### Exhibits

#### Exhibit 3—Estimated Total and Annualized Cost—Continued

<table>
<thead>
<tr>
<th>Cost component</th>
<th>Total cost</th>
<th>Annualized cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Data Processing and Analysis</td>
<td>31,220</td>
<td>10,406</td>
</tr>
<tr>
<td>Overhead</td>
<td>22,034</td>
<td>7,345</td>
</tr>
<tr>
<td>Total</td>
<td>178,137</td>
<td>59,379</td>
</tr>
</tbody>
</table>

### Request for Comments

In accordance with the Paperwork Reduction Act, comments on AHRQ’s information collection are requested with regard to any of the following: (a) Whether the proposed collection of information is necessary for the proper performance of AHRQ health care research and health care information dissemination functions, including whether the information will have practical utility; (b) the accuracy of AHRQ’s estimate of burden (including hours and costs) of the proposed collection(s) of information; (c) ways to enhance the quality, utility, and clarity of the information to be collected; and (d) ways to minimize the burden of the collection of information upon the respondents, including the use of automated collection techniques or other forms of information technology.

Comments submitted in response to this notice will be summarized and included in the Agency’s subsequent request for OMB approval of the proposed information collection. All comments will become a matter of public record.

Dated: September 26, 2011.

Carolyn M. Clancy,
Director.

[FR Doc. 2011–25693 Filed 10–4–11; 8:45 am]

BILLING CODE 4160–90–M

### Department of Health and Human Services

#### Food and Drug Administration

[Docket No. FDA–2011–N–0708]

**Agency Information Collection Activities; Proposed Collection; Comment Request; FDA Form 3728, Animal Generic Drug User Fee Act Cover Sheet**

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

**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA) is announcing an opportunity for public comment on the proposed collection of certain information by the Agency. Under the Paperwork Reduction Act of 1995 (the PRA), Federal Agencies are required to publish a notice in the Federal Register concerning each proposed collection of information, including each proposed extension of an existing collection of information and to allow 60 days for public comment in response to the notice. This notice solicits comments on the information collection burden of the Animal Generic Drug User Fee Cover Sheet Form FDA 3728 that further implements certain provisions of the Animal Generic Drug User Fee Act of 2008 (AGDUFA).

**DATES:** Submit either electronic or written comments on the collection of information by December 5, 2011.

**ADDRESSES:** Submit electronic comments on the collection of information to http://www.regulations.gov. Submit written comments on the collection of information to the Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852. All comments should be identified with the docket number found in brackets in the heading of this document.

For further information contact: Juanmanuel Vilela, Office of Information Management, Food and Drug Administration, 1350 Piccard Dr., PI50–400B, Rockville, MD 20850, 301–796–7651, Juanmanuel.vilela@fda.hhs.gov.

**Supplementary information:** Under the PRA (44 U.S.C. 3501–3520), Federal Agencies must obtain approval from the Office of Management and Budget (OMB) for each collection of information they conduct or sponsor. “Collection of information” is defined in 44 U.S.C. 3502(3) and 5 CFR 1320.3(c) and includes Agency requests or requirements that members of the public submit reports, keep records, or provide information to a third party. Section 3506(c)(2)(A) of the PRA (44 U.S.C. 3506(c)(2)(A)) requires Federal Agencies to provide a 60-day notice in the Federal Register concerning each proposed collection of information, including each proposed extension of an existing collection of information, before submitting the collection to OMB for approval. To comply with this requirement, FDA is publishing notice of the proposed collection of information set forth in this document.

With respect to the following collection of information, FDA invites comments on these topics: (1) Whether the proposed collection of information is necessary for the proper performance of FDA’s functions, including whether the information will have practical utility; (2) the accuracy of FDA’s estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques, when appropriate, and other forms of information technology.

FDA Form 3728, Animal Generic Drug User Fee Act Cover Sheet—21 U.S.C. 379j–21 (OMB Control Number 0910–0632)—Extension

Section 741 of the Federal Food, Drug, and Cosmetic Act (FD&C Act) (21 U.S.C. 379j–21) establishes three different kinds of user fees: (1) Fees for certain types of abbreviated applications for generic new animal drugs, (2) annual fees for certain generic new animal drug products, and (3) annual fees for certain sponsors of abbreviated applications for generic new animal drugs and/or investigational submissions for generic new animal drugs (21 U.S.C. 379j–21(a)). Because the submission of user fees concurrent with applications is required, the review of an application cannot begin until the fee is submitted. FDA Form 3728 is the AGDUFA Cover Sheet, which is designed to provide the minimum necessary information to determine whether a fee is required for review of an application, to determine the amount of the fee required, and to account for and track user fees. FDA estimates the burden of this collection of information as follows:
Respondents to this collection of information are generic animal drug applicants. Based on FDA’s data base system, there are an estimated 20 sponsors of new animal drugs potentially subject to AGDUFA.

Dated: September 30, 2011.

Leslie Kux,
Acting Assistant Commissioner for Policy.

FOR FURTHER INFORMATION CONTACT: Juanmanuel Vilela, Office of Information Management, Food and Drug Administration, 1350 Piccard Dr., PI50–400B, Rockville, MD 20850, 301–796–7651, juanmanuel.vilela@fda.hhs.gov.

SUPPLEMENTARY INFORMATION: In compliance with 44 U.S.C. 3507, FDA has submitted the following proposed collection of information to OMB for review and clearance.

Requirements for Submission of Bioequivalence Data—21 CFR Parts 314 and 320—(OMB Control Number 0910–0630)—Extension

In the Federal Register of January 16, 2009 (74 FR 2849), the Agency published a final rule revising FDA regulations to require applicants to submit data on all bioequivalence (BE) studies, including studies that do not meet passing BE criteria, which are performed on a drug product formulation submitted for approval under an abbreviated new drug application (ANDA), or in an amendment to an ANDA that contains BE studies. In the final rule, FDA amended §§314.94(a)(7)(ii), 314.96(a)(1), 314.97, and 320.21(b)(1), to require an ANDA applicant to submit information from all BE studies, both passing and nonpassing, conducted by the applicant on the same drug product formulation as that submitted for approval under an ANDA, amendment, or supplement.

In Table 1 of this document, FDA has estimated the reporting burden associated with each section of the rule. FDA believes that the majority of additional BE studies will be reported in ANDAs (submitted under § 314.94), rather than supplements (reported in §314.97), because it is unlikely than an ANDA holder will conduct BE studies with a drug after the drug has been approved. With respect to the reporting of additional BE studies in amendments (submitted under § 314.96), this should also account for a small number of reports, because most BE studies will be conducted on a drug prior to the submission of the ANDA, and will be reported in the ANDA itself.

FDA estimates it will require approximately 120 hours of staff time to prepare and submit each additional complete BE study report, and approximately 60 hours of staff time for each additional BE summary report. The Agency believes that a complete report will be required approximately 20 percent of the time, while a summary will suffice approximately 80 percent of the time. Based on a weighted-average calculation using the information presented above, the submission of each additional BE study is expected to take 72 hours of staff time [(120 × 0.2) + (60 × 0.8)].

In the Federal Register of June 10, 2011 (76 FR 34081), FDA published a 60-day notice requesting public comment on the proposed collection of information. No comments were received.

FDA estimates the burden of this collection of information as follows: