

Accenture LLP 1160 West Swedesford Road Berwyn, PA 19312 www.accenture.com

August 20, 2015

Food and Drug Administration Center for Drug Evaluation and Research Central Document Room Drug Master File Staff 5901-B Ammendale Road Beltsville, MD 20705-1266

Re: DMF #: 027320

Holder: McKesson Specialty Health (McKesson)

DMF Subject: Transmucosal Immediate Release Fentanyl (TIRF) Access Program

Re: REMS Shared Program

DMF Type: V

DMF Submission Information: DMF Annual Report

**REMS Submission Identifier: Not Applicable** 

eCTD Sequence Number: 0017

Dear Drug Master File Staff:

This Type V DMF contains the Risk Evaluation and Mitigation Strategy (REMS) for Transmucosal Immediate Release Fentanyl for the Shared System REMS program.

Reference is made to the above DMF, which was initially submitted on August 20, 2013. Enclosed, please find the DMF Annual Report (Reporting Period: August 21, 2014 – August 20, 2015). The Annual Report includes an administrative information page, a summary of the amendments that have been submitted during this reporting period, and a list of authorized persons to incorporate by reference.

McKesson states that information provided in this Master File is current and assures that the material furnished will meet the specifications described herein. McKesson also confirms that the Holder obligations are observed.

We request that all information in this file be treated as confidential commercial information to the Food and Drug Administration pursuant to 21 C.F.R. §20.61, and that no information from this file be provided to any unauthorized persons without written consent.

If you have any questions or concerns, please do not hesitate to contact Jann Kochel, U.S. Agent for McKesson, at 610-407-1738 or alternatively via email at jann.a.kochel@accenture.com.

Sincerely,

Jann A. Kochel, U.S. Agent

Accenture, LLP

Attachments: Table of Contents for the submission

**Electronic Submission Specifications** 

### ANNUAL REPORT Reporting Period: August 21, 2014 – August 20, 2015

Module Section	Description
1.2 Cover Letter	Cover Letter w/ Attachments
1.11.3 – Efficacy Information Amendment	Annual Report - Summary of Changes
1.4.3 – List of Authorized Persons to Incorporate by Reference	List of Authorized Persons

## **Electronic Submission Specifications**

This submission is compliant with FDA's Guideline for Industry: Providing Regulatory Submissions in Electronic Format - Human Pharmaceutical Product Applications and Related Submissions Using the eCTD Specifications (June 2008).

All files were checked and verified to be free of viruses prior to transmission through the electronic submission gateway. This eCTD has been generated by Accenture, LLP (formerly Octagon Research Solutions Inc.), who has filed an acceptable eCTD pilot with the Center (Pilot Number 900777).

Anti-Virus Program	Symantec Endpoint Protection Edition
<b>Program Version</b>	11.0.5002.333
Virus Definition Date	8/16/2015 rev. 21
Submission Size	Approx. 1 MB

The IT point of contact for this submission is:

Name	Matt Francis
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#### ADMINISTRATIVE INFORMATION

Holder's Name: McKesson Specialty Health

**Holder's Address:** 4343 N. Scottsdale Road

Suite 150

Scottsdale, AZ 85251

Holder's Contact Person: Laura Baloun

Contact's Address: 4343 N. Scottsdale Road

Suite 150

Scottsdale, AZ 85251

**Contact's Phone:** 480-663-4009

**Contact's Fax:** 480-663-4973

Contact's Email address: laura.baloun@mckesson.com

**Statement of Commitment:** Attached, please find a signed statement of commitment. The statement certifies that the DMF is current and that McKesson will comply with the statements made in it.

**Agent's Name:** Accenture, LLP

**Agent's Address:** 1160 West Swedesford Road

Berwyn, PA 19312

Agent's Contact Person: Jann A. Kochel

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**Contact's Phone:** 610-407-1738

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Contact's E-mail address: jann.a.kochel@accenture.com

# 3. List of Authorized Persons to Incorporate by Reference

(Annual Report - Reporting Period: August 21, 2014 – August 20, 2015)

The following is a complete list of persons authorized to incorporate information in the DMF by reference:

- Actavis Laboratories Inc. June 17, 2015
- BioDelivery Sciences International, Inc. March 20, 2015
- Cephalon, Inc. August 28, 2013
- Depomed, Inc. August 28, 2013
- Galena BioPharma, Inc. August 28, 2013
- Insys Therapeutics, Inc. August 28, 2013
- Mallinckrodt August 28, 2013
- Mylan August 28, 2013
- Par Pharmaceutical, Inc. August 28, 2013

Please note that during the reporting period, McKesson submitted letters of authorization for Actavis Laboratories Inc. and BioDelivery Sciences International, Inc., and withdrew authorization for Meda Pharmaceuticals, Inc.

## **ANNUAL REPORT - SUMMARY OF CHANGES**

Reporting Period: August 21, 2014 – August 20, 2015

Date / Sequence	Description
November 25, 2014 / 0012	REMS Modification 3 - Draft
December 10, 2014 / 0013	REMS Modification 3 - Final
December 26, 2014 / 0014	Assessment 4 at 3 Years
March 20, 2015 / 0015	Letter of Authorization for BioDelivery Sciences International, Inc.
	Letter of Withdrawal for Meda Pharmaceuticals, Inc.
June 17, 2015 / 0016	Letter of Authorization for Actavis Laboratories Inc.

For further details regarding the above REMS amendments, please see the REMS History (Sequence 0014).