as part of the Preventing and Addressing Intimate Violence when Engaging Dads (PAIVED) study. Since 2006, the Healthy Marriage and Responsible Fatherhood (HMRF) initiative has funded programs that play a key role in helping the Office of Family Assistance (OFA) achieve its goals to foster economically secure households and communities for the well-being and long-term success of children and families. The purpose of the PAIVED study is to better understand the prevalence of intimate partner violence (IPV) experienced by the population of fathers served by Responsible Fatherhood (RF) programs, and the services that federally- and non-federally funded RF programs are providing to address and contribute to the prevention of IPV among its participants.

The proposed data collection will include whether IPV content is included in RF programs, the types of activities they are using to address IPV, and related successes and challenges. Other collected data will include barriers to addressing IPV in RF programs, the relevance of addressing IPV with fathers, fathers' reactions to this programming, and what types of partnerships RF programs have with other agencies to address IPV. This information will be collected through interviews conducted over the phone and in-person with RF program staff and community partners. This information will be critical to inform future efforts to address and contribute to the prevention of IPV through RF programming.

Respondents: Responsible Fatherhood program staff (e.g., program directors and facilitators) and community partners.

### 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>Annual burden hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF program/community partner screening and participant recruitment</td>
<td>50</td>
<td>1</td>
<td>1</td>
<td>50</td>
</tr>
<tr>
<td>RF program staff semi-structured interview</td>
<td>25</td>
<td>1</td>
<td>1.5</td>
<td>38</td>
</tr>
<tr>
<td>Community partner semi-structured interview</td>
<td>15</td>
<td>1</td>
<td>1.5</td>
<td>23</td>
</tr>
</tbody>
</table>

Estimated Total Annual Burden Hours: 111.

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 (OPRE) proposes to collect information

DEPARTMENT OF HEALTH AND HUMAN SERVICES

U.S. National Authority for the WHO Global Code of Practice on the International Recruitment of Health Personnel; Notice of Public Meeting

**Time and date:** Monday, July 2, 2018, 2:00 p.m.–4:00 p.m. EST.

**Place:** Hubert H. Humphrey Building, Room 325A, 200 Independence Ave. SW, Washington, District of Columbia 20201.

**Status:** Open, but requiring RSVP to us.who.irhp@hhs.gov by Monday, June 25, 2018.

Purpose: The purpose of the World Health Organization (WHO) Global Code of Practice on International Recruitment of Health Personnel (Global Code) is “to establish and promote voluntary principles and practices for the ethical international recruitment of health personnel and to facilitate the strengthening of health systems.” The United States Government has designated the Office of Global Affairs (OGA) and the Health Resources and Services Administration (HRSA) as co-National Authorities to be the point of contact for implementation activities. The Global Code encourages WHO Member States to cooperate with all relevant stakeholders in their implementation efforts. This meeting is intended to provide an update to all interested stakeholders on U.S. Global Code implementation efforts to date and to provide a forum for questions on activities related to implementation of the Global Code.

The meeting will be open to the public as indicated above, with attendance limited to space available. Individuals who plan to attend and need special assistance, such as sign language interpretation or other reasonable accommodations, should notify us within their RSVP at least 10 business days prior to the meeting. Foreign nationals planning to attend the session in person will require additional paperwork for security clearance and that this clearance process requires a minimum of 10 business days.

**RSVP:** Due to security restrictions for entry into the HHS Humphrey Federal Building, we will need to receive RSVPs for this event. Please send your full name and organization to us.who.irhp@hhs.gov. If you are not a U.S. citizen, you must RSVP no later than Monday, June 18, 2018. Please note this in the subject line of your RSVP, and our office will contact you to gain additional biographical information for your
DEPARTMENT OF HEALTH AND HUMAN SERVICES

Findings of Research Misconduct

AGENCY: Office of the Secretary, HHS.

ACTION: Notice.

SUMMARY: Findings of research misconduct have been made on the part of Maria Cristina Miron Elqutub, Research Interviewer, University of Texas MD Anderson Cancer Center (MDACC). Dr. Elqutub engaged in research misconduct in research supported by National Institute of Dental and Craniofacial Research (NIDCR), National Institutes of Health (NIH), grant U01 DE019765–01. The administrative actions, including three (3) years of supervision, were implemented beginning on April 26, 2018, and are detailed below.

FOR FURTHER INFORMATION CONTACT: Wanda K. Jones, Dr.P.H., Interim Director, Office of Research Integrity, 1101 Wootton Parkway, Suite 750, Rockville, MD 20852, (240) 453–8200.

SUPPLEMENTARY INFORMATION: Notice is hereby given that the Office of Research Integrity (ORI) has taken final action in the following case:

Maria Cristina Miron Elqutub, University of Texas MD Anderson Cancer Center: Based on Respondent’s admission, the report of an inquiry conducted by MDACC, and analysis conducted by ORI in its oversight review, ORI found that Ms. Maria Cristina Miron Elqutub, Research Interviewer, MDACC, engaged in research misconduct in research supported by NIDCR, NIH, grant U01 DE019765–01.

ORI found that Respondent engaged in research misconduct by intentionally and knowingly falsifying and/or fabricating data that were included in the following two (2) published papers and two (2) grant progress reports submitted to NIDCR, NIH:

  • 5 U01 DE019765–04.
  • 5 U01 DE019765–05.

Specifically, ORI found that Respondent engaged in research misconduct by recording dates and providing her own blood samples to cause these samples to be falsely labeled as samples from ninety-eight (98) study subjects in a cancer genetics study involving human blood samples. This resulted in the reporting of false data in Tables 1, 2, 3, and 4 in PLoS One 2015, in Figure 1 and Tables 1, 2, 3, and 4 in Cancer 2015, and in the Results sections of Project 2 progress reports for NIDCR, NIH, grants 5 U01 DE019765–04 and 5 U01 DE019765–05.

Ms. Elqutub entered into a Voluntary Settlement Agreement and voluntarily agreed, beginning on April 26, 2018:

(1) To have her research supervised for a period of three (3) years; Respondent agreed to ensure that prior to the submission of an application for U.S. Public Health Service (PHS) support for a research project on which Respondent’s participation is proposed and prior to Respondent’s participation in any capacity on PHS-supported research, the institution employing her must submit a plan for supervision of Respondent’s duties to ORI for approval; the supervision plan must be designed to ensure the scientific integrity of Respondent’s research contribution; Respondent agreed that she will not participate in any PHS-supported research until a supervision plan is submitted to and approved by ORI; Respondent agreed to maintain responsibility for compliance with the agreed upon supervision plan;

(2) that for a period of three (3) years, any institution employing her must submit, in conjunction with each application for PHS funds, or report, manuscript, or abstract involving PHS-supported research in which Respondent is involved, a certification to ORI that the data provided by Respondent are based on actual experiments or are otherwise legitimately derived and that the data, procedures, and methodology are accurately reported in the application, report, manuscript, or abstract;

(3) if no supervisory plan is provided to ORI, to provide certification to ORI on an annual basis for a period of three (3) years that she has not engaged in, applied for, or had her name included on any application, proposal, or other request for PHS funds without prior notification to ORI;

(4) to exclude herself voluntarily from serving in any advisory capacity to PHS including, but not limited to, service on any PHS advisory committee, board, and/or peer review committee, or as a consultant for a period of three (3) years; and


Wanda K. Jones, Interim Director, Office of Research Integrity.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Center for Scientific Review; Notice of Closed Meetings

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended, notice is hereby given of the following meetings.

The meetings will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title 5 U.S.C., as amended. The grant applications and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: Population Sciences and Epidemiology Integrated Review Group; Social Sciences and Population Studies A Study Section.

Date: June 6, 2018.

Time: 8:30 a.m. to 6:00 p.m.

Agenda: To review and evaluate grant applications.

Place: Melrose Hotel, 2430 Pennsylvania Ave, NW, Washington, DC 20037.

Contact Person: Suzanne Ryan, Ph.D., Scientific Review Officer, Center for Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 3139, MSC 7770, Bethesda, MD 20892, (301) 435–1712, ryans@csr.nih.gov.

Name of Committee: Cell Biology Integrated Review Group; Development—1 Study Section.

Date: June 12, 2018.

Time: 10:00 a.m. to 6:00 p.m.

Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892 (Virtual Meeting).

Contact Person: Thomas Beres, Ph.D., Scientific Review Officer, Center for Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 5148, MSC 7840, Bethesda, MD 20892, 301–435–1175, berestm@mail.nih.gov.