ADORE PEPPERMINT HAND SANITIZER- alcohol gel K7 Design Group Inc.

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.

Adore Peppermint Hand Sanitizer (Green)

Drug Facts

Active ingredient

Alcohol 69% v/v

Purpose

Antiseptic

Use

for hand-washing to decrease bacteria on the skin, only when water is not available

Warnings

Flammable, keep away from fire and flames

For external use only

When using this product

- do not get into eyes.
- if contact occurs, rinse eyes thoroughly with water

Stop use and ask a doctor if

• irritation and redness develop

Keep out of reach of children.

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

Directions

wet hands thoroughly with product and allow to dry without wiping

Inactive ingredients

Water, Glycerin, Propylene Glycol, Carbomer, Aloe Barbadensis Leaf Extract, Aminomethyl Propanol, Fragrance, Tocopheryl Acetate, Denatonium Benzoate, Yellow 5, Blue 1.

Company Information

MANUFACTURED FOR & DISTRIBUTED BY K7 DESIGN GROUP LLC NEW YORK, NY 10016

Product Packaging







MANUFACTURED FOR & DISTRIBUTED BY K7 DESIGN GROUP LLC. NEW YORK, NY 10016 EXPIRATION: 01/15/2024 LOT: 0Y012021 ORIGIN: CHINA

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Drug Facts (continued)

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ADORE PEPPERMINT HAND SANITIZER

alcohol gel

Product Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:74177-985

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient Name Basis of Strength Strength
ALCOHOL (UNII: 3K9958 V90M) (ALCOHOL - UNII:3K9958 V90M)
ALCOHOL (UNII: 3K9958 V90M) ALCOHOL 69 mL in 100 mL

Ingredient Name Strength ALOE VERA LEAF (UNII: ZY81Z83H0X) AMINOMETHYLPRO PANOL (UNII: LU49E6626Q) WATER (UNII: 059QF0K00R) CARBOMER HOMO POLYMER, UNSPECIFIED TYPE (UNII: 0A5MM307FC) FD&C BLUE NO. 1 (UNII: H3R47K3TBD) GLYCERIN (UNII: PDC6A3C0OX) PROPYLENE GLYCOL (UNII: 6DC9Q167V3) .ALPHA.-TO CO PHEROL ACETATE (UNII: 9E8X80D2L0) DENATONIUM BENZO ATE (UNII: 4YK5Z54AT2) FD&C YELLOW NO. 5 (UNII: 1753WB2F1M)

Packaging					
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:74177-985- 01	30 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	0 1/0 5/20 21		

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
OTC monograph not final	part333A	01/05/2021			

Labeler - K7 Design Group Inc. (080357784)

Revised: 1/2021 K7 Design Group Inc.