ORIGINAL RESEARCH

Safety and Efficacy of a New Posterior Chamber Phakic Intraocular Lens in Cases of High Myopia: Early Results

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submit your manuscript | www.dovepress.com DovePress **Purpose:** To demonstrate the visual outcomes of a foldable, hydroxy ethyl-methacrylate, single-piece, posterior chamber phakic intraocular lens (pIOL).

Study Type: Retrospective study.

Materials and Methods: Patients presenting with moderate to high myopia who underwent surgical correction with a posterior chamber phakic IOL (refractive intraocular lens, phakic intraocular lens, Appasamy Associates, Chennai, India) were retrospectively reviewed. Only patients with at least one-year follow-up were included. Manifest refraction, uncorrected distance visual acuity (UDVA), corrected distance visual acuity (CDVA), endothelial cell density (ECD), and pIOL vault were analyzed 1, 3, 6, and 12 months after surgery. Intraoperative and postoperative events were recorded in all cases.

Results: The study included 30 eyes from 15 patients. The mean patient age was 25.8 ± 3 years. The spherical equivalent of manifest refraction was -11.47 ± 4.38 D preoperatively and -0.44 ± 0.55 D postoperatively. The preoperative CDVA was 0.17 ± 0.12 logMAR. The postoperative UDVA was 0.053 ± 0.11 logMAR (min: -0.17 and max: 0.2) and 0.019 ± 0.091 logMAR (min: -0.17 and max: 0.2) at the end of 1 month and 6 months, respectively. At the end of the 12-month visit, the postoperative UDVA was 0.032 ± 0.094 , and the safety index was 2.42. The mean ECD was 2639 cells/mm² (min: 2389 and max: 2993 with SD: 139.53) at the preoperative visit and 2445 cells/mm² (min: 2050 and max: 2701) at the 12-month visit (5.8% loss, p less than 0.001). ECD loss from 6 months to 12 months was not statistically significant. No significant cataract formation, significant endothelial cell loss, glaucoma, uveitis, or any other vision-threatening complication were observed.

Conclusion: Based on postoperative experience, we found that RIL phakic IOLs are safe and effective for treating high myopia at short-term follow-up.

Keywords: refractive intraocular lens, RIL, myopia, phakic lens

Phakic Intraocular lens (IOL) have become the basic procedure for high degrees of refractive errors.^{1–3} Patients with myopia beyond -10 D prefer to undergo phakic IOLs instead of keratorefractive procedures globally.⁴ Refractive surgeons have determined that keratorefractive procedures for high myopes (>-8.0 D) and corneas with >40% tissue alteration weaken the cornea, significantly increasing the risk of regression and ectasia.⁵ In contrast, phakic IOLs that bypass the cornea provide excellent postoperative vision, maintain accommodation, and avoid the anterior pull of the peripheral vitreous, thus obviating the risk of peripheral retinal tears, and they are reversible.⁶ The most common types of lenses implanted during the last two decades, and considered safe and effective are the iris-claw and collamer posterior chamber

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Materials and Methods

This retrospective clinical study included the medical records of 30 eyes in 15 patients who underwent RIL at Ruby Eye Hospital, Berhampur, Odisha, India. All patients provided informed written consent in congruence with the tenets of the Declaration of Helsinki. The study was taken approval from the Ruby Eye Hospital ethics board, reference number REH-2017-3.

The exclusion criteria were age less than 18 years, endothelial density of less than 2300 cells/mm², endothelial guttae, macular scars, previous corneal surgery, an anterior chamber depth calculated from the endothelium of less than 2.8 mm, corneal astigmatism greater than 3

diopter (D), keratoconus and systemic endocrinal diseases (diabetes mellitus and thyroid disorders).

Description of the Refractive Intraocular Lens Phakic IOL (RIL) (Figure 1A)

The RIL Phakic Intraocular Lens used in this study had a biconcave monofocal design. It was made of 2-hydroxyethyl methacrylate, which is a monomer based on methacrylic acid. It was a single-piece hydrophilic pIOL with an optic diameter of 5.75 mm. It had a central vault of 1000 to 1600 microns, and the overall length ranged from 11.50 mm to 14.00 mm, with increments of 0.25 mm. The lens's thickness at the center varied from 150 to 450 microns, and the peripheral haptic thickness ranged from 100 to 140 microns. It had four holes in the optic region with a diameter of 0.55 mm for the smooth flow of aqueous fluid from posterior chamber to anterior chamber. It had two holes along the haptic plates for the determination of the vault, loading of the lens in the cartridge and implantation of the IOL in the posterior chamber. It had two curves in the haptic plates that had blended and polished edges.

Preoperative Examination

Before RIL implantation, patients underwent comprehensive ophthalmologic examinations including

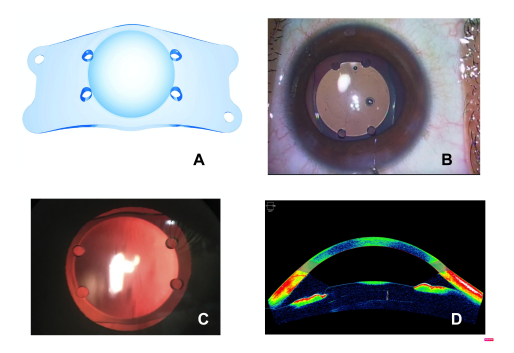


Figure I (A) Computer graphic of Refractive intraocular lens. (B) Intra-operative View of RIL under surgical microscope. (C) Retro-illumination view with slit lamp Biomicroscope. (D) Vault Height assessment with anterior segment OCT (Zeiss Inc., Jena, Germany).

refraction (objective, manifest, cycloplegic); uncorrected (UDVA) and corrected (CDVA) distance logMAR visual acuities; slit-lamp evaluation; Goldmann applanation tonometry; pupil size, anterior chamber depth and corneal topography (pentacam, Oculus); corneal endothelial cell count (Noncon Robo, Konan Medical), optical biometry (IOLMaster, Carl Zeiss Meditec AG), and measurements of the white-to-white (WTW) distance by digital calipers (Figure 2); binocularity evaluation by performing TNO stereopsis test; and fundoscopy (by indirect ophthalmoscopy).

RIL Power Calculation

The RIL with appropriate power and diameter was calculated by furnishing three parameters to the company. The white-to-white distance was measured with digital calipers and the anterior chamber depth recorded from corneal endothelium with Pentacam (OCULUS Optikgeräte GmbH, Wetzlar, Germany) and subjective refractive error.

YAG Peripheral Iridotomy

All patients underwent Nd.YAG peripheral iridotomy with Visulas YAG III lasers (Zeiss inc., Jena, Germany) to prevent postoperative pupillary block glaucoma.

Surgical Technique. (Video I)

Pupillary dilatation was performed by instilling 10% topical tropicamide-phenylephrine every 10 minutes for 1 hour prior to the procedure. One drop of ketorolac eye drops was administered 1 hour prior to implantation. Surgery was performed under topical anesthesia by administering proparacaine eye drops every 5 minutes half an hour before the procedure. Loading of the RIL into the butterfly cartridge was performed initially. Hydroxy-propyl methylcellulose viscoelastics were injected into the cannula and base of the butterfly cartridge. The RIL with the vault facing up and the hole of the leading haptic plate located on the top right side was placed over the base of the butterfly cartridge. With the help of plain-angled forceps, the haptic and optic plates were tapped for accommodation



Figure 2 Demonstrating Digital callipers to measure white to white diameter of cornea.

along the base, followed by the closure of the wings, ensuring that the lens was not engaged in the wings. Next, it was inserted into the plunger and was pushed forward to observe the smooth forward movement of the RIL. Eventually, a self-sealing 2.8 mm self-sealing clear corneal incision was made in the temporal quadrant of the eye. Two side ports were made at diagonally opposite quadrants to guide the haptic plate into the sulcus. Hydroxy-propyl methyl cellulose was injected into the anterior chamber to create a space between the cornea and capsule of the anterior lens. The bore of the butterfly cartridge was introduced into the 2.8 mm incision, and the plunger was pushed forward to observe the smooth unfolding of the RIL in the anterior chamber, ensuring that the vault was facing up by observing the leading haptic hole, which always remained on the top right-hand side. Once 50% of the RIL underwent the desired unfolding, the rest of the lens was pushed forward into the anterior chamber. With the help of smooth guidance, the haptic plates were initially pushed behind the iris, with the leading haptic plate followed by the trailing haptic plate. Finally, the complete removal of viscoelastics was performed using a Simcoe cannula/irrigation and aspiration cannula (Figure 1B). The viscoelastic behind the RIL was removed by gentle tapping of the centre of the optic. At the end, moxifloxacin intracameral preservative-free was introduced.

Postoperative Follow-Up

Follow-up visits were scheduled on the days 1, 7 and 30 after surgery, at 6 months and yearly thereafter. Evaluations included uncorrected distance visual acuity (UCVA) and corrected distance visual acuity (CDVA) in logMAR units, slit-lamp examinations of the position of RIL (Figure 1C), intraocular pressure and fundus examinations. Vault assessment was performed at each visit with the help of anterior segment OCT (Figure 1D).

Outcome Parameters

The primary outcome measure was uncorrected distance visual acuity, and the secondary outcome measures were vault height and endothelial cell count.

Statistical Analysis

Statistical analysis was performed using SPSS software. The normality of the data was checked by the Kolmogorov–Smirnov test and analyzed using paired t tests, Wilcoxon rank-sum tests, or analysis of variance Table I Demographics of the Study

	Demographics	Mean ± SD	Min/ Max
1	Age (Yrs)	25.8 ± 2.6	19/30
2	Spherical equivalent (D)	-11.4 ± 4.3	-25/-6
3	White to white diameter (mm)	11.8 ± 0.4	11/12.5
4	Axial length (mm)	26.4 ± 1.5	24.4/31.3
5	Anterior chamber depth from corneal	3.2 ± 0.2	2.8/3.5
	endothelium (mm)		
6	Flat Keratometry (D)	44.5 ± 1.4	42/48.3
7	Steep keratometry (D)	45.7 ± 1.9	43/50.3

with multiple comparisons, where appropriate, to determine significant differences in refractive, visual, and adverse outcomes. The results are presented as the mean and standard deviation. Differences were considered statistically significant when the p value was less than 0.05.

Results

A total of 30 eyes from 15 patients were included in the study. Ten patients were male, and 15 patients were female. All patients underwent bilateral RIL implantation. The patient demographics are described in Table 1.

Preoperative and Postoperative Visual Acuity (Figures 3D and 4B)

The preoperative mean uncorrected visual acuity (UCVA) recorded was 1.266 logMAR (max: 1 and min: 1.6, std dev: 0.17 and variance: 0.029), and the corrected distance visual acuity (CDVA) recorded was 0.173 logMAR (max: 0, min: 0.4, std dev: 0.128 and variance: 0.016). The postoperative mean UCVA at 1 month was 0.053 logMAR (SD: 0.11, max: -0.17 and min: 0.2, p value<0.0001), at 6 months was 0.019 logMAR (SD: 0.09, max: -0.17 and min: 0.2, p value <0.0001) and at 12 months was 0.032 logMAR (SD 0.09, max: -0.17 and min: 0.3, p-value: <0.0001) (Figure 3). Considerable improvement in post-operative uncorrected visual acuity (UCVA) was observed compared to preoperative corrected distance visual acuity (CDVA) and it was highly significant (p<0.05). (Figure 4A) Cohen's d =[mean postoperative UCVA (0.0320) - mean preoperative CDVA (0.17)] / 0.106066 = 1.301076, Glass's delta = (0.-(0.032-0.17) / 0.12 = 1.15, and Hedges' g = (0.032-0.17) / 0.12 = 1.150.106066 = 1.301076 were calculated at the end of 12 months of the study to determine the success of procedure, which was found to be successful.

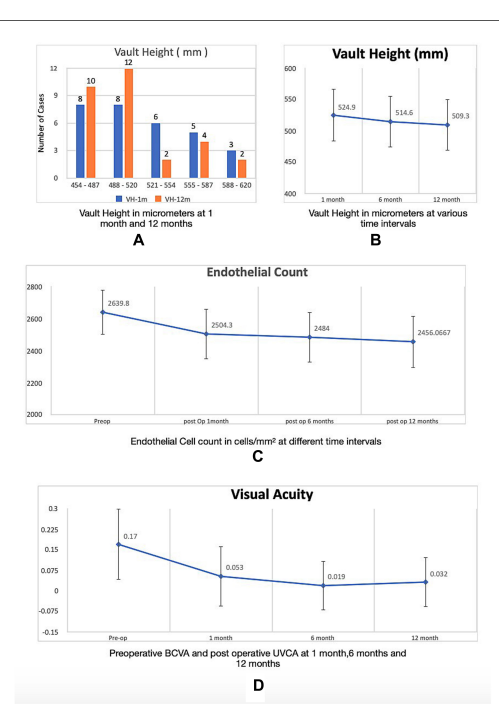


Figure 3 (A) Vault height (microns) at various time points. (B) Vault Height in micro-metres Comparison at 1 month and 12 months. (C) Endothelial Cell Count in cells/ mm² at different time zones. (D) Preoperative CDVA and Post-Operative UCVA at 1 month, 6 months and 12 months.

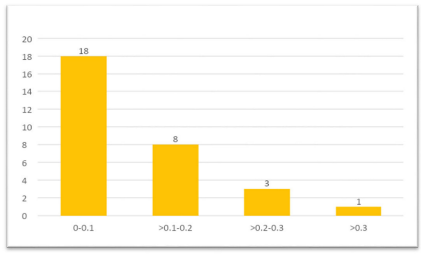
Vault Height (Figure 3A and B)

Vault height was measured by anterior segment OCT (Zeiss Inc., Jena, Germany). The mean vault height at 1 month postoperatively was 524.9 microns (min: 470 and max: 611, SD: 41.5). It was 514.6 microns (min: 454 and max: 602, SD: 41) and 509.36 microns (min: 455 and max: 598, SD: 40.68), with p values of 0.1623

and 0.074 at the end of 6 months and 12 months, respectively.

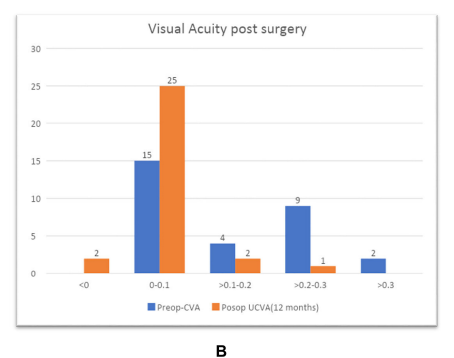
Specular Microscopy (Figure 3C)

The mean preoperative endothelial cell count (ECC) was 2639 cells/mm² (min: 2389 and max: 2993 with SD: 139.53), which decreased by 5.15% to 2504 cells/mm²



Α

Number of eyes showing difference between post-operative UCVA and preoperative BCVA in Logmar Units



Preoperative BCVA vs Post-operative UCVA in Logmar Units

Figure 4 (A) Difference between post-operative UCVA and preoperative BCVA in Logmar units showing significant improvement with p value <0.05. (B) Preoperative CDVA vs Post-Operative UCVA at 12 months in Logmar units.

(min: 2185 cells/mm² and max: 2786 cells/mm² p=0.00038) at 1 month POD. The mean endothelial cell count at 12 months showed a further reduction of 2.3% to 2445 cells/mm² (min: 2050 cells/mm² and max: 2701 cells/mm² p value

0.1215). There was a significant reduction in the endothelial cell count in comparison to the preoperative endothelial cell count at the end of 1 month; however, it remained stable with no significant reduction at the end of 12 months.

Safety Index

The safety index was calculated by dividing the mean postoperative best-corrected visual acuity by the mean preoperative best-corrected visual acuity. The safety index at the end of the final visit was 2.42.

Complications

None of the eyes developed cataracts, pupillary block glaucoma, iris pigment dispersion, retinal detachment or any other vision-threatening complication.

Discussion

In cases of myopia >8 D and those with insufficient corneal thickness, posterior chamber phakic intraocular lens implantation is capable of working successfully, providing accurate spectacle-free vision.¹⁸ This study focuses on the safety of a new intraocular phakic IOL. This is a retrospective observational case series of refractive intraocular lenses showing good results. Recent studies of a novel phakic IOL (IPCL with peripheral holes) by Care Group Inc. achieved a visual acuity of 20/40 or better in 90% of eyes.¹⁷ In our study, 100% of eyes achieved an unaided visual acuity of 20/40 or better. In addition, 10% of eyes had a vision of 20/16 or better.

Similar to previous studies, our study demonstrated improvement in the mean uncorrected visual acuity from 1.266 logmar units to 0.053 units and showed no significant change for 12 months.

Furthermore, the postoperative residual refraction recorded value was similar to that of previously published studies. Alfonso et al, in a study of 182 eyes, demonstrated that 96.8% of eyes were within \pm 1.0 D of attempted correction at the 1-month follow-up.¹¹ Sachdev et al, in a study of 134 eyes, demonstrated that 96.06% of eyes achieved a manifest refractive error within \pm 1.0 D of intended correction.¹⁶ Vasavada et al demonstrated residual refraction within \pm 1D in 92% and 100% of eyes after 1 month and 6 months, respectively.¹⁹ Along similar lines, our study demonstrated that 96.6% of eyes had residual refraction within \pm 1D.

Postoperative vault height, rotational stability and preservation of the endothelium determine the success of phakic IOLs. Vault heights within the range of 250 to 750 microns are considered ideal to prevent cataract formation.²⁰ In our study, the mean vault height was 524 microns, which showed a marginal reduction at the end of 12 months; however, it was

safe enough to prevent cataract formation. In our study, none of the cases had any form of postoperative intraocular phakic IOL rotation, maintaining visual stability. Furthermore, preservation of endothelial cells is crucial and critical because it maintains corneal health throughout the life of a patient.²¹ Endothelial cell loss needs to be minimal by the judicious use of ophthalmic viscosurgical devices during lens implantation. Vasavada et al showed an 8.1% reduction in the endothelial cell count in the immediate postoperative period and showed only a 4% further reduction at the end of 3 years.¹⁹ The highest evidence of endothelial cell loss postphakic IOL implantation is from the multicentric US FDA trial (Visian implantable Collamer lens, STAAR Surgical, Nidau, Switzerland), which showed a reduction of $3.3 \pm$ 7.6% at 1 year (90% confidence limits: 2.4% to 4.3%) and $9.7 \pm 9.3\%$ at 4 years.²² In our study, the endothelial cell loss was 5.1%, 5.8% and 7.3% at 1 month, 6 months and 12 months, respectively.

Cataract, glaucoma and infection are the three most important risks following phakic IOL implantation.

White to white (wtw) measurement is the most critical step in accurate phakic IOL calculation, improper measurement of the wtw diameter generates a faulty length of the phakic IOL, especially if the phakic IOL is smaller in length than the actual wtw diameter, it may lead to lower vault height and intraocular instability coaxing to cataract.²³ In our study, none of the eyes had cataracts, we assume that the accurate calculation of the white-to-white diameter (wtw) maintained safe vault height and peripheral holes allowed a smooth flow of aqueous. However, long-term results are awaited.

Acute rise in intraocular pressure in the immediate post-op period is commonly attributed to nonpatent peripheral iridotomy.²⁴ Hence, in our study, all patients were ensured of patent peripheral iris opening by performing retroilluminationtests with a visible red glow from peripheral opening in addition to pupillary glow prior to surgery.

Finally, post-operative intraocular infection is the most feared complication to any intraocular procedure, and prevention is the most crucial step in this regard.^{25–28} In our study, all patients were injected with preservative-free intracameral moxifloxacin to prevent any intraocular infection.²⁸

The main limitations of the study are the short followup time, the small sample size and its retrospective nature. However, these preliminary results found that Refractive Intraocular Lens (Appasamy Associates Inc.) are safe and effective for treating cases of high myopia at short time follow-up (1 year), and similar to other posterior chamber phakic IOLs that have shown to be safe over much longer periods of time. Additional medium and long-term studies are warranted.

Conclusion

With the emerging trends in the field of refractive surgery, the phakic intraocular lens plays a significant role in spectacle-free vision. The short-term visual outcomes of RILs are promising despite the lower number of lenses implanted. However, RILs represent an important component of novel phakic lenses.

Disclosure

The authors report no conflicts of interest for this work.

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