CBD CRYOTHERAPY PAIN RELIEF ROLL-ON- menthol gel Global Widget, LLC

Drug Facts

Active Ingredients

Menthol USP 4%

Purpose

Topical Analgesic

Uses

For the temporary relief of pain

Warnings

FOR EXTERNAL USE ONLY

Do not use:

- On eyes or on mucous membranes
- On wounds, damaged or irritated skin
- If you are allergic to Menthol or any of the ingredients listed below

When using this product:

- Use only as directed
- Do not bandage or cover with any type of wrap except clothing
- Do not use with heating pad or devices, or apply external heat

Stop use and ask a doctor if

- Localized skin reactions occur, such as rash, itching, redness, irritation, pain, swelling and blistering
- Conditions worsen
- Symptoms persist for more than 7 days or clear up and occur again within a few days

If pregnant or breastfeeding: Do not use this product.

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

Directions

- Shake well before use
- Adults 18 years & over rub a thin layer into affected areas up to 4 times daily.

Other Information:

Store in cool, dry place away from direct sunlight

Inactive Ingredients

Water, Isopropyl Alcohol, Carbomer, Pure CBD Extract, Triethanolamine

Distributed by Global Widget, LLC 8419 Sunstate Street, Tampa, FL 33634

Principal Display Panel NATURE'S SCRIPT CBD Cryotherapy Pain Relief Roll-On MENTHOL 4% 3 FL OZ (89 mL)



CBD CRYOTHERAPY PAIN RELIEF ROLL-ON

menthol gel

Product Information				
Product Type	HUMAN OTC DRUG Item Code (Source)		NDC:73423-005	
Route of Administration	TOPICAL			
A . I I I' I / A . I .	Na . 1 . 1			
Active Ingredient/Active	Molety			
Ingredi		Basis of Strength	n Strength	
MENTHOL (UNII: L7T10EIP3A) (MENTHOL - UNII:L7T10EIP3A)			MENTHOL	40 mg in 1 mL
Inactive Ingredients				
Inactive Ingredients	Ingredient Name	9		Strength
Inactive Ingredients ISOPROPYL ALCOHOL (UNII: ND2	-	1		Strength
ISOPROPYL ALCOHOL (UNII: ND2	2M416302)	9		Strength
-	2M416302))		:С)	Strength

Packaging					
#	ltem Code	Package Description	Marketing Start Date	Marketing End Date	
	NDC:73423-005- 01	6 in 1 PACKAGE	12/11/2020	11/30/2025	
1	89 mL in 1 BOTTLE; Type 0: Not a Combination Product				
M	arketing	Information			
Μ	arketing Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	

Labeler - Global Widget, LLC (089584863)

Establishment					
Name	Address	ID/FEI	Business Operations		
Global Widget, LLC		089584863	manufacture(73423-005)		

Establishment

Name	Address	ID/FEI	Business Operations
Global Widget LLC		118504011	manufacture(73423-005)

Revised: 12/2023

Global Widget, LLC