

ADVANCED HAND SANITIZER- ethyl alcohol gel
Harris Teeter

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.

Advanced Hand Sanitizer with Aloe
439

Active ingredient

Ethyl Alcohol 70%

Purpose

Antiseptic

Uses

- to decrease bacteria on the skin that could cause disease
- recommended for repeated use

Warnings

For external use only-hands

Flammable. Keep away from heat and flame.

When using this product

- keep out of eyes. In case of contact with eyes, flush thoroughly with water
- avoid contact with broken skin
- do not inhale or ingest

Stop use and ask a doctor if

skin irritation develops

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
- for children under 6, use only under adult supervision
- not recommended for infants

Other information

- do not store above 105° F
- may discolor some fabrics
- harmful to wood finishes and plastics

Inactive ingredients

water, aloe, barbadensis leaf juice, glyceryl, caprylate/caprates, glycerin, isopropyl myristate, tocopheryl acetate, acrylates/C10-30 alkyl acrylate crosspolymer, fragrance, benzophenone-4, blue 1, yellow 5

*Effective at eliminating more than 99.99% of many common harmful germs and bacteria in as little as 15 seconds

**This product is not manufactured or distributed by GOJO Industries, Inc. distributor of Purell Refreshing Aloe Advanced Hand Sanitizer.

Proudly Distributed By: Harris Teeter, LLC

Matthews, NC 28105

1-800-432-6111 or harristeeter.com

SDS-MO-15036, SDA-WI-2486, DSP-MO-28, DSP-MO-34

439.000/439AB

Principal Display Panel

Harris Teeter

Advanced Hand Sanitizer With Aloe

Kills more than 99.99% of germs*

More Effective Formula

Compare to Purell® Refreshing Aloe Advanced Hand Sanitizer**

8 FL OZ (236 mL)



ADVANCED HAND SANITIZER

ethyl alcohol gel

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:72036-439	
Route of Administration	TOPICAL			
Active Ingredient/Active Moiety				
Ingredient Name		Basis of Strength	Strength	
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)		ALCOHOL	700 mg in 1 mL	
Inactive Ingredients				
Ingredient Name				Strength
WATER (UNII: 059QF0KO0R)				
ALOE VERA LEAF (UNII: ZY81Z83H0X)				
GLYCERYL CAPRYLATE/CAPRATE (UNII: G7515SW10N)				
GLYCERIN (UNII: PDC6A3C0OX)				
ISOPROPYL MYRISTATE (UNII: 0RE8K4LNJS)				
.ALPHA.-TOCOPHEROL ACETATE (UNII: 9E8X80D2L0)				
CARBOMER INTERPOLYMER TYPE A (ALLYL SUCROSE CROSSLINKED) (UNII: 59TL3WG5CO)				
SULISOBENZONE (UNII: 1W6L629B4K)				
FD&C BLUE NO. 1 (UNII: HBR47K3TBD)				
FD&C YELLOW NO. 5 (UNII: I753WB2F1M)				
Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:72036-439-16	59 mL in 1 BOTTLE, DISPENSING; Type 0: Not a Combination Product	07/18/2014	
2	NDC:72036-439-34	236 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product	07/18/2014	
Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC monograph not final	part333A	07/18/2014		

Labeler - Harris Teeter (047279351)

Registrant - Vi-Jon (790752542)

Establishment

Name	Address	ID/FEI	Business Operations
Vi-Jon		088520668	manufacture(72036-439)