

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Submission for OMB Review; Comment Request

Title: State Plan for Child Support under Title IV–D of the Social Security Act (OCSE–100 and OCSE–21–U4).

OMB No.: 0970–0017.

Description: The State plan preprint pages and amendments serve as a contract between the Office of Child Support Enforcement and State and Territory IV–D agencies. These State plan preprint pages and amendments outline the activities States and Territories will perform as required by law, in Section 454 of the Social Security Act, in order for States and

Territories to receive Federal funds to meet the costs of child support enforcement.

Respondents: State and Territory IV–D Agencies.

ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden hours
State Plan (OCSE–100)	54	8	0.5	216
OCSE–21–U4	54	8	0.25	108

Estimated Total Annual Burden Hours: 324.

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.

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: 202–395–6974, Attn: Desk Officer for the

Administration for Children and Families.

Dated: April 28, 2008.
Robert Sargis,
Reports Clearance Officer.
 [FR Doc. E8–9751 Filed 5–2–08; 8:45 am]
BILLING CODE 4184–01–M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Submission for OMB Review; Comment Request

Title: Reunification Procedures for Unaccompanied Alien Children.
OMB No.: 0970–0278.

Description: Following the passage of the 2002 Homeland Security Act (Pub. L. 107–296), the Administration for Children and Families (ACF), Office of Refugee Resettlement (ORR), is charged with the care and placement of

unaccompanied alien children in Federal custody, and implementing a policy for the release of these children, when appropriate, upon the request of suitable sponsors while awaiting immigration proceedings. In order for ORR to make determinations regarding the release of these children, the potential sponsors must meet certain conditions pursuant to section 462 of the Homeland Security Act and the *Flores v. Reno Settlement Agreement* No. CV85 4544–RJK (C.D. Cal. 1997).

The proposed information collection requests information to be utilized by ORR for determining the suitability of a sponsor/respondent for the release of a minor from ORR custody. The proposed instruments are the Sponsor’s Agreement to Conditions of Release, Verification of Release, Family Reunification Packet, and the Authorization for Release of Information.

Respondents: Sponsors requesting release of unaccompanied alien children to their custody.

ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden hours
Sponsor’s Agreement to Conditions of Release	4,288	2	.0835	716
Verification of Release	4,288	1	.167	716
Family Reunification Packet	4,288	18	.0416	3,211
Authorization for Release of Information	4,288	15	.0222	1,428

Estimated Total Annual Burden Hours: 6,071.

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.

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: 202-395-6974, *Attn:* Desk Officer for the Administration for Children and Families.

Dated: April 28, 2008.

Janean Chambers,

Reports Clearance Officer.

[FR Doc. E8-9762 Filed 5-2-08; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2008-N-0257]

Draft Prescription Drug User Fee Act IV Drug Safety Five-Year Plan; Availability for Comment

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing the availability for public comment of the draft drug safety 5-year plan entitled "Prescription Drug User Fee Act (PDUFA) IV Drug Safety Five-Year Plan." This plan is intended to communicate FDA's strategy for meeting the commitments for enhancing and modernizing the drug safety system within the context of the PDUFA IV program.

DATES: Submit written or electronic comments on the draft drug safety 5-year plan by June 19, 2008.

ADDRESSES: Submit written requests for single copies of the draft plan to the Office of Executive Programs, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 51, rm. 6100, Silver Spring, MD 20993-0002. Send one self-addressed adhesive label to assist that office in processing your requests. Submit written comments on the draft drug safety 5-year plan to the Division of Dockets Management (HFA-305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. Submit electronic comments to <http://www.regulations.gov>. See the **SUPPLEMENTARY INFORMATION** section for electronic access to the document.

FOR FURTHER INFORMATION CONTACT: Jayne C. Ware, Food and Drug

Administration, Center for Drug Evaluation and Research, Office of Executive Programs, 10903 New Hampshire Ave., Bldg. 51, rm. 6100, Silver Spring, MD 20993-0002, 301-796-3200.

SUPPLEMENTARY INFORMATION:

I. Background

On September 27, 2007, President Bush signed into law the Food and Drug Administration Amendments Act of 2007, which includes the reauthorization and expansion of PDUFA. The reauthorization of PDUFA will significantly broaden and modernize the agency's drug safety program and facilitate more efficient development of safe and effective new medications for the American public. During the user fee negotiation process leading up to the renewal of PDUFA, FDA and the relevant regulated industries mutually agreed to certain commitments that the FDA will carry out during fiscal years 2008 through 2012. Congress signaled its agreement with the commitments by authorizing PDUFA funds for them. Among those commitments is the responsibility of the FDA to develop and periodically update a 5-year plan describing activities that will lead to enhancing and modernizing FDA's drug safety system.

FDA is announcing for public comment the availability of the draft drug safety 5-year plan entitled "Prescription Drug User Fee Act (PDUFA) IV Drug Safety Five-Year Plan." This plan is intended to communicate FDA's strategy for meeting the commitments for enhancing and modernizing the drug safety system within the context of the PDUFA IV program. The plan describes the agency's strategy for achieving the commitments defined in section VIII, Enhancement and Modernization of the FDA Drug Safety System, and section IX, Review of Proprietary Names to Reduce Medication Errors, of the PDUFA IV Performance Goals (<http://www.fda.gov/oc/pdufa4/pdufa4goals.html>). At the end of the comment period, FDA will review the comments, update the "PDUFA IV Drug Safety Five-Year Plan," and publish the final version.

II. Comments

Interested persons may submit to the Division of Dockets Management (see **ADDRESSES**) written or electronic comments regarding this document. Submit a single copy of electronic comments or two paper copies of any mailed comments, except that individuals may submit one paper copy. Comments are to be identified with the

docket number found in brackets in the heading of this document. Received comments may be seen in the Division of Dockets Management between 9 a.m. and 4 p.m., Monday through Friday.

Please note that on January 15, 2008, the FDA Division of Dockets Management Web site transitioned to the Federal Dockets Management System (FDMS). FDMS is a Government-wide, electronic docket management system. Electronic comments or submissions will be accepted by FDA only through FDMS at <http://www.regulations.gov>.

III. Electronic Access

Persons with access to the Internet may obtain the document at http://www.fda.gov/cder/pdufa/PDUFA_IV_5yr_plan_draft.pdf.

Dated: April 25, 2008.

Jeffrey Shuren,

Associate Commissioner for Policy and Planning.

[FR Doc. E8-9726 Filed 5-2-08; 8:45 am]

BILLING CODE 4160-01-S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

The 11th Annual Food and Drug Administration-Orange County Regulatory Affairs Educational Conference

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice of meeting.

The Food and Drug Administration (FDA) is announcing the following conference: 11th Annual Educational Conference co-sponsored with the Orange County Regulatory Affairs Discussion Group (OCRA). The conference is intended to provide the Drug, Device, and Biologics industries with an opportunity to interact with FDA reviewers and compliance officers from the Centers and District Offices, as well as other industry experts. The main focus of this interactive conference will be product approval, compliance, and risk management in the three medical product areas. Industry speakers, interactive questions and answers, and workshop sessions will also be included to assure open exchange and dialogue on the relevant regulatory issues.

Date and Time: The conference will be held on June 11 and 12, 2008, from 7:30 a.m. to 5 p.m.

Location: The conference will be held at the Irvine Marriott Hotel, 18000 Von Karman Ave., Irvine, CA 92612.