

**ICY HOT KIDS PAIN RELIEF ROLL-ON- menthol liquid
Chattem, Inc.**

Icy Hot Kids Pain Relief Roll-On

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Drug Facts

Active ingredient

Purpose

Menthol
4%.....Topical
analgesic

Uses

temporarily relieves minor aches and pains of muscles and joints associated with: ■
strains
■ sprains ■ bruises ■ simple backache

Warnings

For external use only

Do not use

- on wounds or on irritated or damaged skin
- with a heating pad

When using this product

- use only as directed
- do not bandage tightly
- avoid contact with eyes and mucous membranes
- do not expose the area to local heat or to direct sunlight
- rare cases of serious burns have been reported with products of this type
- a transient burning sensation may occur upon application but generally disappears in several days
- avoid applying into skin folds

Stop use and ask a doctor if

- condition worsens or symptoms persist for more than 7 days
- symptoms clear up and occur again within a few days
- redness is present or excessive skin irritation occurs
- you experience severe burning pain, swelling, or blistering where the product was applied

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

- children 2 years of age and older:
 - apply a thin layer to affected area not more than 3 to 4 times daily
 - massage until thoroughly absorbed into skin
 - wash hands thoroughly with soap and water after each use
- children under 2 years of age: ask a doctor

Inactive ingredients

allantoin, aloe barbadensis leaf juice, carbomer, citric acid, diisopropyl adipate, ethylhexylglycerin, glycerin, hydroxyacetophenone, potassium hydroxide, steareth-2, steareth-21, water

PRODUCT SHOULD BE APPLIED UNDER ADULT SUPERVISION. Close cap tightly after use.

Keep carton as it contains important information.

PRINCIPAL DISPLAY PANEL

Icy Hot

Kids Pain Relief

Roll-On

1.5 FL OZ (44.3 mL) ROLL-ON



ICY HOT KIDS PAIN RELIEF ROLL-ON

menthol liquid

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:41167-0799
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
MENTHOL (UNII: L7T10EIP3A) (MENTHOL - UNII:L7T10EIP3A)	MENTHOL	4 g in 100 mL

Inactive Ingredients

Ingredient Name	Strength
ALLANTOIN (UNII: 344S277G0Z)	
ALOE VERA LEAF (UNII: ZY81Z83H0X)	
CARBOMER HOMOPOLYMER, UNSPECIFIED TYPE (UNII: 0A5MM307FC)	
CITRIC ACID MONOHYDRATE (UNII: 2968PHW8QP)	
DIISOPROPYL ADIPATE (UNII: P7E6YFV72X)	
ETHYLHEXYLGLYCERIN (UNII: 147D247K3P)	
GLYCERIN (UNII: PDC6A3C0OX)	
HYDROXYACETOPHENONE (UNII: G1L3HT4CMH)	
POTASSIUM HYDROXIDE (UNII: WZ3C48M4T)	
STEARETH-2 (UNII: V56DFE46J5)	
STEARETH-21 (UNII: 53J3F32P58)	

WATER (UNII: 059QF0KO0R)

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:41167-0799-0	1 in 1 CARTON	01/15/2023	
1		44.3 mL in 1 BOTTLE; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M017	01/15/2023	

Labeler - Chattem, Inc. (003336013)

Revised: 10/2023

Chattem, Inc.