Safety and efficacy of fixed-combination travoprost/timolol in patients with open-angle glaucoma or ocular hypertension not controlled with timolol monotherapy

Marcelo Lopes da Silva Jordão¹
Marcelo Hatanaka²
Abayomi Ogundele³
Maria Rosa Bet de Moraes Silva⁴
Roberto Murad Vessani⁵

¹Faculty of Medicine of Ribeirão Preto, University of São Paulo, São Paulo, Brazil; ²University of São Paulo School of Medicine, São Paulo, Brazil; ³Global Medical Affairs, Alcon Laboratories, Inc., Fort Worth, TX, USA; ⁴Faculty of Medicine of Botucatu, Universidade Estadual Paulista (UNESP), São Paulo, Brazil; ⁵General Hospital of Itapecerica da Serra, Serviço Social da Construção Civil do Estado de São Paulo (SECONCI-SP), São Paulo, Brazil

Objective: To assess the intraocular pressure (IOP)-lowering effect of travoprost 0.004%/timolol 0.5% fixed-dose combination (TRAV/TIM–FC) in patients not achieving the target IOP of ≤18 mmHg while on timolol 0.5% (TIM) monotherapy.

Methods: A multicenter, prospective, open-label study (NCT01336569) was conducted in patients with open-angle glaucoma or ocular hypertension. Eligible patients were receiving TIM monotherapy with a screening/baseline IOP of 19–35 mmHg in ≥1 eye. TIM was discontinued on the baseline visit day (no washout period) and TRAV/TIM–FC was initiated and administered once daily at 8 pm for 4–6 weeks. The primary efficacy variable was mean change in IOP from TIM-treated baseline to study end, measured by Goldmann applanation tonometry. Results were analyzed by analysis of variance and paired samples t-test (5% significance).

Results: A total of 49 patients were enrolled (mean age, 63 [range, 42–82] years; 55.1% White; 73.5% women), and 45 were included in the intent-to-treat (ITT) population. Mean duration of treatment with TRAV/TIM–FC was 31 days. Mean ± standard deviation IOP reduction from baseline (TIM) to the follow-up visit (TRAV/TIM–FC) was −5.0±3.6 mmHg. IOP decreased significantly (P<0.0001) from baseline (22.1±2.6 mmHg) to study end (17.1±3.9 mmHg) in the ITT population, with a mean IOP reduction of 22.3%. Most patients (n=33/45; 73.3%) achieved IOP ≤18 mmHg. Two patients experienced a total of four adverse events (AEs), including a patient who reported one serious AE (enterorrhagia) that was considered unrelated to treatment, and a patient who reported one event each of drug-related redness, pruritus, and foreign body sensation. Most patients (n=47/49; 95.9%) reported no AEs.

Conclusions: TRAV/TIM–FC lowered IOP in patients who were not at target IOP while receiving TIM monotherapy, with most patients achieving an IOP ≤18 mmHg with TRAV/TIM–FC. TRAV/TIM–FC was well tolerated in this population.

Keywords: DuoTrav®, intraocular pressure, primary open-angle glaucoma, time since diagnosis

Introduction

Glaucoma and ocular hypertension are associated with elevated intraocular pressure (IOP) and progressive visual field deterioration. An analysis of published data estimated the number of people with glaucoma worldwide to be more than 60 million in 2010 and nearly 80 million by 2020, with 74% of these individuals diagnosed with open-angle glaucoma. Elevated IOP is a causal risk factor for optic nerve damage and vision loss and is one of the only readily treatable causes or symptoms of glaucoma and ocular hypertension.
Topical administration of pharmacologic IOP-lowering agents is the first line of therapy for glaucoma and ocular hypertension. β-blockers like the nonselective β-adrenergic receptor antagonist timolol maleate have historically been the first choice of treatment. Prostaglandin analogs (PGAs), such as travoprost, reduce IOP in patients with glaucoma or ocular hypertension more effectively than timolol, and PGAs are generally well tolerated. Pharmacologic therapy is typically initiated with a single agent, but many patients require one or more additional agents to maintain sufficient IOP reduction after the first year of treatment. Compared with multiple agents in individual bottles or administered at different times of day, fixed-combination medications are associated with better treatment adherence. Additionally, fixed combinations do not introduce risk of drug washout, additive exposure to preservatives, or increased treatment complexity, which can occur when multiple individual ocular hypotensive therapies are administered concomitantly. A meta-analysis of 18 clinical trials that compared the efficacy and tolerability of fixed combinations of PGAs and β-blockers versus their components in unfixed combinations or as monotherapies found that the fixed combinations resulted in lower risk of hyperemia than unfixed combinations or PGA monotherapies. Fixed combination therapies were found to be more effective in reducing IOP compared with monotherapies. Fixed-combination therapies were not as effective in reducing IOP as unfixed combinations; however, the analysis of fixed versus unfixed combinations was limited to five studies, and dosing times of the components may have been different (eg, once daily versus twice daily).

As discussed elsewhere, travoprost and timolol have complementary ocular hypotensive mechanisms, similar pharmacokinetics, and compatible physicochemical properties; furthermore, the safety and IOP-lowering efficacy of a fixed combination of travoprost and timolol has been demonstrated. Fixed-combination travoprost 0.004% (TRAV)/timolol 0.5% (TIM) is currently available in formulations preserved with benzalkonium chloride (BAK; TRAV/TIM–FC [DuoTrav®; Alcon Laboratories, Fort Worth, TX, USA]) or polyquaternium-1 (POLYQUAD® [Alcon Laboratories]). These two formulations were demonstrated to be equally effective in reducing IOP in patients with open-angle glaucoma or ocular hypertension. It is not yet clear whether disease duration influences the IOP-lowering efficacy of TRAV/TIM–FC or its components.

The objective of this study was to assess the safety and IOP-lowering efficacy of changing to BAK-preserved TRAV/TIM–FC in patients with open-angle glaucoma or ocular hypertension uncontrolled with TIM monotherapy.

Methods
Study design and treatment
This was a prospective, multicenter, phase IV, open-label, noncomparative, single-arm study (www.ClinicalTrials.gov identifier, NCT01336569) conducted at four study centers in Brazil between February 2011 and March 2012. The purpose of this study was to assess the safety and IOP-lowering efficacy of changing to TRAV/TIM–FC in patients with open-angle glaucoma or ocular hypertension not sufficiently controlled by TIM monotherapy (ie, with IOP ≥19 mmHg) and requiring additional IOP reduction. The study consisted of two visits: the screening/baseline visit (visit 1), and a follow-up visit conducted at the end of treatment, 4–6 weeks after screening (visit 2). Before participation in the current study, patients were receiving TIM monotherapy. At the conclusion of visit 1, patients discontinued TIM monotherapy and initiated once-daily TRAV/TIM–FC treatment, with no washout period. One eye from each patient was chosen as the study eye; only the study eye was used in the efficacy analysis. If only one eye received medication, that eye was the study eye; if both eyes received medication, the eye with higher IOP at visit 1 was selected as the study eye. If baseline IOP was equal in both eyes, the right eye was chosen. Patients were instructed to instill one drop of TRAV/TIM–FC in the eye(s) to be treated once per day at 8 pm for 4–6 weeks.

The study was conducted in accordance with the Declaration of Helsinki, and all study protocols and consent forms were reviewed and approved by the ethics committees of all study centers. Participating patients gave written informed consent before enrollment.

Patients
Eligible participants were aged ≥18 years and were diagnosed with primary open-angle glaucoma, pigmentary glaucoma, or ocular hypertension. Additional inclusion criteria were a stable IOP-lowering medication regimen for ≥1 week before screening; IOP considered safe in both eyes to ensure clinical stability of vision and optic nerve throughout the study; IOP between 19–35 mmHg in ≥1 eye; IOP in the non-study eye able to be controlled by either the study drug or no treatment; and best corrected visual acuity (BCVA) better than 20/200 (Snellen) or <1.0 logMAR in both eyes. Key exclusion criteria were history of allergy, hypersensitivity, or low tolerance to components of TRAV/TIM–FC; abnormalities preventing applanation tonometry or fundus/anterior chamber examination; corneal dystrophy; concomitant conjunctivitis, keratitis, or uveitis; conventional or laser surgery in either eye within
3 months before screening; progressive retinal or optic nerve disease; history or risk of uveitis or cystoid macular edema; history of *Herpes simplex* eye infection; heart conditions presenting a risk to administration of topical β-blockers (ie, sinus bradycardia, sinoatrial or atrioventricular block, heart failure, or cardiogenic shock); asthma or severe chronic obstructive pulmonary disease; severe allergic rhinitis or bronchial hyper-responsiveness; women who were pregnant, nursing, or planning to become pregnant; or participation in another clinical study within 30 days before the screening visit. Patients receiving systemic treatments that could affect IOP were required to be on a stable regimen for ≥7 days before screening without requiring a dosage change during the study period.

**Efficacy assessments**

The primary efficacy endpoint was mean IOP change from baseline (patients receiving TIM) at visit 2 (patients receiving TRAV/TIM–FC), measured at approximately the same time of day. IOP was assessed by Goldmann applanation tonometry twice consecutively at each visit; if the two measurements for the same eye differed by >2 mmHg, a third measurement was taken and the two IOP measurements closest to each other were averaged. If the three measurements differed by equal amounts, the three measurements were averaged. All IOP assessments for any one patient were performed by the same operator using the same standard tonometer. Overall IOP percent change from baseline, as well as mean IOP levels and percent IOP reduction from baseline in patients diagnosed with glaucoma for ≤4 years versus >4 years, were assessed as additional efficacy endpoints. An arbitrary treatment duration cutoff of 4 years was chosen to allow assessment of whether time since diagnosis might affect IOP response to TRAV/TIM–FC.

**Safety assessments**

Adverse events (AEs) were recorded at visit 2, or as necessary, and assessed for causality by study investigators. BCVA was measured using the Snellen visual acuity chart at visits 1 and 2; if more than one error was recorded on a given line, the results were rounded up. Slit lamp fundus biomicroscopy of both eyes, including the cornea, sclera, eyelids, conjunctiva, anterior chamber, and iris, was performed for each patient at visits 1 and 2, before IOP assessment.

**Statistical analysis**

Patient demographics and baseline characteristics were summarized using descriptive statistics. Mean IOP change from baseline was evaluated in the intent-to-treat (ITT) population (ie, all patients receiving study medication and attending visits 1 and 2) and the per-protocol (PP) population (ie, all patients receiving study medication, attending visits 1 and 2, and meeting medication compliance criteria) by paired t-tests. Efficacy endpoints, stratified by time since diagnosis, were evaluated by analysis of variance and t-test. AEs were evaluated in the safety population (ie, all patients receiving study medication) and were summarized descriptively. BCVA and biomicroscopy observations were analyzed in the safety population; biomicroscopy observations at visits 1 and 2 were compared by McNemar’s test. Statistical analyses were performed using Statistica (version 5.1/97; StatSoft, Inc., Tulsa, OK, USA) with significance established at the 5% level.

**Results**

**Patients**

A total of 49 patients were enrolled, received TRAV/TIM–FC at visit 1, and were included in the safety population. At screening, patients had a mean age ± standard deviation (SD) of 63±11 years; most (73.5%, n=36/49) were women, and 55.1% (n=27/49) were White (Table 1). Most patients (79.6%, n=39/49) were diagnosed with primary open-angle glaucoma, and all patients had IOPs ranging from 19–28 mmHg while receiving TIM. Four patients were lost to follow-up because of non-attendance at visit 2; 45 patients attended visits 1 and 2 and were included in the ITT population. Mean time ± SD on TRAV/TIM–FC treatment was 31.1±4.7 days (range, 27–50 days; median, 30 days).

**Efficacy**

Efficacy data for the ITT and PP data sets were similar; therefore, results for the ITT population are presented. Mean ± SD IOP reduction from baseline to visit 2 (primary endpoint) was −5.0±3.6 mmHg (Figure 1A); IOP decreased from 22.1±2.6 (range, 19–28) mmHg at visit 1 to 17.1±3.9 (range, 10–27) mmHg at visit 2 (P<0.0001). At visit 1, 0/45 ITT patients had IOP ≤18 mmHg; at visit 2, 33/45 (73.3%) patients had IOP ≤18 mmHg (Figure 1B). Individual IOP changes from baseline ranged from +3 to −13 mmHg (Figure 2).

Baseline IOP (mean ± SD) was not significantly different for patients diagnosed for ≤4 years (21.6±2.1 mmHg; n=19) compared with patients diagnosed for >4 years (23.5±3.4 mmHg, n=13; P=0.0585). At visit 2, mean IOP levels were significantly lower in patients diagnosed with glaucoma for ≤4 years (15.4 mmHg, n=19) compared with patients diagnosed for >4 years (19.2 mmHg, n=13; P<0.05; Figure 3A). Mean IOP reduction from baseline was 28.9% for patients diagnosed with glaucoma for ≤4 years compared with 18.4% for patients diagnosed with glaucoma...
for >4 years (Figure 3B). Overall, mean percentage IOP reduction ± SD was −22.3%±15.7%.

**Safety**

Treatment with TRAV/TIM–FC was generally well tolerated. Most patients (95.9%, n=47/49) reported no AEs. A total of four AEs, including one serious AE (SAE), were reported. One event each of redness, pruritus, and foreign body sensation was reported by the same patient; these AEs were considered related to the study drug. Pruritus was reported as severe, and redness and foreign body sensation were moderate. One patient reported enterorrhagia, which was classified as an SAE and considered unrelated to the study drug. No patient discontinued because of an AE or SAE.

### Table 1 Patient demographics and baseline characteristics (safety population)

| Demographic                        | Total n=49 |
|------------------------------------|------------|
| Age                                | 63±11 (42–82) |
| 18–65 y, n (%)                     | 30 (61.2) |
| >65 y, n (%)                       | 19 (38.8) |
| Sex, n (%)                         |            |
| Female                             | 36 (73.5) |
| Male                               | 13 (26.5) |
| Race, n (%)                        |            |
| White                              | 27 (55.1) |
| Black                              | 9 (18.4) |
| Mixed race                         | 13 (26.6) |
| Diagnosis, n (%)                   |            |
| Primary open-angle glaucoma        | 39 (79.6) |
| Pigmentary glaucoma                | 1 (2.0) |
| Ocular hypertension                | 9 (18.4) |
| Time since glaucoma diagnosis a    | 4.0±3.7 (0.02–13) |
| ≤6 y, n (%)                        | 21 (58.3) |
| >6 y, n (%)                        | 15 (41.7) |
| Baseline IOP, mean ± SD (range), mmHg | 22.2±2.6 (19–28) |
| Study eye selected, n (%)          |            |
| OD                                 | 31 (63.3) |
| OS                                 | 18 (36.7) |
| Prior ocular medication, n (%)     |            |
| Timolol b                           | 49 (100.0) |
| Brimonidine                        | 3 (6.1) |
| Brinzolamide                       | 1 (2.0) |
| Latanoprost                        | 1 (2.0) |
| Travoprost                         | 1 (2.0) |

**Notes:** a) n=36 patients; b) achieved with timolol 0.5%; c) current medication at screening.

**Abbreviations:** IOP, intraocular pressure; TIM, timolol maleate 0.5%; TRAV/TIM–FC, travoprost 0.004%/timolol maleate 0.5% fixed combination; SD, standard deviation; n, number of patients.
Distribution of BCVA was similar between visits 1 and 2. The percentage of patients with BCVA better than 20/80 was unchanged between visits (visit 1, n=42/49 [85.7%]; visit 2, 39/45 [86.7%]). Biomicroscopy observations were similar at visits 1 and 2 (Table 2). The only statistically significant finding for this safety endpoint was redness of eyelids or conjunctiva, which was observed in 24.4% of patients at visit 2 compared with 2.0% of patients at visit 1 (P=0.0026).

**Table 2** Biomicroscopy observations by visit (safety population)

| Parameter, n (%)                      | Visit 1 | Visit 2 |
|---------------------------------------|---------|---------|
|                                      | (n=49)  | (n=45)  |
| Cornea                                |         |         |
| Normal                                | 48 (98.0) | 43 (95.6) |
| Subepithelial infiltrate              | 1 (2.0)  | 1 (2.2)  |
| Discrete opacities                    | 0        | 1 (2.2)  |
| Irised anterior chamber               |         |         |
| Normal                                | 47 (95.9) | 43 (95.6) |
| Iridectomy                            | 2 (4.1)  | 2 (4.4)  |
| Lens                                  |         |         |
| No opacity in the lens                | 26 (53.1) | 26 (57.8) |
| Any opacity in the lens               | 15 (30.6) | 12 (26.7) |
| Pseudophakia                          | 8 (16.3) | 7 (15.6) |
| Eyelids and conjunctiva               |         |         |
| Normal                                | 38 (77.6) | 28 (62.2) |
| Temporal pingeueula                   | 2 (4.1)  | 1 (2.2)  |
| Pterygium/nasal pterygium             | 4 (8.2)  | 4 (8.9)  |
| Dermatochalasis                       | 3 (6.1)  | 2 (4.4)  |
| Filtering blebs                       | 1 (2.0)  | 1 (2.2)  |
| Redness                               | 1 (2.0)  | 11 (24.4) |
| Palpebral pigmentation                | 0        | 1 (2.2)  |

**Notes:** Baseline; ‘4- to 6-week follow-up; ‘McNemar’s test, χ²=9.09, P=0.0026.

**Abbreviation:** n, number of patients.
TRAV/TIM–FC was well tolerated. A total of three non-serious AEs were considered related to study treatment, and 96% of patients reported no AEs throughout the study. The most frequent change from baseline in biomicroscopy observations was an increase in redness, which was observed in 2% of patients at visit 1 and 24% of patients at visit 2.

The findings of this study are consistent with previous reports of the IOP-lowering safety and efficacy of TRAV/TIM–FC. In prior studies, as in the current work, redness or hyperemia was the most frequently observed AE associated with TRAV/TIM–FC and was reported in approximately 6%–14% of patients; TRAV/TIM–FC was generally well tolerated and associated with minimal changes in ocular signs and biomicroscopy observations.18–21 In three non-interventional, open-label, multicenter studies, patients with glaucoma or ocular hypertension receiving once-daily TRAV/TIM–FC for 4–6 weeks achieved significant IOP reductions of 4.6–5.5 mmHg, or 21%–25%, from TIM-treated baseline.19,20,22 Furthermore, two double-masked, randomized, active-controlled, parallel-group trials demonstrated that, after 6 weeks of treatment, once-daily TRAV/TIM–FC reduced IOP to a significantly greater extent than twice-daily TIM.18,21 In the current study, IOP change from baseline ranged from +3 mmHg to −13 mmHg with TRAV/TIM–FC. Most patients (n=36/45, 80%) achieved IOP reduction of at least 2 mmHg from baseline TIM-treated levels. Seven patients experienced an IOP change from baseline of 0 or 1 mmHg, suggesting that these patients may not have been responsive to TRAV. IOP was increased from baseline in two patients (+2 mmHg and +3 mmHg, respectively); these patients may have been non-responsive to TRAV and experienced increased IOP because of a potential reduction in TIM dosing (ie, twice-daily before enrollment versus once-daily during the study). However, because TIM dosing schedules prior to study enrollment were not evaluated at screening, support for this hypothesis is limited.

Interestingly, in the current study, patients diagnosed with glaucoma or ocular hypertension for ≤4 years achieved significantly greater IOP reductions compared with patients diagnosed for >4 years. IOP reductions from baseline were nearly 30% in patients who had been diagnosed within 4 years, but were <20% in those who were diagnosed over 4 years. Baseline IOP in patients diagnosed for >4 years was slightly higher than in those diagnosed for ≤4 years; however, this difference was of small magnitude and was not statistically significant. Taken together, these data suggest that patients switched to TRAV/TIM–FC therapy earlier may respond better to treatment.

Previous studies have shown that TRAV/TIM–FC is also more effective than TRAV monotherapy in reducing IOP; furthermore, the two drugs have similar safety profiles.18,23,24 In a prospective, open-label, historical-controlled, single-arm trial of patients in Brazil who were switched from PGA monotherapy (travoprost, bimatoprost, or latanoprost) to TRAV/TIM–FC because of insufficient IOP reduction, 4 weeks of TRAV/TIM–FC reduced IOP by an additional ~19% from baseline values maintained with PGA monotherapy.23 Similarly, in a randomized, double-masked, 8-week trial of TRAV/TIM–FC versus TRAV, TRAV/TIM–FC maintained significantly lower mean IOP than TRAV, with less 24-hour fluctuation.24 These findings are supported by the results of a meta-analysis that demonstrated greater IOP-lowering efficacy of PGA/β-blocker fixed combinations compared with monotherapy with one of their components.13 Compared with fixed and unfixed combinations of latanoprost/TIM, TRAV/TIM–FC produced similar IOP reduction from baseline25–27 with significantly lower mean IOP 24 hours after dosing.27 Treatments had similar safety profiles. Hyperemia has been observed with TRAV, TRAV/TIM–FC, and fixed and unfixed combinations of latanoprost/TIM; in these studies, rates of hyperemia with TRAV/TIM–FC varied from 2.5%–15%.24,26,27 Additional studies are needed to establish hyperemia rates of fixed-combination, polyquaternium-1-preserved TRAV/TIM compared with TRAV/TIM–FC. It has been hypothesized that BAK-free TRAV/TIM–FC may have less ocular surface toxicity without compromising IOP-lowering efficacy and therefore may be a preferred option for patients with ocular surface disease.14

The relatively small study population and single-arm, historical-control design are potential limitations of the current study. Compliance with TIM treatment before study initiation may have been lower than with TRAV/TIM–FC during the study period because of potential effects of participation in a clinical trial.28 Typical patient adherence to glaucoma treatment has been low; depending on the stringency of the definition of “non-adherent” used, rates of non-adherence to ocular hypotensive medication have ranged from 5%–80%.29 Additional studies are needed to investigate the effect of increasing duration after diagnosis on magnitude of IOP reduction with TRAV/TIM–FC.

**Conclusion**

This study demonstrated the safety and IOP-lowering efficacy of changing from TIM monotherapy to TRAV/TIM–FC in
patients with open-angle glaucoma or ocular hypertension uncontrolled with TIM. A significant decrease in mean IOP was observed after 4–6 weeks of treatment, demonstrating that TRAV/TIM–FC was effective in controlling IOP for patients with glaucoma who could not achieve target IOP with TIM monotherapy alone. The magnitude of IOP reduction was greater in patients diagnosed with glaucoma for ≤4 years compared with patients diagnosed for >4 years. TRAV/TIM–FC was generally well tolerated, with only four AEs reported in two patients. No differences in BCVA were observed between visits 1 and 2. The only safety measure that was significantly more frequent at visit 2 (with TRAV/TIM–FC treatment) versus visit 1 (with TIM treatment) was redness of the eyelids or conjunctiva.

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