§ 352.20 Permitted combinations of active ingredients.

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(a) Combinations of sunscreen active ingredients. (1) Two or more sunscreen active ingredients identified in §352.10(a), (c), (e), (f), (g), and (i) through (r) may be combined with each other in a single product when used in the concentrations established for each ingredient in §352.10. The concentration of each active ingredient must be sufficient to contribute a minimum SPF of not less than 2 to the finished product. The finished product must have a minimum SPF of not less than the number of sunscreen active ingredients used in the combination multiplied by 2.

(2) Two or more sunscreen active ingredients identified in §352.10(b), (c), (e), (g), (j) through (m), (o), and (q) may be combined with each other in a single product when used in the concentrations established for each ingredient in §352.10. The concentration of each active ingredient must be sufficient to contribute a minimum SPF of not less than 2 to the finished product. The finished product must have a minimum SPF of not less than the number of sunscreen active ingredients used in the combination multiplied by 2.

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Subpart C—Labeling

§ 352.50 Principal display panel of all sunscreen drug products.

In addition to the statement of identity required in §352.52, the following labeling statements shall be prominently placed on the principal display panel:

(a) For products that do not satisfy the water resistant or very water resistant sunscreen product testing procedures in §352.76. (1) For products with SPF values up to 30. “SPF (insert tested SPF value of the product up to 30).”

(2) For products with SPF values over 30. “SPF 30” (select one of the following: “plus” or “+”). Any statement accompanying the marketed product that states a specific SPF value above 30 or similar language indicating a person can stay in the sun more than 30 times longer than without sunscreen will cause the product to be misbranded under section 502 of the Federal Food, Drug, and Cosmetic Act (the act).

(b) For products that satisfy the water resistant sunscreen product testing procedures in §352.76. (1) (Select one of the following: “Water,” “Water/Sweat,” or “Water/Perspiration”) “Resistant.”

(2) “SPF (insert SPF value of the product, as stated in paragraph (a)(1) or (a)(2) of this section, after it has been tested using the water resistant sunscreen product testing procedures in §352.76).”

(c) For products that satisfy the very water resistant sunscreen product testing procedures in §352.76.

(1) “Very” (select one of the following: “Water,” “Water/Sweat,” or “Water/Perspiration”) “Resistant.”

(2) “SPF (insert SPF value of the product, as stated in paragraph (a)(1) or (a)(2) of this section, after it has been tested using the very water resistant sunscreen product testing procedures in §352.76).”

§ 352.52 Labeling of sunscreen 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 a “sunscreen.”

(b) Indications. The labeling of the product states, under the heading “Uses,” all of the phrases listed in paragraph (b)(1) of this section that are applicable to the product and may contain any of the additional phrases listed in paragraph (b)(2) of this section, as appropriate. Other truthful and non-misleading statements, describing only the uses that have been established and listed in this paragraph (b), may also be used, as provided in §330.1(c)(2) of this chapter, subject to the provisions of section 502 of the act relating to misbranding and the prohibition in section 301(d) of the act against the introduction or delivery for introduction into interstate commerce of unapproved new drugs in violation of section 505(a) of the act.

(1) For products containing any ingredient in §352.10. (1) “[bullet] helps prevent sunburn” [bullet] higher SPF gives more sunburn protection”.

(2) For products that satisfy the water resistant testing procedures identified in §352.76. “[bullet] retains SPF after 40