GUARD- alcohol gel Apollo Health and Beauty Care

NDC- 63148-502, Guard Hand Sanitizer

Active Ingredient

Ethyl Alcohol 70%

Purpose

Antiseptic

Uses

For personal hand hygine to help prevent the spread of bacteria, and can be used in place of hand washing if soap and water are not available.

Warnings

For external use only.

- Flammable
- Keep away from source of heat or fire.

Do not use

- On infants less than 2 months of age.
- On open skin wounds
- On broken or damaged skin.

When using this product avoid

- contact with eyes. If contact occurs, rinse eyes thoroughly with water.
- Do not inhale.

Stop use and ask a doctor

if irritation or redness develops and lasts.

Keep out of reach of children

In case of accidental ingestion, get medical help or contact Poison Control Center immediately.

Directions

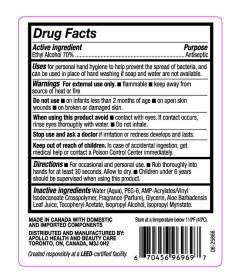
- For occassional and Personal use.
- Rub thoroughly into hands for at least 30 seconds. Allo to dry.
- Children under 6 years should be supervised when using this product.

Inactive Ingredients

Water (Aqua), PEG-6, AMP-Acrylates/vinyl Isodecanoate Crosspolymer, Fragrance (Parfum), Glycerin, Aloe Barbadensis Leaf Juice, Tocopheryl Acetate, Isopropyl Alcohol, Isopropyl Myristate.

Package Label







THIS IS NOT A COLOUR ACCURATE PROOF. Please refer to process and pantone colour charts for accurate colour representation.

Every effort has been made to ensure the accuracy of this proof. However, the client is responsible for ensuring that label size, copy, graphics and colour separations are accurate, and to notify Perflex

Label Inc. of any discrepancies prior to film, plate or label production. The client indemnifies Perflex Label Inc. against any liability related to costs incurred due to errors on a customer signed proof.



PLEASE NOTE: A SIGNED PROOF IS REQUIRED FOR PRODUCTION. EMAIL APPROVAL TO YOUR CONTACT OR FAX TO 416 321-2267.

alcohol gel

Product Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:63148-502

Route of Administration TOPICAL

Active Ingredient/Active Moiety

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	Ingredient Name	Basis of Strength	Strength	
	ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	70 mL in 100 mL	

Inactive Ingredients			
Ingredient Name	Strength		
ACRYLATES/VINYL ISODECANOATE CROSSPOLYMER (10000 MPA.S NEUTRALIZED AT 0.5%) (UNII: 2N8MDB79NA)			
POLYETHYLENE GLYCOL 300 (UNII: 5655G9Y8AQ)			
GLYCERIN (UNII: PDC6A3C0OX)			
WATER (UNII: 059QF0KO0R)			
ISOPROPYL MYRISTATE (UNII: 0RE8K4LNJS)			
ISOPROPYL ALCOHOL (UNII: ND2M416302)			
ALOE VERA LEAF (UNII: ZY81Z83H0X)			
FRAGRANCE CLEAN ORCO600327 (UNII: 329LCV5BTF)			
.ALPHATOCOPHEROL ACETATE (UNII: 9E8X80D2L0)			

l	P	Packaging				
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
	1	NDC:63148- 502-08	227 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	06/01/2023		

Marketing Information					
Application Number or Monograph Citation	Marketing Start Date	Marketing End Date			
05G(a)(3)	06/01/2023				
	Application Number or Monograph Citation	Application Number or Monograph Citation Date			

Labeler - Apollo Health and Beauty Care (201901209)

Registrant - Apollo Health and Beauty Care (201901209)

Establishment					
Na me	Address	ID/FEI	Business Operations		
Apollo Health and Beauty Care		201901209	manufacture(63148-502)		