

ALCOHOL-FREE HAND SANITIZING FOAM- benzalkonium chloride solution
United Laboratories Inc.

Alcohol-Free Hand Sanitizing Foam 63998-764

Drug Facts

Active ingredient

Benzalkonium Chloride 0.1%

Purpose

Antimicrobial

Uses

For hand sanitizing to decrease bacteria on the skin. Recommended for repeated use.

Warnings

For external use only

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

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

Keep out of the reach of children. If swallowed, get medical help or contact a Poison Control Center right away.

Directions

Pump a small amount of foam into palm of hand. Rub thoroughly over all surfaces of both hands. Rub hands together briskly until dry.

Inactive ingredients

Water, cetrimonium chloride, laurtrimonium chloride, dihydroxyethyl cocamine oxide, glycereth-17 cocoate, citric acid and fragrance.

PRINCIPAL DISPLAY PANEL - 2 Ounce Bottle Label

UNITED
LABORATORIES

United 764

MICROMOUSSE

Alcohol-Free
Hand Sanitizing Foam

2 FLUID OUNCES

Eliminates 99.99% of common
disease-causing germs.

Made in USA
1018

UNITED LABORATORIES, INC.
320 37th Avenue • St. Charles, IL 60174
1-800-323-2594 • www.unitedlabsinc.com



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ALCOHOL-FREE HAND SANITIZING FOAM

benzalkonium chloride solution

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:63998-764
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7) (BENZALKONIUM - UNII:7N6JUD5X6Y)	BENZALKONIUM CHLORIDE	1 mg in 1 mL

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
CETRIMONIUM CHLORIDE (UNII: UC9PE95IBP)	
LAURTRIMONIUM CHLORIDE (UNII: A81MSI0FIC)	
DIHYDROXYETHYL COCAMINE OXIDE (UNII: 8AR51R3BL5)	
GLYCERETH-17 COCOATE (UNII: 3057VPT0KC)	
CITRIC ACID MONOHYDRATE (UNII: 2968PHW8QP)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:63998-764-01	59 mL in 1 BOTTLE; Type 0: Not a Combination Product	01/01/2019	
2	NDC:63998-764-02	750 mL in 1 BOTTLE; Type 0: Not a Combination Product	01/01/2019	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	505G(a)(3)	01/01/2019	

Labeler - United Laboratories Inc. (001759737)

Establishment

Name	Address	ID/FEI	Business Operations
Woodbine Products Company		004220323	manufacture(63998-764)

Revised: 11/2024

United Laboratories Inc.