TUSSIN CF- dextromethorphan hydrobromide, guaifenesin, phenylephrine hydrochloride solution Publix Super Markets Inc

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.

Publix Super Markets, Inc. Tussin CF Drug Facts

Active ingredients (in each 10 mL)

Dextromethorphan HBr, USP 20 mg Guaifenesin, USP 200 mg Phenylephrine HCl, USP 10 mg

Purposes

Cough suppressant Expectorant Nasal decongestant

Uses

- helps loosen phlegm (mucus) and thin bronchial secretions to drain bronchial tubes
- temporarily relieves these symptoms occurring with a cold:
- nasal congestion
- cough due to minor throat and bronchial irritation

Warnings

Do not use

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 product.

Ask a doctor before use if you have

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

When using this product

do not use more than directed

Stop use and ask a doctor if

- you get nervous, dizzy, or sleepless
- symptoms do not get better within 7 days or are accompanied by fever
- cough lasts more than 7 days, comes back, or is accompanied by fever, rash, or persistent headache. A persistent cough may be a sign of a serious condition.

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 take more than 6 doses in any 24-hour period
- measure only with dosing cup provided
- keep dosing cup with product
- mL = milliliter
- this adult product is not intended for use in children under 12 years of age

age	dose
adults and children 12 years and over	10 mL every 4 hours
children under 12 years	do not use

Other information

- each 10 mL contains: sodium 5 mg
- store at 20-25°C (68-77°F). Do not refrigerate.

Inactive ingredients

anhydrous citric acid, edetate disodium, FD&C red no. 40, flavor, glycerin, propylene glycol, purified water, sodium benzoate, sodium citrate, sorbitol solution, sucralose

Principal Display Panel

ADULT tussin CF COUGH SUPPRESSANT (DEXTROMETHORPHAN HBr) EXPECTORANT (GUAIFENESIN) NASAL DECONGESTANT (PHENYLEPHRINE HCl) Relieves: Cough Nasal congestion

Mucus

Compare to the active ingredients in Robitussin® Multi-Symptom Cold FOR AGES 12 & OVER NON-DROWSY

4 FL OZ (118 mL)



TUSSIN CF

Product Informa	tion						
Product Type		HUMAN OTC DRUG	Item Code	em Code (Source)		NDC:56062-516	
Route of Administra	ation	ORAL					
		0.000					
Active Ingredien	t/Active M	oiety					
	Ingredient Name			Basis of Stre	ngth	Strength	
DEXTROMETHORPHAN HYDROBROMIDE (UNII: 9D2RTI9KYH) (DEXTROMETHORPHAN - UNII:7355X3ROTS)				DEXTROMETHORPHAN 20 m		20 mg in 10 mL	
GUAIFENESIN (UNII: 495W7451VQ) (GUAIFENESIN - UNII:495W7451VQ)				GUAIFENESIN		200 mg in 10 mL	
PHENYLEPHRINE HYDRO CHLORIDE (UNII: 04JA59TNSJ) (PHENYLEPHRINE - PHENYLEPHRINE UNII: 1WS297W6 MV) HYDROCHLORIDE						10 mg in 10 mL	
Inactive Ingredie	ents						
		Ingredient Name			Sti	rength	
ANHYDRO US CITRIC	C ACID (UNII:)	KF417D3PSL)					
EDETATE DISODIUM	I (UNII: 7FLD9	IC86K)					
D&C RED NO.40 (U	NII: WZB91272	XOA)					
G LYCERIN (UNII: PDC	C6A3C0OX)						
PROPYLENE GLYCO	L (UNII: 6DC9	Q167V3)					
WATER (UNII: 059QF	0KO0R)						
SO DIUM BENZO ATE	(UNII: OJ245F	E5EU)					
		· · · · · · · · · · · · · · · · · · ·					
	JNII: 1Q73Q2JU						
SODIUM CITRATE (U							
SODIUM CITRATE (U SORBITOL (UNII: 506 SUCRALOSE (UNII: 96	6T60A25R)						
SODIUM CITRATE (U SORBITOL (UNII: 506 SUCRALOSE (UNII: 96	5 T6 0 A25 R) 6 K6 UQ 3Z D4)						
GODIUM CITRATE (U GORBITOL (UNII: 506 GUCRALOSE (UNII: 96 Product Characte	5 T6 0 A25 R) 6 K6 UQ 3Z D4)		Score				
GODIUM CITRATE (U GORBITOL (UNII: 506 GUCRALOSE (UNII: 96 Product Characte Color	5 T6 0 A25 R) 6 K6 UQ 3Z D4)	JLR)	Score Size				
GODIUM CITRATE (U GORBITOL (UNII: 506 GUCRALOSE (UNII: 96 Product Characte Color Shape	5 T6 0 A25 R) 6 K6 UQ 3Z D4)	JLR)		e			
SODIUM CITRATE (U SORBITOL (UNII: 506	5 T6 0 A25 R) 6 K6 UQ 3Z D4)	JLR) RED	Size	e			
SODIUM CITRATE (U SORBITOL (UNII: 506 SUCRALOSE (UNII: 90 Product Characte Color Shape Flavor Contains	5 T6 0 A25 R) 6 K6 UQ 3Z D4)	JLR) RED	Size	e			
SODIUM CITRATE (U SORBITOL (UNII: 506 SUCRALOSE (UNII: 90 Product Characte Color Shape Flavor Contains Packaging	5 T6 0 A25 R) 6 K6 UQ 3Z D4)	JLR) RED	Size Imprint Code	e Marketing Start Date	Marketi	ng End Dat	
ODIUM CITRATE (U ORBITOL (UNII: 506 UCRALOSE (UNII: 90 Product Characte Color Shape Packaging Item Code	6T60A25R) 6K6UQ3ZD4) eristics	JLR) ILR) RED CHERRY Backage Description	Size Imprint Code		Marketin	ng End Dat	
SODIUM CITRATE (USORBITOL (UNII: 506) SUCRALOSE (UNII: 90) Product Characte Color Shape Flavor Contains Packaging Item Code NDC:56062-516-26	6T60A25R) 6K6UQ3ZD4) eristics	JLR) ILR) RED CHERRY Backage Description	Size Imprint Code N 0	Marketing Start Date	Marketin	ng End Dat	
SODIUM CITRATE (US) SORBITOL (UNII: 506 SUCRALOSE (UNII: 96 SODIUCRALOSE (UNII: 96 Solor Shape Flavor Contains Packaging I tem Code NDC:56062-516-26	6 T60 A25R) 6 K6 UQ 3ZD4) eristics 1 in 1 CARTO 118 mL in 1 B	JLR) RED CHERRY Package Description N OTTLE; Type 0: Not a Combin	Size Imprint Code Mation Product	Marketing Start Date	Marketin	ng End Dat	
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Labeler - Publix Super Markets Inc (006922009)

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