

QUALITY CHOICE CAMPHOR SPIRIT- camphor 10% liquid

Chain Drug Market Association

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.

Quality Choice Camphor Spirit

Drug Facts

Active Ingredient

Camphor 10%

Purpose

External Anagesic

Indication

For the temporary relief of minor aches, muscle and joint pain associated with arthritis, strains, brusies and simple backache.

Warnings

For external use only.

Do not drink. If swallowed, immediately give 3 or 4 glases of water. Do not induce vomiting. If vomiting occurs, give fluids again. Do not give anything by mouth to an unconscious or convulsing person. Get medical attention immediately.

Avoid contact with eyes or mucous membranes.

Do not apply to irritated skin.

Do not use undiluted product.

When using this product

Do not bandage tightly.

Discontinue use and consult a doctor if

condition worsens, or if excessive irritation develops.

symptoms persit for more than 7 days, or clear up and occur again within a few days.

Keep out of reach of children

In case of accidental ingestion, seek professional assistance or contact a Poison Control Center right away. In case of eye contact, flush eyes with running water for 15 minutes, get medical attention.

Directions

(dilute 3 parts olive oil and mix well)

Adults and children 2 yrs. and older. apply to th affected area not more than 3 or 4 times daily.

Children under 2 yrs of age: consult a doctor befere use.

Other Information

Flammable: Keep away from spark, heat and flame.

Inactive ingredients

Alcohol 84%, Purified Water

Principal Display Panel

Camphor Spirits, USP

Drug Facts

Active ingredient	Purpose
Camphor 10%.....	External analgesic

Indication ■ for temporary relief of minor aches, muscle and joint pain associated with arthritis, strains, bruises and simple backache.

Warnings For external use only.

- Do not drink. If swallowed, immediately give 3 or 4 glasses of water. Do not induce vomiting. If vomiting occurs, give fluids again. Do not give anything by mouth to an unconscious or convulsing person. Get medical attention immediately.
- Avoid contact with eyes or mucous membranes.
- Do not apply to irritated skin. ■ Do not use undiluted product.

When using this product do not bandage tightly. **Discontinue use and consult a doctor if** ■ condition worsens, or if excessive irritation develops. **Continued ▶**

(Drug facts continued)

- symptoms persist for more than 7 days, or clear up and occur again within a few days.

Keep out of reach of children. In case of accidental ingestion, seek professional assistance or contact a Poison Control Center right away. In case of eye contact, flush eyes with running water for 15 minutes, get medical attention.

Directions (Dilute 3 parts olive oil and mix well) ■ Adults and children 2 yrs. and older: apply to the affected area not more than 3 or 4 times daily. ■ Children under 2 yrs. of age: consult a doctor before use.

Other information ■ **Flammable:** Keep away from spark, heat and flame.

Inactive ingredients Alcohol 84%, Purified Water.

MADE IN USA

R071514RLG

6 35515 96711 3

2 fl OZ (59mL)

Satisfaction 100% Guaranteed

Distributed by C.D.M.A., Inc.®
43157 W. Nine Mile
Novi, MI 48376-0995
www.qualitychoice.com
Questions: 248-449-9300

QUALITY CHOICE CAMPHOR SPIRIT

camphor 10% liquid

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:63868-490
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CAMPHOR (NATURAL) (UNII: N20HL7Q941) (CAMPHOR (NATURAL) - UNII:N20HL7Q941)	CAMPHOR (NATURAL)	100 mg in 1 mL

Inactive Ingredients

Ingredient Name	Strength
-----------------	----------

ALCOHOL (UNII: 3K9958V90M)	
WATER (UNII: 059QF0KO0R)	

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:63868-490-02	473 mL in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	12/12/2017	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph not final	part348	01/01/2008	

Labeler - Chain Drug Market Association (011920774)

Registrant - Humco Holding Group, Inc. (825672884)

Establishment

Name	Address	ID/FEI	Business Operations
Humco Holding Group, Inc.		825672884	manufacture(63868-490) , analysis(63868-490) , pack(63868-490) , label(63868-490)

Revised: 12/2020

Chain Drug Market Association