

Consistency in Standalone Canaloplasty Outcomes Using the iTrack Microcatheter

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Purpose: To study the consistency in outcomes of standalone canaloplasty performed via an ab-interno surgical technique in reducing intraocular pressure (IOP) and number of medications in uncontrolled open-angle glaucoma (OAG) eyes over a 12-month period.

Methods: This retrospective multicenter case series included patients who underwent standalone canaloplasty via an ab-interno surgical technique using the iTrack microcatheter (Nova Eye, Inc., Fremont, USA) and had preoperative uncontrolled OAG (IOP $\geq$ 18mmHg) along with no previous glaucoma surgery. The iTrack microcatheter is used to circumnavigate 360° and viscodilate Schlemm's canal. Consistency of IOP and medications reduction on an eye-by-eye basis were evaluated to understand the outcomes in each single eye.

Results: Sixty-four eyes of 60 patients (age 71.5 $\pm$ 13.4 years) were included. Six eyes (9%) that underwent additional glaucoma surgery were considered a failure and were subsequently excluded from analysis. At 12 months, IOP was reduced in 57 of the 58 (89%) remaining eyes; one eye had the same IOP with a reduced number of medications. Of the 57/58 eyes with a reduced IOP: 44 eyes (69%) required fewer medications; 12 eyes (19%) required the same number of medications. Of these 58 eyes, 78% of eyes had a $\geq$ 20% reduction in IOP compared to baseline; 69% eyes had a postoperative IOP $\leq$ 15 mmHg, and 86% eyes $\leq$ 18 mmHg at 12 months. Forty percent of the eyes were medication-free at 12 months compared to none at baseline.

Conclusion: Canaloplasty performed via an ab-interno surgical technique as a standalone procedure consistently reduced IOP and glaucoma medications in almost all eyes.

Keywords: canaloplasty, iTrack, microcatheter, viscodilation, intraocular pressure, open-angle glaucoma



Introduction

Glaucoma has been identified as the leading cause of irreversible but preventable blindness in the world, with the number of cases expected to rise to 112 million by 2040, exerting a huge pressure on healthcare systems worldwide.^{1,2}

Intraocular pressure (IOP) is known to be the only known modifiable risk factor in glaucoma.³ According to the Early Manifest Glaucoma Trial, every 1 mmHg decrease in IOP can reduce the progression of glaucoma by almost 10%.^{4,5}

Introduced to bridge the gap between topical antiglaucoma treatment and invasive filtration surgery, minimally invasive glaucoma surgery (MIGS) has demonstrated utility in lowering IOP in a safe and significant manner. However, the results are usually reported as a mean outcome of a cohort and, still today, little is known regarding how consistent MIGS procedures are on an eye-by-eye basis.

Several MIGS devices are currently available, differing mostly in their mechanism of action and target location, acting on either the trabecular/conventional outflow, the suprachoroidal area, or the subconjunctival area.⁶ The majority of MIGS are focal in their approach, i.e. the iStent Trabecular Micro-Bypass Stent System (Glaukos Corporation, San Clemente, CA) and Hydrus Microstent (Ivantis, Inc, Irvine, CA). With focal-based MIGS, the clinical outcomes may vary depending on the placement of the stent and whether this correlates with the point(s) of outflow resistance. Consequently, results may therefore

fluctuate among eyes, with some eyes experiencing a large IOP reduction and other eyes with no IOP reduction at all.⁷ With this perspective, the mean IOP reduction of a cohort may not help predictability on an individual basis.

Since the variety of MIGS devices and procedures enables surgeons to take a more nuanced treatment approach that is better tailored to the needs of each patient, their effectiveness should also be evaluated with respect to each individual patient rather than relying only on mean outcomes.

The authors decided to investigate whether canaloplasty would yield a consistent eye-by-eye reduction in IOP. Canaloplasty performed via an ab-interno surgical technique is a type of implant-free MIGS that targets all aspects of the complete 360° of the conventional outflow pathway (trabecular meshwork, Schlemm's canal, and the distal collector channels), offering a comprehensive approach to lower IOP, unlike other MIGS that are mainly focal in their approach. This understanding of the disease pathology may be further translated into a consistent reduction in IOP in all patients of primary open-angle glaucoma (POAG).^{8,9} The authors decided to investigate its efficacy when performed as a standalone procedure in excluding the confounding effect of phacoemulsification in reducing IOP.^{10,11}

This study assesses the consistency of results (the IOP lowering effect on each single eye) obtained with standalone canaloplasty via an ab-interno surgical technique in reducing IOP and number of medications in uncontrolled open-angle glaucoma patients over a 12-month period using the iTrack canaloplasty device (Nova Eye, Inc., Fremont, California).

Materials and Methods

Study Design

This study was a multicenter, multi-surgeon, retrospective review of a consecutive case series of eyes treated with canaloplasty as a standalone procedure. This study was conducted according to the tenets of the Declaration of Helsinki, and written consent was obtained from patients. Data meeting inclusion criteria (below) was extracted and collated from three cohorts whose overall results are already published and available in the scientific literature and had received Institutional Review Board (IRB) approval or waiver.^{12–14}

Patient Selection

Inclusion criteria included adult POAG patients (18 years of age or older) with a diagnosis of mild–moderate or severe glaucoma, as per the Hodapp–Parrish–Anderson classification system, uncontrolled IOP with medications (defined as baseline IOP ≥ 18 mmHg), as well as healthy angle structures on gonioscopy with a homogeneously pigmented trabecular meshwork without variegation or excessive pigmentation. Disease severity was determined based on mean deviation (MD) from Humphrey visual field (HVF) testing: MD less than -6 dB (mild), -6 dB to -12 dB (moderate), and greater than -12 dB (severe).

Patients with peripheral anterior synechiae, goniosynechiae, or angle recession, were excluded from the study. Patients with neovascular disease, uveitis, peripheral anterior synechiae, and developmental or other forms of secondary glaucoma, such as steroid-induced glaucoma, were also excluded.

The authors (MK, NK, SO, MG) have collated three different cohorts of eyes with open-angle glaucoma that was uncontrolled by medications at baseline (IOP ≥ 18 mmHg), underwent canaloplasty as a standalone procedure (not concomitant with cataract surgery), reached at least the 12-month follow-up, and did not undergo any additional glaucoma surgery. The outcomes of the entire cohort of eyes (ie, including the eyes treated with other glaucoma surgery and that were controlled at baseline) were previously published in the literature (Gallardo 2022,¹² Khaimi 2021,¹³ Koerber et al 2022¹⁵).

Endpoints

Success was defined either as a reduction in IOP and number of medications, or a reduction in IOP with the same number of medications. An ambiguous outcome was defined as when either the IOP decreased but the number of medications also increased, when IOP and number medications did not change, or when IOP increased and number of medications decreased. Failure was assigned when eyes underwent additional glaucoma surgery and when IOP remained the same or

increased and the number of medications increased or remained the same. Success was also reported according to the Guidelines on Design & Reporting Glaucoma Trials.¹⁶

Surgical Technique

A small 1–2 mm incision is made in the trabecular meshwork, and the microcatheter is inserted into Schlemm's canal, circumnavigating the entire 360°. If an obstruction to the passage of the microcatheter is encountered, the microcatheter is withdrawn and re-inserted into the canal in the opposite direction. After the microcatheter completes 360° catheterization of the canal, it is slowly withdrawn. Simultaneously, precisely regulated microquantities of high-molecular-weight hyaluronic acid (HA)-based ophthalmic viscosurgical device (OVD) is delivered into Schlemm's canal, with a mean of 36 clicks per procedure, resulting in an average of 100µL delivered over the entirety of Schlemm's canal.

The postoperative care included a topical steroid like Loteprednol, and a fourth-generation fluoroquinolone. Anti-glaucoma medications were stopped postoperatively.

A description of the surgical steps is described in the literature.^{12,13,15}

Device

All patients underwent canaloplasty performed with the same device: iTrack (Nova Eye, Inc., Fremont, USA). The iTrack is a 200-micron microcatheter with an illuminated fiber optic tip that provides continuous location while performing surgery in the eye, designed for canaloplasty. It performs 360° catheterization and pressurized viscodilation of Schlemm's canal.

Statistical Analysis

The comparisons between changes in IOP and number of medications between two time points were analyzed using a commercially available software (Excel, Microsoft) with, where applicable, non-parametric tests (Wilcoxon) (Jamovi). Descriptive statistics (mean, standard deviation, and range) were calculated for IOP and number of medications at each visit. A p-value of <0.05 was required for a value to be considered as statistically significant and p-values are indicated where applicable. Graphs were produced according to the Guidelines on Design & Reporting Glaucoma Trials.¹⁶

Results

Demographics

In total, the entire canaloplasty cohort of uncontrolled OAG eyes included 85 eyes. Of those, several eyes were excluded from the analysis for the following reasons: 1 eye was administered Avastin prior to the 6-month follow-up; 6 eyes were lost at follow-up before 12 months. Fourteen additional eyes that were excluded had a preoperative IOP below 18 mmHg.

The remaining 64 eyes met the inclusion criteria (standalone canaloplasty, uncontrolled glaucoma) and were enrolled in the study. Thirty-eight percent of eyes had mild glaucoma, 27% had moderate, and 27% had severe glaucoma, while 9% of the eyes were not classified. Table 1 presents patient demographics while Table 2 shows preoperative measurements and postoperative outcomes at 12 months.

Six eyes (9%) that underwent additional glaucoma surgery prior to the 12-month follow-up (5 Express shunts, Alcon, and 1 cyclophotocoagulation) were categorized as complete failures and were not included in the data analysis.

Mean IOP and Medications

Mean reductions in IOP and number of medications were statistically significant ($p < 0.001$) in the remaining cohort that was analyzed (58 eyes; 91% of the entire cohort): mean IOP decreased from 22.94 ± 7.03 mmHg (median: 20; range: 18, 58) at baseline to 15.03 ± 2.9 mmHg (median: 15; range: 9, 26) at 12 months postop and mean number of medications decreased from 2.74 ± 0.8 (median: 3; range: 1, 4) to 1.14 ± 1.1 (median: 1; range: 1, 3) at the 12-months postoperative time point. Figure 1 shows a bar diagram of the number of medications at baseline and 12 months.

Table 1 Patient Demographics

Parameters	Mean ± SD or n (%)
Number of patients	60
Age (years)	71.5±13.4
Gender	
Male	23 (38%)
Female	36 (60%)
Missing	1 (2%)
Ethnicity	
White	25 (42%)
Black	6 (10%)
Hispanic	26 (43%)
Missing	3 (5%)
Number of eyes	64
Eye	
Right	35 (55%)
Left	29 (45%)
Glaucoma severity	
Mild	24 (37.5%)
Moderate	17 (26.55%)
Severe	17 (26.55%)
Missing	6 (9.4%)

Table 2 Preoperative Mean Measurements and Postoperative Mean Outcomes

Parameters	Preoperative	12 Months
IOP (mmHg)	22.9±7.03 (n=64)	15.0±2.90 (n=58)
Medications (no.)	2.78±0.83 (n=64)	1.14±1.12 (n=58)
CDVA (logMAR)	0.33±0.46 (n=61)	0.32±0.48 (n=52)
VF MD	-8.74±8.25 (n=46)	–

IOP and Medications Reduction on an Eye-by-Eye Basis (Baseline Vs 12 Months) Success and Failures

Figure 2 shows a scatterplot of the 58 eyes that reached the 12-month follow-up with canaloplasty alone. Each point represents an eye’s outcomes at preop and postop. Notably, the reduction in IOP was consistent, with no increase in IOP observed.

Fifty-seven (57) out of those 58 eyes (89% of the entire cohort) demonstrated a reduced IOP at 12 months postop, while one eye had no change in IOP. Of the 57/58 eyes that noted a reduced IOP, 44 eyes (69%) also had a reduced medication burden, while in 12 eyes (19%) the number of medications remained the same. As such, 56 eyes (88%) were categorized as success and 2 eyes (3%) as ambiguous. Six eyes (9%) were categorized as complete failure because they underwent additional glaucoma surgery (Table 3).

When using the Guidelines on Design & Reporting of Glaucoma Surgical Trials (World Glaucoma Association),¹⁶ 87.5% of the eyes were either a complete or qualified success and 12.5% were failures (Table 4).

Stratification of IOP Reduction

Out of the 58 eyes that were analyzed, 28 eyes (48%) had an IOP reduction of ≥7 mmHg and 25 eyes (43%) had an IOP reduction between 3 and 6 mmHg. Only 5 eyes (9%) had an IOP reduction between 0 and 3 mmHg (Figure 3). Forty-five eyes (77.6%) of eyes had a ≥20% reduction in IOP compared to baseline; 69% eyes had a postoperative IOP ≤15 mmHg

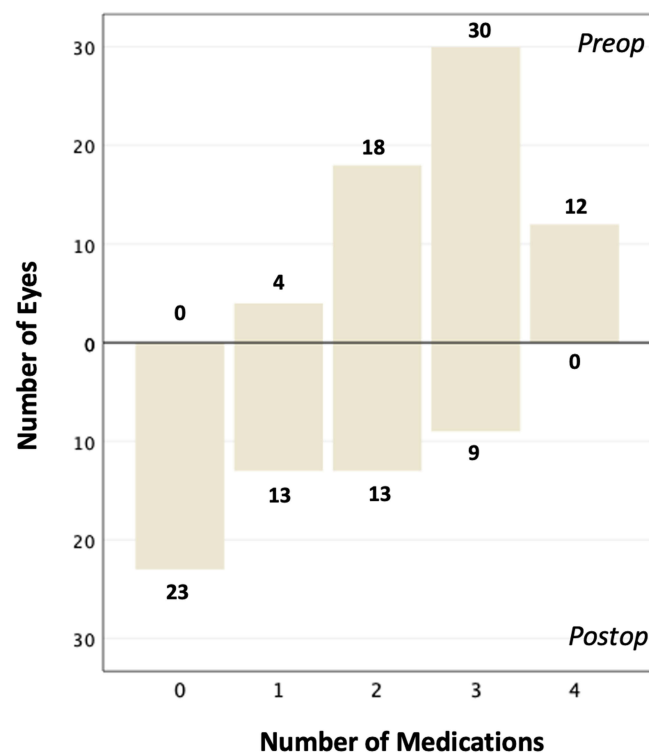


Figure 1 Bar diagram showing number of eyes with medications at preop and postop (12 months).

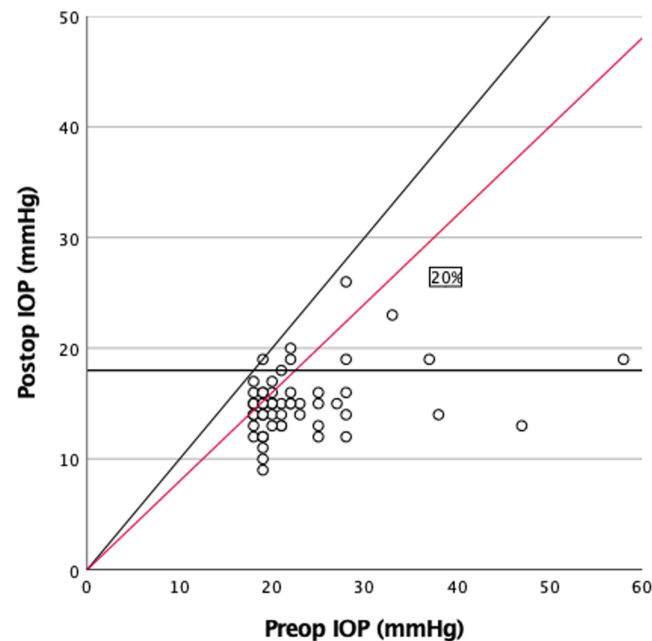


Figure 2 Scatterplot of intraocular pressure (IOP) outcomes at preop and postop (12 months). Points on the diagonal line indicate eyes with no change in IOP; points to the right of the red line indicate eyes with at least 20% reduction in IOP. Horizontal line indicates IOP of 18 mmHg.

compared to none at preop, and 86.2% eyes had an IOP ≤ 18 mmHg at 12 months, compared to 17.2% at preop (10 eyes had an IOP=18mmHg at baseline).

Forty percent of the eyes were medication-free at 12 months compared to zero at baseline (Table 5). Thirty-four percent of the eyes had a postoperative IOP ≤ 18 mmHg without any medications, compared to zero at baseline.

Table 3 Number and Percentage of Eyes That Succeeded, Failed, or Had Ambiguous IOP and Medication Outcomes (Baseline Vs 12 Months Data) as per Methods Section

Surgical Outcomes	n (%)
Success	56 (87.5%)
IOP decreased and meds decreased	44 (68.75%)
IOP decreased and meds remained the same	12 (18.75%)
Ambiguous	2 (3.1%)
IOP decreased and meds increased	1 (1.56%)
IOP same and meds decreased	1 (1.56%)
IOP increased and meds decreased	0%
Failure	0 (0%)
IOP remained the same and meds decreased	0%
IOP increased and meds increased	0%
IOP increased and meds remained the same	0%
Complete failure	6 (9.4%)
Additional glaucoma surgery required	6 (9.38%)

Table 4 Surgical Outcomes (Baseline Vs 12 Months). Success Range Reported as per the Guidelines on Design and Reporting of Glaucoma Surgical Trials (World Glaucoma Association)

Surgical Outcomes	n (%)
Complete success	23 (35.9%)
IOP ≤ 21 mmHg, without medications at postop	3 (4.69%)
IOP ≤ 18 mmHg, without medications at postop	20 (31.3%)
Qualified success	33 (51.6%)
IOP ≤ 21 mmHg, with medications at postop	3 (4.69%)
IOP ≤ 18 mmHg, with medications at postop	30 (46.9%)
Failure	2 (3.1%)
IOP > 21 mmHg, with $\geq 20\%$ reduction from preop	1 (1.56%)
IOP > 21 mmHg, with $< 20\%$ reduction from preop	1 (1.56%)
Complete failure	6 (9.4%)
Additional glaucoma surgery required	6 (9.4%)

Figure 4 shows the mean pre- and postoperative outcomes and the range of the IOP distribution of the 58 eyes, being distributed widely at baseline (range 18–58 mmHg; median 20 mmHg) while much narrower at 12 months postoperatively (range 9–26 mmHg; median 15 mmHg).

Preoperatively, 17% of the eyes were ≤ 18 mmHg and 45% ≤ 21 mmHg; postoperatively they were 86% and 10% respectively, with only 4% above 21 mmHg.

Excluded Eyes Due to Additional Glaucoma Surgery

Six eyes were excluded because they received additional glaucoma surgery for a various of reasons and the IOP and medication outcomes could not be evaluated with canaloplasty only at 12 months. The eyes (glaucoma stage mild $n=3$; moderate $n=2$; severe $n=1$) had a mean preoperative IOP of 23.5 mmHg and were on 3.2 medications.

One of the 6 eyes was excluded because it underwent cyclophotocoagulation at 6 months: this eye belonged to a 90-year-old patient with very low visual acuity (0.10 logMAR) and while this eye IOP was under control, it was not possible to reduce the medications. One eye registered an IOP reduction of 20% which was not enough to bring the glaucoma

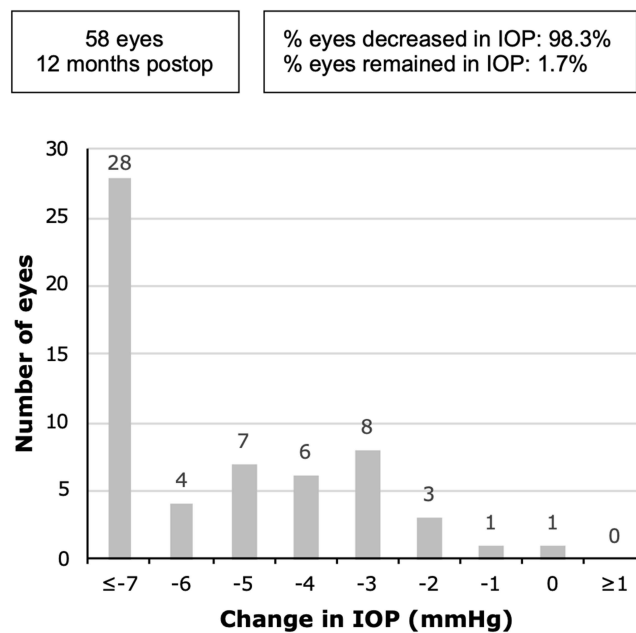


Figure 3 Changes in intraocular pressure outcomes of 58 eyes (baseline vs 12 months).

under control (from 25mmHg to 20mmHg). IOP did not reduce in the remaining 4 eyes (6% of the entire cohort; 2 mild, 1 moderate, 1 severe).

Safety

There were no serious complications recorded. Postoperative micro-hyphema and cells, as well as flare, were observed in some patients, but they were transient and resolved without sequelae. Vision on the logMAR chart remained stable (Table 2).

Discussion

The literature published by the authors in 2021/2022 has supported the safety and effectiveness of canaloplasty via an ab-interno surgical technique for POAG in lowering mean IOP and reducing the mean number of medications^{12,13,15,17} and several other studies have also confirmed these findings.^{18–21} Riaz et al²² showed that the tissue-sparing, minimally invasive nature of canaloplasty, which aims to work with patient physiology, can also effectively and safely reduce IOP in post-keratoplasty glaucoma eyes, where topical hypotensives may pose a risk factor for corneal graft failure and

Table 5 IOP and Medications Outcomes (Baseline Vs 12 Months) of the 58 Eyes Reaching the 12-Month Follow-Up with Canaloplasty Alone

	Preop	12 Months
<i>Intraocular Pressure</i>		
Percentage of eyes with IOP ≤15 mmHg	0 (0%)	40 (69.0%)
Percentage of eyes with IOP ≤18 mmHg	10 (17.2%)	50 (86.2%)
Percentage of eyes with ≥20% reduction in IOP	–	45 (77.6%)
Percentage reduction in mean IOP	–	34.3%
<i>Medications</i>		
Percentage of medication-free eyes	0 (0%)	23 (39.7%)
Percentage of medication-free eyes (eyes with IOP ≤18 mmHg only)	0 (0%)	20 (34.5%)
Percentage of eyes on ≤1 medication	4 (6.90%)	36 (62.1%)
Percentage reduction in mean number of medications	–	58.5%

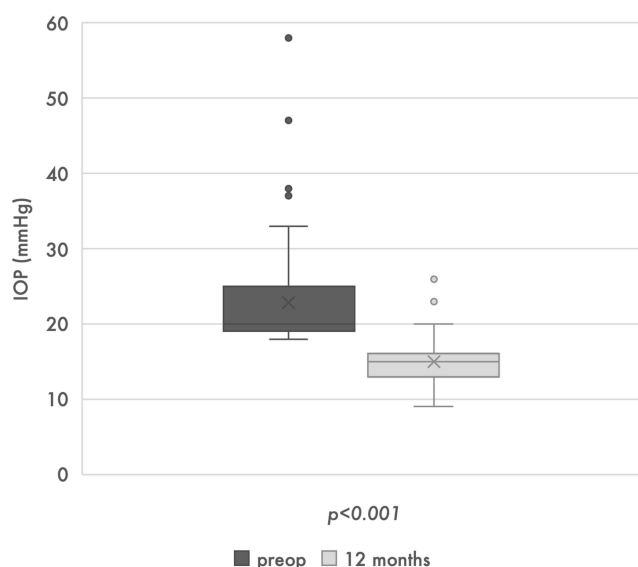


Figure 4 Mean preoperative and postoperative intraocular pressure outcomes of 58 eyes.

further filtration glaucoma surgery may be too invasive for such fragile corneas.²³ The mean IOP and medication reduction of canaloplasty are proven in this study and in the literature.

However, the purpose of this study was not to investigate the mean IOP reduction but the consistency of IOP reduction in each individual eye. It was observed that almost all eyes that reached the 12-month follow-up (57/58 eyes) with canaloplasty alone (and no other glaucoma surgery) have obtained an IOP reduction that, in most cases, was associated with a reduction in the number of medications. It is not possible to compare the results of this cohort in the literature given that, to our knowledge, this is the first study to investigate this kind of eye-by-eye outcome, either employing canaloplasty or other MIGS procedures.

The reason that canaloplasty appears to be, to varying degrees, effective in most eyes (91% of the cohort) may be related to its comprehensive approach. Canaloplasty improves outflow facility via a combination of mechanical, hydrostatic, and biophysical mechanisms: the 360° catheterization of Schlemm's canal mechanically breaks adhesions within the canal while pushing herniations of the trabecular meshwork out of the collector channel ostia, thus improving outflow facility. The hydrostatic pressure caused by the delivery of OVD stretches the trabecular meshwork, possibly creating microperforations into the anterior chamber, while also dilating Schlemm's canal and the collector channels.^{24,25} These obstructions within the trabecular meshwork may account for up to 75% of the total outflow resistance in cases of POAG, and herniations of the trabecular meshwork into the collector channels are a significant contributor to outflow resistance.^{9,26} Research by Gong et al has shown that up to 90% of the collector channels in POAG eyes are blocked by herniations of the trabecular meshwork²⁷ and thus represents another important cause of outflow resistance, in addition to Schlemm's canal, which is significantly shorter, narrower, and often collapsed in POAG eyes.^{28,29} This can be attributed to a better understanding of the disease pathology along with the fact that, since outflow obstruction can reside across all levels of the conventional outflow pathway, including the trabecular meshwork, Schlemm's canal, and the collector channels, glaucoma treatments which employ a more comprehensive approach may offer more significant utility with respect to treatments that are focal in their approach.^{24,25,30}

Indeed, other MIGS such as Trabectome (NeoMedix Corporation, Tustin, USA), the iStent, and the Hydrus Microstent, target the trabecular meshwork in isolation.^{31,32} The Trabectome ablates the trabecular meshwork and inner wall of Schlemm's canal using an electrosurgical pulse while the iStent works as a trabecular microbypass from the anterior chamber into Schlemm's canal.³³ The Hydrus is inserted into Schlemm's canal to improve outflow from the anterior chamber to Schlemm's canal.³⁴

A retrospective study conducted by Arnljots et al on patients undergoing iStent implantation versus Kahook Dual Blade (KDB) goniotomy found that standalone iStent implantation could reduce IOP to <19 mmHg in 57% of cases (8/

14), while the KDB goniotomy successfully reduced IOP to <19 mmHg in 77% of cases (10/13).⁷ In the COMPARE study, 64.4% of eyes were implanted with the Hydrus device (Ivantis) and 57.3% of the eyes implanted with 2 iStent (Glaukos) were ≤ 18 mmHg postoperatively (41.3% and 44.2% respectively preoperatively).³⁵ In the current study, 78% of eyes (50/64) which underwent canaloplasty had a postoperative IOP ≤ 18 mmHg at 12 months with canaloplasty alone – compared to 16% preoperatively. Furthermore, in this study, canaloplasty achieved a $\geq 20\%$ reduction in IOP in 78% of eyes that reached the 12-month follow-up (45/58). In the COMPARE study, at 12 months, the percentage of eyes ≤ 18 mmHg without medications was 30.1% in the Hydrus group and 9.3% in the iStent group.³⁵ In this study with canaloplasty 34% of eyes had IOP of ≤ 18 mmHg without medications.

This suggests that canaloplasty can return consistent results and that its postoperative IOP outcomes are grouped in a more homogenous manner—below the “safe” limit of 18 mmHg and regardless of the mean IOP reduction of the cohort.

Theoretically, the reason may lie in the design of the procedure: a focal-based MIGS procedure such as an iStent, which bypasses the trabecular meshwork, will be effective in overcoming outflow resistance localized in the trabecular meshwork.³³ In those patients where outflow resistance is located distally to the trabecular meshwork, either in Schlemm’s canal or the collector channels, stenting the eye may not be effectual. As stated earlier, the outflow obstruction in glaucoma occurs at all levels in the proximal as well as distal outflow pathway^{24,25,30} and bypassing only the trabecular meshwork may lead to inconsistent results despite a successful procedure on the surgical table.

In addition, common postoperative complications of stent mispositioning or stent occlusion may hinder the effectiveness of stent-based MIGS on a per patient basis. According to a 2019 meta-analysis, the reported rate of further surgical intervention was in the range of 4.5–11.3% of study subjects.³⁶ Another meta-analysis reported that 22.5% of the eyes that received iStent implantation experienced adverse events, of which the most common were IOP elevation or spikes, stent blockage, obstruction, and malpositioning.³⁷ Comparatively, in this study, 6 eyes (6/64; 9%) were excluded from the analysis because they received additional glaucoma surgery: although some of them achieved an IOP reduction, the overall diagnosis recommended further glaucoma surgery.

This is not to suggest that stents are not effective in reducing IOP, because they are and there is a solid foundation of literature that proves significant and sustained IOP reduction in the long term, but that their effectiveness may vary greatly from eye to eye due to the multi-level nature of the outflow obstruction in glaucomatous eyes, which can occur at the trabecular meshwork, Schlemm’s canal, and the collector channels.

A comparison of study results regarding stents^{35,38–47} shows a similar trend, with a consistent reduction in mean IOP being observed in all studies while the standard deviation in postoperative IOP indicates a wide degree of variability, ranging from as low as ± 1.8 to as high as ± 5.2 , while in the present cohort a postoperative standard deviation of ± 2.9 was observed. A higher standard deviation points to the fact that reduction in IOP is not consistent in all patients, leading to variability in outcomes. This also shows that mean IOP reduction may not be a reliable approach to measure repeatability and reliability in treatment outcomes, especially when it comes to IOP control.

Every MIGS has its place in the glaucoma paradigm and the true value of MIGS is that it enables the surgeon to tailor a procedure that is best suited to each individual patient. Regarding canaloplasty for instance, this is not always preferable, namely when the trabecular meshwork is too fibrotic, or when the surgeon expects complicated cataract surgery.

The major limitation of this study is in the retrospective design and selection bias, as well as the lack of randomization. Another limitation is the lack of a control group to compare the results.

In conclusion, glaucoma is a multifactorial disease that requires a comprehensive treatment approach. Canaloplasty performed via an ab-interno surgical technique offers a minimally invasive procedure that targets and treats the complete outflow pathway and across the full 360 degrees, rather than treating an isolated point of the outflow pathway. This allows for results that are more predictable and consistent across all stages and categories of POAG patients.

Data Sharing Statement

All data generated or analysed for this study are included in the article. Further enquiries can be directed to the corresponding author.

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Disclosure

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