manuscripts, online resources, and other avenues.

To monitor changes in attitudes and practices regarding provision of contraception among family planning providers and clinics, we initiated a multi-phase assessment. In 2009–2010, CDC carried out the first phase of the assessment, collecting information before the release of the US MEC (OMB No. 0920–0008). In 2013–2014, CDC, in collaboration with OPA, carried out the second phase of the assessment, collecting information before the release of the US SPR and QFP (OMB No. 0920–0969). These information collections provided useful knowledge about attitudes and practices of family planning providers. CDC and OPA used the findings to develop educational materials and opportunities for health care providers.

In 2018, in collaboration with OPA, CDC plans to request a reinstatement of OMB No. 0920–0969, “Monitoring Changes in Attitudes and Practices among Family Planning Providers and Clinics” to carry out the third phase of the assessment. As in the previous phases, the information collection will allow CDC and OPA to improve family planning-related practice by: (1) Understanding the current use of contraception guidance in practice, including awareness and use of the US MEC, US SPR and QFP; (2) describing current attitudes and practices among family planning providers and clinics related to recommendations included in the US MEC, US SPR, and QFP; and (3) identifying training needs in use of guidance and family planning service delivery (e.g., provider tools, continuing education modules).

As in previous phases of data collection, CDC plans to administer surveys to private and public sector family planning providers and clinic administrators in the United States. The design, methodology, and analytic approach that CDC plans to implement are based on methods previously approved for the 2013–2014 survey, with different instruments being administered to providers and clinic administrators. Minor changes to survey content will be made to eliminate unnecessary questions, add new questions of interest, and improve formatting, usability, and data quality. The estimated burden per response for providers is 15 minutes and has not changed since the previous OMB approval. The estimated burden per response for administrators will be reduced from 40 minutes to 35 minutes. The total burden for participants is estimated at 1,916 hours. Participation is voluntary and there are no costs to respondents other than their time. OMB approval is requested for one year.

### 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>
</tr>
</thead>
<tbody>
<tr>
<td>Office-based physicians (private sector) ...................</td>
<td>2018–2019 Survey of Health Care Providers about Family Planning Attitudes and Practices.</td>
<td>1,000</td>
<td>1</td>
<td>15/60</td>
</tr>
<tr>
<td>Title X clinic providers (public sector) .................</td>
<td>2018–2019 Survey of Health Care Providers about Family Planning Attitudes and Practices.</td>
<td>1,000</td>
<td>1</td>
<td>15/60</td>
</tr>
<tr>
<td>Non-Title X clinic providers (public sector) ..........</td>
<td>2018–2019 Survey of Health Care Providers about Family Planning Attitudes and Practices.</td>
<td>1,000</td>
<td>1</td>
<td>15/60</td>
</tr>
<tr>
<td>Title X clinic administrators (public sector) ..........</td>
<td>2018–2019 Survey of Administrators of Health Centers that Provide Family Planning.</td>
<td>1,000</td>
<td>1</td>
<td>35/60</td>
</tr>
<tr>
<td>Non-Title X clinic administrators (public sector) ......</td>
<td>2018–2019 Survey of Administrators of Health Centers that Provide Family Planning.</td>
<td>1,000</td>
<td>1</td>
<td>35/60</td>
</tr>
</tbody>
</table>

Jeffrey M. Zirger,

[FR Doc. 2018–23862 Filed 10–31–18; 8:45 am]
BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day-19–0488]

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 Report of Illness or Death: Interstate Travel of Persons (42 CFR part 70) 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 August 21, 2018 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 or send an email to omb@cdc.gov. 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

Report of Illness or Death: Interstate Travel of Persons (42 CFR part 70)—Revision—National Center for Emerging and Zoonotic Infectious Diseases (NCEZID), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

Section 361 of the Public Health Service Act (42 U.S.C. 264) authorizes the Secretary of the Department of Health and Human Services to make and enforce regulations necessary to prevent the introduction, transmission, or spread of communicable diseases from foreign countries into the United States, or from one State or possession into any other State or possession. CDC administers regulations pertaining to interstate control of communicable diseases (42 CFR part70), and sections 42 CFR parts 70.4 and 70.11 include requirements for reports of ill persons or death if occurring during interstate travel.

The intended use of the information is to ensure that CDC can assess and respond to reports of ill persons or death that occur on conveyances engaged in interstate travel, and assist state and local health authorities if an illness or death occurs that poses a risk to public health. Generally, the primary source of this information is aircraft traveling within the United States.

In 2017, CDC finalized the Control of Communicable Disease regulations (42 CFR 70 and 71). With this new provision, CDC divided the total anticipated reporting burden between 70.11 and 70.4 in the accompanying Paperwork Reduction Act section of the rule, assuming that aircraft would report most cases of ill people and deaths to CDC, with some airlines and other conveyances reporting still to local public health authorities. For reports of ill persons or death on a conveyance engaged in interstate traffic, the requested burden is approximately 23 hours. This total is estimated from 200 respondents submitting domestic reports of death or communicable disease a year, with an average burden of 7 minutes per report. The only requested change to the approved data collection is a change in title from “Restriction on Travel of Persons (42 CFR part 70)” to “Report of Illness or Death: Interstate Travel of Persons (42 CFR part 70)”. This results in two rows in the burden table, but with no additional burden. The estimated annual Burden Hours are 23. There is no cost to respondents other than their time.

### 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>
</tr>
</thead>
<tbody>
<tr>
<td>Pilot in command</td>
<td>42 CFR 70.11 Report of death or illness onboard aircraft operated by airline.</td>
<td>190</td>
<td>1</td>
<td>7/60</td>
</tr>
<tr>
<td>Pilot in command</td>
<td>42 CFR 70.4 Report by the master of a vessel or person in charge of conveyance of the incidence of a communicable disease occurring while in interstate travel.</td>
<td>10</td>
<td>1</td>
<td>7/60</td>
</tr>
</tbody>
</table>

Jeffrey M. Zirger

[FR Doc. 2016–23861 Filed 10–31–18; 8:45 am]
BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration
(Docket No. FDA–2012–N–0873)

Agency Information Collection Activities; Proposed Collection; Comment Request; Bar Code Label Requirement for Human Drug and Biological Products

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA or Agency) is announcing an opportunity for public comment on the proposed collection of certain information by the Agency. 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 of an existing collection of information, and to allow 60 days for public comment in response to the notice. This notice solicits comments on bar code label requirements for human drug and biological products.

DATES: Submit either electronic or written comments on the collection of information by December 31, 2018.

ADDRESSES: You may submit comments as follows. Please note that late, untimely filed comments will not be considered. Electronic comments must be submitted on or before December 31, 2018. The https://www.regulations.gov electronic filing system will accept comments until 11:59 p.m. Eastern Time at the end of December 31, 2018. Comments received by mail/hand delivery/courier (for written/paper submissions) will be considered timely if they are postmarked or the delivery service acceptance receipt is on or before that date.

Electronic Submissions

Submit electronic comments in the following way:

- Federal eRulemaking Portal: https://www.regulations.gov. Follow the instructions for submitting comments. Comments submitted electronically, including attachments, to https://www.regulations.gov will be posted to the docket unchanged. Because your comment will be made public, you are solely responsible for ensuring that your comment does not include any confidential information that you or a third party may not wish to be posted, such as medical information, your or anyone else’s Social Security number, or confidential business information, such