

**PRIOR AUTHORIZATION REQUEST FORM**
**PULMONARY ARTERIAL HYPERTENSION (PAH) MEDICATIONS**

 Adempas®, Flolan®, Letairis®, Opsumit®, Orenitram®, Remodulin®, Tracleer®, Tyvaso®, Uptravi®, Veletri®,  
 Ventavis®

**For authorization, please answer each question and fax this form PLUS chart notes back to the U of U Health Plans Prior Authorization Department. Failure to submit clinical documentation to support this request will result in delay and/or denial of the request.**

- For **Medical Pharmacy** please fax requests to 801-213-1547
- For **Retail Pharmacy** requests please fax requests to: Commercial Groups, Individual & Family Plans please fax request to 888-509-8142. For Healthy U Medicaid, University of Utah Health Employees please fax request to 844-316-3655

If you have prior authorization questions, please call for assistance: Healthy U: 855-856-5694, University of Utah Health Employees: 855-856-5690, Individual & Family Plans: 855-869-4769, Commercial Groups: 855-859-4892, MHC: 855-885-7695, Advantage U Part B: 888-605-0858

Date:	Member Name:	ID#:
DOB:	Gender:	Physician:
Office Phone:	Office Fax:	Office Contact:
Height/Weight:	HCPCS Code:	

**Member must try formulary preferred drugs before a request for a non-preferred drug may be considered. If treatment with preferred products has not been successful, you must submit which preferred products have been tried, dates of treatment, and reason for failure. Reasons for failure must meet the Health Plan medical necessity criteria.**

**Preferred:**  Adempas® (riociguat),  ambrisentan,  epoprostenol,  Opsumit® (macitentan)  Orenitram® (treprostinil),  
 Remodulin (treprostinil),  treprostinil intravenous,  Uptravi® (selexipag)

**Non-preferred:**  bosentan,  Tyvaso® (treprostinil),  Ventavis® (iloprost)

*\*bosentan and Opsumit® require trial and failure of ambrisentan*

Dosing/Frequency: \_\_\_\_\_

**If the request is for reauthorization, proceed to reauthorization section**

Questions	Yes	No	Comments/Notes
1. Does the member have a diagnosis of Pulmonary Arterial Hypertension (PAH)?	<input type="checkbox"/>	<input type="checkbox"/>	<b>Please provide documentation</b>
2. Is the member classified as WHO (World health Organization) Group 1 pulmonary arterial hypertension? If not, please provide the WHO group classification.	<input type="checkbox"/>	<input type="checkbox"/>	<b>Please provide documentation</b>
3. Is the requesting provider a cardiologist or pulmonologist specializing in pulmonary hypertension?	<input type="checkbox"/>	<input type="checkbox"/>	
4. Does the member have regular follow up visits with the prescriber?	<input type="checkbox"/>	<input type="checkbox"/>	<b>Please provide documentation</b>
5. Has the member demonstrated at least 80% compliance with pulmonary hypertension medications?	<input type="checkbox"/>	<input type="checkbox"/>	<b>Please provide documentation</b>

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6. If the member has a positive vasoreactivity test, have they had a trial and failure of oral calcium channel blocker therapy with dihydropyridine or diltiazem?	<input type="checkbox"/>	<input type="checkbox"/>	<b>Please provide documentation</b>
7. Has the member performed a baseline 6-minute walk test?	<input type="checkbox"/>	<input type="checkbox"/>	<b>Please provide documentation</b>
8. Is the member currently smoking or vaping?	<input type="checkbox"/>	<input type="checkbox"/>	
9. For member with a history of stimulant drug abuse, has a recent (within the past 30 days) clean urine drug screen (UDS) been provided?	<input type="checkbox"/>	<input type="checkbox"/>	<b>Please provide documentation</b>
<b>ENDOTHELIN RECEPTOR ANTAGONISTS: AMBRISENTAN, BOSENTAN, OPSUMIT®</b>			
1. Will the medication be used in combination with a phosphodiesterase inhibitor?	<input type="checkbox"/>	<input type="checkbox"/>	<b>Please provide documentation</b>
2. If the request is for bosentan or Opsumit®, has ambrisentan been trialed and failed?	<input type="checkbox"/>	<input type="checkbox"/>	<b>Please provide documentation</b>
<b>PROSTACYCLIN PATHWAY AGONISTS: ORENITRAM®, TREPROSTINIL IV, TREPROSTINIL SQ, REMODULIN®, TYVASO® UPTRAVI®, VENTAVIS®</b>			
1. Does the following apply: <ul style="list-style-type: none"> <li>• Has the member tried and failed combination treatment with a PDE5 inhibitor with ambrisentan or does clinical documentation show a medical reason why the member cannot?</li> <li>• Is the member in WHO functional class III or IV?</li> </ul>	<input type="checkbox"/>	<input type="checkbox"/>	<b>Please provide documentation</b>
2. For Tyvaso® and Ventavis® only, has the member had a trial and failure to treprostinil IV or SQ?	<input type="checkbox"/>	<input type="checkbox"/>	<b>Please provide documentation</b>
<b>GUANYLATE CYCLASE STIMULATOR: ADEMPAS®</b>			
1. Is the member in WHO functional class II, III or IV?	<input type="checkbox"/>	<input type="checkbox"/>	<b>Please provide documentation</b>
2. If the member has a clinical diagnosis of WHO group 1 PAH, have they tried and failed combination therapy with a PDE5 inhibitor with ambrisentan?	<input type="checkbox"/>	<input type="checkbox"/>	<b>Please provide documentation</b>
3. Does the member have a clinical diagnosis of WHO Group 4 PAH after surgical treatment OR have confirmed inoperable chronic thromboembolic pulmonary hypertension?	<input type="checkbox"/>	<input type="checkbox"/>	<b>Please provide documentation</b>
<b>REAUTHORIZATION</b>			
1. Is the request for reauthorization of therapy?	<input type="checkbox"/>	<input type="checkbox"/>	
2. Does documentation show disease improvement or stabilization (e.g. improvement in 6 minute walk test, functional class, pulmonary arterial pressure, cardiac index, etc.)?	<input type="checkbox"/>	<input type="checkbox"/>	<b>Please provide documentation</b>
<b>What medications and/or treatment modalities have been tried in the past for this condition? Please document name of treatment, reason for failure, treatment dates, etc.</b>			

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Additional information:

Physician's Signature:

**\*\*Failure to submit clinical documentation to support this request will result in delay and/or denial of the request\*\***

Policy PHARM- 063  
Origination Date: 05/02/2018  
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