Agreement will be required to receive a copy of any pending patent application.

Dated: August 21, 2006.

James D. Seligman,

Chief Information Officer, Centers for Disease Control and Prevention.

[FR Doc. E6–14423 Filed 8–29–06; 8:45 am] BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

Advisory Committee on Special Studies Relating to the Possible Long-Term Health Effects of Phenoxy Herbicides and Contaminants (Ranch Hand Advisory Committee); Notice of Meeting

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

This notice announces a forthcoming meeting of a public advisory committee of the Food and Drug Administration (FDA). The meeting will be open to the public.

Name of Committee: Advisory Committee on Special Studies Relating to the Possible Long-Term Health Effects of Phenoxy Herbicides and Contaminants (Ranch Hand Advisory Committee).

General Function of the Committee:
To advise the Secretary and the
Assistant Secretary for Health
concerning its oversight of the conduct
of the Ranch Hand study by the U.S. Air
Force and provide scientific oversight of
the Department of Veterans Affairs (VA)
Army Chemical Corps Vietnam Veterans
Health Study, and other studies in
which the Secretary or the Assistant
Secretary for Health believes
involvement by the committee is
desirable.

Date and Time: The meeting will be held on September 7, 2006, from 8:30 a.m. to 4 p.m.

Location: Food and Drug Administration, 5630 Fishers Lane, rm. 1066, Rockville, MD 20857.

Contact Person: Leonard Schechtman, National Center for Toxicological Research (HFT–10), Food and Drug Administration, 5600 Fishers Lane, rm. 16–85, Rockville, MD 20857, 301–827–6696, or FDA Advisory Committee Information Line, 1–800–741–8138 (301–443–0572 in the Washington, DC area), code 3014512560. Please call the Information Line for up-to-date information on this meeting.

Agenda: The committee will discuss the following items: (1) Summary of

Dioxin 2006 presentations; (2) summary of Technical Reports and manuscripts; (3) summary of transition activities.

Procedure: Interested persons may present data, information, or views, orally or in writing, on issues pending before the committee. Written submissions may be made to the contact person on or before September 4, 2006. Oral presentations from the public will be scheduled between approximately 11:30 a.m. to 12:30 p.m. Time allotted for each presentation may be limited. Those desiring to make formal oral presentations should notify the contact person and submit a brief statement of the general nature of the evidence or arguments they wish to present, the names and addresses of proposed participants, and an indication of the approximate time requested to make their presentation on or before September 4, 2006.

Persons attending FDA's advisory committee meetings are advised that the agency is not responsible for providing access to electrical outlets.

FDA welcomes the attendance of the public at its advisory committee meetings and will make every effort to accommodate persons with physical disabilities or special needs. If you require special accommodations due to a disability, please contact Leonard Schechtman at least 7 days in advance of the meeting.

FDA regrets that it was unable to publish this notice 15 days prior to the September 7, 2006, Advisory Committee on Special Studies Relating to the Possible Long-Term Health Effects of Phenoxy Herbicides and Contaminants (Ranch Hand Advisory Committee) meeting. Because the agency believes there is some urgency to bring these issues to public discussion and qualified members of the Advisory Committee on Special Studies Relating to the Possible Long-Term Health Effects of Phenoxy Herbicides and Contaminants (Ranch Hand Advisory Committee) were available at this time, the Commissioner of Food and Drugs concluded that it was in the public interest to hold this meeting even if there was not sufficient time for the customary 15-day public notice.

Notice of this meeting is given under the Federal Advisory Committee Act (5 U.S.C. app. 2).

Dated: August 23, 2006.

Randall W. Lutter,

Associate Commissioner for Policy. [FR Doc. E6–14371 Filed 8–29–06; 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 the Office of Management and Budget (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) The proposed collection of information for the proper performance of the functions of the agency; (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: Children's Hospitals Graduate Medical Education Payment Program (CHGME PP) (OMB No. 0915– 0247)—Revision

The CHGME PP was enacted by Public Law 106–129 to provide Federal support for graduate medical education (GME) to freestanding children's hospitals. This legislation attempts to provide support for GME comparable to the level of Medicare GME support received by other, non-children's hospitals. The legislation indicates that eligible children's hospitals will receive payments for both direct and indirect medical education. Direct payments are designed to offset the expenses associated with operating approved graduate medical residency training programs and indirect payments are designed to compensate hospitals for expenses associated with the treatment of more severely ill patients and the additional costs relating to teaching residents in such programs.

Data are collected on the number of full-time equivalent residents in applicant children's hospitals' training programs to determine the amount of direct and indirect medical education payments to be distributed to participating children's hospitals. Indirect medical education payments will also be derived from a formula that requires the reporting of discharges, beds, and case mix index information from participating children's hospitals. Hospitals will be requested to submit such information in an annual application. Hospitals will also be requested to submit data on the number

of full-time equivalent residents a second time during the Federal fiscal year to participate in the reconciliation payment process.

The estimated annual burden is as follows:

Form	Number of respondents	Responses per respondent	Total number of responses	Hours per response	Total burden hours
HRSA 99–1 (Initial Application)	60	1	60	26	1,560
HRSA 99-1 (Reconciliation Application)	60	1	60	8	480
HRSA 99–2 (Initial Application)	60	1	60	15	900
HRSA 99–2 (Reconciliation Application)	60	1	60	5	300
HRSA 99–3 (Initial Application)	60	1	60	.25	15
HRSA 99–3 (Reconciliation Application)	60	1	60	.25	15
HRSA 99-4 (Reconciliation Application)	60	1	60	14	840
HRSA 99-5 (Initial Application)	60	1	60	.25	15
HRSA 99-5 (Reconciliation Application)	60	1	60	.25	15
Total	60		60		4,140

Send comments to Susan G. Queen, Ph.D., HRSA Reports Clearance Officer, Room 10–33, Parklawn Building, 5600 Fishers Lane, Rockville, MD 20857. Written comments should be received within 60 days of this notice.

Dated: August 23, 2006.

Cheryl R. Dammons,

Director, Division of Policy Review and Coordination.

[FR Doc. E6–14411 Filed 8–29–06; 8:45 am] BILLING CODE 4165–15–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Submission for OMB Review; Comment Request; National Network of Tobacco Cessation Quitlines Evaluation

Summary: Under the provisions of Section 3507(a)(1)(D) of the Paperwork Reduction Act of 1995, the National Cancer Institute, the National Institutes of Health has submitted to the Office of Management and Budget (OMB) a request to review and approve the information collection listed below. This proposed information collection was previously published in the Federal Register on January 27, 2006 (page 4595) and allowed 60 days for public comment. No public comments were received. The purpose of this notice is to allow an additional 30 days for public comment. The National Institutes of Health may not conduct or sponsor, and the respondent is not required to respond to, an information collection that has been extended, revised, or

implemented on or after October 1, 1995, unless it displays a currently valid OMB control number.

Proposed Collection

Title: Evaluation of the HHS National Network of Tobacco Cessation Quitlines Initiative.

Type of Information Collection Request: New.

Need and Use of Information Collection: In February 2004, the U.S. Department of Health and Human Services announced plans for a national network of tobacco cessation quitlines to provide all smokers in the United States access to the support and latest information to help them quit. To provide the highest level of assistance to smokers across the country who wants to quit, NCI established a new toll-free telephone number (1-800-QUIT-NOW) on November 8, 2004. The aim of the National Network of Tobacco Cessation Quitlines (NNTCQ) initiative (the Initiative) is to strengthen service delivery; provide a mechanism for integration and implementation of state, regional, and national campaigns; and increase healthcare utilization by minority and medically underserved populations. NCI, CDC, and other state, private industry, and partner organizations (the North American Ouitline Consortium) have created the infrastructure and a coordinated mechanism to offer cessation services to the American public. The Initiative seeks to enhance existing state-managed quitlines and to encourage the establishment of quitlines in states without them. It is expected that successful implementation of the Initiative will foster partnerships across

state quitlines for technology transfer, sharing of effective practices, and understanding patterns of use and reach to special populations, thereby ensuring a sustained level of effectiveness over time. The goal of this evaluation is to monitor the implementation of the Initiative, assess its impact on key stakeholders, and examine its implications for public health. To that end, this study will conduct a series of in-depth key informant telephone interviews and selected site visits with state tobacco control officers, quitline administrators and counseling staff. Representatives of organizations and individuals that partner with quitlines, such as community health organizations or health care providers, will also be interviewed. The findings will provide valuable information concerning the development and implementation of the NNTQC initiative as a potential model for Federal-State partnerships, the impact on building and enhancing state quitline capacity, and implications for the state tobacco control community.

The annual reporting burden is presented in exhibit 1, below.

Frequency of Response: One occasion.

Affected Public: State agencies,
businesses or other for-profit, non-profit
associations.

Type of Respondents: Federal and state employees, health services providers, administrators and researchers.

The annual reporting burden is as follows:

Estimated Number of Respondents: 228

Estimated Number of Responses per Respondent: 1.