DONE HAND ANTIBACTERIAL ALOE VERA- benzalkonium chloride liquid Olein Recovery 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.

Done Hand Soap Antibacterial Aloe Vera

Drug Facts

Active ingredient

Benzalkonium Chloride 0.13%

Purpose

Antibacterial

Use

for handwashing to decrease bacteria on ths skin.

Warnings

For external use only

When using this product,

avoid contact with eyes. In case of contact flush with water.

Stop use and ask a doctor if

irritation and redness develops.

Keep out of reach of children.

If swallowed, get medical help or contact a Poison Control Center right away.

Directions

wash skin, rinse and dry thoroughly

Inactive ingredients

purified water, sodium laureth sulfate, sodium C14-16 olefin sulfonate, cocamidorpyopyl betaine, sodium chloride, glycerine, fragrance, tetrasodium EDTA, methylchloroisothiazolinone, Methylisothiazolinone, D&C orange No.4

Package Labeling:









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Drug .	Facts
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DONE HAND ANTIBACTERIAL ALOE VERA

TOPICAL

benzalkonium chloride liquid

Product Information

Route of Administration

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:77142-008
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Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
BENZALKO NIUM CHLO RIDE (UNII: F5UM2KM3W7) (BENZALKO NIUM - UNII:7N6 JUD5X6 Y)	BENZALKONIUM CHLORIDE	1.3 mg in 1 mL

Inactive Ingredients		
Ingredient Name	Strength	
WATER (UNII: 059QF0KO0R)		
SODIUM LAURETH SULFATE (UNII: BPV390 UAPO)		
SODIUM C14-16 OLEFIN SULFONATE (UNII: O9W3D3YF5U)		
COCAMIDO PRO PYL BETAINE (UNII: 50 CF30 11KX)		
SO DIUM CHLO RIDE (UNII: 451W47IQ8X)		
GLYCERIN (UNII: PDC6A3C0OX)		
EDETATE SO DIUM (UNII: MP1J8420 LU)		
METHYLCHLORO ISO THIAZO LINO NE (UNII: DEL7T5QRPN)		
METHYLISOTHIAZOLINONE (UNII: 229 D0 E1QFA)		
D&C ORANGE NO. 4 (UNII: Q1LIY3BO0U)		

l	Packaging				
ı	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
ı	1	NDC:77142-008-00	296 mL in 1 BOTTLE; Type 0: Not a Combination Product	05/15/2020	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part333E	05/15/2020	

Labeler - Olein Recovery Corporation (188543446)

Revised: 5/2020 Olein Recovery Corporation