disclosure. Comments will be posted on https://www.regulations.gov. Therefore, do not include any information in your comment or supporting materials that you consider confidential or inappropriate for public disclosure. If you include your name, contact information, or other information that identifies you in the body of your comments, that information will be on public display. ATSDR will review all submissions and may choose to redact, or withhold, submissions containing private or proprietary information such as Social Security numbers, medical information, inappropriate language, or duplicate/near duplicate examples of a mass-mail campaign. ATSDR will carefully consider all comments submitted in preparation of the final Toxicological Profile and may revise the profile as appropriate.

Legislative Background

The Superfund Amendments and Reauthorization Act of 1986 (SARA) [42 U.S.C. 9601 et seq.] amended the Comprehensive Environmental Response, Compensation, and Liability Act of 1980 (CERCLA or Superfund) [42 U.S.C. 9601 et seq.] by establishing certain requirements for ATSDR and the U.S. Environmental Protection Agency (EPA) regarding the hazardous substances most commonly found at facilities on the CERCLA National Priorities List (NPL). Among these statutory requirements is a mandate for the Administrator of ATSDR to prepare toxicological profiles for each substance included on the priority list of hazardous substances [also called the Substance Priority List (SPL)]. This list identifies 275 hazardous substances that ATSDR and EPA have determined pose the most significant potential threat to human health. The SPL is available online at www.atsdr.cdc.gov/spl.

In addition, CERCLA provides ATSDR with the authority to prepare toxicological profiles for substances not found on the SPL. ATSDR authorizes ATSDR to establish and maintain an inventory of literature, research, and studies on the health effects of toxic substances (CERCLA Section 104(i)(1)(B); 42 U.S.C. 9604(i)(1)(B)); to respond to requests for health consultations (CERCLA Section 104(i)(4); 42 U.S.C. 9604(i)(4)); and to support the site-specific response actions conducted by the agency.

Availability

The Draft Toxicological Profile for Ethylene Oxide will be available online at http://www.atsdr.cdc.gov/ToxProfiles and at www.regulations.gov, Docket No. ATSDR–2016–0004.

Donata Green,
Acting Director, Office of Policy, Planning, and Partnerships, Agency for Toxic Substances and Disease Registry.

[FR Doc. 2020–21619 Filed 9–29–20; 8:45 am]
BILLING CODE 4183–70–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day–20–1242; Docket No. CDC–2020–0099]

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 and/or continuing information collection, as required by the Paperwork Reduction Act of 1995. This notice invites comment on a proposed information collection project titled Strengthening U.S. Response to Resistant Gonorrhea, which is to intended to enhance U.S. state and local public health surveillance and program infrastructure, build capacity to support rapid detection and public health response to antibiotic-resistant gonorrhea (an urgent public health threat), and advance the understanding of epidemiological factors contributing to antibiotic-resistant gonorrhea. CDC is requesting a three-year approval.

DATES: CDC must receive written comments on or before November 30, 2020.

ADDRESSES: You may submit comments, identified by Docket No. CDC–2020–0099 by any of the following methods:

• Federal eRulemaking Portal: 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–D74, Atlanta, Georgia 30329.

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

Please note: Submit all comments through the Federal eRulemaking portal (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–D74, 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; and

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.

5. Assess information collection costs.

Proposed Project

Strengthening U.S. Response to Resistant Gonorrhea (SURRG) (OMB Control No. 0920–1242, Exp. 9/30/2025)
N. gonorrhoeae culture specimens for providers at participating clinics collect overview of SURRG, healthcare and will participate voluntarily. As an applied as part of a competitive process because healthcare providers rarely healthcare activities, such as SURRG, enhanced surveillance and public capacity to rapidly detect, monitor, and respond to emerging antibiotic-resistant gonorrhea (and get actionable information to local health departments), (2) understand trends in and factors contributing to antibiotic-resistant gonorrhea, and (3) build a robust evidence base for public health action. This information collection is important because: (1) Effective treatment of gonorrhea is critical to gonorrhea control and prevention, (2) untreated or inadequately treated gonorrhea can cause serious reproductive health complications, such as infertility, (3) Neisseria gonorrhoeae (the bacterium that causes gonorrhea) has consistently demonstrated the ability to develop antibiotic resistance and may be developing resistance to the last remaining treatment option recommended by the Centers for Disease Control and Prevention (CDC), and (4) antibiotic-resistant gonorrhea is extremely difficult to detect without enhanced surveillance and public health activities, such as SURRG, because healthcare providers rarely perform or have access to culture and resistance testing for individual patients.

Jurisdictions participating in SURRG applied as part of a competitive process and will participate voluntarily. As an overview of SURRG, healthcare providers at participating clinics collect specimens for N. gonorrhoeae culture testing. Specimens that demonstrate N. gonorrhoeae (called “isolates”) rapidly undergo antibiotic resistance testing at the local public health laboratory. Detection of resistance is rapidly communicated by laboratory staff to the healthcare provider and health department. The patient (from whom the resistant specimen was collected) is interviewed by local health department staff about risk factors and recent contacts, and will be re-tested to ensure that they were cured. Recent contacts are interviewed by the health department (contact tracing) and tested for gonorrhea. The participating health departments collect and transmit to CDC demographic and clinical data about persons tested for and diagnosed with gonorrhea in the participating clinics, results of local antibiotic resistance testing, and information about field investigations. None of the data transmitted to CDC contains any personally identifiable information. These data are used by CDC to monitor and better understand resistance and identify effective approaches to prevent resistance spread. Data are transmitted to CDC through a secure encrypted file transfer application and stored in a secure CDC server with strictly controlled and restricted access rights.

In processes that take approximately 16 hours every two months (plus an annual cumulative datafile), local SURRG data managers abstract STD clinic data for patients tested for gonorrhea and field investigation data, receive gonorrhea data from non-STD clinic healthcare sites and resistance testing results from local public health laboratories, and clean and transmit data to CDC. 

Other data managers at each participating non-STD clinic health center abstract, clean, and transmit data (approximately three hours every two months). Microbiologists at public health laboratories from each funded jurisdiction conduct resistance testing on ~700 N. gonorrhoeae isolates each year (600 clinical isolates and 100 control strains; each test ~10 minutes). Laboratory data managers take about one hour every two months to abstract, clean, and transmit data.

Health department staff will interview: Any person diagnosed with antibiotic-resistant gonorrhea or have a case of gonorrhea of public health significance index case, and their sexual contacts. On average, each jurisdiction will identify four drug-resistant isolates each month; these isolates will spur field investigations and six additional interviews monthly. We estimate a total of 120 interviews annually at each site, for a total across the eight sites of 960 interviews each year. Each interview will take ~20 minutes.

The total estimated annual burden hours is 2,665. This burden represents a decrease from the burden of the initial submission. The number of jurisdictions decreased from nine to eight. So the number of local data managers decreased from nine to eight (and the burden hours decreased from 1008 to 896), the number of public health microbiologists decreased from nine to eight (burden hours decreased from 1050 to 933), the number of lab data managers decreased from nine to eight (burden hours decreased from 54 to 48), and the number of gonorrhea and contacts decreased from 1080 to 960 (burden hours decreased from 540 to 480). The number of clinic sites will increase from 18 to 26. Respondents receive federal funds to participate in this project. There are no additional costs to respondents other than their time.

### Estimated Annualized Burden Hours

<table>
<thead>
<tr>
<th>Type of respondent</th>
<th>Form name</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Average burden per response (hours)</th>
<th>Total burden hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>Local SURRG data manager</td>
<td>STD Clinic Facility Data Elements ...</td>
<td>8</td>
<td>7</td>
<td>16</td>
<td>896</td>
</tr>
<tr>
<td>Data manager at non-STD clinic health centers</td>
<td>Non-STD Clinic Facility Data Elements.</td>
<td>26</td>
<td>6</td>
<td>3</td>
<td>468</td>
</tr>
<tr>
<td>Public Health Laboratory Microbiologist.</td>
<td>Laboratory Data Elements ................</td>
<td>8</td>
<td>700</td>
<td>10/60</td>
<td>933</td>
</tr>
<tr>
<td>Public Health Laboratory Data Manager.</td>
<td>Laboratory Testing Data Elements</td>
<td>8</td>
<td>6</td>
<td>1</td>
<td>48</td>
</tr>
<tr>
<td>Gonorrhea Patients, Social and Sexual Contacts.</td>
<td>Investigation Data Elements ..........</td>
<td>960</td>
<td>1</td>
<td>0.33</td>
<td>320</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>2,665</td>
</tr>
</tbody>
</table>
DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Notice of Closed Meeting

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended, notice is hereby given of the following meeting.

The meeting will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title 5 U.S.C., as amended, and the Determination of the Director, Strategic Business Initiatives Unit, Office of the Chief Operating Officer, CDC, pursuant to Public Law 92–463. The grant applications and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: Disease, Disability, and Injury Prevention and Control Special Emphasis Panel (SEP)—Development (SEED) Follow up Studies.

Date: January 12–13, 2021.

Time: 10:00 a.m.–6:00 p.m., EST.

Place: Teleconference.

Agenda: To review and evaluate grant applications.

For Further Information Contact: Jaya Raman Ph.D., Scientific Review Officer, CDC, 4770 Buford Highway, Mailstop S107–8, Atlanta, Georgia 30341, Telephone: (770) 488–6511, JRaman@cdc.gov.

The Director, Strategic Business Initiatives Unit, Office of the Chief Operating Officer, Centers for Disease Control and Prevention, has been delegated the authority to sign Federal Register notices pertaining to announcements of meetings and other committee management activities, for both the Centers for Disease Control and Prevention and the Agency for Toxic Substances and Disease Registry.

Kalwant Smagh,
Director, Strategic Business Initiatives Unit, Office of the Chief Operating Officer, Centers for Disease Control and Prevention.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Submission for OMB Review: National Advisory Committee on the Sex Trafficking of Children and Youth in the United States (NAC) Recommendations and State Self-Assessment Survey (NEW)

AGENCY: Office on Trafficking in Persons, Administration for Children and Families, HHS.

ACTION: Request for public comment.

SUMMARY: The Office on Trafficking in Persons (OTIP), Administration for Children and Families (ACF), U.S. Department of Health and Human Services (HHS), is proposing to collect data for a new survey, the National Advisory Committee on the Sex Trafficking of Children and Youth in the United States (NAC) Recommendations and State Self-Assessment Survey.

DATES: Comments due within 30 days of publication. OMB is required to make a decision concerning the collection of information between 30 and 60 days after publication of this document in the Federal Register. Therefore, a comment is best assured of having its full effect if OMB receives it within 30 days of publication.

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” or by using the search function.

SUPPLEMENTARY INFORMATION:

Description: The Preventing Sex Trafficking and Strengthening Families Act of 2014 mandated the NAC to develop a report describing how each state and territory has implemented its recommendations to address sex trafficking in children and youth. The NAC proposes to administer a survey allowing states to assess their progress in implementing NAC recommendations. Submissions will allow states to document their efforts in the following sections: Multidisciplinary Response, Screening and Identification, Child Welfare, Service Provision, Housing, Law Enforcement and Prosecution, Judiciary, Demand Reduction, Prevention, Legislation and Regulation, Research and Data, and Funding. Each state will have the opportunity to provide a self-assessed tier ranking for each recommendation, a justification of their assessment, sources for their assessment, and the public or private nature of those sources.

Respondents: State Governors, Child Welfare Agencies, Local Law Enforcement, and Other Local Agencies.

Annual Burden Estimates:

**Estimated Opportunity Costs for Respondents**

<table>
<thead>
<tr>
<th>Instrument</th>
<th>Total number of respondents contributing for 50 states</th>
<th>Total number of responses per respondent</th>
<th>Average burden hours per response</th>
<th>Total/annual burden hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>NAC Recommendations and State Self-Assessment Survey</td>
<td>250</td>
<td>1</td>
<td>6.85</td>
<td>1,713</td>
</tr>
</tbody>
</table>

Estimated Total Annual Burden Hours: 1,713.