Purpose: A percentage tissue altered (PTA) score of ≥40% has been advocated as an independent indicator of post-operative ectasia risk following laser in situ keratomileusis (LASIK). This study was performed to test the hypothesis that refractive procedures, such as laser-assisted sub-epithelial keratectomy (LASEK) or small incision lenticule extraction (SMILE), may alter the range of PTA, within which refractive corneal surgery can be safely performed.

Setting: Retrospective observational study.

Methods: Review of case notes was performed for patients who presented for refractive surgeries, other than LASIK. To determine the risk of corneal ectasia for each patient prior to refractive surgery, we estimated what each patient’s PTA would have been if they had undergone LASIK. The Randleman Ectasia Risk Score System (ERSS) was also calculated.

Results: 114 eyes (66 patients) were included. 94 eyes underwent SMILE. 20 eyes underwent LASIK. A significant proportion of eyes had PTA ≥40% – SMILE eyes: up to 31.9%, LASEK eyes: up to 60.0% (at presumed LASIK flap of 120 μm). The maximum calculated PTA was up to 47.9% in the SMILE group and up to 51.5% in the LASEK group. Using ERSS, 12.8–16% of SMILE eyes and 15.0–80.0% of LASEK eyes would have been considered to have moderate-to-high ectasia risk. No post-surgical ectasia was observed at 3 years.

Conclusion: SMILE and LASEK alter the range of PTA, within which corneal refractive surgery may be performed with a lower risk of developing post-operative corneal ectasia; a safe PTA threshold needs to be determined for these procedures before recommendations for clinical practice can be made.

Keywords: percentage tissue altered, ectasia, refractive surgery, laser in situ keratomileusis, laser-assisted sub-epithelial keratectomy, small incision lenticule extraction

Introduction

Despite significant technological advancements in refractive surgery, laser in situ keratomileusis (LASIK) remains the predominant procedure performed for the correction of myopia. It involves the creation of a corneal flap using a mechanical microkeratome or femtosecond laser; this flap is then reflected back, and an excimer laser is used to reshape the underlying corneal stromal bed. LASIK provides rapid post-operative visual improvement with minimal patient discomfort. It has been shown to produce predictable and stable visual outcomes.

Corneal ectasia is one of the most feared complications following corneal refractive surgery as it can result in an irreversible loss of both uncorrected and best-corrected visual acuity. Photoablation of the cornea results in significant changes in its structure and biomechanical properties. As the anterior corneal stroma is 25% stronger than the...
posterior stroma, it contributes more to the overall cohesive tensile strength.\(^4\)\(^7\) Depending on the planned flap thickness, disrupting the anterior lamellar fibres of the corneal stroma during flap creation in LASIK can, therefore, weaken the cornea’s biomechanical stability. Corneal ectasia occurs when the biomechanical integrity of the cornea is compromised with tissue alteration beyond a safe threshold required to maintain its shape and curvature, and there is consequent, irreversible corneal thinning and steepening.\(^8\) The risk of corneal ectasia following LASIK is well described.\(^9\)\(^11\) It has been reported to occur in 0.03% to 0.9% of eyes undergoing LASIK,\(^9\)\(^12\)\(^14\) but this is likely to be an underestimate of its true incidence. It can occur days to years following surgery.\(^9\)\(^10\)\(^15\)\(^17\)

The percentage tissue altered (PTA) metric, representing the percentage of anterior corneal tissue that is modified during refractive surgery, has been described as an independent indicator of post-operative corneal ectasia risk, for patients undergoing LASIK.\(^18\)\(^21\) The PTA takes into account the integrated relationship between pre-operative corneal thickness, ablation depth, and flap thickness and provides a measure of the amount of biomechanical change that occurs following a LASIK procedure. Among patients with normal pre-operative corneal topographies, a PTA of ≥40% was identified as the single, most significant independent indicator for predicting the development of corneal ectasia (OR 223.3, CI 28.8 to 1729.7).\(^18\) Among patients with suspicious pre-operative corneal topographies, PTA has still been reported to provide the highest discriminative capability for identifying those at risk of corneal ectasia. In addition, for patients with remarkable signs of pre-operative topographic abnormality, it has been found that a lower threshold value of PTA may predict significant risk.\(^19\)

Alternative laser refractive procedures to LASIK for the surgical correction of myopia include: advanced surface ablation (ASA) procedures such as laser-assisted sub-epithelial keratectomy (LASEK) and small incision lenticule extraction (SMILE). In the absence of creating stromal flaps, ASA procedures are well established and are considered to offer better biomechanical stability than LASIK.\(^22\) SMILE is the latest technique introduced for flapless corneal refractive surgery. It involves the creation of a lenticule with a femtosecond laser and the removal of this lenticule through a small incision.\(^23\)\(^24\) Since its introduction in 2011, there has been a steady increase in the number of SMILE surgeries performed for the treatment of myopia, as an alternative to excimer-based laser refractive surgeries. One of the postulated benefits of SMILE is better post-operative corneal biomechanical integrity. This postulation is based on mathematical modeling and finite-element analyses.\(^25\)\(^26\) Clinically, however, this has been more difficult to show. Two studies demonstrated that for eyes with myopia of >6.0 D, there was less disruption of biomechanical stability, measured with an ocular response analyzer (ORA; Reichert Ophthalmic Instruments, Buffalo, NY, USA), after SMILE compared to LASIK.\(^27\)\(^28\) A third study used both the ORA and Corvis ST (Oculus, Wetzlar, Germany), also arrived at the same conclusion.\(^29\) However, other studies have reported no significant differences between SMILE and LASIK corneal biomechanics, evaluated using either the ORA\(^30\) or Corvis ST.\(^30\)\(^32\) Thus, the purported benefit of SMILE offering better biomechanical stability than LASIK has not yet been determined formally in clinical practice. Indeed, some cases of corneal ectasia following SMILE have been reported; albeit, many of these cases had preoperative topographical abnormalities.\(^33\)\(^38\)

Given that less corneal stromal fibres are disrupted in flapless procedures such as ASA or SMILE (compared to LASIK), we hypothesized that these procedures may alter the range of PTA within which corneal refractive surgery may be performed safely. Like ASA, there is likely to be a “grey area” where a patient may potentially be at higher risk of ectasia with LASIK, but may be able to undergo SMILE safely. To test this hypothesis, we retrospectively reviewed the case notes of patients who presented for corneal refractive surgeries other than LASIK (i.e. SMILE or LASEK). Based on their examination findings at presentation, we determined the Randleman Ectasia Risk Score System (ERRS) grades and calculated the PTA scores they would have had hypothetically if they had undergone LASIK. Post-operatively, each patient was monitored for signs of corneal ectasia.

**Methods**

This retrospective study reviewed case notes of patients who presented for refractive surgery other than LASIK, from January 2012 to January 2014 at the Singapore National Eye Centre, an ophthalmic tertiary referral centre. The study protocol was approved by the SingHealth Institutional Review Board (CIRB reference number: 2011/109/A), and the study adhered to the tenets of the Declaration of Helsinki. All participants consented to the review of their medical records.
Participants
All patients were adults (≥21 years of age) who had a diagnosis of myopia with a manifest refraction of ≤10.00 dioptres (D) spherical equivalent prior to refractive surgery. Patients with keratoconus or suspected keratoconus on clinical examination or corneal tomography, including suspicious or abnormal Belin–Ambrósio enhanced ectasia display (BAD), were excluded, as were patients with previous ocular surgery, autoimmune conditions, severe dry eyes, active ocular, or systemic disease likely to affect corneal wound healing and patients using any systemic or ocular medication.

Data Collection
Data collected from case notes included patient demographics, pre- and post-operative manifest refraction, pre-operative and post-operative uncorrected distance visual acuity (UDVA), pre-operative and post-operative best-corrected distance visual acuity (CDVA), pre-operative central corneal thickness (CCT), and indices from Scheimpflug colour corneal tomography (Pentacam, Oculus).

Pre-Operative Assessment
All patients had a standard pre-operative evaluation including slit-lamp and dilated fundus examinations, logMAR UDVA assessment, subjective manifest refractions with CDVA assessment, CCT measurement using ultrasound corneal pachymetry, and Scheimpflug colour corneal tomography (Pentacam, Oculus).

Post-Operative Assessment
All patients had a standard post-operative evaluation including slit-lamp examination, logMAR UDVA, subjective manifest refractions with CDVA, and Scheimpflug colour corneal tomography (Pentacam, Oculus). Post-operative evaluations were performed at the following time points: 1 day, 1 week, and 1, 3, 12, and 36 months. All adverse events were documented.

For the peri-operative assessments, a single optometrist (MF) obtained all visual acuities and refraction measurements. Measurements were conducted in a single room under consistent photopic lighting conditions. Being a single-centred study, similar device platforms to obtain ultrasound pachymetry and corneal tomography were used for all patients.

Risk Of Corneal Ectasia If Lasik Were To Be Performed
To determine the risk of corneal ectasia for each patient prior to refractive surgery, we estimated what each patient’s PTA would have been hypothetically if they had undergone LASIK. To further stratify the ectasia risk, we also calculated the ERSS scores.

PTA described for LASIK is calculated with the formula: PTA = (FT + AD)/CCT (FT = flap thickness, AD = ablation depth, and CCT = pre-operative central corneal thickness). Using this formula, the PTA was estimated for all eyes, using standard LASIK ablation depths and flap thicknesses (100 μm, 110 μm, and 120 μm). The ERSS score was determined using methods described by Randleman et al. Briefly, a point score was assigned to each of the following variables: (i) pre-operative topographic pattern, (ii) RSB thickness, (iii) patient age, (iv) pre-operative CCT, and (v) pre-operative myopia. The ERSS is the sum of these scores. Risk of corneal ectasia following LASIK was classified as: low (ERSS = 0 to 2), moderate (ERSS = 3), or high (ERSS ≥4). RSB thickness was estimated as pre-operative CCT (flap thickness + calculated central ablation depth thickness).

Post-Operative Corneal Ectasia
Post-operative corneal ectasia was defined as: (i) increasing myopia or astigmatism (change in manifest refraction of 2 dioptres or more in either sphere or cylinder) and (ii) progressive abnormality on corneal topographies not present in pre-operative scans. Topographic abnormalities include: inferior topographic steepening of 3 dioptres or more compared with the immediate postoperative appearance and asymmetric patterns (e.g. asymmetric bowtie, inferior steep or skewed radial axis, and increase in posterior elevation).

Surgical Procedures
SMILE
All SMILE procedures were performed by a single surgeon (JSM) using a 500-kHz femtosecond laser platform (VisuMax, Carl Zeiss Meditec, Germany). A previously described standard technique with the following parameters was used: 120 μm cap thickness, 7.5 mm cap diameter, 6.0 to 6.5 mm lenticule optical zone, and 145 nJ power with side-cut angles at 90 degrees.

LASEK
All LASEK procedures were performed by a single surgeon (JSM) using an excimer laser platform (Wavelight EX500, Alcon, Texas, USA). Corneal epithelium was
debrided following a 60 s 20% alcohol application. Mitomycin C 0.02% was applied for 60 s to the stromal bed, following excimer laser treatment.

Statistical Analysis

The programme, Statistical Package for the Social Sciences software (version 17.0, SPSS, Inc.), was used for statistical analysis. Means and standard deviations were calculated for all continuous variables. Frequency distribution and percentages were determined for categorical variables. Comparisons between post-operative and pre-operative values, for manifest refractions and visual acuities, were carried out using the paired sample t-test. Differences in the distributions between groups were analysed using the independent t-test. Differences in the distributions of categorical variables between groups were analysed using the Chi-squared test. A p value <0.05 was considered significant. As variance between eyes from the same subject is usually less than the variance between different subjects, for pre-operative and post-operative data, statistical adjustments were made in accordance to previously published guidelines to correct for potential confounding effects.

Results

The study population included 114 eyes (66 patients) (Table 1). Of these, 94 eyes (82.5%) from 56 patients had SMILE procedures. 20 eyes (17.5%) from 10 patients had LASEK. The mean age of patients was 30.0 ± 7.0 years, and 35/66 (53.0%) were male. The mean pre-operative spherical equivalent refractive error treated was −5.71 ± 2.07 D.

Table 2 presents the pre-operative assessments and post-operative outcomes of all recruited patients. As this study includes patients with 3-years follow-up, the visual and refractive outcomes of SMILE were from procedures performed before 2013. This reflects data from our previous paper published in 2015. Since optimizing centration and nomogram adjustment, our current SMILE visual outcomes have improved. However, as this study evaluates pre-operative corneal ectasia risks from PTA and ERSS, rather than post-operative visual or refractive outcomes, results of risks determined from in this study would not have been different even if data of eyes that had more recently undergone SMILE were used.

Percentage Tissue Altered (PTA) Of All 114 Eyes In Study, Had LASIK Been Performed

For a presumed LASIK flap thickness of 120 μm, the mean PTA score of all study eyes would have been 37.4 ± 6.2%. A total of 42/114 (36.8%) eyes would have had PTA ≥40% (Figure 1A). For a presumed LASIK flap thickness of 110 μm, the mean PTA score would have been 35.5 ± 6.1%. A total of 29/114 (25.4%) eyes would have had PTA ≥40%. For a presumed LASIK flap thickness of 100 μm, the mean PTA would have been 33.7 ± 6.0%, and 18/114 (15.8%) eyes would have had PTA ≥40%.

Results For Patients Who Had Small Incision Lenticule Extraction (SMILE) [n=94]

The mean pre-operative spherical equivalent refractive error of eyes was −5.76 ± 2.17 D.

PTA Scores And ERSS Calculations For Eyes Which Had SMILE (Figure 1B, Table 1)

Had these eyes undergone LASIK, for a presumed flap thickness of 120 μm, the mean PTA score, of eyes that had SMILE, would have been 36.3 ± 5.8%. A total of 30/94 (31.9%) eyes would have had PTA ≥40% and the maximum PTA score would have been 47.9%. Also, 31/94 (33.0%) eyes would have had ERSS = 0, 26/94 (27.7%) eyes would have had ERSS = 1, 22/94 (23.4%) eyes would have had ERSS = 2, 11/94 (11.7%) eyes would have had ERSS = 3, and 4/94 (4.3%) eyes would have had ERSS ≥4. Thus, 15/94 (16.0%) eyes would have been considered to have moderate-to-high risk of ectasia following LASIK.

For a presumed LASIK flap thickness of 110 μm, the mean PTA score of these eyes would have been 34.5 ± 5.8%. A total of 18/94 (19.1%) eyes would have had PTA ≥40% and the maximum PTA score would have been 46.1%. A total of 32/94 (34.0%) eyes would have had ERSS = 0, 30/94 (31.9%) eyes would have had ERSS = 1, 20/94 (21.3%) eyes would have had ERSS = 2, 8/94 (8.5%) eyes would have had ERSS = 3, and 4/94 (4.3%) eyes would have had ERSS ≥4. Thus, 12/94 (12.8%) eyes would have been considered to have moderate-to-high risk of ectasia following LASIK.

For a presumed LASIK flap thickness of 100 μm, the mean PTA score of these eyes would have been 32.7 ± 5.8%. A total of 10/94 (10.6%) eyes would have had PTA ≥40% and the maximum calculated PTA score would have been 44.2%. Also, 33/94 (35.1%) eyes would have had ERSS = 0, 29/94 (30.9%) eyes would have had ERSS = 1, 20/94 (21.3%) eyes would have had ERSS = 2, 9/94 (9.6%) eyes would have had ERSS = 3, and 3/94 (3.2%) eyes would have had ERSS = 4. Thus, 12/94 (12.8%) eyes would have been considered to have moderate-to-high risk of developing ectasia following LASIK.
At 3 months following SMILE, the mean post-operative LogMAR UDV A was 0.04 ± 0.10. This was 0.0 ± 0.08 at 3 years. The efficacy index (ratio of postoperative UDV A to mean preoperative CDVA) was 0.92 ± 0.20 and 0.99 ± 0.21 at 3 months and 3 years, respectively (Figure 2A, B).

Predictability
At 3 months, 84.0% of eyes were within ± 1.00D of target refraction (Figure 2C). The attempted and achieved corrections are shown in Figure 2D.

Stability
Three-year follow-up data were available for 16 patients (31 eyes). Changes in the spherical equivalent of manifest refraction, as a function of time, are shown in Figure 2E. For all 31 eyes, the post-operative refraction remained stable. Between the 3 months and 3 years follow-up time points, the difference in the mean spherical equivalent of manifest refraction was: −0.32 ± 0.39 D; 14.5% of eyes had a change in SE by >0.50 D.

Safety And Adverse Events
At 3 months follow-up, the mean post-operative logMAR CDVA was −0.04 ± 0.07. This was −0.01 ± 0.06 at 3 years. The safety index (ratio of mean postoperative CDVA to mean preoperative CDVA) was 1.04 ± 0.16 and 1.17 ± 0.18, at 3 months and 3 years, respectively (Figure 2F). At 3 months follow-up, 21/94 (22.3%) eyes gained one line in CDVA and 5/94 (5.3%) eyes lost one line. At 3 years follow-up, 16/31 (51.6%) eyes gained one line in CDVA and 1/31 (3.2%) eyes lost one line. None of the 31 eyes, that had 3-year follow-up data, were found to have post-surgical corneal ectasia. Of these eyes, for a presumed LASIK flap thickness of 120 μm, 110 μm, and 100 μm, the proportion of eyes with PTA ≥40% would have been 9/31 (29.0%), 8/31 (25.8%), and 4/31 (12.9%) respectively, had LASIK been performed.

**Table 1 Pre-Operative Patient Demographics Of All Eyes In Study Who Subsequently Underwent Small Incision Lenticule Extraction (SMILE) Or Laser-assisted sub-epithelial keratectomy (LASEK)**

<table>
<thead>
<tr>
<th>Variables</th>
<th>Total Eyes</th>
<th>SMILE*</th>
<th>LASEK</th>
<th>p-value †‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number (%)</td>
<td>114 (100%) of 66 patients</td>
<td>94 (82.5%) of 56 patients</td>
<td>20 (17.5%) of 10 patients</td>
<td>0.278 †‡</td>
</tr>
<tr>
<td>Age (years) (mean [M] ± standard deviation [SD], range [R])</td>
<td>30.0 ± 7.0 to 53.1</td>
<td>30.7 ± 7.3 to 53.1</td>
<td>27.4 ± 4.2 to 33.4</td>
<td>0.278 †‡</td>
</tr>
<tr>
<td>Male gender (n, %)</td>
<td>35 (53.0%)</td>
<td>28 (50.0%)</td>
<td>7 (70.0%)</td>
<td>0.226 †§</td>
</tr>
<tr>
<td>Central corneal thickness (M ± SD, R)</td>
<td>541.1 ± 39.1, 437 to 646</td>
<td>553.6 ± 28.9, 504 to 646</td>
<td>482.25 ± 24.4, 467 to 528</td>
<td>&lt;0.0001 †‡</td>
</tr>
<tr>
<td>Sphere (D) (M ± SD, R)</td>
<td>−5.42 ± 2.06, 0.00 to −9.50</td>
<td>−5.47 ± 2.15, 0.00 to −9.50</td>
<td>−5.20 ± 1.55, −3.00 to −7.75</td>
<td>0.748 †‡</td>
</tr>
<tr>
<td>Cylinder (D) (M ± SD, R)</td>
<td>1.00 ± 0.65, 0.00 to 3.50</td>
<td>1.00 ± 0.69, 0.00 to 3.50</td>
<td>0.99 ± 0.46, 0.00 to 2.00</td>
<td>0.467 †‡</td>
</tr>
<tr>
<td>Spherical equivalent (D) (M ± SD, R)</td>
<td>−5.71 ± 2.07, −1.25 to −10.00</td>
<td>−5.76 ± 2.17, −0.75 to −9.75</td>
<td>−5.47 ± 1.52, −3.00 to −7.63</td>
<td>0.883 †‡</td>
</tr>
<tr>
<td>Percentage tissue altered (PTA)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LASIK-100 (M ± SD, R)</td>
<td>33.7 ± 6.0, 17.2 to 46.9</td>
<td>32.7 ± 5.8, 17.2 to 44.2</td>
<td>38.1 ± 5.4, 30.3 to 46.9</td>
<td>&lt;0.0001 †‡</td>
</tr>
<tr>
<td>LASIK-110 (M ± SD, R)</td>
<td>35.5 ± 6.1, 18.7 to 49.2</td>
<td>34.5 ± 5.8, 18.7 to 46.1</td>
<td>40.1 ± 5.5, 32.2 to 49.2</td>
<td>&lt;0.0001 †‡</td>
</tr>
<tr>
<td>LASIK-120 (M ± SD, R)</td>
<td>37.4 ± 6.2, 20.3 to 51.5</td>
<td>36.3 ± 5.8, 20.3 to 47.9</td>
<td>42.2 ± 5.5, 34.1 to 51.5</td>
<td>&lt;0.0001 †‡</td>
</tr>
</tbody>
</table>

Notes: *31 eyes had 3-year followup data. †Comparing SMILE and LASEK eyes. ‡Comparing SMILE and LASEK eyes with 3-year follow-up data. §Paired sample t-test. ††Chi-square test.

Abbreviations: D, dioptries; LASIK-100, PTA based on flap thicknesses of 100 μm; LASIK-110, PTA based on flap thicknesses of 110 μm; LASIK-120, PTA based on flap thicknesses of 120 μm.
Results For Patients Who Had Laser-assisted sub-epithelial keratectomy (LASEK) [n=20]

20 eyes (10 patients) underwent LASEK for myopia during the same time period. The mean pre-operative spherical equivalent refractive error of eyes was $-5.47 \pm 1.52$ D. Three-year follow-up data were available for all 20 eyes. Pre-operative assessments and post-operative outcomes are presented in Table 2.

Baseline characteristics such as age, sex, and pre-operative refraction were similar to the SMILE eyes that had 3-year follow-up data (Table 1).

PTA Scores And ERSS Calculations For Eyes That Had LASEK (Figure 1B, Table 1)

Had these eyes undergone LASIK, for a presumed flap thickness of 120 μm, the mean PTA score of eyes that had LASEK would have been 42.2 ± 5.5%. A total of 12/20 (60.0%) eyes would have had PTA ≥40% and the maximum calculated PTA score would have been 51.5%. Also, 1/20 (5.0%) eyes would have had ERSS = 0, 3/20 (15.0%) eyes would have had ERSS = 2, 1/20 (5.0%) eyes would have had ERSS = 3, and 15/20 (75.0%) eyes would have had ERSS ≥4. Thus, 16/20 (80.0%) eyes would have been considered to have moderate to high risk of ectasia following LASIK.

For a presumed LASIK flap thickness of 110 μm, the mean PTA score of these eyes would have been 40.1 ± 5.5%. A total of 11/20 (55.5%) eyes would have had PTA ≥40% and the maximum calculated PTA score would have been 49.2%. Also, 1/20 (5.0%) eyes would have had ERSS = 0, 4/20 (20.0%) eyes would have had ERSS = 2, and 15/20 (75.0%) eyes would have had ERSS ≥4. Thus, 15/20 (75.0%) eyes would have been considered to have moderate-to-high risk of ectasia following LASIK.

For a presumed LASIK flap thickness of 100 μm, the mean PTA score of these eyes would have been 38.1 ± 5.4%. A total of 8/20 (40.0%) eyes would have had PTA ≥40% and the maximum calculated PTA score would have been 46.9%. 8/20 (40.0%) eyes would have had ERSS = 0, 5/20 (25.0%) eyes would have had ERSS = 1, 4/20 (20.0%) eyes would have had ERSS = 2, 2/20 (10.0%) eyes would have had ERSS = 3, and 1/20 (5.0%) eyes would have had ERSS = 4. Thus, 3/20 (15.0%) eyes would have been considered to have moderate-to-high risk of ectasia following LASIK.

Table 2 Pre-Operative And 3 Months Post-Operative Patient Demographics In Eyes Undergoing SMILE Procedures (n = 94) And LASEK Procedures (n = 20) For Treatment Of Myopia

<table>
<thead>
<tr>
<th>Variables</th>
<th>SMILE</th>
<th>LASEK</th>
<th>p-value</th>
<th>SMILE</th>
<th>LASEK</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sphere (D)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean ± SD</td>
<td>$-5.47 \pm 2.15$</td>
<td>$-5.20 \pm 1.55$</td>
<td>&lt;0.0001*</td>
<td>$-5.47 \pm 1.52$</td>
<td>$-5.20 \pm 1.55$</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>Range</td>
<td>0.00 to $-9.50$</td>
<td>$-3.00$ to $-7.75$</td>
<td></td>
<td>0.00 to 1.00</td>
<td>0.00 to 1.00</td>
<td></td>
</tr>
<tr>
<td>Cylinder (D)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean ± SD</td>
<td>$1.00 \pm 0.69$</td>
<td>$0.99 \pm 0.46$</td>
<td>&lt;0.0001*</td>
<td>$0.99 \pm 0.46$</td>
<td>$0.19 \pm 0.23$</td>
<td>0.001*</td>
</tr>
<tr>
<td>Range</td>
<td>0.00 to 3.50</td>
<td>0.00 to $-2.00$</td>
<td></td>
<td>0.00 to 1.00</td>
<td>0.00 to $-0.50$</td>
<td></td>
</tr>
<tr>
<td>Spherical equivalent (D)</td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean ± SD</td>
<td>$-5.76 \pm 2.17$</td>
<td>$-5.47 \pm 1.52$</td>
<td>&lt;0.0001*</td>
<td>$-5.76 \pm 2.17$</td>
<td>$-5.47 \pm 1.52$</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>Range</td>
<td>$-0.75$ to $-9.75$</td>
<td>$-3.00$ to $-7.63$</td>
<td></td>
<td>$-0.75$ to $-9.75$</td>
<td>$-3.00$ to $-7.63$</td>
<td></td>
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<tr>
<td>UDVA (LogMAR)</td>
<td></td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>Mean ± SD</td>
<td>1.57 ± 0.37</td>
<td>1.45 ± 0.42</td>
<td>&lt;0.0001*</td>
<td>1.57 ± 0.37</td>
<td>1.45 ± 0.42</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td>Range</td>
<td>0.40 to 1.85</td>
<td>0.54 to 1.85</td>
<td></td>
<td>0.40 to 1.85</td>
<td>0.54 to 1.85</td>
<td></td>
</tr>
<tr>
<td>CDVA (LogMAR)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean ± SD</td>
<td>$-0.01 \pm 0.03$</td>
<td>$0.0 \pm 0.00$</td>
<td>0.001*</td>
<td>$-0.01 \pm 0.03$</td>
<td>$0.0 \pm 0.00$</td>
<td>0.168*</td>
</tr>
<tr>
<td>Range</td>
<td>$-0.15$ to 0.10</td>
<td>$-0.12$ to 0.00</td>
<td></td>
<td>$-0.15$ to 0.10</td>
<td>$-0.12$ to 0.00</td>
<td></td>
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Note: *Paired sample t-test (corrected for correlation between eyes from same subjects).

Abbreviations: SD, standard deviation; D, dioptres; UDVA, uncorrected distance visual acuity; CDVA, corrected distance visual acuity.
Stability

Post-operative refraction remained stable within the 3-year follow-up period. Between the 3-month and 3-year follow-up time points, the difference in the mean spherical equivalent of manifest refraction was: $-0.30 \pm 0.48$ D; 25.0% of eyes had a change in SE of $>0.50$ D.

Figure 1 Percentage tissue altered (PTA) of eyes in study had laser in-situ keratomileusis (LASIK) been performed; these eyes subsequently underwent small incision lenticule extraction (SMILE) or laser-assisted sub-epithelial keratectomy (LASEK) procedures. (A) Scatter plot of PTA based on a LASIK flap thickness of 120μm highlighting patients with 36 months follow-up showing a significant proportion of eyes having PTA $\geq 40%$; (B) bar chart showing the proportion of patients in the study with PTA above fixed thresholds of 30%, 40%, and 45% assuming different LASIK flap thicknesses.
Safety And Adverse Events

None of the 20 eyes that underwent LASEK had postsurgical corneal ectasia at 3-year follow-up.

Discussion

In current clinical practice, several surgical options are available to myopic patients seeking refractive surgery.
Laser-based refractive surgeries including LASIK, ASA procedures, and SMILE are currently the most popular. It is important for both clinicians and patients to understand the risks associated with the different procedures offered. Although most patients who have developed corneal ectasia following corneal refractive surgery have been shown to have identifiable risk factors pre-operatively, patients with normal pre-operative assessments are not necessarily insusceptible. This latter group poses specific challenges to the refractive surgeon.

Various nomograms and imaging-based technologies have been introduced to screen for and identify patients at risk of developing post-operative ectasia after LASIK. An example of a commonly used screening nomogram is the Ectasia Risk Score System (ERSS). This is based on known risk factors for the development of post-LASIK corneal ectasia, including: young age, high myopia, thin corneas, suspicious topographic patterns, and low residual stromal bed (RSB) thickness. Randleman et al validated this weighted risk stratification scale in 2008 and concluded that it has high sensitivity and specificity in predicting ectasia following LASIK.

Imaging-based screening tools are predominantly based on indices derived from topographic parameters. Examples include the videokeratography-derived keratoconus prediction index (KPI) and the associated keratoconus index (KCI) by Maeda et al. Likewise, Rabinowitz et al introduced the central keratometric (K) index, the inferior-superior (I-S) index, and the keratomy, inferior–superior value, simulated astigmatism (KISA) index, as screening indicators for ectasia. Other more recent imaging-based tools are based on corneal elevation profiles, corneal thickness profiles, difference corneal reference surfaces, aberrometry, or combinations of these parameters (such as the cone location and magnitude index algorithm). These parameters are reported to have better sensitivities for detecting subclinical keratoconus. Software using some of these parameters, such as the SCORE Analyzer and BAD, have been incorporated within corneal tomography systems to screen for patients at risk of developing postoperative ectasia.

More recently, devices that screen for ectasia risk through assessing the in vivo biomechanics of patient corneas have gained popularity. Such devices include the ocular response analyzer (ORA; Reichert Ophthalmic Instruments) and the Corvis ST (Oculus). Some parameters derived from the ORA waveform, such as keratoconus match index (KMI) and the keratoconus match probability (KMP), have been shown to have good predictive accuracy for corneal ectasia risk, although other parameters derived using ORA applanation pressures, such as corneal hysteresis and corneal resistance factors, have not. The Corvis ST visualizes and monitors corneal deformation using an ultra-high-speed Scheimpflug camera. The Corvis biomechanical index (CBI), calculated using the deformation corneal response and the horizontal corneal thickness profile, has been shown to have high sensitivity and specificity in detecting corneal ectasia.

The tomographic/biomechanical index (TBI) is another parameter that has been shown to have high accuracy in screening for subclinical corneal ectasia in patients with normal corneal topographies. It is derived by combining biomechanical data from the Corvis ST with data obtained from Pentacam HR Scheimpflug-based corneal tomography (Oculus).

PTA has been found to be a reliable measure for assessing the risk of post-operative ectasia, following LASIK, for eyes with normal pre-operative corneal topographies. Santinago et al have shown that PTA had higher odds ratio and higher predictive capabilities for ectasia risk than moderate-to-high ERSS values, or individual factors from the ERSS system (CCT, RSB thickness, spherical equivalent refraction, ablation depth, and age of patients). With a sensitivity of 97% and specificity of 89%, a PTA score ≥40% was recommended as a robust screening indicator for risk of corneal ectasia for patients with normal pre-operative corneal topography, planning to undergo LASIK. Furthermore, Santinago et al calculated the PTA for several case reports of ectasia following LASIK, with normal pre-operative corneal topographies, and found that PTA scores were >40% for all these cases.

Nonetheless, PTA does have its limitations. Accurate estimation of PTA depends on the actual thickness of the LASIK flaps created. It has been reported that the standard deviation of the thickness of flaps cut by mechanical microkeratomes can range from 12.4 μm to 24.0 μm. For LASIK flaps cut by femtosecond lasers, standard deviations can range from 5.6 μm to 16.0 μm. Consequently, even when thin LASIK flaps are planned, thicker flaps may be created, resulting in increased risk of post-LASIK corneal ectasia. Such variation in flap thickness may explain situations where patients develop unexpected post-LASIK corneal ectasia.

In this study, using PTA and ERSS, we determined the risk of post-operative ectasia for eyes that presented for corneal refractive surgeries other than LASIK with normal...
corneal topographies and found that a significant number of eyes would have been considered to be “at risk” had LASIK been performed. Depending on presumed LASIK flap thickness, 10.6% to 31.9% of eyes that subsequently underwent SMILE, and 40.0% to 60.0% of eyes that subsequently underwent LASEK, had PTA ≥40%. Again, depending on presumed LASIK flap thickness, the maximum calculated PTA scores ranged from 44.2% to 47.9% in the SMILE group and 46.9% to 51.5% in the LASEK group (Table 1). Data from previous studies suggest a PTA ≥40% has a specificity of 89% in screening for corneal ectasia. By taking this into account, had these eyes undergone LASIK, we would have expected to observe post-operative corneal ectasia in at least 8 eyes in the SMILE group and 7 eyes in the LASEK group. When the ERSS was applied, 12.8% to 16.0% of SMILE eyes and 15.0% to 80.0% of LASEK eyes would have been considered to have moderate-to-high risk of post-LASIK ectasia. None of the eyes in this cohort were found to have post-operative corneal ectasia after a follow-up period of 3 years.

We can thus infer from our results that, compared to LASIK, the range of PTA, within which corneal refractive surgery can be performed safely on eyes with normal corneal topography, is likely to be wider when either SMILE or LASEK is performed. Opting for these procedures over LASIK may have minimized the risk of ectasia for these patients. None of our patients had pre-operative abnormal corneal topography. A recent study looking at PTA scores for patients with post-operative corneal ectasia after SMILE suggests that the threshold PTA value, below which surgery is safe, is significantly lower in patients with pre-operative topographic abnormality. Another similar study looking at patients with post-operative ectasia after photorefractive keratectomy also arrived at similar conclusions. Additionally, from our data, compared to SMILE eyes, a higher proportion of LASEK eyes would have had PTA scores above set thresholds (Table 1, Figure 1B); still, none of these eyes developed corneal ectasia. It is therefore probable that the “safe” PTA range is wider for LASEK than for SMILE.

As PTA represents the percentage of anterior corneal tissue that is modified during refractive surgery, for both SMILE and LASEK procedures, we can deduce that the alteration of a higher amount of corneal tissue is tolerated, before corneal ectasia develops. Furthermore, we can infer from these results that eyes with similar pre-operative parameters may be more likely to develop post-operative corneal ectasia with LASIK than with either SMILE or LASEK. This study, therefore, provides some clinical evidence to support the hypothesis that SMILE procedure is associated with better biomechanical corneal stability than LASIK.

Compared to LASIK, the anterior lamellar fibres are relatively preserved in SMILE. Through human ex-vivo studies, it has been shown that vertical cuts through corneal lamellae, which occurs during LASIK flap creation, contribute to the loss of corneal structural integrity. This correlates with the depth of incisions. However, such changes in corneal biomechanics are not seen when horizontal delamination incisions are made. Often, fewer vertical incisions are made in current SMILE techniques compared to LASIK flap creation. This may explain why higher PTA scores may still be considered safer when SMILE is used in place of LASIK. As corneal structural integrity appears to be weakened when any side cuts are made, the absence of these incisions in LASEK may explain why it may be possible that LASEK may have a wider safe range of PTA than SMILE.

One potential limitation of this study is that we did not calculate PTA for these eyes using parameters that would have been more relevant for SMILE (e.g., lenticule thickness). As our aim was to compare the range of PTA values within which SMILE/LASEK was safe to established LASIK values. We standardized the method of PTA calculation to one where we supposed all patients would undergo LASIK. Taking into account the differences between LASIK and SMILE procedures, Santhiago et al and Moshirfar et al have recently proposed possible alternative calculations for PTA in SMILE. However, the use of these alternative calculations for PTA for eyes undergoing SMILE and their correlation with ectasia risk requires further evaluation. Indeed, the inaccuracies of the PTA metric, used in eyes undergoing refractive procedures other than LASIK, may explain the reason for the absence of ectasia even with high risk scores in our cohort.

Another limitation of this study is that, as no cases in our cohort developed postoperative ectasia, we are not able to determine a safe upper limit of PTA score for either SMILE or LASEK. Furthermore, the follow-up duration for this study was limited to 3 years. We acknowledge that corneal ectasia can occur at any time following corneal refractive surgery. However, it often presents within the first 12 post-operative months. Previous studies investigating post-LASIK ectasia have also used similar follow-up durations for identifying its incidence. Lastly, this study was conducted in a cohort of myopic...
patients in Singapore, a predominantly Asian population, with a high incidence of myopia. As differences in corneal biomechanical properties have been found in different ethnic groups,\textsuperscript{24} results from this study may not be applicable to all study populations.

In conclusion, both SMILE and LASEK are likely to alter the range of PTA within which corneal refractive surgery may be performed with lowered risks of post-operative ectasia. Eyes with PTA above the threshold of >40% and considered to be at high risk of post-LASIK ectasia could potentially have either SMILE or LASEK. There may be patients with normal topography who, when found to have PTA >40%, elect to have non-corneal procedures such as implantable contact lenses. For these patients, we may be able to offer either SMILE or LASEK as safe, less-invasive alternatives. However, it was beyond the remit of this study to provide recommended guidance of PTA scores that SMILE or LASEK can be performed safely. To better ascertain this, further evaluation of the use of PTA as a tool in assessing ectasia risk following SMILE or LASEK is required; and, a safe upper limit of PTA for these procedures needs to be established.

Ethics
SingHealth Centralised Institutional Review Board.

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Disclosure
Dr Marcony R Santhiago is a consultant at Ziemer, Alcon, and Zeiss. The authors report no other conflicts of interest in this work.

References


