that each assisted reproductive technology (ART) program shall annually report to the Secretary through the Centers for Disease Control and Prevention: (1) Pregnancy success rates achieved by such ART program, and (2) the identity of each embryo laboratory used by such ART program and whether the laboratory is certified or has applied for such certification under the Act. The required information is currently reported by ART programs to CDC as specified in the Assisted Reproductive Technology (ART) Program Reporting System (OMB No. 0920-0556, exp. 9/ 30/2009). CDC seeks to extend OMB approval for a period of three years and incorporate a minor change in wording to one question. In addition, the revised total burden estimate includes an anticipated increase in the number of respondents and a slight decrease in the average number of responses per respondent. The burden estimate per response has also been revised to

include an adjustment for data validation procedures.

The currently approved program reporting system, also known as the National ART Surveillance System (NASS), includes information about all ART cycles initiated by any of the ART programs in the United States. An ART cycle is considered to begin when a woman begins taking ovarian stimulatory drugs or starts ovarian monitoring with the intent of having embryos transferred. The system also collects information about the pregnancy outcome of each cycle, as well as a number of data items deemed important to explain variability in success rates across ART programs and across individuals. Data elements and definitions currently in use reflect CDC's consultations with representatives of the Society for Assisted Reproductive Technology (SART), the American Society for Reproductive Medicine, and RESOLVE, the National Infertility Association (a

national, nonprofit consumer organization), as well as a variety of individuals with expertise and interest in this field.

Respondents are the 480 ART programs in the United States. Approximately 420 clinics are expected to report an average of 286 ART cycles each. Ten percent of responding clinics will be randomly selected to participate in full validation of selected ART cycle records and an abbreviated validation of selected cycles resulting in live birth. All information is collected electronically. Respondents have the option of entering data directly into a Web-based NASS interface or of transmitting system-compatible files extracted from other record systems. The ART program reporting system allows CDC to publish an annual report to Congress as specified by the FCSRCA and to provide information needed by consumers.

There are no costs to respondents other than their time.

ESTIMATED ANNUALIZED BURDEN HOURS

Respondents	Form name	Number of respondents	Number of responses per respondent	Average burden per response (in hours)	Total burden (in hours)
ART Programs	NASS	420	286	38/60	76,076

Dated: January 2, 2009.

Maryam I. Daneshvar,

Acting Reports Clearance Officer, Centers for Disease Control and Prevention.

[FR Doc. E9–405 Filed 1–12–09; 8:45 am]

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Subcommittee on Procedures Reviews, Advisory Board on Radiation and Worker Health (ABRWH), National Institute for Occupational Safety and Health (NIOSH)

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

Time and Date: 9:30 a.m.-5 p.m., January 28, 2009.

Place: Cincinnati Airport Marriott, 2395 Progress Drive, Hebron, Kentucky 41018. Telephone (859) 334–4611, Fax (859) 334–4619.

Status: Open to the public, but without a public comment period. To access by

conference call dial the following information 1(866) 659–0537, Participant Pass Code 9933701.

Background: The Advisory Board was established under the Energy Employees Occupational Illness Compensation Program Act of 2000 to advise the President on a variety of policy and technical functions required to implement and effectively manage the new compensation program. Key functions of the Advisory Board include providing advice on the development of probability of causation guidelines that have been promulgated by the Department of Health and Human Services (HHS) as a final rule; advice on methods of dose reconstruction which have also been promulgated by HHS as a final rule; advice on the scientific validity and quality of dose estimation and reconstruction efforts being performed for purposes of the compensation program; and advice on petitions to add classes of workers to the Special Exposure Cohort (SEC).

In December 2000, the President delegated responsibility for funding, staffing, and operating the Advisory Board to HHS, which subsequently delegated this authority to CDC. NIOSH implements this responsibility for CDC. The charter was issued on August 3, 2001, renewed at appropriate intervals, and will expire on August 3, 2009.

Purpose: The Advisory Board is charged with (a) Providing advice to the Secretary, HHS, on the development of guidelines under Executive Order 13179; (b) providing advice to the Secretary, HHS, on the

scientific validity and quality of dose reconstruction efforts performed for this program; and (c) upon request by the Secretary, HHS, advise the Secretary on whether there is a class of employees at any Department of Energy facility who were exposed to radiation but for whom it is not feasible to estimate their radiation dose, and on whether there is reasonable likelihood that such radiation doses may have endangered the health of members of this class. The Subcommittee on Procedures Reviews was established to aid the Advisory Board in carrying out its duty to advise the Secretary, HHS, on dose reconstruction. It will be responsible for overseeing, tracking, and participating in the reviews of all procedures used in the dose reconstruction process by the NIOSH Office of Compensation Analysis and Support (OCAS) and its dose reconstruction contractor.

Matters To Be Discussed: The agenda for the Subcommittee meeting includes: a discussion of proposed new versions of the computer-assisted telephone interview scripts and procedures NIOSH uses to interview claimants at the outset of the dose reconstruction process; a discussion of ORAUT-OTIB-0054 ("Fission and Activation Product Assignment for Internal Dose-Related Gross Beta and Gross Gamma Analyses'') and ORAUT-OTIB-0066 ("Calculation of Dose from Intakes of Special Tritium Compounds"); and, a continuation of the comment-resolution process for other dose reconstruction procedures under review by the Subcommittee.

The agenda is subject to change as priorities dictate.

In the event an individual cannot attend, written comments may be submitted. Any written comments received will be provided at the meeting and should be submitted to the contact person below well in advance of the meeting.

Contact Person for More Information: Theodore Katz, Executive Secretary, NIOSH, CDC, 1600 Clifton Road, Mailstop E–20, Atlanta, Georgia 30333, Telephone (513) 533–6800, Toll Free 1 (800) CDC–INFO, email ocas@cdc.gov.

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: January 7, 2009.

Elaine L. Baker,

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

[FR Doc. E9–524 Filed 1–12–09; 8:45 am]

BILLING CODE 4163-18-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Administration for Children and Families

Submission for OMB Review; Comment Request

Title: Child Care Quarterly Case Record Report—ACF–801. OMB No.: 0970–0167.

Description: Section 658K of the Child Care and Development Block Grant Act of 1990 (Pub. L. 101–508, 42 U.S.C. 9858) requires that States and Territories submit monthly case-level data on the children and families receiving direct services under the Child Care and Development Fund. The implementing regulations for the statutorily required reporting are at 45 CFR 98.70. Case-level reports, submitted quarterly or monthly (at grantee option), include monthly sample or full population case-level data. The data elements to be included in these reports

are represented in the ACF-801. ACF uses disaggregate data to determine program and participant characteristics as well as costs and levels of child care services provided. This provides ACF with the information necessary to make reports to Congress, address national child care needs, offer technical assistance to grantees, meet performance measures, and conduct research. Consistent with the statute and regulations, ACF requests extension of the ACF-801. With this extension, ACF is proposing several changes and clarifications to the reporting requirements and instructions.

Respondents: States, the District of Columbia, and Territories including Puerto Rico, Guam, the Virgin Islands, American Samoa, and the Northern Mariana Islands.

ANNUAL BURDEN ESTIMATES

Instrument	Number of re- spond- ents	Number of re- sponses per re- spondent	Average burden hours per response	Total bur- den hours
ACF-801	56	4	20	4,480

Estimated Total Annual Burden Hours: 4.480

Additional Information: Copies of the proposed collection may be obtained by writing to the Administration for Children and Families, Office of Administration, Office of Information Services, 370 L'Enfant Promenade, SW., Washington, DC 20447, Attn: ACF Reports Clearance Officer. All requests should be identified by the title of the information collection. E-mail address: infocollection@acf.hhs.gov.

OMB Comment: OMB is required to make a decision concerning the collection of information between 30 and 60 days after publication of this document in the Federal Register. Therefore, a comment is best assured of having its full effect if OMB receives it within 30 days of publication. Written comments and recommendations for the proposed information collection should be sent directly to the following: Office of Management and Budget, Paperwork Reduction Project, Fax: 202-395-6974, Attn: Desk Officer for the Administration for Children and Families

Dated: January 8, 2009.

Janean Chambers,

Reports Clearance Officer.

[FR Doc. E9–447 Filed 1–12–09; 8:45 am]

BILLING CODE 4184-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration [Docket No. FDA-2008-N-0038]

Advisory Committees; Tentative Schedule of Meetings for 2009

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug
Administration (FDA) is announcing a
tentative schedule of forthcoming
meetings of its public advisory
committees for 2009. During 1991, at the
request of the Commissioner of Food
and Drugs (the Commissioner), the
Institute of Medicine (the IOM)
conducted a study of the use of FDA's
advisory committees. In its final report,
one of the IOM's recommendations was
for the agency to publish an annual

tentative schedule of its meetings in the **Federal Register**. This publication implements the IOM's recommendation.

FOR FURTHER INFORMATION CONTACT:

Theresa L. Green, Advisory Committee Oversight and Management Staff (HF– 4), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301–827–1220.

SUPPLEMENTARY INFORMATION: The IOM, at the request of the Commissioner, undertook a study of the use of the FDA's advisory committees. In its final report in 1992, one of the IOM's recommendations was for FDA to adopt a policy of publishing an advance yearly schedule of its upcoming public advisory committee meetings in the Federal Register; FDA has implemented this recommendation. The annual publication of tentatively scheduled advisory committee meetings will provide both advisory committee members and the public with the opportunity, in advance, to schedule attendance at FDA's upcoming advisory committee meetings. Because the schedule is tentative, amendments to this notice will not be published in the Federal Register. However, changes to the schedule will be posted on the FDA