are represented in the ACF–801. Disaggregate data is used to determine program and participant characteristics as well as costs and levels of child care services provided. This provides ACF with the information necessary to make reports to Congress, address national

child care needs, offer technical assistance to grantees, meet performance measures, and conduct research. Consistent with the statute and regulations, ACF requests extension of the ACF–801.

Respondents: States, the District of Columbia, and Territories including Puerto Rico, Guam, the Virgin Islands, American Samoa, and the Northern Marianna Islands.

#### ANNUAL BURDEN ESTIMATES

Instrument	Number of re- spondents	Number of re- sponses per respondent	Average bur- den hours per response	Total burden hours
ACF-801 Estimated Total Annual Burden Hours	56	D4	20	4,480 4,480

In compliance with the requirements of Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995, the Administration for Children and Families is soliciting public comment on the specific aspects of the information collection described above. Copies of the proposed collection of information can be obtained and comments may be forwarded 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.

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: September 26, 2002.

#### Robert Sargis,

Reports Clearance Officer.
[FR Doc. 02–25003 Filed 10–1–02; 8:45 am]
BILLING CODE 4184–01–M

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Administration for Children and Families

# Proposed Information Collection Activity; Comment Request

## **Proposed Projects**

*Title:* Online Interstate Referral Guide (IRG).

OMB No.: 0970-0209.

Description: The IRG is an essential reference maintained by OCSE that provides States with an effective and efficient way of viewing and updating State profile, address, and FIPS code information by consolidating data available through numerous discrete sources into a single centralized, automated repository.

Respondents: State IV–D Child Support Programs.

## ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents	Number of responses per respondents	Average bur- den hours per response	Total burden hours
Online IRG Estimated Total Annual Burden Hours	54	18	.3	292 292

In compliance with the requirements of Section 3056(c)(2)(A) of the Paperwork Reduction Act of 1995, the Administration for Children and Families is soliciting public comment on the specific aspects of the information collection described above. Copies of the proposed collection of information can be obtained and comments may be forwarded 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: AFC Reports Clearance

Officer. 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: September 26, 2002.

#### Robert Sargis,

 $Reports\ Clearance\ Officer.$ 

[FR Doc. 02–25004 Filed 10–1–02; 8:45 am]

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# DEPARTMENT OF HEALTH AND HUMAN SERVICES

## Food and Drug Administration

[Docket No. 02N-0063]

Agency Information Collection Activities; Proposed Collection; Comment Request: Consumer Surveys on Food and Dietary Supplement Labeling Issues

**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 surveys to study consumers' understanding of specific label statements for conventional foods and dietary supplements and the impact of such labeling on consumer practices, knowledge levels, and attitudes. In the Federal Register of July 8, 2002 (67 FR 45128), FDA published a notice announcing OMB's approval of this collection of information (OMB control number 0910-0492). Because this was an emergency approval that will expire on December 31, 2002, FDA in this notice is following the normal PRA clearance procedures by issuing this notice.

**DATES:** Submit written or electronic comments on the collection of information by December 2, 2002.

ADDRESSES: Submit electronic comments on the collection of information to http://
www.accessdata.fda.gov/scripts/oc/
dockets/edockethome.cfm. Submit
written comments on the collection of information to the Dockets Management
Branch (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:

Peggy Schlosburg, Office of Information Resources Management (HFA–250), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301–827–1223.

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: (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

## Consumer Surveys on Food and Dietary Supplement Labeling Issues (OMB Control Number 910–0492)—Extension

FDA is requesting an extension of the OMB approval of consumer surveys to help FDA's Center for Food Safety and Applied Nutrition formulate decisions and policies affecting the labeling of conventional foods and dietary supplements. Determining how consumers are likely to interpret various kinds of claims, disclaimers, warnings, caution statements, and notice statements that might appear in labeling is critical to agency decisionmaking under the Federal Food, Drug, and Cosmetic Act and the first amendment. It is often necessary to test actual or proposed labeling statements in realistic situations with typical consumers to determine what these label statements are communicating to consumers.

FDA or its contractor will collect and use information gathered from telephone, mail, shopping mall intercept or Internet surveys to evaluate how consumers understand and respond to existing label statements, label statements proposed by industry or consumers, and other label statements that are under consideration as part of FDA's policy development process. Potential respondents to the surveys will be individual consumers either randomly chosen to represent specified populations or randomly assigned to experimental treatment conditions to control for the effects of individual differences in the population on the interpretation of label statements. In all instances, FDA will strive to collect a representative sample of individuals from the overall population or from relevant population groups, as appropriate. FDA's general selection method will use stratification, with random sampling within the strata, to achieve representativeness for both overall populations and sensitive subpopulations, such as at-risk individuals or user segments. In the rare cases where geography is a limiting factor, FDA will use population-based cluster sampling to limit Government expense while preserving the statistical properties of the sample.

Respondents will provide background information and respond to package labels that contain the variations of label statements to be tested. Measures will include both self-reported comprehension and acceptance as well as direct behavioral measures of consumer use and understanding of the package labeling.

FDA will use the information from the surveys in evaluating regulatory and policy options with respect to labeling. The agency often lacks empirical data about how consumers understand and respond to statements they might see in product labeling. The information gathered from such surveys can be used to test consumer comprehension and behavioral impact of various label statements and formats, taking into account the existing distribution of behavior, knowledge and attitudes in the population that provides the context for understanding such statements. The surveys will help FDA assess consumer reactions to existing and proposed label statements.

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