

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](#).

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

KEEP OUT OF REACH OF CHILDREN SECTION

Uses

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

Product Information

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: 25X5118 RD4) (NIACINAMIDE - UNII:25X5118 RD4)	Niacinamide	0.09 g in 3 mL

Inactive Ingredients

Ingredient Name	Strength
Water (UNII: 059QF0KO0R)	
Tranexamic Acid (UNII: 6T84R30 KC1)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:62041-280-02	5 in 1 CARTON	02/01/2020	
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.