iStent inject Trabecular Micro-Bypass with or Without Cataract Surgery Yields Sustained 5-Year Glaucoma Control

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Received: November 7, 2021 / Accepted: December 23, 2021 / Published online: February 3, 2022
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ABSTRACT

Introduction: This study evaluated the 5-year effectiveness and safety of iStent inject® trabecular micro-bypass with or without cataract surgery (Combined or Standalone, respectively) in patients with open-angle glaucoma (OAG).

Methods: This prospective longitudinal case series included consecutive iStent inject cases from a single surgeon at a large German academic hospital. Intraocular pressure (IOP), medications, safety, and indicators of disease stability through 5 years were assessed in the Overall cohort and in subgroup analyses stratified by usage (Combined or Standalone).

Results: Preoperative mean IOP in the Overall cohort (n = 125) was 23.5 ± 6.2 mmHg on 2.68 ± 1.02 mean medications, reducing to 14.1 ± 1.8 mmHg on 0.77 ± 0.82 medications at 5 years (40% and 71% reductions, respectively; both p < 0.001). All but 1 eye (>99%) were on medication(s) preoperatively, but 46% were medication-free at 5 years (p < 0.001). In Combined eyes (n = 81), mean IOP decreased by 39% (22.6 mmHg to 13.8 mmHg, p < 0.001) and medications by 69% (2.52 to 0.78, p < 0.001). In Standalone eyes, mean IOP reduced by 42% (25.3 mmHg to 14.6 mmHg, p < 0.001) and medications by 75% (2.98 to 0.74, p < 0.001). At final follow-up, 83% of eyes had achieved ≥ 20% IOP reduction, and all but 1 eye (>99%) had the same or lower IOP versus preoperative; all eyes (100%) maintained or reduced their medication burden versus preoperative. Favorable safety included 0 intraoperative complications and 0 filtration surgeries through 5 years. Long-term indicators of disease stability (visual fields, retinal nerve-fiber layer thickness, and cup:disc ratio) were unchanged over the course of 5-year follow-up.

Conclusions: iStent inject produced significant and durable 5-year reductions in IOP (nearly 10-mmHg reduction) and medications (nearly 2-medication reduction), with stable disease parameters over time. Combined and Standalone subgroups had similar outcomes.

Keywords: Microinvasive glaucoma surgery (MIGS); iStent inject; Trabecular bypass; Cataract; Long-term; Intraocular pressure; Second-generation; Stent
Key Summary Points

Why carry out this study?
Safe and efficacious long-term surgical treatments are needed to prevent or minimize vision loss from glaucoma, a major cause of global blindness.

This prospective study assessed real-world 5-year outcomes following implantation of two second-generation trabecular micro-bypass stents either with or without concomitant cataract surgery in eyes with various severities of open-angle glaucoma.

The study hypothesized that this treatment modality could effectively and safely reduce intraocular pressure (IOP) and medication burden over the long term in these glaucomatous eyes, while maintaining stability of long-term disease measures.

What was learned from the study?
The data demonstrated substantial 5-year reductions in IOP and medications, along with favorable safety, in eyes with a considerable preoperative medication burden and frequent history of glaucoma surgery. Long-term indicators of disease stability (visual fields, retinal nerve-fiber layer thickness, and cup:disc ratio) were unchanged over the course of 5-year follow-up. Results were similarly favorable in combined and standalone usage.

Drawing upon long-term, clinically heterogeneous, real-world patient data, this study provides directly relevant information for clinicians and patients evaluating glaucoma treatment options.

INTRODUCTION
The development of micro-invasive glaucoma surgery (MIGS) has fundamentally altered the way glaucoma is treated worldwide. Constituting virtually no procedures in the United States (US) in 2011, these surgeries became the most commonly-performed incisional glaucoma procedures by 2018 and are expected to further increase over time [1]. Among the various MIGS options, the iStent trabecular micro-bypass (the first US Food and Drug Administration-approved device, in 2012) and the second-generation iStent inject trabecular micro-bypass (containing 2 stents, each with modified design vs. the original) have accumulated the most diverse and long-term MIGS evidence base yet available [2]. The stents have been studied in various races/ethnicities, glaucoma types, severities, and usages (combined or standalone). Studies and subsequent publications have been completed in at least 20 countries, and to date there are at least 15 publications with 4-year or longer follow-up data [2]. Regarding iStent inject specifically, Germany was one of several countries around the world to rapidly adopt the technology, thereby allowing for some of the largest and longest-term datasets yet available on iStent inject effectiveness and safety.

Prospective 3-year findings from the present cohort have been previously published for both Combined and Standalone eyes [3, 4]. At 3 years postoperative, mean IOP and medication reductions in Combined eyes were 37% and 68%, respectively, and were 42% and 88% in Standalone eyes, respectively. In the current paper, we report the 5-year outcomes for the entire cohort, with subgroup analysis of Combined and Standalone eyes.

This report provides some of the first 5-year real-world iStent inject data yet available in the literature. By including consecutive patients (i.e. all-comers) rather than pre-selecting a narrow slice of the population, the results are broadly applicable to surgeons’ actual practices. The inclusion of subgroup stratification by combined-versus-standalone usage may further enhance generalizability and relevance. Subgroup analyses also allow us to discern the independent effect of the stents themselves, apart from those of cataract extraction. And, finally, the prospective longitudinal design and high accountability through 5 years help ensure the reliability of the findings.
Table 1 Demographics and preoperative characteristics \( (n = 125) \)

| Mean ± SD (unless noted otherwise) | Combined iStent inject + Phaco (81 eyes) | Standalone iStent inject (44 eyes) |
|-----------------------------------|----------------------------------------|-----------------------------------|
| Age (years)                       | 72.7 ± 7.5                             | 71.3 ± 10.5                       |
| Range (years)                     | 54–86                                  | 40–88                             |
| Gender                            | 56% male/44% female                    | 52% male/48% female               |
| Cup:disc ratio                    | 0.74 ± 0.17                            | 0.78 ± 0.18                       |
| VF MD                             | −6.6 ± 2.8 dB                          | −7.0 ± 3.5 dB                     |
| Retinal nerve fiber layer thickness (µm) | 81.4 ± 12.3                           | 82.1 ± 13.6                       |
| Race/ethnicity                    | 100% non-Hispanic white                | 100% non-Hispanic white           |
| Eyes with prior glaucoma procedures (%) | 26 (32%)                             | 22 (50%)                          |
| Prior glaucoma procedures         | 31 procedures in 26 eyes               | 35 procedures in 22 eyes          |
| Trabeculectomy                    | 12                                     | 13                                |
| CPC                               | 10                                     | 14                                |
| ALT/SLT                           | 5                                      | 3                                 |
| Laser iridotomy                   | 2                                      | 2                                 |
| Surgical iridectomy               | 1                                      | 0                                 |
| Single first-gen iStent           | 0                                      | 1                                 |
| XEN implantation                  | 1                                      | 2                                 |
| Type of glaucoma \( n \) (%)      | \( n \) (%)                            | \( n \) (%)                       |
| POAG                              | 60 (74%)                               | 38 (86%)                          |
| PXG                               | 15 (19%)                               | 4 (9%)                            |
| Combined-Mechanism                | 4 (5%)                                 | 1 (2%)                            |
| Pigmentary                        | 1 (1%)                                 | 0 (0%)                            |
| Secondary (neovascular/CRVO)      | 1 (1%)                                 | 1 (2%)                            |
| Mean medicated IOP                | 22.6 ± 6.2 mmHg                        | 25.3 ± 6.0 mmHg                   |
| Mean # medications                | 2.52 ± 1.06                            | 2.98 ± 0.88                       |
| Medicated IOP (mmHg) \( n \) (%)  | \( n \) (%)                            | \( n \) (%)                       |
| ≤ 15 mmHg                         | 1 (1%)                                 | 1 (2%)                            |
| ≤ 18 mmHg                         | 22 (27%)                               | 4 (9%)                            |
| Medication burden                 | \( n \) (%)                            | \( n \) (%)                       |
| Eyes med-free                      | 1 (1%)                                 | 0 (0%)                            |
METHODS

Study Design and Participants

In this prospective, longitudinal, single-surgeon study, all consecutive patients undergoing iStent inject implantation with or without cataract surgery were enrolled and followed for 5 years at an academic eye hospital in Germany. Patient recruitment occurred from March 2013 to December 2015; surgeries were completed from May 2013 to February 2016; and final data collection and analysis were completed by March 2021. To be enrolled in the study, patients were required to have a diagnosis of glaucoma [including primary open-angle glaucoma, pseudoexfoliative glaucoma (PXG), combined-mechanism glaucoma, pigmentary glaucoma, or secondary (neovascular) glaucoma]; have a history of inadequate response to prior surgical and medical glaucoma therapy; and to be in need of cataract surgery (for the Combined subgroup). Exclusion criteria consisted of corneal opacities preventing gonioscopic visualization, angle-closure glaucoma, pregnancy, congenital glaucoma, age under 18 years, and active ocular inflammation. Visits followed the surgeon’s standard postoperative schedule which included, at minimum, a preoperative visit and postoperative visits at Day 1, Week 1, Month 1, and Months 3, 6, 12, 24, 36, 48, and 60.

Study Outcomes and Data Analysis

Effectiveness measures included mean and percent reduction in IOP and medications versus preoperative. Proportional analyses for effectiveness included percent of eyes achieving the following outcomes: ≥ 20% IOP reduction versus preoperative; final IOP of ≤ 18 mmHg and ≤ 15 mmHg; same or lower IOP versus preoperative; same or lower medication burden versus preoperative; and treatment with 0 medications or ≥ 3 medications. Safety endpoints included intraoperative and postoperative adverse events, secondary surgical interventions, and proportion of eyes with traditional filtration surgery (trabeculectomy or tube shunt) over 5 years. In addition, long-term indicators of disease stability were measured and compared between the preoperative and 5-year visits; this included visual field mean deviation (VF MD), retinal nerve fiber layer thickness (RNFL), and C:D ratio.

All IOP readings were measured by Goldmann applanation. Preoperative IOP was based on readings at two or more preoperative visits; postoperative IOP was based on three measurements per time point. The study was approved by the institutional review board of the University of Heidelberg, and complied with the Helsinki Declaration. All participants provided informed consent prior to enrollment.

Changes in IOP or medications from preoperative were assessed via paired t test (for continuous variables) or Chi-square test (for categorical variables). Results were calculated and reported for the Overall cohort and for the Combined and Standalone subgroups.

| Table 1 continued |
|------------------|
| Mean ± SD (unless noted otherwise) | CombinediStent inject + Phaco(81 eyes) | StandaloneiStent inject(44 eyes) |
| Eyes on 3–5 preop meds | 44 (54%) | 33 (75%) |

IOP intraocular pressure, SD standard deviation, C:D cup:disc, VF MD visual field mean deviation, Preop preoperative, POAG primary open-angle glaucoma, PXG pseudoexfoliative glaucoma, CRVO central retinal vein occlusion, CPC cyclophotocoagulation, ALT argon laser trabeculoplasty, SLT selective laser trabeculoplasty; Med medication, Preop preoperative, Phaco phacoemulsification
iStent inject Device and Surgical Implantation Technique

The iStent inject device is comprised of two biocompatible titanium stents preloaded on a single-use sterile injector. Each stent contains four lateral outlet lumens to facilitate multidirectional outflow. The stents are placed ab internis through the trabecular meshwork into Schlemm's Canal at approximately 2–3 h apart, providing access to up to 6 h of aqueous outflow [5, 6]. Implantation is completed either following phacoemulsification (in combined-cataract cases) or as a standalone procedure, as described in the respective prior publications [3, 4]. Following implantation, stent position and patency are confirmed via gonioscopy, viscoelastic is removed, and surgical wound

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**Fig. 1**  
a Mean intraocular pressure (IOP) through 5 years postoperative in overall cohort.  
b (left) Mean IOP at 5 years versus preoperative: combined subgroup.  
c (right) Mean IOP at 5 years versus preoperative: standalone subgroup

*M = months; Preop = preoperative; IOP = intraocular pressure. Vertical bars represent standard deviation.*
(a) Preop vs Postop

\[ p < 0.001 \]

Preop (n=125)

Postop (n=121)

IOP ≤ 18 mmHg

% of Eyes

21% 98%

M = months; Preop = preoperative; Postop = postoperative; IOP = intraocular pressure.

(b) Preop vs Postop

\[ p < 0.001 \]

Preop (n=81)

Postop (n=79)

IOP ≤ 18 mmHg

% of Eyes

27% 100%

M = months; Preop = preoperative; Postop = postoperative; IOP = intraocular pressure.

(c) Preop vs Postop

\[ p < 0.001 \]

Preop (n=44)

Postop (n=42)

IOP ≤ 18 mmHg

% of Eyes

9% 95%

M = months; Preop = preoperative; Postop = postoperative; IOP = intraocular pressure.
closure is ensured. Postoperatively, patients were prescribed 1 week of topical antibiotic ciprofloxacin (Floxal®, Bausch&Lomb, Berlin, Germany) and four weeks of topical anti-inflammatory medication (Dexamethasone; Bausch&Lomb).

RESULTS

Subject Accountability, Demographics, and Preoperative Parameters

This prospective consecutive cohort included 125 total eyes (81 Combined, 44 Standalone) of a single glaucoma surgeon at an academic hospital in Germany. Nearly all eyes (121/125 or 97%) completed 5 years of follow-up. Preoperative parameters are shown in Table 1. These were patients carrying a relatively substantial preoperative treatment burden, with overall mean preoperative IOP of 23.5 ± 6.2 mmHg on 2.68 ± 1.02 mean medications. All but 1 eye (>99%) were on medication(s) preoperatively; 38% of eyes had undergone prior glaucoma surgery. In comparison to Combined eyes, Standalone eyes had slightly higher mean IOP and medications, and higher proportions of eyes with history of prior glaucoma surgery (50% vs. 32%, respectively) and eyes on ≥3 glaucoma medications (75% vs. 54%, respectively).

Intraocular Pressure

In the Overall cohort (n = 125), mean IOP decreased to 14.1 ± 1.8 mmHg at 5 years (40% reduction vs. preoperative; p < 0.001) (Fig. 1a). A ≥ 20% IOP reduction versus preoperative was achieved in 83% of eyes, with all but 1 eye (>99%) maintaining or reducing IOP versus preoperative (the one remaining eye increased from 15 to 16 mmHg). The proportion of eyes with IOP ≤ 18 mmHg rose to 98% (vs. 21% preoperatively; p < 0.001); and eyes with IOP ≤ 15 mmHg rose to 81% (vs. <2% preoperatively; p < 0.001) (Fig. 2a).

The IOP outcomes in the Combined and Standalone subgroups were similar to those of the Overall cohort. In Combined eyes, mean IOP decreased from 22.6 mmHg preoperatively to 13.8 mmHg at 5 years (39% reduction; p < 0.001) (Fig. 1b). All Combined eyes (100%) achieved IOP ≤ 18 mmHg (vs. 27% preoperatively; p < 0.001), and 86% of eyes had IOP ≤ 15 mmHg (vs. 1% preoperatively; p < 0.001) (Fig. 2b).

In Standalone eyes, mean IOP reduced from 25.3 mmHg to 14.6 mmHg (42% reduction, p < 0.001) (Fig. 1c). There was a significant rise in the proportion of eyes with IOP ≤ 18 mmHg (95% at 5 years vs. 9% preoperatively, p < 0.001), and with IOP ≤ 15 mmHg (71% at 5 years vs. 2% preoperatively, p < 0.001) (Fig. 2c).

Medication Use

In the Overall cohort, mean number of medications at 5 years decreased to 0.77 ± 0.82 medications (71% reduction vs. preoperative; p < 0.001) (Fig. 3a). At 5 years, 46% of eyes were medication-free (vs. 1 eye or <1% preoperatively; p < 0.001), while <2% of eyes were using ≥3 medications (versus 62% preoperatively; p < 0.001). All eyes (100%) were able to maintain or reduce their medication regimen versus preoperative (Fig. 4a).

Within the Combined and Standalone subgroups, the 5-year medication reductions were similar to those of the Overall cohort. In Combined eyes, mean number of medications reduced from 2.52 to 0.78 medications (69% reduction; p < 0.001) (Fig. 3b). The proportion of medication-free eyes rose to 44% (vs. 1 eye or 1% preoperatively; p < 0.001); inversely, only 1
eye (1%) at 5 years was on ≥3 medications (vs. 54% of eyes preoperatively; \( p < 0.001 \)) (Fig. 4b). All Combined eyes maintained or reduced medication burden versus preoperative.

In Standalone eyes, mean number of medications reduced from 2.98 preoperatively to 0.74 medications at 5 years (75% reduction; \( p < 0.001 \)) (Fig. 3c). The proportion of medication-free eyes rose to 50% (vs. 0 eyes preoperatively; \( p < 0.001 \)); inversely, only 1 eye (2%) at 5 years was on ≥3 medications (vs. 75% of eyes preoperatively; \( p < 0.001 \)) (Fig. 4c). All Standalone eyes achieved the same or lower medication burden versus preoperative.

**Safety Assessment**

All patients underwent successful implantation of 2 iStent inject stents either as a sole procedure...
(a) 100% of eyes had the same or lower medication burden vs preop

M = months; Preop = preoperative; Postop = postoperative IOP = intraocular pressure.

(b) 100% of eyes had the same or lower medication burden vs preop

M = months; Preop = preoperative; Postop = postoperative IOP = intraocular pressure.

(c) 100% of eyes had the same or lower medication burden vs preop

M = months; Preop = preoperative; Postop = postoperative IOP = intraocular pressure.
or following standard phacoemulsification cataract surgery. One intraoperative event occurred: a Standalone case with mild intraoperative hyphema which resolved without intervention or sequelae by Week 1. Postoperative adverse events during the first 3 years have been described in prior publications; these events were generally mild, transient, and self-resolved or were treated conservatively without sequelae [3, 4]. They included 2 Combined eyes with mild corneal edema on Day 1 which resolved by Day 5 postoperatively; 2 eyes of one Standalone patient with cataract progression at Month 3, and 1 Standalone eye with uveitis at Month 24, which resolved within 3 weeks with topical anti-inflammatory medication. During 2 additional years of follow-up, no new adverse events occurred. Throughout follow-up there were no stent-related complications; no signs of chronic inflammation such as peripheral anterior synechiae (PAS); and no cases of hypotony, endophthalmitis, corneal compromise, stent obstruction, myopic shift, or choroidal detachment.

Over the course of 5-year follow-up, a total of 5 eyes (3 Combined, 2 Standalone) underwent an additional glaucoma procedure due to the need for additional IOP-lowering beyond what had been achieved with stent implantation. The procedures, all of which occurred at Month 1 or Month 6 postoperatively, included cyclophotocoagulation (CPC) in 3 eyes and XEN implantation in 2 eyes. No filtering surgeries were performed in any eye throughout the 5 years.

**Long-term Disease Stability**

Long-term indicators of disease stability were measured and compared between the preoperative and 5-year visits. No significant differences were observed in visual fields, retinal nerve fiber layer thickness, or C:D ratio in either the Combined or Standalone subgroups over the course of the study. Regarding VF stability over time, average VF MD in the Combined subgroup was \(-6.6 \pm 2.8\) db preoperatively versus \(-6.7 \pm 2.6\) db at 5 years \((p = 0.462)\). In the Standalone subgroup, average VF MD was \(-7.0 \pm 3.5\) db preoperatively versus \(-7.1 \pm 3.3\) db at 5 years \((p = 0.502)\). Regarding RNFL stability over time, mean RNFL thickness in the Combined subgroup was \(81.4 \pm 12.3\) µm preoperatively versus \(80.2 \pm 13.5\) µm at 5 years \((p = 0.497)\). In the Standalone subgroup, mean RNFL thickness was \(82.1 \pm 13.6\) µm preoperatively versus \(80.9 \pm 14.7\) µm at 5 years \((p = 0.194)\). Regarding C:D stability over time, mean C:D ratio in the Combined subgroup was \(0.74 \pm 0.17\) preoperatively versus \(0.76 \pm 0.14\) at 5 years \((p = 0.324)\). In the Standalone subgroup, mean C:D ratio was \(0.78 \pm 0.18\) preoperatively versus \(0.79 \pm 0.16\) at 5 years \((p = 0.386)\).

**DISCUSSION**

This prospective consecutive longitudinal cohort, with 97% of the 125 eyes followed through 5 years, constitutes one of the largest and longest-term datasets to date on iStent inject. Given that Germany was one of the first countries to adopt this technology, the present dataset provides a valuable window on iStent inject outcomes over the long term. These outcomes included stable and significant reductions in IOP and medications, together with excellent safety and 100% of eyes avoiding filtration surgery. Although the cohort had relatively high glaucoma medication burden preoperatively, nearly half of eyes were medication-free by the end of 5 years, while mean IOP decreased by nearly 10 mmHg. The observed reductions in IOP and medication were similar in magnitude and consistency over time regardless of whether stenting was performed as a standalone surgery or in combination with phacoemulsification.

A number of noteworthy attributes may be highlighted in this study. First, it provides some...
of the first 5-year data on iStent inject in a real-world clinical population, as noted above. Patient accountability was nearly 100% over the entirety of follow-up, supporting the credibility and strength of the findings. The cohort included Combined and Standalone subgroups within the hands of a single surgeon at a single site and drawn from the same patient base. Such uniformity helps reduce confounding when comparing across subgroups and when distinguishing the IOP-lowering effect of the stents versus that of phacoemulsification.

The percent reductions in IOP and medications were similar for both Combined and Standalone subgroups, even though Standalone eyes had higher preoperative IOP, medication burden, and history of prior glaucoma surgery. This is a strong confirmation that the stents—rather than the cataract extraction—are producing the change. The independent effect of the stents is also validated by the consistency of mean IOP and medications throughout 5 years of follow-up (in contrast to the typically temporary ocular hypotensive effect of phacoemulsification). In Standalone eyes, as in combined eyes, the IOP and medication reductions at 5 years were similar to those observed at 3 years, suggesting no diminution of treatment effect over time. Such stability is noteworthy, given the natural history of glaucoma as an otherwise progressive, steadily advancing disease.

In comparison to the majority of existing literature, the postoperative medication burden (both mean and percent reduction vs. baseline) was comparable to what has been reported previously in combined and standalone iStent inject cohorts [7–32]. The postoperative mean IOP was also in the range of prior iStent inject reports, although the current study’s pre-versus-postoperative percent reduction was greater than in many other studies. Given the overall similarity and consistency of postoperative IOP ranges across the reports, the greater postoperative percent reduction in the present study is possibly a function of higher starting pressure, which is known to allow for greater percent IOP reductions postoperatively [8–12].

Among the publications on iStent inject, two in particular may be informative against which to compare our study. For the Combined subgroup, we can consider our findings in the context of the largest multicenter real-world study to-date on combined iStent inject implantation [15]. That study, by Clement et al., included 340 eyes from 9 surgeons across Australia. They reported final IOP of 13.7 mmHg at 2 years postoperative (a 16% reduction vs. a relatively low baseline IOP of 16.4 mmHg), with 77% of eyes achieving IOP ≤ 15 mmHg and mean medication burden reducing by 67% (from 1.49 to 0.49 mean medications) [15]. In comparison, the Combined subgroup in our cohort experienced a final postoperative IOP of 13.8 mmHg (a 39% reduction vs. a baseline IOP of 22.6 mmHg), with 86% of eyes achieving IOP ≤ 15 mmHg and mean medication burden reducing by 69% (from 2.52 to 0.78 mean medications). For the Standalone subgroup in our cohort, the IOP outcomes are closely aligned with the largest and most comprehensive systematic review and meta-analysis to-date of standalone iStent and iStent inject implantation [7]. That publication, by Healey et al., analyzed 13 different studies and reported that they all centered around a final mean IOP of 13.0–16.5 mmHg through up to 5 years postoperative, starting from a mean preoperative IOP of 19.1–25.3 mmHg. Similarly, the Standalone subgroup in our cohort experienced a final IOP of 14.6 mmHg at 5 years postoperative, starting from a mean preoperative IOP of 25.3 mmHg.

As is characteristic of the iStent family of devices, safety outcomes in this study were highly favorable. Notably, there were no intraoperative complications and no filtration surgery in any eye over 5 years of follow-up. Given their substantial preoperative disease burden (mean IOP 23.5 mmHg on 2.68 medications) and history of prior glaucoma surgery (38% of eyes), many of these patients would possibly have needed higher-risk filtration surgery at some point during the 5 years of follow-up if they had not stabilized their IOP with iStent inject implantation.

The study was limited by its unmasked, single-arm, non-randomized design. Patients comprised a consecutive sample of all cases of a single surgeon; this increases generalizability...
but also makes it more difficult to account for non-quantifiable differences in preoperative characteristics. Patients did not complete medication washouts, as this was not indicated nor appropriate in the patients' regular clinical care. All patients were non-Hispanic white and of German nationality, so the reader must exercise discernment when extrapolating outcomes to other populations. Within the Combined subgroup, the IOP-lowering effect of cataract extraction could not be separated from that of the concurrent stent implantation. However, post-phacoemulsification IOP reduction in medicated glaucoma patients is known to be modest (less than 2 mmHg on average) [33–35], and IOP typically starts returning to baseline levels after the first year postoperative [35–38]. Therefore, our study’s findings of consistent IOP reduction through 5 years, and similar reductions in Combined and Standalone patients, both corroborate that phacoemulsification is no longer exerting an effect on IOP outcomes. Future areas for investigation could include expanded analyses of specific glaucoma subtypes (such as PXG), stratification by preoperative glaucoma severity or surgical history, and inclusion of the newest iStent inject model, the wide-flanged iStent inject W (released in Germany in 2020).

CONCLUSIONS

This study contributes valuable 5-year data on iStent inject implantation either in combination with cataract surgery or as a sole procedure. The same cohort was prospectively identified and followed longitudinally, with nearly all patients reaching their 5-year visit. Although patients had a relatively high glaucoma treatment burden preoperatively, nearly half of eyes were medication-free by 5 years, while mean IOP reduced by nearly 10 mmHg. The degree of IOP and medication reduction was similar in both Combined and Standalone subgroups, validating the stents’ independent effect apart from that of cataract extraction. The treatment effect was also consistent visit-to-visit throughout follow-up, confirming the sustained 5-year effectiveness of the stents. Disease stability was also confirmed, with no significant change in VF, RNFL, or C:D ratio at 5 years compared to preoperative. And, finally, since outcomes were recorded in a real-world patient population as part of routine clinical care, they can be applicable to surgeons making treatment decisions with their patients in combined and standalone settings.

ACKNOWLEDGEMENTS

Funding. Open Access funding enabled and organized by Projekt DEAL. An unrestricted research grant and editorial assistance were provided by Glaukos Corporation. Glaukos Corporation also funded the Journal’s Rapid Service and Open Access Fees. All authors had full access to all of the data in this study and take complete responsibility for the integrity of the data and accuracy of the data analysis.

Authorship. All authors meet the International Committee of Medical Journal Editors (ICMJE) criteria for authorship for this article, take responsibility for the integrity of the work as a whole, and have given their approval for this version to be published. Editorial assistance in the preparation of this article was provided by Dr. Dana M. Hornbeak, MD, MPH of Glaukos Corporation.

Author Contributions. Fritz H. Hengerer: concept and design, completeness of surgeries, preoperative/postoperative management, data collection, statistical analysis, manuscript drafting. Gerd U. Auffarth: concept and design, preoperative/postoperative management, data collection, manuscript drafting. Ina Conrad-Hengerer: concept and design, preoperative/postoperative management, statistical analysis, manuscript drafting.

Disclosures. Fritz H. Hengerer: travel support, research grants, and lecture fees from Glaukos Corporation. Gerd U. Auffarth: travel support, research grants, and lecture fees from Glaukos Corporation. Ina Conrad-Hengerer:
research grants, and lecture fees from Glaukos Corporation.

**Compliance with Ethics Guidelines.** All procedures performed were in accordance with the ethical standards of the institutional and/or national research committee (the Institutional Review Board (IRB) of the University of Heidelberg), and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards. Informed consent was obtained from all individual participants included in the study.

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

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