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 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.

Sharon Arnold,

Deputy Director.

[FR Doc. 2015–21719 Filed 9–1–15; 8:45 am]

BILLING CODE 4160-90-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Agency for Toxic Substances and Disease Registry

[Docket No. ATSDR-2015-0004]

Availability of Draft Toxicological Profile; Perfluoroalkyls

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

ACTION: Notice of availability, and request for comment.

SUMMARY: The Agency for Toxic Substances and Disease Registry (ATSDR) located in the Department of Health and Human Services (HHS) announces the availability of the Toxicological Profile for Perfluoroalkyls for review and comment. Comments can include additional information or reports on studies about the health effects of perfluoroalkyls. Although ATSDR considered key studies for this substance during the profile development process, this Federal **Register** notice solicits any relevant, additional studies, particularly unpublished data. ĀTSDR will evaluate the quality and relevance of such data or studies for possible inclusion into the profile. ATSDR remains committed to providing a public comment period for this document as a means to best serve public health and our clients.

DATES: To be considered, comments on the draft Toxicological Profile for Perfluoroalkyls must be received not later than December 1, 2015. Comments received after close of the public comment period will be considered solely at the discretion of ATSDR, based upon what is deemed to be in the best interest of the general public.

ADDRESSES: You may submit comments, identified by the docket number ATSDR-2015-0004, by any of the following methods:

- Federal eRulemaking Portal: http://www.regulations.gov/#!home. Follow the instructions for submitting comments.
- Mail: Division of Toxicology and Human Health Sciences, 1600 Clifton Rd. NE., F57, Atlanta, GA 30329–4027.

FOR FURTHER INFORMATION CONTACT: Ms. Delores Grant, Division of Toxicology and Human Health Sciences, Agency for Toxic Substances and Disease Registry, 1600 Clifton Road NE., MS F-57, Atlanta, GA 30329; telephone number (800) 232-4636 or (770) 488-3351.

SUPPLEMENTARY INFORMATION: The Superfund Amendments and Reauthorization Act of 1986 (SARA) (42 U.S.C. 9601 et seq.) amended the Comprehensive Environmental Response, Compensation, and Liability Act of 1980 (CERCLA or Superfund) (42 U.S.C. 9601 *et seq.*) by establishing certain requirements for ATSDR and the U.S. Environmental Protection Agency (U.S. EPA) regarding hazardous substances that are most commonly found at facilities on the CERCLA National Priorities List (NPL). Among these statutory requirements is a mandate for the Administrator of ATSDR to prepare toxicological profiles for each substance included on the priority list of hazardous substances (also called the Substance Priority List). This list identifies 275 hazardous substances that ATSDR (in cooperation with EPA) has determined pose the most significant potential threat to human health. The availability of the revised list of the 275 priority substances was announced in the Federal Register on May 28, 2014 (79 FR 30613) and is available at www.atsdr.cdc.gov/spl.

In addition, ATSDR has the authority to prepare toxicological profiles for substances not found at sites on the National Priorities List, in an effort to "establish and maintain inventory of literature, research, and studies on the health effects of toxic substances" under CERCLA Section 104(i)(1)(B), to respond to requests for consultation under section 104(i)(4), and as otherwise necessary to support the site-specific response actions conducted by ATSDR.

On November 6, 2008, ATSDR announced the availability of a draft toxicological profile for Set 22 Toxicological Profiles for public comment (73 FR 66047). The Set 22 Toxicological Profiles included Perfluoroalkyls and ATSDR announced that the Perfluoroalkyls profile was on

a modified schedule pending additional review.

On July 23, 2009 ATSDR published a second notice of the availability of the toxicological profile for Perfluoroalkyls in draft form for public review and comment (74 FR 36492). The 90-day comment period ended October 30, 2009. Following the close of the comment period, chemical-specific comments were addressed, and, where appropriate, changes were incorporated into the profile. Given the plethora of new data that have been published since 2009, and the resulting extensive revision to the profile, the agency has determined that it would be in the best interest of public health to release the perfluoroalkyls profile for another public comment period. The public comments and other data submitted in response to the Federal Register notices are available for inspection from Monday through Friday, except for legal holidays, from 9 a.m. until 5 p.m., Eastern Time, at 4770 Buford Hwy NE., Atlanta, Georgia 30341. Please call ahead to 1-800-232-4636 and ask for a representative in the Division of Toxicology and Human Health Sciences to schedule your visit.

Availability

The Toxicological Profile for Perfluoroalkyls prepared by ATSDR will be made available to the public on or about August 31, 2015 at the ATSDR Web site: www.atsdr.cdc.gov/ toxprofiles/index.asp and at the Federal eRulemaking Portal: http:// www.regulations.gov/#!home.

Sascha Chaney,

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

[FR Doc. 2015–21544 Filed 9–1–15; 8:45 am]

BILLING CODE 4163-70-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day-15-0214; Docket No. CDC-2015-

Proposed Data Collection Submitted for Public Comment and Recommendations

AGENCY: Centers for Disease Control and Prevention (CDC), Department of Health and Human Services (HHS).

ACTION: Notice with comment period.

SUMMARY: The Centers for Disease Control and Prevention (CDC), as part of

its continuing efforts to reduce public burden and maximize the utility of government information, invites the general public and other Federal agencies to take this opportunity to comment on proposed and/or continuing information collections, as required by the Paperwork Reduction Act of 1995. This notice invites comment on the proposed revision of the National Health Interview Survey (NHIS). The annual National Health Interview Survey is a major source of general statistics on the health of the U.S. population.

DATES: Written comments must be received on or before November 2, 2015.

ADDRESSES: You may submit comments, identified by Docket No. CDC-2015-0076 by any of the following methods:

- Federal eRulemaking Portal: Regulation.gov. Follow the instructions for submitting comments.
- Mail: Leroy A. Richardson, Information Collection Review Office, Centers for Disease Control and Prevention, 1600 Clifton Road NE., MS— D74, Atlanta, Georgia 30329.

Instructions: All submissions received must include the agency name and Docket Number. All relevant comments received will be posted without change to Regulations.gov, including any personal information provided. For access to the docket to read background documents or comments received, go to Regulations.gov.

FOR FURTHER INFORMATION CONTACT: To request more information on the proposed project or to obtain a copy of the information collection plan and instruments, contact the Information Collection Review Office, Centers for Disease Control and Prevention, 1600 Clifton Road NE., MS–D74, Atlanta, Georgia 30329; phone: 404–639–7570; Email: omb@cdc.gov.

SUPPLEMENTARY INFORMATION: Under the Paperwork Reduction Act of 1995 (PRA) (44 U.S.C. 3501–3520), Federal agencies must obtain approval from the Office of Management and Budget (OMB) for each collection of information they conduct or sponsor. In addition, the PRA also requires Federal agencies to provide a 60-day notice in the Federal Register concerning each proposed collection of information, including each new proposed collection, each proposed extension of existing collection of information, and each reinstatement of previously approved information collection before submitting the collection to OMB for approval. To comply with this requirement, we are publishing this notice of a proposed data collection as described below.

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; (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; and (e) estimates of capital or start-up costs and costs of operation, maintenance, and purchase of services to provide information. Burden means the total time, effort, or financial resources expended by persons to generate, maintain, retain, disclose or provide information to or for a Federal agency. This includes the time needed to review instructions; to develop, acquire, install and utilize technology and systems for the purpose of collecting, validating and verifying information, processing and maintaining information, and disclosing and providing information; to train personnel and to be able to respond to a collection of information, to search data sources, to complete and review the collection of information; and to transmit or otherwise disclose the information.

Proposed Project

National Health Interview Survey (NHIS), (OMB No. 0920–0214, expires 12/31/2017)—Revision—National Center for Health Statistics (NCHS), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

Section 306 of the Public Health Service (PHS) Act (42 U.S.C. 242k), as amended, authorizes that the Secretary of Health and Human Services (DHHS), acting through NCHS, shall collect data on the extent and nature of illness and disability of the population of the United States. The annual National Health Interview Survey is a major source of general statistics on the health of the U.S. population and has been in the field continuously since 1957. Clearance is sought for three years, to collect data from 2016 to 2018. This voluntary and confidential householdbased survey collects demographic and health-related information from a nationally representative sample of noninstitutionalized, civilian persons and households throughout the country. Personal identification information is requested from survey respondents to

facilitate linkage of survey data with health-related administrative and other records. In 2016 the NHIS will collect information from approximately 45,000 households, which contain about 112,000 individuals.

Information is collected using computer assisted personal interviews (CAPI). A core set of data is collected each year that remains largely unchanged, whereas sponsored supplements vary from year to year. The core set includes socio-demographic characteristics, health status, health care services, and health behaviors. For 2016, supplemental questions will be cycled in pertaining to balance, blood donation, chronic pain, diabetes, and vision. Supplemental topics that continue or are enhanced from 2015 pertain to family food security, heart disease and stroke, inflammatory bowel disease, hepatitis B and C screening, children's mental health, disability and functioning, smokeless tobacco and ecigarettes, and immunizations. Questions from 2015 on cancer control, epilepsy, and occupational health have been removed. In addition to these core and supplemental modules, a followback survey will be conducted on previous NHIS respondents to collect additional health related information using alternative question wording and data collection modes as a testbed for the intended 2018 redesign of the NHIS questionnaire. In addition, a subsample of NHIS respondents may be identified to participate in a pilot test to assess the feasibility of integrating wearable devices into the NHIS data collection process. The aim is to directly track health measurements, to compare those measurements to the self-reported health information provided by respondents, and to assess the role of devices in reducing respondent burden.

A new sampling strategy is being implemented in 2016 and for the foreseeable future. This new sampling design is necessitated by the prior 2006-2015 sample being exhausted, and will take into account demographic shifts in the U.S. civilian noninstitutionalized population. It will also be more flexible allowing for additions and contractions to reflect funding availability and to meet estimation goals. As in previous years, the base sample will remain at approximately 35,000 completed household interviews annually. To balance the precision of national and state-based estimates, most of the sample (approximately 25,000 completed interviews) will be allocated proportionally to the state population to maximize the precision of national-level estimates. A smaller portion of the sample (approximately 10,000

completed interviews) will be shifted to increase sample in the 10 least populous states, enabling state-level estimates of key variables to be produced for all 50 states and DC by pooling 3 years of data. This flexibility embedded in the new sampling plan reflects. Additional funding to improve state-level estimates will increase the sample by almost 10,000 completed interviews in midsize states bringing the total expected sample size in 2016 to 45,000 households.

Whereas the sampling frame for the NHIS has traditionally used field listing by the Census Bureau, in order to contain costs, the new frame will use a commercially available address list that covers residential addresses within all 50 states and the District of Columbia. Some field listing will be undertaken to improve coverage in rural areas, in high density areas, and of university housing units. This represents a substantial reduction in the number of listings performed annually.

It is anticipated that this new sampling plan will not affect estimates generated using NHIS data. To monitor the new design's performance, NHIS analysts will perform monthly checks in line with the ones currently performed as part of routine data review. NCHS receives raw data files monthly from the Census Bureau for processing and quality review. Each year, results from the January sample are compared to the previous year to determine whether the results consistent. In addition to comparing the unweighted and weighted frequencies, the input and output specifications are reviewed, and the flowcharts are compared to the skip instructions and universes for each question. If a difference is found, steps are taken to determine whether the change is legitimate or whether there is a factor other than the programming of the questionnaire such as the location or context of the question in the questionnaire. If a difference persists, the paradata are reviewed to determine whether there are changes in the mean or median time spent on that question, whether interviewers had a high rate of backing up to return to that question, and whether other questions in that battery were similarly affected. Persistent differences will be examined to determine whether there is any other interviewer effect such as results comparing newly hired and experienced interviewers and newly added primary sampling units compared to continuing primary sampling units. In addition, national estimates on the key set of indicators that are released in a quarterly report as part of the Early Release program will be monitored by NHIS analysts.

In accordance with the 1995 initiative to increase the integration of surveys within the DHHS, respondents to the NHIS serve as the sampling frame for the Medical Expenditure Panel Survey conducted by the Agency for Healthcare Research and Quality. The NHIS has long been used by government, academic, and private researchers to evaluate both general health and specific issues, such as smoking, diabetes, health care coverage, and access to health care. It is a leading source of data for the Congressionallymandated "Health US" and related publications, as well as the single most important source of statistics to track progress toward the National Health Promotion and Disease Prevention Objectives, "Healthy People 2020."

There is no cost to the respondents other than their time.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondent	Form name	Number of respondents	Number of responses per respondent	Average burden per response (in hours)	Total burden (in hours)
Adult Family Member	Family CoreAdult Core	10,000 45,000 36,000	1 1 1	5/60 23/60 15/60 10/60	833 17,250 9,000 2,333
Adult Family MemberAdult Family Member	Supplements	14,000 45,000 15,000 5,000	1 1	20/60 20/60 20/60 5/60	2,333 15,000 5,000 417
Total	Treinterview Survey				49,833

Leroy A. Richardson,

Chief, Information Collection Review Office, Office of Scientific Integrity, Office of the Associate Director for Science, Office of the Director, Centers for Disease Control and Prevention.

[FR Doc. 2015–21708 Filed 9–1–15; 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: National Youth in Transition Database and Youth Outcome Survey. OMB No.: 0970–0340.

Description: The Foster Care Independence Act of 1999 (42 U.S.C. 1305 et seq.) as amended by Public Law 106–169 requires State child welfare agencies to collect and report to the Administration on Children and Families (ACF) data on the characteristics of youth receiving

independent living services and information regarding their outcomes. The regulation implementing the National Youth in Transition Database, listed in 45 CFR 1356.80, contains standard data collection and reporting requirements for States to meet the law's requirements. ACF will use the information collected under the regulation to track independent living services, assess the collective outcomes of youth, and potentially to evaluate State performance with regard to those outcomes consistent with the law's mandate.

Respondents: State agencies that administer the John H. Chafee Foster Care Independence Program.