

NUMB SKIN - lidocaine cream cream

Seenext Venture Ltd

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

Drug Facts

Active Ingredient

Lidocaine 5%

Purpose

Anorectal (Hemorrhoidal)

Uses

For temporary relief of local discomfort, itching, pain, soreness or burning in the perianal area associated with anorectal disorders.

Warnings

For external use only

When using this product

- do not exceed the recommended daily dosage unless directed by a doctor
- do not put into the rectum by using fingers or any mechanical device or applicator

Stop the use and consult doctor if:

- allergic reaction occurs
- rectal bleeding occurs
- redness, irritation, swelling, pain, or other symptoms begin or increase
- condition worsens or does not improve within 7 days

Keep out of reach of children

If swallowed, get medical help or contact a Poison Control Center right away

Directions

- Adults: When practical, cleanse the affected area with mild soap and warm water and rinse thoroughly. Gently dry by patting or blotting with tissue or a soft cloth before application of product.
- when first opening the tube, puncture foil seal with top end of cap
- apply externally to the affected area up to 6 times daily

- children under 12 years of age: consult a doctor

Other information

- always keep the tube tightly closed
- store at temperatures not exceeding 15 °C - 30 °C
- protect from freezing

Inactive ingredients

Benzyl Alcohol, Carbomer 934, Lecithin (Soybean), Propylene Glycol, Tocopheryl Acetate, Water

Question or comments

Call weekdays 9 AM to 6 PM PST at 1-844-700-6862 or email us at support@numbskin.com

Principal Display Panel

NDC 70907-001-15

FOR EXTERNAL USE ONLY

NumbSkin Cream[®]

5% Lidocaine Cream

Maximum Strength Pain Reliever

NET WT. 15g

NDC XXXXX-XXX-XX

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5% Lidocaine Cream

NUMBSKIN[®]
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Drug Facts (continued)

Directions

- Adults: When practical, cleanse the affected area with mild soap and warm water and rinse thoroughly. Gently dry by patting or blotting with tissue or a soft cloth before application of product
- when first opening the tube, twist the cap and peel off the foil seal
- apply externally to the affected area up to 6 times daily
- children under 12 years of age: consult a doctor

Other information

- always keep the tube tightly closed
- store at temperatures not exceeding 59°F = 86°F (15°C = 30°C)
- protect from freezing

Inactive ingredients Benzyl Alcohol, Carbomer 934, Lecithin (Soybean), Propylene Glycol, Tocopheryl Acetate, Water

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NUMBSKIN[®] CREAM

Imported/Distributed by:
Seenext Venture Ltd.
1124 Fir Avenue, Blaine, WA 98230, United States
www.numbskin.care



Batch No.:
Exp. Date:

NDC 70907-001-30

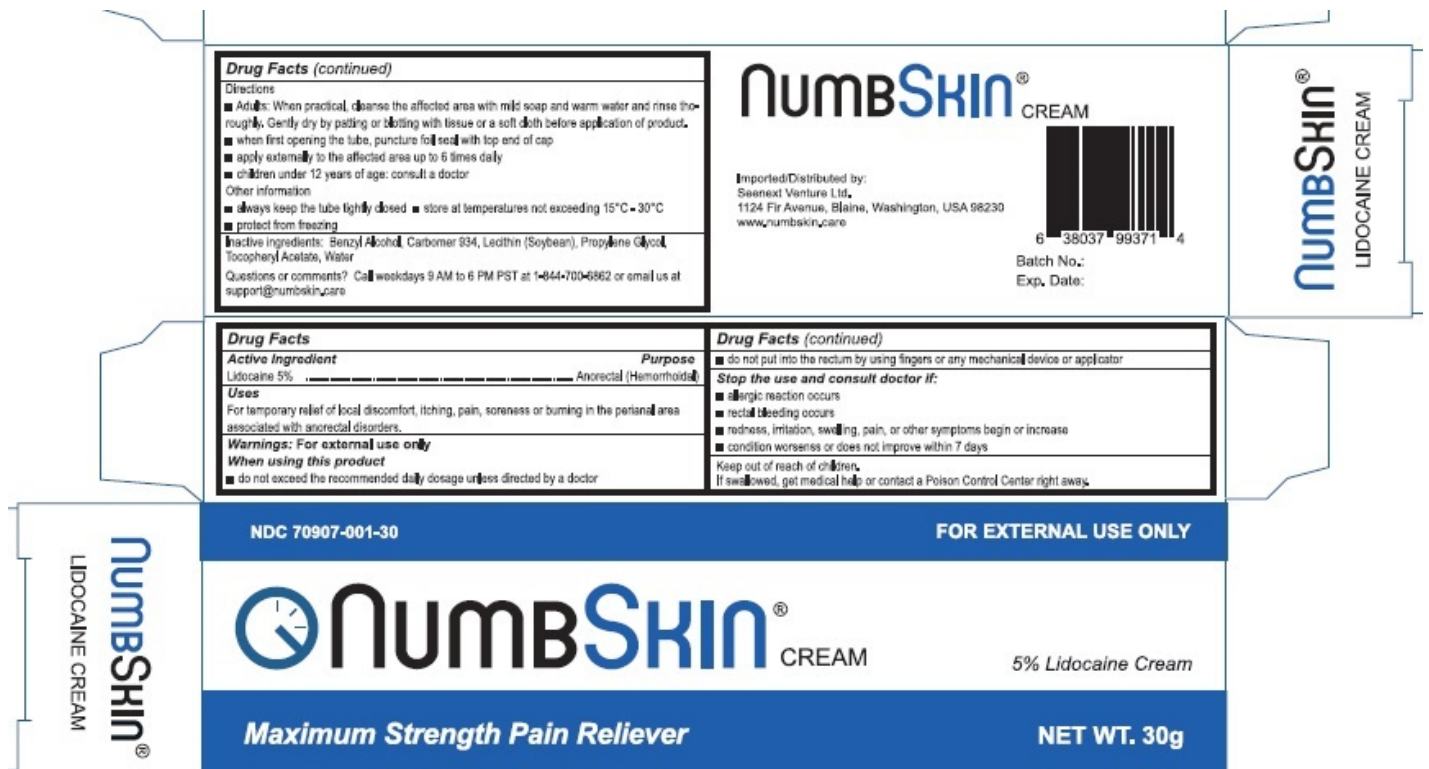
FOR EXTERNAL USE ONLY

NumbSkin Cream[®]

5% Lidocaine Cream

Maximum Strength Pain Reliever

NET WT. 30g



NUMB SKIN

lidocaine cream cream

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:70907-001
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
LIDOCAINE (UNII: 98PI200987) (LIDOCAINE - UNII:98PI200987)	LIDOCAINE	5 mg in 30 g

Inactive Ingredients

Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
CARBOMER 934 (UNII: Z135WT9208)	
PROPYLENE GLYCOL BUTYL ETHER (UNII: 6X8776AP5Z)	
TOCOPHERYL RETINOATE (UNII: 0WN694NBMM)	
BENZYL ALCOHOL (UNII: LKG8494WBH)	
LECITHIN, SOYBEAN (UNII: 1DI56QDM62)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	NDC:70907-001			

1	NDC:70907-001-30	1 in 1 CARTRIDGE	09/01/2016	
1		30 g in 1 TUBE; Type 0: Not a Combination Product		
2	NDC:70907-001-15	1 in 1 CARTRIDGE	04/29/2020	
2		15 g in 1 TUBE; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph final	part346	09/01/2016	

Labeler - Seenext Venture Ltd (203416862)

Registrant - Seenext Venture Ltd (203416862)

Revised: 11/2022

Seenext Venture Ltd