LEVOCABASTINE is a highly potent and selective H₁-receptor antagonist specifically developed for topical administration by ocular and nasal routes. The clinical effects of levocabastine occur rapidly and are predominantly due to local antihistaminic effects at the site of application. Clinically, levocabastine is well tolerated with an adverse effect profile comparable with that of sodium cromoglycate and placebo. As might be expected from the route of drug administration, local irritation is the most frequent adverse event seen with levocabastine eye drops and nasal spray with an incidence comparable with that in placebo-treated controls. Intranasal application of levocabastine has been shown to have no adverse effect on ciliary activity both in vitro and in vivo, while ocular administration has not been shown to have any significant or consistent adverse effect in both animal and human studies. At therapeutic doses, levocabastine appears to be devoid of significant systemic activity producing no apparent effects on cardiovascular, psychomotor and cognitive function. Since levocabastine undergoes little hepatic metabolism, and only low plasma levels of the drug are attained following topical administration, drug interactions are unlikely.

Key words: H₁-receptor antagonist, Levocabastine, Tolerability, Topical antihistamine

Introduction

Histamine H₁-receptor antagonists are widely considered to be a primary treatment option for the clinical management of allergic rhinoconjunctivitis. However, although the clinical efficacy of second generation non-sedating oral antihistamines is well documented, topical therapy may be preferable due to the reduced potential for systemic adverse effects.

Topical therapy available for the treatment of allergic conjunctivitis includes corticosteroids, sodium cromoglycate and vasoconstrictors, either administered alone or in combination with an H₁-receptor antagonist. Intranasal corticosteroids provide effective relief from nasal symptoms and are generally well tolerated. However, ocular steroid administration should generally be avoided due to an associated risk of glaucoma, cataracts and opportunistic infections. Sodium cromoglycate is a well tolerated agent for the prophylaxis of allergic rhinoconjunctivitis and is suitable for both intranasal and ocular administration. The therapeutic efficacy of this drug is highly variable and maintenance therapy throughout the period of allergen exposure is essential. Topical vasoconstrictors provide effective relief only for nasal obstructive symptoms and are only suitable for short-term therapy due to the potential for rebound vasodilatation during prolonged use.

Until recently, topical antihistamines have not been sufficiently potent to enable single agent therapy, necessitating co-administration of a vasoconstrictor. While reports suggest that such combinations may be more effective than vasoconstrictor monotherapy, they are not suitable for long-term use due to the adverse reactions associated with topical vasoconstrictors.

Levocabastine

Levocabastine is an extremely potent and highly selective H₁-receptor antagonist which has been specifically developed for topical ocular and nasal administration. Levocabastine is the most potent antihistamine available to date expressing antihistaminic activity at doses lower than 0.002 mg/kg, making it some 15 000 times more potent than chlorpheniramine. In addition, animal studies have shown that this agent has a very high specificity for H₁-receptors being devoid of anticholinergic, antiserotonergic or antidopaminergic activity at concentrations
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considerably greater than effective antihistaminic doses. This suggests that adverse reactions resulting from a lack of receptor specificity are unlikely. Levocabastine is a potent inhibitor of experimentally induced rhinoconjunctivitis providing effective relief from symptoms within minutes of instillation,\textsuperscript{8-13} with comparative clinical trials demonstrating that this agent is at least as effective as oral antihistamines in patients with clinical disease.\textsuperscript{14-17}

Pharmacokinetic analysis suggests that the potential for systemic adverse effects is very low. Levocabastine is incompletely absorbed after intranasal and ocular administration with systemic availability typically ranging from 30–60% for the eye drops and 60–80% for the nasal spray.\textsuperscript{18} Moreover, as the amount of drug applied topically is small, the plasma concentration of levocabastine attained via these routes of administration is extremely low with \( C_{\text{max}} \) values in the range of 0.2–0.29 mg/l for levocabastine eye drops and 1.4–2.2 mg/l for the nasal spray. Detailed pharmacokinetic–pharmacodynamic modelling confirms those findings, indicating that the clinical benefits seen with this agent are predominantly due to local antihistaminic effects at the site of application.\textsuperscript{19} Coupled with the fact that levocabastine is subject to minimal hepatic metabolism, this suggests that the potential for systemic adverse reactions is minimal.

The present review aims to summarize the data available to date concerning the tolerability and safety of levocabastine eye drops and nasal spray with particular reference to implications for the clinical management of patients with allergic rhinoconjunctivitis.

Tolerability

The tolerability of levocabastine eye drops and nasal spray has been extensively evaluated in clinical trials. The available data suggests that topical levocabastine is well tolerated with an adverse effect profile comparable with that of sodium cromoglycate or placebo. Analysis of data from a total of 1628 patients treated with levocabastine eye drops reveals that local irritation is the most frequently reported adverse event following ocular administration with an incidence of 14%. Of 1758 patients who have received levocabastine nasal spray, headache (4%), nasal irritation (3%), somnolence (3%) and fatigue (2%) have been reported. The overall incidence of adverse experiences on levocabastine eye drops

![Nasal spray diagram]

![Eye drops diagram]

FIG. 1. Incidence of adverse experiences with levocabastine nasal spray and eye drops compared with sodium cromoglycate and placebo.
is 24%. In separate studies the incidence has been comparable for levocabastine and placebo (27 versus 31%), levocabastine and cromoglycate (36 versus 31%) and levocabastine and terfenadine (40 versus 41%). Figure 1 shows the tolerability of levocabastine compared with sodium cromoglycate and placebo.

In addition, no clinically significant changes in biochemical or haematological parameters have been reported during treatment with this topical antihistamine in clinical trials that have performed routine laboratory tests to date. Significantly, the type and frequency of adverse experiences appears to be unrelated to the number of daily applications. Moreover, adverse effects do not appear to be increased by concomitant use of the eye drops and nasal spray compared with use of either formulation alone.

Drug tolerability is a key factor determining choice of therapy in children. Importantly, a study involving 53 children aged between 6 and 15 years reported that levocabastine nasal spray and eye drops are well tolerated in this patient population with type and incidence of adverse reactions similar to those occurring in sodium-cromoglycate-treated children.

Local tolerance: It is well documented that intranasal administration of certain drugs and additives, in particular decongestants, can influence ciliary activity of the upper airways. The influence of levocabastine on ciliary beat frequency and mucociliary clearance has been evaluated in vitro using human adenoid tissue and in vivo following single- and multiple-dose administration in healthy human volunteers. Levocabastine was seen to have only a small, clinically insignificant effect on ciliary beat frequency in human adenoid tissue. No significant effects on ciliary function were observed following single- and multiple-dose administration in healthy volunteers.

Ocular tolerance of levocabastine has been evaluated extensively. Animal studies have failed to reveal any adverse effects resulting from topical administration of levocabastine eye drops. In an eye irritation study in rabbits, application of levocabastine eye drops four to eight times daily for 4 weeks did not produce ocular changes, and in another study, normal intraocular pressure was found to be maintained in rabbits following application of levocabastine. Levocabastine did not induce dermal sensitization when tested in guinea-pigs using the Magnusson maximization technique.

The local tolerance of levocabastine eye drops has also been extensively evaluated in volunteers and atopic patients during clinical trials using a wide range of objective ophthalmological tests including slit lamp evaluation, tonometry, fluorescein and rose bengal staining, ophthalmoscopy and determination of intraocular pressure, visual acuity and visual fields. These studies have failed to reveal any significant or consistent adverse effects attributable to the topical antihistamine at doses of up to 4 times daily and treatment durations of up to 8 weeks.

As many traditional antihistamines have been associated with anticholinergic activity, which may be expected to influence accommodative capacity, the effects of ocular administration of levocabastine on accommodation have been investigated in healthy volunteers and in patients with glaucoma. Administration of a single dose of levocabastine (0.5 mg/ml) was found to have no effect on accommodative capacity in healthy volunteers. Similarly, accommodative capacity and intraocular pressure were unchanged in patients with glaucoma during twice daily treatment with ocular levocabastine over a period of 2 weeks. These findings are in keeping with the highly selective pharmacological profile of the drug.

Psychomotor effects of levocabastine: Many traditional H1-receptor antagonists cross the blood-brain barrier producing unwanted central nervous system effects. Increased daytime somnolence and reduced alertness, are frequent adverse reactions particularly encountered with many first-generation oral antihistamines.

Pharmacokinetic studies indicate that sedative effects would be extremely unlikely in patients receiving intranasal or ocular levocabastine due to the low plasma concentrations of the drug obtained by these routes of administration. Specific studies of psychomotor and cognitive function following topical administration of levocabastine support these findings. Objective assessment of psychomotor performance and subjective assessment of sedation have failed to reveal any central nervous system effects in healthy volunteers treated with levocabastine eye drops (0.5 mg/ml, two drops in each eye, four times daily) over a period of 1 week. Potential central nervous system effects of levocabastine have also been investigated following single- and multiple-dose administration of levocabastine eye drops and nasal spray, and compared with those of oral triprolidine. Performance was assessed following medication by a battery of cognitive and psychomotor tests considered as reliable measures of the sedative effects of psychoactive drugs. In contrast to the significant sedative effects of triprolidine, topical administration of levocabastine eye drops and nasal spray at doses up to 2.0 mg/ml, a dose level four times greater
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than normal therapeutic dosages, had no demonstrable effect on central nervous system function in healthy volunteers.25

Potential pharmacokinetic and psychomotor interactions between levocabastine nasal spray and alcohol or diazepam have also been investigated in healthy volunteers. No evidence of significant psychomotor interactions between levocabastine and either ethanol or diazepam are apparent.

Cardiovascular effects of levocabastine: Possible cardiovascular effects of levocabastine have been assessed in vitro on isolated Purkinje fibres of dogs and in vivo in animal studies and in human volunteers following oral, ocular and nasal administration. In vitro studies on isolated Purkinje fibres revealed no effect of levocabastine on action potential amplitude and action potential duration at 50% and 90% repolarization at doses of up to 2.5 mg/l. Similarly, oral administration of doses of up to 0.16 mg/kg have revealed no significant effect on key cardiovascular parameters in dogs.

More relevantly, levocabastine does not appear to produce significant ECG effects in man. Several studies in healthy volunteers (Fig. 2) have revealed no significant effects on QTc intervals after treatment with levocabastine in single or repeated doses, even when the eye drops and the nasal spray are used in combination four times daily (1.2 mg/day) or after administration of oral levocabastine 3 mg daily (for 3 days).27

Drug interactions: As levocabastine plasma concentrations are extremely low following topical administration and hepatic metabolism is negligible, clinically significant drug interactions are unlikely. But since binding studies have revealed that levocabastine binds to plasma proteins, particularly albumin, the potential for drug interactions involving binding site displacement exists. However, in vitro analysis of potential pharmacokinetic drug interactions have identified levocabastine does not alter the plasma protein binding of imipramine, propranolol, diphenylhydantoin, diazepam, cimetidine, indomethacin or ketoconazole although slight increases in the proportion of unbound levocabastine can be identified with drugs which are highly protein-bound such as sulphamethazine (4.5%), tolbutamide (6%) and warfarin (8.1%). Although such changes are minor they should be recognized. Obviously this interaction is of no clinical relevance for levocabastine as its plasma protein binding is only 55%.18

Conclusion

Extensive clinical investigations have shown levocabastine eye drops and nasal spray to be highly effective for the treatment of allergic rhinoconjunctivitis. Levocabastine eye drops and nasal spray are well tolerated with an adverse-effect profile comparable with that of sodium cromoglycate and placebo. Local irritation following administration is the most common adverse reaction reported in levocabastine-treated patients to date with an incidence similar to that seen in placebo- or cromoglycate-treated controls. Importantly, the potential for systemic
adverse effects is negligible. As would be expec-
ted from its pharmacological profile, levocabas-
tine is devoid of significant systemic activity
having no apparent effects on cardiovascular
function, psychomotor performance, cognitive
function and minimal interaction potential.
Coupled with its well documented therapeutic
efficacy, these findings indicate that topical levo-
cabastine is an attractive therapeutic option for
the treatment of allergic conjunctivitis.

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