FOR FURTHER INFORMATION CONTACT:

Karen Shields, Grants Policy Specialist, Department of Health and Human Services, Administration for Children and Families, OA/Division of Grants Policy, 370 L'Enfant Promenade, SW., Aerospace Building, 6th Floor East, Washington, DC 20447. Email: karen.shields@acf.hhs.gov. Fax: (202) 205–6400.

Dated: October 21, 2011.

Jason Donaldson,

Deputy Assistant Secretary for Administration, Administration for Children and Families.

[FR Doc. 2011–27878 Filed 10–26–11; 8:45 am]

BILLING CODE 4184-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2011-N-0528]

Food Safety Modernization Act
Domestic and Foreign Facility
Reinspections, Recall, and Importer
Reinspection User Fee Rates for Fiscal
Year 2012; Extension of Comment
Period

AGENCY: Food and Drug Administration, HHS

ACTION: Notice; extension of comment period.

SUMMARY: The Food and Drug Administration (FDA) is extending the comment period to November 30, 2011, for the notice entitled, "Food Safety Modernization Act Domestic and Foreign Facility Reinspections, Recall, and Importer Reinspection User Fee Rates for Fiscal Year 2012" that appeared in the Federal Register of August 1, 2011 (76 FR 45820). In that document, FDA announced the establishment of a docket to obtain comments that would be considered in establishing the fee rates for fiscal year (FY) 2013. In particular, the Agency provided the current FY 2012 fees and requested public comments to the document and intends to consider such comments, as well as experience and additional data gained in implementing these fees in FY 2012, in establishing the fee rates for FY 2013. The Agency is taking this action in response to requests for an extension to allow interested persons additional time to submit comments.

DATES: Submit either electronic or written comments by November 30, 2011.

ADDRESSES: Submit electronic comments to *http://*

www.regulations.gov. Submit written comments to the Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852.

FOR FURTHER INFORMATION CONTACT:

Amy Waltrip, 12420 Parklawn Dr., rm. 2012, Rockville, MD 20857, (301) 796–8811, email: Amy.Waltrip@fda.hhs.gov.

SUPPLEMENTARY INFORMATION:

I. Background

In the **Federal Register** of August 1, 2011 (76 FR 45820), FDA published a notice with a 90-day comment period to request comments on the establishment of domestic and foreign facility reinspections, non-compliance with recall order, and importer reinspection FY 2012 user fees. The FDA Food Safety Modernization Act provides the Agency with authority under section 743 of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 379j-31) to assess and collect fees, including those for costs associated with certain domestic and foreign facility reinspections, failure to comply with a recall order, and importer reinspections. The Agency is seeking public comment on the established FY 2012 user fees. In particular, the Agency is seeking public comments intending to consider such comments, as well as experience and additional data gained in implementing these user fees in FY 2012, in establishing the fee rates for FY 2013. The Agency has received a request for an extension of the comment period. The request conveyed concern that the current 90-day comment period does not allow sufficient time to develop a meaningful or thoughtful response to the notice.

FDA has considered the request and is extending the comment period for the notice for 30 days until November 30, 2011. The Agency believes that this extension allows adequate time for interested persons to submit comments.

II. Request for Comments

Interested persons may submit to the Division of Dockets Management (see ADDRESSES) either electronic or written comments on this document. It is only necessary to send one set of comments. It is no longer necessary to send two copies of mailed comments. Identify comments with the docket number found in brackets in the heading of this document. Received comments may be seen in the Division of Dockets Management between 9 a.m. and 4 p.m., Monday through Friday.

Dated: October 24, 2011.

Leslie Kux,

Acting Assistant Commissioner for Policy.
[FR Doc. 2011–27845 Filed 10–26–11; 8:45 am]
BILLING CODE 4160–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Submission for OMB Review; Comment Request: National Institutes of Health Construction Grants

Summary: Under the provisions of Section 3507(a)(1)(D) of the Paperwork Reduction Act of 1995, the National Institutes of Health (NIH) has submitted to the Office of Management and Budget (OMB) a request to review and approve the information collection listed below. This proposed information collection was previously published in the Federal Register on August 17, 2011, pages 51042-51043, and allowed 60 days for public comment. No public comments were received. The purpose of this notice is to allow an additional 30 days for public comment. The NIH may not conduct or sponsor, and the respondent is not required to respond to, information that has been extended, revised or implemented on or after October 1, 2008, unless it displays a currently valid OMB control number.

Proposed Collection: Title: National Institutes of Health Construction GrantsB42 CFR part 52b (Final Rule). *Type of Information Collection Request:* Extension of No. 0925-0424, expiration date 8/31/2008. Need and Use of the *Information Collection:* This request is for OMB review and approval of an extension for the information collection and recordkeeping requirements contained in the regulation codified at 42 CFR part 52b. The purpose of the regulation is to govern the awarding and administration of grants awarded by NIH and its components for construction of new buildings and the alteration, renovation, remodeling, improvement, expansion, and repair of existing buildings, including the provision of equipment necessary to make the buildings (or applicable part of the buildings) suitable for the purpose for which it was constructed. In terms of reporting requirements: Section 52b.9(b) of the regulation requires the transferor of a facility which is sold or transferred, or owner of a facility, the use of which has changed, to provide written notice of the sale, transfer or change within 30 days. Section 52b.10(f) requires a grantee to submit an approved copy of the construction

schedule prior to the start of construction. Section 52b.10(g) requires a grantee to provide daily construction logs and monthly status reports upon request at the job site. Section 52b.11(b) requires applicants for a project involving the acquisition of existing facilities to provide the estimated cost of

the project, cost of the acquisition of existing facilities, and cost of remodeling, renovating, or altering facilities to serve the purposes for which they are acquired. In terms of recordkeeping requirements: Section 52b.10(g) requires grantees to maintain daily construction logs and monthly

status reports at the job site. Frequency of Response: On occasion. Affected Public: Non-profit organizations and Federal agencies. Type of respondents: Grantees. The estimated respondent burden is as follows:

ESTIMATED ANNUAL REPORTING AND RECORDKEEPING BURDEN

| | Number of respondents | Frequency of response | Average time per response | Annual hour burden |
|------------------|-----------------------|-----------------------|---------------------------|--------------------|
| Reporting: | | | | |
| Section 52b.9(b) | 1 | 1 | .50 | .50 |
| Section 2b.10(f) | 60 | 1 | 1.0 | 60 |
| Section 2b.10(g) | 60 | 12 | 1.0 | 720 |
| Section 2b.11(b) | 100 | 1 | 1.0 | 100 |
| Recordkeeping. | | | | |
| Section 2b.10(g) | 60 | 260 | 1.0 | 15,600 |
| Totals | 281 | | | 16,480.5 |

The annualized cost to the public, based on an average of 60 active grants in the construction phase, is estimated at: \$576,818. There are no Capital Costs to report. There are no operating or Maintenance Costs to report.

Request for Comments: Written comments and/or suggestions from the public and affected agencies should address one or more of the following points: (1) Evaluate whether the proposed collection of information and recordkeeping are necessary for the proper performance of the function of the agency, including whether the information will have practical utility; (2) Evaluate the accuracy of the agency=s estimate of the burden of the proposed collection of information and recordkeeping, including the methodology and assumptions used; (3) Enhance the quality, utility, and clarity of the information to be collected and the recordkeeping information to be maintained; and (4) 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 and recordkeeping techniques of other forms of information technology.

Direct Comments to OMB: Written comments and/or suggestions regarding the item(s) contained in this notice, especially regarding the estimated public burden and associated response time, should be directed to the Office of Regulatory Affairs.

OĪRA_submission@omb.eop.gov or by fax to (202) 395–6974, Attention: Desk Officer for NIH. To request more information on the proposed project or to obtain a copy of the data collection plans and instruments, contact Jerry Moore, NIH Regulations Officer, Office of Management Assessment, Division of Management Support, National Institutes of Health, 6011 Executive Boulevard, Room 601, MSC 7669, Rockville Maryland 20852; call (301) 496–4607 (this is not a toll free number) or email your request to *jm40z@nih.gov*.

Comments Due Date: Comments regarding this information collection are best assured of having their full effect if received within 30 days of the date of this publication.

Dated: October 20, 2011.

Jerry Moore,

NIH Regulations Officer, National Institutes of Health.

[FR Doc. 2011–27850 Filed 10–26–11; 8:45 am]

BILLING CODE 4140–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Submission for OMB Review; Comment Request: New Proposed Collection, Neuropsychosocial Measures Formative Research Methodology Studies for the National Children's Study

Summary: Under the provisions of Section (3507(a)(1)(D)) of the Paperwork Reduction Act of 1995, the National Institutes of Health has submitted to the Office of Management and Budget (OMB) a request to review and approve the information collection listed below. This proposed information collection was previously published in the Federal Register on May 2, 2011, pages 24497—24498, and allowed 60 days for public comment. Two written comments and

two verbal comments were received. The verbal comments expressed support for the broad scope of the study. The written comments were identical and questioned the cost and utility of the study. The purpose of this notice is to allow an additional 30 days for public comment. The National Institutes of Health may not conduct or sponsor, and the respondent is not required to respond to, an information collection that has been extended, revised, or implemented on or after October 1, 1995, unless it displays a currently valid OMB control number.

Proposed Collection: Title: Neurodevelopmental and Psycho-Social Measures Formative Research Studies for the National Children's Study (NCS). Type of Information Request: New. Need and Use of Information Collection: The Children's Health Act of 2000 (Pub. L. 106–310) states:

(a) Purpose.—It is the purpose of this section to authorize the National Institute of Child Health and Human Development* to conduct a national longitudinal study of environmental influences (including physical, chemical, biological, and psychosocial) on children's health and development.

(b) In General.—The Director of the National Institute of Child Health and Human Development* shall establish a consortium of representatives from appropriate Federal agencies (including the Centers for Disease Control and Prevention, the Environmental Protection Agency) to—

(1) Plan, develop, and implement a prospective cohort study, from birth to adulthood, to evaluate the effects of both chronic and intermittent exposures on child health and human development; and