ANTIBACTERIAL FOAMING CUCUMBER MELON- triclosan liquid MEIJER DISTRIBUTION 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.

DRUG FACTS

ACTIVE INGREDIENT

TRICLOSAN 0.6%

PURPOSE

ANTIBACTERIAL

USES

HELPS REDUCE BACTERIA ON THE SYSTEM

WARNINGS

FOR EXTERNAL USE ONLY

WHEN USING THIS PRODUCT

AVOID CONTACT WITH EYES. IF CONTACT OCCURS, RINSE EYES THOROUGHLY WITH WATER.

STOP USING THIS PRODUCT AND ASK DOCTOR

IF IRRITATION AND REDNESS DEVELOP AND LAST

KEEP OUT OF REACH OF CHILDREN

IN CASE OF ACCIDENTAL INGESTION, GET MEDICAL HELP OR CONTACT POISON CONTROL CENTER IMMEDIATELY.

DIRECTIONS

PUMP ONTO DRY HANDS, WORK INTO A LATHER AND RINSE THOROUGHLY.

OTHER INFORMATION

STORE AT ROOM TEMPERATURE

INACTIVE INGREDIENTS

WATER (AQUA), SODIUM LAURETH SULFATE, DIPROPYLENE GLYCOL, SODIUM XYLENESULFONATE, COCAMIDOPROPYL BETAINE, FRAGRANCE (PARFUM), POLYSORBATE 20, DISODIUM PHOSPHATE, CITRIC ACID, METHYLISOTHIAZOLINOE, METHYLISOTHIAZOLINONE, BLUE 1 (CI 42090), YELLOW 5 (CI 19140), RED 40 (CI 16035)

LABEL COPY



ANTIBACTERIAL FOAMING CUCUMBER MELON

triclosan liquid

Product Information					
Product Type	HUMAN OTC DRUG	Item Code	NDC:41250-187		
Route of Administration	TOPICAL				
Active Ingredient/Active M	Ioiety				
Ing	redient Name		Basis of Strength	Strength	
TRICLOSAN (UNII: 4NM5039Y5X) (TRICLOSAN - UNII:4NM5039Y5X) TRICLOSAN 0.					
TRICLOSAN (UNII: 4NM5039Y5X)					
				0.6 mL in 100 mL	
	(TRICLOSAN - UNII:4NM5039Y5X)			0.6 mL in 100 mL	
Inactive Ingredients	(TRICLOSAN - UNII:4NM5039Y5X) Ingredient Name			0.6 mL in 100 mL	
Inactive Ingredients WATER (UNII: 059QF0KO0R) SODIUM LAURETH SULFATE (UM	(TRICLOSAN - UNII:4NM5039Y5X) Ingredient Name NII: BPV390UAP0)			0.6 mL in 100 mL	
Inactive Ingredients WATER (UNII: 059QF0KO0R) SODIUM LAURETH SULFATE (UM	(TRICLOSAN - UNII:4NM5039Y5X) Ingredient Name NII: BPV390UAP0) 107L85C40)			0.6 mL in 100 mL	
Inactive Ingredients WATER (UNII: 059QF0K00R) SODIUM LAURETH SULFATE (UN DIPROPYLENE GLYCOL (UNII: E	(TRICLOSAN - UNII:4NM5039Y5X) Ingredient Name NII: BPV390UAP0) 107L85C40) INII: G4LZF950UR)		<u> </u>	0.6 mL in 100 mL	

SOD	IUM PHO SPHATE (U	NII: 5E33/5VY3/)							
CITR	IC ACID MONOHYD	RATE (UNII: 2968PHW8QP)							
MET	HYLCHLOROISOTH	IAZOLINONE (UNII: DEL7T5QRPN)							
MET	HYLISOTHIAZOLIN	ONE (UNII: 229 D0 E1QFA)							
Packaging									
#	Item Code	Package Description	Marketing Start Date N		Ma	larketing End Date			
1 ND	C:41250-187-08	221 mL in 1 BOTTLE, PUMP							
Ma	rketing Inform	mation							
Ma	rketing Category	Application Number or Monogra	ph Citation	Marketing Start	Date	Marketing End Dat			
отс	monograph not final	part333E		11/17/2011					
		-							

Labeler - MEIJER DIST RIBUTION INC (006959555)

Registrant - APOLLO HEALTH AND BEAUTY CARE (201901209)

Establishment

Name	Address	ID/FEI	Business Operations
APOLLO HEALTH AND BEAUTY CARE		201901209	manufacture

Revised: 11/2011

MEIJER DISTRIBUTION INC