# CELLEXOSOME SB- niacinamide liquid PROSTEMICS Co., Ltd.

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.

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### ACTIVE INGREDIENT

Active ingredients: Niacinamide 3.0%

## INACTIVE INGREDIENT

Inactive ingredients:

[Powder] HUMAN ADIPOSE DERIVED STEM CELL CONDITIONED MEDIA, Panax Ginseng Root Extract

[Solvent] Water, Tranexamic Acid, Water, 1,2-Hexanediol, Hydroxyacetophenone, Ascorbyl Glucoside, Sodium Hyaluronate

#### **PURPOSE**

Purpose: Skin Brightening

#### WARNINGS

Warnings:

For external use only

- 1. Discontinue use if signs of irritation or rashes appear. If symptoms get worse, consult with a dermatologist. 1) In case of swelling, itching, or other side effects while or after using this product
- 2. Do not apply to open wounds.
- 3. Avoid contact with eyes.

Storage and handling

- 4. Replace the cap after use
- 5. Keep out of reach of children.
- 6. Avoid direct sunlight.

## KEEP OUT OF REACH OF CHILDREN SECTION

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Uses:

Helps brighten skin tone.

#### **Directions**

Directions:

■ Put solvent into the powder ampoule and shake gently enough to dissolve the mixture

■ Take proper amount and gently apply onto the skin

## PACKAGE LABEL.PRINCIPAL DISPLAY PANEL



## **CELLEXOSOME SB**

niacinamide liquid

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Product Type HUMAN OTC DRUG Item Code (Source) NDC:62041-280

Route of Administration TOPICAL

## **Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
Niacinamide (UNII: 25X5118RD4) (NIACINAMIDE - UNII:25X5118RD4)	Niacinamide	0.09 g in 3 mL

Inactive Ingredients			
Ingredient Name	Strength		
Water (UNII: 059QF0KO0R)			
Tranexamic Acid (UNII: 6T84R30KC1)			

	Packaging			
l	# Item Code	Package Description	<b>Marketing Start Date</b>	<b>Marketing End Date</b>
l	1 NDC:62041-280-02	5 in 1 CARTON	02/01/2020	
l	1 NDC:62041-280-01	$3\ mL$ in $1\ CONTAINER;$ Type $0$ : Not a Combination Product		

## **Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		02/01/2020	

# Labeler - PROSTEMICS Co., Ltd. (689605919)

## Registrant - PROSTEMICS Co., Ltd. (689605919)

Establishment				
Name	Address	ID/FEI	Business Operations	
Prostemics Co., Ltd. Factory		695687674	manufacture(62041-280)	

Revised: 2/2020 PROSTEMICS Co., Ltd.