ORIGINAL RESEARCH

Clinical Outcomes and Patient Satisfaction with a New Diffractive-Refractive Trifocal Intraocular Lens – A 12 Month Prospective Study

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Purpose: To evaluate the clinical outcomes and patient satisfaction after implantation of Optiflex Trio, a new trifocal intraocular lens (IOL) following cataract surgery.

Methods: Patients undergoing phacoemulsification for age-related cataracts and who satisfy the eligibility criteria underwent bilateral implantation with Optiflex Trio trifocal IOL. At follow -up visits of 1, 3, 6, and 12 months, binocular uncorrected and corrected distance, intermediate and near visual acuity, reading performance, contrast sensitivity (CS) and patient satisfaction for dysphotopsia and spectacle independence were evaluated using questionnaires.

Results: A total of 54 eyes from 27 patients with mean age of 66.30±7.48 years were included in the study. At 12 months, 78% (n = 21) patients had binocular cumulative UDVA of 20/20 or better. Post-op SE refraction accuracy was within ± 0.50 D for 93% (n = 50) eyes, and refractive cylinder accuracy was within ≤ 0.50 D in 94% (n = 51) eves. The mean binocular UNVA was 0.01±0.05 LogMAR, and the mean UIVA at 60 and 80 cm was 0.07 ± 0.06 and 0.03 ± 0.05 LogMAR, respectively, at 12 months. Reading speeds at 40, 60 and 80 cm showed improvement overtime. No patient had complained of severe dysphotopsia, and none of the patients required glasses for any activity. No eye underwent YAG-laser capsulotomy for significant PCO at the end of mean follow-up.

Conclusion: After 12 months, Optiflex Trio trifocal IOL provided a complete visual restoration with good visual quality outcomes in terms of uncorrected distance, intermediate and near visual acuity. The incidence of dysphotopsia was low, and spectacle independence was high, resulting in good patient satisfaction.

Trial Registry: CTRI/2019/10/021647 (www.ctri.nic.in). Keywords: Optiflex, trifocal, IOL, spectacle independence, intraocular lens

Introduction

Trifocal intraocular lenses (IOLs) were developed to bridge the gap between monofocal and bifocal multifocal IOLs in order to provide significant advantages in terms of post-operative unaided intermediate vision, due to the addition of a third foci in the diffractive platform of the optic of the lens.^{1–3}

Ever since the introduction of the first trifocal IOL from Finevison,⁴ various trifocal IOLs evaluated in the past were shown to successfully reduce spectacle dependence without compromising the quality of vision (QOV) in patients desiring freedom from glasses following cataract surgery for age-related cataracts.^{5–7}

Optiflex Trio trifocal IOL (Biotech Healthcare Holding GmbH, Obergrundstrasse 17, 6002 Luzern, Switzerland), is a relatively recent introduction in the field of trifocal

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Received: 14 May 2021 Accepted: 5 July 2021 Published: 3 August 2021 intraocularIOL technology. The IOL is a single piece, hydrophobic acrylic, aspheric, IOLs containing natural chromophore, with a 360° square edge. According to the best of our knowledge, the clinical outcomes with this model of trifocal IOL have not been evaluated yet. In this study, we have reported one-year clinical outcomes related to safety, efficacy, predictability, contrast sensitivity (CS), reading performance, patient satisfaction, complications and overall results with this new trifocal IOL.

Materials and Methods

This prospective, single-centre single arm study was approved by the institutional ethics committee of Nethradhama Superspeciality Eye Hospital, Bangalore, and was conducted in accordance with the principles of the Declaration of Helsinki. All patients provided written informed consent.

Inclusion criteria were healthy eyes besides senile cataract, corneal astigmatism ≤ 0.75 dioptres (D), IOL power calculation resulting in dioptres between +7.0 D to +30.00 D, capsular bag IOL implantation, and ability to read English language fluently.

Exclusion criteria were patients with corneal astigmatism of >0.75 D, irregular astigmatism due to keratoconus, pellucid marginal degeneration, or corneal scars, corneal dystrophy, severe ocular surface disorders, pupillary abnormalities, history of glaucoma, intraocular inflammation, macular degenerations or retinopathies potentially affecting visual outcome, vulnerable subjects, neuroophthalmic diseases, intraoperative complications such as posterior capsule rupture, nucleus drop or capsular bag loss precluding the implantation of the planned IOL, and unassured follow-ups.

Pre-operatively, all patients underwent complete ophthalmologic examination including measurement of uncorrected and best distance corrected visual acuity (ETDRS charts, Precision Vision, LaSalle, IL, USA), manifest refraction, slit lamp biomicroscopy, non-contact tono-(Tomey NCT, NishiKu, Nagoya, metry Japan), tomography using elevation based Scheimpflug imaging device Pentacam HR (Oculus Optikgeräte GmbH, Wetzlar, Germany) to rule out irregular astigmatism, HD Analyzer (Visiometrics, Spain) to assess ocular dryness, specular microscopy (Tomey, Japan), macular OCT (Optovue, Fremont, USA) and dilated fundus examination. Biometric assessments were performed using a swept source OCT based optical biometer, IOL Master-700 (Carl Zeiss Meditec, Jena, Germany) using Barrett TK Universal II formula. All eyes were targeted at emmetropia. An optimized A-constant of 118.5 was used for IOL power calculation.

Description of the Study IOL

The Optiflex Trio trifocal IOL is a single piece, hydrophobic acrylic, aspheric, diffractive-refractive trifocal IOL containing natural chromophore, with a 360° square edge for prevention of posterior capsular opacification (Table 1, Figure 1). Lens material with natural yellow chromophore prevents the risk of ARMD, does not disturb the circadian rhythm and does not attribute to altered color perception. The optic of the lens is aspheric with negative spherical aberrations (S.A.) of -0.2μ . The optic of the IOL is diffractive-refractive trifocal with a near add of +3.5 D, and

	Table I	Description	of Optiflex	Trio IOL	Specifications
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Material	Hydrophobic Acrylic Containing Natural Chromophore
Optic type	Single piece, 360° square edge with aspheric optic, Diffractive- Refractive, Negative aspheric
Optic size	6.00 mm
Haptic design	Optimized C haptic design
Overall size	13.00 mm
Angulation	0°
Diffractive surface	Anterior
Zone	4mm Diffractive, Peripheral – Refractive Zone
Near Addition	+3.5D
Intermediate Addition	+1.85D
ACD	5.28 mm
Refractive index	1.48
Dioptre range	+7.00 to +30.00 D in 0.5 D steps
A constant (Optical)	118.5
Light distribution	45% for Far, 27% for Intermediate and 28% for near distances
Abbe Number	49
Injector	Optiject BES28 Injector system
Implantation site	Capsular bag
Sterilization	Irradiation

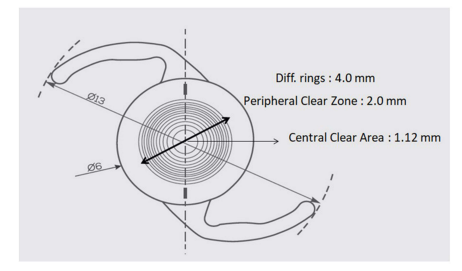


Figure I Representative image showing the design of the Optiflex Trio trifocal IOLs. Notes: Reproduced with permission from Biotech Healthcare Holding GmbH.

an intermediate add of +1.85 D on the IOL plane. Optimum Asymmetric and balanced light distribution of 45% for far, 27% for intermediate and 28% for near, is designed to provide good unaided vision at all distances.

Surgical Procedure

All surgeries were performed by a single experienced surgeon (S.G.), using a standard phacoemulsification technique under topical anaesthesia, using the Centurion Precision system (Alcon Laboratories, Fort Worth, TX, USA). Through a temporal clear corneal incision of 2.8 mm, a 5.0 -5.5 mm capsulorhexis was aimed and direct chop technique was used for nuclear deployment. After irrigation and aspiration of the cortex, the left side port was hydrated and BSS injected from the main wound to inflate the bag and form the anterior chamber. Followed by this, the Optiflex Trio trifocal IOL was injected into the capsular bag using its dedicated injector system (Optiject BES28 system). Supplementary Video S1 shows the surgical video of loading and implantation of the Optiflex Trio trifocal IOL, in the right eye of one of the study participants. Any device related to intra-operative complications such as haptic or optic breakage, or explantation of the IOL due to device damage or incorrect IOL power, were recorded.

Post-operative topical therapy included topical prednisolone (1%, Pred Forte, Allergan) 6 times for 6 weeks tapering weekly, moxifloxacin (0.5%, Vigamox, Alcon) 4 times for 2 weeks, nepafenac (0.1%, Nevanac, Alcon) 3 times for 4 weeks and lubricants 4 times or SOS for 4 weeks or more.

Follow-up examinations were performed at 1 day, 1 week, 1 month, 3 months, 6 months, and 12 months after surgery. Dilated slit-lamp examination was performed on post-op day 1 to assess the corneal clarity, anterior chamber inflammation and IOL position. From one month onwards, in addition to the above, assessment of manifest refraction, uniocular and binocular uncorrected and corrected distance visual acuity (UDVA, CDVA), uniocular and binocular uncorrected and corrected near visual acuity (UNVA, CNVA), photopic CS using CSV-1000 (Vector Vision, Greenville, Ohio), reading performance using Salzburg Reading Desk (SRD, University Eye Clinic, Paracelsus Medical University of Salzburg, Austria), defocus curve charting from +2 to -4 D. From 3 months onwards, and a QOV and patient satisfaction questionnaire regarding dysphotopsia symptoms and spectacle independence was also obtained.

Statistical Analysis

SPSS software for Windows version 17.0.0 (IBM Corp., Armonk, NY) was used for statistical analysis. All values were expressed as mean \pm standard deviation (SD). Data was checked for normality before subjecting to analysis. A p-value of 0.05 or less was considered statistically significant. Outcome analysis was performed according to the Standard Graphs for Reporting Refractive Outcomes following Intraocular Lens-Based Refractive Surgery.

Results

Fifty-four eyes of 27 patients were evaluated in the study. Table 2 shows the demographic profile and baseline pre-

	Table 2 Demographics and Baseline Pre-	Operative Parameters
,	of All Eyes (n = 54) Included in the Study	

Parameter	Mean± SD
Age (years)	66.30±7.48
Male: Female	12 Males, 15 Females
KI (D)	43.51±1.39
K2(D)	44.26±1.47
Astigmatism (D)	0.74±0.38
Axial length (mm)	23.35±0.75
ECD (cells/mm ²)	2618.59±186.19
IOL Power (D)	21.17±1.611

Abbreviations: K, keratometry; D, dioptre; ECD, endothelial cell density; IOL, intraocular lens; SD, standard deviation.

operative parameters of the eyes included in the study. Table 3 shows the visual and refractive outcomes evaluated for distance and near vision at all post-operative visits of 1 week, 1, 3, 6 and 12 months.

Visual Outcomes

At 12 months, the mean binocular UDVA and CDVA were -0.00 ± 0.07 and -0.06 ± 0.04 LogMAR respectively. There was no significant difference in the binocular UDVA and CDVA at 12 months compared to post-op visits of 1 week, 1 month, 3 months, and 6 months (p-values for change in UDVA and CDVA >0.05), Table 3.

Uni-ocularly, 59% (32) eyes had UDVA of 20/20 or better, while all eyes (54) had a minimum UDVA of 20/32 (Supplementary file 1). Binocularly, 78% (21) patients had UDVA of 20/20 or better, while all patients (27), had a minimum UDVA of 20/25 (Figure 2). Fifty-two percent (28) of eyes had post-op UDVA same as post-op CDVA, whereas 35% (19) of eyes had post-op UDVA better than post-op CDVA by 1 line and 13% (7) eyes had post-op UDVA better than post-op CDVA by 2 lines (Supplementary file 2).

The mean binocular UNVA at 12 months was 0.01 ± 0.05 LogMAR and mean binocular DCNVA was -0.01 ± 0.03 LogMAR, both the parameters were not statistically significantly different from the post-op values at 1 week, 1 month, 3 months, and 6 months (Table 3). Ninety-seven percent (n = 26) of patients had binocular uncorrected near vision of N6 or better, while all patients had a minimum UNVA of N8 (Figure 3).

Refractive Outcome

At 12 months, the mean sphere, cylinder and SE was -0.07 D±0.19, -0.18 D±0.23, and -0.16 D±0.21, which was not significantly different from their respective values at post-op 1 week, 1 month, 3 months and 6 months (p-values> 0.05 for all parameters), Table 3. Ninety-three percent (50) of eyes were within ±0.50 D of SE correction, while 94% (51) eyes were within ±0.50 D of cylinder correction (Figure 4).

Intermediate Visual Outcomes at 60, 80 Cm Using ETDRS Charts

At 12 months, the mean uncorrected visual acuity at 60 cm was 0.07 ± 0.06 LogMAR and at 80 cm was 0.03 ± 0.05 LogMAR. No significant differences were found between uncorrected and corrected intermediate values at 60 and 80 cm, and also when these were compared to the previous follow-ups, i.e., 1 week, 1 month, 3 months, and 6 months (Table 4).

Reading Speeds

The binocular uncorrected reading speeds (words per minute) assessed with Salzburg Reading Desk (SRD) at 40, 60 and 80 cm showed improvement from 1 week to 12 months, however, it was not statistically significant. A similar trend was observed for corrected reading speeds for all distances (Table 4).

Parameter (Mean± SD)	I-Week	I-Month	3-Months	6-Months	12 -Months	p-value
UDVA (LogMAR)	0.06±0.09	0.06±0.09	0.04±0.09	0.04±0.08	0.04±0.08	0.54
CDVA (LogMAR)	-0.00±0.04	-0.01±0.04	-0.02±0.04	-0.02±0.04	-0.02±0.04	0.56
Sph (D)	-0.07±0.28	-0.07±0.22	-0.07±0.19	-0.07±0.19	-0.07±0.19	0.99
Cyl (D)	-0.26±0.32	-0.24±0.28	-0.21±0.27	-0.19±0.24	-0.18±0.23	0.52
SE (D)	-0.20±0.31	-0.19±0.26	-0.17±0.23	-0.17±0.22	-0.16±0.21	0.95
Binocular UNVA (LogMAR)	0.02±0.05	0.01±0.05	0.01±0.05	0.01±0.05	0.01±0.05	0.97
Binocular DCNVA (LogMAR)	-0.00±0.03	-0.01*±0.03	-0.01±0.03	-0.01±0.03	-0.01±0.03	0.88
Binocular UDVA (LogMAR)	0.00±0.07	0.00±0.07	0±0.07	0±0.07	-0.00±0.07	0.95
Binocular CDVA (LogMAR)	-0.05±0.05	-0.05±0.05	-0.06±0.05	-0.06±0.04	-0.06±0.04	0.96

Abbreviations: UDVA, uncorrected distance visual acuity; CDVA, corrected distance visual acuity; Sph, sphere; Cyl, cylinder; SE, spherical equivalent; IOL, intraocular lens; UNVA, uncorrected near visual acuity; DCNVA, distance corrected near visual acuity; D, dioptre; SD, standard deviation.

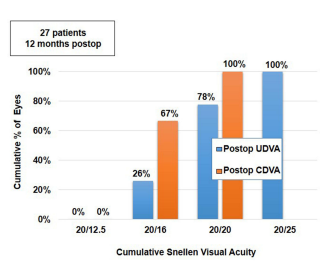


Figure 2 Histogram showing binocular results for UDVA and CDVA obtained following implantation of Optiflex Trio IOL at 12 months post-operatively.

Contrast Sensitivity

Photopic CS evaluated binocularly at 1 week, 1 month, 3 months, 6 months, and 12 months showed improvement in the log values of all spatial frequencies over time; however, this was not statistically significant (Table 4, Figure 5).

Defocus Curve

Binocular defocus curves were charted with correction at 1 week, 1 month, 3 months, 6 months, and 12 months using defocusing lenses from +2.00 to -4.00. At 12 months postop, the defocus curve showed two peaks corresponding to 0.00 D and -2.5D, with a gradual decline of the slope in the intermediate range of vision (-1.50 D) (Figure 6). Overall, a full range of functional vision (20/32 or better), was maintained through a defocus of 4.5D.

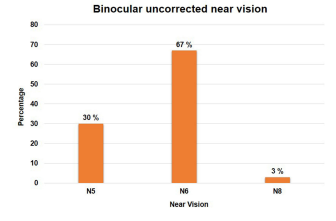


Figure 3 Binocular uncorrected near vision results at 12 months post-op.

Endothelial Cell Count

Endothelial cell count (ECC) was calculated using specular microscopy. Mean ECC at pre-op, 6 months and 1 year were 2618 ± 186 , 2591 ± 362 , 2576 ± 96 cells/mm², respectively, and were found to be non-significant (p value: 0.20).

Patient Satisfaction & Spectacle Independence

A QOV questionnaire was obtained for all patients from 3 months onwards. The mean score of dysphotopsia symptoms (graded from 0–10, 0 being minimal and 10 being severe) reduced significantly at 12 months (0.61 ± 0.49) when compared to 3 months (3.11 ± 0.43), p = 0.00. Spectacle independence scores (graded from 0–10, 0 being dependent on glasses and 10 being completely glass free) for distance, intermediate and near vision were 9.56 ± 0.09 , 9.55 ± 0.09 and 9.25 ± 0.36 respectively at 12 months. Overall patient satisfaction score (graded in percentages from 0–100%, 0 being not satisfied at all and 100 being fully satisfied) was 97.07 ± 2.23 at the end of 12 months (Supplementary file 3).

Adverse Effects and Complications

Dilated clinical examination was performed at 3, 6, and 12 months to assess optical clarity of the IOL in terms of glistenings, opacification, calcification, or posterior capsule opacification (PCO), and IOL centration. All IOLs were found to be well-centered in the bag with 360° overlap of capsulorhexis and without any significant tilt or decentration. None of the eyes IOL glistening, calcification, or PCO affecting the visual outcomes or patient satisfaction at the last follow-up. No other vision threatening complications such as cystoid macular oedema, postoperative uveitis, secondary glaucoma or endophthalmitis occurred in any of the eyes included in the study.

Discussion

Several studies have evaluated the clinical outcomes of various trifocal IOLs.^{4,5,7,8} In the present study, we evaluated the results of Optiflex Trio, which is a recently introduced trifocal IOLs in the market. Table 5 shows the comparative analysis of the design and clinical outcomes between Optiflex Trio and published results of currently available trifocal IOLs.

Ribeiro et al compared the clinical outcomes of 3 different models of diffractive trifocal IOLs: FineVision POD F (PhysIOL), RayOne Trifocal (Rayner IOL, Ltd.), or AcrySof IQ PanOptix (Alcon Laboratories, Inc.) IOL in

Mean±SD			l Week	l Month	Months	6 Months	l Year	p-value*
ETDRS (LogMAR)	40 cm	Uncorrected Corrected p-value**	0.02±0.05 -0.00±0.03 0.08	0.01±0.05 -0.01±0.03 0.06	0.01±0.05 -0.01±0.03 0.06	0.01±0.05 -0.01±0.03 0.06	0.01±0.05 -0.01±0.03 0.06	0.97 0.88
	60cm	Uncorrected Corrected p-value**	0.08±0.07 0.07±0.06 0.54	0.08±0.07 0.07±0.06 0.54	0.07±0.06 0.06±0.05 0.49	0.07±0.06 0.06±0.05 0.49	0.07±0.06 0.06±0.05 0.49	0.93 0.85
	80 cm	Uncorrected Corrected p-value**	0.04±0.05 0.03±0.05 0.58	0.04±0.05 0.03±0.05 0.58	0.03±0.05 0.02±0.04 0.55	0.03±0.05 0.02±0.04 0.55	0.03±0.05 0.02±0.04 0.55	0.94 0.93
Reading speeds (wpm)	40cm	Uncorrected Corrected p-value**	162±40 163±40 0.86	163±40 165±40 0.85	164±40 166±40 0.85	64±40 66±40 0.85	165±40 167±40 0.86	0.99 0.99
	60cm	Uncorrected Corrected p-value**	67±3 69±3 0.76	168±33 170±33 0.76	169±33 171±33 0.77	169±33 171±33 0.77	170±33 172±33 0.76	0.98 0.98
	80cm	Uncorrected Corrected p-value**	70±3 72±3 0.78	171±33 173±34 0.78	172±34 174±34 0.82	172±34 174±34 0.82	173±34 175±34 0.85	0.99 0.98
CSV 1000 (LogMAR)		A(3.0) B(6.0) C(12.0) D(18.0)	1.55±0.13 1.90±0.08 1.54±0.08 1.05±0.06	1.56±0.13 1.91±0.08 1.54±0.08 1.06±0.06	1.57±0.13 1.92±0.08 1.55±0.09 1.07±0.05	1.57±0.13 1.93±0.07 1.56±0.08 1.07±0.05	1.58±0.14 1.94±0.07 1.57±0.06 1.08±0.05	0.94 0.15 0.43 0.09

Table 4 Binocular Near and Intermediate Visual Acuity with ETDRS Charts and Reading Speeds at 40, 60, and 80 Cm and CS with
CSV-1000 at Post-Operative Visits from 1-Week Through 12 Months

Notes: P value** calculated using Independent t-test. P value* calculated using Anova single factor test.

Abbreviations: SRD, Salzburg Reading Desk; wpm, words per minute; CSV, contrast sensitivity (Vector Vision); SD, standard deviation.

terms of visual acuity, refraction, CS, and visual quality.⁸ The study did not find any statistically significant differences between groups in distance, intermediate, and near visual acuity, and postoperative refraction. Postoperative binocular uncorrected intermediate VA of 0.10 logMAR or better was found in 14 (93.33%) patients in the 3 groups. In the present study, however, the postoperative binocular uncorrected intermediate VA of 0.10 logMAR or better

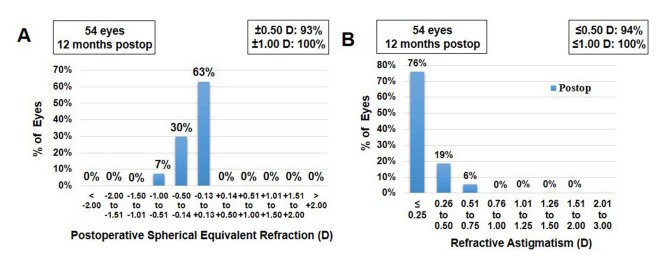


Figure 4 Histogram showing the accuracy to the intended spherical equivalent refraction (A) and refractive astigmatism (B) at 12 months post-op.

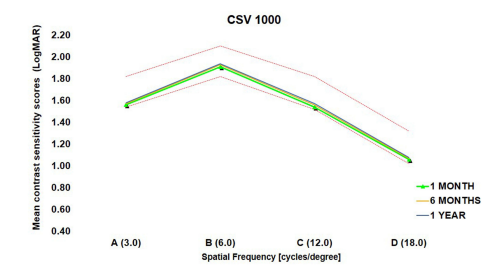
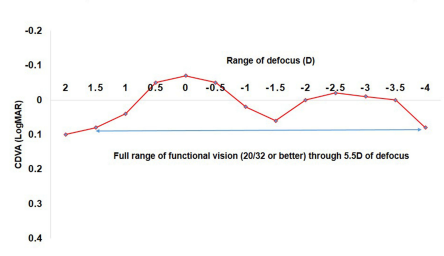


Figure 5 Photopic CS evaluated binocularly (with correction) at 12 months.

was found in 24 (88.8%) patients. The mean UIVA in our study was 0.07±0.06 logMAR, which was slightly lower than the 3 trifocal IOLs evaluated in the study by Ribeiro et al, wherein the RayOne trifocal showed maximum mean UIVA of 0.00 logMAR.

However, this may be explained by the fact that in our study intermediate visual acuity was evaluated at 60 cm versus 66 cm in their study. Despite this, none of the patients in our study complained of dissatisfaction with intermediate vision or required glasses for the same. However, when compared to studies by Kim⁹ and Asena⁷ et al, which evaluated outcomes with AT LISA 839 MP IOL, the intermediate VA results appear to be far superior with Optiflex Trio at a 12 months post-op.

In terms of uncorrected near vision results, Ribeiro⁸ et al found that a postoperative binocular uncorrected near VA of 0.10 logMAR or better was found in 13 (86.67%), 14 (93.33%), and 13 (86.67%) patients in the POD F, RayOne, and PanOptix IOLs groups, respectively. In contrast, a binocular uncorrected near VA of 0.10 logMAR or better was found in all (100%) patients in the present study. Since the near vision was evaluated at same distance in both our study and their study (40 cm), this suggests that Optiflex Trio trifocal IOL may provide uncorrected near vision as good as or better than the currently available trifocal IOLs. Although, the near addition for Optiflex Trio IOL (+3.5 D at 38 cm), which appears to be almost similar to that of the POD F and RayOne trifocal IOLs (Table 5),



Binocular corrected defocus curve at 12 months (+2 to -4D)

Figure 6 Binocular distance corrected defocus curve evaluated from +2 to -4 D defocus at 12 months.

Table 5 Comparis	Table 5 Comparison of the Design and Clinical Outcomes	I Clinical Outcomes Between Optifl	lex Trio and Published Results of $C_{\rm U}$	Between Optiflex Trio and Published Results of Currently Available Trifocal Intraocular Lenses	Lenses
	Optifiex Trio (Biotech Healthcare Holding GmbH, Luzren, Switzerland)	Zeiss AT LISA Tri 839MP (Carl Zeiss Meditec, Germany)	AcrySof IQ PanOptix (Alcon Laboratories, Switzerland)	RayOne Trifocal (Rayner Intraocular Lenses Ltd, England, UK)	Fine Vision POD F/ Micro F (PhysIOL, BVI, Belgium)
Optical Design	Diffractive- Refractive	Diffractive	Diffractive- Refractive	Diffractive- Refractive	Diffractive
Optic material	Hydrophobic acrylic containing natural chromophore	Hydrophilic acrylic with hydrophobic surface properties	Hydrophobic acrylic	Hydrophilic acrylic	Hydrophilic acrylic
Abbe number	49	58	37	56	58
Spherical aberrations	—0.20µт	—0.18 µm	—0.10 µm	Neutral SA	—0.11 µm
Number of diffractive rings	12 rings on anterior surface. Monofocal beyond 4mm	29 rings anteriorly on entire optic surface. Bifocal beyond 4.34 mm	15 rings on 4.5mm anterior optic surface. Monofocal beyond 4.5mm	16 rings on 4.5mm anterior optic surface. Monofocal beyond 4.5mm	26 rings on entire optic surface
Addition	+1.85D l(72cm)/ +3.5D N(38 cm)	+1.66D l(80cm)/ +3.33D N (40cm)	+2.17D (62cm)/+3.25D N(42cm)	+1.75D 1 (75cm)/+3.50D N (37.5cm)	+1.75D 1 (75cm) /+3.5D N (37.5 cm)
Light Utilization	88.3%	85.7%	88%	89%	86%
Light distribution	45% D/27% I/28% N	50% D/20%//30% N	42% D/ 24% I/ 22% N	52% D/22% I /26% N	49% D/ 17% I/ 34% N
			RESULTS		
Binocular UDVA (LogMAR)	Present study: -0.00±0.07	Ganesh et al' ¹⁸ : 0.01±0.05 Mencucci et al ¹⁷ : 0.00±0.02 Doroodgar et al ²² : 0.03±0.06	Ribeiro et al ⁸ . –0.02±0.09 Mencucci et al ¹⁷ . –0.02±0.08 Doroodgar et al ²² . 0.04±0.06	Ribeiro et al ⁸ : -0.02±0.08 Ferreira et al ²³ : -0.02±0.08	Ribeiro et al ⁸ : -0.01±0.06 Ferreira et al ²³ : -0.01±0.06 Marques et al ²⁴ : 0.02±0.02
Binocular CDVA (LogMAR)	Present study: -0.06±0.04	Ganesh et al ¹⁸ : -0.02±0.05 Mencucci et al ¹⁷ : -0.01±0.02 Doroodgar et al ²² : 0.02±0.05	Ribeiro et al ⁸ 0.01±0.07 Mencucci et al ¹⁷ : 0.03±0.08 Doroodgar et al ²² : 0.02±0.04	Ribeiro et al ⁸ : -0.04±0.10 Ferreira et al ²³ : -0.04±0.10	Ribeiro et al ⁸ : -0.02±0.06 Ferreira et al ²³ : -0.02±0.06 Marques et al ²⁴ : -0.02±0.04

Binocular UIVA	Present	Present study: 1)	Doroodgar et al ²² : 60cm 0.01±0.05	Ribeiro et al ⁸ : 66cm 0.06±0.06	Ribeiro et al ⁸ : 66cm 0.00±0.10	Ribeiro et al ⁸ . 66cm 0.04±0.07
(LOBITAR)	2) 80cm	60 cm: 0.07 ±0.06 2) 80cm: 0.03±0.05	Kim et al ⁹ : 80cm 0.22±0.10 Asena et al ⁷ : 60cm 0.25±0.12	Mencucci et al ¹⁷ 60cm: 0.07±0.04 Doroodgar et al ²² : 60cm 0.02±0.04	Ferreira et al ¹³ : 80cm 0.00±0.10	Ferreira et al ²³ : 80cm 0.04±0.07 Marques et al ²⁴ : 80cm 0.03±0.05
Binocular UNVA (LogMAR)	Present study: 40 cm 0.01±0.	Present study: 40 cm 0.01±0.05	Ganesh et al ¹⁸ . 40cm 0.02±0.04 Doroodgar et al ²² : 40cm 0.02±0.05 Kim et al ⁹ : 40cm 0.05±0.09	Ribeiro et al ⁸ . 40 cm 0.03±0.09 Doroodgar et al ²² : 40cm 0.02±0.04 Asena et al ⁷ : 40cm 0.03±0.08	Ribeiro et al ⁸ : 40 cm 0.01±0.12 Ferreira et al ²³ : 40cm 0.01±0.12	Ribeiro et al ⁸ : 40 cm 0.02 \pm 0.11 Ferreira et al ²³ : 40cm 0.02 \pm 0.11 Marques et al ²⁴ : 40cm 0.02 \pm 0.02
Reading speed (wpm, uncorrected)	40cm	40cm Present study: 165 ±40	-			,
	60cm	Present study: 170 ±33	Mencucci et al: ¹⁷ : 177.30±21.68	Mencucci et al: ¹⁷ : 176.28±15.04		
	80cm	80cm Present study: 173 ±34	Ganesh et al: ¹⁸ : 169.0±74.85		-	1
Abbreviations: D, distance; I, i	ance; l, inte	rmediate distan	ce; N, near distance; UDVA, uncorrected dist:	ance visual acuity; CDVA, corrected distance vi	Abbreviations: D, distance; I, intermediate distance; NN near distance; UDVA, uncorrected distance visual acuity; UNVA, uncorrected intermediate visual acuity; UNVA, uncorrected near visual	ual acuity; UNVA, uncorrected near visu

visual acuity; wpm, words per minute. its high percentage of overall light utilization (88.8%), combined with high light distribution for near (28%), may theoretically result in effective near visual acuity. However, future prospective, comparative studies between Optiflex Trio and different trifocal IOLs are needed to establish this preliminary observation.

Various studies have reported reduction in CS following implantation of multifocal IOLs.¹⁰⁻¹² However, modern multifocal IOLs based upon trifocal and EDOF technologies did not show a significant reduction in CS and visual quality.^{13–15} The CS with trifocal IOLs has been shown to provide conflicting results when compared to found to extended depth of focus lenses. Gunderson¹⁶ et al suggested comparable results between the two technologies, while Mencucci¹⁷ et al showed that Symfony ERV IOL resulted in significantly better CS than AT LISA Tri 839 MP trifocal IOL. However, while comparing 3 different trifocal IOLs, Ribiero⁸ et al did not find any statistically significant differences between groups for any of the spatial frequencies evaluated for photopic CS without glare $(P > 0.05)^{8}$. In the present study, Optiflex Trio achieved CS results within the physiologic CS range set for the measuring device for normal subjects of similar age (Figure 5).

Few studies have evaluated reading performance following implantation of trifocal IOLs. Compared to the study by Mencucci¹⁷ et al, the reading speed with Optiflex Trio was slightly less than that with AT LISA Tri and AcrySof IQ Panoptix trifocal IOL. This may be due to the difference on the tools used for evaluating reading performance in both studies, which was SRD in the present and MNREAD charts in Mencucci¹⁷ et al study. However, using the same tool, i.e., SRD, the mean reading speeds were found to be higher with Optiflex Trio versus Zeiss Trifocal IOL at the end of 12 months, Table 5. The relatively higher amount of light distribution dedicated for intermediate in Optiflex Trio (27% vs 20% in Zeiss Trifocal), and nil PCO incidence at 12 months may possibly explain relatively better reading speeds with Optiflex Trio trifocal IOL.

When compared to our published results of AT LISA TRI trifocal IOL, wherein, 5 eyes underwent YAG laser capsulotomy for early PCO at a mean follow-up of 7.2 ± 2.9 months, no eye developed PCO following Optiflex Trio IOL implantation in the present study.¹⁸ This may be explained by the fact that Optiflex Trio is a hydrophobic acrylic foldable IOL, whereas the material of AT LISA Tri is hydrophilic acrylic, and it is well-known that

hydrophilic IOLs have been shown to have significantly higher rates of PCO formation.^{19–21}

In conclusion, Optiflex Trio trifocal IOLs provided good visual outcomes at all distances at 12 months, which were comparable with the published results of currently available trifocal IOLs. Patients achieved high level of satisfaction in terms of spectacle independence as well as comfort with their vision due to minimal dysphotopsia symptoms and nil incidence of posterior capsular opacification at the end of 12 months. To our knowledge, this is the first study reporting the outcomes of Optiflex Trio trifocal IOL. Further prospective, randomised comparative studies with larger sample sizes are suggested to compare the long-term results of this lens with other available trifocal IOLs.

Disclosure

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