RITE AID RENEWAL- selenium sulfide liquid RITE AID 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.

DRUG FACTS

ACTIVE INGREDIENT

SELENIUM SULFIDE 1%

PURPOSE

ANTI-DANDRUFF

USES

FOR RELIEF OF FLAKING AND ITCHING DUE TO 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 THE EYES. IF CONTACT OCCURS, RINSE EYES THOROUGHLY WITH WATER. FOR USE ON COLOR-TREATED OR PERMED HAIR, RINSE THOROUGHLY.

STOP USE 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

IF SWALLOWED, GET MEDICAL HELP OR CONTACT A POISON CONTROL CENTER RIGHT AWAY.

DIRECTIONS

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

QUESTIONS/COMMENTS?

1-866-695-3030

INACTIVE INGREDIENTS:

WATER (AQUA), SODIUM LAURETH SULFATE, DIHYDROGENATED TALLOW PHTHALIC ACID AMIDE, SODIUM CHLORIDE, TITANIUM DIOXIDE, COCAMIDOPROPYL BETAINE, SODIUM STEAROYL LACTYLATE, FRAGRANCE (PARFUM), DIMETHICONE, CITRIC ACID, DMDM HYDANTOIN, SODIUM CITRATE, ALOE BARBADENSIS LEAF JUICE, HYDROXYPROPYL METHYLCELLULOSE, BLUE 1 (CI 42090).

LABEL COPY



RITE AID RENEWAL

selenium sulfide liquid

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Product	Inform	ation
Promici		411011

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:11822-6211
110ddct 1 ypc	IIONEIN OTO DICO	nem code (Source)	1120111011

Route of Administration TOPICAL

Active Ingredient/Active Moiety

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	Ingredient Name	Basis of Strength	Strength
	SELENIUM SULFIDE (UNII: Z69D9E381Q) (SELENIUM SULFIDE - UNII:Z69D9E381Q)	SELENIUM SULFIDE	10 mg in 1 mL

Inactive Ingredients Ingredient Name Strength

 $\textbf{WATER} \; (\text{UNII:} \; 0\,59\,\text{QF0}\,\text{KO0}\,\text{R})$

SODIUM LAURETH SULFATE (UNII: BPV390 UAP0)

DIHYDRO GENATED TALLO W PHTHALIC ACID AMIDE (UNII: 1R8 1RPY10 G)

SODIUM CHLORIDE (UNII: 451W47IQ8X)

TITANIUM DIO XIDE (UNII: 15FIX9 V2JP)

COCAMIDO PRO PYL BETAINE (UNII: 50 CF3 O 11 KX)

SODIUM STEAROYL LACTYLATE (UNII: IN99IT31LN)

DIMETHICO NE (UNII: 92RU3N3Y10)

CITRIC ACID MONOHYDRATE (UNII: 2968 PHW8 QP)

DMDM HYDANTO IN (UNII: BYR0546 TOW)

SODIUM CITRATE (UNII: 1Q73Q2JULR)

ALOE VERA LEAF (UNII: ZY81Z83H0X)

HYPROMELLOSES (UNII: 3NXW29V3WO)

FD&C BLUE NO. 1 (UNII: H3R47K3TBD)

Packaging

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ı	# Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	1 NDC:11822-6211-1	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	06/20/2013	

Labeler - RITE AID CORPORATION (014578892)

Registrant - APOLLO HEALTH AND BEAUTY CARE (201901209)

Establishment

Name	Address	ID/FEI	Business Operations
APOLLO HEALTH AND BEAUTY CARE		201901209	manufacture(11822-6211)

Revised: 6/2013 RITE AID CORPORATION