# DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Administration for Children and Families

Submission for Office of Management and Budget Review; Community Services Block Grant (CSBG) Model Tribal Plan Applications (New Collection)

**AGENCY:** Office of Community Services, Administration for Children and Families, U.S. Department of Health and Human Services.

**ACTION:** Request for public comments.

SUMMARY: The Office of Community Services (OCS), Administration for Children and Families (ACF), requests an approval of the Community Services Block Grant (CSBG) Model Tribal Plan. DATES: Comments due within 30 days of publication. OMB must make a decision about 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. You can also obtain copies of the proposed collection of information by emailing infocollection@acf.hhs.gov. Identify all emailed requests by the title of the information collection.

### SUPPLEMENTARY INFORMATION:

## ANNUAL BURDEN ESTIMATES

Description: Section 677 of the CSBG Act requires Indian tribes or tribal organizations to submit an application and plan (CSBG Model Tribal Plan). The CSBG Model Tribal Plan must meet statutory requirements prior to OCS awarding CSBG tribal grant recipients with CSBG funds. Tribal grant recipients have the option to submit a detailed plan annually or biannually. Tribal grant recipients that submit a biannual plan must provide an abbreviated plan the following year if substantial changes to the initial plan will occur. The CSBG Model Tribal Plan has been used in previous years without OMB approval. To come into compliance with the PRA, ACF is submitting the CSBG Model Tribal Plan as a new request to OMB.

Respondents: Tribal grant recipients (tribes and tribal organizations)

Instrument	Total number of respondents	Annual number of responses per respondent	Average burden hours per response	Annual burden hours
CSBG Model Tribal Plan	66	1	10	660

Authority: Sec. 677, Pub. L. 105–285, 112 Stat. 2742 (42 U.S.C. 9911).

### Mary C. Jones,

ACF/OPRE Certifying Officer.
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# DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration [Docket No. FDA-2022-N-0622]

Advisory Committee; Pulmonary-Allergy Drugs Advisory Committee; Renewal

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

**ACTION:** Notice; renewal of Federal advisory committee.

SUMMARY: The Food and Drug
Administration (FDA or the Agency) is
announcing the renewal of the
Pulmonary-Allergy Drugs Advisory
Committee by the Commissioner of
Food and Drugs (the Commissioner).
The Commissioner has determined that
it is in the public interest to renew the
Pulmonary-Allergy Drugs Advisory
Committee for an additional 2 years
beyond the charter expiration date. The

new charter will be in effect until the May 30, 2026, expiration date.

**DATES:** Authority for the Pulmonary-Allergy Drugs Advisory Committee will expire on May 30, 2026, unless the Commissioner formally determines that renewal is in the public interest.

## FOR FURTHER INFORMATION CONTACT:

Takyiah Stevenson, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 31, Rm. 2417, Silver Spring, MD 20993–0002, 240– 402–2507, PADAC@fda.hhs.gov.

**SUPPLEMENTARY INFORMATION:** Pursuant to 41 CFR 102-3.65 and approval by the Department of Health and Human Services and by the General Services Administration, FDA is announcing the renewal of the Pulmonary-Allergy Drugs Advisory Committee (the Committee). The Committee is a discretionary Federal advisory committee established to provide advice to the Commissioner. The Committee advises the Commissioner or designee in discharging responsibilities as they relate to helping to ensure safe and effective drugs for human use and, as required, any other product for which FDA has regulatory responsibility.

The Committee reviews and evaluates available data concerning the safety and effectiveness of marketed and investigational human drug products for use in the treatment of pulmonary disease and diseases with allergic and/or immunologic mechanisms and makes appropriate recommendations to the Commissioner of Food and Drugs.

Pursuant to its Charter, the Committee shall consist of a core of 11 voting members, including the Chair. Members and the Chair are selected by the Commissioner or designee from among authorities knowledgeable in the fields of pulmonary medicine, allergy, clinical immunology, and epidemiology or statistics. Members will be invited to serve for overlapping terms of up to 4 years. Non-Federal members of this committee will serve as Special Government Employees or representatives. Federal members will serve as Regular Government Employees or Ex-Officios. The core of voting members may include one technically qualified member, selected by the Commissioner or designee, who is identified with consumer interests and is recommended by either a consortium of consumer-oriented organizations or other interested persons. In addition to the voting members, the Committee may include one non-voting representative member who is identified with industry interests. There may also be an alternate industry representative.