

SANITAXE ANTIBACTERIAL- benzalkonium chloride gel
American Consumer Products Corp

Sanitaxe Hand Soap

Active Ingredient

Active Ingredient

Benzalkonium Chloride .13%

Purpose

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Antiseptic

Uses

Uses ● Helps reduce bacteria on the skin that could cause disease. Recommended for repeated use.

Warnings

Warnings - For external use only. Do not ingest or swallow.

Do not apply around eyes. Do not use in ears & mouth.

When using this product

When using this product, avoid contact with eyes. In case of contact, flush eyes with water.

Stop use and ask a doctor

Stop use and ask a doctor if redness or irritation develops and persists for more than 72 hours.

Keep out of reach of children

Keep out of reach of children. Do not use on children less than 2 months of age.

Supervise use in children under 6 years of age to prevent accidental swallowing.

If swallowed, get medical help or contact a Poison Control Center right away.

Directions

Directions ● Pump onto wet hands ● Work into lather ● Wash hands, rinse and dry thoroughly.

Inactive ingredients

Inactive ingredients: Water (Aqua), Cocamidopropyl Betaine, Sodium Cocoamphoacetate, PEG-150 Disterate, Fragrance, Citric Acid, Methylisothiazolinone, Iodopropynyl Butylcarbamate, FD&C Yellow No. 5, D&C Red No. 33.

Sanitaxe Hand Soap



SANITAXE ANTIBACTERIAL

benzalkonium chloride gel

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:72197-033
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7) (BENZALKONIUM - UNII: 7N6JUD5X6Y)	BENZALKONIUM CHLORIDE	0.13 g in 100 mL

Inactive Ingredients

Ingredient Name	Strength
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FD&C YELLOW NO. 5 (UNII: I753WB2F1M)	
FD&C RED NO. 40 (UNII: WZB9127XOA)	
WATER (UNII: 059QF0KO0R)	
METHYLISOTHIAZOLINONE (UNII: 229D0E1QFA)	
IODOPROPYNYL BUTYLCARBAMATE (UNII: 603P14DHEB)	
PEG-150 DISTEARATE (UNII: 6F36Q0I0AC)	
ANHYDROUS CITRIC ACID (UNII: XF417D3PSL)	
COCAMIDOPROPYL BETAINE (UNII: 5OCF3O11KX)	
SODIUM COCOAMPHOACETATE (UNII: W7Q5E87674)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:72197-033-27	800 mL in 1 BOX; Type 0: Not a Combination Product	07/15/2020	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M003	07/15/2020	

Labeler - American Consumer Products Corp (081101181)

Revised: 1/2025

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