

HAND SANITIZER, GLITTER FEVER- alcohol gel

Merci Handy Corporation

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.

Hand Sanitizer, Glitter Fever

Drug Facts

Active ingredient

Alcohol 67%

Purpose

Antiseptic

Use

- for handwashing to decrease bacteria on the skin

Warnings

For external use only. Flammable, keep away from fire or flame.

Do not use

in the eyes. In case of contact, flush eyes with water.

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.

Directions

wet hands thoroughly with product and allow to dry without wiping

Other information

store at a temperature below 110° F (43° C)

Inactive ingredients

Water (Aqua), Fragrance (Parfum), Aloe Vera Leaf Juice Powder, Glycerin, Propylene Glycol, Acrylates/C10-30 Alkyl Acrylate Crosspolymer, Aminomethyl Propanol, Mannitol, Microcrystalline Cellulose, Sucrose, Corn (Zea Mays) Starch, Denatonium Benzoate, Tocopheryl Acetate, Maltodextrin, Hydroxypropyl Methylcellulose, Potassium Sorbate, Sodium Benzoate, Fluorophlogopite, Coumarin, Limonene, Alpha-Isomethyl Ionone, Cinnamyl Alcohol, Isoeugenol, Titanium

Dioxide, FD&C Yellow No. 5, Iron Oxides FD&C Red No. 4.

QUESTIONS OR COMMENTS?

(646)-358-3432

Package Labeling:



HAND SANITIZER, GLITTER FEVER

alcohol gel

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:72866-009
Route of Administration	TOPICAL		
Active Ingredient/Active Moiety			
Ingredient Name		Basis of Strength	Strength
ALCOHOL (UNII: 3K9958V90M) (ALCOHOL - UNII:3K9958V90M)		ALCOHOL	670 mg in 1 mL
Inactive Ingredients			
Ingredient Name			Strength
WATER (UNII: 059QF0KO0R)			
ALOE VERA LEAF (UNII: ZY81Z83H0X)			
GLYCERIN (UNII: PDC6A3C0OX)			
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)			
CARBOMER INTERPOLYMER TYPE A (ALLYL SUCROSE CROSSLINKED) (UNII: 59TL3WG5CO)			
AMINOMETHYLPROPANOL (UNII: LU49E6626Q)			
MANNITOL (UNII: 3OWL53L36A)			
MICROCRYSTALLINE CELLULOSE (UNII: OP1R32D61U)			
SUCROSE (UNII: C151H8M554)			
CORN (UNII: 0N8672707O)			
DENATONIUM BENZOATE (UNII: 4YK5Z54AT2)			
.ALPHA.-TOCOPHEROL ACETATE (UNII: 9E8X80D2L0)			
MALTODEXTRIN (UNII: 7CVR7L4A2D)			
HYPROMELLOSE, UNSPECIFIED (UNII: 3NXW29V3WO)			
POTASSIUM SORBATE (UNII: 1VPU26JZZ4)			
SODIUM BENZOATE (UNII: OJ245FE5EU)			
COUMARIN (UNII: A4VZ22K1WT)			

LIMONENE, (+)- (UNII: GFD7C86Q1W)	
ISOMETHYL-.ALPHA.-IONONE (UNII: 9XP4LC555B)	
CINNAMYL ALCOHOL (UNII: SS8YOP444F)	
ISOEUGENOL (UNII: 5M0MWY797U)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	
FD&C YELLOW NO. 5 (UNII: I753WB2F1M)	
FERRIC OXIDE RED (UNII: 1K09F3G675)	
FD&C RED NO. 4 (UNII: X3W0AM1JLX)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:72866-009-30	30 mL in 1 BOTTLE; Type 0: Not a Combination Product	03/10/2020	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333E	03/01/2020	

Labeler - Merci Handy Corporation (116958007)

Revised: 3/2020

Merci Handy Corporation