NUMOTIZINE- menthol ointment Hobart Laboratories, Inc.

NUMOTIZINE OINTMENT

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Active Ingredients

Menthol 1.25%

Purpose

Topical Analgesic

Inactive ingredients

Clay, Color, Fragrance of Guaiacol, Methyl Guaiacol and Oil of Wintergreen, Polyols.

Directions

Stir in any liquid at top of jar. Keep sealed when not in use.

Spread 1/8" to 1/4" of ointment to the skin. Cover the ointment with a cloth or bandage to protect clothing. Remove with warm water before totally dry (usually 8 to 12 hours). Application may be repeated every 12 hours as needed.

Uses

For use as a topical analgesic

- Provides temporary relief of muscle pain, soreness and stiffness
- Temporary pain relief on strains, sprains, ligament and tendon injuries
- Arthritis

Stop use and ask a doctor if

- Excessive irritation of the skin occurs.
- Persistent swellings.

Keep out of the reach of children

Warnings:

For external use only.

Use only as directed. Avoid contact with eyes and mucous memebranes.

Do not apply to irritated or broken skin or to large areas of the body.

Representative Labeling For - Numotizine Ointment 3.5oz/99g (10546-100-35) | Numotizine Ointment 8oz/228g (10546-100-08)





WARNINGS

For external use only.
Use only as directed. Avoid contact with eyes and mucous membranes.
Do not apply to irritated or broken skin or to large areas of the body.



NDC 10546-100-08 NET WT. 8 OZ (228 GRAMS)

NUMOTIZINE, Trade Mark Reg. Manufactured in U.S.A.

HOBART LABORATORIES, INC., Bernidji, MN 56601, U.S.A. Phone 218-751-9505 www.rumotizine.com

NUMOTIZINE[®]

PROVIDES TEMPORARY RELIEF OF PAIN AND SORENESS



NET WT. 8 OZ (228 GRAMS)

03/13 4 50386-01





Package Labeling:

3.5 OZ. NET WT. 99 GRAMS Drug Facts Warnings: For external use only. Use only as directed. Avoid contact with eyes and mucous membranes. Do not apply to irritated or broken skin or to large areas of the body. **NUMOTIZINE** Active Ingredients Purpose Topical analgesic Menthol 1.25% Directions Spread 1/8" to 1/4" of ointment to the skin. Cover Stop use and ask a doctor if Excessive irritation of the skin occurs, Persistent swellings. Keep aut of the reach of children the ointment with a cloth or bandage to protect For use as a topical analgesic clothing. Remove with warm water before totally Provides temporary relief of muscle pain. soreness and stiffness dry (usually 8 to 12 hours). Application may be repeated every 12 hours as needed. Temporary pain relief on strains, sprains, ligament and tendon injuries. Stir in any liquid at top of jar. Keep sealed when not in use. Arthritis NUMOTIZINE T.M. Reg. Made in U.S.A. HOBART LABORATORIES, INC., BEMIDJI, MN 56601 U.S.A. PHONE 1-218-751-9505 www.numotizine.com Inactive ingredients Clay, Color, Fragrance of Guaiacol, Methyl Guaiacol and Oil of Wintergreen, Polyols.



NUMOTIZINE

menthol ointment

Product Information

Product Type HUMAN OTC DRUG Item Code (Source) NDC:10546-100

Route of Administration TOPICAL

Active Ingredient/Active Moiety

Ingredient Name
Basis of Strength
MENTHOL (UNII: L7T10EIP3A) (MENTHOL - UNII:L7T10EIP3A)

MENTHOL
1.25 g in 100 g

Inactive Ingredients

Ingredient Name	Strength
KAOLIN (UNII: 24H4NWX5CO)	
GUAIACOL (UNII: 6JKA7MAH9C)	
CREOSOL (UNII: W9GW1KZG6N)	
METHYL SALICYLATE (UNII: LAV5U5022Y)	
PROPYLENE GLYCOL (UNII: 6DC9Q167V3)	

Product Characteristics			
Color	pink	Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

P	Packaging					
#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
1	NDC:10546-100- 35	99 g in 1 JAR; Type 0: Not a Combination Product	11/30/2011			
2	NDC:10546-100- 08	228 g in 1 JAR; Type 0: Not a Combination Product	11/30/2011			

Marketing Information				
Marketing Application Number or Monograph Category Citation		Marketing Start Date	Marketing End Date	
OTC Monograph Drug	M017	11/30/2011		

Labeler - Hobart Laboratories, Inc. (005111786)

Registrant - Hobart Laboratories, Inc. (005111786)

Establishment				
Name	Address	ID/FEI	Business Operations	
Hobart Laboratories, Inc.		005111786	manufacture(10546-100)	

Revised: 10/2023 Hobart Laboratories, Inc.