MHS PHARMACY BENEFIT PULMONARY ANTIHYPERTENSIVES PRIOR AUTHORIZATION REQUEST FORM

MHS 550 N. Meridian St. Suite 101 Indianapolis, IN, 46204-1208 Phone: (877) 647-4848 Fax: (866) 399-0929



Today's	Dat	te			
	/		/		

Note: This form must be completed by the prescribing provider.

******All sections must be completed or the request will be returned**

Patient's Medicaid #	Date of Birth
Patient's Name	Prescriber's Name
Prescriber's IN License #	Specialty
Prescriber's NPI #	Prescriber's Signature
Return Fax # - - -	Return Phone # - - -
Check box if requesting retro-active PA	Date(s) of service requested for retro-active eligibility (if applicable):

Note: Submit PA requests for retroactive claims (dates of service prior to eligibility determination, but within established eligibility timelines) with dates of service prior to 30 calendar days of submission separately from current PA requests (dates of service 30 calendar days or less and going forward).

Requested Medication	Strength	Quantity	Dosage Regimen

General information applicable to all products:

Pulmonary Antihypertensive PA Requirements for ALL agents:
1. Member has a diagnosis of pulmonary hypertension \Box Yes \Box No
2. Member has a diagnosis of pulmonary hypertension associated with interstitial lung disease (only
applicable to Tyvaso/Tyvaso DPI) 🗌 Yes 🗌 No
3. Member has a diagnosis of chronic thromboembolic pulmonary hypertension (CTEPH) (only
applicable to Adempas) 🗌 Yes 🗌 No
4. Requested agent has been prescribed by, or in consultation with, a pulmonologist or cardiologist

Product specific information:

l lf i	the request is for Adempas (riociguat):
	For those of childbearing potential, a negative pregnancy test obtained in the past 30 days has been submitted
2.	Member is currently receiving one of the following: nitrate therapy, PDE5 inhibitor, nonspecific PDE inhibitor (dipyridamole; theophylline; aminophylline), vericiguat \Box Yes \Box No
3.	Member is enrolled in the riociguat REMS program if meeting eligibility requirement Yes No Not applicable to member
4.	Requested dose is 7.5mg per day or less $\ \square$ Yes $\ \square$ No
	If no, please explain:
If	the request is for Adcirca (tadalafil):
	Member is currently receiving one of the following: nitrate therapy, PDE-5 inhibitor (other than the one being requested), riociguat
2.	Dose requested is 40 mg per day or less $\ \square$ Yes $\ \square$ No
	Note: 'Alyq' requires trial and failure of generic tadalafil or medical justification for use
If	the request is for Letairis (ambrisentan):
	Member is enrolled in the ambrisentan or PS-ambrisentan REMS program if meeting eligibility requirement
2.	
	For those of childbearing potential, a negative pregnancy test obtained in the past 30 days has been submitted
3.	been submitted
3. 4.	been submitted
	been submitted Yes No Not applicable to member Date of negative pregnancy test (include documentation):
4. 5.	been submitted Yes No Not applicable to member Date of negative pregnancy test (include documentation): Member is currently receiving cyclosporine therapy (requires dose reduction) Yes No Note: dose of Letairis (ambrisentan) must be adjusted to max: 5 mg/day Member has had a previous trial and failure of Tracleer (bosentan) Yes No If no, please explain Dose requested is 10 mg per day or less Yes No
4. 5.	been submitted Yes No Not applicable to member Date of negative pregnancy test (include documentation):
4. 5.	been submitted Yes No Not applicable to member Date of negative pregnancy test (include documentation): Member is currently receiving cyclosporine therapy (requires dose reduction) Yes No Note: dose of Letairis (ambrisentan) must be adjusted to max: 5 mg/day Member has had a previous trial and failure of Tracleer (bosentan) Yes No If no, please explain Dose requested is 10 mg per day or less Yes No The request is for Ligrev (sildenafil) oral suspension:

- PDE-5 inhibitor (other than the one being requested) \Box Yes \Box No
- 4. Dose requested is 60 mg per day or less $\ \square$ Yes $\ \square$ No
- 5. Member has had a previous trial and failure of sildenafil suspension □ Yes □ No If no, please explain_____

If the request is for Opsumit (macitentan):

- 1. Member is enrolled in the macitentan REMS program if meeting eligibility requirement □ Yes □ No □ Not applicable to member
- For those of childbearing potential, a negative pregnancy test obtained in the past 30 days has been submitted □ Yes □ No □ Not applicable to member Date of negative pregnancy test (include documentation): ______
- 3. Member has had a previous trial and failure of Tracleer (bosentan)
 Yes No If no, please explain

4. Dose requested is 10 mg per day or less \Box Yes \Box No

If the request is for Orenitram (treprostinil):

Does the member have severe hepatic impairment (Child-Pugh class C)?
 Yes
 No
 Note: members with Child-Pugh class C hepatic impairment will be denied; Orenitram titration packs will be limited to 1 pack per 90 days

If the request is for Revatio (sildenafil) tablets or injection:

- Member is currently receiving one of the following: nitrate therapy, riociguat, atazanavir, darunavir, fosamprenavir, indinavir, lopinavir/ritonavir, nelfinavir, ritonavir, saquinavir, tipranavir, PDE-5 inhibitor (other than the one being requested) □ Yes □ No
- 2. Dose requested is 60 mg per day or less \Box Yes \Box No

If the request is for Revatio (sildenafil) oral suspension:

- 1. Member is under 12 years of age \Box Yes \Box No
- 2. Member is unable to swallow tablet formulation \Box Yes \Box No
- 3. Member is currently receiving one of the following: nitrate therapy, riociguat, atazanavir, darunavir, fosamprenavir, indinavir, lopinavir/ritonavir, nelfinavir, ritonavir, saquinavir, tipranavir, PDE-5 inhibitor (other than the one being requested) □ Yes □ No
- 4. Dose requested is 60 mg per day or less \Box Yes \Box No

If the request is for Tadliq (tadalafil) oral suspension:

- 1. Member is under 12 years of $age \square Yes \square No$
- 2. Member is unable to swallow tablet formulation \Box Yes \Box No
- 3. Member is currently receiving one of the following: nitrate therapy, PDE-5 inhibitor (other than the one being requested), riociguat □ Yes □ No
- 4. Dose requested is 40 mg per day or less \Box Yes \Box No
- 5. Member has had a previous trial and failure of sildenafil oral suspension \Box Yes \Box No
 - If no, please explain____

If the request is for Uptravi (selexipag):
1. Member has had a previous trial and failure of Orenitram (treprostinil) \Box Yes \Box No
If no, please explain
2. Will the member be utilizing a CYP2C8 inhibitor (e.g., gemfibrozil) concurrently with selexipag? □ Yes □ No
Note: members planning to use CYP2C8 inhibitors concurrently with selexipag will be denied
If the request is for Tracleer (bosentan):
Request is for:
 Tracleer tablet Tracleer dispersible tablet
\square bosentan tablet [*]
1. Member is enrolled in the bosentan REMS program (Note: ALL members must be enrolled in the
bosentan REMS program) 🗌 Yes 🗌 No
 For those of childbearing potential, a negative pregnancy test obtained in the past 30 days has been submitted
3. Will the member be utilizing cyclosporine-A or glyburide therapy concurrently with bosentan? □ Yes □ No
Note: members planning to use cyclosporine-A or glyburide concurrently with bosentan will be denied
4. Member age: weight: LB/KG (circle one)
5. Does the requested dose exceed 250mg per day OR dose limits based on age/weight listed in
criteria? 🗌 Yes 🗌 No
If yes, please explain:
Note: Tracleer tablets are brand preferred. Authorization for generic bosentan tablets is contingent upon medical necessity for use instead of the branded agent.

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