GERMA CAMPHOR ANALGESIC- camphor 10% ointment Germa Products, LLC

Camphor Ointment

Active Ingredient

Camphor 10%

Uses

Temporary relief of minor pain • Rheumatism • Muscular aches • Arthritis • Backache

Do not use • on children • on irritated skin • in or near eyes or mucous membranes • on wounds or damaged skin • if prone to allergic reaction from any of the ingredient • on large areas of the body. • if you are taking any other medication or drinking alcohol • if safety seal is torn, broken or missing.

Purpose

Topical Analgesic

Warnings

For external use only

Do not use • on children • on irritated skin • in or near eyes or mucous membranes • on wounds or damaged skin • if prone to allergic reaction from any of the ingredient • on large areas of the body. • if you are taking any other medication or drinking alcohol • if safety seal is torn, broken or missing.

Warnings

KEEP OUT OF THE REACH OF CHILDREN

In case of accidental ingestion, seek professional assistance or contact a Poison Control Center immediately.

If pregnant or breastfeeding • ask a health professional before use.

Stop use and ask a doctor • if condition worsens or cleans up and occurs again • if symptoms persist for more than 7 days or irritation or severe burning occurs.

Other Information

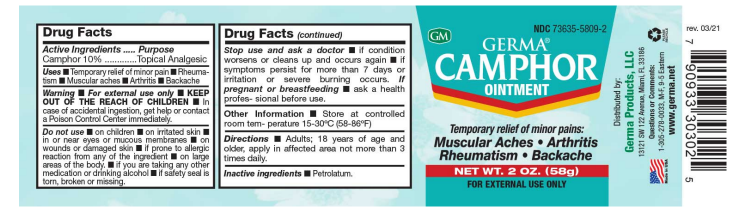
Store at controlled room temperature 15-30°C (59-86°F)

Directions

Adults; 18 years of age and older, apply in affected area not more than 3 times daily.

Inactive Ingredients

Petrolatum



GERMA CAMPHOR ANALGESIC

camphor 10% ointment

Product Type HUMAN OTC DRUG Item Code (Source) NDC:73635-5809

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CAMPHOR (SYNTHETIC) (UNII: 5TJD82A1ET) (CAMPHOR (SYNTHETIC) - UNII:5TJD82A1ET)	CAMPHOR (SYNTHETIC)	100 mg in 1 mL

Inactive Ingredients

Ingredient Name	Strength
ingredient Name	Strength

WHITE PETROLATUM (UNII: B6E5W8RQJ4)

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#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:73635- 5809-2	56.69 mL in 1 JAR; Type 0: Not a Combination Product	03/29/2019	

Marketing	Information
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Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
	0.10.10.1		

	OTC Monograph Drug	M017	03/29/2019	
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Labeler - Germa Products, LLC (116626935)

Revised: 10/2023 Germa Products, LLC