# HEMPVANA COLD AS ICE PAIN RELIEF- menthol gel Telebrands Corp

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**Hempvana Cold As Ice Pain Relief** 

# **Drug Facts**

## **Active Ingredient:**

Menthol USP 8% w/w

## **Purpose**

Menthol USP 8% w/w.....Topical Analgesic

#### Uses

temporarily relief of minor aches and pains of muscles and joints associated with:

• simple backache • arthritis • strains • bruises • sprains

# Warnings

# For external use only

Flammable: Keep away from excessive heat or open flame

When using this product, avoid contact with eyes.

- Do nut use in large quantities, particularly over raw surfaces or blistered areas.
- Do not apply to wounds or damaged skin
- Do not bandage tightly

# Stop use and ask a doctor if

- Condition worsens
- Symptoms persist for more than 7 days or clear up and occur again within a few days

**Keep out of reach of children.** If product is swallowed, get medical help or contact a Poison Control Center right away.

#### **Directions**

**Adults**: Apply to affected are not more than 3 to 4 times daily. If produce comes in contact with hands, wash with soap and water.

For children under 18 years of age: Consulta a physician.

Store tightly closed in a cool dry place.

**Inactive Ingredients** 

Arctium Lappa Root Extract, Aloe Barbadensis Leaf Extract, Arnica Montana Flower Extract, Boswellia Carterii Resin Extract, Calendula Officinalis Extract, Camellia Sinensis Leaf Extract, Camphor, Cannabis Sativa Seed Oil, Carbomer, FD&C Blue No 1, FD&C Yellow No 5, Glycerin, Ilex Paraguariensis Leaf Extract, Isopropyl Alcohol, Isopropyl Myristate, Melissa Officinalis (Lemon Balm) Extract, Silica, Tocopheryl Acetate, Triethanolamine, Water

## Questions?

Call (855) 877-4503 (M-F, 9am-5pm EST)

# **Packaging**



#### **HEMPVANA COLD AS ICE PAIN RELIEF**

menthol gel

## **Product Information**

Product Type HUMAN OTC DRUG Item Code (Source) NDC:73287-004

Route of Administration TOPICAL

# Active Ingredient/Active Moiety Ingredient Name Basis of Strength MENTHOL (UNII: L7T10EIP3A) (MENTHOL - UNII:L7T10EIP3A) MENTHOL 8 mg in 100 mL

Inactive Ingredients	
Ingredient Name	Strength
WATER (UNII: 059QF0KO0R)	
ETHYLHEXYL STEARATE (UNII: EG3PA2K3K5)	
ARCTIUM LAPPA ROOT (UNII: 597E9BI3Z3)	
ALOE VERA LEAF (UNII: ZY81Z83H0X)	
ILEX PARAGUARIENSIS LEAF (UNII: 1Q953B4O4F)	
ARNICA MONTANA FLOWER (UNII: OZ0E5Y15PZ)	
FRANKINCENSE (UNII: R9XLF1R1WM)	
CALENDULA OFFICINALIS WHOLE (UNII: PFR03EBU0H)	
GREEN TEA LEAF (UNII: W2ZU1RY8B0)	
CAMPHOR (NATURAL) (UNII: N20HL7Q941)	
CANNABIS SATIVA SEED OIL (UNII: 69VJ1LPN1S)	
CARBOMER HOMOPOLYMER, UNSPECIFIED TYPE (UNII: 0A5MM307FC)	
FD&C BLUE NO. 1 (UNII: H3R47K3TBD)	
FD&C YELLOW NO. 5 (UNII: I753WB2F1M)	
GLYCERIN (UNII: PDC6A3C0OX)	
ISOPROPYL ALCOHOL (UNII: ND2M416302)	
ISOPROPYL MYRISTATE (UNII: 0RE8K4LNJS)	
MELISSA OFFICINALIS (UNII: YF70189L0N)	
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
.ALPHATOCOPHEROL ACETATE (UNII: 9E8X80D2L0)	
TROLAMINE (UNII: 903K93S3TK)	
D&C GREEN NO. 5 (UNII: 8J6RDU8L9X)	
SODIUM HYDROXIDE (UNII: 55X04QC32I)	
PHENOXYETHANOL (UNII: HIE492ZZ3T)	

Packaging						
#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
1	NDC:73287- 004-01	1 in 1 CARTON	06/11/2019			
1		74 mL in 1 BOTTLE, WTH APPLICATOR; 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	06/11/2019	

# Labeler - Telebrands Corp (177266558)

Establishment					
Name	Address	ID/FEI	<b>Business Operations</b>		
A.I.G. TECHNOLOGIES, INC.		086365223	manufacture(73287-004)		

Revised: 11/2024 Telebrands Corp