

EXHIBIT 1.—ESTIMATE OF COST BURDEN TO RESPONDENTS

Data collection effort	Number of estimated respondents	Estimated time per respondent in hours	Estimated total burden hours	Average hourly wage rate	Estimated annual cost burden to respondents
Draft narrative response to RFP by Collaborative Manager	50	8	400	\$34.67	\$13,868
Narrative reviews by 2 members of Collaborative executive committee	100	1	100	57.90	5,790
Narrative revisions by Collaborative Manager	50	8	400	34.67	13,868
Assembly of narrative with any supporting documents by Collaborative Assistant	50	2	100	12.58	1,258
Total	250	1,000	34,784

This information collection will not impose a cost burden on the respondent beyond that associated with the above estimates of the time needed to provide the application-requested information. No additional costs to respondents are anticipated, e.g., for capital equipment, software, etc.

Estimated Costs to the Federal Government

The total cost to the government for its proposal review activity is estimated to be \$500,000 annually.

Request for Comments

In accordance with the above-cited legislation, comments on the AHRQ's information collection are requested with regard to any of the following: (a) Whether the proposed collection of information is necessary for the proper performance of health care improvement and information dissemination functions of AHRQ, including whether the information requested will have practical utility; (b) the accuracy of AHRQ's estimate of burden (including hours and costs) of the proposed collection(s) of information; (c) ways to enhance the quality, utility, and clarity of information to be collected; and, (d) ways to minimize the burden of the collection of information upon the respondents, including the use of automated collection techniques or other forms of information technology.

Comments submitted in response to this notice will be summarized and included in the request for OMB approval of the proposed information collection. All comments will become a matter of public record.

Dated: May 1, 2007

Carolyn M. Clancy,
Director.

[FR Doc. 07-2268 Filed 5-7-07; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Agency for Healthcare Research and Quality

Notice of Meetings

In accordance with section 10(d) of the Federal Advisory Committee Act as amended (5 U.S.C., Appendix 2), the Agency for Healthcare Research and Quality (AHRQ) announces meetings of scientific peer review groups. The subcommittees listed below are part of the Agency's Health Services Research Initial Review Group Committee.

The subcommittee meetings will be closed to the public in accordance with the Federal Advisory Committee Act, section 10(d) of 5 U.S.C., Appendix 2 and 5 U.S.C. 552b(c)(6). Grant applications are to be reviewed and discussed at these meetings. These discussions are likely to involve information concerning individuals associated with the applications, including assessments of their personal qualifications to conduct their proposed projects. This information is exempt from mandatory disclosure under the above-cited statutes.

1. *Name of Subcommittee:* Health Care Research Training.

Date: June 14-15, 2007 (Open from 8 a.m. to 8:15 a.m. on June 14 and closed for remainder of the meeting).

Place: Agency for Healthcare Research and Quality (AHRQ), John Eisenberg Conference Center, 540 Gaither Road, Rockville, Maryland 20850.

2. *Name of Subcommittee:* Health Systems Research.

Date: June 14-15, 2007 (Open from 8 a.m. to 8:15 a.m. on June 14 and closed for remainder of the meeting).

Place: Agency for Healthcare Research and Quality (AHRQ), John Eisenberg Conference Center, 540 Gaither Road, Rockville, Maryland 20850.

3. *Name of Subcommittee:* Health Care Quality and Effectiveness Research.

Date: June 21-22, 2007 (Open from 8 a.m. to 8:15 a.m. on June 21 and closed for remainder of the meeting).

Place: Agency for Healthcare Research and Quality (AHRQ), John Eisenberg Conference Center, 540 Gaither Road, Rockville, Maryland 20850.

4. *Name of Subcommittee:* Health Care Technology and Decision Sciences.

Date: June 28-29, 2007 (Open from 8 a.m. to 8:15 a.m. on June 28 and closed for remainder of the meeting).

Place: Agency for Healthcare Research and Quality (AHRQ), John Eisenberg Conference Center, 540 Gaither Road, Rockville, Maryland 20850.

Contact Person: Anyone wishing to obtain a roster of members, agenda or minutes of the nonconfidential portions of the meetings should contact Mrs. Bonnie Campbell, Committee Management Officer, Office of Extramural Research, Education and Priority Populations, AHRQ, 540 Gaither Road, Suite 2000, Rockville, Maryland 20850, Telephone (301) 427-1554.

Agenda items for these meetings are subject to change as priorities dictate.

Dated: April 23, 2007.

Carolyn M. Clancy,

Director.

[FR Doc. 07-2240 Filed 5-7-07; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Agency for Toxic Substances and Disease Registry

[ATSDR-230]

Public Health Assessments and Health Consultations Completed January 2007-March 2007

AGENCY: Agency for Toxic Substances and Disease Registry (ATSDR), Department of Health and Human Services (HHS).

ACTION: Notice.

SUMMARY: This notice announces those sites for which ATSDR has completed public health assessments and health consultations during the period from January 1, 2007 through March 31, 2007. This list includes sites that are on or proposed for inclusion on the National

Priorities List (NPL) and includes sites for which assessments or consultations were prepared in response to requests from the public.

FOR FURTHER INFORMATION CONTACT:

William Cibulas, Jr., PhD, Director, Division of Health Assessment and Consultation, Agency for Toxic Substances and Disease Registry, 1600 Clifton Road, NE., Mailstop E-32, Atlanta, Georgia 30333, telephone (404) 498-0007.

SUPPLEMENTARY INFORMATION: The most recent list of completed public health assessments and health consultations was published in the **Federal Register** on March 20, 2007 [72 FR 13115]. This announcement is the responsibility of ATSDR under the regulation "Public Health Assessments and Health Effects Studies of Hazardous Substances Releases and Facilities" [42 CFR Part 90]. This rule sets forth ATSDR's procedures for the conduct of public health assessments under section 104(i) of the Comprehensive Environmental Response, Compensation, and Liability Act (CERCLA), as amended by the Superfund Amendments and Reauthorization Act (SARA) [42 U.S.C. 9604(i)].

Availability

The completed public health assessments and health consultations are available for public inspection at the ATSDR Records Center, 1825 Century Boulevard, Atlanta, Georgia (not a mailing address), between 8 a.m. and 4:30 p.m., Monday through Friday except legal holidays. Public health assessments and health consultations are often available for public review at local repositories such as libraries in corresponding areas. Many public health assessments and health consultations are available through ATSDR's Web site at <http://www.atsdr.cdc.gov/HAC/PHA/>. In addition, the completed public health assessments are available by mail through the U.S. Department of Commerce, National Technical Information Service (NTIS), 5285 Port Royal Road, Springfield, Virginia 22161, or by telephone at (800) 553-6847. NTIS charges for copies of public health assessments. The NTIS order numbers are listed in parentheses following the site names.

Public Health Assessments Completed or Issued

Between January 1, 2007, and March 31, 2007, public health assessments were issued for the sites listed below:

NPL and Proposed NPL Sites

Georgia

Alternate Energy Resources, Inc.—(PB2007-106100); January 25, 2007.

Idaho

Bunker Hill Mining and Metallurgical Complex Operable Unit 3 (a/k/a Coeur D'Alene River Basin)—(PB2007-107841); March 26, 2007.

Illinois

Hegeler Zinc—(PB2007-106152); February 2, 2007.

New Hampshire

Former Chlor Alkali Facility Below Saw Mill Dam—(PB2007-106545); February 7, 2007.

Tennessee

Evaluation of Current (1990-2003) and Future Chemical Exposures in the Vicinity of the Oak Ridge Reservation, U.S. Department of Energy—(PB2007-106119); January 31, 2007.

Texas

Sandy Beach Road Groundwater Plume—(PB2007-104896); January 17, 2007.

West Virginia

Ravenswood PCE Ground Water Plume Site—(PB2007-107220); March 14, 2007.

Non-NPL Petitioned Sites

Florida

Coronet Industries, Incorporated (a/k/a Borden Feed Phosphate Complex)—(PB2007-104895); January 18, 2007.

Massachusetts

Evaluation of Cancer Incidence, 1982-2000, and Environmental Concerns Related to the Bird Landfill—(PB2007-106569); February 15, 2007.

Texas

East Kelly Air Force Base—(PB2007-106614); February 27, 2007.
Base de Fuerza Aerea, East Kelly (Spanish Version)—(PB2007-107191); February 27, 2007.

Health Consultations Completed or Issued

Between January 1, 2007, and March 31, 2007, health consultations were issued for the sites listed below:

Arizona

Evaluation of Primary Metals in Private Drinking Water Wells in the Walker Area; March 6, 2007.
Lone Butte Industrial Park—Perchlorate; March 8, 2007.

Lone Butte Industrial Park—TCE; January 10, 2007.

North Indian Bend Wash, Area 7 Groundwater Extraction and Treatment Facility; March 8, 2007.

California

Evaluation of Indoor Air Migration in Building On-Site and Adjacent to the Omega Chemical Site; March 20, 2007.

Klau/Buena Vista Mines—Evaluation of Fish Consumption from Lake Nacimiento—Exposure Investigation Report; February 6, 2007.

Colorado

A1 Stop Laundry and Dry Cleaners—Indoor Air Quality Assessment of a Residential Neighborhood Overlying a Tetrachloroethylene (PCE) Groundwater Plume; February 20, 2007.

Connecticut

Raymark Industries, Inc.—Evaluation of Soils at Selected School/Daycare Properties Located Near Raymark Industries Waste Disposal Areas; February 13, 2007.

Somers Plating, Inc.—Public Health Evaluation of Soil Sampling Data for Lagoon 3; February 13, 2007.

Florida

Former Royal Oaks Charcoal Facility—Air Testing; January 17, 2007.

Lincoln Park Complex, Durrs Neighborhood (Off-Site) Soil; March 27, 2007.

Georgia

Miller Bottom Road Municipal Solid Waste Landfill; March 28, 2007.

Idaho

Sunnyside Area Groundwater Contamination—Evaluation of Antibiotic, Steroid Hormone & Nitrate Compounds in Groundwater Near a Confined Animal Feeding Operation (CAFO); March 19, 2007.

Indiana

Rumpke Medora Landfill—Exposure Investigation Report; February 1, 2007.

Louisiana

Bayou Bonfouca—Post-Hurricane Evaluation; January 24, 2007.

Michigan

Kingsford Middle School; March 30, 2007.

Ontonagon High School Mercury Release; March 30, 2007.

Nebraska

Arsenic in Soil in East Omaha, Nebraska; March 20, 2007.

New Jersey

Celotex Corporation; January 10, 2007.

North Carolina

APAC Carolina Inc. and Associated Asphalt Inc., Jake Alexander Boulevard; February 14, 2007.
Weyerhaeuser Pulp and Paper Mill—Exposure Investigation Report; March 22, 2007.

Pennsylvania

Ivy Industrial Park Site—Public Health Evaluation of Residential Indoor Air and Well Water Sample Results; March 5, 2007.

Remacor Site; January 10, 2007.

Tennessee

Mr. Zip Convenience Store; March 14, 2007.

Texas

Former Delroc Oil Refinery/Woodwind Lakes Subdivision; February 23, 2007.

Utah

Vermiculite Intermountain and Intermountain Products, Inc.—Epidemiological Investigation of Human Exposure to a Contaminated Vermiculite Ore Processing Site in Utah; March 1, 2007.

Washington

Home Heating Oil Release, Technical Review of the Site Hazard Assessment; March 29, 2007.

Wisconsin

Amery-Dresser Trail; January 23, 2007.

Dated: May 2, 2007.

Kenneth Rose,

Acting Director, Office of Policy, Planning, and Evaluation, National Center for Environmental Health/Agency for Toxic Substances and Disease Registry.

[FR Doc. E7-8758 Filed 5-7-07; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES
Centers for Medicare & Medicaid Services
Privacy Act of 1974; Report of a Modified or Altered System of Records

AGENCY: Department of Health and Human Services (HHS), Centers for Medicare & Medicaid Services (CMS).

ACTION: Notice of a Modified or Altered System of Records (SOR).

SUMMARY: In accordance with the requirements of the Privacy Act of 1974, CMS is proposing to modify or alter existing system of records titled “Complaints Against Health Insurance Issuers and Health Plans (CAHII),” System No. 09-70-9005, established at 66 FR 9858, (February 12, 2001). We propose to assign a new CMS identification number to this system to simplify the obsolete and confusing numbering system originally designed to identify the Bureau, Office, or Center that maintained information in the Health Care Financing Administration systems of records. The new assigned identifying number for this system should read: System No. 09-70-0516.

We propose to modify existing routine use number 1 that permits disclosure to agency contractors and consultants to include disclosure to CMS grantees who perform a task for the agency. CMS grantees, charged with completing projects or activities that require CMS data to carry out that activity, are classified separate from CMS contractors and/or consultants. The modified routine use will remain as routine use number 1. We will delete routine use number 2 authorizing disclosure to support constituent requests made to a congressional representative. If an authorization for the disclosure has been obtained from the data subject, then no routine use is needed. The Privacy Act allows for disclosures with the “prior written consent” of the data subject.

We propose to add 2 new routine uses authorizing disclosure to support a CMS contractor, consultant, or a grantee of a CMS-administered grant program, when disclosure is deemed reasonably necessary by CMS to combat fraud, waste, and abuse in certain health care programs. The new routine use will be published as routine use number 6. We will add a second new routine use to support another Federal agency or to an instrumentality of any governmental jurisdiction within or under the control of the United States, when disclosure is deemed reasonably necessary by CMS to combat fraud, waste, and abuse in certain health care programs. This new routine use will be published as routine use number 7. We will broaden the scope of this system by including the section titled “Additional Circumstances Affecting Routine Use Disclosures,” that addresses “Protected Health Information (PHI)” and “small cell size.” The requirement for compliance with HHS regulation “Standards for Privacy of Individually Identifiable Health Information” apply when ever the system collects or maintain PHI. This system may contain

PHI. In addition, our policy to prohibit release if there is a possibility that an individual can be identified through “small cell size” will apply to the data disclosed from this system.

We are modifying the language in the remaining routine uses to provide a proper explanation as to the need for the routine use and to provide clarity to CMS’s intention to disclose individual-specific information contained in this system. The routine uses will then be prioritized and reordered according to their usage. We will also take the opportunity to update any sections of the system that were affected by the recent reorganization or because of the impact of the Medicare Prescription Drug, Improvement, and Modernization Act of 2003 (MMA) (Pub. L. 108-173) provisions and to update language in the administrative sections to correspond with language used in other CMS SORs.

The primary purpose of this system is to collect and maintain information initiated by consumers complaints/reports to CMS that their health insurance issuers and/or non-Federal governmental health plans are in violation of one or more of the following statutes: §§ 2722 and 2761 of the Public Health Service (PHS) Act; the Mental Health Parity Act of 1996 (MHPA); the Newborns’ and Mothers’ Health Protection Act of 1996 (NMHPA); and, the Women’s Health and Cancer Rights Act of 1998 (WHCRA). Information maintained in this system will also be disclosed to: (1) Support regulatory, reimbursement, and policy functions performed within the Agency or by a contractor, consultant or grantee; (2) assist another Federal or state agency, agency of a state government, an agency established by state law, or its fiscal agent; (3) assist third party contacts in situations where the party to be contacted has, or is expected to have information relating to the individual’s capacity to manage his or her affairs or to his or her eligibility for, or an entitlement to benefits under the Medicare program; (4) inform a health insurance issuer and/or health plan who has been named in a complaint/inquiry and is believed to be potentially in violation of relevant portions of the PHS; (5) support litigation involving the Agency; and (6) combat fraud, waste, and abuse in certain health benefits programs. We have provided background information about this new system in the **SUPPLEMENTARY INFORMATION** section below. Although the Privacy Act requires only that CMS provide an opportunity for interested persons to comment on the proposed routine uses, CMS invites comments on