Pursuant to the May 9, 2023, Proclamation, and the President's revocation of the vaccination requirements contained in Proclamation 10294, CDC has reviewed its Amended Order Implementing Presidential Proclamation on Safe Resumption of Global Travel During the COVID-19 Pandemic and has determined that termination of this Amended Order is warranted. CDC's Amended Order, which implemented Proclamation 10294's vaccination requirements, is terminated and no longer remains in effect as of 12:01 a.m. eastern daylight time on May 12, 2023.

This means that as of 12:01 a.m. eastern daylight time on May 12, 2023, noncitizen, nonimmigrant air passengers no longer need to show proof of being fully vaccinated with an accepted COVID–19 vaccine to board a flight to the United States.¹

Kathryn L. Wolff,

Chief of Staff, Centers for Disease Control and Prevention.

[FR Doc. 2023–10276 Filed 5–10–23; 11:15 am]

BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day-23-22DI; Docket No. CDC-2023-0036]

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 information collection, as required by the Paperwork Reduction Act of 1995. This notice invites comment on a proposed information collection project titled Noise Exposures and Hearing Loss in the Oil and Gas Extraction Industry. This information collection is designed to evaluate oil and gas extraction workers' noise and chemical exposures and hearing.

DATES: CDC must receive written comments on or before July 11, 2023.

ADDRESSES: You may submit comments, identified by Docket No. CDC-2023-0036 by either of the following methods:

- Federal eRulemaking Portal: www.regulations.gov. Follow the instructions for submitting comments.
- Mail: Jeffrey M. Zirger, Information Collection Review Office, Centers for Disease Control and Prevention, 1600 Clifton Road NE, MS H21–8, Atlanta, Georgia 30329.

Instructions: All submissions received must include the agency name and Docket Number. CDC will post, without change, all relevant comments to www.regulations.gov.

Please note: Submit all comments through the Federal eRulemaking portal (www.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 Jeffrey M. Zirger, Information Collection Review Office, Centers for Disease Control and Prevention, 1600 Clifton Road NE, MS H21–8, 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:
- 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; and
 - 5. Assess information collection costs.

Proposed Project

Noise Exposures and Hearing Loss in the Oil and Gas Extraction Industry— New—National Institute for Occupational Safety and Health (NIOSH), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

Oil and gas extraction (OGE) workers play an important role in supporting the United States economy and help fulfill the energy needs of Americans and American businesses. OGE workers have significant risks for a variety of exposures at oil and gas well sites, and there has been no significant occupational noise exposure research in the United States onshore upstream OGE sector. This proposed project will characterize relationships between noise exposure, chemical exposures, hearing loss, and hearing loss prevention practices within the onshore OGE industry.

Primary data will be collected using three approaches. First, researchers will collect direct measurements of noise and ototoxic chemicals on job sites, including personal exposure assessments of OGE workers. Second, researchers will use a questionnaire to collect information on noise and chemical exposures, hearing loss, and associated factors among OGE workers. Third, audiometry tests performed by NIOSH will be offered to industry partners to further understand extent of hearing loss amongst OGE workers.

Data will be used to understand noise exposures, ototoxic chemical exposures, self-reported hearing loss, and hearing loss prevention practices in the OGE industry. Subsequently, the data and

¹ This Notice, like CDC's April 2022 Amended Order that no longer will be in effect as of 12:01 a.m. eastern daylight time on May 12, is not a substantive rule within the meaning of the Administrative Procedure Act (APA) because it implements the President's revocation of the vaccination requirements contained in the October 2021 Proclamation (which in turn was the basis for the CDC's Amended Order). In any event, the APA's requirement of a 30-day delay in the effective date of certain "substantive rule[s]," 5 U.S.C. 553(d), would not apply to this notice, as this notice "relieves a restriction" contained in the Amended Order, id. Section 553(d)(1). Independently, were the APA applicable, CDC finds good cause for its termination of the April 2022 Amended Order to take effect at 12:01 a.m. on May 12, 2023, which coincides with the end of the COVID-19 public health emergency, given the latest public health conditions and the latest guidance from public health experts. See 5 U.S.C. 553(b), (d).

analysis will be used to create evidencebased interventions and recommendations, which will be communicated to the spectrum of OGE industry stakeholders.

CDC requests OMB approval for an estimated 65 annual burden hours.

There is no cost to respondents other than their time to participate.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondents	Form name	Number of respondents	Number of responses per respondent	Average burden per response (in hours)	Total burden (in hours)
Oil and gas workers	Noise and Hearing Questionnaire Audiometry Testing Exposure Monitoring Results Notification Form.	167 33 40	1 1 1	17/60 30/60 2/60	47 17 1
Total					65

Jeffrey M. Zirger,

Lead, Information Collection Review Office, Office of Public Health Ethics and Regulations, Office of Science, Centers for Disease Control and Prevention.

[FR Doc. 2023–10188 Filed 5–11–23; 8:45 am] BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day-23-1243]

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 "Rapid Response Suicide Investigation Data Collection' 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 October 28, 2022 to obtain comments from the public and affected agencies. There were no comments to the 60-day Federal Register 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

 (\bar{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

Rapid Response Suicide Investigation Data Collection (OMB Control No. 0920–1243, Exp. 5/31/2023)— Extension—National Center for Injury Prevention and Control (NCIPC), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

CDC is frequently called upon to respond to urgent requests from one or more external partners (e.g., local, State, Territory, and Tribal health authorities; other Federal agencies; local and State leaders; schools; or other partner organizations) to conduct investigations of suicide. Supporting rapid investigations to inform the implementation of effective suicide prevention strategies is one of the most important ways CDC can serve to protect and promote the health of the public.

Rapid Response Suicide Investigation Data Collections are specifically designed to inform the implementation of prevention strategies in a state, county, community, or vulnerable population where a possible suicide cluster or increasing trend has been observed. This Generic Clearance will not be used to conduct research studies or to collect data designed to draw conclusions about the United States or areas beyond the defined geographic location or vulnerable population that is the focus of the investigation. CDC, in collaboration with external partners, will identify the respondent universe for each Rapid Response Suicide Investigation Data Collection. The respondent universe will be determined based on the information needed to understand potential suicide clusters, significant increases in suicidal behavior and suicide, risk and protective factors, and vulnerable populations, in order to inform the implementation of suicide prevention strategies.

CDC requests OMB approval for an estimated 1,000 annual burden hours. There are no costs to respondents other than their time to participate.