

6 January 2025

Original: English

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Committee on Sanitary and Phytosanitary Measures

NOTIFICATION

1.	Notifying Member: UNITED STATES OF AMERICA
	If applicable, name of local government involved:
2.	Agency responsible: Food and Drug Administration (FDA)
3.	Products covered (provide tariff item number(s) as specified in national schedules deposited with the WTO; ICS numbers should be provided in addition, where applicable): Mushrooms and truffles, prepared or preserved otherwise than by vinegar or acetic acid (HS code(s): 2003); Food technology (ICS code(s): 67)
4.	Regions or countries likely to be affected, to the extent relevant or practicable:
	[X] All trading partners
	[] Specific regions or countries:
5.	Title of the notified document: Monterey Mushrooms, LLC; Filing of Food Additive Petition; Notification of Petition. Language(s): English. Number of pages: 2
	https://www.federalregister.gov/d/2024-30362 https://members.wto.org/crnattachments/2025/SPS/USA/25_00040_00_e.pdf
6.	Description of content: The Food and Drug Administration (FDA or we) is announcing that we have filed a petition, submitted by Monterey Mushrooms, LLC, proposing that the food additive regulations for vitamin D_2 mushroom powder be amended to provide for an additional method for producing the additive.
	The food additive petition was filed on 11 December 2024.
7.	Objective and rationale: [X] food safety, [] animal health, [] plant protection, [] protect humans from animal/plant pest or disease, [] protect territory from other damage from pests.
8.	Is there a relevant international standard? If so, identify the standard:
	[] Codex Alimentarius Commission (e.g. title or serial number of Codex standard or related text):
	[] World Organization for Animal Health (OIE) (e.g. Terrestrial or Aquatic Animal Health Code, chapter number):
	[] International Plant Protection Convention (e.g. ISPM number):
	[X] None
	Does this proposed regulation conform to the relevant international standard?
	[]Yes []No
	If no, describe, whenever possible, how and why it deviates from the international standard:

- 9. Other relevant documents and language(s) in which these are available:
- 10. Proposed date of adoption (dd/mm/yy): Not applicable
 Proposed date of publication (dd/mm/yy): Not applicable
- 11. Proposed date of entry into force: [] Six months from date of publication, and/or (dd/mm/yy): Not applicable
 - [X] Trade facilitating measure
- 12. Final date for comments: [] Sixty days from the date of circulation of 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 access to the docket to read background documents or 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.

For further information contact: Katie Overbey, Human Foods Program, Food and Drug Administration, 5001 Campus Dr., College Park, MD 20740, +(240) 402 7536.

13. 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 Federal Register, Vol. 86, No. 172, Page 50496 or on the Internet at: https://www.federalregister.gov/d/2024-30362