

ACROPASS ACNE CURE- acropass acne cure patch
RAPHAS CO., LTD

Active Ingredients Purpose

Salicylic Acid 2%..... Acne Treatment

Drug Facts	
Active Ingredients	Purpose
Salicylic Acid 2%	Acne Treatment
Uses For the treatment of acnes	
Warnings For external use only	
When using this product · Skin irritation and dryness may occur if you use another topical acne medication at the same time or immediately following. · If irritation occurs, only use one topical acne medication at a time unless directed by doctor.	
Stop use and ask a doctor If irritation becomes severe.	
Keep out of reach of children If swallowed, get medical help or contact a Poison Control Center right away.	
Directions 1. Open the packaging and take out the patch. 2. Carefully remove the white film, ensuring not to touch the center of the patch. 3. Apply the patch directly to the affected area and press vertically with your fingers. 4. Leave on for 2 hours or overnight, then remove.	
Other information Store at room temperature. Avoid use if you have very sensitive skin or are allergic to Salicylic acid.	
Inactive Ingredients Sodium Hyaluronate, Niacinamide, Oligopeptide-10, Phytosphingosine HCl Patch : Hydrocolloid	

Uses: For the treatment of acnes

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Sodium Hyaluronate, Niacinamide, Oligopeptide-10, Phytosphingosine HCl

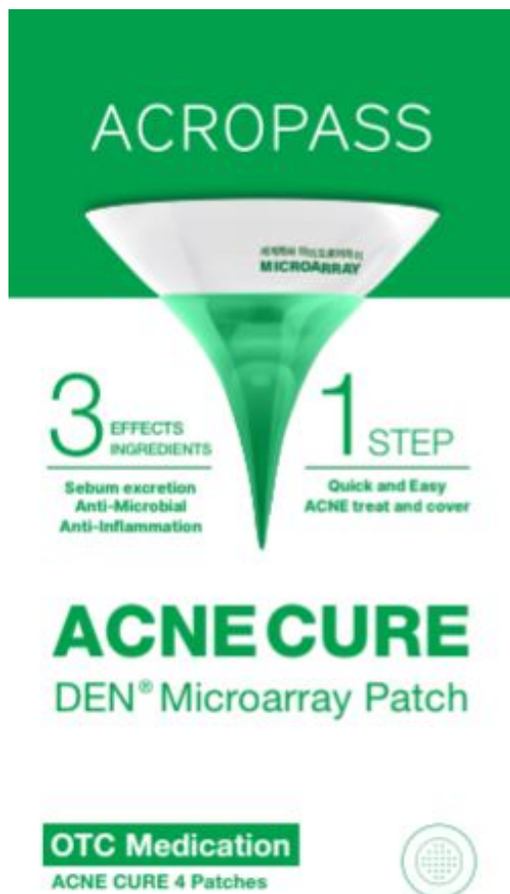
Patch : Hydrocolloid

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Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:71134-241
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
SALICYLIC ACID (UNII: O414PZ4LPZ) (SALICYLIC ACID - UNII:O414PZ4LPZ)	SALICYLIC ACID	0.012 mg in 0.6013 mg

Inactive Ingredients

Ingredient Name	Strength
PHYTOSPHINGOSINE HYDROCHLORIDE (UNII: TT871XV7TU)	
NIACINAMIDE (UNII: 25X51I8RD4)	
HYALURONATE SODIUM (UNII: YSE9PPT4TH)	
OLIGOPEPTIDE-10 (UNII: Q46328TRNK)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:71134-241-01	4 mg in 1 BOX; Type 0: Not a Combination Product	08/28/2024	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M006	08/28/2024	

Labeler - RAPHAS CO., LTD (689054529)

Establishment

Name	Address	ID/FEI	Business Operations
RAPHAS CO.,LTD		695914964	manufacture(71134-241)

Revised: 8/2024

RAPHAS CO., LTD