

Food and Drug Administration Silver Spring MD 20993

BLA 125274/S-102

SUPPLEMENT APPROVAL FULFILLMENT OF POSTMARKETING COMMITMENT RELEASE FROM POSTMARKETING COMMITMENT

Ipsen Biopharm, Limited Attention: James Adams, MS Manager, Regulatory Affairs 106 Allen Road, 3rd Floor Basking Ridge, NJ 07920

Dear Mr. Adams:

Please refer to your Biologics License Application (BLA) dated August 28, 2014, received September 15, 2014, submitted under section 351(a) of the Public Health Service Act for Dysport (abobotulinumtoxinA) injection.

We also refer to our approval letter dated July 15, 2015, which contained the following error: agreed-upon labeling that accompanied the letter contained a copyright protected

This replacement approval letter and enclosed, agreed-upon labeling incorporate the correction of the error. The effective approval date will remain July 15, 2015, the date of the original approval letter.

We acknowledge receipt of your amendments listed below.

October 28, 2014	December 10, 2014	May 26, 2015	July 2, 2015
October 31, 2014	December 18, 2014	June 1, 2015	July 14, 2015
November 6, 2014	January 13, 2015	June 4, 2015	
November 6, 2014	February 12, 2015	June 23, 2015	
November 7, 2014	May 22, 2015	July 1, 2015	

This Prior Approval supplemental biologics application proposes the addition of the following indication: treatment of upper limb spasticity in adult patients, to decrease the severity of increased muscle tone in the elbow, wrist, and finger flexors.

APPROVAL & LABELING

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

WAIVER OF HIGHLIGHTS SECTION

Please note that we have previously granted a waiver of the requirements of 21 CFR 201.57(d)(8) regarding the length of Highlights of prescribing information.

CONTENT OF LABELING

As soon as possible, but no later than 14 days from the date of this letter, submit, via the FDA automated drug registration and listing system (eLIST), the content of labeling [21 CFR 601.14(b)] in structured product labeling (SPL) format, as described at http://www.fda.gov/ForIndustry/DataStandards/StructuredProductLabeling/default.htm, that is identical to the enclosed labeling (text for the package insert and Medication Guide) and include the labeling changes proposed in any pending "Changes Being Effected" (CBE) supplements. Information on submitting SPL files using eLIST may be found in the guidance for industry titled "SPL Standard for Content of Labeling Technical Qs and As" at http://www.fda.gov/downloads/Drugs/GuidanceComplianceRegulatoryInformation/Guidances/UCM072392.pdf.

The SPL will be accessible via publicly available labeling repositories.

Also within 14 days, amend all pending supplemental applications that include labeling changes for this BLA, including pending "Changes Being Effected" (CBE) supplements, for which FDA has not yet issued an action letter, with the content of labeling [21 CFR 601.12(f)] in MS Word format that includes the changes approved in this supplemental application.

CARTON AND IMMEDIATE CONTAINER LABELS

Submit final printed carton and container labels that are identical to the carton and immediate container labels submitted on July 14, 2015, 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 (June 2008)". 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 "**Product Correspondence – Final Printed Carton and Container Labels for approved BLA 125274/102**." Approval of this submission by FDA is not required before the labeling is used.

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

REQUIRED PEDIATRIC ASSESSMENTS

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.

We are waiving the pediatric study requirement for children less than 2 years of age because necessary studies are impossible or highly impracticable. This is because the diagnosis of cerebral palsy, the most frequent cause of spasticity in children, is not reliable until children are two years of age or older.

We are deferring submission of your pediatric studies for ages 2 to 17 years for this application because this product is ready for approval for use in adults and the pediatric studies have not been completed.

Your deferred pediatric studies required by section 505B(a) of the Federal Food, Drug, and Cosmetic Act are required postmarketing studies. The status of these postmarketing studies must be reported annually according to 21 CFR 601.28 and section 505B(a)(3)(B) of the Federal Food, Drug, and Cosmetic Act. These required studies are listed below:

A juvenile rat toxicology study is required to identify the unexpected serious risk of adverse effects on postnatal growth and development. The study should utilize animals of an age range and stage(s) of development that are comparable to the intended pediatric population; the duration of dosing should cover the intended length of treatment in the pediatric population. In addition to the usual toxicological parameters, this study should evaluate effects of Dysport (abobotulinumtoxinA) on growth, reproductive development, and neurological and neurobehavioral development.

Final Report Submission: 08/15

A randomized, double-blind, adequately controlled, multiple fixed doses, parallel group clinical study of Dysport (abobotulinumtoxinA) in botulinum toxin-naive children age 2-17 years with upper extremity spasticity. The minimum duration of the study is 12 weeks. The study should be submitted to the FDA for special protocol assessment. The timetable you submitted on June 4, 2015, states that you will conduct this study according to the following schedule:

Study Completion:05/18Final Report Submission:10/18

Reports of these required pediatric postmarketing studies must be submitted as a BLA or as a supplement to your approved BLA with the proposed labeling changes you believe are warranted based on the data derived from these studies. When submitting the reports, please clearly mark your submission "SUBMISSION OF REQUIRED PEDIATRIC ASSESSMENTS" in large font, bolded type at the beginning of the cover letter of the submission.

We note that PMR 2933-1 above, may be addressed by submission of the final report intended to address PMR 2564-1 (identified as postmarketing requirement #1) listed in the April 29, 2009 approval letter for BLA 125274 and sBLA 125274/S001. When submitting the final report please be sure to reference both PMRs 2933-1 and 2564-1 in the cover letter.

RELEASE OF POSTMARKETING COMMITMENT

We remind you of the following postmarketing commitment (PMC) listed in the April 29, 2009, approval letter for BLA 125274 and sBLA125274/S-001.

2564-7 (identified as PMC #6 in the letter of April 29, 2009):
A randomized, double-blind, adequately controlled, multiple fixed doses, parallel group clinical study of Dysport (abobotulinumtoxinA) in botulinum toxin-naïve children age 2-17 years with upper extremity spasticity. The minimum duration of the study is 12 weeks. The study should be submitted to the FDA for special protocol assessment.

We have determined that you are released from the above commitment because it is being replaced by PMR 2933-2 above.

FULFILLMENT OF POSTMARKETING COMMITMENT

We have received your submission dated June 13, 2014, received June 16, 2014, containing the final report for the following postmarketing commitment (PMC) listed in the April 29, 2009, approval letter for BLA 125274 and sBLA125274/S-001.

2564-9 (identified as PMC #8 in the letter of April 29, 2009):
A randomized, double-blind, adequately controlled, multiple fixed doses, parallel group clinical study of Dysport (abobotulinumtoxinA) in botulinum toxin-naïve adults with upper extremity spasticity. The minimum duration of the study is 12 weeks. The study should be submitted to the FDA for special protocol assessment.

We have reviewed your submission and conclude that the above commitment was fulfilled.

We remind you that there are postmarketing requirements and commitments listed in the April 29, 2009, approval letter that are still open.

PROMOTIONAL MATERIALS

You may request advisory comments on proposed introductory advertising and promotional labeling. To do so, submit, in triplicate, a cover letter requesting advisory comments, the proposed materials in draft or mock-up form with annotated references, and the package insert(s) to:

OPDP Regulatory Project Manager Food and Drug Administration Center for Drug Evaluation and Research Office of Prescription Drug Promotion 5901-B Ammendale Road Beltsville, MD 20705-1266

Alternatively, you may submit a request for advisory comments electronically in eCTD format. For more information about submitting promotional materials in eCTD format, see the draft Guidance for Industry (available at:

http://www.fda.gov/downloads/Drugs/GuidanceComplianceRegulatoryInformation/Guidances/U CM443702.pdf).

As required under 21 CFR 601.12(f)(4), you must submit final promotional materials, and the package insert(s), at the time of initial dissemination or publication, accompanied by a Form FDA 2253. Form FDA 2253 is available at

<u>http://www.fda.gov/downloads/AboutFDA/ReportsManualsForms/Forms/UCM083570.pdf</u>. Information and Instructions for completing the form can be found at

<u>http://www.fda.gov/downloads/AboutFDA/ReportsManualsForms/Forms/UCM375154.pdf</u>. For more information about submission of promotional materials to the Office of Prescription Drug Promotion (OPDP), see <u>http://www.fda.gov/AboutFDA/CentersOffices/CDER/ucm090142.htm</u>.

REPORTING REQUIREMENTS

We remind you that you must comply with reporting requirements for an approved BLA (in 21 CFR 600.80 and in 21 CFR 600.81).

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If you have any questions, contact Taura Holmes, RPh, Regulatory Project Manager, via email or telephone at <u>Taura.Holmes@fda.hhs.gov</u> or (301) 796-1932.

Sincerely,

{See appended electronic signature page}

Eric Bastings, MD Deputy Director Division of Neurology Products Office of Drug Evaluation I Center for Drug Evaluation and Research

ENCLOSURE: Content of Labeling

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

/s/

ERIC P BASTINGS 07/15/2015