

(3) train state and local personnel in health work. The BSC, NCEH/ATSDR provides advice and guidance to the Secretary, HHS; the Director, CDC and Administrator, ATSDR; and the Director, NCEH/ATSDR, regarding program goals, objectives, strategies, and priorities in fulfillment of the agency's mission to protect and promote people's health. The board provides advice and guidance that will assist NCEH/ATSDR in ensuring scientific quality, timeliness, utility, and dissemination of results. The board also provides guidance to help NCEH/ATSDR work more efficiently and effectively with its various constituents and to fulfill its mission in protecting America's health.

Matters for Discussion: The agenda items for the BSC Meeting will include NCEH/ATSDR Office of the Director updates; CDC Winnable Battles: Food Safety; consideration of a subcommittee to the BSC for childhood lead poisoning prevention; vote on recommendation regarding a subcommittee to the BSC for childhood lead poisoning prevention; radiation preparedness planning; NCEH/ATSDR Strategic Planning and Priorities; NCEH/ATSDR Priority: Water Safety; updates from the National Institute for Environmental Health Services, National Institute for Occupational Safety and Health, U.S. Department of Energy and the U.S. Environmental Protection Agency; NCEH/ATSDR Response to Prior BSC Guidance; discussion of future BSC agenda topics and action items.

Agenda items are subject to change as priorities dictate.

SUPPLEMENTARY INFORMATION: The public comment period is scheduled on Thursday, May 22, 2014 from 3:00 p.m. until 3:15 p.m., and on Friday, May 23, 2014 from 10:45 a.m. until 11:00 a.m.

Contact Person for More Information: Sandra Malcom, Committee Management Specialist, NCEH/ATSDR, 4770 Buford Highway, Mail Stop F-61, Chamblee, Georgia 30345; Telephone 770/488-0575 or 770/488-0755, Fax: 770/488-3377; Email: smalcom@cdc.gov. The deadline for notification of attendance is May 16, 2014.

The Director, Management Analysis and Services Office, has been delegated the authority to sign **Federal Register** notices pertaining to announcements of meetings and other committee management activities for both the Centers for Disease Control and

Prevention and the Agency for Toxic Substances and Disease Registry.

Gary J. Johnson,

Acting Director, Management Analysis and Services Office Centers for Disease Control and Prevention.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2014-N-0487]

Agency Information Collection Activities; Proposed Collection; Comment Request; Generic Clearance for the Collection of Qualitative Feedback on Agency Service Delivery

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing an opportunity for public comment on the proposed collection of certain information by the Agency. Under the Paperwork Reduction Act of 1995 (the PRA), Federal Agencies are required to publish notice in the **Federal Register** concerning each proposed collection of information, including each proposed extension of an existing collection of information, and to allow 60 days for public comment in response to the notice. This notice solicits comments on the collection of information involving a generic clearance for qualitative feedback on Agency service delivery.

DATES: Submit either electronic or written comments on the collection of information by June 30, 2014.

ADDRESSES: Submit electronic comments on the collection of information to <http://www.regulations.gov>. Submit written comments on the collection of information to the Division of Dockets Management (HFA-305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. All comments should be identified with the docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT: FDA PRA Staff, Office of Operations, Food and Drug Administration, 1350 Piccard Dr., PI50-400B, Rockville, MD 20850, PRAStaff@fda.hhs.gov.

SUPPLEMENTARY INFORMATION: Under the 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. "Collection of information" is defined in 44 U.S.C. 3502(3) and 5 CFR 1320.3(c) and includes Agency requests or requirements that members of the public submit reports, keep records, or provide information to a third party. Section 3506(c)(2)(A) of the PRA (44 U.S.C. 3506(c)(2)(A)) requires Federal Agencies to provide a 60-day notice in the **Federal Register** concerning each proposed collection of information, including each proposed extension of an existing collection of information, before submitting the collection to OMB for approval. To comply with this requirement, FDA is publishing notice of the proposed collection of information set forth in this document.

With respect to the following collection of information, FDA invites comments on these topics: (1) Whether the proposed collection of information is necessary for the proper performance of FDA's functions, including whether the information will have practical utility; (2) the accuracy of FDA'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 respondents, including through the use of automated collection techniques, when appropriate, and other forms of information technology.

Generic Clearance for the Collection of Qualitative Feedback on Agency Service Delivery (OMB Control Number 0697)—Extension

The information collection activity will garner qualitative customer and stakeholder feedback in an efficient, timely manner, in accordance with the Administration's commitment to improving service delivery. By qualitative feedback we mean information that provides useful insights on perceptions and opinions, but are not statistical surveys that yield quantitative results that can be generalized to the population of study. This feedback will provide insights into customer or stakeholder perceptions; experiences and expectations; provide an early warning of issues with service; or focus attention on areas where communication, training, or changes in operations might improve delivery of products or services. These collections will allow for ongoing collaborative and actionable communications between the Agency and its customers and stakeholders. It will also allow feedback

to contribute directly to the improvement of program management. Feedback collected under this generic clearance will provide useful information, but it will not yield data that can be generalized to the overall population. This type of generic clearance for qualitative information will not be used for quantitative information collections that are designed to yield reliably actionable results, such as monitoring trends over

time or documenting program performance. Such data uses require more rigorous designs that address the following: The target population to which generalizations will be made, the sampling frame, the sample design (including stratification and clustering), the precision requirements or power calculations that justify the proposed sample size, the expected response rate, methods for assessing potential non-response bias, the protocols for data

collection, and any testing procedures that were or will be undertaken prior to fielding the study. Depending on the degree of influence the results are likely to have, such collections may still be eligible for submission for other generic mechanisms that are designed to yield quantitative results. FDA estimates the burden of this collection of information as follows:

TABLE 1—ESTIMATED ANNUAL REPORTING BURDEN¹

| Activity | Number of respondents | Number of responses per respondent | Total annual responses | Average burden per response | Total hours |
|-------------------------------------|-----------------------|------------------------------------|------------------------|-----------------------------|-------------|
| Focus groups | 725 | 1 | 725 | 1 hour, 45 minutes | 1,269 |
| Customer comment cards/forms | 1,200 | 1 | 1,200 | 15 minutes | 300 |
| Small discussion groups | 725 | 1 | 725 | 1 hour, 45 minutes | 1,269 |
| Customer satisfaction surveys | 6,450 | 1 | 6,450 | 20 minutes | 2,129 |
| Total | 4,967 | | | | |

¹ There are no capital costs or operating and maintenance costs associated with this collection of information.

Dated: April 23, 2014.
Leslie Kux,
Assistant Commissioner for Policy.
 [FR Doc. 2014-09693 Filed 4-28-14; 8:45 am]
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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration
 [Docket No. FDA-2010-N-0623]

Agency Information Collection Activities; Submission for Office of Management and Budget Review; Comment Request; Voluntary Cosmetic Registration Program

AGENCY: Food and Drug Administration, HHS.
ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA or we) is announcing that a proposed collection of information has been submitted to the Office of Management and Budget (OMB) for review and clearance under the Paperwork Reduction Act of 1995.

DATES: Fax written comments on the collection of information by May 29, 2014.

ADDRESSES: To ensure that comments on the information collection are received, OMB recommends that written comments be faxed to the Office of Information and Regulatory Affairs, OMB, Attn: FDA Desk Officer, FAX: 202-395-7285, or emailed to oir_submission@omb.eop.gov. All comments should be identified with the

OMB control number 0910-0027. Also include the FDA docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT: FDA PRA Staff, Office of Operations, Food and Drug Administration, 1350 Piccard Dr., PI50-400B, Rockville, MD 20850, PRAStaff@fda.hhs.gov.

SUPPLEMENTARY INFORMATION: In compliance with 44 U.S.C. 3507, FDA has submitted the following proposed collection of information to OMB for review and clearance.

Voluntary Cosmetic Registration Program—21 CFR Parts 710 and 720 (OMB Control Number 0910-0027)—Extension

The Federal Food, Drug, and Cosmetic Act (the FD&C Act) provides us with the authority to regulate cosmetic products in the United States. Cosmetic products that are adulterated under section 601 of the FD&C Act (21 U.S.C. 361) or misbranded under section 602 of the FD&C Act (21 U.S.C. 362) may not be distributed in interstate commerce. We have developed the VCRP to assist us in carrying out our responsibility to regulate cosmetics.

In 21 CFR part 710, we request that establishments that manufacture or package cosmetic products register with us on Form FDA 2511 entitled “Registration of Cosmetic Product Establishment.” The term “Form FDA 2511” refers to both the paper and electronic versions of the form. The electronic version of Form FDA 2511 is available on our VCRP Web site at

<http://www.fda.gov/Cosmetics/RegistrationProgram/OnlineRegistration/ucm090947.htm>. We strongly encourage electronic registration of Form FDA 2511 because it is faster and more convenient. A registering facility will receive confirmation of electronic registration, including a registration number, by email, usually within 7 business days. The online system also allows for amendments to past submissions.

Because registration of cosmetic product establishments is not mandatory, voluntary registration provides us with the best information available about the locations, business trade names, and types of activity (manufacturing or packaging) of cosmetic product establishments. We place the registration information in a computer database and use the information to generate mailing lists for distributing regulatory information and for inviting firms to participate in workshops on topics in which they may be interested. We also use the information for estimating the size of the cosmetic industry and for conducting onsite establishment inspections. Registration is permanent, although we request that respondents submit an amended Form FDA 2511 if any of the originally submitted information changes.

In part 720 (21 CFR part 720), we request that firms that manufacture, pack, or distribute cosmetics file with us an ingredient statement for each of their products. Ingredient statements for new submissions (§§ 720.1 through 720.4) are reported on Form FDA 2512,