

SULFO LO- sulfur 3% soap bar soap
Crown Laboratories

Sulfo-Lo Cleansing Bar Soap

Active ingredients

Sulfur 3%

Purpose

Acne Treatment

Uses

Deep cleansing antibacterial soap for management of acne, blackheads and comedones.

Warnings

- **For external use only** • Avoid contact with eyes, wash with water if exposed
- **Do no use on**
 - broken skin • large areas of skin
- **When using this product**
 - apply only to areas with acne • skin irritation and dryness is more likely to occur if you use another topical acne medication at the same time. • If irritation occurs, only use one topical acne medication at a time • If undue irritation persists consult a physician
- **Keep out of reach of children.**
 - If swallowed, get medical help or contact a Poison Control Center.

Directions

- Work into a lather and rinse with warm water • Use daily as skin cleanser or as directed by a physician.

Other information

Store at 20° -25°C (68° -77°F) [see USP Controlled Room Temperature].

Inactive ingredients

Sodium Tallowate, Sodium Cocoate/Sodium Palm Kernelate, Water (Aqua), Glycerin,

Fragrance (Parfum), Sodium Chloride, Butyrospermum parkii (Shea) Butter, Yellow 5 (CI19140), Titanium Dioxide (CI77891), Pentasodium Pentetate, Tetrasodium Etidronate, Iron Oxides (CI77492), Tocopheryl Acetate

NDC 0316-0118-35

sulfo lo®

Cleansing Bar Soap

Helps Treat and Manage Acne, Blackheads and Comedones

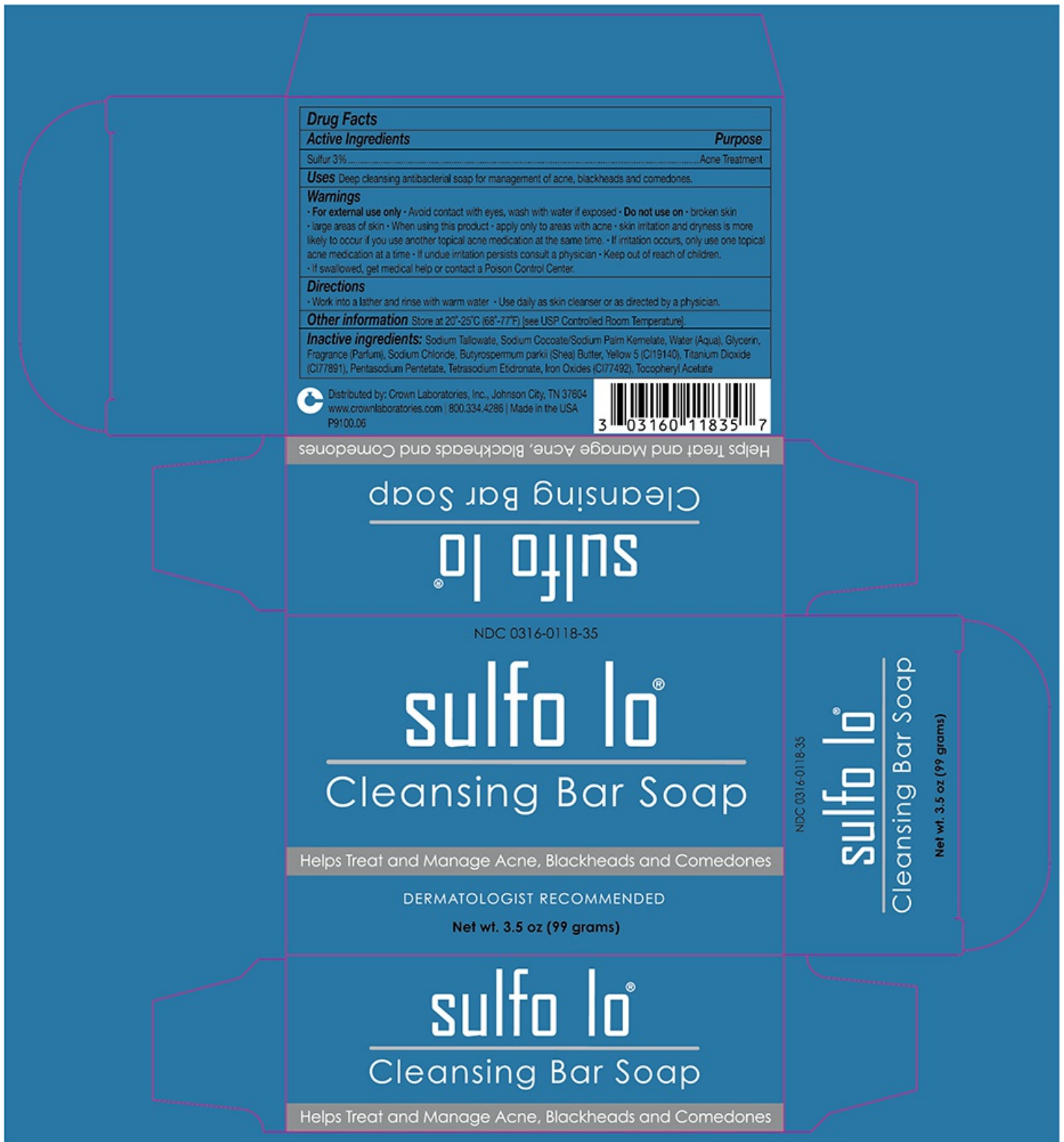
DERMATOLOGIST RECOMMENDED

Net wt. 3.5oz (99 grams)

Distributed by: Crown Laboratories, Inc., Johnson City, TN 37604

www.crownlaboratories.com | 800.334.4286 | Made in the USA

P9100.06



Drug Facts

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SULFO LO
 sulfur 3% soap bar soap

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:0316-0118
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SULFUR (UNII: 70FD1KFU70) (SULFUR - UNII:70FD1KFU70)	SULFUR	30 mg in 1 g

Inactive Ingredients

Ingredient Name	Strength
.ALPHA.-TOCOPHEROL ACETATE, D- (UNII: A7E6112E4N)	
SODIUM TALLOWATE, BEEF (UNII: 07RIK6QMEW)	
SODIUM COCOATE (UNII: R1TQH25F4I)	
SODIUM PALM KERNELATE (UNII: 6H91L1NXTW)	
WATER (UNII: 059QF0KO0R)	
GLYCERIN (UNII: PDC6A3C0OX)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
PENTASODIUM PENTETATE (UNII: 961TOZ5L7T)	
ETIDRONATE TETRASODIUM (UNII: CZZ9T1T1X4)	
BROWN IRON OXIDE (UNII: 1N032N7MFO)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	
SHEA BUTTER (UNII: K49155WL9Y)	
BASIC YELLOW 5 (UNII: 07BP340B4T)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0316-0118-35	99 g in 1 CARTON; Type 0: Not a Combination Product	11/01/1965	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M006	11/01/1965	

Labeler - Crown Laboratories (079035945)**Registrant** - Crown Laboratories (079035945)

Revised: 11/2023

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