Clinical Outcomes After Second-Generation Trabecular Microbypass Stents (iStent inject®) with Phacoemulsification in Korean Patients

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ABSTRACT

Introduction: To evaluate the intraocular pressure (IOP)-lowering effect of second-generation trabecular microbypass stents (iStent inject®) with cataract extraction (combination group) and compare refractive changes in the combination group and the control (phacoemulsification only) group.

Methods: This retrospective case–control study included 36 eyes with cataract and medically controlled open-angle glaucoma with IOP < 21 mmHg and 100 nonglaucomatous eyes with cataract. Data were collected preoperatively and for 6 months postoperatively. Data included IOP, number of glaucoma medications, corrected distance visual acuity, and mean absolute error (MAE) from target refraction, and astigmatic vector analysis. Surgical success for the combination group was defined according to three criteria: (A) IOP < 15 mmHg without medication, (B) IOP < 18 mmHg with or without medication, and (C) IOP < 18 mmHg without medication.

Results: In the combination group, mean IOP was reduced from 15.1 ± 2.9 mmHg to 12.5 ± 2.0 mmHg, and the mean number of medications decreased from 1.9 ± 1.0 to 0.4 ± 0.8 at postoperative 6 months (both P < 0.001). Surgical success rates were 77.8%, 83.3%, and 97.2% at 6 months by criteria A, B, and C, respectively. Mean IOP was reduced from 14.3 ± 2.7 mmHg to 13.1 ± 2.1 mmHg at 1 month in the control group (P < 0.001). The MAE was 0.33 ± 0.26 D, and 83.3% of eyes had spherical equivalent difference within 0.50 D in the combination group (0.38 ± 0.33 D and 76.0% in the control group; P = 0.309 and P = 0.363, respectively). Preoperative and postoperative centroid values were 0.51 D @ 176° and 0.66 D @ 178°, respectively (0.23 D @ 176° and 0.66 D @ 1° in the control group). There were no statistical differences between the two groups with respect to preoperative and postoperative mean absolute values (P = 0.154 and P = 0.322, respectively).

Conclusions: On the basis of our results using Korean real-world interim experience, iStent inject with cataract extraction has favorable IOP-lowering effects and minimally impacts refractive outcomes.
Although many previous studies have demonstrated favorable efficacy of second-generation trabecular microbypass stents (iStent® inject) in patients with high intraocular pressure (IOP), there have been no published studies in Korean patients.

Moreover, few studies comparing refractive outcomes after iStent® inject with cataract surgery (combination procedure) to cataract surgery alone have been conducted.

At the 6-month follow-up, 77.8% of eyes had IOP < 15 mmHg without medication, 83.3% of eyes had IOP < 18 mmHg without medication, and 83.3% of eyes were medication-free (vs. mean 1.9 medications preoperatively).

The mean absolute error was 0.33 ± 0.26 D, and 83.3% of eyes had spherical equivalent difference within 0.5 D.

Trabecular microbypass stent (iStent® inject) with cataract extraction affords beneficial reductions in IOP and glaucoma medications and excellent refractive outcomes without severe refractive surprise in our interim experience in Korea.

INTRODUCTION

Glaucoma is one of the leading causes of blindness worldwide [1]. Many randomized clinical trials have demonstrated that intraocular pressure (IOP) is the primary modifiable risk factor for “diagnosis” and “disease progression” of glaucoma [2–4]. Thus, the focus of glaucoma treatment aims to reduce IOP, which is the only treatment strategy proven to reduce its progression [2, 4–6].

Although standard glaucoma surgeries, including trabeculectomy and glaucoma drainage devices (GDD), have favorable IOP-lowering efficacy, they also have risks of serious vision-threatening complications including hypotony, inflammation, bleb-related complications, and loss of vision [5, 7]. The goal of glaucoma treatment is to maintain visual function and quality of life at a sustainable cost [8]. In this context, the number of micro-invasive glaucoma surgery (MIGS) procedures has increased because they offer safer interventional glaucoma treatment earlier in the course of the disease [6, 9–11]. In particular, canal-based MIGS provides advantages such as quicker visual recovery and sparing of conjunctival tissue for future glaucoma surgeries [10, 12–14].

The single trabecular microbypass stent with an internal diameter of 120 μm (iStent®) and the second-generation iStent® inject (containing two stents, each with an internal diameter of 80 μm) were designed to allow direct aqueous flow into Schlemm’s canal, bypassing the trabecular meshwork [12, 13]. Both devices have been studied in a variety of surgical treatment approaches, including stand-alone and cataract combination treatment; with single or multiple stents; and in the spectrum of glaucoma severities from mild to advanced stages, or even failed trabeculectomy [12, 15–19]. Although many previous studies have demonstrated favorable efficacy in patients with uncontrolled IOP or high IOP, studies in patients with normal tension glaucoma are limited [13, 14] and no studies have been published on second-generation trabecular microbypass stents in Korean eyes [15]. In addition, many studies with conventional glaucoma surgeries, such as trabeculectomy and glaucoma drainage device surgery, have shown decreased axial length, significant astigmatic changes, and increased refractive surprise [20–22]. Few studies, however, focused on refractive outcomes after phaco-combined surgery using iStent® or iStent® inject (combination group) [9, 23].

The purposes of this study are (1) to analyze IOP-lowering effects after combination
procedures in primary open-angle glaucoma (POAG) and (2) to analyze and compare refractive outcomes between combination procedures in a glaucoma group and phaco-only procedures in a control group without glaucoma.

METHODS

Patients

This was a single-center, retrospective, case control study conducted at Daegu Veterans Health Service Medical Center. This comparative study on refractive outcomes consisted of 36 patients undergoing second-generation trabecular microbypass stent with phacoemulsification and 100 nonglaucomatous patients who underwent cataract surgery only from September 2020 to January 2021. The study protocol was approved by the Institutional Review Board of Daegu Veterans Health Service Medical Center. All participants provided signed informed consent. The study adhered to the tenets of the Declaration of Helsinki. The inclusion and exclusion criteria were similar to those in our previous study [17].

Inclusion criteria for the combination group were (1) IOP $< 21$ mmHg with anti-glaucoma medications, (2) presence of cataract, and (3) need for additional reduction of IOP (poor compliance, drug allergy, or disease progression despite use of ocular hypotensive medications). The first eye undergoing the procedure was chosen for cases of bilateral disease. The cataract surgery control group included 100 consecutive eyes without severe ocular disease, such as glaucoma, macular degeneration, or ocular surface diseases, and that underwent phacoemulsification cataract surgery.

Exclusion criteria for the combination group included (1) eyes with less than 6 months of follow-up, (2) patients who underwent another procedure at the same time, (3) angle-closure glaucoma or secondary glaucoma such as inflammatory glaucoma or neovascular glaucoma, or (4) history of previous glaucoma surgery, such as trabeculectomy or glaucoma drainage device implantation.

Baseline Examination

All patients underwent comprehensive ophthalmic examinations, including best corrected visual acuity, Goldmann applanation tonometry, gonioscopy, dilated fundus examination, and retinal nerve fiber layer (RNFL) measurement using spectral domain optical coherence tomography (SD-OCT; Cirrus6000, Zeiss, Jena, Germany). Automated perimetry was performed using a Humphrey Visual Field Analyzer (740i, 840, Carl Zeiss Meditec Inc., Dublin, CA).

Glaucomatic visual field defects were defined as follows: (1) those corresponding to the optic nerve head or RNFL changes; (2) glaucoma hemifield test (GHT) results outside of the normal limits; and (3) a cluster of three or more non-edge contiguous points not crossing the horizontal meridian, with $P$ value < 0.01. The Hodapp-Anderson criterion was used to classify the severity of glaucoma [24].

Biometry was performed using an IOL Master 500 instrument (Zeiss, Jena, Germany). On the basis of surgeon choice, the SRK/T power calculation formula was used to determine the postoperative refraction target.

Surgical Technique

All surgeries were performed by a single glaucoma specialist (S.H.L.) who completed a wet-lab training program and glaucoma fellowship. Briefly, under topical anesthesia, phacoemulsification was performed with a 2.4 mm keratome. A single-piece, hydrophobic, acrylate, intraocular lens was inserted in both groups. Then, trabecular microbypass stents (iStent® inject) were implanted after routine cataract procedures. The iStent® inject is comprised of two, preloaded, heparin-coated, biocompatible, titanium stents in a single-use injector. Each stent was inserted ab interno through the nasal trabecular meshwork into Schlemm’s canal using a Swan Jacob goniprism. A second stent was placed 2–3 hours away from the first stent. The ophthalmic viscodevice (OVD) was removed, and the eye was filled with a balanced salt solution.
Postoperative Medications and Follow-up

Postoperatively, topical 0.5% gatifloxacin and 0.1% fluorometholone acetate eye drops were prescribed four times daily for 1 week and then tapered according to the resolution of inflammation. Topical bromfenac 0.07% for 4 weeks also was prescribed.

Each patient’s best corrected visual acuity (BCVA), IOP, number of anti-glaucoma medications, and complications were recorded at 1 day, 1 week, 1 month, 2 months, 3 months, and 6 months, in accordance with the World Glaucoma Association’s Guideline [25]. Refractive data were collected 4 weeks postoperatively. The postoperative medication dose was increased if there was a clinical judgment that the IOP level would likely cause progression of glaucoma. In general, glaucoma medication was started if IOP exceeded 18 mmHg, because the Advanced Glaucoma Intervention Study (AGIS) demonstrated that progression was delayed when IOP was maintained below 18 mmHg [4].

Data Analysis

Surgical successes based on IOP control were defined according to three criteria: (A) IOP < 15 mmHg without medication; (B) IOP < 18 mmHg without medication; and (C) IOP < 18 mmHg with or without medication. Fixed combination medications were reported as two medications.

The spherical equivalent (SE) refractive outcome was calculated as the difference between the postoperative SE and target SE and divided into the percentage within 0.25, 0.50, and 1.00 D of the target. Astigmatism outcomes were measured using the American Society of Cataract and Refractive Surgery Astigmatism Double Angle Plot Tool version 1.3.2 [26]. This tool was used to analyze preoperative corneal astigmatism and postoperative refractive astigmatism (https://ascrs.org/tools/astigmatism-double-angle-plot-tool).

The sample size calculation (non-inferiority, dichotomous outcomes) was confirmed on the basis of the following assumptions: type I error rate, α = 0.05; power, 1 − β = 0.80; ratio of case to control, 1:3; drop rate = 10%; proportion of the outcome (SE from target < 1 D) in the control group = 0.85 (considering The National Health Service data) [27, 28]; and margin on risk difference scale = 0.20. As a result, calculated sample sizes were 33 cases and 99 controls.

Statistical analyses were performed using SPSS for Windows software (version 18.0, SPSS Inc., Chicago, IL). A paired t test was used to determine the significance of the mean change in IOP from baseline to 1, 3, and 6 months. Independent t tests and chi-square tests were used to compare the control group. P < 0.05 was defined as statistical significance.

RESULTS

Baseline Characteristics

A total of 36 eyes from 36 patients with open-angle glaucoma who underwent concomitant cataract surgery with two iStent® inject were analyzed retrospectively. Four of the initially enrolled 40 eyes were excluded: two eyes with missing postoperative refraction and two eyes with suboptimal implantation (one eye with under-implantation, one eye with over-implantation). We excluded the suboptimal cases to minimize the confounding factors (corneal properties, refractive outcomes, and IOP). Table 1 summarizes the demographic data of the participants. The mean age was 74.4 ± 4.1 years in the combination group and 73.9 ± 4.9 years in the control group (P = 0.554). Preoperative baseline IOP was 15.1 ± 2.9 mmHg on 1.9 ± 1.0 glaucoma medications in the combination group and 14.3 ± 2.7 mmHg on no glaucoma medication in the control group (P = 0.132). Most of the participants in the combination group had mild-to-moderate OAG, and the remaining eight eyes (22.2%) had severe disease. Two eyes had previously undergone selective laser trabeculoplasty (SLT). None of the patients in the treatment group had previously undergone incisional glaucoma surgery.
Intraocular Pressure Reduction

In the combination group, significant IOP reduction was observed at postoperative 1 week, 1 month, 3 months, and 6 months compared to the preoperative baseline IOP. Compared to the baseline mean IOP (15.1 ± 2.9 mmHg), the mean IOP reduction was 13.5 ± 4.2 mmHg at 1 week, 12.6 ± 3.0 mmHg at 1 month, 12.2 ± 2.4 mmHg at 3 months, and 12.5 ± 2.0 mmHg at 6 months in the combination group (P = 0.011, P < 0.001, P < 0.001, P < 0.001, respectively). Regarding hypotensive medication use, statistically significant reductions were found during the entire study period (all P < 0.001). In summary, IOP decreased by 2.6 mmHg (17.2%) at 6 months and the mean number of medications decreased from 1.9 ± 1.0 to 0.4 ± 0.9 at 1 month, 0.3 ± 0.8 at 3 months, and 0.4 ± 0.9 at 6 months in the combination group. These results are summarized in Fig. 1. The mean IOP

| Table 1 Demographic and baseline patient characteristics |
|---------------------------------------------------------|
| Combination group (N = 36)                              | Control group (N = 100) | P value |
| Trabecular microbypass stents (iStent® inject) with cataract extraction | Phacoemulsification alone |
| Age (years)    | 74.4 ± 4.1 | 73.9 ± 4.9 | 0.554 |
| Eye (right/left) | 18/18      | 48/52      | 0.848 |
| Preoperative IOP | 15.1 ± 2.9 mmHg | 14.3 ± 2.7 mmHg | 0.132 |
| Proportion of preoperative IOP |                      |          |
| < 15 mmHg | 20 (55.6%) | 59 (59.0%) | 0.844 |
| < 18 mmHg | 28 (77.8%) | 85 (85.0%) | 0.314 |
| < 21 mmHg | 36 (100%) | 100 (100.0%) | 1.000 |
| Visual field |                        |          |
| Mean deviation (dB) | −8.67 ± 6.67 (range 0.48–25.00) | NA |
| Pattern standard deviation (dB) | 6.41 ± 3.41 (range 1.98–13.09) | NA |
| Visual field index (VFI, %) | 76.90 ± 22.33 (range 23–98) | NA |
| Optical coherence tomography |                                |          |
| Retinal nerve fiber layer thickness (μm) | 73.47 ± 12.75 (range 48–101) | NA |
| Vertical cup-to-disc ratio | 0.79 ± 0.09 (range 0.61–0.94) | NA |
| Glaucoma stage |                      |          |
| Early | 15 (41.7%) | NA |
| Moderate | 13 (36.1%) | NA |
| Advanced | 8 (22.2%) | NA |

Values in the table are presented as mean ± standard deviation (SD) or as a percentage. IOP intraocular pressure, NA Not applicable
was 14.3 ± 2.7 mmHg at baseline, 13.3 ± 3.0 mmHg at 1 week, and 13.1 ± 2.1 mmHg at 1 month in the control group (%0.001, %0.001, respectively). Proportional Analysis of IOP Reduction

A greater proportion of eyes with lower IOP at follow-up vs. baseline was observed in the combination group (Fig. 2). The proportion of patients was 77.8% for criterion A (versus 55.6% of patients with preoperative IOP < 15 mmHg, %0.046), 83.3% for criterion B (versus 77.8% with preoperative IOP < 18 mmHg, %0.551), and 97.2% for criterion C (versus 77.8% with preoperative IOP < 18 mmHg, %0.028). Moreover, 83.3% of the combination patients were medication-free at 6 months.

Refractive Outcome

The corrected distance visual acuity (CDVA) significantly improved from 0.47 ± 0.21 logMAR to 0.06 ± 0.07 logMAR at 1 month in the combination group (%0.001). Mean error is an unreliable value because it can be positive or negative values and, therefore, might be affected by outliers. Thus, we used the mean absolute error (MAE) from a refractive target. Regarding MAE from a refractive target using spherical equivalent (SE) outcomes, 44.4% (16 of 36 eyes) had SEs within 0.25 D (47% in the control group, %0.792), 83.3% had SEs within 0.50 D (76% in the control group, %0.363), and 97.2% had SEs within 1.00 D of the target (97% in the control group, %1.000). The MAE was 0.33 ± 0.26 D in the combination group and 0.38 ± 0.33 D in the control group (%0.309).

Figure 3 shows the magnitude of preoperative corneal astigmatism and postoperative refractive astigmatism. For magnitude of astigmatism at postoperative 1 month, 64% of participants were within 1.00 D, 83% in 1.5 D, and 94% within 2.0 D in the combination group compared to 70%, 86%, and 99%, respectively.
in the control group. There were no significant differences between the combination and control groups ($P = 0.328$, $0.652$, and $0.111$, respectively).

Figure 4 demonstrates the double-angle plot showing the preoperative and postoperative astigmatic vectors. The preoperative centroid was $0.51 \text{ D at } 168^\circ$ with a mean absolute $1.02 \pm 0.66 \text{ D}$, and the postoperative centroid was $0.66 \text{ D at } 178^\circ$ with a mean absolute $1.06 \pm 0.54 \text{ D}$ in the combination group. The preoperative centroid was $0.23 \text{ D at } 176^\circ$ with a mean absolute $0.85 \pm 0.49 \text{ D}$, and the postoperative centroid was $0.66 \text{ D at } 1^\circ$ with a mean absolute $0.97 \pm 0.81 \text{ D}$ in the control group. There were no statistical significant differences between the two groups with respect to preoperative mean absolute and postoperative mean absolute. ($P = 0.164$, $P = 0.322$, respectively).

**Safety**

There were no severe intraoperative complications that required additional procedures. The most frequently reported side effects were hyphema in three eyes (8.3%) and transient IOP spike ($\text{IOP} > 25 \text{ mmHg}$) in two eyes (5.6%) within the first week. No vision-threatening adverse events, such as endophthalmitis, hypotony, choroidal detachment, or corneal decompensation, occurred.

**DISCUSSION**

In the present study, the combination of two second-generation trabecular microbypass stents (iStent® inject) and phacoemulsification demonstrated favorable IOP control, including mean IOP reduction and a significant decrease in number of medications. Our study showed a mean IOP reduction of $2.6 \text{ mm Hg}$ (17.2%) at month 6 without washout periods. Furthermore
77.8% of eyes maintained an IOP < 15 mmHg, and 83.3% of eyes maintained an IOP < 18 mmHg without medication. A total of 97.2% of eyes had an IOP < 18 mmHg with or without medication. Moreover, 83.3% of eyes were medication-free (all medication was
eliminated). These results were similar to those of previous studies.

Guedes and Gravina [12] reported that mean IOP decreased from 17.3 ± 3.0 mmHg on 2.3 medications to 12.7 ± 1.8 mmHg on 0.4 medications in those with the iStent® inject at 6 months. In their study, proportional analysis showed that the percentage of eyes with IOP < 18 mmHg, < 16 mmHg, and < 14 mmHg was 100.0%, 88.6%, and 71.4%, respectively.

Fig. 4 Double-angle plot showing preoperative and postoperative astigmatic vectors, mean, and spread in eyes with trabecular microbypass stent (iStent® inject) implantation with cataract extraction (a) and in the control group (b) using the American Society of Cataract and Refractive Surgery’s Astigmatism Double Angle Plot Tool version 1.3.2.
Manning reported IOP reduction from 20.4 ± 5.6 mmHg on 1.3 ± 0.9 medications to 14.1 ± 2.1 mmHg on 0.1 ± 0.3 medications. He also observed 92.5% of patients had IOP ≤ 18 mmHg, and 71.6% of patients had IOP ≤ 15 mmHg at 12 months [16]. Our baseline IOP (15.1 mmHg) was somewhat lower than that reported in the aforementioned studies (17.3 and 20.4 mm Hg, respectively) because we only included eyes with preoperative IOP less than 21 mmHg with medication, and Asian eyes involve a higher proportion of normal-tension glaucoma. Given the well-known pattern of less dramatic IOP reduction in eyes with lower baseline IOP, our result that IOP reduction was less than in previous studies of iStent® inject with phacoemulsification in patients with predominantly high-tension primary open-angle glaucoma (POAG) is reasonable. Most previous studies regarding the efficacy of iStent implant of iStent® inject included patients with uncontrolled IOP or mean medicated IOP above 21 mmHg, and there are limited studies on outcomes in controlled glaucoma [10, 11, 13, 17]. The majority of open-angle glaucoma cases in Korea and Japan are low-tension or normal-tension glaucoma. To the best of our knowledge, the present study is the first study that describes the clinical efficacy (IOP-lowering effect and refractive changes) of iStent® inject in a South Korean population. Thus, our results have strength in management of glaucoma in subjects in East-Asians, who are considered to have lower baseline IOP with or without medications compared to other populations.

Considering these factors, our results are similar to the study by Salimi and colleagues [13]. In their 1-year multicenter study, they reported that postoperative IOP declined by 22% from 15.8 mmHg on 1.5 ± 1.28 medications to 12.3 mmHg on 0.4 ± 0.86 medications [13]. They also reported that all eyes (100%) measured ≤ 18 mmHg postoperatively (versus 74% preoperatively), and 92% had IOP ≤ 15 mmHg. In addition, 73% of the eyes were medication-free at 1 year. In summary, the results from previous studies and the present study demonstrated a significant reduction in medication use, and a significant proportion of patients ceased their medication after combination procedures [12, 13, 18]. These results are valuable in the context of disease progression, quality of life, secondary ocular surface disease, medication costs, and psychological aspects [9, 10, 15, 23].

The refractive impact of MIGS-associated phacoemulsification (combination procedure) is expected to be minimal; however, there are a limited number of studies on refractive outcomes [9]. Regarding spherical equivalent (SE) outcomes, 44.4% had SEs within 0.25 D, 83.3% had SEs within 0.50 D, and 97.2% had SEs within 1.00 D of the target in combination group in our data. There was no statistical difference between the combination and control groups (P = 0.792, P = 0.363, and P = 1.000, respectively). Our results are favorable and similar to those of previous studies regarding phaco-iStent® inject and those of a large cohort registry in standard cataract surgery. Scott et al. reported 80% and 95% of eyes within 0.50 D and 1.00 D, respectively, after iStent® implant and cataract surgery [9]. Recently, Ioannidis et al. reported that 73.8% of eyes had SEs ≤ 0.50 D and 98.9% of eyes had SEs ≤ 1.00 D [23]. The National Health Service data suggested a benchmark standard of 55% within 0.50 D and 85% within 1.00 D; 62.3% of patients within 0.50 D of the refractive target should represent a minimum level of efficiency following cataract surgery [27, 28]. MAE was 0.33 ± 0.26 D in our results, a finding similar to that of Ioannidis et al. (0.36 ± 0.25 D) [23].

For the magnitude of astigmatism at postoperative 1 month, there were no statistical significant differences in the proportions of ≤ 1.00 D, ≤ 1.50 D, and ≤ 2.00 D between both groups (all P values > 0.05). Moreover, the double-angle plot demonstrated that the postoperative centroid in the combination group was similar to that in the control group (centroid, 0.66 @ 178° ± 1.00 D vs. 0.66 D @ 1° ± 0.87 D; mean absolute, 1.06 ± 0.54 vs. 0.97 ± 0.51 D, P = 0.322). These results were similar to those of a previous phaco-iStent® implant study by Scott et al. (75% within 1.00 D, 87% within 1.50 D, 89% within 2.00 D; centroid, 0.8 D @ 92° ± 1.05 D; mean absolute, 0.8 D) [9]. Moreover, our results demonstrate

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that the difference between preoperative centroid and postoperative centroid was not clinically significant in the combination group (0.51 D @ 168° ± 1.12 D vs. 0.66 @ 178° ± 1.00 D).

Because astigmatism has both magnitude and direction, and the analysis is complicated [29], there are two widely used categories of information: (1) certain scalar values: mean ± SD, independent of angle, and percentages of eyes within certain ranges; and (2) vector analysis. However, vector of astigmatism is actually bidirectional because the axis or meridian is made up of two semi-meridians [29, 30]. As a result, certain univariate analyses of astigmatism are unpredictably erroneous and misleading. In these contexts, recommended vector analyses should include double-angle plots with centroids and confidence ellipse of preoperative and postoperative astigmatism, along with means and standard deviations of these vector magnitudes [30]. Considering this background, we analyzed (1) the mean absolute error, (2) proportion within certain ranges, and (3) double-angle plot analysis.

The results of this study suggest that combination procedures—iStent® inject with phacoemulsification—are a good option in maintaining refractive outcomes with minimal risks for unanticipated refractive surprise. Our results add to the literature supporting the minimal impact of MIGS devices on refractive outcomes and surgically induced astigmatism effects [9, 10, 23].

This study had some limitations. First, there was no washout period because of the retrospective design. We could not explain the exact proportion of normal-tension glaucoma because most patients were referred with glaucoma medications and we could not compare the long-term IOP changes in both groups. Thus, the “case–control” aspect of this study was limited on refractive outcomes. Second, cataract surgery itself could strengthen the IOP-lowering efficacy as observed in the Ocular Hypertension Treatment Study (OHTS) study [31]. Third, the study had a small sample size. Despite these drawbacks, this is the first study evaluating IOP-lowering efficacy and refractive outcomes in Korean eyes. Despite the lack of a washout period, the results represent real-world clinical outcomes [12]. Moreover, Baek and colleagues reported a mean IOP reduction of 0.78 mmHg at 1 year after cataract surgery in patients with normal tension glaucoma, suggesting that the IOP-lowering effects of cataract surgery itself might be limited in low-tension or medically controlled glaucoma given that eyes in this study showed a greater IOP response with iStent inject [32]. Future prospective studies with larger sample sizes would be valuable to confirm our results.

**CONCLUSIONS**

Combination procedures with second-generation trabecular microbypass stents seem to be effective in lowering IOP and do not compromise refractive outcomes. On the basis of our favorable results using Korean real-world clinical data, combination procedures have modest IOP-lowering effect and ability to reduce the glaucoma drug burden with no serious complications, and minimally impact the refractive outcome in patients with medically controlled glaucoma and coexisting cataracts.

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Compliance with Ethics Guidelines. The study protocol was approved by the Institutional Review Board of Daegu Veterans Health Service Medical Center. All participants provided signed informed consent, and this study adhered to the tenets of the 1964 Declaration of Helsinki.

Data Availability. The datasets generated and/or analyzed during the current study are available from the corresponding author on reasonable request.

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