

work, the focus of their work and the population groups and geographic areas served.

The evaluation instruments are used to assess training and capacity-building outcomes (knowledge, confidence, intention to use information, actual changes made as a result of training) immediately after and again 90 days

after training events. The evaluation instruments vary based on the type of training offered and take between approximately 16 minutes to complete (for intensive multi-day trainings) to 2 minutes to complete (for short didactic or webinar sessions).

The CDC's Funding Opportunity Announcement PS 14-1407, NNPTC,

requires the collection of national demographic information on grantees' trainees and national evaluation outcomes.

There are no costs to respondents other than their time. The estimated annualized burden hours for this data collection are 502 hours.

ESTIMATES OF ANNUALIZED BURDEN

Type of respondent	Form name	Number of respondents	Number responses per respondent	Average burden per response (in hours)	Total burden hours
Healthcare Professionals	NNPTC Abbreviated Health Professional Application for Training (HPAT).	4,500	1	3/60	225
Healthcare Professionals	Intensive Complete Post-Course Evaluation.	116	1	16/60	31
	Intensive Complete Long-Term Evaluation.	36	1	10/60	6
Healthcare Professionals	Intensive-Didactic Post-Course Evaluation.	166	1	10/60	28
	Intensive-Didactic Long-Term Evaluation.	58	1	7/60	7
Healthcare Professionals	Practicum Post-Course Evaluation ..	70	1	4/60	5
	Practicum Long-Term Evaluation	20	1	3/60	1
Healthcare Professionals	Wet Mount Post-Course Evaluation	40	1	3/60	2
	Wet Mount Long-Term Evaluation ...	15	1	2/60	1
Healthcare Professionals	STD Tx Guidelines Complete Post-Course Evaluation.	548	1	6/60	55
	STD Tx Guidelines Complete Long-Term Evaluation.	180	1	5/60	15
Healthcare Professionals	Short Guidelines Post-Course Evaluation.	500	1	3/60	25
	Short Guidelines Long-Term Evaluation.	160	1	3/60	8
Healthcare Professionals	Basic Post-Course Evaluation	150	1	2/60	5
	Basic Long-Term Evaluation	50	1	2/60	2
Healthcare Professionals	Immediate Post-Course email invitation.	4,500	1	1/60	75
Healthcare Professionals	3 Month Long-Term email invitation	660	1	1/60	11
Total					502

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. 2016-17603 Filed 7-25-16; 8:45 am]

BILLING CODE 4163-18-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[60Day-16-0612; Docket No. CDC-2016-0070]

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.

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 Well-Integrated Screening and Evaluation for Women Across the Nation (WISEWOMAN) Reporting System. The WISEWOMAN program aims to reduce cardiovascular disease in women ages 40-64 by providing screening services, referrals to medical care, and lifestyle intervention programs.

DATES: Written comments must be received on or before September 26, 2016.

ADDRESSES: You may submit comments, identified by Docket No. CDC-2016-0070 by any of the following methods:

• *Federal eRulemaking Portal: Regulations.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*. *Please note: All public comment should be submitted through the Federal eRulemaking portal (Regulations.gov) or by U.S. mail to the address listed above.*

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

Well-Integrated Screening and Evaluation for Women across the Nation (WISEWOMAN) Reporting System (OMB No. 0920-0612, exp. 12/31/2016)—Extension—National Center for Chronic Disease Prevention and Health Promotion (NCCDPHP), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

The WISEWOMAN program (Well-Integrated Screening and Evaluation for Women Across the Nation), sponsored by the CDC, was established to examine ways to improve the delivery of services for women who have limited access to health care and elevated risk factors for cardiovascular disease (CVD). The program focuses on reducing CVD risk factors and provides screening services for selected risk factors such as elevated blood cholesterol, hypertension, and abnormal blood glucose levels. The program also provides women with referrals to lifestyle programs and medical care. The WISEWOMAN program provides services to women who are jointly enrolled in the National Breast and Cervical Cancer Early Detection Program (NBCCEDP), also administered by CDC.

The WISEWOMAN program is administered by state health departments and tribal programs. In

2013, new cooperative agreements were awarded under Funding Opportunity Announcement DP13-1302. These awards are currently in the final year of funding, but may be extended by CDC for one additional year, subject to the availability of funds.

CDC collects two types of information from WISEWOMAN awardees. The hardcopy Annual Progress Report provides a narrative summary of each awardee's objectives and the activities undertaken to meet program goals. The estimated burden per response is 16 hours.

In addition, each WISEWOMAN awardee submits an electronic data file to CDC twice per year. The Minimum Data Elements (MDE) file contains de-identified, client-level information about the cardiovascular disease risk factors of women served by the program, and the number and type of lifestyle program sessions they attend. The estimated burden per response for the MDE file is 24 hours.

CDC seeks a one-year extension to enable reporting for the final year of activities funded under the current cooperative agreement and the option year, subject to the availability of funds. There are no changes to the information collected, the burden per response, reporting frequency, the number of awardees, or the total annualized burden hours.

CDC will continue to use the information collected from WISEWOMAN awardees to support program monitoring and improvement activities, evaluation, and assessment of program outcomes. The overall program evaluation is designed to demonstrate how WISEWOMAN can obtain more complete health data on vulnerable populations, promote public education about disease incidence, cardiovascular disease risk-factors, health promotion, improve the availability of screening and diagnostic services for underserved women, ensure the quality of services provided to underserved women, and develop strategies for improved interventions.

OMB approval is requested for one year. Participation in this information collection is required as a condition of cooperative agreement funding. There are no costs to respondents other than their time. The total annualized burden hours are 1,344.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondents	Form name	Number of respondents	Number of responses per respondent	Average burden per response (in hrs.)	Total burden (in hrs.)
WISEWOMAN Awardees	Screening and Assessment and Lifestyle Program MDEs.	21	2	24	1,008
	Annual Progress Report	21	1	16	336
Total					1,344

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. 2016-17602 Filed 7-25-16; 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: Phase II Evaluation Activities
 for Implementing a Next Generation

Evaluation Agenda for the Chafee Foster
 Care Independence Program.

OMB No.: New Collection.

Description: The Administration for
 Children and Families (ACF), Office of
 Planning Research and Evaluation
 (OPRE) is proposing an information
 collection activity as part of the Phase
 II Evaluation Activities for
 Implementing a Next Generation
 Evaluation Agenda for the Chafee Foster
 Care Independence Program. The
 proposed information collection
 consists of site visits by staff from the
 Urban Institute and Chapin Hall at the
 University of Chicago to conduct
 formative evaluations of programs
 serving transition-age foster youth. The
 evaluations will include preliminary

visits to discuss the evaluation process
 with program administrators. Then, the
 research team will conduct site visits to
 each program to speak with program
 leaders, partners and key stakeholders,
 front-line staff, and participants. These
 formative evaluations will determine
 programs' readiness for more rigorous
 evaluation in the future. The activities
 and products from this project will help
 ACF to fulfill their ongoing legislative
 mandate for program evaluation
 specified in the Foster Care
 Independence Act of 1999.

Respondents: Program leaders,
 partners and stakeholders, and frontline
 staff as well as young adults being
 served by the programs.

ANNUAL BURDEN ESTIMATES

Instrument	Total number of respondents	Annual number of respondents	Number of responses per respondent	Average burden hours per response	Annual burden hours
Outreach email for discussion with program admin and staff	16	8	1	1	8
Outreach email for Focus Group Recruiters	16	8	1	8	64
Informed Consent and Discussion Guide for program leaders	48	24	4	1	96
Informed Consent and Discussion Guide for program partners and stakeholders	80	40	2	1	80
Informed Consent and Discussion Guide for program front-line staff	128	64	1	1	64
Informed Consent and Focus Group Guide for program participants	200	100	1	2	200
Compilation and Submission of Administrative Data	24	12	2	12	288

*Estimated Total Annual Burden
 Hours:* 800.

Additional Information: Copies of the
 proposed collection may be obtained by
 writing to the Administration for
 Children and Families, Office of
 Planning, Research and Evaluation, 330
 C Street SW., Washington, DC 20201,
 Attn: OPRE Reports Clearance Officer.
 All requests should be identified by the
 title of the information collection. Email
 address: [OPREinfocollection@](mailto:OPREinfocollection@acf.hhs.gov)
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, Email: [OIRA_](mailto:OIRA_SUBMISSION@OMB.EOP.GOV)
SUBMISSION@OMB.EOP.GOV, Attn:

Desk Officer for the Administration for
 Children and Families.

Robert Sargis,
ACF Certifying Officer.

[FR Doc. 2016-17618 Filed 7-25-16; 8:45 am]

BILLING CODE 4184-01-P