NOTIFICATION

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| **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):** HS Code(s): 2003; 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:** Monaghan Mushrooms Ireland Unlimited Company; Filing of Food Additive Petition.**Language(s):** English. **Number of pages:** 1<https://www.govinfo.gov/content/pkg/FR-2021-09-09/pdf/2021-19409.pdf><https://members.wto.org/crnattachments/2021/SPS/USA/21_5786_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 Monaghan Mushrooms Ireland Unlimited Company, proposing that the food additive regulations be amended to provide for the safe use of vitamin D2 mushroom powder produced by exposing dried and powdered edible cultivars of Agaricus bisporus to ultraviolet light. The food additive petition was filed on 8 June 2021. |
| **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:** ADDRESSES: 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, Center for Food Safety and Applied Nutrition, 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.govinfo.gov/content/pkg/FR-2021-09-09/pdf/2021-19409.pdf> |