the animal, where the revenue from the sale of apparel or fur products is not the primary source of income of such person.

13. Amend § 301.41 by removing paragraph (a)(7) and by revising paragraph (a)(4) to read as follows:

### § 301.41 Maintenance of Records.

(a) \* \* \*

(4) That the fur product is composed in whole or in substantial part of paws, tails, bellies, gills, ears, throats, heads, scrap pieces, or waste fur, when such is the fact;

\* \* \* \* \*

By direction of the Commission.

### Donald S. Clark,

RIN 0910-AG31

Secretary.

[FR Doc. 2012-22568 Filed 9-14-12; 8:45 am]

BILLING CODE 6750-01-P

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

## **Food and Drug Administration**

21 CFR Parts 16, 801, 803, 806, 810, 814, 820, 821, 822, and 830

[Docket No. FDA-2011-N-0090]

Agency Information Collection Activities; Proposed Collection; Unique Device Identification System; Extension of Comment Period

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Notification; extension of comment period.

SUMMARY: The Food and Drug Administration (FDA) is extending the comment period pertaining to information collection issues under the Paperwork Reduction Act of 1995 (the PRA) associated with the proposed rule, Unique Device Identification System, that appeared in the Federal Register of July 10, 2012 (77 FR 40736). 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 on the proposed collection of information by October 25, 2012

ADDRESSES: Submit electronic comments on the collection of information to the Office of Regulatory Affairs, Office of Management and Budget (OMB) at FAX: 202–395–7285, or email comments to OIRA submissions@omb.eop.gov. Please

mark your comment to the FDA desk officer and reference this rule.

FOR FURTHER INFORMATION CONTACT: Jay Crowley, Center for Devices and Radiological Health, Food and Drug Administration, 10903 New Hampshire Ave., Silver Spring, MD 20993, 301–796–5995, email: cdrhudi@fda.hhs.gov. SUPPLEMENTARY INFORMATION:

#### I. Background

In the **Federal Register** of July 10, 2012 (77 FR 40736), FDA published a notice of proposed rulemaking with a 60-day comment period concerning the proposed information collection. Comments on the proposed rulemaking will inform FDA's rulemaking to establish regulations for Unique Device Identification System.

The Agency has received requests for a 45-day extension of the comment period for the information collection. Each request conveyed concern that the current 60-day comment period does not allow sufficient time to develop a meaningful or thoughtful response to the information collection.

FDA has considered the requests and is extending the comment period for the information collection for 45 days, until October 25, 2012. The Agency believes that a 45-day extension allows adequate time for interested persons to submit comments without significantly delaying rulemaking on these important issues.

Dated: September 12, 2012.

### Leslie Kux,

Assistant Commissioner for Policy. [FR Doc. 2012–22821 Filed 9–14–12; 8:45 am]

BILLING CODE 4160-01-P

# DEPARTMENT OF HEALTH AND HUMAN SERVICES

## **Food and Drug Administration**

## 21 CFR Chapter I

[Docket No. FDA-2012-N-0780]

Regulatory New Drug Review: Solutions for Study Data Exchange Standards; Notice of Meeting; Request for Comments; Correction

AGENCY: Food and Drug Administration,

**ACTION:** Announcement of meeting; request for comments; correction.

SUMMARY: The Food and Drug Administration (FDA) is correcting a document that appeared in the **Federal Register** of August 14, 2012 (77 FR 48491). The document announced a meeting entitled "Regulatory New Drug Review: Solutions for Study Data Exchange Standards." The document was published with an incorrect email address. This document corrects that error.

FOR FURTHER INFORMATION CONTACT: Ron Fitzmartin, Office of Planning & Informatics, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 51, Rm. 1160, Silver Spring, MD 20993–0002, 301–796–5333, FAX: 301–847–8443, email: CDERDataStandards@fda.hhs.gov.

**SUPPLEMENTARY INFORMATION:** In FR Doc. 2012–19748, appearing on page 48491 in the **Federal Register** of August 14, 2012, the following corrections are made:

1. On page 48491, in the first column, in the FOR FURTHER INFORMATION CONTACT section, the email address "CDERDataStandards@hhs.fda.gov" is corrected to read "CDERDataStandards@fda.hhs.gov."

2. On page 48491, in the second column, in the SUPPLEMENTARY INFORMATION section, under "Registration," the email address "CDERDataStandards@hhs.fda.gov" is corrected to read "CDERDataStandards@fda.hhs.gov."

Dated: September 11, 2012.

#### Leslie Kux.

Assistant Commissioner for Policy.
[FR Doc. 2012–22793 Filed 9–14–12; 8:45 am]
BILLING CODE 4160–01–P

## **DEPARTMENT OF THE TREASURY**

## Office of the Secretary

31 CFR Part 10

[REG-138367-06]

RIN 1545-BF96

# Regulations Governing Practice Before the Internal Revenue Service

**AGENCY:** Office of the Secretary, Treasury.

**ACTION:** Withdrawal of notice of proposed rulemaking; notice of proposed rulemaking and notice of public hearing.

**SUMMARY:** This document proposes modifications of the regulations governing practice before the Internal Revenue Service (IRS). These proposed regulations affect individuals who practice before the IRS. These proposed regulations modify the standards governing written advice and update certain provisions as appropriate. This document also provides notice of a