BASIC CARE MUCUS ER MAX- guaifenesin tablet, multilayer, extended release Amazon.com Services LLC

Amazon Mucus-ER Max Drug Facts

Active ingredient (in each extended-release tablet)

Guaifenesin 1200 mg

Purpose

Expectorant

Uses

• helps loosen phlegm (mucus) and thin bronchial secretions to rid the bronchial passageways of bothersome mucus and make coughs more productive

Warnings

Do not use

• for children under 12 years of age

Ask a doctor before use if you have

- persistent or chronic cough such as occurs with smoking, asthma, chronic bronchitis, or emphysema
- cough accompanied by too much phlegm (mucus)

Stop use and ask a doctor if

 cough lasts more than 7 days, comes back, or occurs with fever, rash, or persistent headache.

These could be signs of a serious illness.

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 (1-800-222-1222).

Directions

- do not crush, chew, or break extended-release tablet
- take with a full glass of water
- this product can be administered without regard for timing of meals
- adults and children 12 years of age and over: 1 extended-release tablet every 12 hours. Do not exceed 2 extended-release tablets in 24 hours.
- children under 12 years of age: do not use

Other information

• store at 20-25°C (68-77°F)

Inactive ingredients

carbomer homopolymer type B, copovidone, FD&C blue #1 aluminum lake, hypromellose, magnesium stearate, microcrystalline cellulose, silicon dioxide, sodium starch glycolate

Questions or comments?

1-800-719-9260

Package/Label Principal Display Panel

amazon

basic care

Maximum Strength

Compare to Maximum Strength Mucinex[®] active ingredient

Mucus-ER Max

Guaifenesin Extended-Release Tablets, 1200 mg

Expectorant

12 Hour

- Relieves Chest Congestion
- Thins and Loosens Mucus
- Immediate and Extended Release actual size

42 Extended-Release Tablets



BASIC CARE MUCUS ER MAX

guaifenesin tablet, multilayer, extended release

Product Information						
Product Type	HUMAN OTC DRUG	Item Code (Source) NDC:		NDC:722	72288-613	
Route of Administration	ORAL					
Active Ingredient/Active	Molety					
Ingre		Basis of Strength		Strength		
GUAIFENESIN (UNII: 495W7451VQ)	7451VQ)	GUAIFENESIN		1200 mg		
In a stine In one dis sta						
Inactive Ingredients						
	Ingredient Nam	e			Strength	
CARBOMER HOMOPOLYMER TYP HHT01ZNK31)	PE B (ALLYL PENTAERYTH	IRITOL CROSSL	INKED) (UNII:			
COPOVIDONE K25-31 (UNII: D9C3	330MD8B)					
FD&C BLUE NO. 1 ALUMINUM LA	KE (UNII: J9EQA3S2JM)					

M	PROMELLOSE, AGNESIUM STEA	RATE (UNII:	•	-			
М	CROCRYSTALLI	NE CELLULO	DSE (UNII: C	P1R32D61U)			
SI	LICON DIOXIDE	(UNII: ETJ7Z6	6XBU4)				
sc	DIUM STARCH	GLYCOLATE	TYPE A (U	NII: H8AV0SQX4D)			
D	roduct Chara	etorictio	-				
			-	-		no score	
Color		BLUE		Score			
Shape			OVAL		Size		
	avor			Imprint Code	Imprint Code		
Сс	ontains						
Pa	ackaging						
#	ltem Code	I	Package Description		Marketing Start Date	: Marketing End Date	
1	NDC:72288-613- 55	6 in 1 CART	RTON		04/03/2025		
		7 in 1 BLISTER PACK; Type 0: Not a Combination Product					
1		Product	,,,				
1 2	NDC:72288-613- 74				04/03/2025		
		Product 2 in 1 CART	ON	ype 0: Not a Combination	04/03/2025		
2		Product 2 in 1 CART 7 in 1 BLIST	ON		04/03/2025		
2		Product 2 in 1 CART 7 in 1 BLIST	ON		04/03/2025		
2		Product 2 in 1 CART 7 in 1 BLIST Product	ON ER PACK; T <u>i</u>		04/03/2025		
2	74	Product 2 in 1 CART 7 in 1 BLIST Product	ON ER PACK; Ty ation cation Nu		04/03/2025	t Marketing End Date	

Labeler - Amazon.com Services LLC (128990418)

Revised: 7/2025

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