including training needs and information dissemination on health care quality and (C) the role of the Agency in light of private sector activity and opportunities for public-private partnerships. The Council is composed of members of the public, appointed by the Secretary, and Federal ex-officio members specified in the authorizing legislation. The Council did not have a quorum for the meeting scheduled for March 24th. Therefore, AHRQ is cancelling the meeting. The next meeting of the NAC is planned for July 26th.

Sharon B. Arnold, Acting Director.

[FR Doc. 2017–05588 Filed 3–21–17; 8:45 am]
BILLING CODE 4160–90–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

[60-Day—17–17XR; Docket No. CDC–2017–0027]

Proposed Data Collection Submitted for Public Comment and Recommendations

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

ACTION: Notice with comment period.

SUMMARY: The Centers for Disease Control and Prevention (CDC), as part of its continuing efforts to reduce public burden and maximize the utility of government information, invites the general public and other Federal agencies to take this opportunity to comment on proposed and/or continuing information collections, as required by the Paperwork Reduction Act of 1995. This notice invites comment on the donor registration form in support of the project titled “Acquisition of Freshly Drawn Whole Blood/Blood Products for Reference Diagnostic and Research Use.”

DATES: Written comments must be received on or before May 22, 2017.

ADDRESSES: You may submit comments, identified by Docket No. CDC–2017–0027 by any of the following methods:
• Federal eRulemaking Portal: Regulations.gov. Follow the instructions for submitting comments.
• Mail: Leroy A. Richardson, Information Collection Review Office, Centers for Disease Control and Prevention, 1600 Clifton Road NE., MS–D74, Atlanta, Georgia 30329.

Instructions: All submissions received must include the agency name and Docket Number. All relevant comments received will be posted without change to Regulations.gov, including any personal information provided. For access to the docket to read background documents or comments received, go to Regulations.gov.

Please note: All public comment should be submitted through the Federal eRulemaking portal (Regulations.gov) or by U.S. mail to the address listed above.

FOR FURTHER INFORMATION CONTACT: To request more information on the proposed project or to obtain a copy of the information collection plan and instruments, contact the Information Collection Review Office, Centers for Disease Control and Prevention, 1600 Clifton Road NE., MS–D74, Atlanta, Georgia 30329; phone: 404–639–7570; Email: obm@cdc.gov.

SUPPLEMENTARY INFORMATION: Under the Paperwork Reduction Act of 1995 (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. In addition, the PRA also requires Federal agencies to provide a 60-day notice in the Federal Register concerning each proposed collection of information, including each new proposed collection, each proposed extension of existing collection of information, and each reinstatement of previously approved information collection before submitting the collection to OMB for approval. To comply with this requirement, we are publishing this notice of a proposed data collection as described below.

Comments are invited on:
(a) Whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have practical utility; (b) the accuracy of the agency’s estimate of the burden of the proposed collection of information; (c) ways to enhance the quality, utility, and clarity of the information to be collected; (d) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques or other forms of information technology; and (e) estimates of capital or start-up costs and costs of operation, maintenance, and purchase of services to provide information. Burden means the total time, effort, or financial resources expended by persons to generate, maintain, retain, disclose or provide information to or for a Federal agency. This includes the time needed to review instructions; to develop, acquire, install and utilize technology and systems for the purpose of collecting, validating and verifying information, processing and maintaining information, and disclosing and providing information; to train personnel and to be able to respond to a collection of information, to search data sources, to complete and review the collection of information; and to transmit or otherwise disclose the information.

Proposed Project

Acquisition of Freshly-Drawn Whole Blood/Blood Products for Reference Diagnostic and Research Use—Existing Information Collection in Use Without an OMB Control Number—National Center for Emerging and Zoonotic Infectious Diseases (NCEZID), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

The CDC seeks a three-year OMB approval to collect information in support of fresh blood/blood products for laboratory programs.

The CDC regularly requires freshly drawn whole blood, serum, plasma, mononuclear white cell and platelet concentrates for research purposes, for reagents, and as “normal” control materials. To enhance the safety of CDC personnel handling these materials, the blood/blood products, or the donors thereof, must be screened for evidence of possible infections by specific testing. At the same time, donor confidentiality must be assured and adequate counseling must be available, in case any specimens or donors test positive for certain transmissible infections.

The donor registration form referenced by this request is a brief, 11-question form that establishes the availability of volunteer donors to participate in the donor program to fill this need for fresh blood/blood products for CDC. The registration form captures donors’ availability to donate, interest in various types of donations, smoking history, exercise background, alcohol consumption, measles vaccination history, cholesterol test history, and medications background.

Donors required to maintain the CDC donor pool are recruited by contract program managers often by referral of current donors, directed outreach for new donors by email, occasional posting of notices in areas frequented by CDC personnel, or at local universities for possible student populations.

All donor information is collected and protected by medical professionals with donor/patient confidentiality protected. Information from this form is only used to determine donor eligibility for blood product requests made by CDC laboratory programs. Approximately 25 volunteer donors are enrolled annually.
There is no cost to respondents other than the time to participate. Authorizing legislation comes from Section 301 of the Public Health Service Act (42 U.S.C. 241).

### Table 1: Estimated Annualized Burden Hours

<table>
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<th>Type of respondents</th>
<th>Form name</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Avg. burden per response (in hrs.)</th>
<th>Total burden (in hrs.)</th>
</tr>
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<td>Total</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

![Image of the table with the data filled in]

### Estimate of Annual Burden

- **Total burden (in hrs.):** 7

#### DEPARTMENT OF HEALTH AND HUMAN SERVICES

**Centers for Disease Control and Prevention**

[Docket Number NIOSH–294]

**World Trade Center Health Program: Request for Nominations of Scientific Peer Reviewers of Proposed Additions to the List of WTC-Related Health Conditions**

**AGENCY:** National Institute for Occupational Safety and Health (NIOSH) of the Centers for Disease Control and Prevention (CDC), Department of Health and Human Services (HHS).

**ACTION:** Request for scientific peer reviewers.

**SUMMARY:** The CDC is soliciting nominations, including self-nominations, for scientific peer reviewers of proposed additions of conditions to the List of World Trade Center (WTC)-Related Health Conditions (List).

Title I of the James Zadroga 9/11 Health and Compensation Act of 2010, Public Law 111–347 (Jan. 2, 2011), amended by Public Law 114–113 (Dec. 18, 2015), added Title XXXIII to the Public Health Service Act (PHS Act), establishing the WTC Health Program within HHS (42 U.S.C. 300mm to 300mm–61). When the Administrator proposes to add a condition to the List, he must publish the proposed rule in accordance with the Administrative Procedure Act (5 U.S.C. 553). Additionally, as required by the James Zadroga 9/11 Health and Compensation Reauthorization Act in section 3312(a)(6)(F), prior to issuing a final rule to add a health condition to the List, the Administrator must provide for an independent peer review of the scientific and technical evidence that would be the basis for issuing such final rule.

#### Table of Contents:

- Dates:
- Addresses:
- For Further Information Contact:
- Supplementary Information:

**DATES:** Nominations must be submitted (postmarked or electronically received) by February 1, 2019.

**ADDRESSES:** You may submit a nomination identified by NIOSH Docket 294, by any of the following methods.

- Electronic nominations, including attachments to nioshdocket@cdc.gov.
  - Regular, Express, or Overnight Mail: Written nominations may be submitted (one original and two copies) to the following address only: NIOSH Docket 294, c/o Kiana Harper, National Institute for Occupational Safety and Health, Centers for Disease Control and Prevention, Patriots Plaza 1, 95 E Street SW, Suite 9200, Washington, DC 20201. Telephone and facsimile submissions cannot be accepted.

**FOR FURTHER INFORMATION CONTACT:** Paul Middendorf, Ph.D., Deputy Associate Director for Science, 1600 Clifton Rd. NE., MS: E–20, Atlanta, GA 30329; telephone (404)498–2500 (this is not a toll-free number); email pmiddendorf@cdc.gov.

**Restrictions:** Nominations of peer reviewers must be accompanied by:

- Name
- Occupation
- Employer
- Contact information including mailing address, email, and phone number
- Listing of scientific credentials including academic degrees and specialized training
- Area of competencies (e.g., medical, epidemiology, exposure assessment, industrial hygiene)
- Area of specialty (e.g., Cardiovascular, Integumentary, Gastrointestinal, Endocrine, Urinary, Immune, Lymphatic, Muscular, Nervous, Reproductive, Respiratory, Skeletal) Publication list
  - Other materials to support the nominee’s ability to perform scientific peer review
  - For third-party nominations, affirmation from the nominee that they are aware of and agree to the nomination

**A Curriculum vitae that includes all of the above information may alternatively be submitted.**

**SUPPLEMENTARY INFORMATION:** The James Zadroga 9/11 Health and Compensation Reauthorization Act in section 3312(a)(6)(F) requires the Administrator to provide for an independent peer review of the scientific and technical evidence that would be the basis for issuing a final rule to add a health condition to the List prior to issuing the final rule. To assist in accomplishing independent peer review in a timely manner, the Administrator has determined that he will develop a standing pool of persons with the scientific, technical, and medical background to potentially serve in this role to provide their individual input to the Administrator based on the health condition in the proposed rule under consideration. The peer reviewers will not meet as a group, provide consensus advice or recommendations to the Administrator, or produce a collective work product(s). Therefore, the Administrator is requesting nominations of persons to serve as scientific peer reviewers.

All persons who have the necessary minimum qualifications will be included in the standing pool of potential peer reviewers. These persons will be included in the standing pool of potential peer reviewers for 3 years unless they request in writing to be removed. After 3 years persons may be nominated again and will be required to update their information.

The Administrator will select peer reviewers for any proposed rule by matching the nature of the proposed conditions with the expertise of the scientific peer reviewers.