Estimated Total Annual Burden Hours: 15,114.

In compliance with the requirements of the Paperwork Reduction Act of 1995 (Pub. L. 104–13, 44 U.S.C. Chap 35), the Administration for Children and Families is soliciting public comment on the specific aspects of the information collection described above. Copies of the proposed collection of information can be obtained and comments may be forwarded by writing to the Administration for Children and Families, Office of Planning, Research and Evaluation, 330 C Street SW., Washington, DC 20201. Attn: ACF Reports Clearance Officer. Email address: infocollection@acf.hhs.gov. All requests should be identified by the title of the information collection.

The Department specifically requests comments on: (a) Whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information shall have practical utility; (b) the accuracy of the agency’s estimate of the burden of the proposed collection of information; (c) the quality, utility, and clarity of the information to be collected; and (d) ways to minimize the burden of the collection of information on respondents, including through the use of automated collection techniques or other forms of information technology.

Consideration will be given to comments and suggestions submitted within 60 days of this publication.

Robert Sargis,
Reports Clearance Officer.
[FR Doc. 2017–10150 Filed 5–18–17; 8:45 am]
BILLING CODE 4184–40–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Submission for OMB Review; Comment Request

Title: Community Services Block Grant (CSBG) Model State Plan (Revision).
OMB No.: 0970–0382.
Description: Section 676 of the Community Services Block Grant (CSBG) Act requires States, including the District of Columbia and the Commonwealth of Puerto Rico, and U.S. territories applying for CSBG funds to submit an application and plan (Model State Plan). The CSBG State Plan submitted by States must meet statutory requirements prior to being funded with CSBG funds. Applicants have the option to submit a detailed plan annually or biannually. Entities that submit a biannual plan must provide an abbreviated plan the following year if substantial changes to the initial plan will occur.

In 2015, the Model State Plan was substantially revised by automating the form, streamlining the information, and incorporating accountability measures that include customer satisfaction information from eligible entities that receive a proportional share of CSBG funding through State CSBG lead agencies along with technical assistance, monitoring, and other programmatic support.

In fall 2015, the Office of Community Services (OCS) used the American Customer Satisfaction Index (ACSI) to obtain feedback from CSBG eligible entities about services provided by the state CSBG Lead Agencies, as detailed in the new State Accountability Measures. OCS also obtained feedback from state CSBG Lead Agencies on services provided by the federal agency, as outlined in the new Federal Accountability Measures. Both OCS and state CSBG Directors received their state survey results in February 2016.

To support ongoing implementation of state accountability measures related to customer satisfaction from eligible entities, OCS plans to survey eligible entities using the ACSI survey instrument. No changes are planned from the content of the 2015 survey.

Respondents: CSBG eligible entities.

ANNUAL BURDEN ESTIMATES

<table>
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<tr>
<th>Instrument</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Average burden hours per response</th>
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DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Submission for OMB Review; 30-Day Comment Request: Application Process for Clinical Research Training and Medical Education at the NIH Clinical Center and Its Impact on Course and Training Program Enrollment and Effectiveness (Clinical Center)

AGENCY: National Institutes of Health, HHS.

ACTION: Notice.

SUMMARY: In compliance with the Paperwork Reduction Act of 1995, the National Institutes of Health (NIH) has submitted to the Office of Management and Budget (OMB) a request for review and approval of the information...
collection listed below. This proposed information collection was previously published in the Federal Register on March 9, 2017, and allowed 60 days for public comment. No public comments were received. The purpose of this notice is to allow an additional 30 days for public comment.

DATES: Comments regarding this information collection are best assured of having their full effect if received within 30-days of the date of this publication.

ADDRESSES: Written comments and/or suggestions regarding the item(s) contained in this notice, especially regarding the estimated public burden and associated response time, should be directed to the: Office of Management and Budget, Office of Regulatory Affairs, OIRA_submission@omb.eop.gov or by fax to 202–395–6974, Attention: Desk Officer for NIH.

FOR FURTHER INFORMATION CONTACT: To request more information on the proposed project or to obtain a copy of the data collection plans and instruments, contact: Dr. Robert M. Lembo, Deputy Director, Office of Clinical Research Training and Medical Education, NIH Clinical Center.

Dated: May 9, 2017.

Laura M. Lee,
Project Clearance Liaison, NIH Clinical Center, National Institutes of Health.

[FR Doc. 2017–10205 Filed 5–18–17; 8:45 am]

BILLING CODE 4140–01–P
DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Prospective Grant of Exclusive Patent License: Chimeric L1/L2 Protein and Virus-Like Particles Based Human Papillomavirus Vaccines

AGENCY: National Institutes of Health, Department of Health and Human Services.

ACTION: Notice.

SUMMARY: The National Cancer Institute, an institute of the National Institutes of Health, Department of Health and Human Services, is contemplating the grant of an exclusive patent license to practice the inventions embodied in the U.S. Patents and Patent Applications listed in the Supplementary Information section of this notice to PathoVax, LLC located in Baltimore, MD.

DATES: Only written comments and/or applications for a license which are received by the National Cancer Institute's Technology Transfer Center on or before June 5, 2017 will be considered.

ADDRESSES: Requests for copies of the patent application, inquiries, and comments relating to the contemplated exclusive license should be directed to: Kevin W. Chang, Ph.D., Senior Technology Transfer Manager, NCI Technology Transfer Center, 9609 Medical Center Drive, RM 1E530 MSC 9702, Bethesda, MD 20892–9702 (for business mail), Rockville, MD 20850–9702 (for facsimile: (240)–276–6910; Facsimile: (240)–276–5504 Email: changke@mail.nih.gov.

SUPPLEMENTARY INFORMATION:

Intellectual Property


The prospective exclusive license territory may be worldwide and the field of use may be limited to the use of Licensed Patent Rights for the following: “Use of Human Papillomavirus Virus (HPV) L1/L2 chimeric proteins and Virus Like Particles (VLPs) for the prevention and/or treatment of cutaneous, mucosal HPV infections and diseases.”

The subject technologies are papillomavirus L2 capsid protein based vaccines against HPV. The L2 protein is the minor papillomavirus capsid protein for papillomaviruses. It is known that antibodies to this protein can neutralize homologous infection. Furthermore, L2 proteins can induce cross-neutralizing antibodies. Specifically, epitopes at the N-terminus of L2 shared by cutaneous and mucosal types of papillomavirus types and by types that infect divergent species are broadly cross-neutralizing. These epitopes at the N-terminus of L2 can be used to elicit cross-neutralizing antibodies against different types of HPV.

This notice is made in accordance with 35 U.S.C. 209 and 37 CFR part 404. The prospective exclusive license will be royalty bearing, and the prospective exclusive license may be granted unless within fifteen (15) days from the date of this published notice, the National Cancer Institute receives written evidence and argument that establishes that the grant of the license would not be consistent with the requirements of 35 U.S.C. 209 and 37 CFR part 404.

Complete applications for a license in the prospective field of use that are filed in response to this notice will be treated as objections to the grant of the contemplated Exclusive Patent License Agreement. Comments and objections submitted to this notice will not be made available for public inspection and, to the extent permitted by law, will not be released under the Freedom of Information Act, 5 U.S.C. 552.