

COMPETITIVE PROBLEMS IN THE DRUG INDUSTRY 10373

DEFENSE MEDICAL PURCHASE DESCRIPTION		NUMBER 9	DATE 11 August 1971
FEDERAL STOCK NO.	ITEM IDENTIFICATION		UNIT
6505-130-1500	NIACINAMIDE TABLETS, USP, 50 mg, 100s		Bottle

Shall be Niacinamide Tablets, U.S.P., and shall be in accordance with all applicable requirements of Federal Standard Fed. Std. No. 140a, dated 30 October 1966, and Amendment-1, dated 25 March 1970, and as specified herein:

S2. Classification. Shall be type I, class 1.

S5.2 The following additional requirements and tests are added to this paragraph:

Shall be tablets containing 50 mg of Niacinamide per tablet, within the applicable assay limits for the tablets.

S6.4.2 Color. Uncoated tablets shall be white.

PREPARATION FOR DELIVERY

Shall be in accordance with all applicable requirements of Interim Federal Specification PPP-C-00186a, dated 15 May 1969, and Amendment-1, dated 27 October 1969, and as specified herein:

Immediate containers. Shall comply with the following classification:

GROUP A	CLASS 1	TYPE e	STYLE 2	GRADE 1
CLOSURE A, B, or F				SEAL A or B

Labeling. Labeling shall be in accordance with the requirements of the Federal Food, Drug, and Cosmetic Act, and shall include the information required below:

Immediate containers. Each immediate container label shall bear the following information. However, the information is not required to appear in the sequence indicated:

- (a) the item name designated as "NIACINAMIDE TABLETS, U.S.P."
- (b) the quantity of active ingredient designated as "50 mg"
- (c) the phrase "100 tablets" or a similar phrase
- (d) the Federal Stock No.

(See additional label information on page 2)

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