To obtain copies of the supporting statement and any related forms for the proposed paperwork collections referenced above, access CMS' Web site at *http://www.cms.hhs.gov/ PaperworkReductionActof1995*, or Email your request, including your address, phone number, OMB number, and CMS document identifier, to *Paperwork@cms.hhs.gov*, or call the Reports Clearance Office on (410) 786– 1326.

In commenting on the proposed information collections please reference the document identifier or OMB control number. To be assured consideration, comments and recommendations must be submitted in one of the following ways by *December 28, 2010:*

1. *Electronically*. You may submit your comments electronically to *http:// www.regulations.gov*. Follow the instructions for "Comment or Submission" or "More Search Options" to find the information collection document(s) accepting comments.

2. *By regular mail.* You may mail written comments to the following address:

CMS, Office of Strategic Operations and Regulatory Affairs, Division of Regulations Development, Attention: Document Identifier/OMB Control Number, Room C4–26–05, 7500 Security Boulevard, Baltimore, Maryland 21244–1850.

Dated: October 26, 2010.

Martique Jones,

Director, Regulations Development Group, Division B, Office of Strategic Operations and Regulatory Affairs.

[FR Doc. 2010–27470 Filed 10–28–10; 8:45 am]

BILLING CODE 4120-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Submission for OMB Review; Comment Request; Cancer Trials Support Unit (CTSU) Public Use Forms and Customer Satisfaction Surveys (NCI)

SUMMARY: Under the provisions of Section 3507(a)(1)(D) of the Paperwork Reduction Act of 1995, the National Cancer Institute (NCI), 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 July 13, 2010 (75 FR 39950) and allowed 60-days for public comment. There have been no public comments. Additionally, the 30-day Federal Register was published on September 13, 2010. The purpose of this notice is to allow an additional 30 days for public comment to the revisions. 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: Cancer Trial Support Unit (CTSU). Type of Information Collection Request: Existing Collection in Use Without an OMB Number. Need and Use of Information Collection: CTSU collects annual surveys of customer satisfaction for clinical site staff using the CTSU Help Desk and the CTSU Web site. An ongoing user satisfaction survey is in place for the Oncology Patient

Enrollment Network (OPEN). User satisfaction surveys are compiled as part of the project quality assurance activities and used to direct improvements to processes and technology. In addition, the CTSU collects standardized forms to process site regulatory information, changes to membership, patient enrollment data, and routing information for case report forms. This questionnaire adheres to The Public Health Service Act, Section 413 (42 U.S.C. 285a-2) authorizes CTEP to establish and support programs to facilitate the participation of qualified investigators on CTEP-supported studies, and to institute programs that minimize redundancy among grant and contract holders, thereby reducing overall cost of maintaining a robust treatment trials program. Based on a conversation with the Office of Management and Budget on October 17, 2010, the burden table has been revised to take into account future submissions of a generic data transmittal forms (see Attachment 1gg in the Table below). It was agreed that the generic forms will be finalized and submitted in the future as non-substantive change requests for OMB clearance as needed. Frequency of *Response:* The help desk and Web site survey are collected annually. The OPEN survey is ongoing. The form submissions vary depending on the purpose of the form and the activity of the local site. Affected Public: CTSU's target audience is staff members at clinical sites and CTEP-supported programs. Respondent and burden estimates are listed in the Table below. The annualized burden is estimated to be 34,802 hours and the annualized cost to respondents is estimated to be \$946, 601. There are no Capital Costs, Operating Costs, and/or Maintenance Costs to report.

Attach No.	Section/form or survey title	Use metrics/ month— # respond	Estimated time for site to complete (minutes)	Estimated burden (minutes/ hours)	Frequency of response	Total annual usage/annual burden hours
Regulatory/Roster						
1a	CTSU IRB/Regulatory Approval Transmittal Form.	9,000	2	0.03	12.00	3,600
1b	CTSU IRB Certification Form	8,500	10	0.17	12.00	17,000
1c	CTSU Acknowledgement Form	500	5	0.08	12.00	500
1d	Optional Form 1—Withdrawal from Protocol Participation Form.	50	5	0.08	12.00	50
Roster Forms						
1e	CTSU Roster Update Form	50	2–4	0.07	12.00	40
1f	CTSU Radiation Therapy Facilities Inventory Form.	20	30	0.50	12.00	120
Drug shipment						

Attach No.	Section/form or survey title	Use metrics/ month— # respond	Estimated time for site to complete (minutes)	Estimated burden (minutes/ hours)	Frequency of response	Total annual usage/annual burden hours
1g 1h	CTSU IBCSG Drug Accountability Form CTSU IBCSG Transfer of Investigational Agent Form.	11 3	5–10 20	0.17 0.33	12.00 12.00	22 12
Data Mana	agement					
1i 1j 1k	Site Initiated Data Update Form (generic) N0147 CTSU Data Transmittal Form Site Intimated Data Update Form (DUF), Pro- tocol: NCCTG N0147*.	100 1000 75	5–10 5–10 5–10	0.17 0.17 0.17	12.00 12.00 12.00	200 2,000 150
11	TAILORX/PACCT 1 CTSU Data Transmittal Form.	2100	5–10	0.17	12.00	4,200
1m 1n	Data Clarification Form Unsolicited Data Modification Form (UDM), Protocol: TAILORx/PACCT1.	650 75	15–20 5–10	0.33 0.17	12.00 12.00	2,600 150
10 1p 1q	Z4032 CTSU Data Transmittal Form Z1031 CTSU Data Transmittal Form Z1041 CTSU Data Transmittal Form	50 50 50	5–10 5–10 5–10	0.17 0.17 0.17	12.00 12.00 12.00	100 100 100
1r 1s 1t	Z6051 CTSU Data Transmittal Form RTOG 0834 CTSU Data Transmittal Form* CTSU 7868 Data Transmittal Form Site Initiated Data Lindets Form protocol 7869	75 60 50	5–10 5–10 5–10	0.17 0.17 0.17	12.00 12.00 12.00	150 120 100
1u 1v 1w 1x	Site Initiated Data Update Form, protocol 7868 MC0845(8233) CTSU Data Transmittal* 8121 CTSU Data Transmittal Form* Site Initiated Data Update Form, Protocol	10 50 100 10	5–10 5–10 5–10 5–10	0.17 0.17 0.17 0.17	12.00 12.00 12.00 12.00	20 100 200 20
1y	8121. USMCI 8214/Z6091: CTSU Data Transmittal *In Development	50	5–10	0.17	12.00	100
1z	USMCI 8214/Z6091 Crossover Request/ Checklist Transmittal Form.	5	5–10	0.17	12.00	10
Patient Er	nrollment					
1aa 1bb 1cc	CTSU Patient Enrollment Transmittal Form CTSU P2C Enrollment Transmittal Form CTSU Transfer Form	600 30 40	5–10 5–10 5–10	0.17 0.17 0.17	12.00 12.00 12.00	1,200 60 80
Administr	ative					
1dd 1ee 1ff 1gg	CTSU System Account Request Form CTSU Request for Clinical Brochure CTSU Supply Request Form CTSU Generic Data Transmittal Form	50 35 130 500	15–20 10 5–10 5–10	0.33 0.17 0.17 0.17	12.00 12.00 12.00 12.00	200 70 260 1000.00
Surveys/V	Veb Forms					
2 3 4	CTSU Web Site Customer Satisfaction Survey CTSU Helpdesk Customer Satisfaction Survey CTSU OPEN Survey	250 300 120	10–15 10–15 10–15	0.2500 0.2500 0.2500	1.00 1.00 1.00	63 75 30
Annual To	• • • 21,770					34,802
						-

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 is 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, including the validity of the methodology and assumptions used; (3) Enhance the quality, utility, and clarity of the information to be collected; 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 techniques or 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 Attention: NIH Desk Officer, Office of Management and Budget, at *OIRA_submission@omb.eop.gov* or by fax to 202–395–6974. To request more information on the proposed project or to obtain a copy of the data collection plans and instruments, contact Michael Montello, Pharm. D., CTEP, 6130 Executive Blvd., Rockville, MD 20852, call non-toll-free number 301–435–9206 or e-mail your request, including your address to: *montellom@mail.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 21, 2010.

Vivian Horovitch-Kelley,

NCI Project Clearance Liaison, National Institutes of Health.

[FR Doc. 2010–27330 Filed 10–28–10; 8:45 am] BILLING CODE 4140–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2010-D-0378]

Draft Compliance Policy Guide Sec. 690.800 Salmonella in Animal Feed; Availability; 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 to December 31, 2010, the comment period for a notice of availability of a draft compliance policy guide (CPG) that appeared in the Federal Register of August 2, 2010 (75 FR 45130). In the document, FDA requested comments on its proposal that certain criteria should be considered in recommending enforcement action against animal feed or feed ingredients that are adulterated due to the presence of *Salmonella*. 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 December 31, 2010.

ADDRESSES: Submit electronic comments on the draft CPG to *http:// www.regulations.gov.* Submit written comments on the draft CPG to the Division of Dockets Management (HFA– 305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852.

FOR FURTHER INFORMATION CONTACT: Kim Young, Center for Veterinary Medicine (HFV–230), Food and Drug Administration, 7519 Standish Pl., MPN–4, rm. 106, Rockville, MD 20855, 240–276–9200, e-mail: *Kim.young@fda.hhs.gov.*

SUPPLEMENTARY INFORMATION:

I. Background

In the Federal Register of August 2, 2010 (75 FR 45130), FDA published a notice of availability of a draft CPG with a 90-day comment period to request comments on its proposal that certain criteria should be considered in recommending enforcement action against animal feed or feed ingredients that are adulterated due to the presence of Salmonella. The Agency has received a request for a 60-day extension of the comment period for the draft CPG. 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 draft CPG. FDA has considered the request and is extending the comment period for the draft CPG for 60 days, until December 31, 2010.

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 26, 2010.

Dara Corrigan,

Associate Commissioner for Regulatory Affairs.

[FR Doc. 2010–27448 Filed 10–28–10; 8:45 am] BILLING CODE 4160–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Centers for Disease Control and Prevention

Advisory Board on Radiation and Worker Health (ABRWH or Advisory Board), National Institute for Occupational Safety and Health (NIOSH)

In accordance with section 10(a)(2) of the Federal Advisory Committee Act (Pub. L. 92–463), and pursuant to the requirements of 42 CFR 83.15(a), the Centers for Disease Control and Prevention (CDC), announces the following meeting of the aforementioned committee:

Board Public Meeting Times and Dates (All Times Are Mountain Time)

8:15 a.m.–5:15 p.m., November 16, 2010.

8:15 a.m.–5:15 p.m., November 17, 2010.

8:15 a.m.–12 p.m., November 18, 2010

Public Comment Times and Dates (All Times Are Mountain Time)

5:30 p.m.–7 p.m.,* November 16, 2010.

5:30 p.m.–6:30 p.m.,* November 17, 2010.

*Please note that the public comment periods may end before the times indicated, following the last call for comments. Members of the public who wish to provide public comments should plan to attend public comment sessions at the start times listed.

Place: Hilton Santa Fe Historic Plaza, 100 Sandoval Street, Santa Fe, New Mexico; Phone: 505–988–2811; Fax: 505–986–6439. Audio Conference Call via FTS Conferencing. The USA toll-free dial-in number is 1–866–659–0537 with a pass code of 9933701.

Status: Open to the public, limited only by the space available. The meeting space accommodates approximately 150 people.

Background: The Advisory Board was established under the Energy Employees Occupational Illness Compensation Program (EEOICP) Act of 2000 to advise the President on a variety of policy and technical functions required to implement and effectively manage the new compensation program. Kev functions of the Advisory Board include providing advice on the development of probability of causation guidelines which have been promulgated by the Department of Health and Human Services (HHS) as a final rule, advice on methods of dose reconstruction which have also been promulgated by HHS as a final rule, advice on the scientific validity and quality of dose estimation and reconstruction efforts being performed for purposes of the compensation program, and advice on petitions to add classes of workers to the Special Exposure Cohort (SEC).

In December 2000, the President delegated responsibility for funding, staffing, and operating the Advisory Board to HHS, which subsequently delegated this authority to the CDC. NIOSH implements this responsibility for CDC. The charter was issued on August 3, 2001, renewed at appropriate intervals, and will expire on August 3, 2011.

Purpose: This Advisory Board is charged with (a) Providing advice to the Secretary, HHS, on the development of guidelines under Executive Order 13179; (b) providing advice to the Secretary, HHS, on the scientific validity and quality of dose