

PRODUCT MONOGRAPH

CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH
Modified-Release Caplets
(loratadine 10 mg/pseudoephedrine sulfate 240 mg)

Histamine H₁ receptor antagonist/Sympathomimetic amine

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PRODUCT MONOGRAPH

NAME OF DRUG

CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH
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(loratadine 10 mg/pseudoephedrine sulfate 240mg)

PART I: HEALTH PROFESSIONAL INFORMATION

THERAPEUTIC CLASSIFICATION

Histamine H₁ receptor antagonist/Sympathomimetic Amine

ACTIONS AND CLINICAL PHARMACOLOGY

Loratadine is a long-acting tricyclic antihistamine with selective peripheral H₁ receptor antagonistic activity. It exhibits a dose-related inhibition of the histamine-induced skin wheal and flare response in humans which is rapid in onset, is apparent at two hours and persists throughout the 24 hour observation period (Roman et al. 1986). Single oral doses up to 160 mg and repeat daily doses of 40 mg for up to 13 weeks were well tolerated with the incidence of sedation and dry mouth being no different from placebo.

¹⁴C-loratadine is rapidly absorbed reaching C_{max} values (4.7, 10.8 and 26.1 ng/mL) at 1.5, 1.0 and 1.3 hours for the 10, 20 and 40 mg dose, respectively. The loratadine elimination half-life (T-1/2 β) ranged from 7.8-11.0 hours.

Descarboethoxy-loratadine, the major active metabolite, reached C_{max} values (4.0, 9.9 and 16.0 ng/mL) at 3.7, 1.5 and 2.0 hours after a dose of 10, 20 and 40 mg, respectively. Its T-1/2 β ranged from 17 to 24 hours. The accumulation indices, calculated by C_{max} and the area under the curve (AUC) ratios did not change after the 5th day, indicating little or no accumulation of

either loratadine or its metabolite after a multiple once per day dosage regimen. The T-1/2 β at steady state levels for loratadine and its active metabolite were 14.4 and 18.7 hours, respectively, similar to that reported following a single oral dose (Hilbert et al. 1987).

Approximately 82% of the ¹⁴C-loratadine dose is excreted in the urine (40%) and feces (42%) over a 10-day period. Approximately 27% of the dose is eliminated in the urine during the first 24 hours largely in the conjugated form. Unchanged drug is present only in trace quantities in the urine and the active metabolite descarboethoxyloratadine represents only 0.4 to 0.6% of the administered loratadine dose.

Pseudoephedrine, one of the naturally occurring alkaloids of Ephedra and an orally administered vasoconstrictor, produces a gradual but sustained decongestant effect facilitating shrinkage of congested mucosa in upper respiratory areas. The mucous membrane of the respiratory tract is decongested through the action of the sympathetic nerves.

A drug interaction cross-over study was performed to compare CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplets to the individual components (loratadine 10 mg and pseudoephedrine sulfate 240 mg). Coadministration of loratadine did not affect the bioavailability of pseudoephedrine. Similarly, coadministration of pseudoephedrine did not affect the pharmacokinetics of descarboethoxyloratadine although it resulted in the slightly higher (8%) bioavailability of loratadine: C_{max}=2.79 ng/mL when administered in combination versus C_{max}=2.55 ng/mL when administered alone. This is not considered to be of clinical significance.

Another study was conducted to characterize and compare the pharmacokinetic profile of loratadine, descarboethoxyloratadine and pseudoephedrine following oral administration of CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH given once daily and CLARITIN[®] ALLERGY + SINUS (loratadine 5 mg / pseudoephedrine sulfate 120 mg) given every 12 hours. The results of this study show that after multiple doses to steady state, CLARITIN[®] ALLERGY

+ SINUS EXTRA STRENGTH and the CLARITIN[®] ALLERGY + SINUS comparator were equivalent with respect to the bioavailability of loratadine and descarboethoxyloratadine (based on AUC), and bioequivalent for pseudoephedrine.

INDICATIONS AND CLINICAL USES

CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH (loratadine 10 mg/pseudoephedrine sulfate 240 mg) modified-release caplets are indicated for the fast and long-lasting relief of symptoms associated with allergic rhinitis, including nasal congestion, sinus pressure and sinus congestion, sneezing, postnasal drip/discharge and tearing and redness of the eyes. They are intended for short-term use only unless taken under medical supervision.

CONTRAINDICATIONS

CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH (loratadine 10 mg/pseudoephedrine sulfate 240 mg) modified-release caplets are contraindicated in those patients who have shown sensitivity or idiosyncrasy to their components including the metabolite Descarboethoxyloratadine, to adrenergic agents or to other drugs of similar chemical structures. They are also contraindicated in patients receiving MAO inhibitor therapy or within 14 days of discontinuing such treatment and in patients with narrow-angle glaucoma, urinary retention, hypertension, severe coronary artery disease and hyperthyroidism.

PRECAUTIONS

General

Sympathomimetics should be used with caution in patients with stenosing peptic ulcer, pyloroduodenal obstruction, prostatic hypertrophy or bladder neck obstruction, cardiovascular disease, increased intraocular pressure or diabetes mellitus.

Sympathomimetics should be used with caution in patients receiving digitalis.

Sympathomimetics may cause central nervous system (CNS) stimulation and convulsions or cardiovascular collapse with accompanying hypotension.

Patients with severe liver impairment should be administered a lower initial dose because they may have reduced clearance of loratadine. For CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH, an initial dose of one caplet every other day is recommended.

Patients who have a history of difficulty in swallowing tablets or who have upper gastrointestinal narrowing or abnormal esophageal peristalsis should not use this product.

Acute generalized exanthematous pustulosis (AGEP), a form of severe skin reaction, may occur with pseudoephedrine-containing products in isolated cases. If signs and symptoms such as the sudden occurrence of small (generalized) pustules, erythema, or fever are observed, patients should discontinue using the drug.

Use in Elderly

In patients 60 years of age or older, sympathomimetics are also more likely to cause adverse reactions such as confusion, hallucination, convulsions, CNS depression and death. Consequently, caution should be exercised when administering a long-acting formulation to this patient group.

Dependence Liability

There are no data available to indicate that abuse or dependency occurs with loratadine.

Pseudoephedrine sulfate, like other CNS stimulants, has been abused. At high doses, subjects commonly experience mood elevation, decreased appetite and a sense of increased energy, physical strength, mental capacity and alertness. Anxiety, irritability and loquacity also have been reported. With continued use, tolerance develops; the user increases the dose and ultimately toxicity occurs. Depression may follow rapid withdrawal.

Use in Children

Safety and efficacy of CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplets in children younger than 12 years of age have not yet been established.

Use in Obstetrics

The safe use of CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplets during pregnancy or lactation has not been established and is therefore not recommended for use in pregnant or nursing mothers.

Nursing Mothers

Loratadine and its active metabolite are eliminated in the breast milk of lactating women with milk concentrations being similar to plasma concentrations. Through 48 hours after dosing, only 0.029% of the loratadine dose is eliminated in the milk as unchanged loratadine and its active metabolite, descarboethoxyloratadine.

Pseudoephedrine has been reported to be excreted into breast milk of lactating women. The use of CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplets in nursing mothers is therefore not recommended.

Long-Term use

Because of the lack of experience with long-term use of this drug, its use should be limited to three months unless recommended by a physician.

Drug Interactions

When administered concomitantly with alcohol, loratadine has no potentiating effect as measured by psychomotor performance studies.

When sympathomimetic drugs are given to patients receiving monoamine oxidase inhibitors (MAO), hypertensive reactions, including hypertensive crises, may occur. The antihypertensive effects of methyldopa, mecamylamine, reserpine and veratrum alkaloids may be reduced by sympathomimetics. Beta-adrenergic blocking agents may also interact with sympathomimetics. Increased ectopic pacemaker activity can occur when pseudoephedrine sulfate is used concomitantly with digitalis. Antacids increase the rate of pseudoephedrine sulfate absorption; kaolin decreases it. The antibacterial agent, furazolidone, is known to cause a dose-related inhibition of MAO. Although there are no reports of a hypertensive crisis caused by the concurrent administration of pseudoephedrine and furazolidone, they should not be taken together. Care should be taken in the administration of CLARITIN® ALLERGY + SINUS EXTRA STRENGTH concomitantly with other sympathomimetic amines because the combined effects on the cardiovascular system may be harmful to the patient.

Increase in plasma concentrations of loratadine have been reported after concomitant use with ketoconazole, erythromycin or cimetidine in controlled clinical trials, but without clinically significant changes (including electrocardiographic). Other drugs known to inhibit hepatic metabolism should be coadministered with caution until definitive interaction studies can be completed.

Drug/Laboratory Test Interactions:

Loratadine should be discontinued approximately 48 hours prior to skin testing procedures since antihistamines may prevent or diminish otherwise positive reactions to dermal reactivity indicators.

ADVERSE REACTIONS

Adverse experiences reported during the study with CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH (loratadine 10 mg/pseudoephedrine sulfate 240 mg) modified-release caplets, administered once daily, were similar to those previously encountered during treatment with CLARITIN[®] ALLERGY + SINUS tablets (loratadine 5 mg/pseudoephedrine sulfate 120 mg), administered twice daily. No unusual or unexpected adverse events were reported.

In clinical studies, the most frequently reported adverse events associated with CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplets were headache, dry mouth, insomnia and somnolence.

Table 1

Number (%) of Patients Reporting Adverse Experiences (probably or possibly related to treatment) $\geq 5\%$ incidence during treatment with CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplets, Loratadine, Pseudoephedrine sulfate or placebo in clinical studies.

Adverse Experience	<u>CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH</u>	<u>Loratadine</u>	<u>Pseudoephedrine Sulfate</u>	<u>Placebo</u>
	(N=583)	(N=217)	(N=220)	(N=370)
Dry Mouth	55 (9)	7(3)	16(7)	11(3)
Headache	53 (9)	21(10)	21(10)	39(11)
Insomnia	38 (7)	2(1)	17(8)	4(1)
Somnolence	47 (8)	9(4)	9(4)	14(4)

Rarely reported events in decreasing order of frequency included dizziness, fatigue, anorexia, nervousness, nausea, epistaxis, rhinitis, lacrimal gland disorder, asthenia, hyperkinesia, constipation, dyspepsia, palpitation, tachycardia, thirst, agitation, irritability, coughing, dyspnea, nasal irritation, and pharyngitis.

With exception of headache, which was occasionally severe, most of the adverse events associated with CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplets were mild to moderate in severity.

During the marketing of loratadine, alopecia, anaphylaxis (including angioedema), abnormal hepatic function, dizziness, palpitations and tachycardia have been reported rarely.

Very rare adverse events include convulsions or seizures which have been reported during the post-marketing of loratadine.

There were rare postmarketing reports of mechanical upper gastrointestinal tract obstruction in patients taking the original round tablet formulation of CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH. In many of these cases, patients have had a history of difficulty in swallowing tablets, or had known upper gastrointestinal narrowing or abnormal esophageal peristalsis.

SYMPTOMS AND TREATMENT OF OVERDOSAGE

In the event of overdose, treatment, which should be started immediately, is symptomatic and supportive. Discontinuation of use, gastric lavage and support vital functions are advised.

Manifestations

Somnolence, tachycardia and headache have been reported with overdoses of loratadine.

Symptoms associated with overdoses of sympathomimetics may vary from CNS depression (sedation, apnea, diminished mental alertness, cyanosis, coma, cardiovascular collapse) to stimulation (insomnia, hallucination, tremors or convulsions) to death. Other signs and symptoms may be euphoria, excitement, tachycardia, palpitations, thirst, perspiration, nausea, dizziness, tinnitus, ataxia, blurred vision and hypertension or hypotension. Stimulation is

particularly likely in children, as are atropine-like signs and symptoms (dry mouth; fixed, dilated pupils; flushing; hyperthermia; and gastrointestinal symptoms).

In large doses sympathomimetics may give rise to giddiness, headache, nausea, vomiting, sweating, thirst, tachycardia, precordial pain, palpitations, difficulty in micturition, muscular weakness and tenseness, anxiety, restlessness and insomnia. Many patients can present a toxic psychosis with delusions and hallucinations. Some may develop cardiac arrhythmias, circulatory collapse, convulsions, coma and respiratory failure.

Treatment

Adsorption of any drug in the stomach may be attempted by the administration of activated charcoal as a slurry with water. Gastric lavage should be considered. Physiologic saline solution is the lavage solution of choice, particularly in children. In adults, tap water can be used; however, as much as possible of the amount administered should be removed before the next instillation. Saline cathartics draw water into the bowel by osmosis and therefore may be valuable for their action in rapid dilution of bowel content. Loratadine is not removed by hemodialysis, it is not known if loratadine is removed by peritoneal dialysis. After emergency treatment, the patient should continue to be medically monitored.

Treatment of the signs and symptoms of overdose is symptomatic and supportive. Stimulants (analeptic agents) should not be used. Vasopressors may be used to treat hypotension. Short-acting barbiturates, diazepam or paraldehyde may be administered to control seizures. Hyperpyrexia, especially in children, may require treatment with tepid water sponge baths or hypothermic blanket. Apnea is treated with ventilatory support.

DOSAGE AND ADMINISTRATION

Adults and children 12 years of age and over:

One CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH (loratadine 10 mg/pseudoephedrine sulfate 240mg) modified-release caplet once daily taken whole with a glass

of water, preferably upon waking. CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH may be taken without regard to mealtime.

Patients who have a history of difficulty in swallowing tablets or who have upper gastrointestinal narrowing or abnormal esophageal peristalsis should not use this product (see PRECAUTIONS and ADVERSE REACTIONS).

PART II: SCIENTIFIC INFORMATION

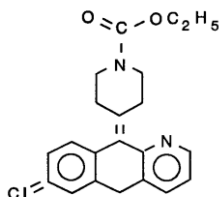
PHARMACEUTICAL INFORMATION

Drug substance

Proper name: loratadine (INN, USAN)

Chemical name: 1-Piperidinecarboxylic acid, 4-(8-chloro-5,6-dihydro-11H-benzo[5,6]cyclohepta[1,2-β]pyridin-11-ylidene)-, ethyl ester.

Structural formula:



Molecular formula: C₂₂H₂₃ClN₂O₂

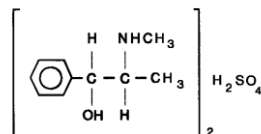
Molecular weight: 382.89

Appearance: White to off-white powder which melts between 131° and 137° C.

Proper name: pseudoephedrine sulfate (USP)

Chemical name: Benzenemethanol, α-[1-(methylamino)-ethyl]-, [S-(R*,R*)]-, sulfate (2:1) (salt)

Structural formula:



Molecular formula: $(C_{10}H_{15}NO)_2 \cdot H_2SO_4$

Molecular weight: 428.54

Appearance: White to off-white crystals or powder.

Composition

CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH is an modified-release caplet. Each CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH caplet contains 10 mg loratadine in the caplet coating and 240 mg pseudoephedrine sulfate in the modified-release core. The loratadine component is released immediately, whereas the pseudoephedrine sulfate component is released slowly from the core allowing for once daily administration.

The nonmedicinal ingredients in CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplet coating include: hydroxypropyl methylcellulose 2910, polyethylene glycol 3350, polyethylene glycol 400, sucrose, carnauba wax and Opaspray White*. The non-medical ingredients in CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplet core include: hydroxypropyl methylcellulose 2208, dibasic calcium phosphate dihydrate, ethylcellulose, povidone, silicon dioxide, magnesium stearate.

*contains titanium dioxide and hydroxypropyl methylcellulose.

Storage conditions

Store between 15° and 30° C. Protect from exposure to excessive moisture.

AVAILABILITY OF DOSAGE FORM

Description: White, oval, biconvex, coated caplet.

Packaging: Blister packaged in 15's.

PHARMACOLOGY

Human Pharmacology

Pharmacokinetics

A randomised, single-dose, open-label, five-way crossover study was conducted in order to evaluate the bioavailability of loratadine and desloratadine following administration of five different CLARITIN[®] formulations.

This study demonstrated that the mean C_{max} and AUC values for loratadine and desloratadine were bioequivalent between the CLARITIN[®] 10mg immediate release tablet and CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplets.

Table 2

Arithmetic mean pharmacokinetic parameters of loratadine and its major metabolite, desloratadine, for the CLARITIN[®] 10 mg immediate release tablet and CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplet.

	CLARITIN[®] 10 mg immediate release tablet	CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplet
Loratadine		
C _{max} (ng/ml)	3.12	3.68
AUC (ng . hr/ml)	8.76	9.34
T _{max} (hr)	1.52	1.23
Desloratadine		
C _{max} (ng/ml)	3.40	3.54
AUC (ng . hr/ml)	48.6	50.3
T _{max} (hr)	2.06	1.79

A multiple-dose study was conducted to determine the steady-state bioequivalence of CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplets administered

once daily as compared to a reference standard consisting of loratadine 10 mg tablet given once daily and pseudoephedrine sulfate 120 mg CLARITIN[®] ALLERGY + SINUS tablet in twice-a-day administration.

The results of the study showed that CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplets and the reference standard were equivalent with respect to the bioavailability of pseudoephedrine, although the comparator in this study was a delayed-release pseudoephedrine sulfate formulation intended for twice-a-day dosing. CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplets and the reference standard gave similar mean plasma concentrations for loratadine; however, no statistical conclusion regarding the bioequivalency could be made due to the low plasma drug concentrations and high intersubject variability.

A single-dose study was conducted to evaluate and compare the effect of food on the oral bioavailability of pseudoephedrine when administered as a CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplet or a pseudoephedrine sulfate SR (sustained release) tablet. The bioavailability of pseudoephedrine from the CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplet or the pseudoephedrine SR (sustained release) tablet was not affected significantly by food intake.

When administered with a high-fat meal, as compared with administration in a fasting state, the C_{max} of pseudoephedrine was 22% higher from the CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplet and 25% higher from the pseudoephedrine sulfate SR tablet: C_{max}=304.5 ng/mL when pseudoephedrine was given after a 10-hour fast, C_{max}=382.5 ng/mL when pseudoephedrine was given with breakfast and C_{max}=376.6 ng/mL when pseudoephedrine + loratadine was given with breakfast. However, this difference is not considered to be clinically relevant.

A second food effect study was conducted with CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplet. In this study, a standardized high-calorie, high-fat breakfast significantly increased the C_{max} and AUC of loratadine by a mean of 53% and 76%, respectively, compared to the administration of CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH under fasted conditions (p<0.05). In contrast to loratadine, there was a very small and non-significant increase in descarboethoxyloratadine C_{max} and AUC values when CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH was given with food. Concomitant food slightly (7%) but significantly (p<0.05) increased the mean peak plasma pseudoephedrine concentrations, without significantly affecting the rate or extent of pseudoephedrine absorption.

Considering the magnitude of changes, the pharmacodynamics and safety of pseudoephedrine and loratadine, the increases in the plasma concentrations of these compounds that may occur when CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH is given with food are not expected to be clinically important.

A single-dose study was conducted to characterize the pharmacokinetic profile of pseudoephedrine following oral administration of three specially formulated loratadine/pseudoephedrine sulfate tablets with different in vitro release profiles and the standard CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplet formulation. In vitro release profiles were characterized by "very fast", "fast", or "slow" dissolution tablets of loratadine/pseudoephedrine sulfate formulations. A positive correlation was obtained between in vitro dissolution rates and in vivo bioavailability of pseudoephedrine for varying formulations of loratadine/ pseudoephedrine sulfate.

Clinical Studies:

The efficacy of once-daily CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH was shown by its consistent pattern of superiority when compared with placebo in reducing the symptoms of seasonal allergic rhinitis. Total, total nasal, total nonnasal, rhinorrhea and nasal stuffiness symptom scores were significantly reduced (p<0.05) in patients treated with CLARITIN[®]

ALLERGY + SINUS EXTRA STRENGTH modified-release caplets compared to placebo. When compared with its individual components, improvement in symptom scores in patients treated with CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplets was consistently greater, numerically, than that seen in patients treated with either loratadine or pseudoephedrine alone.

The comparability of CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH modified-release caplets to CLARITIN[®] ALLERGY + SINUS was shown in two studies.

In one study, CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH and CLARITIN[®] ALLERGY + SINUS were generally comparable in improving total nasal, total nonnasal and total symptom scores. CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH and CLARITIN[®] ALLERGY + SINUS were significantly more effective than placebo in reducing composite scores at Day 4 and Endpoint ($p < 0.05$).

In another study, CLARITIN[®] ALLERGY + SINUS EXTRA STRENGTH and CLARITIN[®] ALLERGY + SINUS improvement in composite symptom scores was not different at key time points. Investigator evaluations of therapeutic response were consistent with symptom score evaluations.

TOXICOLOGY

Preclinical Data - Toxicity:

Loratadine/pseudoephedrine sulfate: In acute and single-dose studies, loratadine/pseudoephedrine sulfate tablets exhibited a low order of toxicity. Acute oral LD50 values ranged from approximately 600 mg/kg in mice to about 2000 mg/kg in rats. Cynomolgus monkeys tolerated single doses up to 240 mg/kg. Loratadine/pseudoephedrine sulfate tablets were no more toxic than either their individual components, and observed effects were generally related to the pseudoephedrine component.

Loratadine/pseudoephedrine sulfate tablets were administered orally for 3 months to rats and monkeys. Loratadine/pseudoephedrine sulfate tablets were well tolerated in rats at doses up to 200 mg/kg/day, which is 40 times the proposed maximum clinical dose. In monkeys, daily doses up to 50 mg/kg/day were also well tolerated. Severe toxicity was observed in monkeys at a dose of 125 mg/kg/day and was attributed to the effects of the pseudoephedrine component.

Loratadine

In acute and single-dose toxicity studies, loratadine exhibits a low order of toxicity. It is relatively well tolerated in rats and monkeys treated for periods up to 2 years. In these studies, rats received oral doses of loratadine ranging from 2 to 240 mg/kg/day while monkeys were given doses ranging from 0.4 to 90 mg/kg/day.

Pseudoephedrine sulfate

This sympathomimetic agent is known to be less toxic and to produce less side effects than the ephedrine isomers, while being as potent as ephedrine as a bronchodilator and nasal decongestant.

Teratogenicity, mutagenicity and carcinogenicity

The combination loratadine/pseudoephedrine sulfate tablets was not teratogenic when administered orally to rats and rabbits during the period of organogenesis. The course of pregnancy or embryo/fetal viability of rats was not affected at doses up to 150 mg/kg/day (30 times the proposed clinical dose).

Loratadine/pseudoephedrine sulfate tablets did not directly affect embryo/fetal viability or offspring development of rabbits at doses up to 120 mg/kg/day.

Carcinogenicity, mutagenicity and teratology studies demonstrate that loratadine is not carcinogenic, mutagenic or teratogenic.

Likewise, pseudoephedrine sulfate is not considered to be carcinogenic, mutagenic or teratogenic. Therefore, loratadine/pseudoephedrine sulfate tablets are no more toxic than loratadine or pseudoephedrine sulfate alone.

REFERENCES

1. Barnett A, Iorio LC, Kreutner W, Tozzi S, Ahn HS, Gulbenkian A. Evaluation of the CNS properties of SCH 29851, a potential non-sedating antihistamine. *Agents and Actions* 1984; 14: 590-597.
1. Dockhorn RJ, Bergner A, Connell JT, Falliers CJ, Grabiec SV, Weiler JM, Shellenberger MK. Safety and efficacy of loratadine (SCH 29851): A new non-sedating antihistamine in seasonal allergic rhinitis. *Annals of Allergy* 1987; 58: 407-411.
3. Dockhorn RJ, Shellenberger MK, Hassanien R, Trachelman L. Efficacy of SCH 434 (loratadine plus pseudoephedrine) versus components and placebo in seasonal allergic rhinitis. *Journal of Allergy and Clinical Immunology* 1988; 81: 178.
4. Ercoli N, Schachter RJ, Hueper WC, Lewis MN. The toxicologic and antihistaminic properties of N,N'-dimethyl-N'-phenyl-N'-(2-thienylmethyl) ethylene-diamine hydrochloride (diatrim). *Journal of Pharmacology and Experimental Therapeutics* 1948; 93: 210-222.
5. Grossman J, Schenkel EC, Bronsky E, Linzmayer I, Selner J, Lanier BQ, Moss BA. Double-blind control study of loratadine 5 mg plus pseudoephedrine 120 mg (SCH 434) in seasonal allergic rhinitis. *Annals of Allergy* 1987; 58: 280.
6. Hassan AB, Ayoub MM, Doghaim REM, Youssef HI, Eid MA. Studies on the effect of some antihistamines on the reproductive organs of male rats. *Veterinary Medical Journal* 1983; 31(2): 227-286.
7. Hébert J, Bédard P-M, Del Carpio J, Drouin M, Gutkowski A, Kabbash LG, Nedilski MM, Prévost M, Schulz JI, Turenne Y, Yang WH. Loratadine and pseudoephedrine sulfate: A double-blind, placebo-controlled comparison of a combination tablet (SCH 434) and its individual components in seasonal allergic rhinitis. *American Journal of Rhinology* 1988; 2(2): 71-75.
8. Hilbert J, Radwanski E, Weglein R, Van L, Perentesis G, Symchowicz S, Zampaglione N. Pharmacokinetics and dose proportionality of loratadine. *Journal of Clinical Pharmacology* 1987; 27: 694-698.
9. Horak F, Bruttmann G, Pedrali P, Weeke B, Frolung D, Wolff HH, Christophers E. A multicentric study of loratadine, terfenadine and placebo in patients with seasonal allergic rhinitis. *Arzneim-Forsch/Drug Research* 1988; 38: 124-128.

10. Kreutner W, Chapman RW, Gulbenkian A, Siegel MI. Antiallergic activity of loratadine, a non-sedating antihistamine. *Allergy* 1987; 42: 57-63.
11. Roman IJ, Kassem N, Gural RP, Herron J. Suppression of histamine-induced wheal response by loratadine (SCH 29851) over 28 days in man. *Annals of Allergy* 1986; 57: 253-256.
12. Roth T, Roehrs T, Koshorek G, Sickelsteel J, Zorick F. Sedative effects of antihistamines. *Journal of Allergy and Clinical Immunology* 1987; 80: 94-98.

PART III: CONSUMER INFORMATION
CLARITIN® ALLERGY + SINUS EXTRA
STRENGTH

(10 mg loratadine/240 mg pseudoephedrine sulfate modified-release caplets)

This leaflet is part III of a three part "Product Monograph" published when CLARITIN® ALLERGY + SINUS EXTRA STRENGTH was approved for sale in Canada and is designed specifically for Consumers. This leaflet is a summary and will not tell you everything about CLARITIN® ALLERGY + SINUS EXTRA STRENGTH. Contact your doctor or pharmacist if you have any questions about the drug.

ABOUT THIS MEDICATION

What the medication is used for:

CLARITIN® ALLERGY + SINUS EXTRA STRENGTH provides:

- fast and long-lasting relief of symptoms from indoor (dust mites, pet dander, moulds) and outdoor (pollen, trees, grass, ragweed) allergies (allergic rhinitis) including itchy, watery, red burning eyes; sneezing, runny nose, itchy nose, post-nasal drip/discharge; nasal congestion, sinus pressure and sinus congestion.

What it does:

CLARITIN® ALLERGY + SINUS EXTRA STRENGTH is a fast acting and long-lasting antihistamine + decongestant that contains:

- an antihistamine (loratadine), which blocks the action of histamine and relieves allergy symptoms. When you are exposed to indoor or outdoor allergens your body responds by releasing histamine. Histamine causes allergy symptoms such as itchy, watery, red burning eyes; sneezing, runny nose, itchy nose, and post-nasal drip/discharge.
- a decongestant (pseudoephedrine sulfate) that relieves nasal congestion, sinus pressure and sinus congestion due to allergies by constricting the blood vessels in the lining of the nose and sinuses.

One dose of CLARITIN® ALLERGY + SINUS EXTRA STRENGTH provides 24 hour relief of allergy symptoms.

When it should not be used:

CLARITIN® ALLERGY + SINUS EXTRA STRENGTH should not be used if you are/have:

- allergic to loratadine, desloratadine, pseudoephedrine or to any of the product ingredients (see What the important non-medicinal ingredients are);
- taking a monoamine oxidase inhibitor (MAOI) (drugs for depression or Parkinson's disease) or for 2 weeks after stopping the MAOI drug;
- narrow-angle glaucoma (increased pressure in the eye);
- difficulty urinating due to enlargement of the prostate gland;
- high blood pressure;
- hyperthyroidism (overactive thyroid);
- heart disease.

What the medicinal ingredients are:

- Loratadine
- Pseudoephedrine sulfate

What the important non-medicinal ingredients are:

CLARITIN® ALLERGY + SINUS EXTRA STRENGTH (in alphabetical order): carnauba wax, dibasic calcium phosphate dihydrate, ethylcellulose, hydroxypropyl cellulose, hydroxypropyl methylcellulose, magnesium stearate, polyethylene glycol, povidone, silicon dioxide, sucrose, and titanium dioxide.

What dosage forms it comes in:

Modified-release caplets: containing 10 mg loratadine/240 mg pseudoephedrine sulfate.

WARNINGS AND PRECAUTIONS

BEFORE you use CLARITIN® ALLERGY + SINUS EXTRA STRENGTH talk to your doctor or pharmacist if you are/have:

- pregnant or breastfeeding;
- elderly;
- taking other medications;
- liver or kidney disease as you may require a dose adjustment;
- diabetes;
- difficulty in swallowing caplets;
- stomach problems.

Stop use and ask a doctor if:

- symptoms do not improve within 7 days or are accompanied by skin blister, redness, rash or fever.
- nervousness, dizziness or sleeplessness occurs.

Stop taking CLARITIN® ALLERGY + SINUS EXTRA STRENGTH 48 hours prior to any skin testing procedures.

INTERACTIONS WITH THIS MEDICATION

The drugs that may interact with CLARITIN® ALLERGY + SINUS EXTRA STRENGTH: monoamine oxidase inhibitors, methyl dopa, mecamylamine, reserpine and veratrum alkaloids, beta-adrenergic blocking agents, digitalis, antacids, kaolin, furazolidone, ketoconazole, erythromycin, cimetidine or other sympathomimetic amines, it is important to ask your doctor or pharmacist before taking CLARITIN® ALLERGY + SINUS EXTRA STRENGTH.

PROPER USE OF THIS MEDICATION

Usual dose:

Adults and children (12 years of age and older): one caplet daily.

- Do not crush, break or chew the caplet. Swallow whole with water.
- May be taken with or without food.
- Limited to 3 months of use unless recommended by a doctor.

Overdose:

In case of drug overdose, contact your Poison Control Centre, doctor or pharmacist as soon as possible, even if there are no symptoms.

Missed Dose:

If you miss taking your dose on time, do not worry; take your dose when you remember. Do not exceed more than one dose in 24 hours.

SIDE EFFECTS AND WHAT TO DO ABOUT THEM

Along with its desired effects, CLARITIN® ALLERGY + SINUS EXTRA STRENGTH may cause undesirable effects.

Side effects that may occur include dizziness, dry mouth, fatigue, headache, sleeplessness, nervousness, nausea, stomach discomfort and sleepiness. Taking more than directed may cause drowsiness. If these side effects do not go away or worsen, stop use and call your doctor or pharmacist.

SERIOUS SIDE EFFECTS, HOW OFTEN THEY HAPPEN AND WHAT TO DO ABOUT THEM

Symptom / effect	Talk with your doctor or pharmacist		Stop taking drug and call your doctor or pharmacist
	Only if severe	In all cases	
Rarely	Allergic reaction (rash, swelling, difficulty in breathing)		√
	Fast heart rate or heart palpitations		√
	Liver dysfunction		√
Very Rare	Convulsions or Seizures		√

This is not a complete list of side effects. For any unexpected effects while taking CLARITIN® ALLERGY + SINUS EXTRA STRENGTH, contact your doctor or pharmacist.

HOW TO STORE IT

Store between 15° and 30°C. Protect from exposure to excessive moisture.

Keep this and all medication stored in a safe place and out of reach of children.

Reporting Side Effects

You can report any suspected side effects associated with the use of health products to Health Canada by:

- Visiting the Web page on Adverse Reaction Reporting (<http://www.hc-sc.gc.ca/dhp-mps/medeff/report-declaration/index-eng.php>) for information on how to report online, by mail or by fax; or
- Calling toll-free at 1-866-234-2345.

NOTE: Contact your health professional if you need information about how to manage your side effects. The Canada Vigilance Program does not provide medical advice.

MORE INFORMATION

This document plus the full product monograph, prepared for health professionals, can be found at: www.bayer.ca

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