HEAD AND SHOULDERS CLINICAL STRENGTH ADVANCED OIL CONTROLselenium sulfide shampoo The Procter & Gamble Manufacturing Company

Head and Shoulders ®

Clinical Strength Advanced Oil Control Shampoo

Drug Facts

Active ingredient

Selenium Sulfide 1%

Purpose

Anti-dandruff, anti-seborrheic dermatitis

Uses

helps prevent recurrence of flaking, itching, irritation, scaling and redness associated with dandruff and seborrheic dermatitis.

Warnings

For external use only.

Ask a doctor before use if you have a condition that covers a large area of the body.

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 this and all drugs out of reach of children. If swallowed, get medical help or contact a Poison Control Center right away.

Directions

- for best results use <u>at least</u> twice a week or as directed by a doctor.
- for maximum dandruff control, use every time you shampoo.
- shake before use.
- wet hair, massage onto scalp, rinse, repeat if desired.
- caution: if used on bleached, tinted, grey, or permed hair, rinse for 5 minutes.

Inactive ingredients

Water, sodium lauryl sulfate, glycol distearate, sodium chloride, fragrance,

cocamidopropyl betaine, sodiumcitrate, sodium xylenesulfonate, dimethicone, citric acid, sodium benzoate, tetrasodium EDTA

Questions (or comments)?

1-800-723-9569

Dist. by PROCTER & GAMBLE, CINCINNATI, OH 45202

PRINCIPAL DISPLAY PANEL - BOTTLE LABEL

head & shoulders ®

selenium sulfide dandruff & seborrheic dermatitis shampoo Clinical Strength Advanced Oil Control 13.5 FL OZ (400 mL)



HEAD AND SHOULDERS CLINICAL STRENGTH ADVANCED OIL CONTROL

selenium sulfide shampoo

Product Information

Product Type

HUMAN OTC DRUG

Item Code (Source)

NDC:84126-075

Route of Admi	nistration	TOPICAL					
Active Ingree	dient/Active	e Moiety					
Ingredient Name				Basis of Strength Str		Strength	
SELENIUM SULFIDE (UNII: Z69D9E381Q) (SELENIUM SULFIDE - UNII:Z69D9E381Q)			: -	SELENIUM SULFIDE		g n 100 mL	
0MII:269D9E381Q)		IN 100 M		n 100 mL		
Inactive Ingr	redients						
		Ingredient Name			Str	ength	
TETRASODIUM E	DTA (UNII: MP1	J8420LU)					
WATER (UNII: 059	QF0KO0R)						
SODIUM LAURYL	SULFATE (UNI	l: 368GB5141J)					
GLYCOL DISTEARATE (UNII: 13W7MDN21W)							
SODIUM CHLORIDE (UNII: 451W47IQ8X)							
SODIUM XYLENESULFONATE (UNII: G4LZF950UR)							
DIMETHICONE (UNII: 92RU3N3Y1O)							
SODIUM BENZOATE (UNII: OJ245FE5EU)							
SODIUM CITRATE (UNII: 1Q73Q2JULR)							
CITRIC ACID (UNII: 2968PHW8QP)							
COCAMIDOPROP	PYL BETAINE (U	JNII: 50CF3011KX)					
Packaging							
# Item Code	F	Package Description	Marl	ceting Start Date		eting End Date	
1 NDC:84126- 075-40	400 mL in 1 BC Combination P	OTTLE, PLASTIC; Type 0: Not roduct	a 11/01/	2024			
2 NDC:84126- 075-01	2 in 1 CARTON		11/01/	2024			
2	400 mL in 1 BC Combination P	OTTLE, PLASTIC; Type 0: Not roduct	а				

Marketing Information

Category Citation Date	art Marketing End Date
OTC Monograph Drug M032 11/01/2024	

Labeler - The Procter & Gamble Manufacturing Company (004238200)

Revised: 1/2025

The Procter & Gamble Manufacturing Company