

EMETROL CHEWABLES MIXED BERRY- sodium citrate dihydrate tablet, chewable

WellSpring Pharmaceutical Corporation

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

Emetrol Chewables Rapid Nausea Relief

Active ingredients

- Sodium citrate dihydrate USP 230 mg

Purpose

Upset stomach reliever/antacid

Uses

For the relief of nausea associated with upset/sour stomach, including that due to overindulgence in food and drink.

Warnings

Do not use if you have

- Hereditary Fructose Intolerance (HFI). This product contains fructose.

Ask a doctor before use if you

- have diabetes because this product contains sugar
- are on a sodium-restricted diet

Ask a doctor or pharmacist before use if

- you are taking any other medications. This product may interact with certain prescription drugs.

When using this product,

- do not take more than 24 tablets in a 24-hour period.

Stop use and ask a doctor if

- nausea lasts more than two weeks or recurs frequently.

If pregnant or breast-feeding,

- ask a health professional before use. In case of overdose, get medical help or contact a Poison Control Center right away.

Keep out of reach of children

Directions

- Adults: 2-4 tablets.
- Children: Consult a doctor for appropriate dosage.
- Chew tablets completely. Do not swallow whole
- Dosage may be repeated after 15 minutes, not to exceed 24 tablets in a 24-hour period unless advised by a doctor.
- Read all package directions and warnings before use and use only as directed.
- These tablets are intended for use by normally healthy persons only.
- Persons under 18 years of age should use only as directed by a doctor.

Other information

- **Each table contains:** Sodium 60mg
- Store at room temperature
- Protect product from excessive heat, cold and moisture
- Note: This product is not intended as a substitute for a balanced nutritional diet or as an electrolyte replenishment.
- Do not use if printed seal under cap is broken or missing.

Inactive ingredients

dextrose, flavor, fructose, hydroxypropyl cellulose, magnesium stearate, maltodextrin, silica, sodium carboxymethylcellulose, sucralose.

Questions or Comments?

1-844-241-5454 or Emetrol.com

Distributed By

WellSpring Pharmaceutical Corporation
Sarasota, FL 34243
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PACKAGE LABEL

1 Pharmacist Recommended Brand

New

Non-Drowsy

Emetrol Chewable Rapid Nausea Relief

Treats the source of Nausea*

*Due to overindulgence in food and drink

Powerful, Soothing Relief

Mixed Berry flavored
60 Chewable Tablets



Emetrol Chewables Mixed Berry Flavor

Side Panels

Emetrol Chewables Rapid Nausea Relief

Non-Drowsy

Treats the source of Nausea*

Specially Formulated for Adults

Great-tasting Lemon Chewables

*Due to overindulgence in food and drink

Emetrol Chewables relieves stomach discomfort with:

No Acetaminophen

No Aspirin

No Caffeine

EMETROL CHEWABLES MIXED BERRY			
sodium citrate dihydrate tablet, chewable			
Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:65197-420
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
TRISODIUM CITRATE DIHYDRATE (UNII: B22547B95K) (ANHYDROUS CITRIC ACID - UNII:XF417D3PSL)	ANHYDROUS CITRIC ACID	230 mg

Inactive Ingredients

Ingredient Name	Strength
DEXTROSE (UNII: IY9XDZ35W2)	
FRUCTOSE (UNII: 6YSS42VSEV)	
HYDROXYPROPYL CELLULOSE, UNSPECIFIED (UNII: 9XZ8H6N6OH)	
MAGNESIUM STEARATE (UNII: 70097M6I30)	
MALTODEXTRIN (UNII: 7CVR7L4A2D)	
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
CARBOXYMETHYLCELLULOSE SODIUM, UNSPECIFIED FORM (UNII: K679OBS311)	
SUCRALOSE (UNII: 96K6UQ3ZD4)	

Product Characteristics

Color	white	Score	no score
Shape	ROUND	Size	16mm
Flavor	BERRY	Imprint Code	
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:65197-420-60	1 in 1 BOX	07/01/2023	
1		60 in 1 BOTTLE; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved drug other		07/01/2023	

Labeler - WellSpring Pharmaceutical Corporation (110999054)

Revised: 3/2023

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