DAYLOGIC DANDRUFF CLASSIC CLEAN- pyrithione zinc 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

Pyrithione Zinc 1%

Purpose

Anti-dandruff

Uses

to help prevent recurrence of flaking and itching associated with dandruff.

Warnings

For external use only.

When using this product

avoid contact with eyes. If contact occurs, rinse eyes thoroughly with water.

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.

In case of accidental ingestion, get medical help or contact a Poison Control Center immediately.

Directions

- for maximum dandruff control, use every time you shampoo
- wet hair, massage onto scalp and rinse
- repeat if desired.

Inactive ingredients

Water (Aqua), Sodium Laureth Sulfate, Sodium Chloride, Acrylates Copolymer, Glycol Distearate, Cocamidopropyl Betaine, Cocamide MEA, Laureth-4, Fragrance (Parfum), Sodium Hydroxide, Tetrasodium EDTA, Methylchloroisothiazolinone, Methylisothiazolinone, Blue 1 (CI 42090), Red 33 (CI 17200).

Label Copy



DAYLOGIC DANDRUFF CLASSIC CLEAN

pyrithione zinc liquid

Product Information								
Product Type	HUMAN OTC DRUG	Item Code (Sour	ce) NDC:118	322-4272				
Route of Administration	TOPICAL							
Active Ingredient/Active M	Ioiety							
0	Ioiety Ingredient Name		Basis of Strength	Strength				
Active Ingredient/Active M PYRITHIONE ZINC (UNII: R95302	Ingredient Name	R953O2RHZ5)	Basis of Strength PYRITHIONE ZINC	Streng				

Ingredient Name						
WATER (UNII: 059QF0KO0R)						
SO DIUM LAURETH	SULFATE (UNII: BPV390UA	?0)				
SODIUM CHLORID	E (UNII: 451W47IQ8X)					
METHACRYLIC AC	D - METHYL METHACRYLA	TE COPOLYMER (1:1) (UNII:	74G4R6TH13)			
GLYCOL DISTEARATE (UNII: 13W7MDN21W)						
CO CAMIDO PRO PY	L BETAINE (UNII: 50CF3011	KX)				
COCO MONOETHA	NOLAMIDE (UNII: C8068414	46 D)				
SO DIUM HYDRO XI	DE (UNII: 55X04QC32I)					
EDETATE SO DIUM (UNII: MP1J8420LU)						
METHYLCHLOROI	GOTHIAZOLINONE (UNII: D	EL7T5QRPN)				
METHYLISOTHIAZOLINONE (UNII: 229 D0 E1QFA)						
METHYLISOTHIAZ	DLINONE (UNII: 229 D0 E1QF	(A)				
		A)				
FD&C BLUE NO. 1 (D&C RED NO. 33 (U	UNII: H3R47K3TBD)	A)				
FD&C BLUE NO.1(UNII: H3R47K3TBD)	A)				
FD&C BLUE NO.1(UNII: H3R47K3TBD)	A)				
FD&C BLUE NO. 1 (D&C RED NO. 33 (U	UNII: H3R47K3TBD) NII: 9DBA0SBB0L)	A) Description	Marketing Start Date	Marketing End Date		
FD&C BLUE NO. 1 (D&C RED NO. 33 (U Packaging # Item Code	UNII: H3R47K3TBD) NII: 9DBA0SBB0L) Package		-	<u> </u>		
FD&C BLUE NO. 1 (D&C RED NO. 33 (U Packaging Item Code NDC:11822-4272-	UNII: H3R47K3TBD) NII: 9DBA0SBB0L) Package 701 mL in 1 BOTTLE, PLASTI	Description	Date	<u> </u>		
D&C BLUE NO. 1 (D&C RED NO. 33 (U Packaging Item Code NDC:11822-4272-	UNII: H3R47K3TBD) NII: 9DBA0SBB0L) Package 701 mL in 1 BOTTLE, PLASTI	Description	Date	<u> </u>		
FD&C BLUE NO. 1 (D&C RED NO. 33 (U Packaging Item Code NDC:11822-4272-3	UNII: H3R47K3TBD) NII: 9DBA0SBB0L) Package 701 mL in 1 BOTTLE, PLASTI Product	Description	Date	<u> </u>		
FD&C BLUE NO. 1 (D&C RED NO. 33 (U Packaging Item Code NDC:11822-4272-	UNII: H3R47K3TBD) NII: 9DBA0SBB0L) Package 701 mL in 1 BOTTLE, PLAST Product formation	Description C; Type 0: Not a Combination	Date	<u> </u>		

Labeler - Rite Aid Corporation (014578892)

Registrant - Apollo Health and Beauty Care Inc. (201901209)

Establishment						
Name	Address	ID/FEI	Business Operations			
Apollo Health and Beauty Care Inc.		201901209	manufacture(11822-4272)			

Revised: 5/2017

Rite Aid Corporation