RELIEVEIT GEL MAX- arnica montana, calendula officinalis, camphora (camphor), capsaicum oleoresin (capsaicin), mentha piperita (peppermint) gel Cosmetic Specialty Labs, 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.

RelieveIt Gel Max

Active Ingredient

Arnica montana 1X: 7%

Calendula officinalis 6X: 2%

Camphora (Camphor): 7%

Capsaicum Oleoresin (Capsaicin): 0.02%

Mentha piperita (Peppermint): 7%)

Purpose

Temporary pain relief, reduces swelling and alleviates inflammation.

Uses:

Arthritis • Osteoarthritis • Backaches • Bursitis •

Fibromyalgia • Tendonitis • Joint pain/discomfort

• Sciatica • Swelling

Warnings:

for external use only

When using this product:

use only as directed • avoid contact with eyes or mucous membranes • do not apply to open wounds, damaged, or very sensitive skin • do not use if you are allergic to any of this product's active or inactive ingredients • do not apply bandage tightly or use heating pad • do not resume normal activity without the advice of a • medical professionall

Keep out of reach of children.

If swallowed, get medical help or contact the Poison Control Center right away.

Directions:

Adults and children 2 years and over:

•Shake well before using •Apply a sufficient amount of RelieveIt® Gel Max to cover the affected area • Repeat as needed • After applying, wash hands with soap and water Children under 2: consult a doctor

Other ingredients:

Carbomer 934P, Cyclomethicone, Ethyl Alcohol, Eucalyptus Oil, Glycerin, Hydroxypropyl Cellulose, C13/14 Isoparaffin, Isopropyl Myristate, Juniper Oil, Laureth-7, Polyacrylamide, Polyethylene Glycol 3350, Polysorbate 80, Propyl Gallate, Propylene Glycol, Purified Water, Resin, Sodium Hydroxide, Sorbic Acid, Sorbitan Monooleate, Xanthan Gum.

*These "Uses" have not been evaluated by the Food & Drug Administration.

Stop use and ask a doctor if:

excessive redness or irritation is present • condition worsens • pain persists for more than 7 days

If pregnant or breast feeding:

ask a health professional before use

Other Information

Store at 68-77°F (20-25°C)

Principal Display Panel and Drug Facts



RELIEVEIT GEL MAX

arnica montana, calendula officinalis, camphora (camphor), capsaicum oleoresin (capsaicin), mentha piperita (peppermint) gel

Product Information				
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:58133-953	
Route of Administration	TOPICAL			

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Ingredient Name Basis of Strength Strength

CAMPHOR (NATURAL) (UNII: N20 HL7Q941) (CAMPHOR (NATURAL) - UNII:N20 HL7Q941)	CAMPHOR (NATURAL)	7 [hp_M] in 1 mL
ARNICA MONTANA (UNII: O80TY208ZW) (ARNICA MONTANA - UNII:O80TY208ZW)	ARNICA MONTANA	7 [hp_M] in 1 mL
MENTHA PIPERITA (UNII: 79M2M2UDA9) (MENTHA PIPERITA - UNII:79M2M2UDA9)	MENTHA PIPERITA	7 [hp_M] in 1 mL
CAPSAICIN (UNII: S07O44R1ZM) (CAPSAICIN - UNII:S07O44R1ZM)	CAPSAICIN	0.02 [hp_M] in 1 mL
CALENDULA OFFICINALIS SEED OIL (UNII: 9JS8DS42SV) (CALENDULA OFFICINALIS SEED OIL - UNII:9JS8DS42SV)	CALENDULA OFFICINALIS SEED OIL	2 [hp_M] in 1 mL

Inactive Ingredients	
Ingredient Name	Strength
CYCLOMETHICONE (UNII: NMQ347994Z)	
METHYL ALCOHOL (UNII: Y4S76JWI15)	
EUCALYPTUS OIL (UNII: 2R04ONI662)	
LOW-SUBSTITUTED HYDROXYPROPYL CELLULOSE, UNSPECIFIED (UNII: 2165RE0K14)	
JUNIPER BERRY O IL (UNII: SZH16H44UY)	
POLYSORBATE 80 (UNII: 6OZP39ZG8H)	
BENZO IN RESIN (UNII: GK21SBA74R)	
SO DIUM HYDRO XIDE (UNII: 55X04QC32I)	
PROPYLENE GLYCOL 2-METHYLBUTYRATE (UNII: QH216IX8SV)	
POLYETHYLENE GLYCOL 3350 (UNII: G2M7P15E5P)	
PROPYL GALLATE (UNII: 8 D4SNN7V92)	
GLYCERIN (UNII: PDC6A3C0OX)	
ISOPROPYL MYRISTATE (UNII: 0 RE8 K4LNJS)	
POLYACRYLAMIDE (10000 MW) (UNII: E2KR9C9V2I)	
SORBIC ACID (UNII: X045WJ989B)	
XANTHAN GUM (UNII: TTV12P4NEE)	
WATER (UNII: 059QF0KO0R)	
AMMO NIUM LAURETH-7 SULFATE (UNII: 9LPV636QCV)	
CARBO MER 934 (UNII: Z135WT9208)	
SORBITAN MONOOLEATE (UNII: 06 XEA2VD56)	
C13-14 ISOPARAFFIN (UNII: E4F12ROE70)	

I	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:58133-953- 02	70 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product	08/15/2018		
2	NDC:58133-953- 16	474 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product	08/15/2018		
3	NDC:58133-953- 37	3780 mL in 1 BOTTLE, PUMP; Type 0: Not a Combination Product	08/15/2018		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
unapproved homeopathic		08/15/2018		
unapproved homeopathic		08/15/2018		

Labeler - Cosmetic Specialty Labs, Inc. (032973000)

Registrant - Cosmetic Specialty Labs, Inc. (032973000)

Establishment				
Name	Address	ID/FEI	Business Operations	
Cosmetic Specialty Labs, Inc.		032973000	manufacture(58133-953)	

Revised: 10/2018 Cosmetic Specialty Labs, Inc.