

Prior Authorization Criteria Pulmonary Arterial Hypertension (PAH) Injectable Agents

All requests for injectable Pulmonary Arterial Hypertension (PAH) agents require a prior authorization and will be screened for medical necessity and appropriateness using the criteria listed below.

For all requests for injectable Pulmonary Arterial Hypertension (PAH) agents all of the following criteria must be met:

- Treatment is prescribed by, or in consultation with, a cardiologist or pulmonologist
- The requested dose and frequency is in accordance with FDA-approved labeling, nationally recognized compendia, and/or evidence-based practice guidelines. If a requested dose is above these recommendations, medical rationale must be submitted. For infused products, must provide member's weight, dose, frequency and titration schedule.
- Request meets diagnostic and drug criteria outlined in sections A and B
- If member is new to the plan and requests a continuation of therapy, must provide chart documentation indicating member is currently on requested therapy

A. Diagnostic Criteria

Coverage may be provided with a <u>diagnosis</u> of Pulmonary Arterial Hypertension (PAH) WHO Group I and the following criteria is met:

- Member has a diagnosis of PAH WHO Group I (refer below to Appendix I) confirmed by chart documentation of right-heart catheterization (RHC) or echocardiography if the provider indicates RHC is not recommended. RHC documentation must contain the following hemodynamic values:
 - o Mean pulmonary arterial pressure > 20 mmHg
 - o Pulmonary capillary wedge pressure ≤ 15 mmHg
 - o Pulmonary vascular resistance > 2 Wood units.
- Documentation of member's vasoreactivity test and one of the following, unless member has a contraindication to vasoreactivity testing (e.g. low systemic blood pressure, low cardiac index, or the presence of severe (functional class IV) symptoms):
 - Member had a positive response (pulmonary artery pressure decreases at least 10 mmHg and to a value less than or equal to 40 mmHg, with an increased or unchanged cardiac output, and a minimally reduced or unchanged systemic blood pressure) and had inadequate response, contraindication or intolerance to calcium channel blocker therapy with diltiazem or a dihydropyridine
 - o Member did not have a positive response to the vasoreactivity test
- Member has functional class II, III or IV symptoms at baseline prior to initiating therapy with the requested drug(s) (refer below to Appendix II)
- One of the following:



- o The requested drug will be used as monotherapy (except Winrevair (sotatercept-csrk) see drug specific criteria below.
- The requested drug will be used for add-on therapy to existing monotherapy or dual therapy in addition to **both** of the following:
 - The medications must be from different therapeutic classes
 - The member must have unresponsive or progressive disease despite established PAH-specific therapies

B. Drug Criteria

- Prostanoids/prostacyclin therapies
 - o Infused agents: epoprostenol, Flolan, Veletri, Remodulin
 - For Flolan, Veletri, or Remodulin requests in members with functional class III or IV symptoms, must provide documentation of inadequate response, contraindication or intolerance to generic epoprostenol or clinical rationale for why generic epoprostenol cannot be used
 - Must meet **one** of the following criteria:
 - Documentation of WHO functional class IV symptoms or functional class III symptoms with **any** of the following:
 - o Evidence of progression of their disease
 - o Any marker of poor clinical prognosis defined as:
 - Clinical signs of right heart failure
 - Repeated episodes of syncope, even with little or regular physical activity
 - <165 meter 6-minute walking distance (6MWD)</p>
 - Peak oxygen consumption (VO₂) <11ml/min/kg (<35% predicted)
 - Ventilatory equivalents of CO₂ (VE/VCO₂) slope >45
 - BNP >300 ng/1
 - NT-proBNP >1400ng/l
 - Right atrium area $\geq 26 \text{ cm}^2$
 - Presence of pericardial effusion
 - Right atrial pressure >14 mmHg
 - Cardiac index <2.0 l/min/m²
 - Mixed venous oxygen saturation (SvO₂) <60%
 - Documentation of WHO functional class II symptoms and all of the following:
 - Request is for Remodulin
 - Inadequate response, contraindication or intolerance to combination therapy of generic ambrisentan and tadalafil
 - Inadequate response, contraindication or intolerance to one additional PAH medication therapy (could have been in combination or monotherapy)

• Activin Signaling Inhibitor:

• Winrevair (sotatercept-crsk) for new starts only: the medication will be taken in combination with other PAH therapies and **ONE** of the following:



 Member is currently receiving at least 2 other PAH therapies from different pharmacologic categories

■ The member is currently receiving at least one other PAH therapy and the prescriber attests the member is unable to tolerate combination therapy with a phosphodiesterase type 5 inhibitors (PDE5i), endothelin receptor antagonists (ERAs), soluble guanylate cyclase stimulator (sGCs), or prostacyclin

Initial Duration of Approval: 6 months

Reauthorization criteria

• Must provide documentation that demonstrates member is tolerating and receiving clinical benefit from treatment.

Reauthorization Duration of Approval: 12 months

Coverage may be provided for any non-FDA labeled indication if it is determined that the use is a medically accepted indication supported by nationally recognized pharmacy compendia or peer-reviewed medical literature for treatment of the diagnosis(es) for which it is prescribed. These requests will be reviewed on a case by case basis to determine medical necessity.

When criteria are not met, the request will be forwarded to a Medical Director for review. The physician reviewer must override criteria when, in their professional judgment, the requested medication is medically necessary.



PULMONARY ARTERIAL HYPERTENSION INJECTABLE AGENTS PRIOR AUTHORIZATION FORM

Please complete and fax all requested information below including any progress notes, laboratory test results, or chart documentation as applicable to Highmark Wholecare Pharmacy Services. **FAX:** (888) 245-2049

1.1	• •	esentative. PHONE : (80	00) 392-1147 Mon – Fri 8:30am to 5:00pm		
	PROVIDER I	NFORMATION			
Requesting Provider:			Provider NPI:		
Provider Specialty:			Office Contact:		
State license #:			Office NPI:		
Office Address:			Office Phone:		
			Office Fax:		
MEMBER INFORMATION					
Member Name: DOB:					
Member ID: Member			Height:		
	REQUESTED DR	UG INFORMATION			
Medication: Streng					
		Quantity:	Refills:		
Is the member currently receiving rec	quested medication? Yes		Medication Initiated:		
Billing Information					
This medication will be billed: at a pharmacy OR medically, JCODE:					
		per's home Other			
Place of Service Information					
Name:		NPI:			
Address:		Phone:			
	MEDICAL HISTORY (Complete for ALL req	uests)		
Diagnosis:		ICD Code:			
Has chart documentation of right-heart catheterization or echocardiography been provided confirming the diagnosis					
of pulmonary arterial hypertensi	on? Yes No				
Mean Pulmonary Arterial Pressure: Pulmonary Capillary Wedge Pressure: _			Pressure:		
Left Atrial Pressure: Left Ventricular End-diastolic Pressure:					
Pulmonary Vascular Resistance: Date of Exam:					
Please select the World Health Organization (WHO) Classification of Pulmonary Hypertension:					
Group 1 Group 2 Group 3 Group 4 Group 5					
Please indicate WHO functional			lass III 🔲 Class IV		
Will the requested medication be	used as monotherapy or	Monotherapy			
combination therapy?		Combination			
If combination therapy, please list other drug(s):					
CURRENT or PREVIOUS THERAPY					
Medication Name	Strength/ Frequency	Dates of Therapy	Status (Discontinued & Why/Current)		



PULMONARY ARTERIAL HYPERTENSION (PAH) PRIOR AUTHORIZATION FORM (CONTINUED)- PAGE 2 of 2

Please complete and fax all requested information below including any progress notes, laboratory test results, or chart documentation

1.1	ole to Highmark Wholecare	•	X: (888) 245-2049 00) 392-1147 Mon – Fri 8:30am to 5:00pm			
if fleeded, you may can to speak to	· ·	NFORMATION	30, 372 1147 Wolf 111 0.30am to 3.00pm			
Member Name:		DOB:				
Member ID:		Member weight:	Height:			
MEDICAL HISTORY (Complete for ALL requests)						
Please check any boxes applicable to the member: ☐ Clinical signs of right heart failure ☐ Repeated episodes of syncope, even with little or regular physical activity ☐ Presence of pericardial effusion ☐ Peak oxygen consumption (VO2) <11ml/min/kg (<35% predicted) ☐ Ventilatory equivalents of CO2 (VE/VCO2) slope ≥45 ☐ <165 meter 6-minute walking distance (6MWD) ☐ Mixed venous oxygen saturation (SvO2) <60% ☐ Cardiac index <2.0 l/min/m2 ☐ BNP >300 ng/l ☐ NT-proBNP >1400ng/l ☐ Right atrium area ≥26 cm2 ☐ Right atrial pressure >14 mmHg						
Medication Name	Strength/ Frequency	Dates of Therapy	Status (Discontinued & Why/Current)			
	DEALITH	ODIZATION				
REAUTHORIZATION Has the member experienced a significant improvement with treatment? Yes No Please describe:						
SUPPORTING INFORMATION or CLINICAL RATIONALE						
D :: D ::	G: /					
Prescribing Provide	er Signature		Date			