A comment filed in paper form should include the reference "Food Industry Marketing to Children and Adolescents Study: Paperwork Comment; Project No. P094511" both in the text and on the envelope, and should be mailed or delivered to the following address: Federal Trade Commission, Office of the Secretary, Room H-135 (Annex J), 600 Pennsylvania Avenue, N.W., Washington, D.C. 20580. The FTC is requesting that any comment filed in paper form be sent by courier or overnight service, if possible, because U.S. postal mail in the Washington area and at the Commission is subject to delay due to heightened security precautions.

Comments on the proposed reporting requirements, which are subject to OMB review under the PRA, should additionally be submitted to: Office of Information and Regulatory Affairs, Office of Management and Budget, Attention: Desk Officer for Federal Trade Commission. Comments should be submitted via facsimile to (202) 395-5167 because U.S. postal mail at the OMB is subject to delays due to heightened security precautions.

The FTC Act and other laws the Commission administers permit the collection of public comments to consider and use in this proceeding as appropriate. The Commission will consider all timely and responsive public comments that it receives. whether filed in paper or electronic form. Comments received will be available to the public on the FTC Website, to the extent practicable, at (http://www.ftc.gov/os/ publiccomments.shtm). As a matter of discretion, the FTC makes every effort to remove home contact information for individuals from the public comments it receives before placing those comments on the FTC Website. More information, including routine uses permitted by the Privacy Act, may be found in the FTC's privacy policy, at (http://www.ftc.gov/ ftc/privacy.shtm).

David C. Shonka,

Acting General Counsel.
[FR Doc. 2010–12511 Filed 5–24–10; 8:45 am]
BILLING CODE 6750–01–S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60 Day-10-10DT]

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 and send comments to Maryam Daneshvar, CDC 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

Monitoring and Reporting System for Chronic Disease Prevention and Control Programs—New—National Center for Chronic Disease Prevention and Health Promotion (NCCDPHP), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

Chronic diseases are the leading causes of death and disability in the United States, accounting for seven of every ten deaths and affecting the quality of life for 90 million Americans. Chronic diseases represent 83% of all U.S. health care spending.

The National Center for Chronic Disease Prevention and Health Promotion (NCCDPHP) provides funding to health departments in States, territories, and the District of Columbia to implement and evaluate chronic

disease prevention and control programs. Traditionally, support has been provided through cooperative agreements that are specific to a chronic disease or condition. In 2009, CDC announced a new cooperative agreement program for collaborative chronic disease prevention and health promotion programs (RFA DP09-901; authorized under sections 301, 307, 310, and 311 of the Public Health Service Act [42 U.S.C. 241 and 247(b)(k)). The new program streamlines funding, communication and collaboration in four areas that have previously been funded and evaluated independently: Tobacco control, diabetes prevention and control, State-based surveillance through the Behavioral Risk Factor Surveillance System (BRFSS), and the Healthy Communities initiative. Awardees are the 50 States, the District of Columbia, Puerto Rico, and the Virgin Islands. The new cooperative agreement encourages each awardee to collaborate with partners (both internal and external to the State health department) to develop and implement a multi-year, statewide strategic plan.

CDC requests OMB approval to collect information from the four specified State programs through a new, electronic Management Information System (MIS). Upon approval, the new information collection system will replace two previously approved information collections for tobacco control programs (OMB No. 0920-0601, exp. 5/31/2010), and diabetes prevention and control programs (OMB No. 0920-0479, exp. 4/30/2013), and harmonize their content. In addition, the new MIS will provide a common progress reporting framework for Statebased BRFSS programs and Healthy Community programs, which have previously reported progress information to CDC using standard progress reporting forms for cooperative agreement awardees (OMB No. 4040-0004, exp. 3/31/2012).

Information will be collected on each program's objectives, planning activities, resources, partnerships, policy and environmental strategies for preventing or controlling chronic diseases, and progress toward meeting goals. The increased emphasis on partnership and collaboration is intended to identify priorities, gaps in chronic disease prevention and health promotion activities, and opportunities to leverage CDC and State (Federal and non-Federal) resources. Information will be collected electronically through a new Management Information System (MIS). The collection of information, in a uniform and efficient manner, will reduce duplicative reporting

requirements for awardees, minimize respondent burden, facilitate collaborative efforts and provide common performance metrics across program areas. The collaborative cooperative agreement is part of an initiative within NCCDPHP to standardize and streamline the funding and performance monitoring processes for programs funded through the Center; to promote more efficient ways to use resources; and to achieve greater health impact. The objectives for awardees, and the performance indicators defined for them, reflect CDC's support for more integrated approaches to the prevention and control of chronic conditions.

Awardees will use the information collection to manage and coordinate

their activities and to improve their efforts to prevent and control chronic diseases. The MIS will allow awardees to fulfill their reporting obligations under the cooperative agreements in an efficient manner by employing a single instrument to collect necessary information for both progress reports and continuation applications including work plans.

CDC will use the information collected in the MIS to monitor each awardee's progress and to make adjustments, as needed, in the type and level of technical assistance provided to them. The information collection will allow CDC to oversee the use of Federal funds, and identify and disseminate information about successful prevention

and control strategies implemented by awardees. CDC will also use the information collection to respond to Congressional and stakeholder inquiries about chronic disease control activities, program implementation, and program impact. Finally, the information collection will allow CDC to evaluate the success of the collaborative funding model which places increased emphasis on partnerships as well as policy and environmental strategies for preventing and controlling chronic diseases.

Information will be collected from each State-based program twice per year. There are no costs to respondents other than their time.

ESTIMATED ANNUALIZED BURDEN TABLE

Type of respondents	Number of respondents	Number of responses per respondent	Average burden per response (in hours)	Total burden
State Diabetes Program State Tobacco Program State BRFSS Program State Healthy Communities Program	53 53 53 53	2 2 2 2	6 6 6 6	636 636 636 636
Total				2,544

Carol Walker,

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

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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, Pub. L. 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, e-mail

paperwork@hrsa.gov or call the HRSA Reports Clearance Officer at (301) 443–

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: Healthcare Integrity and Protection Data Bank for Final Adverse Information on Health Care Providers, Suppliers and Practitioners (45 CFR 61) (OMB No. 0915–0239)— [Extension]

This is a request for extension of OMB approval of the information collections contained in regulations found in 45 CFR part 61 governing the Healthcare Integrity and Protection Data Bank (HIPDB) and the forms to be used in reporting information to and requesting information from the HIPDB cleared under OMB No. 0915–0239. The HIPDB

is authorized by section 1128E of the Social Security Act (hereinafter referred to as section 1128E), as added by section 221(a) of the Health Insurance Portability and Accountability Act of 1996. Section 1128E directs the Secretary of Health and Human Services (the Secretary) to establish a national health care fraud and abuse data collection program for the reporting and disclosing of certain final adverse actions (excluding settlements in which no findings of liability have been made) taken against health care providers, suppliers, or practitioners. It also directs the Secretary to maintain a database of final adverse actions taken against health care providers, suppliers, or practitioners. The regulations implementing section 1128E governing the operation of the HIPDB are codified at 45 CFR part 61. The HIPDB became operational November 22, 1999.

Approval is requested to continue the following reporting data collection and disclosure requirements and the ensuing HIPDB forms along with the instructions. The recordkeeping, reporting, and disclosure requirements are specified in the regulations to implement the HIPDB. The annual estimate of burden is as follows: