

SO TOTALLY CLEAN DEEP PORE CLEANSER- salicylic acid liquid
ASPIRE BRANDS INC

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.

ASPIRE BRANDS - SO TOTALLY CLEAN DEEP PORE CLEANSER (70626-104)

ACTIVE INGREDIENTS

SALICYLIC ACID 0.5%

PURPOSE

ACNE TREATMENT

USE

ACNE TREATMENT TO HELP CLEAR UP AND PREVENT NEW ACNE BLEMISHES, BLACKHEADS, AND WHITEHEADS.

WARNINGS

- FOR EXTERNAL USE ONLY.
- FLAMMABLE - KEEP AWAY FROM OPEN FIRE OR FLAME.

WHEN USING THIS PRODUCT

- AVOID CONTACT WITH EYES. IF CONTACT OCCURS, FLUSH THOROUGHLY WITH WATER.
- 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.

KEEP OUT OF REACH OF CHILDREN. IF SWALLOWED, GET MEDICAL HELP OR CONTACT A POISON CONTROL CENTER IMMEDIATELY.

DIRECTIONS

- CLEAN THE SKIN THOROUGHLY BEFORE APPLYING THIS PRODUCT.
- MOISTEN A COTTON BALL. COVER THE ENTIRE AFFECTED AREA WITH A THIN LAYER ONE TO THREE TIMES DAILY.
- BECAUSE EXCESSIVE DRYING OF THE SKIN MAY OCCUR, START WITH ONE APPLICATION DAILY, THEN GRADUALLY INCREASE TO TWO OR THREE TIMES DAILY IF NEEDED OR AS DIRECTED BY A DOCTOR.
- IF BOTHERSOME DRYNESS OR PEELING OCCURS, REDUCE APPLICATION TO ONCE A DAY OR EVERY OTHER DAY.

INACTIVE INGREDIENTS

ALCOHOL DENAT., WATER, SODIUM PCA, LACTIC ACID, FRAGRANCE, ALLANTOIN, SODIUM CARBONATE, RED 4, YELLOW 5, YELLOW 6.



FORMULA 10.0.6[®]

**SO
TOTALLY
CLEAN**

**DEEP PORE CLEANSER
ORIGINAL FORMULA
WITH SALICYLIC ACID
ACNE TREATMENT**

HELPS CLEAR ACNE PIMPLES,
BLACKHEADS AND WHITEHEADS.
HELPS PREVENT THE DEVELOPMENT
OF NEW ACNE BLEMISHES.

2.0 fl oz (60 mL)

DISTRIBUTED BY:
ASPIRE BRANDS
WESTLAKE OH 44145
1-800-843-3636
WWW.FORMULA10.0.6.COM

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Drug Facts

ACTIVE INGREDIENTS	PURPOSE
Salicylic Acid 0.5%	Acne treatment

USE:
• Acne treatment to help clear up and prevent new acne blemishes, blackheads, and whiteheads.

WARNINGS:
• For external use only.
• Flammable: keep away from open fire or flame.

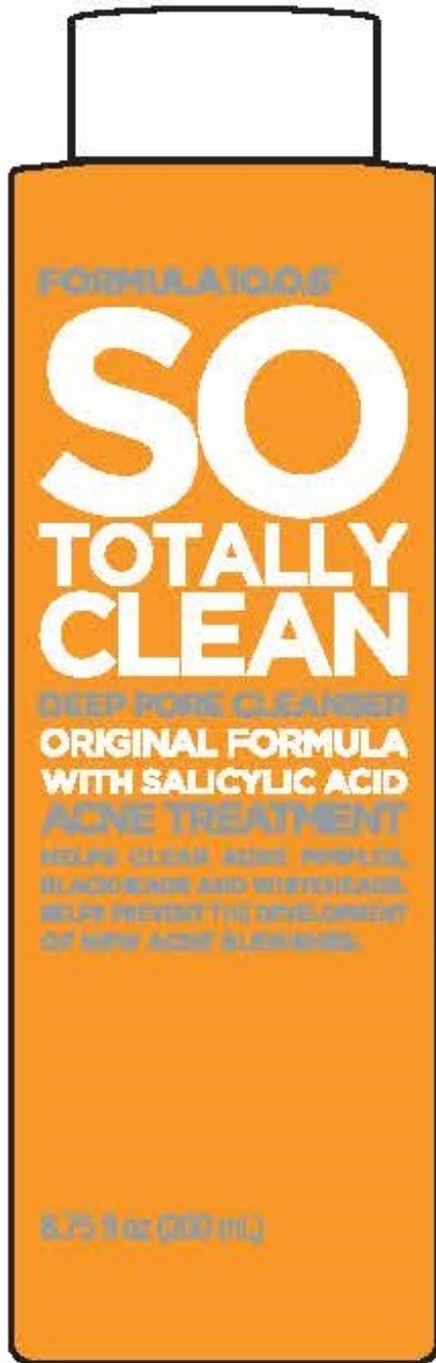
When using this product:
• Avoid contact with eyes. If contact occurs, flush thoroughly with water.
• Skin irritation and dryness is more likely to occur if you use another topical acne medication at the same time. If irritation occurs, only use one topical acne medication at a time.

Keep out of the reach of children. If swallowed, get medical help or contact a Poison Control Center immediately.

DIRECTIONS:
• Cleanse the skin thoroughly before applying.
• Moisten a cotton ball. Cover the entire affected area with a thin layer one to three times daily.
• Because excessive drying of the skin may occur, start with one application daily, then gradually increase to two or three times daily if needed or as directed by a doctor.
• If bothersome dryness or peeling occurs, reduce application to once a day or every other day.

INACTIVE INGREDIENTS: ALCOHOL DENAT., WATER, SODIUM PCA, LACTIC ACID, FRAGRANCE, ALLANTOIN, SODIUM CARBONATE, RED 4, YELLOW 5, YELLOW 6. NMS-14

QUESTIONS/COMMENTS: 1-866-984-1433



SO TOTALLY CLEAN DEEP PORE CLEANSER

salicylic acid liquid

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:70626-104
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SALICYLIC ACID (UNII: O414PZ4LPZ) (SALICYLIC ACID - UNII:O414PZ4LPZ)	SALICYLIC ACID	0.5 g in 100 mL

Inactive Ingredients

Ingredient Name	Strength
ALCOHOL (UNII: 3K9958V90M)	
WATER (UNII: 059QF0KO0R)	
SODIUM PYRROLIDONE CARBOXYLATE (UNII: 469OTG57A2)	
LACTIC ACID (UNII: 33X04XA5AT)	
ALLANTOIN (UNII: 344S277G0Z)	
SODIUM CARBONATE (UNII: 45P3261C7T)	
FD&C RED NO. 4 (UNII: X3W0AM1JLX)	
FD&C YELLOW NO. 5 (UNII: I753WB2F1M)	
FD&C YELLOW NO. 6 (UNII: H77VEI93A8)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:70626-104-21	200 mL in 1 BOTTLE; Type 0: Not a Combination Product	05/02/2016	
2	NDC:70626-104-22	60 mL in 1 BOTTLE; Type 0: Not a Combination Product	05/17/2017	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph final	part333D	05/02/2016	

Labeler - ASPIRE BRANDS INC (832462811)

Revised: 10/2023

ASPIRE BRANDS INC