NOTIFICATION

Addendum

The following communication, received on 7 June 2024, is being circulated at the request of the Delegation of the United States of America.

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| Laboratory Accreditation for Analyses of Foods; Program Implementation; Determination of Sufficient Laboratory Capacity for Import-Related Food Testing Covered by the Regulation; Notification |
| The Food and Drug Administration (FDA or we) has determined that there is sufficient laboratory capacity in the Laboratory Accreditation for Analyses of Foods (LAAF) program for the import-related food testing covered by the LAAF regulation for mycotoxins. As sufficient capacity is reached for additional analytes covered under the import- related food testing provisions of the LAAF regulation, those specific analytes and compliance dates will be posted on the LAAF Dashboard. Owners and consignees of imported food subject to the LAAF regulation must use a LAAF- accredited laboratory to conduct covered import-related food testing starting on the applicable compliance date, which is six months from the date a specific analyte is listed on a public registry, based on FDA's determination that sufficient laboratory capacity has been achieved for such analyte. FDA has not yet made a capacity determination for the other food testing circumstances covered by the LAAF regulation.  Compliance dates: A LAAF- accredited laboratory must conduct certain import-related food testing covered by the LAAF regulation (21 CFR 1.1107(a)(4), (5)) beginning six months from the date a specific analyte is posted on the LAAF Dashboard at <https://datadashboard.fda.gov/ora/fd/laaf.htm>.  <https://www.federalregister.gov/documents/2024/06/03/2024-12027/laboratory-accreditation-for-analyses-of-foods-program-implementation-determination-of-sufficient>  <https://members.wto.org/crnattachments/2024/SPS/USA/24_03655_00_e.pdf> |
| **This addendum concerns a:** |
| [ ] Modification of final date for comments |
| [**X**] 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)*: Not applicable |
| **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:** |
| For further information contact: Stacie Hammack, Chemist, Food and Feed Laboratory Operations, Office of Regulatory Affairs, Food and Drug Administration, 60 8th St. NE, Atlanta, GA 30309, +(301) 796 5817; [Stacie.Hammack@fda.hhs.gov](mailto:Stacie.Hammack@fda.hhs.gov) |
| **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. 89, No. 107, page 47463 or on the Internet at: <https://www.govinfo.gov/content/pkg/FR-2024-06-03/pdf/2024-12027.pdf> |

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