EXHIBIT 1.—ESTIMATED ANNUALIZED BURDEN HOURS

Data collection	Number of respondents	Number of responses per respondent	Hours per response	Total burden hours
In-person interviews	41	1	1.5	61.5
Total	43	na	na	61.5

EXHIBIT 2.—ESTIMATED ANNUALIZED COST BURDEN

Data collection	Number of respondents	Total burden hours	Average hourly wage rate*	Total cost burden
In-person interviews	41	61.5	\$32.13	\$1,976
Total	41	na	na	1,976

^{*}Based upon the actual site personnel wages. Clinical personnel averages are weighted by the number of physicians, nurses and medical assistants in the sample. Administrative personnel averages are weighted by the number of administrators, lab, IT and other support personnel. Total average is weighted by relative number of administrative and clinical personnel being interviewed.

Estimated Annual Costs to the Federal Government

The total cost to the Federal Government for this project is \$393,457 over a two-year period. The average annual cost is \$196,728. the following is a breakdown of average annual costs:

Direct Costs:	
Personnel	\$108,320
Consultancies	24,400
Data support	5,000
Travel	2,575
Supplies	100
IRB review	125
Indirect Costs:	
Indirect costs 40%	56,208

Request for Comments

In accordance with the above-cited Paperwork Reduction Act legislation, comments on AHRQ's information collection are requested with regard to any of the following: (a) Whether the proposed collection of information necessary for the proper performance of AHRO's health care research and health care information dissemination functions, including whether the information 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 the 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 Agency's subsequent request for OMB approval of the proposed information collection. All comments will become a matter of public record.

Dated: February 6, 2008.

Carolyn M. Clancy,

Director.

[FR Doc. 08–660 Filed 2–14–08; 8:45 am]
BILLING CODE 4160–90–M

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day-08-0493]

Proposed Data Collections Submitted for Public Comment and Recommendations

In compliance with the requirement of Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995 for opportunity for public comment on proposed data collection projects, the Centers for Disease Control and Prevention (CDC) will publish periodic summaries of proposed projects. To request more information on the proposed projects or to obtain a copy of the data collection plans and instruments, call 404-639-5960, send comments to Maryam I. Daneshvar, CDC Acting Reports Clearance Officer, 1600 Clifton Road, MS-D74, Atlanta, GA 30333 or send an e-mail to omb@cdc.gov.

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. Written comments should be received within 60 days of this notice.

Proposed Project

2009 and 2011 National Youth Risk Behavior Surveys (YRBS) (OMB No. 0920–0493)—Reinstatement—National Center for Chronic Disease Prevention and Health Promotion (NCCDPHP), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

The purpose of this request is to obtain OMB approval to continue data collection for the National Youth Risk Behavior Survey (YRBS), a school-based survey that has been conducted biennially since 1991. OMB approval for the 2005 YRBS and 2007 YRBS expired November 30, 2007 (OMB No. 0920-0493). CDC seeks a three-year approval to conduct the YRBS in Spring 2009 and Spring 2011. Minor changes incorporated into this reinstatement request include: An updated title for the information collection, to accurately reflect the years in which the survey will be conducted; minor changes to the burden estimate; and minor changes to the data collection instrument.

The YRBS assesses priority health risk behaviors related to the major preventable causes of mortality, morbidity, and social problems among both youth and young adults in the United States. Data on health risk behaviors of adolescents are the focus of approximately 40 national health

objectives in Healthy People 2010, an initiative of the U.S. Department of Health and Human Services (HHS). The YRBS provides data to measure at least 10 of the health objectives and 3 of the 10 Leading Health Indicators established by Healthy People 2010. In addition, the YRBS can identify racial and ethnic disparities in health risk behaviors. No other national source of

data measures as many of the Healthy People 2010 objectives addressing adolescent behaviors as the YRBS. The data also will have significant implications for policy and program development for school health programs nationwide.

In Spring 2009 and Spring 2011, the YRBS will be conducted among nationally representative samples of

students attending public and private schools in grades 9–12. Information supporting the YRBS also will be collected from school administrators and teachers. The table below reports the number of respondents annualized over the 3-year project period.

There are no costs to respondents except their time.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondents	Form name	Number of respondents	Number of responses per respondent	Average burden per response (in hours)	Total burden (in hours)
Administrators	Recruitment Script for the Youth Risk Behavior Survey.	230	1	30/60	115
Teachers	Data Collection Checklist for the Youth Risk Behavior Survey.	400	1	15/60	100
Students	Youth Risk Behavior Survey	8,000	1	45/60	6,000
Total					6,215

Dated: February 7, 2008.

Maryam I. Daneshvar,

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

[FR Doc. E8–2832 Filed 2–14–08; 8:45 am] BILLING CODE 4163–18–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day-08-0337]

Proposed Data Collections Submitted for Public Comment and Recommendations

In compliance with the requirement of Section 3506(c)(2)(A) of the Paperwork Reduction Act of 1995 for opportunity for public comment on proposed data collection projects, the Centers for Disease Control and Prevention (CDC) will publish periodic summaries of proposed projects. To request more information on the proposed projects or to obtain a copy of the data collection plans and instruments, call 404-639-5960, send comments to Maryam I. Daneshvar, CDC Acting Reports Clearance Officer, 1600 Clifton Road, MS–D74, Atlanta, GA 30333 or send an e-mail to omb@cdc.gov.

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. Written comments should be received within 60 days of this notice.

Proposed Project

National Blood Lead Surveillance System (OMB No. 0920–0337)— Revision—National Center for Environmental Health (NCEH), Coordinating Center for Environmental Health and Injury Prevention (CCEHIP), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

The National Blood Lead Surveillance System (NBLSS) would like to continue its effort to collect information related to lead exposure among children less than six years old. The overarching goal of this system is to establish Childhood Lead Surveillance Systems at the state and national levels. This is a revision request in addition to a 3-year revision with an increase in the burden hours and inclusion of the adult blood lead surveillance system. As part of this effort we would like to revise this application to include 3 additional State and local Childhood Lead Poisoning Prevention Programs (CLPPP) who report to the NBLSS. These three programs were added to help provide a more comprehensive picture of

childhood lead poisoning in the United States.

The objectives for developing this system are three-fold. First, we would like to use surveillance data to estimate the extent of elevated blood-lead levels (BLLs) among children less than 6 years old. This is important because it will allow us to systematically track the management and follow-up of those children found to be poisoned with lead.

Our next objective for the development of this system is to examine potential sources of lead exposure. Although we've been successful in eliminating atmospheric lead with the use of unleaded gasoline and have continued to make strides in the elimination of household sources of lead commonly found in paint and dust, recent events have highlighted other potentially hidden sources of lead. This system will allow us to track the burden of such hidden sources and will help us eliminate such threats with the establishment of laws aimed at preventing the importation of such goods into our nation. The establishment of such laws will of course be a joint effort between several federal agencies; however, this surveillance system will help facilitate our efforts.

The final objective of this system is to facilitate the allocation of resources for lead poison prevention activities. The allocation of federal resources to State surveillance systems are based on reports of blood-lead tests from laboratories. Ideally, laboratories report results of all lead tests to the state health department. State health departments