ANTI-BACTERIAL FRESH ALOE HAND - triclos an soap BB17, LLC

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if the comply with applicable regulations and policies. FDA has not evaluated whether this product complies.				
Active ingredient: Triclosan 0.3%				
Purpose Antimicrobial				
KILLS MORE THAN 99.99% OF COMMON GERMS				
FOR EXTERNAL USE ONLY. DO NOT USE IN THE EYES.				
DISCONTINUE USE IF IRRITATION AND REDNESS DEVELOP. IF CONDITION PERSISTS FOR MORE THAN 72 HOURS, CONSULT A DOCTOR.				
Keep out of reach of children. CHILDREN CAN ONLY USE THIS PRODUCT WITH ADULT SUPERVISION.				
DIRECTIONS: WET HANDS.APPLY PALMFUL TO HANDS. SCRUB THOROUGHLY. RINISE. RECOMMENDED FOR REPEATED USE.				

INACTIVE INGREDIENTS:Water, Sodium Alkyl Ether Sulphate, Ammonium Lauryl Sulphate, Cocamidopropyl Betaine, Cocamide DEA, Disodium Laureth Sulfosuccinate, Sodium Chloride, Glycol Distearate, Fragrance, Methylchloroisothiazolinone, Methylisothiazolinone, Citric Acid, Disodium EDTA, FD&C Yellow NO. 5, FD&C Blue NO. 1



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DRUG FACTS

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ANTI-BACTERIAL FRESH ALOE HAND

triclosan soap

Product Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:53603-2015

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient NameBasis of StrengthStrengthTRICLO SAN (UNII: 4NM5039 Y5X) (TRICLOSAN - UNII:4NM5039 Y5X)TRICLOSAN0.3 g in 100 mL

Inactive Ingredients Ingredient Name Strength WATER (UNII: 059QF0KO0R) SODIUM C12-15 PARETH-3 SULFATE (UNII: 19Q4RW8UWP) AMMONIUM LAURYL SULFATE (UNII: Q7AO2R1M0B) COCAMIDO PRO PYL BETAINE (UNII: 5OCF3O11KX) COCO DIETHANOLAMIDE (UNII: 92005F972D) DISODIUM LAURETH SULFOSUCCINATE (UNII: D6 DH1DTN7E) SODIUM CHLORIDE (UNII: 451W47IQ8 X) GLYCOL DISTEARATE (UNII: 13W7MDN21W) METHYLCHLORO ISOTHIAZOLINO NE (UNII: DEL7T5QRPN)

METHYLISOTHIAZOLINONE (UNII: 229 D0 E1QFA)	
CITRIC ACID MO NO HYDRATE (UNII: 2968 PHW8 QP)	
EDETATE DISO DIUM (UNII: 7FLD91C86K)	
FD&C YELLOW NO. 5 (UNII: I753WB2F1M)	
FD&C BLUE NO. 1 (UNII: H3R47K3TBD)	

P	Packaging						
#	Item Code	Package Description	Marketing Start Date	Marketing End Date			
1	NDC:53603-2015-8	236.6 mL in 1 BOTTLE, SPRAY					

Marketing Information						
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date			
OTC monograph not final	part333E	08/20/2013				

Labeler - BB17, LLC (828378294)

Revised: 8/2013 BB17, LLC