development and use of the Licensed Patent Rights in combination with Licensee's proprietary nanosphere encapsulation technology for the treatment, diagnosis and imaging of cancer tumors and metastases as well as their respective pre-cursor dysplasia states. Licensee's proprietary nanosphere encapsulation technology is understood to consist of: (1) Methods for manipulating the outer proteins of human papillomavirus-derived nanoparticles to create particles targeted to solid tumors and distant metastases; and (2) enhancements for the delivery of particles created by Licensee's proprietary technology.

DATES: Only written comments and/or applications for a license which are received by the NIH Office of Technology Transfer on or before April 25, 2013 will be considered.

ADDRESSES: Requests for copies of the patent application, inquiries, comments, and other materials relating to the contemplated exclusive license should be directed to: Jennifer Wong, M.S., Senior Licensing and Patenting Manager, Cancer Branch, Office of Technology Transfer, National Institutes of Health, 6011 Executive Boulevard, Suite 325, Rockville, MD 20852–3804; Telephone: (301) 435–4633; Facsimile: (301) 402–0220; Email: wongje@mail.nih.gov.

SUPPLEMENTARY INFORMATION: There is extensive literature on the use of viral vectors, particularly those based on the adenovirus, to increase the potency of anti-tumor gene therapy. However, these approaches have had limited success because of limited anti-tumor effects and unacceptable toxicity. This invention describes the use of human papillomavirus pseudoviruses (PsV) as a cancer diagnostic and therapeutic. Preliminary studies showed that PsVs bind to ovarian tumor cells while normal tissues were not affected. PsVs does not infect several other normal intact tissues but continues to selectively infect additional cancer cells. This technology could be an effective anti-tumor therapy because it has shown increased infection of cancer cells with an inability to infect normal cells thereby reducing potential toxicity to patients. In addition to a potential anti-cancer therapeutic, this technology could also be used as a diagnostic tool in the detection of tumor masses. Detection can be achieved through the use of fluorescent dye coupled particles of PsVs that have preferential binding to tumor tissues and not normal tissues.

The prospective exclusive license will be royalty bearing and will comply with the terms and conditions of 35 U.S.C. 209 and 37 CFR Part 404.7. The prospective exclusive license may be granted unless within thirty (30) days from the date of this published notice, the NIH 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 404.7.

Applications for a license in the field of use filed in response to this notice will be treated as objections to the grant of the contemplated exclusive license. 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.

Dated: March 18, 2013.

Richard U. Rodriguez,

Director, Division of Technology Development and Transfer, Office of Technology Transfer, National Institutes of Health.

[FR Doc. 2013-06837 Filed 3-25-13; 8:45 am]

BILLING CODE 4140-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Prospective Grant of Start-Up Exclusive License: Manual Device for Constructing Tissue Micro Arrays and Methods for Making Cryo Arrays for Use in Association With the Device

AGENCY: National Institutes of Health, Public Health Service, HHS.

ACTION: Notice.

SUMMARY: This is notice, in accordance with 35 U.S.C. 209(c)(1) and 37 CFR 404.7(a)(1)(i), that the National Institutes of Health (NIH), Department of Health and Human Services, is contemplating the grant of a start-up exclusive license to practice the inventions embodied in U.S. Patent No. 7,854,899, (E-098-2004/0) filed 08/26/2004 and issued 12/20/2010 entitled "Template Methods and Device for Preparing Sample Arrays"; by Hewitt et al. (NCI); and U.S. Patent No. 6,951,761 9 (E-064-2001/0) filed 08/30/ 2002 and issued 11/04/2005 "Measurements of Multiple Molecules Using a Cryoarray" by Star et al. (NIDDK) to Micatu, Inc. having a place of business at 231 West Water Street, Elmira, NY 14901. The patent rights in this invention have been assigned to the

DATES: Only written comments and/or application for a license that are received by the NIH Office of

United States of America.

Technology Transfer on or before April 10, 2013 will be considered.

ADDRESSES: Requests for a copy of the patent application, inquiries, comments and other materials relating to the contemplated license should be directed to: Tedd Fenn, Office of Technology Transfer, National Institutes of Health, 6011 Executive Boulevard, Suite 325, Rockville, MD 20852–3804; Email: fennea@mail.nih.gov; Telephone: 301–435–5031; Facsimile: 301–402–0220.

SUPPLEMENTARY INFORMATION: The prospective worldwide exclusive license will be royalty bearing and will comply with the terms and conditions of 35 U.S.C. 209 and 37 CFR 404.7. The prospective exclusive license may be granted unless, within fifteen (15) days from the date of this published Notice, NIH 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 404.7.

The patents relate to a device for tissue microarray construction having a block of embedding medium, a platform configured to retain the block, a templates secured to the platform and aligned to guide needles into the embedding block; and methods of making a block containing liquid biological samples that can be frozen and sectioned to make tissue microarray.

The field of use may be limited to the field of devices for construction of tissue microarrays.

Properly filed competing applications for a license filed in response to this notice will be treated as objections to the contemplated license. Comments and objections submitted in response 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.

Dated: March 18, 2013.

Richard U. Rodriguez,

Director, Division of Technology Development & Transfer, Office of Technology Transfer, National Institutes of Health.

[FR Doc. 2013-06835 Filed 3-25-13; 8:45 am]

BILLING CODE 4140-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Substance Abuse and Mental Health Services Administration

Fiscal Year (FY) 2013 Funding Opportunity

AGENCY: Substance Abuse and Mental Health Services Administration, HHS.

ACTION: Notice of intent to award a single source grant to the state of Idaho for a Strategic Prevention Framework State Incentive Grant.

SUMMARY: This notice is to inform the public that the Substance Abuse and Mental Health Services Administration (SAMHSA) intends to award \$1.5 million (total costs) for up to five years to the state of Idaho for a Strategic Prevention Framework State Incentive Grant. This is not a formal request for applications. Assistance will be provided only to the state of Idaho based on the receipt of a satisfactory application that is approved by an independent review group

Funding Opportunity Title: SP-13-

Catalog of Federal Domestic Assistance (CFDA) Number: 93.243. Authority: Section 516 of the Public Health Service Act, as amended.

Justification: Eligibility for this SPF SIG award is limited to the state of Idaho, the only state receiving a Substance Abuse Prevention and Treatment Block Grant (SABG) that has never been awarded a SPF SIG grant from SAMHSA. The SPF SIG grant has already allowed 49 states to strengthen and consolidate their prevention systems and build greater capacity in their communities. SAMHSA/CSAP believes that every state must build prevention capacity and infrastructure to prevent the onset and reduce the progression of substance abuse, including childhood and underage drinking, and to reduce substance abuse-related problems across the nation. Following the SPF five-step process, the state of Idaho will have the opportunity to use SPF SIG funds to develop a comprehensive prevention plan at the state level and support a broad range of sub-recipient communities to implement effective programs, policies and practices to reduce substance abuse and its related problems. By giving a SPF SIG to every state, including Idaho, SAMHSA will have effected nationwide, systemic change in preventing the onset and reducing the progression of substance abuse and substance abuse-related problems nationwide.

Contact: Cathy Friedman, Substance Abuse and Mental Health Services Administration, 1 Choke Cherry Road, Room 8-1097, Rockville, MD 20857; telephone: (240) 276-2316; email: cathy.friedman@samhsa.hhs.gov.

Cathy Friedman,

SAMHSA Public Health Analyst. [FR Doc. 2013-06897 Filed 3-25-13; 8:45 am]

BILLING CODE 4162-20-P

DEPARTMENT OF HOMELAND SECURITY

U.S. Citizenship and Immigration Services

[OMB Control Number 1615-0010]

Agency Information Collection Activities: Nonimmigrant Petition Based on Blanket L Petition; Form I-129S; Revision of a Currently **Approved Collection**

ACTION: 30-Day Notice.

SUMMARY: The Department of Homeland Security (DHS), U.S. Citizenship and Immigration Services (USCIS) will be submitting the following information collection request to the Office of Management and Budget (OMB) for review and clearance in accordance with the Paperwork Reduction Act of 1995. The information collection notice was previously published in the **Federal** Register on January 8, 2013, at 78 FR 1218, allowing for a 60-day public comment period. USCIS did not receive any comment in connection with the 60day notice.

DATES: The purpose of this notice is to allow an additional 30 days for public comments. Comments are encouraged and will be accepted until April 25, 2013. This process is conducted in accordance with 5 CFR 1320.10.

ADDRESSES: Written comments and/or suggestions regarding the item(s) contained in this notice, especially regarding the estimated public burden and associated response time, must be directed to the OMB USCIS Desk Officer via email at

oira submission@omb.eop.gov. The comments submitted to the OMB USCIS Desk Officer may also be submitted to DHS via the Federal eRulemaking Portal Web site at http://www.regulations.gov under e-Docket ID number USCIS-2006-0050 or via email at uscisfrcomment@uscis.dhs.gov. All submissions received must include the agency name and the OMB Control Number 1615–0010.

SUPPLEMENTARY INFORMATION:

Comments

Regardless of the method used for submitting comments or material, all submissions will be posted, without change, to the Federal eRulemaking Portal at www.regulations.gov, and will include any personal information you provide. Therefore, submitting this information makes it public. You may wish to consider limiting the amount of personal information that you provide in any voluntary submission you make

to DHS. For additional information please read the Privacy Act notice that is available via the link in the footer of www.regulations.gov.

Note: The address listed in this notice should only be used to submit comments concerning this information collection. Please do not submit requests for individual case status inquiries to this address. If you are seeking information about the status of your individual case, please check "My Case Status'' online at: https://egov.uscis.gov/cris/ Dashboard.do, or call the USCIS National Customer Service Center at 1-800-375-5283.

Written comments and suggestions from the public and affected agencies should address one or more of the following four points:

(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 submission of responses.

Overview of This Information Collection

(1) Type of Information Collection Request: Revision of a Currently Approved Collection.

(2) Title of the Form/Collection: Nonimmigrant Petition Based on Blanket L Petition.

(3) Agency form number, if any, and the applicable component of the DHS sponsoring the collection: I-129S; **ŪSCIS**.

(4) Affected public who will be asked or required to respond, as well as a brief abstract: Primary: Business or others for profit. This form is used by an employer to classify employees as L-1 nonimmigrant intracompany transferees under a blanket L petition approval. USCIS will use the data on this form to determine eligibility for the requested immigration benefit.

(5) An estimate of the total number of respondents and the amount of time estimated for an average respondent to respond: 75,000 responses at 1.5 hours (1 hour and 30 minutes) per response.