Long-term Clinical Outcomes for Overnight Corneal Reshaping in Children and Adults

Michael J. Lipson, O.D., F.A.A.O.

Purpose. To retrospectively evaluate outcomes of overnight corneal reshaping (OCR) in children aged 12 years or younger compared to children older than 12 years and adults at one practice to establish the efficacy and safety of OCR during a period of 51 months. Methods. Examination records of OCR patients were reviewed for pretreatment data, including manifest refraction, keratometric readings, topography, corneal staining, and age at beginning OCR. Posttreatment records were reviewed for manifest refraction, unaided visual acuity, keratometric readings, topography, corneal staining, adverse events, and duration of OCR lens wear. Results. Records of 296 OCR patients were evaluated. One hundred fifty-four (52.0%) patients were 12 years old or younger. Sixty-eight percent of all patients in the study were Asian, and almost 95% of the patients aged 12 years or younger were Asian. The patients aged 12 years or younger had a mean original spherical equivalent refractive error of −3.50 ± 1.50 diopters (D). The patients older than 12 years had a mean original spherical equivalent refractive error of −3.20 ± 1.50 D. Refractive changes were similar between the group aged 12 years or younger and the group older than 12 years (3.30 ± 1.40 D vs. 3.10 ± 1.40 D) (P = 0.14). The mean unaided, binocular logMAR visual acuity was 0.03 ± 0.06 (i.e., 20/20) for the group aged 12 years or younger and 0.02 ± 0.07 (i.e., 20/20) for the group older than 12 years. There were three adverse events during the study that did not result in a loss of best-corrected visual acuity. A total of 507 patient-years of wear was represented in the study. Conclusions. OCR resulted in comparable safety and efficacy in temporarily reducing myopia in children younger than 12 years as it is for children older than 12 years and adults. Key Words: Corneal refractive therapy—Myopia reduction in children—Overnight corneal reshaping—Refractive error—Retrospective study.

Overnight corneal reshaping (OCR) is an alternative method to correct myopia in adults and children. OCR involves specially designed, reverse-geometry, rigid gas-permeable contact lenses worn while sleeping to temporarily reduce myopia by reshaping the corneal epithelium. The lenses are made of highly permeable materials in specific designs that have been approved by the Food and Drug Administration for overnight wear. OCR has been studied to analyze cellular changes of the cornea, evaluated for changes in visual performance and optical quality, corneal curvature, refraction, and unaided visual acuity, and evaluated for its effect on vision-related quality of life. One crossover study with OCR and soft disposable lenses showed that 85% of patients with myopia between −1.00 and −3.00 diopters (D) preferred OCR. Studies have also shown that the corneal and refractive changes made during OCR are reversible. Furthermore, studies have looked specifically at the efficacy of OCR for children and have shown the process to be effective in temporarily changing the cornea to reduce myopia. OCR has also been compared to LASIK for effectiveness and patient satisfaction. These cited studies have shown OCR to be effective in temporarily reducing myopia with little or no mention of any adverse events. Studies have shown that wearing soft contact lenses while sleeping is associated with an increased risk of microbial keratitis. To date, there are no studies to evaluate such a risk with OCR lens wear. There are a number of case reports of serious complications, such as microbial keratitis in children and adult patients using OCR lenses.

The purpose of this study was to retrospectively examine the results of OCR by a single doctor for efficacy and safety. The findings are reported for adults and children during a 4-year period and compare pretreatment data with outcomes for myopia reduction, unaided visual acuity, corneal staining, and adverse events. Previous studies on OCR have been conducted with a small study population (i.e., 50 or fewer subjects) and for short periods (i.e., less than 1 year). This study has a significantly greater population monitored for a longer period. Retrospective analyses such as this have been recommended by concerned groups of eye care professionals and regulators. It is not known at this time whether the risk of adverse events with OCR is different with children versus adults. As the process of contact lens corneal reshaping becomes more widely used, information on efficacy and safety is critical for doctors, patients, and parents to assess individual patient characteristics that influence potential success and risks before commencing treatment.

MATERIALS AND METHODS

Records of 296 OCR patients were reviewed for data before and during treatment between May 2002 and August 2006. Included in this study were all patients of any age who were seen for fitting or care involving corneal reshaping. Approval from the University of Michigan’s Institutional Review Board was obtained to conduct this retrospective review of records. Most patients were first-time wearers fitted and followed up by the author. The population included patients who were previously fitted by other practitioners who were seen for refitting, modification, or continuation of care.
Baseline examination included a comprehensive pretreatment vision and eye health evaluation, including unaided visual acuity, acuity with current spectacles, keratometry, topography, manifest refraction, phoria and cover testing, tonometry, slitlamp examination, and fundus examination. If patients were good candidates for OCR and they chose to wear OCR lenses, a diagnostic evaluation with OCR lenses was performed. After a successful diagnostic evaluation, patients or their parents signed an OCR informed consent and fitting agreement, and lenses were ordered. OCR lenses were dispensed during a session of careful instruction on insertion, removal, and care of lenses.

Initial follow-up visits were routinely performed after the first night of wear, 1 week, 1 month, 2 months, 4 months, and 6 months. After the 6-month visit, follow-up care included an annual comprehensive examination and a checkup every 6 months or additional visits as necessary for symptomatic reports of discomfort, change in vision, or damaged or lost lenses. The during-treatment data used in the study included manifest refraction, best-corrected visual acuity, slitlamp observations with grading of findings, and unaided visual acuity. All visual acuity measurements were performed monocularly and binocularly with a high-contrast acuity chart with Snellen notation converted to logMAR for statistical purposes. Refractive data for refractive change was calculated from baseline to the most recent follow-up visit. Evaluations made for slitlamp findings of corneal staining were recorded for degree and location. Baseline (pretreatment) evaluation data were available for 282 patients. Grade changes were defined as a change from baseline grade to any other visit of "none to trace, 1+" or a change from "trace to 1+ or 2+." Data were also tracked on age at commencement of OCR, months of OCR lens wear, whether patients were still currently using OCR lenses, and reporting of adverse events or complications during OCR lens wear. Corneal topography was obtained at each follow-up visit to ensure proper fitting and corneal integrity but was not used for data analysis.

Overnight corneal reshaping lenses used in the study were Paragon CRT (Paragon Vision Sciences, Mesa, AZ) or custom-designed lenses made from Boston XO material (Polymer Technology/Bausch & Lomb, Rochester, NY) manufactured by Art Optical Contact Lens, Inc. (Grand Rapids, MI). Recommended care solutions were Unique pH (Alcon, Fort Worth, TX) or Boston Simplus (Bausch & Lomb, Rochester, NY).

Definition of an Adverse Event

In this study, an adverse event included any of the following: microbial keratitis or a corneal ulcer, a corneal abrasion requiring medical treatment, loss of best-corrected visual acuity, or a corneal scar.

Data were analyzed by using SAS 9.1 statistical software (SAS Institute, Cary, NC). Comparisons between age groups made use of independent, two-tailed Student t tests. Comparisons of change from baseline findings to post-OCR findings were performed with paired Student t tests. Comparisons of duration of OCR lens wear were performed with Mann-Whitney U tests. Comparisons of proportions (e.g., ethnicity percentage) made use of the chi-square test.

RESULTS

Data from the records of 296 OCR patients were analyzed in the study. The mean age of all patients at the commencement of OCR was 17.7 ± 13.2 years. One hundred fifty-four (52.0%) patients were 12 years and 0 months old or younger at the commencement of OCR, and 142 (47.9%) patients were 12 years and 1 month old or older (Fig. 1). Of the 296 total patients, 146 (49.3%) were male and 150 (50.3%) were female. The mean duration of wear was 21.6 ± 18.1 months. For patients aged 12 years or younger, the mean duration of wear was 20.8 ± 14.7 months, whereas the mean duration of wear for those patients older than 12 years was 22.5 ± 21.2 months. The difference in duration of wear between the two groups was not significant (P = 0.6462). This represents a total of 506.95 patient-years of wear during the 51 months (i.e., 4.25 years) of the study period.

Ethnicity

Two hundred one (68%) patients were Asian; 90 (30%) were white; three (1%) were black; and two (<1%) were Eastern Indian. Of the 154 patients aged 12 years or younger, 146 (94.8%) were Asian and eight (5.2%) were of other ethnicity. Of the 142 patients older than 12 years, 55 (38.7%) were Asian and 87 (61.3%) were of other ethnicity. The difference in ethnic distribution between the two groups was statistically significant (P < 0.0001).

Termination of OCR Lens Wear

During the study, 32 (10.8%) patients discontinued OCR lens wear (Table 1). Of those 32, only six were aged 12 years or younger (6 [3.9%] of the 154), whereas 26 were older than 12 years (26 [18.3%] of the 142). It is statistically significant that 96.1% of those aged 12 years or younger are still using OCR lenses, whereas 81.7% of those older than 12 years are still wearing OCR lenses (P < 0.0001).

<table>
<thead>
<tr>
<th>Reason</th>
<th>No.</th>
<th>Aged 12 years or younger</th>
<th>Older than 12 years</th>
</tr>
</thead>
<tbody>
<tr>
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<td>1</td>
<td>3</td>
</tr>
<tr>
<td>Comfort</td>
<td>7</td>
<td>1</td>
<td>6</td>
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<tr>
<td>High refractive error (&gt; −4.75 D)</td>
<td>6</td>
<td>3</td>
<td>3</td>
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<tr>
<td>Solution reaction</td>
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<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Handling or inconvenience</td>
<td>3</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>Dry eye</td>
<td>1</td>
<td>0</td>
<td>1</td>
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<tr>
<td>Underwent LASIK</td>
<td>2</td>
<td>0</td>
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</tr>
<tr>
<td>Preferred soft contact lenses</td>
<td>5</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>Moved</td>
<td>3</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Total</td>
<td>32</td>
<td>6</td>
<td>26</td>
</tr>
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</table>
Refractive Error
The mean original spherical equivalent refractive error for all
patients was $-3.40 \pm 1.5 \text{ D}$, and the mean original
cylinder was $-0.42 \pm 0.46 \text{ D}$. In the group aged 12 years or younger,
the mean original spherical equivalent refractive error was $-3.50 \pm
1.5 \text{ D}$, and the mean original cylinder was $-0.33 \pm 0.36 \text{ D}$. In
the group older than 12 years, the mean original spherical equivalent
refractive error was $-3.20 \pm 1.5 \text{ D}$, and the mean original cylinder
was $-0.49 \pm 0.55 \text{ D}$ (Fig. 2). At the latest examination, the mean
spherical equivalent for all eyes had reduced to $-0.20 \pm 0.40 \text{ D},$
resulting from a mean change in the spherical equivalent of $3.20 \pm
1.4 \text{ D}$. Because the target result was a spherical equivalent of $0.00 \text{ D}$,
the mean change in spherical equivalent was greater for the group
with the higher mean original spherical equivalent. The mean spher-
ical equivalent change for the group aged 12 years or younger was
$3.30 \pm 1.4 \text{ D} \text{ versus } 3.10 \pm 1.4 \text{ D} \text{ for the group older than 12 years}
(P=0.1385).

The mean original cylinder amount was $-0.40 \pm 0.47 \text{ D}$, which
was reduced at the latest examination by $0.07 \pm 0.41 \text{ D} \text{ to } -0.33
\pm 0.41 \text{ D} (P=0.0039)$. For the group aged 12 years or younger, the
mean original cylinder was $-0.32 \pm 0.34 \text{ D}$, which decreased at
the latest examination by $0.01 \pm 0.35 \text{ D} \text{ to } -0.31 \pm 0.35 \text{ D}$.
For the group older than 12 years, the mean original cylinder was
$-0.48 \pm 0.57 \text{ D}$, which decreased at the latest examination by
$0.13 \pm 0.46 \text{ D} \text{ to } -0.35 \pm 0.33 \text{ D}$. The difference between the
amount of cylinder change in the two age groups was significant
($P=0.02$).

Keratometric Readings
The mean flat keratometric reading for all eyes was $43.37 \pm 1.4$
D. The mean flat keratometric reading was $43.12 \pm 1.5 \text{ D}$ for
the group aged 12 years or younger and $43.59 \pm 1.4 \text{ D}$ for the
group older than 12 years. Comparison of the flat keratometric reading
at baseline and unaided visual acuity or refractive change at the latest
examination showed no significant relationship.

Visual Acuity
At the latest examination, the mean unaided distance visual
acuity for all patients was $0.08 \pm 0.10 \text{ (i.e., } 20/25^{-1}) \text{ for the right}
eye, $0.08 \pm 0.10 \text{ for the left eye, } 0.03 \pm 0.07 \text{ (i.e., } 20/20^{-1}) \text{ for both}
eyes, and $0.11 \pm 0.12 \text{ (i.e., } 20/25 \text{) for the worse eye.}$
The mean unaided distance visual acuity for the group aged 12 years or
younger was $0.08 \pm 0.08 \text{ for the right eye, } 0.08 \pm 0.08 \text{ for the left}
eye, $0.03 \pm 0.06 \text{ for both eyes, and } 0.10 \pm 0.08 \text{ for the worse eye.}$
For the group older than 12 years, the mean unaided distance
visual acuity was $0.07 \pm 0.11 \text{ (i.e., } 20/20^{-3}) \text{ for the right eye,
0.09 \pm 0.13 \text{ (i.e., } 20/25^{-1}) \text{ for the left eye, } 0.02 \pm 0.07 \text{ (i.e., }
20/20^{-1}) \text{ for both eyes, and } 0.11 \text{ (i.e., } 20/25^{-1}) \text{ for the worse eye.}$
For the group aged 12 years or younger, 93.6% had a monocular
unaided visual acuity of 0.19 or less (i.e., better than 20/30) in each
eye (Table 2). For the group older than 12 years, 89.8% had a
monocular visual acuity of 0.19 or less. For the group aged 12
years or younger, 98.7% had a visual acuity of 0.19 or better, and
97.2% of the group older than 12 years had a visual acuity of 0.19
or better.

Slitlamp Findings
Corneal staining was evident in 25 (8.8%) of 282 of the patients
at the baseline examination. All were graded trace; 14 cases were
peripheral (i.e., exposure or incomplete blinking); and 11 were the
result of misdirected eyelashes (i.e., trichiasis). Corneal staining
was observed in 90 (32%) of the 282 patients at the 1-day
examination, in 45 (16%) of the patients at the 1-week visit, in 15
(5.3%) of the patients at the 1-month visit, and in 75 (25.3%) of
296 patients at any other visit during the study (Table 3).

Multivariable analysis showed some significant findings. When
comparing baseline findings to all follow-up visits, 58.2% of
patients had no change in or less staining and 41.8% had increased
staining at one or more visits. Patients who showed an increase in
staining versus baseline had a higher original spherical equivalent
($-3.70 \text{ D vs. } -3.11 \text{ D} (P=0.0013)$) than those who showed no
change in staining. In comparing increased grade of staining between
lenses of different designs, a higher percentage of patients
showed an increased grade of staining with the custom-designed
lenses than the CRT lenses (52% vs. 36%) ($P=0.0104$).

Trends observed in this same analysis were that younger
children showed a trend toward more staining increase than older
children and that there was no central staining observed at base-
line. Central staining was most often observed after the first night
of wear (26.6%) and least often at the 1-month visit (2.1%).

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**TABLE 2. Unaided Acuity Distribution**

<table>
<thead>
<tr>
<th></th>
<th>Aged 12 years or younger</th>
<th>Other than 12 years</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Right eye, no. (%)</td>
<td>Left eye, no. (%)</td>
</tr>
<tr>
<td></td>
<td>Both eyes, no. (%)</td>
<td>Both eyes, no. (%)</td>
</tr>
<tr>
<td>Unaided logMAR</td>
<td></td>
<td></td>
</tr>
<tr>
<td>visual acuity</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$-0.01 \text{ to } -0.10$</td>
<td>3 (2.0)</td>
<td>1 (0.7)</td>
</tr>
<tr>
<td>$0.0 \text{ to } 0.09$</td>
<td>87 (56.5)</td>
<td>83 (54.3)</td>
</tr>
<tr>
<td>$0.1 \text{ to } 0.19$</td>
<td>54 (35.1)</td>
<td>59 (38.6)</td>
</tr>
<tr>
<td>$0.2 \text{ to } 0.29$</td>
<td>9 (5.8)</td>
<td>6 (3.9)</td>
</tr>
<tr>
<td>$0.3 \text{ to } 0.39$</td>
<td>1 (0.7)</td>
<td>4 (2.6)</td>
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**TABLE 3. Corneal Staining**

<table>
<thead>
<tr>
<th></th>
<th>Baseline (n = 282)</th>
<th>1-day visit (n = 282)</th>
<th>1-week visit (n = 282)</th>
<th>1-month visit (n = 282)</th>
<th>Other visit (n = 296)</th>
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<tbody>
<tr>
<td>Central trace</td>
<td>0</td>
<td>57</td>
<td>26</td>
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<td>37</td>
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<tr>
<td>Central 1+</td>
<td>0</td>
<td>15</td>
<td>2</td>
<td>8</td>
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<tr>
<td>Central 2+</td>
<td>0</td>
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<td>0</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>Peripheral trace</td>
<td>14</td>
<td>6</td>
<td>11</td>
<td>5</td>
<td>13</td>
</tr>
<tr>
<td>Peripheral 1+</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>1</td>
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<tr>
<td>Peripheral 2+</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Trichiasis trace</td>
<td>11</td>
<td>6</td>
<td>5</td>
<td>3</td>
<td>8</td>
</tr>
<tr>
<td>Trichiasis 1+</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>2</td>
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<tr>
<td>Trichiasis 2+</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
</tbody>
</table>
Safety and Adverse Events

During the study, three patients had an incident of an adverse event, as defined earlier. Each of the adverse events was central, in children aged 12 years or younger, and after a minimum of 6 months of OCR lens wear. None of the events resulted in a loss of best-corrected visual acuity and each of the patients has continued to wear OCR lenses. One patient had total resolution, and the other two have central scarring remaining.

Lens Type

Of the two different lens designs, Paragon CRT and custom-designed reverse-geometry lenses in Boston XO material, 192 (65%) patients wore the CRT lenses and 104 (35%) wore the custom lenses. By age group, 111 (72%) of the group aged 12 years or younger wore the CRT lenses, whereas 43 (28%) wore the custom lenses. In the group older than 12 years, 81 (57%) wore the CRT lenses and 61 (43%) wore the custom lenses. These are significant differences between the two age groups (P=0.0068). Patients wearing the custom lenses showed a higher original spherical equivalent (−4.00 ± 1.7 D vs. −3.05 ± 1.3 D for the CRT wearers [P<0.0001]). The custom lens–wearing patients also had a statistically significantly higher original cylinder (0.56 ± 0.59 D vs. 0.33 ± 0.35 D for the CRT wearers [P<0.0001]). Because of the targeted refractive change in OCR, analysis also showed a significantly higher amount of spherical equivalent change in the custom lens wearers than in the CRT wearers (3.70 ± 1.6 D vs. 2.90 ± 1.3 D). However, there was also a greater cylinder change (i.e., decreased cylinder power) in the custom lens wearers (0.17 ± 0.48 D vs. 0.03 ± 0.36 D for the CRT wearers).

Mean binocular unaided visual acuity was 0.02 ± 0.07 for the CRT wearers and 0.04 ± 0.07 for the custom lens wearers. Of the CRT wearers, 98.5% had an unaided binocular visual acuity of 0.19 or less (i.e., 20/30 or better). Of the custom lens wearers, 97.1% had an unaided binocular visual acuity of 0.19 or less.

DISCUSSION

Use of OCR has increased in the United States and worldwide in recent years. This study objectively evaluated clinical outcomes of refractive changes, visual acuity, and corneal health for safety and efficacy for more than 4 years. The findings establish that vision correction is good and that there were isolated minor complications. The results show some important findings regarding differences in refractive changes and age.

Age Differences

As noted in the results, 52% of the study patients were aged 12 years or younger. Because there was slightly higher original refractive error in the younger patients, there was a correspondingly higher refractive change in the group aged 12 years or younger. It is also noteworthy that this group had a slightly flatter flat keratometric reading at baseline, but showed a slightly higher degree of myopia. This suggests that the mostly Asian population of patients aged 12 years or younger had a greater axial length of the eye (not evaluated in this study). Patients in both age groups achieved targeted vision improvement, and unaided visual acuity was not significantly different between the two age groups. Overall, unaided visual acuity was excellent, and patients often commented about their good quality of vision.

Corneal Staining

Corneal staining was a common observation in patients during the study. Low grade corneal staining has also been documented in as many as 79% of non–contact lens wearers and in 33% to 55% of asymptomatic soft lens wearers.33–36 The incidence of corneal staining observed in this study is similar to that in soft lens studies and shows that OCR wear may increase staining after the first night of wear, but not more than soft lenses on a longer-term basis. It is noteworthy that corneal staining was not significantly different between the two age groups. This finding suggests that the slitlamp observation of corneal response to OCR is not different in patients aged 12 years or younger compared to those older than 12 years. These data suggest that increased corneal staining compared to baseline is more related to the degree of original myopia than to age and is not predictive of adverse events. Corneal staining observed in this study was not significantly different than what has been reported with soft lenses worn on a daily-wear basis. It is noteworthy that corneal staining was recorded least often at the 1-month visit. At 1 month in the OCR process, patients have progressed through the initial adaptation but, in general, lens deposits have not become a factor for increased corneal staining. This emphasizes that careful lens maintenance and cleaning are important factors in long-term success with OCR.

Ethnicity

Ethnicity was a significant factor in this study. Of the patients in this study, 68% were Asian and they accounted for almost 95% of the patients aged 12 years or younger. Studies have shown that a high percentage of the Asian population become myopic at a young age.37 Similar studies in non-Asians from the United States, Scandinavia, and Australia showed myopia with less prevalence and a smaller degree.38 One recently published study39 showed the prevalence of myopia in children in rural southern China to be almost 37% by age 13 years and to increase to 54% by age 17 years. Another study of Canadian-Chinese children showed the prevalence of myopia to be 25% at age 6 years and to increase to 71% by age 12 years.40 Parents of these children are concerned about the high degree of myopia and the rate of progression in their children and look for alternatives to correction with spectacles or traditional contact lenses. The population reported in this study is similar to that in a large corneal reshaping practice in New Jersey (N. Despotidis, oral communication, 2006). The Asian population in this study showed a high degree of myopia development before the age of 12 years and interest in OCR treatment.

Lens Type

Paragon CRT lenses and custom reverse-geometry lenses were used in the study. Differences between results in using one lens versus the other are attributable to the fact that the original spherical equivalent and the original cylinder were higher for the patients who wore the custom lenses. As such, there was a greater change in spherical equivalent for the custom group versus the CRT group. Unaided acuity was insignificantly different with each lens-wearing group, but there was a significant decrease in cylinder in the custom lens group. Minor differences noted in the corneal staining between the two types of lenses also may be the result of effecting a larger change in corneal curvature and refraction with the custom-designed lenses.
Adverse Events

Case reports of adverse events with OCR have been published in the last few years. In this study, none of the adverse events resulted in a loss of best-corrected acuity. From the current data and the case reports, a generalized statement regarding safety cannot be made. It can be said that in this setting with the given patient population, there were three adverse events during the 4.25 years observed and that these events did not cause a loss of best-corrected visual acuity. Also, the three patients involved are currently still wearing OCR lenses. After resolution of these events, modifications were made to lens design, cleaning and care regimes, and the follow-up schedule in an effort to avoid recurrence of the event. These events are serious complications and can threaten vision. They are of major concern for doctors, patients, and parents when deciding to start OCR or to continue the process once started.

Retrospectively, the treatment and care provided to these OCR patients suggests other topics of interest. First, are there axial length differences among ethnic groups in children, and secondly, does wearing OCR lenses affect changes in axial length compared to spectacle or soft lens wearers? Finally, experience from this study suggests that measurement of axial length should be standard before starting OCR. Continuing follow-up for this study population is planned to monitor longer-term corneal health, safety, and unaided visual acuity. Future study of OCR may include axial length, whether lens parameter changes are necessary over time and further evaluation of higher-order aberrations with OCR lens use.

In summary, this study showed OCR to be at least as safe and effective at temporarily reducing myopia for children aged 12 years or younger as it is for those older than 12 years. OCR is an alternative to spectacle correction or traditional contact lens correction for children and adults. OCR is of special interest to the Asian population, which develops significant myopia at a young age. It is hoped that this study, in combination with increasing experience, may help practitioners select the right patients for OCR, remain aware of potential problems and limitations, and stimulate additional research that continues to make OCR even safer and better.

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REFERENCES

31. Saviola JP. The current FDA view on overnight orthokeratology: How we got here and where we are going. Cornea 2005;24:770–771.


