state child support enforcement agencies with potential reprogramming at varying times due to future changes in either Part A or Part B, the Administration for Children and Families is resubmitting an unchanged information collection package and requesting an extension to the current OMB approval of NMSN Part A to synchronize with the expiration date of NMSN Part B.

Respondents: State child support enforcement agencies, employers, and health plan administrators.

### ANNUAL BURDEN ESTIMATES

<table>
<thead>
<tr>
<th>Instrument</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Average burden hours per response</th>
<th>Total burden hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>National Medical Support Notice</td>
<td>54</td>
<td>97,775</td>
<td>0.17</td>
<td>897,574.50</td>
</tr>
</tbody>
</table>

Estimated Total Annual Burden Hours: 897,574.50.

In compliance with the requirements of Section 506(c)(2)(A) of the Paperwork Reduction Act of 1995, 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, 370 L'Enfant Promenade SW., Washington, DC 20447, 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. 2012–16029 Filed 6–29–12; 8:45 am]

**BILLING CODE 4184–01–P**

### DEPARTMENT OF HEALTH AND HUMAN SERVICES

#### Food and Drug Administration

[Docket No. FDA–2012–N–0063]

**Sami Arshak Yanikian: Debarment Order**

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

**ACTION:** Notice.

**SUMMARY:** The Food and Drug Administration (FDA) is issuing an order under the Federal Food, Drug, and Cosmetic Act (the FD&C Act) debarring Sami Arshak Yanikian for 10 years from providing services in any capacity to a person that has an approved or pending drug product application. FDA bases this order on a finding that Mr. Yanikian was convicted of two counts of introducing unapproved new drugs into interstate commerce, which relates to the development or approval, including the process for development or approval, of drug products and to the regulation of drug products under the FD&C Act. In addition, the type of conduct that served as the basis for Mr. Yanikian’s convictions undermine the process for the regulation of drugs. Mr. Yanikian was given notice of the proposed debarment and an opportunity to request a hearing within the timeframe prescribed by regulation. Mr. Yanikian failed to respond. Mr. Yanikian’s failure to respond constitutes a waiver of his right to a hearing concerning this action.

**DATES:** This order is effective July 2, 2012.

**ADDRESSES:** Submit applications for termination of debarment to the Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.

**FOR FURTHER INFORMATION CONTACT:** Kenny Shade, Office of Regulatory Affairs, Food and Drug Administration, 12420 Parklawn Dr., Rockville, MD 20857, 301–796–4640.

### SUPPLEMENTARY INFORMATION:

**I. Background**

Section 306(b)(2)(B)(i)(I) of the FD&C Act (21 U.S.C. 335a(b)(2)(B)(i)(I)) permits FDA to debar an individual if it finds that the individual has been convicted of a misdemeanor under Federal law for conduct relating to the development or approval, including the process for development or approval, of any drug product or otherwise relating to the regulation of drug products under the FD&C Act, and if FDA finds that the type of conduct that served as the basis for the conviction undermines the process for the regulation of drugs. On June 29, 2011, Mr. Yanikian was found guilty of two counts of introduction of an unapproved drug in interstate commerce, in violation of sections 301(d), 505(a), and 303(a)(1) of the FD&C Act (21 U.S.C. 331(d), 355(a), 333(a)(1)) and of aiding and abetting, in violation of 18 U.S.C. 2(b), and the U.S. District Court for the Central District of California entered judgment against Mr. Yanikian for the misdemeanor offenses of introduction of an unapproved drug in interstate commerce and aiding and abetting.

The FDA’s finding that debarment is appropriate is based on the misdemeanor convictions referenced herein. The factual basis for the conviction is as follows: On March 17, 2005, FDA sent Mr. Yanikian a warning letter regarding his marketing and sale of Novil natural formulation for atrial fibrillation, Super Nasal Drops, and Sams No Tinnitus Formulation. The warning letter described the claims Mr. Yanikian’s Web site was making and informed him that his claims caused the products to be “drugs” as defined by the FD&C Act because they were intended to cure, mitigate, treat, or prevent disease. Mr. Yanikian was informed that his products were “new drugs” and that a new drug could not be introduced or delivered for introduction into interstate commerce unless an FDA-approved application was in effect for it. The...
warning letter additionally noted that none of the products described had an approved application and that their introduction or delivery for introduction into interstate commerce violated section 301(d) of the FD&C Act. Mr. Yanikian was advised to immediately correct the violations.

In response, on April 11, 2005, Mr. Yanikian wrote a reply letter to FDA in which he stated that the products were mailed for sale outside the United States to hospitals that deal with natural health products. He further noted that his products were not intended for sale as over-the-counter or for single individuals in the United States until they were approved by FDA.

Despite knowing that he was not allowed to sell these unapproved new drugs in the United States without FDA approval, and despite his repeated representations to FDA that he was not selling his products to customers in the United States, Mr. Yanikian subsequently sold his unapproved new drug products to an undercover agent in November 2005, and again in November 2006, in violation of sections 301(d), 505(a), and 303(a)(1) of the FD&C Act and 18 U.S.C. 2(b).

As a result of his conviction, on April 3, 2012, FDA sent Mr. Yanikian a notice by certified mail proposing to debar him for 10 years from providing services in any capacity to a person that has an approved or pending drug product application. FDA subsequently confirmed that Mr. Yanikian personally received the notice on April 11, 2012. The proposal was based on a finding, under section 306(b)(2)(B)(i)(I) of the FD&C Act that Mr. Yanikian was convicted of two counts of a misdemeanor under Federal law. The proposal was based on a finding, under section 306(b)(2)(B)(i)(II) of the FD&C Act that Mr. Yanikian was convicted of two counts of a misdemeanor under Federal law. In the notice, FDA found that the conduct underlying these Federal misdemeanor convictions relates to the development or approval, including the process for development or approval, of drug products and relates to the regulation of drug products under the FD&C Act and 18 U.S.C. 2(b).

As a result of the foregoing finding, Mr. Yanikian is debarred for 10 years from providing services in any capacity to a person with an approved or pending drug product application under sections 505, 512, or 502 of the FD&C Act (21 U.S.C. 355, 360b, or 382), or under section 351 of the Public Health Service Act (42 U.S.C. 262), effective (see DATES), (see sections 306(c)(1)(B), (c)(2)(A)(ii), and 201(dd) of the FD&C Act (21 U.S.C. 335(a)(1)(B), (c)(2)(A)(ii), and 321(dd)). Any person with an approved or pending drug product application who knowingly employs or retains as a consultant or contractor, or otherwise uses the services of Mr. Yanikian, in any capacity during Mr. Yanikian’s debarment, will be subject to civil money penalties (section 307(a)(6) of the FD&C Act (21 U.S.C. 335(a)(6))). If Mr. Yanikian provides services in any capacity to a person with an approved or pending drug product application during his period of debarment he will be subject to civil money penalties (section 307(a)(7) of the FD&C Act). In addition, FDA will not accept or review any abbreviated new drug applications submitted by or with the assistance of Mr. Yanikian during his period of debarment (section 306(c)(1)(B) of the FD&C Act).

Any application by Mr. Yanikian for termination of debarment under section 306(d)(1) of the FD&C Act should be identified with Docket No. FDA–2012–N–0063 and sent to the Division of Dockets Management (see ADDRESSES). All such submissions are to be filed in four copies. The public availability of information in these submissions is governed by 21 CFR 10.20(j).

Publicly available submissions may be seen in the Division of Dockets Management between 9 a.m. and 4 p.m., Monday through Friday.


Armando Zamora,
Acting Director, Office of Enforcement, Office of Regulatory Affairs.

[FR Doc. 2012–16156 Filed 6–29–12; 8:45 am]

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.

Name of Committee: Center for Scientific Review Special Emphasis Panel; Review of Behavioral and Social HIV/AIDS Applications.

Date: July 20, 2012.

Time: 2 p.m. to 5 p.m.

Agenda: To review and evaluate grant applications.

Place: St. Gregory Hotel, 2033 M Street NW, Washington, DC 20036.

Contact Person: Mark P Rubert, Ph.D., Scientific Review Officer, Center for Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 5218, MSC 7852, Bethesda, MD 20892, 301–435–1775, rubertm@csr.nih.gov.

Name of Committee: Center for Scientific Review Special Emphasis Panel; PAR 12–010: NIH Competitive Revision Applications for Research Relevant to the Family Smoking Prevention and Tobacco Control Act (R01).

Date: July 20, 2012.

Time: 3 p.m. to 4 p.m.

Agenda: To review and evaluate grant applications.

Place: St. Gregory Hotel, 2033 M Street NW., Washington, DC 20036.

Contact Person: Mark P Rubert, Ph.D., Scientific Review Officer, Center for Scientific Review, National Institutes of Health, 6701 Rockledge Drive, Room 5218, MSC 7852, Bethesda, MD 20892, 301–435–1775, rubertm@csr.nih.gov.

Name of Committee: Center for Scientific Review Special Emphasis Panel; Muscle Small Business.

Date: July 23, 2012.