

MINUTES OF THE MEETING OF THE COLORADO MEDICAID P&T COMMITTEE

Department of Health Care Policy and Financing Virtual Meeting via Zoom

April 4, 2023

1. Call to Order

A quorum being present, G. ATHEY officially called the meeting to order at 13:07 MT.

2. Roll Call

Board introductions were made. There were sufficient members for a quorum with ten members participating and no members excused. Members Present

A. Members Present

Morgan Alonzo, PharmD George Athey, MD (Chairperson) Emily Kosirog, PharmD Daralyn Morgenson, PharmD Kelet Robinson, MD (Vice-Chairperson) Marisa Wiktor, MD

B. Members Excused

Gwen Black, PharmaD James Feinstein, MD Thuy McKitrick, PharmD

C. Members Absent

Paulette Campbell



D. Staff Present

HCPF Pharmacy Office

Mohamed Duklef, RPh Jim Leonard, PharmD

Magellan RX Management

Jessica Czechowski, PharmD Erik Hamel, PharmD Jessica Bacon

3. Approval of Minutes

G. ATHEY asked for approval of the minutes from the January 10, 2023, meeting. The minutes were approved with no audible dissent.

4. Department Updates:

M. DUKLEF reviewed updates from the January 10, 2023, P&T meeting.

- Non-Opioid Analgesia Agents Oral & Topical
- Opioids Short-Acting, Fentanyl Preparations, Long-Acting
- Anticonvulsants, Oral
- Newer Generation Antidepressants
- Atypical Antipsychotics Oral/Topical
- Neurocognitive Disorder Agents
- Sedative Hypnotics
- Skeletal Muscle Relaxants
- Stimulants and Related Agents
- Triptans, Ditans, and Other Migraine Treatments Oral & Non-Oral
- Multiple Scleroisis Therapies Disease Modifying & Symptom Management
- Ophthalmics, Immunomodulators
- Mass review drug classes
 - Non-Steroidal Anti-Inflammatory Drugs (NSAIDs) Oral & Non-Oral
 - Monoamine Oxidase Inhibitors (MAOIs)
 - Tricyclic Antidepressants (TCAs)
 - Anti-Parkinson's Agents
 - Benzodiazepines (Non-Sedative Hypnotic)
 - o Anxiolytics, Non-Benzodiazepine
 - Calcitonin Gene-Related Peptide Inhibitors (CGRPIs)
 - Lithium Agents
 - Ophthalmics, Allergy
 - Ophthalmics, Anti-Inflammatories



o Ophthalmics, Glaucoma

5. NEW BUSINESS

- A. M. DUKLEF reviewed updates from the Prior Authorization Call Center.
 - Prior authorization requests for Pharmacy benefits can be faxed or called-in, in most cases. Also, the new prescriber tool, accessible through the EHR, allows for real time benefit check, electronic eprescribing, and electronic 'e-PAs'.
 - 1st Quarter of 2023
 - 73% approvals and 20% denials, 7% change in therapy
 - Average hold time for the call center for the past quarter was 3 minutes and 13 seconds
 - Average call length was 7 minutes
 - 27,698 ePAs were initiated, with 76% approvals. ePA made up 40% of all PAs initiated
- B. M. DUKLEF mentioned there are still three positions open on the committee.

6. Rules

- G. ATHEY presented rules for drug classes that are up for review and will contain public testimony, class updates and market share, and Committee discussion.
- Each review will contain the following:
 - Opportunity for disclosures by Committee members and speakers.
 - Oral presentations by manufacturers, providers and public.
 - Overview for each Drug Class including market share and FDA updates.
 - Committee Discussion and Recommendations for each Class.
- > Mass review Drug classes will only include:
 - Overview for each Drug Class including market share and FDA updates.
- > Rules for presentation:
 - Oral presentations are restricted to products that are being reviewed for PDL status.
 - Presentations will be limited to 3 minutes per representative per drug product.
 - Representatives will be called to present in the order in which they signed in by drug class.
 - Presentations will be limited by verbal comments.
 - No visual aids other than designated handouts are permitted.
 - Presentations should follow the one-page summary that was submitted to the Department.
 - Stakeholders' comments are to:
 - Be limited to clinical information only;



- ♦ Exclude any reference to cost;
- ♦ Exclude anecdotal content;
- Exclude general drug or disease specific economic information.
- > The audience will be considered a reference tool for the Committee.
- > The Committee will discuss topics and audience participation will be allowed if P&T members ask for clarification.
- > The Department disseminated recently received public comments to the Committee members prior to the meeting.
- G. ATHEY presented Committee Discussion and Recommendations for each Class should address the following questions:
- Do the agents differ in efficacy or effectiveness?
- Do the agents differ in safety or adverse effects?
- Are there subgroups for which one agent is associated with either differences in efficacy or effectiveness, or differences in safety or adverse effects?

Factual Inaccuracy:

G. ATHEY presented Factual Inaccuracy. During a Committee meeting, if a stakeholder believes that a factual inaccuracy has been stated by a Committee member, the stakeholder may hand a note or email the Department representative. The stakeholder must provide the factual inaccuracy or a summary of the inaccuracy on the note. The Department representative will forward any comment to the Chair or Vice Chair. The Committee Chair/Vice Chair will then determine if there is need to publicly hear the inaccuracy prior to moving forward with motions and discussion. The Chair/Vice Chair will state the purported factual inaccuracy and will ask the Committee if they want to hear testimony regarding the factual inaccuracy. When providing testimony, the stakeholder must provide evidence to support the claim of inaccuracy and cannot provide opinions on the drug class being considered.

A. DRUG CLASSES FOR REVIEW

- G. ATHEY moved to discuss Drug Classes for Review.
- M. DUKLEF asked for any disclosures for all classes to be reviewed. No disclosures noted. M. DUKLEF asked for all speakers to provide disclosures before speaking.
- 1. G. ATHEY moved to discuss **PAH Therapies (PDEIs, Endothelin Antagonists, Prostanoids, & Guanylate Cyclase Stimulators).** KIMBERLY SIMPSON from UNITED THERAPEUTICS spoke on Tyvaso. ADRIAN LAU from JANSSEN spoke on Uptravi. E. HAMEL reviewed utilization and updates. (1) K. ROBINSON made a motion that at least one from each of the four classes (endothelin antagonists, prostanoids, guanylate cyclase stimulator (sGC) and phosphodiesterase inhibitors) be



- preferred. D. MORGENSON seconded. The motion passed with no audible dissent. (2) M. WIKTOR made a motion that at least one agent with pediatric indication be preferred. E. KOSIROG seconded. The motion passed with no audible dissent. (3) D. MORGENSON made a motion that all available routes of administration across all classes be available as preferred. R. ATHEY seconded. The motion passed with no audible dissent.
- 2. G. ATHEY moved to discuss **Anti-Psoriatics Oral & Topical.** No speakers. E. HAMEL reviewed utilization and updates. (1) E. KOSIROG made a motion that various topical formulations be available as preferred based on application site. K. ROBINSON seconded. The motion passed with no audible dissent.
- 3. G. ATHEY moved to discuss Immunomodulators, Topical (Atopic Dermatitis, Antineoplastics, & Other Agents). No speakers. E. HAMEL reviewed utilization and updates. Antineoplastics (1) M. WIKTOR made a motion that at least one agent be preferred for the treatment of cutaneious T-cell lymphoma (CTCL). E. KOSIROG seconded. The motion passed with no audible dissent. (2) K. ROBINSON made a motion that at least one agent be preferred for actinic keratosis. M. WIKTOR seconded. The motion passed with no audible dissent. *Atopic Dermatitis* (3) G. ATHEY made a motion that at least one agent with a pediatric indication for children two years and above be preferred. E. KOSIROG seconded. The motion passed with no audible dissent. (4) E. KOSIROG made a motion that at least one agent with an indication for mild-to-moderate atopic dermatitis and one agent with an indication for moderate-to-severe dermatitis be preferred. G. ATHEY seconded. The motion passed with no audible dissent. (5) E. KOSIROG made a motion that at least one cream and one ointment be preferred. D. MORGENSON seconded. The motion passed with no audible dissent. Other Agents (6) K. ROBINSON made a motion that at least one type of each formulation (cream, gel, ointment, etc.) be available as preferred. G. ATHEY seconded. The motion passed with no audible dissent. (7) E. KOSIROG made a motion that at least one agent with an FDA indication for the treatment of genital warts be preferred. K. ROBINSON seconded. The motion passed with no audible dissent.
- 4. G. ATHEY moved to discuss **Bile Salts.** No speakers. E. HAMEL reviewed utilization and updates. (1) M. WIKTOR made a motion that at least one agent that can be administered via feeding tube be preferred. E. KOSIROG seconded. The motion passed with no audible dissent. (2) K. ROBINSON made a motion to have one available agent to treat cholestasis of pregnancy. G. ATHEY seconded. The motion passed with no audible dissent.
- 5. G. ATHEY moved to discuss **Anti-Emetics**, **Oral & Non-Oral**. No speakers. E. HAMEL reviewed utilization and updates. (1) G. ATHEY made a motion that at least one agent with pediatric indication from oral and non-oral subcategory be



- preferred. D. MORGENSON seconded. The motion passed with no audible dissent. (2) K. ROBINSON made a motion that alternate dosage forms for all ages be available, such as liquid, ODT, and patch. E. KOSIROG seconded. The motion passed with no audible dissent. (3) M. WIKTOR made a motion that multiple mechanisms of action in multiple dosage forms be preferred. G. ATHEY seconded. The motion passed with no audible dissent. (4) M. WIKTOR made a motion that at least one agent for the prevention of delayed nausea and vomiting be available. K. ROBINSON seconded. The motion passed with no audible dissent. (5) M. WIKTOR made a motion that one agent be preferred for the indication of nausea and vomiting associated with pregnancy. E. KOSIROG seconded. The motion passed with no audible dissent.
- 6. G. ATHEY moved to discuss **GI Motility, Chronic.** No speakers. E. HAMEL reviewed utilization and updates. (1) D. MORGENSON made a motion that at least one agent that has a non-oral route be preferred. G. ATHEY seconded. The motion passed with one opposed. (2) E. KOSIROG made a motion that at least one preferred product be available for each of the indications (IBS-C, IBS-D, CIC, and OIC). D. MORGENSON seconded. The motion passed with no audible dissent.
- 7. G. ATHEY move to discuss Hemorrhoidal, Anorectal, and Related Topical Anesthetic Agents. No speakers. E. HAMEL reviewed utilization and updates. (1) E. KOSIROG made a motion that preferred agents include multiple formulations of administration. K. ROBINSON seconded. The motion passed with no audible dissent. (2) D. MORGENSON made a motion that at least one agent be preferred for anal fissures. E. KOSIROG seconded. The motion passed with no audible dissent. (3) E. KOSIROG made a motion that at least one agent be preferred with an anesthetic. M. WIKTOR seconded. The motion passed with no audible dissent.
- 8. G. ATHEY moved to discuss Anticoagulants (Oral & Parenteral). KANONDA BAYO from BristolMyersSquibb spoke on Eliquis. E. HAMEL reviewed utilization and updates. Oral (1) E. KOSIROG made a motion that at least two DOACs be preferred as a first-line agent. D. MORGENSON seconded. The motion passed with no audible dissent. (2) M. WIKTOR made a motion that at least one agent with a lower risk of bleeding be preferred. G. ATHEY seconded. The motion passed with no audible dissent. (3) K. ROBINSON made a motion that just agents with higher safety profiles be preferred. E. KOSIROG seconded. The motion passed with no audible dissent. Parenteral (4) E. KOSIROG made a motion that at least one low molecular weight heparin be preferred as a first line agent. K. ROBINSON seconded. The motion passed with no audible dissent. (5) E. KOSIROG made a motion that at least one agent for the indication of HIT be preferred. D. MORGENSON seconded. The motion passed with no audible dissent.
- 9. G. ATHEY moved to discuss **Anti-Platelets**. SUNNY HIRPARA from AstraZeneca spoke on Brilinta. E. HAMEL reviewed utilization and updates. (1) E. KOSIROG



- made a motion that multiple agents be available due to varying levels of efficacy and safety. D. MORGENSON seconded. The motion passed with no audible dissent.
- 10. G. ATHEY moved to discuss **Colony Stimulating Factors**. No speakers. E. HAMEL reviewed utilization and updates. (1) K. ROBINSON made a motion that one long-acting and one short-acting CSF agent be preferred. M. ALONZO seconded. The motion passed with no audible dissent.

Break at 14:32 MST and meeting resumed at 14:42 MST.

- 11. G. ATHEY moved to discuss Mass Review Drug Classes and reviewed the rules for Mass Review Drug Classes.
 - Tetracyclines
 - 1. At least one agent with a pediatric indication be preferred.
 - 2. At least two different tetracyclines be preferred on the PDL.
 - 3. At least one agent used to treat chlamydia be preferred.
 - 4. At least on agent that can be administered via feeding tube be preferred.
 - Alpha-Blockers
 - 1. No motions given.
 - Beta-Blockers Single Agent, Anti-Arrhythmics, & Combinations
 - 1. At least one agent be available with the indication for treatment of heart failure.
 - 2. At least one agent that is Beta-1 selective be available.
 - 3. At least one agent with a pediatric indication be preferred.
 - 4. At least one agent be included that is considered acceptable during pregnancy.
 - 5. At least one dosage form be preferred that can be administered through a feeding tube.
 - Calcium Channel Blockers DHPs & Non-DHPs
 - 1. At least one agent be included that is considered acceptable during pregnancy.
 - 2. At least one agent be preferred in each subcategory.
 - 3. At least one agent with a pediatric indication be preferred in each subcategory.
 - Angiotensin Converting Enzyme (ACE) Inhibitors & Combinations
 - 1. At least two ACEIs and two ACEI Combinations be preferred.
 - 2. At least one liquid formulation that can be administered via a feeding tube be preferred.
 - 3. At least one ACEI for pediatric indications be preferred.
 - Angiotensin Receptor Blockers (ARBs) & Combinations
 - 1. At least two ARBs and two ARB combinations be preferred.
 - 2. At least one combination, including a neprilysin inhibitor, be preferred for the treatment of heart failure.
 - 3. At least one ARB for pediatric indications be preferred.



- Renin Inhibitors & Combinations
 - 1. No motions given.
- Lipotropics Bile Acid Sequestrants, Fibrates, & Other Agents
 - 1. Agents be preferred in each drug class that are capable of being administered through a feeding tube, where available.
- Statins & Combinations
 - 1. At least one agent with a pediatric indication be preferred.
 - 2. One product with reduced drug interaction risk be included as preferred.
 - 3. Two high potency statins defined as >50% reduction in LDL should be included.
- Acne Agents, Topical
 - 1. At least one product be available as preferred in each topical category (antibiotics, antibiotic combinations, retinoids, retinoid combinations, other).
 - 2. At least one agent that is recommended for the treatment of inflammatory acne be preferred.
 - 3. At least one product from the antibiotic combination subclass and one product from the retinoid combination subclass be preferred.
- Acne Agents, Oral Isotretinoins
 - 1. No motions given.
- Rosacea Agents
 - 1. Preferred agents should include cream, gel, foam, and lotion.
 - 2. One formulation be available for those individuals who are of child-bearing potential.
- Topical Steroids Low, Medium, High, & Very High Potency
 - 1. At least one agent from each potency category with pediatric indication be preferred.
 - 2. At least two preferred agents be available for each potency category.
 - 3. Consideration be given for multiple formulations to account for application site across potency categories.
- H. Pylori Treatments
 - 1. No motions given.
- Pancreatic Enzymes
 - 1. Two or more agents be preferred due to the variability in patient response.
- Proton Pump Inhibitors
 - 1. At least one agent with a pediatric indication be preferred.
 - 2. Consideration be given to a variety of formulations for people with special needs (such as trouble swallowing and feeding tube.)
- Non-Biologic Ulcerative Colitis Oral & Rectal
 - 1. Product formulation (oral and non-oral, foam, suppositories, enema, capsule, tablet, and a product that can be opened and poured onto applesauce) be considered for preferred status.
 - 2. At least one agent with pediatric indication be preferred.



- 3. At least one preferred agent be available for treatment as well as maintenance therapy.
- Erythropoiesis Stimulating Agents
 - 1. No motions given.
- G. ATHEY made a motion to approve the mass review drug classes. K. ROBINSON seconded. The motion passed with no audible dissent.
- M. DUKLEF announced the next meeting for July 11, 2023.
- G. ATHEY made a motion to adjourn. K. ROBINSON seconded. The motion passed with no audible dissent. The meeting adjourned at 14:53 MST.

By:	George Athey, MD	
Date: _	07/11/2023	

Reasonable accommodations will be provided upon request for persons with disabilities. Please notify the Committee Coordinator at 303-866-6371 or Mohamed.duklef@state.co.us or the 504/ADA Coordinator hcpf504ada@state.co.us at least one week prior to the meeting.

