DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Center for Scientific Review; Notice of Closed Meetings

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is hereby given of the following meetings.

The meetings will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title 5 U.S.C., as amended. The grant applications and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.


Date: January 20, 2016.
Time: 1:00 p.m. to 5:30 p.m.
Agenda: To review and evaluate grant applications.
Place: National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892, (Virtual Meeting).

Name of Committee: Center for Scientific Review Special Emphasis Panel; PAR Panel: High Throughput Screening

Date: January 21, 2016.
Time: 8:00 a.m. to 6:00 p.m.
Agenda: To review and evaluate grant applications.

Name of Committee: National Advisory Dental and Craniofacial Research Council.

Date: January 22, 2016.
Time: 8:30 a.m. to 10:10 a.m.
Open: 8:30 a.m. to 10:10 a.m.
Agenda: Report to the Director, NIDCR.
Place: National Institutes of Health, Building 31, 6th Floor, Conference Room 10, 31 Center Drive, Bethesda, MD 20892.

Time: 10:25 a.m. to 12:30 p.m.
Agenda: Special session on Health Disparities.

Place: National Institutes of Health, Building 31, 6th Floor, Conference Room 10, 31 Center Drive, Bethesda, MD 20892.

Time: 2:00 p.m. to Adjournment.
Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, Building 31, 6th Floor, Conference Room 10, 31 Center Drive, Bethesda, MD 20892.

Contact Person: Alicia J. Dombroski, Ph.D., Director, Division of Extramural Activities, National Institute of Dental and Craniofacial Research, National Institutes of Health, Bethesda, MD 20892, 301–594–4805, adombroski@nidcr.nih.gov.

In the interest of security, NIH has instituted stringent procedures for entrance onto the NIH campus. All visitor vehicles, including taxicabs, hotel, and airport shuttles will be inspected before being allowed on campus. Visitors will be asked to show one form of identification (for example, a government-issued photo ID, driver’s license, or passport) and to state the purpose of their visit.

Information is also available on the Institute’s/Center’s home page: http://www.nidcr.nih.gov/about, where an agenda and any additional information for the meeting will be posted when available.

Program Nos. 93.121, Oral Diseases and Disorders Research, National Institutes of Health, HHS

Dated: December 21, 2015.
Natasha M. Copeland,
Program Analyst, Office of Federal Advisory Committee Policy.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Institute of Dental & Craniofacial Research; Notice of Meeting

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. App.), notice is hereby given of a meeting of the National Advisory Dental and Craniofacial Research Council.

The meeting will be open to the public as indicated below, with attendance limited to space available. Individuals who plan to attend and need special assistance, such as sign language interpretation or other reasonable accommodations, should notify the Contact Person listed below in advance of the meeting.

The meeting will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title 5 U.S.C., as amended. The grant applications and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning individuals associated with the grant applications, the disclosure of which would constitute a clearly unwarranted invasion of personal privacy.

Name of Committee: National Advisory Dental and Craniofacial Research Council.

Date: January 22, 2016.

Place: National Institutes of Health, 6701 Rockledge Drive, Bethesda, MD 20892.

Time: 8:00 a.m. to 10:10 a.m.
Open: 8:30 a.m. to 10:10 a.m.
Agenda: Report to the Director, NIDCR.
Place: National Institutes of Health, Building 31, 6th Floor, Conference Room 10, 31 Center Drive, Bethesda, MD 20892.

Time: 10:22 a.m. to 12:30 p.m.
Agenda: Special session on Health Disparities.

Place: National Institutes of Health, Building 31, 6th Floor, Conference Room 10, 31 Center Drive, Bethesda, MD 20892.

Time: 2:00 p.m. to Adjournment.
Agenda: To review and evaluate grant applications.

Place: National Institutes of Health, Building 31, 6th Floor, Conference Room 10, 31 Center Drive, Bethesda, MD 20892.

Contact Person: Alicia J. Dombroski, Ph.D., Director, Division of Extramural Activities, National Institute of Dental and Craniofacial Research, National Institutes of Health, Bethesda, MD 20892, 301–594–4805, adombroski@nidcr.nih.gov.

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Program Nos. 93.846–93.878, 93.892, 93.893, National Institute of Dental and Craniofacial Research

Dated: December 21, 2015.
Natasha M. Copeland,
Program Analyst, Office of Federal Advisory Committee Policy.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Substance Abuse and Mental Health Services Administration

Agency Information Collection Activities; Proposed Collection; Comment Request

In compliance with Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995 concerning opportunity for public comment on proposed collections of information, the Substance Abuse and Mental Health Services Administration (SAMHSA) will publish periodic summaries of proposed projects. To request more information on the proposed projects or to obtain a copy of the information collection plans, call the SAMHSA Reports Clearance Officer on (240) 276–1243.

Comments are invited on: (a) Whether the proposed collections of information are 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) ways to enhance 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.

Section 1926 of the Public Health Service Act [42 U.S.C. 300x–26] stipulates that funding Substance Abuse Prevention and Treatment Block Grant (SABG) agreements for alcohol and drug abuse programs for fiscal year 1994 and subsequent fiscal years require states to have in effect a law providing that it is unlawful for any manufacturer, retailer, or distributor of tobacco products to sell or distribute any such product to any individual under the age of 18. This section further requires that states conduct annual, random, unannounced inspections to ensure compliance with the law; that the state submit annually a report describing the results of the inspections; that the activities carried out by the state to enforce the required law, the success the state has achieved in reducing the availability of tobacco products to individuals under the age of 18, and the strategies to be utilized by the state for enforcing such law during the fiscal year for which the grant is sought.

Before making an award to a State under the SABG, the Secretary must make a determination that the state has maintained compliance with these requirements. If a determination is made that the state is not in compliance, penalties shall be applied. Penalties ranged from 10 percent of the Block Grant in applicable year 1 (FFY 1997 SABG Applications) to 40 percent in applicable year 4 (FFY 2000 SABG Applications) and subsequent years. Respondents include the 50 states, the District of Columbia, the Commonwealth of Puerto Rico, the U.S. Virgin Islands, Guam, American Samoa, the Commonwealth of the Northern Mariana Islands, Palau, Micronesia, and the Marshall Islands.

Regulations that implement this legislation are at 45 CFR 96.130, are approved by OMB under control number 0930–0163, and require that each state submit an annual Synar report to the Secretary describing their progress in complying with section 1926 of the PHS Act. The Synar report, due December 31 following the fiscal year for which the state is reporting, describes the results of the inspections and the activities carried out by the state to enforce the required law; the success the state has achieved in reducing the availability of tobacco products to individuals under the age of 18; and the strategies to be utilized by the state for enforcing such law during the fiscal year for which the grant is sought.

SAMHSA’s Center for Substance Abuse Prevention will request OMB approval of revisions to the current report format associated with Section 1926 (42 U.S.C. 300x–26). The report format is not changing significantly. Any changes in either formatting or content are being made to simplify the reporting process for the states and to clarify the information as the states report it; both outcomes will facilitate consistent, credible, and efficient monitoring of Synar compliance across the states. All of the information required in the new report format is already being collected by the states. Specific changes are listed below:

Clarification Changes

To decrease the need for supplemental questions and reporting, additional instruction has been included in 3 portions of the report.

In Section I (Compliance Progress), the following clarification changes are being made with respect to the Annual Synar Report:

**Question 1b: Changes to state law**—This question asks about changes in state laws that impact the state’s protocol for conducting Synar inspections and has been edited to include an option for changes to state law concerning changes in the definition of tobacco products. Many states are changing the definition of tobacco products in their state laws to include electronic nicotine delivery systems, which would impact the types of products that could be included in Synar surveys.

**Question 1c: Changes to state law**—This question asks about changes to state youth access to tobacco laws and has been edited to include an option for changes to state law concerning additional product categories to their youth tobacco access law. While some states have changed the definition in the state law to include electronic nicotine delivery systems, smokeless tobacco, and other tobacco products, other states have added these products as additional product categories in addition to tobacco products.

**Question 2: Describe how the Annual Synar Report and the state plan were made public prior to submission of the ASR.** This question asks states to describe how they make their ASR public prior to submission. States have been asked to provide a Web address and the date the ASR was posted to that Web address if they choose to post the ASR on an agency Web site. The ASR format has been clarified to provide a separate text box to enter both of these pieces of information.

**Questions 4d–j:** Coordination with Agency that Receives the FDA State Enforcement Contract—These close-ended questions ask the state to list the agency that is under contract to the FDA to enforce federal youth access laws, to describe the relationship between the state’s Synar program and this agency, and to identify if the state uses data from the FDA enforcement inspections for the Synar survey. This question has been edited to include skip logic and response options if a state does not have a current contract with the FDA.

**Questions 5b, 5c, 5d, 5e, 5f:** Enforcement Agencies, Evidence of Enforcement and Frequency of Enforcement—These questions have been clarified so it is clear that they refer to enforcement of state youth access laws, and not federal or local youth access laws. In addition, these questions have been re-ordered (but the wording has not been changed) to improve logical flow of the questions. In addition, question 5e has been edited to include separate response options to allow states to describe each of the additional activities listed in the question stem to encourage states to describe each of those activities fully.

In Section II (Intended Use), the following clarification change is being made:

**Question 3—State Challenges:** This question asks states to identify and describe their challenges in implementing the Synar program. This question has been edited to include separate response options to allow states to describe each of the challenges listed in the question stem to encourage states to describe each of the challenges fully and to make targeted technical assistance requests.

In Appendix C (Synar Survey Inspection Protocol Summary), the following change is being made:

**Title:** The title of this Appendix has been edited to reflect that it is the summary of the state’s inspection protocol and that the Appendix itself is not detailed enough to serve as the entirety of the state’s inspection protocol.

**Questions 4—Type of Tobacco Products—**These questions, which ask the state to define the type of tobacco products requested during Synar inspections and to describe the protocol for tobacco type selection, have been edited to separate the options of including small cigars and cigarillos and to add the option of including electronic nicotine delivery systems or electronic cigarettes.

**Questions 5a and b—**The previous question 5 has been separated into two sections to provide ensure states are
able to fully describe the methods used to recruit, select and train adult supervisors for the survey separately from the methods used to recruit, select, and train youth inspectors.

Content Changes

The content of the Synar Report has changed little. The content changes that have been made address the need to (1) clarify the intent of information requested via the addition of clarifying questions, (2) reduce the need for State Project Officers to ask additional questions to supplement the originally submitted Report. These additions and changes are essential to SAMHSA’s ability to adequately assess state and jurisdictional compliance with the Synar regulation.

In Section I (Compliance Progress), the following changes are being made with respect to the Annual Synar Report:

Question 6: Changes to the sampling methodology—This question asks states if their sampling methodology has changed from the previous year. If there has been a change, a sub-question has been added to document how that change was communicated to SAMHSA. Since this change requires prior approval, a state who has not received prior approval will have the opportunity to discuss the process that was used to determine a change needed to be made. Existing questions 9a, 9b, and 9c have been renumbered to account for this new sub-question.

In Appendix B (Synar Survey Sampling Methodology), the following changes are being made:

Question 4—Vending machine inclusion in Synar Survey—This question, which asks if vending machines are included in the Synar survey and the reasons for their elimination if they are not included. Because many states have a contract with the FDA and is actively enforcing the vending machine requirements of the Family Smoking Prevention and Tobacco Control Act, some states that include vending machines in their sampling protocols do not sample any because there are few eligible vending machines remaining on their list frame. A second part has been added to this question to determine how vending machines are sampled.

There are no changes to Forms 1–5 or Appendix D.

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¹ Red Lake Indian Tribe is not subject to tobacco requirements.

Send comments to Summer King, SAMHSA Reports Clearance Officer, Room 2–1057, One Choke Cherry Road, Rockville, MD 20857 OR email a copy to summer.king@samhsa.hhs.gov. Written comments should be received by February 26, 2016.

Summer King, Statistician.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Substance Abuse and Mental Health Services Administration

Agency Information Collection Activities: Proposed Collection; Comment Request

In compliance with Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995 concerning opportunity for public comment on proposed collections of information, the Substance Abuse and Mental Health Services Administration (SAMHSA) will publish periodic summaries of proposed projects. To request more information on the proposed projects or to obtain a copy of the information collection plans, call the SAMHSA Reports Clearance Officer on (240) 276–1243.

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Proposed Project: Now Is the Time (NITT)—Minority Fellowship Program (MFP) Evaluation—New

SAMHSA is conducting a national evaluation of the No Is the Time (NITT) initiative, which includes separate programs—the Minority Fellowship Program-Youth (MFP–Y), the Minority Fellowship Program-Addiction Counselors (MFP–AC), Project AWARE (Advancing Wellness and Resilience in Education)—State Educational Agency, and Healthy Transitions. These programs are united by their focus on capacity building, system change, and workforce development.

The NITT–MFP (Youth and Addiction Counselors) programs, which are the focus of this data collection, represent a response to the fourth component of President Obama’s NITT Initiative: increasing access to mental health/behavioral health services. The purpose of the NITT–MFP programs is to improve behavioral health care outcomes for underserved racially and ethnically diverse populations by