document are, therefore, $83,000.
Respondents are already required to disclose the quantitative amount and the percentage of the daily value of a dietary ingredient on a per serving basis as part of the nutrition information for dietary supplements. Respondents may also provide such information on a per unit basis. The information provided for under the proposed rule would be generated by simple extrapolation from that information.

Margaret M. Dotzel,
Associate Commissioner for Policy.

[FR Doc. 01–28105 Filed 11–8–01; 8:45 am]

BILLING CODE 4160–01–S

DEPARTMENT OF HEALTH AND
HUMAN SERVICES

Food and Drug Administration

[Docket No. 01F–0484]

Anitox Corp.; Filing of Food Additive Petition (Animal Use); Formaldehyde

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA) is announcing that Anitox Corp. has filed a petition proposing that the food additive regulations be amended to allow a variable usage rate of 2.0 to 5.4 pounds (lb) of formaldehyde (CAS No. 50–00–0; 37 percent aqueous solution) per ton of animal feeds for feed ingredients.

DATES: Submit written or electronic comments on the petitioner’s environmental assessment by January 23, 2002. Two copies of any comments are to be submitted, except that individuals may submit one copy. Comments are to be identified with the docket number found in brackets in the heading of this document. Received comments may be seen in the Dockets Management Branch (address above) for public review and comment.

Interested persons may submit to the Dockets Management Branch (address above) written or electronic comments to January 23, 2002. Two copies of any comments are to be submitted, except that individuals may submit one copy. Comments are to be identified with the docket number found in brackets in the heading of this document. Received comments may be seen in the Dockets Management Branch (address above) for public review and comment. A regulation, the potential environmental impact of this action is being reviewed. To encourage public participation consistent with regulations issued under the National Environmental Policy Act (40 CFR 1501.4(b)), the agency is placing the environmental information submitted with the petition that is the subject of this notice on public display at the Dockets Management Branch (address above) for public review and comment.

The potential environmental impact of this action is being reviewed. To encourage public participation consistent with regulations issued under the National Environmental Policy Act (40 CFR 1501.4(b)), the agency is placing the environmental information submitted with the petition that is the subject of this notice on public display at the Dockets Management Branch (address above) for public review and comment.

The purpose of this guidance is to identify for the industry fumonisin levels that FDA considers adequate to protect human and animal health and that are achievable in human foods and animal feeds with the use of good agricultural and good manufacturing practices. FDA considers this guidance to be a prudent public health measure during the development of a long-term risk management policy and program by the agency for the control of fumonisins in human foods and animal feeds. The agency is also announcing the availability of the final supporting documents entitled “Background Paper in Support of Fumonisin Levels in Corn and Corn Products Intended for Human Consumption,” and “Background Paper in Support of Fumonisin Levels in Animal Feed.”

DATES: Submit written or electronic comments concerning the final guidance and the final supporting documents at any time.

ADDRESSES: Submit written comments on the final guidance to the Dockets Management Branch (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. Submit electronic comments to http://www.fda.gov/dockets/ecomments.

FOR FURTHER INFORMATION CONTACT: Henry E. Ekperigin, Center for Veterinary Medicine (HFV 222), Food and Drug Administration, 7500 Standish Pl., Rockville, MD 20855, 301–827–0174.

SUPPLEMENTARY INFORMATION: Under the Federal Food, Drug, and Cosmetics Act (sec. 409(b)(5) (21 U.S.C. 348(b)(5))), notice is given that a food additive petition (FAP 2245) has been filed by Anitox Corp., 1055 Progress Circle, P.O. Box 490310, Lawrenceville, GA 30043. The petition proposes to amend the food additive regulations in part 573—Food Additives Permitted in Feed and Drinking Water of Animals (21 CFR part 573) to allow a variable usage rate of 2.0 to 5.4 lb of formaldehyde (CAS No. 50–00–0; 37 percent aqueous solution) per ton of animal feeds for feed ingredients.

The potential environmental impact of this action is being reviewed. To encourage public participation consistent with regulations issued under the National Environmental Policy Act (40 CFR 1501.4(b)), the agency is placing the environmental information submitted with the petition that is the subject of this notice on public display at the Dockets Management Branch (address above) for public review and comment.

ADDRESSES: Submit written comments to the Dockets Management Branch (address above) or Communications Staff (HFV–12), Center for Veterinary Medicine (CVM), 7500 Standish Pl., Rockville, MD 20855, 301–594–1755. Send one self-adhesive address label to assist that office in processing your request. See the SUPPLEMENTARY INFORMATION section for electronic access to these documents.

For further information contact: Henry Kim, Center for Food Safety and Applied Nutrition (CFSAN) (address below), or Communications Staff (HFV–12), Center for Veterinary Medicine (CVM), 7500 Standish Pl., Rockville, MD 20855, 301–594–1755. A regulation, the potential environmental impact of this action is being reviewed. To encourage public participation consistent with regulations issued under the National Environmental Policy Act (40 CFR 1501.4(b)), the agency is placing the environmental information submitted with the petition that is the subject of this notice on public display at the Dockets Management Branch (address above) for public review and comment.

The purpose of this guidance is to identify for the industry fumonisin levels that FDA considers adequate to protect human and animal health and that are achievable in human foods and animal feeds with the use of good agricultural and good manufacturing practices. FDA considers this guidance to be a prudent public health measure during the development of a long-term risk management policy and program by the agency for the control of fumonisins in human foods and animal feeds. The agency is also announcing the availability of the final supporting documents entitled “Background Paper in Support of Fumonisin Levels in Corn and Corn Products Intended for Human Consumption,” and “Background Paper in Support of Fumonisin Levels in Animal Feed.”

DATES: Submit written or electronic comments concerning the final guidance and the final supporting documents at any time.

ADDRESSES: Submit written comments on the final guidance to the Dockets Management Branch (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. Submit electronic comments to http://www.fda.gov/dockets/ecomments.

FOR FURTHER INFORMATION CONTACT: Henry Kim, Center for Food Safety and Applied Nutrition (CFSAN) (address below), or Communications Staff (HFV–12), Center for Veterinary Medicine (CVM), 7500 Standish Pl., Rockville, MD 20855, 301–594–1755. Send one self-adhesive address label to assist that office in processing your request. See the SUPPLEMENTARY INFORMATION section for electronic access to these documents.

SUMMARY: The Food and Drug Administration (FDA) is announcing the availability of a final guidance document entitled “Guidance for Industry: Fumonisin Levels in Human Foods and Animal Feeds.” The purpose of this guidance is to identify for the industry fumonisin levels that FDA considers adequate to protect human and animal health and that are achievable in human foods and animal feeds with the use of good agricultural and good manufacturing practices. FDA considers this guidance to be a prudent public health measure during the development of a long-term risk management policy and program by the agency for the control of fumonisins in human foods and animal feeds. The agency is also announcing the availability of the final supporting documents entitled “Background Paper in Support of Fumonisin Levels in Corn and Corn Products Intended for Human Consumption,” and “Background Paper in Support of Fumonisin Levels in Animal Feed.”

I. Background

On June 6, 2000, FDA issued a draft guidance document that presented recommended levels of fumonisins in corn used for production of human
foods and animal feeds. The purpose of the draft guidance was to identify for the industry fumonisin levels that FDA considers adequate to protect human and animal health and that are achievable in human foods and animal feeds with the use of good agricultural and good manufacturing practices. In the Federal Register notice of June 6, 2000 (65 FR 35945) announcing the availability of the draft guidance document, FDA provided a 60-day period for comment on the recommended fumonisin levels. FDA received 12 comments in response to the June 6, 2000, draft guidance. The comments represented the views of seven trade associations representing manufacturers of dry and wet milled corn products, popcorn, snack foods, processed grain and feed products, food and other consumer products, and pet foods; a snack food company; a dry miller of corn; a food and food ingredient company; a State health department; and a life science society. The majority of the comments stated that they supported the recommended fumonisin levels in corn used for production of human foods and animal feeds. A number of comments suggested changes or modification to the various recommended fumonisin levels. FDA has considered the submitted comments and has revised the supporting documents as appropriate.

II. Conclusion

The agency is adopting the recommended fumonisin levels in human foods and animal feeds as presented in the draft guidance document. The majority of the comments that the agency received supported the recommended fumonisin levels. Further, after considering carefully the comments that the agency received that suggested modification or opposition to aspects of the recommended levels in the draft guidance, the agency has determined that no changes are warranted. The final supporting documents explain the agency’s reasoning in selecting the recommended levels.

FDA considers the final guidance to be a prudent public health measure during the development of a long-term risk management policy and program by the agency for the control of fumonisins in human foods and animal feeds. Based on information obtained from future national and international workshops on the risk from exposure to fumonisins, FDA will consider whether to establish tolerances, regulatory limits, or action levels, as appropriate, for fumonisins in human foods and animal feeds, respectively, under 21 CFR part 109—Unavoidable Contaminants in Food for Human Consumption and Food-Packaging Material and under 21 CFR part 509—Unavoidable Contaminants in Animal Food and Food-Packaging Material.

The final guidance document is being issued as a level 1 guidance, consistent with FDA’s good guidance practices regulation (21 CFR 10.115). The final guidance represents the agency’s current thinking on the control of fumonisins in human foods and animal feeds as a prudent public health measure. It does not create or confer any rights for or on any person and does not operate to bind FDA or the public. An alternative approach may be used if such approach satisfies the requirements of the applicable statute and regulations.

III. Comments

Interested persons may submit to the Dockets Management Branch (address above) written or electronic comments regarding the final guidance at any time. Two copies of any comments are to be submitted, except that individuals may submit one copy. Comments are to be identified with the docket number found in brackets in the heading of this document. The final guidance, the final supporting documents entitled “Background Paper in Support of Fumonisin Levels in Corn and Corn Products Intended for Human Consumption,” and “Background Paper in Support of Fumonisin Levels in Animal Feed,” and received comments are available in the Dockets Management Branch between 9 a.m. and 4 p.m. Monday through Friday.

IV. Electronic Access

The final guidance, as well as the final supporting documents entitled “Background Paper in Support of Fumonisin Levels in Corn and Corn Products Intended for Human Consumption,” and “Background Paper in Support of Fumonisin Levels in Animal Feed,” may be accessed from the home pages of CFSAN and CVM on the Internet at http://www.cfsan.fda.gov and http://www.fda.gov/cvm, respectively.

Dated: November 1, 2001.

Margaret M. Dotzel,
Associate Commissioner for Policy.

[FR Doc. 01–28104 Filed 11–8–01; 8:45 am]

BILLING CODE 4160–01–S

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Health Resources and Services Administration

Advisory Committee; Notice of Meeting

Pursuant to section 10(d) of the Federal Advisory Committee Act, as amended (5 U.S.C. Appendix 2), notice is hereby given of the following Health Professions and Nurse Education Special Emphasis Panel Meetings.

Name: Minority Faculty Fellowship Program Peer Review Group.


Place: Holiday Inn Silver Spring, 8777 Georgia Avenue, Silver Spring, MD 20910.

Open on: November 27, 2001, 8:00 a.m. to 10:00 a.m.

Closed on: November 27, 2001, 10:00 a.m. to 6:00 p.m.; November 28–30, 2001, 8:00 a.m. to 6:00 p.m.

Name: Nursing Faculty Development in Geriatrics Program Peer Review Group.

Date and Time: January 14–17, 2002.

Place: Holiday Inn Silver Spring, 8777 Georgia Avenue, Silver Spring, MD 20910.

Open on: January 14, 2002, 8:00 a.m. to 10:00 a.m.

Closed on: January 14, 2002, 10:00 a.m. to 6:00 p.m.; January 15–17, 2002, 8:00 a.m. to 6:00 p.m.

Name: Geriatric Nursing Knowledge and Experience in Long Term Care Facilities for Baccalaureate Nursing Students Program Peer Review Group.

Date and Time: January 14–17, 2002.

Place: Holiday Inn Silver Spring, 8777 Georgia Avenue, Silver Spring, MD 20910.

Open on: January 14, 2002, 8:00 a.m. to 10:00 a.m.

Closed on: January 14, 2002, 10:00 a.m. to 6:00 p.m.; January 15–17, 2002, 8:00 a.m. to 6:00 p.m.

Name: Centers of Excellence Program Peer Review Group.

Date and Time: January 22–25, 2002.

Place: Holiday Inn Silver Spring, 8777 Georgia Avenue, Silver Spring, MD 20910.

Open on: January 22, 2002, 8:00 a.m. to 10:00 a.m.

Closed on: January 22, 2002, 10:00 a.m. to 6:00 p.m.; January 23–25, 2002, 8:00 a.m. to 6:00 p.m.

Name: Nursing Workforce Diversity Program Peer Review Group.

Date and Time: January 26–31, 2002.

Place: Holiday Inn Silver Spring, 8777 Georgia Avenue, Silver Spring, MD 20910.

Open on: January 28, 2002, 8:00 a.m. to 10:00 a.m.

Closed on: January 28, 2002, 10:00 a.m. to 6:00 p.m.; January 29–31, 2002, 8:00 a.m. to 6:00 p.m.

Name: Model State-Supported Area Health Education Centers Program Peer Review Group.


Place: Holiday Inn Silver Spring, 8777 Georgia Avenue, Silver Spring, MD 20910.

Open on: February 4, 2002, 8:00 a.m. to 10:00 a.m.