LOHIST DM- brompheniramine maleate, dextromethorphan hydrobromide and phenylephrine hydrochloride liquid Larken Laboratories, Inc.

LoHist DM

Drug Facts

Active ingredients

(In each 5 mL teaspoonful)

Brompheniramine Maleate, USP 2 mg

Dextromethorphan HBr, USP 10 mg

Phenylephrine HCl, USP 5 mg

Purpose

Brompheniramine Maleate Antihistamine

Dextromethorphan HBr Antitussive (cough suppressant)

Phenylephrine HCI Nasal decongestant

Uses

Temporarily relieves these symptoms due to hay fever (allergic rhinitis):

- cough due to minor throat and bronchial irritation
- runny nose
- sneezing
- itchy, watery eyes
- itching of the nose or throat
- Temporarily relieves nasal congestion due to the common cold, hay fever or other upper respiratory allergies, or associated with sinusitis.
- Temporarily restores freer breathing through the nose

Warnings

Do not use

- to sedate a child or make a child sleepy
- if you are now taking a prescription monoamine oxidase inhibitor (MAOI) (certain drugs for depression, psychiatric, or emotional conditions, or Parkinson's disease), or for 2 weeks after stopping the MAOI drug. If you do not know if your prescription drug contains an MAOI, ask a doctor or pharmacist before taking this drug.

Ask a doctor before use if you have

- heart disease
- high blood pressure
- thyroid disease
- diabetes
- glaucoma
- trouble urinating due to an enlarged prostate gland
- a breathing problem such as emphysema or chronic bronchitis
- a cough that lasts or is chronic such as occurs with smoking, asthma or emphysema
- a cough that occurs with too much phlegm (mucus)

Ask a doctor or pharmacist before use if you are

- taking any other nasal decongestant or stimulant
- taking sedatives or tranquilizers

When using this product

Do not exceed recommended dosage.

- marked drowsiness may occur
- avoid alcoholic beverages
- alcohol, sedatives, and tranquilizers may increase drowsiness
- be careful when driving a motor vehicle or operating machinery
- excitability may occur, especially in children

Stop use and ask a doctor if

- nervousness, dizziness, or sleeplessness occur.
- cough or nasal congestion persists for more than 1 week, tends to recur, or is accompanied by a fever, rash or persistent headache. These could be signs of a serious condition.

If pregnant or breast-feeding

ask a health professional before use.

Keep out of the reach of children

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

Directions

Do not exceed 6 doses in a 24-hour period

AgeDoseAdults and children2 teaspoonsful (10over 12 years ofmL) every 4 hours

age Children 6 to under 1 teaspoonful (5 12 years of age mL) every 4 hours Children under 6 Ask your doctor years of age

Other Information

- store at 20°-25°C (68°-77°F)
- very low sodium, contains 1 mg sodium per teaspoonful (5 mL)

Inactive Ingredients

Benzoic acid, edetate disodium, FD&C Red #40, propylene glycol, purified water, saccharin sodium, sorbitol solution, and strawberry flavoring

Questions or Comments

Call 1-601-855-7678 weekdays from 9:00 am to 4:00 pm CST or go to http://www.larkenlabs.com.

Principal Display Panel

16 oz Bottle Label



Drug Facts (continue	d)
Stop use and ask a doctor ■ nervousness, dizziness, o ■ cough or nasal congestion week, tends to recur, or is a rash or persistent headache serious condition.	r sleeplessness occur. n persists for more than 1 ccompanied by a fever,
If pregnant or breast-feedir before use.	ng, ask a health professional
Keep out of reach of childre get medical help or contact immediately.	
<i>Directions</i> do not exceed 6 doses in a 2 Age	24-hour period Dose
Adults and children over 12 years of age	2 teaspoonsful (10 mL) every 4 hours
Children 6 to under 12 years of age	1 teaspoonful (5 mL) every 4 hours
Children under 6 years of age	Ask your doctor
Other information ∎store at 20°- 25°C (68°- 7 ∎very low sodium, contains 1	7°F) mg sodium per teaspoonful (5 mL)
Inactive ingredients Benzoic acid, edetate disodi glycol, purified water, sacch and strawberry flavoring	um, FD&C Red #40, propylene arin sodium, sorbitol solution
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LOHIST DM

7

brompheniramine maleate, dextromethorphan hydrobromide and phenylephrine hydrochloride liquid

Product Information					
Product Type	HUMAN OTC DRUG	Item Code (Source)		NDC:68047-129	
Route of Administration	ORAL				
	Mainter.				
Active Ingredient/Active	моюту				
Ingredient Name			Basis of Strength		Strength
DEXTROMETHORPHAN HYDROBROMIDE (UNII: 9D2RTI9KYH) (DEXTROMETHORPHAN - UNII:7355X3ROTS)			DEXTROMETHORPHAN HYDROBROMIDE		10 mg in 5 mL
PHENYLEPHRINE HYDROCHLOR UNII:1WS297W6MV)	IDE (UNII: 04JA59TNSJ) (PHE	NYLEPHRINE -	PHENYLEPHRINE HYDROCHLORIDE		5 mg in 5 mL
BROMPHENIRAMINE MALEATE (UNII:H57G17P2FN)	UNII: IXA7C9ZN03) (BROMPH	IENIRAMINE -	BROMPHENIRAMIN MALEATE	IE	2 mg in 5 mL

Ingredient Name

Strength

EDETATE DISODIUI	M (UNII: 71	⁻ LD91C86K)				
FD&C RED NO. 40	(UNII: WZE	39127XOA)				
PROPYLENE GLYCO)L (UNII: 6	DC9Q167V3)				
WATER (UNII: 059QF	OKOOR)					
SACCHARIN SODIU	M (UNII: S	B8ZUX40TY)				
SORBITOL (UNII: 50	6T60A25R)				
BENZOIC ACID (UN	II: 85 KN0B	OMIM)				
Product Chara	cterist	ics				
Color		red	Sco	ore		
Shape			Siz	e		
Flavor	avor STRAWBERRY Imprint		orint Code			
Contains						
Packaging						
5						
		Package Description		Marketing Start Date	Marketin Date	
# Item Code	473 mL in Product	Package Description 1 BOTTLE; Type 0: Not a Combin	nation			
# Item Code			nation	Date		
 # Item Code 1 NDC:68047-129- 16 	Product	1 BOTTLE; Type 0: Not a Combin	nation	Date		
# Item Code	Product	1 BOTTLE; Type 0: Not a Combin	nation	Date		
 # Item Code 1 NDC:68047-129- 16 	Product nform	1 BOTTLE; Type 0: Not a Combin		Date		e ng End
 # Item Code 1 NDC:68047-129-16 Marketing I Marketing 	Product nform App	1 BOTTLE; Type 0: Not a Combin Nation lication Number or Monog		Date 06/01/2012 Marketing Start	Dato	e ng End

Labeler - Larken Laboratories, Inc. (149484540)

Registrant - Larken Laboratories, Inc. (149484540)

Revised: 10/2024

Larken Laboratories, Inc.