SANA SANA MENTHOL EXTERNAL ANALGESIC- menthol gel Alquimia Najulam S de RL de CV

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.

SANA SANA Menthol External Analgesic

Drug Facts

Active ingredient

Menthol 15%

Purpose

External Analgesic

Uses

for the temporary relief of minor aches and pains of muscles and joints associated with

- simple backache
- arthritis
- strains
- bruises
- sprains

Warnings

For external use only

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

When using this product

- avoid contact with eyes
- do not apply to wounds or damaged skin
- do not bandage tightly

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

Package Labeling:

Menthol External Analgesic FANASANA

Are you tired of being in pain?



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Inactive ingredients

ionized water, glycerin, triethanolamine, carbopol

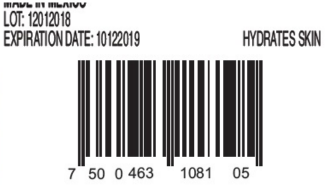
MANUFACTURED BY ALQUIMIA NAJULAM S. DE R.L. DE C.V. AVENIDA ALLENDE NUMERO 63 INTERIOR 3. COLONIA CENTRO ORIENTE, TORREON COAHUILA, MEXICO

TO REPORT A SERIOUS ADVERSE EVENT, CONTACT (512) 567-3598 1413 BARONET'S TRAIL, AUSTIN, TEXAS 78753

MADE IN MEXICO

HECHO EN







SANA SANA MENTHOL EXTERNAL ANALGESIC

menthol gel

Product Information

Product Type

HUMAN OTC DRUG

Item Code (Source)

NDC:72367-000

Route	of Administration	
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Active Ingredient/Active Moiety						
Ingredient Name		Basis of Strengt	th Strength			
MENTHOL (UNII: L7T10 EIP3A) (MENTHOL - UNII:L7T10 EIP3A) MENTHOL			150 mg in 1 g			
Inactive Ingredients						
	inclute ingreate	Strength				
w	ATER (UNII: 059QF0					
G	LYCERIN (UNII: PDC					
Т	ROLAMINE (UNII: 90					
С	ARBO XYPO LYMET					
Packaging						
#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
1	NDC:72367-000-00	1 in 1 BOX	06/22/2018			
1		240 g in 1 BOTTLE; Type 0: Not a Combination Product				
Marketing Information						
	Marketing Categor	y Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
OTC monograph not final		al part348	06/22/2018			

Labeler - Alquimia Najulam S de RL de CV (812839238)

Revised: 6/2018

Alquimia Najulam S de RL de CV