ARMIYOU HYDROGEN TONER- adenosine liquid Armiyou

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.

ACTIVE INGREDIENT

Active ingredients: Adenosine 0.04%

INACTIVE INGREDIENT

Inactive ingredients: Water, Butylene Glycol, Glycerin, Alcohol, PEG-60 Hydrogenated Castor Oil, Phenoxyethanol, Allantoin, Carbomer, Panthenol, Sodium Benzoate, Triethanolamine, Flavor, Centella Asiatica Extract, Ethylhexylglycerin, Polygonum Cuspidatum Root Extract, Tocopheryl Acetate, Scutellaria Baicalensis Root Extract, Polyquaternium-51, Camellia Sinensis Leaf Extract, Glycyrrhiza Glabra (Licorice) Root Extract, Rosmarinus Officinalis (Rosemary) Leaf Extract, Anthemis Nobilis Flower Extract, Sodium Hyaluronate, Pueraria Mirifica Root Extract

PURPOSE

Purpose: Anti wrinkle

WARNINGS

Warnings: For external use only Avoid contact with eyes. Discontinue use if signs of irritation or rashes appear. Replace the cap after use. Keep Out of Reach of Children. Avoid use in areas affected by wounds.

KEEP OUT OF REACH OF CHILDREN

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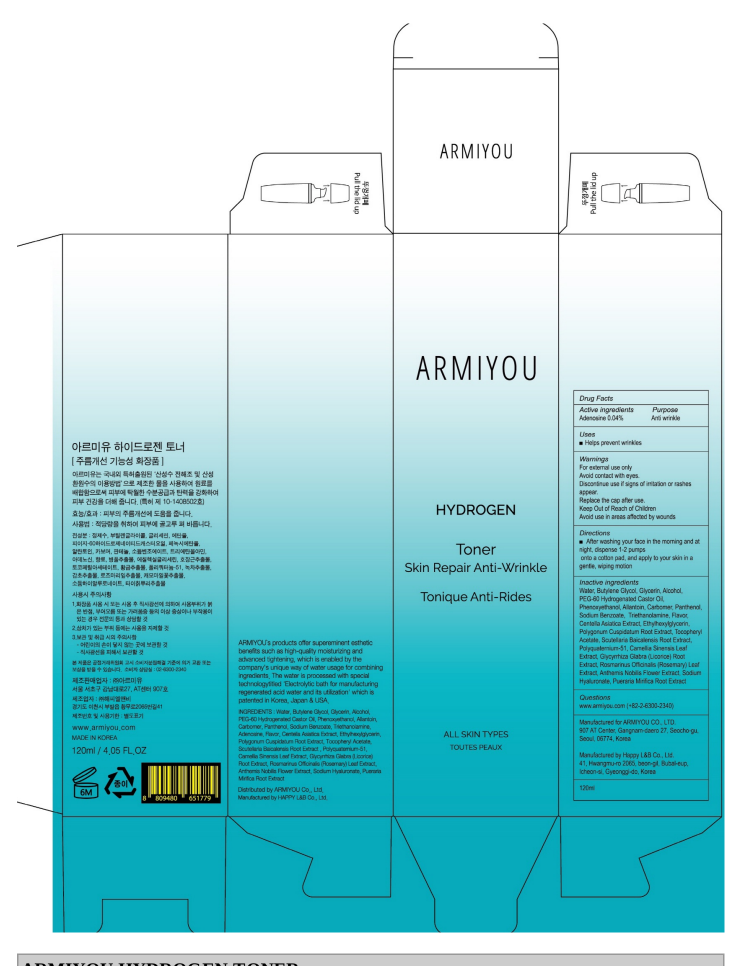
Uses

Uses: Helps prevent wrinkles

Directions

Directions: After washing your face in the morning and at night, dispense 1-2 pumps onto a cotton pad, and apply to your skin in a gentle, wiping motion.

PACKAGE LABEL.PRINCIPAL DISPLAY PANEL



ARMIYOU HYDROGEN TONER

adenosine liquid

Product Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:71878-010

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
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Adenosine (UNII: K72T3FS567) (ADENOSINE - UNII:K72T3FS567)
Adenosine 0.04 g in 120 mL

Inactive Ingredients

Ingredient Name	Strength

Water (UNII: 059QF0KO0R)

Butylene Glycol (UNII: 3XUS85K0RA)

Packaging

#	# Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:71878-010- 02	1 in 1 CARTON	10 /0 1/20 17	
1	NDC:71878-010- 01	120 mL in 1 CONTAINER; Type 0: Not a Combination Product		

Marketing Information

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Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		10 /0 1/20 17	

Labeler - Armiyou (694891814)

Registrant - Armiyou (694891814)

Establishment			
Name	Address	ID/FEI	Business Operations
Armiyou		694891814	manufacture(71878-010)

Revised: 11/2017 Armiyou