SMART SENSE ICEBERG BLUE- eucalyptol, menthol, methyl salicylate, thymol liquid Kmart 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 INGREDIENTS

Eucalyptol 0.092%, Menthol 0.042%, Methyl Salicylate 0.060%, Thymol 0.064%

PURPOSE

Antiplaque/Antigingivitis

USES

To help reduce and prevent plaque and gingivitis

WARNINGS

Do not use for children under 12 years of age.

Keep out of reach of children

If more than used for rinsing accidently swallowed, get medical help or contact a Poison Control Center immediately (1-800-222-1222)

Directions

Rinse full strength for 30 seconds with 20 mL (2/3 fluid ounce) or 4 teaspoons ful) mornings and night. Do not swallow

OTHER INFORMATION

Store at room temperature. Cold weather may cloud this product. Its antiseptic properties are not affected.

INACTIVE INGREDIENTS

Water (Aqua), Alcohol (21.6%), Sorbitol Solution, Flavor, Poloxamer 407, Benzoic Acid, Zinc Chloride, Sodium Benzoate, Sucralose, Sodium Saccharin, Blue 1 (CI 42090)

Questions or comments?

1-800-842-7886

Label Copy



	Drug Facts Active ingredients Euolypin 0.092% Menthol 0.042% Methyl 3.16/24% Methyl 3.16/24% Thymol 0.064%
	Uses m To help reduce and prevent plaque and ginglyitts.
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SMART SENSE ICEBERG BLUE

eucalyptol, menthol, methyl salicylate, thymol liquid

Product Information					
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:49738-551		
Route of Administration	ORAL				

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
EUCALYPTOL (UNII: RV6J6604TK) (EUCALYPTOL - UNII:RV6J6604TK)	EUCALYPTOL	0.92 mg in 1 mL		
MENTHOL (UNII: L7T10EIP3A) (MENTHOL - UNII:L7T10EIP3A)	MENTHOL	0.42 mg in 1 mL		
METHYL SALICYLATE (UNII: LAV5U5022Y) (SALICYLIC ACID - UNII:O414PZ4LPZ)	METHYL SALICYLATE	0.6 mg in 1 mL		
THYMOL (UNII: 3J50XA376E) (THYMOL - UNII:3J50XA376E)	THYMOL	0.64 mg in 1 mL		

Inactive Ingredients				
Ingredient Name	Strength			
WATER (UNII: 059QF0KO0R)				
ALCOHOL (UNII: 3K9958V90M)				
SORBITOL (UNII: 506T60A25R)				
POLOXAMER 407 (UNII: TUF2IVW3M2)				
BENZOIC ACID (UNII: 8 SKN0 B0 MIM)				
ZINC CHLORIDE (UNII: 86Q357L16B)				
SODIUM BENZOATE (UNII: OJ245FE5EU)				
SUCRALOSE (UNII: 96K6UQ3ZD4)				
SACCHARIN SODIUM (UNII: SB8ZUX40TY)				
FD&C BLUE NO. 1 (UNII: H3R47K3TBD)				

Packaging						
#	Item Code	Package Description	Market	ing Start Date	Market	ing End Date
1	NDC:49738-551-34	1000 mL in 1 BOTTLE, PLASTIC				
Marketing Information						
I	Marketing Category	Application Number or Monograph	Citation	Marketing Start Da	ite Marl	keting End Date

04/22/2013

Labeler - Kmart Corporation (008965873)

OTC monograph not final part356

Registrant - Apollo Health and Beauty Care (201901209)

Establishment				
Name	Address	ID/FEI	Business Operations	
Apollo Health and Beauty Care		201901209	manufacture (49 738 - 551)	

Revised: 4/2013 Kmart Corporation