

(c) \* \* \*

(5) \* \* \* For purposes of § 416.2090 of this part, which discusses the rules for limitation on fiscal liability of States (hold harmless), these retroactive adjustments are State expenditures when made and shall be counted as a State expenditure in the fiscal year in which the adjustments are made.

\* \* \* \* \*

**PART 422—ORGANIZATION AND PROCEDURES**

**Subpart B—[Amended]**

1. The authority citation for subpart B of part 422 continues to read as follows:

**Authority:** Secs. 205, 232, 702(a)(5), 1131, and 1143 of the Social Security Act (42 U.S.C. 405, 432, 902(a)(5), 1320b-1, and 1320b-13).

2. Section 422.125 is amended by revising paragraph (c) to read as follows:

**§ 422.125 Statements of earnings; resolving earnings discrepancies.**

\* \* \* \* \*

(c) *Detailed earnings statements.* A more detailed earnings statement will be furnished upon request, generally without charge, where the request is program related under § 402.170 of this part. If the request for a more detailed statement is not program related under § 402.170 of this part, a charge will be imposed according to the guidelines set out in § 402.175 of this part.

\* \* \* \* \*

3. Section 422.130 is amended by revising the third sentence of paragraph (a) to read as follows:

**§ 422.130 Claim procedure.**

(a) \* \* \* See § 404.614 of this chapter for offices at which applications may be filed.

\* \* \* \* \*

4. Section 422.135 is amended by revising the last sentence of paragraph (a) to read as follows:

**§ 422.135 Reports by beneficiaries.**

(a) \* \* \* (See §§ 404.415 *et seq.* and 404.1571 of this chapter.)

\* \* \* \* \*

[FR Doc. 00-7639 Filed 3-29-00; 8:45 am]

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**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Food and Drug Administration**

**21 CFR Part 177**

[Docket No. 97F-0157]

**Indirect Food Additives: Polymers**

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Final rule.

**SUMMARY:** The Food and Drug Administration (FDA) is amending the food additive regulations to provide for the safe use of 2-propenoic acid, polymer with 2-ethyl-2-(((1-oxo-2-propenyl)oxy)methyl)-1,3-propanediyl di-2-propenoate and sodium 2-propenoate (CAS Reg. No. 76774-25-9) as a fluid absorbent material intended for use in contact with food. This action responds to a petition filed by Japan Vilene Co., Ltd.

**DATES:** This rule is effective March 30, 2000. Submit written objections and requests for a hearing by May 1, 2000.

**ADDRESSES:** Submit written objections to the Dockets Management Branch (HFA-305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852.

**FOR FURTHER INFORMATION CONTACT:** Andrew J. Zajac, Center for Food Safety and Applied Nutrition (HFS-215), Food and Drug Administration, 200 C St. SW., Washington, DC 20204, 202-418-3095.

**SUPPLEMENTARY INFORMATION:** In a notice published in the **Federal Register** of April 22, 1997 (62 FR 19580), FDA announced that a food additive petition (FAP 7B4537) had been filed by Japan Vilene Co., Ltd., c/o Center for Regulatory Services, 2347 Paddock Lane, Reston, VA 20191 (current 5200 Wolf Run Shoals Rd., Woodbridge, VA 22192). The petition proposed to amend the food additive regulations to provide for the safe use of 2-propenoic acid, polymer with 2-ethyl-2-(((1-oxo-2-propenyl)oxy)methyl)-1,3-propanediyl di-2-propenoate and sodium 2-propenoate (CAS Reg. No. 76774-25-9) as a fluid absorbent material intended for use in contact with food.

FDA has evaluated data in the petition and other relevant material. Based on this information, the agency concludes that: (1) The proposed use of the additive is safe, (2) the additive will achieve its intended technical effect, and therefore, (3) the regulations in 21 CFR 177.1211 should be amended as set forth below in this document.

In accordance with § 171.1(h) (21 CFR 171.1(h)), the petition and the

documents that FDA considered and relied upon in reaching its decision to approve the petition are available for inspection at the Center for Food Safety and Applied Nutrition by appointment with the information contact person listed above. As provided in § 171.1(h), the agency will delete from the documents any materials that are not available for public disclosure before making the documents available for inspection.

The agency has carefully considered the potential environmental effects of this action. FDA has concluded that the action will not have a significant impact on the human environment, and that an environmental impact statement is not required. The agency's finding of no significant impact and the evidence supporting that finding, contained in an environmental assessment, may be seen in the Dockets Management Branch (address above) between 9 a.m. and 4 p.m., Monday through Friday.

This final rule contains no collections of information. Therefore, clearance by the Office of Management and Budget under the Paperwork Reduction Act of 1995 is not required.

Any person who will be adversely affected by this regulation may at any time file with the Dockets Management Branch (address above) written objections by May 1, 2000. Each objection shall be separately numbered, and each numbered objection shall specify with particularity the provisions of the regulation to which objection is made and the grounds for the objection. Each numbered objection on which a hearing is requested shall specifically so state. Failure to request a hearing for any particular objection shall constitute a waiver of the right to a hearing on that objection. Each numbered objection for which a hearing is requested shall include a detailed description and analysis of the specific factual information intended to be presented in support of the objection in the event that a hearing is held. Failure to include such a description and analysis for any particular objection shall constitute a waiver of the right to a hearing on the objection. Three copies of all documents are to be submitted and are to be identified with the docket number found in brackets in the heading of this document. Any objections received in response to the regulation may be seen in the Dockets Management Branch between 9 a.m. and 4 p.m., Monday through Friday.

**List of Subjects in 21 CFR Part 177**

Food additives, Food packaging.

Therefore, under the Federal Food, Drug, and Cosmetic Act and under

authority delegated to the Commissioner of Food and Drugs and redelegated to the Director, Center for Food Safety and Applied Nutrition, 21 CFR part 177 is amended as follows:

**PART 177—INDIRECT FOOD ADDITIVES: POLYMERS**

1. The authority citation for 21 CFR part 177 continues to read as follows:

**Authority:** 21 U.S.C. 321, 342, 348, 379e.

2. Section 177.1211 is amended by revising paragraphs (a), (d), and the last sentence in paragraph (c) to read as follows:

**§ 177.1211 Cross-linked polyacrylate copolymers.**

\* \* \* \* \*

(a) *Identity.* For the purpose of this section, the cross-linked polyacrylate copolymers consist of:

(1) The grafted copolymer of cross-linked sodium polyacrylate identified as 2-propenoic acid, polymers with *N,N*-di-2-propenyl-2-propen-1-amine and hydrolyzed polyvinyl acetate, sodium salts, graft (CAS Reg. No. 166164-74-5); or

(2) 2-propenoic acid, polymer with 2-ethyl-2-(((1-oxo-2-propenyl)oxy)methyl)-1,3-propanediyl di-2-propenoate and sodium 2-propenoate (CAS Reg. No. 76774-25-9).

\* \* \* \* \*

(c) *Extractive limitations.* \* \* \* The solvent used shall be at least 60 milliliters aqueous sodium chloride solution per gram of copolymer.

(d) *Conditions of use.* The copolymers identified in paragraph (a)(1) of this section are limited to use as a fluid absorbent in food-contact materials used in the packaging of frozen or refrigerated poultry. The copolymers identified in paragraph (a)(2) of this section are limited to use as a fluid absorbent in food-contact materials used in the packaging of frozen or refrigerated meat and poultry.

Dated: March 20, 2000.

**L. Robert Lake,**

*Director of Regulations and Policy, Center for Food Safety and Applied Nutrition.*

[FR Doc. 00-7930 Filed 3-29-00; 8:45 am]

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**DEPARTMENT OF HEALTH AND HUMAN SERVICES**

**Food and Drug Administration**

**21 CFR Part 524**

**Ophthalmic and Topical Dosage Form New Animal Drugs; Triamcinolone Acetonide Cream**

**AGENCY:** Food and Drug Administration, HHS.

**ACTION:** Final rule.

**SUMMARY:** The Food and Drug Administration (FDA) is amending the animal drug regulations to reflect approval of an abbreviated new animal drug application (ANADA) filed by Med-Pharmex, Inc. The ANADA provides for veterinary prescription use of triamcinolone acetonide cream on dogs for topical treatment of allergic dermatitis and summer eczema.

**DATES:** This rule is effective March 30, 2000.

**FOR FURTHER INFORMATION CONTACT:** Lonnie W. Luther, Center for Veterinary Medicine (HFV-102), Food and Drug Administration, 7500 Standish Pl., Rockville, MD 20855, 301-827-0209.

**SUPPLEMENTARY INFORMATION:** Med-Pharmex, Inc., 2727 Thompson Creek Rd., Pomona, CA 91767-1861, filed ANADA 200-275 that provides for veterinary prescription use of triamcinolone acetonide cream on dogs for topical treatment of allergic dermatitis and summer eczema. Med-Pharmex's ANADA 200-275 MEDALOG cream is approved as a generic copy of Fort Dodge Animal Health's NADA 46-146 VETALOG® cream. ANADA 200-275 is approved as of February 4, 2000, and 21 CFR 524.2481(b) is amended to reflect the approval. The basis for approval is discussed in the freedom of information summary.

In accordance with the freedom of information provisions of 21 CFR part 20 and 514.11(e)(2)(ii), a summary of safety and effectiveness data and information submitted to support approval of this application may be seen in the Dockets Management Branch (HFA-305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852, between 9 a.m. and 4 p.m., Monday through Friday.

The agency has determined under 21 CFR 25.33(a)(1) that this action is of a type that does not individually or cumulatively have a significant effect on the human environment. Therefore, neither an environmental assessment nor an environmental impact statement is required.

This rule does not meet the definition of "rule" in 5 U.S.C. 804(3)(A) because it is a rule of "particular applicability." Therefore, it is not subject to the congressional review requirements in 5 U.S.C. 801-808.

**List of Subjects in 21 CFR Part 524**

Animal drugs.

Therefore, under the Federal Food, Drug, and Cosmetic Act and under authority delegated to the Commissioner of Food and Drugs and redelegated to the Center for Veterinary Medicine, 21 CFR part 524 is amended as follows:

**PART 524—OPHTHALMIC AND TOPICAL DOSAGE FORM NEW ANIMAL DRUGS**

1. The authority citation for 21 CFR part 524 continues to read as follows:

**Authority:** 21 U.S.C. 360b.

**§ 524.2481 [Amended]**

2. Section 524.2481 *Triamcinolone acetonide cream* is amended in paragraph (b) by adding after "No." the phrase "051259 and".

Dated: March 20, 2000.

**Stephen F. Sundlof,**

*Director, Center for Veterinary Medicine.*

[FR Doc. 00-7931 Filed 3-29-00; 8:45 am]

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**DEPARTMENT OF HOUSING AND URBAN DEVELOPMENT**

**24 CFR Part 982**

**[Docket No. FR-4459-C-07]**

**RIN 2577-AB96**

**Renewal of Expiring Annual Contributions Contracts in the Tenant-Based Section 8 Program; Formula for Allocation of Housing Assistance; Correction**

**AGENCY:** Office of Public and Indian Housing, HUD.

**ACTION:** Final rule.

**SUMMARY:** On October 21, 1999, HUD published a final rule that specified the method HUD will use in allocating housing assistance available to renew expiring contracts with public housing agencies (PHAs) for Section 8 tenant-based housing assistance. As required by statute, the October 21, 1999 final rule was the product of a negotiated rulemaking, following implementation, as further required by statute, of a HUD notice on this subject. The purpose of this document is to correct two typographical errors contained in the October 21, 1999 final rule.