Table 2 of this unit includes the names and addresses of record for all the registrants of the products in Table 1 of this unit, in sequence by EPA company number. This number corresponds to the first part of the EPA registration numbers of the products listed in Table 1 of this unit.

III. Summary of Public Comments Received and Agency Response to Comments

During the public comment period provided, EPA received no comments in response to the March 13, 2013 Federal Register (78 FR 15949) notice announcing the Agency’s receipt of the request for voluntary cancellation of products listed in Table 1 of Unit II.

IV. Cancellation Order

Pursuant to FIFRA section 6(f) (7 U.S.C. 136d(f)), EPA hereby approves the requested cancellation of the registrations identified in Table 1 of Unit II. Accordingly, the Agency hereby orders that the product registrations identified in Table 1 of Unit II are canceled. The effective date of the cancellations are the subject of this notice is August 19, 2015. Any distribution, sale, or use of existing stocks of the products identified in Table 1 of Unit II in a manner inconsistent with any of the provisions for disposition of existing stocks set forth in Unit VI will be a violation of FIFRA.

V. What is the Agency’s authority for taking this action?

Section 6(f)(1) of FIFRA (7 U.S.C. 136d(f)(1)) provides that a registrant of a pesticide product may at any time request that any of its pesticide registrations be canceled or amended to terminate one or more uses. FIFRA further provides that, before acting on the request, EPA must publish a notice of receipt of any such request in the Federal Register. Thereafter, following the public comment period, the EPA Administrator may approve such a request. The notice of receipt for this action was published for comment in the Federal Register on March 13, 2013 (78 FR 15949). The comment period closed on April 12, 2013.

VI. Provisions for Disposition of Existing Stocks

Existing stocks are those stocks of canceled pesticide products that are in the United States and that were appropriately packaged, labeled, and released for shipment prior to the effective date of cancellation of the underlying registration. The existing stocks provisions for the products subject to this order are as follows.

The registrant is prohibited from selling or distributing existing stocks above as of August 19, 2015, except for export in accordance with FIFRA section 17 (7 U.S.C. 136o), or proper disposal. Persons other than the registrant may sell, distribute, or use existing stocks of products listed above until existing stocks are exhausted, provided that such sale, distribution, or use is consistent with the terms of the previously approved labeling on, or that accompanied, the canceled products.

Authority: 7 U.S.C. 136 et seq.


Richard P. Keigwin, Jr.,
Director, Pesticide Re-Evaluation Division,
Office of Pesticide Programs.

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

An amendment changes...
which is designed to assess insurance coverage, employment status and out-of-pocket health care expenses among young women diagnosed with breast cancer and to look at the relationship between these variables and treatment decisions.

DATES: Written comments must be received on or before October 19, 2015.

ADDRESSES: You may submit comments, identified by Docket No. CDC–2015–0070 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 Regulation.gov, including any personal information provided. For access to the docket to read background documents or comments received, go to Regulation.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 existing data sources, to complete and review the collection of information; and to transmit or otherwise disclose the information.

Proposed Project

Insurance Coverage, Employment Status, and Copayments/Deductibles Faced by Young Women Diagnosed with Breast Cancer—New—National Center for Chronic Disease Prevention and Health Promotion (NCCDPHP), Centers for Disease Control and Prevention (CDC).

Background and Brief Description

The Education and Awareness Requires Learning Young (EARLY) Act of 2009, which is outlined in section 10413 of the Patient Protection and Affordable Care Act, authorizes the CDC to fund research and initiatives that increase knowledge of breast health and breast cancer among women, particularly among those under the age of 40. The EARLY Act along with section 301 of the Public Health Service Act authorizes the CDC to conduct research that will inform the prevention of physical and mental diseases such as breast cancer, and serves as the main basis for this data collection activity.

Research indicates that young women diagnosed with breast cancer face many barriers accessing high-quality breast cancer care and treatment. These barriers are compounded by the multiple roles that these young women serve in society including parenting young children, developing a career, and completing their education. Treatment decisions can be complicated for young women with breast cancer. Some research indicates that employment status, financial stability, and insurance coverage are variables that affect treatment compliance, access to quality care, and ultimately quality of life for young women with breast cancer. However, to date, no comprehensive assessment has been conducted to examine breast cancer care and treatment for young women.

CDC propose to address this gap by answering the following two research questions: (1) What are young, female breast cancer survivors experiencing after their diagnosis in terms of (a) continuation of insurance coverage, access to care, and quality of care; (b) changes in employment status after breast cancer diagnosis; and (c) out-of-pocket medical costs? (2) What factors affect young breast cancer survivors’ access to comprehensive, high quality care?

To answer these research questions, CDC is sponsoring a study to collect information from two groups of breast cancer survivors: One randomly drawn from state-based cancer registries (Sample 1), the other a self-selected convenience sample drawn from two advocacy organizations (Sample 2).

Sample 1 will include up to 1,750 young (diagnosed between the ages of 18 and 39), female breast cancer survivors diagnosed for the first time with breast cancer 12 months before the survey is fielded. Respondents will be recruited through approximately four state-based central cancer registries. These respondents will be asked to complete a mail-in or web-based questionnaire. Self-reported survey data from Sample 1 will be supplemented by data maintained by their state’s cancer registry, including information about tumor characteristics, date of diagnosis, and stage. The linked survey and cancer registry data will be used to answer research question #2 (What factors affect young breast cancer survivors’ access to comprehensive, high quality care?).

Sample 2 will include a nation-wide convenience sample of 2,000 female breast cancer survivors diagnosed between the ages of 18 and 49 who are associated with one of two breast cancer advocacy groups (Living Beyond Breast Cancer and Young Survival Coalition). This cohort will exclude individuals from Sample 1 and will not be linked to any other data source.

Comparing results between Sample 1 and Sample 2 will help us address these additional research questions: (1) How generalizable are the results from the convenience Sample 2? (2) Are there...
differences between young breast cancer survivors based on the length of time that has elapsed from cancer diagnosis? (3) Do the experiences and barriers faced by women diagnosed between 18 and 39 years of age (Samples 1 and 2) differ from those of women diagnosed between 40 and 44 years of age and 45 and 49 years of age (Sample 2)? This comparison will also help CDC explore whether drawing a convenience sample from survivorship groups will be a methodologically legitimate, less expensive method to recruit respondents for future breast cancer survivor surveys.

The target number of responses for the overall study will result in up to 3,750 completed surveys. Respondents will be asked to complete a questionnaire, which is estimated to take about 22 minutes. Sample 1 respondents will have the option of completing a hardcopy questionnaire or an online questionnaire. Sample 2 respondents will complete the questionnaire online. Demographic information will be collected from all patients who participate in the study.

Findings from this study will be used to identify interventions to ameliorate or eliminate existing barriers to treatment so that young women have access to high quality breast cancer treatment and care. Study findings will be disseminated through reports, presentations, and publications. Results will also be used by participating sites, CDC, and other federal agencies to improve care and services provided to young women diagnosed with breast cancer.

OMB approval is requested for three years and the burden table presents annualized estimates. CDC’s data collection contractor will securely maintain identifiable information from respondents recruited from state registries (Sample 1). No identifiable information will be collected by CDC. Participation is voluntary and there are no costs to respondents other than their time.

### ESTIMATED ANNUALIZED BURDEN HOURS

<table>
<thead>
<tr>
<th>Type of respondent</th>
<th>Form name</th>
<th>Number of respondents</th>
<th>Number of responses per respondent</th>
<th>Average burden per response (in hours)</th>
<th>Total burden (in hours)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample 1—Breast Cancer survivors included in one of as many as four state registries.</td>
<td>Breast Cancer in Young Women Survey (Mail or web-based version questionnaire)</td>
<td>583</td>
<td>1</td>
<td>22/60</td>
<td>214</td>
</tr>
<tr>
<td>Sample 2—Breast Cancer survivors associated with advocacy groups.</td>
<td>Breast Cancer in Young Women Survey (Web-based questionnaire)</td>
<td>667</td>
<td>1</td>
<td>22/60</td>
<td>244</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>458</td>
</tr>
</tbody>
</table>

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. 2015–20479 Filed 8–18–15; 8:45 am]
BILLING CODE 4163–18–P

---

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

[30Day–15–15NR]

Agency Forms Undergoing Paperwork Reduction Act Review

The Centers for Disease Control and Prevention (CDC) has submitted the following information collection request to the Office of Management and Budget (OMB) for review and approval in accordance with the Paperwork Reduction Act of 1995. The notice for the proposed information collection is published to obtain comments from the public and affected agencies.

Written comments and suggestions from the public and affected agencies concerning the proposed collection of information are encouraged. Your comments should address any of the following: (a) Evaluate whether the proposed collection of information is necessary for the proper performance of the functions of the agency, including whether the information will have practical utility; (b) Evaluate the accuracy of the agencies estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used; (c) Enhance the quality, utility, and clarity of the information to be collected; (d) Minimize the burden of the collection of information on those who are to respond, including through the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology, e.g., permitting electronic submission of responses; and (e) Assess information collection costs.

To request additional information on the proposed project or to obtain a copy of the information collection plan and instruments, call (404) 639–7570 or send an email to ombr@cdc.gov. Written comments and/or suggestions regarding the items contained in this notice should be directed to the Attention: CDC Desk Officer, Office of Management and Budget, Washington, DC 20503 or by fax to (202) 295–5806. Written comments should be received within 30 days of this notice.

Proposed Project


Background and Brief Description

The CDC is requesting the Office of Management and Budget (OMB) to grant a three-year approval to collect data that comprises Health Professional Application for Training (HPAT), the Training Follow-up Instrument, the Technical Assistance (TA) Satisfaction Instrument, and the Capacity Building Assistance (CBA) Key Informant Interview. The purpose of this information collection is to assess the degree to which the CDC's CBA program meets the needs of its consumers in order to enhance its capacity building strategy over time. The HPAT serves as the official application form for training and technical activities conducted by the Sexually Transmitted Disease (STD)/Human immunodeficiency virus (HIV) Prevention Training Centers’ (PTCs) grantees and the HIV Capacity Building Assistance (CBA) providers grantees funded by the CDC. The HPAT form is currently approved under OMB Control Number 0920–0995 and expires on October 31, 2016.