BRONCOMAR DM- dextromethorphan hbr, guaifenes in liquid Dannso Corp./d.b.a. Essential Products

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.

Broncomar DM

Active Ingredients:(in each 5 ml.)	Purpose
Dextromethorphan Hydrobromide 10	mgCough Suppressant
Guaifenesin 100 mg	Expectorant

Uses:

- Helps loosen phlegm (mucus) and thin bronchial secretions to drain bronchial tubes and make cough more productive.
- Temporarily relieves cough due to minor throat and bronchial irritation as may occur with the common cold.

Warnings

Do not exceed recommended dosage

Do not use

- If you are now taking a prescription monoamine oxidase inhibitor (MAOI) (certain drugs for depression, psychiatric conditions, or Parkison's disease) or for 2 weeks after stopping MAOI drug.
- If you do not know if your prescription drug contains MAOI as your doctor or pharmacist before taking this product.
- If you have a chronic pulmonary disease or shortness of breath unless directed by a doctor.
- Avoid alcoholic beverage while taking this product.

Stop use and ask a doctor

- Nervousness, dizziness or sleeplessness occurs.
- Cough persists more than 1 week, tends to recur or is accompanied by a fever, rash or persistent headache. A persistent cough may be signserious condition.

Ask doctor before use if you have

- Cough that occurs with too much phlegm(mucus)
- Cough that last or is a chronic such as occurs with smoking, asthma, chronic bronchitis or emphysema.

If pregnant or breast-feeding ask a health professional before use.

Keep out of reach of children.

In case of overdose, get medical help or contact a Poison Control Center right away.

Directions

- Do not exceed 6 doses in any 24 hour period.
- Shake well before use.

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Adults and Children 12 years and over	10 ml (2 tsps) every 4 hours
Children 6 to under 12 years of age	5 ml (1 tsp) every 4 hours
Children under 6 years of age	Do not use

Other Information:

- Each 5 mls contains: sodium 8 mg
- Store between 15 30 degrees Celsius (59 86 Fahrenheit).
- Tamper Evident Feature:Do not use if seal under cap is torn, broken or missing.

Inactive Ingredient

Blue Cohosh, Citric Acid, Echinacea, Eucalyptus Oil, Ginkgo Biloba, Golden Seal Root, Honey Pure, Honey flavor, Horehound Herb, Licorice Root, Menthol, Mullein, Myrrh, Potassium Sorbate, Slippery Elm Bark, Sodium Benzoate, Propylene Glycol, Purified Water, Sodium Chloride, Sucralose, Wild Cherry bark and Zinc Sulfate.

Questions or Comments

Call Weekdays from 9:30 AM to 5PM EST at Tel 305-261-762

Distributed by

Essential Products Miami FL 33126

www.jjjdistributors.com

Made in U.S.A.



BRONCOMAR DM dextromethorphan hbr, guaifenesin liquid Product Information Product Type HUMAN OTC DRUG Item Code (Source) NDC:70242-101

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
DEXTROMETHORPHAN HYDROBROMIDE (UNII: 9 D2RTI9 KYH) (DEXTROMETHORPHAN - UNII:7355X3ROTS)	DEXTROMETHORPHAN HYDROBROMIDE	10 mg in 5 mL	
GUAIFENESIN (UNII: 495W7451VQ) (GUAIFENESIN - UNII:495W7451VQ)	GUAIFENESIN	100 mg in 5 mL	

Inactive Ingredients	
Ingredient Name	Strength
CAULOPHYLLUM THALICTROIDES ROOT (UNII: JTJ6 HH6 YEH)	
CITRIC ACID MONOHYDRATE (UNII: 2968PHW8QP)	
ECHINACEA (UNII: 4N9 P6 CC1DX)	
EUCALYPTUS OIL (UNII: 2R04ONI662)	
GINKGO (UNII: 19FUJ2C58T)	
GOLDENSEAL (UNII: ZW3Z11D0JV)	
HO NEY (UNII: Y9 H1V576 FH)	
HOREHOUND (UNII: K08036XEJV)	
LICORICE (UNII: 61ZBX54883)	
MENTHOL (UNII: L7T10EIP3A)	
VERBASCUM THAPSUS (UNII: C9TD27U172)	
MYRRH (UNII: JC71GJ1F3L)	
POTASSIUM SORBATE (UNII: 1VPU26JZZ4)	
ULMUS RUBRA BARK (UNII: 91QY4PXU8Q)	
SODIUM BENZOATE (UNII: OJ245FE5EU)	
PROPYLENE GLYCOL (UNII: 6 DC9 Q167V3)	
WATER (UNII: 059QF0KO0R)	
SODIUM CHLORIDE (UNII: 451W47IQ8X)	
SUCRALOSE (UNII: 96K6UQ3ZD4)	
PRUNUS SEROTINA BARK (UNII: 5D48 E975HA)	
ZINC SULFATE (UNII: 89 DS0 H96 TB)	

Packaging			
# Item Code	Package Description	Marketing Start Date	Marketing End Date
1 NDC:70242-101-06	1 in 1 CARTON	0 1/0 1/20 15	
1	177 mL in 1 BOTTLE; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
OTC monograph final	part341	10/01/2001	

Labeler - Dannso Corp./d.b.a. Essential Products (059741071)

Registrant - Dannso Corp./d.b.a. Essential Products (059741071)

Establishment				
Name	Address	ID/FEI	Business Operations	
All Pharma LLC		078572520	MANUFACTURE(70242-101)	

Revised: 1/2016 Dannso Corp./d.b.a. Essential Products