

Proposed Rules

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This section of the FEDERAL REGISTER contains notices to the public of the proposed issuance of rules and regulations. The purpose of these notices is to give interested persons an opportunity to participate in the rule making prior to the adoption of the final rules.

DEPARTMENT OF AGRICULTURE

Animal and Plant Health Inspection Service

9 CFR Part 77

[Docket No. 99-038-4]

Tuberculosis in Cattle, Bison, Goats, and Captive Cervids; State and Zone Designations

AGENCY: Animal and Plant Health Inspection Service, USDA.

ACTION: Proposed rule; reopening and extension of comment period; notice of public hearings.

SUMMARY: We are reopening and extending the comment period for our proposed rule that would amend the bovine tuberculosis regulations to establish new levels of tuberculosis risk classifications to be applied to States and zones within States. The proposed rule would also classify States and zones according to their tuberculosis risk with regard to captive cervids. Additionally, it would amend the regulations to specify that the regulations apply to goats as well as to cattle, bison, and captive cervids, and would increase the amount of testing that must be done before certain cattle, bison, and goats may be moved interstate. This action will allow interested persons additional time to prepare and submit comments. We are also advising the public that we are hosting two public hearings on the proposed rule.

DATES: We invite you to comment on Docket No. 99-038-1. We will consider all comments that we receive by June 16, 2000. We will also consider comments made at public hearings to be held in Albuquerque, NM, on June 14, 2000, from 8 a.m. to noon, and in Lansing, MI, on June 15, 2000, from 6 p.m. to 9 p.m.

ADDRESSES: Please send your comment and three copies to: Docket No. 99-038-1, Regulatory Analysis and Development, PPD, APHIS, Suite 3C03,

4700 River Road, Unit 118, Riverdale, MD 20737-1238.

Please state that your comment refers to Docket No. 99-038-1.

You may read any comments that we receive on this docket in our reading room. The reading room is located in room 1141 of the USDA South Building, 14th Street and Independence Avenue, SW., Washington, DC. Normal reading room hours are 8 a.m. to 4:30 p.m., Monday through Friday, except holidays. To be sure someone is there to help you, please call (202) 690-2817 before coming.

APHIS documents published in the **Federal Register**, and related information, including the names of organizations and individuals who have commented on APHIS dockets, are available on the Internet at <http://www.aphis.usda.gov/ppd/rad/webrepor.html>.

The public hearings will be held at the following locations:

- (1) Albuquerque, NM: Racing Commission Conference Room, Lower Level, 300 San Mateo Boulevard NE, Albuquerque, NM.
- (2) Lansing, MI: The Forum, 1st Floor, Michigan Library and Historical Center, 717 West Allegan Street, Lansing, MI.

FOR FURTHER INFORMATION CONTACT: Dr. Joseph Van Tiem, Senior Staff Veterinarian, VS, APHIS, USDA, 4700 River Road Unit 43, Riverdale, MD 20737-1231; (301) 734-7716.

SUPPLEMENTARY INFORMATION: On March 7, 2000, we published in the **Federal Register** (65 FR 11912-11940, Docket No. 99-038-1) a proposal to amend the bovine tuberculosis regulations, contained in 9 CFR part 77. We proposed to: (1) Establish several new levels of tuberculosis risk classifications to be applied to States and zones within States; (2) classify States and zones according to their tuberculosis risk with regard to captive cervids; (3) apply the regulations to goats as well as to cattle, bison, and captive cervids; and (4) increase the amount of testing required for the interstate movement of certain cattle, bison, and goats.

Comments on the proposed rule were required to be received on or before April 21, 2000. On March 24, 2000, we published in the **Federal Register** (65 FR 15877-15878, Docket No. 99-038-2) a correction to Docket No. 99-038-1. Comments on the proposed rule as

corrected were required to be received on or before April 21, 2000.

In response to requests from commenters that we extend the comment period on Docket No. 99-038-1, we published a notice in the **Federal Register** on May 1, 2000 (65 FR 25292, Docket No. 99-038-3), that we were reopening and extending the comment period until May 8, 2000.

During the extended comment period, a number of commenters requested that we further extend the comment period to allow additional time for members of the public to review the proposed rule and to submit comments. In response to these requests, we are reopening and extending the comment period on Docket No. 99-038-1 until June 16, 2000. This action will allow interested persons additional time to prepare and submit comments.

Public Hearings

We are advising the public that we are hosting two public hearings on Docket No. 99-038-1. The first public hearing will be held in Albuquerque, NM, on Wednesday, June 14, 2000, in the Racing Commission Conference Room, Lower Level, 300 San Mateo Boulevard NE. The second public hearing will be held in Lansing, MI, on Thursday, June 15, 2000, at The Forum, 1st Floor, Michigan Library and Historical Center, 717 West Allegan Street.

A representative of the Animal and Plant Health Inspection Service (APHIS) will preside at the public hearings. Any interested persons may appear and be heard in person, by an attorney, or by another representative. Written statements may be submitted and will be made part of the hearing record. Persons who wish to speak at either of the public hearings will be asked to sign in with their name and organization to establish a record for the hearing. We ask that anyone who reads a statement provide two copies to the presiding officer at the hearing.

The public hearing in Albuquerque will begin at 8 a.m. and is scheduled to end at noon, local time. The public hearing in Lansing will begin at 6 p.m. and is scheduled to end at 9 p.m., local time. However, the hearings may be terminated at any time if all persons desiring to speak have been heard.

If the number of speakers at a hearing warrants it, the presiding officer may limit the time for each presentation so

that everyone wishing to speak has the opportunity.

The purpose of the hearings is to give interested persons an opportunity for oral presentation of data, views, and arguments. Questions about the content of the proposed rule may be part of the commenters' oral presentations. However, neither the presiding officer nor any other representative of APHIS will respond to comments at the hearing, except to clarify or explain provisions of the proposed rule.

Authority: 21 U.S.C. 111, 114, 114a, 115–117, 120, 121, 134b, and 134f; 7 CFR 2.22, 2.80, and 371.2(d).

Done in Washington, DC, this 25th day of May 2000.

Bobby R. Acord,

Acting Administrator, Animal and Plant Health Inspection Service.

[FR Doc. 00–13589 Filed 5–30–00; 8:45 am]

BILLING CODE 3410–34–P

DEPARTMENT OF AGRICULTURE

Animal and Plant Health Inspection Service

9 CFR Part 112

[Docket No. 96–034–2]

Viruses, Serums, Toxins, and Analogous Products; Packaging and Labeling

AGENCY: Animal and Plant Health Inspection Service, USDA.

ACTION: Proposed rule; withdrawal.

SUMMARY: We are withdrawing a proposed rule to amend the regulations regarding the packaging and labeling of veterinary biological products. The proposed rule would have required the Animal and Plant Health Inspection Service product code number as well as an appropriate consumer contact telephone number to appear on labeling. In addition, the proposed rule would have clarified label requirements with respect to overshadowing the true name of the product and requirements for products shipped to a foreign country. The proposed rule also contained label requirements concerning the minimum age for product administration and the potential for maternal antibody interference. We are withdrawing the proposed rule due to the comments we received following its publication.

FOR FURTHER INFORMATION CONTACT: Dr. Albert P. Morgan, Chief Staff Officer, Operational Support Section, Center for Veterinary Biologics, Licensing and Policy Development, APHIS, 4700 River Road Unit 148, Riverdale, MD 20737–1231; (301) 734–8245.

SUPPLEMENTARY INFORMATION:

Background

The regulations in 9 CFR part 112 set forth packaging and labeling requirements for veterinary biological products. On March 18, 1999, we published in the **Federal Register** (64 FR 13365–13368, Docket No. 96–034–1) a proposed rule to amend the regulations. First, we proposed to require labels for veterinary biological products to include the Animal and Plant Health Inspection Service (APHIS) product code number and a consumer contact telephone number. Second, we proposed to require labels for veterinary biological products to bear the true name of the product in a prominent fashion and more prominently than the trade name. Third, we proposed to amend the requirements for labels for exported products to state that labels that do not conform to the regulations may be used with an exported product if the labels do not contain false or misleading information and are acceptable to the appropriate regulatory officials of the foreign country to which the products are exported. We proposed that verification of foreign regulatory acceptance of the labels could be supplied to APHIS through the submission of a label mounting prepared as described in § 112.5(d)(2) that bears a stamp or other mark of approval of the appropriate foreign regulatory agency. Finally, we proposed to require labels for veterinary biological products, as described in the proposed rule, to consider the potential for maternal antibody interference with product efficacy and to specify a minimum age for product administration that is consistent with the efficacy and safety data developed for the product.

We solicited comments concerning our proposal for 60 days ending May 17, 1999. We received 11 comments by that date. The comments were from licensed veterinary biologics manufacturers, a national trade association representing U.S. manufacturers of animal health products, an organization representing veterinarians, and a university. Most of the commenters expressed concerns and opposition regarding certain provisions of the proposed rule, including concerns regarding the economic effects of the proposed provisions on veterinary biologics manufacturers and the estimated burden for information collection that was provided in the Paperwork Reduction Act section of the proposed rule.

After considering all of the comments we received, we have concluded that we must reevaluate the provisions of the

proposed rule. Therefore, we are withdrawing the March 18, 1999, proposed rule referenced above. The concerns and recommendations of all of the commenters will be considered if any new proposed regulations regarding the packaging and labeling of veterinary biological products are developed.

Authority: 21 U.S.C. 151–159; 7 CFR 2.22, 2.80, and 371.2(d).

Done in Washington, DC, this 24th day of May 2000.

Bobby R. Acord,

Acting Administrator, Animal and Plant Health Inspection Service.

[FR Doc. 00–13549 Filed 5–30–00; 8:45am]

BILLING CODE 3410–34–U

NUCLEAR REGULATORY COMMISSION

10 CFR Part 50

[Docket No. PRM–50–71]

Nuclear Energy Institute; Receipt of Petition for Rulemaking

AGENCY: Nuclear Regulatory Commission.

ACTION: Petition for rulemaking; Notice of receipt.

SUMMARY: The Nuclear Regulatory Commission (NRC) has received and requests public comment on a petition for rulemaking filed by the Nuclear Energy Institute. The petition was docketed on April 12, 2000, and has been assigned Docket No. PRM–50–71. The petitioner requests that the NRC amend its regulations to allow nuclear power plant licensees to use zirconium-based cladding materials other than zircaloy or ZIRLO, provided the cladding materials meet the requirements for fuel cladding performance and have received approval by the NRC staff. The petitioner believes the proposed amendment would improve the efficiency of the regulatory process by eliminating the need for individual licensees to obtain exemptions to use advanced cladding materials which have already been approved by the NRC.

DATES: Submit comments by August 14, 2000. Comments received after this date will be considered if it is practical to do so, but the Commission is able to assure consideration only for comments received on or before this date.

ADDRESSES: Mail comments to: Secretary, U.S. Nuclear Regulatory Commission, Washington, DC 20555–0001, Attention: Rulemakings and Adjudications Staff.