- Healthcare utilization (use and length of hospice care, hospitalizations, advance directive documentation) and costs and resource use (use of outpatient clinician services, including palliative care)
- Adverse effects
  - Medication side effects
  - Dropouts
- Timing
  - Any timing
- Settings
  - Ambulatory primary and specialty care, including geriatrics, nephrology, pulmonology, cardiology, and neurology
  - U.S.-based studies, as systems of care differ in other countries

Dated: January 15, 2020.

#### Virginia L. Mackay-Smith,

Associate Director, Office of the Director, AHRQ.

[FR Doc. 2020–00903 Filed 1–21–20; 8:45 am]

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

## Centers for Disease Control and Prevention

[0Day-20-0006; Docket No. CDC-2019-0118]

# 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 Statements in Support of Application of Waiver of Inadmissibility (0920-0006). CDC uses the information collected in 0920-0006 to review Class A medical waiver applications for prospective

immigrants to the United States. CDC assists DHS/USCIS in determining whether or not a prospective immigrant with a Class A mental health designation may be admitted into the United States.

**DATES:** CDC must receive written comments on or before March 23, 2020.

**ADDRESSES:** You may submit comments, identified by Docket No. CDC-2019-0118 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, of the 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**

Statements in Support of Application of Waiver of Inadmissibility (OMB Control No. 0920–0006 Exp. 6/30/2020)—Revision—National Center for Emerging and Zoonotic Infectious Diseases (NCEZID), Centers for Disease Control and Prevention (CDC).

### **Background and Brief Description**

Section 212(a)(1) of the Immigration and Nationality Act states that aliens with specific health related conditions are ineligible for admission into the United States. The Attorney General may waive application of this inadmissibility on health-related grounds if an application for waiver is filed and approved by the consular office considering the application for visa. CDC uses this application primarily to collect information to establish and maintain records of waiver applicants in order to notify the U.S. Citizenship and Immigration Services when terms, conditions and controls imposed by waiver are not met.

The purpose of this Revision is to remove information collections for form 4.422–1a, because CDC does not receive information about the evaluation report of an applicant who received a waiver. This results in a reduction of 67 burden hours. CDC requests approval for 33 annual burden hours.

#### **ESTIMATED ANNUALIZED BURDEN HOURS**

Type of respondent	Form	Number of respondents	Number of responses per respondent	Average burden per response (in hours)	Total burden hours
Physician	CDC 4.422-1	200	1	10/60	33
Total					33

### Jeffrey M. Zirger,

Lead, Information Collection Review Office, Office of Scientific Integrity, Office of Science, Centers for Disease Control and Prevention. [FR Doc. 2020–01051 Filed 1–21–20; 8:45 am]

BILLING CODE 4163-18-P

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Administration for Children and Families

[OMB No. 0970-0492]

### Submission for OMB Review; Community Services Block Grant Annual Report

**AGENCY:** Office of Community Services; Administration for Children and Families; HHS.

**ACTION:** Request for public comment.

**SUMMARY:** The Administration of Children and Families (ACF), Office of Community Services (OCS) is requesting a three-year extension with minor

changes of the Community Services Block Grant (CSBG) Annual Report (OMB No.: 0970–0492, expiration 1/31/ 2020). This request will support the currently utilized CSBG Annual Report, comprised of Modules 1–4, and incorporates performance management. **DATES:** Comments due within 30 days of publication. OMB is required to make a

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 directly to the following: Office of Management and Budget, Paperwork Reduction Project, Email: OIRA\_SUBMISSION@OMB.EOP.GOV, Attn: Desk Officer for the Administration for Children and Families.

Copies of the proposed collection may be obtained by emailing *infocollection*@

acf.hhs.gov. Alternatively, copies can also be obtained by writing to the Administration for Children and Families, Office of Planning, Research, and Evaluation (OPRE), 330 C Street SW, Washington, DC 20201, Attn: OPRE Reports Clearance Officer. All requests, emailed or written, should be identified by the title of the information collection.

#### SUPPLEMENTARY INFORMATION:

Description: Module 1 includes minor edits to align with the updated, and OMB approved, CSBG State Plan. Module 2, Module 3, and Module 4 include only technical and grammatical updates for ease and clarity of current reporting. Copies of the proposed collection of information can be obtained by visiting: http://www.acf.hhs.gov/programs/ocs/programs/csbg.

Respondents: State governments, including the District of Columbia and the Commonwealth of Puerto Rico, and U.S. territories and CSBG eligible entities (Community Action Agencies).

Annual Burden Estimates:

Instrument	Annual number of respondents	Annual number of responses per respondent	Average burden hours per response	Annual burden hours
CSBG Annual Report (States)	52 1,009	1 1	198 697	10,296 703,273

Estimated Total Annual Burden Hours: 713,569.

**Authority:** 112 Stat. 2729; 42 U.S.C. 9902(2).

### Mary B. Jones,

 $\label{eq:acf-opre-operator} ACF/OPRE\ Certifying\ Officer. \\ \mbox{[FR Doc. 2020–00928 Filed 1–21–20; 8:45 am]}$ 

BILLING CODE 4184-27-P

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Food and Drug Administration

[Docket Nos. FDA-2018-E-3053 and FDA-2018-E-4226]

### Determination of Regulatory Review Period for Purposes of Patent Extension; MAVYRET

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA or the Agency) has determined the regulatory review period for MAVYRET and is publishing this notice of that determination as required by law. FDA has made the determination because of the submission of applications to the Director of the U.S. Patent and Trademark Office (USPTO), Department of Commerce, for the extension of patents which claims that human drug product.

DATES: Anyone with knowledge that any of the dates as published (see SUPPLEMENTARY INFORMATION) are incorrect may submit either electronic or written comments and ask for a redetermination by March 23, 2020. Furthermore, any interested person may petition FDA for a determination regarding whether the applicant for extension acted with due diligence during the regulatory review period by July 20, 2020. See "Petitions" in the