Background and Brief Description

The Centers for Disease Control and Prevention’s (CDC), National Center for Emerging and Zoonotic Infectious Diseases (NCEZID), Division of Global Migration and Quarantine (DGMQ), Immigrant, Refugee, and Migrant Health Branch (IRMH), requests approval for a revision of an existing information collection. This project pertains to collecting annual reports on certain tuberculosis data from U.S. panel physicians.

The respondents are panel physicians. More than 760 panel physicians from 336 panel sites perform overseas pre-departure medical examinations in accordance with requirements, referred to as technical instructions, provided by DGMQs Quality Assessment Program (QAP). The role of QAP is to assist and guide panel physicians in the implementation of the Technical Instructions; evaluate the quality of the overseas medical examination for U.S.-bound immigrants and refugees; assess potential panel physician sites; and provide recommendations to the U.S. Department of State in matters of immigrant medical screening.

To achieve DGMQ’s mission, the Immigrant, Refugee and Migrant Health branch (IRMH) works with domestic and international programs to improve the health of U.S.-bound immigrants and refugees to protect the U.S. public by preventing the importation of infectious disease. These goals are accomplished through IRMH’s oversight of medical exams required for all U.S.-bound immigrants and refugees who seek permanent residence in the U.S. IRMH is responsible for assisting and training the international panel physicians with the implementation of medical exam Technical Instructions (TI). Technical Instructions are detailed requirements and national policies regarding the medical screening and treatment of all U.S.-bound immigrants and refugees.

Screening for tuberculosis (TB) is a particularly important component of the immigration medical exam and allows panel physicians to diagnose active TB disease prior to arrival in the United States. As part of the Technical Instructions requirements, panel physicians perform chest x-rays and laboratory tests that aid in the identification of tuberculosis infection (Class B1 applicants) and diagnosis of active tuberculosis disease (Class A, inadmissible applicants). CDC uses these classifications to report new immigrant and refugee arrivals with a higher risk of developing TB disease to U.S. state and local health departments for further follow-up. Some information that panel physicians collect as part of the medical exam is not reported on the standard Department of State forms (DS-forms), thereby preventing CDC from evaluating TB trends in globally mobile populations and monitoring program effectiveness.

Currently, CDC is requesting this data be sent by panel physicians once per year. The consequences of reducing this frequency would be the loss of monitoring program impact and TB burdens in mobile populations and immigrants and refugees coming to the United States on an annual basis. The total hours requested is 1,008. There is no cost to the respondents other than their time.

Estimated annual burden is being reduced by 1,640 hours per year. The number of respondents is being reduced by 17. Reductions are due to revised estimates on burden time per response, and the removal of four variables from the data collection form and improved IT capacity at most panel sites.

<table>
<thead>
<tr>
<th>Type of respondents</th>
<th>Form name</th>
<th>Number of responses per respondent</th>
<th>Average burden per response (in hours)</th>
<th>Total burden (in hours)</th>
</tr>
</thead>
<tbody>
<tr>
<td>International panel physicians</td>
<td>TB Indicators Excel Spreadsheet</td>
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<td>1</td>
<td>3</td>
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<tr>
<td>Total</td>
<td></td>
<td></td>
<td></td>
<td>1,008</td>
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</table>

Jeffrey M. Zirger,

[FR Doc. 2018–18054 Filed 8–21–18; 8:45 am]
BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day–18–0338; Docket No. CDC–2018–0076]

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 efforts to reduce public burden and maximize the utility of government information, invites the general public and other Federal agencies to take this opportunity to comment on proposed work and/or continuing information collections, as required by the Paperwork Reduction Act of 1995. This notice invites comment on a continuing information collection project titled Annual Submission of the Ingredients Added to, and the Quantity of Nicotine Contained in, Smokeless Tobacco Manufactured, Imported, or Packaged in the U.S.

DATES: CDC must receive written comments on or before October 22, 2018.

ADDRESSES: You may submit comments, identified by Docket No. CDC–2018–0076 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
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

Annual Submission of the Ingredients Added to, and the Quantity of Nicotine Contained in, Smokeless Tobacco Manufactured, Imported, or Packaged in the U.S.—Extension—National Center for Chronic Disease Prevention and Health Promotion (NCDCPH), Centers for Disease Control and Prevention (CDC).

### Background and Brief Description

Smokeless tobacco products (SLT) are associated with many health problems. Using smokeless tobacco: Can lead to nicotine addiction; causes cancer of the mouth, esophagus, and pancreas; is associated with diseases of the mouth; can increase risks for early delivery and stillbirth when used during pregnancy; can cause nicotine poisoning in children; and, may increase the risk for death from heart disease and stroke.

The CDC’s Office on Smoking and Health (OSH) has the primary responsibility for the HHS smoking and health program. As required by the Comprehensive Smokeless Tobacco Health Education Act of 1986 (CSTHEA, 15 U.S.C. 4401 et seq., Pub. L. 99–252), CDC collects a list of ingredients added to tobacco in the manufacture of smokeless tobacco products and a specification of the quantity of nicotine contained in each product. HHS has delegated responsibility for implementing the required information collection to CDC’s OSH. Respondents are manufacturers, packagers, or importers (or their representatives) of smokeless tobacco products.

Respondents are not required to submit specific forms; however, they are required to meet reporting guidelines and to submit the ingredient report by chemical name and Chemical Abstract Service (CAS) Registration Number, consistent with accepted reporting practices for other companies that are required to report ingredients added to other consumer products. Typically, respondents submit a summary report to CDC with the ingredient information for multiple products, or a statement that there are no changes to their previously submitted ingredient report. Respondents may submit the required information to CDC through a designated representative. The information collection is subject to strict confidentiality provisions.

Ingredient reports for new SLT products are due at the time of first importation. Thereafter, ingredient reports are due annually on March 31. Information is submitted to CDC by mailing a written report on the respondent’s letterhead, by CD, three-inch floppy disk, or thumb drive. Electronic mail submissions are not accepted. Annual submission reports are mailed to Attention: FCLAA Program Manager, Office on Smoking and Health, National Center for Chronic Disease Prevention and Health Promotion, Centers for Disease Control and Prevention, 4770 Buford Highway NE, MS S107–7, Atlanta, GA 30341–3717.

Upon receipt and verification of the annual nicotine and ingredient report, CDC issues a Certificate of Compliance to the respondent. As deemed appropriate by the Secretary of HHS, HHS is authorized to use the information to report to Congress the health effects of ingredients, research activities related to the health effects of ingredients, and other information that the Secretary determines to be of public interest. The estimated annual Burden Hours are 18,843. There are no costs to respondents other than their time. OMB approval is requested for three years.

### 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>
</tr>
</thead>
<tbody>
<tr>
<td>Smokeless Tobacco Manufacturers, Packagers, and Importers.</td>
<td>SLT Nicotine and Ingredient and Report.</td>
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<td>1</td>
<td>1,713</td>
<td>18,843</td>
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</table>
DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Administration for Native Americans; Notice of Meeting

AGENCY: Administration for Children and Families, Department of Health and Human Services.

ACTION: Notice of Tribal Consultation.

SUMMARY: The Department of Health and Human Services, Administration for Children and Families (ACF) will host a Tribal Consultation to consult on ACF programs and tribal priorities.

DATES: September 13, 2018.

ADDRESS: Capital Skyline Hotel, 10 “T” (eye) Street SW, Washington, DC 20024.

FOR FURTHER INFORMATION CONTACT: Jeannie Hovland, Commissioner, Administration for Native Americans and Deputy Assistant Secretary for Native American Affairs at 202–401–5156, by email at anacommissioner@acf.hhs.gov or by mail at 330 C Street SW, MS–4126, Washington, DC 20201.

SUPPLEMENTARY INFORMATION: In accordance with the ACF Tribal Consultation Policy, ACF announces tribal consultation with tribal leaders operating ACF programs.

The consultation will be conducted with elected or appointed leaders of tribal governments and their designated representatives. Designees must have a letter from the tribal government authorizing them to represent the tribe. Tribal governments must submit the designee letter at least 3 days in advance of the consultation session to the Administration for Native Americans at anacommissioner@acf.hhs.gov. Other representatives of tribal organizations and Native non-profit organizations are welcome to attend as observers. A report of the consultation session will be prepared and made available at the following address within 45 days after the closing of the consultation session. Tribes wishing to submit written testimony should send it to anacommissioner@acf.hhs.gov either prior to the consultation session or within 30 days after the meeting. ACF will summarize oral testimony and comments from the consultation session along with topics of concern and recommendations.

ACF has identified the following topics for consultation:

Family First Services Act

Title IV–E Planning Grants—What barriers are preventing Tribes from applying for the grant

Office of Head Start annual consultation

TANF and Welfare reform

The ACF Tribal Consultation Session will begin at 9:00 a.m. on September 13 and continue throughout the day until all discussions have been completed. To help both you and the ACF Principals prepare for this consultation, planning teleconference calls will be held:

Wednesday, August 22, 2018 @ 4:00 p.m.–4:30 p.m. (EST)

Thursday, August 23, 2018 @ 4:00 p.m.–4:30 p.m. (EST)

Tuesday, August 28, 2018 @ 4:00 p.m.–4:30 p.m. (EST)

The call-in number and passcode are: 866–769–9393 passcode: 4494949#.

The purpose of the planning calls will be to identify individuals who will provide oral testimony to ACF, solicit for tribal moderators and identify specific topics of interest so we can ensure that all appropriate individuals are present.

Any tribe unable to attend in person, ACF will provide a webinar link. Please contact our 1–877–922–9262 (TANF) or 1–877–922–9262 ( Infant-Family First Services Act) for the webinar information.

We have set up a registration for all participants whether attending in person or by webinar. The registration address is: www.regonline.com/

2018acftribalconsultation. If you plan on providing testimony, please include the name of the office you wish to address.

Jeannie Hovland,
Deputy Assistant Secretary for Native American Affairs.

[FR Doc. 2018–18053 Filed 8–21–18; 8:45 am]

BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families


AGENCY: Unaccompanied Alien Children’s (UAC) Program, Office of Refugee Resettlement (ORR), Administration for Children and Families (ACF), U.S Department of Health and Human Services (HHS).

ACTION: Notice of intent to issue one OPDIV-Initiated Supplement to BCFS Health and Human Services, San Antonio, TX under the UAC Program.

SUMMARY: ACF, ORR, announces the intent to issue one OPDIV-Initiated Supplement to BCFS Health and Human Services, San Antonio, TX in the amount of $28,003,926. ORR has been identifying additional capacity to provide shelter for potential increases in apprehensions of Unaccompanied Children at the U.S. Southern Border. Planning for increased shelter capacity is a prudent step to ensure that ORR is able to meet its responsibility, by law, to provide shelter for Unaccompanied Alien Children referred to its care by the Department of Homeland Security (DHS). To ensure sufficient capacity to provide shelter to unaccompanied children referred to HHS, BCFS proposed to provide ORR with 850 beds in an expedited manner.

DATES: Supplemental award funds will support activities through September 13, 2018.

FOR FURTHER INFORMATION CONTACT: Jallyn Sualog, Director, Division of Children’s Services, Office of Refugee Resettlement, 330 C Street SW, Washington, DC 20447. Phone: 202–401–4997. Email: DCSProgram@acf.hhs.gov.

SUPPLEMENTARY INFORMATION: ORR is continuously monitoring its capacity to shelter the unaccompanied children referred to HHS, as well as the information received from interagency partners, to inform any future decisions or actions.

ORR has specific requirements for the provision of services. Award recipients must have the infrastructure, licensing, experience, and appropriate level of trained staff to meet those requirements. The expansion of the existing program and its services through this