

program priorities, geographic balance, and the achievement of national objectives.

c. The overall balance of the ICRC program in addressing the three phases of injury control (prevention, acute care, and rehabilitation); the control of injury among populations who are at increased risk, including racial/ethnic minority groups, the elderly and children; the major causes of intentional and unintentional injury; and the major disciplines of injury control (such as biomechanics, epidemiology, and behavioral science).

d. Budgetary considerations. The ACIPC will recommend annual funding levels as detailed under the heading, Availability of Funds.

3. Continued Funding

Continuation awards within the project period will be made on the basis of the availability of funds and the following criteria:

a. The accomplishments of the current budget period show that the applicant's objectives as prescribed in the yearly work plans are being met;

b. The objectives for the new budget period are realistic, specific, and measurable;

c. The methods described will clearly lead to achievement of these objectives;

d. The evaluation plan allows management to monitor whether the methods are effective by having clearly defined process, impact, and outcome objectives, and the applicant demonstrates progress in implementing the evaluation plan; and

e. The budget request is clearly explained, adequately justified, reasonable, and consistent with the intended use of grant funds.

H. Other Requirements

Technical Reporting Requirements

Provide CDC with original plus two copies of

1. progress report annually;
2. financial status report, no more than 90 days after the end of the budget period; and
3. final financial status report and performance report, no more than 90 days after the end of the project period.

Send all reports to the Grants Management Specialist identified in the Where to Obtain Additional Information section of this announcement.

The following additional requirements are applicable to this program. For a complete description of each see Addendum 1 in the application kit.

AR-1 Human Subjects Certification

AR-2 Requirements for inclusion of Women and Racial and Ethnic Minorities in Research

AR-3 Animal Subjects Requirements

AR-9 Paperwork Reduction Act Requirements

AR-10 Smoke-Free Workplace Requirement

AR-11 Healthy People 2010

AR-12 Lobbying Restrictions

AR-13 Prohibition on Use of CDC funds for Certain Gun Control Activities

AR-20 Conference Activities within Grants/Cooperative Agreements

I. Authority and Catalog of Federal Domestic Assistance Number

This program is authorized under Sections 301, 391, 392, 393, and 394 of the Public Health Service Act, [42 U.S.C. 241, 280b, 280b-1, 280b-1a, and 280b-2] as amended. Catalog of Federal Domestic Assistance number is 93.136.

J. Where To Obtain Additional Information

For this announcement and other CDC announcements, see the CDC home page on the Internet: <http://www.cdc.gov>.

To receive additional written information and to request an application kit, call 1-888-GRANTS4 (1-888-472-6874). You will be asked to leave your name and address and will be instructed to identify the Announcement number of interest. A complete program description and information on application procedures are contained in the application package.

If you have questions after reviewing the contents of all the documents, business management assistance may be obtained from: Joanne Wojcik, Grants Management Specialist, Grants Management Branch, Procurement and Grants Office, Centers for Disease Control and Prevention (CDC), Colgate Building, Room 3000, 2920 Brandywine Road, Atlanta, GA 30341-4146, Telephone 770-488-2717, Internet address: jcw6@cdc.gov

Programmatic assistance may be obtained from: Tom Voglesonger, Program Manager, Office of Research Grants, National Center for Injury Prevention and Control, Centers for Disease Control and Prevention (CDC), 4770 Buford Highway, NE., (K58), Atlanta, GA 30341-3724, Telephone 770-488-4265, Internet address: tdv1@cdc.gov.

Please refer to Announcement 01007 when requesting information and submitting an application.

Dated: June 7, 2000.

Henry S. Cassell, III,

Acting Director, Procurement and Grants Office, Centers for Disease Control and Prevention (CDC).

[FR Doc. 00-14831 Filed 6-12-00; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Advisory Committee for Energy-Related Epidemiologic Research: Conference Call Meeting

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92-463), the Centers for Disease Control and Prevention (CDC) announces the following conference call meeting.

Name: Advisory Committee for Energy-Related Epidemiologic Research (ACERER), Subcommittee for Management Review of the Chernobyl Studies (SMRCS).

Time and Date: 1 p.m.-1:30 p.m., June 26, 2000.

Place: The conference call will originate at the National Center for Environmental Health (NCEH), CDC, in Atlanta, Georgia. Please see **SUPPLEMENTARY INFORMATION** for details on accessing the conference call.

Status: Open to the public, limited only by the availability of telephone ports.

Purpose: This subcommittee is charged with providing guidance to the scientific reviewers and staff, and reporting back to the full ACERER on the charge from the Department and Congress to assess the management, goals, and objectives of the National Cancer Institute Chernobyl studies.

Matters To Be Discussed: The conference call agenda is to reach consensus on the review and report submitted by the SRMCS.

Agenda items are subject to change as priorities dictate.

SUPPLEMENTARY INFORMATION: This conference call is scheduled to begin at 1 p.m., Eastern Time. To participate in the conference call, please dial 1-877-322-9654 and enter conference code 970943. You will then be automatically connected to the call.

CONTACT PERSON FOR MORE INFORMATION: Michael J. Sage, Executive Secretary, ACERER, and Acting Deputy Director, NCEH, CDC, 4770 Buford Highway, NE, (F-28), Atlanta, Georgia 30341-3724, telephone 770/488-7002, fax 770/488-7015.

The Director, Management Analysis and Services Office, has been delegated the authority to sign **Federal Register** notices pertaining to announcements of meetings and other committee management activities for both CDC and the Agency for Toxic Substances and Disease Registry.

Dated: June 8, 2000.

Carolyn J. Russell,

Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

[FR Doc. 00-14945 Filed 6-12-00; 8:45 am]

BILLING CODE 4163-18-M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Advisory Committee on Immunization Practices: Meeting

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92-463), the Centers for Disease Control and Prevention (CDC) announces the following committee meeting:

Name: Advisory Committee on Immunization Practices (ACIP).

Times and Dates: 8:30 a.m.-6:30 p.m., June 21, 2000. 8 a.m.-4:30 p.m., June 22, 2000.

Place: Four Points Hotel by Sheraton, 1850 Cotillion Drive, Atlanta, Georgia 30338.

Status: Open to the public, limited only by the space available.

Purpose: The Committee is charged with advising the Director, CDC, on the appropriate uses of immunizing agents. In addition, under 42 U.S.C. § 1396s, the Committee is mandated to establish and periodically review and, as appropriate, revise the list of vaccines for administration to vaccine-eligible children through the Vaccines for Children (VFC) program, along with schedules regarding the appropriate periodicity, dosage, and contraindications applicable to the vaccines.

Matters To Be Discussed: The agenda will include a discussion on the ACIP policies and procedures; ACIP recommendations for the pneumococcal conjugate vaccine; vaccine additive: aluminum update; vaccine additive: thimerosal; vaccines and autism; bioterrorism working group; general recommendations; anaphylaxis after MMR due to gelatin; progress report on vaccine identification standards initiative; status of high-speed needle-free jet injectors for mass vaccination campaigns; update on Geneva meeting on rotavirus vaccination; Vaccines for Children program update; adult working group: pneumococcal polysaccharide update; CDC/FDA report on two dose schedule for hepatitis B for adolescent; update on influenza vaccine supply; Global Alliance for Vaccines and Immunization: progress in supporting global immunization programs and introduction of new vaccines; Nabi an

update from the Food and Drug Administration; update from the National Center for Infectious Diseases; update from the National Immunization Program; update from the Vaccine Injury Compensation Program; update from the National Vaccine Program. Other matters of relevance among the committee's objectives may be discussed.

Agenda items are subject to change as priorities dictate.

Contact Person for More Information:

Gloria A. Kovach, Program Analyst, Epidemiology and Surveillance Division, National Immunization Program, CDC, 1600 Clifton Road, NE, m/s E61, Atlanta, Georgia 30333. Telephone 404/639-8096.

The Director, Management Analysis and Services office has been delegated the authority to sign **Federal Register** notices pertaining to announcements of meetings and other committee management activities for both the Centers for Disease Control and Prevention and the Agency for Toxic Substances and Disease Registry.

Dated: June 8, 2000.

Carolyn J. Russell,

Director, Management Analysis and Services Office, Centers for Disease Control and Prevention.

[FR Doc. 00-14944 Filed 6-12-00; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 92N-0412]

Rajaram K. Matkari; Conviction Reversal; Final Order Terminating Debarment

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 act), terminating the debarment of Rajaram K. Matkari, 1304 Riverglen Way, Berthoud, CO 80513. FDA is issuing this order because the U.S. District Court for the District of Maryland issued a Writ of Error Coram Nobis, reversing Mr. Matkari's conviction and Mr. Matkari applied for termination of his debarment on this basis.

EFFECTIVE DATE: June 13, 2000.

ADDRESSES: Submit written comments to the Dockets Management Branch (HFA-305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. Comments are to be identified with the docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT: Leanne Cusumano, Center for Drug

Evaluation and Research (HFD-7), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301-594-2041.

SUPPLEMENTARY INFORMATION: In a notice published in the **Federal Register** of October 20, 1993 (58 FR 54156), Rajaram K. Matkari was permanently debarred from providing services in any capacity to a person with an approved or pending drug product application (sections 306(c)(1)(B) and (c)(2)(A)(ii) and 201(dd) of the act (21 U.S.C. 335a(c)(1)(B) and (c)(2)(A)(ii) and 21 U.S.C. 321(dd)). The debarment was based on FDA's finding that Mr. Matkari was convicted of a felony under Federal law for conduct relating to the development or approval of a drug product, or otherwise relating to the regulation of a drug product (section 306(a)(2)(A) and (a)(2)(B) of the act). Mr. Matkari, the former Vice President for Regulatory Affairs and Product Development of Pharmaceutical Basics, Inc. (PBI), pled guilty to and was sentenced on July 28, 1989, for giving an unlawful gratuity, a felony offense under 18 U.S.C. 201(c)(1)(A). The basis for this conviction was Mr. Matkari's payment of approximately \$2,000 to an FDA chemistry review branch chief who was responsible for supervising the chemists who reviewed PBI's applications to determine whether those applications met certain statutory standards for approval.

On February 22, 2000, the U.S. District Court for the District of Maryland issued an order granting Mr. Matkari's petition for a Writ of Error Coram Nobis in his criminal case. A copy of the court's order is available in Docket No. 92N-0412. By this order, the court reversed Mr. Matkari's conviction. On April 18, 2000, Mr. Matkari petitioned for termination of debarment under section 306(d)(3)(B)(i) of the act, as amended by the Generic Drug Enforcement Act. Section 306(d)(3)(B)(i) of the act states that "If the conviction which served as the basis for the debarment of an individual under subsection (a)(2) * * * is reversed, the Secretary shall withdraw the order of debarment."

Accordingly, the Senior Associate Commissioner for Policy, Planning, and Legislation, under section 306(d)(3)(B)(i) of the act and under authority delegated to him (21 CFR 5.20), is issuing this order withdrawing the order of permanent debarment of Rajaram K. Matkari, thereby allowing him to provide services in any capacity to a person with an approved or pending drug product application. Rajaram K. Matkari's debarment is terminated