A. Purpose
Past performance information regarding a contractor’s actions under previously awarded contracts is relevant information for future source selection purposes. The information collection requirements at FAR 15.304 and 42.15 remain the same; however, the public burden has been adjusted downward, based on the total annual responses. The estimated responses used to calculate the burden is based on the availability of data on Fiscal Year (FY) 2017 awards from existing systems (the Federal Procurement Data System and the Contractor Performance Assessment Reporting System).

B. Annual Reporting Burden
Responses during Source Selection:
- **Respondents:** 7,055.
- **Responses per Respondent:** 4.
- **Annual Responses:** 28,220.
- **Hours per Response:** 2.
- **Total Burden Hours:** 56,440.

Responses in CPARS:
- **Respondents:** 63,444.
- **Responses per Respondent:** 1.
- **Annual Responses:** 63,444.
- **Hours per Response:** 2.
- **Total Burden Hours:** 126,888.
- **Total Annual Burden:** 183,328.

C. Public Comments
A public notice published in the Federal Register at 82 FR 57270 on December 4, 2017. No comments were received. 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.

**Instructions:** Please submit comments only and cite “Information Collection 9000–0142, Past Performance Information”, in all correspondence related to this collection. 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 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:** Mr. Curtis E. Glover, Sr., Procurement Analyst, Acquisition Policy Division, at GSA 202–501–1448 or email curtis.glover@gsa.gov.

**SUPPLEMENTARY INFORMATION:**

**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Centers for Disease Control and Prevention**

[60Day–18–0530; Docket No. CDC–2018–0016]

**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 effort to reduce public burden and maximize the utility of government information, invites the general public and other Federal agencies the opportunity to comment on a proposed and/or continuing information collection, as required by the Paperwork Reduction Act of 1995. This notice invites comment on a proposed information collection project titled EEOICPA Dose Reconstruction Interviews and Forms. This data collection permits claimants under the Energy Employees Occupational Illness Compensation Program Act (EEOICPA) to provide information potentially useful in reconstructing radiation doses, and to confirm that they have no further information to submit.

**DATES:** CDC must receive written comments on or before April 23, 2018.

**ADDRESSES:** You may submit comments, identified by Docket No. CDC–2018–0016 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. CDC will post, without change, all relevant comments to Regulations.gov.

Please note: Submit all Federal comments 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 Leroy A. Richardson, Information Collection Review Office, Centers for Disease Control and Prevention, 1600 Clifton Road NE, MS–D74, Atlanta, Georgia 30329; phone: 404–639–7570; Email: omb@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 the OMB for approval. To comply with this requirement, we are publishing this notice of a proposed data collection as described below.

The OMB is particularly interested in comments that will help:
1. Evaluate whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information will have practical utility;
2. Evaluate the accuracy of the agency’s estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used;
3. Enhance the quality, utility, and clarity of the information to be collected; and
4. Minimize the burden of the collection of information on those who are to respond, including through the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology, e.g., permitting electronic submissions of responses.

5. Assess information collection costs.

Proposed Project

EEOICPA Dose Reconstruction Interviews and Forms, OMB No. 0920–0530, expires 04/30/2018—Extension—National Institute for Occupational Safety and Health (NIOSH), Centers for Disease Control and Prevention (CDC), Department of Health and Human Services (DHHS).

Background and Brief Description

On October 30, 2000, the Energy Employees Occupational Illness Compensation Program Act of 2000 (42 U.S.C. 7384–7385) was enacted. This Act established a federal compensation program for employees of the Department of Energy (DOE) and certain of its contractors, subcontractors and vendors, who have suffered cancers and other designated illnesses as a result of exposures sustained in the production and testing of nuclear weapons.

Executive Order 13179, issued on December 7, 2000, delegated authorities assigned to “the President” under the Act to the Departments of Labor, Health and Human Services, Energy and Justice. The Department of Health and Human Services (DHHS) was delegated the responsibility of establishing methods for estimating radiation doses received by eligible claimants with cancer applying for compensation. NIOSH is applying the following methods to estimate the radiation doses of individuals applying for compensation.

In performance of its dose reconstruction responsibilities, under the Act, NIOSH is providing voluntary interview opportunities to claimants (or their survivors) individually and providing them with the opportunity to assist NIOSH in documenting the work history of the employee by characterizing the actual work tasks performed. In addition, NIOSH and the claimant may identify incidents that may have resulted in undocumented radiation exposures, characterizing radiological protection and monitoring practices, and identify co-workers and other witnesses as may be necessary to confirm undocumented information. In this process, NIOSH uses a computer assisted telephone interview (CATTI) system, which allows interviews to be conducted more efficiently and quickly as opposed to a paper-based interview instrument. Both interviews are voluntary and failure to participate in either or both interviews will not have a negative effect on the claim, although voluntary participation may assist the claimant by adding important information that may not be otherwise available.

NIOSH uses the data collected in this process to complete an individual dose reconstruction that accounts, as fully as possible, for the radiation dose incurred by the employee in the line of duty for DOE nuclear weapons production programs. After dose reconstruction, NIOSH also performs a brief, voluntary final interview with the claimant to explain the results and to allow the claimant to confirm or question the records NIOSH has compiled. This will also be the final opportunity for the claimant to supplement the dose reconstruction record.

At the conclusion of the dose reconstruction process, the claimant submits a form to confirm that the claimant has no further information to provide to NIOSH about the claim at this time. The form notifies the claimant that signing the form allows NIOSH to forward a dose reconstruction report to DOL and to the claimant, and closes the record on data used for the dose reconstruction. Signing this form does not indicate that the claimant agrees with the outcome of the dose reconstruction. The dose reconstruction results will be supplied to the claimant and to the DOL, the agency that will utilize them as one part of its determination of whether the claimant is eligible for compensation under the Act.

There is no cost to respondents other than their time.

ESTIMATED ANNUALIZED BURDEN HOURS

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<th>Type of respondent</th>
<th>Form name</th>
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<th>Number of responses per respondent</th>
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</table>

Leroy A. Richardson,
Chief, Information Collection Review Office, Office of Scientific Integrity, Office of the Associate Director for Science, Office of the Director, Centers for Disease Control and Prevention.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day–18–1048]

Agency Forms Undergoing Paperwork Reduction Act Review

In accordance with the Paperwork Reduction Act of 1995, the Centers for Disease Control and Prevention (CDC) has submitted the information collection request titled “Assessing Education Agency Staff Perceptions of School Climate and Youth Access to Services.” This study provides in-depth assessment of HIV and STD prevention efforts in three local education agencies funded by CDC’s Division of Adolescent and School Health to the Office of Management and Budget (OMB) for review and approval. CDC previously published a “Proposed Data Collection Submitted for Public Comment and Recommendations” notice on August 17, 2017, to obtain comments from the public and affected agencies. CDC received four comments related to the previous notice. This notice serves to