

**BLACKHEAD SOLUTIONS 7 DAY DEEP PORE CLEANSE AND SCRUB ACNE
MEDICATION- salicylic acid cream
CLINIQUE LABORATORIES LLC**

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.

blackhead solutions 7 day deep pore cleanse and scrub acne medication

Drug Facts

Active ingredient

Salicylic Acid 1.0%

Purpose

Acne Treatment

Uses

- reduces the number of blackheads
- helps keep skin clear of new blackheads
- helps prevent development of new blackheads

Warnings

For external use only

When using this product skin irritation and dryness are more likely to occur if you use another topical acne medication at the same time. If irritation occurs, only one topical acne medication should be used unless directed by a doctor.

Keep out of reach of children. If product is swallowed, get medical help or contact a Poison Control Center right away. Keep out of eyes.

Directions

- Apply to face daily, wet and massage, rinse off.

Inactive ingredients

water\aquaeau • kaolin • glycerin • butylene glycol • magnesium aluminum silicate • glyceryl stearate • peg-100 stearate • silica • sodium lauroyl sarcosinate • allyl methacrylates crosspolymer • gentiana lutea (gentian) root extract • laminaria saccharina extract • sucrose • acetyl glucosamine • caprylyl glycol • 1,2-hexanediol • tocopheryl acetate • sodium hydroxide • xanthan gum • disodium edta • phenoxyethanol • titanium dioxide (ci 77891) • blue 1 (ci 42090) • yellow 5 (ci 19140)

[iln43442]

PRINCIPAL DISPLAY PANEL - 15 ml Bottle Carton

CLINIQUE

blackhead
solutions

7 day deep pore
cleanser & scrub

SALICYLIC ACID
ACNE MEDICATION

ALL SKIN TYPES

.5 FL.OZ.LIQ./15 ml e

007873

CULTECH INC
CAD# 2103592© CLINIQUE LABORATORIES, DIST.
NEW YORK, N.Y. 10022
WLB LTD. GU32 3DD, UK • PARIS
MADE IN U.S.A. ZRON-40

clinique.com

Drug Facts (continued)**Warnings** For external use only

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Drug Facts (continued)**Directions**

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**Drug Facts** (continued)

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CLINIQUE
blackhead solutions7 day deep pore
cleanse & scrubSALICYLIC ACID
ACNE MEDICATION

ALL SKIN TYPES

.5 FL.OZ./15 ml **Drug Facts****Active ingredient**

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Purpose

Acne Treatment

Uses • reduces the number of blackheads

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ALL SKIN TYPES**Allergy Tested. 100% Fragrance Free.**

Multi-functional pore-clearing formula effectively treats and prevents blackheads. Use daily to cleanse & scrub and twice a week as a 5-minute mask.

Not For Sale

ZRON-40-1111

BLACKHEAD SOLUTIONS 7 DAY DEEP PORE CLEANSE AND SCRUB ACNE MEDICATION

salicylic acid cream

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:49527-062
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SALICYLIC ACID (UNII: O414PZ4LPZ) (SALICYLIC ACID - UNII:O414PZ4LPZ)	SALICYLIC ACID	10 mg in 1 mL

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
KAOLIN (UNII: 24H4NWX5CO)	
GLYCERIN (UNII: PDC6A3C0OX)	
BUTYLENE GLYCOL (UNII: 3XUS85K0RA)	
MAGNESIUM ALUMINUM SILICATE (UNII: 6M3P64V0NC)	
GLYCERYL MONOSTEARATE (UNII: 230OU9XXE4)	
PEG-100 STEARATE (UNII: YD01N1999R)	
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
SODIUM LAUROYL SARCOSINATE (UNII: 632GS99618)	
SUCROSE (UNII: C151H8M554)	
N-ACETYLGLUCOSAMINE (UNII: V956696549)	
CAPRYLYL GLYCOL (UNII: 00YIU5438U)	
1,2-HEXANEDIOL (UNII: TR046Y3K1G)	
SODIUM HYDROXIDE (UNII: 55X04QC32I)	
XANTHAN GUM (UNII: TTV12P4NEE)	
EDETATE DISODIUM ANHYDROUS (UNII: 8NLQ36F6MM)	
PHENOXYETHANOL (UNII: HIE492ZZ3T)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	
FD&C BLUE NO. 1 (UNII: H3R47K3TBD)	
FD&C YELLOW NO. 5 (UNII: I753WB2F1M)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:49527-062-01	1 in 1 CARTON	08/30/2018	11/30/2022
1		20 mL in 1 BOTTLE; Type 0: Not a Combination Product		
2	NDC:49527-062-02	1 in 1 CARTON	08/01/2022	

2	125 mL in 1 TUBE; Type 0: Not a Combination Product		
Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph final	M006	08/30/2018	

Labeler - CLINIQUE LABORATORIES LLC (044475127)

Registrant - Estee Lauder Companies Inc. (790802086)

Establishment

Name	Address	ID/FEI	Business Operations
PALC		078364654	pack(49527-062) , label(49527-062)

Establishment

Name	Address	ID/FEI	Business Operations
Estee Lauder Cosmetics Ltd.		204132062	pack(49527-062) , label(49527-062)

Establishment

Name	Address	ID/FEI	Business Operations
Estee Lauder N.V.		370151326	manufacture(49527-062) , pack(49527-062) , label(49527-062)

Establishment

Name	Address	ID/FEI	Business Operations
The Estee Lauder Inc		802599436	manufacture(49527-062) , pack(49527-062) , label(49527-062)

Establishment

Name	Address	ID/FEI	Business Operations
Northtec LLC		943871157	pack(49527-062) , label(49527-062)

Establishment

Name	Address	ID/FEI	Business Operations
PADC 1		949264774	pack(49527-062) , label(49527-062)