

0938–1187); *Frequency*: Annually; *Affected Public*: State, Local, or Tribal Governments, Private Sector (Business or other for-profits); *Number of Respondents*: 2,892; *Number of Responses*: 2,892; *Total Annual Hours*: 68,666. (For questions regarding this collection contact Joshua Annas at 301–492–4407.)

Dated: February 22, 2019.

William N. Parham, III,

Director, Paperwork Reduction Staff, Office of Strategic Operations and Regulatory Affairs.

[FR Doc. 2019–03459 Filed 2–27–19; 8:45 am]

BILLING CODE 4120–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifier: CMS–10065/10066]

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

AGENCY: Centers for Medicare & Medicaid Services (CMS), HHS.

ACTION: Correction of notice.

SUMMARY: This document corrects the information provided for [Document Identifier: CMS–10065/10066] titled “Hospital Notices: IM/DND.”

FOR FURTHER INFORMATION CONTACT: William N. Parham, III, (410) 786–4669.

SUPPLEMENTARY INFORMATION:

I. Background

In the February 22, 2019, issue of the *Federal Register* (84 FR 5690), we published a Paperwork Reduction Act notice requesting a 60-day public comment period for the information collection request identified under CMS–10065/10066, OMB control number 0938–1019, and titled “Hospital Notices: IM/DND.”

II. Explanation of Error

In the February 22, 2019, notice, the information provided in the second column in the middle of the notice, on page 5691, was published with incorrect information at the end of the notice. This notice corrects the language found at the end of the notice, under the second column in the middle of the notice, on page 5691 of the February 22nd notice. The related public comment period remains in effect and ends April 23, 2019.

III. Correction of Error

In FR Doc. 2019–03015 of February 22, 2019 (84 FR 5690), page 5691, the language in the second column, in the middle of the notice that begins with “[For policy questions regarding” and ends with “1799.]” is corrected to read as follows:

[(For policy questions regarding this collection contact Janet Miller at Janet.Miller@cms.hhs.gov)]

Dated: February 22, 2019.

William N. Parham, III,

Director, Paperwork Reduction Staff, Office of Strategic Operations and Regulatory Affairs.

[FR Doc. 2019–03462 Filed 2–27–19; 8:45 am]

BILLING CODE 4120–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Submission for OMB Review; Descriptive Study of the Unaccompanied Refugee Minors Program (New Collection)

AGENCY: Office of Planning, Research, and Evaluation; Administration for Children and Families; HHS.

ACTION: Request for public comment.

SUMMARY: The Office of Planning, Research, and Evaluation (OPRE) at the Administration for Children and Families (ACF), U.S. Department of Health and Human Services (HHS) is proposing data collection activities as part of a project to better understand the range of child welfare services and benefits provided through the Unaccompanied Refugee Minors (URM) Program.

DATES: *Comments due within 30 days of publication.* 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.

ADDRESSES: 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.

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: OPREinfocollection@acf.hhs.gov.

SUPPLEMENTARY INFORMATION:

Description: The proposed information collection activities to be submitted in the Descriptive Study of the Unaccompanied Refugee Minors Program package include:

1. Survey of State Refugee Coordinators (SRCs) from the 15 states with URM programs.
2. Survey of URM Program Directors from all 22 URM programs.
3. Survey of Private Custody Child Welfare Agency Administrators from nine states with private custody arrangements.
4. Interviews with URM Program Managers from six URM programs.
5. Interviews with URM Program Staff (e.g., case managers, data managers) from six URM programs.
6. Interviews with Child Welfare Agency Administrators who have contact with six URM programs.
7. Interviews with Community Partners including leadership and line staff from local organizations, such as health care and mental health care providers, legal aid organizations, and faith-based groups serving the URM population at six URM program sites.
8. Interviews with Community Partners in the field of education, such as school administrators and counselors, and organizations providing English language education and support at six URM program sites.
9. Focus Groups for URM Youth from six URM programs.
10. Focus Groups for URM Foster Families from six URM programs.

Respondents: State Refugee Coordinators and supporting staff, URM Program Directors and supporting staff, Child Welfare Agency Administrators and supporting staff, URM program staff (case workers, data managers, and other staff), staff from community partner organizations (e.g., health and mental health service providers, education service providers, faith-based organizations), URM youth, and URM foster families.

ANNUAL BURDEN ESTIMATES

Instrument	Total number of respondents	Number of responses per respondent	Average burden hours per response	Annual burden hours
Survey of State Refugee Coordinators	38	1	0.67	25
Survey of URM Program Directors	55	1	1	55
Survey of Private Custody Child Welfare Agency Administrators	21	1	0.67	14
Interviews with URM Program Managers	9	1	1.5	14
Interviews with URM Program Staff	36	1	1.5	54
Interviews with Child Welfare Agency Administrators	26	1	1	26
Interviews with Community Partners [General]	48	1	1	48
Interviews with Community Partners [Education]	12	1	1	12
Focus Groups with URM Youth	54	1	1.5	81
Focus Groups with URM Foster Families	54	1	1.5	81

Estimated Total Annual Burden Hours: 410.

Authority: Section 1110 of the Social Security Act.

Mary B. Jones,

ACF/OPRE Certifying Officer.

[FR Doc. 2019-03489 Filed 2-27-19; 8:45 am]

BILLING CODE 4184-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2019-N-0639]

Science Advisory Board to the National Center for Toxicological Research Advisory Committee; Notice of Meeting

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA or Agency) announces a forthcoming public advisory committee meeting of the Science Advisory Board (SAB) to the National Center for Toxicological Research (NCTR). The general function of the committee is to provide advice and recommendations to the Agency on research being conducted at the NCTR. At least one portion of the meeting will be closed to the public.

DATES: The meeting will be held on March 19, 2019, from 8 a.m. to 5:45 p.m., and on March 20, 2019, from 8 a.m. to 11:30 a.m.

ADDRESSES: Heifer Village, 1 World Avenue, Little Rock, AR 72202. 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> and [https://www.heifer.org/what-you-can-](https://www.heifer.org/what-you-can-do/experience-heifer/heifer-village/index.html)

[do/experience-heifer/heifer-village/index.html](https://www.heifer.org/what-you-can-do/experience-heifer/heifer-village/index.html).

FOR FURTHER INFORMATION CONTACT:

Donna Mendrick, National Center for Toxicological Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 32, Rm. 2208, Silver Spring, MD 20993-0002, 301-796-8892; or FDA Advisory Committee Information Line, 1-800-741-8138 (301-443-0572 in the Washington, DC area). A notice in the **Federal Register** about last minute modifications that impact a previously announced advisory committee meeting cannot always be published quickly enough to provide timely notice. Therefore, you should always check the Agency's website at <https://www.fda.gov/AdvisoryCommittees/default.htm> and scroll down to the appropriate advisory committee meeting link, or call the advisory committee information line to learn about possible modifications before coming to the meeting.

SUPPLEMENTARY INFORMATION:

Agenda: On March 19, 2019, the SAB Chair will welcome the participants, and the NCTR Director will provide a Center-wide update on scientific initiatives and accomplishments during the past year. The SAB will be presented with an overview of the SAB Subcommittee Site Visit Report and a response to this review. There will be updates from the NCTR Research Divisions and a public comment session.

On March 20, 2019, there will be a statement given by the FDA Chief Scientist. The Center for Biologics and Evaluation and Research, Center for Drug Evaluation and Research, Center for Devices and Radiological Health, Center for Food Safety and Applied Nutrition, and Center for Tobacco Products will each briefly discuss their center-specific research strategic needs and potential areas of collaboration.

Following an open discussion of all the information presented, the open

session of the meeting will close so the SAB members can discuss personnel issues at NCTR at the end of each day.

FDA intends to make background material available to the public no later than 2 business days before the meeting. If FDA is unable to post the background material on its website prior to the meeting, the background material will be made publicly available at the location of the advisory committee meeting, and the background material will be posted on FDA's website after the meeting. Background material is available at <https://www.fda.gov/AdvisoryCommittees/Calendar/default.htm>. Scroll down to the appropriate advisory committee meeting link.

Procedure: On March 19, 2019, from 8 a.m. to 5:45 p.m., and March 20, 2019, from 8 a.m. to 11:30 a.m., the meeting is open to the public. Interested persons may present data, information, or views, orally or in writing, on issues pending before the committee. Written submissions may be made to the contact person on or before March 12, 2019. Oral presentations from the public will be scheduled between approximately 1:15 p.m. and 2:15 p.m. Those individuals interested in making formal oral presentations should notify the contact person and submit a brief statement of the general nature of the evidence or arguments they wish to present, the names and addresses of proposed participants, and an indication of the approximate time requested to make their presentation on or before March 14, 2019. Time allotted for each presentation may be limited. If the number of registrants requesting to speak is greater than can be reasonably accommodated during the scheduled open public hearing session, FDA may conduct a lottery to determine the speakers for the scheduled open public hearing session. The contact person will notify interested persons regarding their request to speak by March 14, 2019.