<table>
<thead>
<tr>
<th>Attach No.</th>
<th>Section/form or survey title</th>
<th>Use metrics/month—# respond</th>
<th>Estimated time for site to complete (minutes)</th>
<th>Estimated burden (minutes/hours)</th>
<th>Frequency of response</th>
<th>Total annual usage/annual burden hours</th>
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<tr>
<td>1dd</td>
<td>CTSU System Account Request Form</td>
<td>10</td>
<td>15–20</td>
<td>0.33</td>
<td>12.00</td>
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<td>1ee</td>
<td>CTSU Request for Clinical Brochure.</td>
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<td>10</td>
<td>0.17</td>
<td>12.00</td>
<td>71</td>
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<td>1ff</td>
<td>CTSU Supply Request Form</td>
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<td>5–10</td>
<td>0.17</td>
<td>12.00</td>
<td>265</td>
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<td>Surveys/Web Forms:</td>
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<td>2</td>
<td>CTSU Web Site Customer Satisfaction Survey.</td>
<td>250</td>
<td>10–15</td>
<td>0.2500</td>
<td>1.00</td>
<td>63</td>
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<td>3</td>
<td>CTSU Helpdesk Customer Satisfaction Survey.</td>
<td>300</td>
<td>10–15</td>
<td>0.2500</td>
<td>1.00</td>
<td>75</td>
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<td>4</td>
<td>CTSU OPEN Survey</td>
<td>120</td>
<td>10–15</td>
<td>0.2500</td>
<td>1.00</td>
<td>30</td>
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<td>Annual Totals</td>
<td></td>
<td></td>
<td></td>
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<td>21,770</td>
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**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Food and Drug Administration**

**[Docket No. FDA–2010–N–0344]**

**Agency Information Collection Activities; Proposed Collection; Comment Request; Testing Communications on Medical Devices and Radiation-Emitting Products**

**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 and to allow 60 days for public comment in response to the notice. This notice solicits comments on communication studies involving medical devices and radiation-emitting products regulated by FDA. This information will be used to explore concepts of interest and assist in the development and modification of communication messages and campaigns to fulfill the agency’s mission to protect the public health.

**DATES:** Submit either electronic or written comments on the collection of information by September 13, 2010.

**ADDRESSES:** Submit electronic comments on the collection of information to http://www.regulations.gov. Submit written comments on the collection of information to the Division of Dockets Management (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:** Daniel Gittleson, Office of Information Management, Food and Drug Administration, 1350 Piccard Dr., P150–400B, Rockville, MD 20850, 301–796–5156, Daniel.Gittleson@fda.hhs.gov.

**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 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 these topics: (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.
Testing Communications on Medical Devices and Radiation-Emitting Products—(OMB Control Number 0910–New)

FDA is authorized by section 1003(d)(2)(D) of the Federal Food, Drug, and Cosmetic Act (the act) (21 U.S.C. 393(d)(2)(D)) to conduct educational and public information programs relating to the safety of regulated medical devices and radiation-emitting products. FDA must conduct needed research to ensure that such programs have the highest likelihood of being effective. Improving communications about medical devices and radiation-emitting products will involve many research methods, including individual indepth interviews, mall-intercept interviews, focus groups, self-administered surveys, gatekeeper reviews, and omnibus telephone surveys.

The information collected will serve three major purposes. First, as formative research it will provide critical knowledge needed about target audiences to develop messages and campaigns about medical device and radiation-emitting product use. Knowledge of consumer and health care professional decisionmaking processes will provide the better understanding of target audiences that FDA needs to design effective communication strategies, messages, and labels. These communications will aim to improve public understanding of the risks and benefits of using medical devices and radiation-emitting products by providing users with a better context in which to place risk information more completely.

Second, as initial testing, it will allow FDA to assess 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 will be asked to give their reaction to the messages in either individual or group settings.

Third, as evaluative research, it will allow FDA to ascertain the effectiveness of the messages and the distribution method of these messages in achieving the objectives of the message campaign. Evaluation of campaigns is a vital link in continuous improvement of communications at FDA.

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

<table>
<thead>
<tr>
<th>Section of the act</th>
<th>No. of Respondents</th>
<th>Annual Frequency per Response</th>
<th>Total Annual Responses</th>
<th>Hours per Response</th>
<th>Total Hours</th>
<th>Total Operating &amp; Maintenance Costs1</th>
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<tbody>
<tr>
<td>1003(d)(2)(D)</td>
<td>16,448</td>
<td>1</td>
<td>16,448</td>
<td>0.1739</td>
<td>2,860</td>
<td>$25,239</td>
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<tr>
<td>Total</td>
<td>16,448</td>
<td>1</td>
<td>16,448</td>
<td>0.1739</td>
<td>2,860</td>
<td>$25,239</td>
</tr>
</tbody>
</table>

1 There are no capital costs associated with this collection of information.

Annually, FDA projects about 30 studies using a variety of research methods, and lasting an average of 0.17 hours each (varying from 0.08–1.5 hours). The operating and maintenance costs include contractor expenses for designing and conducting information collection activities, specifically, drawing samples, training interviewers, collecting and analyzing information, and reporting and disseminating findings. FDA estimates the burden of this collection of information based on prior recent experience with the various types of data collection methods described earlier. FDA is requesting this burden so as not to restrict the agency’s ability to gather information on public sentiment for its proposals in its regulatory and communications programs.


Leslie Kux,
Acting Assistant Commissioner for Policy.

[FR Doc. 2010–16973 Filed 7–12–10; 8:45 am]

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Modification to the Basic Center Program Funding Opportunity Announcement

Program Office: Administration on Children, Youth, & Families—Family & Youth Services Bureau.

Funding Opportunity Title: Basic Center Program.

Announcement Type: Modification.


CFDA Number: 93.623.

Due Date for Applications: 07/19/2010.

This is a Modification to the Basic Center Program Funding Opportunity Announcement (FOA), HHS–2010–ACF–ACYF–CY–0002, published to the ACF Grant Opportunities webpage on June 2, 2010, http://www.acf.hhs.gov/grants/open/foa/view/HHS-2010-ACF-ACYF-CY-0002. A modified FOA that incorporates the following changes was published to the ACFGrant Opportunities webpage on June 25, 2010. The application procedures are hereby modified.

SUMMARY: The Family and Youth Services Bureau (FYSB) is accepting applications for the Basic Center Program (BCP), which is authorized by the Runaway and Homeless Youth Act to address Runaway and Homeless Youth (RHY) problems. BCPs provide an alternative for runaway and homeless youth who might otherwise end up with law enforcement or in the child welfare, mental health, or juvenile justice systems. Each BCP must provide runaway and homeless youth with a safe and appropriate shelter; individual, family, and group counseling, as appropriate; and aftercare.

The purpose of the modification is to correct information appearing in Section IV.2 Content and Form of Application Submission regarding application formatting and point deduction for noncompliance with FOA instructions.

Modification to the Published Announcement

Please delete the following under Section IV.2 Content and Form of Application Submission:

“Applicants that do not adhere to the prescribed format will have points deducted from the overall total after the grant review: Program narrative (which includes Objective and Need for Assistance, Results and Benefits, Approach, Organizational Profile, Staff and Position Data, and Budget