#### **ROCKSAUCE ICY COLD PAIN RELIEF- menthol gel** Implus Footcare, LLC

Disclaimer: Most OTC drugs are not reviewed and approved by FDA, however they may be marketed if they comply with applicable regulations and policies. FDA has not evaluated whether this product complies.

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RockSauce® ICY COLD PAIN RELIEF GEL 120Z

# **ACTIVE INGREDIENT**

Menthol6%

# PURPOSE

Topical Analgesic

# **USES:**

Provides soothing cold to sore muscles and joints.

### WARNINGS:

For external use only. Avoid contact with eyes, mucous membrane or broken skin.

### When using this product, do not:

heat, microwave, add to hot water or any container where heating water may cause splattering and result in burns, use in eyes or directly on mucus membrane, take by mouth or place in nostrils, apply to wounds or damaged skin.

### Do not use otherwise than as directed.

Consult a doctor and discontinue use if irritation occurs.

Ask a health professional before use if pregnant or breastfeeding.

Ask a doctor before using if you have sensitive skin.

### Keep out of reach of children:

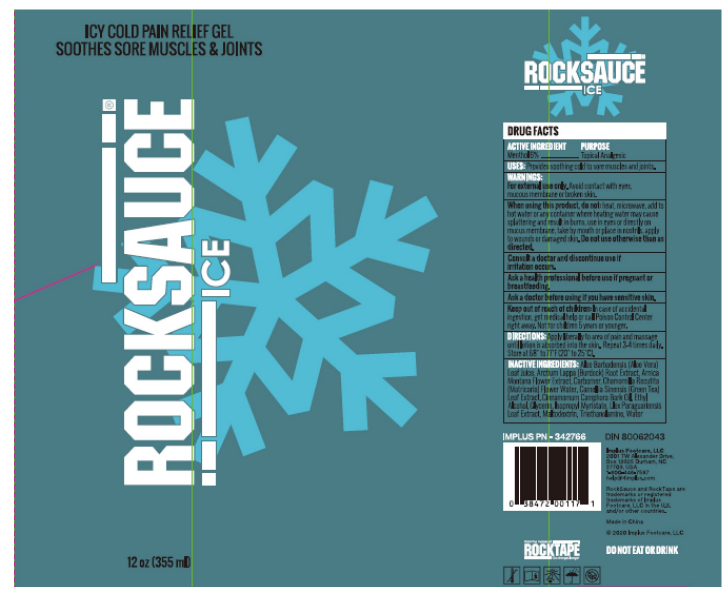
In case of accidental ingestion, get medical help or call Poison Control Center right away. Not for children 5 years or younger.

# **DIRECTIONS:**

Apply liberally to area of pain and massage until lotion is absorbed into the skin. Repeat 3-4 times daily. Store at 68° to 77 °F(20° to 25°C).

### **INACTIVE INGREDIENTS:**

Aloe Barbadensis(Aloe Vera) Leaf Juice, Arctium Lappa(Burdock)Root Extract, Arnica Montana Flower Extract, Carbomer, Chamomilla Recutita (Matricaria) Flower Water, Camellia Sinensis(Green Tea) Leaf Extract, Cinnamomum Camphora Bark Oil, Ethyl Alcohol, Glycerin, Isopropyl Myristate, Ilex Paraguariensis Leaf Extract, Maltodextrin, Triethanolamine, Water



# **ROCKSAUCE ICY COLD PAIN RELIEF**

menthol gel

#### Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:81683-012
Route of Administration	TOPICAL		

Active Ingred	ient/Active Moiety		
	Ingredient Name	Basis of Strength	Strength
MENTHOL (UNII: L	7T10EIP3A) (MENTHOL - UNII:L7T10EIP3A)	MENTHOL	33.2 g in 355 mL
Inactive Ingre	edients		
Ingredient Name			Strength
LEX PARAGUARIE	NSIS LEAF (UNII: 1Q953B4O4F)		
ARCTIUM LAPPA F			
CHAMOMILE FLO			
CARBOMER 940 (	UNII: 4Q93RCW27E)		
ALOE VERA LEAF	(UNII: ZY81Z83H0X)		
CAMPHOR OIL (UI			
ISOPROPYL MYRI	STATE (UNII: ORE8K4LNJS)		
MALTODEXTRIN (	UNII: 7CVR7L4A2D)		
WATER (UNII: 0590	QF0KO0R)		
GLYCERIN (UNII: P	DC6A3C0OX)		
TROLAMINE (UNII:	903K93S3TK)		
	A FLOWER (UNII: OZ0E5Y15PZ)		
ALCOHOL (UNII: 31			
GREEN TEA LEAF	(UNII: W2ZU1RY8B0)		
Packaging			
# Item Code	Package Description	Marketing Start Date	Marketing End Date
<b>1</b> NDC:81683-012- 01	355 mL in 1 BOTTLE; Type 0: Not a Combir Product	ation 03/18/2021	
Marketing	Information		
Marketing Category	Application Number or Monog Citation	raph Marketing Start Date	Marketing End Date
DTC monograph no	part348	03/18/2021	

Labeler - Implus Footcare, LLC (361531585)

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Implus Footcare, LLC