#### MINERAL ICE PAIN RELIEVING- menthol gel Crown Laboratories

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Mineral Ice

#### Active ingredient

Menthol 2%

## Purpose

Topical analgesic

#### Uses

- temporarily relieves minor aches and pains of muscles and joints associated with:
- arthritis simple backache strains
- • bruises sprains
- provides cooling penetrating relief

## Warnings

## For external use only

#### Do not use

- with other topical pain relievers
- with heating pads or heating devices

## When using this product

- do not use in or near the eyes
- do not apply to wounds or damaged skin
- do not bandage tightly

## Stop use and ask a doctor if

- 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**

- clean affected area before applying product
- adults and children 2 years of age and older: apply to affected area not more than 3 to 4 times daily
- children under 2 years of age: ask a doctor

#### Other information

- store at 20 ° 25 °C (68 ° 77 °F) [see USP Controlled Room Temperature], in a tightly closed container
- store in a cool place
- do not use, pour, spill or store near heat or open flame

#### Inactive ingredients

ammonium hydroxide, carbomer, cupric sulfate, FD&C blue no. 1, isopropyl alcohol, magnesium sulfate, purified water, sodium hydroxide, thymol

#### Questions or comments?

call **1-833-279-6522** 

#### **Additional Information Listed on Other Panels**

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## **Principal Display**

NDC 0316-0226-08

ORIGINAL THERAPEUTIC

Mineral Ice®

Menthol Pain Relieving Gel

Greaseless with DEEPCOLD® Pain Reliever

Net wt. 8 oz (226.8 g)

P11529.00



## MINERAL ICE PAIN RELIEVING

menthol gel

#### **Product Information**

Product Type HUMAN OTC DRUG Item Code (Source) NDC:0316-0226

Route of Administration TOPICAL

## **Active Ingredient/Active Moiety**

Ingredient Name	Basis of Strength	Strength
MENTHOL (UNII: L7T10EIP3A) (MENTHOL - UNII:L7T10EIP3A)	MENTHOL	20 mg in 1 g

Inactive Ingredients				
Ingredient Name	Strength			
AMMONIA (UNII: 5138Q19F1X)				
CARBOMER HOMOPOLYMER TYPE B (UNII: HHT01ZNK31)				
CUPRIC SULFATE (UNII: LRX7AJ16DT)				
FD&C BLUE NO. 1 (UNII: H3R47K3TBD)				
ISOPROPYL ALCOHOL (UNII: ND2M416302)				
MAGNESIUM SULFATE HEPTAHYDRATE (UNII: SK47B8698T)				
SODIUM HYDROXIDE (UNII: 55X04QC32I)				
THYMOL (UNII: 3J50XA376E)				
WATER (UNII: 059QF0KO0R)				

P	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:0316-0226- 35	99.2 g in 1 JAR; Type 0: Not a Combination Product	12/01/2018		
2	NDC:0316-0226- 08	226.8 g in 1 JAR; Type 0: Not a Combination Product	12/01/2018		
3	NDC:0316-0226- 16	453.6 g in 1 JAR; Type 0: Not a Combination Product	12/01/2018		

## **Marketing Information**

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M017	11/01/2011	

# **Labeler -** Crown Laboratories (079035945)

Revised: 11/2023 Crown Laboratories