Agenda items are subject to change as priorities dictate.

Contact Person for More Information: Price Connor, PhD, NIOSH Health Scientist, 1600 Clifton Road, NE., Mailstop E–20, Atlanta, Georgia 30333, Telephone (404) 498–2511, Fax (404) 498–2571.

The Director, Management Analysis and Services Office, has been delegated the authority to sign **Federal Register** notices pertaining to announcements of meetings and other committee management activities for both the Centers for Disease Control and Prevention, and the Agency for Toxic Substances and Disease Registry.

Dated: August 12, 2011.

Elizabeth Millington,

Acting Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

[FR Doc. 2011-21410 Filed 8-19-11; 8:45 am]

BILLING CODE 4163-18-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Proposed Information Collection Activity; Comment Request

Proposed Projects

Title: Annual Report/ACF 204 (State MOE)—1 collection.

OMB No.: 0970–0248.

Description

The Administration for Children and Families (ACF) is requesting a three-year extension of the ACF–204 (Annual MOE Report). The report is used to collect descriptive program characteristics information on the programs operated by States and Territories in association with their Temporary Assistance for Needy Families (TANF) programs. All State and Territory expenditures claimed toward States and Territories MOE

requirements must be appropriate, i.e., meet all applicable MOE requirements. The Annual MOE Report provides the ability to learn about and to monitor the nature of State and Territory expenditures used to meet States and Territories MOE requirements, and it is an important source of information about the different ways that States and Territories are using their resources to help families attain and maintain selfsufficiency. In addition, the report is used to obtain State and Territory program characteristics for ACFs annual report to Congress, and the report serves as a useful resource to use in Congressional hearings about how TANF programs are evolving, in assessing State the Territory MOE expenditures, and in assessing the need for legislative changes.

Respondents

The 50 States of the United States, the District of Columbia, Guam, Puerto Rico, and the Virgin Islands.

ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden hours
ACF-204	54	1	128	6,912

Estimated Total Annual Burden Hours: 6.912.

In compliance with the requirements of Section 506(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 Planning, Research and Evaluation, 370 L'Enfant Promenade, SW., Washington, DC 20447, Attn: ACF Reports Clearance Officer. E-mail address: 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 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.

Robert Sargis,

Reports Clearance Officer.
[FR Doc. 2011–21317 Filed 8–19–11; 8:45 am]
BILLING CODE 4184–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Submission for OMB Review; Comment Request

Title: State Plan for the Temporary Assistance for Needy Families (TANF). *OMB No.:* 0970–0145.

Description

The State plan is a mandatory statement submitted to the Secretary of the Department of Health and Human Services by the State. It consists of an outline specifying how the state's TANF program will be administered and operated and certain required certifications by the State's Chief Executive Officer. It is used to provide the public with information about the program.

Authority to require States to submit a State TANF plan is contained in section 402 of the Social Security Act, as amended by Public Law 104–193, the Personal Responsibility and Work Opportunity Reconciliation Act of 1996. States are required to submit new plans periodically (*i.e.*, within a 27-month period).

We are proposing to continue the information collection without change.

Respondents

The 50 States of the United States, the District of Columbia, Guam, Puerto Rico, and the Virgin Islands.

ANNUAL BURDEN ESTIMATES

Instrument	Number of respondents	Number of responses per respondent	Average burden hours per response	Total burden
Title Amendments	18	1	3	54
	18	1	30	540

Estimated Total Annual Burden Hours: 594.

Additional Information

Copies of the proposed collection may be obtained by writing to the Administration for Children and Families, Office of Planning, Research and Evaluation, 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– 395–7285. E-mail: OIRA_SUBMISSION@OMB.EOP.GOV. Attn: Desk Officer for the Administration for Children and Families.

Robert Sargis,

Reports Clearance Officer. [FR Doc. 2011–21316 Filed 8–19–11; 8:45 am] BILLING CODE 4184–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2011-N-0591]

Agency Information Collection Activities; Submission for Office of Management and Budget Review; Comment Request; Data To Support Communications Usability Testing, as Used by the Food and Drug Administration

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. DATES: Fax written comments on the collection of information by September 21, 2011.

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: 202-395-7285, or e-mailed to oira submission@omb.eop.gov. All comments should be identified with the OMB control number 0910-New and title: "Data to Support Communications Usability Testing, as Used by the Food and Drug Administration." Also include the FDA 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: In compliance with 44 U.S.C. 3507, FDA has submitted the following proposed collection of information to OMB for review and clearance.

Data To Support Communications Usability Testing, as Used by the Food and Drug Administration—(OMB Control Number 0910–New)

FDA plans to use the data collected under this generic clearance to inform its communications campaigns on a variety of topics related to products that FDA regulates. FDA expects the data to help staff message developers achieve FDA communication objectives. FDA also plans to use the data to help tailor print, broadcast, and electronic media communications in order for them to have powerful and desired impacts on

target audiences. The data will not be used for the purposes of making policy or regulatory decisions.

The information collected will serve two major purposes. First, as formative research it will provide the critical knowledge needed about target audiences. FDA must explore audiences' beliefs, perceptions, and decisionmaking processes on specific topics in order to meet the basic objectives of its risk communication campaigns. Such knowledge will provide the needed target audience understanding to design effective communication strategies, messages, and product labels. These communications will aim to improve public understanding of the risks and benefits of using various FDA-regulated products by providing users with a better context in which to place risk information more completely.

Second, as pretesting, it will give FDA some information about the potential effectiveness of messages and materials in reaching and successfully communicating with their intended audiences. Testing messages with a sample of the target audience will allow FDA to refine messages while still in the developmental stage. Respondents may be asked to give their reaction to the messages in person or on-line.

FDA's Centers and Offices will use this mechanism to test the usability of messages about FDA-regulated products for consumers, patients, industry representatives, or health care professionals. The data will not be used for the purposes of making policy or regulatory decisions.

In the **Federal Register** of June 10, 2011 (76 FR 34083), 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: