providing clinical services (currently 7, previously 8) related to Healthy People 2020 and:
(7) removal of the outdated project specific Health Improvement Special Project measure (1 measure removed).

In total, the proposed changes reflect the addition of 43 measures and the removal of 5 measures for an increase in measures by a total of 38 measures. Of these measures, 17 are required and 26 are optional. All additional measures proposed are project specific (only applicable to anticipated total ranging from 15–16 out of 61 awardees). All measures will not be applicable to all 61 respondents. Project specific measures will remain applicable only to Outreach Awardees focusing on the respective project specific topic.

Likely Respondents: The respondents would be award recipients of the Rural Health Care Services Outreach Program.

Burden Statement: Burden in this context means the time expended by persons to generate, maintain, retain, disclose or provide the information requested. This includes the time needed to review instructions; develop, acquire, install, and utilize technology and systems for the purpose of collecting, validating, and verifying information, processing and maintaining information, and disclosing and providing information; train personnel and to be able to respond to a collection of information; to search data sources; complete and review the collection of information; and transmit or otherwise disclose the information. The total annual burden hours estimated for this ICR are summarized in the table below.

### Total Estimated Annualized Burden Hours:

<table>
<thead>
<tr>
<th>Form name</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Total responses</th>
<th>Average burden per response (in hours)</th>
<th>Total burden hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rural Health Care Services Outreach PIMS</td>
<td>61</td>
<td>1</td>
<td>61</td>
<td>3.5</td>
<td>213.5</td>
</tr>
<tr>
<td></td>
<td>61</td>
<td></td>
<td>61</td>
<td></td>
<td>213.5</td>
</tr>
</tbody>
</table>

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.

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The surveys have been developed to capture the following: (1) Readiness of state public health newborn screening programs to expand newborn screening to include the target condition, (2) specific requirements of screening for a condition that could hinder or facilitate implementation in each state, and (3) estimated timeframes needed for each state to complete major milestones toward full implementation of newborn screening for the condition.

The following is a summary of proposed changes to the Committee's Public Health System Assessment Surveys:

- Proposed change: (strike "Secretary’s") "INITIAL Survey of the Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:")
  - Survey title:
    - Current title: "INITIAL Survey of the Secretary’s Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
    - Proposed change: (strike "Secretary’s") "INITIAL Survey of the Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:")
  - Rationale: Per the charter signed November 10, 2020, the Advisory Committee on Heritable Disorders in Newborns and Children is the correct name of the Committee.

- Proposed change: (strike "Secretary’s") "INITIAL Survey of the Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
  - Survey title:
    - Current title: "INITIAL Survey of the Secretary’s Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
    - Proposed change: (strike "Secretary’s") "INITIAL Survey of the Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
  - Rationale: Change made to correct grammatical error.

- Proposed change: (strike "Secretary’s") "INITIAL Survey of the Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
  - Survey title:
    - Current title: "INITIAL Survey of the Secretary’s Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
    - Proposed change: (strike "Secretary’s") "INITIAL Survey of the Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
  - Rationale: Change made to correct grammatical error.

- Proposed change: (strike "Secretary’s") "INITIAL Survey of the Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
  - Survey title:
    - Current title: "INITIAL Survey of the Secretary’s Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
    - Proposed change: (strike "Secretary’s") "INITIAL Survey of the Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
  - Rationale: Change made to correct grammatical error.

- Proposed change: (strike "Secretary’s") "INITIAL Survey of the Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
  - Survey title:
    - Current title: "INITIAL Survey of the Secretary’s Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
    - Proposed change: (strike "Secretary’s") "INITIAL Survey of the Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
  - Rationale: Change made to correct grammatical error.

- Proposed change: (strike "Secretary’s") "INITIAL Survey of the Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
  - Survey title:
    - Current title: "INITIAL Survey of the Secretary’s Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
    - Proposed change: (strike "Secretary’s") "INITIAL Survey of the Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
  - Rationale: Change made to correct grammatical error.

- Proposed change: (strike "Secretary’s") "INITIAL Survey of the Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
  - Survey title:
    - Current title: "INITIAL Survey of the Secretary’s Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
    - Proposed change: (strike "Secretary’s") "INITIAL Survey of the Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment:"
  - Rationale: Change made to correct grammatical error.

The data gathered informs the Committee on the following: (1) Feasibility of implementing population-based screening for the target condition, (2) readiness of state newborn screening programs to adopt screening for the condition, (3) gaps or limitations related to the feasibility or readiness of states to screen for a condition, and (4) areas of technical assistance and resources needed to facilitate screening for conditions with low feasibility or readiness.

Likely Respondents: The respondents to the survey will be state and territorial newborn screening programs.

Burden Statement: Burden in this context means the time expended by persons to generate, maintain, retain, disclose or provide the information requested. This includes the time needed to review instructions; to develop, acquire, install and utilize technology and systems for the purpose of collecting, validating and verifying information, processing and maintaining information, and disclosing and providing information; to train personnel and to be able to respond to a collection of information; to search data sources; to complete and review the collection of information; and to transmit or otherwise disclose the information. The total annual burden hours estimated for this ICR are summarized in the table below.

**Total Estimated Annualized Burden Hours**

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<th>Average burden per response (in hours)</th>
<th>Total burden hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>INITIAL Survey of the Advisory Committee on Heritable Disorders in Newborns and Children's Public Health System Assessment</td>
<td>159</td>
<td>2</td>
<td>118</td>
<td>10.0</td>
<td>1,180</td>
</tr>
<tr>
<td>FOLLOW–UP Survey of the Advisory Committee on Heritable Disorders in Newborns and Children’s Public Health System Assessment</td>
<td>230</td>
<td>2</td>
<td>60</td>
<td>2.0</td>
<td>120</td>
</tr>
<tr>
<td>Total</td>
<td>89</td>
<td></td>
<td>178</td>
<td></td>
<td>1,300</td>
</tr>
</tbody>
</table>

*It is anticipated that the proposed revisions will not impact the estimated annualized burden hours.

† The respondents to the survey will be state and territorial newborn screening programs.
HRSA specifically requests comments on (1) the necessity and utility of the proposed information collection for the proper performance of the agency's functions, (2) the accuracy of the estimated burden, (3) ways to enhance the quality, utility, and clarity of the information to be collected, and (4) the 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. 2021–15598 Filed 7–21–21; 8:45 am]  
BILLING CODE 4165–15–P  

DEPARTMENT OF HEALTH AND HUMAN SERVICES  

Health Resources and Services Administration  

Meeting of the Advisory Committee on Heritable Disorders in Newborns and Children  

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

ACTION: Notice.  

SUMMARY: In accordance with the Public Health Service Act and the Federal Advisory Committee Act, this notice announces that the Advisory Committee on Heritable Disorders in Newborns and Children (ACHDNC or Committee) has scheduled a public meeting to be held on Thursday, August 12, 2021, and Friday, August 13, 2021. Information about the ACHDNC and the agenda for this meeting can be found on the ACHDNC website at https://www.hrsa.gov/advisory-committees/heritable-disorders/index.html.  

DATES: Thursday, August 12, 2021, from 10:00 a.m. to 2:15 p.m. Eastern Time (ET) and Friday, August 13, 2021, from 10:00 a.m. to 2:00 p.m. ET.  

ADDRESSES: This meeting will be held via webinar. While this meeting is open to the public, advance registration is required.  

Please register online at https://www.achdncmeetings.org/registration/ by the deadline of 12:00 p.m. ET on August 11, 2021. Instructions on how to access the meeting via webcast will be provided upon registration.  

FOR FURTHER INFORMATION CONTACT: Alaina Harris, Maternal and Child Health Bureau, HRSA, 5600 Fishers Lane, Rockville, Maryland 20857; 301–443–0721; or ACHDNC@hrsa.gov.  

SUPPLEMENTARY INFORMATION: ACHDNC provides advice and recommendations to the Secretary of HHS (Secretary) on the development of newborn screening activities, technologies, policies, guidelines, and programs for effectively reducing morbidity and mortality in newborns and children having, or at risk for, heritable disorders. The ACHDNC reviews and reports regularly on newborn and childhood screening practices, recommends improvements in the national newborn and childhood screening programs, and fulfills requirements stated in the authorizing legislation. In addition, ACHDNC’s recommendations regarding inclusion of additional conditions for screening, following adoption by the Secretary, are evidence-informed preventive health services provided for in the comprehensive guidelines supported by HRSA through the Recommended Uniform Screening Panel pursuant to section 2713 of the Public Health Service Act (42 U.S.C. 300gg–13). Under this provision, non-grandfathered group health plans and health insurance issuers offering group or individual health insurance are required to provide insurance coverage without cost-sharing (a co-payment, co-insurance, or deductible) for preventive services for plan years (i.e., policy years) beginning on or after the date that is one year from the Secretary’s adoption of the condition for screening.  

During the August 12–13, 2021, meeting, ACHDNC will hear from experts in the fields of public health, medicine, heritable disorders, rare disorders, and newborn screening. Agenda items include the following:  

(1) Overview of the Committee’s review of its evidence-review processes and proposed updates,  

(2) A presentation on phase one of the mucopolysaccharidosis type II evidence review,  

(3) Guanidinoacetate methyltransferase (GAMT) deficiency nomination summary,  

(4) Possible Committee vote on whether to move GAMT deficiency forward to a full evidence review,  

(5) Committee discussion on emerging issues for newborn screening,  

(6) A panel presentation on national registries followed by Committee discussion,  

(7) A panel presentation on emerging issues facing the newborn screening workforce followed by Committee discussion, and  

(8) Public comments on any newborn screening related topic.  

The public is also encouraged to provide public comment on the proposed updates to the Committee’s evidence review processes. For reference, a summary of questions for public consideration is on the ACHDNC website. We request that public participants providing oral comments on the review of the Committee’s evidence review process also submit a written version of their remarks.  

The agenda for this meeting does not include any vote or decision to recommend a condition for inclusion in the Recommended Uniform Screening Panel. As noted in the agenda items, the Committee may hold a vote on whether or not to recommend a nominated condition (GAMT deficiency) to full evidence review, which may lead to such a recommendation at a future time. Agenda items are subject to change as priorities dictate. Information about the ACHDNC, including a roster of members and past meeting summaries, is available on the ACHDNC website listed above.  

As previously noted, members of the public will have the opportunity to provide comments. Public participants providing general oral comments may submit written statements in advance of the scheduled meeting. Oral comments will be honored in the order they are requested and may be limited as time allows. Requests to provide a written statement or make oral comments to the ACHDNC must be submitted via the registration website by 10:00 a.m. ET on Monday, August 9, 2021.  

Individuals who need special assistance or another reasonable accommodation should notify Alaina Harris at the address and phone number listed above at least 10 business days prior to the meeting.  

Maria G. Button,  
Director, Executive Secretariat.  
[FR Doc. 2021–15569 Filed 7–21–21; 8:45 am]  
BILLING CODE 4165–15–P