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Approval Package for:

APPLICATION NUMBER:

22-387

Trade Name:	TYVASO INHALATION SOLUTION
Generic Name:	Treprostinil
Sponsor:	United Therapeutics Corporation
Approval Date:	July 30, 2009
Indications:	For the treatment of pulmonary arterial hypertension (WHO Group I) in patients with NYHA Class III

symptoms, to increase walk distance.

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APPLICATION NUMBER: 22-387

APPROVAL LETTER

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Food and Drug Administration Silver Spring MD 20993

NDA 22-387

NDA APPROVAL

United Therapeutics Corporation Attention: Mr. Dean Bunce P.O. Box 14186 55 TW Alexander Drive Research Triangle Park, NC 27709

Dear Mr. Bunce:

Please refer to your June 27, 2008 new drug application (NDA) submitted under section 505(b)(1) of the Federal Food, Drug, and Cosmetic Act for Tyvaso (treprostinil) Inhalation Solution.

We acknowledge receipt of your submissions dated July 3, August 14 and 26, September 26, October 1, 9, 24, 29, and 31, November 12 and 13, and December 22, 2008, and January 2, 22, and 29, February 9, 19, and 25, March 12, April 1 (two), 3 (two), 8, 16, 17, 24, and 29, May 7, June 5, 17, and 25 (two), July 2, 7, 9, 22, and 28, 2009.

This new drug application provides for the use of Tyvaso for the treatment of pulmonary arterial hypertension (WHO Group I) in patients with NYHA Class III symptoms, to increase walk distance.

We have completed our review of this application, as amended. It is approved, effective on the date of this letter, for use as recommended in the enclosed agreed-upon labeling text.

CONTENT OF LABELING

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As soon as possible, but no later than 14 days from the date of this letter, please submit the content of labeling [21 CFR 314.50(1)] in structured product labeling (SPL) format as described at http://www.fda.gov/ForIndustry/DataStandards/StructuredProductLabeling/default.htm that is identical to the enclosed agreed-upon labeling text. Upon receipt, we will transmit that version to the National Library of Medicine for public dissemination. For administrative purposes, please designate this submission, "SPL for approved NDA 22-387."

Submit final printed carton and container labels that are identical to the enclosed carton and immediate container labels as soon as they are available, but no more than 30 days after they are printed. Please submit these labels electronically according to the guidance for industry titled *Providing Regulatory Submissions in Electronic Format – Human Pharmaceutical Product Applications and Related Submissions Using the eCTD Specifications (October 2005).* Alternatively, you may submit 12 paper copies, with 6 of the copies individually mounted on heavy-weight paper or similar material. For administrative purposes, designate this submission "**Final Printed Carton and Container Labels for approved NDA 22-387**." Approval of this submission by FDA is not required before the labeling is used.

Marketing the product with FPL that is not identical to the approved labeling text may render the product misbranded and an unapproved new drug.

PROPRIETARY NAME

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The Division of Medication Error Prevention and Analysis (DMEPA) and the Division of Cardiovascular and Renal Products do not object to the use of the proprietary name, Tyvaso, for this product.

PEDIATRIC RESEARCH EQUITY ACT (PREA)

Under the Pediatric Research Equity Act (PREA) (21 U.S.C. 355c), all applications for new active ingredients, new indications, new dosage forms, new dosing regimens, or new routes of administration are required to contain an assessment of the safety and effectiveness of the product for the claimed indication(s) in pediatric patients unless this requirement is waived, deferred, or inapplicable.

Because this drug product for this indication has an orphan drug designation, you are exempt from this requirement.

POSTMARKETING REQUIREMENTS UNDER 505(0)

Section 505(o) of the Federal Food, Drug, and Cosmetic Act (FDCA) authorizes FDA to require holders of approved drug and biological product applications to conduct postmarketing studies and clinical trials for certain purposes, if FDA makes certain findings required by the statute (section 505(o)(3)(A)).

We have sufficient concerns that treprostinil by the inhaled route of administration may be irritating to the respiratory tract. This concern is based on:

- Adverse events observed during the clinical experience
- Observations of oropharyngeal and other respiratory tract lesions in rats and dogs during the preclinical assessments of inhaled treprostinil
- The finding that when administered by the subcutaneous route, treprostinil is extremely irritating at the site of administration.

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We have determined that an analysis of spontaneous postmarketing adverse events reported under subsection 505(k)(1) of the FDCA will not be sufficient to assess a signal of a serious risk of oropharyngeal or pulmonary toxicity in patients using Tyvaso (treprostinil).

Furthermore, the new pharmacovigilance system that FDA is required to establish under section 505(k)(3) of the FDCA has not yet been established and is not sufficient to assess this serious risk.

Therefore, based on appropriate scientific data, FDA has determined that you are required to conduct the following:

1. A long-term observational study in the US that will include another 1000 patient-years of follow-up in Tyvaso-treated patients, and 1000 patient-years of follow-up in matched controls receiving other treatments for pulmonary hypertension, to evaluate the potential association between Tyvaso (treprostinil) and oropharyngeal and pulmonary toxicity.

The timetable you submitted on July 28, 2009 states that you will conduct this study according to the following timetable:

Draft Protocol Submission:	September 15, 2009
Final Protocol Submission:	December 15, 2009
Interim Study Report #1:	December 31, 2011
Interim Study Report #2:	September 30, 2012
Study Completion Date:	June 30, 2013
Final Report Submission/Supplement:	December 15, 2013

Submit the protocol to your IND, with a cross-reference letter to NDA 22-387. Submit all final report(s) to NDA 22-387. Prominently identify the submission with the following wording in bold capital letters at the top of the first page of the submission, as appropriate:

- **REQUIRED POSTMARKETING PROTOCOL UNDER 505(0)**
- REQUIRED POSTMARKETING FINAL REPORT UNDER 505(0)
- REQUIRED POSTMARKETING CORRESPONDENCE UNDER 505(0)

Section 505(0)(3)(E)(ii) of the FDCA requires you to report periodically on the status of any study or clinical trial required under this section. This section also requires you to periodically report to FDA on the status of any study or clinical trial otherwise undertaken to investigate a safety issue. Section 506B of the FDCA, as well as 21 CFR 314.81(b)(2)(vii) requires you to report annually on the status of any postmarketing commitments or required studies or clinical trials.

FDA will consider the submission of your annual report under section 506B and 21 CFR 314.81(b)(2)(vii) to satisfy the periodic reporting requirement under section 505(o)(3)(E)(ii) provided that you include the elements listed in 505(o) and 21 CFR 314.81(b)(2)(vii). We remind you that to comply with 505(o), your annual report must also include a report on the status of any study or clinical trial otherwise undertaken to investigate a safety issue. Failure to

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