

PET SCHOOLED

The Educated Community

August 8, 2022

Via Electronic Submission

Division of Dockets Management
Food and Drug Administration
Department of Health and Human Services
5630 Fishers Lane, Room 1061 (HFA-305) Rockville, MD 20852

Re: Citizen Petition- Asking FDA-CVM To Publish On The Public FDA FOIA Reading Room Website, *All reports of adverse events and/or reactions dog and/or cats have had to dog and cat food and dog and cat treats for the month of July 2021*, In Compliance With FOIA Law.

To whom it may concern:

The undersigned submits this petition under the Freedom of Information Act ("FOIA"), 5 U.S.C. § 552, and pursuant to 21 C.F.R. §§ 25.30 and 25.34 and 21 C.F.R. § 10.30(b), requesting both the FDA-CVM and FDA FOIA Department comply with FOIA law and post publicly on the FDA FOIA reading room all documents requested through various FOIA requests pertaining to *All reports of adverse events and/or reactions dog and/or cats have had to dog and cat food and dog and cat treats for the month of July 2021*.

A. Action Requested

Petition requests FDA-CVM to comply with FOIA law and post publicly on the FDA FOIA reading room all documents requested through various FOIA requests pertaining to *All reports of adverse events and/or reactions dog and/or cats have had to dog and cat food and dog and cat treats for the month of July 2021*.

B. Statement of Grounds

FDA-CVM is a subagency of the FDA. The FDA is an agency within the Department of Health and Human Services, and is subject to FOIA laws.

The FOIA requires each agency to make available for public inspection, in electronic format, copies of all records that have previously been released under FOIA and "(I) that because of the nature of their subject matter, the agency determines have become or are likely to become the subject of subsequent requests for substantially the same records; or (II) that have been requested 3 or more times." 5 U.S.C. § 552(a)(2)(D).

1. All reports of adverse events and/or reactions dog and/or cats have had to dog and cat food and dog and cat treats [for the month of July 2021.]

a. Please include the type of animal, the reaction, the brand name and the product name of the food item to which the dog or cat reacted, the symptoms displayed and the outcome, if available.

b. Please include whether any of the reports of adverse events triggered any sort of follow up by the FDA.

The following information has been requested through FOIA “3 or more times” and FDA is now required to comply with 5 U.S.C. § 552(a)(2)(D) and publish these records on the FDA FOIA reading room.

This citizen petition specifically requests FDA to comply with law 5 U.S.C. § 552(a)(2)(D), and publish on the public FDA FOIA reading room website, *All reports of adverse events and/or reactions dog and/or cats have had to dog and cat food and dog and cat treats for the month of July 2021.*

C. Environmental Impact

A claim for categorical exclusion of the requirement for submission of an environmental assessment or environmental impact statement is made pursuant to 21 C.F.R. §§ 25.30 and 25.34.

D. Economic Impact

In accordance with 21 C.F.R. § 10.30(b), economic impact information is to be submitted only when requested by FDA following review of the petition. The undersigned will promptly provide this information if requested.

E. Certification

The undersigned certifies, that, to the best knowledge and belief of the undersigned, this petition includes all information and views on which the petition relies, and that it includes representative data and information known to the petitioner which are unfavorable to the petition.

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