



ANDA 077312

Barr Laboratories, Inc.
Attention: Nicholas Tantillo
Senior Director, Regulatory Affairs
400 Chestnut Ridge Road
Woodcliffe Lake, NJ 07677

Dear Sir:

This is in reference to your abbreviated new drug application (ANDA) dated October 1, 2004, submitted pursuant to section 505(j) of the Federal Food, Drug, and Cosmetic Act (the Act), for Oral Transmucosal Fentanyl Citrate, 200 mcg, 400 mcg, 600 mcg, 800 mcg, 1200 mcg and 1600 mcg.

Reference is also made to your amendments dated August 5, November 11, November 29, and December 16, 2005; March 17, July 13, July 19, October 13, and October 16, 2006; June 29, October 5, and October 24, 2007; January 4, June 23, June 27, July 15, and October 31, 2008; and February 19, March 30, May 11, June 18, and July 21, 2009.

We have completed the review of this ANDA, including the Risk Management Plan (RMP), as amended, and have concluded that adequate information has been presented to demonstrate that the drug is safe and effective for use as recommended in the submitted labeling.

The following elements of the Risk Management Plan have been identified as being essential to the safe and effective use of your drug product.

- Medication Guide
- Plan to Monitor
- Welcome Kit: Fanny Pack
Lock & Keys
Child Safety Lock
Child Resistant Temporary Storage Container
Home Warning Stickers
Daily Diary
Brightly Colored Warning Flyers

- The following components will not be approved as part of the Risk Management Plan. However, you may distribute them as promotional materials, to the extent that the content of the materials is in compliance with the Act and implementing regulations.
 - Refrigerator Magnets
 - Children's Booklet
 - Diary Marker
 - Patient Safety Video

Accordingly the ANDA is approved, effective on the date of this letter. The Division of Bioequivalence has determined your Oral Transmucosal Fentanyl Citrate, 200 mcg, 400 mcg, 600 mcg, 800 mcg, 1200 mcg and 1600 mcg to be bioequivalent and, therefore, therapeutically equivalent to the reference listed drug, Actiq, 200 mcg, 400 mcg, 600 mcg, 800 mcg, 1200 mcg and 1600 mcg, of Cephalon, Inc. Your dissolution testing should be incorporated into the stability and quality control program using the same method proposed in your application.

Under section 506A of the Act, certain changes in the conditions described in this ANDA require an approved supplemental application before the change may be made. Additionally, any proposed change in the RMP must be discussed with FDA prior to institution. FDA will determine whether the proposed change is subject to FDA approval before implementation.

In accordance with section 505-1(i) of the FDCA, an abbreviated new drug application (ANDA) is required to have a Risk Evaluation & Mitigation Strategy (REMS) if the applicable listed drug has an approved REMS. The reference listed drug, Actiq, in addition to other oral transmucosal drug products containing fentanyl, is being required to implement a REMS. Pursuant to section 505-1(i) of the FDCA, a drug that is the subject of an ANDA and the listed drug it references must use a single shared system for elements to assure safe use unless FDA waives that requirement. We suggest that you contact Cephalon to pursue a single shared system for the elements to assure safe use in your future REMS.

Postmarketing reporting requirements for this ANDA are set forth in 21 CFR 314.80-81 and 314.98. The Office of Generic Drugs should be advised of any change in the marketing status of this drug.

Promotional materials may be submitted to FDA for comment prior to publication or dissemination. Please note that these submissions are voluntary. If you desire comments on proposed launch promotional materials with respect to compliance with applicable regulatory requirements, we recommend you submit, in draft or mock-up form, two copies of both the promotional materials and package insert(s) directly to:

Food and Drug Administration
Center for Drug Evaluation and Research
Division of Drug Marketing, Advertising, and Communications
5901-B Ammendale Road
Beltsville, MD 20705

We call your attention to 21 CFR 314.81(b)(3) which requires that all promotional materials be submitted to the Division of Drug Marketing, Advertising, and Communications with a completed Form FDA 2253 at the time of their initial use.

Within 14 days of the date of this letter, submit updated content of labeling [21 CFR 314.50(1)] in structured product labeling (SPL) format, as described at <http://www.fda.gov/oc/datacouncil/spl.html>, that is identical in content to the approved labeling. Upon receipt and verification, we will transmit that version to the National Library of Medicine for public dissemination. For administrative purposes, please designate this submission as "**Miscellaneous Correspondence - SPL for Approved ANDA 077312**".

Sincerely yours,

{See appended electronic signature page}

Gary Buehler
Director
Office of Generic Drugs
Center for Drug Evaluation and Research

Attachment: Critical Elements of the Risk Management Plan (RMP)

Application
Type/Number

Submission
Type/Number

Submitter Name

Product Name

ANDA-77312

ORIG-1

BARR
LABORATORIES
INC

FENTANYL CITRATE

This is a representation of an electronic record that was signed electronically and this page is the manifestation of the electronic signature.

/s/

ROBERT L WEST

10/30/2009

Deputy Director, for Gary Buehler