

that most closely resembles an in-person advisory committee meeting.

Procedure: Interested persons may present data, information, or views, orally or in writing, on issues pending before the committee. Written submissions may be made to the contact person on or before October 20, 2020. Oral presentations from the public will be scheduled on November 9, 2020, between approximately 1 p.m. Eastern Time and 2 p.m. Eastern Time. Those individuals interested in making formal oral presentations should notify the contact person (see **FOR FURTHER INFORMATION CONTACT**). The notification should include a brief statement of the general nature of the evidence or arguments they wish to present, the names and addresses of proposed participants, and an indication of the approximate time requested to make their presentation on or before October 13, 2020. Time allotted for each presentation may be limited. If the number of registrants requesting to speak is greater than can be reasonably accommodated during the scheduled open public hearing session, FDA may conduct a lottery to determine the speakers for the scheduled open public hearing session. The contact person will notify interested persons regarding their request to speak by October 14, 2020.

For press inquiries, please contact the Office of Media Affairs at fdadoma@fda.hhs.gov or 301-796-4540.

FDA welcomes the attendance of the public at its advisory committee meetings and will make every effort to accommodate persons with disabilities. If you require accommodations due to a disability, please contact Artair Mallett at artair.mallett@fda.hhs.gov or 301-796-9638 at least 7 days in advance of the meeting.

FDA is committed to the orderly conduct of its advisory committee meetings. Please visit our website at <https://www.fda.gov/AdvisoryCommittees/AboutAdvisoryCommittees/ucm111462.htm> for procedures on public conduct during advisory committee meetings.

Notice of this meeting is given under the Federal Advisory Committee Act (5 U.S.C. app. 2).

Dated: August 26, 2020.

Lauren K. Roth,

Associate Commissioner for Policy.

[FR Doc. 2020-19482 Filed 9-2-20; 8:45 am]

BILLING CODE 4164-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2020-D-1530]

Control of Nitrosamine Impurities in Human Drugs; Guidance for Industry; Availability

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice of availability.

SUMMARY: The Food and Drug Administration (FDA, the Agency, or we) is announcing the availability of a final guidance for industry, entitled “Control of Nitrosamine Impurities in Human Drugs.” This guidance recommends steps manufacturers of active pharmaceutical ingredients and drug products should take to detect and prevent objectionable levels of nitrosamine impurities in pharmaceutical products. The guidance also describes conditions that may introduce nitrosamine impurities. The recent unexpected finding of nitrosamine impurities, which are probable human carcinogens, in drugs such as angiotensin II receptor blockers, ranitidine, nizatidine, and metformin, has made clear the need for a risk assessment strategy to identify and minimize nitrosamines in any pharmaceutical product at risk for their presence.

DATES: The announcement of the guidance is published in the **Federal Register** on September 3, 2020.

ADDRESSES: You may submit either electronic or written comments on Agency guidances at any time as follows:

Electronic Submissions

Submit electronic comments in the following way:

- *Federal eRulemaking Portal:* <https://www.regulations.gov>. Follow the instructions for submitting comments. Comments submitted electronically, including attachments, to <https://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 confidential information that you or a third party may not wish to be posted, such as medical information, your or anyone else’s Social Security number, or confidential business information, such as a manufacturing process. Please note that if you include your name, contact information, or other information that identifies you in the body of your

comments, that information will be posted on <https://www.regulations.gov>.

- If you want to submit a comment with confidential information that you do not wish to be made available to the public, submit the comment as a written/paper submission and in the manner detailed (see “Written/Paper Submissions” and “Instructions”).

Written/Paper Submissions

Submit written/paper submissions as follows:

- *Mail/Hand delivery/Courier (for written/paper submissions):* Dockets Management Staff (HFA-305), Food and Drug Administration, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852.

- For written/paper comments submitted to the Dockets Management Staff, FDA will post your comment, as well as any attachments, except for information submitted, marked and identified, as confidential, if submitted as detailed in “Instructions.”

Instructions: All submissions received must include the Docket No. FDA-2020-D-1530 for “Control of Nitrosamine Impurities in Human Drugs.” Received comments will be placed in the docket and, except for those submitted as “Confidential Submissions,” publicly viewable at <https://www.regulations.gov> or at the Dockets Management Staff between 9 a.m. and 4 p.m., Monday through Friday, 240-402-7500.

- **Confidential Submissions—**To submit a comment with confidential information that you do not wish to be made publicly available, submit your comments only as a written/paper submission. You should submit two copies total. One copy will include the information you claim to be confidential with a heading or cover note that states “THIS DOCUMENT CONTAINS CONFIDENTIAL INFORMATION.” The Agency will review this copy, including the claimed confidential information, in its consideration of comments. The second copy, which will have the claimed confidential information redacted/blacked out, will be available for public viewing and posted on <https://www.regulations.gov>. Submit both copies to the Dockets Management Staff. If you do not wish your name and contact information to be made publicly available, you can 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: <https://www.govinfo.gov/content/pkg/FR-2015-09-18/pdf/2015-23389.pdf>.

Docket: For access to the docket to read background documents or the electronic and written/paper comments received, go to <https://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 Dockets Management Staff, 5630 Fishers Lane, Rm. 1061, Rockville, MD 20852, 240-402-7500.

You may submit comments on any guidance at any time (see 21 CFR 10.115(g)(5)).

Submit written requests for single copies of the guidance to the Office of Communications, Division of Drug Information, Center for Drug Evaluation and Research, Food and Drug Administration, 10001 New Hampshire Ave., Hillandale Bldg., 4th Floor, Silver Spring, MD 20993-0002. Send two self-addressed adhesive labels to assist that office in processing your requests. See the **SUPPLEMENTARY INFORMATION** section for electronic access to the guidance document.

FOR FURTHER INFORMATION CONTACT: Dongmei Lu, Center for Drug Evaluation and Research, Food and Drug Administration, 10903 New Hampshire Ave., Bldg. 75, Rm. 6649, Silver Spring, MD 20993, 240-402-7966.

SUPPLEMENTARY INFORMATION:

I. Background

We are announcing the availability of a guidance for industry entitled "Control of Nitrosamine Impurities in Human Drugs." We are issuing this guidance consistent with our good guidance practices (GGP) regulation (§ 10.115 (21 CFR 10.115)). We are implementing this guidance without prior public comment because we have determined that prior public participation is not feasible or appropriate (§ 10.115(g)(2)). We made this determination because of the importance of providing timely information to manufacturers regarding risk assessments, testing, and other appropriate actions they should take to reduce and mitigate nitrosamine impurities in active pharmaceutical ingredients (APIs) and drug products. Although this guidance document is immediately in effect, it remains subject to comment in accordance with FDA's GGP regulation (§ 10.115(g)(3)(D)).

Nitrosamines have been classified as probably carcinogenic to humans by the World Health Organization. This guidance recommends steps

manufacturers of APIs and drug products should take to detect and prevent objectionable levels of nitrosamine impurities in pharmaceutical products. The guidance also describes conditions that may introduce nitrosamine impurities.

The recent discovery of nitrosamine impurities in some types of drug products, including angiotensin II receptor blockers, ranitidine, nizatidine, and metformin, led FDA and other international regulators to conduct a detailed analysis of these impurities in affected APIs and drug products.

Recently, preliminary results from FDA stability testing raised concerns that N-Nitrosodimethylamine (NDMA) levels in some ranitidine products stored at room temperature can increase with time to unacceptable levels. Results from other tests FDA conducted suggest that the NDMA levels increase with storage time. On April 1, 2020, FDA requested that all ranitidine products be withdrawn from the U.S. market.

Based on the testing results and the Agency's current understanding of the chemistry, FDA has developed this guidance to provide API and drug product manufacturers information on the potential root causes of nitrosamine formation. It recommends ways API and drug product manufacturers can conduct risk assessments of their products, whether approved, marketed, or with pending applications. The guidance also suggests actions they should take to reduce or prevent the presence of nitrosamines in APIs and drug products.

API and drug product manufacturers should assess the risk of nitrosamine contamination or formation in their drugs. These risk assessments should be conducted in a timely manner. Manufacturers do not need to submit risk assessment documents to the Agency, but they should retain them so that they are available if requested. FDA may request an expedited risk assessment, confirmatory testing, or other regulatory action based on information available to the Agency.

For products at the pre-submission stage, FDA recommends that applicants conduct a risk assessment for nitrosamine impurities in APIs and proposed drug products and conduct confirmatory testing as needed prior to submission of an original application.

However, the risk assessment and submission of any confirmatory testing or changes to the drug master file or application may be submitted in an amendment if they are not available at the time of the original submission filing. For applications that are pending with the Agency, applicants should

conduct the risk assessment expeditiously and inform FDA if confirmatory testing finds nitrosamine levels above the recommended acceptable daily intake (ADI) limit. If a nitrosamine impurity is detected above the limit of quantitation but is within the ADI limit, the applicant should amend the application as appropriate. The Agency will work with the applicant to resolve issues during the review cycle or immediately after approval, and before distribution, if determined to be necessary by the Agency.

This guidance is being issued consistent with FDA's good guidance practices regulation (§ 10.115). The guidance represents the current thinking of FDA on the "Control of Nitrosamine Impurities in Human Drugs." 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.

II. Paperwork Reduction Act of 1995

This guidance refers to previously approved FDA collections of information. 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. 3501-3521). The collections of information in 21 CFR parts 210 and 211 have been approved under OMB control number 0910-0139; the collections of information in 21 CFR part 312 have been approved under OMB control number 0910-0014; the collections of information in 21 CFR part 314 have been approved under OMB control number 0910-0001; the collections of information in 21 CFR part 601 have been approved under OMB control number 0910-0338; the collections of information for the permanent discontinuation or interruption in manufacturing of certain drug and biological products have been approved under OMB control number 0910-0759; the collections of information pertaining to the guidance for industry entitled "Controlled Correspondence Related to Generic Drug Development" have been approved under OMB control number 0910-0797.

III. Electronic Access

Persons with access to the internet may obtain the document at <https://www.fda.gov/RegulatoryInformation/Guidances/default.htm> or <https://www.regulations.gov>. Use the FDA website listed in the previous sentence to find the most current version of the guidance.

Dated: August 28, 2020.
Lowell J. Schiller,
Principal Associate Commissioner for Policy.
 [FR Doc. 2020–19519 Filed 9–2–20; 8:45 am]
BILLING CODE 4164–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

National Institutes of Health

Submission for OMB review; 30-Day Comment Request; Specimen Resource Locator (National Cancer Institute)

AGENCY: National Institutes of Health, HHS.
ACTION: Notice.

SUMMARY: In compliance with the Paperwork Reduction Act of 1995, the National Institutes of Health (NIH) has submitted to the Office of Management and Budget (OMB) a request for review and approval of the information collection listed below.

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

ADDRESSES: Written comments and recommendations for the proposed information collection should be sent within 30 days of publication of this notice to www.reginfo.gov/public/do/PRAMain. Find this particular

information collection by selecting “Currently under 30-day Review—Open for Public Comments” or by using the search function.

FOR FURTHER INFORMATION CONTACT: Joanne Demchok, Program Director, Cancer Diagnosis Program, Division of Cancer Treatment and Diagnosis, 9609 Medical Center Drive, Rockville, MD 20892 or call non-toll-free number 240–276–5959 or Email your request, including your address to: peterjo@mail.nih.gov.

SUPPLEMENTARY INFORMATION: This proposed information collection was previously published in the **Federal Register** on June 18, 2020, page 36871 (Vol. 85, No. 118 FR 36871) 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 National Cancer Institute (NCI), 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.

In compliance with 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 for review and approval of the information collection listed below.

Proposed Collection: Specimen Resource Locator, OMB #0925–0703: Expiration Date 11/30/2020, EXTENSION, National Cancer Institute (NCI), National Institutes of Health (NIH).

Need and Use of Information Collection: The availability of specimens and associated data is critical to increase our knowledge of cancer biology, and to translate important research discoveries to clinical application. The discovery and validation of cancer prevention markers require access, by researchers, to quality clinical biospecimens. In response, to this need, the National Cancer Institute’s (NCI) Cancer Diagnosis Program has developed, and is expanding, a searchable database: Specimen Resource Locator (SRL). The SRL allows scientist in the research community and the NCI to locate specimens needed for their research. The SRL will list all NCI supported repositories and their links. This administrative submission is an on-line form that will collect information to manage and improve a program and its resources for the use of all scientists. This submission does not involve any analysis.

OMB approval is requested for 3 years. There are no costs to respondents other than their time. The total estimated annualized burden hours are 105.

ESTIMATED ANNUALIZED BURDEN HOURS

| Type of respondent | Form name | Number of respondents | Number of responses per respondent | Average burden per response (in hours) | Total burden hour |
|--------------------------|-----------------------|-----------------------|------------------------------------|--|-------------------|
| Private Sector | Initial Request | 70 | 1 | 30/60 | 35 |
| State Government | | 70 | 1 | 30/60 | 35 |
| Federal Government | | 60 | 1 | 30/60 | 30 |
| Private Sector | Annual Update | 20 | 1 | 5/60 | 2 |
| State Government | | 20 | 1 | 5/60 | 2 |
| Federal Government | | 10 | 1 | 5/60 | 1 |
| Total | | | 250 | | 105 |

Dated: August 26, 2020.
Diane Kreinbrink,
Project Clearance Liaison, National Cancer Institute, National Institutes of Health.
 [FR Doc. 2020–19446 Filed 9–2–20; 8:45 am]
BILLING CODE 4140–01–P

DEPARTMENT OF HOMELAND SECURITY

Coast Guard

[Docket No. USCG–2020–0185; OMB Control Number 1625–0102]

Collection of Information Under Review by Office of Management and Budget

AGENCY: Coast Guard, DHS.

ACTION: Thirty-day notice requesting comments.

SUMMARY: In compliance with the Paperwork Reduction Act of 1995 the U.S. Coast Guard is forwarding an Information Collection Request (ICR), abstracted below, to the Office of Management and Budget (OMB), Office of Information and Regulatory Affairs (OIRA), requesting an extension of its approval for the following collection of information: 1625–0102, National Response Resource Inventory; without