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should be used and any or all of the additional indications for sunscreen drug products may be used.

(c) Warnings. The labeling of the product states, under the heading ‘‘Warnings,’’ the warning(s) for each ingredient in the combination, as established in the warnings section of the applicable OTC drug monographs unless otherwise stated in this paragraph (c).

(1) For combinations containing a skin protectant and a sunscreen identified in §§ 347.20(d) and 352.20(b). The warnings for sunscreen drug products in § 352.60(c) of this chapter are used.

(2) [Reserved]

(d) Directions. The labeling of the product states, under the heading ‘‘Directions,’’ directions that conform to the directions established for each ingredient in the combination in the directions sections of the applicable OTC drug monographs, unless otherwise stated in this paragraph (d). When the time intervals or age limitations for administration of the individual ingredients differ, the directions for the combination product may not contain any dosage that exceeds those established for any individual ingredient in the applicable OTC drug monograph(s), and may not provide for use by any age group lower than the highest minimum age limit established for any individual ingredient.

(1) For combinations containing a skin protectant and a sunscreen identified in §§ 347.20(d) and 352.20(b). The directions for sunscreen drug products in § 352.60(d) of this chapter are used.

(2) [Reserved]

PART 348—EXTERNAL ANALGESIC DRUG PRODUCTS FOR OVER-THE-COUNTER HUMAN USE

Subpart C—Labeling

§ 348.50 Labeling of external analgesic drug products.


SOURCE: 57 FR 27656, June 19, 1992, unless otherwise noted.

Subpart A—General Provisions

§ 348.1 Scope.

(a) An over-the-counter external analgesic drug product in a form suitable for topical administration is generally recognized as safe and effective and is not misbranded if it meets each condition in this part and each general condition established in §330.1 of this chapter.

(b) References in this part to regulatory sections of the Code of Federal Regulations are to chapter I of title 21 unless otherwise noted.

Subpart B—Active Ingredients

§ 348.10 Analgesic, anesthetic, and antipruritic active ingredients.

The active ingredient of the product consists of any of the following within the specified concentration established for each ingredient:

(a) Male genital desensitizers. (1) Benzocaine, 3 to 7.5 percent in a water-soluble base.

(2) Lidocaine in a metered spray with approximately 10 milligrams per spray.

(b) [Reserved]

Subpart C—Labeling

§ 348.50 Labeling of external analgesic drug products.

(a) Statement of identity. The labeling of the product contains the established name of the drug. If any, and identifies the product as follows:

(1) For products containing any ingredient identified in §348.10(a). “Male genital desensitizer.”