

PRODUCT MONOGRAPH
INCLUDING PATIENT MEDICATION INFORMATION

^{Pr}**SIMBRINZA**®

Brinzolamide/Brimonidine Tartrate Ophthalmic Suspension
Suspension, 1%/0.2% w/v, ophthalmic

Elevated Intraocular Pressure Therapy
Topical Carbonic Anhydrase Inhibitor and Topical Alpha-2 Agonist Agent
ATC code S01EC54

Novartis Pharmaceuticals Canada Inc.
700 Saint-Hubert St., Suite 100
Montreal, Quebec H2Y 0C1
www.novartis.ca

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RECENT MAJOR LABEL CHANGES

7 WARNINGS AND PRECAUTIONS; Hypersensitivity; Renal; Skin

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PART I: HEALTH PROFESSIONAL INFORMATION

1 INDICATIONS

SIMBRINZA (brinzolamide / brimonidine tartrate ophthalmic suspension) is indicated for:

- the reduction of intraocular pressure (IOP) in adult patients with open-angle glaucoma or ocular hypertension for whom monotherapy provides insufficient IOP reduction AND when the use of SIMBRINZA is considered appropriate.

1.1 Pediatrics

Paediatrics (< 18 years of age):

The safety and efficacy of SIMBRINZA has not been established in pediatric patients; therefore, Health Canada has not authorized an indication for pediatric use. Several serious adverse reactions have been reported in association with the administration of brimonidine tartrate ophthalmic solution 0.2% in pediatric populations. (see [2 CONTRAINDICATIONS](#) and [7.1.3, Pediatrics](#)).

1.2 Geriatrics

Geriatrics (> 65 years of age): No overall differences in safety and effectiveness have been observed between elderly and other adult patients.

2 CONTRAINDICATIONS

Brinzolamide / Brimonidine Tartrate ophthalmic suspension is contraindicated in:

- Patients with hypersensitivity to brinzolamide, brimonidine, or to any ingredient in the formulation or component of the container. For a complete listing, see [6 DOSAGE FORMS, STRENGTHS, COMPOSITION AND PACKAGING](#).
- Patients with hypersensitivity to sulfonamides.
- Patients receiving monoamine oxidase (MAO) inhibitor therapy.
- Patients on antidepressants that affect noradrenergic transmission (e.g. tricyclic antidepressants and mianserin).
- Patients with severe renal impairment.
- Patients with hyperchloraemic acidosis.
- Neonates and infants under the age of 2 years.

SIMBRINZA is not recommended in patients with hepatic impairment. SIMBRINZA has not been studied in patients with hepatic impairment.

SIMBRINZA is contraindicated in patients with severe renal impairment ($\text{CrCl} < 30 \text{ mL/min}$). SIMBRINZA has not been studied in patients with severe renal impairment ($\text{CrCl} < 30 \text{ mL/min}$) or in patients with hyperchloraemic acidosis. Since the brinzolamide component of SIMBRINZA and its metabolite are excreted predominantly by the kidney, SIMBRINZA is contraindicated in such patients.

4 DOSAGE AND ADMINISTRATION

4.1 Dosing Considerations

- SIMBRINZA is a carbonic anhydrase inhibitor formulated for topical ophthalmic use. With any evident signs of hypersensitivity or discomfort, SIMBRINZA must be discontinued.
- Caution is advised when using SIMBRINZA in patients with mild to moderate renal impairment. (see [7 WARNING AND PRECAUTIONS](#))

4.2 Recommended Dose and Dosage Adjustment

The recommended dose is one drop of SIMBRINZA in the affected eye(s) two times daily.

Health Canada has not authorized an indication for pediatric use. (see [1.1 Pediatrics](#) and [7.1.3 Pediatrics](#).)

4.4 Administration

Patients should be instructed to shake the bottle well before use.

Nasolacrimal occlusion or gently closing the eyelid after instillation for up to one minute is recommended. This may reduce the systemic absorption of medications administered via the ocular route and result in a decrease in systemic adverse events.

SIMBRINZA may be used concomitantly with other topical ophthalmic drug products to lower intraocular pressure. If more than one topical ophthalmic drug is being used, the drugs should be administered at least five minutes apart.

Do not allow the dropper tip of the bottle to touch the eye or other surrounding structures, because this could cause eye injury or contaminate the tip with common bacteria known to cause eye infections. Serious damage to the eye with subsequent loss of vision may result from using contaminated eye drop solutions. Do not use suspension if the bottle is cracked or damaged in any way.

4.5 Missed Dose

If a dose is missed, treatment should be continued with the next dose as planned. The dose should not exceed one drop in the affected eye(s) two times daily.

5 OVERDOSAGE

There were no cases of overdose reported in the clinical development program for SIMBRINZA.

Although no human data are available, symptoms of brinzolamide overdose may include hypotension, electrolyte imbalance, development of an acidotic state, and possible nervous system effects may occur. Serum electrolyte levels (particularly potassium) and blood pH levels must be monitored.

Symptoms of brimonidine overdose such as apnea, bradycardia, coma, hypotension, hypothermia, hypotonia, lethargy, pallor, respiratory depression, and somnolence have been reported in neonates, infants, and children receiving brimonidine as part of medical treatment of congenital glaucoma or by accidental oral ingestion. Brimonidine toxicity resembles that of clonidine, which has been reported to cause fatigue, apnea, bradycardia, hypothermia, dizziness, hypotension and somnolence.

Oral overdoses of other alpha-2-agonists have been reported to cause symptoms such as hypotension, asthenia, vomiting, lethargy, sedation, bradycardia, arrhythmias, miosis, apnoea, hypotonia, hypothermia, respiratory depression and seizure.

Treatment of oral overdose includes supportive and symptomatic therapy; patient's airway should be maintained.

For management of a suspected drug overdose, contact your regional poison control centre.

6 DOSAGE FORMS, STRENGTHS, COMPOSITION AND PACKAGING

Table 1 – Dosage Forms, Strengths, Composition and Packaging

Route of Administration	Dosage Form / Strength/Composition	Non-medicinal Ingredients
Ophthalmic (topical)	Suspension/ brinzolamide 1% w/v and brimonidine tartrate 0.2% w/v	Benzalkonium chloride (as a preservative), Boric Acid, Carbomer 974P, Hydrochloric acid (to adjust pH), Mannitol, Propylene glycol, purified water, Sodium chloride, Sodium hydroxide (to adjust pH), Tyloxapol.

Description

SIMBRINZA is a white to off-white uniform suspension containing 10 mg/mL brinzolamide and 2 mg/mL brimonidine tartrate.

SIMBRINZA is supplied in a round, white, low-density polyethylene (LDPE) bottles with a natural LDPE dispensing plug and white polypropylene screw cap containing 10 mL of suspension.

7 WARNINGS AND PRECAUTIONS

General

FOR TOPICAL OPHTHALMIC USE ONLY.

Like other topically applied ophthalmic agents, brinzolamide and brimonidine tartrate, the active ingredients of SIMBRINZA, are absorbed systemically. However, brinzolamide significantly accumulates in the blood due to extremely long half-life (see [10. CLINICAL PHARMACOLOGY](#)).

Acid-base disturbances have been reported with oral carbonic anhydrase inhibitors.

Carcinogenicity and Mutagenesis

See [16 NON-CLINICAL TOXICOLOGY, Carcinogenicity and Genotoxicity](#).

Cardiovascular

Alpha 2 agonists as a class may reduce pulse and blood pressure. In clinical trials SIMBRINZA is associated with mild reduction in blood pressure in some patients. Caution is advised when using SIMBRINZA concomitantly with antihypertensives and/or cardiac glycosides or in patients with severe or unstable and uncontrolled cardiovascular disease.

Brimonidine tartrate, a component of SIMBRINZA, may potentiate syndromes associated with vascular insufficiency, therefore, caution is advised when using SIMBRINZA in patients with depression (especially those taking tricyclic antidepressants as these agents may blunt the hypotensive response), cerebral or coronary insufficiency, Raynaud's phenomenon, orthostatic hypotension or thromboangiitis obliterans.

Driving and Operating Machinery

SIMBRINZA can cause fatigue and/or drowsiness in some patients. Patients who engage in hazardous activities should be cautioned about the potential for a decrease in mental alertness.

SIMBRINZA may also cause blurred vision or visual disturbance in some patients. The patient should wait until these symptoms have cleared before driving or using machinery.

Endocrine and Metabolism

SIMBRINZA significantly increases blood glucose level and decreases blood cholesterol level when topically administered twice daily (BID) in the eye of rabbits. These reactions have been observed with rabbits topically dosed with brimonidine alone in the eyes (see 16 NON-CLINICAL TOXICOLOGY).

Hepatic/Biliary/Pancreatic

Brinzolamide and brimonidine tartrate, components of the active ingredients of SIMBRINZA, have not been studied in patients with hepatic impairment. Use of SIMBRINZA is not recommended in patients with hepatic impairment.

Hypersensitivity

SIMBRINZA contains brinzolamide which is a sulfonamide, and although administered topically is absorbed systemically. Therefore, the same types of adverse reactions that are attributable to sulfonamides may occur with the topical administration of SIMBRINZA. Fatalities have occurred due to severe reactions to sulfonamides including Stevens Johnson Syndrome (SJS), Toxic Epidermal Necrolysis (TEN), fulminant hepatic necrosis, agranulocytosis, aplastic anemia, and other blood dyscrasias (see 8 ADVERSE REACTIONS section). Rechallenge irrespective of the route of administration should not be undertaken in patients with hypersensitivity syndrome and SJS/TEN (see 7 Warning and Precautions, Skin).

Neurologic

SIMBRINZA may cause fatigue and somnolence. Carbonic anhydrase inhibitors can impair ability to perform tasks requiring mental alertness and/or physical coordination. As SIMBRINZA is absorbed systemically, caution is advised when using SIMBRINZA in patients requiring mental alertness and/or physical coordination.

Ophthalmologic

SIMBRINZA is not recommended in patients with acute or narrow-angle glaucoma. SIMBRINZA has not been studied in patients with acute or narrow-angle glaucoma.

Carbonic anhydrase activity has been observed in both the cytoplasm and around the plasma membranes of the corneal epithelium. There is an increased potential for developing corneal edema in patients with low endothelial cell counts.

The possible role of brinzolamide on corneal endothelial function has not been investigated in patients with compromised corneas (particularly in patients with low endothelial cell count). Specifically, patients

wearing contact lenses have not been studied and careful monitoring of these patients when using brinzolamide is recommended, since carbonic anhydrase inhibitors may affect corneal hydration and wearing contact lenses might increase the risk for the cornea. Careful monitoring of patients with compromised corneas, such as patients with diabetes mellitus or corneal dystrophies, is recommended.

SIMBRINZA contains benzalkonium chloride as preservative, which may cause eye irritation and is known to discolour soft contact lenses. Contact with soft contact lenses is to be avoided. Patients must be instructed to remove contact lenses prior to the instillation of SIMBRINZA and wait at least 15 minutes after dosing before contact lenses are reinserted.

Benzalkonium chloride has also been reported to cause punctate keratopathy and/or toxic ulcerative keratopathy. Close monitoring is required with frequent or prolonged use.

SIMBRINZA may cause temporary blurred vision or other visual disturbances that can affect the ability to drive or use machines. If blurred vision occurs at instillation, the patient must wait until the vision clears before driving or using machinery.

Animal studies show that brinzolamide and brimonidine significantly accumulates in iris-ciliary body, choroid and/or retina during BID topical ocular administration of SIMBRINZA (see 16 NON-CLINICAL TOXICOLOGY, Animal Pharmacokinetics).

Brimonidine tartrate may cause ocular allergic reactions. If allergic reactions are observed, treatment should be discontinued.

Psychiatric

Caution is advised with using SIMBRINZA in patients with depression.

Renal:

SIMBRINZA is contraindicated in patients with severe renal impairment. Caution is advised when using SIMBRINZA in patients with mild to moderate renal impairment because of the possible risk of metabolic acidosis.

Reproductive Health: Female and Male Potential

- **Fertility**

The effect of SIMBRINZA on human fertility is unknown. Nonclinical data do not suggest an effect of brinzolamide or brimonidine on fertility. In animals, developmental toxicity was observed with brinzolamide at doses that induced maternal toxicity. (see 16 NON-CLINICAL TOXICOLOGY, Reproductive and Developmental Toxicology).

Skin

SIMBRINZA should be discontinued immediately at the appearance of a skin rash, as the rash may be, in some instances, followed by dermatological reactions/hypersensitivity syndrome including SJS and TEN. At the time of prescription, patients should be informed of the signs and symptoms, and advised to monitor closely for skin reactions.

7.1 Special Populations

7.1.1 Pregnant Women

SIMBRINZA is not recommended during pregnancy or in women of child-bearing potential not using contraception.

Developmental toxicity studies with brinzolamide in rabbits at oral doses of 1, 3, and 6 mg/kg/day (43, 129, and 258 times the recommended human ophthalmic dose) produced maternal toxicity at 6 mg/kg/day and a significant increase in the number of fetal variations, such as accessory skull bones, which was only slightly higher than the historic value at 1 and 6 mg/kg. In rats, statistically decreased body weights of fetuses from dams receiving oral doses of 18 mg/kg/day (783 times the recommended human ophthalmic dose) during gestation were proportional to the reduced maternal weight gain, with no statistically significant effects on organ or tissue development. Increases in unossified sternebrae, reduced ossification of the skull, and unossified hyoid that occurred at 6 and 18 mg/kg were not statistically significant. No treatment-related malformations were seen. Following oral administration of ¹⁴C-brinzolamide to pregnant rats, radioactivity was found to cross the placenta and was present in the fetal tissues and blood.

In animal studies, brimonidine crossed the placenta and entered into the fetal circulation to a limited extent. Drug derived material was eliminated from fetal tissues by 24 hours post dose.

(See 16 NON-CLINICAL TOXICOLOGY, Reproductive and Developmental Toxicology).

7.1.2 Breast-feeding

SIMBRINZA should not be used by women nursing neonates/infants.

It is not known whether topical SIMBRINZA is excreted in human milk; however, a risk to the nursing child cannot be excluded.

Available pharmacodynamic/toxicological data in animals have shown that following oral administration, brinzolamide and brimonidine are excreted in breast milk.

7.1.3 Pediatrics

Pediatrics (< 18 years of age): SIMBRINZA is contraindicated in neonates and infants under the age of 2 years. Several serious adverse reactions have been reported in association with the administration of brimonidine tartrate to infants. (see 2. CONTRAINDICATIONS)

SIMBRINZA is not recommended in children between 2 and 18 years.

7.1.4 Geriatrics

Geriatrics (> 65 years of age): No overall differences in safety or effectiveness of SIMBRINZA have been observed between elderly and adult patients.

8 ADVERSE REACTIONS

8.1 Adverse Drug Reaction Overview

The individual active components SIMBRINZA are currently marketed as AZOPT (brinzolamide 10 mg/mL) and ALPHAGAN (brimonidine tartrate 2 mg/mL). These individual component products have well-characterized safety profiles obtained via clinical trials and post-marketing surveillance.

Adverse drug reactions (ADRs) observed with the use of SIMBRINZA in clinical trials were consistent with ADRs that have been observed with the use of one or both of the individual components. The most frequent ocular ADRs observed with the use of SIMBRINZA were nonserious local ocular side effects (e.g., hyperemia of the eye, ocular allergic type reactions, blurred vision, and ocular discomfort). Common systemic ADRs reported with its use were nonserious and included dysgeusia (bad taste), oral dryness and fatigue/drowsiness.

The majority of ADRs leading to patient discontinuation from study participation were for nonserious local ocular side effects (e.g., ocular discomfort, ocular hyperemia, and ocular allergic type reactions). The incidence of patients discontinuing due to these events was similar between SIMBRINZA and concomitant dosing with the individual components (brinzolamide 10 > mg/mL+ brimonidine tartrate 2mg/mL).

Overall, the safety profile of SIMBRINZA was similar to that of the individual components (brinzolamide 10 mg/mL and brimonidine 2 mg/mL) and did not result in additional risk to patients relative to the known risks of the individual components.

8.2 Clinical Trial Adverse Reactions

Clinical trials are conducted under very specific conditions. Therefore, the adverse reaction rates observed in the clinical trials, may not reflect the rates observed in practice and should not be compared to the rates in the clinical trials of another drug. Adverse reaction information from clinical trials may be useful in identifying and approximating rates of adverse drug reactions in real-world use.

The evaluation of the safety of a fixed combination brinzolamide/ brimonidine dosed twice daily (BID) included 2 phase III clinical studies of 6 months duration. During these clinical studies 645 patients were exposed to SIMBRINZA.

The Table below summarizes adverse events assessed by the examining physician as related to the use of SIMBRINZA e.g., ADR reported at an incidence of $\geq 1\%$ in clinical trials (data pooled) along with the corresponding incidence of the ADRs reported for the comparator groups.

Table 2 - Adverse Drug Reactions $\geq 1\%$ Reported With the use of SIMBRINZA (Brinz/Brim BID) and the Corresponding Incidences Observed in the Comparator Groups C 10 040/C 10 041

Coded Adverse Event	Brinz/Brim BID N = 645		Brinz+Brim BID N = 436		Brinz BID N = 192		Brim BID N = 175	
	N	%	N	%	N	%	N	%
Eye Disorders								
Ocular hyperemia	27	4.2	17	3.9	1	0.5	8	4.6
Vision blurred	18	2.8	13	3.0	1	0.5	2	1.1
Eye pain	18	2.8	8	1.8	3	1.6	-	-
Conjunctivitis allergic	17	2.6	9	2.1	-	-	3	1.7
Eye pruritus	14	2.2	8	1.8	3	1.6	4	2.3
Eye irritation	14	2.2	7	1.6	4	2.1	3	1.7
Conjunctivitis	12	1.9	5	1.1	-	-	1	0.6
Conjunctival hyperaemia	10	1.6	13	3.0	3	1.6	4	2.3
Eye allergy	8	1.2	6	1.4	-	-	2	1.1
Gastrointestinal Disorders								
Dry mouth	18	2.8	14	3.2	2	1.0	9	5.1
Nervous System Disorders								
Dysgeusia	22	3.4	16	3.7	4	2.1	2	1.1
Somnolence	14	2.2	15	3.4	-	-	4	2.3

8.3 Less Common Clinical Trial Adverse Reactions

The list below summarizes ADRs reported for the fixed combination brinzolamide/brimonidine as well as the individual components dosed either concomitantly or as monotherapy at an incidence of $< 1\%$ in clinical trials [C-10-041] and [C-10-040] (data pooled). Adverse drug reactions are presented in alphabetical order within the System Organ Classification.

Ear and labyrinth disorders: vertigo.

Eye disorders: abnormal sensation in eye, asthenopia, blepharitis allergic, blepharitis, cataract cortical, conjunctival disorder, conjunctival follicles, conjunctival irritation, conjunctival oedema, corneal deposits, corneal erosion, corneal oedema, dry eye, ectropion, eczema eyelids, erythema of eyelid, eye discharge, eyelid margin crusting, eyelid oedema, foreign body sensation in eyes, hypermetropia, increased lacrimation, keratitis, keratoconjunctivitis sicca, meibomianitis, ocular discomfort, photophobia, punctate keratitis, visual acuity reduced, visual impairment.

Gastrointestinal disorders: abdominal discomfort, dyspepsia, hypoaesthesia oral, oral discomfort, paraesthesia oral.

General disorders and administration site conditions: asthenia, drug intolerance, exercise tolerance decreased, fatigue, medication residue, mucosal dryness.

Immune system disorders: drug hypersensitivity.

Injury, poisoning and procedural complications: excoriation.

Investigations: blood pressure decreased, corneal staining.

Nervous system disorders: balance disorder, dizziness, headache, hypersomnia, hypotonia, sedation.

Psychiatric disorders: insomnia, libido increased.

Respiratory, thoracic and mediastinal disorders: dry throat, nasal dryness, nasal congestion, postnasal drip, rhinalgia, rhinorrhoea.

Skin and subcutaneous tissue disorders: blister, dermatitis, dermatitis allergic, dermatitis contact.

Vascular disorders: hypertension, hypotension.

SIMBRINZA contains the individual active components brinzolamide and brimonidine tartrate. Additional adverse events associated with the use of the individual components that may potentially occur with SIMBRINZA include:

Brinzolamide:

Adverse reactions associated with brinzolamide include gastrointestinal, nervous system, haematological, renal and metabolic effects. The same type of adverse reactions attributable to oral carbonic anhydrase inhibitors may occur with topical administration. Fatalities have occurred due to severe reactions to sulphonamides, including Stevens-Johnson syndrome, toxic epidermal necrolysis, fulminant hepatic necrosis, agranulocytosis, aplastic anemia, and other blood dyscrasias. Sensitization may recur when a sulphonamide is re-administered irrespective of the route of administration.

Brimonidine tartrate:

Adverse reactions associated with the brimonidine tartrate include the development of ocular allergic type reactions, fatigue and/or drowsiness, and dry mouth. The use of brimonidine has been associated with minimal decreases in blood pressure.

Adverse reactions associated with the individual components of SIMBRINZA are presented within the Product Monographs for brinzolamide and brimonidine tartrate, respectively.

Abnormal Hematologic and Clinical Chemistry Findings

Clinical laboratory evaluations for the analysis of safety were performed during the pharmacokinetic clinical trial (C-10-010) involving healthy volunteers who were dosed with 1 mg of oral brinzolamide BID for 2 weeks prior to dosing topically with SIMBRINZA, dosed BID or three times a day (TID), for 13 weeks. No clinically relevant change in hematology, blood chemistry, or urinalysis parameter was reported with the use of SIMBRINZA, dosed BID or TID.

8.5 Post-Market Adverse Reactions

The following adverse reactions have been identified during post marketing use of SIMBRINZA in clinical practice. As these events are reported voluntarily from a population of unknown size, estimates of frequency cannot be made:

Blurred vision,

Dizziness,

Dry mouth,

Eye irritation, eye pain, eye pruritus,

Fatigue, foreign body sensation in eyes,

Ocular hyperaemia, and somnolence.

Overall, these events are consistent with clinical trial experience with SIMBRINZA and/or with the individual components (i.e., brinzolamide 10 mg/ml and brimonidine tartrate 2 mg/ml).

The following reactions have been identified during post marketing use of AZOPT® (brinzolamide 1%). Because these events are reported voluntarily from a population of unknown size, estimates of frequency cannot be made:

-Ageusia, arrhythmia, arthralgia, asthma, blood pressure increased, chest pain, increased heart rate
corneal disorder, dermatitis, erythema, erythema of eyelid, eye allergy, eyelid disorder,
hypersensitivity, hypertension, hypoaesthesia, madarosis, malaise, medication residue,
pain in extremity, peripheral oedema, pollakiuria,
rhinitis, tachycardia, tremor, vertigo, and visual disturbance.

-Skin and subcutaneous tissue disorders: Stevens-Johnson syndrome (SJS), Toxic epidermal necrolysis (TEN), alopecia, erythema, rash

The following reactions have been identified during post marketing use of brimonidine tartrate ophthalmic solutions in clinical practice. Because these events are reported voluntarily from a population of unknown size, estimates of frequency cannot be made:

Hypersensitivity, iritis, iridocyclitis (anterior uveitis), conjunctivitis, miosis

-Syncope and hypotension

-Vision blurred, conjunctivitis, fatigue, dizziness, hypersensitivity and somnolence have also been reported in infants receiving brimonidine tartrate ophthalmic solutions.

9 DRUG INTERACTIONS

9.2 Drug Interactions Overview

No specific drug interaction studies have been performed with SIMBRINZA.

9.3 Drug-Behavioural Interactions

Not applicable.

9.4 Drug-Drug Interactions

SIMBRINZA is contraindicated in patients receiving monoamine oxidase inhibitors (which can affect the metabolism and uptake of circulating amines) and patients on antidepressants that affect noradrenergic transmission (e.g., tricyclic antidepressants and mianserin) (see [2 CONTRAINDICATIONS](#)). Tricyclic antidepressants have been reported to blunt the hypotensive effect of systemic clonidine. It is unknown whether the concurrent use of these agents with SIMBRINZA in humans can lead to resultant interference with the IOP lowering effect of the product. Such drugs may theoretically interfere with the metabolism of brimonidine tartrate and potentially result in increased systemic side effects, such as hypotension.

The possibility of an additive or potentiating effect with CNS depressants (i.e., alcohol, barbiturates, opiates, sedatives, or anaesthetics) should be considered.

No data are available on the level of circulating catecholamines after SIMBRINZA administration; however, caution is advised in patients taking medication that can affect the metabolism and uptake of circulating amines (e.g., chlorpromazine, methylphenidate, and reserpine).

Alpha adrenergic agonists (e.g., brimonidine tartrate), as a class, may reduce pulse and blood pressure. Following administration of SIMBRINZA, small decreases in blood pressure were observed in some patients. Caution is advised when using drugs such as antihypertensives and/or cardiac glycosides concomitantly with SIMBRINZA.

Caution is advised when using a concomitant systemic agent (irrespective of pharmaceutical form) that may interact with alpha adrenergic agonists or interfere with their activity (e.g. isoprenaline, prazosin).

Brinzolamide, a component of SIMBRINZA, is a carbonic anhydrase inhibitor and, although administered topically, is absorbed systemically. Acid-base disturbances have been reported with oral carbonic anhydrase inhibitors. The potential for interactions must be considered in patients receiving SIMBRINZA. There is a potential for an additive effect on the known systemic effects of carbonic anhydrase inhibition in patients receiving an oral carbonic anhydrase inhibitor and topical brinzolamide. The concomitant administration of SIMBRINZA and oral carbonic anhydrase inhibitors is not recommended.

Concomitant use of salicylates (e.g., acetylsalicylic acid) with SIMBRINZA is not recommended, especially with high dose therapy (>1 g daily) as this may lead to decreased efficacy of the salicylate, CNS toxicity, metabolic acidosis, and other adverse reactions. These alterations were not observed in clinical trials with brinzolamide ophthalmic suspension 1%; however, in patients treated with oral carbonic anhydrase inhibitors, rare instances of acid–base alterations have occurred with high dose salicylate therapy.

The cytochrome P-450 isozymes responsible for metabolism of brinzolamide include CYP3A4 (main), CYP2A6, CYP2B6, CYP2C8 and CYP2C9. It is expected that inhibitors of CYP3A4 such as ketoconazole, itraconazole, clotrimazole, ritonavir and troleandomycin will inhibit the metabolism of brinzolamide by CYP3A4. Caution is advised if CYP3A4 inhibitors are given concomitantly. Brinzolamide is not an inhibitor of cytochrome P-450 isozymes.

9.5 Drug-Food Interactions

Interactions with food have not been established.

9.6 Drug-Herb Interactions

Interactions with herbal products have not been established.

9.7 Drug-Laboratory Interactions

Interactions with laboratory tests have not been established.

10 CLINICAL PHARMACOLOGY

10.1 Mechanism of Action

SIMBRINZA contains two active substances: brinzolamide and brimonidine tartrate. Although both brinzolamide and brimonidine lower IOP by suppressing aqueous humour formation, their mechanisms of action are different.

Brinzolamide acts by inhibiting the enzyme carbonic anhydrase (CA-II) in the ciliary epithelium, which reduces the formation of bicarbonate ions and results in a subsequent reduction in sodium and fluid transport across the ciliary epithelium, resulting in decreased aqueous humour formation.

Brimonidine tartrate, an alpha-2 adrenergic receptor agonist, inhibits the enzyme adenylate cyclase and suppresses the cAMP-dependent formation of aqueous humour. Additionally, administration of brimonidine results in an increase in uveoscleral outflow.

10.2 Pharmacodynamics

Not Available.

10.3 Pharmacokinetics

Brinzolamide pharmacokinetics are inherently non-linear due to saturable binding to carbonic anhydrase in whole blood and various tissues. Steady-state exposure does not increase in a dose-proportional manner. Brimonidine pharmacokinetics are linear and systemic exposure increases with increasing dose in an approximately proportional manner.

The pharmacokinetics and disposition of brinzolamide and brimonidine in humans have been characterized and are reported in the Product Monographs for AZOPT® (brinzolamide 10 mg/mL eye drops, suspension) and ALPHAGAN® (brimonidine tartrate 2 mg/mL eye drops, solution).

Table 3: Steady-state [trough] red blood cell and plasma concentrations of brinzolamide and N-desethyl brinzolamide following administration of brinzolamide 1%/brimonidine tartrate 0.2% (BID) in healthy subjects [Day 107]

	Red Blood Cells C_{ss} (μ M)	Plasma C_{ss} (ng/mL)
Brinzolamide	15.2 (3.49, 24.5)	2.52 (1.19, 6.67)
N-desethyl-brinzolamide	1.81 (0.592, 4.89)	ND

C_{ss} and $AUC_{15-107days}$ data are presented as geometric means. Values in parentheses are (minimum, maximum) values for respective parameter. ND – Not detected (below 1 ng/mL limit of detection, if present). Steady-state RBC AUC not calculated since AUC would be directly proportional to C_{ss} due to rectangular concentration versus time profile across a given dosing interval.

Table 4: Brimonidine steady-state plasma pharmacokinetic parameters following administration of brinzolamide 1%/brimonidine tartrate 0.2% (BID) in healthy subjects [Day 21]

	C_{max} (ng/mL)	T_{max} (h)	$AUC_{0-\infty}$ (ng*h/mL)	$t_{1/2}$ (h)
Brimonidine	0.072 (0.023, 0.179)	0.50 (0.25, 1.00)	0.196 (0.058, 0.408)	2.57 (1.37, 4.69)

C_{max} , $AUC_{0-t_{last}}$, and $t_{1/2}$ data are presented as geometric means. T_{max} data is presented as median. Values in parentheses are (minimum, maximum) values for respective parameter.

Absorption:

Brinzolamide is absorbed through the cornea following topical ocular administration. The drug is also absorbed into the systemic circulation where it binds strongly to carbonic anhydrase in red blood cells (RBCs). Plasma drug concentration is low.

Whole blood elimination half-life is prolonged (>100 days) in humans due to RBC carbonic anhydrase binding, resulting in significant accumulation of brinzolamide in the blood. Plasma brimonidine levels (mean C_{max} =0.06 ng/mL after 10 days treatment) peak within 1 – 4 hours and decline with a systemic half-life of approximately 3 hours.

In a topical ocular clinical study comparing the systemic pharmacokinetics of brinzolamide/brimonidine

tartrate ophthalmic suspension to brinzolamide and brimonidine administered individually, the steady-state whole blood brinzolamide and N-desethylbrinzolamide pharmacokinetics were similar between the combination product and brinzolamide administered alone. Likewise, the steady-state plasma pharmacokinetics of brimonidine from the combination was similar to that observed for brimonidine administered alone.

Distribution:

Studies in rabbits showed that during a course of topical ocular BID administration, brinzolamide significantly accumulates in iris-ciliary body, choroid and especially retina, while brimonidine significantly accumulates in choroid, retina and especially in iris-ciliary body (see [Animal Pharmacokinetics](#)). Accumulation of brimonidine in iris, ciliary body, choroid/retina was also reported in cynomolgus monkeys when 0.5% brimonidine was administered BID topically in the eye. Binding to carbonic anhydrase may be a reason for prolonged ocular retention of brinzolamide.

Metabolism:

Brinzolamide is metabolized by hepatic cytochrome P450 isozymes, specifically CYP3A4, CYP2A6, CYP2B6, CYP2C8 and CYP2C9. The primary metabolite is N-desethylbrinzolamide followed by the N-desmethoxypropyl and O-desmethyl metabolites as well as an N-propionic acid analog formed by oxidation of the N-propyl side chain of O-desmethyl brinzolamide. Brinzolamide and N-desethylbrinzolamide do not inhibit cytochrome P450 isozymes at concentrations at least 100-fold above maximum systemic levels.

Brimonidine is primarily metabolized by liver, most likely by cytochrome P450 and aldehyde oxidase. The principle metabolic pathways of brimonidine are α (N)-oxidation to 2-oxobrimonidine, 3-oxobrimonidine and 2,3-dioxobrimonidine and oxidative cleavage of the imadazoline ring to yield 5-bromo-6-guanidinoquinoxaline.

Elimination:

Brinzolamide is primarily eliminated in urine as unchanged drug. In humans, urinary brinzolamide and N-desethylbrinzolamide accounted for about 60 and 6% of the dose, respectively.

Brimonidine tartrate is eliminated rapidly via extensive systemic metabolism; there is no marked systemic accumulation after multiple dosing. Urinary excretion is the major route of elimination of the drug and its metabolites. Approximately 87% of an orally-administered radioactive dose was eliminated within 120 hours, with 74% found in the urine in the first 96 hours.

Special Populations and Conditions

Studies to determine the effects of age, race, and renal or hepatic impairment have not been conducted with the brinzolamide/brimonidine fixed combination.

A study of brinzolamide in Japanese versus non-Japanese subjects showed similar systemic pharmacokinetics between the two groups. In a study of brinzolamide in subjects with renal impairment, a 1.6- to 2.8-fold increase in the systemic exposure to brinzolamide and N-desethylbrinzolamide between normal and moderately renally-impaired subjects was demonstrated. This increase in steady-state RBC concentrations of drug-related material did not inhibit RBC carbonic anhydrase activity to levels that are associated with systemic side effects. However, the combination product is not recommended for patients with severe renal impairment (creatinine clearance < 30 mL/minute).

- **Pediatrics** The systemic pharmacokinetics of brinzolamide and brimonidine, alone or in combination, in paediatric patients have not been studied.
- **Geriatrics** No overall differences in pharmacokinetics and safety have been observed between elderly and other adult patients. The C_{max} and apparent half-life of brimonidine tartrate were similar in elderly subjects (65 years or older) and younger adults, indicating that its systemic absorption and elimination are not significantly affected by age.

11 STORAGE, STABILITY AND DISPOSAL

Store SIMBRINZA at 2° to 25°C (36° - 77°F). Discard 125 days after opening the 10 mL bottle. Keep out of sight and reach of children.

12 SPECIAL HANDLING INSTRUCTIONS

Patients should be advised to avoid touching the tip of the bottle to the eye or any surface, as this may contaminate the solution. See [4.4 Administration](#) for more detailed information.

PART II: SCIENTIFIC INFORMATION

13 PHARMACEUTICAL INFORMATION

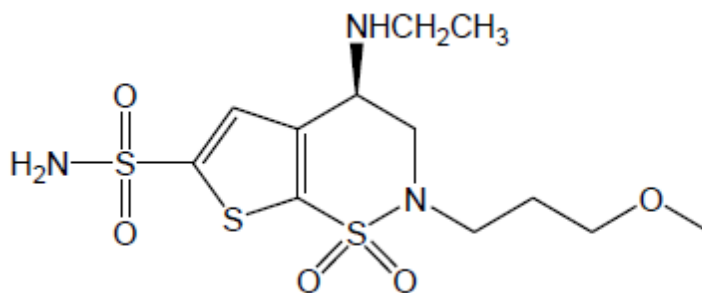
Drug Substance

Proper name: Brinzolamide

Chemical name: (R)-4-(Ethylamino)-3,4-dihydro-2-(3-methoxypropyl)-2H-thieno[3,2-e]-1,2-thiazine-6-sulfonamide 1,1-dioxide

Molecular formula and molecular mass: $C_{12}H_{21}N_3O_5S_3$; 383.51 g/mol

Structural formula:



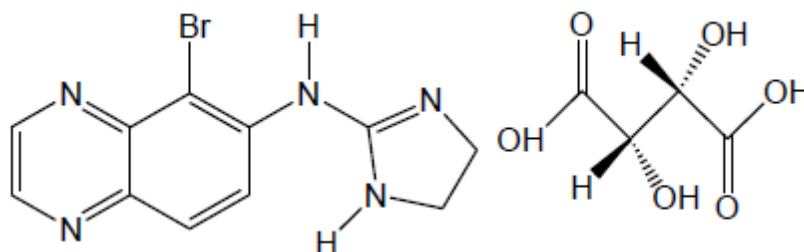
Physicochemical properties: White to off-white powder or crystals. Very slightly soluble in water at pH 7, sparingly soluble in methanol and slightly soluble in ethanol; melting point of about 131°C

Proper name: Brimonidine / Brimonidine tartrate

Chemical name: 5-Bromo-6-(2-imadazolin-2-ylamino) quinoxaline L-tartrate

Molecular formula and molecular mass: $C_{11}H_{10}BrN_5 \cdot C_4H_6O_6$; 442.22 g/mol

Structural formula:



Physicochemical properties: White to off-white, pale yellow to yellow powder or crystals. Soluble in water at pH 6.35, slightly soluble in water at pH 7.49 and in methanol; melting point 200°C

14 CLINICAL TRIALS

14.1 Clinical Trials by Indication

Intraocular Pressure (IOP) Reduction

Table 5: Summary of patient demographics for clinical trials in IOP

Study	Study design	Dosage, route of administration and duration	Study subjects (N)	Mean age ^a (±SD)	Sex (M/F)	Baseline IOP ^b (mean ± SE) [mmHg]
C-10-040 / Phase 3	Multicenter, randomized, double-masked, parallel-group, active-controlled, 3-arm, parallel-group, superiority study	<u>Brinzolamide/ Brimonidine 1%/0.2%</u> : 1 drop in each eye BID; topical ocular; 6 months	N = 193	64.6 (± 12.18)	87/106	Diurnal: 25.9 ± 0.19 9 AM: 27.0 ± 0.18 11 AM: 25.9 ± 0.21 4 PM: 24.8 ± 0.23
		<u>Brinzolamide (1%)</u> : 1 drop in each eye BID; topical ocular; 6 months	N = 192	64.2 (± 11.26)	90/102	Diurnal: 25.9 ± 0.20 9 AM: 27.0 ± 0.18 11 AM: 25.9 ± 0.22 4 PM: 24.8 ± 0.24
		<u>Brimonidine (0.2%)</u> : 1 drop in each eye BID; topical ocular; 6 months	N = 175	64.3 (± 11.61)	73/102	Diurnal: 26.0 ± 0.19 9 AM: 27.0 ± 0.19 11 AM: 26.2 ± 0.22 4 PM: 24.9 ± 0.21
C-10-041 / Phase 3	Multicenter, randomized, double-masked, parallel-group, active-controlled, 2-arm, parallel-group, non-inferiority study	<u>Brinzolamide/ Brimonidine 1%/0.2%</u> : 1 drop in each eye BID; topical ocular; 6 months	N = 452	63.2 (± 11.87)	200/252	Diurnal: 26.4 ± 0.13 9 AM: 27.0 ± 0.13 11 AM: 25.8 ± 0.14
		<u>Brinzolamide + Brimonidine (1% + 0.2%)</u> : 1 drop in each eye BID; topical ocular; 6 months	N = 436	63.4 (± 12.28)	190/246	Diurnal: 26.5 ± 0.13 9 AM: 27.0 ± 0.13 11 AM: 25.9 ± 0.15

^a Subjects and age for C-10-040 and C-10-041 are derived from the safety population

^b Baseline IOP values for C-10-040 and C-10-041 are derived from the intent-to-treat (ITT) and per protocol (PP) populations, respectively

Two multicenter, randomized, double-masked, parallel-group, active-controlled clinical studies of 6 months duration with twice daily dosing of SIMBRINZA (brinzolamide / brimonidine tartrate ophthalmic suspension) 1% / 0.2% w/v were conducted in 1450 patients with open-angle glaucoma or ocular hypertension who, in the opinion of the investigator, were insufficiently controlled on monotherapy or already using multiple IOP-lowering medications (Table 5).

Primary and supportive efficacy endpoints:

The primary efficacy endpoint for both C-10-040 and C-10-041 was an assessment of mean diurnal IOP change from baseline at 3 months, with safety and supportive efficacy evaluated through 6 months.

The C-10-040 supportive efficacy endpoints were inclusive of 3 time points (i.e., 9am, +2hr, +7hr) whereas the C 10 041 supportive efficacy endpoints were inclusive of 2 time points (i.e., 9am, +2hr).

Select supportive efficacy endpoints included:

- Mean diurnal change from baseline at Week 2, Week 6 and Month 6
- Mean IOP change from baseline at Week 2, Week 6, Month 3, and Month 6 for each assessment time point.
- Mean IOP percent change from baseline at Week 2, Week 6, Month 3, and Month 6 for each assessment time point.

Study C-10-040

Based on clinical study C-10-040, brinzolamide/brimonidine fixed combination was superior to brinzolamide and brimonidine with respect to mean diurnal IOP reduction from baseline at all visits. The mean difference in diurnal IOP change from baseline favoured the brinzolamide/brimonidine fixed combination compared to brinzolamide or brimonidine, respectively, at all study visits. The observed magnitude of mean difference in change at all visits was statistically significant (Table 6).

Table 6: Comparison of Mean Diurnal^a IOP (mmHg) Change from Baseline by Visit (C-10-040) – Intent-to-treat Population

Visit	Brinz/Brim [BID]		Brinz [BID]		Brim [BID]		Mean Difference ^b (95% CI) [mmHg]	p-value
	N	Mean (SE) [mmHg]	N	Mean (SE) [mmHg]	N	Mean (SE) [mmHg]		
2 weeks	191	-7.6 (0.22)	191	-6.1 (0.22)	--	--	-1.5 (-2.1, -1.0)	p<0.0001
			--	--	174	-6.0 (0.23)	-1.6 (-2.1, -1.0)	p<0.0001
6 weeks	188	-7.8 (0.22)	189	-6.2 (0.22)	--	--	-1.6 (-2.1, -1.0)	p<0.0001
			--	--	172	-6.2 (0.23)	-1.6 (-2.2, -1.1)	p<0.0001
3 months ^c	176	-7.9 (0.22)	182	-6.5 (0.23)	--	--	-1.4 (-1.9, -0.8)	p<0.0001
			--	--	161	-6.4 (0.24)	-1.5 (-2.0, -0.9)	p<0.0001
6 months	160	-7.8 (0.23)	178	-6.7 (0.23)	--	--	-1.1 (-1.7, -0.6)	p=0.0001
			--	--	145	-6.4 (0.24)	-1.4 (-1.9, -0.8)	p<0.0001

Brinz/Brim [BID] = Brinzolamide/Brimonidine 10 mg/mL + 2 mg/mL Eye Drops, Suspension [BID]

Brinz [BID] = Brinzolamide 10 mg/mL Eye Drops, Suspension [BID]

Brim [BID] = Brimonidine Tartrate 2 mg/mL Eye Drops, Solution [BID]

SE = Standard Error; CI = Confidence Interval

^a Diurnal IOP change averaged over 9AM, +2 hr, and +7 hr time points.

^b Estimates based on the least squares means derived from a statistical model that accounts for correlated IOP measurements within patient where site and actual 9 AM baseline IOP stratum are in the model.

^c Primary efficacy endpoint.

The difference in mean IOP change from baseline (Table 7) at each time point favoured the brinzolamide/brimonidine fixed combination compared to brinzolamide and brimonidine. The observed magnitude of difference in change at all time points was statistically significant, favouring brinzolamide/brimonidine over brimonidine by at least 1 mmHg at all (12/12) time points and over brinzolamide at a majority (8/12) of time points. Mean IOP reductions from baseline with brinzolamide/brimonidine fixed combination ranged from 6.0 to 8.8 mmHg.

Table 7: Comparison of Mean IOP (mmHg) Change from Baseline by Visit and Time Point (C-10-040) - Intent-to-treat Population

Visit	Brinz/Brim [BID]		Brinz [BID]		Brim [BID]		Mean Difference ^a (95% CI) [mmHg]	p-value
	N	Mean (SE) [mmHg]	N	Mean (SE) [mmHg]	N	Mean (SE) [mmHg]		
2 weeks								
9AM	191	-6.0 (0.25)	191	-5.0 (0.25)	--	--	-0.9 (-1.6, -0.3)	p=0.0036
			--	--	174	-4.3 (0.26)	-1.7 (-2.3, -1.0)	p<0.0001
+2 hr	189	-8.4 (0.25)	191	-5.9 (0.25)	--	--	-2.5 (-3.1, -1.8)	p<0.0001
			--	--	174	-7.0 (0.26)	-1.4 (-2.1, -0.8)	p<0.0001
+7 hr	188	-7.3 (0.25)	191	-5.9 (0.25)	--	--	-1.4 (-2.0, -0.8)	p<0.0001
			--	--	173	-5.5 (0.26)	-1.8 (-2.5, -1.2)	p<0.0001
6 weeks								
9AM	188	-6.2 (0.25)	189	-5.0 (0.25)	--	--	-1.2 (-1.8, -0.5)	p=0.0003
			--	--	172	-4.7 (0.26)	-1.5 (-2.2, -0.9)	p<0.0001
+2 hr	185	-8.6 (0.25)	188	-6.3 (0.25)	--	--	-2.3 (-3.0, -1.7)	p<0.0001
			--	--	170	-7.0 (0.26)	-1.6 (-2.3, -1.0)	p<0.0001
+7 hr	184	-7.6 (0.25)	188	-6.3 (0.25)	--	--	-1.4 (-2.0, -0.7)	p<0.0001
			--	--	170	-5.8 (0.26)	-1.9 (-2.5, -1.2)	p<0.0001
3 months								
9AM	176	-6.2 (0.26)	182	-5.4 (0.26)	--	--	-0.8 (-1.4, -0.1)	p=0.0188
			--	--	161	-4.8 (0.27)	-1.5 (-2.1, -0.8)	p<0.0001
+2 hr	173	-8.8 (0.25)	182	-6.4 (0.25)	--	--	-2.3 (-3.0, -1.7)	p<0.0001
			--	--	159	-7.4 (0.27)	-1.4 (-2.0, -0.7)	p<0.0001
+7 hr	172	-7.7 (0.25)	180	-6.4 (0.25)	--	--	-1.3 (-1.9, -0.7)	p<0.0001
			--	--	159	-5.8 (0.27)	-1.9 (-2.6, -1.3)	p<0.0001
6 months								
9AM	160	-6.1 (0.26)	178	-5.4 (0.26)	--	--	-0.7 (-1.4, -0.0)	p=0.0363
			--	--	145	-4.9 (0.28)	-1.3 (-1.9, -0.6)	p=0.0003
+2 hr	160	-8.6 (0.26)	178	-6.6 (0.25)	--	--	-2.0 (-2.6, -1.3)	p<0.0001
			--	--	145	-7.0 (0.27)	-1.6 (-2.3, -0.9)	p<0.0001
+7 hr	160	-7.5 (0.26)	178	-6.7 (0.25)	--	--	-0.8 (-1.5, -0.2)	p=0.0137
			--	--	144	-6.0 (0.27)	-1.5 (-2.1, -0.8)	p<0.0001

Brinz/Brim [BID] = Brinzolamide/Brimonidine 10 mg/mL + 2 mg/mL Eye Drops, Suspension [BID]

Brinz [BID] = Brinzolamide 10 mg/mL Eye Drops, Suspension [BID]

Brim [BID] = Brimonidine Tartrate 2 mg/mL Eye Drops, Solution [BID]

SE = Standard Error; CI = Confidence Interval

^a Estimates based on the least squares means derived from a statistical model that accounts for correlated IOP measurements within patient where site and actual 9 AM baseline IOP stratum are in the model.

The difference in mean percent IOP change from baseline at each time point favoured the brinzolamide/brimonidine fixed combination compared to brinzolamide and brimonidine (Table 8). Mean percent IOP reductions from baseline with brinzolamide/brimonidine fixed combination ranged from 23 to 34%.

Table 8: Comparison of Mean IOP Percent Change from Baseline (mmHg) by Visit and Time Point (C-10-040) - Intent-to-treat Population

Visit	Brinz/Brim [BID]		Brinz [BID]		Brim [BID]		Mean Difference ^a (95% CI) [%]	p-value
	N	Mean (SE) [%]	N	Mean (SE) [%]	N	Mean (SE) [%]		
2 weeks	191	-22.9 (0.95)	191	-19.4 (0.96)	--	--	-3.5 (-5.9, -1.1)	p=0.0043
9AM			--	--	174	-16.6 (1.00)	-6.3 (-8.8, -3.9)	p<0.0001
+2 hr	189	-32.4 (0.94)	191	-22.9 (0.94)	--	--	-9.5 (-11.9, -7.1)	p<0.0001
			--	--	174	-26.6 (0.99)	-5.8 (-8.3, -3.4)	p<0.0001
+7 hr	188	-28.2 (0.94)	191	-22.4 (0.94)	--	--	-5.7 (-8.1, -3.3)	p<0.0001
			--	--	173	-20.9 (0.99)	-7.2 (-9.7, -4.7)	p<0.0001
6 weeks	188	-23.8 (0.96)	189	-19.3 (0.96)	--	--	-4.5 (-6.9, -2.1)	p=0.0003
9AM			--	--	172	-18.0 (1.00)	-5.8 (-8.3, -3.3)	p<0.0001
+2 hr	185	-33.1 (0.95)	188	-23.9 (0.95)	--	--	-9.2 (-11.6, -6.8)	p<0.0001
			--	--	170	-26.7 (1.00)	-6.4 (-8.9, -3.9)	p<0.0001
+7 hr	184	-29.5 (0.95)	188	-23.7 (0.95)	--	--	-5.8 (-8.3, -3.4)	p<0.0001
			--	--	170	-22.1 (0.99)	-7.4 (-9.9, -4.9)	p<0.0001
3 months	176	-23.9 (0.97)	182	-20.9 (0.97)	--	--	-3.0 (-5.5, -0.6)	p=0.0153
9AM			--	--	161	-18.4 (1.02)	-5.5 (-8.0, -3.0)	p<0.0001
+2 hr	173	-33.8 (0.97)	182	-24.6 (0.96)	--	--	-9.2 (-11.7, -6.8)	p<0.0001
			--	--	159	-28.5 (1.02)	-5.3 (-7.9, -2.8)	p<0.0001
+7 hr	172	-29.6 (0.96)	180	-24.2 (0.96)	--	--	-5.5 (-7.9, -3.0)	p<0.0001
			--	--	159	-22.1 (1.01)	-7.6 (-10.1, -5.0)	p<0.0001
6 months	160	-23.6 (1.00)	178	-20.8 (0.98)	--	--	-2.7 (-5.2, -0.2)	p=0.0318
9AM			--	--	145	-18.8 (1.05)	-4.8 (-7.4, -2.2)	p=0.0003
+2 hr	160	-33.2 (0.99)	178	-25.4 (0.96)	--	--	-7.8 (-10.3, -5.3)	p<0.0001
			--	--	145	-26.7 (1.04)	-6.4 (-9.0, -3.8)	p<0.0001
+7 hr	160	-29.0 (0.98)	178	-25.4 (0.96)	--	--	-3.5 (-6.0, -1.0)	p=0.0055
			--	--	144	-23.0 (1.04)	-5.9 (-8.5, -3.3)	p<0.0001

Brinz/Brim [BID] = Brinzolamide/Brimonidine 10 mg/mL + 2 mg/mL Eye Drops, Suspension [BID]

Brinz [BID] = Brinzolamide 10 mg/mL Eye Drops, Suspension [BID]

Brim [BID] = Brimonidine Tartrate 2 mg/mL Eye Drops, Solution [BID]

SE = Standard Error; CI = Confidence Interval

^a Estimates based on the least squares means derived from a statistical model that accounts for correlated IOP measurements within patient where site and actual 9 AM baseline IOP stratum are in the model.

Study C-10-041

Based on clinical study C-10-041, brinzolamide/brimonidine fixed combination was non-inferior to the unfixed combination of brinzolamide plus brimonidine administered concomitantly (i.e., brinzolamide+brimonidine) with respect to mean diurnal IOP change from baseline at all visits. The upper bound of the 95% confidence interval for the treatment group difference was well below the pre-specified non-inferiority margin of 1.5 mmHg (Table 9).

Table 9: Comparison of Mean Diurnal^a IOP (mmHg) Change from Baseline by Visit – Study (C-10-041) – Per Protocol Population

Visit	Brinz/Brim [BID]		Brinz+Brim [BID]		Mean Difference ^b (95% CI) [mmHg]
	N	Mean (SE) [mmHg]	N	Mean (SE) [mmHg]	
2 weeks	394	-8.4 (0.16)	384	-8.4 (0.16)	-0.0 (-0.4, 0.3)
6 weeks	384	-8.5 (0.16)	377	-8.4 (0.16)	-0.1 (-0.4, 0.2)
3 months ^c	384	-8.5 (0.16)	373	-8.3 (0.16)	-0.1 (-0.5, 0.2)
6 months	346	-8.1 (0.16)	330	-8.2 (0.16)	0.1 (-0.3, 0.4)

Brinz/Brim [BID] = Brinzolamide/Brimonidine 10 mg/mL + 2 mg/mL Eye Drops, Suspension [BID]

Brinz+Brim [BID] = Brinzolamide 10 mg/mL Eye Drops, Suspension [BID] + Brimonidine Tartrate 2 mg/mL Eye Drops, Solution [BID]

SE = Standard Error; CI = Confidence Interval

^a Diurnal IOP change averaged over 9AM and +2 hr time points.

^b Estimates based on the least squares means derived from a statistical model that accounts for correlated IOP measurements within patient where site and actual 9 AM baseline IOP stratum are in the model.

^c Primary efficacy endpoint.

Mean IOP change from baseline at each time point was similar between brinzolamide/brimonidine fixed combination and the unfixed combination of brinzolamide plus brimonidine administered concomitantly (i.e., Brinz+Brim) (Table 10). Mean IOP reductions from baseline with brinzolamide/brimonidine fixed combination ranged from 6.5 to 9.7 mmHg.

Table 10: Comparison of Mean IOP (mmHg) Change from Baseline by Visit and Time Point (C-10-041) - Per Protocol Population

Visit	Brinz/Brim [BID]		Brinz+Brim [BID]		Mean Difference ^a (95% CI) [mmHg]
	N	Mean (SE) [mmHg]	N	Mean (SE) [mmHg]	
2 weeks					
9AM	394	-6.9 (0.17)	384	-7.0 (0.17)	0.1 (-0.3, 0.5)
+2 hr	392	-9.6 (0.16)	383	-9.4 (0.16)	-0.2 (-0.6, 0.2)
6 weeks					
9AM	384	-6.9 (0.17)	377	-6.9 (0.17)	-0.0 (-0.4, 0.3)
+2 hr	383	-9.6 (0.16)	372	-9.5 (0.16)	-0.1 (-0.5, 0.3)
3 months					
9AM	384	-6.9 (0.17)	373	-6.8 (0.17)	-0.1 (-0.5, 0.3)

+2 hr	380	-9.7 (0.16)	363	-9.6 (0.17)	-0.2 (-0.6, 0.2)
6 months					
9AM	345	-6.5 (0.18)	330	-6.7 (0.18)	0.2 (-0.2, 0.6)
+2 hr	344	-9.3 (0.17)	328	-9.2 (0.17)	-0.1 (-0.5, 0.3)

Brinz/Brim [BID] = Brinzolamide/Brimonidine 10 mg/mL + 2 mg/mL Eye Drops, Suspension [BID]

Brinz+Brim [BID] = Brinzolamide 10 mg/mL Eye Drops, Suspension [BID] + Brimonidine Tartrate 2 mg/mL Eye Drops, Solution [BID]

SE = Standard Error; CI = Confidence Interval

^a Estimates based on the least squares means derived from a statistical model that accounts for correlated IOP measurements within patient where site and actual 9 AM baseline IOP stratum are in the model.

Mean IOP percent change from baseline at each time point was similar between Brinzolamide/brimonidine fixed combination and the unfixed combination of brinzolamide plus brimonidine administered concomitantly (ie, Brinz+Brim) (Table 11). Mean percent IOP reductions from baseline with brinzolamide/brimonidine fixed combination ranged from 25 to 37%.

Table 11: Comparison of Mean IOP (mmHg) Percent Change from Baseline by Visit and Time Point (C-10-041) - Per Protocol Population

Visit	Brinz/Brim [BID]		Brinz+Brim [BID]		Mean Difference ^a (95% CI) [mmHg]
	N	Mean (SE) [mmHg]	N	Mean (SE) [mmHg]	
2 weeks					
9AM	394	-26.3 (0.63)	384	-26.7 (0.62)	0.4 (-1.0, 1.8)
+2 hr	392	-36.5 (0.60)	383	-35.6 (0.60)	-0.9 (-2.3, 0.6)
6 weeks					
9AM	384	-26.5 (0.63)	377	-26.2 (0.63)	-0.3 (-1.7, 1.2)
+2 hr	383	-36.6 (0.61)	372	-36.0 (0.61)	-0.5 (-2.0, 0.9)
3 months					
9AM	384	-26.5 (0.63)	373	-26.0 (0.63)	-0.5 (-1.9, 1.0)
+2 hr	380	-36.8 (0.61)	363	-36.1 (0.61)	-0.7 (-2.2, 0.8)
6 months					
9AM	345	-24.9 (0.65)	330	-25.7 (0.65)	0.8 (-0.7, 2.3)
+2 hr	344	-35.2 (0.62)	328	-35.0 (0.63)	-0.2 (-1.7, 1.3)

Brinz/Brim [BID] = Brinzolamide/Brimonidine 10 mg/mL + 2 mg/mL Eye Drops, Suspension [BID]

Brinz+Brim [BID] = Brinzolamide 10 mg/mL Eye Drops, Suspension [BID] + Brimonidine Tartrate 2 mg/mL Eye Drops, Solution [BID]

SE = Standard Error; CI = Confidence Interval

^a Estimates based on the least squares means derived from a statistical model that accounts for correlated IOP measurements within patient where site and actual 9 AM baseline IOP stratum are in the model.

15 MICROBIOLOGY

No microbiological information is required for this drug product.

16 NON-CLINICAL TOXICOLOGY

General Toxicology:

Acute Toxicity:

Brinzolamide

Single-dose toxicity studies with brinzolamide included a 1-day topical ocular irritation evaluation in rabbits and acute oral toxicity studies in rats and mice. Exaggerated topical ocular dosing studies with a 2.0% formulation of brinzolamide indicated that ocular irritation and comfort scores were consistent with those normally observed with eye drop suspensions, and no significant clinical findings were noted.

Single-dose oral toxicity studies were conducted in rats and mice to assess the acute toxicity of brinzolamide. The oral LD₅₀ of brinzolamide in mice was estimated to be 1,400 mg/kg, with the oral LD₅₀ in rats estimated at 1,000 to 2,000 mg/kg.

Brimonidine

Single-dose toxicity studies showed that intravenous and oral LD₅₀ of brimonidine was 50 mg/kg in mice and 100 mg/kg in rats. The most frequently observed clinical signs in the acute/single dose toxicity studies included sedation, ataxia, prostration, ptosis, reduced/loss of blink reflex, opacification of the cornea, hypotension, bradycardia, hypothermia, respiratory depression, respiratory arrest and circulatory collapse.

Repeat-dose Toxicity:

Brinzolamide/brimonidine fixed combination

Toxicity of brinzolamide/brimonidine tartrate was studied in rabbits that were topically administered in the TID up to nine months.

These studies demonstrated that there was no significant ocular irritation when brinzolamide/brimonidine tartrate was administered topically. Irritation scores were unremarkable and similar to controls. Compared to the vehicle control, brinzolamide/brimonidine tartrate significantly increased corneal thickness which was more prominent in male rabbits. This finding has been observed in animal studies with 1% brinzolamide.

Brinzolamide /brimonidine tartrate caused sedation, penile erection and female urogenital swelling, significant increase in blood glucose level by about 2- to 3- fold and significant decrease in blood cholesterol level. These findings were also observed in rabbits topically administered in the eyes with 0.2% brimonidine tartrate alone. Reduced food consumption and decreased body weight gain were consistently observed in brinzolamide/brimonidine tartrate treated rabbits compared to those in the vehicle control group in a nine-month long-term study. These effects were more prominent in male rabbits. Furthermore, mean liver weight and the mean ratio of liver weight/body weight in male rabbits were significantly higher in the brinzolamide/brimonidine tartrate study group at Day 91, but not on Day 274. Hepatocellular cytoplasmic vacuolization and concurrent hepatocellular glycogen accumulation were found in the rabbits treated with a brimonidine/brinzolamide fixed combination formulation, including brinzolamide/brimonidine tartrate, but not in the vehicle control group.

The severity and incidence of these changes increased with treatment length and were more prominent in male rabbits. There were no pathological changes (such as hepatocellular degeneration or necrosis) identified in the liver. Minimal to mild diffuse islet cell hyperplasia in pancreas was observed in male rabbits treated with brinzolamide/brimonidine tartrate.

Brinzolamide

Five repeat-dose topical ocular studies were conducted in rabbits, ranging in duration from 1 to 6 months, and a 1-year topical ocular study was conducted in nonhuman primates. These studies demonstrated that there was no significant ocular toxicity or irritation when the drug was administered topically. Irritation scores were unremarkable and similar to controls. Concentrations of brinzolamide ophthalmic suspension as high as 4.0% were administered chronically up to 4 times a day in rabbits and three times a day in monkeys without significant toxicological findings.

In repeated-dose oral toxicity studies in rats, pharmacological effects on urine volume, specific gravity, and electrolytes were observed. Increase in relative kidney weights, chronic nephritis, minimal to mild nephropathy, with crystalline material in the urine, was also observed at 8 mg/kg/day and higher. Reduction in food consumption and decrease in body weight gain were observed in higher doses. Increased incidence of chronic nephritis was observed in mice.

Brimonidine

Repeat-dose toxicity studies with brimonidine tartrate were conducted in mice, rats, rabbits, dogs and monkeys for durations of up to one year. The most notable effects seen in these studies were related to the known pharmacological effect of brimonidine.

There were no observable adverse effects in oral dosing in mice, rats, rabbits, dogs and monkeys at approximately 165, 80, 25, 55 and 33 times the recommended ocular human dose, respectively.

Orally dosed at levels of approximately 3000 times the recommended human ocular dose in chronic studies, mice showed goblet cell hyperplasia and depletion in the rectum and colon, hypertrophy of the tunica muscularis of small and large intestine, and hyperplasia of the non-glandular epithelium of the stomach. Rats dosed orally at approximately 1500 times the human ocular dose showed thickening of muscularis mucosa of small intestine, and a dose related incidence of ileal intussusception was observed in all rats, but no associated lesions or morphological changes were observed. Evidence of toxicity characterised by decreased body weight gain and/or decreased food consumption was often seen at the higher oral doses in the mouse, rat and monkey. The most notable effects seen in the subacute studies was sedation, ataxia, hypoactivity, ptosis, decreased muscle tone, hypotension and bradycardia.

Topically dosed in the eyes with a brimonidine solution (0.15% to 0.2%) three time daily for up to three months, rabbits showed sedation, penile erection, female urogenital swelling, significant increase in blood glucose level (about 2 to 3 folds) and/or significant decrease in blood cholesterol level. There was no significant effect on body weight gain and no test-article-related abnormal findings in pathological examination in all organs after three-month of the treatment.

Carcinogenicity:

Brinzolamide

An initial cell proliferation study in rats showed an absence of proliferation potential with brinzolamide. No test article-related carcinogenicity was observed in 2-year oral dosing studies in mice and rats.

Brimonidine

Carcinogenicity studies with brimonidine were conducted in mice and rats orally dosed at 2.5 mg/kg/day and 1 mg/kg/day, respectively, for 21 to 24 months. No test article-related carcinogenicity was observed.

Genotoxicity:

Brinzolamide

Two *in vitro* and two *in vivo* mutation assays were conducted with brinzolamide in order to evaluate the genotoxicity potential of the drug substance. Results of the *in vitro* bacterial mutation and the two *in vivo* assays unequivocally demonstrate a lack of mutagenicity. The *in vitro* mammalian cell mutation assay indicated a potential for mutagenicity. However, carcinogenic studies did not show brinzolamide-related carcinogenicity (see [Carcinogenicity](#)).

Brimonidine

Brimonidine tartrate was not mutagenic or cytogenic in a series of *in vitro* and *in vivo* studies including the Ames/Salmonella mutagenicity assay, chromosomal aberration assay in Chinese Hamster Ovary (CHO) cells, cytogenetic assay, host-mediated assay and dominant lethal assay in mice.

Reproductive and Developmental Toxicology:

Brinzolamide

Brinzolamide when given orally demonstrated no effect on male or female fertility. Brinzolamide increased the incidence of unossified sternebrae or hyoid and reduced ossification of the skull in rats at 18 mg/kg/day (783 times the recommended human ophthalmic dose) given orally. Reduced ossification was not dose-dependent. In rabbits, no malformations were observed and ossification appeared to be unaffected. In a peri- and postnatal effect study, F1 pup body weights were significantly reduced, as compared with controls, throughout the lactation period, at the 15 mg/kg/day dose level. These effects are comparable with other drugs of this class.

Brimonidine

Reproductive toxicology studies conducted with brimonidine tartrate in rats and rabbits showed that brimonidine tartrate had no adverse effects on fertility and general reproductive performance and showed no evidence of embryo lethality or teratogenicity at the dosages administered.

There were no treatment-related reproductive and teratological effects observed in the F1 rat pup group, although a reduction in body weight was observed at a dose level of 1.65 mg base/kg/day, after 14 days. Dose related reduction in body weight gains were observed in rat dams at dose levels of 0.66 and 1.65 mg base/kg/day after 15 days.

There was no evidence of treatment-related embryotoxicity, fetal toxicity, or teratogenicity in rabbits at brimonidine oral dose levels up to 3.3 mg base/kg/day at which spontaneous abortions occurred. A decrease in weight gain during the dosing period was observed.

Special Toxicology:

Brinzolamide is considered to have little or no potential to induce contact sensitisation based on a guinea pig maximisation test.

Animal Pharmacology

The pharmacology, safety pharmacology, pharmacokinetic and toxicological attributes of brinzolamide and brimonidine tartrate in nonclinical models have been extensively characterized. The nonclinical assessment of brinzolamide/brimonidine tartrate is based upon the established nonclinical profiles of the individual active drug substances and evaluation of important aspects of the combination product.

Brinzolamide has high affinity for, and potent inhibitory activity against, human carbonic anhydrase II with a K_i of 0.13 nM and an IC_{50} of 3.2 nM. Carbonic anhydrase is an enzyme found in many tissues of the body, including the eye. It catalyzes the reversible reaction involving the hydration of carbon

dioxide and the dehydration of carbonic acid. Inhibition of carbonic anhydrase in the ciliary processes of the eye decreases aqueous humour secretion, presumably by slowing the formation of bicarbonate ions with subsequent reduction of sodium and fluid transport. The result is a reduction in intraocular pressure (IOP). Alone, brinzolamide has a peak ocular hypotensive effect occurring at 2 to 3 hours post-dosing.

Brinzolamide lowered IOP by 20–30% in a dose-dependent manner in lasered ocular hypertensive cynomolgus monkeys and Dutch-belted rabbits following topical ocular delivery.

Brimonidine is a potent, selective alpha-2 adrenoceptor agonist. Alpha-2 adrenoceptors have been localized to prejunctional loci in the eye where they inhibit the release of norepinephrine from nerve endings. High levels of alpha-2 adrenoceptors have been found in the iris and ciliary epithelia and lower levels in ciliary muscle and retina.

Topical ocular treatment of normal, unanesthetized cats, New Zealand albino rabbits and capucin monkeys (*Cebus paella*) with 0.5 µg to 5000 µg of brimonidine tartrate produces dose-related decreases in IOP and in pupil diameter. A dose of 200 µg brimonidine tartrate promotes a bilateral maximal reduction in IOP at 1 to 2 hours, followed by a gradual return to baseline after unilateral, topical ocular treatment in ketamine-anesthetized normal cynomolgus monkeys (*Macaca fascicularis*). The IOP-lowering effect was also observed in cynomolgus monkeys that had undergone unilateral superior cervical sympathectomy. Treatment of glaucomatous cynomolgus monkeys (rendered ocular hypertensive as a consequence of laser trabeculoplasty) for 5 days with brimonidine tartrate 5 mg/mL two times daily decreased IOP by approximately 50%. Topical ocular treatment of normal cynomolgus monkeys with brimonidine tartrate promotes a dose-dependent reduction (up to 67% below baseline for 5 mg/mL brimonidine tartrate) in aqueous humor flow without affecting total outflow measured by fluorophotometry. This suggests that the drug's mechanism of action to lower IOP in monkeys is by reducing the production of aqueous humor.

Fluorophotometric studies in animals and humans suggest that brimonidine tartrate (alpha-2 adrenergic receptor agonist) has a dual mechanism of action by reducing aqueous humor production and increasing uveoscleral outflow. Brimonidine tartrate has a peak ocular hypotensive effect occurring at two hours post-dosing. The result is a reduction in intraocular pressure (IOP).

Nonclinical primary pharmacology studies utilizing brinzolamide/brimonidine have not been conducted in animal models.

Animal Pharmacodynamics

No non-clinical ocular or systemic pharmacodynamic studies were conducted with brinzolamide/brimonidine tartrate since the pharmacology of each active component has been well established previously in the medical and scientific literature.

Other Safety Pharmacology

An *in vitro* hERG channel assay was conducted with the brinzolamide/ brimonidine tartrate combination. Treatment with 0.3/0.1, 1/0.3, and 3/1 µg/mL brinzolamide/ brimonidine combinations produced no inhibition of hERG tail current in HEK293 cells stably transfected with hERG cDNA. No other safety pharmacology studies were conducted using the brinzolamide/ brimonidine combination.

Animal Pharmacokinetics

In vivo Studies

Two ocular tissue uptake studies in male rabbits with brinzolamide/brimonidine tartrate were conducted to assess potential pharmacokinetic interactions when administered by the topical ocular route.

Brinzolamide

Following a single topical ocular dose of brinzolamide/brimonidine tartrate in male rabbits, brinzolamide is readily absorbed into all internal ocular structures with T_{max} in aqueous humor (AH) about 2 hours and distributed more in iris-ciliary body and choroid than other ocular structures. In a rabbit study with radiolabeled brinzolamide, half-lives ($T_{1/2}$) of the radioactivity in iris-ciliary body and choroid was about 38 days and 50 days, respectively, while the $T_{1/2}$ in AH was about 3.8 hours. During BID topical ocular administration in rabbits, brinzolamide significantly accumulates in iris-ciliary body, choroid and especially retina, resulting in a higher level of AUC_{0-12h} and similar level of C_{max} of brinzolamide in these ocular structures compared to those at the site of administration (i.e. Bulbar conjunctiva (BC) and cornea) (Table 12). Brinzolamide also significantly accumulates in the blood.

Similar observations are obtained for 1% brinzolamide therapy. However, the level of exposure (AUC_{0-12h} and C_{max}) to brinzolamide was somewhat lower in ICB but moderately higher in choroid after 14 days of BID topical ocular administration of brinzolamide/brimonidine tartrate compared to that in the 1% brinzolamide therapy (Table 12).

Table 12: Mean AUC_{0-12h} and C_{max} for brinzolamide in male rabbits following a single bilateral dose at Day 1, and following the last bilateral dose at Day 14 of BID topical ocular administration with 1% brinzolamide/0.2% brimonidine, or 1% brinzolamide

Study P-11-511 (PKDM 1688)		1% Brinzolamide/ 0.2% Brimonidine		1% Brinzolamide	
		AUC_{0-12h} (ng*h/g or mL)	C_{max} (ng/g or mL)	AUC_{0-12h} (ng*h/g or mL)	C_{max} (ng/g or mL)
Day 1	Bulbar Conjunctiva	15800	11000	16500	3410
	Cornea	16200	4950	18700	4890
	Aqueous Humor	1350	277	1740	389
	Iris Ciliary Body	19400	2540	20900	2390
	Choroid	36700	5030	25400	3000
	Lens	BLQ	BLQ	374	45.5
	Retina	3770	442	3210	333
	Blood	16400	1760	16500	1640
Day 14	Bulbar Conjunctiva	26600	8980	28500	3850
	Cornea	24000	5500	24300	5950
	Aqueous Humor	1800	354	2300	681

Study P-11-511 (PKDM 1688)		1% Brinzolamide/ 0.2% Brimonidine		1% Brinzolamide	
		AUC _{0-12h} (ng*h/g or mL)	C _{max} (ng/g or mL)	AUC _{0-12h} (ng*h/g or mL)	C _{max} (ng/g or mL)
	Iris Ciliary Body	29600	3300	40400	5580
	Choroid	89900	9260	55500	7630
	Lens	4880	460	5490	493
	Retina	90600	8220	87500	7760
	Blood	62500	6020	64100	6140

Due to its high affinity for CA-II, brinzolamide distributes extensively into the RBCs and exhibits a long half-life in whole blood (approximately 111 days). N-desethyl brinzolamide also binds to CA and accumulates in RBCs. This metabolite binds mainly to CA-I in the presence of brinzolamide. Brinzolamide is eliminated predominantly in the urine as unchanged drug. N-Desethyl brinzolamide is also found in the urine along with lower concentrations of the N-desmethoxypropyl and O-desmethyl metabolites.

Brimonidine

Following topical ocular administration of brinzolamide/brimonidine tartrate in male rabbits, brimonidine is rapidly absorbed into all internal ocular structures with a $T_{max} \leq 0.5$ hour in AH and retina. Brimonidine in the eye mostly distributes into iris-ciliary body and choroid following a single dose. The apparent $T_{1/2}$ in BC, cornea and AH is 1 to 3 hours, while the apparent $T_{1/2}$ in iris-ciliary body, choroid and retina is more than 10 hours. During BID topical ocular administration, brimonidine significantly accumulates in choroid, retina and especially iris-ciliary body, resulting in higher AUC_{0-12h} and C_{max} in these ocular structures compared to that at the site of administration (BC and cornea). There is no significant accumulation of brimonidine in the plasma (Table 13). Similar observations were obtained for 0.2% brimonidine monotherapy. However, compared to 0.2% brimonidine, brinzolamide/brimonidine tartrate appears to result in a somewhat higher brimonidine levels in BC, cornea, AH, choroid and lens (Table 13).

Table 13: Mean AUC_{0-12h} and C_{max} for brimonidine in male rabbits following a single bilateral dose at Day 1 and following the last bilateral dose at Day 14 of BID topical ocular administration with 1% brinzolamide/0.2% brimonidine, or 0.2% brimonidine.

Study P-11-511 (PKDM 1688)		1% Brinzolamide/ 0.2% Brimonidine		1% Brinzolamide	
		AUC _{0-12h} (ng*h/g or mL)	C _{max} (ng/g or mL)	AUC _{0-12h} (ng*h/g or mL)	C _{max} (ng/g or mL)
Day 1	Bulbar Conjunctiva	7470	2900	2570	802
	Cornea	3680	2840	2220	1480
	Aqueous Humor	572	396	339	237
	Iris Ciliary Body	98800	9890	56500	5940
	Choroid	30000	2930	10000	1010
	Lens	77.6	10	48.8	6.11
	Retina	742	91.7	851	104
	Plasma	2.03	1.71	2.47	1.43
Day 14	Bulbar Conjunctiva	11400	3900	4720	1010
	Cornea	6470	3480	3420	1630
	Aqueous Humor	1170	575	557	264
	Iris Ciliary Body	1260000	136000	1110000	105000
	Choroid	752000	75200	280000	28200
	Lens	685	74.8	350	45.3
	Retina	13300	2130	14900	1980
	Plasma	3.82	2.77	3.35	2.67

PATIENT MEDICATION INFORMATION

READ THIS FOR SAFE AND EFFECTIVE USE OF YOUR MEDICINE

Pr**SIMBRINZA®**

Brinzolamide / Brimonidine Tartrate Ophthalmic Suspension

Read this carefully before you start taking **SIMBRINZA®** and each time you get a refill. This leaflet is a summary and will not tell you everything about this drug. Talk to your healthcare professional about your medical condition and treatment and ask if there is any new information about **SIMBRINZA**.

What is SIMBRINZA used for?

SIMBRINZA is used to treat high pressure in the eyes of adults with open-angle glaucoma or ocular hypertension (conditions where there is increased pressure in the eyes).

How does SIMBRINZA work?

SIMBRINZA contains brinzolamide and brimonidine tartrate. These reduce the production of fluid in the eye and increases the amount of fluid coming out of the eye. This lowers high pressure in the eye.

What are the ingredients in SIMBRINZA?

Medicinal ingredients: brinzolamide and brimonidine tartrate.

Non-medicinal ingredients: Benzalkonium chloride (as preservative 0.003%), Boric acid, Carbomer 974P, Hydrochloric acid (to adjust pH), Mannitol, Propylene glycol, purified water, Sodium chloride, Sodium hydroxide (to adjust pH) and Tyloxapol.

SIMBRINZA comes in the following dosage forms:

Liquid suspension.

Do not use SIMBRINZA:

- If you are allergic to brinzolamide or brimonidine tartrate, sulfonamides, any of the other ingredients of SIMBRINZA or any parts of the container (see **What are the ingredients in SIMBRINZA?**).
- if you are taking monoamine oxidase (MAO) inhibitors or certain antidepressants (used to treat depression). You must tell your healthcare professional if you are taking any antidepressant drug.
- if you have severe kidney problems.
- if you have too much acidity in your blood (a condition called hyperchloraemic acidosis).
- in babies and infants aged less than 2 years and in children 2 to 17 years of age.

To help avoid side effects and ensure proper use, talk to your healthcare professional before you take SIMBRINZA. Talk about any health conditions or problems you may have, including if you have or have had:

- dry eyes or problems with your cornea
- kidney problems
- previous treatment for glaucoma (a condition where there is too much pressure in the eye)
- history of using contact lens

- coronary heart disease (a condition where there is a narrowing or blocking of the blood vessels that carry blood and oxygen to the heart)
- heart failure
- high or low blood pressure
- liver problems
- depression

Other warnings you should know about:

While you are using SIMBRINZA suspension, talk to your healthcare professional immediately if you:

- develop an eye infection, swelling, red or irritated eyelid.
- suffer any eye injury or have eye surgery.

Pregnancy

If you are pregnant, or think you may be pregnant, or are planning to have a baby, talk to your healthcare professional before you use SIMBRINZA.

Women who may become pregnant are advised to use birth control while using SIMBRINZA.

SIMBRINZA use is not recommended during pregnancy. Do not use SIMBRINZA unless your healthcare professional tells you to.

Breast-feeding

If you are breast-feeding, SIMBRINZA may pass into your milk. The use of SIMBRINZA is not recommended during breast-feeding.

Contact Lenses

There is a preservative in SIMBRINZA (benzalkonium chloride) that may be absorbed by contact lenses. It can discolour soft contact lenses and may cause eye irritation. Therefore, do not wear contact lenses while using SIMBRINZA.

Wait 15 minutes after using SIMBRINZA before putting your lenses back in.

Driving and Using Machines

Using SIMBRINZA may cause fatigue, sleepiness, drowsiness and blurred vision. Do NOT drive or engage in hazardous activities until these symptoms are gone.

Tell your healthcare professional about all the medicines you take, including any drugs, vitamins, minerals, natural supplements or alternative medicines.

There have been no specific drug interaction studies done for SIMBRINZA.

Tell your healthcare professional about all medicines, including eye drops, that you are using or plan to use, including those without a prescription.

The following may interact with SIMBRINZA:

- Other eye drops for the treatment for glaucoma
- Medicines to lower blood pressure
- Medicines used to treat heart conditions (for example, digoxin)
- Medicines used to treat glaucoma and altitude sickness (for example, acetazolamide, Methazolamide and dorzolamide)
- Medicines used to treat viral infections (antivirals)
- Medicines used to treat bacterial infections (antibiotics)
- Acetylsalicylic acid, used to treat pain, fever and inflammation
- Medicines used to treat depression such as monoamine oxidates (MAO) inhibitors, or antidepressants (for example, amitriptyline, nortriptyline, clomipramine and mianserin)
- Medicine used to treat schizophrenia like chlorpromazine
- Medicine used to treat attention deficit hyperactivity disorder (ADHD) like methylphenidate
- Medicine used to treat high blood pressure like reserpine
- Medicines used to treat fungal infections (for example, ketoconazole, itraconazole and clotrimazole)
- Medicines used to treat HIV/AIDS like ritonavir
- Medicine used to treat slow heart rates like isoprenaline
- Medicine used to treat high blood pressure like prazosin
- Opiates or opioids that are used to treat pain
- Sedatives, used to treat sleep disorders
- Anaesthetics, used to block pain

How to take SIMBRINZA:

Always use SIMBRINZA exactly as your healthcare professional has told you. Check with them if you are not sure.

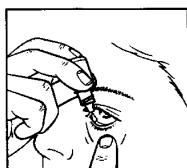
Only use SIMBRINZA for your eyes. Do not swallow or inject SIMBRINZA.

Usual dose:

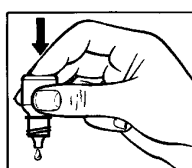
One drop in the eye or eyes, twice a day, in the morning and night. Use at the same times each day.

Only use SIMBRINZA in both eyes if your healthcare professional told you to. Take it for as long as your healthcare professional told you to.

How to use:



1



2



3

- Get the SIMBRINZA bottle and a mirror.
- Wash your hands before use
- Shake well before use.
- Twist off the bottle cap. Remove security-ring if loose.
- Hold the bottle, pointing down, between your thumb and fingers.
- Tilt your head back. Pull down your eyelid with a clean finger, until there is a 'pocket' between the eyelid and your eye. The drop will go in here (**Picture 1**).
- Bring the bottle tip close to the eye. Use the mirror if it helps.
- Do NOT touch your eye or eyelid, surrounding areas or other surfaces with the dropper.
 - If the tip of the bottle touches the eye or surrounding structures, it can become contaminated by common bacteria known to cause eye infections. It could contaminate the drops.
 - Serious damage to the eye and loss of vision may result from using contaminated eye drops.
- Gently press on the base of the bottle to release one drop of SIMBRINZA at a time.
- Do NOT squeeze the bottle: it is designed so that a gentle press on the bottom is all that it needs (**Picture 2**).
- After using SIMBRINZA, press a finger into the corner of your eye, by the nose (**Picture 3**) for up to 1 minute. This helps to stop SIMBRINZA getting into the rest of the body.
- If you use drops in both eyes, repeat the steps for your other eye.
- Close the bottle cap firmly immediately after use.
- Use up one bottle before opening the next bottle.
- Do NOT use **for more than 125 days** after opening the bottle.
- If solution changes color, do NOT use.
- Do NOT use the product after the expiration date marked on the bottle.

If a drop misses your eye, try again.

If you are using other eye drops, wait at least 5 minutes between using SIMBRINZA and the other drops.

SIMBRINZA contains a preservative called benzalkonium chloride which may discolour soft contact lenses. If you wear contact lenses, remove them before using SIMBRINZA. Wait 15 minutes after using the drops before you put your lenses back in.

Overdose:

If you use more SIMBRINZA than you should, rinse your eye(s) with warm water. Do not put in any more drops until it is time for your next regular dose.

If you think you, or person you are caring for have taken too much SIMBRINZA, contact a healthcare professional, hospital emergency department or regional poison control centre immediately, even if there are no symptoms.

Missed Dose:

If you forget to use SIMBRINZA, continue with the next dose as planned. Do not use a double dose to make up for the missed dose.

Do not use more than one drop in the affected eye(s) two times a day.

If you have any further questions on the use of this medicine, ask your healthcare professional.

What are possible side effects from using SIMBRINZA?

These are not all the possible side effects you may have when taking SIMBRINZA. If you experience any side effects not listed here, tell your healthcare professional.

If you get any severe allergic reaction while you are using SIMBRINZA, contact your healthcare professional immediately.

- allergic conjunctivitis (eye allergy)
- eye surface inflammation
- eye pain, eye discomfort
- blurred vision
- eye redness
- eye surface damage with loss of cells
- inflammation of the eyelid
- deposits on the eye surface
- sensitivity to light
- eye swelling
- dry eye
- eye discharge
- increased tear production
- eyelid redness
- abnormal sensation in eye
- tired eye
- Corneal oedema (a condition where there is a build up of fluid in the cornea)
- Corneal staining
- Miosis (a condition where there is excessive shrinking of the pupil)
- drowsiness
- bad taste in mouth
- dry mouth
- difficulty sleeping (insomnia)
- headache
- dizziness

- dry nose or throat
- runny nose
- stuffy nose
- indigestion, stomach-ache
- abnormal sensation in mouth
- Vertigo (a condition where this is a sensation of feeling off balance)
- Ectropion (a condition where the lower eyelid turns outward)
- Hypermetropia (a condition where it is hard to see up close)
- Hypoaesthesia oral (a condition where you have a loss of sensation in the mouth)
- Drug intolerance
- Medication residue
- Excoriation (a condition where you cannot stop picking at your skin)
- Balance disorder
- Hypersomnia (a condition that causes you to be very sleepy during the day)
- Hypotonia (a condition where you have low muscle tone)
- Sedation
- Libido increased
- Rhinalgia (a condition where you have pain in the nose)
- Hypertension (a condition where you have increased blood pressure)
- Ageusia (a condition where you have a loss of taste)
- Arrhythmia (a condition where you have an irregular heartbeat)
- Arthralgia (a condition where the joints are stiff)
- Asthma (a condition where your airways get narrow and swollen making it difficult to breathe)
- Chest pain
- Madarosis (a condition where you lose your eyebrows or eyelashes)
- Peripheral oedema (a condition where you have swelling in your lower legs or hands)
- Pollakiuria (a condition where you urinate frequently)
- Rhinitis (a condition where the passages in your nose are inflamed)
- Tachycardia (a condition where you have an increased heart rate)
- Tremor (a condition where your body shakes without your control)
- Iritis (an eye condition where your iris is swollen or irritated)
- Iridocyclitis (an eye condition where your iris and the muscles in your eye are inflamed)
- Syncope (a condition where you faint or pass out)

Serious side effects and what to do about them			
Symptom / effect	Talk to your healthcare professional		Stop taking drug and get immediate medical help
	Only if severe	In all cases	
UNCOMMON			
Bradycardia (heart rate decreased): fainting, dizziness, weakness, changes in breath		✓	
Hypotension (blood pressure decreased): fainting, dizziness, weakness, changes in breath		✓	
Unusual eye symptoms: severe eye pain or sensitivity to light, or sudden decrease or loss of vision		✓	
Hypersensitivity (allergic reaction): trouble breathing, tightness in throat or severe skin rash			✓
UNKNOWN			
Stevens-Johnson syndrome (severe skin rash): redness, blistering and/or peeling of the skin and/or inside of the lips, eyes, mouth, nasal passages or genitals, accompanied by fever, chills, headache, cough, body aches or swollen glands.			✓
Toxic epidermal necrolysis (severe skin reaction): redness, blistering and/or peeling of large areas of the skin.			✓

If you have a troublesome symptom or side effect that is not listed here or becomes bad enough to interfere with your daily activities, tell your healthcare professional.

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 (<https://www.canada.ca/en/health-canada/services/drugs-health-products/medeffect-canada.html>) 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.

Storage:

Store at 2°C to 25°C.

Keep out of reach and sight of children.

Do NOT use SIMBRINZA after the expiry date which is stated on the bottle and the carton after EXP.

The expiry date refers to the last day of that month.

Throw away the bottle 125 days after first opening to prevent infections; and use a new bottle.

Write down the date of opening on the carton label in the space provided.

Do not throw away any medicines via wastewater or household waste. Ask your healthcare professional how to throw away medicines you no longer use. These measures will help protect the environment.

If you want more information about SIMBRINZA:

- Talk to your healthcare professional
- Find the full product monograph that is prepared for healthcare professionals and includes this Patient Medication Information by visiting the Health Canada website: (<https://www.canada.ca/en/health-canada/services/drugs-health-products/drug-products/drug-product-database.html>); the manufacturer's website <http://www.novartis.ca>, or by calling 1-800-363-8883)

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SIMBRINZA is a registered trademark.

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