DE LA CRUZ CAMPHOR- camphor ointment **DLC Laboratories**, Inc.

CAMPHOR Ointment 11%

Active Ingredient

Camphor, USP 11%

Purpose

External analgesic

Uses

for the temporary relief of minor aches and pains of muscles and joints associated with simple backache, arthritis, strains, bruises and sprains

Warnings

For external use only

When using this product

avoid contact with eyes or mucous membranes do not apply to wounds or damaged skin do not bandage tightly or use with a heating pad

Stop use and consult a doctor if

condition worsens or if symptoms persist for more than 7 days symptoms clear up and occur again within a few days. excessive skin irritation develops

If pregnant or breastfeeding,

consult a doctor before use.

KEEP OUT OF THE REACH OF CHILDREN.

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

Directions

adults and children 2 years of age or older, apply to affected area not more than 3 to 4 times daily

children under 2 years of age: consult a doctor

Inactive ingredient

polyethylene glycol

Questions?

1-800-858-3889 or www.dlclabs.com

De La Cruz

CAMPHOR
Ointment 11%

Pain relieving rub

2.5 OZ (70.9g)

FAST, PENETRATING RELIEF FOR:

Muscle and joint pain

Backaches and arthritis

Strains and sprains

Itching

NON-IRRITATING

WATER WASHABLE

NO PARABENS OR ARTIFICIAL FRAGRANCES OR COLORS

Manufactured by:

De La Cruz Products

A Division of DLC Laboratories, Inc.

Paramount, CA 90723 USA

Questions: 1-800-858-3889

www.dlclabs.com (c) DLC

PRINCIPAL DISPLAY PANEL - 70.9 g Jar Label

De La Cruz ®

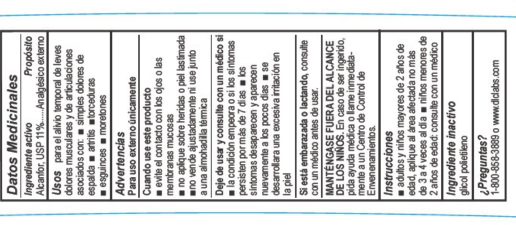
Clinically Tested*

CAMPHOR
OINTMENT 11%

Pain Relieving Rub

2.5 OZ (70.9 g)





Purpose persist for more than 7 days - symptoms External analgesic If pregnant or breast-feeding, consult a days - excessive skin irritation develops **Uses** for the temporary relief of minor CHILDREN. If swallowed, get medical help or contact a Poison Control Center membranes do not apply to wounds adults and children 2 years of age or than 3 to 4 times daily **a** children under aches and pains of muscles and joints older, apply to affected area not more ■arthritis ■ strains ■ sprains ■ bruises clear up and occur again within a few avoid contact with eyes or mucous 1-800-858-3889 or www.dlclabs.com or damaged skin

do not bandage tightly or use with a heating pad condition worsens or if symptoms associated with: simple backache Stop use and consult a doctor if KEEP OUT OF THE REACH OF 2 years of age: consult a doctor When using this product Inactive ingredient For external use only Camphor, USP 11% polyethylene glycol Active ingredient doctor before use. Questions? Directions Warnings right away

Drug Facts

camphor ointment

Product	Intorm	ation
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Product Type HUMAN OTC DRUG Item Code (Source) NDC:242
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Route of Administration TOPICAL

Active Ingredient/Active Moiety

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	Ingredient Name	Basis of Strength	Strength			
	CAMPHOR (SYNTHETIC) (UNII: 5TJD82A1ET) (CAMPHOR (SYNTHETIC) - UNII:5TJD82A1ET)	CAMPHOR (SYNTHETIC)	11 g in 100 g			

Inactive Ingredients

Ingredient Name	Strength

POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:24286- 1521-2	70.9 g in 1 JAR; Type 0: Not a Combination Product	07/26/2012	
2	NDC:24286- 1521-5	155.9 g in 1 JAR; Type 0: Not a Combination Product	03/10/2021	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M017	07/26/2012	

Labeler - DLC Laboratories, Inc. (093351930)

Establishment

Name	Address	ID/FEI	Business Operations	
DLC Laboratories, Inc.		093351930	manufacture(24286-1521) , label(24286-1521)	

Revised: 10/2024 DLC Laboratories, Inc.