Efficacy of a Gas Permeable Contact Lens to Induce Peripheral Myopic Defocus

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ABSTRACT

Purpose. The purpose of this work was to evaluate the potential of a novel custom-designed rigid gas permeable (RGP) contact lens to modify the relative peripheral refractive error in a sample of myopic patients.

Methods. Fifty-two right eyes of 52 myopic patients (mean ±SD age, 21 ±2 years) with spherical refractive errors ranging from −0.75 to −8.00 diopters (D) and refractive astigmatism of 1.00 D or less were fitted with a novel experimental RGP (ExpRGP) lens designed to create myopic defocus in the peripheral retina. A standard RGP (StdRGP) lens was used as a control in the same eye. The relative peripheral refractive error was measured without the lens and with each of two lenses (StdRGP and ExpRGP) using an open-field autorefractometer from 30 degrees nasal to 30 degrees temporal, in 5-degree steps. The effectiveness of the lens design was evaluated as the amount of relative peripheral refractive error difference induced by the ExpRGP compared with no lens and with StdRGP conditions at 30 degrees in the nasal and temporal (averaged) peripheral visual fields.

Results. Experimental RGP lens induced a significant change in relative peripheral refractive error compared with the no-lens condition (baseline), beyond the 10 degrees of eccentricity to the nasal and temporal side of the visual field (p < 0.05). The maximum effect was achieved at 30 degrees. Wearing the ExpRGP lens, 60% of the eyes had peripheral myopia exceeding −1.00 D, whereas none of the eyes presented with this feature at baseline. There was no significant correlation (r = 0.04; p = 0.756) between the degree of myopia induced at 30 degrees of eccentricity of the visual field with the ExpRGP lens and the baseline refractive error.

Conclusions. Custom-designed RGP contact lenses can generate a significant degree of relative peripheral myopia in myopic patients regardless of their baseline spherical equivalent refractive error.

Key Words: myopia progression, gas permeable contact lens, peripheral refraction
might also provide some peripheral myopic defocus. Other investigators recently reported a similar observation.6

The purpose of the current study was to evaluate the effectiveness of an experimental RGP (ExpRGP) lens to create peripheral myopic defocus in a cohort of myopic patients compared with a standard RGP (StdRGP) design and determine if the axial refraction affects the outcomes.

METHODS

Subjects and Lenses

Fifty-two neophyte myopic subjects were enrolled in the study. Measurements were obtained from the right eye only, initially without a lens (baseline), and then with an StdRGP and an ExpRGP lens intended to change the pattern of the RPRE. The two lenses were worn in randomized order on two different days between 9:00 and 11:00 AM and at least 2 hours after awakening.

After receiving an explanation of the nature of the study, each patient signed a consent form before enrollment. The research followed the tenets of the Declaration of Helsinki, and the Ethics Committee for Clinical Research at Clínica Teknon, Barcelona, Spain, reviewed and approved the protocol. The inclusion criteria required that the subjects had up to −8.00 diopeters (D) of myopia, had astigmatism less than −1.00 diopter of cylinder (DC), did not have a current ocular disease or injury, were not taking any ocular or systemic medications, and had no contraindications to contact lens wear.

The RGP lenses were made of Boston XO2 (hexafocon B) material with nominal properties of oxygen permeability of 141 barrers, according to the polymer manufacturer (Polymer Technology, MA). The central thicknesses varied depending on the back vertex power of the lenses according to the vertex-corrected spherical equivalent of the noncycloplegic subjective refraction of each eye. The overall diameter was 10.60 mm for the ExpRGP lens and 10.50 mm for the StdRGP lens. The base curve radii varied from 7.50 to 8.20 mm across the subject group, and the same base curve radius was used for both the StdGP and ExpGP lens for each subject. An alignment fit was attempted in all eyes by selecting the base curve radius as a function of the flat keratometric reading and corneal eccentricity.

Experimental RGP lens was designed using Zemax-EE software v.6 (Radiant ZEMAX, WA). Parameters for theoretical eyes were obtained from the study of Atchison7 assuming a standard corneal eccentricity of 0.50. Experimental RGP lens has 2.2 mm aspheric front and back optic zones. This design affords +1.50 D add increase at 2 mm from center (4 mm chord diameter) corresponding to about 30 degrees of retinal eccentricity and achieving around +6.5 D at the edge of the optic zone (9.0 mm chord diameter) as described in Spanish Patent Application P-201030694.

Standard RGP lenses were commercial lenses PRE AS (Precilens, France) with a central spherical surface followed by a peripheral aspheric surface.

Standard RGP and ExpRGP lenses were fitted according to the topographical information (simK readings measured over the 3 mm of the central cornea and eccentricity over a chord diameter of 9.0 mm). Trial lenses were used to achieve optimal fitting in a prestudy visit. Lenses were ordered based on the vertexed spherical equivalent refraction. Overrefraction was done at the trial visit over the contact lenses and a new lens was ordered if discrepancies greater than ±0.25D were found. Standard RGP and ExpRGP had the same central distance power.

Peripheral Refraction

Measurements of the central and peripheral (off-axis) refraction were obtained with an open-field Auto-Refractometer/Keratometer WAM-5500 (Grand Seiko Co, Ltd, Hiroshima, Japan) up to 30 degrees in the nasal and temporal visual field along the horizontal meridian in 5-degree increments. After lens insertion, peripheral refraction was evaluated after cessation of excessive tearing and once the lens was centered in the interblink interval. This took between 5 and 15 minutes after insertion for all eyes evaluated. Each automated refraction measurement was obtained only 1 to 2 seconds after a blink to allow the lens to center and settle. If a blink occurred in the middle of a measurement, the measurement was repeated. To minimize lens decentration during peripheral fixation, head rotation instead of ocular rotation has been used. We used a previously reported method.8,9 A laser pointer positioned over the patient’s head was oriented toward the primary gaze position. The patient rotated his or her head, avoiding lateral displacement, until the laser pointer reached the given eccentric location while the eyes remained in the primary gaze position.

Statistical Analysis

SPSS software package version 19 (SPSS Inc, Chicago, IL) was used for statistical analysis. The Kolmogorov-Smirnov test was applied to assess the normality of data distribution. Considering that we are evaluating different conditions on the same subjects (repeated measures), we used repeated-measures analysis of variance and the Friedman nonparametric test to compare the outcomes between the three conditions (baseline, StdRGP, and ExpRGP values) for normally or non–normally distributed variables, respectively. The Bonferroni post hoc correction for multiple comparisons was applied.

The Spearman rho correlation was applied when normality could not be assumed, and the Pearson correlation was used when normal distribution of data was verified to evaluate the relationship between the RPRE change and baseline refractive error.

For statistical purposes, p value less than 0.05 was considered significant.

RESULTS

The mean (±SD) central baseline spherical equivalent was −3.22 (±1.66) D (range, −0.75 to −8.00 D) of sphere with less than 1.00 DC of refractive astigmatism in the spectacle plane. The mean (±SD) flat keratometric reading was 7.85 (±0.25) mm.

Statistical analysis showed an interaction between the type of correction and the eccentricity for M and J0 (p < 0.05) but not for J45. For the most peripheral eccentricities, the M and J0 components of refraction became significantly different between the
FIGURE 1.

(A) Spherical equivalent component (M) of the RPRE without lenses (baseline) and with StdRGP and ExpRGP lenses. (B, C) Astigmatic component (J₀ and J₄₅) of the RPRE without lenses (baseline) and with StdRGP and ExpRGP lenses. Lines represent the second-order polynomial fitting. The error bars represent the SEM.
FIGURE 2. The correlation between the baseline axial refraction and the amount of change in RPRE achieved with the ExpRGP lens compared with baseline for 30 degrees of eccentricity in the nasal (30 degrees nasal; \( y = -0.0715x - 2.585; r = 0.05; p < 0.05; \) black diamonds, black line) and temporal (30 degrees temporal; \( y = 0.1902x - 2.0931; r = 0.16; p < 0.001; \) gray circles, gray line) visual fields.

Fig. 2 shows the relationship between the spherical equivalent axial refraction without lens and the averaged RPRE at 30 degrees of eccentricity (average of 30 degrees nasal and 30 degrees temporal) for the StdRGP and ExpRGP lenses. Correlations were very weak \((r = 0.05, r = 0.19, \) and \( r = 0.04 \) for no lens, StdRGP, and ExpRGP, respectively) and nonsignificant. The same results were confirmed when the nasal or temporal RPRE was evaluated independently, instead of averaged.

Fig. 3 shows the relationship between the amount of RPRE induced by the ExpRGP lens (ExpRGP minus baseline) at 30 degrees of the temporal and nasal visual field as a function of the baseline spherical equivalent. There was negligible or no relationship between central refractive error and RPRE with the ExpRGP lens at 30 degrees nasal \((r = 0.05; p < 0.05)\) or 30 degrees temporal \((r = 0.16; p < 0.001)\) eccentricities of the visual field. Despite the statistical significance of the correlation, both showed poor correlation. This means that the RPRE induced was independent of the central refractive error. Overall, in 60% of the eyes, the ExpRGP lens provided a change in RPRE of at least \(-2.00\) D of myopia at 30 degrees with a mean \((\pm SD)\) change in this group of \(-2.83 \pm 0.56\) D (range, \(-2.01\) to \(-3.69\) D).

Fig. 4 shows the averaged nasal and temporal RPRE values for the no-lens condition (baseline) as well as with StdRGP and ExpRGP lenses. Without a lens, only 12% of the eyes presented some degree of relative peripheral refractive myopia. This increased to 29% with the StdRGP lens, with most of the eyes showing myopic RPRE below \(1.00\) D (21%) and a few cases (8%) between \(1.00\) and \(2.00\) D. With the ExpRGP lens, up to 89% of the eyes presented some degree of myopic RPRE. In 60% of the eyes, the ExpRGP lens provided at least \(1.00\) D of relative peripheral myopia at the 30-degree eccentricity. The total proportion of eyes with RPRE greater than \(-1.00\) was

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Correlation between the axial refraction at baseline and the amount of change in RPRE achieved with the ExpRGP lens compared with baseline for 30 degrees of eccentricity in the nasal (30 degrees nasal; \( y = -0.0715x - 2.585; r = 0.05; p < 0.05; \) black diamonds, black line) and temporal (30 degrees temporal; \( y = 0.1902x - 2.0931; r = 0.16; p < 0.001; \) gray circles, gray line) visual fields.

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significantly higher with the ExpRGP compared with StdRGP and no-lens conditions \((p < 0.05\) for both comparisons, Cochran \(Q\)). When the analysis was conducted for the 30-degree eccentricity in the nasal and temporal fields separately, it was evident that the proportion of eyes with relative peripheral myopia greater than \(-1.00\) was higher in the temporal visual field (79%) compared with the nasal visual field (43%).

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DISCUSSION

Rigid gas permeable materials are excellent platforms for contact lens design because of their optical properties and resistance to flexure forces while on the eye. They are considered safe because they are associated with low risks of infection and other adverse events.\(^1\) Despite this, they represent a small part of the contact lens market because of their initial discomfort and the need for longer periods of adaptation compared with soft contact lenses.\(^1\)

The current study showed for the first time that a custom RGP lens might effectively induce relative peripheral myopic defocus across a wide range of values of axial myopia. Contrary to the relationship between induced peripheral myopic defocus and central refractive error in orthokeratology treatments,\(^1\) the amount of myopia induced in the periphery of the visual field with the lens evaluated in the present study is not limited by, or related to, the axial refraction.

Queiros et al.\(^1\) reported that the relationship between the amount of myopia corrected in the corneal center and the amount of peripheral myopia induced after treatment follows an almost 1:1 pattern, such that for 1.00 D of myopia to be compensated for, there

![Figure 4](image-url)

**FIGURE 4.**

The proportion of eyes with a given RPRE (average between 30 degrees in the nasal and temporal visual fields) at baseline and with the StdRGP and ExpRGP lenses. Negative values indicate myopic RPRE and positive values indicate hyperopic RPRE.
is the potential to induce 1.00 D of RPRE at 30 degrees in the extremes of the horizontal visual field. Cho et al. 12 showed that for myopes with less than 2.00 D of myopia, the progression of the axial elongation was very similar to subjects wearing spectacles. Kakita et al. 13 found a similar trend in Japanese myopic children, but this result was not observed in two other recently published studies from Spain14,15 and Hong Kong.16

Soft contact lenses currently in use to address myopia regulation incorporate treatment zones of +2.00 D including multifocal center-distance, peripheral gradient,17 and dual-focus lens designs.18 The mechanism underlying myopia control through manipulation of the peripheral optics is not fully understood; hence, it is not possible to establish the minimal effective level of peripheral myopia to reduce myopia progression. However, our results showed that if at least 2.00 D of relative peripheral myopic defocus might effectively be used to manipulate the relative peripheral myopic defocus, such an effect could be achieved in 60% of eyes evaluated in the current study.

Although soft contact lenses might result in faster adaptation associated with better comfort and centration with the pupil,19 RGPs appear to offer the best platform to create a given optical design and maintain such properties while being worn on the eye. Such an effect cannot be provided in the same way using soft contact lens materials. In our previous study, Proclear center-distance multifocal contact lenses with +4.00 D of add did not show significantly different peripheral myopic defocus compared with the +3.00-D add lenses; similarly, +1.00 D of add did not show any measurable benefit compared with the naked eye.3 We reported a similar effect with the same lens in myopic subjects.20 Recently, Ticak and Walline21 did not find a significant peripheral myopic defocus effect with the Proclear multifocal lenses compared with the effect produced by orthokeratology. However, they used an add power of +2.00 D, which, in our studies, showed only a slight peripheral myopic defocus effect compared with the more effective +3.00 D add. This means that a given increase in the peripheral power addition on a soft contact lens material does not necessarily increase the effect in the same proportion when measured by peripheral refraction. This has been confirmed recently in a pilot study that compared the peripheral refraction between the current ExpRGP design and the same design manufactured in a hydrogel material.5 In contrast, it appears that the effect of the RGP lens might be stronger (average change in RPRE, −2.00 D) than the one intended to be reproduced (−1.50 D as expected from the lens design at 30 degrees). In the present work, it has been found that the change induced in the RPRE can be greater than 2.00 or even 3.00 D in a significant proportion of eyes with the ExpRGP lens. This would not be expected as the lens is designed to induce up to 1.5 D of myopic defocus. To find an explanation to the higher spherical equivalent obtained compared with the theoretical peripheral myopic defocus induced by the lens design, we have to consider the astigmatism induced by the oblique incidence of light. When computed as spherical equivalent, the astigmatism induced by the optical surfaces for oblique incidence of light into the eye contributes to an overall increase in the myopic defocus that is beyond the change expected attributed solely to the increase in spherical power by the design of the lens. Additionally, the use of an autorefractor, which averages refraction over a 2.3-mm-diameter measurement ring, is likely to give highly myopic readings as the power add continues increasing up to the edge of the optic zone as explained in METHODS.

Ticak and Walline21 suggested in their recent study that more accurate instruments are needed to measure the power at discrete points in lenses with strong changes in power across their surface while the lenses are on the eye to better understand the refractive
effect of orthokeratology treatment and multifocal contact lenses in the context of myopia progression. However, previous studies have shown similar effects of the Proclear center-distance lens using an aberrometer that measures the full pupillary area.22 Furthermore, because previous studies of orthokeratology and other lenses also have been conducted with the Grand Seiko or the Shin-Nippon instruments, the current results were comparable to those previously reported.

Our results show that the defocus effect might be different at different retinal areas. As an example, although more than 80% of the eyes will experience a change in RPRE of at least −1.00 D, this amount reduces to 45% when we consider the nasal field. For 29% of the subjects, the amount of the RPRE with the ExpRGP lens ranged from 0.25 to 1.00 D of myopia. The peripheral refraction and probably the retinal shape are not the same between the horizontal and vertical meridians,21,23 not even between the nasal and temporal sides.24 Our results show that with the present lenses, higher myopic defocus is usually achieved in the temporal visual field, compared with the nasal visual field. We cannot justify these outcomes by the decentration of the lens with gaze changes. However, slight temporal decentration of the RGP lenses is usually observed, even when the lens remains within the accepted levels of decentration (0.50 mm).25 In the present study, lens centration was improved using the large-diameter lens (10.60 mm for the ExpRGP lens and 10.50 mm for the StdRGP lens). This is critical when fitting RGP lenses to maximize the symmetry in the myopic defocus induced around the foveal area. To maximize centration during the peripheral measurements, head turn instead of eye turn was used to fixate peripheral targets.

Results presented in this study show that the asymmetry between refraction in the nasal and temporal retinal eccentricity was approximately constant without and with lenses. Any significant decentration of the lens during measurements would result in reduction or exaggeration of the asymmetry found. This asymmetry in baseline data has been previously documented.8,20,26,27 This might be related to the asymmetry in the nasal and temporal eye length and corresponding posterior retinal contour in myopia.24,25 It could be also related to the measuring technique as we centered our measurements on the pupillary center. Asymmetries between nasal and temporal corneal curvature could also account for some differences. Kwok et al.8 found that asymmetries between nasal and temporal peripheral refractive error would be minimized if measurements were referred to the optical axis of the eye instead of the foveal axis. Overall, it can be concluded that lens decentration does not contribute to a change in the asymmetry of the nasal and temporal refraction and the factors previously mentioned should be considered.

Considering that treatments attempting to slow myopia progression should be more effective in younger children, safety becomes a major concern. Different authors have reported that RGP lenses worn overnight during orthokeratology treatment are well tolerated by children,13,14,29 which is probably related to the fast adaptation to this modality.30 Children also have been shown to successfully adapt to RGP lenses on a daily wear basis in different clinical trials.19,31 This makes RGP lenses a viable option for controlling myopia progression even if they have to be worn on a daily wear basis. One relevant aspect is the higher dropout rate that might be present with RGP compared with soft contact lenses. In a randomized clinical trial addressing the potential efficacy of conventional RGP lenses for myopia progression, Walline et al.31 reported a 70% retention rate in the RGP arm compared with 93% in the soft contact lens control arm. Although the tolerance must be a limitation for patients who previously wore soft contact lenses, this might not be the case for most young children wearing lenses for the first time to control myopia. Finally, the data summarized in the Tear Film & Ocular Surface Society International Workshop on Contact Lens Discomfort showed a trend for RGP lenses with larger diameters being more comfortable.32,33 As in the present study, RGP lenses intended for myopia regulation should be manufactured in larger-than-average diameters to improve centration and provide a sufficiently large treatment zone over the larger pupil areas of children. Furthermore, to overcome the short-term discomfort issues with corneal RGP lenses, other strategies might be considered in the future, such as large-diameter scleral or hybrid lenses. These lenses can also provide more lens stability and centration against misalignments related to ocular rotation and blinking.

In summary, the present study showed that RGP lenses can be custom designed to provide myopic RPRE in most of the myopic eyes irrespective of the degree of axial myopia within the range from −1.00 to −8.00 D of spherical equivalent included in this study. Comfort-related aspects of this modality compared with soft contact lenses need to be considered. Finally, although the efficacy of this device in providing myopic RPRE has been shown, the efficacy to inhibit myopia progression still needs to be investigated with the appropriate clinical trial.

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AQ1 = “TFOS CLD workshop” was expanded as “Tear Film & Ocular Surface Society International Workshop on Contact Lens Discomfort”. Is this correct?

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