INDIANA HEALTH COVERAGE PROGRAMS (IHCP) PHARMACY BENEFIT PULMONARY ANTIHYPERTENSIVES PRIOR AUTHORIZATION REQUEST FORM



MDwise Fax to: (858) 790-7100 c/o MedImpact Healthcare Systems, Inc. Attn: Prior Authorization Department 10181 Scripps Gateway Court, San Diego, CA 92131 Phone: (800) 788-2949



Today's Date										
Note: This form must be completed by the pres **All sections mus			e request will be	returned	**					
Patient's Medicaid #		Date of Birth / / / /								
Patient's Name	Presci	Prescriber's Name								
Prescriber's IN License #	Specia	Specialty								
Prescriber's NPI #		Prescr	iber's Signature							
Return Fax #		Retur	Phone #	-			-			
Check box if requesting retro-active PA			Date(s) of service requested for retro-active eligibility (if applicable):							
timelines) with dates of service prior to 30 calendar d days or less and going forward).	ays of subi	mission sepc	rately from current	PA request	ts (date	es of s	ervic	:e 30	calen	dar
Requested Medication Stren	gth C	Quantity	Dosage Regimen							
General information applicable to a Pulmonary Antihypertensive PA R										
1. Member has a diagnosis of pulmonar			Yes ☐ No							
2. Member has a diagnosis of pulmonar	y hypert	tension as	sociated with	interstitia	al lun	g dis	eas	se (c	only	
applicable to Tyvaso/Tyvaso DPI) \Box	Yes □ N	No								
Note: A diagnosis of pulmonary hypertensi	ion is red	quired for	plan approval,	excluding	Ade	mpas				
3. Requested agent has been prescribe ☐ Yes ☐ No	d by, or	in consul	ation with, a p	ulmonolo	ogist	or ca	ardi	olog	gist	

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Product specific information:

lf t	he request is for Adempas (riociguat):
1.	Please select member's diagnosis
	□ Pulmonary hypertension
	☐ Chronic thromboembolic pulmonary hypertension (CTEPH)
2.	Member has had a negative pregnancy test in the past 30 days ☐ Yes ☐ No ☐ Not applicable to member Date of negative pregnancy test (include documentation):
3.	Member is currently receiving one of the following: nitrate therapy, PDE5 inhibitor, nonspecific PDE inhibitor (dipyridamole; theophylline; aminophylline), vericiguat \square Yes \square No
4.	Member is enrolled in the riociguat REMS program if meeting eligibility requirement ☐ Yes ☐ No ☐ Not applicable to member
5.	Requested dose is 7.5mg per day or less $\ \square$ Yes $\ \square$ No
	If no, please explain:
lf t	he request is for Adcirca (tadalafil):
1.	Member is currently receiving one of the following: nitrate therapy, PDE-5 inhibitor (other than the one being requested), riociguat \Box Yes \Box No
2.	Dose requested is 40 mg per day or less ☐ Yes ☐ No
	Note: 'Alyq' requires trial and failure of generic tadalafil or medical justification for use
lt t	he request is for Letairis (ambrisentan):
1.	Member is enrolled in the ambrisentan or PS-ambrisentan REMS program if meeting eligibility requirement ☐ Yes ☐ No ☐ Not applicable to member
2.	Member has had a negative pregnancy test in the past 30 days ☐ Yes ☐ No ☐ Not applicable to member Date of negative pregnancy test (include documentation):
3.	Member is currently receiving cyclosporine therapy (requires dose reduction) ☐ Yes ☐ No Note: dose of Letairis (ambrisentan) must be adjusted to max: 5 mg/day
4.	Member has had a previous trial and failure of Tracleer (bosentan) $\ \square$ Yes $\ \square$ No
	If no, please explain
5.	Dose requested is 10 mg per day or less ☐ Yes ☐ No

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If t	the request is for Opsumit (macitentan):
1.	Member is enrolled in the macitentan REMS program if meeting eligibility requirement \Box Yes \Box No \Box Not applicable to member
2.	Member has had a negative pregnancy test in the past 30 days ☐ 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) $\ \square$ Yes $\ \square$ No
	If no, please explain
4.	Dose requested is 10 mg per day or less ☐ Yes ☐ No
lf 1	the request is for Orenitram (treprostinil):
1.	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
If t	the request is for Revatio (sildenafil) tablets or injection:
1.	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) \square Yes \square No
2.	Dose requested is 60 mg per day or less ☐ Yes ☐ No
lf t	the request is for Revatio (sildenafil) oral suspension:
1.	Member is under 18 years of age ☐ Yes ☐ No
2.	Member is unable to swallow tablet formulation $\ \square$ Yes $\ \square$ 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) \square Yes \square No
4.	Dose requested is 60 mg per day or less $\ \square$ Yes $\ \square$ No
	Note: Revatio Suspension is brand preferred. Authorization for generic sildenafil oral suspension is contingent upon medical necessity for use instead of the branded agent.

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If the request is for Tadliq (tadalafil) oral suspension:					
1. Member is under 18 years of age ☐ Yes ☐ No					
2. Member is unable to swallow tablet formulation $\ \square$ Yes $\ \square$ 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 ☐ Yes ☐ No					
5. Member has had a previous trial and failure of Revatio (sildenafil) oral suspension ☐ Yes ☐ No If no, please explain					
If the request is for Uptravi (selexipag):					
1. Member has had a previous trial and failure of Orenitram (treprostinil) ☐ Yes ☐ 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 ☐ Bosentan tablet					
1. Member is enrolled in the bosentan REMS program (<i>Note: ALL members must be enrolled in the bosentan REMS program</i>) ☐ Yes ☐ No					
2. Member has had a negative pregnancy test in the past 30 days ☐ Yes ☐ No ☐ Not applicable to member Date of negative pregnancy test (include documentation):					
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					

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