understanding of the technical regulation of these systems is the essential to the such purposes. Essential to the system manufacturers to help ensure the images produced for intended over the last decade. FDA regulates WSI systems, have accelerated the adoption of digital imaging in pathology, similar to digital microscopy, in particular the development of whole slide scanning systems, have accelerated the adoption of digital imaging in pathology, similar to the digital transformation that radiology departments have experienced over the last decade. FDA regulates WSI system manufacturers to help ensure that the images produced for intended clinical uses are safe and effective for such purposes. Essential to the regulation of these systems is the understanding of the technical performance of the WSI system and the components in the imaging chain—from image acquisition to image display, and their effect on pathologist’s diagnostic performance and workflow. This guidance provides industry and Agency staff with recommendations regarding the technical performance assessment for regulatory evaluation of a digital WSI system. This document does not cover the clinical submission data that may be necessary to support approval or clearance. The guidance provides suggestions on how to best characterize the technical aspects that are relevant to WSI performance for their intended use and determine any possible limitations that might affect their safety and effectiveness. In the Federal Register of February 25, 2015 (80 FR 10122), FDA announced the availability of the draft guidance and interested persons were invited to comment by May 25, 2015. II. Significance of Guidance This guidance is being issued consistent with FDA’s good guidance practices regulation (21 CFR 10.115). The guidance represents the current thinking of FDA on technical performance assessment of digital pathology whole slide imaging devices. It does not establish any rights for any person and is not binding on FDA or the public. You can use an alternative approach if it satisfies the requirements of the applicable statutes and regulations. III. Electronic Access Persons interested in obtaining a copy of the guidance may do so by downloading an electronic copy from the Internet. A search capability for all Center for Devices and Radiological Health guidance documents is available at http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/Guidance Documents/default.htm. Guidance documents are also available at http://www.regulations.gov. Persons unable to download an electronic copy of “Technical Performance Assessment of Digital Pathology Whole Slide Imaging Devices” may send an email request to CDRH-Guidance@fda.hhs.gov to receive an electronic copy of the document. Please use the document number 1400053 to identify the guidance you are requesting. IV. Paperwork Reduction Act of 1995 This guidance refers to previously approved collections of information found in FDA regulations. These collections of information are subject to review by the Office of Management and Budget (OMB) under the Paperwork Reduction Act of 1995 (44 U.S.C. 3501–3520). The collections of information in 21 CFR part 807, subpart E (premarket notification) have been approved under OMB control number 0910–0120; the collections of information in 21 CFR part 814 (premarket approval) have been approved under OMB control number 0910–0231; and the collections of information in 21 CFR part 801 and 21 CFR 809.10 (labeling) have been approved under OMB control number 0910–0485.

Dated: April 13, 2016.

Leslie Kux, Associate Commissioner for Policy.

[FR Doc. 2016–09140 Filed 4–19–16; 8:45 am]

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2015–D–3056]

Distributor Labeling for New Animal Drugs; Guidance for Industry; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice of availability.

SUMMARY: The Food and Drug Administration (FDA or Agency) is announcing the availability of guidance for industry #231 entitled “Distributor Labeling for New Animal Drugs.” This guidance discusses FDA’s current thinking with respect to the factors it considers in determining whether to take regulatory action against distributor labeling for a new animal drug that differs from the labeling approved as part of a new animal drug application or abbreviated new animal drug application in ways other than those permitted by regulation.

DATES: Submit either electronic or written comments on Agency guidances at any time.

ADDRESSES: You may submit comments as follows:

Electronic Submissions

Submit electronic comments in the following way:

• Federal eRulemaking Portal: http://www.regulations.gov. Follow the instructions for submitting comments. Comments submitted electronically, including attachments, to http://www.regulations.gov will be posted to the docket unchanged. Because your comment will be made public, you are solely responsible for ensuring that your comment does not include any...
provide this information on the cover sheet and not in the body of your comments and you must identify this information as “confidential.” Any information marked as “confidential” will not be disclosed except in accordance with 21 CFR 10.20 and other applicable disclosure law. For more information about FDA’s posting of comments to public dockets, see 80 FR 56469, September 18, 2015, or access the information at: http://www.fda.gov/regulatoryinformation/dockets/default.htm.

Docket: For access to the docket to read background documents or the electronic and written/paper comments received, go to http://www.regulations.gov and insert the docket number, found in brackets in the heading of this document, into the “Search” box and follow the prompts and/or go to the Division of Dockets Management, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.

Submit written requests for single copies of the guidance to the Policy and Regulations Staff (HFV–6), Center for Veterinary Medicine, Food and Drug Administration, 7519 Standish Pl., Rockville, MD 20855. Send one self-addressed adhesive label to assist that office in processing your requests. See the SUPPLEMENTARY INFORMATION section for electronic access to the guidance document.

FOR FURTHER INFORMATION CONTACT: Dorothy McAdams, Center for Veterinary Medicine, Division of Surveillance (HFV–210), Food and Drug Administration, 7519 Standish Pl., Rockville, MD 20855, 240–402–5763, email: dorothy.mcadams@fda.hhs.gov.

SUPPLEMENTARY INFORMATION:

I. Background

In the Federal Register of September 10, 2015 (80 FR 54568), FDA published the notice of availability for a draft guidance entitled “Distributor Labeling for New Animal Drugs” giving interested persons until November 9, 2015, to comment on the draft guidance. FDA received no comments on the draft guidance. The guidance announced in this notice finalizes the draft guidance dated September 2015.

II. Significance of Guidance

This level 1 guidance is being issued consistent with FDA’s good guidance practices regulation (21 CFR 10.115). The guidance represents the current thinking of FDA on distributor labeling for new animal drugs. It does not establish new rights for any person and is not binding on FDA or the public. You can use an alternative approach if it satisfies the requirements of the applicable statutes and regulations.

III. Paperwork Reduction Act of 1995

This guidance refers to previously approved collections of information found in FDA regulations. These collections of information are subject to review by the Office of Management and Budget (OMB) under the Paperwork Reduction Act of 1995 (44 U.S.C. 3501–3520). The collections of information in 21 CFR 514.80 have been approved under OMB control number 0910–0284.

IV. Electronic Access

Persons with access to the Internet may obtain the guidance at either http://www.fda.gov/AnimalVeterinary/GuidanceComplianceEnforcement/GuidanceforIndustry/default.htm or http://www.regulations.gov.

Dated: April 13, 2016.

Leslie Kux,
Associate Commissioner for Policy.

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DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA–2016–N–1024]

Preparation for International Cooperation on Cosmetics Regulation

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice of meeting.

The Food and Drug Administration (FDA or we) is announcing a public meeting entitled “International Cooperation on Cosmetics Regulation (ICCR)—Preparation for ICCR–10 Meeting.” The purpose of the meeting is to invite public input on various topics pertaining to the regulation of cosmetics. We may use this input to help us prepare for the ICCR–10 meeting that will be held July 12–14, 2016, in Bethesda, MD.

Date and Time: The public meeting will be held on June 15, 2016, from 2 p.m. to 4 p.m.

Location: This meeting will be held at the Food and Drug Administration, Center for Food Safety and Applied Nutrition, 5100 Paint Branch Pkwy., Wiley Auditorium, College Park, MD 20740.

Contact Person: Maria Rossana (Rosemary) Cook, Office of Cosmetics and Colors, Food and Drug Administration, 4300 River Rd., College Park, MD 20740, maria.cook@fda.hhs.gov, or FAX: 301–436–2975.