Section 1334 of the Small Business Jobs and Credit Act of 2010 (Pub. L. 111–240) and the Small Business Administration’s Final Rule at 78 FR 42391, Small Business Subcontracting, published on July 16, 2013, and effective August 15, 2013, requires the prime contractor to self-report to the contracting officer when the prime contractor makes late or reduced payments to small business subcontractors. In addition, the contracting officer is required to record the identity of contractors with a history of late or reduced payments to small business subcontractors. In accordance with the clause at 52.242–XX, Payments to Small Business Subcontractors, that are determined by the contracting officer to be unjustified. A notice was published in the Federal Register at 81 FR 3087, on January 20, 2016, as part of a proposed rule under FAR Case 2014–004. Two comments were received on the information collection.

B. Discussion and Analysis

Comment: Two respondents stated that the Councils had underestimated the public burden in regards to the proposed rule. One respondent commented that the FAR Councils had underestimated the implementation burden on commercial item and construction item contractors, especially considering the broad definition of “subcontractor” that applies to the proposed rule. The other respondent believed that the estimate of reporting time of only two hours per respondent is grossly underestimated. This negligible amount of time assumes that all contractors can easily identify from their payment systems which subcontractors are small businesses. The respondent believed that this is often not the case, and that the small business size status of a subcontractor may be unknown to the contractor’s other accounting systems. The other respondent commented that since the Small Business Jobs Act of 2010 does not specifically require that the subcontractor payment clause apply to commercial contracts, the respondent recommended that the FAR Council seek additional information about the burden on contractors before a determination is made to apply the payment of subcontractor requirements to commercial item acquisitions. The respondent did not find that the availability of limited information indicated that the burden may not be significant, as described in the proposed rule. Rather, initial feedback from contractors suggested that the burdens associated with reporting under the rule will have a significant impact.

Response: The respondents do not offer data with which to support changing the current estimated public burden hours. However, since this is a new rule without an empirical frame of reference, the public reporting burden is reviewed every three years and can be adjusted as necessary.

Public comments are particularly invited on: Whether this collection of information is necessary for the proper performance of functions of the FAR, and whether it will have practical utility; whether our estimate of the public burden of this collection of information is accurate, and based on valid assumptions and methodology; ways to enhance the quality, utility, and clarity of the information to be collected; and ways in which we can minimize the burden of the collection of information on those who are to respond, through the use of appropriate technological collection techniques or other forms of information technology.

C. Annual Reporting Burden

Respondents: 5,457.
Responses per Respondent: 1.
Total Annual Responses: 5,457.
Hours per Response: 2.
Total Burden Hours: 10,914.
Obtaining Copies of Proposals: Requesters may obtain a copy of the information and documents from the General Services Administration, Regulatory Secretariat Division (MVCB), 1800 F Street NW., Washington, DC 20405, telephone 202–501–4755. Please cite OMB Control Number 9000–0196, Payment of Subcontractors, in all correspondences.

Dated: November 15, 2016.

Lorin S. Curit,
Director, Federal Acquisition Policy Division, Office of Governmentwide Acquisition Policy, Office of Acquisition Policy, Office of Governmentwide Policy.

[FR Doc. 2016–11130 Filed 12–23–16; 8:45 am]
BILLING CODE 9520–EP–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Disease, Disability, and Injury Prevention and Control Special Emphasis Panel (SEP): Initial Review

In accordance with Section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92–463), the Centers for Disease Control and Prevention (CDC) announces a meeting for the initial review of applications in response to Funding Opportunity Announcement (FOA) GH17–001, Evaluations to Improve Prevention Interventions Under the President’s Emergency Plan for AIDS Relief (PEPFAR).

Time and Date: 9:00 a.m.–2:00 p.m., EST, January 25, 2017 (Closed).

Place: Teleconference.

Status: The meeting will be closed to the public in accordance with provisions set forth in Section 552b(c)(4) and (6), Title 5 U.S.C., and the Determination of the Director, Management Analysis and Services Office, CDC, pursuant to Public Law 92–463.

Matters for Discussion: The meeting will include the initial review, discussion, and evaluation of applications received in response to FOA GH17–001, Evaluations to Improve Prevention Interventions Under the President’s Emergency Plan for AIDS Relief (PEPFAR).

Contact Person for More Information: Hylan Shoob, Scientific Review Officer, Center for Global Health (CGH) Science Office, CGH, CDC, 1600 Clifton Road, NE., Mailstop D–69, Atlanta, Georgia 30329, Telephone: (404) 639–4796.

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
Department of Health and Human Services
Administration for Children and Families

[CFDA Number: 93.676]

Announcing the Intent To Award a Single-Source Program Expansion Supplements to Cooperative Agreements Within the Office of Refugee Resettlement’s Unaccompanied Children’s (UC) Program

AGENCY: Office of Refugee Resettlement, ACF, HHS.

ACTION: This notice announces the intent to award a single-source expansion supplement grant to existing grantees’, BCFS Health and Human Services (90ZU0075) and the U.S. Committee for Refugees and Immigrants (90ZU0081), Cooperative Agreement within the Office of Refugee Resettlement’s Unaccompanied Children’s (UC) Program.

SUMMARY: The Administration for Children and Families (ACF), Office of Refugee Resettlement (ORR), announces its intent to award a cooperative agreement of up to $3,311,087 as a single-source expansion supplements to the Post Release Services Programs within the Unaccompanied Children’s (UC) Program.

The expansion supplement grants will support the immediate need for additional post-release services to accommodate the increasing number of UCs being referred by DHS, and as a result, the increase of UCs referred for post-release services. The increase in the UC population necessitates the need for expansion of services to expedite the release of UC. The Flores v. Reno settlement agreement requires that requires the timely release of children and youth to qualified parents, guardians, relatives or other adults, referred to as “sponsors.”

DATES: Supplemental award funds will support activities from September 30, 2015 through September 29, 2016.

FOR FURTHER INFORMATION CONTACT: Jallyn Sualog, Director, Division of Children’s Services, Office of Refugee Resettlement, 330 C Street SW., Washington, DC 20201. Email: DCSProgram@acf.hhs.gov

SUPPLEMENTARY INFORMATION: ORR is continuously monitoring its capacity to provide post-release services to the unaccompanied children in HHS custody.

ORR has specific requirements for the provision of services. Award recipients must have the infrastructure, licensing, experience, and appropriate level of trained staff to meet those requirements. The expansion of the existing post-release services program through this supplemental award is a key strategy for ORR to be prepared to meet its responsibility of safe and timely release of Unaccompanied Children referred to its care by DHS and so that the US Border Patrol can continue its vital national security mission to prevent illegal migration, trafficking, and protect the borders of the United States.

Statutory Authority: This program is authorized by—
(A) Section 462 of the Homeland Security Act of 2002, which in March 2003, transferred responsibility for the care and custody of Unaccompanied Alien Children from the Commissioner of the former Immigration and Naturalization Service (INS) to the Director of ORR of the Department of Health and Human Services (HHS).
(B) The Flores Settlement Agreement, Case No. CV85–4544RJK (C.D. Cal. 1996), as well as the William Wilberforce Trafficking Victims Protection Reauthorization Act of 2008 (Pub. L. 110–457), which authorizes post release services under certain conditions to eligible children. All programs must comply with the Flores Settlement Agreement, Case No. CV85–4544RJK (C.D. Cal. 1996), pertinent regulations and ORR policies and procedures.

Christopher Beach,
Senior Grants Policy Specialist, Division of Grants Policy, Office of Administration, Administration for Children and Families.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2016–D–1495]

Factors to Consider Regarding Benefit-Risk in Medical Device Product Availability, Compliance, and Enforcement Decisions: Guidance for Industry and Food and Drug Administration Staff; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice of availability.

SUMMARY: The Food and Drug Administration (FDA or Agency) is announcing the availability of the guidance entitled “Factors to Consider Regarding Benefit-Risk in Medical Device Product Availability, Compliance, and Enforcement Decisions.” This guidance is intended to provide clarity for FDA staff and industry regarding the benefit and risk factors FDA may consider in prioritizing resources for compliance and enforcement efforts to maximize medical device quality and patient safety. Although product availability and other medical device compliance and enforcement decisions are generally fact-specific, FDA believes that explaining how we consider the factors listed in the guidance will improve the consistency and transparency of these kinds of decisions. A common understanding of how FDA considers benefit and risk may better align industry’s and FDA’s focus on actions that maximize benefit to patients, improve medical device quality, and reduce risk to patients. This guidance is in effect at this time.

DATES: Submit either electronic or written comments on this guidance at any time. General comments on Agency guidance documents are welcome at any time.

ADDRESSES: 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.
• If you want to submit a comment with confidential information that you do not wish to be made available to the public, submit the comment as a written/paper submission and in the manner detailed (see “Written/Paper Submissions” and “Instructions”).