**Matters To Be Discussed:** The panel will gather information regarding the use of vaccinia (smallpox) vaccine.

**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 for both the Centers for Disease Control and Prevention and the Agency for Toxic Substances and Disease Registry.

**Dated:** April 29, 2002.

**John Burckhardt,**

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

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**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Centers for Disease Control and Prevention**

**Citizens Advisory Committee on Public Health Service Activities and Research at Department of Energy (DOE) Sites: Savannah River Site Health Effects Subcommittee (SRSHES)**

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), and the Agency for Toxic Substances and Disease Registry (ATSDR) announce the following meeting.

**Name:** Citizens Advisory Committee on Public Health Service Activities and Research at Department of Energy (DOE) Sites: Savannah River Site Health Effects Subcommittee (SRSHES).

**Time and Date:** 8:30 a.m.–4:45 p.m., June 6, 2002.

**Place:** Radisson Riverfront Hotel Augusta, 2 10th Street, Augusta, Georgia 30901, telephone (706) 722–8900, fax (706) 724–0044, www.radisson.com.

**Status:** Open to the public, limited only by the space available. The meeting room accommodates approximately 50 people.

**Background:** Under a Memorandum of Understanding (MOU) signed in December 1990 with DOE, and replaced by MOUs signed in 1996 and 2000, the Department of Health and Human Services (HHS) was given the responsibility and resources for conducting analytic epidemiologic investigations of residents of communities in the vicinity of DOE facilities, workers at DOE facilities, and other persons potentially exposed to radiation or to potential hazards from non-nuclear energy production use. HHS delegated program responsibility to CDC.

In addition, a memo was signed in October 1990 and renewed in November 1992, 1996, and in 2000, between ATSDR and DOE. The MOU delineates the responsibilities and procedures for ATSDR’s public health activities at DOE sites required under sections 104, 105, 107, and 120 of the Comprehensive Environmental Response, Compensation, and Liability Act (CERCLA or “Superfund”). These activities include health consultations and public health assessments at DOE sites listed on, or proposed for, the Superfund National Priorities List and at sites that are the subject of petitions from the public; and other health-related activities such as epidemiologic studies, health surveillance, exposure and disease registries, health education, substance-specific applied research, emergency response, and preparation of toxicological profiles.

**Purpose:** This subcommittee is charged with providing advice and recommendations to the Director, CDC, and the Administrator, ATSDR, regarding community concerns pertaining to CDC’s and ATSDR’s public health activities and research at this DOE site. The purpose of this meeting is to provide a forum for community interaction and to a vehicle for community concerns to be expressed as advice and recommendations to CDC and ATSDR.

**Matters to be Discussed:** Agenda items include: Update on Department of Labor/Department of Energy Workers Occupational Illness Compensation Program; Overview of Process on Risk Based Screening Criteria and Where it is Going; International Atomic Energy Administration Recommendations; ATSDR Needs Assessment; American College of Preventive Medicine Initiatives Update; and ATSDR/Tritium Health Consult on Potential Tritium Exposures at SRSHES. Agenda items are subject to change as priorities dictate.

**Contact Person for More Information:** Phillip Green, Executive Secretary, SRSHES, Radiation Studies Branch, Division of Environmental Hazards and Health Effects, NCEH, CDC, 1600 Clifton Road, NE, (E–39), Atlanta, Georgia 30333, telephone (404) 498–1800, fax (404) 498–1811.

The Food and Drug Administration (FDA) is announcing two public workshops, both entitled “An FDA/Industry Dialog on the Application Submission Process: Public Workshop.” The purpose of the public workshops is to discuss common application deficiencies and strategies to avoid these deficiencies leading to faster approval times. Staff from the Center for Biologics Evaluation and Research (CBER) will provide general information on the review process and options to consider. CBER staff will also lead discussion groups designed to respond to your general issues and questions on submission requirements. These discussion groups will be established based on the input provided to CBER on your issues relative to the purpose of this workshop.

**Date and Time:** Send registration and issues by May 17, 2002, for the May 29, 2002, workshop and by June 14, 2002, for the June 26, 2002, workshop. See table 1 of this document.

**Location:** See table 1 of this document.

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**TABLE 1**

<table>
<thead>
<tr>
<th>Meeting address</th>
<th>Dates and local time</th>
<th>FDA contact person</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hyatt Regency Bethesda, 1 Bethesda Metro Center, Bethesda, MD 20814, 301–657–1234.</td>
<td>May 29, 2002, from 9 a.m. to 5 p.m.</td>
<td>Kathy Eberhart.</td>
</tr>
</tbody>
</table>
Contact Persons:

For information about this notice: Michael D. Anderson, Center for Biologics Evaluation and Research (CBER) (HFM–17), Food and Drug Administration, 1401 Rockville Pike, Rockville, MD 20852, 301–827–6210, FAX 301–594–1944, e-mail: Andersonm@cber.fda.gov.

For information about the workshop and registration: Kathy Eberhart, Center for Biologics Evaluation and Research (HFM–42), Food and Drug Administration, 1401 Rockville Pike, Rockville, MD 20852, 301–827–2000, FAX 301–827–3079, e-mail eberhart@cber.fda.gov.

Procedure: Mail or fax your registration information (including name, professional degree, title, e-mail address, firm name, address, telephone, and fax number) to Kathy Eberhart, Center for Biologics Evaluation and Research (HFM–42), Food and Drug Administration, 1401 Rockville Pike, Rockville, MD 20852, 301–827–2000, FAX 301–827–3079, by May 17, 2002, for the May 29, 2002, workshop, and by June 14, 2002, for the June 26, 2002, workshop. There is no registration fee for the public workshops. Space is limited, therefore interested parties are encouraged to register early. There will be no onsite registration.

If you need special accommodations due to a disability, please contact Kathy Eberhart (see Contact Persons) at least 7 days in advance.

CBER is requesting that you submit your issues related to application deficiencies and approval times before the workshop. There will be an opportunity to submit additional issues and questions at the end of the morning sessions. Mail or fax your issues to Kathy Eberhart (see Contact Persons) by May 17, 2002, for the May 29, 2002, workshop and by June 14, 2002, for the June 26, 2002, workshop.

Dated: April 26, 2002.

Margaret M. Dotzel,
Associate Commissioner for Policy.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Health Resources and Services Administration

Emergency Medical Services for Children; National Trauma Registry for Children Demonstration Project

AGENCY: Health Resources and Services Administration, HHS.

ACTION: Notice of availability of funds.

SUMMARY: The Health Resources and Services Administration (HRSA) announces that up to $400,000 in fiscal year (FY) 2002 funds is available to fund two grants to assess the feasibility and value of developing a National Trauma Registry for Children (NTRC) as part of the Emergency Medical Services for Children (EMSC) program. Two grant awards are planned for 2002. One award of up to $150,000 will be made for a Data Identification, Collection and Use Planning Project to identify and prioritize data elements necessary for a pediatric trauma registry, develop sustainable data collection procedures that take into account the needs of all potential users, including but not limited to, State Emergency Medical Services (EMS) Offices, pediatric trauma surgeons, pre-hospital and hospital health care providers; and develop procedures for accessing the data. A second award of up to $250,000 will be for a Registry Design and Technology Planning Project to assess existing trauma registries, assess the feasibility of obtaining and linking information from them, and assess the feasibility of collecting additional data elements identified by the Data Identification, Collection and Use Planning Project. A pediatric trauma registry could be useful to clinicians and program planners for understanding the epidemiology and improving the clinical management of pediatric trauma. These demonstration project grants will be awarded under the program authority of the Public Health Service Act, Title XIX, Section 1910 (42 U.S.C. 300w–9), and will be administered by the Maternal and Child Health Bureau (MCHB), HRSA. Projects will be approved for up to a 2-year period. However, funding beyond FY 2002 is contingent upon the availability of funds.

DATES: Applicants are expected to notify MCHB’s Division of Child, Adolescent, and Family Health of their intent by June 14, 2002. The deadline for receipt of applications is July 15, 2002. Applications will be considered “on time” if they are either received on or before the deadline date or postmarked on or before the deadline date. The projected award date is September 3, 2002.

ADDRESSES: To receive a complete application kit, applicants may telephone the HRSA Grants Application Center at 1–877–477–2123 (1–877–HRSAS–123) or register on-line at: http://www.hrsa.gov/g_order3.htm directly. The Emergency Medical Services for Children program uses the standard Form PHS 5161–1 (rev. 7/00) for applications (approved under OMB No. 0920–0428). Applicants must use Catalog of Federal Domestic Assistance (CFDA) #93.127J when requesting application kits. The CFDA is a Government wide compendium of enumerated Federal programs, project services, and activities that provide assistance. All applications must be mailed or delivered to Grants Management Officer, MCHB: HRSA Grants Application Center, 901 Russell Avenue, Suite 450, Gaithersburg, MD 20879: telephone 1–877–477–2123: E-mail: hrsagac@hrsa.gov.

Necessary application forms and an expanded version of this Federal Register notice may be downloaded in either Microsoft Office 2000 or Adobe Acrobat format (.pdf) from the MCHB Home Page at http://www.mchb.hrsa.gov. Please contact Joni Johns, at 301–443–2088, or jjohns@hrsa.gov, if you need technical assistance in accessing the MCHB Home Page via the Internet.


LETTER OF INTENT: Notification of intent to apply can be made in one of three ways: Cindy Doyle, RN, telephone, 301–443–3888; email: cdoyle@hrsa.gov, mail,