CYCLOBENZAPRINE HYDROCHLORIDE- cyclobenzaprine hydrochloride tablet RedPharm Drug, Inc.

DESCRIPTION

Cyclobenzaprine hydrochloride is a white, crystalline tricyclic amine salt. It has a melting point of 217°C, and a pKa of 8.47 at 25°C. It is freely soluble in water and alcohol, sparingly soluble in isopropanol, and insoluble in hydrocarbon solvents. If aqueous solutions are made alkaline, the free base separates. Cyclobenzaprine HCl is designated chemically as 3-(5H- dibenzo[a,d]cyclohepten-5-ylidene)- N, N -dimethyl-1- propanamine hydrochloride, and has the following structural formula:

[cyclobenzaprine hydrochloride structural formula]

C20H21N•HCl0000000000M.W. 311.9

Cyclobenzaprine hydrochloride tablets, USP are available for oral administration as 5 mg, 7.5 mg and 10 mg tablets. Cyclobenzaprine hydrochloride 5 mg, 7.5 mg and 10 mg tablets contain the following inactive ingredients: colloidal silicon dioxide, croscarmellose sodium, dibasic calcium phosphate, hydroxypropyl cellulose, hypromellose, polyethylene glycol, magnesium stearate, microcrystalline cellulose, and titanium dioxide.

CLINICAL PHARMACOLOGY

Cyclobenzaprine HCl relieves skeletal muscle spasm of local origin without interfering with muscle function. It is ineffective in muscle spasm due to central nervous system disease.

Cyclobenzaprine reduced or abolished skeletal muscle hyperactivity in several animal models. Animal studies indicate that cyclobenzaprine does not act at the neuromuscular junction or directly on skeletal muscle. Such studies show that cyclobenzaprine acts primarily within the central nervous system at brain stem as opposed to spinal cord levels, although its action on the latter may contribute to its overall skeletal muscle relaxant activity. Evidence suggests that the net effect of cyclobenzaprine is a reduction of tonic somatic motor activity, influencing both gamma (γ) and alpha (α) motor systems.

Pharmacological studies in animals showed a similarity between the effects of cyclobenzaprine and the structurally related tricyclic antidepressants, including reserpine antagonism, norepinephrine potentiation, potent peripheral and central anticholinergic effects, and sedation. Cyclobenzaprine caused slight to moderate increase in heart rate in animals.

Pharmacokinetics

Estimates of mean oral bioavailability of cyclobenzaprine range from 33% to 55%. Cyclobenzaprine exhibits linear pharmacokinetics over the dose range 2.5 mg to 10 mg, and is subject to enterohepatic circulation. It is highly bound to plasma proteins. Drug accumulates when dosed three times a day, reaching steady-state within 3-4 days at plasma concentrations about four-fold higher than after a single dose. At steady state in healthy subjects receiving 10 mg t.i.d. (n=18), peak plasma concentration was 25.9 ng/mL (range, 12.8-46.1 ng/mL), and area under the concentration-time (AUC) curve over an 8-hour dosing interval was 177 ng.hr/mL (range, 80-319 ng.hr/mL).

Cyclobenzaprine is extensively metabolized, and is excreted primarily as glucuronides via the kidney. Cytochromes P-450 3A4, 1A2, and, to a lesser extent, 2D6, mediate N-demethylation, one of the oxidative pathways for cyclobenzaprine. Cyclobenzaprine is eliminated quite slowly, with an effective half-life of 18 hours (range 8-37 hours; n=18); plasma clearance is 0.7 L/min.

The plasma concentration of cyclobenzaprine is generally higher in the elderly and in patients with hepatic impairment. (See PRECAUTIONS, USE IN THE ELDERLY and PRECAUTIONS, IMPAIRED HEPATIC FUNCTION.)

Elderly

In a pharmacokinetic study in elderly individuals (\geq 65yrs old), mean (n=10) steady-state cyclobenzaprine AUC values were approximately 1.7 fold (171.0 ng.hr/mL, range 96.1 to 255.3) higher than those seen in a group of eighteen younger adults (101.4 ng.hr/mL, range 36.1 to 182.9) from another study. Elderly male subjects had the highest observed mean increase, approximately 2.4 fold (198.3 ng.hr/mL, range 155.6 to 255.3 versus 83.2 ng.hr/mL, range 41.1 to 142.5 for younger males) while levels in elderly females were increased to a much lesser extent, approximately 1.2 fold (143.8 ng.hr/mL, range 96.1 to 196.3 versus 115.9 ng.hr/mL, range 36.1 to 182.9 for younger females).

In light of these findings, therapy with cyclobenzaprine HCl in the elderly should be initiated with a 5 mg dose and titrated slowly upward.

Hepatic Impairment

In a pharmacokinetic study of sixteen subjects with hepatic impairment (15 mild, 1 moderate per Child-Pugh score), both AUC and Cmax were approximately double the values seen in the healthy control group. Based on the findings, cyclobenzaprine HCl should be used with caution in subjects with mild hepatic impairment starting with the 5 mg dose and titrating slowly upward. Due to the lack of data in subjects with more severe hepatic insufficiency, the use of cyclobenzaprine HCl in subjects with moderate to severe impairment is not recommended.

No significant effect on plasma levels or bioavailability of cyclobenzaprine HCl or aspirin was noted when single or multiple doses of the two drugs were administered concomitantly. Concomitant administration of cyclobenzaprine HCl and naproxen or diflunisal was well tolerated with no reported unexpected adverse effects. However combination therapy of cyclobenzaprine HCl with naproxen was associated with more side effects than therapy with naproxen alone, primarily in the form of drowsiness. No well-controlled studies have been performed to indicate that cyclobenzaprine HCl enhances the clinical effect of aspirin or other analgesics, or whether analgesics enhance the clinical effect of cyclobenzaprine HCl in acute musculoskeletal conditions.

Clinical Studies

Eight double-blind controlled clinical studies were performed in 642 patients comparing cyclobenzaprine HCl 10 mg, diazepam**, and placebo. Muscle spasm, local pain and tenderness, limitation of motion, and restriction in activities of daily living were evaluated. In three of these studies there was a significantly greater improvement with cyclobenzaprine HCl than with diazepam, while in the other studies the improvement following both treatments was comparable.

Although the frequency and severity of adverse reactions observed in patients treated with cyclobenzaprine HCl were comparable to those observed in patients treated with diazepam, dry mouth was observed more frequently in patients treated with cyclobenzaprine HCl and dizziness more frequently in those treated with diazepam. The incidence of drowsiness, the most frequent adverse reaction, was similar with both drugs.

The efficacy of cyclobenzaprine HCl 5 mg was demonstrated in two seven-day, double-blind, controlled clinical trials enrolling 1405 patients. One study compared cyclobenzaprine HCl 5 and 10 mg t.i.d. to placebo; and a second study compared cyclobenzaprine HCl 5 and 2.5 mg t.i.d. to placebo. Primary endpoints for both trials were determined by patient-generated data and included global impression of change, medication helpfulness, and relief from starting backache. Each endpoint consisted of a score on a 5-point rating scale (from 0 or worst outcome to 4 or best outcome). Secondary endpoints included a physician's evaluation of the presence and extent of palpable muscle spasm.

Comparisons of cyclobenzaprine HCl 5 mg and placebo groups in both trials established the statistically significant superiority of the 5 mg dose for all three primary endpoints at day 8 and, in the study comparing 5 and 10 mg, at day 3 or 4 as well. A similar effect was observed with cyclobenzaprine HCl 10 mg (all endpoints). Physician-assessed secondary endpoints also showed that cyclobenzaprine HCl 5

mg was associated with a greater reduction in palpable muscle spasm than placebo.

Analysis of the data from controlled studies shows that cyclobenzaprine HCl produces clinical improvement whether or not sedation occurs.

**VALIUM® (diazepam, Roche)

Surveillance Program

A postmarketing surveillance program was carried out in 7607 patients with acute musculoskeletal disorders, and included 297 patients treated with cyclobenzaprine HCl 10 mg for 30 days or longer. The overall effectiveness of cyclobenzaprine HCl was similar to that observed in the double-blind controlled studies; the overall incidence of adverse effects was less (see ADVERSE REACTIONS).

INDICATIONS AND USAGE

Cyclobenzaprine HCl is indicated as an adjunct to rest and physical therapy for relief of muscle spasm associated with acute, painful musculoskeletal conditions.

Improvement is manifested by relief of muscle spasm and its associated signs and symptoms, namely, pain, tenderness, limitation of motion, and restriction in activities of daily living.

Cyclobenzaprine HCl should be used only for short periods (up to two or three weeks) because adequate evidence of effectiveness for more prolonged use is not available and because muscle spasm associated with acute, painful musculoskeletal conditions is generally of short duration and specific therapy for longer periods is seldom warranted.

Cyclobenzaprine HCl has not been found effective in the treatment of spasticity associated with cerebral or spinal cord disease, or in children with cerebral palsy.

CONTRAINDICATIONS

Hypersensitivity to any component of this product.

Concomitant use of monoamine oxidase (MAO) inhibitors or within 14 days after their discontinuation. Hyperpyretic crisis seizures, and deaths have occurred in patients receiving cyclobenzaprine (or structurally similar tricyclic antidepressants) concomitantly with MAO inhibitor drugs.

Acute recovery phase of myocardial infarction, and patients with arrhythmias, heart block or conduction disturbances, or congestive heart failure.

Hyperthyroidism.

WARNINGS

Serotonin Syndrome

The development of a potentially life-threatening serotonin syndrome has been reported with Cyclobenzaprine Hydrochloride when used in combination with other drugs, such as selective serotonin reuptake inhibitors (SSRIs), serotonin norepinephrine reuptake inhibitors (SNRIs), tricyclic antidepressants (TCAs), tramadol, bupropion, meperidine, verapamil, or MAO inhibitors. The concomitant use of Cyclobenzaprine Hydrochloride with MAO inhibitors is contraindicated (seeCONTRAINDICATIONS). Serotonin syndrome symptoms may include mental status changes (e.g., confusion, agitation, hallucinations), autonomic instability (e.g., diaphoresis, tachycardia, labile blood pressure, hyperthermia), neuromuscular abnormalities (e.g., tremor, ataxia, hyperreflexia, clonus, muscle rigidity), and/or gastrointestinal symptoms (e.g., nausea, vomiting, diarrhea). Treatment with Cyclobenzaprine Hydrochloride and any concomitant serotonergic agents should be discontinued

immediately if the above reactions occur and supportive symptomatic treatment should be initiated. If concomitant treatment with Cyclobenzaprine Hydrochloride and other serotonergic drugs is clinically warranted, careful observation is advised, particularly during treatment initiation or dose increases (seePRECAUTIONS, DRUG INTERACTIONS).

Cyclobenzaprine is closely related to the tricyclic antidepressants, e.g., amitriptyline and imipramine. In short term studies for indications other than muscle spasm associated with acute musculoskeletal conditions, and usually at doses somewhat greater than those recommended for skeletal muscle spasm, some of the more serious central nervous system reactions noted with the tricyclic antidepressants have occurred (see WARNINGS, below, and ADVERSE REACTIONS).

Tricyclic antidepressants have been reported to produce arrhythmias, sinus tachycardia, prolongation of the conduction time leading to myocardial infarction and stroke.

Cyclobenzaprine HCl may enhance the effects of alcohol, barbiturates, and other CNS depressants.

PRECAUTIONS

General

Because of its atropine-like action, cyclobenzaprine HCl should be used with caution in patients with a history of urinary retention, angle-closure glaucoma, increased intraocular pressure, and in patients taking anticholinergic medication.

Impaired Hepatic Function

The plasma concentration of cyclobenzaprine is increased in patients with hepatic impairment (seeCLINICAL PHARMACOLOGY, PHARMACOKINETICS, HEPATIC IMPAIRMENT).

These patients are generally more susceptible to drugs with potentially sedating effects, including cyclobenzaprine. Cyclobenzaprine HCl should be used with caution in subjects with mild hepatic impairment starting with a 5 mg dose and titrating slowly upward. Due to the lack of data in subjects with more severe hepatic insufficiency, the use of cyclobenzaprine HCl in subjects with moderate to severe impairment is not recommended.

Information for Patients

Cyclobenzaprine HCl, especially when used with alcohol or other CNS depressants, may impair mental and/or physical abilities required for performance of hazardous tasks, such as operating machinery or driving a motor vehicle. In the elderly, the frequency and severity of adverse events associated with the use of cyclobenzaprine, with or without concomitant medications, is increased. In elderly patients, cyclobenzaprine HCl should be initiated with a 5 mg dose and titrated slowly upward.

Patients should be cautioned about the risk of serotonin syndrome with the concomitant use of Cyclobenzaprine Hydrochloride and other drugs, such as SSRIs, SNRIs, TCAs, tramadol, bupropion, meperidine, verapamil, or MAO inhibitors. Patients should be advised of the signs and symptoms of serotonin syndrome, and be instructed to seek medical care immediately if they experience these symptoms (see WARNINGS, and see PRECAUTIONS, DRUG INTERACTIONS).

Drug Interactions

Cyclobenzaprine HCl may have life-threatening interactions with MAO inhibitors (seeCONTRAINDICATIONS). Postmarketing cases of serotonin syndrome have been reported during combined use of Cyclobenzaprine Hydrochloride and other drugs, such as SSRIs, SNRIs, TCAs, tramadol, bupropion, meperidine, verapamil, or MAO inhibitors. If concomitant treatment with Cyclobenzaprine Hydrochloride and other serotonergic drugs is clinically warranted, careful observation is advised, particularly during treatment initiation or dose increases (see WARNINGS).

Cyclobenzaprine HCl may enhance the effects of alcohol, barbiturates, and other CNS depressants.

Tricyclic antidepressants may block the antihypertensive action of guanethidine and similarly acting

compounds.

Tricyclic antidepressants may enhance the seizure risk in patients taking tramadol.⁺

† ULTRAM® (tramadol HCl tablets, PriCar, Division of Ortho-McNeil-Janssen Pharmaceuticals, Inc.)

[†]ULTRACET[®] (tramadol HCl and acetaminophen tablets, PriCara, Division of Ortho-McNeil-Janssen Pharmaceuticals, Inc.)

Carcinogenesis, Mutagenesis, Impairment of Fertility

In rats treated with cyclobenzaprine HCl for up to 67 weeks at doses of approximately 5 to 40 times the maximum recommended human dose, pale, sometimes enlarged, livers were noted and there was a dose-related hepatocyte vacuolation with lipidosis. In the higher dose groups this microscopic change was seen after 26 weeks and even earlier in rats which died prior to 26 weeks; at lower doses, the change was not seen until after 26 weeks.

Cyclobenzaprine did not affect the onset, incidence or distribution of neoplasia in an 81-week study in the mouse or in a 105-week study in the rat.

At oral doses of up to 10 times the human dose, cyclobenzaprine did not adversely affect the reproductive performance or fertility of male or female rats. Cyclobenzaprine did not demonstrate mutagenic activity in the male mouse at dose levels of up to 20 times the human dose.

Pregnancy

Pregnancy Category B

Reproduction studies have been performed in rats, mice and rabbits at doses up to 20 times the human dose, and have revealed no evidence of impaired fertility or harm to the fetus due to cyclobenzaprine HCl. There are, however, no adequate and well-controlled studies in pregnant women. Because animal reproduction studies are not always predictive of human response, this drug should be used during pregnancy only if clearly needed.

Nursing Mothers

It is not known whether this drug is excreted in human milk. Because cyclobenzaprine is closely related to the tricyclic antidepressants, some of which are known to be excreted in human milk, caution should be exercised when cyclobenzaprine HCl is administered to a nursing woman.

Pediatric Use

Safety and effectiveness of cyclobenzaprine HCl in pediatric patients below 15 years of age have not been established.

Use in the Elderly

The plasma concentration of cyclobenzaprine is increased in the elderly (see CLINICAL PHARMACOLOGY, PHARMACOKINETICS, ELDERLY). The elderly may also be more at risk for CNS adverse events such as hallucinations and confusion, cardiac events resulting in falls or other sequelae, drug-drug and drug-disease interactions. For these reasons, in the elderly, cyclobenzaprine should be used only if clearly needed. In such patients cyclobenzaprine HCl should be initiated with a 5 mg dose and titrated slowly upward.

ADVERSE REACTIONS

Incidence of most common adverse reactions in the 2 double-blind \ddagger , placebo-controlled 5 mg studies (incidence of > 3% on cyclobenzaprine HCl 5 mg):

Cyclobenzaprine HCl Tablets Cyclobenzaprine HCl Tablets Placebo

5 mg 10 mg N=464 N=249 N=469 Drowsiness 29% 38% 10% Dry Mouth 21% 32% 7% Fatigue 6% 6% 3% Headache 5% 5% 8% ‡Note: Cyclobenzaprine HCl Tablets 10 mg data are from one clinical trial. Cyclobenzaprine HCl Tablets 5 mg and placebo data are from two studies.

Adverse reactions which were reported in 1% to 3% of the patients were: abdominal pain, acid regurgitation, constipation, diarrhea, dizziness, nausea, irritability, mental acuity decreased, nervousness, upper respiratory infection, and pharyngitis.

The following list of adverse reactions is based on the experience in 473 patients treated with cyclobenzaprine HCl 10 mg in additional controlled clinical studies, 7607 patients in the postmarketing surveillance program, and reports received since the drug was marketed. The overall incidence of adverse reactions among patients in the surveillance program was less than the incidence in the controlled clinical studies.

The adverse reactions reported most frequently with cyclobenzaprine HCl were drowsiness, dry mouth and dizziness. The incidence of these common adverse reactions was lower in the surveillance program than in the controlled clinical studies:

Clinical Studies with Surveillance Program with Cyclobenzaprine HCl Tablets 10 mg Cyclobenzaprine HCl Tablets 10 mg Drowsiness 39% 16% Dry mouth 27% 7% Dizziness 11% 3%

Among the less frequent adverse reactions, there was no appreciable difference in incidence in controlled clinical studies or in the surveillance program. Adverse reactions which were reported in 1% to 3% of the patients were: fatigue/tiredness, asthenia, nausea, constipation, dyspepsia, unpleasant taste, blurred vision, headache, nervousness, and confusion.

The following adverse reactions have been reported in postmarketing experience or with an incidence of less than 1% of patients in clinical trials with the 10 mg tablet:

Body as a Whole: Syncope; malaise.

Cardiovascular: Tachycardia; arrhythmia; vasodilatation; palpitation; hypotension.

Digestive: Vomiting; anorexia; diarrhea; gastrointestinal pain; gastritis; thirst; flatulence; edema of the tongue; abnormal liver function and rare reports of hepatitis, jaundice and cholestasis.

Hypersensitivity: Anaphylaxis; angioedema; pruritus; facial edema; urticaria; rash.

Musculoskeletal: Local weakness.

Nervous System and Psychiatric: Seizures, ataxia; vertigo; dysarthria; tremors; hypertonia; convulsions; muscle twitching; disorientation; insomnia; depressed mood; abnormal sensations; anxiety; agitation; psychosis, abnormal thinking and dreaming; hallucinations; excitement; paresthesia; diplopia, serotonin syndrome.

Skin: Sweating.

Special Senses: Ageusia; tinnitus.

Urogenital: Urinary frequency and/or retention.

Causal Relationship Unknown

Other reactions, reported rarely for cyclobenzaprine HCl under circumstances where a causal

relationship could not be established or reported for other tricyclic drugs, are listed to serve as alerting information to physicians:

Body as a Whole: Chest pain; edema.

Cardiovascular: Hypertension; myocardial infarction; heart block; stroke.

Digestive: Paralytic ileus, tongue discoloration; stomatitis; parotid swelling.

Endocrine: Inappropriate ADH syndrome.

Hematic and Lymphatic: Purpura; bone marrow depression; leukopenia; eosinophilia; thrombocytopenia.

Metabolic, Nutritional and Immune: Elevation and lowering of blood sugar levels; weight gain or loss.

Musculoskeletal: Myalgia.

Nervous System and Psychiatric: Decreased or increased libido; abnormal gait; delusions; aggressive behavior; paranoia; peripheral neuropathy; Bell's palsy; alteration in EEG patterns; extrapyramidal symptoms.

Respiratory: Dyspnea.

Skin: Photosensitization; alopecia.

Urogenital: Impaired urination; dilatation of urinary tract; impotence; testicular swelling; gynecomastia; breast enlargement; galactorrhea.

To report SUSPECTED ADVERSE REACTIONS, contact Actavis at 1-800-272-5525 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

DRUG ABUSE AND DEPENDENCE

Pharmacologic similarities among the tricyclic drugs require that certain withdrawal symptoms be considered when cyclobenzaprine HCl is administered, even though they have not been reported to occur with this drug. Abrupt cessation of treatment after prolonged administration rarely may produce nausea, headache, and malaise. These are not indicative of addiction.

OVERDOSAGE

lthough rare, deaths may occur from overdosage with cyclobenzaprine HCl. Multiple drug ingestion (including alcohol) is common in deliberate cyclobenzaprine overdose. As management of overdose is complex and changing, it is recommended that the physician contact a poison control center for current information on treatment. Signs and symptoms of toxicity may develop rapidly after cyclobenzaprine overdose; therefore, hospital monitoring is required as soon as possible. The acute oral LD50 of cyclobenzaprine HCl is approximately 338 and 425 mg/kg in mice and rats, respectively.

Manifestations

The most common effects associated with cyclobenzaprine overdose are drowsiness and tachycardia. Less frequent manifestations include tremor, agitation, coma, ataxia, hypertension, slurred speech, confusion, dizziness, nausea, vomiting, and hallucinations. Rare but potentially critical manifestations of overdose are cardiac arrest, chest pain, cardiac dysrhythmias, severe hypotension, seizures, and neuroleptic malignant syndrome. Changes in the electrocardiogram, particularly in QRS axis or width, are clinically significant indicators of cyclobenzaprine toxicity.

Other potential effects of overdosage include any of the symptoms listed under ADVERSE REACTIONS.

Management General As management of overdose is complex and changing, it is recommended that the physician contact a poison control center for current information on treatment.

In order to protect against the rare but potentially critical manifestations described above, obtain an ECG and immediately initiate cardiac monitoring. Protect the patient's airway, establish an intravenous line and initiate gastric decontamination. Observation with cardiac monitoring and observation for signs of CNS or respiratory depression, hypotension, cardiac dysrhythmias and/or conduction blocks, and seizures is necessary. If signs of toxicity occur at any time during this period, extended monitoring is required. Monitoring of plasma drug levels should not guide management of the patient. Dialysis is probably of no value because of low plasma concentrations of the drug.

Gastrointestinal Decontamination

All patients suspected of an overdose with cyclobenzaprine HCl should receive gastrointestinal decontamination. This should include large volume gastric lavage followed by activated charcoal. If consciousness is impaired, the airway should be secured prior to lavage and emesis is contraindicated.

Cardiovascular

A maximal limb-lead QRS duration of ≥ 0.10 seconds may be the best indication of the severity of the overdose. Serum alkalinization, to a pH of 7.45 to 7.55, using intravenous sodium bicarbonate and hyperventilation (as needed), should be instituted for patients with dysrhythmias and/or QRS widening. A pH > 7.60 or a pCO2< 20 mmHg is undesirable. Dysrhythmias unresponsive to sodium bicarbonate therapy/hyperventilation may respond to lidocaine, bretylium or phenytoin. Type 1A and 1C antiarrhythmics are generally contraindicated (e.g., quinidine, disopyramide, and procainamide).

CNS

In patients with CNS depression, early intubation is advised because of the potential for abrupt deterioration. Seizures should be controlled with benzodiazepines or, if these are ineffective, other anticonvulsants (e.g. phenobarbital, phenytoin). Physostigmine is not recommended except to treat life-threatening symptoms that have been unresponsive to other therapies, and then only in close consultation with a poison control center.

Psychiatric Follow-Up

Since overdosage is often deliberate, patients may attempt suicide by other means during the recovery phase. Psychiatric referral may be appropriate.

Pediatric Management

The principles of management of child and adult overdosages are similar. It is strongly recommended that the physician contact the local poison control center for specific pediatric treatment.

DOSAGE & ADMINISTRATIONS

For most patients, the recommended dose of cyclobenzaprine HCl is 5 mg three times a day. Based on individual patient response, the dose may be increased to either 7.5 mg or 10 mg three times a day. Use of cyclobenzaprine HCl for periods longer than two or three weeks is not recommended. (SeeINDICATIONS AND USAGE.)

Less frequent dosing should be considered for hepatically impaired or elderly patients (seePRECAUTIONS, IMPAIRED HEPATIC FUNCTION, and USE IN THE ELDERLY). CLOSE

HOW SUPPLIED

Cyclobenzaprine Hydrochloride Tablets USP, 5 mg are round, white, film-coated tablets imprintedWATSON and 3256 supplied in bottles of 30, 100 and 1000.

Cyclobenzaprine Hydrochloride Tablets, USP 7.5 mg are round, white, film-coated tablets imprintedWATSON and 3330 supplied in bottles of 100.

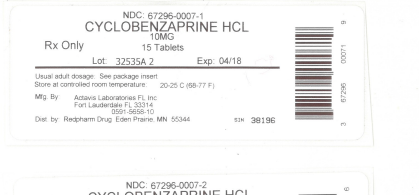
Cyclobenzaprine Hydrochloride Tablets USP, 10 mg are round, white, film-coated tablets imprinted DANand 5658 supplied in bottles of 100, 500 and 1000.

Dispense in a well-closed container with child-resistant closure. Store at 20° to 25°C (68° to 77°F) [See USP Controlled Room Temperature].

Manufactured by: Actavis Laboratories FL, Inc. Fort Lauderdale, FL 33314 USA

Distributed by: Actavis Pharma, Inc. Parsippany, NJ 07054 USA

Revised: August 2016







Product Informat	tion							
Product Type		UMAN PRESCRIPTION RUG	Item Code (So	Code (Source)		NDC:67296-0007(NDC:0		
Route of Administration ORAL								
Active Ingredient	t/Active Moiety	ý						
						sis of Strength Strengt		
CYCLOBENZAPRINE HYDROCHLORIDE (UNII: 0 VE05JYS2P) (CYCLOBENZAPRINE - UNII:6905WQQ5TI) CYCLOBENZAPRINE HYDROCHLORIDE						-	10 mg	
Inactive Ingredie	nts							
		Ingredient Name				1	Strength	
MAGNESIUM STEARA								
CELLULOSE, MICRO								
		E H) (UNII: RFW2ET671P)						
CALCIUM PHO SPHAT	E, DIBASIC, ANHY	YDROUS (UNII: L11K75P92J)						
HYPROMELLOSES (U	JNII: 3NXW29V3WC))						
	UNII: 15FIX9V2JP)							
TTTANIUM DIO XIDE (01111 101 110 (201)							
TITANIUM DIO XIDE (SILICON DIO XIDE (U								
SILICON DIO XIDE (U CROSCARMELLOSE	NII: ETJ7Z6XBU4) SODIUM (UNII: M2							
SILICON DIO XIDE (U	NII: ETJ7Z6XBU4) SODIUM (UNII: M2							
SILICON DIO XIDE (U CROSCARMELLOSE	NII: ETJ7Z6XBU4) SODIUM (UNII: M2							
SILICON DIOXIDE (U CROSCARMELLOSE POLYETHYLENE GLY	NII: ETJ7Z6XBU4) SODIUM (UNII: M2 YCOLS (UNII: 3WJ0							
SILICON DIO XIDE (U CROSCARMELLOSE POLYETHYLENE GLY Product Characte	NII: ETJ7Z6XBU4) SODIUM (UNII: M2 YCOLS (UNII: 3WJ0	Q0SDW1A)	score with	1 unever	1 pieces			
SILICON DIOXIDE (U CROSCARMELLOSE POLYETHYLENE GLY Product Characte Color	NII: ETJ7Z6XBU4) SODIUM (UNII: M2 YCOLS (UNII: 3WJ0 Pristics white	QOSDW1A) Score	score with 8mm	1 unever	1 pieces			
SILICON DIO XIDE (U CROSCARMELLOSE POLYETHYLENE GLY Product Characte Color Shape	NII: ETJ7Z6XBU4) SODIUM (UNII: M2 YCOLS (UNII: 3WJ0 Pristics	QOSDW1A) Score Size	8 mm		ı pieces			
SILICON DIO XIDE (U CROSCARMELLOSE POLYETHYLENE GLY Product Characte Color Shape Flavor	NII: ETJ7Z6XBU4) SODIUM (UNII: M2 YCOLS (UNII: 3WJ0 Pristics white	QOSDW1A) Score			1 pieces			
SILICON DIO XIDE (U CROSCARMELLOSE POLYETHYLENE GLY Product Characte Color Shape Flavor	NII: ETJ7Z6XBU4) SODIUM (UNII: M2 YCOLS (UNII: 3WJ0 Pristics white	QOSDW1A) Score Size	8 mm		1 pieces			
SILICON DIOXIDE (U CROSCARMELLOSE POLYETHYLENE GLY Product Characte Color Shape Flavor Contains	NII: ETJ7Z6XBU4) SODIUM (UNII: M2 YCOLS (UNII: 3WJ0 Pristics white	QOSDW1A) Score Size	8 mm		n pieces			
SILICON DIO XIDE (U CROSCARMELLOSE POLYETHYLENE GLY Product Characte Color Shap e Flavor Contains Packaging	NII: ETJ7Z6 XBU4) SODIUM (UNII: M2 YCOLS (UNII: 3WJ0 Pristics white ROUND	QOSDW1A) Score Size	8 mm DAN;56 58	3		Marketin	ıg End Date	
SILICON DIO XIDE (U CROSCARMELLOSE POLYETHYLENE GLY Product Characte Color Shape Flavor Contains Packaging # Item Code	NII: ETJ7Z6 XBU4) SODIUM (UNII: M2 YCOLS (UNII: 3WJ0 Pristics white ROUND	QOSDW1A) Score Size Imprint Code	8 mm DAN;5658 Marke	3 ting St		Marketin	ng End Date	
SILICON DIO XIDE (U CROSCARMELLOSE POLYETHYLENE GLY Polyethylene GLY Color Shape Flavor Contains Packaging # Item Code 1 NDC:67296-0007-1	NII: ETJ7Z6 XBU4) SODIUM (UNII: M2 YCOLS (UNII: 3WJ0 Pristics white ROUND ROUND Pa 15 in 1 BOTTLE; 7	QOSDW1A) Score Size Imprint Code	8 mm DAN;5658 Marke duct 09/28/20	3 ting St 0 16		Marketin	ıg End Date	
SILICON DIO XIDE (U) CROSCARMELLOSE POLYETHYLENE GLY Polyethylene GLY Color Shape Flavor Contains Packaging I tem Code NDC:67296-0007-1 2 NDC:67296-0007-2	NII: ETJ7Z6 XBU4) SODIUM (UNII: M2 YCOLS (UNII: 3WJ0 eristics white ROUND 15 in 1 BOTTLE; 7 20 in 1 BOTTLE; 7	QOSDW1A) Score Size Imprint Code Ckage Description Gype 0: Not a Combination Processor	8 mm DAN;5658 Marke duct 09/28/20	3 ting S 1 0 16 0 16		Marke tin	ıg End Date	
SILICON DIO XIDE (U) CROSCARMELLOSE POLYETHYLENE GLY Polyethylene GLY Cortains Contains Packaging # Item Code 1 NDC:67296-0007-1 2 NDC:67296-0007-3	NII: ETJ7Z6 XBU4) SODIUM (UNII: M2 COLS (UNII: 3WJ0 ristics white ROUND 15 in 1 BOTTLE; 7 20 in 1 BOTTLE; 7 30 in 1 BOTTLE; 7	Score Size Imprint Code Ckage Description Type 0: Not a Combination Proc	8 mm DAN;5658 Marke duct 09/28/20	3 ting S 1 0 16 0 16		Marke tin	ng End Date	
SILICON DIO XIDE (U CROSCARMELLOSE POLYETHYLENE GLY Product Characte Color Shape Flavor Contains Packaging	NII: ETJ7Z6 XBU4) SODIUM (UNII: M2 COLS (UNII: 3WJ0 ristics white ROUND 15 in 1 BOTTLE; 7 20 in 1 BOTTLE; 7 30 in 1 BOTTLE; 7	Score Size Imprint Code Ckage Description Type 0: Not a Combination Proc	8 mm DAN;5658 Marke 1007 09/28/20 1007 09/28/20 1007 09/28/20 1007 09/28/20	ting S 1 0 16 0 16 0 16			ıg End Date	

Establishment								
Name	Address	ID/FEI	Business Operations					
RedPharm Drug, Inc.		828374897	relabel(67296-0007), repack(67296-0007)					

Revised: 9/2016

RedPharm Drug, Inc.