

to the Administration for Children and Families, Office of Administration, Office of Information Services, 370 L'Enfant Promenade, SW., Washington, DC 20447, *Attn:* ACF Reports Clearance Officer. *E-mail address:* [infocollection@acf.hhs.gov](mailto:infocollection@acf.hhs.gov). All requests should be identified by the title of the information collection.

The Department specifically requests comments on: (a) Whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have practical utility; (b) the accuracy of the agency's estimate of the burden of the proposed collection of information; (c) the quality, utility, and clarity of the information to be collected; and (d) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques or other forms of information technology. Consideration will be given to comments and suggestions submitted within 60 days of this publication.

Dated: June 8, 2009.  
**Janean Chambers,**  
*Reports Clearance Officer.*  
[FR Doc. E9-13684 Filed 6-10-09; 8:45 am]  
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**DEPARTMENT OF HEALTH AND HUMAN SERVICES**  
**Food and Drug Administration**  
[Docket No. FDA-2008-N-0572]  
**Agency Information Collection Activities; Submission for Office of Management and Budget Review; Comment Request; Implementation of the Animal Generic Drug User Fee Act of 2008; User Fee Cover Sheet Form FDA 3728**  
**AGENCY:** Food and Drug Administration, HHS.  
**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA) is announcing that a proposed collection of information has been submitted to the Office of Management and Budget (OMB) for review and clearance under the Paperwork Reduction Act of 1995 (the PRA).  
**DATES:** Fax written comments on the collection of information by July 13, 2009.  
**ADDRESSES:** To ensure that comments on the information collection are received, OMB recommends that written comments be faxed to the Office of Information and Regulatory Affairs, OMB, Attn: FDA Desk Officer, FAX: 202-395-6974, or e-mailed to [oira\\_submission@omb.eop.gov](mailto:oira_submission@omb.eop.gov). All comments should be identified with the OMB control number 0910-0632. Also include the FDA docket number found in brackets in the heading of this document.  
**FOR FURTHER INFORMATION CONTACT:** Denver Presley Jr., Office of Information Management (HFA-710), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-796-3793.

**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.  
**Implementation of the Animal Generic Drug User Fee Act of 2008; User Fee Cover Sheet Form FDA 3728—21 U.S.C. 379j-21 (OMB Control Number 0910-0632)—Extension**  
This collection of information is currently approved under the emergency processing provisions of the PRA for 180 days. FDA is now seeking a 3-year clearance.

Section 741 of the 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. Because the submission of user fees concurrently with applications is required, the review of an application cannot begin until the fee is submitted. Form FDA 3728, the Animal Generic Drug User Fee Cover Sheet, is designed to provide the minimum necessary information in order to do the following: (1) Determine whether a fee is required for review of an application, (2) determine the amount of fee required, and (3) account for and track user fees.  
In the **Federal Register** of March 11, 2009 (74 FR 10596), FDA published a 60-day notice requesting public comment on the information collection provisions. No comments were received.  
FDA estimates the burden of this collection of information as follows:

TABLE 1.—ESTIMATED ANNUAL REPORTING BURDEN<sup>1</sup>

21 U.S.C. 379j-21	No. of Respondents	Annual Frequency per Response	Total Annual Responses	Hours per Response	Total Hours
Form FDA 3728	20	2	40	.08	3.2

<sup>1</sup> There are no capital costs or operating and maintenance costs associated with this collection of information.

Respondents to this collection of information are generic animal drug applicants. Based on FDA's database system, there are an estimated 20 sponsors of new animal drugs potentially subject to the Animal Generic Drug User Fee Act of 2008.

Dated: June 3, 2009.  
**Jeffrey Shuren,**  
*Associate Commissioner for Policy and Planning.*  
[FR Doc. E9-13716 Filed 6-10-09; 8:45 am]  
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**DEPARTMENT OF HEALTH AND HUMAN SERVICES**  
**Administration for Children and Families**  
**Submission for OMB Review; Comment Request**  
*Title:* Performance Progress Report.  
*OMB No.:* 0970-0334.

**Description:** The Performance Progress Report (SF-PPR) is a set of uniform reporting formats used for standard reporting on performance under grants and cooperative agreements.

In addition to allowing for uniformity of information collection, these formats will support systematic electronic collection and submission of information. These formats will provide interim and final performance progress information as required by OMB Circulars A-102 and 2 CFR part 215.

The SF-PPR consists of a cover page and six optional formats. The cover page contains identifying data elements and a section for a performance narrative. Use of the cover page is required, and programs may require their respondents to submit only this page and/or attach a performance narrative. Alternatively, programs may opt to require the cover page and one or more of the six optional formats: Performance Measures, Program Indicators, Benchmark Evaluations, Table of Activity Results, Activity-Based Expenditures, and Program/Project Management

The SF-PPR has been successfully piloted at the Administration for Children and Families (ACF). All discretionary programs (starting with FY09 awards) are to submit the SF-PPR to the ACF Office of Grants Management. Program offices with expiring data collections are required to migrate to the SF-PPR format.

Additionally, a number of program offices have voluntarily migrated their collections to the SF-PPR format in anticipation of government-wide standardization. ACF, with its Online Data Collection tool (OLDC), has provided program offices with the capability to collect SF-PPR data electronically.

ACF and the Grants Center of Excellence (CoE) is sponsoring this collection on behalf of the Grants Policy Committee, other Federal grant-making agencies, and the CoE partners.

CoE Partners are defined as:

Corporation for National and Community Service,  
Denali Commission,  
Department of State,  
DHHS/Administration on Aging,

DHHS/Centers for Medicare Services,  
DHHS/Health Research and Services Administration,  
DHHS/Indian Health Services,  
DHHS/Office of Public Health Services,  
DOT/Federal Air Administration,  
DOT/Federal Highway Administration,  
DOT/Federal Motor Carrier Safety Administration,  
DOT/Federal Railroad Administration,  
DOT/Federal Transport Administration,  
DOT/Pipeline and Hazardous Materials Safety Administration,  
Environmental Protection Agency,  
Institute of Museum and Library Services,  
Social Security Administration,  
Department of Treasury,  
USDA/Food Safety and Inspection Service,  
Veterans Administration.

The revised burden estimates are based on grant projects and awards for ACF and its CoE partners for FY2008 as reported by internal ACF reporting systems and *USASpending.gov*.

**Respondents:** Federal government grantees

#### ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden hours
Performance Progress Report (SF-PPR) .....	131,281	1	0.42	55,138.02
Cover Page Continuation (SF-PPR-2) .....	86	1	0.33	28.38
Performance Measures (SF-PPR-A) .....	430	1	0.75	322.50
Program Indicators (SF-PPR-B) .....	8,961	1	3	26,883
Benchmark Evaluations (SF-PPR-C) .....	248	1	1.50	372
Table of Activity Results (SF-PPR-D) .....	4,238	1	0.75	3,178.50
Activity Based Expenditures (SF-PPR-E) .....	2,616	1	0.33	863.28
Program/Project Management (SF-PPR-F) .....	45	1	0.50	22.50

**Estimated Total Annual Burden Hours:** 86,808.18

#### **Additional Information:**

Copies of the proposed collection may be obtained by writing to the Administration for Children and Families, Office of Administration, Office of Information Services, 370 L'Enfant Promenade, SW., Washington, DC 20447, *Attn:* ACF Reports Clearance Officer. All requests should be identified by the title of the information collection. *E-mail address:* [infocollection@acf.hhs.gov](mailto:infocollection@acf.hhs.gov).

#### **OMB Comment:**

OMB is required to make a decision concerning the collection of information between 30 and 60 days after publication of this document in the **Federal Register**. Therefore, a comment is best assured of having its full effect if OMB receives it within 30 days of publication. Written comments and recommendations for the proposed

information collection should be sent directly to the following: Office of Management and Budget, Paperwork Reduction Project, *Fax:* 202-395-7245, *Attn:* Desk Officer for the Administration for Children and Families.

Dated: June 8, 2009.

**Janean Chambers,**

*Reports Clearance Officer.*

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#### **DEPARTMENT OF HEALTH AND HUMAN SERVICES**

#### **Food and Drug Administration**

[Docket No. FDA-2009-D-0260]

#### **Draft Guidance for Industry: Questions and Answers Regarding the Reportable Food Registry as Established by the Food and Drug Administration Amendments Act of 2007; Availability; Announcement of Further Delay in Implementation of the Food and Drug Administration Amendments Act of 2007**

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

**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA) is announcing the availability of a draft guidance entitled "Guidance for Industry: Questions and