INTOMEDI AMPLANCE HYDROCREAM- adenosine cream Janytree Inc.

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.

INACTIVE INGREDIENTS

water, arginine, tocopherol etc.

ACTIVE INGREDIENTS

adenosine

intensive hydrating ampoule for healthy, balanced skin

WARNINGS

For external use only When using this product

■ If following abnormal symptoms occurs after using the product, stop using the product and consult with a skin specialist. Symptoms : Red specks, swelling, itching

■ Do not use on the skin parts affected by wound, eczema, or dermatitis. Keep out of reach of children.

■ If swallowed, get a medical help or contact a person in control center immediately.

■ Avoid contact with eyes.

KEEP OUT OF REACH OF CHILDREN

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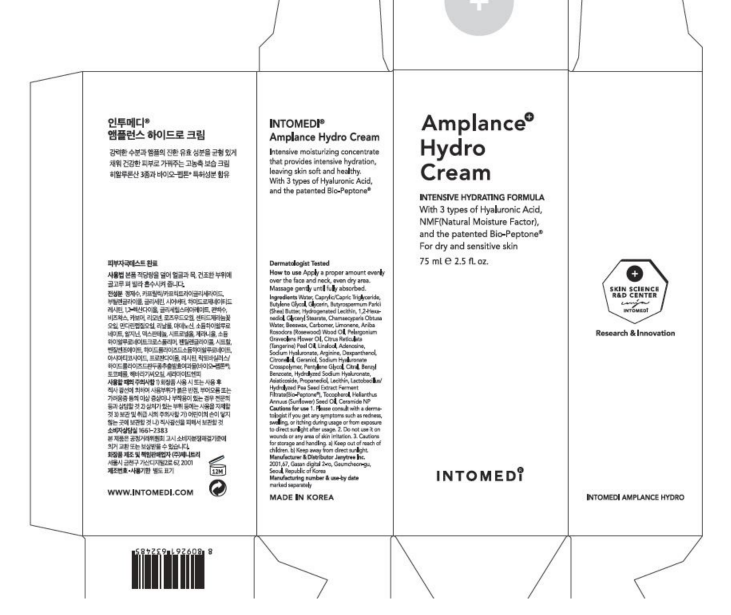
If swallowed, get a medical help or contact a person in control center immediately.

apply a properamount to the skin, and massage gently until fully absorbed

for external use only

Amplance^o Hydro

Cream



INTOMEDI AMPLANCE HYDROCREAM								
adenosine cream								
Product Information								
Product Type	HUMAN OTC DRUG	Item Code (Source)		NDC	NDC:82879-0002			
Route of Administration	TOPICAL							
Active Ingredient/Active	Moiety							
Active Ingredient/Active Moiety								
Ingredient Name			Basis of Strength		Strength			
ADENOSINE (UNII: K72T3FS567) (ADENOSINE - UNII:K72T3FS567)			ADENOSINE		0.1 g in 100 mL			

In	active Ingre	dients						
Ingredient Name					Strength			
W	WATER (UNII: 059QF0KO0R)							
AR	ARGININE (UNII: 94Z LA3W45F)							
Packaging								
#	ltem Code	Package Description	Marketing Date	Start	Marketing End Date			
1	NDC:82879- 0002-1	75 mL in 1 TUBE; Type 0: Not a Combination Product	07/13/2022					
Marketing Information								
	Marketing Category	Application Number or Monograph Citation	Marketing Date		Marketing End Date			
	approved drug ner		07/13/2022					

Labeler - Janytree Inc. (688403840)

Registrant - Janytree Inc. (688403840)

Establishment									
Name	Address	ID/FEI	Business Operations						
Janytree Inc.		688403840	manufacture(82879-0002)						

Revised: 7/2022

Janytree Inc.