Orenitram® (treprostinil) Extended-Release Tablets Referral Form

HOW TO GET STARTED



Follow these 3 steps to complete the referral form.

- 1. Obtain all the necessary documentation from your patient to fill out the Patient Information (**A and B**), and have your patient sign (**C**).
 - Let your patient know that an Access Solutions and Support Team (ASSIST) representative will be calling to verify insurance coverage or to obtain additional information. It is very important he or she answers or returns the call in a timely manner or the approval process could be delayed
 - Obtain a copy of the patient's insurance cards (front and back) to submit with their referral form

2. Complete and sign the following forms:

- Prescriber Information (**D**)
- Medical Information/Patient Evaluation/Supporting Documentation (E)
- Prescription Information (**F**)
- Prescriber Signature (G)

SUPPORT FOR YOU AND YOUR PATIENTS

United Therapeutics Support

ASSIST is a centralized referral service that helps simplify the referral process by providing support until your patients receive their first shipment of medication.



ASSIST will:

- Review referral forms and work with your patients to help determine the best coverage for their medication
- Reach out to your patients directly and screen for financial program eligibility*
- Refer to the Specialty Pharmacy Service best suited to provide medication to each patient based on insurance coverage
- Facilitate processing of patients' referrals and keep you informed of the progress

If you or your patients have any questions about completing the referral forms, the ASSIST Financial Assistance Programs, or program eligibility, please contact ASSIST at 1-877-864-8437.

*Patients must meet certain eligibility criteria to qualify for financial assistance.

Specialty Pharmacy Services (SPS)

SPS providers are available to answer questions from your patients or your practice regarding treatment with Orenitram. SPS nurses provide in-home medication education for patients new to therapy, as well as ongoing support throughout their treatment.

SPS providers will also work with your patient's insurance company and your office to obtain any necessary Prior Authorization. Once the insurance company approves, the SPS will be contacting your patient to review his or her financial responsibility and apply any financial assistance programs offered by United Therapeutics for which the patient qualified.

Think of SPS as a resource to help your patients get the information and support they need to understand the treatment process and manage their condition.

Orenitram is a prostacyclin vasodilator indicated for the treatment of pulmonary arterial hypertension (PAH) (WHO Group 1) to improve exercise capacity.



^{3.} Use the Fax Cover Sheet included in this PDF to fax the completed Referral Form (2 pages) to ASSIST. Include any comments in the section provided on the Cover Sheet.

Orenitram® (treprostinil) Extended-Release Tablets Referral Form

Please complete, sign, and fax Steps 1 and 2 to ASSIST using the following Fax Cover Sheet.



STEP 1 - PATIENT INFORMATION AND AUTHORIZATION

A PATIENT INFORMATION			
Name: First	Middle	Last	
Date of Birth	Gender	SSN	
Home Address			
City	State	Zip	
Shipping Address	(if not home address)		
City	State	Zip	
Telephone	Alternate Telephone	Best Time to Call	
E-mail Address			
Caregiver/Family Member	Telephone	Alternate Telephone	

B INSURANCE INFORMATION

Pharmacy Benefits Manager:

Thanhacy benefits manager.		
Subscriber ID #	Group #	Telephone #
Primary Medical Insurance:		Policy Holder/Relationship
Subscriber ID #	Group #	Telephone #
Secondary Medical Insurance:		Policy Holder/Relationship
Subscriber ID #	Group #	Telephone #

Please include copies of the front and back of the Patient's Insurance Card(s).

PATIENT AUTHORIZATION FOR THE USE AND DISCLOSURE OF PROTECTED HEALTH INFORMATION

I authorize the use and/or disclosure of my private health information, described below, which may include "Protected Health Information" or "PHI" as defined by the Health Insurance Portability and Accountability Act of 1996 (as amended, "HIPAA"). In general terms, I understand that Protected Health Information is health information that identifies me or that could reasonably be used to identify me. I understand that this authorization is voluntary.

I authorize my health care providers, including my pharmacies, and my health plan(s), to disclose information about me as described below to the United Therapeutics Corporation/Lung Biotechnology Inc. Access Solutions and Support Team (ASSIST), its authorized Program Administrator, and its Financial Assistance Partners (collectively, "United Therapeutics") for the purposes stated below.

This information may include:

- · Information about my health benefits, health insurance coverage or other third-party payers
- · Relevant information about my medical condition and history
- Financial information about me
- · Contact information, such as my physical and email address and telephone number
- · Information about my circumstances, such as my marital, veteran, employment, disability and citizenship status
- Identifying information, such as my name, birth date and social security number

This information may be disclosed to United Therapeutics in order for it to: (1) contact me to discuss its various available services; (2) determine my initial and continuing eligibility for the assistance program(s); (3) administer the assistance program(s); (4) identify sources of payment for the provision of medications to me; (5) help me find education and therapy support services; (6) review the success of the services and look at whether patients are happy with them; (7) comply with law and; (8) conduct limited commercial and sales activities.

I understand that once my health care providers, including my pharmacies, and my health plan(s) share information about me to United Therapeutics, the information is no longer protected by federal health privacy laws and may be given out (re-disclosed) to others by United Therapeutics if permitted by laws that apply to United Therapeutics. I know that I may refuse to sign this authorization, and that this refusal will not affect my treatment, payment for treatment, enrollment in a health plan or eligibility for benefits. However, if I do not sign it, I may not be eligible to receive the education and therapy patient support services provided by United Therapeutics.

This authorization will expire ten (10) years after the date it is signed unless a shorter period is mandated by State Law or I revoke or cancel (i.e., take back) my authorization before then. I understand that I may cancel this authorization at any time by fax at 1-800-380-5294 or by writing to: United Therapeutics Corporation/Lung Biotechnology Inc., ASSIST, 1130 S. Harbor City Blvd., Suite 103, Melbourne, Florida 32901, but that the cancellation will not apply to information that my health care providers, including my pharmacies, and health plans have already given out based on this authorization and before they learn about my cancellation. I understand I am entitled to receive a copy of this authorization once signed.

I understand that certain of my health care providers, such as my pharmacy may receive compensation in connection with their disclosure of my information to United Therapeutics for the purposes I allow through this authorization.

I have read this authorization and or had its contents read to me. I have had an opportunity to ask questions about the uses and disclosures of PHI described above and all of my questions have been answered to my satisfaction.

Patient Name (Print)	Patient Signature	Date
If the patient cannot sign, Patient's Representative must sign here. Patient Representative must sign here.	ntative Signature	Date
Describe relationship to patient and authority to sign this form for patient:		

Please note: United Therapeutics cannot guarantee payment for United Therapeutics products and directs patients to discuss treatment options with their healthcare provider.



Please see Indication and Important Safety Information on page 5. Please click links for Full Prescribing Information and Patient Information for Orenitram.

US/ORE/JAN15/070a

Orenitram® (treprostinil) Extended-Release Tablets Referral Form

Please complete, sign, and fax Steps 1 and 2 to ASSIST using the following Fax Cover Sheet.



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		Date of Birth:
EP 2 - PRESCRIBER, MEDICAL AND P	RESCRIPTION INFORMATION	
PRESCRIBER INFORMATION		
Prescriber: First	Last	
NPI #	State License #	
Facility Name	Group NPI # (if applicable)	
Address		
City	State	Zip
Office Contact Name		
elephone		Fax
-mail Address	Preferred Method of Communicati	 on
MEDICAL INFORMATION / PATIENT I	EVALUATION / SUPPORTING DOCUME	ITATION
Interview of the second sec	ested drug Current Specialty Pharmacy	Patient Status Allergies
	Transition	
IO Group NYHA Functional Class		
gnosis - The following ICD-9/ICD-10 codes do not sug	IV Weight	
-9 416.0/ICD-10 I27.0 Primary pulmonary hypertension		rt diseases: pulmonary arterial hypertension, secondary Other ICD-9/ICD-10
Idiopathic PAH		al Heart Disease 🔲 Portal Hypertension
· <u> </u>	Drugs/Toxins Induced HIV	Other
	days u	Intil goal ofmg TID is achieved CTIONS HERE (above fields may be left blank if preferred)
RECTIONS: Take tablets by mouth with food SPENSE: Quantity sufficient for up to maximum all Orenitram dosing and titration information, pleas ecialty Pharmacy to contact Prescriber for adju	se see the Dosage and Administration section of	fills12 Months OR Refills Time
ecify any palliative measure to be taken visit to provide education on self-administration of	of Oronitram to include desing and titration as n	er prescriber order: Location: 🔲 Home 🔲 Hospital 🔲 Outpatient clinic
		faxed and applicable, must be on state specific form.
M DDFCCDIRED CIGNATIIDE: DDFCCDIL	TION AND STATEMENT OF MEDICAE N	supervising the care of this patient. I authorize United Therapeutics ASSIST
ertify that the medication ordered above is n he HUB) to act on my behalf for the limited put	urposes of transmitting this prescription to th	he appropriate pharmacy designated by the Patient utilizing their benefit plan.
ertify that the medication ordered above is n ne HUB) to act on my behalf for the limited pu HYSICIAN SIGNATURE REQUIRED TO VALIDAT ysician's signature	urposes of transmitting this prescription to the PRESCRIPTIONS.	ne appropriate pharmacy designated by the Patient utilizing their benefit plan.
certify that the medication ordered above is n	urposes of transmitting this prescription to the prescriptions.	ne appropriate pharmacy designated by the Patient utilizing their benefit plan.

indication and important Safety Information on page links for <u>Full Prescribing Information</u> and <u>Patient Information</u> for Orenitram.



Please complete, sign, and fax Steps 1 and 2 to ASSIST using this Fax Cover Sheet.



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Date:	
To: Access Solutions and Support Team	Fax Number 1-800-380-5294 Phone Number 1-877-864-8437
From:	
Facility Name:	
Fax:	
Included in this fax: Completed UT PAH Th	erapy Referral Form including
Step 1 - Patient Inf	ormation and Authorization
Step 2 - Prescriber,	Medical, and Prescription Information
Copy of Insurance C	Card(s)
Number of Pages:	
Comments:	



ORENITRAM® (treprostinil) Extended-Release Tablets

INDICATION

Orenitram is a prostacyclin vasodilator indicated for treatment of pulmonary arterial hypertension (PAH) (WHO Group 1) to improve exercise capacity. The study that established effectiveness included predominately patients with WHO functional class II-III symptoms and etiologies of idiopathic or heritable PAH (75%) or PAH associated with connective tissue disease (19%).

When used as the sole vasodilator, the effect of Orenitram on exercise is about 10% of the deficit, and the effect, if any, on a background of another vasodilator is probably less than this. Orenitram is probably most useful to replace subcutaneous, intravenous, or inhaled treprostinil, but this use has not been studied.

IMPORTANT SAFETY INFORMATION for Orenitram

CONTRAINDICATIONS

• Orenitram is contraindicated in patients with severe hepatic impairment (Child Pugh Class C)

WARNINGS AND PRECAUTIONS

- Abrupt discontinuation or sudden large reductions in dosage of Orenitram may result in worsening of PAH symptoms
- Orenitram inhibits platelet aggregation and increases the risk of bleeding
- Orenitram should not be taken with alcohol as release of treprostinil from the tablet may occur at a faster rate than intended
- The Orenitram tablet shell does not dissolve. In patients with diverticulosis (blind-end pouches), Orenitram tablets can lodge in a diverticulum

DRUG INTERACTIONS/SPECIFIC POPULATIONS

- Concomitant administration of Orenitram with diuretics, antihypertensive agents, or other vasodilators increases the risk of symptomatic hypotension
- Orenitram inhibits platelet aggregation; there is an increased risk of bleeding, particularly among patients receiving anticoagulants
- Co-administration of Orenitram and the CYP2C8 enzyme inhibitor gemfibrozil increases exposure to treprostinil; therefore, Orenitram dosage reduction may be necessary in these patients
- Pregnancy Category C. Animal reproductive studies with Orenitram have shown an adverse effect on the fetus. There are no adequate and well-controlled studies in humans
- Safety and effectiveness in patients under 18 years of age have not been established
- There is a marked increase in the systemic exposure to treprostinil in hepatically impaired patients

ADVERSE REACTIONS

 In the 12-week placebo-controlled monotherapy study, adverse reactions with rates at least 5% higher on Orenitram than on placebo included headache, diarrhea, nausea, flushing, pain in jaw, pain in extremity, hypokalemia, and abdominal discomfort

Please see accompanying Full Prescribing Information and Patient Information for Orenitram.

For additional information about Orenitram, visit Orenitram.com or call 1-877-UNITHER (1-877-864-8437).