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Implantable collamer lens for the correction of post – keratoplasty myopia and astigmatism

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后房型人工晶状体植入用于角膜移植术后近视 和散光的矫正

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摘要

目的:评估角膜移植术 (KP) 后植入后房型人工晶状体 (ICL) 的安全性、屈光效果和视力。

方法:单中心回顾性研究。包括 32 例 35 眼接受后房型人 工晶状体植入以矫正角膜移植术后近视和散光。患者在 无法配戴眼镜、隐形眼镜或准分子激光术治疗不当的情况 下接受后房型人工晶状体植入术。观察患者裸眼远视力 (UDVA)、最佳矫正远视力 (CDVA), 屈光度及并发症。 对术前术后随访资料进行分析(16.7±13mo)(P<0.05)。

结果:术前,患者等效球镜和柱镜分别为-4.00~-20.00 D 和-2.00~-9.00 D。平均等效球镜由术前-11.41± 3.62 D显著下降至术后-1.95±1.78 D(P<0.0001)。平均 UDVA 由术前 20/400 显著增加至术后 20/25 (P < 0.0001)。与术前相比,术后 CDVA 平均提高1.5行,其中 37%提高2行或更多。1眼(2.8%)CDVA降低≥1行。 术中或术后无并发症。

结论:后房型人工晶状体植入对于无法配戴眼镜、隐形眼 镜或准分子激光术治疗不当的患者是一种安全有效的治 疗手段。

关键词:后房型人工晶状体植入;近视;散光;角膜移植 术后

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Abstract

• AIM: To evaluate the safety, refractive outcomes and vision after phakic posterior chamber implantable collamer lens (ICL) after keratoplasty (KP).

 METHODS: This retrospective single center study evaluated 32 (35 eyes) patients who received an ICL for myopia and/or astigmatism after keratoplasty. Patients underwent ICL surgery if they were unable to wear glasses or contact lenses and excimer laser surgery was contraindicated. Data were collected on uncorrected distance visual acuity (UDVA), corrected distance visual acuity (CDVA), refraction and complications. Data were analyzed for the preoperative and last postoperative visits (16.7±13mo) (*P*<0.05).

• RESULTS: Preoperatively, spherical equivalent (SE) ranged from -4.00 to -20.00 D and cylinder from -2.00 to -9.00 D. The mean SE decreased statistically significantly from - 11. 41 ± 3. 62 D preoperatively to - 1. 95 ± 1. 78 D postoperatively (P < 0. 0001). Mean UDVA increased statistically significantly from 20/400 preoperatively to 20/ 25 postoperatively (P < 0.0001). There was a mean improvement in postoperative CDVA of 1. 5 lines compared to preoperatively, 37% of eyes had an increase of 2 or more lines. One eye (2.8%) lost ≥ 1 line of CDVA. There were no intraoperative or postoperative complications.

• CONCLUSION: Posterior chamber phakic intraocular lens implantation is a safe and effective treatment for post-keratoplasty myopia and astigmatism in patients who are unable to wear spectacles or contact lenses and where corneal refractive surgery is contraindicated.

 KEYWORDS: implantable collamer lens; myopia; astigmatism; post-keratoplasty

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INTRODUCTION

P ostoperative refractive error and anisometropia can limit visual performance after keratoplasty^[1]. Moderate to high postoperative refractive error have been reported after keratoplasty^[2-3]. Postoperative refractive error may preclude adequate optical or excimer laser correction due to anisometropia, high astigmatism, dry eye or abnormal corneal topography^[3]. Hence, alternate secondary surgical procedures are often required for postoperative functional vision.

Various procedures have been used to treat post-keratoplasty refractive error including intraocular lens implantation (IOL), piggyback IOL implantation, limbal relaxing incisions and phakic IOL (pIOL) implantation^[4-6].

The implantable collamer lens (ICL) is a posterior chamber phakic IOL that is safe and efficacious for the treatment of moderate to high myopia and astigmatism^[7-8]. However, there is a relative paucity of publications on the treatment of postkeratoplasty refractive error with ICL surgery^[6,9-10]. Recent publications are limited to case reports, case seriesor outcomes of 15 eyes^[11]. In the current study, we present the safety, visual and refractive outcomes of a relatively large cohort of patients who underwent ICL surgery for correction of post-keratoplasty refractive error.

SUBJECTS AND METHODS

This study evaluated the eyes of patients with residual refractive error after keratoplasty that underwent implantation of a myopic or toric ICL at the King Khalid Eye Specialist Hospital (KKESH), Riyadh, Saudi Arabia. All surgeries were performed between January 2008 and June 2012. An institutional review board approved this study and this study adhered to the tenets of the Declaration of Helsinki.

Post-keratoplasty patients were included if they were unable to wear spectacles or contact lenses, the excimer laser surgery was contraindicated, and eyes had a stable refraction for at least 3mo after suture removal. Exclusion criteria were as follows, anterior chamber depth (ACD) less than 2.8 mm, cataract, a history of glaucoma or ocular inflammation, retinal detachment, neuro-ophthalmic disease, macular degeneration and retinal disease.

Before ICL implantation, all patients underwent a complete ophthalmologic examination, including measurement of uncorrected distance visual acuity (UDVA), corrected distance visual acuity (CDVA), manifest and cycloplegic refractions, keratometry, corneal tomography, pachymetry, slit lamp biomicroscopy, Goldmann applanation tonometry, specular microscopy and a dilated fundus examination. Postoperative examinations were performed at 1d, 2wk and 3, 6mo, 12mo and yearly thereafter.

Surgical Procedure All patients received 2 peripheral iridotomies with a neodymium: YAG laser 1wk prior to ICL surgery. To control for potential cyclotorsion in the supine position, the horizontal axis was marked with the patient sitting at the slit lamp just prior to ICL surgery. The patient was taken to the operating room and the eve was prepared in a sterile fashion and draped to isolate the lids and lashes. A topical anesthetic, dilating and cycloplegic drops were instilled in the operative eye. A lid speculum was inserted followed by instillation of an additional drop of anesthetic. A Mendez ring was used for intraoperative measurement of rotation from the horizontal axis. A viscoelastic device was placed into the anterior chamber and an ICL was inserted through a 3 - mm clear corneal incision using an injector cartridge. The ICL was rotated with a manipulator to ensure proper placement in the posterior chamber. Subsequently, the remaining viscoelastic material was completely washed out of the anterior chamber with balanced salt solution and a miotic agent was instilled. All surgeries were uneventful and there were no intraoperative complications. Postoperatively, the patients were prescribed topical steroid and an antibiotic (0.5% levofloxacin, CravitTM, Santen, Osaka, Japan) to be instilled 4 times daily for 2wk. The topical steroid was slowly tapered after 2wk.

Implantable collamer lens (**ICL**) **size and power calculation** All eyes were targeted for emmetropia. The ICL diameter was based on the horizontal white-to-white (WTW) diameter using a caliper and anterior chamber depth (ACD) measured with the Pentacam tomographer (Oculus Gmbh, Wetzlar, Germany). The modified vertex formula was used for calculating the ICL power with software provided by the manufacturer.

Outcome Measures and Statistics Data are presented for the preoperative and last postoperative visit. Data were collected on refraction, UDVA, CDVA, and complications. Data analysis was performed using SPSS for Windows version 20.0 (IBM Corp., New York, NY, USA). Visual acuity was converted to logMAR values for statistical analysis and the means and standard deviations were back – calculated to Snellen acuity. The efficacy index was calculated as:

Equation 1 mean postoperative UDVA/mean preoperative CDVA

The safety index was calculated as:

Equation 2 mean postoperative CDVA/mean preoperative CDVA

The paired t-test was used evaluating the difference between preoperative and postoperative visits. A P value less than 0.05 was considered statistically significant.

RESULTS

Patient Population The study cohort was comprised of 35 eves of 32 patients with a mean patient age of 31.12 ± 7.86 (range from: 22 to 49) y. Twenty-five (78%) of the 32 patients were male. Preoperatively, the mean anterior chamber depth was 3.83 ± 0.458 (range from: 3.02 to 5.10) mm. The mean preoperative corneal thickness was $516.63 \pm$ 60.44 (range from: 323 to 605) µm. The mean preoperative endothelial cell count was 1867.52±696 (range from: 507 to 2900) cells/mm². The mean time between keratoplasty and ICL implantation was 4.12±1.63y. The mean follow up was 16.7 \pm 4 (range from: 3 to 39) mo. The mean preoperative sphere was -8.60 ± 3.51 (range from: -2.00 to -18.75) D. The mean cylinder decreased statistically significantly from -4.84 ± 2.01 (range from: -2.00 to -9.00) D preoperatively to -3.63 ± 2.01 (range from: 0.00 to -8.00) D postoperatively (P=0.023).

The mean spherical equivalent decreased statistically significantly from -11.41 ± 3.62 (range from: -4.00 to -20.00) D preoperatively to -1.95 ± 1.78 (range from: 1.00 to -7.00) D postoperatively (P < 0.0001). At baseline, no eyes (0) were within ± 1.00 D MRSE compared with 28.57% at the last postoperative visit.



Figure 1 Preoperative (Preop) uncorrected distance visual acuity (UDVA) versus last postoperative (Postop) visit UDVA for 35 eyes that underwent implantable collamer lens surgery for myopia and/or astigmatism.



Figure 2 Preoperative (Preop) corrected distance visual acuity (CDVA) versus last postoperative visit uncorrected distance visual acuity (UDVA) for 35 eyes that underwent implantable collamer lens surgery forpost-keratoplasty myopia and/or astigmatism.

There was a statistically significant increase in mean UDVA from 20/400 preoperatively to 20/25 at the last postoperative visit (P < 0.0001) (Figure 1).

At the last postoperative visit, mean UDVA improved by 8.5 lines. The UDVA improved to 20/25 or better in 28.57% of patients and 57.14% of eyes were 20/40 or better. The UDVA was 20/100 or worse in 11.42%. No eyes lost UDVA.

Figure 2 presents the preoperative CDVA versus postoperative UDVA.

Preoperatively 11% of eyes had 20/20 or better acuity CDVA compared to 11. 42% of eyes with this level of UDVA postoperatively (Figure 2).

Postoperative UDVA was better than or equal to preoperative CDVA in 51. 42% of eyes. The proportion of eyes that presented with preoperative CDVA < 20/20 yet had postoperative UDVA $\geq 20/20$ was 5.71%. The efficacy index was 1.18(According to Equation 1).

The mean improvement in postoperative CDVA was 1.5 lines compared to preoperatively. Postoperative CDVA was higher than or equal to preoperative CDVA in 80% eyes. There was a statistically significant increase of 28.57% in the proportion of eyes with CDVA greater than or equal to 20/20 from preoperatively to postoperatively (P < 0.0001). No eyes had vision worse than 20/100 postoperatively (Figure 3).

At the last postoperative visit there was an increase in CDVA of 2 or more lines in 37.14% of eyes and one eye (2.8%) lost 3 lines of CDVA due to posterior subcapsular cataract (Figure 4). The safety index was 0.46 (According to Equation 2).



Figure 3 Preoperative (Preop) corrected distance visual acuity (CDVA) versus last postoperative (Postop) visit CDVA for 35 eyes that underwent implantable collamer lens surgery for myopia and/or astigmatism.



Figure 4 Change in the corrected distancevisual acuity for 35 eyes that underwent implantable collamer lens surgery for myopia and/or astigmatism.

DISCUSSION

The outcomes of this study of pIOL implantation for post – keratoplasty myopia or astigmatism indicated statistically significant increases in vision and reduced refractive error postoperatively. For example, there was a statistically significant increase in UDVA from 20/400 preoperatively to 20/25 postoperatively (P<0.0001). There was a statistically significant reduction in spherical equivalent of -9.46 D (P<0.0001).

To our knowledge, this is one of the largest series of patients in a publication of the ICL for post-keratoplasty refractive error. Our study evaluated patients who had exhausted all other optical or surgical options for treatment of the post keratoplasty refractive error. For example, anisometropia, dry eye, irregular topography or inadequate residual stromal bed precluded spectacle wear, contact lens wear and excimer laser refractive surgery. The outcomes of the current study indicate that ICL implantation in this subset of patients results in good refractive outcomes with 37.14% of eyes gaining 2 or more lines of CDVA and minimal loss (2.8% of eyes) of CDVA. The refractive outcomes of this study are consistent with previous studies of pIOL for post-keratoplasty refractive error. The statistical decrease in spherical equivalent of -9.46 D is slightly greater than a study with smaller sample size (15 eyes) that reported a decrease of $-8.85 \text{ D}^{[6]}$. A study of 7 eyes reported a change in spherical equivalent of -7.76 D after ICL implantation for post-keratoplasty anisometropia^[11]. The differences in enrollment criteria, range of post keratoplasty refractive error and sample size between studies may account for the difference in outcomes reported in our study and previous studies.

The visual outcomes in the current study are well within the range reported for previous studies of ICL implantation for post-keratoplasty ametropia. For example, 7 eyes had a mean gain in UDVA of 7.6±1.9 lines 12.8±8.8mo after ICL implantation for post – keratoplasty ametropia^[11]. Our outcomes are similar with a mean increase in UDVA of 8.5 lines. UDVA was 20/40 or better in 7 eyes (46.6%) in a study^[6] of 15 eyes after ICL implantation for post-keratoplasty ametropia which is somewhat lower that our outcome of 57.14%.

Loss of CDVA was minimal in our study with 1 eye (2.8%) losing 1 (or more) lines of CDVA at the last postoperative visit due to the development of posterior subcapsular cataract. This outcome concurs with a similar study that reported no loss of CDVA in 15 eyes^[6].

In summary, ICL implantation is efficacious for patients with post – keratoplasty myopia and astigmatism who are not candidates for optical treatments and in whom other surgical options are contraindicated. ICL implantation resulted in minimal loss of CDVA.

Myopia and astigmatismare common after keratoplasty and the patients may not be candidates for optical or excimer laser correction. Hence, alternate intraocular procedure may be required for correcting the post-keratoplasty refractive error.

Inappropriate candidates with post – keratoplasty ametropia, phakic intraocular lens implantation in a reasonable alternative to decrease refractive error and increase vision postoperatively.

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