HAND SANITIZER- alcohol aerosol, spray Cosmetic Specialty Labs, 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.

This is a hand sanitizer manufactured according to the Temporary Policy for Preparation of Certain Alcohol-Based Hand Sanitizer Products During the Public Health Emergency (CoViD-19); Guidance for Industry.

The hand sanitizer is manufactured using only the following United States Pharmacopoeia (USP) grade ingredients in the preparation of the product (percentage in final product formulation) consistent with World Health Organization (WHO) recommendations:

- a. Alcohol (ethanol) (USP or Food Chemical Codex (FCC) grade) (75%, volume/volume (v/v)) in an aqueous solution denatured according to Alcohol and Tobacco Tax and Trade Bureau regulations in 27 CFR part 20.
- b. Aloe Barbadenis Leaf Juice.
- c. Caprylyl Glycol.
- d. Glycerin.
- e. Phenoxyethanol.
- f. Polysorate-20.
- g. Purified Water.
- h. Tocopheryl Acetate.

The firm does not add other active or inactive ingredients. Different or additional ingredients may impact the quality and potency of the product.

Active Ingredient

SD Alcohol 40 75%

Antimicrobial

Use

Hand Sanitizer to help reduce bacteria on the skin.

Warnings

Flammable. Keep away from fire or flame. For external use only.

When using this product do not use in or near the eyes. In case of contact, rinse eyes thoroughtly with water.

Stop use and ask a doctor if irritation or rash appears and lasts.

Keep out of reach of children. If swallowed, get medical help or contact a Poison Center right away.

Directions

- Use enough product in your palm to cover hands and rub hands together biskly until dry.
- Children under 6 years of age should be supervised when using this product.

Other information

- Store below 110F (43C)
- May discolor fabrics or surfaces

Inactive ingredients

Aloe Barbadensis Leaf Juice, Caprylyl Glycol, Fragrance, Glycerin, Phenoxyethanol, Polysorbate-20, Purified Water, Tocopheryl Acetate.

Package Label - Principal Display Panel

	ND SANITI					
	or acrosol, spi	ay				
Pro	duct Inform	ation				
Prod	uct Type	ype HUMAN OTC DRUG Item Code (Source)		e (Source)	NDC:58133-959	
Rout	e of Administra					
Acti	ve Ingredie	at/Activ	e Mojety			
ACU	ve ingredier	Strength				
ALCO	HOL (UNII: 3K995	Basis of Strength	75 mL in 100 mL			
Inac	tive Ingredi	ents				
	Strength					
ALPH	A-TOCOPHEROL	ACETATE (UNII: 9E8X80D2L0)			
POLY	SORBATE 20 (UN	II: 7T1F30V	5YH)			
PHEN	IOXYETHANOL (U	NII: HIE492	ZZ3T)			
WATE	R (UNII: 059QF0K	00R)				
CAPR	YLYL GLYCOL (U	NII: 00YIU54	138U)			
GLYC	ERIN (UNII: PDC64	A3C0OX)				
ALOE	VERA LEAF (UNI	I: ZY81Z83H	10X)			
Pac	kaging					
#	Item Code		Package Description		Marketing Start Date	Marketing End Date
1 ND	C:58133-959-67	178 mL in	1 BOTTLE, PUMP; Type 1: Convenience	Kit of Co-Package	08/30/2021	
NA -		.				
	rketing In					
Ma	arketing Catego	-	Application Number or Monogr	aph Citation	Marketing Start Date	Marketing End Date
	OTC monograph not final		part333A		08/30/2021	

Labeler - Cosmetic Specialty Labs, Inc. (032973000)

Establishment							
Name	Address	ID/FEI	Business Operations				
Cosmetic Specialty Labs, Inc.		032973000	manufacture(58133-959) , label(58133-959) , pack(58133-959) , analysis(58133-959)				

Revised: 8/2021

Cosmetic Specialty Labs, Inc.