NOTIFICATION

Addendum

The following communication, received on 21 December 2020, is being circulated at the request of the Delegation of the United States of America.

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| Requirements for Additional Traceability Records for Certain Foods; Extension of Comment Period |
| The Food and Drug Administration (FDA, the Agency, or we) is extending the comment period for the proposed rule and reopening the comment period for the information collection related to the proposed rule entitled "Requirements for Additional Traceability Records for Certain Foods" that appeared in the Federal Register of 23 September 2020. We are taking this action in response to a request from stakeholders to extend the comment period to allow additional time for interested persons to submit comments on the proposed rule. We also are taking this action to keep the comment period for the information collection provisions associated with the rule consistent with the comment period for the proposed rule.DATES: FDA is extending the comment period on the proposed rule published 23 September 2020 (85 FR 59984). Submit either electronic or written comments on the proposed rule by 22 February 2021. Submit comments on information collection issues under the Paperwork Reduction Act of 1995 (PRA) by 22 February 2021.<https://members.wto.org/crnattachments/2020/SPS/USA/20_7841_00_e.pdf> |
| **This addendum concerns a:** |
| [**X**] Modification of final date for comments |
| [ ] Notification of adoption, publication or entry into force of regulation |
| [ ] Modification of content and/or scope of previously notified draft regulation |
| [ ] Withdrawal of proposed regulation |
| [ ] Change in proposed date of adoption, publication or date of entry into force |
| [ ] Other:  |
| **Comment period: *(If the addendum extends the scope of the previously notified measure in terms of products and/or potentially affected Members, a new deadline for receipt of comments should be provided, normally of at least 60 calendar days. Under other circumstances, such as extension of originally announced final date for comments, the comment period provided in the addendum may vary.)*** |
| [ ] Sixty days from the date of circulation of the addendum to the notification and/or *(dd/mm/yy)*: Electronic comments must be submitted on or before 22 February 2021. The <https://www.regulations.gov> electronic filing system will accept comments until 11:59 p.m. Eastern Time at the end of 22 February 2021. Comments received by mail/hand delivery/courier (for written/paper submissions) will be considered timely if they are postmarked or the delivery service acceptance receipt is on or before that date. |
| **Agency or authority designated to handle comments: [ ]****National Notification Authority, [ ]** **National Enquiry Point. Address, fax number and e-mail address (if available) of other body:** |
| 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>.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-2014-N-0053 for "Requirements for Additional Traceability Records for Certain Foods". Received comments, those filed in a timely manner (see ADDRESSES), 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. |
| **Text(s) available from: [ ]****National Notification Authority, [ ]** **National Enquiry Point. Address, fax number and e-mail address (if available) of other body:** |
| Text can be found in the US Federal Register, Vol. 85, No. 244, page 82393 or on the internet at: <https://www.govinfo.gov/content/pkg/FR-2020-12-18/pdf/2020-27829.pdf>. |

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