

**CENTER FOR DRUG EVALUATION AND  
RESEARCH**

*APPLICATION NUMBER:*

**208276Orig1s000**

**PRODUCT QUALITY REVIEW(S)**

Recommendation: **APPROVAL**

**NDA 208276  
Resubmission After 2017 Complete Response  
Review #1**

Drug Name/Dosage Form	Treprostinil Injection
Strength	10 mg/mL
Route of Administration	Intravenous
Rx/OTC Dispensed	Rx
Applicant	United Therapeutics Corporation
US agent, if applicable	n/a

SUBMISSION(S) REVIEWED	DOCUMENT DATE	DISCIPLINE(S) AFFECTED
<i>Amendment</i>	<i>19-JUN-2018</i>	<i>Quality Labeling</i>
<i>Resubmission</i>	<i>30-JAN-2018</i>	<i>All</i>

**Quality Review Team**

DISCIPLINE	REVIEWER	OPQ OFFICE
Drug Substance	Thomas Wong	ONDP
Drug Product		
Process		
Environmental Analysis		
Microbiology	John Metcalfe	OPF
Facility	Christina Capacci-Daniel	OPF
Regulatory Business Process Manager	Grafton Adams	OPRO
Application Technical Lead	Wendy Wilson-Lee	ONDP

## Quality Review Data Sheet

### 1. RELATED/SUPPORTING DOCUMENTS

#### A. DMFs:

None.

#### B. Other Documents: *IND, RLD, or sister applications*

DOCUMENT	APPLICATION NUMBER	DESCRIPTION
NDA	21272	Treprostinil Injection
PMA	(b) (4)	Implantable Pump Device

### 2. CONSULTS

No consults requested.

## Executive Summary

### I. Recommendations and Conclusion on Approvability

OPQ recommends **APPROVAL** of NDA 208276 for Remodulin (treprostinil) Injection, 10 mg/mL drug product with the implantable infusion pump.

### II. Summary of Quality Assessments

#### A. Product Overview

<b>Proposed Indication(s) including Intended Patient Population</b>	Treatment of pulmonary arterial hypertension
<b>Duration of Treatment</b>	16 weeks
<b>Maximum Daily Dose</b>	2.5 ng/kg/min per week
<b>Alternative Methods of Administration</b>	Initial administration will be via external infusion system until implantation.

NDA 208276 seeks approval of the use of Remodulin (treprostinil) Injection with a fully implanted, programmable infusion system for the chronic intravenous delivery of treprostinil. The NDA is a sister application to Medtronic, Inc. Premarket Approval Application (PMA) P140032 for the device components of the implantable system. The applicant references approved NDA 21272 for all relevant chemistry, manufacturing, and controls (CMC) information for the treprostinil drug substance and drug product. The chemical and physical compatibility of the drug product with the implantable system was reviewed under PMA P140032.

This NDA was initially submitted in February 2015. Due to Refuse to File status, the applicant resubmitted this NDA in December 2015. The December 2015 resubmission was recommended for Complete Response due to insufficient data to evaluate the risk of potential patient exposure to microbial contaminants when the treprostinil injection drug product is used in the proposed implantable pump system.

In December 2016, the applicant resubmitted this NDA, addressing all deficiencies. Changes provided in the December 2016 resubmission were in response to the Complete Response Letter deficiencies (October 2016) and the elimination of the 2.5 mg/mL and 5.0 mg/mL drug product strengths listed in the initial filing and first resubmission (December 2015). The applicant retained only the 10.0 mg/mL strength for proposed commercialization and use with the implantable pump in the December 2016 resubmission. There are no changes made to any of the CMC information filed in the referenced NDA 21272. An approval recommendation was made by OPQ at the end of the review cycle. However, due to deficiencies identified under the PMA and insufficient human factors data, a Complete Response letter was issued (June 2017).

The 2018 resubmission did not include any new CMC information. All referenced CMC information under NDA 21272 remained unchanged as well. **Therefore, OPO recommends approval of NDA 208276.**

### B. Quality Assessment Overview

Treprostinil is a tricyclic benzindene analog of prostacycline (PGI<sub>2</sub>). It is a white to cream-colored powder, and is insoluble in water. There are five chiral centers in treprostinil and the drug substance is a single enantiomer. **The retest date of the drug substance is** <sup>(b)</sup><sub>(4)</sub> **months** <sup>(b)</sup><sub>(4)</sub>

<sup>(b)</sup><sub>(4)</sub>

:

The drug product, Remodulin® (treprostinil injection), is a sterile, 10 mg/mL, ready to use, solution packaged in a 20-mL vial for intravenous injection administered by continuous infusion. The vial and carton labels for the 10 mg/mL strength include the following statement: “When used with the Implantable System for Remodulin, no dilution is required.” Other strengths of Remodulin are approved but will not be indicated for use with the implantable pump system. Since the drug in the newly proposed drug-device combination product is the identical drug as the approved Treprostinil injection in NDA 021272, a Biopharmaceutics review is not needed. **An expiration dating period of** <sup>(b)</sup><sub>(4)</sub> **months will be granted for the drug product when stored at 25°C. The categorical exclusion is granted based on compliance with both 21 CFR 25.31(a) and 25.15 (d).**

There appears to be no significant or outstanding risks to the manufacturing process or final product based on the individual and composite evaluation of the listed facility’s inspection results, inspectional history, and relevant experience. There is no change to the proposed facilities, responsibilities, or assessment outcomes in the resubmission. **All facilities are acceptable to support approval of NDA 208276.**

### C. Special Product Quality Labeling Recommendations (NDA only)

None.

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