ACP HAND SANITIZER- ethyl alcohol gel Swanrose, 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.

ACP Hand Sanitizer

Drug Facts

Active Ingredient

Ethyl Alcohol 73%

Purpose

Antimicrobial

Uses

- Hand sanitizer to reduce microorganisms on the skin.
- Use this product when soap and water are not available.

Warnings

- **For external use only.** Avoid contact with eyes. If contact occurs, rinse thoroughly with water. **FLAMMABLE. This product contains ethyl alcohol.Keep away from sources of ignition.** Discontinue use if irritation or redness develops. If irritation persists for more than 72 hours, consult a physician.
- KEEP OUT OF REACH OF CHILDREN.
- If swallowed, get medical help or contact a Poison Control Center right away.

Directions

- Read the entire label before using this product.
- Place enough product on your palm to thoroughly cover your hands.
- Rub hands together briskly until dry.

Inactive Ingredients

Aqua, Glycerin, Carbomer, Triethanolamine

73% Alcohol Self Drying Gel Rub well until dry

Distributed & Marketed by Swanrose Inc. SWANROSE, INC. 751 N. SPAULDING AVENUE LOS ANGELES, CA 90046

PHONE: +1 (310) 266-5812

Made in China

Packaging



73% Alcohol Self Drying Gel Rub well until dry

500ML

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ACP HAND SANITIZER

ethyl alcohol gel

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:78287-160
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)	ALCOHOL	73 mL in 100 mL	

Inactive Ingredients		
Ingredient Name	Strength	
WATER (UNII: 059QF0KO0R)		
GLYCERIN (UNII: PDC6A3C0OX)		
CARBOMER HOMOPOLYMER, UNSPECIFIED TYPE (UNII: 0 A5MM307FC)		
TROLAMINE (UNII: 903K93S3TK)		

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:78287-160-01	29 mL in 1 BOTTLE; Type 0: Not a Combination Product	06/24/2020	
2	NDC:78287-160-02	60 mL in 1 BOTTLE; Type 0: Not a Combination Product	06/24/2020	
3	NDC:78287-160-03	80 mL in 1 BOTTLE; Type 0: Not a Combination Product	06/24/2020	
4	NDC:78287-160-04	100 mL in 1 BOTTLE; Type 0: Not a Combination Product	06/24/2020	
5	NDC:78287-160-05	250 mL in 1 BOTTLE; Type 0: Not a Combination Product	06/24/2020	
6	NDC:78287-160- 06	260 mL in 1 BOTTLE; Type 0: Not a Combination Product	06/24/2020	

7	NDC:78287-160-07	$300\ mL$ in 1 BOTTLE; Type 0: Not a Combination Product	06/24/2020	
8	NDC:78287-160- 08	500 mL in 1 BOTTLE; Type 0: Not a Combination Product	06/24/2020	
9	NDC:78287-160- 09	980 mL in 1 BOTTLE; Type 0: Not a Combination Product	06/24/2020	
10	NDC:78287-160-10	3785 mL in 1 BOTTLE; Type 0: Not a Combination Product	06/24/2020	
11	NDC:78287-160-11	5000 mL in 1 CAN; Type 0: Not a Combination Product	06/24/2020	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333A	06/24/2020	

Labeler - Swanrose, Inc. (117523391)

Revised: 6/2020 Swanrose, Inc.