# CARE ONE MEDICATED- selenium sulfide liquid AMERICAN SALES COMPANY

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.

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# **DRUG FACTS**

# **ACTIVE INGREDIENT**

SELENIUM SULFIDE 1%

#### PURPOSE

ANTIDANDRUFF

#### USE

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.

# **OTHER INFORMATION**

STORE AT ROOM TEMPERATURE.

# **INACTIVE INGREDIENTS:**

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

# **QUESTIONS/COMMENTS?**

1-866-695-3030

# LABEL COPY



# CARE ONE MEDICATED

selenium sulfide liquid

Product Informat	1011						
Product Type	HUMAN OTC DRUGItem Code (Source)		NDC:41520-620				
Route of Administra	tion	TOPICAL					
Active Ingredient	Active Moi	ety					
Ingredient Name Basis of Strengt						Strength	
SELENIUM SULFIDE (UNII: Z69D9E381Q) (SELENIUM SULFIDE - UNII:Z69D9E381Q) SELENIUM SULFIDE							
Inactive Ingredie	nts						
		Ingredient Na	ame			Strength	
WATER (UNII: 059QF0	KO0R)						
SODIUM LAURETH S	ULFATE (UNII: H	3PV390UAP0)					
METHACRYLIC ACID	- METHYL MET	HACRYLATE COPOLY	<b>MER (1:1)</b> (UN	II: 74G4R6TH13)			
TEA-LAURYL SULFA	<b>ΓΕ</b> (UNII: E8458	C1KAA)					
COCAMIDOPROPYL	BETAINE (UNII:	50CF3011KX)					
CITRIC ACID MONOF	IYDRATE (UNII:	2968PHW8QP)					
AMMONIUM CHLORI	<b>DE</b> (UNII: 01Q9F	C255D)					
SO DIUM HYDRO XIDE	(UNII: 55X04Q0	C32I)					
DMDM HYDANTO IN (	UNII: BYR0546T	OW)					
MAGNESIUM ALUMIN	<b>UM SILICATE</b>	(UNII: 6 M3P6 4 V0 NC)					
HYPROMELLOSES (U	NII: 3NXW29V3	WO)					
MENTHOL (UNII: L7T1	0EIP3A)						
FD&C BLUE NO.1 (U							
D&C RED NO. 33 (UN	II: 9 DBA0 SBB0 I	.)					
Packaging							
# Item Code	Pac	kage Description	Market	ing Start Date	Marketin	Marketing End Date	
1 NDC:41520-620-11	325 mL in 1	BOTTLE, PLASTIC					
Marketing Info	ormation						
8		N 1 NC	nh Citatian	Marketing Start	Data Marka	ting End Date	
Marketing Category	Applicatio	n Number or Monogra	ph Chauon	Markeung Start	Date Maine	ung Enu Date	

# Labeler - AMERICAN SALES COMPANY (809183973)

# **Registrant** - APOLLO HEALTH AND BEAUTY CARE (201901209)

Name	Address	ID/FEI	Business Operations
APOLLO HEALTH AND BEAUTY CARE		201901209	manufacture(41520-620)

Revised: 7/2013