DRUGGIST CREAM- cream cream Shenzhen Xiaomai Manufacturing Co., Ltd.

Active ingredients

Arbutin

Purpose

Anti-wrinkle, moisturizes

Uses

Anti-wrinkle, moisturizes, nourishes and repairs skin, keeping skin hydrated, soft and smooth.

Warnings

For external use only
Do not use on damaged or broken skin
When using this product keep out of eyes. Rinse with water to remove.
Stop use and ask a doctor if rash occurs
Keep out of reach of children. If swallowed, get medical help or contacta Poison Control Center right away.

Do not use

Do not use on damaged or broken skin

When using this product

When using this product keep out of eyes. Rinse with water to remove.

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Ask Doctor

Directions

- 1. After cleansing the face, take an appropriate amount of spot lightening cream.
- 2. Apply evenly to the area on the face that needs to lighten spots, and massage gently until completely absorbed.
- 3. Use once in the morning and evening. Continuous use will provide better results.

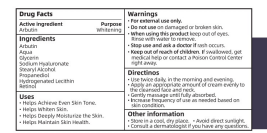
Other information

Please store in a cool, dry place away from direct sunlight.

Inactive ingredients

Arbutin, Aqua, Glycerin, Sodium Hyaluronate, Stearyl Alcohol, Propanediol, Hydrogenated Lecithin, Retinol

PRINCIPAL DISPLAY PANEL





4)

DRUGGIST CREAM

cream cream

Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:83872-531	
Route of Administration	TOPICAL			

Active Ingredient/Active Moiety				
	Ingredient Name	Basis of Strength	Strength	
	ARBUTIN (UNII: C5INA23HXF) (ARBUTIN - UNII:C5INA23HXF)	ARBUTIN	4 g in 50 mL	

Inactive Ingredients			
Ingredient Name	Strength		
GLYCERIN (UNII: PDC6A3C0OX)			
STEARYL ALCOHOL (UNII: 2KR89I4H1Y)			
AQUA (UNII: 059QF0KO0R)			

SODIUM HYALURONATE (UNII: YSE9PPT4TH)	
PROPANEDIOL (UNII: 5965N8W85T)	
RETINOL (UNII: G2SH0XKK91)	

l	Packaging				
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:83872- 531-01	50 mL in 1 BOTTLE, UNIT-DOSE; Type 0: Not a Combination Product	11/26/2024	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	505G(a)(3)	11/26/2024	

Labeler - Shenzhen Xiaomai Manufacturing Co., Ltd. (712999147)

Establishment				
Name	Address	ID/FEI	Business Operations	
Shenzhen Xiaomai Manufacturing Co., Ltd.		712999147	manufacture(83872-531)	

Revised: 11/2024 Shenzhen Xiaomai Manufacturing Co., Ltd.