Prospective comparative study of tolerance to refractive errors after implantation of extended depth of focus and monofocal intraocular lenses with identical aspheric platform in Korean population

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Abstract

Background: To evaluate the clinical outcomes of extended depth of focus (EDOF) and monofocal intraocular lenses (IOLs) that share identical aspheric platform and compare their visual acuity tolerance to postoperative refractive errors.

Methods: This non-randomized, prospective comparative study included 120 eyes undergoing cataract surgery with implantation of either Tecnis ZCB00 IOL (Abbott Medical Optics Inc., Santa Ana, CA) (monofocal group: 60 eyes of 30 patients) or Tecnis Symfony IOL (Abbott Medical Optics, Inc.) (EDOF group: 60 eyes of 30 patients). Monocular and binocular visual outcomes, changes in refraction, defocus curve, contrast sensitivity, and perception of photic phenomena (Halo & Glare Simulator; Eyeland Design Network, Vreden, Germany) were evaluated 3 months postoperatively. To compare the refractive tolerance, each group was divided into three subgroups according to the postoperative uncorrected distance visual acuity (UDVA) and postoperative spherical equivalent (SE).

Results: In the EDOF group, the mean 3-months postoperative monocular UDVA, intermediate (UIVA), and near (UNVA) visual acuities were 0.03 ± 0.07, 0.09 ± 0.15, and 0.24 ± 0.16 logMAR, respectively. A total of 100, 96.55, and 68.97% of eyes in the EDOF group achieved binocular UDVA, UIVA, and UNVA values of 0.20 logMAR or better, respectively. In respect to refractive tolerance, the EDOF group showed higher SE values and statistically significantly better mean UDVA than the monofocal group in all subgroups, with UDVA of −0.013 and 0.028 logMAR for EDOF and monofocal groups (p = 0.037), respectively, in the subgroup where SE was within ±0.50 D, UDVA of 0.004 and 0.048 logMAR for EDOF and monofocal groups (p = 0.046), respectively, in the subgroup where SE was within −1.00 D, and UDVA of 0.020 and 0.083 logMAR for EDOF and monofocal groups (p = 0.026), respectively, in the subgroup where SE was more than −1.00 D. The mean patient satisfaction scores for spectacle-free distance, intermediate, and near visual acuities were 86.0, 85.0, and 66.0, respectively.

Conclusions: The EDOF IOL provided excellent postoperative visual outcomes in far and intermediate distances, with high patient satisfaction rate. Regarding the postoperative refractive tolerance to SE, the Tecnis Symfony IOL showed better tolerance to residual postoperative refractive error than the monofocal IOL with the same material and optical platform.

Keywords: Extended depth of focus IOL, Symfony, ZXR00, Visual acuity tolerance to postoperative refractive errors
**Background**

Today, multifocal intraocular lenses (IOLs) seem to offer the most promising treatment option for presbyopic patients [1]. Despite numerous advantages of multifocal IOLs, factors such as residual refractive error can lead to high dissatisfaction rate [2]. Minimum postoperative refractive error is required to achieve optimal visual outcomes, with even minor levels of astigmatism significantly undermining the patients’ postoperative visual acuity [3]. Refractive error, therefore, needs to be corrected as much as possible in order to fully exploit the benefits of multifocal IOLs [4]. The estimated percentage of enhancement procedures performed to reduce residual astigmatism after implantation of multifocal lenses varies from 5.24 to 23.66% depending on the study. For example, Gundersen et al. [5] observed considerable retreatment rates (10.8%), most of which were due to decrease in visual acuity (VA) secondary to residual astigmatism.

An extended depth of focus (EDOF) IOL provides significantly increased range of vision with minimal optical side effects of multifocality [6, 7]. Currently, there are no studies comparing the refractive error tolerance of an EDOF IOL to that of a monofocal one. Therefore, the aim of this study was to assess the clinical performance of an EDOF IOL (Tecnis® Symfony ZXR00) and compare its visual acuity tolerance to postoperative refractive errors of a monofocal IOL (Tecnis® ZCB00).

**Patient and methods**

In this prospective comparative study, 60 patients who underwent cataract surgery with implantation of either EDOF IOLs or monofocal IOLs that share the same material and aspheric platform were included. The EDOF group included 30 patients with bilateral implantation of the Tecnis® Symfony IOL (Abbott Medical Optics, Inc.), while the monofocal group included 30 patients with bilateral implantation of the aspheric Tecnis® ZCB00 IOL (Abbott Medical Optics, Inc.). Every patient was informed about the inclusion in the study and provided a written consent. The study procedure complied with the tenets of the Declaration of Helsinki and was approved by the local institutional review board.

Inclusion criteria consisted of patients with cataract and preexisting corneal astigmatism of 1.25 diopters (D) or less. Patients with a history of ocular pathology, trauma, contact lens wear, pregnancy, systemic or local medication, and ocular surgeries other than laser refractive surgery for myopia were excluded.

**Examination protocol**

Preoperative ophthalmological examination included manifest refraction, monocular corrected distance visual acuity (CDVA), Goldmann applanation tonometry, slit-lamp examination, corneal topography (Galilei G6; Ziemer Ophthalmic Systems AG, Port, Switzerland), optical biometry (IOLMaster 500; Carl Zeiss Meditec, Jena, Germany), funduscopy, and pupil size measurement (KR-1 W, Wavefront Analyser; Topcon, Tokyo, Japan).

Postoperative follow-up examinations were performed 1 day after surgery as well as 1 week, 1 month, and 3 months postoperatively. The postoperative examination at 1 month and 3 months included measurements of monocular and binocular uncorrected distance visual acuity (UDVA), CDVA, uncorrected intermediate visual acuity (UIVA) at 66 cm, binocular near visual acuity (UNVA) at 40 cm, contrast sensitivity under photopic (85 cd/m²) and mesopic conditions (3 cd/m²) (CSV-1000, VectorVision, Greenville, OH), and halo and glare using a computer-based software (Halo & Glare Simulator; Carl Zeiss Meditec). The simulator utilizes a numerical scale to quantify the size and intensity of halos and glare, ranging from 0 (none), 25 (mild), 50 (moderate), 75 (severe) to 100 (very severe). The simulator also classifies the halos perceived by patients into three different types: T1 (diffuse halo ring), T2 (starburst type), and T3 (distinct halo ring). In addition, at 3 months postoperatively, the pupil size was measured again using the same measurement device as noted above.

All patients were asked about their usage of spectacles after surgery with the question: “How often do you need spectacles to see at far/intermediate/near distances?” The answer choices were classified as 0, 25, 50, 75, and 100% of time. Furthermore, patients were asked about their satisfaction with the postoperative results with: “How satisfied are you with your spectacle-free vision at far/intermediate/near distances?” The answer choices ranged from 0 (not at all satisfied) to 100 (very satisfied). In addition, they were asked following questions: “Would you choose the same lens again?” and “Would you recommend this lens to your relatives and friends?”

Uncorrected monocular and binocular defocus curves were recorded from +1.00 D to −4.00 D in 0.50-D steps. All data were computed into a Cartesian graphic display, with the x-axis indicating the level of defocus and the y-axis the visual acuity values.

For comparison of the visual acuity tolerance to postoperative refractive errors in EDOF and monofocal IOL groups, each IOL group was subdivided into three groups depending on the measured postoperative uncorrected distance visual acuity: less than 0.0 (1.0 in decimal), 0.1 (0.8 in decimal), and more than 0.2 logMAR (0.63 in decimal). Parallelly, another subdivision was made in which each IOL group was divided according to the achieved postoperative spherical equivalent (SE) values: within ±0.50 D, within −1.00 D, and larger than −1.00 D.

**Surgical procedure**

One experienced surgeon (C.Y.C.) performed all surgeries using standard phacoemulsification via sutureless
2.2-mm incision. All incisions were placed at the steepest corneal meridian and topical anesthesia as well as mydriatic drops were instilled prior to the surgical procedure. After performing capsulorhexis and phacoemulsification, the study IOL was placed into the capsular bag using the Unfolder Platinum 1 series screw-style inserter (Abbott Laboratories, Inc.) through the main incision. In both groups, emmetropia or minimal myopia was aimed as target refraction in IOL power calculation using the Haigis formula.

Statistical analysis
A statistical software SPSS (Version 24.0 for Windows; IBM, Armonk, NY) was used for statistical analysis. The Kolmogorov–Smirnov test was used to check the normality of the data distribution. When parametric analysis was possible, the Student’s t test for paired data was performed for parameter comparisons between preoperative and postoperative results. In cases when parametric analysis was not possible, the Mann–Whitney U test was applied to assess the significance of differences between the results. An independent two sample T-test was performed to assess the significance of differences between the monofocal and the EDOF IOL groups. When parametric analysis was not possible, the Mann-Whitney U test was applied to assess the significance of differences between the results.

Results
The analysis included 60 patients who completed the 3-month follow-up examinations. The results for the 3-month timeline were stratified based on the available patients in this as well as patients with EDOF IOLs (EDOF group) and patients with monofocal IOLs in both eyes (monofocal group). Table 1 gives an overview of the patients’ demographics. There were no significant differences in preoperative visual acuity values between the monofocal and the EDOF IOL groups.

Visual and refractive outcomes and spectacle independence
Table 2 demonstrates the preoperative and postoperative visual outcomes of all patients. Overall, statistically significant improvements in monocular and binocular CDVA, UDVA, UIVA and UNVA were found after surgery ($P < 0.001$). When a comparison was made between monocular and binocular visual acuities at 3-months postoperatively, binocular CDVA ($P = 0.020$) and UDVA ($P = 0.005$) showed statistically significant superiority to monocular CDVA and UDVA. However, there were no statistically significant differences between binocular and monocular UIVA ($P = 0.174$) and between binocular and monocular UNVA ($P = 0.066$). Changes in refractive sphere ($P < 0.001$) and cylinder ($P = 0.017$) values also reached statistical significance at 3 months postoperatively, while changes in spherical equivalent ($P = 0.253$) did not.

Table 2 summarizes the postoperative monocular visual and refractive data of the two groups. As shown, the EDOF group achieved significantly better CDVA than the monofocal group ($P = 0.008$). Monocular UDVA of 0.20 logMAR or better was found in 98 and 96% of eyes in the EDOF and the monofocal group, respectively. Mean postoperative spherical equivalent was $−0.81\ D (−1.75\ to +0.50\ D)$ and $−0.40\ D (−1.50\ to +0.25\ D)$ in the EDOF and the monofocal group, respectively ($P < 0.001$), and postoperative spherical equivalent was within $±1.00\ D$ in 82.5 and 90.8% of eyes in the EDOF and the monofocal groups, respectively. Mean postoperative cylinder was $−0.59\ D (−1.75\ to 0.00\ D)$ and $−0.58\ D (−1.75\ to 0.00\ D)$ in the EDOF and the monofocal group, respectively ($P = 0.896$).

Figure 1 demonstrates the distribution of 3-month postoperative monocular and binocular visual acuity values in the EDOF group. A total of 98.28, 91.38, and 50.00% of eyes achieved monocular UDVA, UIVA, and UNVA of 0.20 logMAR or better. Binocularly, UDVA, UIVA, and UNVA

### Table 1 Patients demographics

| Parameter         | EDOF group | Monofocal group | $P$ value |
|-------------------|------------|-----------------|-----------|
| Eyes, n           | 58         | 60              | 0.283     |
| Patients, n       | 29         | 30              |          |
| Age (years)       |            |                 | 0.192     |
| Mean ± SD         | 64.59 ± 10.00 | 66.35 ± 8.71    |          |
| Range             | 45 to 84   | 48 to 82        |          |
| Male gender, n (%)| 10 (34.5)  | 14 (46.7)       | 0.192     |

2.2-mm incision. All incisions were placed at the steepest corneal meridian and topical anesthesia as well as mydriatic drops were instilled prior to the surgical procedure. After performing capsulorhexis and phacoemulsification, the study IOL was placed into the capsular bag using the Unfolder Platinum 1 series screw-style inserter (Abbott Laboratories, Inc.) through the main incision. In both groups, emmetropia or minimal myopia was aimed as target refraction in IOL power calculation using the Haigis formula.

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**Results**
The analysis included 60 patients who completed the 3-month follow-up examinations. The results for the 3-month timeline were stratified based on the available patients in this as well as patients with EDOF IOLs (EDOF group) and patients with monofocal IOLs in both eyes (monofocal group). Table 1 gives an overview of the patients’ demographics. There were no significant differences in preoperative visual acuity values between the monofocal and the EDOF IOL groups.

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| Variable  | Preoperative | Monocular | Binocular | $P$ value\footnote{For monocular comparison} |
|-----------|--------------|-----------|-----------|------------------------------------------|
| UDVA (logMAR) | Mean ± SD 0.46 ± 0.43 0.03 ± 0.07 −0.01 ± 0.05 | 0.005 \footnote{For monocular comparison} |
| CDVA (logMAR) | Mean ± SD 0.21 ± 0.31 −0.02 ± 0.07 −0.06 ± 0.06 | 0.001 \footnote{For monocular comparison} |
| UIVA (logMAR) | Mean ± SD 0.60 ± 0.36 0.09 ± 0.15 0.04 ± 0.13 | 0.001 \footnote{For monocular comparison} |
| UNVA (logMAR) | Mean ± SD 0.70 ± 0.37 0.24 ± 0.16 0.17 ± 0.14 | 0.001 \footnote{For monocular comparison} |

$D$ diopeters, $EDOF$ Extended depth of focus, $SD$ standard deviation, $SE$ spherical equivalent

\footnote{For monocular comparison}

\footnote{$P$ value is statistically significant ($P < 0.05$)
achieved values of 0.20 logMAR or better in 100.00, 96.55,
and 68.97% of eyes, respectively. And more than 90% of all
patients achieved 0.40 logMAR and 0.5 logMAR in mon-
ocular and binocular UNVA, respectively. Binocular post-
operative CDVA values were 0.10 logMAR or better in all
cases.

The level of spectacle independence reported by pa-
tients is demonstrated in Table 4. It shows that 96, 92
and 75% of patients with the EDOF IOL never or only
occasionally requires spectacles for distance, interme-
diate, and near vision, respectively.

Defocus curve outcomes
Figure 2 shows the mean monocular and binocular de-
focus curve results of the two groups. Under monocular
and binocular conditions, no significant differences were
found between the two groups for the defocus levels of −
0.50 D (monocular, \( P = 0.298 \); binocular, \( P = 0.978 \)), 0.00D
(monocular, \( P = 0.530 \); binocular, \( P = 0.874 \)), and + 0.50 D
(monocular, \( P = 0.502 \); binocular, \( P = 0.578 \)). For the rest
of the defocus levels, monocular and binocular visual acu-
ity values were superior in the EDOF group (\( P \leq 0.05 \)).

Photic phenomena outcomes
In EDOF group, halos were reported by 11 patients (37%)
and glare by 4 patients (13%) (Table 5). 14 patients (47%)
indicated to experience no or mild levels of halos, glare,
starbursts, and other types of dysphotopsia. At the 3-
month postoperative assessment, 5 patients (16%) re-
ported to experience severe visual symptoms. In contrast,
in monofocal group, there were much less patients that experienced halos, glare, starbursts, and other types of photic phenomena ($P = 0.956$, $P = 0.557$, $P = 0.046$, and $P = 1.000$, respectively).

**Contrast sensitivity outcomes**

Figure 3 shows the binocular contrast sensitivity results of the EDOF group under photopic and mesopic conditions. At all spatial frequencies, binocular contrast sensitivity values were within the normal ranges reported by the CSV-1000 [8].

**Tolerance to postoperative residual refractive errors**

In subgroups consisting of patients with postoperative UDVA of 0.1 logMAR or postoperative UDVA of less than 0.0 logMAR, EDOF group had significantly myopic spherical equivalent than the monofocal group ($P = 0.049$ and $P = 0.039$, respectively) (Table 6).

When the subgroups were divided according to the postoperative refractive errors, all EDOF subgroups

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### Table 4 Postoperative spectacle independence data 3 months after surgery in EDOF group

| Spectacle independence | Percent (%) |
|------------------------|-------------|
| Distance (%)           |             |
| Never/occasionally     | 96          |
| 50% of time            | 4           |
| Frequently             | 0           |
| Intermediate (%)       |             |
| Never/occasionally     | 92          |
| 50% of time            | 8           |
| Frequently             | 0           |
| Near (%)               |             |
| Never/occasionally     | 75          |
| 50% of time            | 14          |
| Frequently             | 11          |

*EDOF* extended depth of focus

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### Table 5 Incidence and level of photic phenomena 3 months after surgery

| Photic phenomenon | EDOF group | Monofocal group | $P$ value |
|-------------------|------------|-----------------|-----------|
| Type 1 Halo       |            |                 | 0.956     |
| No                | 28 (94)    | 29 (97)         |           |
| Mild              | 1 (3)      | 1 (3)           |           |
| Moderate          | 1 (3)      | 0               |           |
| Severe            | 0          | 0               |           |
| Very severe       | 0          | 0               |           |
| Type 2 Halo       |            |                 | 0.046*    |
| No                | 24 (70)    | 28 (94)         |           |
| Mild              | 3 (10)     | 1 (3)           |           |
| Moderate          | 2 (7)      | 1 (3)           |           |
| Severe            | 3 (10)     | 0               |           |
| Very severe       | 1 (3)      | 0               |           |
| Type 3 Halo       |            |                 | 1.000     |
| No                | 30 (100)   | 30 (100)        |           |
| Mild              | 0          | 0               |           |
| Moderate          | 0          | 0               |           |
| Severe            | 0          | 0               |           |
| Very severe       | 0          | 0               |           |

*P value is statistically significant ($P < 0.05$)
had significantly better mean UDVA than the monofocal subgroups ($P = 0.037$, $P = 0.046$ and $P = 0.026$, respectively) (Table 7).

**Pupil size**
The mean preoperative pupil sizes were 3.92 ± 0.53 mm and 3.90 ± 0.49 mm in the EDOF and monofocal group, respectively. At 3 months postoperatively, the mean pupil sizes were 3.86 ± 0.45 mm and 3.76 ± 0.56 mm in the EDOF and monofocal group, respectively. In both pre- ($p = 0.779$) and postoperative ($p = 0.270$) measurements, there were no statistically significant differences in the mean pupil sizes between the two groups.

**Patient satisfaction**
The mean patient satisfaction scores for distance, intermediate, and near visual acuities were 86.0, 85.0, and 66.0, respectively. 27 patients (93.1%) stated that they would recommend the same treatment to their friends and family. Also, 26 patients (89.6%) said they would choose the same IOL again.

**Complications**
One eye (1.7%) in EDOF group and three eyes (5.0%) in monofocal group developed posterior capsule opacification which required neodymium: YAG capsulotomy. No complication led to IOL explantation.

**Discussion**
Multifocal IOLs can successfully restore both near and distance visual acuities and yield satisfactory outcomes in the majority of patients [9, 10]. Such design can offer both cataract and presbyopic treatment, but bears disadvantages that are ascribable to its inherent optical design such as perception of photic phenomena, reduced contrast sensitivity, and decreased visual function in dim light settings [11–16]. Furthermore, residual refractive error can considerably undermine the postoperative visual performance, causing high dissatisfaction rate [2].

An EDOF IOL constitutes the most recent form of multifocal technology and has been reported to provide significantly increased range of vision with minimal optical side effects [6, 7]. In previous studies, the EDOF IOLs were able to restore excellent far and intermediate visual acuity with functional near vision compared to other multifocal IOL designs [17–21]. Furthermore, the EDOF IOLs demonstrated superior range of vision and spectacle independence than the monofocal lenses that were targeted to achieve emmetropia [22]. However, in a recent study, Cochener et al. [23] reported that while both
trifocal and EDOF IOLs provided good visual acuity at all distances, near vision was statistically better in the trifocal lenses compared to the EDOF ones. The current study evaluated the clinical performance of an EDOF IOL and compared its visual acuity tolerance to postoperative refractive error to that of a monofocal IOL that uses the same aberration-correcting optical platform.

In our study, distance visual outcomes were excellent in all EDOF patients, with mean binocular UDVA and CDVA values of \(-0.01 \pm 0.05\) logMAR and \(-0.06 \pm 0.06\) logMAR, respectively. This confirmed the ability of the EDOF IOL to successfully restore distance visual function, as it has also been reported for other models of multifocal IOLs [16, 24–39]. The logMAR postoperative monocular and binocular UDVA of 0.00 or better was achieved by 70.69 and 93.10% of eyes in the EDOF group, respectively. These results are similar to [23, 40] or even better than the outcomes observed in previously published data on other types of multifocal lenses [29, 41, 42]. As expected, the visual outcomes for intermediate vision was also excellent, with mean monocular and binocular UIVA of 0.09 ± 0.15 and 0.04 ± 0.13, respectively. The logMAR postoperative monocular and binocular UIVA of 0.20 or better was achieved by 91.38 and 96.55% of eyes in the EDOF group, respectively, which again, were comparable or superior to those reported for other models of refractive and diffractive multifocal IOLs [16, 24–39]. Similar to previous studies [19, 23], the performance of the EDOF IOL for distance or intermediate vision was better than for near vision. The noted differences in CDVA, UDVA, UIVA, and UNVA among these studies may be ascribable to factors such as variances in visual acuity measurement methods, residual refractive errors, and study populations.

In our study, the excellent visual outcomes at far and intermediate distances were consistent with the high levels of spectacle independency. In the EDOF group, about 90% of the patients indicated to not at all need spectacles or only occasionally need them for performing tasks at far and intermediate distances. For near vision, 10.3% of patients reported to require spectacles frequently. According to the Concerto study, micro-monovision method (\(-0.50\) to \(-0.75\) residual myopia in the nondominant eye) improved UDVA, spectacle independency as well as satisfaction rate for near vision [43]. Such low levels of spectacle dependence are comparable to those reported for other multifocal lenses [33, 39]. Law et al. [33] found that a limited number of patients with trifocal IOLs experienced difficulties in performing near and intermediate visual tasks such as reading newspapers or working with a computer. In another study that assessed a different trifocal lens model, Kohnen et al. [31] noted that 100% of the patients were independent of spectacles for distance and intermediate vision, while 12% occasionally required correction for near-vision. After implantation of another trifocal IOL that combines bifocal diffractive profiles, Jonker et al. [30] found that 80% of patients were independent from spectacles.

Our dysphotopsia assessment showed that starburst was the most commonly perceived optical phenomenon (reported by 38% of patients), followed by halo (8% of patients) and glare (7% of patients). In multifocal IOLs, one image is in-focus, while the out-of-focus image is neuronally suppressed (simultaneous vision) yet still produces such unwanted dysphotopsia [44]. In contrast, the Tecnis Symfony IOL provided consistent and excellent visual acuity at all distances, with minimal levels of visual disturbances compared to other multifocal IOLs [43, 45]. Halos are not less expected with such lens design as it generates an elongated focal depth rather than one or more fixed foci. In other studies reporting bilateral implantation of multifocal IOLs, 25 to 60% of patients reported difficulties due to perception of photic phenomena postoperatively [31, 33, 34]. In this study, a computer-based halo and glare simulator was implemented for assessment of occurrence of photic phenomena in order to avoid suggestive triggering of patients’ answers. As explained earlier, the Tecnis Symfony IOL does not create an out-of-focus image that would generate halos, which may help explain the low incidence rate of photic phenomena in our study. In contrast, in the monofocal IOL group, less patients were disturbed by photic phenomena. Previous studies also established that patients who received multifocal IOLs experienced dysphotopsia more often than those with monofocal IOLs [14, 46].

In this study, the contrast sensitivity values measured with the CSV-1000 showed good levels under both photopic and mesopic conditions. Though the values decreased at high spatial frequencies, this was still within the normal range and consistent with previous reports. Ruiz-Mesa et al. [21] observed no significant differences

| Post-operative spherical equivalent | EDOF IOL, Decimal (logMAR) | Monofocal IOL, Decimal (logMAR) | \(P\) value |
|------------------------------------|---------------------------|-------------------------------|------------|
| Within ±0.50 D                     | 1.04 (−0.013)             | 0.94 (0.028)                  | 0.037*     |
| Within −1.00 D                     | 0.99 (0.004)              | 0.91 (0.048)                  | 0.046*     |
| More than −1.00 D                  | 0.96 (0.020)              | 0.84 (0.083)                  | 0.026*     |

EDOF extended depth of focus, IOL intraocular lens, logMAR logarithm of the minimum angle of resolution, SE spherical equivalent, UDVA uncorrected distance visual acuity, VA visual acuity

\(*\) \(P\) value is statistically significant (\(P < 0.05\) )
in levels of contrast sensitivity between a group of eyes implanted with Tecnis Symfony IOLs and another group of eyes with PanOptix IOLs. de Medeiros et al. [17] showed that under photopic conditions, patients who received Tecnis Symfony IOL on one eye and a bifocal lens on the other (Tecnis’ ZMB80) showed better results at low spatial frequencies. Nevertheless, it has been reported that in patients with multifocal lenses, the performance of contrast sensitivity decreases at high spatial frequencies [25, 47]. Further research is required to evaluate the contrast sensitivity of the patients with bilateral implantation of Tecnis Symfony IOLs.

According to our results, the residual refractive error after implantation of the studied EDOF IOL had a very limited impact on monocular UDVA compared to the monofocal IOL with the same platform. In all sub-division groups, the EDOF group always showed better UDVA than the monofocal IOL. It is also important to note that we found no statistically significant differences in the mean pupil sizes measured pre- and postoperatively between the EDOF and the monofocal groups. Cochener et al. [48] observed that the Tecnis Symfony lens demonstrates an excellent tolerance to unexpected refractive errors and that monocular and binocular UDVA did not greatly change, with a mere difference of 0.05 between postoperative SE of ±0.25 and ± 2.00 D. In another study, Carones et al. compared the impact of induced astigmatism on the visual acuity in four different types of multifocal lenses and observed the highest tolerance with the Symfony IOL, which retained good visual acuity (0.7, decimal scale) even with an induced astigmatism of ~1.50 D [49]. Multifocal IOLs require emmetropia as target refraction to achieve the best visual outcomes and small amounts of refractive error may considerably degrade their visual performance [3]. Thus, refractive error must be fully corrected to achieve the maximum visual potential of a multifocal IOL [4]. However, residual refractive errors can be related to a variety of factors and it is not possible to predict absolute postoperative refractive errors. Besides, residual refractive error is the main identifiable cause of blurred vision [50] and can lead to high levels of dissatisfaction [2]. Such dissatisfaction can lead to lens explantation, IOL exchange, or laser refractive surgery postoperatively [51, 52]. Regarding the postoperative SE, the Tecnis Symfony IOL showed a stable tolerance to unexpected postoperative refractive errors. Such characteristic adds an additional value to the optical property of this lens and renders it versatile for different clinical situations, which is a key factor for high satisfaction rate. Further study is yet required to assess the visual acuity tolerance to the postoperative SE of patients with bilateral implantation of Symfony IOLs.

The study has several notable limitations. First, the refractive tolerances were only tested with an EDOF and a monofocal IOL. Ideally, more than three groups (other type of diffractive-refractive, EDOF, or monofocal IOLs) would have to be included to minimize confounding factor. Additionally, the EDOF lens was targeted for emmetropia, which may result in better intermediate vision, not for near vision. Lastly, our study did not analyze the optical performance of different IOL designs on an optical bench to characterize their behavior. Future studies should address these limitations. As certain multifocal IOLs remain unknown with respect to visual acuity tolerance to postoperative refractive errors, future randomized comparative studies with a larger cohort of the Tecnis Symfony IOL and other types of multifocal IOLs should be conducted to confirm its superiority in terms of residual refractive error and visual acuity tolerance to the postoperative SE.

To summarize, the Tecnis Symfony EDOF IOL is a promising means for producing excellent visual rehabilitation in patients receiving presbyopic treatment. Although previous studies showed that near vision was better in trifocal IOLs compared to the EDOF IOLs, EDOF IOLs provided superior far and intermediate vision and high levels of postoperative visual acuity tolerance than other types of monofocal or multifocal IOLs that are currently available. To minimize the optical side effects of multifocality, EDOF IOLs represent new-generation multifocal IOLs that provide a better alternative to patients who desire a spectacle-free lifestyle postoperatively. Further studies with longer follow-up periods may help provide quantitative long-term information on the optical properties of EDOF IOLs in comparison to the currently gold standard monofocal IOL and ultimately assist surgeons in choosing the appropriate IOL design for individual patients.

Abbreviations
CDVA: Corrected distance visual acuity; EDOF: Extended depth of focus; IOLs: Intraocular lenses; SE: Spherical equivalent; UDVA: Uncorrected distance visual acuity; UIVA: Uncorrected intermediate visual acuity; UNVA: Uncorrected near visual acuity; VA: Visual acuity

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Authors’ contributions
Involved in conception, design, conduct, collection and management of the study (SHK, CYC); analysis and interpretation of data (HSS, SHK, GUA, CYC); writing the article (SHK, CYC); draft and critical revision of the manuscript (HSS, SHK, GUA, CYC); final approval (CYC). HSS contributed equally to the manuscript as the first author. CYC contributed to the manuscript as the corresponding author. All authors read and approved the final manuscript.

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Every patient was informed about the inclusion in the study and provided a written consent. The study procedure complied with the tenets of the Declaration of Helsinki and was approved by the Institutional Review Board of Kangbuk Samsung Hospital.

Consent for publication
Not Applicable.

Competing interests
The authors declare that they have no competing interests.

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