg: 6 per day; 50 mg: 4 per day

Mektovi® – Prefer with clinical criteria in the PDL class: Oncology, Oral – Skin (Oral Oncology, Skin Cancer)

Length of Authorization: 1 year

• Mektovi® (binimetinib) is a kinase inhibitor indicated, in combination with encorafenib, for the treatment of patients with unresectable or metastatic melanoma with a BRAF V600E or V600K mutation, as detected by an FDA-approved test.

Criteria for Approval:

- Diagnosis of unresectable or metastatic melanoma with a BRAF V600E or V600K mutation, as detected by an FDA-approved test; AND
- Used in combination with encorafenib.

Renewal Criteria:

- Meet initial approval criteria; AND
- Evidence of tumor response or lack of disease progression.

Age Limit: ≥ 18 years Quantity Limit: 6 per day

Drug Class	Preferred Agents	Non-Preferred Agents
Oral Oncology, Skin	Erivedge™ CC, QL	Cotellic™ CC, QL
Cancer	Braftovi™ ^{CC, QL}	Zelboraf™ ^{CC,QL}
	Mekinist™ CC, QL	
	Mektovi ^{® CC, QL}	
	Odomzo® CC, QL	
	Tafinlar® CC, QL	





Doptelet® – Non-prefer in the PDL class: *Thrombopoiesis Stimulating Agents* **Length of Authorization:** Date of Service; 1 fill per procedure

• Doptelet® (avatrombopag), a thrombopoietin (TPO) receptor agonist, is indicated for the treatment of thrombocytopenia in adult patients with chronic liver disease (CLD) who are scheduled to undergo a procedure.

Criteria for Approval:

- Diagnosis of chronic liver disease; AND
- Documentation of platelet count < 50 x 10⁹/L; AND
- Dosed per FDA-approved labeling (10 tablets per 5 days for platelets \geq 40 x 10 9 /L or 15 tablets per 5 days for platelets < 40 x 10 9 /L); AND
- Confirmation of a scheduled invasive procedure occurring 5 to 8 days following the last dose of avatrombopag.

Age Limit: ≥18 years

Quantity Limit: 15 tablets per fill

Mulpleta® – Non-prefer in the PDL class: *Thrombopoiesis Stimulating Agents* Length of Authorization: Date of Service; 1 fill per procedure

• Mulpleta® (lusutrombopag), a thrombopoietin (TPO) receptor agonist, is indicated for the treatment of thrombocytopenia in adult patients with chronic liver disease (CLD) who are scheduled to undergo a procedure.

Criteria for Approval:

- Diagnosis of chronic liver disease (CLD); AND
- Documentation of platelet count < 50 x 10⁹/L; AND
- NOT have severe hepatic impairment (Child-Pugh class C), absence of hepatopetal blood flow, a prothrombotic condition other than CLD or a history of splenectomy, partial splenic embolization, or thrombosis; AND
- Confirmation of a scheduled invasive procedure occurring 2 to 8 days following the last dose of lusutrombopag.

Age Limit: ≥18 years

Quantity Limit: 7 tablets per fill

Drug Class	Preferred Agents	Non-Preferred Agents
Thrombopoiesis	Promacta® ^{CC}	Doptelet ^{® CC, QL}
Stimulating Agents		<mark>Mulpleta® ^{cc, qL}</mark>
		Nplate™ ^{CC}
		Tavalisse™ ^{CC, QL}





Criteria Review

Movement Disorders: Austedo® (deutetrabenazine)

Austedo® (deutetrabenazine) is a vesicular monoamine transporter 2 (VMAT2) inhibitor approved for the treatment of chorea associated with Huntington's disease and the treatment of tardive dvskinesia.

Current criteria: Trial and failure of a preferred agent, unless contraindicated.

Recommended criteria (in addition to current criteria):

Length of Authorization: 1 year

Criteria for Approval:

- Patient is not concurrently using monoamine oxidase (MAO) inhibitors (e.g., isocarboxazid, phenelzine, rasagiline, safinamide, selegiline, tranylcypromine, etc within 14 days) OR reserpine (within 20 days) OR another VMAT2 inhibitor (e.g., tetrabenazine, valbenazine);
 AND
- Patient is not pregnant; **AND**
- Patient does not have hepatic impairment (e.g., Child-Pugh A-C); AND
- Patient meets the following criteria for either Huntington's chorea or tardive dyskinesia:

Huntington's Chorea

- Patient is diagnosed with chorea related to Huntington's disease; AND
- Patient is able to swallow; AND
- Patient does not have the following conditions:
 - History of, or current, untreated or inadequately treated depression; OR
 - o Suicidal ideation.

Tardive Dyskinesia

- Diagnosis of tardive dyskinesia; AND
- Patient is able to swallow; **AND**
- Documentation that AIMS test has been completed (e.g., score or copy of AIMS assessment);
 AND
- Prescribed by or in consultation with a neurologist or psychiatrist (or other mental health provider), provided patient has reasonable access; **AND**
- Documentation or claims history of current or former chronic patient use of a dopamine antagonist (e.g., antipsychotic, metoclopramide, prochlorperazine, droperidol, promethazine, etc.).

Renewal Criteria:

- Patient continues to meet criteria defined for initial approval; AND
- Documentation of improvement in symptoms associated with respective condition (e.g., tardive dyskinesia or Huntington's chorea).

Age Limit: ≥ 18 years **Quantity Limit**: 4 per day

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Movement Disorders: Ingrezza™ (valbenazine)

Ingrezza[™] (valbenazine) is a vesicular monoamine transporter 2 (VMAT2) inhibitor indicated for the treatment of adults with tardive dyskinesia (TD). Tardive dyskinesia is a side effect that can be seen in patients on long treatments of antipsychotic medications and medications used for gastrointestinal disease.

Current criteria: Trial and failure of a preferred agent, unless contraindicated.

Recommended criteria (in addition to current criteria):

Length of Authorization: 1 year

Criteria for Approval:

- Diagnosis of tardive dyskinesia; AND
- Documentation that AIMS test has been completed (e.g., score or copy of AIMS assessment);
 AND
- Prescribed by or in consultation with a neurologist or psychiatrist (or other mental health provider), provided patient has reasonable access; AND
- Documentation or claims history of current or former chronic use of a dopamine antagonist (e.g., antipsychotic, metoclopramide, prochlorperazine, droperidol, promethazine, etc.); AND
- NO concurrent use of MAO inhibitors (e.g., isocarboxazid, phenelzine, rasagiline, safinamide, selegiline, tranylcypromine, etc.) or strong CYP3A4 inducers (e.g., carbamazepine, phenytoin, phenobarbital, rifampin and related agents, St. John's wort, etc.).

Renewal Criteria:

- Patient continues to meet criteria defined for initial approval; AND
- Attestation or documentation of improvement in TD symptoms.

Age Limit: ≥ 18 years **Quantity Limit**: 1 per day

Full Class Reviews

Acne Agents, Topical

Class Selection & Guidelines

Topical Acne Agents

- DMS to select preferred agent(s) based on economic evaluation; however, at least 5 products unique chemical entities or combinations should be preferred.
- Agents not selected as preferred will be considered non-preferred and require PA.
- For any new chemical entity in the *Topical Acne Agents* class, require PA until reviewed by the P&T Advisory Committee.





Drug Class	Preferred Agents	Non-Preferred Agents
Topical Acne Agents	clindamycin solution	Acanya™
	clindamycin/benzoyl peroxide (generic for	Aczone™
	BenzaClin® or Duac®; excluding pump)	adapalene cream, gel
	Differin® cream, gel	adapalene/benzoyl peroxide
	erythromycin solution	Atralin™
	Retin-A® cream, gel	Avar™
		Avar E™
		Avar E LS™
		Avar LS™
		Avita®
		<mark>BenzaClin®</mark>
		Benzamycin®
		BenzePro™
		benzoyl peroxide cleanser, kit,
		microspheres, gel, foam, medicated
		pad, towlette
		BP 10-1®
		BPO®
		BPO-5®
		BPO-10®
		BP Wash™
		Brevoxyl®
		Cleocin-T®
		Clindacin PAC™
		Clindagel®
		clindamycin gel, foam, lotion, medicated
		swab
		clindamycin/benzoyl peroxide <mark>pump</mark>
		clindamycin/tretinoin
		dapsone gel
		DermaPak Plus Kit
		Differin® lotion
		Duac®
		Effaclar Duo®
		Epiduo™
		Epiduo Forte™
		Erygel®
		Erythromycin gel, medicated swab
		erythromycin/benzoyl peroxide
		Fabior®
		Inova™
		Inova™ 4/1
		Inova™ 8/2
		Klaron®
		Neuac®
		Pacnex®





Drug Class	Preferred Agents	Non-Preferred Agents
		Panoxyl®
		Persa-Gel®
		PR benzoyl peroxide
		OC8®
		Onexton™
		Ovace®
		Ovace Plus®
		Retin-A Micro®
		Rosula®
		sodium sulfacetamide 10% CLNSG
		sodium sulfacetamide/sulfur 10-4% pad
		sodium sulfacetamide/sulfur cleanser
		sodium sulfacetamide/sulfur/urea
		SSS 10-5®
		sulfacetamide cleanser
		sulfacetamide/urea
		Sumadan™
		Sumadan™ XLT
		Sumaxin®
		Tazorac®
		tazarotene
		Tretin-X™
		tretinoin
		tretinoin (generic Atralin™)
		tretinoin microsphere
		Vanoxide-HC®
		Ziana™

Anticholinergics/ Antispasmodics

Class Selection & Guidelines

Anticholinergics/ Antispasmodics

- DMS to select preferred agent(s) based on economic evaluation; however, at least 4 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 *Anticholinergics/Antispasmodics* class, require PA until reviewed by the P&T Advisory Committee.

Drug Class	Preferred Agents	Non-Preferred Agents
Antispasmodics/	dicyclomine	Anaspaz®
Anticholinergics	glycopyrrolate	Bentyl®
	hyoscyamine	chlordiazepoxide/clidinium
	methscopolamine	Cuvposa®
		Donnatal®





Drug Class	Preferred Agents	Non-Preferred Agents
		Hyosyne®
		Levbid®
		Levsin®
		Librax®
		Oscimin®
		Phenohytro®
		propantheline
		Robinul®
		Robinul Forte®
		Symax®

Antiemetics & Antivertigo Agents

Class Selection & Guidelines

Anti-Emetics: Other

- DMS to select preferred agent(s) based on economic evaluation; however, at least 5 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 *Anti-Emetics: Other* class, require PA until reviewed by the P&T Committee.

Oral Anti-Emetics: 5-HT3 Antagonists

- DMS to select preferred agent(s) based on economic evaluation; however, at least 1 unique chemical entity should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the *Oral Anti-Emetics:* 5-HT3 Antagonists class, require PA until reviewed by the P&T Committee.

Oral Anti-Emetics: Delta-9-THC Derivatives

- DMS to select preferred agent(s) based on economic evaluation; however, at least 1 unique chemical entity should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the *Oral Anti-Emetics: Delta-9-THC Derivatives* class, require PA until reviewed by the P&T Committee.

Oral Anti-Emetics: NK-1 Antagonists

- DMS to select preferred agent(s) based on economic evaluation; however, at least 1 unique chemical entity should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the *Oral Anti-Emetics: NK-1 Antagonists* class, require PA until reviewed by the P&T Committee.

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Drug Class	Preferred Agents	Non-Preferred Agents
Anti-Emetics: Other	meclizine	Compazine®
	metoclopramide oral solution, tablets	Compro®
	prochlorperazine	Bonjesta® ^{CC, QL}
	promethazine syrup, tablets	Diclegis™ ^{CC, QL}
	promethazine 12.5, 25 mg suppositories	metoclopramide ODT
	Transderm-Scop®	Phenadoz®
		Phenergan®
		promethazine 50 mg suppositories
		Reglan®
		scopolamine transdermal system
		Tigan®
		<mark>trimethobenzamide</mark>
Oral Anti-Emetics: 5-	ondansetron	Aloxi® QL
HT3 Antagonists		Anzemet®
		granisetron
		Sancuso® cc, QL
		Zofran®
		Zuplenz®
Oral Anti-Emetics:	Emend® capsules QL	Akynzeo® QL
NK-1 Antagonists		aprepitant ^{QL}
-		Emend® powder packet ^{QL}
		Varubi® ^{CC, QL}
Oral Anti-Emetics: Δ-	dronabinol ^{CC, QL}	Cesamet® CC, QL
9-THC Derivatives		Marinol® CC, QL
		Syndros™ ^{CC, QL}

Antifungals, Topical

Class Selection & Guidelines

Topical Antifungal Agents

- DMS to select preferred agent(s) based on economic evaluation; however, at least 4 unique chemical entities or combinations should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the *Topical Antifungal Agents* class, require PA until reviewed by the P&T Committee.





Drug Class	Preferred Agents	Non-Preferred Agents
Topical Antifungal	clotrimazole cream, solution	Ciclodan® cream, kit, solution
Agents	clotrimazole/betamethasone cream	ciclopirox
	ketoconazole cream, shampoo	clotrimazole/betamethasone lotion
	nystatin cream, ointment, powder	econazole
	nystatin/triamcinolone cream, ointment	Ertazczo®
		Exelderm®
		Extina®
		Jublia® ^{cc}
		Kerydin™ ^{CC}
		ketoconazole foam
		Ketodan™
		Loprox®
		Lotrimin®
		Lotrisone®
		luliconazole
		Luzu®
		Mentax®
		naftifine
		Naftin®
		Nizoral Shampoo®
		Nyamyc®
		nystatin/triamcinolone cream
		Nystop®
		Oxistat®
		oxiconazole
		Penlac®
		Therazole Pak™ ^{QL}
		Vusion® ^{cc}

Antiparasitics, Topical

Class Selection & Guidelines

Topical Antiparasitic Agents

- 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 *Topical Antiparasitic Agents* class, require PA until reviewed by the P&T Committee.





Drug Class	Preferred Agents	Non-Preferred Agents
Topical Antiparasitic	Natroba®	Crotan™
Agents	permethrin 5% cream	Elimite™
	Sklice [®]	Eurax®
		lindane
		malathion
		Ovide®
		spinosad
		Ulesfia®

Bile Salts

Class Selection & Guidelines

- DMS to select preferred agent(s) based on economic evaluation; however, at least generic ursodiol capsules and tablets should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the *Bile Salts* class, require PA until reviewed by the P&T Advisory Committee.

Drug Class	Preferred Agents	Non-Preferred Agents
Bile Salts	ursodiol capsules, tablets	Actigall®
		Chenodal®
		Cholbam®
		Ocaliva®
		Urso®/Urso Forte®

Cytokine and CAM Antagonists

Class Selection & Guidelines

Immunomodulators

- 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 *Immunomodulators* class, require PA until reviewed by the P&T Advisory Committee.

New agent in the class: IlumyaTM

Non-prefer in the PDL class: Cytokine and CAM Antagonists (Immunomodulators)

Length of Authorization: 1 year

• Ilumya[™] (tildrakizumab-asmn), a high affinity, humanized IgG1 kappa monoclonal antibody that targets the p19 subunit of interleukin 23 (IL-23), is indicated for the treatment of adults with moderate-to-severe plaque psoriasis (PSO) who are candidates for systemic therapy or phototherapy.





Criteria for Approval:

- Diagnosis of moderate to severe plaque psoriasis; AND
- Symptoms persistent for ≥ 6 months with at least 1 of the following:
 - o Involvement of at least 10% of body surface area (BSA); OR
 - o Psoriasis Area and Severity Index (PASI) score of 12 or greater; OR
 - o Incapacitation due to plaque location (i.e., head and neck, palms, soles or genitalia); AND
- Negative tuberculosis (TB) screening prior to initiating treatment; AND
- Trial and failure of 2 of the following therapies:
 - Methotrexate
 - Cyclosporine
 - o Oral retinoid (e.g., Soriatane®, acitretin)
 - Topical corticosteroids
 - o Phototherapy/UV light
 - Coal tar preparations; AND
- Trial and failure of, or contraindication to, a preferred immunomodulator (i.e., Enbrel® or Humira®); AND
- NOT to be used in combination with a TNF inhibitor, anakinra, abatacept, apremilast or other biologic response modifier.

Renewal Criteria:

- Patient continues to meet criteria identified above; AND
- Ongoing monitoring for TB; AND
- Disease response as indicated by improvement in signs and symptoms compared to baseline, such as redness, thickness, scaliness, and/or the amount of surface area involvement.

Age Limit: ≥ 18 years

Quantity Limit: 1 syringe per fill

Drug Class	Preferred Agents	Non-Preferred Agents
Immunomodulators	Enbrel® CC QL	Actemra® ^{CC, QL}
	Cosentyx ^{® cc, QL}	Cimzia® CC, QL
	Humira® CC, QL	Entyvio™ CC, QL
		llumya™ ^{CC, QL}
		Kevzara® CC, QL
		Kineret® CC, QL
		Orencia® CC, QL
		Otezla® CC, QL
		Siliq™ CC, QL
		Simponi™ ^{CC, QL}
		Stelara™ ^{CC, QL}
		Taltz® CC, QL
		Tremfya™ ^{CC, QL}
		Xeljanz® ^{CC, QL}
		Xeljanz® XR ^{cc, QL}





Multiple Sclerosis Agents

Class Selection & Guidelines

- DMS to select preferred agent(s) based on economic evaluation; however, at least 5 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 *Multiple Sclerosis Agents* class, require PA until reviewed by the P&T Advisory Committee.

Criteria review: GilenyaTM (fingolimod)

Current criteria and PDL status: Preferred with clinical PA

• Requires a step through an injectable agent (e.g., Avonex®, Betaseron®, Copaxone®, Rebif®).

Recommended PDL status: Preferred

• Clinical step edit is removed from Gilenya and is available without a PA.

Drug Class	Preferred Agents	Non-Preferred Agents
Multiple Sclerosis	Avonex® QL	Ampyra™ ^{QL, CC}
Agents	Avonex Administration Pack® QL	Aubagio® ^{QL}
	Betaseron® QL	Copaxone® 40 mg ^{QL}
	Copaxone® 20 mg QL	Extavia® QL
	Gilenya™ ^{QL}	glatiramer acetate ^{QL}
	Rebif® QL	Glatopa™ ^{QL}
		Plegridy®
		Tecfidera™ ^{QL}

Ophthalmics for Allergic Conjunctivitis

Class Selection & Guidelines

Ophthalmic Antihistamines

- DMS to select preferred agent(s) based on economic evaluation; however, at least 1 unique chemical entity should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the *Ophthalmic Antihistamines* class, require PA until reviewed by the P&T Committee.

Ophthalmic Mast Cell Stabilizers

- DMS to select preferred agent(s) based on economic evaluation; however, at least 1 unique chemical entity should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the *Ophthalmic Mast Cell Stabilizers* class, require PA until reviewed by the P&T Committee.





Drug Class	Preferred Agents	Non-Preferred Agents
Ophthalmic	olopatadine 0.1% (generic for Patanol®)	azelastine
Antihistamines	Pataday™	Bepreve™
	Pazeo™	Elestat™
		Emadine®
		epinastine
		Lastacaft™
		olopatadine <mark>0.2% (generic for Pataday™)</mark>
		Optivar®
		Patanol®
Ophthalmic Mast	cromolyn sodium	Alocril®
Cell Stabilizers		Alomide®

Ophthalmic Antibiotics

Class Selection & Guidelines

Ophthalmic Antibiotics, Non-Quinolones

- DMS to select preferred agent(s) based on economic evaluation; however, at least 5 unique chemical entities or combinations should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the *Ophthalmic Antibiotics, Non-Quinolones* class, require PA until reviewed by the P&T Committee.

Ophthalmic Antifungals

- DMS to select preferred agent(s) based on economic evaluation.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the *Ophthalmic Antifungals* class, require PA until reviewed by the P&T Committee.

Ophthalmic Macrolides

- DMS to select preferred agent(s) based on economic evaluation; however, at least 1 unique chemical entity should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the *Ophthalmic Macrolides* class, require PA until reviewed by the P&T Committee.

Ophthalmic Quinolones

- DMS to select preferred agent(s) based on economic evaluation; however, at least 3 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 *Ophthalmic Quinolones* class, require PA until reviewed by the P&T Committee.

Drug Class	Preferred Agents	Non-Preferred Agents
Ophthalmic	bacitracin	Bleph®-10
Antibiotics, Non-	bacitracin/polymyxin B	Garamycin®
Quinolones	gentamicin solution/ointment	Neocidin®
	polymyxin B/trimethoprim	neomycin/polymyxin B/bacitracin
	sulfacetamide solution	neomycin/polymyxin B/gramicidin
	tobramycin solution	Neosporin®
		Polytrim®
		sulfacetamide ointment
		Tobrex®
Ophthalmic Antifungals	N/A	<mark>Natacyn®</mark>
Ophthalmic	erythromycin 0.5% ointment	AzαSite™
Macrolides		llotycin®
Ophthalmic	ciprofloxacin ophthalmic solution	Besivance™
Quinolones	Moxeza™	Ciloxan®
	ofloxacin	gatifloxacin
	Vigamox™	levofloxacin 0.5%
		moxifloxacin (generic Vigamox™)
		Ocuflox®
		Quixin®
		Zymaxid™

Otic Antibiotics

Class Selection & Guidelines

- DMS to select preferred agent(s) based on economic evaluation; however, at least 3 unique chemical entities or combinations should be preferred.
- Agents not selected as preferred will be considered non-preferred and will require PA.
- For any new chemical entity in the *Otic Antibiotics* class, require PA until reviewed by the P&T Committee.

Drug Class	Preferred Agents	Non-Preferred Agents
Otic Antibiotics	CiproDex® Otic	Cipro HC® Otic
	ciprofloxacin	Coly-mycin® S
	<u>ofloxacin</u>	Floxin™
	hydrocortisone/neomycin sulfate/polymyxin B solution,	Otovel™
	suspension	





Steroids, Topical (Low Potency)

Class Selection & Guidelines

Topical Steroids (Low Potency)

- DMS to select preferred agent(s) based on economic evaluation; however, at least 3 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 *Steroids, Topical (Low Potency)* class, require PA until reviewed by the P&T Committee.





Drug Class	Preferred Agents	Non-Preferred Agents
Topical Steroids	alclometasone dipropionate	Aqua Glycolic®
	betamethasone valerate cream, ointment	Aqua Glycolic HC®
	clobetasol propionate cream, gel, ointment,	amcinonide
	solution	ApexiCon®/ApexiCon E®
	Clobex® shampoo	Balneol®
	Derma-Smoothe/FS®	betamethasone dipropionate
	fluocinonide solution	betamethasone dipropionate augmented
	fluticasone propionate cream, ointment	betamethasone valerate foam, lotion
	halobetasol propionate	Capex® Shampoo
	hydrocortisone cream, gel, lotion, ointment	clobetasol emollient
	mometasone furoate cream, ointment, solution	clobetasol propionate foam, lotion,
	triamcinolone acetonide cream, lotion, ointment	shampoo, spray
		Clobex® lotion, spray
		clocortolone
		Clodan®
		Cloderm®
		Cordran® Tape
		Cutivate®
		DermacinRx® Silapak
		DermacinRx® Silazone PharmPak
		Dermatop®
		Desonate®
		desonide
		desoximetasone
		diflorasone diacetate
		Diprolene®
		Diprolene AF®
		fluocinolone acetonide oil
		fluocinonide emollient
		fluocinonide cream, gel, ointment
		fluocinolone acetonide
		flurandrenolide
		fluticasone propionate lotion
		Halog®
		hydrocortisone-aloe
		hydrocortisone butyrate
		hydrocortisone butyrate/emollient
		hydrocortisone valerate
		hydrocortisone-urea
		Kenalog®
		Locoid®
		Locoid Lipocream®
		Luxiq®
		Micort-HC®
		Olux®/Olux-E®





Pandel®
prednicarbate
Psorcon®
Sernivo™
Silazone-II™
Synalar®

Classes Reviewed by Consent Agenda

No change in PDL status:

- Acne Agents, Oral
- Anti-Ulcer Protectants
- Antibiotics, Topical
- Antidiarrheals
- Antipsoriatics, Oral
- Antipsoriatics, Topical
- Antivirals, Topical
- GI Motility, Chronic
- H. Pylori Treatment
- Histamine II Receptor Blockers
- Immunomodulators, Atopic Dermatitis
- Immunosuppressives, Oral
- Laxatives and Cathartics

- Ophthalmic Immunomodulators
- Ophthalmics, Antibiotic-Steroid Combinations
- Ophthalmics, Anti-inflammatories
- Ophthalmics, Antivirals
- Ophthalmics, Glaucoma Agents
- Ophthalmics, Mydriatic
- Ophthalmics, Vasoconstrictors
- Otic Anti-Infectives and Anesthetics
- Otics, Anti-Inflammatory
- Proton Pump Inhibitors
- Rosacea Agents, Topical
- Steroids, Topical (Medium, High, Very High)
- Ulcerative Colitis Agents