FROSTED MINT- hand sanitizer spray Bell International Laboratories, 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.

Touchland Frosted Mint Hand Sanitizer Spray

Active Ingredient

Ethyl Alcohol 70% v/v

Purpose

Antiseptic

Uses

To decrease bacteria on the skin

Recommended for repeated use

Warnings

For external use only

Flammable, keep away from fire or flame

Do not use in or near the eyes. In case of contact, rinse eyes thoroughly with water. Avoid contact with broken skin

Stop use and ask a doctor

- if irritation and redness develop
- 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

Other Information

Protect the product in this container from excessive heat and direct sun

Store below 104F (40C)

May discolor certain fabrics

Directions

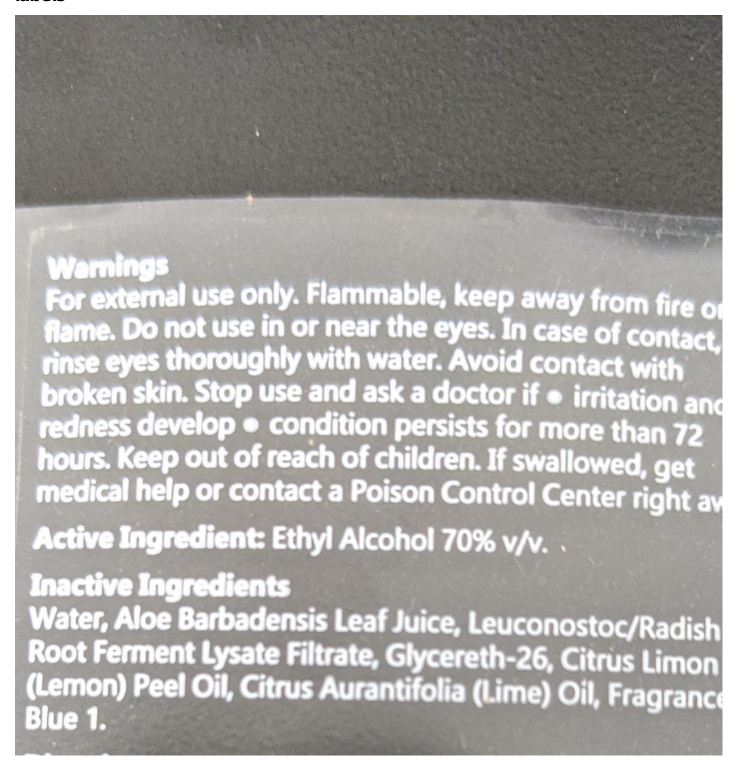
Place enough product in your palm to thoroughly cover your hands

Children under 6 years of age should be supervised when using this product Rub hands together briskly until dry

Inactive Ingredients

Water, Aloe Barbadensis Leaf Juice, Leuconostoc/Radish Root Ferment Lysate Filtrate, Glycereth-26, Citrus Limon (Lemon) Peel Oil, Citrus Aurantifolia (Lime) Oil, Fragrance, Blue 1

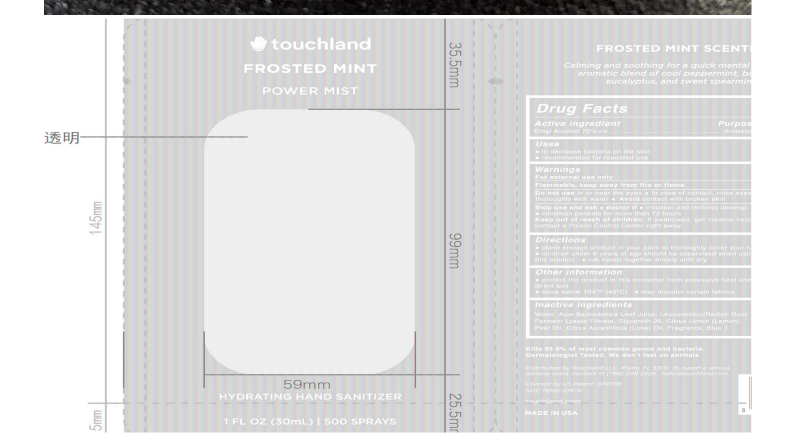
labels



Directions Spray onto hands and rub together until dry. Children under 6 years of age should be supervised when using this product. FROSTED MINT SCENTED

Distributed by Touchland LLC. Miami, FL 33131.
To report a serious adverse event,
contact +1 (786) 209 2328
hello@touchland.com

MADE IN USA



FROSTED MINT

hand sanitizer spray

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Product Type HUMAN OTC DRUG Item Code (Source) NDC:76150-278

Route of Administration TOPICAL

Active Ingredient/Active Moiety Ingredient Name Basis of Strength

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

Inactive Ingredients

mactive ingredients		
Ingredient Name	Strength	
GLYCERETH-26 (UNII: NNE56F2N14)		
FD&C BLUE NO. 1 (UNII: H3R47K3TBD)		
ALOE VERA LEAF (UNII: ZY81Z83H0X)		
WATER (UNII: 059QF0KO0R)		
LIME OIL (UNII: UZH29XGA8G)		
LEUCONOSTOC/RADISH ROOT FERMENT FILTRATE (UNII: D2QHA03458)		
LEMON OIL (UNII: 19GRO824LL)		

Packaging

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	#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
		NDC:76150- 278-51	30 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product	07/01/2021		

Marketing Information

OTC monograph not part333A 07/01/2021	Marketing	Application Number or Monograph	Marketing Start	Marketing End
	Category	Citation	Date	Date
Illidi	OTC monograph not final	part333A	07/01/2021	

Labeler - Bell International Laboratories, Inc. (967781555)

Revised: 8/2021 Bell International Laboratories, Inc.