MUSCLE CRAMPS- artemisia vulgaris root pellet Natural Health Supply

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.

Indications

Indications: To be used for acute self-limiting conditions according to standard homeopathic indications

Active Ingredient

ARTEMISIA VULGARIS

Directions

Take at onset of symptoms. Repeat every 2 hours or as needed until relieved. If condition persists or worsens discontinue use and consult a practitioner.

Adults: dissolve 5-10 pellets in 1 oz. of filtered water or take dry by mouth. Children and infants: 1-5 pellets.

Keep Out of Reach of Children

Keep these and all medications out of the reach of children.

Warning

If pregnant or nursing, consult a practitioner before using.

Inactive Ingredients

Inactive Ingredients - Lactose, Sucrose

Purpose

Purpose: MUSCLE CRAMPS

Package Label

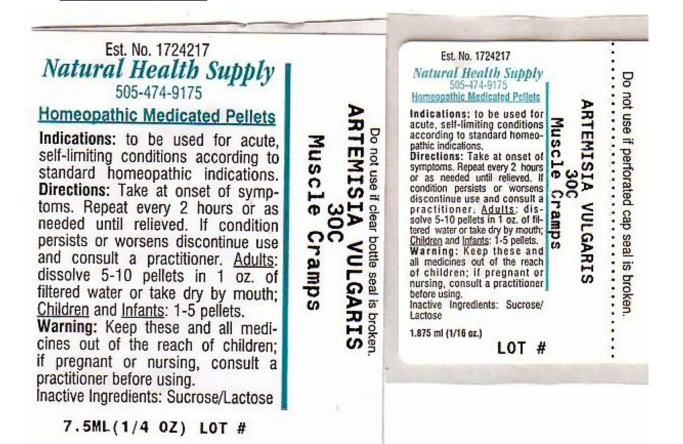
Est. No. 1724217

Natural Health Supply

505-474-9175

Homeopathic Medicated Pellets

Do not use if clear bottle seal is broken ARTEMISIA VULGARIS 30CC 7.5ML (1/4 OZ)



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Lot #_

MUSCLE CRAMPS					
artemisia vulgaris root pellet					
Product Information					
Product Type	HUMAN OTC DRUG	Item Code (Source)		NDC:64117-209	
Route of Administration	ORAL				
Active Ingredient/Active Mo	iety				
Ingredient Name			Basis of Strength		Strength
ARTEMISIA VULGARIS ROOT (UNII: 32MP823R8S) (ARTEMISIA VULGARIS ROOT - UNII:32MP823R8S)					30 [hp_C] in 1 mL
Inactive Ingredients					
Inactive Ingredients	Ingredient Name			Strei	ıgth
	Ingredient Name			Strei	ngth
LACTOSE (UNII: J2B2A4N98G)	Ingredient Name			Strei	ıgth
Inactive Ingredients LACTOSE (UNII: J2B2A4N98G) SUCROSE (UNII: C151H8M554)	Ingredient Name			Strei	ıgth
LACTOSE (UNII: J2B2A4N98G)	Ingredient Name			Strei	ngth

#	Item Code	Package Description	Marketing Start Date	Marketing End Date			
1	NDC:64117-209- 01	1.875 mL in 1 BOTTLE, GLASS; Type 0: Not a Combination Product	06/04/1998				
12	NDC:64117-209- 02	C:64117-209- 7.5 mL in 1 BOTTLE, GLASS; Type 0: Not a Combination Product					
Marketing Information							
	Marketing Categ	ory Application Number or Monograph Citation	Marketing Start Date	Marketing End Date			
un	approved homeopa	athic	06/04/1998				

Labeler - Natural Health Supply (018504618)

Registrant - Natural Health Supply (018504618)

Establishment								
Name	Address	ID/FEI	Business Operations					
Natural Health Supply		0 18 50 46 18	manufacture(64117-209)					

Revised: 4/2019

Natural Health Supply