



Healthcare Services **Medical & Pharmacy Policy Alerts**

Number 84 July 1, 2023 This is the July 1, 2023 issue of the Providence Health Plans, Providence Health Assurance and Providence Plan Partners, Medical and Pharmacy Policy Alert to our providers. The focus of this update is to communicate to providers' new or revised Medical or Pharmacy policy changes. The Health Plan has a standard process to review all Medical & Pharmacy Policies annually. Policies will be available for review on ProvLink and via the PHP website at: https://healthplans.providence.org/providers/provider-support/medical-

policy-pharmacy-policy-and-provider-information/

The Provider Alert, Prior Authorization Requirements, and Medical policies are all available on ProvLink and through the link above.

NOTE: For Oregon Medicaid requests, services which do not require prior authorization will process against the Prioritized List. To determine which services require prior-authorization, please see the current PHP prior authorization list here.

EXTERNAL PROVIDER REVIEW OPPORTUNITY

PHP Medical Policy Committee is seeking feedback from providers to serve as clinical subject matter experts (SMEs) through the policy development and annual review processes. This review process allows providers to offer their expertise and discuss relevant research in their field that will be used to support how these policy decisions are made. This will allow providers an opportunity to offer valuable insight that will help shape policies that affect provider reimbursement and patient care.

If interested, please email us at PHPmedicalpolicyinquiry@providence.org with your name, specialty, and preferred email address.





MEDICAL POLICY COMMITTEE

MEDICAL

COMPANY POLICIES

Effective 8/1/2023

Pellet Implant	Codes/PA: No changes to codes or configuration.				
MP109	OHP: These changes do not apply to OHP. The Prioritized List and the Oregon Administrative Rules will be followed.				
Glycated Hemoglobin and	Policy Updates: No recommended changes to criteria.				
Glycated Protein Testing	Codes/PA: Add dx code Z7985 to pair to pay with CPT codes, based on CMS updates.				
MP267					
	OHP: These changes do not apply to OHP. The Prioritized List and the Oregon Administrative Rules will be followed.				
Genetic Testing:	Policy Updates: Update criteria to allow for testing of women's reproductive partner for cystic fibrosis (CF) if test is positive in woman				
Reproductive Planning and					
Prenatal Testing	Codes/PA: No changes				
MP78	OHP: These changes do not apply to OHP. The Prioritized List and the Oregon Administrative Rules will be followed.				
Genetic Testing: Gene	Policy Updates: Moved code 0155U from Circulating Tumor Cell and DNA Assays for Cancer Management. Will remain PA. Updated				
Expression Profile Testing	cross reference policy section.				
for Breast Cancer	Codes/PA: 0155U added- will remain PA				
MP47	OHP: These changes do not apply to OHP. The Prioritized List and the Oregon Administrative Rules will be followed.				





Circulating Tumor Cell and DNA Assays for Cancer Management	Policy Updates: Remove 0155U as it utilizes tumor tissue. Code will be moved to Genetic Testing: Gene Expression Profile Testing for Breast Cancer policy. Will remain PA. Policy reference listed.
MP122	Reformatting Codes/PA:
	O155U- remove and place on Genetic Testing: Gene Expression Profile Testing for Breast Cancer policy (will remain PA)
	OHP: These changes do not apply to OHP. The Prioritized List and the Oregon Administrative Rules will be followed.

Effective 9/1/2023

Blood Brain Barrier	Policy Updates: Change denial type from "investigational" to "not medically necessary."				
Disruption and Bypass	Codes/PA: No changes; codes will continue to require PA.				
MP147	OHP: These changes do not apply to OHP. The Prioritized List and the Oregon Administrative Rules will be followed.				
Cardiac: Disease Risk	Policy Updates: Change denial type from "investigational" to "not medically necessary" for measurement of biomarkers for the				
Screening	assessment of cardiovascular disease risk, and cardiovascular disease risk panels. Add GlycA to the list of non-covered biomarkers per consideration.				
MP148	Codes/PA: Change denial type from "investigational" to "not medically necessary" for the following codes: 0052U, 0119U, 83876, 0308U, 0309U, 0310U, 0377U, 0389U, 0401U, 0024U.				
	OHP: These changes do not apply to OHP. The Prioritized List and the Oregon Administrative Rules will be followed.				
Cardiac: Transcatheter	Policy Updates:				
Aortic Valve Replacement	Change denials from investigational to not medically necessary				
MP77	Add Portico TAVR System to table of FDA approved systems				
	Move FDA table to Regulatory section of policy				
	Codes/PA: No coding changes				





	OHP: These changes do not apply to OHP. The Prioritized List and the Oregon Administrative Rules will be followe				
Eye: Corneal Collagen	Policy Updates: Change denial language in criterion III to 'not medically necessary'.				
Cross-Linking	Codes/PA: No changes to coding or PA.				
MP158	OHP: These changes do not apply to OHP. The Prioritized List and the Oregon Administrative Rules will be followed.				
Peroral Endoscopic	Policy Updates: Change denial language in criterion III to 'not medically necessary'.				
Myotomy (POEM)	Codes/PA: No changes to coding or PA.				
MP191					
	OHP: These changes do not apply to OHP. The Prioritized List and the Oregon Administrative Rules will be followed.				
Allergy Testing	Policy Updates: Change non coverage position from investigational to not medically necessary. Codes/PA:				
MP153	Non coverage position changed from investigational to not medically necessary (86001, 86343, 95060, 95065, 0165U)				
	Change denial on 86005 from u31 (Deny, Not covered per Medical Policy) to u21 (Deny, not medically necessary per the Medical Policy) Policy)				
	OHP: These changes do not apply to OHP. The Prioritized List and the Oregon Administrative Rules will be followed.				
Blood Counts	Policy Updates: Annual Update				
MP208	Recommendation: No recommended changes to criteria				
WIF 200	Codes/PA: Update coding configuration for all CPT codes, adding a number of dx codes to pair to deny, based on CMS updates				
	OHP: These changes do not apply to OHP. The Prioritized List and the Oregon Administrative Rules will be followed.				
Intraoperative Monitoring	Interim Update: Update billing guidelines to state that CPT codes for EMG (95860, 95861, 95863, 95864) cannot be billed together with 95938 (SSEP) for the same episode of intraoperative monitoring.				
MP295	Codes/PA: Set up EMG (95860-4) and SSEP (95938) codes to deny t07 when billed together.				
	OHP: These changes do not apply to OHP. The Prioritized List and the Oregon Administrative Rules will be followed.				
Next Generation Sequencing for Minimal Residual Disease Detection	Policy Updates: • Reformat criteria; add acute myeloid leukemia (AML) to list of indications appropriate for MRD detection.				





MP110	Change denial type from "investigational" to "not medically necessary" for not medically necessary indications for MRD detection.
	Codes/PA:
	 Change denial type from "investigational" to "not medically necessary" for the following codes: 0306U, 0307U, 0340U.
	 Add 0171U to policy (MyMRD NGS Panel) – this code is specific to MRD detection in AML; currently requires PA per two other policies.
	OHP: These changes do not apply to OHP. The Prioritized List and the Oregon Administrative Rules will be followed.
Electrical Stimulation: Non-	Policy Updates:
Covered Therapies	Unselected Medicare for coverage- Medicare has own policy.
MP331	Update non-coverage position from investigational to not medically necessary.
	 Update language in peripheral nerve stimulation (PNS) to highlight some of the potential indications. Codes/PA: Investigational denial changed to NMN
	OHP: These changes do not apply to OHP. The Prioritized List and the Oregon Administrative Rules will be followed.

MEDICARE

Effective 8/1/23

Surgical Site of Service	New Medicare Advantage medical policy
MP395	Policy Update: New Medicare Advantage medical policy, separating by line of business. No change to criteria, continue to use Company criteria. Codes/PA: No changes to codes or configuration.
Subcutaneous Hormone Pellet Implant (Medicare)	Policy Update: Archive. Service will be managed by Pharmacy. Codes/PA: Remove medical policy configuration. Any related configuration will be managed by Pharmacy.
MP336	





Diabetes: Blood Glucose Monitor and Supplies	Policy Update: No change to criteria. Codes/PA: Code and configuration changes:		
MP276	 Change configuration on codes that currently bundle into K0553. HCPCS code K0553 (and K0554) were termed 1/1/2023, so configuration connected to these codes needs to be updated with the replacement HCPCS codes and the above-the-login (ATL) non-covered list (NCL) needs further details added to explain when the code may be determined "not covered." No changes to any codes which are unaffected by this bundling edit update. 		

Effective 9/1/23

Allergy Testing MP152	Policy Updates: No change to criteria. The Company policy criteria changing from INV to NMN changes some of the generic language found in the Medicare version. Codes/PA: CPT 86005, remove u31 (Deny, Not covered per Medical Policy) and add u21 (NMN denial). No configuration changes to any other code in the policy.
Cardiac: Disease Risk Screening MP132	Policy Update: Add the GlycA test to the criteria table, as well as an additional LCD, but no change to existing criteria. Codes/PA: Add 0024U to the policy with NMN edit. No changes to other codes currently in the policy.
Intraoperative Monitoring	Policy Update: No change to criteria. Update billing guidelines to state that CPT codes for EMG (95860, 95861, 95863, 95864) cannot be billed together with 95938 (SSEP) for the same episode of intraoperative monitoring.
MP296	Codes/PA: Set up EMG (95860, 95861, 95863, and 95864) codes to deny t07 (no separate reimbursement per MP) when billed with and SSEP (95938). No changes to other codes in this policy.





REIMBURSEMENT POLICIES

Effective 7/1/2023

Reimbursement Methodologies and All- Inclusive Rates RP4	New Reimbursement policy Recommendation: Converting Coding Policy 02.0 to a Reimbursement Policy since the policy is primarily a reimbursement-related topic. This particular policy is specific to all-inclusive rates. Will now be under the reimbursement policy team for continued policy management.
Transfers Between	Annual Review
Hospitals	Recommendation: No recommended changes to the reimbursement methodology for transfers. Continue to follow Medicare rules for commercial and Medicare members and OARs for OHP members.
RP75	Reimbursement Methodology: No change to 2022 presentation of policy guidelines for the transferring hospitals reimbursement rate.
	• <u>Commercial and Medicare</u> : transferring facility reimbursement graduated per diem rate (if there is no specific transfer payment language in the facility contract)
	OHP: transferring facility reimbursement based on OARs per diem interhospital transfer payment rate
	All LOBs: full payment rate is made to the final discharging hospital
	Relevant References/CMS Guidance:
	 42 CFR § 412.4 - discharges and transfers. Legal Information Institute. https://www.law.cornell.edu/cfr/text/42/412.4.
	 Medicare Claims Processing Manual; Chapter 3 - Inpatient Hospital Billing. https://www.cms.gov/Regulations-and-guidance/Guidance/Manuals/Downloads/clm104c03.pdf. Section 20.1.2.4 Transfers
	 Review of hospital compliance with Medicare's transfer policy. https://www.cms.gov/files/document/se21001.pdf.
	OAR 410-125-0165 - Transfers and Reimbursement - Oregon Administrative Rules. https://oregon.public.law/rules/oar-410-125-0165 .





VENDOR UPDATES

Medicare Echo Scans and Nuclear Medicine

Effective 8/1/2023, for Medicare only, the below services no longer require prior authorization through Carelon:

Nuclear Cardiology	СРТ
Myocardial Perfusion Imaging	78451
	78452
	78453
	78454
Infarct Imaging	78466
	78468
	78469
Cardiac Blood Pool Imaging	78472
	78473
	78481
	78483
	78494

Cardiac Services	СРТ
Stress Echo (SE)	93350
	93351
Resting Trans Echo (TTE)	93303
	93304
	93306
	93307
	93308
Transesophageal Echo (TEE)	93312
	93313
	93314
	93315
	93316
	93317





Here's what's new from the following policy committees:

Pharmacy & Therapeutics (P&T) Committee

Oregon Region P&T Committee Meeting June 2, 2023 Go-Live Date: Wednesday, August 02, 2023, unless otherwise noted

Table of Contents:

- New Drugs or Combinations
- New Strengths or Formulations
- New Indications Monitoring
- Drug Safety Monitoring
- Other Formulary Changes
- New Generic Medications
- Clinical Policy Changes

New Drugs or Combinations:

- 1. Tremelimumab-actl (Imjudo) Vial
 - a. Indication:
 - 1. In combination with durvalumab, for the treatment of adult patients with unresectable hepatocellular carcinoma (uHCC)
 - 2. In combination with durvalumab and platinum-based chemotherapy for the treatment of adult patients with metastatic non-small-cell lung cancer (NSCLC) with no sensitizing EGFR mutations or ALK genomic tumor aberrations
 - b. **Decision**:

Commercial	Medicaid	Medicare





Formulary Status*	Medical	Medical	Part D: Non-formulary Part B: Medical
Tier**	N/A	N/A	N/A
Affordable Care Act Eligible	N/A; Non-Formulary	N/A	N/A
Utilization Management Edits	Prior Authorization	Prior Authorization	Prior Authorization
Quantity Limit	N/A	N/A	N/A

^{*} Recommendations for placement may differ between lines of business due to regulatory requirements.

Formulary Alternatives:

HCC: atezolizumab (Tecentriq®) + bevacizumab (Zirabev®, Mvasi®)

NSCLC: First line- pembrolizumab (Keytruda®), alternative therapies: nivolumab (Opdivo®)/ipilimumab (Yervoy®), cemiplimab-rwlc (Libtayo®)

c. Prior Authorization Criteria for Commercial/Medicaid/Medicare Part B: Added to Injectable Anti-Cancer Medications Policy

2. Elacestrant hydrochloride (Orserdu) Tablet

a. **Indication**: For the treatment of postmenopausal women or adult men, with ER-positive, HER2-negative, ESR1-mutated advanced or metastatic breast cancer with disease progression following at least one line of endocrine therapy.

b. **Decision**:

	Commercial	Medicaid	Medicare
Formulary Status*	Farmulan	Formulary	Part D: Formulary
Formulary Status	Formulary		Part B: N/A
Tier**	Tier 6 - Non-Preferred Specialty	N/A	Specialty
Affordable Care Act Eligible	No	N/A	N/A
Utilization Management Edits	Prior Authorization	Prior Authorization	Prior Authorization
Quantity Limit			
* Pacamendations for placement may differ between lines of business due to regulatory requirements			

^{*} Recommendations for placement may differ between lines of business due to regulatory requirements.

^{**} Medications will be placed on recommended tier above for the base formulary; tier placement for custom formulary(ies) will be based on designation above. For example, Commercial Tier 6 designation above means that the medication will be placed on the highest cost-sharing tier on the respective formulary(ies).





** Medications will be placed on recommended tier above for the base formulary; tier placement for custom formulary(ies) will be based on designation above. For example, Commercial Tier 6 designation above means that the medication will be placed on the highest cost-sharing tier on the respective formulary(ies).

Formulary Alternatives: N/A

- c. Prior Authorization Criteria for Commercial/Medicaid: Added to Oral Anti-Cancer Medications Policy
- d. Prior Authorization Criteria for Medicare Part D: Added to Anti-Cancer Agents Program
- 3. Velmanase alfa-tycv (Lamzede) Vial
 - a. **Indication**: For the treatment of non-CNS manifestations of alpha-mannosidosis in adults and pediatric patients.
 - b. **Decision**:

	Commercial	Medicaid	Medicare
Formulary Status*	Medical	I Medical I	Part D: Non-formulary
Torritalary Status	Wedical		Part B: Medical
Tier**	N/A	N/A	Choose an item.
Affordable Care Act Eligible	N/A; Non-Formulary	N/A	N/A
Utilization Management Edits	Prior Authorization	Prior Authorization	Prior Authorization
Quantity Limit	N/A	N/A	N/A

^{*} Recommendations for placement may differ between lines of business due to regulatory requirements.

Formulary Alternatives: None

c. Prior Authorization Criteria for Commercial/Medicaid/Medicare Part B: Added to Enzyme Replacement Therapy Policy

PA PROGRAM NAME	Enzyme Replacement Therapy	
MEDICATION NAME	Velmanase alfa-tycv vial (Lamzede [®])	
REQUIRED MEDICAL INFORMATION	For initial authorization all the following must be met:	

^{**} Medications will be placed on recommended tier above for the base formulary; tier placement for custom formulary(ies) will be based on designation above. For example, Commercial Tier 6 designation above means that the medication will be placed on the highest cost-sharing tier on the respective formulary(ies).





	4. December of FDA labeled to the first of the control of the control of
	Documentation of FDA-labeled indication for the requested product
	AND
	2. Dosing is within FDA-labeled guidelines
	3. For velmanase alpha only, the following additional criteria apply:
	a. Confirmed diagnosis of alpha-mannosidosis as defined by alpha-mannosidase activity
	less than 10% of normal activity in blood leukocytes
	b. Documented baseline serum oligosaccharide level
	c. Documented baseline value of either 6-minute walk test, 3-minute stair climb or
	forced vital capacity. Note: This may be waved for children under the age of three.
	Improvement or stabilization is required for reauthorization.
	d. Therapy is being used to treat non-central nervous system manifestations of alpha
	mannosidosis such as skeletal abnormalities, myopathy, motor function disturbances, immune deficiency
	e. No prior history of bone marrow transplant
	Note: If request is for a non-FDA approved dose, medical rationale must be submitted in support of
	therapy with a higher dose for the intended diagnosis (such as high-quality peer reviewed literature,
	accepted compendia or evidence-based practice guidelines) and exceptions will be considered on a
	case-by-case basis.
	case-by-case basis.
	REAUTHORIZATION:
	Both of the following must be met:
	1. Documentation of successful response to therapy (e.g., disease stability or improvement in
	symptoms).
	2. Dosing is within FDA-labeled guidelines
	3. For velmanase alpha only,
	a. For initial reauthorization: a decrease of serum oligosaccharides of 3 micromoles per
	liter or at least 30%
	b. For subsequent reauthorizations: stabilization or improvement in either the 6-minute
	walk test, 3-minute stair climb or forced vital capacity
	Note: If request is for a non-FDA approved dose, medical rationale must be submitted in support of
	therapy with a higher dose for the intended diagnosis (such as high-quality peer reviewed literature,
	accepted compendia or evidence-based practice guidelines) and exceptions will be considered on a
	case-by-case basis.
COVERAGE DURATION	Initial and reauthorization will be approved for one year.





4. Omaveloxolone (Skyclarys) Capsule

a. **Indication**: For treatment of Friedreich's ataxia.

b. **Decision**:

	Commercial	Medicaid	Medicare
Formulary Status*	Non-formulary	Non-formulary	Part D: Non-formulary Part B: N/A
Tier**	N/A	N/A	N/A
Affordable Care Act Eligible	N/A; Non-Formulary	N/A	N/A
Utilization Management Edits	Prior Authorization	Prior Authorization	N/A
Quantity Limit			

^{*} Recommendations for placement may differ between lines of business due to regulatory requirements.

Formulary Alternatives: None

c. Prior Authorization Criteria for Commercial/Medicaid: Add to Medications For Rare Indications Policy

5. Trofinetide (Daybue) Solution

a. Indication: For the treatment of Rett syndrome (RTT) in adults and pediatric patients 2 years of age and older.

b. **Decision**:

	Commercial	Medicaid	Medicare
Formulary Status*	Non-formulary	Non-formulary	Part D: Non-formulary Part B: N/A
Tier**	N/A	N/A	N/A
Affordable Care Act Eligible	No	N/A	N/A

^{**} Medications will be placed on recommended tier above for the base formulary; tier placement for custom formulary(ies) will be based on designation above. For example, Commercial Tier 6 designation above means that the medication will be placed on the highest cost-sharing tier on the respective formulary(ies).





Utilization Management Edits	Prior Authorization	Prior Authorization	N/A
Quantity Limit	N/A	N/A	N/A

^{*} Recommendations for placement may differ between lines of business due to regulatory requirements.

Formulary Alternatives: None

c. Prior Authorization Criteria for Commercial/Medicaid

PA PROGRAM NAME	Medications for Rare Indications
MEDICATION NAME	trofinetide oral solution (Daybue®)
COVERAGE DURATION	For Nulibry®: Initial authorization will be approved for three months. Reauthorization will be approved for 12 months. For Daybue®: Initial authorization will be approved for 6 months. Reauthorization will be approved for 12 months.
	For all other medications: Initial authorization will be approved for one year and reauthorization will be approved until no longer eligible with the plan, subject to formulary or benefit changes.

New Drug Strengths or Formulations:

- 1. Pegcetacoplan-PF (Syfovre) Vial
 - a. Indication: For the treatment of geographic atrophy (GA) secondary to age-related macular degeneration (AMD).
 - b. **Decision**:

	Commercial	Medicaid	Medicare
Formulary Status*	Madical	Medical	Part D:
	Medical		Part B: Medical
Tier**	N/A	N/A	N/A
Affordable Care Act Eligible	No	N/A	N/A
Utilization Management Edits	Prior Authorization	Prior Authorization	Prior Authorization

^{**} Medications will be placed on recommended tier above for the base formulary; tier placement for custom formulary(ies) will be based on designation above. For example, Commercial Tier 6 designation above means that the medication will be placed on the highest cost-sharing tier on the respective formulary(ies).





Quantity Limit	N/A	N/A	N/A
* Recommendations for placement	may differ between lines of business	due to regulatory requirements.	
** Medications will be placed on re	** Medications will be placed on recommended tier above for the base formulary; tier placement for custom formulary(ies) will be based on		
designation above. For example, Commercial Tier 6 designation above means that the medication will be placed on the highest cost-sharing		ed on the highest cost-sharing tier	
on the respective formulary(ies).			
Formulary Alternatives: N/A			

c. Prior Authorization Criteria for Commercial/Medicaid/Medicare Part B:

PA PROGRAM NAME	Syfovre		
MEDICATION NAME	Pegcetacoplan-pf vial		
PA INDICATION INDICATOR	1 - All FDA-Approved Indications		
OFF-LABEL USES	N/A		
EXCLUSION CRITERIA	 History of or active choroidal neovascularization (CNV), associated with AMD or any other cause History of ocular or periocular infections History of endophthalmitis, retinal detachments, or increased intraocular pressure 		
	For initial authorization, all of the following criteria must be met:		
REQUIRED MEDICAL	 Documentation of diagnosis of geographic atrophy (GA) confirmed by clinical exam or diagnostic imaging (such as Color Fundus Photography, Fundus Autofluorescence, Near Infrared Reflectance Imaging, Optical Coherence Tomography) Documentation that GA is secondary to age-related macular degeneration (AMD) 		
INFORMATION	For reauthorization: Documentation of response to therapy defined as one of the following: 1. Reduction in GA growth lesion		
	Documentation of improvement in visual function through visual function assessment test (such as normal luminance best-correct visual acuity [BCVA], maximum reading speed, Functional Reading Independence Index, microperimetry)		
AGE RESTRICTIONS	Age equal to or greater than 60 years of age.		
PRESCRIBER RESTRICTIONS	Must be prescribed by, or in consultation with, an ophthalmologist.		
COVERAGE DURATION	Initial authorization and reauthorization will be approved for one year.		





New Indications Monitoring:

The following information is gathered from the United States Food and Drug Administration (FDA) Approved Drug Products database from 2/1/2023–3/31/2023

Therapies with Prior Authorization Policies (Non-oncology)

- 1. Synjardy (empagliflozin and metformin hydrochloride)
 - a. Previous Indication(s):
 - a. Adjunct to diet and exercise to improve glycemic control in adults with type 2 diabetes mellitus.
 - b. Reduce the risk of cardiovascular death in adults with type 2 diabetes mellitus and established cardiovascular disease.
 - b. New indication approved 02/06/2023:
 - a. In adults with type 2 diabetes mellitus to reduce the risk of cardiovascular death and hospitalization for heart failure in adults with heart failure.
 - c. **RECOMMENDATION:** Inform prescribers via Medical Policy Alert. Update Medicaid policy with indication.
- 2. Kevzara (sarilumab)
 - a. Previous Indication(s):
 - a. Treatment of adult patients with moderately to severely active rheumatoid arthritis who have had an inadequate response or intolerance to one or more disease-modifying antirheumatic drugs (DMARDs).
 - b. New indication approved 02/28/2023:
 - a. Adult patients with polymyalgia rheumatica (PMR) who have had an inadequate response to corticosteroids or who cannot tolerate corticosteroid taper.
 - c. **RECOMMENDATION:** Inform prescribers via Medical Policy Alert. Update policy with new indication and add new criteria.

Prior Authorization for Commercial/Medicaid:

(Note: PMR is on line 106 on the current OHP Prioritized List and is a coverable diagnosis without disease severity requirements)

PA PROGRAM NAME	THERAPEUTIC IMMUNOMODULATORS
	(TIMs)
MEDICATION NAME	Kevzara (sarilumab)
COVERED USES	1 - All FDA-Approved Indications
EXCLUSION CRITERIA	Combination therapy with another therapeutic immunomodulator (TIM) agent
REQUIRED MEDICAL	For polymyalgia rheumatica (PMR), sarilumab (Kevzara®) may be covered if the following criteria are met:
INFORMATION	1. Diagnosis of PMR and documentation of the following:





	 Age 50 years or older at disease onset AND One of the following: Bilateral shoulder or pelvic aching or stiffness lasting longer than 45 minutes and persisting for at least two weeks OR If younger than 50 years of age and having asymmetric shoulder or pelvic pain, documentation of PMR with atypical features Documentation that similar disorders have been ruled out (such as giant cell arteritis rheumatoid arthritis, drug-induced maylgias, fibromyalgia, other muscoloskeletal disease, or other bone disease). One of the following: Indadequate response to full dose systemic systemic corticosteroid Documented PMR flare while attempting to taper systemic corticosteroid Intolerance or contraindication to systemic corticosteroids
AGE RESTRICTIONS	Age must be appropriate based on FDA-approved indication
PRESCRIBER RESTRICTIONS	Rheumatoid arthritis, ankylosing spondylitis, non-radiographic axial spondyloarthritis, juvenile idiopathic arthritis, polymyalgia rheumatica: must be prescribed by, or in consultation with, a rheumatologist
COVERAGE DURATION	 Prior Authorization: Authorization will be approved until no longer eligible with the plan, subject to formulary or benefit changes Quantity Limitation: Initial authorization will be approved for six months, and reauthorization will be approved for one year. FDA-labeled dosing will be approved until no longer eligible with the plan, subject to formulary and/or benefit changes

3. Eylea (Aflibercept)

- a. Previous Indication(s):
 - a. treatment of patients with:
 - 1. Neovascular (Wet) Age-Related Macular Degeneration (AMD)
 - 2. Macular Edema Following Retinal Vein Occlusion (RVO)
 - 3. Diabetic Macular Edema (DME)
 - 4. Diabetic Retinopathy (DR)
- b. New indication approved 02/08/2023:
 - a. Retinopathy of Prematurity (ROP)
- c. **RECOMMENDATION:** Inform prescribers via Medical Policy Alert. This drug is ophthalmic vascular endothelial growth factor (VEGF) inhibitors policy was updated as part of annual review and is included in the policy review section of the consent agenda vote.





- 4. Evkeeza (evinacumab-dgnb)
 - a. Previous Indication(s):
 - i. Adjunct to other low-density lipoprotein-cholesterol (LDL-C) lowering therapies for the treatment of adult and pediatric patients, aged 12 years and older, with homozygous familial hypercholesterolemia (HoFH) (DR)
 - b. New indication approved 03/21/2023:
 - a. Adjunct to other low-density lipoprotein-cholesterol (LDL-C) lowering therapies for the treatment of adult and pediatric patients, aged 5 years and older, with homozygous familial hypercholesterolemia (HoFH)
 - c. **RECOMMENDATION:** Inform prescribers via Medical Policy Alert. Update policy with new indication.

Therapies with Prior Authorization Policies (Oncology)

- 5. **Verzenio** (abemaciclib)
 - a. New indication(s) approved 03/03/2023:
 - a. In combination with endocrine therapy (tamoxifen or an aromatase inhibitor) for the adjuvant treatment of adult patients with hormone receptor (HR)-positive, human epidermal growth factor receptor 2 (HER2)-negative, node-positive, early breast cancer at high risk of recurrence. (Requirement of a Ki-67 score ≥20% as determined by an FDA approved test is no longer a part of the indication)
 - b. in combination with an aromatase inhibitor as initial endocrine based therapy for the treatment of adult patients with hormone receptor (HR)-positive, human epidermal growth factor receptor 2 (HER2)-negative advanced or metastatic breast cancer. (Indication previously was limited to men and postmenopausal women)
 - b. **RECOMMENDATION:** Inform prescribers via Medical Policy Alert. Prior authorization policy coverage criteria are based on recommendations from the National Comprehensive Cancer Network (NCCN); no updates to the policy are warranted.
- 6. Tafinlar (dabrafenib)
 - a. New indication(s) approved 03/16/2023:
 - a. Treatment of pediatric patients 1 year of age and older with low-grade glioma (LGG) with a BRAF V600E mutation who require systemic therapy.
 - b. **RECOMMENDATION:** Inform prescribers via Medical Policy Alert. Prior authorization policy coverage criteria are based on recommendations from the National Comprehensive Cancer Network (NCCN); no updates to the policy are warranted.
- 7. **Mekinist (**tramitinib)
 - a. New indication(s) approved 03/16/2023:





- a. Treatment of pediatric patients 1 year of age and older with low-grade glioma (LGG) with a BRAF V600E mutation who require systemic therapy.
- b. **RECOMMENDATION:** Inform prescribers via Medical Policy Alert. Prior authorization policy coverage criteria are based on recommendations from the National Comprehensive Cancer Network (NCCN); no updates to the policy are warranted.
- 8. Lutrate Depot (leuprolide acetate)
 - a. Previous Indication(s):
 - a. Palliative treatment of advanced prostate cancer
 - b. New indication approved 02/06/2023:
 - a. Treatment of advanced prostate cancer.
 - c. **RECOMMENDATION:** Inform prescribers via Medical Policy Alert. Prior authorization policy coverage criteria are based on recommendations from the National Comprehensive Cancer Network (NCCN); no updates to the policy are warranted. Update policy with indication.
- 9. Keytruda (pembrolizumab)
 - a. Full indication approved 03/28/2023:
 - a. For the treatment of adult and pediatric patients with unresectable or metastatic microsatellite instability-high (MSI-H) or mismatch repair deficient (dMMR) solid tumors, as determined by an FDA-approved test, that have progressed following prior treatment and who have no satisfactory alternative treatment options
 - b. **RECOMMENDATION:** Inform prescribers via Medical Policy Alert. Prior authorization policy coverage criteria are based on recommendations from the National Comprehensive Cancer Network (NCCN); no updates to the policy are warranted.
- 10. Jemperli (pembrolizumab)
 - a. Full indication approved 02/09/2023:
 - a. Treatment of adult patients with mismatch repair deficient (dMMR) recurrent or advanced endometrial cancer, as determined by an FDA-approved test, that has progressed on or following prior treatment with a platinum-containing regimen in any setting and are not candidates for curative surgery or PD-1/PD-L1-blocking antibody radiation
 - b. **RECOMMENDATION:** Inform prescribers via Medical Policy Alert. Prior authorization policy coverage criteria are based on recommendations from the National Comprehensive Cancer Network (NCCN); no updates to the policy are warranted.
- 11. **Obdivo** (nivolumab)
 - a. Full indication approved 02/15/2023:
 - a. adult and pediatric (12 years and older) patients with unresectable or metastatic melanoma, as a single agent or in combination with ipilimumab.





- b. adult and pediatric (12 years and older) patients with melanoma with lymph node involvement or metastatic disease who have undergone complete resection, in the adjuvant setting
- b. **RECOMMENDATION:** Inform prescribers via Medical Policy Alert. Prior authorization policy coverage criteria are based on recommendations from the National Comprehensive Cancer Network (NCCN); no updates to the policy are warranted.

12. Yervoy (ipilimumab)

- a. Full indication approved 02/15/2023:
 - a. Adult and pediatric (12 years and older) patients with unresectable or metastatic melanoma, as a single agent or in combination with nivolumab.
- b. **RECOMMENDATION:** Inform prescribers via Medical Policy Alert. Prior authorization policy coverage criteria are based on recommendations from the National Comprehensive Cancer Network (NCCN); no updates to the policy are warranted.

Therapies Without Prior Authorization Policies

- 13. Illuccix (kit for the preparation of gallium Ga 68 gozetotide injection)
 - a. Previous Indication(s):
 - a. LLUCCIX, after radiolabeling with Ga 68, is a radioactive diagnostic agent indicated for positron emission tomography (PET) of prostate-specific membrane antigen (PSMA) positive lesions in men with prostate cancer:
 - b. with suspected metastasis who are candidates for initial definitive therapy.
 - c. with suspected recurrence based on elevated serum prostate specific antigen (PSA) level indication
 - b. New indication(s) approved 03/15/2023:
 - a. For selection of patients with metastatic prostate cancer, for whom
 - b. lutetium Lu 177 vipivotide tetraxetan PSMA-directed therapy is indicated
 - c. **RECOMMENDATION:** Inform prescribers via Medical Policy Alert.

Drug Safety Monitoring:

The following information is gathered from the United States Food and Drug Administration (FDA) database from 2/1/2023 – 3/31/2023

FDA Drug Safety Communications

There were no drug safety communications reported during this period.

Drug Recalls/Market Withdrawals





- 1. **Drug Name:** TIROSINT®-SOL (levothyroxine sodium)
 - Date of Recall: 02/01/2023
 - Reason for recall: Possible sub potency in 27 lots
 - Link to more information: https://www.fda.gov/safety/recalls-market-withdrawals-safety-alerts/ibsa-pharma-inc-issues-voluntary-nationwide-recall-select-lots-tirosintr-sol-levothyroxine-sodium
 - Health Plan Recommendation: Notify providers via Medical Policy Alert.
- 2. Drug Name: Artificial Tears Lubricant Eye Drops from EzriCare & Delsam Pharma
 - Date of Recall: 02/02/2023
 - Reason for recall: Potential microbial contamination in all lots
 - Link to more information: https://www.fda.gov/safety/recalls-market-withdrawals-safety-alerts/global-pharma-healthcare-issues-voluntary-nationwide-recall-artificial-tears-lubricant-eye-drops-due
 - Health Plan Recommendation: Notify providers via Medical Policy Alert.
- 3. Drug Name: PrimeZEN Black 6000 male enhancement capsules
 - Date of Recall: 02/13/2023
 - Reason for recall: Product contains undeclared tadalafil and sildenafil
 - Link to more information: https://www.fda.gov/safety/recalls-market-withdrawals-safety-alerts/volt-candy-issues-voluntary-nationwide-recall-primezen-black-6000-capsules-due-presence-sildenafil
 - Health Plan Recommendation: Notify providers via Medical Policy Alert.
- 4. Drug Name: Artificial Eye Ointment from Delsam Pharma
 - Date of Recall: 02/24/2023
 - Reason for recall: Potential microbial contamination of batch
 - Link to more information: https://www.fda.gov/safety/recalls-market-withdrawals-safety-alerts/global-pharma-healthcare-issues-voluntary-nationwide-recall-delsam-pharma-artificial-eye-ointment
 - Health Plan Recommendation: Notify providers via Medical Policy Alert.
- 5. Drug Name: Brimonidine Tartrate Ophthalmic Solution, 0.15%
 - Date of Recall: 03/02/2023





- Reason for recall: Potential lack of sterility due to cracked lids in six lots
- Link to more information: https://www.fda.gov/safety/recalls-market-withdrawals-safety-alerts/apotex-corp-issues-voluntary-nationwide-recall-brimonidine-tartrate-ophthalmic-solution-015-due
- Health Plan Recommendation: Notify providers via Medical Policy Alert.

6. Drug Name: 15% MSM Drops

• Date of Recall: 03/03/2023

• Reason for recall: Non-sterility in two lots

• Link to more information: https://www.fda.gov/safety/recalls-market-withdrawals-safety-alerts/pharmedica-usa-llc-issues-voluntary-worldwide-recall-purely-soothing-15-msm-drops-due-non-sterility

• Health Plan Recommendation: Notify providers via Medical Policy Alert.

7. Drug Name: Dabigatran Etexilate Capsules, USP

• Date of Recall: 03/22/2023

• Reason for recall: Detection of N-nitroso-dabigatran (NDAB) impurity in multiple lots

• Link to more information: https://www.fda.gov/safety/recalls-market-withdrawals-safety-alerts/ascend-laboratories-llc-issues-voluntary-nationwide-recall-dabigatran-etexilate-capsules-usp-75-mg

• Health Plan Recommendation: Notify providers via Medical Policy Alert.

8. Drug Name: Atovaquone Oral Suspension

• Date of Recall: 03/31/2023

• Reason for recall: Potential Bacillus cereus contamination of one lot

• Link to more information: https://www.fda.gov/safety/recalls-market-withdrawals-safety-alerts/camber-pharmaceuticals-inc-issues-voluntary-nationwide-recall-atovaquone-oral-suspension-usp

• Health Plan Recommendation: Notify providers via Medical Policy Alert.

Other Formulary Changes:

Drug Name	Recommendation	Policy Name
 Codeine phosphate/guaifenesin 	Remove from Commercial formulary	N/A
(Guaifenesin AC) 10-100mg/5 Liquid	Effective 11/01/2023	





• Codeine Phosphate/Guaifenesin (Guaifenesin-Codeine) 10-100mg/5; 20-		
200/10 Liquid		
Naloxone hcl (Kloxxado) Spray	Commercial: Move to Tier 4	N/A
	Medicaid: Remove from formulary	
	Effective 09/01/2023	
Omeprazole/sodium bicarbonate	New dosage form (susp recon) and strength (2-	Commercial/Medicaid: New Medications
(Konvomep) Susp Recon	84mg/ml);	and Formulations without Established
	Commercial/Medicaid: Non-Formulary,	Benefit
	Prior Authorization	Medicare Part D: N/A
	Medicare Part D: Non-Formulary	
Mifepristone 200 mg Tablet	Commercial/Medicaid: Non-formulary, Prior	Mifepristone
	Authorization	
Naloxone hcl (Narcan) Spray	Commercial: Move brand to Tier 4	N/A
	• Medicare Part D: Down tier generic to Tier 2	
Dabigatran etexilate mesylate (Pradaxa)	New Dosage Form (Pellet Pack) and Strength;	N/A
Pellet Pack	Commercial/Medicaid: Non-Formulary,	
	Quantity Limit (2 packs per day)	
	Medicare Part D: Non-Formulary	
Ropinirole ER 2 mg Tab ER 24H	Commercial: Add Quantity Limit (1 tab per	N/A
	day)	
	Effective 09/01/2023	
Ropinirole ER Tab ER 24H (4 mg, 6 mg)	Commercial Standard: Add to Formulary,	N/A
	Tier 2, Quantity Limit (1 tab per day)	
	Commercial Dynamic: Add to Formulary,	
	Tier 3, Quantity Limit (1 tab per day)	
	Medicaid: Add to Formulary, Quantity Limit	
	(1 tab per day)	
	Effective 09/01/2023	
Ropinirole ER Tab ER 24H (8 mg, 12 mg)	Commercial Standard: Add to Formulary,	N/A
	Tier 2, Quantity Limit (2 tabs per day)	
	Commercial Dynamic: Add to Formulary,	
	Tier 3, Quantity Limit (2 tabs per day)	





	Medicaid: Add to Formulary, Quantity Limit (2 tabs per day) Effective 09/01/2023	
Pseudoephed/codeine/guaifen (Virtussin Dac) Syrup	Remove from Commercial formulary	N/A
Naloxone hcl (Zimh) Syringe	 Commercial: Move brand to Tier 4 Medicaid: Remove from Formulary Effective 09/01/2023 	N/A
Dapsone (Aczone) Gel (Gram)/Gel w/Pump	Commercial: Add to Formulary, Tier 4	N/A
Ipratropium bromide (Atrovent) Spray	Remove from Medicaid formulary	Intranasal Allergy Medications – Medicaid
Epinephrine (Auvi-Q) Auto Injct	Remove from Commercial and Medicaid formularies Effective 09/01/2023	N/A
Roflumilast (Daliresp) Tablet	 Commercial Dynamic formulary: Down tier generic to Tier 3, add Quantity Limit (1 tablet per day) Medicaid: Add Quantity Limit (1 tablet per day) Effective 09/01/2023 	N/A
Erythromycin/benzoyl peroxide 3%/5% Gel	Add to Medicaid formulary	Acne Medications – Medicaid
Somatropin (Genotropin) Cartridge/Disp Syrin	Add to Commercial Formulary, Tier 5	Human Growth Hormones for Adults
 Ambrisentan (Letairis) Tablet Bosentan (Tracleer) Tablet 	Down-tier generic for Commercial: Standard formulary: Tier 2Dynamic formulary: Tier 3	Pulmonary Hypertension
Somatropin (Nutropin AQ Nuspin) Cartridge	Remove from Medicaid Formulary	Human Growth Hormones for Adults
Macitentan (Opsumit) Tablet	 Commercial: Move to Tier 6 from Tier 5 Medicaid: Remove from formulary Effective 09/01/2023 	Pulmonary Hypertension
Becaplermin (Regranex) Gel (Gram)	Commercial/Medicaid: Add Quantity Limit (15 grams per 6 months) Effective 09/01/2023	Regranex





Netarsudil mesylate (Rhopressa) Drops	Commercial/Medicaid: Add Quantity Limit (2.5 ml per 25 days) Effective 09/01/2023	Anti-Glaucoma Agents Step Therapy Policy
Tretinoin 0.025 % Cream	Add to Medicaid formulary	Acne Medications – Medicaid
Oxymetazoline hcl/pf (Upneeq) Droperette	 Commercial/Medicaid: Add Quantity Limit (2 dropperettes per day) Effective 09/01/2023 	Upneeq
Sinecatechins (Veregen) Ointment	Remove from Commercial and Medicaid formularies	N/A
Tirzepatide (Mounjaro®)	Add to Commercial formulary, Tier 3, Step Therapy, Quantity Limit (2 mL per 28 days)	GLP-1/GIP Receptor Agonists
Desvenlafaxine ER tablets	Add Priori Authorization for Commercial	New Medications and Formulations without Established Benefit
Vibegron (Gemtesa)	Add to Medicare Part D formulary, Tier 3 Effective 07/01/2023	N/A

The formulary status for the following drugs was line extended in accordance with Providence Health Plan Pharmacy Operational Policy ORPTCOPS062

Drugs released between March 4, 2023 and April 1, 2023

NEW DRUGS / COMBINATIONS / STRENGTHS / DOSAGE FORMS		
Drug Name	Action Taken	Policy Name
Sodium	New strength (10-3.5/175). Line extend with Clenpiq 10-	N/A
picosulfate/magnesium	3.5/160;	
oxide/citric acid (Clenpiq)	Commercial: Formulary, Tier 4	
Solution	Medicaid: Non- Formulary	
	Medicare Part D: Non- Formulary	
Pegfilgrastim-pbbk (Fylnetra)	Biosimilar to Neulasta. Line extend to Neulasta;	N/A
Syringe	Commercial: Formulary, Tier 5 (Covered Medical Benefit)	
	Medicaid: Formulary, Specialty (Covered Medical Benefit)	
	Medicare Part D: Formulary, Tier 5	





	Medicare Part B: Covered Medical Benefit	
Sotorasib (Lumakras) Tablet	 New Strength (320mg). Line extend with Lumakras 120mg; Commercial: Formulary, Tier 6, Prior Authorization Medicaid: Formulary, Specialty, Prior Authorization Medicare Part D: Formulary, Tier 5, Prior Authorization 	Oral Anti-Cancer Medications
Onasemnogene abeparvovec- xioi (Zolgensma) Kit	 New dose kit. Line extend with other Zolgensma; Commercial/Medicaid/Medicare Part B: Medical Benefit, Prior Authorization Medicare Part D: Non- Formulary 	Zolgensma
Methoxy polyethylene glycol- epoetin beta (Mircera) Syringe	New strength (120mcg/0.3ml). Line extend with existing Mircera strengths; Commercial/Medicaid: Medical Benefit, Prior Authorization Medicare Part D: Non-Formulary Medicare Part B: Medical Benefit, Prior Authorization	 Commercial/Medicaid: Erythropoiesis Stimulating Agents Medicare Part B: Erythropoiesis Stimulating Agents (ESAs) - Medicare Part B
Gabapentin (Gralise) Tab ER 24H	New strength (450mg, 750mg, 900mg). Line extend with Gralise (300mg, 600mg); Commercial/Medicaid: Non-Formulary, Prior Authorization Medicare Part D: Non-Formulary	 Commercial/Medicaid: New Medications and Formulations without Established Benefit Medicare Part D: N/A
Leuprolide acetate (Lupron Depot-Ped) SyingeKit	New kit (45mg). Line extend with Lupron Depot-PED (7.5mg, 11.25mg, 15mg); Commercial/Medicaid: Medical Benefit, Prior Authorization Medicare Part D: Formulary, Tier 5, Prior Authorization Medicare Part B: Medical Benefit, Prior Authorization	 Commercial/Medicaid: Gonadotropin Releasing Hormone Agonists Medicare Part D: Lupron Depot Program Medicare Part B: Gonadotropin Releasing Hormone Agonists – Medicare Part B
Elexacaftor/tezacaftor/ivacaft or (Trikafta) Gran PK SQ	 New dosage form (GRAN PK SQ). Line extend with Trikafta tablets; Commercial: Formulary, Tier 5, Prior Authorization, Quantity Limit (2 packets per day) Medicaid: Formulary, Prior Authorization, Quantity Limit (2 packets per day) Medicare Part D: Formulary, Tier 5, Prior Authorization, Quantity Limit (2 packets per day) 	CFTR Modulators





New Generics:

Drug Name	Action Taken	Policy Name
Bismuth-Metronidazole	First generic (Pylera). Line extend as generic;	N/A
Tetracyc Capsule	Commercial Standard: Formulary, Tier 2, Quantity Limit	
	(120 capsules per 28 days)	
	• Commercial Dynamic: Formulary, Tier 4, Quantity Limit (120	
	capsules per 28 days)	
	Medicaid: Non-Formulary, Quantity Limit (120 capsules per	
	28 days)	
	Medicare Part D: Formulary, Tier 4	
Diltiazem hcl (Diltiazem 24HR	First generic (Cardizem LA). Line extend as generic;	N/A
ER (LA)) Tab ER 24H	Commercial Standard: Formulary, Tier 2	
	Commercial Dynamic: Formulary, Tier 4	
	Medicaid: Formulary	
	Medicare Part D: Formulary, Tier 4	
Teriflunomide Tablet	First generic (Aubagio). Line extend as generic;	N/A
	• Commercial: Formulary, Tier 5, Quantity Limit (1 tablet per	
	day)	
	Medicaid: Formulary, Specialty, Quantity Limit (1 tablet per	
	day)	
	Medicare Part D: Formulary, Tier 3	
Vancomycin HCL Soln Recon	NDA authorized generic (Firvanq). Line extend as generic;	N/A
	Commercial Standard: Formulary, Tier 2	
	Commercial Dynamic: Formulary, Tier 4	
	Medicaid: Non-Formulary	
	Formulary, Tier 4	
Baclofen Oral Susp	First generic drug (Fleqsuvy). Line extend as generic;	N/A
	Non-formulary for all lines of business	
Budesonide Foam/Appl	First generic drug (Uceris). Line extend as generic;	Commercial/Medicaid: Uceris
	• Commercial/Medicaid: Non-formulary, Prior Authorization;	Medicare Part D: N/A
	Medicare Part D: Non-formulary	
Naftifine hcl Gel (Gram)	First generic drug (Naftin). Line extend as generic;	N/A
	 Non-formulary for all lines of business 	





Clinical Policy Changes:

MAJOR CHANGES		
Policy Name	Summary of Change	
Acne Medications – Medicaid	Removed prior authorization on preferred products for children under age of 21 years.	
Adbry	 Updated Medicaid criteria to align with updated criteria from the Oregon Health Authority Prioritized List of Healthcare services for inflammatory skin conditions. Removed Commercial requirement for chronic condition of at least one year, clarified quantity limit for initial authorization, and aligned exclusion criteria with primary therapeutic immunomodulator (TIM) policy to exclude combination use with other TIM agents. 	
Anti-Glaucoma Agents Step	Addition of Rhopressa® to policy with quantity limit.	
Therapy Policy		
Benlysta	Age restriction updated to allow coverage in patients five and older for both Systemic lupus erythematosus (SLE) and active lupus nephritis due to FDA label change.	
Bepreve, Zerviate	Updated policy criteria language regarding trial and failure of brand azelastine 0.05% ophthalmic solution to require trial and failure of generic azelastine as brand has been discontinued.	
Camzyos	Updated prerequisite therapy to require trial and failure of two preferred therapies, instead of three.	
CFTR Modulators	Updates age restrictions and added quantity limit for new Trikafta® packet formulation.	
Cibinqo	 Updated Medicaid criteria to align with updated criteria from the Oregon Health Authority Prioritized List of Healthcare services for inflammatory skin conditions. Updated age restriction to align with FDA label. Removed Commercial requirement for chronic condition of at least one year, aligned exclusion criteria with primary therapeutic immunomodulator (TIM) policy to exclude combination use with other TIM agents 	
Compounded Drugs	Updated to require trial of all formulary options to mirror formulary exception policy	
Corlanor	Rearranged guideline-directed therapy criteria.	
Dupixent	 Updated Medicaid criteria to align with updated criteria from the Oregon Health Authority Prioritized List of Healthcare services for inflammatory skin conditions. Updated age restriction to align with FDA label. For Atopic Dermatitis in Commercial: Removed requirement for severe symptoms. For nasal polyps: Removed requirement for previous nasal surgery, lack of candidacy for nasal surgery, or trial of oral systemic corticosteroids. For eosinophilic esophagitis: Added weight requirement of 40 kg per FDA-approved indication. 	





	• For prurigo nodularis: removed requirement for trial and failure of standard anti-itch therapy, removed quantity of lesions required, changed requirement for prurigo nodularis for at least three months to itching for at least six weeks.	
	 Aligned exclusion criteria with primary therapeutic immunomodulator (TIM) policy to exclude combination use with other TIM agents 	
Elidel, Protopic – Medicaid	Updated Medicaid criteria to align with updated criteria from the Oregon Health Authority Prioritized List of	
	Healthcare services for inflammatory skin conditions. Medications will not require prior authorization for children	
	less than 21 years of age.	
Enstilar, Taclonex, Taclonex	Updated Medicaid criteria to align with updated criteria from the Oregon Health Authority Prioritized List of	
Scalp, Wynzora	Healthcare services for inflammatory skin conditions.	
Eucrisa	Updated Medicaid criteria to align with updated criteria from the Oregon Health Authority Prioritized List of	
	Healthcare services for inflammatory skin conditions.	
	• Updated Commercial criteria to only require tacrolimus in patients at least two years of age as allowed by FDA-	
	approved indication	
Fertility and Related	New Policy- Policy was split from Commercial policy due to differences in authorized uses of these medications by	
Medications Prior Authorization	the Centers of Medicare and Medicaid Services (CMS).	
and Step Therapy Policy -		
Medicare Part B		
GIP/GLP-1 Receptor Agonists	Tirzepatide (Mounjaro®) added as preferred therapy. Policy criteria was updated to clarify medications will only be	
Step Therapy Policy	covered with step through metformin and/or diagnosis of type 2 diabetes.	
GIP/GLP-1 Receptor Agonists	Policy criteria was updated to clarify medications will only be covered with step through metformin and/or	
Step Therapy Policy – Medicaid	diagnosis of type 2 diabetes.	
Homozygous Familial	Added additional genetic mutation for diagnosis [LDL receptor adapter protein 1 (LDLRAP1)] and updated	
Hypercholesterolemia (FH)	definition of clinical diagnosis of homozygous familial hypercholesterolemia to align with American Heart	
Agents	Association and European Atherosclerosis Society definitions.	
Homozygous FH Agents -		
Medicare Part B		
Human Growth Hormones for	Combined Pediatric and Adult policy for ease of review	
Adults	Provider restrictions updated to align with respective disease states	
Human Growth Hormones -	New Policy - Separated from Commercial policy due to significant differences in coverage criteria. Updated to align	
Medicaid	with Oregon Health Authority criteria	
IL-5 Inhibitors	For EGPA: Updated diagnostic criteria to align with the American College of Rheumatology and Lanham guidelines.	
	For HES: Updated trial and failure criteria to only apply to commercial line of business to align with Oregon Health	





• II-5 Inhibitors – Medicare	Authority guidance. For nasal polyps: Removed surgery criteria to align with the International Consensus	
Part B	Statement on Rhinology and Allergy. For all indications, updated exclusion criteria to specify which drug classes are	
	not allowed to be used in combination with requested agent. Updated duration of approval.	
Immune Gamma Globulin (IGG)	Adding criteria for Myelin Oligodendrocyte Glycoprotein Antibody Disease	
Immune Gamma Globulin (IGG)	New policy – separated from Commercial policy due to differences in authorized uses of these medications by the	
Prior Authorization and Step	Centers of Medicare and Medicaid Services (CMS).	
Therapy Policy - Medicare Part		
В		
Intranasal Allergy Medications –	Updated criteria to align with Oregon Health Authority criteria.	
Medicaid		
Lidocaine Patch	Added ICD-10 codes that are considered coverable and will set up to pay automatically at point-of-sale if diagnosis	
	code submitted.	
Luxturna	Updated policy coverage duration from 12 weeks to 6 months to align with coverage duration from Oregon Health	
	Authority.	
Mifepristone	New policy – clarify coverage for non-elective termination of pregnancy uses	
• Ophthalmic Vascular Retired prior authorization for Cimerli® (biosimilar for Lucentis), as this is now considered a preferred product.		
Endothelial Growth Factor	Added criteria for non-preferred products to require	
(VEGF) Inhibitors		
Opzelura	Updated Medicaid criteria to align with updated criteria from the Oregon Health Authority Prioritized List of	
	Healthcare services for inflammatory skin conditions.	
Palforzia	Clarified wording of criteria to ensure use is reserved for patients with history of severe-type reaction to peanut	
	containing products.	
PCSK9 Inhibitors – Commercial	1. Updated statin intolerance criteria to include trial of at least two different statins instead of just either	
	rosuvastatin or atorvastatin, 2. Removed "clinically significant multi vessel coronary hear disease" from clinical	
	ASCVD option as it does not align with the American College of Cardiology/American Heart Association guideline	
	definitions of clinical ASCVD, 3. added coverage for primary hypercholesterolemia (LDL-c greater than 190 g/dL)	
	when secondary causes have been ruled out to align with American College of Cardiology/American Heart	
	Association guidelines.	
PCSK9 Inhibitors - Medicare	1. Updated statin intolerance criteria to include trial of at least two different statins instead of just either	
Part B	rosuvastatin or atorvastatin, 2. Removed "clinically significant multi vessel coronary hear disease" from clinical	
	ASCVD option as it does not align with the American College of Cardiology/American Heart Association guideline	
	definitions of clinical ASCVD.	





PCSK9 Inhibitors- Medicaid	1. simplified criteria for familial hypercholesterolemia to state a "possible" diagnosis of FH via Simon Broome criteria or a "probable" diagnosis of FH via Dutch Lipid Clinic Network Criteria, 2. Increased initial authorization to 12 months.
Pulmonary Arterial	Change policy name to Pulmonary Hypertension, requiring trial and failure or medical rationale for use of brand
Hypertension	Tracleer, Opsumit, or Letairis over generic ambrisentan or bosentan.
Regranex	Updated Medicaid criteria to align with Oregon Health Authority criteria. Commercial criteria updated to require only diagnosis and concurrent wound care.
Rituximab	Updated coverage duration to allow long-term authorization for oncologic diagnoses (aligning with oncology
• Rituximab - Medicare Part B	policies).
Second and Third generation	Updated duration of approval.
antihistamines – Medicaid	
• Soliris	Adding step through Ultomiris® for Atypical Hemolytic Uremic Syndrome and Paroxysmal Nocturnal
 Soliris - Medicare Part B 	Hemoglobinuria, removing Uplizna as a trial option for Neuromyelitis Optica Spectrum Disorder.
Tafamidis	Updated radionuclide imaging criteria to include additional acceptable radiotracers.
Testosterone Replacement	Added language around coverage of pellet insertion procedural codes and for coverage of hormone replacement
Therapy (TRT)	for females. Renamed policy to "Hormone Replacement Therapy"
Therapeutic	Added medical necessity criteria for the use of baricitinib (Olumiant®) in the treatment of alopecia areata
Immunomodulators (TIMS)	
TIMS – Medicaid	Updated Medicaid criteria to align with updated criteria from the Oregon Health Authority Prioritized List of
	Healthcare services for inflammatory skin conditions. Added medical necessity criteria for the use of baricitinib (Olumiant®) in the treatment of alopecia areata
Upneeq	Update quantity limit in clinical policy criteria to align with package insert dosing.
Vascepa	Statin intolerance criteria updated to align with other policies for hypercholesterolemia. Modified age cutoff for ASCVD risk factors to match trial evidence.
Vtama, Zoryve	Updated Medicaid criteria to align with updated criteria from the Oregon Health Authority Prioritized List of Healthcare services for inflammatory skin conditions.
Xolair	For urticaria: Updated Medicaid criteria to align with Oregon Health Authority
	For nasal polyps: Removed surgery criteria to align with the International Consensus Statement on Rhinology
	and Allergy.
	For all indications, updated exclusion criteria to specify which drug classes are not allowed to be used in
	combination with requested agent.
	Updated duration of approval.
Xolair – Medicare Part B	For urticaria: Added definition of chronic disease.





	 For nasal polyps: Removed surgery criteria to align with the International Consensus Statement on Rhinology and Allergy. For all indications, updated exclusion criteria to specify which drug classes are not allowed to be used in combination with requested agent.
Zinplava	Removed exclusion criteria to align with label, added dosing criteria and updated risk factor criteria to only apply to
	Commercial and Medicare Part B to align with the Oregon Health Authority.
Zyflo CR	Due to low utilization, simplified policy to step therapy, requiring a trial of both formulary leukotriene modifiers
	(montelukast and zafirlukast).

RETIRED POLICIES		
Policy Name	Summary of Change	
Aczone Step Therapy Policy	Due to low utilization.	
Daliresp	Due to low utilization and to align with market due to generic availability.	
Human Growth Hormones for	Combined with Human Growth Hormones for Adults policy.	
Pediatrics		
Pneumococcal Vaccines / Pneumococcal Vaccines - Medicare Part B	Due to recommendations continuing to change and the Plan does not want to limit access to Pneumococcal vaccines.	
Topical Androgen Receptor Inhibitors	Will manage utilization with criteria as outlined in the "Formulary and Quantity Exceptions" and "Medical Necessity" policies.	
Veregen	Due to low utilization.	