CAMPHOTREX- camphor, menthol gel PureTek Corporation

Camphotrex™

Extra Strength Pain Relieving Gel Roll-on Applicator

Professional Therapy for Muscle & Joint Pain Relief

Active Ingredients (% by weight)

Camphor 4%

Menthol 10%

Purpose

Analgesic (pain relief)

Uses

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

Warnings

For external use only

Do not use on

■ wounds ■ damaged skin

When using this product

■ avoid getting into eyes or mucous membranes ■ do not bandage tightly

Stop use and ask a doctor if

- excessive irritation of the skin develops
 condition worsens
- symptoms last more than 7 days or clear up and occur again within a few days

If pregnant or breast-feeding,

ask a health professional before use.

Keep out of reach of children.

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

Directions

- adults and children 12 years of age or older: using the roll-on applicator massage a liberal amount of gel directly on the affected area, not more than 3 to 4 times daily
- children under the age of 12: do not use, consult a doctor use only as directed

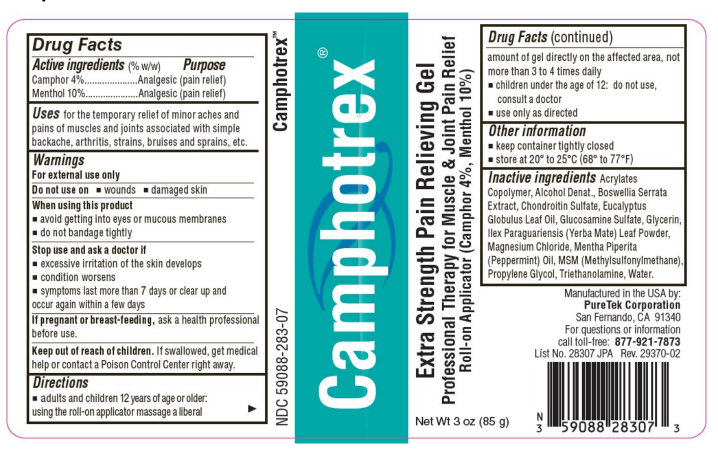
Other information

■ keep container tightly closed ■ store at 20° to 25°C (68° to 77°F)

Inactive ingredients

Acrylates Copolymer, Alcohol Denat., Boswellia Serrata Extract, Chondroitin Sulfate, Eucalyptus Globulus Leaf Oil, Glucosamine Sulfate, Glycerin, Ilex Paraguariensis (Yerba Mate), Magnesium Chloride, Mentha Piperita (Peppermint) Oil, MSM (Methylsulfonylmethane), Propylene Glycol, Triethanolamine, Water.

Camphotrex®



CAMPHOTREX camphor, menthol gel Product Information Product Type HUMAN OTC DRUG Item Code (Source) NDC:59088-283

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
CAMPHOR (SYNTHETIC) (UNII: 5TJD82A1ET) (CAMPHOR (SYNTHETIC) - UNII:5TJD82A1ET)	CAMPHOR (SYNTHETIC)	4 g in 100 g		
MENTHOL (UNII: L7T10EIP3A) (MENTHOL - UNII:L7T10EIP3A)	MENTHOL	10 g in 100 g		

Inactive Ingredients				
Ingredient Name	Strength			
METHACRYLIC ACID - ETHYL ACRYLATE COPOLYMER (4500 MPA.S) (UNII: T967IEU43C)				
ALCOHOL (UNII: 3K9958V90M)				
INDIAN FRANKINCENSE (UNII: 4PW41QCO2M)				
CHONDROITIN SULFATE (BOVINE) (UNII: 6IC1M3OG5Z)				
EUCALYPTUS OIL (UNII: 2R040NI662)				
GLUCOSAMINE SULFATE (UNII: 1FW7WLR731)				
GLYCERIN (UNII: PDC6A3C0OX)				
ILEX PARAGUARIENSIS LEAF (UNII: 1Q953B4O4F)				
MAGNESIUM CHLORIDE (UNII: 02F3473H9O)				
PEPPERMINT OIL (UNII: AV092KU4JH)				
DIMETHYL SULFONE (UNII: 9H4PO4Z4FT)				
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)				
TROLAMINE (UNII: 903K93S3TK)				
WATER (UNII: 059QF0KO0R)				

Packaging					
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:59088- 283-07	85 g in 1 BOTTLE, WTH APPLICATOR; Type 0: Not a Combination Product	04/12/2016		

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
OTC Monograph Drug	M017	04/12/2016			

Labeler - PureTek Corporation (785961046)

Revised: 7/2024 PureTek Corporation