DEPARTMENT OF HEALTH AND HUMAN SERVICES
Administration for Children and Families
Submission for OMB Review; Comment Request

Title: Pre-testing of Evaluation Data Collection Activities.
OMB No.: 0970–0355.
Description: The Office of Planning, Research, and Evaluation (OPRE), in the Administration for Children and Families (ACF) at the U.S. Department of Health and Human Services (HHS) intends to request approval from the Office of Management and Budget (OMB) to renew a generic clearance to pre-test data collection instruments with more than nine participants to identify and resolve any question or procedural problems in survey administration.
OPRE studies ACF programs, and the populations they serve, through rigorous research and evaluation projects. These include evaluations of existing programs, evaluations of innovative approaches to helping low-income children and families, researchsyntheses and descriptive and exploratory studies. To improve the development of its research and evaluation surveys, OPRE uses the pre-testing of evaluation surveys generic clearance to employ a variety of techniques including cognitive and usability laboratory and field techniques, behavior coding, exploratory interviews, respondent debriefing questionnaires, split sample experiments, focus groups, and pilot studies/pre-tests. These activities allow OPRE to identify if and when a survey may be simplified for respondents, respondent burden may be reduced, and other possible improvements.
Following standard OMB requirements, OPRE will submit a change request for each individual data collection activity under this generic clearance. Each request will include the individual instrument(s), a justification specific to the individual information collection, and any supplementary documents. OMB should review within 10 days of receiving each change request.
The information collected in this effort will not be the primary subject of any published ACF reports; however, information may be made public through methodological appendices or footnotes, reports on instrument development, instrument user guides, descriptions of respondent behavior, and other publications or presentations describing findings of methodological interest. When necessary, results will be labeled as exploratory in nature. The results of this pre-testing research may be prepared for presentation at professional meetings or publication in professional journals.
Respondents: Participants in ACF programs being evaluated; participants in ACF demonstrations; recipients of ACF Grants and individuals served by ACF Grantees; comparison group members; and other relevant populations, such as individuals at risk of needing ACF services.

ANNUAL BURDEN ESTIMATES

<table>
<thead>
<tr>
<th>Instrument</th>
<th>Total number of respondents</th>
<th>Number of responses per respondent</th>
<th>Average burden hours per response</th>
<th>Total burden hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>Survey development field tests, respondent debriefing questionnaires, cognitive interviews, split sample experiments, focus groups</td>
<td>3,825</td>
<td>1</td>
<td>1</td>
<td>3,825</td>
</tr>
</tbody>
</table>

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, 330 C Street SW, Washington, DC 20201, Attn: OPRE Reports Clearance Officer. All requests should be identified by the title of the information collection. Email address: OPREinfoacfcollection@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, Email: OIRA_SUBMISSION@OMB.EOP.GOV, Attn: Desk Officer for the Administration, for Children and Families.

Mary Jones,
ACF/OPRE, Certifying Officer.

DEPARTMENT OF HEALTH AND HUMAN SERVICES
Administration for Children and Families
Proposed Information Collection Activity; Comment Request

Proposed Projects: Immediate Disaster Case Management Intake Assessment (hardcopy and electronic versions).
Title: Immediate Disaster Case Management Intake Assessment.
OMB No.: 0970—NEW.
Description: Section 426 of the Robert T. Stafford Disaster Relief and
Emergency Assistance Act (Stafford Act), as amended, 42 U.S.C. 5189d authorizes the Federal Emergency Management Agency (FEMA) and the U.S. Department of Health Services’ Administration for Children and Families (ACF) to provide Immediate Disaster Case Management (IDCM) services under the federal Disaster Case Management Program (DCMP).

The use of the Electronic Case Management Record System (ECMRS) is aligned with Executive Order of the President 13559 and the memorandum to the Heads of Executive Departments and Agencies M–12–12 from the Office of Management and Budget to “Promote Efficient Spending to Support Agency Operations.”

The primary purpose of the information collection pertains to ACF/OHSEP’R’s initiative to improve the intake process and delivery of case management services to individuals and households impacted by a disaster. Further, the information collection will be used to support ACF/OHSEP’R’s goal to quickly identify critical gaps, resources, needs, and services to support State, local and non-profit capacity for disaster case management and to augment and build capacity where none exists. All information gathered will be exclusively used to inform the delivery of disaster case management services and programmatic strategies and improvements.

Respondents: Individuals impacted by a disaster.

ANNUAL BURDEN ESTIMATES

<table>
<thead>
<tr>
<th>Instrument</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Burden hours per response</th>
<th>Total burden hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>IDCM Intake Assessment</td>
<td>3,500</td>
<td>1</td>
<td>40 minutes</td>
<td>2,333</td>
</tr>
</tbody>
</table>

**Estimated Total Annual Burden Hours:** 2,333 hours or 140,000 minutes.

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. Email 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 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. 2018–03105 Filed 2–14–18; 8:45 am]

BILLING CODE 4184–01–P

**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Food and Drug Administration**

[Docket No. FDA–2018–N–0408]

**Oncologic Drugs Advisory Committee; Notice of Meeting; Establishment of a Public Docket; Request for Comments**

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

**ACTION:** Notice; establishment of a public docket; request for comments.

**SUMMARY:** The Food and Drug Administration (FDA) announces a forthcoming public advisory committee meeting of the Oncologic Drugs Advisory Committee. The general function of the committee is to provide advice and recommendations to FDA on regulatory issues. The meeting will be open to the public. FDA is establishing a docket for public comment on this document.

**DATES:** The meeting will be held on March 7, 2018, from 8 a.m. to 12:30 p.m.

**ADDRESSES:** FDA White Oak Campus, 10903 New Hampshire Ave., Bldg. 31 Conference Center, the Great Room (Rm. 1503), Silver Spring, MD 20993–0002. Answers to commonly asked questions including information regarding special accommodations due to a disability, visitor parking, and transportation may be accessed at: https://www.fda.gov/AdvisoryCommittees/AboutAdvisoryCommittees/ucm408555.htm.

FDA is establishing a docket for public comment on this meeting. The docket number is FDA–2018–N–0408. The docket will close on March 6, 2018. Submit either electronic or written comments on this public meeting by March 6, 2018. Please note that late, untimely filed comments will not be considered. Electronic comments must be submitted on or before March 6, 2018. The https://www.regulations.gov electronic filing system will accept comments until midnight Eastern Time at the end of March 6, 2018. Comments received by mail/hand delivery/courier (for written/paper submissions) will be considered timely if they are postmarked or the delivery service acceptance receipt is on or before that date.

Comments received on or before February 26, 2018, will be provided to the committee. Comments received after that date will be taken into consideration by FDA.

You may submit comments as follows:

**Electronic Submissions**

Submit electronic comments in the following way:

- Federal eRulemaking Portal: https://www.regulations.gov. Follow the instructions for submitting comments. Comments submitted electronically, including attachments, to https://www.regulations.gov will be posted to the docket unchanged. Because your comment will be made public, you are solely responsible for ensuring that your comment does not include any confidential information that you or a third party may not wish to be posted, such as medical information, your or anyone else’s Social Security number, or confidential business information, such as a manufacturing process. Please note that if you include your name, contact information, or other information that identifies you in the body of your comments, that information will be posted on https://www.regulations.gov.