



PRODUCT INFORMATION

CLARINEX® (desloratadine) TABLETS, REDITABS® TABLETS

DESCRIPTION: CLARINEX (desloratadine) Tablets are light blue, round, film coated tablets containing 5 mg desloratadine, an antihistamine, to be administered orally. If also contains the following excipients: dibasic calcium phosphate dihydrate USP microcrystalline cellulose NF, corn starch NF, talc USP, carnauba wax NF, white wax NF, coating material consisting of lactose monohydrate, hydroxypropyl methylcelluses, titanium dioxide, polyethylene glycol, and FD&C Blue #2 Aluminum Lake.

The CLARINEX RediTabs* brand of desloratadine orally-disintegrating tablets is a pink colored, round, tablet shaped unit with a "C" debossed on one side. Each RediTabs unit contains 5 mg of desloratadine. It also contains the following inactive ingredients: gelatin Type B NF, mannitol USP, sapartame NF, polarcrillin potassium NF, citric acid USP, red dye and tutti frutti flavoring.

Desloratadine is a white to off-white powder that is slightly soluble in water, but very soluble in ethanol and propylene glycol. It has an empirical formula: C₁₉H₁₉ClN₂ and a molecular weight of 310.8. The chemical name is 8-chloro-6,11-dihydro-11-(4-piperdinylidene)-5*H*-benzo[5,6]cyclohepta[1,2-*b*]pyridine and has the following structure:

CLINICAL PHARPMACOLOGY. Mechanism of Action: Deskoratadine is a long-acting tricyclic histamine anaponist with selective H, receptor histamine anaponist activity. Receptor binding data indicates that at a concentraliation of 2-3 april. (7 nanomalar), desforatadine shows significant interaction with the human histamine H, receptor bending data indicates that at a concentraliation of 2-3 april. (7 nanomalar), desforatadine inhibited histamine release from human mast cells in vitro. Results of a radiolabeled tissue distribution study in rata and a radioligand H, receptor binding study in guines pigs showed that desforatadine cells in vitro. Results of a radiolabeled tissue distribution study in rata and a radioligand H, receptor binding study in guines pigs showed but necess the man time to maximum plasma concentrations (T_{m-m}) occurred at approximately 3 hours post dose and mans steady state peak plasma concentrations (C_{m-m}) and area under the concentration-time curve (AUC) of 4 ng/mL and 56.9 ng hr/mL were observed, respectively, Nelher food nor grapefurly juice had an effect on the bioavailability (C_{mux} and AUC) of desforatadine. Profile of CLARINEX Redifabs Tablets was valuated in a three way crossover study in 30 adult volunteers, 5 anglie CLARINEX Redifabs three way crossover study in 30 adult volunteers, 5 od or water had no effect on the bioavailability (AUC and C_{mux}) of CLARINEX Redifabs Tablets, however, food shifted the desforatadine median T_{mux} value from 2.5 to 4 hr.

Distribution: Desforatadine and 3-hydroxydesloradadine, and 3-hydroxydesloradadine and 3-hydroxydeslor

Drug Interactions: In two controlled crossover clinical pharmacology studies in healthy male (n=12 in each study) and female (n=12 in each study) volunteers, destorated in 7.5 mg (1.5 times the daily dose) once daily was coadministered with erythromycin 500 mg every 8 hours or ketoconazole 200 mg every 12 hours for 10 days. In 3 separate controlled, parallel group clinical pharmacology studies, desloratadine at the clinical dose of 5 mg has been coadministered with azithromycin 500 mg followed by 250 mg once daily for 4 days (n=18) or with fluoxetine 20 mg once daily for 7 days after a 23 day pretreatment period with fluoxetine (n=18) or with cimelidine 600 mg every 12 hours for 14 days (n=18) under steady state conditions to normal healthy male and female volunteers. Although increased plasma concentrations (C_{max} and AUC 0-24 hrs) of desloratadine and 3-hydroxydesloratadine were observed (see Table 1), there were no clinically relevant changes in the safety profile of desloratadine, as assessed by electrocardiographic parameters (including the corrected QT interval), clinical laboratory tests, vital signs, and adverse events.

Table 1
Changes in Desloratadine and 3-Hydroxydesloratadine
Pharmacokinetics in Healthy Male and Female Volunteers 3-Hydroxy-desloratadine Desloratadine AUC 0-24 hrs ALIC 0-24 hrs Erythromycin (500 mg Q8h) +14% +43% +40% +24% Ketoconazole (200 mg Q12h) +39% +43% +72% +45%

Azithromycin

(500 mg day 1,							
250 mg QD x 4 days)	+15%	+5%	+15%	+4%			
Fluoxetine							
(20 mg QD)	+15%	+0%	+17%	+13%			
Cimetidine							
(600 mg Q12h)	+12%	+19%	-11%	-3%			
Pharmacodynamics: Wheal and Flare: Human histamine skin wheal studies following single and repeated 5 mg doses of desloratadine have shown that the drug exhibits an antihistaminic effect by 1 hour: this activity may persist for as long as 24 hours. There was no evidence of histamine-induced skin wheal tachyphylaxis within the desloratadine 5 mg group over the 28 day treatment period. The clinical relevance of histamine wheal skin testing is unknown. Effects on QT; Single dose administration of desloratadine did not alter the corrected QT interval (QT;) in rats (up to 12 mg/kg, oral), or quinea pigs							
(25 mg/kg, intravenous). R) in rats (up to 12 m	ig/kg, oral), or	guinea p	oigs		

rected QT interval (QT_c) in rats (up to 12 mg/kg, oral), or guinea pigs (25 mg/kg, intravenous). Repeated oral administration at doses up to 24 mg/kg for durations up to 3 months in monkeys did not alter the QT_c at an estimated desloratedine exposure (AUC) that was approximately 955 times the mean AUC in humans at the recommended daily oral dose. See **OVERDOSAGE** section for information on human QT_c experience.

Clinical Trials: Seasonal Allergic Rhinitis: The clinical efficacy and safety of CLARINEX Tablets were evaluated in over 2,300 patients 12 to 75 years of age with seasonal allergic rhinitis. A total of 1,838 patients received 2.5-20 mg/day of CLARINEX in 4 double-blind, randomized, placebo-controlled clinical trials of 2 to 4 weeks' duration conducted in the United States. The results of these studies demonstrated the efficacy and safety of CLARINEX 5 mg in the treatment of adult and adolescent patients with seasonal allergic rhinitis. In a dose ranging trial, CLARINEX 2.5-20 mg/day was studied. Doses of 5, 7.5, 10, and 20 mg/day were superior to placebo; and no additional benefit was seen at doses above 5.0 mg. In the same study, an increase in the incidence of somnolence was observed at doses of 10 mg/day and 20 mg/day (5.2% and 7.6%, respectively), compared to placebo (2.3%). In 2 four-week studies of 924 patients (aged 15 to 75 years) with seasonal allergic rhinitis and concomitant asthma, CLARINEX Tablets 5 mg once daily improved rhinitis symptoms, with no decrease in pulmonary function. This supports the safety of administering CLARINEX Tablets to adult patients with seasonal allergic rhinitis with mild to moderate asthma.

CLARINEX Tablets 5 mg once daily significantly reduced the Total Symptom

Table 2 TOTAL SYMPTOM SCORE (TSS) Changes in a 2 Week Clinical Trial in Patients with Seasonal Allergic Rhinitis Treatment Mean Change Placebo Group Baseline* from Comparison (n) (sem) Baseline** (P- value)	CLARINEX Tablets 5 Scores (the sum of indi with seasonal allergic rh	mg once daily si vidual scores of na	gnificantly reduced sal and non-nasal s	I the Total Symptom ymptoms) in patients			
Changes in a 2 Week Clinical Trial in Patients with Seasonal Allergic Rhinitis Treatment Mean Change Placebo Group Baseline* from Comparison (n) (sem) Baseline** (P-value)	with seasonal allergie in		e 2				
Treatment Mean Change Placebo Group Baseline* from Comparison (n) (sem) Baseline** (P-value)							
Group Baseline* from Comparison (n) (sem) Baseline** (P- value)							
(n) (sem) Baseline** (P- value)							
(Serii)	(11)	(Sem)		(P- value)			
			(Sem)				

CLARINEX 5.0 mg (171) 14.2 (0.3) 13.7 (0.3) P<0.01 4.3 (0.3) İ Placebo (173) 2.5 (0.3) * At baseline, a total nasal symptom score (sum of 4 individual symptoms) of at least 6 and a total non-nasal symptom score (sum of 4 individual symptoms) of least 5 (each symptom scored 0 to 3 where 0=no symptom and 3=severe symptoms) was required for trial eligibility. TSS ranges from 0=no symptoms to 24=maximal symptoms.

24-maximal symptoms.

**Mean reduction in TSS averaged over the 2-week treatment period.

There were no significant differences in the effectiveness of CLARINEX Tablets 5 mg across subgroups of patients defined by gender, age, or race.

Perennial Allergic Rhinitis: The clinical efficacy and safety of CLARINEX Tablets 5 mg were evaluated in over 1,300 patients 12 to 80 years of age with perennial allergic rhinitis. A total of 685 patients received 5 mg/day of CLARINEX in 2 double-blind, randomized, placebo-controlled clinical trials of 4 weeks' duration conducted in the United States and internationally. In one of these studies CLARINEX Tablets 5 mg once daily was shown to significantly reduce symptoms of perennial allergic rhinitis (Table 3).

Table 3

TOTAL SYMPTOM SCORE (TSS)

Changes in a 4 Week Clinical Trial in Patients with Perennial Allergic Rhinitis

Treatment Mean Change Placebo

Group Baseline* from Comparison Group compariso (P- value) Bas selin (sem) CLARINEX 5.0 mg (337) Placebo (337) 4.06 (0.21) 12.37 (0.18) 12.30 (0.18) P=0.01

Placebo (337) 12.30 (0.18) -3.27 (0.21)

* At baseline, average of total symptom score (sum of 5 individual nasal symptoms and 3 non-nasal symptoms, each symptom score 0 to 3 where 0-no symptom and 3-severe symptoms) of at least 10 was required for trial eligibility. TSS ranges from 0-no symptoms to 24-maximal symptoms.

**Mean reduction in TSS averaged over the 4-week treatment period.

Chronic Idiopathic Urticaria: The efficacy and safety of CLARINEX Tablets 5 mg once daily was studied in 416 chronic idiopathic urticaria patients 12 to 84 years of age, of whom 211 received CLARINEX. In two double-blind, placebo-controlled, randomized clinical trials of six weeks' duration, at the pre-specified one-week primary time point evaluation, CLARINEX Tablets significantly reduced the severity of pruritus when compared to placebo (Table 4). Secondary endpoints were also evaluated and during the first week of therapy CLARINEX Tablets 5 mg reduced the secondary endpoints, "Number of Hives" and the "Size of the Largest Hive," when compared to placebo.

: Urticaria

Compared to placebo.

Table 4

PRURITUS SYMPTOM SCORE

Changes in the First Week of a Clinical Trial in Patients with Chronic Idiopathic Urt

Treatment Mean Change Placebo

Group Baseline from Comparison

(sem) Baseline* (P- value)

(sem) CLARINEX 5.0 mg (115) 2.19 (0.04) P<0.01 Placebo (110) 2.21 (0.04) -0.52 (0.07)

Pruritus scored 0 to 3 where 0=no symptom to 3=maximal symptom Placebo (110)

*Mean reduction in pruritus averaged over the first week of treatment

INDICATIONS AND USAGE: Allergic Rhinitis: CLARINEX Tablets 5 mg are indicated for the relief of the nasal and non-nasal symptoms of allergic rhinitis (seasonal and perennial) in patients 12 years of age and older.

Chronic Idiopathic Urticaria: CLARINEX Tablets are indicated for the symptomatic relief of pruritus, reduction in the number of hives, and size of hives, in patients with chronic Idiopathic urticaria 12 years of age and older.

CONTRAINDICATIONS: CLARINEX Tablets 5 mg are contraindicated in patients who are hypersensitive to this medication or to any of its ingredients, or to loratadine.

DECALITIONS: Craricareactic Michaelescie Investment of Entilitive The ear.

Chronic Idiopathic Urticaria: CLARINEX Tablets are indicated for the symptomatic relief of puritivis, reduction in the number of hives, and size of hives, in patients with chronic idiopathic urticaria 12 years of age and older.

CONTRAINDICATIONS: CLARINEX Tablets 5 may are contraindicated in patients who are hypersensitive to this medication or to any of its ingredients, or to loratadine. PRECAUTIONS: Carcinogenesis, Mutagenesis, Impairment of Fertility. The carcinogenic potential of desioratadine was assessed using loratadine was administered in the diet at doses up to 40 mg/kg/day in mice (estimated desioratadine was administered in the diet at doses up to 40 mg/kg/day in mice (estimated desioratadine and desioratadine metabolitie exposures were approximately 30 times the AUC in humans at the recommended daily oral dose). Male mice given 40 mg/kg/day loratadine had a significantly higher incidence of hepatocellular tumors (combined adenomas and carcinomas) than concurrent controls. In rats, a significantly higher incidence of hepatocellular tumors (combined adenomas and carcinomas) was observed in males given 10 mg/kg/day and in males and females given 25 mg/kg/day. The estimated desioratadine and desioratadine metabolitie exposures of rats given 10 mg/kg/day and in males and females given 25 mg/kg/day. The estimated desioratadine and desioratadine man at the recommended daily oral dose. The clinical significance of these findings during ingentern use of desioratadine is not known.

In gentotokity studies with desioratadine, there was no evidence of genotoxic potential in a reverse mutation assay (Salmonella/E. colf mammalian microsome bacterial mutagenicity assay) or in two assays for chromosomal abertations (unma peripheral blood lymphocyte clastogenicity assay) and mouse bone marrow the complex of the particular changes, occurred at an oral desioratadine and desioratadine metabolitie exposures were approximately 30 times the AUC in humans at the recommended daily oral dose, and the particular desioratadin

 $\label{eq:continuous} \frac{\text{Table 5}}{\text{Incidence of Adverse Events Reported by } \geq 2\%$ of Allergic Rhinitis Patients in Placebo-Controlled, Multiple-Dose Clinical Trials CLARINEX Tablets 5 m (n=1,655) Placebo n=1,652) Adverse Experience 2.0% 1.9% 1.8% 1.2% 4.1% 3.0% 2.1% 2.1% 2.1% 2.1% Pharyngitis Dry Mouth Myalgia Fatigue Somnolence

1.8% 1.6% Dysmenorrhea The frequency and magnitude of laboratory and electrocardiographic abnormalities were similar in CLARINEX and placebo-treated patients.

There were no differences in adverse events for subgroups of patients as defined by gender, age, or race.
Chronic Idiopathic Urticaria: In multiple-dose, placebo-controlled trials of chronic idiopathic urticaria, 211 patients received CLARINEX Tablets and 205 received placebo. Adverse events that were reported by greater than or equal to 2% o patients who received CLARINEX Tablets and that were more common with CLARINEX than placebo were (rates for CLARINEX and placebo, respectively) headache (14%, 13%), nausea (5%, 2%), fatigue (5%, 1%), dizziness (4%, 3%) pharynglits (3%, 2%), dyspepsia (3%, 1%), and myalgia (3%, 1%). The following spontaneous adverse events have been reported during the mar
keting of decloratedine: tachycardia (palnitations) and rarely hypersensitivity reac

keting of destoratadine: tachycardia palpitations, and rarelý hypersensitívity reac-tions (such as rash, pruritus, urticarla, edema, dyspnea, and anaphylaxis), and elevated liver enzymes including bilirubin.

DRUG ABUSE AND DEPENDENCE: There is no information to indicate that abus or dependency occurs with CLARINEX Tablets. DRUG ABUSE AND DEPENDENCE: Intere is no information to indicate that abuse or dependency occurs with CLARINEX Tablets.

OVERDOSAGE: Information regarding acute overdosage is limited to experience from clinical trials conducted during the development of the CLARINEX product. In a dose ranging trial, at doses of 10 mg and 20 mg/day somnolence was reported. Single daily doses of 45 mg were given to normal male and female volunteers for 0 days. All ECGs obtained in this study were manually read in a blinded fashion by a cardiologist. In CLARINEX-treated subjects, there was an increase in mean heart rate of 9.2 bpm relative to placebo. The QT interval was corrected for heart rate (QT.) by both the Bazett and Fridericia methods. Using the OT, (Bazett) there was a mean increase of 8.1 msec in CLARINEX-treated subjects relative to placebo. Using QT. (Fridericia) there was a mean increase of 0.4 msec in CLARINEX-treated subjects relative to placebo. No clinically relevant adverse events were reported. In the event of overdose, consider standard measures to remove any unabsorbed drug. Symptomatic and supportive treatment is recommended. Desloratadine and 3-hydroxydesloratadine are not eliminated by hemodialysis. Lethality occurred in rats at oral doses of 250 mg/kg or greater (estimated desloratadine and desloratadine metabolite exposures were approximately 120 times the AUC in humans at the recommended daily oral dose). The oral median lethal dose in mice was 353 mg/kg (estimated desloratadine exposures were approximately 290 times the human daily oral dose on a mg/m² basis). No deaths occurred at oral doses up to 250 mg/kg in monkeys (estimated desloratadine exposures were approximately 810 times the human daily oral dose on a mg/m² basis).

approximately a for unlines the number daily of an adults and children 12 years of age and over, the recommended dose of CLARINEX Tablets is 5 mg once daily. In patients with liver or renal impairment, a starting dose of one 5 mg tablet every other day is recommended based on pharmacokinetic data.

Administration of CLARINEX RediTabs Tablets: Place CLARINEX RediTabs Tablet on the tongue. Tablet dishitegration occurs rapidly. Administer with or without water. Take tablet immediately after opening the blister.

NOW SUPPLIED: CLARINEX Tablets: Embossed "C5", light blue film coated tablets: that are packaged in high-density polyethylene plastic bottles of 100 (NDC 0085-1264-02). Also available, CLARINEX Unit-of-Use package of 30 tablets (3 x 10; 10 blisters per card) (NDC 0085-1264-04); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 blisters per card) (NDC 0085-1264-04); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 blisters per card) (NDC 0085-1264-05); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 blisters per card) (NDC 0085-1264-05); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 blisters per card) (NDC 0085-1264-05); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 blisters per card) (NDC 0085-1264-05); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 blisters per card) (NDC 0085-1264-05); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 blisters per card) (NDC 0085-1264-05); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 blisters per card) (NDC 0085-1264-05); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 blisters per card) (NDC 0085-1264-05); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 blisters per card) (NDC 0085-1264-05); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 blisters per card) (NDC 0085-1264-05); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 blisters per card) (NDC 0085-1264-05); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 blisters per card) (NDC 0085-1264-05); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 blisters per card) (NDC 0085-1264-05); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 blisters per card) (NDC 0085-1264-05); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 blisters per card) (NDC 0085-1264-05); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 blisters per card) (NDC 0085-1264-05); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10 tblisters per card) (NDC 0085-1264-05); and Unit Dose-Hospital Pack of 100 Tablets (10 x 10; 10

Protect Unit-of-Use packaging and Unit Dose-Hospital Pack from excessive ofsture. Store between 2° and 25°C (36° and 77°F). Heat Sensitive. Avoid exposure at or above 30°C (86°F).

CLARINEX REDITABS (designated or ally-disintegrating tablets) 5 mg: "C" debossed, pink tablets in foil/foil blisters. Packs of 30 tablets (containing 3 x 10s) NDC 0085-1280-01.

Store REDITABS TABLETS at 25°C (77°F); excursions permitted between 15°-30°C (59°-86°F) [See USP Controlled Room Temperature].

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Schering Corporation Kenilworth, NJ 07033 USA

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TABLETS, REDITABS® TABLETS

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