

SNUGZ HAND SANITIZING SPRY- hand sanitizer spray spray
SnugZ/USA, LLC

ZS - Hand Sanitizer Spray

Active Ingredient: Ethyl Alcohol 70% v/v

Antiseptic

Warnings: For external use only. Flammable, keep away from fire. Do not use on damaged or broken skin. Keep out of eyes. Rinse with water to remove. **Stop use and ask a doctor** if rash occurs. **Keep out of reach of children.** If product is swallowed, get medical help or contact a Poison Control Center right away.

Inactive Ingredients: Aloe Vera Gel, Aminomethyl Propanol, Carbomer, Clycerine, Water

Wet hands thoroughly with product and rub into skin until dry.

Helps reduce bacteria on skin.

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Drug Facts
Active Ingredients: Ethyl Alcohol 70% v/v.....Antiseptic
Uses: Helps reduce bacteria on skin.
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Directions: Wet hands thoroughly with product and rub into skin until dry.
Inactive Ingredients: Aloe Vera Gel, Aminomethyl Propanol, Carbomer, Glycerine, Water.
Other: Store below 110°F (43°C)
Questions? 1.800.611.4270. MADE IN THE USA. DISTRIBUTED BY: (COMPANY NAME) (COMPANY CITY, STATE ZIP)

1.0 fl. oz. (28 mL.) Unscented Hand Sanitizer

SNUGZ HAND SANITIZING SPRY

hand sanitizer spray spray

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:76309-451	
Route of Administration	TOPICAL			
Active Ingredient/Active Moiety				
Ingredient Name		Basis of Strength	Strength	
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)		ALCOHOL	70 mL in 100 mL	
Inactive Ingredients				
Ingredient Name			Strength	
WATER (UNII: 059QF0K00R)				
AMINOMETHYL PROPANEDIOL (UNII: CZ7BU4QZJZ)				
CARBOMER 940 (UNII: 4Q93RCW27E)				
ALOE VERA LEAF POLYSACCHARIDES (UNII: W21O437517)				
GLYCERIN (UNII: PDC6A3C0OX)				
Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:76309-451-71	28 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product	01/01/2019	
2	NDC:76309-451-10	10 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product	01/01/2019	
3	NDC:76309-451-18	18 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product	01/01/2019	
4	NDC:76309-451-03	3 mL in 1 BOTTLE, SPRAY; Type 0: Not a Combination Product	01/01/2019	
Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
OTC Monograph Drug	M003	01/01/2019		

Labeler - SnugZ/USA, LLC (615959228)

Registrant - SnugZ/USA, LLC (615959228)

Establishment

Name	Address	ID/FEI	Business Operations
SnugZ/USA, LLC		615959228	manufacture(76309-451)