DOCTORS KLINE AND GREEN BOARD CERTIFIED DERMATOLOGIST HAND SANITIZER ALCOHOL ANTISEPTIC- alcohol liquid Formulated Solutions, LLC

Doctors Kline & Green Board Certified Dermatologist Hand Sanitizer Alcohol Antiseptic

Drug Facts

Active ingredient

Ethyl Alcohol 77% v/v

Purpose

Antiseptic

Uses(s)

Hand sanitizer to help reduce bacteria that potentially can cause disease. For use when soap and water are not available.

Warnings

For external use only. Extermely flammable, do not use near heat or flame or while smoking.

Do not use

- On children less than 2 months of age
- On open skin wounds

When using this product

- keep out of eyes, ears, and mouth
- In case of contact with eyes, rinse eyes thoroughly with water.
- Do not puncture or incinerate
- Contents under pressure
- Do not expose to heat or store at temperatures above 120°F (49°C)

Stop use and ask a doctor

• If irritation or rash occurs. These may be signs of a serious condition.

Keep out of reach of children.

If swallowed, get medical help or contact a Poison Control Center right away.

Directions

- Place enough product on hands to cover all surfaces. Rub hands together until dry
- Supervise children under 6 years of age when using this product to avoid swallowing.

Other information

- Store between 15-30°C (59-86°F)
- Avoid freezing and excessive heat beyond 40°C (104°F)

Inactive ingredients

Aloe Barbadensis Leaf Juice, Ammonium Acryloyldimethyltaurate/VP Copolymer, HYdroxyethyl Urea, Tocopheryl Acetate, Water

Package Labeling:

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MANUFACTURED FOR Three Seasons Healthcare LLC One North Clematis St., Suite 110 West Palm Beach, FL 33401-5551







BOARD CERTIFIED DERMATOLOGISTS

HAND SANITIZER

Alcohol Antiseptic 77% Topical Solution



NET WT. 5.0 OZ. (141g) / 5.6 FL. OZ. (166 mL)

DOCTORS KLINE AND GREEN BOARD CERTIFIED DERMATOLOGIST HAND SANITIZER ALCOHOL ANTISEPTIC

alcohol liquid

Product Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:23667-102

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Active ingredient/Active Molecy			
Ingredient Name	Basis of Strength	Strength	
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	77 mL in 100 mL	

Inactive Ingredients		
Ingredient Name	Strength	
ALOE VERA LEAF (UNII: ZY81Z83H0X)		
AMMONIUM ACRYLOYLDIMETHYLTAURATE/VP COPOLYMER (UNII: W59H9296ZG)		
HYDROXYETHYL UREA (UNII: MBQ7DDQ7AR)		
.ALPHATOCOPHEROL ACETATE (UNII: 9E8X80D2L0)		
WATER (UNII: 059QF0KO0R)		

Packaging					
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
		NDC:23667-102- 00	166 mL in 1 BOTTLE; Type 0: Not a Combination Product	02/09/2021	

tart Marketing End Date
1

Labeler - Formulated Solutions, LLC (143266687)

Revised: 11/2023 Formulated Solutions, LLC