

WHISKCARE 377- benzalkonium chloride solution
Whisk Products, Inc.

WhiskCare 377

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 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, Cocamidopropyl PG-Dimonium Chloride Phosphate, Dihydroxyethyl Cocamine Oxide, Acetamidoethoxyethanol, Fragrance, Citric Acid



Whisk Products, Inc.
Wentzville, MO 63385 • www.whiskproducts.com

WhiskCare 377
FOAMING Instant Hand Sanitizer
Alcohol Free

WC-377-C-4

Drug Facts	
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Benzalkonium Chloride 0.1%	Antimicrobial
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LOT #: EXP:

Contains: Four CleanShot Bags
Net Contents Each: 33.8 fl. oz. (1 qt. 1.8 fl. oz.) 1000 mL
Total Net Volume: 135.2 fl. oz. (1 gal. 7.2 fl. oz.) 4 L



WHISKCARE 377

benzalkonium chloride solution

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:65585-356
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
ACETAMIDOETHOXYETHANOL (UNII: LVX2APC4XR)	
DIHYDROXYETHYL COCAMINE OXIDE (UNII: 8AR51R3BL5)	
COCAMIDOPROPYL PG-DIMONIUM CHLORIDE PHOSPHATE (UNII: H2KVQ74JM4)	
CITRIC ACID MONOHYDRATE (UNII: 2968PHW8QP)	
WATER (UNII: 059QF0KO0R)	

Product Characteristics

Color	white (water white - colorless, dispensed as a white foam)	Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65585-356-01	50 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product	02/22/2024	08/01/2027
2	NDC:65585-356-02	24 in 1 BOTTLE	02/22/2024	08/01/2027
2		50 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product		
3	NDC:65585-356-03	208198 mL in 1 DRUM; Type 0: Not a Combination Product	02/22/2024	08/01/2027
4	NDC:65585-356-04	4 in 1 BOX	02/22/2024	08/01/2027
4		1000 mL in 1 BAG; Type 0: Not a Combination Product		
5	NDC:65585-356-05	8 in 1 BOX	02/22/2024	08/01/2027
5		1000 mL in 1 BAG; Type 0: Not a Combination Product		
6	NDC:65585-356-06	6 in 1 BOX	02/22/2024	08/01/2027
6		550 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product		
7	NDC:65585-356-07	6 in 1 BOX	02/22/2024	02/22/2024
7		1750 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product		
8	NDC:65585-356-08	3785 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	02/22/2024	08/01/2027

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	505G(a)(3)	02/22/2024	08/01/2027

Labeler - Whisk Products, Inc. (834270639)

Establishment

Name	Address	ID/FEI	Business Operations
Whisk Products, Inc.		834270639	manufacture(65585-356)

Revised: 9/2024

Whisk Products, Inc.