

*Subject:* Amendment of the Amateur Service Rules to Provide for Greater Use of Spread Spectrum Communications Technologies (WT Docket No. 97-12, RM-8737).

*Number of Petitions Filed:* 1.  
Federal Communications Commission.

**Magalie Roman Salas,**  
*Secretary.*

[FR Doc. 00-1463 Filed 1-20-00; 8:45 am]

**BILLING CODE 6712-01-M**

## FEDERAL RESERVE SYSTEM

### Sunshine Act Meeting

**AGENCY HOLDING THE MEETING:** Board of Governors of the Federal Reserve System.

**TIME AND DATE:** 0:00 a.m., Wednesday, January 26, 2000.

**PLACE:** Marriner S. Eccles Federal Reserve Board Building, 20th and C Streets, N.W., Washington, D.C. 20551

**STATUS:** Closed.

**MATTERS TO BE CONSIDERED:** 1. Personnel actions (appointments, promotions, assignments, reassignments, and salary actions) involving individual Federal Reserve System employees.

2. Any matters carried forward from a previously announced meeting.

**CONTACT PERSON FOR MORE INFORMATION:** Lynn S. Fox, Assistant to the Board; 202-452-3204.

**SUPPLEMENTARY INFORMATION:** You may call 202-452-3206 beginning at approximately 5 p.m. two business days before the meeting for a recorded announcement of bank and bank holding company applications scheduled for the meeting; or you may contact the Board's Web site at <http://www.federalreserve.gov> for an electronic announcement that not only lists applications, but also indicates procedural and other information about the meeting.

Dated: January 19, 2000.

**Robert deV. Frierson,**

*Associate Secretary of the Board.*

[FR Doc. 00-1618 Filed 1-19-00; 12:05 pm]

**BILLING CODE 6210-01-P**

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### National Committee on Vital and Health Statistics: Meeting

Pursuant to the Federal Advisory Committee Act, the Department of Health and Human Services announces the following advisory committee meeting.

**NAME:** National Committee on Vital and Health Statistics (NCVHS).

**JOINT MEETING:** Subcommittee on Standards and Security and Workgroup on Computer-based Patient Records.xxx

**TIME AND DATE:** 9 a.m. to 5:30 p.m., January 31, 2000; 8:30 a.m. to 1:30 p.m., February 1, 2000.

**PLACE:** Room 705A, Hubert H. Humphrey Building, 200 Independence Avenue, SW, Washington, DC 20201.

**STATUS:** Open.

**PURPOSE:** The Subcommittee and Working Group will review the first draft of its report to the HHS Secretary on standards for patient medical records information as required by the Health Insurance Portability and Accountability Act of 1996 (HIPAA). The Subcommittee will review the recommendations contained in the report, specify proposed revisions in the draft report, and confirm the work plan for the completion of the report. On the second day, the Subcommittee will be updated on HHS activities related to the implementation of the administrative simplification provisions of HIPAA, will review its overall work plan for the year 2000, and review the first draft of the annual NCVHS report to Congress on implementation of the HIPAA administrative simplification provisions.

**Notice:** In the interest of security, the Department has instituted stringent procedures for entrance to the Hubert H. Humphrey building by non-government employees. Thus, persons without a government identification card will need to have the guard call for an escort to the meeting.

**CONTACT PERSON FOR MORE INFORMATION:** Substantive program information as well as summaries of meetings and a roster of committee members may be obtained from J. Michael Fitzmaurice, Ph.D., Senior Science Advisor for Information Technology, Agency for Health Care Research and Quality, 2101 East Jefferson Street, #600, Rockville, MD 20852, phone: (301) 594-3938; or Marjorie S. Greenberg, Executive Secretary, NCVHS, National Center for Health Statistics, Centers for Disease Control and Prevention, Room 1100, Presidential Building, 6525 Belcrest Road, Hyattsville, Maryland 20782, telephone (301) 458-4245. Information also is available on the NCVHS home page of the HHS website, <http://www.ncvhs.hhs.gov/> where an agenda for the meeting will be posted when available.

Dated: January 13, 2000.

**James Scanlon,**

*Director, Division of Data Policy, Office of the Assistant Secretary for Planning and Evaluation.*

[FR Doc. 1460 File 1-20-00; 8:45 am]

**BILLING CODE 4151-04-M**

## DEPARTMENT OF HEALTH AND HUMAN SERVICES

### Centers for Disease Control and Prevention

[30DAY-13-00]

### Agency Forms Undergoing Paperwork Reduction Act Review

The Centers for Disease Control and Prevention (CDC) publishes a list of information collection requests under review by the Office of Management and Budget (OMB) in compliance with the Paperwork Reduction Act (44 U.S.C. Chapter 35). To request a copy of these requests, call the CDC Reports Clearance Officer at (404) 639-7090. Send written comments to CDC, Desk Officer; Human Resources and Housing Branch, New Executive Office Building, Room 10235, Washington, DC 20503. Written comments should be received within 30 days of this notice.

### Proposed Project

Supplement to HIV/AIDS Surveillance (SHAS) Project—New—The National Center for HIV, STD and TB Prevention (NCHSTP). NCHSTP is proposing revisions to the currently-approved questionnaire for the Supplement to HIV/AIDS Surveillance (SHAS) project (OMB No. 0920-0262). This questionnaire provides detailed information about persons with HIV infection which continues to be of significant interest to public health, community, minority groups and affected groups. Since 1989, the CDC, in collaboration with 12 state and local health agencies, has collected data through the national Supplemental HIV/AIDS Surveillance project. The objective of this project is to obtain increased descriptive information on persons with newly-reported HIV and AIDS infections, including sociodemographic characteristics, risk behaviors, use of health care services, sexual and substance abuse behaviors, minority issues and adherence to therapy. The revised questionnaire will address important emerging surveillance and prevention issues, particularly those related to the recent advances in therapy for HIV infection. This information supplements routine, national HIV/AIDS surveillance and is used to

improve CDC's understanding of minority issues related to the epidemic

of HIV, target educational efforts to prevent transmission, and improve

services for persons with HIV infection. The total annual burden hours are 3500.

DATA FOR CALENDAR YEAR 1998

Respondents	No. of respondents	No. of responses/re-spondent	Avg. burden of response (in hrs.)
Georgia .....	292	1	.75
California .....	301	1	.75
Michigan .....	82	1	.75
New Mexico .....	81	1	.75
Arizona .....	165	1	.75
Colorado .....	139	1	.75
Connecticut .....	229	1	.75
Delaware .....	43	1	.75
Florida .....	430	1	.75
S. Carolina .....	270	1	.75
New Jersey .....	86	1	.75
Washington .....	160	1	.75

Dated: January 14, 2000.

**Nancy Cheal, Ph.D.,**

*Acting Associate Director for Policy, Planning and Evaluation, Centers for Disease Control and Prevention (CDC).*

[FR Doc. 00-1450 Filed 1-20-00; 8:45 am]

**BILLING CODE 4163-18-P**

**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Centers for Disease Control and Prevention**

[30DAY-11-00]

**Agency Forms Undergoing Paperwork Reduction Act Review**

The Centers for Disease Control and Prevention (CDC) publishes a list of information collection requests under review by the Office of Management and Budget (OMB) in compliance with the Paperwork Reduction Act (44 U.S.C. Chapter 35). To request a copy of these requests, call the CDC Reports Clearance Officer at (404) 639-7090. Send written comments to CDC, Desk Officer; Human Resources and Housing Branch, New Executive Office Building, Room 10235, Washington, DC 20503. Written comments should be received within 30 days of this notice.

**Proposed Project**

Specifications and Tests for Approval of Coal Mine Dust Personal Sampler Units—(0920-0148)—Extension—National Institute for Occupational Safety and Health (NIOSH)—Under the Federal Coal Mine Health & Safety Act of 1977, PL91-173 (amended the Federal Coal Mine & Safety Act of 1969), mine operators must periodically sample mine atmospheres and submit the samples to the Mine Safety and Health Administration (MSHA). The Act

states that sampling equipment used must be approved by the Secretaries of the Department of Health and Human Services (DHHS) and the Department of Labor (DOL). Concurrent permissibility approval for electrical intrinsic safety is provided by MSHA while NIOSH certifies the performance under Title 30 CFR Part 74. Under this regulation, certification applicants are required to submit detailed parts lists, drawings, and inspection instructions, along with the personal sampler unit to be tested. These materials are provided to NIOSH along with a letter from the applicant requesting certification. After NIOSH has tested the unit and certifies the performance of the equipment, a certificate of approval is issued to the manufacturer. Should the equipment be disapproved, a letter is sent to the manufacturer outlining the details of the defects resulting in disapproval, with suggestions for possible corrections to the unit. Certificates of approval are accompanied by photographs of designs for approval labels to be affixed to each coal mine dust personal sampler unit. Use of the approval label is authorized only on sampler units which conform strictly with the drawings and specifications upon which the certificate of approval is based. Changes or modifications in the unit after certification will result in the manufacturer requesting extensions of approval through the original certification process.

The information is used by NIOSH to fulfill its legislatively-mandated responsibilities to evaluate and approve coal mine dust personal sampler units (CMDPSU) submitted for certification and approval actions (30 U.S.C. 957 and 961). Before NIOSH grants a certification, it must have sufficient evidence of safety and adequate

performance. The parts listing, engineering drawings, and inspection instructions submitted are used by NIOSH to assure that descriptions of tested units are fully detailed and that future units produced are equivalent to those currently certified. Without the information specified in 30 CFR Part 74, NIOSH will be unable to adequately evaluate CMDPSU safety and efficacy, and to determine if functional changes were made in the manufacture of certified products. The total annual burden hours are 44.

Respondents	No. of re-spondents	No. of re-sponses/respondent	Avg. burden of response (in hrs.)
Manufacturer ....	1	1	44

Dated: January 14, 2000.

**Nancy Cheal, Ph.D.,**

*Acting Associate Director for Policy, Planning and Evaluation, Centers for Disease Control and Prevention (CDC).*

[FR Doc. 00-1451 Filed 1-20-00; 8:45 am]

**BILLING CODE 4163-18-P**

**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Centers for Disease Control and Prevention**

[30DAY-12-00]

**Agency Forms Undergoing Paperwork Reduction Act Review**

The Centers for Disease Control and Prevention (CDC) publishes a list of information collection requests under review by the Office of Management and Budget (OMB) in compliance with the