

MUSCLE PAIN RELIEF GEL- menthol gel
Delon Laboratories (1990) Ltd.

Muscle Pain Relief Gel

Active ingredient

Menthol 2.0%

Purpose

Topical analgesic

Uses

- for the temporary relief of minor aches and pains of muscles and joints

Warnings

For external use only

When using this product

- avoid contact with the eyes
- 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

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

- adults and children 2 years of age and older: apply to affected area not more than 3 to 4 times daily
- children under 2 years of age: consult a doctor

Inactive ingredients

acrylates/C10-30 alkyl acrylate crosspolymer, aminomethyl propanol, denatonium benzoate, ethyl alcohol, FD&C blue no. 1, thymol, water

DELON Muscle Pain Relief Gel 3.17 OZ

DO NOT USE IF SECURITY SEAL IS BROKEN OR MISSING.

Drug Facts

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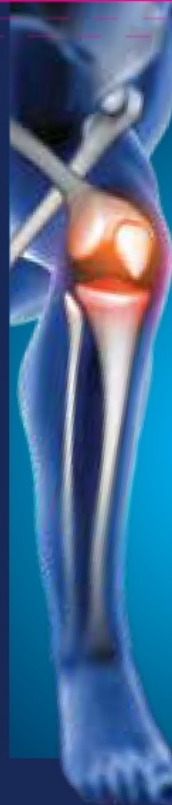
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DELON
NDC 61734-060-01

TEMPORARILY

**RELIEVES MINOR
ACHES AND PAINS
OF MUSCLES AND JOINTS**

**Muscle
Pain Relief
Gel**

Temporary relief of:

- Muscle Pain
- Joint Pain
- Simple Backache
- Arthritis

NET WT. 3.17 OZ

SPACE FOR LOT # AND EXPIRY
DO NOT PRINT

Made in Canada by :
LABORATOIRES DELON (1990) INC.
Pointe-Claire, Québec, Canada, H9R 1E2
www.labdelon.com @labdelon



MUSCLE PAIN RELIEF GEL

menthol gel

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:61734-060
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
MENTHOL (UNII: L7T10EIP3A) (MENTHOL - UNII:L7T10EIP3A)	MENTHOL	2 g in 100 g

Inactive Ingredients

Ingredient Name	Strength
DENATONIUM BENZOATE (UNII: 4YK5Z54AT2)	
ACRYLATES/C10-30 ALKYL ACRYLATE CROSSPOLYMER (60000 MPA.S) (UNII: 8Z5ZAL5H3V)	
ALCOHOL (UNII: 3K9958V90M)	
FD&C BLUE NO. 1 (UNII: H3R47K3TBD)	
AMINOMETHYLPROPANOL (UNII: LU49E6626Q)	
THYMOL (UNII: 3J50XA376E)	
WATER (UNII: 059QF0KO0R)	

Product Characteristics

Color	blue	Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:61734-060-01	90 g in 1 BOTTLE, WTH APPLICATOR; Type 0: Not a Combination Product	09/09/2024	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC Monograph Drug	M017	09/09/2024	

Labeler - Delon Laboratories (1990) Ltd. (248364184)

Establishment

Name	Address	ID/FEI	Business Operations
Laboratoires Delon		208896216	manufacture(61734-060) , pack(61734-060) , label(61734-060)

Revised: 10/2024

Delon Laboratories (1990) Ltd.