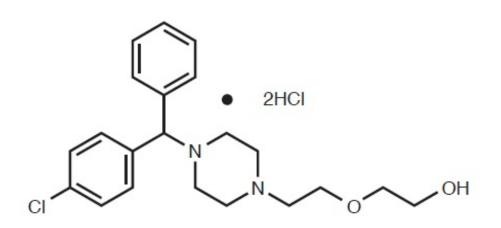
HYDROXYZINE HYDROCHLORIDE- hydroxyzine hydrochloride tablet Bryant Ranch Prepack

Rx only

DESCRIPTION

Hydroxyzine hydrochloride, USP has the chemical name of $(\pm)-2-[2-[4-(p-Chloro-\alpha-phenylbenzyl)-1-piperazinyl]ethoxy]ethanol dihydrochloride.$



Molecular Formula: C₂₁H₂₇ClN₂O₂ · 2HCl Molecular Weight: 447.83

Hydroxyzine hydrochloride, USP occurs as a white, odorless powder which is very soluble in water.

Each tablet for oral administration contains 10 mg, 25 mg, or 50 mg hydroxyzine hydrochloride, USP. Inactive ingredients include anhydrous lactose, colloidal silicon dioxide, crospovidone, hypromellose, magnesium stearate, microcrystalline cellulose, polyethylene glycol, polysorbate and titanium dioxide.

CLINICAL PHARMACOLOGY

Hydroxyzine hydrochloride is unrelated chemically to the phenothiazines, reserpine, meprobamate or the benzodiazepines. Hydroxyzine is not a cortical depressant, but its action may be due to a suppression of activity in certain key regions of the subcortical area of the central nervous system.

Primary skeletal muscle relaxation has been demonstrated experimentally. Bronchodilator activity, and antihistaminic and analgesic effects have been demonstrated experimentally and confirmed clinically. An antiemetic effect, both by the apomorphine test and the veriloid test, has been demonstrated.

Pharmacological and clinical studies indicate that hydroxyzine in therapeutic dosage does not increase gastric secretion or acidity and in most cases has mild antisecretory activity.

Hydroxyzine is rapidly absorbed from the gastrointestinal tract and hydroxyzine's clinical

effects are usually noted within 15 to 30 minutes after oral administration.

INDICATIONS AND USAGE

For symptomatic relief of anxiety and tension associated with psychoneurosis and as an adjunct in organic disease states in which anxiety is manifested.

Useful in the management of pruritus due to allergic conditions such as chronic urticaria and atopic and contact dermatoses and in histamine-mediated pruritus.

As a sedative when used as a premedication and following general anesthesia, **hydroxyzine may potentiate meperidine and barbiturates,** so their use in preanesthetic adjunctive therapy should be modified on an individual basis. Atropine and other belladonna alkaloids are not affected by the drug. Hydroxyzine is not known to interfere with the action of digitalis in any way and it may be used concurrently with this agent.

The effectiveness of hydroxyzine as an antianxiety agent for long term use, that is more than 4 months, has not been assessed by systematic clinical studies. The physician should reassess periodically the usefulness of the drug for the individual patient.

CONTRAINDICATIONS

Oral hydroxyzine hydrochloride is contraindicated in patients with known hypersensitivity to hydroxyzine hydrochloride products, and in patients with known hypersensitivity to cetirizine hydrochloride or levocetirizine hydrochloride.

Hydroxyzine is contraindicated in patients with a prolonged QT interval.

Hydroxyzine, when administered to the pregnant mouse, rat, and rabbit induced fetal abnormalities in the rat and mouse at doses substantially above the human therapeutic range. Clinical data in human beings are inadequate to establish safety in early pregnancy. Until such data are available, hydroxyzine is contraindicated in early pregnancy.

Hydroxyzine is contraindicated for patients who have shown a previous hypersensitivity to any component of this medication.

WARNINGS

Nursing Mothers

It is not known whether this drug is excreted in human milk. Since many drugs are so excreted, hydroxyzine should not be given to nursing mothers.

PRECAUTIONS

THE POTENTIATING ACTION OF HYDROXYZINE MUST BE CONSIDERED WHEN THE DRUG IS USED IN CONJUNCTION WITH CENTRAL NERVOUS SYSTEM DEPRESSANTS SUCH AS NARCOTICS, NON-NARCOTIC ANALGESICS AND BARBITURATES. Therefore, when central nervous system depressants are administered concomitantly with hydroxyzine their dosage should be reduced. QT Prolongation/Torsade de Pointes (TdP): Cases of QT prolongation and Torsade de Pointes have been reported during post-marketing use of hydroxyzine. The majority of reports occurred in patients with other risk factors for QT prolongation/TdP (preexisting heart disease, electrolyte imbalances or concomitant arrhythmogenic drug use). Therefore, hydroxyzine should be used with caution in patients with risk factors for QT prolongation, congenital long QT syndrome, a family history of long QT syndrome, other conditions that predispose to QT prolongation and ventricular arrhythmia, as well as recent myocardial infarction, uncompensated heart failure, and bradyarrhythmias.

Caution is recommended during the concomitant use of drugs known to prolong the QT interval. These include Class 1A (e.g., quinidine, procainamide) or Class III (e.g., amiodarone, sotalol) antiarrhythmics, certain antipsychotics (e.g., ziprasidone, iloperidone, clozapine, quetiapine, chlorpromazine), certain antidepressants (e.g., citalopram, fluoxetine), certain antibiotics (e.g., azithromycin, erythromycin, clarithromycin, gatifloxacin, moxifloxacin); and others (e.g., pentamidine, methadone, ondansetron, droperidol).

Since drowsiness may occur with use of this drug, patients should be warned of this possibility and cautioned against driving a car or operating dangerous machinery while taking hydroxyzine. Patients should also be advised against the simultaneous use of other CNS depressant drugs, and cautioned that the effects of alcohol may be increased.

Acute Generalized Exanthematous Pustulosis (AGEP)

Hydroxyzine may rarely cause acute generalized exanthematous pustulosis (AGEP), a serious skin reaction characterized by fever and numerous small, superficial, non-follicular, sterile pustules, arising within large areas of edematous erythema. Inform patients about the signs of AGEP, and discontinue hydroxyzine at the first appearance of a skin rash, worsening of pre-existing skin reactions which hydroxyzine may be used to treat, or any other sign of hypersensitivity. If signs or symptoms suggest AGEP, use of hydroxyzine should not be resumed and alternative therapy should be considered. Avoid cetirizine or levocetirizine in patients who have experienced AGEP or other hypersensitivity reactions with hydroxyzine, due to the risk of cross-sensitivity.

Geriatric Use

A determination has not been made whether controlled clinical studies of hydroxyzine included sufficient numbers of subjects aged 65 and over to define a difference in response from younger subjects. Other reported clinical experience has not identified differences in responses between the elderly and younger patients. In general, dose selection for an elderly patient should be cautious, usually starting at the low end of the dosing range, reflecting the greater frequency of decreased hepatic, renal or cardiac function, and of concomitant disease or other drug therapy.

The extent of renal excretion of hydroxyzine has not been determined. Because elderly patients are more likely to have decreased renal function, care should be taken in dose selections.

Sedating drugs may cause confusion and over sedation in the elderly; elderly patients generally should be started on low doses of hydroxyzine and observed closely.

ADVERSE REACTIONS

Skin and Appendages: Oral hydroxyzine hydrochloride is associated with <u>Acute</u> <u>Generalized Exanthematous Pustulosis (AGEP) and</u> fixed drug eruptions in post marketing reports.

Side effects reported with the administration of hydroxyzine hydrochloride are usually mild and transitory in nature.

Anticholinergic: Dry mouth.

Central Nervous System: Drowsiness is usually transitory and may disappear in a few days of continued therapy or upon reduction of dose. Involuntary motor activity including rare instances of tremor and convulsions have been reported, usually with doses considerably higher than those recommended. Clinically significant respiratory depression has not been reported at recommended doses.

In post-marketing experience, the following additional undesirable effects have been reported:

Cardiac System: QT prolongation, Torsade de Pointes.

Body as a Whole: Allergic reaction.

Nervous System: Headache.

Psychiatric: Hallucination.

Skin and Appendages: Pruritus, rash, urticaria.

OVERDOSAGE

The most common manifestation of hydroxyzine overdosage is hypersedation. Other reported signs and symptoms were convulsions, stupor, nausea and vomiting. As in the management of overdosage with any drug, it should be borne in mind that multiple agents may have been taken.

If vomiting has not occurred spontaneously, it should be induced. Immediate gastric lavage is also recommended. General supportive care, including frequent monitoring of the vital signs and close observation of the patient, is indicated. Hypotension, though unlikely, may be controlled with intravenous fluids and levarterenol or metaraminol. Do not use epinephrine as hydroxyzine counteracts its pressor action.

Hydroxyzine overdose may cause QT prolongation and Torsade de Pointes. ECG monitoring is recommended in cases of hydroxyzine overdose.

There is no specific antidote. It is doubtful that hemodialysis would be of any value in the treatment of overdosage with hydroxyzine. However, if other agents such as barbiturates have been ingested concomitantly, hemodialysis may be indicated. There is no practical method to quantitate hydroxyzine in body fluids or tissue after its ingestion or administration.

DOSAGE AND ADMINISTRATION

For symptomatic relief of anxiety and tension associated with psychoneurosis and as an

adjunct in organic disease states in which anxiety is manifested: adults, 50 to 100 mg q.i.d.; children under 6 years, 50 mg daily in divided doses; children over 6 years, 50 to 100 mg daily in divided doses.

For use in the management of pruritus due to allergic conditions such as chronic urticaria and atopic and contact dermatoses and in histamine-mediated pruritus: adults, 25 mg t.i.d. or q.i.d.; children under 6 years, 50 mg daily in divided doses; children over 6 years, 50 to 100 mg daily in divided doses.

As a sedative when used as a premedication and following general anesthesia: 50 to 100 mg for adults and 0.6 mg/kg of body weight in children.

When treatment is initiated by the intramuscular route of administration, subsequent doses may be administered orally.

As with all potent medication, the dosage should be adjusted according to the patient's response to therapy.

HOW SUPPLIED

Hydroxyzine Hydrochloride Tablets, USP 25 mg tablets: round, film coated white tablets. Debossed H over 501 on one side and plain on the reverse side. They are available as follows:

- NDC 71335-2340-1: 30 Tablets in a BOTTLE
- NDC 71335-2340-2: 60 Tablets in a BOTTLE
- NDC 71335-2340-3: 20 Tablets in a BOTTLE
- NDC 71335-2340-4: 100 Tablets in a BOTTLE
- NDC 71335-2340-5: 15 Tablets in a BOTTLE
- NDC 71335-2340-6: 90 Tablets in a BOTTLE
- NDC 71335-2340-7: 19 Tablets in a BOTTLE
- NDC 71335-2340-8: 10 Tablets in a BOTTLE

Dispense in a tight container as defined in the USP, with a child-resistant closure (as required).

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

KEEP THIS AND ALL MEDICATIONS OUT OF THE REACH OF CHILDREN.

Repackaged/Relabeled by: Bryant Ranch Prepack, Inc. Burbank, CA 91504

HydrOXYzine Hydrochloride 25mg Tablet



Each film-coated tablet contains: Hydroxyzine Hydrochloride, USP 25 mg.

Keep this and all medication out of the reach hydrOXYzine Hydrochloride of children.

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

Dispense in a tight container as defined in the USP, with a child-resistant closure (as required).



Insert

Product Characteristics

Usual Dosage: Scan Package Insert QR Code for full prescribing information.

NDC 71335-2340-1

Tablets, USP



PHARMACEUTICALS

30 Tablets Repackaged by: Bryant Ranch Prepack, Inc. Burbank, CA 91504 USA Manufactured by: Pharmaceuticals



Rx only

Emcure

Limited

hydroxyzine hydrochloride ta	ROCHLORIDE ablet				
Product Information					
Product Type	HUMAN PRESCRIPTION DRUG	ltem Code (Source)			
Route of Administration	ORAL				
Active Ingredient/Active	e Moiety				
Ingr	Ingredient Name Basis of Stree			is of Strength	Strength
HYDROXYZINE DIHYDROCHLORIDE (UNII: 76755771U3) (HYDROXYZINE - HYDROXYZINE UNII: 30S50YM80G) HYDROCHLORIDE			YZINE	25 mg	
UNII:30S50YM8OG)			DIHYDR	OCHLORIDE	25 mg
			DIHYDR	OCHLORIDE	23 mg
UNII:30550YM8OG)			DIHYDR	OCHLORIDE	25 mg
	Ingredient Name		DIHYDR		Strength
	•		DIHYDR		
Inactive Ingredients	SY5LH9PMK)		DIHYDR		
Inactive Ingredients ANHYDROUS LACTOSE (UNII: 35	SY5LH9PMK) (BU4)		DIHYDR		
Inactive Ingredients ANHYDROUS LACTOSE (UNII: 33 SILICON DIOXIDE (UNII: ETJ7Z6)	SY5LH9PMK) (BU4) (UNII: 3NXW29V3WO)		DIHYDR		
Inactive Ingredients ANHYDROUS LACTOSE (UNII: 33 SILICON DIOXIDE (UNII: ETJ7Z6) HYPROMELLOSE, UNSPECIFIE	SY5LH9PMK) (BU4) • (UNII: 3NXW29V3WO) 0097M6I30)		DIHYDR		
Inactive Ingredients ANHYDROUS LACTOSE (UNII: 33 SILICON DIOXIDE (UNII: ETJ7Z6) HYPROMELLOSE, UNSPECIFIEI MAGNESIUM STEARATE (UNII: 7	SY5LH9PMK) (BU4) (UNII: 3NXW29V3WO) 0097M6I30) SE (UNII: OP1R32D61U)	Α)	DIHYDR		
Inactive Ingredients ANHYDROUS LACTOSE (UNII: 33 SILICON DIOXIDE (UNII: ETJ7Z6) HYPROMELLOSE, UNSPECIFIE MAGNESIUM STEARATE (UNII: 7 MICROCRYSTALLINE CELLULO	SY5LH9PMK) (BU4) (UNII: 3NXW29V3WO) 0097M6I30) SE (UNII: OP1R32D61U) PECIFIED (UNII: 3WJQ0SDW14	A)	DIHYDR		
Inactive Ingredients ANHYDROUS LACTOSE (UNII: 33 SILICON DIOXIDE (UNII: ETJ7Z6) HYPROMELLOSE, UNSPECIFIE MAGNESIUM STEARATE (UNII: 7 MICROCRYSTALLINE CELLULO POLYETHYLENE GLYCOL, UNS	GY5LH9PMK) (BU4) (UNII: 3NXW29V3WO) 0097M6I30) SE (UNII: OP1R32D61U) PECIFIED (UNII: 3WJQ0SDW1A 9ZG8H)	A)	DIHYDR		

Color	WHITE	Score	no score	
Shape	ROUND	Size	7mm	
Flavor		Imprint Code	H;501	
Contains				

Packaging					
#		Packag	e Description	Marketing Start Date	Marketing End Date
1	NDC:71335- 2340-1	30 in 1 BOTTLE; Typ Product	e 0: Not a Combination	06/27/2025	
2	NDC:71335- 2340-2	60 in 1 BOTTLE; Typ Product	e 0: Not a Combination	06/27/2025	
3	NDC:71335- 2340-3	20 in 1 BOTTLE; Typ Product	e 0: Not a Combination	06/27/2025	
4	NDC:71335- 2340-4	100 in 1 BOTTLE; Ty Product	pe 0: Not a Combination	06/27/2025	
5	NDC:71335- 2340-5	15 in 1 BOTTLE; Typ Product	e 0: Not a Combination	06/27/2025	
6	NDC:71335- 2340-6	90 in 1 BOTTLE; Typ Product	e 0: Not a Combination	06/27/2025	
7	NDC:71335- 2340-7	19 in 1 BOTTLE; Typ Product	e 0: Not a Combination	06/27/2025	
8	NDC:71335- 2340-8	10 in 1 BOTTLE; Typ Product	e 0: Not a Combination	06/27/2025	
Marketing Information					
	Marketing Category	Application I	Number or Monograph Citation	Marketing Start Date	Marketing End Date
٨N	IDA	ANDA204279		08/20/2014	

Labeler - Bryant Ranch Prepack (171714327)

Registrant - Bryant Ranch Prepack (171714327)

Establishment					
Name	Address	ID/FEI	Business Operations		
Bryant Ranch Prepack		171714327	REPACK(71335-2340), RELABEL(71335-2340)		

Revised: 6/2025

Bryant Ranch Prepack