

GENERAL SERVICES ADMINISTRATION

[Notice—CECANF—2015—07; Docket No. 2015—0004; Sequence No. 8]

Commission To Eliminate Child Abuse and Neglect Fatalities; Announcement of Meeting

AGENCY: Commission to Eliminate Child Abuse and Neglect Fatalities, GSA.

ACTION: Meeting Notice.

SUMMARY: The Commission to Eliminate Child Abuse and Neglect Fatalities (CECANF), a Federal Advisory Committee established by the Protect Our Kids Act of 2012, will hold conference calls open to the public on the following dates: Wednesday, August 26, 2015; Thursday, September 24, 2015; Friday, October 30, 2015; Thursday, November 12, 2015; and December 3, 2015.

DATES: The meetings will be held on the noted dates from 1:00 p.m. to 3:00 p.m., Eastern Daylight Time (EDT).

ADDRESSES: CECANF will convene these meetings via conference call.

Submit comments identified by “Notice—CECANF—2015—08,” by either of the following methods:

- *Regulations.gov:* <http://www.regulations.gov>.

Submit comments via the Federal eRulemaking portal by searching for “Notice—CECANF—2015—08.” Select the link “Comment Now” that corresponds with “Notice—CECANF—2015—08.” Follow the instructions provided on the screen. Please include your name, organization name (if any), and “Notice—CECANF—2015—08” on your attached document.

- *Mail:* U.S. General Services Administration, 1800 F Street NW., Room 7003D, Washington DC 20405, Attention: Tom Hodnett (CD) for CECANF.

Instructions: Please submit comments only and cite “Notice—CECANF—2015—08, Announcement of Meeting” in all correspondence related to this notice. Comments received generally will be posted without change to <http://www.regulations.gov>, including any personal and/or business confidential information provided. To confirm receipt of your comment(s), please check <http://www.regulations.gov>, approximately two to three days after submission to verify posting (except allow 30 days for posting of comments submitted by mail).

FOR FURTHER INFORMATION CONTACT: Visit the CECANF Web site at <https://eliminatechildabusefatalities.sites.usa.gov/> or contact Patricia Brincefield,

Communications Director, at 202–818–9596, General Services Administration, 1800 F Street NW., Room 7003D, Washington DC 20405, Attention: Tom Hodnett (CD) for CECANF.

SUPPLEMENTARY INFORMATION:

Background: CECANF was established to develop a national strategy and recommendations for reducing fatalities resulting from child abuse and neglect.

Agenda: Commission members will continue discussing the work plans of the Commission subcommittees, the information that they have obtained to date, and emerging recommendations.

Attendance at the Meetings: Individuals interested in participating by teleconference should dial 1–800–870–9004 and then enter *3676137#. Detailed meeting minutes will be posted within 90 days of the meeting. Members of the public will not have the opportunity to ask questions or otherwise participate in the meeting.

However, members of the public wishing to comment should follow the steps detailed under the heading **ADDRESSES** in this publication or contact us via the CECANF Web site at <https://eliminatechildabusefatalities.sites.usa.gov/contact-us/>.

Dated: August 20, 2015.

Karen White,

Executive Assistant.

[FR Doc. 2015–21189 Filed 8–25–15; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Reporting of Pregnancy Success Rates From Assisted Reproductive Technology (ART) Programs

AGENCY: Centers for Disease Control and Prevention (CDC), Department of Health and Human Services (DHHS).

ACTION: Final notice.

SUMMARY: The Centers for Disease Control and Prevention (CDC) in the Department of Health and Human Services (HHS) announces the requirements for reporting of pregnancy success rates from assisted reproductive technology (ART) programs as required by the Fertility Clinic Success Rate and Certification Act of 1992 (FCSRCA). This notice describes who shall report to HHS/CDC, the reporting system (National ART Surveillance System (NASS)); the process for reporting by each ART program; the data to be

reported; and the contents of the published reports. The proposed changes to reporting requirements were published in the **Federal Register** on July 21, 2014 (79 FR. 42328) and February 18, 2015 (80 FR. 8659) in accordance with requirements under the Paperwork Reduction Act; public comments and recommendations were requested. This notice incorporates the comments received from those notices and supersedes the previous notice published in the **Federal Register**, September 1, 2000 (65 FR. 53310).

FOR FURTHER INFORMATION CONTACT: Jeani Chang, Division of Reproductive Health, National Center for Chronic Disease Prevention and Health Promotion, Centers for Disease Control and Prevention, 4770 Buford Highway, MS–74, Atlanta, Georgia 30341. Phone: (770) 488–6370. Email: artinfo@cdc.gov.

SUPPLEMENTARY INFORMATION: Section 2(a) of Public Law 102–493 (42 U.S.C.263a–1(a)), the Fertility Clinic Success Rate and Certification Act of 1992 (FCSRCA) requires that each ART program report annually to the Secretary of the Department of Health and Human Services through the Centers for Disease Control and Prevention (1) pregnancy success rates achieved by such ART program and (2) the identity of each embryo laboratory used by such ART program and whether the laboratory is certified or has applied for such certification under the act.

FCSRCA defines “assisted reproductive technology” (ART) as “all treatments or procedures which include the handling of human oocytes or embryos, including in vitro fertilization, gamete intrafallopian transfer, zygote intrafallopian transfer, and such other specific technologies as the Secretary may include in this definition, after making public any proposed definition in such manner as to facilitate comment from any person (including any federal or other public agency).”

The Secretary is directed in FCSRCA to define pregnancy success rates and “make public any proposed definition in such a manner as to facilitate comment from any person during its development.”

Section 2(c) (42 U.S.C. 263a–1(c)) states “the Secretary shall consult with appropriate consumer and professional organizations with expertise in using, providing, and evaluating professional services and embryo laboratories associated with assisted reproductive technologies.”

Since 1995, HHS/CDC has reported pregnancy success rates of United States ART programs as required by FCSRCA. Changes in ART practice require