UNSCENTED NASALGUARD ANTISEPTIC TOPICAL- benzalkonium chloride gel TRUTEK CORP.

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

Unscented NasalGuard Antiseptic Topical Gel

Active Ingredient

Benzalkonium Chloride 0.13%

Purpose

Antiseptic

Use

To decrease bacteria on the skin.

Directions

Apply to the outside of the nose as indicated below.

Begin by squeezing 1 to 2 pin-sized drops on your pointer finger.

Spread the gel between your pointer finger and thumb to ensure an even application.

Apply directly around the nostrils and above the upper lip.

Apply every 4 to 6 hours or as often as needed.

Reapply if nose or face becomes wet.

SAVE THIS (BOX) FOR REFERENCE

Inactive Ingredients

Dimethicone, Glycerin, Hydroxyethyl Cellulose, Purified Water. Alson contains non-toxic, cosmetic grade preservatives and emulsifiers.

Questions or Comments?

Visit www.NasalGuard.com or call 1-855-NASALGUARD (1-855-627-2548).

NASALGUARD ANTISEPTIC TOPICAL Gel Unscented

For external use only

When using this product keep out of eyes. In case of contact with eyes, rinse eyes thoroughly with water.

Stop use and ask a doctor if

irritation and redness develop condition persists for more than 72 hours

If pregnant or breast-feeding, ask a health professional before use.

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

Center right away.

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DATE: 20.OCT.2022

BRAND: NASALGUARD USA

JOB NAME: UNSCENTED ANTISEPTIC TOPICAL GEL

VERSION: FINAL

Card Stock 360 GSM

Computer Equivalents:











NasalGuard Unscented Tube and Carton

UNSCENTED NASALGUARD ANTISEPTIC TOPICAL

benzalkonium chloride gel

Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:81907-102	
Route of Administration	TOPICAL			

Active Ingredient/Active Moiety			
	Ingredient Name	Basis of Strength	Strength
	, (, ,)	BENZ ALKONIUM CHLORIDE	1.3 mg in 1000 mg

Inactive Ingredients			
Ingredient Name	Strength		
DIMETHICONE 350 (UNII: 2Y53S6ATLU)			
STEARETH-2 (UNII: V56DFE46J5)			
WATER (UNII: 059QF0KO0R)			
BEHENTRIMONIUM CHLORIDE (UNII: X7GNG3S47T)			
MEDIUM-CHAIN TRIGLYCERIDES (UNII: C9H2L21V7U)			
HYDROXYETHYL CELLULOSE, UNSPECIFIED (UNII: T4V6TWG28D)			
STEARETH-21 (UNII: 53J3F32P58)			
GLYCERIN (UNII: PDC6A3C0OX)			
POTASSIUM SORBATE (UNII: 1VPU26JZZ4)			
PHENOXYETHANOL (UNII: HIE492ZZ3T)			

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:81907-102- 01	1 in 1 BOX	02/14/2022	
1	NDC:81907-102- 02	3000 mg in 1 TUBE; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333E	02/14/2022	

Labeler - TRUTEK CORP. (170729235)

Registrant - TRUTEK CORP. (170729235)

Establishment				
Name	Address	ID/FEI	Business Operations	
TRUTEK CORP.		170729235	manufacture(81907-102)	

Revised: 3/2023 TRUTEK CORP.