

DHS SAL 4 OZ.- dhs sal shampoo
Person and Covey

DHS SAL

WARNINGS AND PRECAUTIONS

For external use only. Ask a physician before use if you have Psoriasis or Seborrheic Dermatitis that covers a large area of the body. Do not get into eyes. If contact occurs, rinse eyes thoroughly with water. Stop use and ask a physician if condition worsens or does not improve after regular use.

Dosage and Administration

Wet hair thoroughly. Apply a liberal amount of shampoo and massage into a rich lather. Allow lather to remain on scalp for several minutes. Rinse hair well and repeat application. For best results use at least twice a week or as directed by a physician. Store away from direct sunlight.

Indications and Use

Controls crusty, flaky buildup, while relieving scalp itching and flaking, symptomatic of Psoriasis and Seborrheic Dermatitis.

If swallowed, get medical help or contact a Poison Control Center right away.

PURPOSE

Controls symptoms of Psoriasis, Seborrheic Dermatitis and Dandruff

INACTIVE INGREDIENTS

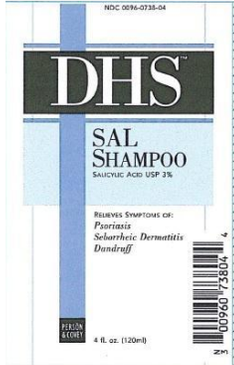
Purified Water
Sodium C14-16 Olefin Sulfonate
Triethanolamine Lauryl Sulfate
Cocamidopropyl Betaine
Cocamidopropyl Hydroxysultaine
PEG 8 Distearate

Active Ingredient

Salicylic Acid 35 mg/g

PACKAGE LABEL.PRINCIPAL DISPLAY PANEL

DHS Sal Shampoo.jpg



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dhs sal shampoo

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:0096-0738
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SALICYLIC ACID (UNII: O414PZ4LPZ) (SALICYLIC ACID - UNII:O414PZ4LPZ)	SALICYLIC ACID	0.035 g in 1 g

Inactive Ingredients

Ingredient Name	Strength
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WATER (UNII: 059QF0KO0R)	
SODIUM C14-16 OLEFIN SULFONATE (UNII: O9W3D3YF5U)	
TRIETHANOLAMINE LAURYL SULFATE (UNII: E8458C1KAA)	
COCAMIDOPROPYL BETAINE (UNII: 5OCF3O11KX)	
COCAMIDOPROPYL HYDROXYSULTAINE (UNII: 62V75NI93W)	
PEG-8 DISTEARATE (UNII: 7JNC8VN07M)	

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0096-0738-04	123.84 g in 1 BOTTLE; Type 0: Not a Combination Product	06/01/1995	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M032	06/01/1995	

Labeler - Person and Covey (008482473)

Establishment			
Name	Address	ID/FEI	Business Operations
Person and Covey		008482473	manufacture(0096-0738)

Revised: 1/2024

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