THERAWORX RELIEF- magnesium sulfate heptahydrate liquid AVADIM HOLDINGS, INC.

Disclaimer: This homeopathic product has not been evaluated by the Food and Drug Administration for safety or efficacy. FDA is not aware of scientific evidence to support homeopathy as effective.

Theraworx Relief

Drug Facts

Active Ingredient

Contains Magnesium Sulfate (Magnesia sulphurica) 6X 0.05% HPUS

The letters H.P.U.S. indicate that the components in this product are officially monographed in the Homeopathic Pharmacopoeia of the United States.

Purpose

Muscle Soreness Relief

Uses

- prevents cramps and spasms
- releases muscle tightness
- relieves muscle soreness

Warnings

For external use only. If eye contact occurs, rinse thoroughly with water.

When using this product

- avoid eye contact
- store between 32°F and 120°F
- use only as directed
- not for ingestion

Stop use and ask a doctor if

unintended effects occur.

If pregnant or breastfeeding,

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.

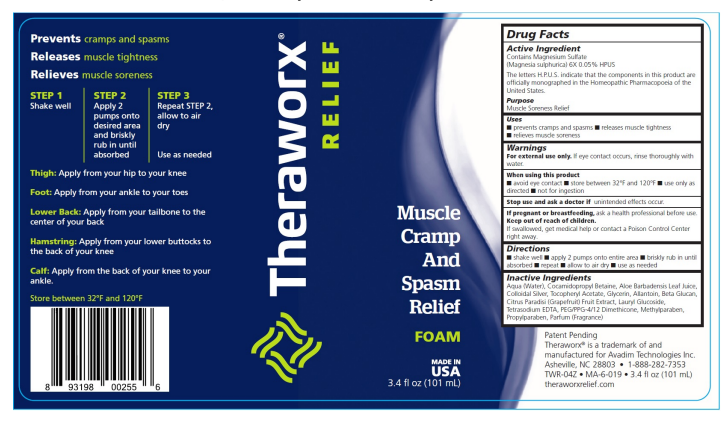
Directions

• shake well • apply 2 pumps onto entire area • briskly rub in until absorbed • repeat • allow to air dry • use as needed

Inactive Ingredients

Aqua (Water), Cocamidopropyl Betaine, Aloe Barbadensis Leaf Juice, Colloidal Silver, Tocopheryl Acetate, Glycerin, Allantoin, Beta Glucan, Citrus Paradisi (Grapefruit) Fruit Extract, Lauryl Glucoside, Tetrasodium EDTA, PEG/PPG-4/12 Dimethicone, Methylparaben, Propylparaben, Parfum (Fragrance)

Theraworx Relief 3.4oz/101mL (61594-006-03)



THERAWORX RELIEF magnesium sulfate heptahydrate liquid **Product Information HUMAN OTC DRUG Product Type** Item Code (Source) NDC:61594-006 **TOPICAL Route of Administration Active Ingredient/Active Moiety Basis of Strength Ingredient Name** Strength MAGNESIUM SULFATE HEPTAHYDRATE (UNII: SK47B8698T) MAGNESIUM SULFATE 6 [hp X]

Inactive Ingredients				
Ingredient Name	Strength			
WATER (UNII: 059QF0KO0R)				
COCAMIDOPROPYL BETAINE (UNII: 50CF3011KX)				
ALOE VERA LEAF (UNII: ZY81Z83H0X)				
SILVER (UNII: 3M4G523W1G)				
GLYCERIN (UNII: PDC6A3C0OX)				
ALLANTOIN (UNII: 344S277G0Z)				
GRAPEFRUIT (UNII: O82C39RR8C)				
LAURYL GLUCOSIDE (UNII: 76LN7P7UCU)				
EDETATE SODIUM (UNII: MP1J8420LU)				
PEG/PPG-4/12 DIMETHICONE (UNII: JAN3585W85)				
METHYLPARABEN (UNII: A218C7HI9T)				
PROPYLPARABEN (UNII: Z8IX2SC10H)				

II	Packaging					
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
	1	NDC:61594-006- 03	101 mL in 1 BOTTLE; Type 0: Not a Combination Product	11/08/2018		

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
unapproved homeopathic		11/08/2018			

Labeler - AVADIM HOLDINGS, INC. (118512488)

Establishment						
Name	Address	ID/FEI	Business Operations			
AVADIM HOLDINGS, INC.		118512488	manufacture(61594-006)			

Revised: 4/2023 AVADIM HOLDINGS, INC.