EQUATE- selenium sulfide liquid WAL-MART STORES 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

SELENIUM SULFIDE 1%

PURPOSE

ANTI-DANDRUFF

USES

FOR RELIEF OF FLAKING AND ITCHING ASSOCIATED WITH DANDRUFF AND SEBORRHEIC DERMATITIS AND TO HELP PREVENT THE CHANCE OF RECURRENCE.

WARNINGS

FOR EXTERNAL USE ONLY.

ASK A DOCTOR BEFORE USING IF YOU HAVE

SEBORRHEIC DERMATITIS IN AREAS OTHER THAN THE SCALP.

WHEN USING THIS PRODUCT

AVOID CONTACT WITH EYES. IF PRODUCT GETS INTO EYES, RINSE THOROUGHLY WITH WATER. FOR USE ON COLOR TREATED OR PERMED HAIR, RINSE THOROUGHLY.

STOP USING THIS PRODUCT AND ASK A DOCTOR IF

CONDITION WORSENS OR DOES NOT IMPROVE AFTER REGULAR USE OF THIS PRODUCT AS DIRECTED.

KEEP OUT OF REACH OF CHILDREN

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

DIRECTIONS

SHAKE WELL, APPLY SHAMPOO, RINSE THOROUGHLY. FOR BEST RESULTS USE AT LEAST TWICE A WEEK OR AS DIRECTED BY A DOCTOR.

OTHER INFORMATION

STORE AT ROOM TEMPERATURE.

INACTIVE INGREDIENTS

WATER (AQUA), SODIUM LAURETH SULFATE, ACRYLATES COPOLYMER, TEA-LAURYL SULFATE, COCAMIDOPROPYL BETAINE, COCAMIDOPROPYL HYDROXYSULTAINE, CITRIC ACID, FRAGRANCE (PARFUM), AMMONIUM CHLORIDE, DMDM HYDANTOIN, MENTHOL, SODIUM HYDROXIDE, MAGNESIUM ALUMINUM SILICATE, HYDROXYPROPYL METHYLCELLULOSE, BLUE 1 (CI 42090), RED 33 (CI 17200).

QUESTIONS?

CALL: 1-800-287-1915

LABEL COPY





selenium sulfide liquid

Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:49035-620	
Route of Administration	TOPICAL			

Active Ingredient/Active Moiety					
Ingredient Name	Basis of Strength	Strength			
SELENIUM SULFIDE (UNII: Z69 D9 E38 1Q) (SELENIUM SULFIDE - UNII: Z69 D9 E38 1Q)	SELENIUM SULFIDE	10 mg in 1 mL			

Inactive Ingredients	
Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
SO DIUM LAURETH SULFATE (UNII: BPV390 UAP0)	
METHACRYLIC ACID - METHYL METHACRYLATE COPOLYMER (1:1) (UNII: 74G4R6TH13)	
TEA-LAURYL SULFATE (UNII: E8458C1KAA)	
COCAMIDO PRO PYL BETAINE (UNII: 50 CF30 11KX)	
COCAMIDO PRO PYL HYDRO XYSULTAINE (UNII: 62V75NI93W)	
CITRIC ACID MO NO HYDRATE (UNII: 2968 PHW8 QP)	
AMMO NIUM CHLO RIDE (UNII: 01Q9 PC255D)	
DMDM HYDANTO IN (UNII: BYR0 546 TOW)	
MENTHOL (UNII: L7T10EIP3A)	
SO DIUM HYDRO XIDE (UNII: 55X04QC32I)	
MAGNESIUM ALUMINUM SILICATE (UNII: 6 M3P6 4V0 NC)	
HYPROMELLOSES (UNII: 3NXW29V3WO)	
FD&C BLUE NO. 1 (UNII: H3R47K3TBD)	
D&C RED NO. 33 (UNII: 9DBA0SBB0L)	

P	ackaging			
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:49035-620-11	325 mL in 1 BOTTLE, PLASTIC		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph final	part358H	06/21/2013	

Labeler - WAL-MART STORES INC. (051957769)

Registrant - APOLLO HEALTH AND BEAUTY CARE (201901209)

Establishment

Name	Address	ID/FEI	Business Operations
APOLLO HEALTH AND BEAUTY CARE		201901209	manufacture(49035-620)

Revised: 6/2013 WAL-MART STORES INC.