Philadelphia, PA 19106, 215–597–2120, ext. 4003, FAX: 215–597–5798, e-mail: *mfalcone@ora.fda.gov.* 

*Registration*: Send registration information (including name, title, firm name, address, telephone, and fax number) and \$460 (member) or \$535 (non-member) registration fee made payable to SoCRA, P.O. Box 101, Furlong, PA 18925. To register via the Internet go to *http://www.socra.org/ FDA\_Conference.htm.* FDA has verified the Web site address, but is not responsible for subsequent changes to the Web site after this document publishes in the **Federal Register**.

Registrar will also accept payment by major credit cards. For more information on the meeting, or for questions on registration, contact 800 -SoCRA92 (800–762–7292), or 215–345– 7369 or via e-mail to *socramail@aol.com*. Attendees are responsible for their own accommodations. To make reservations at the Pittsburgh Marriott Center City Hotel at the reduced conference rate, contact the Pittsburgh Marriott Center City Hotel at 412–471–4000 or 888–456– 6600 or by fax at hotel FAX: 412–281– 4797 before June 3, 2003.

The registration fee will be used to offset the expenses of hosting the conference, including meals, refreshments, meeting rooms, and materials. Space is limited, therefore interested parties are encouraged to register early. Limited onsite registration may be available. Please arrive early to ensure prompt registration.

If you need special accommodations due to a disability, please contact Marie Falcone at least 7 days in advance of the workshop.

**SUPPLEMENTARY INFORMATION:** The "FDA Clinical Trials Statutory and Regulatory Requirements" workshop helps fulfill the Department of Health and Human Services' and FDA's important mission to protect the public health by educating researchers on proper conduct of clinical trials. FDA has made education of the research community a high priority to assure the quality of clinical data and protect research subjects.

The workshop helps to implement the objectives of section 406 of the FDA Modernization Act (21 U.S.C. 393) and the FDA Plan for Statutory Compliance, which includes working more closely with stakeholders and ensuring access to needed scientific and technical expertise. The workshop also furthers the goals of the Small Business Regulatory Enforcement Fairness Act (Public Law 104–121) by providing outreach activities by Government agencies directed to small businesses.

Dated: May 20, 2003.

#### Jeffrey Shuren,

Assistant Commissioner for Policy. [FR Doc. 03–13192 Filed 5–27–03; 8:45 am] BILLING CODE 4160–01–S

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

# Health Resources and Services Administration

#### Agency Information Collection Activities: Proposed Collection: Comment Request

In compliance with the requirement for opportunity for public comment on proposed data collection projects (section 3506(c)(2)(A) of Title 44, United States Code, as amended by the Paperwork Reduction Act of 1995, Public Law 104–13), the Health **Resources and Services Administration** (HRSA) publishes periodic summaries of proposed projects being developed for submission to OMB under the Paperwork Reduction Act of 1995. To request more information on the proposed project or to obtain a copy of the data collection plans and draft instruments, call the HRSA Reports Clearance Officer on (301) 443–1129.

*Comments are invited 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) 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.

#### Proposed Project: Web-based Semi Annual Report (SAR) (OMB No. 0915– 0262)—Revision

The Health Resources and Services Administration (HRSA), Bureau of Primary Health Care (BPHC) plans to collect the annual reporting requirements for the primary care grantees funded by BPHC using the web-based Semi Annual Report (SAR). The SAR includes reporting requirements for grantees of the following primary care programs: State Primary Care Associations and State Primary Care Offices. Authorizing legislation is found in Section 330(m) of the Public Health Service Act, as amended.

BPHC collects data on its programs to ensure compliance with legislative mandates and to report to Congress and policy makers on program accomplishments. To meet these objectives, BPHC requires a core set of information collected semi-annually that is appropriate for monitoring and evaluating performance and reporting on annual trends. The SAR has been a valuable instrument for collecting this information from grantees. The SAR provides data on services, characteristics of populations, leveraged funds, and services that fall within the scope of the grant.

The estimated burden is as follows:

Form	Number of respondents	Responses per respondent	Hours per response	Total burden hours
SAR	103	1	18	1854

Send comments to Susan G. Queen, Ph.D., HRSA Reports Clearance Officer, Room 14–45, Parklawn Building, 5600 Fishers Lane, Rockville, MD 20857. Written comments should be received within 60 days of this notice. Dated: May 20, 2003.

Jane M. Harrison,

Director, Division of Policy Review and Coordination. [FR Doc. 03–13224 Filed 5–27–03; 8:45 am] BILLING CODE 4165–15–P

### DEPARTMENT OF HEALTH AND HUMAN SERVICES

#### Health Resources and Services Administration

### Availability of Funds

**AGENCY:** Health Resources and Services Administration (HRSA), HHS.