ANNUAL BURDEN ESTIMATES—Continued

<table>
<thead>
<tr>
<th>Information collection instrument</th>
<th>Total estimated number of respondents</th>
<th>Total number of responses per respondent</th>
<th>Average burden hours per response</th>
<th>Total burden hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>e-IWO NPO Profile</td>
<td>46</td>
<td>1</td>
<td>0.22</td>
<td>10.12</td>
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<tr>
<td>MSFI–FAST Levy Profile</td>
<td>5</td>
<td>1</td>
<td>0.08</td>
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<tr>
<td>Portal Registration Screens</td>
<td>1,254</td>
<td>1</td>
<td>0.15</td>
<td>188.10</td>
</tr>
</tbody>
</table>

Estimated Total Annual Burden Hours: 964.88.

Comments: The Department specifically requests comments on (a) whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have practical utility; (b) the accuracy of the agency’s estimate of the burden of the proposed collection of information; (c) the quality, utility, and clarity of the information to be collected; and (d) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques or other forms of information technology. Consideration will be given to comments and suggestions submitted within 60 days of this publication.

Authority: 42 U.S.C. 653(m)(2) and 44 U.S.C. 3554.

Mary B. Jones, ACF/OPRE Certifying Officer.

BILLCODE 4184–41–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Health Resources and Services Administration

Agency Information Collection Activities: Proposed Collection: Public Comment Request, Information Collection Request Title: Rural Health Care Services Outreach Program Performance Improvement and Measurement Systems (PIMS) Measures, OMB No. 0906–0009, Revision

AGENCY: Health Resources and Services Administration (HRSA), Department of Health and Human Services.

ACTION: Notice.

SUMMARY: In compliance with the requirement for opportunity for public comment on proposed data collection projects of the Paperwork Reduction Act of 1995, HRSA announces plans to submit an Information Collection Request (ICR), described below, to the Office of Management and Budget (OMB). Prior to submitting the ICR to OMB, HRSA seeks comments from the public regarding the burden estimate, below, or any other aspect of the ICR.

DATES: Comments on this ICR must be received no later than September 20, 2021.

ADDRESSES: Submit your comments to paperwork@hrsa.gov or mail the HRSA Information Collection Clearance Officer, Room 14N136B, 5600 Fishers Lane, Rockville, MD 20857.

FOR FURTHER INFORMATION CONTACT: To request more information on the proposed project or to obtain a copy of the data collection plans and draft instruments, email paperwork@hrsa.gov or call Lisa Wright-Solomon, the HRSA Information Collection Clearance Officer at (301) 443–1984.

SUPPLEMENTARY INFORMATION: When submitting comments or requesting information, please include the information request collection title for reference.

Information Collection Request Title: Rural Health Care Services Outreach Program PIMS Measures OMB No. 0906–0009—Revision

Abstract: The Rural Health Care Services Outreach (Outreach) Program is authorized by Section 330A(e) of the Public Health Service Act (42 U.S.C. 254c(e)) to “promote rural health care services outreach by improving and expanding the delivery of health care services to include new and enhanced services in rural areas, through community engagement and evidence-based or innovative, evidence-informed models.” The goals for the Outreach Program are as follows: (1) Expand the delivery of health care services to include new and enhanced services exclusively in rural communities, (2) deliver health care services through a strong consortium, in which every consortium member organization is actively involved and engaged in the planning and delivery of services, (3) utilize community engagement and evidence-based or innovative, evidence-informed model(s) in the delivery of health care services, and (4) improve population health, and demonstrate health outcomes and sustainability.

Need and Proposed Use of the Information: The PIMS measures for the Outreach Program enable HRSA and the Federal Office of Rural Health Policy to capture awardee-level and aggregate data that illustrate the impact and scope of federal funding. The collection of this information helps further inform and substantiate the focus and objectives of the grant program. The measures encompass the following topics: (a) Access to care, (b) population demographics, (c) consortium/network, (d) sustainability, and (e) project specific domains.

The proposed Outreach PIMS measures reflect an increase in the number of measures including the following: (1) The addition of project-specific measures related to the Healthy Rural Hometown Initiative (includes 17 required and 20 optional measures for a total of 37 additional measures) applicable only to Outreach awardees who apply to be part of the Healthy Rural Hometown Initiative track (anticipated total of 16 out of 61 awardees) to focus on one or more of the five causes of excess death in rural communities (heart disease, cancer, unintentional injury/substance use, chronic lower respiratory disease, and stroke); (2) addition of project-specific measures (3 additional measures) only applicable to Outreach Awardees with a focus on telehealth (anticipated total of 15 out of 61 awardees); (3) the addition of social determinants of health measures (3 additional measures) only applicable to Outreach Awardees addressing social determinants of health as part of their grant funded activities (anticipated total of 15 out of 61 awardees); (4) the consolidation of the access to care measures from singular to composite measure format (currently 14, previously 16) applicable to all awardees (anticipated total of 61 awardees); (5) removal of an outdated project specific measure (1 measure removed) applicable to awardees focused on childhood obesity; (6) removal of an outdated project specific applicable to awardees...
HRSA specifically requests comments on the: (1) Necessity and utility of the proposed information collection for the proper performance of the agency’s functions; (2) accuracy of the estimated burden; (3) ways to enhance the quality, utility, and clarity of the information to be collected; and (4) use of automated collection techniques or other forms of information technology to minimize the information collection burden.

Maria G. Button, Director, Executive Secretariat.

[FR Doc. 2021–15607 Filed 7–21–21; 8:45 am]

BILLING CODE 4165–15–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Health Resources and Services Administration

Agency Information Collection Activities: Proposed Collection: Public Comment Request; Information Collection Request Title: The Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment Surveys, OMB No. 0906–0014, Revision

AGENCY: Health Resources and Services Administration (HRSA), Department of Health and Human Services.

ACTION: Notice.

SUMMARY: In compliance with the requirement for opportunity for public comment on proposed data collection projects of the Paperwork Reduction Act of 1995, HRSA announces plans to submit an Information Collection Request (ICR), described below, to the Office of Management and Budget (OMB). Prior to submitting the ICR to OMB, HRSA seeks comments from the public regarding the burden estimate, below, or any other aspect of the ICR.

DATES: Comments on this ICR should be received no later than September 20, 2021.

ADDRESSES: Submit your comments to paperwork@hrsa.gov or mail the HRSA Information Collection Clearance Officer, Room 14N136B, 5600 Fishers Lane, Rockville, MD 20857.

FOR FURTHER INFORMATION CONTACT: To request more information on the proposed project or to obtain a copy of the data collection plans and draft instruments, email paperwork@hrsa.gov or call Lisa Wright-Solomon, the HRSA Information Collection Clearance Officer at (301) 443–1984.

SUPPLEMENTARY INFORMATION: When submitting comments or requesting information, please include the information request collection title for reference.

Information Collection Request Title: The Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment Surveys, OMB No. 0906–0014—Revision

Abstract: The purpose of the Public Health System Assessment Surveys is to inform the Advisory Committee on Heritable Disorders in Newborns and Children (Committee) on states’ ability to add newborn screening for particular conditions, including the feasibility, readiness, and overall capacity to screen for a new condition.

The Committee was established under the Public Health Service Act, 42 U.S.C. 217a: Advisory councils or committees, and Title XI § 1111 (42 U.S.C. 300b-10).

The purpose of the Committee is to provide the Secretary with recommendations, advice, and technical information regarding the most appropriate application of technologies, policies, guidelines, and standards for:

(a) Effectively reducing morbidity and mortality in newborns and children having, or at risk for, heritable disorders;

(b) Enhancing the ability of state and local health agencies to provide for newborn and child screening, counseling, and health care services for newborns and children having, or at risk for, heritable disorders.

Specifically, the Committee makes systematic evidence-based recommendations on newborn screening for conditions that have the potential to change the health outcomes for newborns.

The Committee tasks an external workgroup to conduct systematic evidence-based reviews for conditions being considered for addition to the Recommended Uniform Screening Panel, and their corresponding newborn screening test(s), confirmatory test(s), and treatment(s). Reviews also include an analysis of the benefits and harms of newborn screening for a selected condition at a population level and an assessment of state public health newborn screening programs’ ability to implement the screening of a new condition.

Need and Proposed Use of the Information: The Committee’s Evidence Review Group administers the surveys to collect data from state newborn screening programs in the United States.