(4) Inform continuous quality improvement activities over the course of the funding period, to include quarterly and annual strategic planning for current and later iterations of this program under future funding. CDC requests OMB approval for an estimated 655 annual burden hours.

### ESTIMATED ANNUALIZED BURDEN HOURS

<table>
<thead>
<tr>
<th>Type of respondents</th>
<th>Form name</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Average burden per response (in hours)</th>
<th>Total burden (in hours)</th>
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<td><strong>Total</strong></td>
<td></td>
<td></td>
<td></td>
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<td><strong>655</strong></td>
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</tbody>
</table>

Jeffrey M. Zirger,

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day–21–21BL]

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 “Evaluation of the Overdose Data to Action Technical Assistance Hub” 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 December 1, 2020 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

Evaluation of the Overdose Data to Action Technical Assistance Hub—New—National Center for Injury Prevention and Control (NCIPC), Centers for Disease Control and Prevention (CDC)

Background and Brief Description

The Division of Overdose Prevention (DOP), at Centers for Disease Control and Prevention (CDC) requests a three-year OMB approval to support the evaluation of technical assistance (TA) provided for the Overdose Data to Action (OD2A) program. OD2A is a cooperative agreement (CDC–RFA–CE19–1904) funded in 2019 to focus on comprehensive and interdisciplinary opioid overdose prevention efforts in 47 state health departments, 16 localities, Puerto Rico, Washington DC, and the North Mariana Islands. This program consists of two required components—a surveillance component and a prevention component. OD2A recipients implement a combination of activities across 10 strategies within these components in order to gain access to high quality, complete, and timelier data on opioid prescribing and overdoses and to use those data to inform prevention and response efforts in their jurisdictions.

Training and technical assistance (TA) is essential to building knowledge and strengthening the capacity of recipients to implement and evaluate OD2A program strategies. CDC will develop and deploy a TA hub (hereafter referred to as the OD2A TA Center) to deliver comprehensive technical assistance and training to support the successful implementation and
evaluation of surveillance and prevention activities. The OD2A TA Center is designed to enhance the efficiency, coordination, and effectiveness of TA efforts by streamlining and centralizing the provision of overdose surveillance and prevention TA. TA to OD2A recipients is divided into four different levels with multiple modes of TA delivery and involves a wide range of TA providers including CDC staff, internal and external subject matter experts (SMEs), and program partners, as well as contract staff.

The evaluation consists of two web-based surveys designed to collect process and outcome measures about TA access, utilization, and outcomes across all 66 OD2A recipient programs. The Technical Assistance Feedback Form will be administered to collect immediate feedback following individual TA encounters and group events such as webinars and in-person trainings. The Annual OD2A TA Survey will be distributed twice (mid-point and final) to assess satisfaction with overall TA provided and the extent to which TA supports implementation of OD2A strategies. The information obtained through this evaluation will allow TA providers to assess OD2A recipients’ experience and utility of knowledge and resources gained through individual TA support, peer-to-peer sessions, and other group trainings. Ultimately, the evaluation data will inform subsequent rounds of TA and allow TA providers to make necessary adjustments to the overall TA strategy for continuous quality improvement.

This will ensure recipients have the support necessary to implement strategies that will improve opioid surveillance and prevention policies and practices within their communities.

The total annualized burden estimate is 222 hours. There is no cost to respondents other than the time to participate.

**ESTIMATED ANNUALIZED BURDEN HOURS**

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<td>Email invitation and reminders for OD2A Survey</td>
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</table>

Jeffrey M. Zirger,
Lead, Information Collection Review Office,
Office of Scientific Integrity, Office of Science, Centers for Disease Control and Prevention.
[FR Doc. 2021–12208 Filed 6–9–21; 8:45 am]
BILLING CODE 4163–18–P

**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

Food and Drug Administration [Docket No. FDA–2018–N–1129]

Agency Information Collection Activities; Submission for Office of Management and Budget Review; Comment Request; National Agriculture and Food Defense Strategy Survey

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing that a proposed collection of information has been submitted to the Office of Management and Budget (OMB) for review and clearance under the Paperwork Reduction Act of 1995.

DATES: Submit written comments (including recommendations) on the collection of information by July 12, 2021.

ADDRESSES: To ensure that comments on the information collection are received, OMB recommends that written comments be submitted to https://www.reginfo.gov/public/do/PRAMain. Find this particular information collection by selecting “Currently under Review—Open for Public Comments” or by using the search function. The OMB control number for this information collection is 0910–0855. Also include the FDA docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT: Ila S. Mizrachi, Office of Operations, Food and Drug Administration, Three White Flint North, 10A–12M, 11601 Landsdown St., North Bethesda, MD 20852, 301–796–7726, PRASstaff@ fda.hhs.gov.

SUPPLEMENTARY INFORMATION: In compliance with 44 U.S.C. 3507, FDA has submitted the following proposed collection of information to OMB for review and clearance.

**National Agriculture and Food Defense Strategy Survey**

**OMB Control Number 0910–0855—Extension**

We are seeking OMB approval of the National Agriculture and Food Defense Strategy (NAFDS) under section 108 of the FDA Food and Safety Modernization Act (FSMA). This is a voluntary survey of State, local, territorial, and/or tribal (SLTT) governments intended to gauge government activities in food and agriculture defense from intentional contamination and emerging threats. The collected information will be included in the mandatory NAFDS followup Report to Congress. The authority for us to collect the information derives from the Commissioner of Food and Drugs’ authority provided in section 1003(d)(2)(C) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 393(d)(2)(C)).

Protecting the nation’s food and agriculture supply against intentional contamination and other emerging threats is an important responsibility shared by SLTT governments as well as private sector partners. On January 4, 2011, the President signed into law FSMA. FSMA focuses on ensuring the safety of the U.S. food supply by shifting the efforts of Federal regulators from response to prevention and recognizes the importance of strengthening existing collaboration among all stakeholders to achieve common public health and security goals. FSMA identifies some key priorities for working with partners in areas such as reliance on Federal, State, and local agencies for inspections; improving foodborne illness surveillance; and leveraging and enhancing State and local food safety and defense capacities. Section 108 of FSMA (NAFDS) requires the Department of Health and Human Services (HHS) and the U.S. Department of Agriculture (USDA), in coordination