SPL response, we estimate that each facility will average approximately 6 products. 

Similarly, we estimate that 6 outsourcing facilities will submit an initial report identifying all drugs repackaged in the facility in the past year. Taking into account that a particular product that is repackaged may come in different strengths and can be reported in a single SPL response, we assume that each facility will average 6 products. Our estimate is based on current product reporting data. We expect that each product report will consist of multiple SPL responses per facility and assume preparing and electronically submitting this information will take up to 2 hours for each initial SPL response.

We also estimate 3 registered outsourcing facilities will submit a report twice each year (June and December) that identifies all drugs repackaged at the facility in the previous 6 months. We also estimate that an outsourcing facility will prepare and submit 3 SPL responses and assume that preparing and submitting this information electronically could also take up to 2 hours per response. If a product was not repackaged during a particular reporting period, outsourcing facilities do not need to send an SPL response for that product during that reporting period. Our figures reflect what we believe to be the average burden among respondents. We expect to receive no waiver requests from the electronic submission process for initial product reports and semiannual reports.

**Biologics Guidance**

We estimate 15 outsourcing facilities annually who mix, dilute, or repackage biological products will each design, test, and produce 5 different labels, for a total of 75 labels that include the information set forth in section III.C “Licensed Allergic Extracts for Subcutaneous Immunotherapy” of the draft guidance (including directions for use), for a total of 1,500 disclosures. We assume the initial process of designing, testing, and producing labeling and attaching to each prescription set each label, package, and/or container will take approximately 30 minutes (0.5 hours), for a total of approximately 750 hours.

Finally, we estimate that annually five outsourcing facilities who repackage biological products and establish a BUD in accordance with Appendix A “Assigning a BUD for Repackaged Biological Products Based On Stability Testing” will maintain 150 records of the testing, as described in Appendix A of the guidance. We assume maintaining the records will take 5 minutes per record, for a total of 12.5 hours.

Our estimated burden for the information collection reflects program changes and adjustments. We are changing the scope of the information collection to include burden attendant to provisions found in the Agency guidance documents discussed in this notice and have adjusted our estimate to reflect a resulting increase of 955 hours and 1,873 responses annually.

Dated: August 5, 2021.

Lauren K. Roth, Acting Principal Associate Commissioner for Policy.

**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Food and Drug Administration**


**Bioequivalence Studies With Pharmacokinetic Endpoints for Drugs Submitted Under an Abbreviated New Drug Application; Draft Guidance for Industry; Availability**

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notice of availability.

**SUMMARY:** The Food and Drug Administration (FDA or Agency) is announcing the availability of a revised draft guidance for industry entitled “Bioequivalence Studies With Pharmacokinetic Endpoints for Drugs Submitted Under an ANDA.” This revised draft supersedes the draft guidance entitled “Bioequivalence Studies With Pharmacokinetic Endpoints for Drug Products Submitted Under an ANDA,” which was announced in the Federal Register on December 5, 2013. This revised draft guidance provides recommendations to applicants planning to include bioequivalence (BE) information in abbreviated new drug applications (ANDAs) and ANDA supplements. In addition, this guidance describes how to meet the BE requirements set forth in the Federal Food, Drug, and Cosmetic Act (FD&C Act) and FDA regulations.

**DATES:** Submit either electronic or written comments on the draft guidance by October 22, 2021 to ensure that the Agency considers your comment on this draft guidance before it begins work on the final version of the guidance.

**ADDRESSES:** You may submit comments on any guidance at any time as follows:

- Electronic Submissions
  - Submit electronic comments in the following way:
    - Federal eRulemaking Portal: https://www.regulations.gov. Follow the instructions for submitting comments. Comments submitted electronically, including attachments, to https://www.regulations.gov will be posted to the docket unchanged. Because your comment will be made public, you are solely responsible for ensuring that your comment does not include any confidential information that you or a third party may not wish to be posted, such as medical information, your or anyone else’s Social Security number, or confidential business information, such as a manufacturing process. Please note that if you include your name, contact information, or other information that identifies you in the body of your comments, that information will be posted on https://www.regulations.gov.

- Written/Paper Submissions
  - If you want to submit a comment with confidential information that you do not wish to be made available to the public, submit the comment as a written/paper submission and in the manner detailed (see “Written/Paper Submissions” and “Instructions”).

**Written/Paper Submissions**

Submit written/paper submissions as follows:

- Mail/Hand Delivery/Courier (for written/paper submissions): Dockets Management Staff (HFA–305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.
  - For written/paper comments submitted to the Dockets Management Staff, FDA will post your comment, as well as any attachments, except for information submitted, marked and identified, as confidential, if submitted as detailed in “Instructions.”

Instructions: All submissions received must include the Docket No. FDA–
2013–D–1464 for “Bioequivalence Studies With Pharmacokinetic Endpoints for Drugs Submitted Under an ANDA.” Received comments will be placed in the docket and, except for those submitted as “Confidential Submissions,” publicly viewable at https://www.regulations.gov or at the Dockets Management Staff between 9 a.m. and 4 p.m., Monday through Friday, 240–402–7500.

• Confidential Submissions—To submit a comment with confidential information that you do not wish to be made publicly available, submit your comments only as a written/paper submission. You should submit two copies total. One copy will include the information you claim to be confidential with a heading or cover note that states “THIS DOCUMENT CONTAINS CONFIDENTIAL INFORMATION.” The Agency will review this copy, including the claimed confidential information, in its consideration of comments. The second copy, which will have the claimed confidential information redacted/blacked out, will be available for public viewing and posted on https://www.regulations.gov. Submit both copies to the Dockets Management Staff. If you do not wish your name and contact information to be made publicly available, you can provide this information on the cover sheet and not in the body of your comments and you must identify this information as “confidential.” Any information marked as “confidential” will not be disclosed except in accordance with 21 CFR 10.20 and other applicable disclosure law. For more information about FDA’s posting of comments to public dockets, see 80 FR 56469, September 18, 2015, or access the information at: https://www.govinfo.gov/content/pkg/FR-2015-09-18/pdf/2015-23389.pdf.

Docket: For access to the docket to read background documents or the electronic and written/paper comments received, go to https://www.regulations.gov and insert the docket number, found in brackets in the heading of this document, into the “Search” box and follow the prompts and/or go to the Dockets Management Staff, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852, 240–402–7500.

You may submit comments on any guidance at any time (see 21 CFR 10.115(g)(5)).

Submit written requests for single copies of the draft guidance to the Division of Drug Information, Center for Drug Evaluation and Research, Food and Drug Administration, 10001 New Hampshire Ave., Bldg. 35, Rm. 5125, Silver Spring, MD 20993–0002. Send one self-addressed adhesive label to assist that office in processing your requests. See the SUPPLEMENTARY INFORMATION section for electronic access to the draft guidance document.

FOR FURTHER INFORMATION CONTACT:

David Coppersmith, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 75, Rm. 1673, Silver Spring, MD 20993–0002, 301–796–9193.

SUPPLEMENTARY INFORMATION:

I. Background

FDA is announcing the availability of a revised draft guidance for industry entitled “Bioequivalence Studies With Pharmacokinetic Endpoints for Drugs Submitted Under an ANDA.” The revised draft guidance supersedes the draft guidance “Bioequivalence Studies With Pharmacokinetic Endpoints for Drugs Submitted Under an ANDA,” which was announced in the Federal Register on December 5, 2013 (78 FR 73199). FDA received nine comments on the draft guidance, which were considered before publication of this revised draft guidance.

This revised draft guidance provides recommendations to applicants planning to include BE information in ANDAs and ANDA supplements. In addition, this guidance describes how to meet the BE requirements set forth in the FD&C Act and FDA regulations. This guidance is generally applicable to dosage forms intended for oral administration and to non-orally administered drug products in which reliance on systemic exposure measures is suitable for documenting BE (e.g., transdermal delivery systems and certain rectal and nasal drug products). This guidance will also be useful to applicants planning BE studies intended to be conducted during the post-approval period for changes to a drug product approved in an ANDA. FDA recommends that applicants consult this revised draft guidance, in conjunction with any relevant product-specific guidances for industry, when considering the appropriate BE study and/or other studies for a proposed drug product.

This revised draft guidance is being issued consistent with FDA’s good guidance practices regulation (21 CFR 10.115). The revised draft guidance, when finalized, will represent the current thinking of FDA on “Bioequivalence Studies With Pharmacokinetic Endpoints for Drugs Submitted Under an ANDA.” It does not establish any rights for any person and is not binding on FDA or the public. You can use an alternative approach if it satisfies the requirements of the applicable statutes and regulations.

II. Paperwork Reduction Act of 1995

FDA tentatively concludes that this draft guidance contains no collection of information. Therefore, clearance by the Office of Management and Budget under the Paperwork Reduction Act of 1995 is not required.

III. Electronic Access


Dated: August 18, 2021.

Lauren K. Roth,
Acting Principal Associate Commissioner for Policy.

[FR Doc. 2021–18073 Filed 8–20–21; 8:45 am]

BILLING CODE 4164–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2021–P–0292]

Determination That ORTHO-CEPT (Desogestrel-Ethinyl Estradiol) 21- and 28-Day Oral Tablets, 0.15 Milligram/0.03 Milligram, Were Not Withdrawn From Sale for Reasons of Safety or Effectiveness

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA, Agency, or we) has determined that ORTHO-CEPT (desogestrel-ethinyl estradiol) 21- and 28-day oral tablets, 0.15 milligram (mg)/0.03 mg, were not withdrawn from sale for reasons of safety or effectiveness. This determination means that FDA will continue to approve ANDAs that refer to this drug product, and it will allow FDA to continue to approve ANDAs that refer to the product as long as they meet relevant legal and regulatory requirements.

FOR FURTHER INFORMATION CONTACT:

Stacy Kane, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 51, Rm. 6236, Silver Spring, MD 20993–0002, 301–796–8363, Stacy.Kane@fda.hhs.gov.