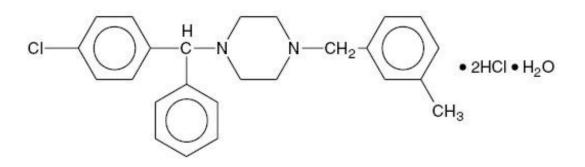
## MECLIZINE HYDROCHLORIDE- meclizine hydrocloride tablet Aidarex Pharmaceuticals LLC

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# MECLIZINE HYDROCHLORIDE TABLETS, USP Rx only

### DESCRIPTION

Chemically, Meclizine HCl is  $1-(p-chloro-\alpha-phenylbenzyl)-4-(m-methylbenzyl)$  piperazine dihydrochloride monohydrate.



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C_{25}H_{27}ClN_2. 2HCl. H_2O 		 M.W.
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481.88

Meclizine HCI Tablets, USP are available in two different strengths: 12.5 mg and 25 mg. In addition each tablet contains the following inactive ingredients: Colloidal Silicon Dioxide, Croscarmellose Sodium, Lactose Monohydrate, Magnesium Stearate, Microcrystalline Cellulose. Also, Meclizine HCI Tablets USP, 12.5 mg contains FD&C Blue #1 Aluminum Lake (11-13%) and Meclizine HCI Tablets USP, 25 mg contains D&C Yellow #10 Aluminum Lake (15-20%).

### **CLINICAL PHARMACOLOGY**

Meclizine Hydrochloride is an antihistamine which shows marked protective activity against nebulized histamine and lethal doses of intravenously injected histamine in guinea pigs. It has a marked effect in blocking the vasodepressor response to histamine, but only a slight blocking action against acetylcholine. Its activity is relatively weak in inhibiting the spasmogenic action of histamine on isolated guinea pig ileum.

### INDICATIONS AND USAGE

For the management of nausea and vomiting, and dizziness associated with motion sickness.

### **CONTRAINDICATIONS**

Meclizine Hydrochloride is contraindicated in individuals who have shown a previous hypersensitivity to it.

### WARNINGS

Since drowsiness may, on occasion, occur with use of this drug, patients should be warned of this possibility and cautioned against driving a car or operating dangerous machinery.

Patients should avoid alcoholic beverages while taking this drug.

Due to its potential anticholinergic action, this drug should be used with caution in patients with asthma, glaucoma or enlargement of the prostate gland.

# PRECAUTIONS

# PREGNANCY, Teratogenic Effects

Pregnancy Category B. Reproduction studies in rats have shown cleft palates at 25-50 times the human dose. Epidemiological studies in pregnant women, however, do not indicate that medicine increases the risk of abnormalities when administered during pregnancy. Despite the animal findings, it would appear that the possibility of fetal harm is remote. Nevertheless, meclizine, or any other medication, should be used during pregnancy only if clearly necessary.

### Pediatric Use

Clinical studies establishing safety and effectiveness in children have not been done; therefore, usage is not recommended in children under 12 years of age.

# **ADVERSE REACTIONS**

Drowsiness, dry mouth and, on rare occasions, blurred vision have been reported.

# DOSAGE AND ADMINISTRATION

### **Motion Sickness**

The initial dose of 25 to 50 mg of Meclizine HCI should be taken one hour prior to travel for protection against motion sickness. Thereafter, the dose may be repeated every 24 hours for the duration of the journey.

# HOW SUPPLIED

Meclizine HCI Tablets, USP are available in the following strengths and package sizes:

25 mg (Yellow, oval-shaped, scored, debossed with TL121)

Bottles of 8	NDC 332610718-08
Bottles of 10	NDC 332610718-10
Bottles of 20	NDC 332610718-20
Bottles of 30	NDC 332610718-30
Bottles of 60	NDC 332610718-60

Store at 20-25°C (68-77°F) (See USP Controlled Room Temperature].

Manufactured By:

Jubilant Cadista Pharmaceuticals Inc. Salisbury, MD 21801, USA.

Repackaged By : Aidarex Pharmaceuticals LLC, Corona, CA 92880

# PRINCIPAL DISPLAY PANEL

Л СН ОF 59-86F)	Packaged and Distributed by:	AIDAREX PH	MECLIZINE 25mg 30	PEEL HER LVEILVA	
BITS DISPENSING WITHOUT NSERT.KEEP OUT OF REACH LED ROOM TEMP 15-30C (59		25mg	CLIZINE EACH TABLET CONTAINS THE FOLLOWING ACTIVE INGREDIENTS: MECLIZIN E	NDC: 33261-0718-30 RXQLS0000 MECLIZINE 25mg 30 NDC: 33261-0718-30 RX QLS0000	E
CAUTION: FEDERAL LAW PROH PRESCRIPTION. SEE PACKAGE CHILDREN. STORE AT CONTRO	T		YELLOW OVAL TABLET W/TL PARTIAL SCORE 121 ON ONE SIDE	MECLIZINE 25mg 30 NDC: 33261-0718-30 RX QLS0000 MECLIZINE 25mg 30 NDC: 33261-0718-30 RX QLS0000	TAAHO LAAN TIIB

### NDC 33261-0718-30

# Meclizine Hydrochloride Tablets, USP

25 mg

**Rx only** 

**30 Tablets** 

Each tablet contains 25 mg of meclizine HCl.

### DOSAGE AND USE

See accompanying prescribing information

# **MOTION SICKNESS:**

25 mg to 50 mg daily.

Dispense in tight, light-resistant containers (USP).

Store at 20-25°C (68-77°F) [See USP Controlled Room temperature].

Jubilant Cadista Pharmaceuticals Inc. Salisbury, MD 21801, USA

Repackaged By : Aidarex Pharmaceuticals LLC, Corona, CA 92880

# Lot No.:

Exp Date:

MECLIZINE HYDROCHLORIDE meclizine hydrocloride tablet									
<b>Product Information</b>									
Product Type		HUMAN PRESCRIPTI	ION DR	UG Item Code (S	ource)	NDC:3320	51-718(NDC:5	59746-121)	
Route of Administration		ORAL							
Active Ingredient/Act					-				
MECLIZINE HYDRO CHLO	-	redient Name				asis of Sta	-	Strength	
Inactive Ingredients									
inactive ingreatents		Ingredient N	ame				Str	ength	
SILICON DIOXIDE (UNII: E'	TJ7Z6 XBU	•							
CROSCARMELLOSE SOD		,							
LACTOSE MONOHYDRAT									
MAGNESIUM STEARATE (	UNII: 7009	7M6I30)							
CELLULOSE, MICROCRYS	STALLINE	C (UNII: OP1R32D61U)							
D&C YELLOW NO. 10 (UN	II: 35SW5U	SQ3G)							
ALUMINUM O XIDE (UNII: L	.MI260693	33)							
Product Characterist			-						
Color	YELLO	W	Score	2			2 pieces		
Shape	OVAL		Size				13mm		
Flavor			Impri	int Code			TL121	121	
Contains									
Packaging									
# Item Code	Pac	kage Description		Marketing Star	t Date	Ma	rketing En	d Date	
1 NDC:33261-718-08	8 in 1 BO	TTLE, PLASTIC					-		
<b>2</b> NDC:33261-718-10	10 in 1 BC	OTTLE, PLASTIC							
<b>3</b> NDC:33261-718-20	20 in 1 BC	OTTLE, PLASTIC							
4 NDC:33261-718-30	30 in 1 BC	OTTLE, PLASTIC							
5 NDC:33261-718-60	60 in 1 BC	DTTLE, PLASTIC							
Marketing Inform	nation								

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
ANDA	ANDA040659	06/04/2010	

# Labeler - Aidarex Pharmaceuticals LLC (801503249)

Revised: 1/2014

Aidarex Pharmaceuticals LLC