

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Administration for Children and Families

[Program Announcement No. ACF/ACYF/CB-98-04]

#### Fiscal Year (FY) 1998 Notice of an Announcement of the Availability of Financial Assistance and Request for Applications To Support Demonstration Projects Under the Adoption Opportunities Program

**AGENCY:** Administration on Children, Youth and Families, ACYF, ACF, DHHS.

**ACTION:** Notice of Fiscal Year (FY) 1998 availability of financial assistance and request for applications to support demonstration projects under the Adoption Opportunities Program, Title II of the Child Abuse Prevention and Treatment Act, as amended, Pub. L. 104-235.

**SUMMARY:** The Children's Bureau, within the Administration on Children, Youth and Families announces the availability of FY 1998 funds for competing new discretionary grants under the Adoption Opportunities Program. Adoption Opportunities Program funds are designed to provide services that facilitate the elimination of barriers to adoption and to provide permanent loving homes for children who would benefit from adoption, particularly children with special needs. Specific priority areas for which grant awards are being solicited include:

- 98.1—National Resource Center on Special Needs Adoption
- 98.2—Administration of the Interstate Compact on Adoption and Medical Assistance
- 98.3—Achieving Increased Adoptive Placements of Children in Foster Care
- 98.4—Effective Collaborations for Timely Adoptions
- 98.5—Overcoming State and Local Barriers to Adoption
- 98.6—Adoption 2002 Support Project
- 98.7—Post-Legal Adoption Services

**DATES:** The date and time deadline for RECEIPT of applications by DHHS for new grants under this announcement 4:30 p.m. (Eastern Time Zone) on July 24, 1998.

**FOR FURTHER INFORMATION CONTACT:** Copies of the program announcement will be automatically sent to all current Adoption Opportunities Program grantees, all organizations that applied for grant awards in FY 97 and all individuals and organizations that have asked to be placed on the mailing list for

FY 1998. Copies of the program announcement can be obtained the ACYF Operations Center at 1-800-351-2293. A copy of this program announcement is also located at the CB website at <http://www.acf.dhhs.gov/programs/CB> under Policy and Funding Announcements.

**SUPPLEMENTARY INFORMATION:** Grant awards of FY 1998 funds will be made by September 30, 1998. The estimated funds available for new awards is \$4.9 million and the approximate number of new grants is estimated at 28.

(*Catalog of Federal Domestic Assistance*. Number 93.652, Adoption Opportunities Grants)

Dated: June 3, 1998.

**James A. Harrell,**

*Deputy Commissioner, Administration on Children, Youth and Families.*

[FR Doc. 98-15284 Filed 6-8-98; 8:45 am]

BILLING CODE 4184-01-M

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Food and Drug Administration

[Docket No. 98N-0373]

#### Agency Information Collection Activities: Proposed Collection; Comment Request

**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 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 FDA Recall Regulations under 21 CFR part 7. Recall guidelines set forth procedures to be used by manufacturers and distributors or other responsible persons in notifying or alerting health professionals or other persons of an unreasonable risk of substantial harm to the public's health and describe the procedures used or required by FDA in the recall process.

**DATES:** Submit written comments on the collection of information by August 10, 1998.

**ADDRESSES:** Submit written comments on the collection of information to the

Dockets Management Branch (HFA-305), Food and Drug Administration, 12420 Parklawn Dr., rm. 1-23, Rockville, MD 20857. All comments should be identified with the docket number found in brackets in the heading of this document.

**FOR FURTHER INFORMATION CONTACT:** JonnaLynn P. Capezzuto, Office of Information Resources Management (HFA-250), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-827-4659.

**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 listed below.

With respect to the following collection of information, FDA invites comments on: (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 Recall Regulations—Part 7 (21 CFR Part 7), Subpart C—(OMB Control Number 0910-0249—Extension)

These regulations were established to provide guidance to manufacturers on recall responsibilities. These responsibilities include development of a recall strategy; providing complete details of the recall reason, risk evaluation, quantity produced, distribution information, firm's recall