### 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 respondent (in hours)</th>
<th>Total annual burden hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>State, tribal, local, territorial health departments.</td>
<td>Component 1 Forms: NWSS Data Dictionary v5.0.0; CDC Seq Manifest Data Dictionary; BioSample WW Template v1.9; SRA Template v5.7 NWSS; NCBI DCIPHER Crosswalk Data Dictionary; NWSS DCIPHER Wastewater Data CSV Upload Template; Component 1–2–3 NWSS DCIPHER CSV Bulk Upload Tool.</td>
<td>55</td>
<td>2,080</td>
<td>139/60</td>
<td>265,026</td>
</tr>
<tr>
<td>Private laboratory</td>
<td>Component 1 Forms: Component; NWSS Data Dictionary v5.0.0; CDC Seq Manifest Data Dictionary; BioSample WW Template v1.9; SRA WW Template v5.7; NCBI DCIPHER Crosswalk Data Dictionary; NWSS DCIPHER Wastewater Data CSV Upload Template v3; Component 1–2–3 NWSS DCIPHER CSV Bulk Upload Tool.</td>
<td>1</td>
<td>52,000</td>
<td>139/60</td>
<td>120,467</td>
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<td>State, tribal, local, territorial health departments.</td>
<td>Component 2 Forms: Sewershed Spatial Files (No Form); Component 1–2–3 NWSS DCIPHER CSV Bulk Upload Tool.</td>
<td>55</td>
<td>20</td>
<td>5/60</td>
<td>92</td>
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<td>Wastewater utilities</td>
<td>Component 2 Forms: Sewershed Spatial Files (No Form); Component 1–2–3 NWSS DCIPHER CSV Bulk Upload Tool.</td>
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<td>1</td>
<td>2</td>
<td>2,200</td>
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<td>State, tribal, local, territorial health departments.</td>
<td>Component 3 Forms: NWSS COVID Case Data Dictionary; NWSS DCIPHER Case Data CSV Upload Template; NWSS DCIPHER Sewershed Name Crosswalk CSV Upload Template; Component 1–2–3 NWSS DCIPHER CSV Bulk Upload Tool.</td>
<td>55</td>
<td>39,977</td>
<td>5/60</td>
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<td>Total</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>571,013</td>
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</tbody>
</table>

Jeffrey M. Zirger,  

BILLING CODE 4163–18–P

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day–23–1204]

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 “Behavioral Risk Factor Surveillance System (BRFSS) Asthma Call-back Survey (ACBS)” 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 March 6, 2023 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 “http://www.reginfo.gov/public/doPRAMain”. 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

Behavioral Risk Factor Surveillance System (BRFSS) Asthma Call-back Survey (ACBS) (OMB Control No. 0920–1204, Expiration Date 11/30/2023)—Revision—National Center for Environmental Health (NCEH), Centers for Disease Control and Prevention (CDC).
Background and Brief Description

The CDC’s National Center for Environmental Health (NCEH) is requesting OMB approval for three years to revise and continue the “Behavioral Risk Factor Surveillance System (BRFSS) Asthma Call-back Survey (ACBS)” (OMB Control No. 0920–1204, Expiration Date 11/30/2023). The ACBS is funded by the NCEH National Asthma Control Program (NACP) in the Asthma and Community Health Branch (ACHB) and is administered on behalf of NCEH by the CDC’s National Center for Chronic Disease Prevention and Health Promotion (NCCDPHP) BRFSS Program. BRFSS coordinators in the health departments in U.S. states, territories, and the District of Columbia (collectively referred to as “states” and “jurisdictions”) are responsible for administering both the BRFSS and the ACBS.

The BRFSS (OMB Control No. 0920–1061, expiration date 12/31/2024) is a nationwide system of customized, cross-sectional telephone health surveys. The BRFSS information collection is conducted in a continuous, three-part telephone interview process: screening, participation in a common BRFSS core survey, and participation in optional question modules that states use to customize survey content. The ACBS is a follow-up survey conducted in households that include an individual who has been diagnosed with asthma. The ACBS is conducted within two days after the BRFSS survey. The purpose of ACBS is to gather state-level asthma data and to make them available to track the burden of the disease, to monitor adherence to asthma guidelines, and to direct and evaluate interventions undertaken by asthma control programs located in state health departments. Federal agencies and other entities also rely on this critical information for planning and evaluating efforts to reduce the burden of this disease. The CDC makes annual ACBS datasets available for public use and provides guidance on statistically appropriate uses of the data.

In response to the contours of 2020 Terms of Clearance, the annual joint response rates from BRFSS and ACBS were reported with the BRFSS annual dataset. To communicate the caveats of state-to-state comparisons, the ACBS nonresponse bias and impact on prevalence estimation are being analyzed and reported as appendix tables of the annual data quality report released with the public use dataset for adult and child participants (https://www.cdc.gov/brfss/acbs/2020/pdf/sdq_report_acbs_20-508.pdf). The first table reports unweighted and weighted demographic distribution percentages for each participating state based on BRFSS-eligible asthma respondents, non-respondents to the ACBS, and ACBS final completes. The second table reports estimated current asthma percentage among individuals who have ever been diagnosed with asthma. These two tables will help communicate the potential impact of nonresponse bias on the ACBS published dataset.

To provide clear communication about the caveats of state-to-state comparisons, the NACP revised the tables of prevalence estimates for asthma risk factors based on ACBS and reduced the number of risk factors prevalence tables from 20 to 13. The NACP also deleted the tables (active asthma related risk factors) that didn’t provide enough information for state-to-state comparisons. A footnote with a hyperlink to the nonresponse report has been incorporated into the footnote for annual ACBS risk factors prevalence tables which can be viewed at: https://www.cdc.gov/brfss/acbs/2019_tables_LLCP.html.

In addition, and also in response to the 2020 Terms of Clearance, the NACP undertook efforts to streamline the ACBS, reduce unnecessary burden, and ensure that the question wording is aligned with more recent studies. The questionnaires were re-evaluated by ACBS questionnaire working groups and the ACBS recipients. Beginning in 2024, the NACP proposes to delete 6 questions from the adult’s questionnaire and 17 questions from child’s questionnaire. The adult’s questionnaire will include nine new questions and the child’s questionnaire will include 10 new questions. The estimated time burden for the interview will remain unchanged from that of the 2021 questionnaire (10 minutes per response). There are no proposed changes to the number of responses per respondent.

The total BRFSS sample size decreased from 476,217 in 2016 to 393,474 in 2020, and as a result of decreasing BRFSS sample size, the number of individuals eligible for the ACBS decreased from 46,100 to 41,444 during that period. The NACP proposes the following changes to the burden estimation from 2021 (based on 2016 ACBS response data) to 2024 (based on 2020 response data):

- The total estimated number of responses is 57,852, which is a decrease of 10,074 from the previously approved 68,846. The total estimated annualized time burden is 6,073 hours, which is a decrease of 542 hours from the previously approved 6,615 hours.
- Participation in the ACBS is voluntary and there are no costs to respondents other than their time.
DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Medicare & Medicaid Services

[Document Identifier: CMS–10488, CMS–10708 and CMS–10846]

Agency Information Collection Activities: Submission for OMB Review; Comment Request

AGENCY: Centers for Medicare & Medicaid Services, Health and Human Services (HHS).

ACTION: Notice.

SUMMARY: The Centers for Medicare & Medicaid Services (CMS) is announcing an opportunity for the public to comment on CMS’ intention to collect information from the public. Under the Paperwork Reduction Act of 1995 (PRA), Federal agencies are required to publish notice in the Federal Register concerning each proposed collection of information, including each proposed extension or reinstatement of an existing collection of information, and to allow a second opportunity for public comment on the notice. Interested persons are invited to send comments regarding the burden estimate or any other aspect of this collection of information, including the necessity and utility of the proposed information collection for the proper performance of the agency’s functions, the accuracy of the estimated burden, ways to enhance the quality, utility, and clarity of the information to be collected, and the use of automated collection techniques or other forms of information technology to minimize the information collection burden.

DATES: Comments on the collection(s) of information must be received by the OMB desk officer by August 7, 2023.

ADDRESSES: Written 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” by using the search function.

To obtain copies of a supporting statement and any related forms for the proposed collection(s) summarized in this notice, please access the CMS PRA website by copying and pasting the following web address into your web browser: https://www.cms.gov/Regulations-and-Guidance/Legislation/PaperworkReductionActof1995/PRA-Listing.

FOR FURTHER INFORMATION CONTACT: William Parham at (410) 786–4669.

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. The term “collection of information” is defined in 44 U.S.C. 3502(3) and 5 CFR 1320.3(c) and includes agency requests or requirements that members of the public submit reports, keep records, or provide information to a third party. Section 3506(c)(2)(A) of the PRA (44 U.S.C. 3506(c)(2)(A)) requires Federal agencies to publish a 30-day notice in the Federal Register concerning each proposed collection of information, including each proposed extension or reinstatement of an existing collection of information, before submitting the collection to OMB for approval. To comply with this requirement, CMS is publishing this notice that summarizes the following proposed collection(s) of information for public comment:

1. Type of Information Collection Request: Revision of currently approved collection; Title of Information Collection: Consumer Experience Survey Data Collection; Use: Section 1311(c)(4) of the Affordable Care Act requires the Department of Health and Human Services (HHS) to develop an enrollee satisfaction survey system that assesses consumer experience with qualified health plans (QHPs) offered through an Exchange. It also requires public display of enrollee satisfaction information by the Exchange to allow individuals to easily compare enrollee satisfaction levels between comparable plans. HHS established the QHP Enrollee Experience Survey (QHP Enrollee Survey) to assess consumer experience with the QHPs offered through the Marketplaces. The survey includes topics to assess consumer experience with the health care system such as communication skills of providers and ease of access to health care services.

CMS developed the survey using the Consumer Assessment of Health Providers and Systems (CAHPS®) principles (https://www.ahrq.gov/cahps/about-cahps/principles/index.html) and established an application and approval process for survey vendors who want to participate in collecting QHP enrollee experience data. The QHP Enrollee Survey, which is based on the CAHPS® Health Plan Survey, will be used to (1) help consumers choose among competing health plans, (2) provide actionable information that the QHPs can use to improve performance, (3) provide information that regulatory and accreditation organizations can use to regulate and accredit plans, and (4) provide a longitudinal database for consumer research. CMS completed two rounds of developmental testing including 2014 psychometric testing and 2015 beta testing of the QHP Enrollee Survey.

The psychometric testing helped determine psychometric properties and...