MOTION SICKNESS RELIEF- dimenhydrinate tablet Chain Drug Marketing Association, Inc.

Quality Choice 44-198

Active ingredient (in each tablet)

Dimenhydrinate 50 mg

Purpose

Antiemetic

Uses

for prevention and treatment of these symptoms associated with motion sickness:

- nausea
- vomiting
- dizziness

Warnings

Do not use

for children under 2 years of age unless directed by a doctor.

Ask a doctor before use if you have

- a breathing problem such as emphysema or chronic bronchitis
- glaucoma
- difficulty in urination due to enlargement of the prostate gland

Ask a doctor or pharmacist before use if you are

taking sedatives or tranquilizers.

When using this product

- marked drowsiness may occur
- alcohol, sedatives, and tranquilizers may increase drowsiness
- use caution when driving a motor vehicle or operating machinery
- avoid alcoholic beverages

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

 to prevent motion sickness, the first dose should be taken one-half to one hour before starting activity

adults and children 12 years and over	1 to 2 tablets every 4-6 hours; do not exceed 8 tablets in 24 hours, or as directed by a doctor
children 6 to under 12 years	1/2 to 1 tablet every 6-8 hours; do not exceed 3 tablets in 24 hours, or as directed by a doctor
children 2 to under 6 years	1/2 tablet every 6-8 hours; do not exceed 1&1/2 tablets in 24 hours, or as directed by a doctor

Other information

- each tablet contains: calcium 35 mg
- TAMPER EVIDENT: DO NOT USE IF OUTER PACKAGE IS OPENED OR BLISTER IS TORN OR BROKEN
- store at 25°C (77°F); excursions permitted between 15°-30°C (59°-86°F)
- protect from moisture
- see end flap for expiration date and lot number

Inactive ingredients

croscarmellose sodium, dibasic calcium phosphate dihydrate, magnesium stearate, microcrystalline cellulose, silicon dioxide, stearic acid

Questions or comments?

1-800-426-9391

Principal Display Panel

NDC 83324-032-12

QC® QUALITY CHOICE

*Compare to the Active Ingredient in Dramamine® Original Formula

Motion Sickness Relief Original Formula Dimenhydrinate 50 mg | Antiemetic Prevents: Nausea, Vomiting & Dizziness

for Children & Adults

12 Tablets

actual size

TAMPER EVIDENT: DO NOT USE IF PACKAGE IS OPENED OR IF BLISTER UNIT IS TORN, BROKEN OR SHOWS ANY SIGNS OF TAMPERING

*This product is not manufactured or distributed by Medtech Products Inc., owner of the registered trademark Dramamine® Original Formula.

50844 REV0518A19802

SATISFACTION GUARANTEED 100% QC

Distributed by CDMA, Inc. Novi, MI 48375 www.qualitychoice.com Questions: 800-935-2362





Motion Sickness Relief

Original Formula

Dimenhydrinate 50 mg | Antiemetic

Prevents: Nausea, Vomiting & Dizziness for Children & Adults

NDC 83324-032-12



Compare to the Active Ingredient in Dramamine Original Formula

Motion Sickness Relief

Original Formula

Dimenhydrinate 50 mg | Antiemetic

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12 Tablets

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children 12 years and over	not exceed 8 tablets in 24 hours, or or as directed by a doctor

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No print/No varnish Lot & Exp date TAMPER EVIDENT: DO NOT USE IF PACKAGE IS OPENED OR IF BLISTER UNIT IS TORN, BROKEN OR SHOWS ANY SIGNS OF TAMPERING

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MOTION SICKNESS RELIEF

dimenhydrinate tablet

Product Information			
Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:83324-032
Route of Administration	ORAL		

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
DIMENHYDRINATE (UNII: JB937PER5C) (DIPHENHYDRAMINE - UNII:8GTS82S83M, 8-CHLOROTHEOPHYLLINE - UNII:GE2UA340FM)	DIMENHYDRINATE	50 mg		

Inactive Ingredients	
Ingredient Name	Strength
CROSCARMELLOSE SODIUM (UNII: M28OL1HH48)	
DIBASIC CALCIUM PHOSPHATE DIHYDRATE (UNII: O7TSZ97GEP)	
MAGNESIUM STEARATE (UNII: 70097M6I30)	
MICROCRYSTALLINE CELLULOSE (UNII: OP1R32D61U)	
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
STEARIC ACID (UNII: 4ELV7Z65AP)	

Product Characteristics				
Color	white	Score	2 pieces	
Shape	ROUND	Size	9mm	
Flavor		Imprint Code	44;198	
Contains				

P	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:83324-032- 12	2 in 1 CARTON	04/04/2024		
1		6 in 1 BLISTER PACK; Type 0: Not a Combination Product			

2	NDC:83324-032- 24	4 in 1 CARTON	04/04/2024	
2		6 in 1 BLISTER PACK; Type 0: Not a Combination Product		

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
OTC Monograph Drug	M009	04/04/2024			

Labeler - Chain Drug Marketing Association, Inc. (011920774)

Establishment			
Name	Address	ID/FEI	Business Operations
LNK International, Inc.		038154464	pack(83324-032)

Establishment			
Name	Address	ID/FEI	Business Operations
LNK International, Inc.		832867837	manufacture(83324-032) , pack(83324-032)

Establishment			
Name	Address	ID/FEI	Business Operations
LNK International, Inc.		117025878	manufacture(83324-032)

Revised: 5/2024 Chain Drug Marketing Association, Inc.