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Lead, Information Collection Review Office, Office of Scientific Integrity, Office of Science, Centers for Disease Control and Prevention. [FR Doc. 2022–27504 Filed 12–19–22; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day-23-1185]

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 "Youth Outreach Generic Clearance for the National Center for Health Statistics (NCGS)" 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 September 23, 2022 to obtain comments from the public and affected agencies. CDC did not receive any comments related to the previous notice. This notice serves to allow an additional 30 days for public and affected agency comments.

CDC will accept all comments for this proposed information collection project. The Office of Management and Budget is particularly interested in comments that:

(a) 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:

(b) Evaluate the accuracy of the agencies estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used;

(c) Enhance the quality, utility, and clarity of the information to be collected;

(d) 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 submission of responses; and

(e) Assess information collection costs.

To request additional information on the proposed project or to obtain a copy of the information collection plan and instruments, call (404) 639-7570. Comments and recommendations for the proposed information collection should be sent within 30 days of publication of this notice to www.reginfo.gov/public/ do/PRAMain. Find this particular information collection by selecting "Currently under 30-day Review—Open for Public Comments" or by using the search function. Direct written comments and/or suggestions regarding the items contained in this notice to the Attention: CDC Desk Officer, Office of Management and Budget, 725 17th Street, NW, Washington, DC 20503 or by fax to (202) 395-5806. Provide written comments within 30 days of notice publication.

Proposed Project

Youth Outreach Generic Clearance for the National Center for Health Statistics (NCHS) (OMB Control No. 0920–1185, Exp. 07/31/2023)—Extension—National Center for Health Statistics (NCHS), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

The National Center for Health Statistics (NCHS) is authorized to collect data under Section 306 of the Public Health Service Act (42 U.S.C. 242k). NCHS has a history of reaching out to young people to encourage their interest in Science, Technology, Engineering and Math (STEM).

Examples of past involvement include adopting local schools, speaking at local colleges, conducting a Statistics Day for high school students, and, most recently, conducting the NCHS Data Detectives Camp for middle school students.

The success of these programs has inspired NCHS leadership and staff to want to look for new and continuing opportunities to positively impact the lives of young people and expand their interest, understanding of, and involvement in the sciences. NCHS requests approval for a New Generic Clearance mechanism to collect information for these youth outreach activities and to inform future NCHS planning activities.

As stated, these activities include hosting the Data Detectives Camp annually; hosting Statistics Day annually; creating youth poster sessions for professional conferences (such as the NCHS National Conference on Health Statistics or the American Statistical Association Conference etc.); hosting a statistical or health sciences fair or other STEM-related competitions; organizing a STEM Career Day or similar activity; developing web-based sites or materials with youth focus, as well as other programs developed to meet future youth outreach needs, particularly activities that encourage STEM.

Information will be collected using a combination of methodologies appropriate to each program. These may include: registration forms, letters of recommendation, evaluation forms, mail surveys, focus groups, automated and electronic technology (e.g. email, Webbased surveys), and telephone surveys.

OMB approval is requested for three years to conduct the Youth Outreach Generic Clearance for the National Center for Health Statistics (NCHS). CDC requests OMB approval for an estimated 1,750 annual burden hours. Participation is voluntary and there are no costs to respondents other than their time.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of survey	Respondent	Number of respondents	Number of responses/ respondent	Average burden/ response (in hours)
Questionnaires/Applications	Student/Youth	800	1	30/60
Applicants Questionnaire/Application	Parents/Guardians of Applicants	800	1	30/60
Applications, Recommendations, and Other applicant-supporting documentation.	School Officials/Community Representatives	1200	1	30/60
Focus Groups	Student/Youth; Parent/Guardian; School Officials; Other.	50	1	60/60
Other Program Surveys	Student/Youth; Parent/Guardian; School Officials; Other.	600	1	30/60

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day-23-0639]

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 Energy Employees Occupational Illness Compensation Program Act (EEOICPA) of 2000 Special Exposure Cohort Petitions 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 January 31, 2022 to obtain comments from the public and affected agencies. CDC did not receive comments related to the previous notice. This notice serves to allow an additional 30 days for public and affected agency comments.

CDC will accept all comments for this proposed information collection project. The Office of Management and Budget is particularly interested in comments that:

- (a) 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;
- (b) Evaluate the accuracy of the agencies estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used;
- (c) Enhance the quality, utility, and clarity of the information to be collected:
- (d) 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 submission of responses; and
- (e) Assess information collection costs.

To request additional information on the proposed project or to obtain a copy of the information collection plan and instruments, call (404) 639-7570. Comments and recommendations for the proposed information collection should be sent within 30 days of publication of this notice to www.reginfo.gov/public/ do/PRAMain. Find this particular information collection by selecting "Currently under 30-day Review—Open for Public Comments" or by using the search function. Direct written comments and/or suggestions regarding the items contained in this notice to the Attention: CDC Desk Officer, Office of Management and Budget, 725 17th Street NW, Washington, DC 20503 or by fax to (202) 395-5806. Provide written comments within 30 days of notice publication.

Proposed Project

Energy Employees Occupational Illness Compensation Program Act of 2000 (EEOICPA) Special Exposure Cohort Petitions. (OMB Control No. 0920–0639, Exp. 01/31/2023)— Extension—National Institute for Occupational Safety and Health (NIOSH), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

On October 30, 2000, the Energy **Employees Occupational Illness** Compensation Program Act of 2000 (EEOICPA), 42 U.S.C. 7384-7385 [1994, supp. 2001] was enacted. The Act established a compensation program to provide a lump sum payment of \$150,000 and medical benefits as compensation to covered employees suffering from designated illnesses incurred as a result of their exposure to radiation, beryllium, or silica while in the performance of duty for the Department of Energy and certain of its vendors, contractors and subcontractors. This legislation also provided for payment of compensation for certain survivors of these covered employees. This program has been mandated to be in effect until Congress ends the funding.

Among other duties, the Department of Health and Human Services (HHS) was directed to establish and implement procedures for considering petitions by classes of nuclear weapons workers to be added to the "Special Exposure Cohort" (the "Cohort"). In brief, EEOICPA authorizes HHS to designate such classes of employees for addition to the Cohort when NIOSH lacks sufficient information to estimate with sufficient accuracy the radiation doses of the employees, and if HHS also finds that the health of members of the class

may have been endangered by the radiation dose the class potentially incurred. HHS must also obtain the advice of the Advisory Board on Radiation and Worker Health (the "Board") in establishing such findings. On May 28, 2004, HHS issued a rule that established procedures for adding such classes to the Cohort (42 CFR part 83). The rule was amended on July 10, 2007.

The HHS rule authorizes a variety of respondents to submit petitions. Petitioners are required to provide the information specified in the rule to qualify their petitions for a complete evaluation by HHS and the Board. HHS has developed two forms to assist the petitioners in providing this required information efficiently and completely. Form A is a one-page form to be used by EEOICPA claimants for whom NIOSH has attempted to conduct dose reconstructions and has determined that available information is not sufficient to complete the dose reconstruction. Form B, accompanied by separate instructions, is intended for all other petitioners. Forms A and B can be submitted electronically as well as in hard copy. Respondent/petitioners should be aware that HHS is not requiring respondents to use the forms. Respondents can choose to submit petitions as letters or in other formats, but petitions must meet the informational requirements stated in the rule. NIOSH expects, however, that all petitioners for whom Form A would be appropriate will actually use the form, since NIOSH will provide it to them upon determining that their dose reconstruction cannot be completed and encourage them to submit the petition. NIOSH expects the large majority of petitioners for whom Form B would be appropriate will also use the form, since it provides a simple, organized format for addressing the informational requirements of a petition.

NIOSH will use the information obtained through the petition for the following purposes: (a) identify the petitioner(s), obtain their contact information, and establish that the petitioner(s) is qualified and intends to petition HHS; (b) establish an initial definition of the class of employees being proposed to be considered for addition to the Cohort; (c) determine whether there is justification to require HHS to evaluate whether or not to designate the proposed class as an addition to the Cohort (such an evaluation involves potentially extensive data collection, analysis, and related deliberations by NIOSH, the Board, and HHS); and (d) target an evaluation by HHS to examine relevant