



# Pulmonary Hypertension (PH) Agents (Oral/Inhalation)

## Medical policy no. 40.12.00-2

**Effective Date: August 1, 2018** 

**Note:** New-to-market drugs included in this class based on the Apple Health Preferred Drug List are non-preferred and subject to this prior authorization (PA) criteria. Non-preferred agents in this class require an inadequate response or documented intolerance due to severe adverse reaction or contraindication to at least TWO preferred agents. If there is only one preferred agent in the class documentation of inadequate response to ONE preferred agent is needed. If a drug within this policy receives a new indication approved by the Food and Drug Administration (FDA), medical necessity for the new indication will be determined on a case-by-case basis following FDA labeling.

To see the list of the current Apple Health Preferred Drug List (AHPDL), please visit: <a href="https://www.hca.wa.gov/assets/billers-and-providers/apple-health-preferred-drug-list.xlsx">https://www.hca.wa.gov/assets/billers-and-providers/apple-health-preferred-drug-list.xlsx</a>

#### **Background:**

Pulmonary hypertension (PH) is a rare, progressive disorder characterized by high blood pressure (hypertension) in the arteries of the lungs (pulmonary artery). The pulmonary arteries are the blood vessels that carry deoxygenated blood from the right side of the heart to the lungs. The World Health Organization (WHO) classifies pulmonary hypertension into five groups based upon etiology. WHO Group I is classified as pulmonary arterial hypertension (PAH), while the other four groups are referred to as pulmonary hypertension (PH)

PH may develop from many different conditions, but the most common type is idiopathic PAH. Common symptoms of PAH include shortness of breath (dyspnea), chest pain, and fainting.

#### **Medical necessity**

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Drug	Medical Necessity
Endothelin Receptor Antagonists	Medications listed in this table may be considered medically
ambrisentan (LETAIRIS®)	necessary when used for the treatment of:
<ul> <li>bosentan (TRACLEER®)</li> </ul>	Pulmanary hyportonsian (DH)
macitentan (OPSUMIT®)	<ul> <li>Pulmonary hypertension (PH)</li> <li>Chronic thromboembolic pulmonary hypertension</li> </ul>
Phosphodiesterase Inhibitors (PDEI)	(CTEPH)
<ul> <li>sildenafil citrate tablets (REVATIO®)</li> </ul>	(612111)
• tadalafil (ADCIRCA®, ALYQ™)	
Prostacyclin Pathway Agonists	
• Iloprost (VENTAVIS®)	
• selexipag ( <b>UPTRAVI</b> ®)	
• treprostinil (ORENITRAM®/TYVASO®)	
Soluble Guanylate Cyclase (SGC)	
Stimulator	
• riociguat (ADEMPAS®)	



#### **Clinical policy:**

#### **Clinical Criteria**

#### **Pulmonary Hypertension (PH)**

# Chronic Thromboembolic Pulmonary Hypertension (CTEPH)

Medications requested for the treatment of PAH may be authorized when **ALL** of the following are met:

- 1. Patient must have **ONE** (either a or b) of the following diagnoses and criteria:
  - a. PH diagnosis WHO Groups III or IV (CTEPH), in which general treatment measures (e.g., anticoagulation) have failed, and PH is thought to be unrelated to underlying lung disease; **OR**
  - b. PAH diagnosis WHO Group I
    - Documentation of PAH WHO Functional class (II, III, or IV);
       AND
    - ii. History of failure, contraindication, or intolerance to amlodipine, diltiazem, or long-acting nifedipine EXCEPT for the following circumstances:
      - (1) Patient had a negative response to acute vasoreactivity testing (AVT); **OR**
      - (2) AVT is not indicated for the patient (PAH due to connective tissue disease, congenital heart disease, HIV, portal hypertension, schistosomiasis, pulmonary veno-occulusive/pulmonary capillary hypertension); **OR**
      - (3) AVT is contraindicated (SBP < 90 mmHg; cardiac index < 2 L/min/m², or PH functional class IV); AND
- 2. Requested therapy is not for ANY of the following:
  - a. A combination of a phosphodiesterase inhibitor and soluble guanylate cyclase stimulator; **OR**
  - b. A combination of selexipag and parenteral prostanoid; OR
- 3. Patient is currently established on requested therapy; **OR**
- 4. <u>For Selexipag</u>: history of failure, contraindication or intolerance to an endothelin receptor antagonist; **AND**
- Prescribed by or in consultation with a specialist in cardiology or pulmonology

If all of the above criteria are met, the request will be **approved for 12** months

If all criteria are not met, but there are documented medically necessary circumstances based on the professional judgement of the clinical reviewer, requests may be approved on a case-by-case basis up to the initial authorization duration.

#### Criteria (Reauthorization)



Medications used for the treatment of PH may be reauthorized when documentation of response (e.g. disease stability or mild progression indicated by a slowing of decline using WHO Functional Class scale) is provided. If all of the above criteria are met, the request will be <b>approved</b> for 12 months
If all criteria are not met, but there are circumstances supported by clinical judgement and documentation, requests may be approved by a clinical reviewer on a case-by-case basis up to the reauthorization duration

### **Coding:**

HCPCS Code	Description
J7686	Treprostinil, inhalation solution, FDA-approved final product, non-compounded, administered through DME, unit dose form, 1.74 mg
Q4074	Iloprost, inhalation solution, FDA-approved final product, non- compounded, administered through DME, unit dose form, up to 20 mcg

## **Dosage and Quantity Limits:**

Drug Name	Dose and Quantity Limits				
Ambristentan	10 mg per day;				
(Letairis)	#60 tablets per 30 day supply (5 mg strength ) <b>OR</b> #30 tablets for 30 day				
	supply (10 mg strength)				
Bosentan	250 mg per day; #60 tablets per 30 day supply				
(Tracleer)					
Macitentan	10 mg per day; #30 tablets per 30 day supply				
(Opsumit)					
Sildenafil citrate	60 mg per day; #90 tablets per 30 day supply				
(Revatio)					
Tadalafil	40 mg per day; #60 tablets per 30 day supply				
(Adcirca)					
Selexipag	3200 mcg per day; can increase to the highest tolerated dose in 200 mcg				
(Uptravi)	twice daily increments at weekly intervals				
lloprost	45 mcg per day;				
(Ventavis)					
Treprostinil	216 mcg per day;				
(Tyvaso)					
Treprostinil diolamine	0.5 mg per day; titrate by 0.25 mg or 0.5 mg twice daily <b>OR</b> 0.125 mg 3				
(Orenitram)	times daily not more than every 3-4 days to the highest tolerated dose				
Riociguat	7.5 mg per day;				
(Adempas)					

## Appendix:

Table 1. WHO Clinical Classification of Pulmonary Hypertension (PH)

WHO Clinical Classification	Description			
Group 1	Pulmonary Arterial Hypertension (PAH)			
	- Idiopathic			
	- Heritable			
	- Drug/toxin induced			



	<ul> <li>Associated with connective tissue disease, HIV infection, portal hypertension, congenital heart disease</li> </ul>
Group 2	PH due to left heart disease
Group 3	PH due to chronic lung disease or hypoxemia
Group 4	Chronic thromboembolic pulmonary hypertension (CTEPH)
Group 5	PH due to unclear multifactorial mechanisms

Table 2. WHO Functional Classification of Patients with PH

WHO Functional Classification	Description		
Class I	Patients with PH without resulting limitation of physical		
	activity. Ordinary physical activity does not cause undue		
	dyspnea or fatigue, chest pain, or near syncope.		
Class II	Patients with PH resulting in slight limitation of physical		
	activity. They are comfortable at rest. Ordinary physical		
	activity causes undue dyspnea or fatigue, chest pain, or		
	near syncope		
Class III	Patients with PH resulting in marked limitation of		
	physical activity. They are comfortable at rest. Less than		
	ordinary activity causes undue dyspnea or fatigue, chest		
	pain, or near syncope.		
Class IV	Patients with PH with inability to carry out any physical		
	activity without symptoms. These patients manifest		
	signs of right-sided heart failure. Dyspnea and/or		
	fatigue may even be present at rest. Discomfort is		
	increased by any physical activity		

#### References

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#### History

Date	Action and Summary of Changes				
10/29/2020	Annual policy update  - Updated preferred/non-preferred status  - Updated PAH clinical criteria  - Updated CTEPH clinical criteria				
10/02/2019	Edit Note				
11/07/2018	HCPCS update				
07/31/2018	Update				
08/16/2017	New Policy				





## **Pulmonary Arterial Hypertension (PAH) Agents**

Please provide the information below, please print your answer, attach supporting documentation, sign, date, and return to our office as soon as possible to expedite this request. Without this information, we may deny the request in seven (7) working days.

Date of request:	Reference #:		MAS:	MAS:		
Patient	Date of birth		ProviderOn	ProviderOne ID		
Pharmacy name	Pharmacy NPI Telepho		ohone number	Fax number		
Prescriber	Prescriber NPI	Tele	ohone number	Fax number	Fax number	
Medication and strength			Directions for use	9	Qty/Days supply	
<ol> <li>Is this request for a continuation of existing therapy?  Yes  No         If yes, is there documentation supporting disease stability Yes  No</li> <li>Indicate the diagnosis:  Pulmonary arterial hypertension (PAH) World Health Organization (WHO) Group I and</li> </ol>						
<ul> <li>WHO Functional class II symptoms</li> <li>WHO Functional class III symptoms</li> <li>WHO Functional class IV symptoms</li> <li>Persistent/recurrent chronic thromboembolic pulmonary hypertension (CTEPH) (WHO group 3 or 4)</li> <li>Other. Specify</li> </ul>						
<ul> <li>3. Has the patient tried a calcium channel blocker?</li></ul>						
<ul> <li>Will the requested therapy be used in combination with any of the following (check all that apply)?</li> <li>Combination of phosphodiesterase inhibitor and soluble guanylate cyclase stimulator</li> <li>Combination of selexipag and parenteral prostanoid</li> <li>None of the above</li> </ul>						
5. For Selexipag: History of failure, contraindication or intolerance to an endothelin receptor antagonist						
6. Is this prescribed by or in consultation with a specialist in one of the following:  Cardiology Pulmonology Other. Specify						
CHART NOTES ARE REQUIRED WITH THIS REQUEST						
Prescriber signature Prescriber specialty			Date			