SUNDROPS 87- benzalkonium chloride soap Sunburst Chemicals, Inc.

SunDrops 87

Active Ingredient

Benzalkonium Chloride 0.1%

Purpose

Skin Antimicrobial

Use

Reduces amount of bacteria on hands

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.
- Scrub thoroughly for at least fifteen seconds.
- Rinse completely and dry.

Inactive Ingredients

Water, Lauramine Oxide, Glycerin, PEG-120 Methyl Glycose Dioleate, Fragrance



Distributed Exclusively By: Sunburst Chemicals, Inc. Minneapolis, MN 55420 www.sunburstresults.com

Contains: Eight Bags

Net Contents Each: 33.8 fl. oz. (1 qt. 1.8 fl. oz.) 1000 mL

Total Net Volume: 270.4 fl. oz. (2 gal. 14.4 fl. oz.) 8 L

Drug Facts

Active Ingredient Purpose Bengalkonium Chloride 0.1% Sidn Antimicrobial

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LOT #: EXP:

SUNDROPS 87

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Product Information

Product Type HUMAN OTC DRUG **Item Code (Source)** NDC:63621-387

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
BENZALKONIUM CHLORIDE (UNII: F5UM2KM3W7) (BENZALKONIUM -	BENZALKONIUM	1 mg

UNII:7N6JUD5X6Y) **CHLORIDE** in 1 mL

Inactive Ingredients		
Ingredient Name	Strength	
WATER (UNII: 059QF0KO0R)		
GLYCERIN (UNII: PDC6A3C0OX)		
LAURAMINE OXIDE (UNII: 4F6FC4MI8W)		
PEG-120 METHYL GLUCOSE DIOLEATE (UNII: YM0K64F20V)		

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

ı	Pa	ckaging			
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date

1	NDC:63621- 387-50	6 in 1 BOX	06/08/2020	
1		500 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product		
2	NDC:63621- 387-14	3785 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	06/08/2020	11/17/2023
3	NDC:63621- 387-65	8 in 1 BOX	07/14/2020	
3		1000 mL in 1 BAG; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	505G(a)(3)	06/08/2020	

Labeler - Sunburst Chemicals, Inc. (006159339)

Revised: 3/2024 Sunburst Chemicals, Inc.