

**29791. Rawleigh beef and sheep premix, and Rawleigh poultry premix.** (F.D.C. No. 49099. S. Nos. 11-850/1 V.)

**QUANTITY:** 2 cases, each containing 4 10-lb. bags of beef and sheep premix, and 8 cases, each containing 4 10-lb bags of poultry premix, at Menands, N.Y.

**SHIPPED:** Prior to 4-5-63, from Freeport, Ill., by W. T. Rawleigh Co.

**LABEL IN PART:** (Tags) "Rawleigh Beef and Sheep Premix \* \* \* Active Drug Ingredients Tetra Alkylammonium Stearate (from Dynafac) 2% Ethylenediamine Dihyriodide\* .65% \* \* \* Manufactured by The W. T. Rawleigh Company Freeport, Illinois"; "Rawleigh Poultry Premix Medicated Active Drug Ingredients Arsanilic Acid 1.98% Growth Stimulant For Poultry For use in poultry feeds in amounts of not more than 10 pounds or less than 5 pounds per ton of complete feed. Analysis Per Pound of Rawleigh Poultry Premix Medicated Procaine Penicillin Not less than 0.5 gms. Equivalent to 0.3 gm. of Crystalline Penicillin G (Master Standard) \* \* \* Manufactured for the W. T. Rawleigh Company Freeport, Illinois."

**ACCOMPANYING LABELING:** Feeding chart entitled "Poultry Premix Ration Chart Direction Card No. 14."

**RESULTS OF INVESTIGATION:** Investigations showed that the label for the poultry premix directed, as one use of the article, that it be used in a complete feed for the intended purpose of growth stimulation at a level of procaine penicillin which was lower than that set forth in the food additive regulations.

**LIBELED:** 7-10-63, N. Dist. N.Y.

**CHARGE:** 402(a) (2) (C)—when shipped and while held for sale, the beef and sheep premix contained the food additives, tetra alkylammonium stearate and ethylenediamine dihyriodide, which in combination were unsafe within the meaning of 409; and the poultry premix contained arsanilic acid and procaine penicillin which were unsafe within the meaning of 409 under the directions for use as a concentrate for continuous feeding (arsanilic acid) and as a complete feed (procaine penicillin), since such food additives and their use and intended use were not in conformity with a regulation or exemption.

The articles were alleged also to be violative of the law applicable to drugs as reported in notices of judgment on drugs and devices, No. 7887.

**DISPOSITION:** 9-9-63. Default—destruction.

**29792. Super Bee vitamin-mineral tablets.** (F.D.C. No. 50113. S. Nos. 44-723 A, 43-842 A.)

**QUANTITY:** 69 90-tablet btl., and 60 45-tablet btl., at Denver, Colo., in possession of Interstate Distributors, Inc.

**SHIPPED:** In March or April 1960, from Los Angeles, Calif., by Aldan Distributors.

**LABEL IN PART:** (Btl.) "No. 90 Super Bee Vitamins-Minerals 10,000 Mcg. 100% Pure Royal Jelly A Food Supplement \* \* \* Aldan Distributors, Inc."

**LIBELED:** 5-8-64, Dist. Colo.

**CHARGE:** 403(a)—when shipped and while held for sale, the labeling of the article contained false and misleading representations that royal jelly was a substance of significance for dietary supplementation or dietary use; 403(a)—the statement "100% Pure Royal Jelly" in the setting in which it appeared in the labeling represented and suggested that the article was 100 percent

royal jelly which was false and misleading; and 403(a)—the prominence given "10,000 Mcg." was misleading since it represented and suggested that the article contained materially more than the figure's equivalence, 10 mg., which is the common term or unit normally used.

DISPOSITION: 6-18-64. Default—destruction.

29793. Vitamin capsules (3 seizure actions). (F.D.C. Nos. 49762, 49764/5. S. Nos. 13-665 A; 13-667 A; 13-668 A.)

QUANTITY: 4 100-capsule btls. of prenatal vitamin capsules, 6 100-capsule btls. of yellow gelatin vitamin capsules, and 62 100-capsule btls. of high potency vitamin capsules, at Fall River, Mass.

SHIPPED: Prior to 1959, from New York, N.Y., and Providence, R.I.

LIBELED: 2-11-64, Dist. Mass.

CHARGE: 402(a)(2)(C)—while held for sale, the articles contained a food additive, folic acid, which was unsafe within the meaning of 409, since it and its use or intended use were not in conformity with a regulation or exemption.

DISPOSITION: 6-29-64; 10-5-64; 10-5-64. Default—destruction.

29794. Vitamin drops and capsules. (F.D.C. No. 49766. S. Nos. 13-669/70 A.)

QUANTITY: 11 30-cc. btls., and 7 60-cc. btls., of vitamin drops, and 8 250-capsule btls., of vitamin capsules, at Fall River, Mass.

SHIPPED: Prior to 1959, from Kalamazoo, Mich.

LABEL IN PART: "Each cc. contains: Folic Acid 1 mg. \* \* \* Dosage Infants and Children 1 to 2 cc. daily \* \* \* Adults 1 to 5 cc. daily," and "Each capsule contains: Folic Acid 0.5 mg. \* \* \* Usual Adult Dosage—1 to 2 capsules daily."

LIBELED: 2-11-64, Dist. Mass.

CHARGE: 402(a)(2)(C)—while held for sale, the article contained a food additive, folic acid, which was unsafe within the meaning of 409 since it and its use or intended use were not in conformity with a regulation or exemption.

DISPOSITION: 6-30-64. Default—destruction.

29795. Vitamin and mineral food supplement capsules. (F.D.C. No. 49665. S. Nos. 39-501 X, 39-718/19 X.)

QUANTITY: 212 cases, each containing 12 boxes of 12 18-capsule pkgs., 25 cases, each containing 72 18-capsule boxes, and 90 cases, each containing 12 boxes of 12 21-capsule pkgs., at Newark, N.J.

SHIPPED: Between 8-9-60 and 11-1-60, from Detroit, Mich.

RESULTS OF INVESTIGATION: Examination showed that the articles contained between 67 percent and 75 percent of the declared amount of vitamin B<sub>1</sub> and between 70 percent and 78 percent of the declared amount of vitamin C.

LIBELED: On or about 1-16-64, Dist. N.J.

CHARGE: 402(b)(1)—while held for sale, the valuable constituents, vitamin B<sub>1</sub> and vitamin C had been in part omitted or abstracted from the articles; and 403(a) the label statements "Each \* \* \* Capsule Contains: \* \* \* Vitamin B-1 \* \* \* 3 mg. [or "2 mg."] \* \* \* Vitamin C \* \* \* 50 mg." were false and misleading.

DISPOSITION: 2-13-64. Default—destruction.

29796. Vitamin and mineral food supplement tablets. (F.D.C. No. 49802. S. Nos. 62-001/2 X.)