EXTRA STRENGTH SKIN REPAIR- lidocaine hcl cream ridge properties

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.

Directions: Test skin to sensitivity prior to procedure. Apply generously to Affected area as needed for pain. Discontinue use if sensitivity occurs. Not for use on face.

Warning - Keep out of reach of children - For external use only - Avoid contact with the eyes

Warning - Keep out of reach of children - For external use only - Avoid contact with the eyes

Purpose:
Topical Anesthetic

Uses: Temporarily relieves pain

Active ingredients: Lidocaine HCL

4%

Inactive Ingredients:

Water, Witch Hazel, Kava kava, Organic Alcohol, Propolis, Yarrow, Nutmeg, Copaiba Balsam, BTMS 50 (A natural extract of the Colza Seed)

C# 69804-051-02 any known allergy to any Lidocaine HCL 4% product. Discontinue use





manufactured by Pain Relief Naturally. For contact info please visit WWW.NATURALLYHL.COM

This product was

Water, Witch Hazel, Organic Alcohol, Kava Kava, Yarrow, Nutmeg, Propolis, (BTMS 50 -A Natural Extract of The Colza Seed)

EXTRA STRENGTH SKIN REPAIR

lidocaine hcl cream

Product Information

HUMAN OTC DRUG Item Code (Source) NDC:69804-051 Product Type

Route of Administration TOPICAL

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
LIDO CAINE HYDRO CHLO RIDE (UNII: V13007Z41A) (LIDOCAINE - UNII:98PI200987)	LIDOCAINE HYDROCHLORIDE ANHYDROUS	40 mg in 1000 mg		

Inactive Ingredients		
Ingredient Name	Strength	
PROPOLIS WAX (UNII: 6 Y8 XYV2NOF)	40 mg in 1000 mg	
PIPER METHYSTICUM WHOLE (UNII: 3P306S300W)	190 mg in 1000 mg	
BEHENTRIMO NIUM METHO SULFATE (UNII: 5SHP745C61)	120 mg in 1000 mg	
ACHILLEA MILLEFO LIUM O IL (UNII: 97P5D0 WG43)	60 mg in 1000 mg	
WITCH HAZEL (UNII: 10 1I4J0 U34)	400 mg in 1000 mg	
COPAIBA OIL (UNII: 64VX45Y68N)	90 mg in 1000 mg	
NUTMEG OIL (UNII: Z1CLM48948)	60 mg in 1000 mg	

F	ackaging			
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:69804-051-06	14200 mg in 1 JAR; Type 0: Not a Combination Product	04/14/2017	
2	NDC:69804-051-05	28500 mg in 1 JAR; Type 0: Not a Combination Product	04/14/2017	
3	NDC:69804-051-02	56700 mg in 1 JAR; Type 0: Not a Combination Product	0 4/14/20 17	
4	NDC:69804-051-03	113400 mg in 1 JAR; Type 0: Not a Combination Product	04/14/2017	

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
OTC monograph not final	part348	0 4/14/20 17			

Labeler - ridge properties (029478762)

Establishment					
Name	Address	ID/FEI	Business Operations		
ridge properties		029478762	manufacture(69804-051)		

Revised: 4/2017 ridge properties