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### Treprostinil Injection is available through the Specialty Pharmacy (SP) provider listed on page 7.

**Complete all sections on this enrollment form.** Let your patient know that the Specialty Pharmacy will be calling to process their prescription and that it is important to answer or return any messages.

Sign the Statement of Medical Necessity on page 3 for the Prescription.

Sign at the bottom of pages 4 and 5.

Fax the enrollment form and signed supporting documents (use Fax Cover Sheet provided on page 7) to the SP.

Information regarding the Centers for Medicare and Medicaid Services (CMS) established and expected coverage criteria for prostacyclin is included for your convenience.

## MEDICARE COVERAGE CRITERIA FOR PROSTACYCLIN

## The current Local Coverage Determination for Prostacyclin is as follows:

The pulmonary hypertension is not secondary to pulmonary venous hypertension (e.g., left sided atrial or ventricular disease, left sided valvular heart disease, etc.) or disorders of the respiratory system (e.g., chronic obstructive pulmonary disease, interstitial lung disease, obstructive sleep apnea or other sleep disordered breathing, alveolar hypoventilation disorders, etc.); and

The patient has idiopathic/heritable pulmonary hypertension or pulmonary hypertension which is associated with one of the following conditions: connective tissue disease, thromboembolic disease of the pulmonary arteries, human immunodeficiency virus (HIV) infection, cirrhosis, diet drugs, congenital left to right shunts, etc.

## If the above conditions are present, the following criteria must be met:

The pulmonary hypertension has progressed despite maximal medical and/or surgica treatment of the identified condition; and
The mean pulmonary artery pressure is greater than 25 mm Hg at rest or greater than 30 mm Hg with exertion; and
The patient has significant symptoms from the pulmonary hypertension (i.e., severe dyspnea on exertion, and either fatigability, angina, or syncope); and
Treatment with oral calcium channel blocking agents has been tried and failed or has been considered and ruled out.

Medicare coverage criteria provided for informational purposes only. Please check with the payer to verify billing requirements. Liquidia and Sandoz do not make any representation or guarantees concerning reimbursement or coverage for any service or item.





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PATIENT INFORMATION	
	Gender:
Patient Name (first, MI, last)	Date of Birth (mm/dd/yyyy)
Address	Email
	☐ Cell ☐ Cell ☐ Other ☐ Other
City State Zip	Phone Alternate Phone
SHIPPING ADDRESS (if different from above):	Preferred contact: Phone Email
Address	Best time to call:
Address	OK to leave message with Caregiver? Yes No
City State Zip	The search message man careginal.
CAREGIVER	
	☐ Cell ☐ Cell ☐ Other ☐ Other
Caregiver Name	Caregiver Phone Alternate Phone
	Preferred contact: Phone Email
Caregiver Email	Best time to call: OMorning Afternoon Night
INSURANCE INFORMATION	
	Please include copies of the front and back of all
Pharmacy Benefits Manager	patient's medical and prescription insurance cards.
, ,	
PRIMARY Medical Insurance Carrier	SECONDARY Medical Insurance Carrier
Policyholder Name	Policyholder Name
Policy ID Number Group No (if applicable)	Policy ID Number Group No (if applicable)
Medical Insurance Phone Relationship to Policyholder	Medical Insurance Phone Relationship to Policyholder



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PRESCRIBER INFORMATION	Patient Name (first, MI, last)  Date of Birth
Prescriber Name (first, MI, last)	NPI # State License # Tax ID #
Office / Clinic / Institution Name  Address  City State Zip	Office Contact Name  Office Contact Email  Phone  Fax
PRESCRIPTION INFORMATION	Preferred method of communication: Phone Email Fax
Sandoz® Treprostinil Injection vial concentration  NDC(s) prescribed:  O 1 mg/mL (20-mL vial) (00781-3420-80) O 2.5 mg/mL (20-mL vial) (00781-3425-80) O 5 mg/mL (20-mL vial) (00781-3427-80) O 10 mg/mL (20-mL vial) (00781-3430-80)  Diluent: (0.9% Sodium Chloride will be used if no box is checked) O 0.9% Sodium Chloride for Injection O Sandoz® Sterile Diluent for Treprostinil Injection O Sterile Water for Injection O Epoprostenol Sterile Diluent for Injection Infusion route and pumps: O Subcutaneous continuous infusion with appropriate ambulatory infusion pump. O Intravenous continuous infusion with appropriate ambulatory infusion pump.	Dosing and titration instructions  Patient dosing weight: Initiation dosage:
I certify that the pulmonary arterial hypertension therapy ordered above is medically necessary and that I am personally supervising the care of this patient.  Dispense As Written (DAW) / Brand Medically Necessary /	Prescriber Full Name (print) Substitution Permitted / May Substitute / Product Selection Permitted
CA, MA, NC & PR: Interchange is mandated unless Prescribe	scription requirements such as e-prescribing, state specific prescription form,

NOTE: The responsibility to determine coverage and reimbursement parameters, and appropriate coding for a particular patient and/or procedure, is the responsibility of the provider. The information provided here is not a guarantee of coverage or reimbursement.





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Patient Name (first, MI, last)	Date of Birth	Prescriber Name (first, MI, last)	NPI #			
NURSING ORDERS						
NURSE VISITS (select one option)						
SP home healthcare RN visit(s) to provide assessment and education on self-administration of Treprostinil to include dose, titration, and						
side effect management OR						
O Prescriber-directed SP home healthcare RN visit(s)	Prescriber-directed SP home healthcare RN visit(s) as detailed below:					
Location: O Home O Outpatient clinic O Hos	nital O Virtual					
	,					
SITE CARE						
O Dressing change every days						
Per standard of care						
CALCIUM CHANNEL BLOCKER STATEMENT						
Indicate whether the patient named above was triale	ed on a calcium chann	el blocker prior to the initiation of therapy a	and provide the results			
A calcium channel blocker was not trialed because:		The following calcium channel blocker wa	•			
Patient has depressed cardiac input		The following calcium channel blocker we	is tridied.			
Patient has depressed cardiac input     Patient has systematic hypotension						
O Patient has known hypersensitivity		The patient had the following response(s)	:			
O Patient is hemodynamically unstable or has a history	ory of	Patient hypersensitive or allergic				
postural hypotension	,	Adverse event				
O Patient did not meet ACCP Guidelines for Vasodila	ator Response	O Patient became hemodynamically uns	stable			
O Patient has documented brachycardia or second or		O Pulmonary arterial pressure continued to rise				
third-degree heartblock		O Disease continued to progress, or pat	cient remained symptomatic			
Other:		Other:				
<u> </u>						
PRESCRIBER SIGNATURE						
SIGN						
HERE	Presci	riber Signature	Date			
Prescriber Full Name (print)		riber Signature				
Prescriber Full Name (print)  NOTE: The responsibility to determine cov	verage and reimbursem	riber Signature  ent parameters, and appropriate coding for a rovided here is not a guarantee of coverage o	particular patient and/or			



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atient Name (first, MI, last)	Date of Birth	Prescriber Name (first, MI, last) NPI #
ATIENT EVALUATION		MEDICAL INFORMATION
Actient Status: Treprostinil Injection Status: Outpatient Naïve / New Inpatient Restart Date To Transition  Allergies: No known drug allergies (NKDA) Yes (specify): Viabetic? Yes WHO Group: No NYHA Functional Class: Current Medications (list all):	kg lb t	REQUIRED: Please select one of the following ICD-10 codes, or CICD-10 code, as applicable. The following ICD-10 codes do not su approval, coverage, or reimbursement for specific uses or indicat  DIAGNOSIS  ICD-10 127.0 Primary pulmonary hypertension  Idiopathic PAH Heritable PAH  ICD-10 127.2 Other secondary pulmonary hypertension Connective tissue disease Drugs/Toxins induced HIV  Other ICD-10:  Code Description
REATMENT HISTORY		TRANSITION STATEMENT (if applicable)
Please indicate treatment history		It is necessary for this patient to transition
dempas® (riociguat) Tablets	Current	O Discontinued from:
poprostenol Sodium for Injection	Current	O Discontinued to:
lolan* (epoprostenol sodium) for Injection	Current	Discontinued
etairis* (ambrisentan) Tablets	Current	O Discontinued Please provide justification for this transitio
Opsumit* (macitentan) Tablets	O Current	O Discontinued
Orenitram® (treprostinil) Extended-Release Tablets	Current	O Discontinued
		O Discontinued
DE-5i (specify drugs):	Current	
DE-5i (specify drugs): emodulin* (treprostinil) Injection	O Current	ODiscontinued
DE-5i (specify drugs): emodulin* (treprostinil) Injection racleer* (bosentan) Tablets	O Current O Current	O Discontinued O Discontinued
DE-5i (specify drugs): emodulin* (treprostinil) Injection racleer* (bosentan) Tablets yvaso* (treprostinil) Inhalation Solution	Current Current Current	O Discontinued O Discontinued O Discontinued
DE-5i (specify drugs):  emodulin* (treprostinil) Injection  racleer* (bosentan) Tablets  yvaso* (treprostinil) Inhalation Solution  yvaso* DPI (treprostinil) Inhalation Powder	Current Current Current Current Current	O Discontinued O Discontinued O Discontinued O Discontinued
DE-5i (specify drugs):  emodulin* (treprostinil) Injection  racleer* (bosentan) Tablets  yvaso* (treprostinil) Inhalation Solution  yvaso* DPI (treprostinil) Inhalation Powder  ptravi* (selexipag) Tablets	Current Current Current Current Current Current	O Discontinued O Discontinued O Discontinued O Discontinued O Discontinued
PDE-5i (specify drugs):  Remodulin* (treprostinil) Injection  Tracleer* (bosentan) Tablets  Tyvaso* (treprostinil) Inhalation Solution  Tyvaso* DPI (treprostinil) Inhalation Powder  Uptravi* (selexipag) Tablets  Veletri* (epoprostenol) for Injection	Current Current Current Current Current Current Current Current	O Discontinued O Discontinued O Discontinued O Discontinued O Discontinued O Discontinued
PDE-5i (specify drugs):  Remodulin* (treprostinil) Injection  Tracleer* (bosentan) Tablets  Tyvaso* (treprostinil) Inhalation Solution  Tyvaso* DPI (treprostinil) Inhalation Powder  Uptravi* (selexipag) Tablets  Veletri* (epoprostenol) for Injection  Ventavis* (iloprost) Inhalation Solution	Current Current Current Current Current Current	O Discontinued O Discontinued O Discontinued O Discontinued O Discontinued
PDE-5i (specify drugs):  Remodulin* (treprostinil) Injection  Tracleer* (bosentan) Tablets  Tyvaso* (treprostinil) Inhalation Solution  Tyvaso* DPI (treprostinil) Inhalation Powder  Uptravi* (selexipag) Tablets  Veletri* (epoprostenol) for Injection  Ventavis* (iloprost) Inhalation Solution  Other:  RESCRIBER SIGNATURE	Current Current Current Current Current Current Current Current Current	O Discontinued
PDE-5i (specify drugs): Remodulin* (treprostinil) Injection Fracleer* (bosentan) Tablets Fyvaso* (treprostinil) Inhalation Solution Fyvaso* DPI (treprostinil) Inhalation Powder Uptravi* (selexipag) Tablets Veletri* (epoprostenol) for Injection Ventavis* (iloprost) Inhalation Solution Other:	Current Current Current Current Current Current Current Current Current	O Discontinued





## **Important Safety Information**

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### **INDICATION**

Treprostinil injection is a prostacyclin mimetic indicated for

- Treatment of pulmonary arterial hypertension (PAH), World Health Organization (WHO) Group 1, to diminish symptoms associated with exercise. Studies establishing effectiveness included patients with New York Heart Association (NYHA) Functional Class II-IV symptoms and etiologies of idiopathic or heritable PAH (58%), PAH associated with congenital systemic-to-pulmonary shunts (23%), or PAH associated with connective tissue diseases (19%).
- Patients who require transition from epoprostenol to reduce the rate of clinical deterioration. The risks and benefits of each drug should be carefully considered prior to transition.

### **IMPORTANT SAFETY INFORMATION**

### **CONTRAINDICATIONS**

None.

#### **WARNINGS AND PRECAUTIONS**

- Chronic intravenous (IV) infusions delivered using an external infusion pump with an indwelling central venous catheter are associated with the risk of bloodstream infections (BSIs) and sepsis, which may be fatal. Therefore, continuous subcutaneous (SC) infusion is the preferred mode of administration.
- Do not abruptly lower the dose or withdraw dosing.
- · Treprostinil injection may cause symptomatic hypotension.
- Titrate slowly in patients with hepatic insufficiency because such patients will likely be exposed to greater systemic concentrations relative to patients with normal hepatic function.
- · Treprostinil injection inhibits platelet aggregation and increases the risk of bleeding.

#### **ADVERSE REACTIONS**

During clinical trials with SC infusion of treprostinil, infusion site pain and infusion site reaction (e.g., erythema, induration, or rash) were the most common adverse events and occurred in majority of those treated with treprostinil. Infusion site reactions were sometimes severe and led to discontinuation of treatment. Rash and hypotension (14% and 4%, respectively) were also commonly reported with SC infusion of treprostinil. Other common adverse events (≥3% more than placebo) included headache, diarrhea, jaw pain, edema, vasodilatation, and nausea, and these are generally considered to be related to the pharmacologic effects of treprostinil, whether administered subcutaneously or intravenously. The adverse reactions reported with treprostinil IV included bloodstream infections, arm swelling, paresthesia, hematoma, and pain.

### **DRUG INTERACTIONS**

Treprostinil injection dosage adjustment may be necessary if inhibitors or inducers of CYP2C8 are added or withdrawn.

## **USE IN SPECIFIC POPULATIONS**

- · Safety and effectiveness of Treprostinil injection in pediatric patients have not been established.
- It is unknown if geriatric patients respond differently than younger patients. Caution should be used when selecting a dose for geriatric patients.
- · There are no adequate and well-controlled studies with Treprostinil injection in pregnant women.
- It is not known whether Treprostinil injection is excreted in human milk.

Please see accompanying full Prescribing Information for additional safety information, also available by clicking here.







Data						
Date						
ТО	Accredo Health Group, Inc.					
	FAX 1-800-711-3526					
	Phone: 1-866-344-4874					
FROM						
	(Name of agent of prescriber transmitting this fax/prescription	) Phone				
	Facility Name	Fax				
RE						
	DOCUMENTATION CHECKLIST					
	Indicate all current, signed and dated documents enclose	d with this fax.				
	Fully completed Treprostinil Enrollment Form,	Echocardiogram				
	including:  — Patient/Insurance Information	6-minute walk test results				
	<ul><li>– Prescriber/Prescription Information</li><li>– Medical Information/Patient Evaluation</li></ul>	O History and physical, including onset of symptoms, PAH clinical signs and symptoms				
	<ul><li>Copy of front and back of Patient's Insurance card(s)</li><li>Right heart catheterization</li></ul>	<ul><li>and course of illness</li><li>Need for specific drug therapy</li></ul>				
	Comments:					