Trifocal diffractive intraocular lens implantation in patients after previous corneal refractive laser surgery for myopia

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Abstract

**Background:** With the difficulties in IOL power calculation and the potential side effects occurring postoperatively, multifocal IOL implantation after previous corneal refractive surgery are rarely reported especially for the trifocal IOL. Herein we report the clinical observation of trifocal IOL implantation in patients with previous myopia excimer laser correction. In this study, a multi-formula average method was performed for the IOLs power calculation to improve the accuracy. Visual and refractive outcomes were analyzed, and the subjective quality of patients’ life was evaluated by questionnaires survey.

**Methods:** This retrospective case series included patients with previous myopia excimer laser correction who underwent femtosecond laser assisted phacoemulsification and trifocal IOL (AT LISA tri 839 MP) implantation. Follow-up was done at 1-day, 1-month and 3-month to assess the visual outcomes. Outcome measures were uncorrected distance, intermediate and near visual acuity (UDVA, UIVA, UNVA), manifest refraction, defocus curve, and subjective quality of vision.

**Results:** 16 patients (14 eyes with previous laser in situ keratomileusis and 7 eyes with previous photorefractive keratectomy) were included. Mean postoperative spherical equivalent (SE) at 3-month was $-0.56 \pm 0.49$ SD, wherein, 48% of eyes were within $\pm 0.50$ D of emmetropia, and 91% were within $\pm 1.0$ D. Mean monocular UDVA, UIVA and UNVA (logMAR) at last visit were $0.02 \pm 0.07$, $0.10 \pm 0.10$, and $0.15 \pm 0.11$ respectively. Three patients (19%) reported halos and glare in postoperative 3 months, two of them needed to use spectacles to improve the intermediate visual acuity. Fifteen patients (94%) reported a satisfaction score of $\geq 3.5$ out of 4.0, and thirteen patients (81%) did not need spectacles at all distances. Mean composite score of the VF-14 questionnaire was $95.00 \pm 7.29$ out of 100.

**Conclusions:** Trifocal IOL implantation after myopia excimer laser correction could safely achieve a full range of adequate vision and accurate refractive outcomes.

**Background**

Corneal refractive laser surgery that began in the 1990s has been well developed and widely applied in ametropia correction for millions of patients. Over time, these patients eventually become presbyopic or cataractous and are more demanding for the freedom from glasses [1]. To that end, a further refractive procedure is required such as multifocal intraocular lens (IOL) implantation. With the continuous renewal and improvement of surgical equipment and implantation material, multifocal IOL implantation after cataract surgery has been increasingly applied in clinical practice and effectively restored visual acuity, thus becoming an alternative of refractive procedure [2−7]. Recently, some studies have shown that the multifocal IOL implantation could be safely and efficiently used for patients who had previously undergone corneal refractive laser surgery [8−11]. Obviously, the modern cataract surgery combining an intraocular lens implantation gradually becomes a refractive procedure rather than a treatment only for blindness. However, due to the difficulties in IOL power calculation and the potential side effects occurring postoperatively such as glare or other visual acuity problems, only few reports of the clinical observation
of multifocal IOL implantation after previous corneal refractive laser surgery exist, and even less for the trifocal IOL implantation.

Currently, bifocal IOLs are used as the major multifocal IOLs with only near and far focus and hence provide good distance and near visual acuity and moderate intermediate vision. As a newly emerging product from IOL multifocality technological development, trifocal IOLs were designed to provide three useful focal distances (near, intermediate and far) and thus improve the intermediate vision without impairing distance and near visual acuity [12−16]. An increasing number of reported clinical observations have demonstrated the positive outcome after trifocal IOL implantation and recent systematic reviews also showed the advantages of trifocal IOLs in comparison with bifocal IOLs [17,18]. Within this trend, trifocal IOLs would become a reliable alternative option for cataract patients with previous corneal refractive laser surgery in the foreseeable future. So far, there are few studies reporting the visual outcomes of trifocal IOL implantation after previous corneal refractive surgery. Wang et. al. recently reported on a trifocal IOL implantation in eyes with nuclear cataract about 15 years after myopic LASIK surgery, which restored vision efficiently [19]. Chow and co-worker thereafter reported a case series of trifocal IOL implantation for post-myopic LASIK patients providing a good visual outcome at both near and distance vision [20]. To further confirm the clinical effect of trifocal IOL implantation after previous corneal refractive surgery, we herein present a retrospective case series of trifocal IOL (AT LISA tri 839MP) implantation in eyes with previous corneal refractive laser surgery for myopia. In this study, a multi-formula average method was performed for the IOLs power calculation to improve the accuracy. The visual and refractive outcomes were analyzed, and patient satisfaction and subjective quality of vision were also evaluated using VF−14 questionnaire and another short questionnaire concerning negative visual symptoms.

Materials And Methods

This retrospective case series included patients who had previously undergone LASIK or PRK for myopia and underwent femtosecond laser assisted phacoemulsification and trifocal IOL (AT LISA tri 839MP, Carl Zeiss Meditec AG, Jena, Germany) implantation in AIER Group’s Eye Hospitals (Beijing Aier-Intech Eye Hospital, Guangzhou Aier Eye Hospital, Chongqing Aier Eye Hospital, Wanzhou Aier Eye Hospital and Shenzhen Aier Eye Hospital) from January 2017 to May 2019. Ethics Committee approval by the Institutional Review Board of the Aier School of Ophthalmology of Central South University was obtained for the present study protocol that adhered to the tenets of the Declaration of Helsinki.

The inclusion criteria consisted of eyes (1) that had surgery indication for cataract surgery without any contraindications of ocular surgical therapy in the preoperative examination, (2) with corneal astigmatism ≤ 1.5 D and the angles of Kappa and Alpha both being < 0.3 mm, and (3) had no complications of posterior capsular rupture or zonular dialysis during the cataract surgery. The exclusion criteria included eyes that (1) had decentered ablation (decentration > 0.5 mm), corneal scar, retinoschisis, haze, myopic retinopathy, retinoschisis or retinal detachment after myopia excimer laser correction, (2) with corneal...
astigmatism ≥ 1.5 D or irregular astigmatism ≥ 0.3 µm, and (3) had inflammation, glaucoma or other diseases that might affect the multifocal IOL implantation.

**Patient examinations**

Enrolled patients all underwent preoperative ophthalmologic examinations including corrected distance visual acuity (CDVA), UDVA, manifest refraction, retinal visual acuity, intraocular pressure, ocular A and B-scan ultrasonography, non-contact specular microscope, optical coherence tomography (OCT) and ray tracing aberrometry. The retinal visual acuity was assessed with Lambda 100 retinometer (Heine, Germany). The axial length, keratometry and anterior chamber depth were measured using the LenStar LS900 (Haag Streit, Switzerland).

The postoperative measurements at 1-day, 1-month and 3-month included CDVA, UDVA, UIVA, UNVA, and the subjective manifest refractions (spherical equivalent) as well as their changes. Through-focus monocular logMAR acuity (defocus curve) was also measured. The spectacle independence rate, satisfaction, and visual symptoms were recorded by VF–14 questionnaire and another short questionnaire concerning negative visual symptoms at 3-month postoperatively.

**Surgical technique**

All the surgeries were performed combining femtosecond laser-assisted phacoemulsification and intraocular lens implantation by experienced surgeons. For the eyes with corneal astigmatism between 0.75 D and 1.5 D (6 eyes), femtosecond laser-assisted corneal relaxing incision was performed for correction. Preoperatively, the surgeons used tropicamide to maintain pupil dilation intraoperatively. Under topical anaesthesia, capsulotomies (diameters were all set as 5.2 mm), lens fragmentation and corneal relaxing incisions were performed subsequently using the LenSx femtosecond laser (Alcon Laboratories, Inc, Fort Worth, Texas, USA). Phacoemulsification was then performed using standard ultrasound technique. The trifocal IOL (AT LISA tri 839MP) was implanted in the capsular bag using an injector. The residual ophthalmic viscosurgical device was removed, and the position of the lens was adjusted. All incisions were hydrated and the patients’ conjunctival sac was treated with dexamethasone-tobramycin ophthalmic ointment.

For lens power calculation, a multi-formula average method was performed, in which 4 formulas of Hagis-L [21], Barrett True K [22], Shammas No-History [23] and ray-tracing methods [24] setting the target refraction as postoperative emmetropia were used. The implanted IOL power was the average of the calculated results from all 4 formulas. For the first treated eye of patients who underwent bilateral implantation and the eyes with a large difference (> 1.0 D) in the IOL power calculations using the 4 formulas, a modified aphakic refraction technique was applied. The surgical procedure was done according to the original aphakic refraction technique reported by Dr. Mackool [25]. At first, the cataract
removal was performed without IOL implantation. 1 Day later, manifest refraction examination was performed for the lens power calculation followed by the IOL insertion.

Postoperative treatment included one drop of tobramycin-dexamethasone eye drops every two hours for 3 days, afterward four times per day until 2 weeks, and then three times per day for another 2 weeks; and one drop of levofloxacin eye drops four times every day for 1 week; pranoprofen and sodium hyaluronate eye drops were administered as appropriate.

Statistical Analysis

Statistical analysis was performed using Microsoft Excel. Mean (± SD) was reported for continuous variables. Normal probability plots and Kolmogorov-Smirnov and Shapiro-Wilk tests were used to check the normality of data in SPSS software. Pre- and post-UDVA, pre- and post-CDVA outcomes were compared with the t test. Nonparametric tests were used to evaluate differences within groups. Differences were considered statistically significant when the P value was less than 0.05.

Results

This study comprised 21 eyes of 16 patients (5 men, 31.3%). The demographics of the study population are summarized in Table 1. The mean patient age at the time of IOL implantation was 48.5 ± 8.9 years (range, 36 to 66 years). Eleven (68.7%) and five patients (31.3%) underwent unilateral and bilateral implantation respectively. No complications developed intraoperatively. No further laser enhancement was performed. The mean logMAR of UDVA and CDVA were 0.83 ± 0.48 (range, 0.22 to 1.70) and 0.37 ± 0.30 (range, 0.00 to 1.00), respectively. The mean axial length was 27.70 mm ± 2.20 (range, 25.13 to 33.00 mm). The mean preoperative spherical equivalent (SE) was −5.49 D ± 5.75 (range, −18.75 to +1.37 D).
## Table 1
Demographics of the patients before IOL implantation

| Item                                                                 | Number or Mean (± SD) |
|----------------------------------------------------------------------|-----------------------|
| Number of eyes (N)                                                   | 21                    |
| Number of patients (n)                                               | 16                    |
| Sex (male/female, n/n)                                               | 5/11                  |
| Refractive surgery procedures (numbers of eyes), (percentage)        |                       |
| PRK                                                                  | 7 (33.3%)             |
| LASIK                                                                | 14 (66.7%)            |
| Symptoms (numbers of patients), (percentage)                         |                       |
| Bilateral                                                            | 5 (31.3%)             |
| Unilateral                                                           | 11 (68.7%)            |
| Eyes with modified aphakic refraction technique                      | 11 (52.4%)            |
| Mean age (Years) ± SD (range)                                        | 48.5 ± 8.9 (36, 66)   |
| Mean pre-UDVA (logMAR) ± SD (range)                                  | 0.83 ± 0.48 (0.22, 1.70) |
| Mean pre-CDVA (logMAR) ± SD (range)                                  | 0.37 ± 0.30 (0.00, 1.00) |
| Mean pre-SE (D) ± SD (range)                                         | −5.49 ± 5.75 (−18.75, +1.37) |
| Mean axial length (mm) ± SD (range)                                  | 27.70 ± 2.20 (25.13, 33.00) |
| Mean keratometry (D) ± SD (range)                                    | 38.36 ± 2.27 (34.34, 41.94) |
| Mean power implanted IOL (D) ± SD (range)                            | 17.64 ± 3.87 (8.0, 23.00) |

IOL = intraocular lens; LASIK = laser in situ keratomileusis; PRK = photorefractive keratectomy; UDVA = uncorrected distance visual acuity; SE = spherical equivalent; pre- = preoperative.

## Refraction

After lens extraction and IOL implantation at 3-month, the mean SE was reduced to −0.56 D ± 0.49. Fig. 1A shows the change of mean SE at the three follow-up intervals (1-day, 1-month and 3-month). The histograms of the postoperative refractive accuracy are presented in Fig. 1B. Ten eyes (48%) were within ± 0.50 D of emmetropia and nineteen eyes (91%) within ± 1.0 D at postoperative 3 months. In contrast, only three eyes (14%) were within ± 1.0 D of emmetropia preoperatively (Fig. 1C). Fig. 1D shows the attempted versus achieved SE refraction. At 3-month follow-up, the overall rate of overcorrection was 0%, while the overall rate of undercorrection was 9.5% (2 eyes).
Visual Acuity

The mean postoperative CDVA was 0.00 ± 0.05 logMAR, the mean postoperative UDVA was 0.10 ± 0.16 logMAR at the first day postoperatively (Table 2). All the surgery eyes (21 eyes, 100%) achieved postoperative CDVA ≥ 20/25, while the percentage for UDVA ≥ 20/25 was 76% (for the postoperative cumulative Snellen visual acuity, see Fig. 2A). Additionally, the mean postoperative UNVA, UIVA and UDVA at 3-month were 0.15 ± 0.11 logMAR, 0.10 ± 0.10 logMAR and 0.02 ± 0.07 logMAR respectively. All obtained monocular visual acuity data at the last visit are summarized in Table 2.

Table 2
Monocular visual acuity (logMAR).

| Parameter | Preoperative | Postoperative at 1-day | P       | Postoperative at 3-months |
|-----------|--------------|------------------------|---------|--------------------------|
|           | Mean ± SD    | Range                  | Mean ± SD | Range          | Mean ± SD | Range |
| CDVA      | 0.37 ± 0.30  | 0.00, 1.00             | 0.00 ± 0.05 | -0.08, 0.10     | < 0.001 | \     |
| UDVA      | 0.83 ± 0.48  | 0.22, 1.70             | 0.10 ± 0.16 | -0.08, 0.52     | < 0.001 | 0.02 ± 0.07 | -0.08, 0.22 |
| UIVA      | \            | \                      | \        | \              | \        | 0.10 ± 0.10 | 0.00, 0.30 |
| UNVA      | \            | \                      | \        | \              | \        | 0.15 ± 0.11 | 0.00, 0.40 |

Defocus curve

Monocular defocus curve at 1-month (Fig. 2B, results of 20 eyes) showed best visual acuity (0.02 logMAR and 0.13 logMAR) with defocus of 0.00 D and −3.00 D, simulating distances of 4 m and 33 cm. In the range between the peaks, the curve transitioned smoothly and reached the bottom as 0.14 logMAR at −2.50 D. Besides the defocus of 1.50 D (with a visual acuity of 0.34 logMAR), all visual acuities tested at other defocus diopters were better than 0.23 logMAR, maintaining a functional range of visual acuity.

Questionnaires

Table 3 and 4 summarized the results of the VF–14 and supplementary questionnaires. Three patients (19%) reported halos giving the least mean scores of all visual symptoms in the questionnaire as 3.42, 3.50 and 3.64 respectively, and the symptoms were relieved unafflicting the life as time went on. Three patients (19%) reported the need to use spectacles only for the near visual applications such as reading...
books and doing handwork. Fifteen patients (94%) reported a satisfaction score of ≥ 3.50, and thirteen patients (81%) did not need spectacles at all distances.

Table 3
Three-month postoperative responses to visual function questionnaire (VF-14) items.*

| Item                                      | Score (mean) ± SD | Score multiplied by 25 | Spectacle independence rate |
|-------------------------------------------|-------------------|------------------------|-----------------------------|
| Reading small print                       | 3.06 ± 0.77       | 77                     | 81%                         |
| Reading a newspaper or a book             | 3.63 ± 0.62       | 91                     | 88%                         |
| Reading a large-print book or numbers on a telephone | 3.88 ± 0.34       | 97                     | 100%                        |
| Recognizing people when they are close to you | 4.00 ± 0.00       | 100                    | 100%                        |
| Seeing steps, stairs, or curbs            | 4.00 ± 0.00       | 100                    | 100%                        |
| Reading traffic, street, or store signs   | 4.00 ± 0.00       | 100                    | 100%                        |
| Doing fine handwork like sewing†         | 3.36 ± 0.50       | 84                     | 87%                         |
| Writing checks or filling out forms       | 3.81 ± 0.40       | 95                     | 100%                        |
| Playing games such as bingo, dominos, card games, mahjong | 4.00 ± 0.00       | 100                    | 100%                        |
| Taking part in sports like bowling, handball, tennis, golf | 3.94 ± 0.25       | 98                     | 100%                        |
| Cooking                                   | 3.94 ± 0.25       | 98                     | 100%                        |
| Watching television                       | 3.88 ± 0.34       | 97                     | 100%                        |
| Driving during the day‡                   | 4.00 ± 0.00       | 100                    | 100%                        |
| Driving at night‡                         | 3.50 ± 0.76       | 88                     | 100%                        |

* Score scale: 4, no difficulty; 3, a little difficulty; 2, moderate difficulty; 1, quite difficult; 0, impossible to perform; † N = 15; ‡ N = 8.
Table 4
Results of the short questionnaire concerning negative visual symptoms in postoperative 3 months

| Negative visual symptoms in postoperative 3 months | Number of patients | Percentage |
|---------------------------------------------------|--------------------|------------|
| Halos                                             | 3                  | 19%        |
| Glare                                             | 3                  | 19%        |
| Flare                                             | 0                  | 0%         |

Discussion

Previous studies of eyes after corneal refractive laser surgery have focused primarily on monofocal IOLs [22,26–28] and bifocal IOLs [8–11]. In our study, we present this retrospective case series of trifocal IOL (AT LISA tri839MP) implantation in patients who underwent a previous corneal refractive laser surgery for myopia. The AT LISA tri839MP used in our study is a monolithic diffractive trifocal intraocular lens which is made of collapsible hydrophilic acrylate (25%) with a hydrophobic surface. Its optical zone has a diameter of 6.0 mm, a total diameter of 11.0 mm, and a four-turn intraocular lens with a zero angle. The intraocular lens combines a bifocal diffraction range from 4.34 to 6.00 mm, and a trifocal diffraction range (over 4.34 mm) on the front surface of the lens, which is achieved based on the asymmetrical light split, that is 30%, 20%, and 50% of the incoming light is split at near focus, intermediate focus, and distant focus respectively. It was developed to overcome the photic phenomena and the poor level of intermediate vision of traditional multifocal IOLs [29] and has already been demonstrated to have good outcomes of visual acuity at near, intermediate and far distance and high postoperative satisfaction [30].

Patients were extensively counselled regarding the possible side effects of the treatment such as night glare, halos and decreased contrast sensitivity, even the possibility of deteriorated visual quality after a refractive corneal laser procedure. The final decision was made based on their wishes to restore vision and hence becoming independent of spectacles.

Intraocular lens implantation in eyes with previous corneal refractive laser surgery is full of challenges due to the difficulty in IOL power calculation [26,31,32]. To date, some reported IOL power calculation methods have been used for IOL implantation in eyes with previous corneal refractive laser surgery providing relatively accurate outcomes [27,33,34]. However, obvious hyperopia is often observed postoperatively [35,36]. For the lens power calculation in the present study, a multi-formula average method was applied which has been described and proven to be accurate in our previous study [37]. Besides that, a modified aphakic refraction technique was used for the first operative eyes of patients who underwent bilateral implantation and the eyes with large difference (> 1.0 D) in the IOL calculated results from multi-formulas. Both these improvements were made based on the previous investigation and to pursue accuracy in IOL power calculation.
In the refraction outcome, the mean postoperative SE was reduced to \(-0.56 \text{ D } \pm 0.49\) while Chow’s study reported a \(-0.92 \text{ D } \pm 0.76\) with a significant myopic shift [20]. In addition, 48% of eyes were within \(\pm 0.50\) D of emmetropia and 91% within \(\pm 1.0 \text{ D}\) at postoperative 3 months in our study. All our cases were targeted for emmetropia, and the results were similar to Vrijman’s study with a lower percentage of eyes with \(\pm 0.50 \text{ D}\). This is because most of the patients included were preoperatively high myopia with either less spherical equivalent (9 eyes, 43%, with \(< -6.00 \text{ D}\)) or longer axial length (16 eyes, 76%, with \(> 26.00 \text{ mm}\)). It is consistent with Vrijman’s finding that results were less predictable in eyes with myopia greater than 6.0 D [10].

The results of vision outcomes show a statically significant improvement after surgery. The mean postoperative CDVA and UDVA was \(0.00 \pm 0.05\) and \(0.10 \pm 0.16\) (logMAR), respectively. Both these two outcomes are significantly better than preoperative outcomes (Table 2, \(P < 0.001\)). The mean value of postoperative CDVA agrees with the result of Chang and colleagues (20/19, Snellen Vision) by converting into Snellen Vision [11]. To accurately evaluate the visual outcomes of trifocal IOLs after previous corneal refractive surgery, the monocular UDVA, UIVA and UNVA were recorded at last follow up in our study, and the mean values (logMAR) were \(0.02 \pm 0.07, 0.10 \pm 0.10\) and \(0.15 \pm 0.11\) respectively. However, among the few reports using multifocal IOL implantation, only Chang’s study recorded the same visual outcomes wherein the mean UIVA was \(0.22 \pm 0.15\) (logMAR) as slightly higher than the value we observed.

As an important indicator for the vision over the entire range, defocus curve was also recorded in this study. In 2015, Jonker et. al. compared the defocus curves after trifocal IOL implantation with that after bifocal IOL implantation in cataract patients [13]. In the comparison, the defocus curves of the trifocal IOL group showed a more continuous performance at the intermediate range under photopic and mesopic conditions. In the defocus curve of our study, the curve transited smoothly in the range between the peaks with defocus of \(0.00 \text{ D}\) and \(-3.00 \text{ D}\) and reached the bottom as \(0.14 \text{ logMAR}\) at \(-2.50 \text{ D}\). Besides the defocus of \(1.50 \text{ D}\) (with a visual acuity of \(0.34 \text{ logMAR}\)), all visual acuities tested at other defocus diopters were better than \(0.23 \text{ logMAR}\), maintaining a functional range of visual acuity. These results further demonstrate the good and wide range visual outcomes of trifocal IOL implantation after previous corneal refractive laser surgery for myopia.

In terms of the questionnaires survey, three patients (19%) reported halos and glare in postoperative 3 months, while no patient reported flare. Two of these three patients needed to use spectacles to read small print or a newspaper or book or to do some handwork. Looking back on the preoperative parameters of the two patients, both of their surgery eyes were with high myopia (preoperative spherical equivalents are \(-7.60\) and \(-12.37\) respectively). It seems that the capsule of high myopic patients is larger, and the stability of intraocular lenses is slightly worse compared to normal patients. The position and functional deviation of intraocular lens might lead to bad visual symptoms such as halos. In the other hand, the preoperative symptoms of halos or glare, and the status of the fellow untreated eye from patients who was performed unilateral surgery, might affect the postoperative visual outcomes and interfere with the clinical analysis of the efficiency of trifocal IOL implantation. Unfortunately, all the information had not been recorded preoperatively. Nevertheless, the total spectacles independence rate is
81% (thirteen patients), and those patients without halos or glare all have scores better than 3.67. Overall, the questionnaires survey showed good quality of life postoperatively in the current study.

The limitation of this study was the small sample size. Undoubtedly, a larger sample size would be more helpful to shed light onto the efficiency of trifocal IOL implantation after previous corneal refractive laser surgery for myopia. However, due to the uncertain side effects of mutifocal IOL implantation after previous corneal surgery, the number of patients who balanced the side effects by their demands for becoming independent of spectacles are rare.

Conclusions

In conclusion, the trifocal intraocular lens can safely provide a full range of adequate vision and accurate refractive outcomes for cataract patients after myopic excimer laser correction. The high spectacle independence rate affords the patients a high satisfaction. Only few side effects such as halos and glare were observed and are possibly caused by extremely high myopia in few of the patients.

Abbreviations

IOL: Intraocular lens; UDVA: Uncorrected distance visual acuity; UIVA: Uncorrected intermediate visual acuity; UNVA: Uncorrected near visual acuity; LASIK: Laser in situ keratomileusis; PRK: Photorefractive keratectomy; SE: Spherical equivalent; D: Diopter; SD: Standard deviation; logMAR: Logarithm of the minimal angle of resolution; CDVA: Corrected distance visual acuity; OCT: Optical coherence tomography.

Declarations

Authors’ contributions

Both QML and FW contributed equally to this work. SWL and KS conceived of and designed this study. SWL, ZMW, ZL, CZ, BHC, JS performed the surgeries and contributed to the data acquisition. QML performed the patient follow-up and data collection. QML and FW conducted independently the data analysis and rechecking. FW drafted the initial manuscript. FW, SWL and KS revised the manuscript critically for important intellectual content. This manuscript has been read and approved by all the authors and each author believes that the manuscript represents honest work.

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Availability of data and materials
The analytic data used to support the findings of this study are included within the article. Other original data used to support the findings of this study are available from the corresponding author upon request.

**Ethics approval and consent to participate**

Ethics Committee approval by the Institutional Review Board of the Aier School of Ophthalmology of Central South University was obtained for the present study protocol that adhered to the tenets of the Declaration of Helsinki.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no conflicts of interest.

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Figures
Figure 1

Refraction outcomes. (A) Changes of mean SE at the three follow-up intervals (1-day, 1-month and 3-month); (B) Histograms of the postoperative refractive accuracy; (C) Percentages of eyes within ± 0.5 D and ± 1.0 D of emmetropia; (D) Spherical equivalent attempted versus achieved.

Figure 2

Visual outcomes. (A) Postoperative cumulative Snellen visual acuity; (B) Monocular distance-corrected defocus curve given in logMAR at 1-month postoperatively (N = 20).