

Kaiser Permanente Health Plan of Mid-Atlantic States, Inc. UPTRAVI (Selexipag), TYVASO (Treprostinil), ORENITRAM (Treprostinil/Diolamine).

Prior Authorization (PA)

Pharmacy Benefits Prior Authorization Help Desk Length of Authorizations: Initial- 12 months; Continuation- 12 months

## **Instructions:**

This form is used by Kaiser Permanente and/or participating providers for coverage **UPTRAVI** (Selexipag), **TYVASO** (**Treprostinil**), **ORENITRAM** (**Treprostinil/Diolamine**). Please complete all sections, incomplete forms will delay processing. Fax this form back to Kaiser Permanente within 24 hours fax: 1-866-331-2104. If you have any questions or concerns, please call 1-866-331-2103. **Requests will not be considered unless all sections are complete. <b>KP-MAS Formulary can be found at:** http://www.providers.kaiserpermanente.org/mas/formulary.html

	1 – Patient Information	
Patient Name:	Kaiser Medical ID#:	Date of Birth:
	2 – Prescriber Information	
Is the prescriber a Pulmonologist or Cardiologis	st? 🗆 No 🗆 Yes	
If consulted with a specialist, specialist name ar	nd specialty:	
Prescriber Name:	Specialty:	NPI:
Prescriber Address:		
Prescriber Phone #:	Prescriber Fax #:	
	3 – Pharmacy Information	
Pharmacy Name:	Pharmacy NPI:	
Pharmacy Phone #	Pharmacy Fax #:	
	4 – Drug Therapy Requested	
Drug 1: Name/Strength/Formulation:		
Drug 2: Name/Strength/Formulation:		
Sig:		

## 5- Diagnosis/Clinical Criteria

	iber Signature: Date:		
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	tify that the information provided is accurate. Supporting documentation is available for State audits.		
	onal Information – Please submit chart notes/medical records for the patient that are applicable to this request. e any additional supporting information that should be taken into consideration:		
	6 – Prescriber Sign-Off		
	□ No □ Yes		
2.	Does member continue to meet initial review criteria?		
	disease improvement? <b>AND</b>		
1.	Is there documentation the member is experiencing clinical benefit from therapy as evidenced by disease stability or		
or Co	ntinuation of Therapy, Please Respond to Additional Questions Below:		
	□ No □ Yes		
	macitentan (Opsumit) <b>OR</b> c. A soluble guanylate cyclase stimulator Riociguat (Adempas®)		
	b. One endothelin receptor antagonist (ERA) (e.g. ambrisentan (Letairis®), or bosentan (Tracleer®) <b>OR</b>		
6.	Is there documentation treatment failure, intolerance, or contraindication to at least two of the following:  a. One phosphodiesterase type (PDE-5) inhibitor (e.g. sildenafil (Revatio®), tadalafil (Adcirca®) OR		
5.	Is member currently receiving a prostanoid/prostacyclin analogue (e.g. treprostinil (Orenitram®, Tyvaso®, Remodulin®)? <b>AND</b>		
_	□ No □ Yes		
	AND		
4.	□ No □ Yes  Does the member have a diagnosed with WHO/New York Heart Association Functional Class II, III or IV symptoms?		
	OR		
	Does the member have a diagnosis of pulmonary arterial hypertension World Health Organization [WHO] Group I?		
2.	Indicate the Member's diagnosis for the requested medication:		
	Is this request for initial or continuing therapy?  □ Initial therapy □ Continuing therapy, State date:		
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