Recent advances in corneal collagen cross-linking

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Corneal collagen cross-linking has become the preferred modality of treatment for corneal ectasia since its inception in late 1990s. Numerous studies have demonstrated the safety and efficacy of the conventional protocol. Our understanding of the cross-linking process is ever evolving, with its wide implications in the form of accelerated and pulsed protocols. Newer advancements in technology include various riboflavin formulations and the ability to deliver higher fluence protocols with customised irradiation patterns. A greater degree of customisation is likely the path forward, which will aim at achieving refractive improvements along with disease stability. The use of cross-linking for myopic correction is another avenue under exploration. Combination of half fluence cross-linking with refractive correction for high errors to prevent post LASIK regression is gaining interest. This review aims to highlight the various advancements in the cross-linking technology and its clinical applications.

Key words: Accelerated cross-linking, corneal collagen cross-linking, customized cross-linking, photorefractive intrastromal cross-linking, pulsed cross-linking

Keratoconus is a degenerative condition associated with progressive corneal ectasia and thinning. Visual loss occurs following progressive myopia, irregular astigmatism, and corneal scarring.[1] Corneal collagen cross-linking was first described by Spoerl et al. as a modality for increasing the corneal biomechanical strength to halt disease progression. [2] Over the years, our understanding of the cross-linking process has evolved. Various advances have highlighted the opportunity to optimize the procedure improving efficacy and refractive outcomes. [3-5] These include new riboflavin formulations, higher ultraviolet A (UV-A) irradiance sources, and programmable UV-A patterns. The aim of this review is to detail upon the various advances and their clinical applications.

Accelerated Cross-linking

In 1998, corneal collagen cross-linking was first proposed as a treatment modality to stabilize the ectatic cornea. [5] The standard Dresden protocol entails UV-A treatment over a central 9 mm zone at an irradiance of 3.0 mW/cm² for 30 min, delivering a fluence of 5.4 J/cm². [6] Although various studies have demonstrated safety and efficacy of this protocol, an increased intraoperative time is a major drawback. Accelerated protocols have evolved in an attempt to overcome the limitations of conventional cross-linking, while maintaining the efficacy of results.

The Bunsen and Roscoe law of reciprocity states that the effect of a photochemical or photobiological reaction is directly proportional to the total irradiation dose, irrespective of the time span over which the dose is administered. [7] Thus, the same effect can be achieved by either applying a higher intensity for a shorter duration or a lower irradiation for a longer period. Stress–strain measurements of porcine corneal strips, comparing conventional with accelerated cross-linking (10 mW/cm² for 9 min), demonstrated similar results in both groups. [8] Wernli et al. demonstrated a failure of the accelerated treatment for higher irradiance protocols (90 mW/cm²) in porcine eyes. [9] A significant difference in Young’s modulus was noted between treatment groups up to 45 mW/cm² and control group. However, treatment groups from 50 mw/cm² up to 90 mW/cm² demonstrated no significant difference. Hence, the Bunsen and Roscoe reciprocity law is valid only for illumination intensity up to 45 mW/cm² and an irradiance time >2 min.

Various clinical studies demonstrated that the accelerated corneal cross-linking (CXL) provided stabilization of disease process along with significant flattening in certain protocols [Table 1].

Refractive and keratometric stability

The first clinical results were described by Kanellopoulos comparing conventional cross-linking to the accelerated protocol (7 mW/cm² for 15 min) in 42 eyes with a mean follow-up of 46 months. [10] The study demonstrated a similar efficacy and refractive effect between the two groups, with no adverse effects.

Since then, numerous studies have been published using varied irradiation intensities and exposure time, with a standard total fluence of 5.4 J/cm².

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Table 1: Clinical results of accelerated cross-linking

| Author             | Number of eyes/duration of follow-up (months) | Irradiation intensity (mW/cm²)/Time (min) | Visual acuity | Keratometry (diopters) |
|--------------------|-----------------------------------------------|-------------------------------------------|---------------|------------------------|
| Kanellopoulos[10]  | 42/46                                         | 7/15 versus 3/30                          | Change in UDVA (preoperative to final visit) Accelerated CXL: 20/60-20/38 Standard CXL: 20/62-20/40 | Preoperative/postoperative Accelerated: 49.5/46.1 Standard: 48.7/45.8 |
| Elbaz et al.[11]   | 16/12                                         | 9/10                                      | Gain in UDVA of 0.13 LogMAR (P=0.012) | Nonstatistically significant difference |
| Cinar et al.[12]   | 26/6                                          | 9/10 Versus 3/30                          | Change in UDVA (LogMAR) Accelerated: 0.25±0.39 Standard: 0.15±0.33 P=0.408 | Change in keratometry Accelerated: 0.65±0.17 Standard: 0.45±0.45 P=0.762 |
| Kymionis et al.[13]| 10/3                                          | 9/10                                      | No significant improvement in CDVA | |
| Ulusoy et al.[14]  | 28/12                                         | 9/10                                      | +0.22 LogMAR improvement in BDVA (P=0.005) | Mean keratometric value dropped by at least 1D or remained stable |
| Shetty et al.[15]  | 30/24                                         | 9/10                                      | Preoperative/postoperative UDVA 0.76±0.26/0.61±0.25 (P=0.005) | 2.04D and 2.07 D flattening in mean K1 and K2 (P=0.001) |
| Hashemi et al.[16] | 62/6                                          | 18/5 versus 3/30                          | Preoperative/postoperative UDVA (LogMAR) Accelerated: 0.72±0.53/0.61±0.49 Conventional: 0.74±0.50/0.72±0.51 (P=0.733) | Preoperative/postoperative Accelerated: 46.39/46.44 Conventional: 47.12/46.87 (P=0.974) |
| Hashemi et al.[17] | 31/18                                         | 18/5 versus 3/30                          | Preoperative/postoperative UDVA (LogMAR) Accelerated: 0.72±0.53/0.72±0.53 Conventional: 0.74±0.50/0.72±0.51 (P=0.745) | Preoperative/postoperative Accelerated: 46.39±3.28/46.18±3.43 Conventional: 47.10±2.84/46.87 (P=0.004) |
| Chow et al.[18]    | 38/12                                         | 18/5 versus 3/30                          | Change in UDVA (LogMAR) Accelerated: 0.20±0.06 Standard: 0.28±0.08 P=0.508 | Change in K maximum Accelerated: 0.47±0.82 Standard: 0.16±0.72 P=0.343 |
| Hashemian et al.[19]| 153/15                                       | 30/3 versus 3/30                          | Change in UDVA (LogMAR) Accelerated: 0.19±0.20 Standard: 0.21±0.13 P=0.64 | Change in K maximum Accelerated: −1.85±0.99 Standard: −1.98±0.93 P=0.36 |
| Sherif[20]         | 25/12                                         | 3/30 versus 30/3                          | BDVA (LogMAR) preoperative/postoperative Accelerated: 0.48±0.17/0.61±0.15 Standard: 0.49±0.19/0.64±0.16 P=0.015 | K maximum preoperative/postoperative Accelerated: 49.43±1.63/48.2±1.43 Conventional: 51.40±1.69/50.24±2.0 |
| Tomita et al.[21]  | 48/12                                         | 3/30 versus 30/3                          | UDVA (LogMAR) preoperative/postoperative Accelerated: 1.08/0.94 Standard: 0.96/0.85 | Mean K preoperative/postoperative Accelerated: 45.51/44.98 Standard: 44.86/43.99 |
| Mita et al.[22]    | 39/6                                          | 30/3                                      | UDVA (LogMAR) preoperative/postoperative 0.11±0.42/0.89±0.53 (P<0.1) | Mean K preoperative/postoperative 49.95±6.11/49.19±5.82 (P<0.01) |

*UDVA/BDVA: Uncorrected distance visual acuity/best corrected distant visual acuity, LogMAR: Logarithm of the minimum angle of resolution, CXL: Corneal cross-linking, CDVA: Corrected distance visual acuity

Corneal stability following a 10-min irradiation at 9 mW/cm² was demonstrated in multiple studies with a significant improvement in visual acuity and keratometric values.[11-13] Similar results were noted in cases of pediatric keratoconus.[14]
Shetty et al. described a posttreatment disease progression in three eyes over a 2-year follow-up.[39]

Irradiance with 18 mW/cm² for 5 min demonstrated efficacy in halting disease progression; however, the topographic flattening was lower as compared to conventional CXL.[16-18]

Similar results were noted with the 3 min-30 mW/cm² protocol.[19-21] Mita et al. reported loss of one line of UDVA and CDVA in two eyes each. The exact cause of visual loss was not explained.[22] Ozgurhan et al. demonstrated significant improvements in visual acuity, keratometry, and aberrations in pediatric patients.[23]

Depth of demarcation line
The demarcation line represents the transition zone between the anterior cross-linked stroma and the posterior untreated stroma. It can be appreciated at a depth of around 300 µm, as early as 2 weeks postoperatively.[24] Studies, comparing the depth of demarcation line on anterior segment optical coherence tomography (AS-OCT) in conventional and accelerated approaches while delivering the same total fluence, demonstrated greater depth in the conventional group.[25,26] As the extent of demarcation line is considered a surrogate marker for the depth of treatment, superior results of the standard protocol in comparison to the accelerated approach may be assumed. With a greater total fluence of 7.5 J/cm² and increased riboflavin presoak, similar depths as conventional CXL were attained.[27-29]

While comparing the results of different accelerated protocols, a deeper and well-defined demarcation line was noted in the 3 mW/cm² and 9 mW/cm² group and was patchy and shallow with higher irradiances.[30] Since the anterior stromal fibers attribute to a majority of the biomechanical strength, a shallower demarcation line may still prove adequate to prevent disease progression. Long-term studies are required to further validate the results.

In conclusion, accelerated cross-linking significantly shortens the procedural time and reduces the patient discomfort. Moreover, as the UV-A irradiation time is significantly lower with the accelerated protocols, it avoids excessive stromal thinning and subsequent endothelial damage intraoperative.[31]

Pulsed Cross-linking
Shetty et al. compared the results following conventional cross-linking and accelerated protocols.[32] They demonstrated greater refractive and keratometric efficacy in the 3 mW/cm² and 9 mW/cm² groups as compared to the higher irradiation protocols. Moreover, a deeper and more well-defined demarcation line was observed.

Sufficient penetration of riboflavin, UV-A irradiation, and presence of oxygen are required for an effective stromal cross-linking.[2] The physiochemical basis of cross-linking lies in the photodynamic Type I and II reactions. The latter mediates cross-link formation via reactive oxygen species. It has been hypothesized that a more rapid oxygen depletion with accelerated protocols leads to a reduced efficacy.[32] Pulsed delivery of UV-A irradiation with a predetermined on and off pattern would enable better diffusion of oxygen into the stroma and a subsequent greater effect. Numerous studies compared the results of continuous versus pulsed irradiance in the accelerated protocols. A deeper demarcation line and higher apoptotic effect were noted with the pulsed approach.[33,34] Peyman et al. demonstrated a significantly deeper demarcation line following the pulsed approach (1 s on 1 s off) as compared to the 4 min of highly accelerated continuous UV-A irradiation.[35] Similar results were demonstrated by Moramarco et al. comparing the results in sixty eyes. The mean depth of demarcation line on AS-OCT in the pulsed group was 213 ± 47.38 µm as against 149.32 ± 36.03 µm in the group receiving continuous irradiation.[36] Mazzotta et al. confirmed similar findings on confocal microscopy.[37]

Although pulsed cross-linking shows promising results, the exact duration of pulsing is a question that remains unanswered. The rate of oxygen depletion in the Type II reaction is 15–20 s.[32] On the other hand, the normal tissue levels of oxygen are restored within 3–4 min of UV-A cessation. Therefore, further studies are required to determine the ideal pulsing approach.

**Laser In situ Keratomileusis Xtra**

**Laser vision correction with concomitant cross-linking**

Laser in situ keratomileusis (LASIK) is one of the more commonly performed ocular surgeries in the world. It has over the past few decades provided rapid visual recovery with improvement in quality of life and adequate patient satisfaction.[37] Despite the excellent immediate postoperative results, the long-term visual outcomes, especially in high refractive errors, have been less satisfactory due to regression of the refractive effect leading to high rate of enhancement procedures.[38] This is in part due to the corneal weakening or reduced biomechanical strength following flap creation and stromal ablation. In rare situations, corneal weakening leads to ectasia with subsequent degradation of vision.[39]

Over the past decade, corneal collagen cross-linking has become a mainstay of treatment for arresting the progression of keratoconus. CXL has shown to enhance corneal stiffening in both animal studies and clinical practice.[40,41] It does appear intuitive to combine the two procedures wherein LASIK improves quality of vision but reduces biomechanical strength of the cornea which could be partially or completely compensated by the CXL induced strengthening.

However, there are some concerns about combining the two techniques. Collagen cross-linking in addition to increasing corneal rigidity also induces a flattening effect.[42] This could limit the refractive accuracy by inducing long-term flattening and resultant overcorrection or hyperopic shift. In addition, the development of post-CXL stromal haze can further deteriorate visual quality. The advent of accelerated protocols has helped reduce treatment time by delivering higher irradiance in shorter duration. Numerous studies demonstrate the efficacy of accelerated cross-linking in stabilizing the cornea with a better safety profile as compared to conventional treatment.[30,32]

**Laser in situ keratomileusis Xtra procedure**

The LASIK XTRA procedure entails the administration of half fluence high irradiance cross-linking subsequent to refractive correction. A higher concentration 0.25% riboflavin is applied on the stromal bed subsequent to excimer laser ablation with a soak time of 90 s [Fig. 1]. The interface is washed thoroughly and the flap is repositioned. UV-A irradiance is delivered as
a homogenous beam of 30 mW/cm² for 90 s to deliver a total fluence of 2.7 J/cm². This is precisely half the energy delivered during conventional cross-linking in the Dresden protocol. The goal of cross-linking in this accelerated half fluence format is to restore or improve corneal strength without inducing a refractive change. The riboflavin is instilled on the stromal bed and not through the flap as it obviates the need for deepithelization to promote riboflavin diffusion. The flap itself does not contribute to the residual biomechanical strength, and therefore, cross-linking the flap would provide no advantage. In addition, cross-linking the flap may lead to subsequent shrinkage with undesirable consequences. AS-OCT is a useful tool to demonstrate thickness of flap and depth of demarcation line [Fig. 2].

Clinical outcomes
Over the last few years, numerous studies have shown promising results with LASIK Xtra.[43]

Majority of the studies demonstrated greater stability of refraction. The incidence of post-LASIK regression is high in hyperopic eyes. Contralateral study comparing the results of LASIK with or without concomitant cross-linking in hyperopic eyes demonstrated a significantly lower regression in the LASIK Xtra group.[44] Encouraging results have been shown in the treatment of high myopia as well. Tan et al. compared the results of LASIK Xtra for high myopic correction (−8.0 D−19.0 D) with a spherical equivalent matched historical cohort.[45] A greater refractive accuracy of the Xtra group was noted at 3 months with 98% of the eyes attaining a UDVA of 20/25 or better against 61% of the LASIK group. A longitudinal observational study of 140 eyes with a 2-year follow-up showed lower refractive shift and greater keratometric stability in the Xtra group.[46]

The application of CXL along with refractive correction also had an influence on epithelial remodeling post-LASIK, especially while treating higher degrees of myopia. A study showed a significantly lower increase in midperipheral thickness when LASIK was combined with CXL (3.79 µ) as compared to LASIK alone (9.32 µ).[47] A greater refractive stability was noted with no progressive flattening. The procedure demonstrated a good safety profile with stable endothelial cell count and no visually significant haze development.[48] In a large study of 601 eyes, a stable uncorrected visual acuity with no significant changes in spherical equivalent or keratometry was noted at 1-year follow-up.[49]

In summary, LASIK with concomitant half fluence cross-linking is a promising treatment modality with significant improvement in refractive stability and possible reduced incidence of post-LASIK regression.

Cross-linking in Thin Corneas

Standard Dresden protocol mandates a minimal corneal thickness of 400 µ following epithelial debridement. This would limit the UV irradiance at the endothelial level to 0.18 mW/cm², which is at least a factor of 2 smaller than the damage threshold level. Unfortunately, a number of patients have thin corneas often below the threshold of safety guidelines, making the disease not amenable to traditional cross-linking.[50]

Various techniques have been described to overcome these limitations. Hyposmolar cross-linking describes the instillation of hypotonic riboflavin to increase the corneal hydration and thickness intraoperatively.[51] Increased intraoperative time and a relatively lower concentration of collagen in the hydrated stoma are some of the limitations of this technique. Transepithelial cross-linking was introduced to prevent the adverse effects associated with epithelial debridement as well as a possible role in thinner corneas.[52] However, long-term studies demonstrated unsatisfactory results perhaps due to limited penetration of riboflavin.[53] Customized pachymetry-guided epithelial debridement entails preservation of the epithelium over the thinnest cone or area of maximal topographical steepening.[54] However, this technique demonstrated limited penetration with a demarcation line at 150 µ.[55] Jacob et al. described the use of a riboflavin-soaked bandage contact lens to augment the corneal thickness by roughly 100 µ.[56] However, the absorption properties of contact lenses differ from that of the corneal stroma. Moreover, inability to customize the contact lens thickness and intraoperative buckling were some of the associated limitations. Long-term results of this procedure are not available. Sachdev et al. described the intraoperative augmentation of stromal thickness, using refractive lenticules obtained from patients undergoing small incision lenticule extraction for myopic correction.[57] In this modified technique, the thickness of the corneal stroma is increased in the most physiological manner. Placement of the central lenticule over the apex of the cone enables one to augment the corneal thickness where required while sparing the remaining stroma to be cross-linked normally. Moreover, the relatively rough host stromal surface allows the lenticule to spread easily and buckling is avoided [Fig. 3].

In addition, the use of hydroxypropyl methylcellulose riboflavin prevents corneal dehydration induced by dextran and is more suitable for thin corneas.[58] The principle of customizing UV-A irradiance to the stromal thickness is evolving as the latest technique for thin corneas (adapted fluence). In conclusion, various advancements in procedures enable safe cross-linking in suboptimal corneal thickness.

Transepithelial Cross-linking

Conventional cross-linking entails epithelial debridement to achieve greater riboflavin penetration, which is otherwise impeded by epithelial tight junctions. Postoperative pain, increased risk of haze, and infection are associated limitations. Transepithelial application of riboflavin with numerous techniques to modify the epithelial permeability has been described including pharmacological cleavage of tight junctions and application via intrastromal pocket.[59-61] Although the transepithelial approach demonstrated fewer complications, efficacy was lower as compared to conventional treatment particularly in stabilizing or improving keratometry.[62]

Iontophoresis is a noninvasive technique that allows transepithelial riboflavin penetration following application of a mild electric current. Riboflavin is a negatively charged, water-soluble molecule with a relatively low molecular weight making it suitable for iontophoresis.[63,64] Numerous studies demonstrated stabilization of the disease process following iontophoresis-assisted CXL (I-CXL).[65] However, the keratometric regression was lower as compared to conventional epi-off cross-linking.[66] Similar results were noted in cases of pediatric keratoconus over a 15-month
follow-up period. The depth of the demarcation line noted was lower as compared to standard cross-linking. In a study by Mastropasqua et al., I-CXL demonstrated deeper saturation of riboflavin with respect to conventional epi-on but did not reach the concentrations with standard epi-off. Although I-CXL-assisted riboflavin delivery is lower than the conventional approach, the effective concentration to halt disease process is not yet established.

I-CXL has the potential to become a valid alternative treatment for keratoconus while reducing treatment time, postoperative patient discomfort, and risk of infection.

Further studies are needed to establish the mechanism and efficacy of this relatively new treatment modality.

**Customized Corneal Collagen Cross-linking for Keratoconus: Improved Corneal Regularization to Maximize Visual Rehabilitation**

Keratoconus is a disease clinically characterized by increased corneal curvature, reduced corneal thickness, and progressive topographic irregularity. However, it has been proposed that the biomechanical modification is focal in nature, rather than a uniform generalized weakening. Roy and Dupps demonstrated differential biomechanical weakening in the area of the cone using three-dimensional finite element analysis model. They additionally concluded greater efficacy of smaller diameter cone-centric treatments for the reduction of corneal curvature and higher order aberrations.

Clinical application of this principle would require UV-A irradiation in customized treatment patterns localized on specific corneal zones. The Mosaic delivery system (KXL II, Avedro Inc., Waltham, MA, USA) uses an advanced pupillary tracking mechanism to offer customized cross-linking (photorefractive intrastromal cross-linking). The Mosaic device has been afforded the CE mark in Europe and Health Canada Approval. Kanellopoulos et al. first described a case of customized cross-linking to achieve refractive results in progressive keratoconus. UV-A irradiation was applied in a customized toric pattern using the transepithelial approach. A mean astigmatic reduction of 0.8D with a subsequent improvement in uncorrected visual acuity from 20/40 to 20/25 was noted at 6 months.

Various other studies have been published since demonstrating the results of the customized approach. The use of varying treatment patterns including customized toric, asymmetric arcuates, and concentric circles has been described. An important aspect is to determine the localization of the irradiation patterns. In the previous studies, the treatment was centered on the area of greatest curvature. However, centration of irradiation around the maximum point of posterior float elevation may be more intuitive, since pachymetry as well as curvature is modulated by epithelial thickness and tear film.

Table 2 summarizes the results of customized CXL.

Customized CXL may offer several advantages over the conventional approach. A greater corneal surface normalization leads to superior visual results. The propensity for haze formation was similar to conventional treatment despite the application of higher fluence. Fig. 4 demonstrates a Gaussian distribution of demarcation line on AS-OCT.

Moreover, a paracentral treatment with subsequent haze will have a significantly lower impact than if the haze was central following conventional treatment. A reduced treatment zone with a smaller area of epithelial debridement would reduce postoperative discomfort and risk of infection.

In conclusion, customized CXL is a promising treatment modality. Studies are required to further customize the treatment approach and optimize outcome.

Figure 1: Instillation of 0.25% riboflavin (VibeX Xtra, Avedro, Inc.) on the stromal bed following excimer laser ablation

Figure 2: Anterior segment optical coherence tomography demonstrating a well-apposed flap and the demarcation line in a case of femtosecond LASIK with half-fluence accelerated cross-linking

Figure 3: Anterior segment optical coherence tomography after placement of stromal lenticule and initial riboflavin soakage
Photorefractive Intrastromal Cross-linking: High-Fluence Corneal Collagen Cross-linking for Low Myopia

Customized cross-linking for refractive correction is an emerging concept. It offers a nonincisional, nonablative treatment approach involving high-fluence irradiation to induce subsequent flattening and refractive correction. The UVA irradiation is delivered by the Mosaic device (Avedro Inc., Waltham, MA, USA) which uses advanced pupil tracking technology to deliver a customized treatment. The Mosaic device has been afforded the CE mark in Europe and the Health Canada Approval.

| Author (year of publication) | Number of eyes | Mean follow-up | Treatment parameters | Clinical finding |
|-----------------------------|----------------|----------------|----------------------|------------------|
| Kanellopoulos (2014)⁷⁰ | Single eye | 6 months | Customized transepithelial toric fluence ranging from 4 J/cm² to 14 J/cm² | Improvement in UDVA from 20/40 to 20/25 Mean astigmatic reduction of 0.8D |
| Nordstrom et al. (2016)⁷¹ | 25 eyes (conventional cross-linking) 25 eyes (PiXL) | 12 months | Asymmetrical arcuate shaped fluence ranging from 7.2 J/cm² to 15 J/cm² | Greater corneal regularization with a mean reduction in maximum keratometry of-1.74D in the PiXL group Improvement in UDVA and BSCVA from 3-month follow-up Asymmetric pattern of densitometry in PiXL group, with a greater increase associated with higher fluence No corneal endothelial cell loss in either group |
| Seiler et al., (2016)⁷² | 20 eyes (conventional cross-linking) 20 eyes (PiXL) | 12 months | Concentric circles with a fluence ranging from 5.4 J/cm² to 10 J/cm² | Two eyes in the standard group versus 7 eyes in the customized group showed a flattening of >2D The regularization index was significantly higher in the customized group (5.2D) versus control group (4.1D). $P<0.03$ No significant variation in maximal depth of demarcation line |
| Mazzotta et al. (2016)⁷³ | 21 eyes (PiXL) | 12 months | Arc patterns for peripheral cones and circular patterns for central cones, with fluence ranging from 7.2 J/cm² to 15 J/cm² | No significant improvement in UDVA or CDVA A significant decrease in topographic astigmatism of 1.41D at 1-year follow-up Significant reduction in coma from baseline An improved corneal symmetry with flattening of steeper areas and compensatory steepening of flat areas was noted No corneal endothelial cell loss Significant decrease in keratometry of 1.07D in PiXL group at 6 months Corneal normalization analysis demonstrated superior results in the PiXL group Deeper demarcation line in cone area than surrounding area in customized group Less stromal damage and faster healing on confocal microscopy |
| Cassagne et al. (2017)⁷⁴ | 30 eyes (conventional cross-linking) 30 eyes (PiXL) | 12 months | Customized patterns with fluence ranging 5.4 J/cm²–15 J/cm² | |
Kanellopoulos et al. described the preliminary results of transepithelial cross-linking for low myopic correction.[59] High-fluence UV-A irradiation of 12 J/cm² was delivered as a customized central treatment using the KXL II system (Avedro, Waltham, MA, USA). Refractive and keratometric changes were demonstrated over a 6-month follow-up period. An average corneal flattening of 2.3 D at 1 week, with subsequent regression and stabilization to 1.4 D was noted at 1 month. No significant endothelial cell loss was noted despite the application of higher fluence.

Lim et al. demonstrated the results in a cohort of 14 eyes with a 1-year follow-up period.[60] High-fluence UV-A irradiation ranging from 10 to 15 J/cm² was delivered over a 4.5 mm central zone. At 12 months' postprocedure, a mean reduction of 0.72 ± 0.43D was noted in the mean residual spherical error (P < 0.001). No significant regression was noted over the 1-year follow-up. Transient corneal haze subsided gradually, with no loss in best-corrected visual acuity.

PiXL for hyperopia has also been reported by Kanellopoulos and Asimellis, with a mean hyperopic correction of +0.85 D using an epithelial-on approach.[61]

Further studies are underway to determine the results of this potentially revolutionary refractive procedure. Possible advances include the development of a nomogram for astigmatic correction, addition of supplemental oxygen to enhance the efficacy of epithelium-on approach, and topographically guided treatment patterns.

Future Directions

Adapted fluence: Crosslinking in thin corneas

The standard Dresden protocol for cross-linking recommends a minimal postdebridement thickness of 400 µ to prevent irradiation damage to the corneal endothelium.[62]

Lack of sufficient thickness for UVA absorption and attenuation is a limiting factor in thin corneas. A number of techniques with varying degrees of efficacy have been described for cross-linking the thin corneas. The three main factors than can be altered in the cross-linking process include stromal thickness, riboflavin concentration, and UV-A irradiation. However, the techniques described thus far employ the principle of increasing corneal thickness through tissue augmentation or stromal hydration.[53,56,78,79]

The UV-A fluence delivered in all techniques was a constant 5.4 J/cm². Hafezi and Kling describe the principle of adapted fluence wherein customized energy is delivered by altering the irradiation time (UVA irradiation of 3 mW/cm² for a customized irradiation time).[80] The process can be tailored based on the corneal thickness to deliver a cross-linking effect with an adequate safety zone. This eliminates the need for varying riboflavin concentrations and overcomes the limitations associated with earlier described techniques. However, the concept of adapted fluence requires validation through clinical trials before widespread use.

Scleral cross-linking for axial myopia

Scleral thinning and subsequent weakening results in axial length elongation and progressive myopia. Scleral cross-linking (SXL) using photosensitizer and blue light to mechanically reinforce the sclera may prevent progression in such cases.

Kwok et al. described the application of flexible optical waveguides around the equatorial region to induce SXL in porcine eyes.[81] They demonstrated a significant increase in Young’s modulus with similar results in proximally and distally treated halves. The elastomer material with the linearly tapering design allowed a more homogenous light delivery and reduced thermal injury in comparison to earlier techniques.[82,84]

In vivo studies are needed to demonstrate the safety and efficacy of this method.

Photoactivated chromophore for infectious keratitis – corneal collagen crosslinking (PACK-CXL)

In addition to corneal strength augmentation, the cross-linking procedure demonstrates significant cytotoxic effect against living cells and microorganisms. UV-A irradiation has a known antimicrobial effect against bacteria, viruses, and microbes. In addition, riboflavin when photoactivated produces reactive oxygen species with subsequent microbicidal effect.[85,86] The combination of both produces cytotoxic effect which is drastically greater than the simple additive action.[87,88]

The application of CXL for keratitis was demonstrated at first for noninfectious and infectious corneal melts resistant to therapy.[89,90] The indications were subsequently extended to infectious keratitis as the first line of management in 2011.[91] Various studies demonstrated promising results for infectious keratitis excluding herpetic etiology. In addition, the results in bacterial and Acanthamoeba were superior to fungal keratitis.[92]

Further studies are required to compare the efficacy of this novel approach with traditional antimicrobial therapy in the treatment of corneal infections.

Conclusion

Cross-linking is still an evolving technology whose full potential is yet to be realized. The newer accelerated protocols and combination treatments have opened up a multitude of avenues with far reaching implications on the way we approach ectasia and refractive surgery. A greater degree of customization of treatments is likely the path forward which will enable us to achieve better refractive outcomes while maintaining a high level of safety.
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There are no conflicts of interest.

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