

DRS. ACNE SPOT TREATMENT- salicylic acid gel
OL PHARMA TECH, LLC Drs. PHARMACY

ACTIVE INGREDIENT

Salicylic Acid 2% w/w.

PURPOSE

Acne treatment

USES

For the treatment of acne. Dries and clears acne pimples, blackheads and whiteheads and allows skin to heal.

DO NOT USE

if you have sensitive skin or are sensitive to salicylic acid

DIRECTIONS

Morning and evening, after a thorough cleansing of the skin, apply acne clear gel locally on cutaneous imperfections. Then apply the usual day or night cream. Renew application 1 to 3 times daily.

WARNINGS

For external use only. Using other topical acne medications at the same time or immediately following use of this product may increase dryness or irritation of the skin. If this occurs, only one medication should be used unless directed by a doctor. Avoid direct contact with the eyes. If product gets into the eyes, rinse liberally with water. Discontinue use if skin irritation develops or increases. If irritation persists, consult a doctor.

WHEN USING THIS PRODUCT

Avoid contact with eyes, lips, and mouth.

KEEP OUT OF REACH OF CHILDREN

if swallowed, get medical help or contact poison control center right away

carbomer interpolymers type A, trolamine, vitamin E, propylene glycol, water, isopropyl alcohol, methyl paraben, EDTA, propyl paraben, DMDM hydantoin

OTHER INFORMATION

- store at 15-30 C (59-86 F)
- close cap tightly after use
- keep away from heat

QUESTIONS

www.drsparmacyusa.com

PACKAGE LABEL



	Client Name - DRS Pharmacy USA Job Name - 100221_Acne Gel_DRs_PHARMACY_USA_28.3g 22 dia 134 mm (+/-2mm) tube length	Artwork's Data Received from Client BRAND NAME: JOB TITLE: FILE NAME: DATE: PROJECT: JOB NO:
	Cyan Black Pantone 381 c Pan 186 c Pantone Reflex Blue C	White Substrate

DRS. ACNE SPOT TREATMENT

salicylic acid gel

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:80489-357
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SALICYLIC ACID (UNII: O414PZ4LPZ) (SALICYLIC ACID - UNII:O414PZ4LPZ)	SALICYLIC ACID	20 mg in 1 g

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	
.ALPHA.-TOCOPHEROL (UNII: H4N855PNZ1)	
METHYLPARABEN (UNII: A2I8C7HI9T)	
EDETATE DISODIUM (UNII: 7FLD91C86K)	
TROLAMINE (UNII: 9O3K93S3TK)	
ISOPROPYL ALCOHOL (UNII: ND2M416302)	
CARBOMER INTERPOLYMER TYPE A (55000 CPS) (UNII: 59TL3WG5CO)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:80489-357-01	1 in 1 CARTON	04/13/2024	
1		20 g in 1 TUBE, WTH APPLICATOR; Type 0: Not a Combination Product		
2	NDC:80489-357-02	1 in 1 CARTON	04/13/2024	
2		30 g in 1 TUBE, WTH APPLICATOR; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M006	02/01/2022	

Labeler - OL PHARMA TECH, LLC Drs. PHARMACY (021170377)

Registrant - OL PHARMA TECH, LLC Drs PHARMACY (021170377)

Establishment

Name	Address	ID/FEI	Business Operations
OL PHARMA TECH, LLC Drs PHARMACY		021170377	manufacture(80489-357)

Revised: 4/2024

OL PHARMA TECH, LLC Drs. PHARMACY