CURITHYS- sodium hyaluronate patch J World

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

Drug Facts

Sodium hyaluronate

Niacinamide

Salicylic Acid

Sh-Oligopepide-1

Sh-Polypepide-1

TetraPeptide-44

Centella Asiatica Extract

Tranexamic acid

Camellia Japonica Flower Extract

Micro-pyramid stimulates the skin and gives moisture and nutrition to sensitive skin.

keep out of reach of the children

Remove the protective film from the patch.

Attach it vertically to the desired site.

Re-press on thea ttached patch every few minutes..

Remove the patch after attaching for more than 1 hour.

Use it for your own use only.

For external use only.

Discontinue use if adverse reaction occurs.

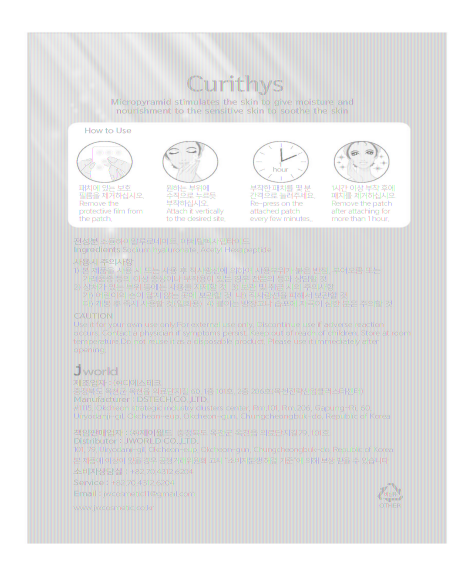
Contact a physician if symptoms persist.

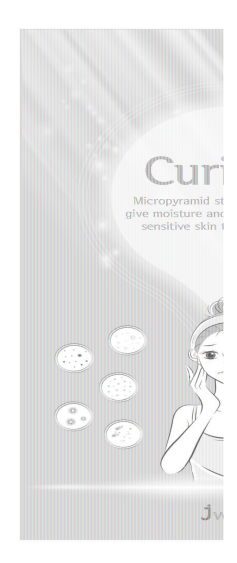
Store at room temperature

.Do not reuse it as a disposable product.

Please use it immediately after opening.

for topical use only





CURITHYS

sodium hyaluronate patch

Product Information Product Type HUMAN OTC DRUG Item Code (Source) NDC:72945-0002 Route of Administration TOPICAL

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
HYALURONATE SODIUM (UNII: YSE9PPT4TH) (HYALURONIC ACID - UNII:S270N0TRQY)	HYALURONATE SODIUM	94.39 mg in 100 mg

Inactive Ingredients		
Ingredient Name	Strength	
NIACINAMIDE (UNII: 25X5118RD4)		

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:72945- 0002-2	6 in 1 POUCH	07/21/2019	
1	NDC:72945- 0002-1	5 mg in 1 PATCH; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		07/21/2019	
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Labeler - J World (690082781)

Registrant - J World (690082781)

Establishment			
Name	Address	ID/FEI	Business Operations
J World		690082781	label(72945-0002) , manufacture(72945-0002)

Revised: 12/2021 J World