

The following tables list the agenda items as well as the Options for Consideration that are scheduled to be presented and reviewed at the April 15, 2025 meeting of the Pharmacy and Therapeutics Advisory Committee.

SINGLE AGENT REVIEWS

Agent	Options for Consideration presented by MedImpact
New Product to Market	Non-PDL
Alyftrek™ (vanzacaftor,	
tezacaftor, and deutivacaftor)	Approval Duration: 6 months
	• Vanzacaftor and tezacaftor work additively to increase the amount of cystic fibrosis transmembrane conductance regulator (CFTR) protein on the cell surface. Deutivacaftor increases the channel open probability of the CFTR protein at the cell surface. The three agents increase CFTR activity.
	Initial Approval Criteria:
	Patient has a documented diagnosis of cystic fibrosis
	with:
	 A genetic profile (e.g., gene mutation) that is considered responsive to the product based on clinical and/or in vitro data contained in the FDA labeling; AND Confirmed by an FDA-approved diagnostic test; AND Patient meets the FDA-approved minimum age; AND
	 Documentation (e.g., progress notes) of baseline
	functional status and baseline predicted FEV1.
	Renewal Criteria:
	 Patient has had disease response, as indicated by one or more of the following:
	 Decreased pulmonary exacerbations, as compared to pre-treatment baseline; OR Improvement or stabilization of lung function, compared to baseline; OR Decrease in decline of lung function; OR Improvement in quality of life, weight gain, or growth.
	Age Limit: 6 years of age or older
	 Quantity Limit: 50-20-4 mg tablets: 3 per day 125-50-10 mg tablets: 2 per day

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Agent	Options for Consideration presented by MedImpact
New Product to Market Sofdra™ (sofpironium)	Non-PDL
	Approval Duration: 6 months
	 Sofpironium competitively inhibits acetylcholine receptors on some peripheral tissues, including sweat glands. With receptor stimulation prevented, the rate of sweating decreases.
	Initial Approval Criteria:
	 Patient has a diagnosis of primary axillary hyperhidrosis; AND
	 Prescriber attests hyperhidrosis is significantly interfering with activities of daily living; AND Patient meets the FDA-approved minimum age.
	 Renewal Criteria: Prescriber attestation of clinically significant improvement in clinical signs and symptoms.
	Age Limit: 9 years of age or older
	Quantity Limit: 1 bottle per 30 days
New Product to Market Ryzumvi™ (phentolamine	Non-PDL
ophthalmic solution)	Approval Duration: Single fill only
	• Phentolamine is a relatively non-selective alpha-1 and alpha-2 adrenergic antagonist. Muscles involved in dilating the pupil are primarily activated by these receptors to directly reduce pupil diameter, therefore reversing the mydriasis induced by specific pharmacological agents.
	Approval Criteria:
	• Patient has a diagnosis of pharmacologically-induced mydriasis produced by adrenergic agonists (e.g., phenylephrine) OR parasympatholytic (e.g., tropicamide) agents; AND
	 Prescriber attests product will be used within 24 hours of the procedure; AND



Agent	 Options for Consideration presented by MedImpact Prescribed by, or in consultation with, an ophthalmologist or other specialist in the treatment of pharmacologically-induced mydriasis Age Limit: 3 years of age or older Quantity Limit: 1 single-patient-use vial per fill
New Product to Market Crenessity™ (crinecerfont)	 Non-PDL Approval Duration: 6 months initial, 1 year renewal Crinecerfont is a selective corticotropin-releasing factor (CRF) type 1 receptor antagonist. Crinecerfont blocks the binding of CRF to CRF type 1 receptors in the pituitary but not CRF type 2 receptors. Crinecerfont binding to CRF type 1 receptors inhibits adrenocorticotropic hormone (ACTH) secretion from the pituitary, thereby reducing ACTH-mediated adrenal androgen production. Initial Approval Criteria: Patient has a diagnosis of classic congenital adrenal hyperplasia (CAH) defined by ≥ 1 of the following: Elevated 17-hydroxyprogesterone (17-OHP) level; OR Confirmed CYP21A2 genotype; OR Positive newborn screening with confirmatory second-tier testing (e.g., liquid chromatography – tandem mass spectrometry); OR Cosyntropin stimulation test; AND Prescribed initially by, or in consultation with an endocrinologist; AND Crenessity (crinecerfont) will be used as an adjunct therapy with chronic glucocorticoid therapy for CAH (e.g., hydrocortisone, prednisone, methylprednisolone, dexamethasone) at a minimum glucocorticoid dose required for cortisol replacement; AND If prescribed concomitantly with a moderate or strong CYP3A4 inducer (e.g., phenytoin, carbamazepine, rifampin, rifabutin, rifapentine, phenobarbital, bosentan, efavirenz, etravirine, and primidone), dosages will be modified as recommended by the package insert; AND



Agent	Options for Consideration presented by MedImpact
	 Patient meets the minimum age 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, as indicated by ≥ 1 of the following:
	 o Reduction in glucocorticoid daily use; OR o Reduction in serum androstenedione (A4) levels.
	Age Limit: 4 years of age or older
	Quantity Limit:
	 25 mg, 50 mg, and 100mg oral capsules: 2 per day 50 mg/mL oral solution: 4 mL per day
New Product to Market	Non-PDL
Journavx™ (suzetrigine)	Approval Duration: 3 months (Limit to 1 fill per approval)
	• Suzetrigine is a selective blocker of the NaV1.8 voltage- gated sodium channel. NaV1.8 is expressed in peripheral sensory neurons including dorsal root ganglion neurons, where its role is to transmit pain signals. By selectively inhibiting NaV1.8 channels, suzetrigine inhibits transmission of pain signals to the spinal cord and brain.
	Initial Approval Criteria:
	 Patient has a diagnosis of moderate to severe acute pain; AND
	 Journavx (suzetrigine) will be used for up to 14 days; AND
	 Prescriber attests that the member's pain is unable to be managed with an NSAID, acetaminophen, or other non- opioid analgesic; AND
	 Journavx (suzetrigine) is not being prescribed to treat chronic pain; AND
	 Journavx (suzetrigine) is not being prescribed to treat pain associated with migraine; AND
	 Patient does not have severe hepatic impairment (Child- Pugh Class C); AND



Agent	Options for Consideration presented by MedImpact
	 Patient has been counseled to avoid food or drink containing grapefruit during treatment with Journavx (suzetrigine); AND Patient is not concurrently taking a strong CYP3A inhibitor; AND Patient is not concurrently taking a moderate or strong CYP3A inducer; AND Patients using hormonal contraceptives containing progestins other than levonorgestrel and norethindrone have been counseled regarding alternative or additional contraception, if appropriate, per product labeling; AND Patient meets the minimum age recommended by the package insert for the provided indication. Age Limit: 18 years of age or older Quantity Limit: 30 tablets per 14 days
New Product to Market Tryngolza™ (olezasren)	 Non-PDL Approval Duration: 1 year Olezarsen is an APOC-III-directed antisense oligonucleotide (ASO) indicated as an adjunct to diet to reduce triglycerides in adults with familial chylomicronemia syndrome (FCS). Initial Approval Criteria: Diagnosis of familial chylomicronemia syndrome (FCS) confirmed by genetic mutations in one of the following: LPL gene APOA5 gene GPIHBP1 gene LMF1 gene APOC2 gene; AND Patient has a fasting triglyceride level greater than or equal to 880 mg/dL; AND Prescribed by, or in consultation with, an endocrinologist, or other specialist in the treatment of familial chylomicronemia syndrome (FCS); AND Patient meets the minimum age recommended by the package insert for the provided indication.



Agent	Options for Consideration presented by MedImpact
ingent	Renewal Criteria:
	Prescriber attestation of clinically significant
	improvement or stabilization in the patient's condition.
	Age Limit: 18 years of age or older
	Quantity Limit: 1 autoinjector per month
New Product to Market Hympavzi™(marstacimab-hncq)	Non-PDL
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	Hympavzi is a non-factor, monoclonal antibody targeting
	and blocking tissue factor pathway inhibitor (TFPI), an
	anti-clotting protein. It is the first non-factor therapy
	approved for both hemophilia A and B but limited to only
	those without inhibitors.
	Initial Approval Criteria:
	Prescribed for the prophylactic treatment in patients with
	one of the following:
	 Hemophilia A without inhibitors to Factor 8
	(FVIII); OR
	• Hemophilia B without inhibitors to Factor 9 (FIX).
	• Documentation (e.g., an inhibitor lab result within the
	past year) demonstrating the absence of one of the
	following:
	 Factor VIII inhibitors for hemophilia A; OR
	 Factor IX inhibitors for hemophilia B; AND
	Patient meets the minimum age recommended by the
	package insert for the provided indication.
	Renewal Criteria:
	 Prescriber attests patient has experienced clinical benefit compared to baseline
	benefit compared to baseline.
	• Documentation (e.g., an inhibitor lab result within the
	past year) demonstrating the absence of one of the
	following:
	 Factor VIII inhibitors for hemophilia A; OR
	 Factor IX inhibitors for hemophilia B.
	Age Limit: 12 years of age or older
	Quantity Limit: 300 mg (2mL) per week
	Contracting Line, 500 mg (2mL) per week
New Product to Market Alhemo® (concizumab-mtci)	Non-PDL
	Approval Duration: 1 year initial, renewal



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Agent	Options for Consideration presented by MedImpact
	 Alhemo is a non-factor, monoclonal antibody targeting and blocking tissue factor pathway inhibitor (TFPI), an anti-clotting protein. Only TFPI antagonist for use in patients WITH inhibitors.
	Initial Approval Criteria:
	 Prescribed for the prophylactic treatment in patients with one of the following: Hemophilia A with inhibitors to FVIII; OR Hemophilia B with inhibitors to FIX. Documentation (e.g., an inhibitor lab result within the past year) demonstrating one of the following: Factor VIII inhibitor for hemophilia A; OR Factor IX inhibitor for hemophilia B; AND Patient meets the minimum age recommended by the package insert for the provided indication.
	Renewal Criteria:
	 Prescriber attests patient has experienced clinical benefit compared to baseline. Documentation (e.g., an inhibitor lab result within the past year) demonstrating one of the following: Factor VIII inhibitor for hemophilia A; OR Factor IX inhibitor for hemophilia B
	Age Limit: 12 years of age or older

FULL CLASS REVIEWS

PDL Class	Options for Consideration presented by MedImpact
Narcotics, Long-Acting	 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 Narcotics, Long-Acting class, require PA until reviewed by the P&T Committee.
Colony Stimulating Factors	 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 Colony Stimulating Factors class, require PA until reviewed by the P&T Committee.



PDL Class	Options for Consideration presented by MedImpact
Erythropoiesis Stimulating Proteins	 DMS to select preferred agent(s) based on economic evaluation; however, at least 2 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 Erythropoiesis Stimulating Proteins class, require PA until reviewed by the P&T Committee.
Phosphate Binders	 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 Phosphate Binders class, require PA until reviewed by the P&T Committee.
Insulins & Related 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 Insulins & Related Agents class, require PA until reviewed by the P&T Committee.
Sodium Glucose Co-Transporter 2 (SGLT2) Inhibitors	 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 Sodium Glucose Co-Transporter 2 (SGLT2) Inhibitors class, require PA until reviewed by the P&T Committee.
Growth Hormones	 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 Growth Hormones class, require PA until reviewed by the P&T Committee.





CONSENT AGENDA ITEMS

Consent Agenda

Options for Consideration presented by MedImpact

For the following therapeutic classes, there are **no recommended changes to the Preferred Drug** List (PDL) status; these may be voted on as a group

- Narcotic Agonist/Antagonists
 Narcotics, Fentanyl Buccal Products
- Narcotics, Short-Acting
- Non-Steroidal Anti-Inflammatory Drugs (NSAIDs)
- Opiate Dependence Treatments
- Antihyperuricemics
- Sickle Cell Anemia Treatments
- Thrombopoiesis Stimulating
 Proteins
- Alpha-Glucosidase Inhibitors
- Dipeptidyl Peptidase-4 (DPP-4)
 Inhibitors

- Glucagon-Like Peptide-1 (GLP-1) Receptor Agonists
- Meglitinides
- Metformins
- Sulfonylureas
- Thiazolidinediones (TZDs)
- Androgenic Agents
- Bone Resorption Suppression & Related Agents
- Glucagon Agents
- Pancreatic Enzymes
- Progestins for Cachexia
- Steroids, Oral
- Uterine Disorder Treatments