Type of respondents	Form name	Number of respondents/ year	Number of responses per respondent	Average burden per response (in hours)	Total annual burden hours
Chief Executives, Medical Scientists, Health Educators, Family/General	Symposium on Global Cancer Research.	150	1	20/60	50
Practitioners, Registered Nurses, Medical and Health Services Man-	Workshop in Cancer Control Planning and Implementation.	140	1	20/60	47
agers.	The Summer Curriculum in Cancer Prevention.	27	1	30/60	14
	Women's Empowerment Cancer Advocacy Network (We-Can).	140	1	20/60	47
	Regional Grant Writing and Peer Review Workshop.	60	1	30/60	30
	Other CGH Workshops	30	1	30/60	15

ESTIMATED ANNUALIZED BURDEN HOURS

Dated: May 1, 2014. Vivian Horovitch-Kelley,

NCI Project Clearance Liaison, National Institutes of Health.

[FR Doc. 2014–10590 Filed 5–7–14; 8:45 am]

BILLING CODE 4140-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Proposed Collection; 60-day Comment Request; NCI Cancer Genetics Services Directory Web-Based Application and Update Mailer

Summary: 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 National Cancer Institute (NCI), National Institutes of Health (NIH), will publish periodic summaries of proposed projects to be submitted to the Office of Management and Budget (OMB) for review and approval.

Written comments and/or suggestions from the public and affected agencies are invited on one or more of the following points: (1) Whether the proposed collection of information is necessary for the proper performance of the function of the agency, including whether the information will have practical utility; (2) The accuracy of the agency's estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used; (3) Ways to enhance the quality, utility, and

clarity of the information to be collected; and (4) Ways to minimize the burden of the collection of information on those who are to respond, including the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology.

To Submit Comments and for Further *Information:* To obtain a copy of the data collection plans and instruments, submit comments in writing, or request more information on the proposed project, contact: Margaret Beckwith. International Cancer Research Databank Branch, Office of Communications and Education, 9609 Medical Center Drive, MSC 9776, Bethesda, MD 20892-9776 or call non-toll-free number 240-376-6593 or Email your request, including your address to: mbeckwit@ mail.nih.gov. Formal requests for additional plans and instruments must be requested in writing.

Comment Due Date: Comments regarding this information collection are best assured of having their full effect if received within 60 days of the date of this publication.

Proposed Collection: NCI Cancer Genetics Services Directory Web-Based Application and Update Mailer, 0925– 0639, Date 08/31/2014, Revision, National Cancer Institute (NCI), National Institutes of Health (NIH).

Need and Use of Information Collection: The Office of Communications and Education International Cancer Research Databank Branch has created the NCI Cancer Genetics Services Directory on NCI's Web site Cancer.gov. This directory is a searchable collection of information about professionals who provide services related to cancer genetics. These services include cancer risk assessment, genetic counseling, and genetic susceptibility testing. The professionals have applied to be in the directory using an online application form and have met basic criteria outlined on the form.

There are currently 587 genetics professionals listed in the directory. Approximately 30–60 new professionals are added to the directory each year. The applicants are nurses, physicians, genetic counselors, and other professionals who provide services related to cancer genetics. The information collected on the application form includes name, professional qualifications, practice locations, and the area of specialization. The information is updated annually using a Web-based update mailer that mirrors the application form.

The NCI Cancer Genetics Services
Directory is a unique resource for cancer
patients and their families who are
looking for information about their
family risk of cancer and genetic
counseling. Collecting applicant
information and verifying it annually by
using the NCI Cancer Genetics Services
Directory Web-based Application Form
and Update Mailer is important for
providing this information to the public
and for keeping it current.

OMB approval is requested for 3 years. There are no costs to respondents other than their time. The total estimated annualized burden hours are 180.

ESTIMATED ANNUALIZED BURDEN HOURS

Form name	Type of respondent	Number of respondents	Number of responses per respondent	Average burden per response (in hours)	Total annual burden hours
Web-based Application Form	Genetics Professional	60 600	1 1	30/60 15/60	30 150

Dated: April 30, 2014.

Karla Bailey,

NCI Project Clearance Liaison, National Institutes of Health.

[FR Doc. 2014–10521 Filed 5–7–14; 8:45 am]

BILLING CODE 4140-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Proposed Collection; 60-Day Comment Request; Food and Drug Administration (FDA) and the National Cancer Institute (NCI) Health Communication Survey (FDA–NCI)

SUMMARY: 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 National Cancer Institute (NCI), National Institutes of Health (NIH), will publish periodic summaries of proposed projects to be submitted to the Office of Management and Budget (OMB) for review and approval.

Written comments and/or suggestions from the public and affected agencies are invited on one or more of the following points: (1) Whether the proposed collection of information is necessary for the proper performance of the function of the agency, including whether the information will have

practical utility; (2) The accuracy of the agency's estimate of the burden of the proposed collection of information, including the validity of the methodology and assumptions used; (3) Ways to enhance the quality, utility, and clarity of the information to be collected; and (4) Ways to minimize the burden of the collection of information on those who are to respond, including the use of appropriate automated, electronic, mechanical, or other technological collection techniques or other forms of information technology.

To Submit Comments and for Further Information: To obtain a copy of the data collection plans and instruments, submit comments in writing, or request more information on the proposed project contact: Bradford W. Hesse, Ph.D., Health Communication and Informatics Research Branch, 9609 Medical Center Drive, MSC 9761, Room 3E610, Rockville, MD 20850 or call nontoll free number 240–276–6721 or Email your request, including your address, to hesseb@mail.nih.gov. Formal requests for additional plans and instruments must be requested in writing.

Comment Due Date: Comments regarding this information collection are best assured of having their full effect if received within 60 days of the date of this publication.

Proposed Collection: Food and Drug Administration (FDA) and the National Cancer Institute (NCI) Health Communication Survey (FDA–NCI), 0925–NEW, National Cancer Institute (NCI), National Institutes of Health (NIH).

Need and Use of Information Collection: This partnership between NCI and FDA will include assessing the public's knowledge of medical devices, communications related to product recalls, nutritional supplement labeling, and topics to inform FDA's regulatory authority over tobacco, such as risk perceptions about new tobacco products, product pack color gradations, perceptions of product harm, and tobacco product claims and labels. This NCI-FDA survey will couple knowledge-related questions with inquiries into the communication channels through which understanding is being obtained, and assessment of FDA-regulated material. This survey will extend the information collected and priorities from the Health Information National Trends Survey (HINTS) which has been to provide a comprehensive assessment of the American public's current access to, and use of, information about cancer across the cancer care continuum from cancer prevention, early detection, diagnosis, treatment, and survivorship.

OMB approval is requested for 1 year. There are no costs to respondents other than their time. The total estimated annualized burden hours are 2,159.

ESTIMATED ANNUALIZED BURDEN HOURS

Type of respondents	Number of respondents	Number of responses per respondent	Average burden per response (in hours)	Total annual burden hour
Individuals	4,318	1	30/60	2,159

Dated: May 1, 2014. Vivian Horovitch-Kellev,

NCI Project Clearance Liaison, National Institutes of Health.

[FR Doc. 2014–10520 Filed 5–7–14; 8:45 am]

BILLING CODE 4140-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

National Center for Advancing Translational Sciences; Notice of Closed Meetings

Pursuant to section 10(d) of the Federal Advisory Committee Act, as

amended (5 U.S.C. App.), notice is hereby given of the following meetings.

The meetings will be closed to the public in accordance with the provisions set forth in sections 552b(c)(4) and 552b(c)(6), Title 5 U.S.C., as amended. The grant applications and the discussions could disclose confidential trade secrets or commercial property such as patentable material, and personal information concerning