ABSTRACT

Corneal cross-linking has proven safety and efficacy in arresting the progression of keratoconus. Its use has been extended to resolution of corneal edema. We present a case report and review of literature on the treatment of corneal edema using corneal cross-linking. Corneal cross-linking seems to be an effective method in reducing corneal edema at least as a temporary measure till definitive solution, such as penetrating keratoplasty (PKP) or Descemet's stripping automated endothelial keratoplasty (DASEK) is performed.

Keywords: Cornea, Corneal edema, Collagen Corneal cross-linking, Collagen, Topography, Corneal dystrophy, Orbscan, Keratoconus, Corneal thickness, Glaucoma, Intraocular Pressure.

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INTRODUCTION

Collagen corneal cross-linking (CXL) is gaining popularity as a treatment for arresting the progression of keratoconus since the first clinical result publication in 2003 by Theo Seiler’s group. CXL increases the corneal biomechanical strength by increasing the interfibrillar covalent bonds. It increases collagen fibrils thickness, decreases the potential space for water accumulation and decreases swelling rate.

Wollensak et al reported on the hydration behavior of the cross-linked cornea. He demonstrated increased resistance of the anterior cross-linked cornea to the diffusion of water from the posterior to the anterior cornea. Recently, there have been various reports on the use of CXL as a treatment for bullous keratopathy (BK) as a result of Fuch’s dystrophy, glaucoma, pseudophakic BK and following penetrating keratoplasty. CXL reduces corneal edema, corneal thickness and epithelial bullae consequently improving symptoms, such as pain and improving visual acuity (VA) and photophobia.

LITERATURE REVIEW

There are various ways of performing CXL for BK.

1. The traditional way reported by Seiler’s group; the epithelium is removed from the central 8 to 9 mm of the cornea. Riboflavin 0.1% with dextran 20% is instilled until the appearance of yellow flare in the anterior chamber (AC). The cornea is irradiated with Ultraviolet A (UVA), (365-370 nm) light source 5 cm from the eye with a light intensity of 3 mW/cm² for 30 minutes. During treatment riboflavin and balanced salt solution (BSS) are instilled into the eye.

2. Glucose 40% is instilled to the eye 1 day prior to the procedure in order to reduce corneal thickness followed by the same procedure as in 1.

3. Glycerin 70% in instilled to the eye till achievement of corneal thickness of 340 to 370 µ prior to the start of the procedure as in 1.

4. Two pockets are created with femtosecond laser. The epithelium is not removed and riboflavin 0.2 ml of 0.1% are injected in the pockets and then the cornea is exposed to UVA (365-370 nm) light source 5 mm from the eye with a light intensity of 15 mW/cm² for 7 minutes. The advantage of this method is that it painless as the epithelium is not removed.

Ehlers et al reported on decreased corneal edema in 10 out of 11 eyes treated with CXL. VA improved in most eyes following CXL. In one eye VA improved from 0.04 to 0.4 on decimal notation after 6 months and this improvement was maintained at the 17 months follow-up (FU). Ghanem et al reported on 14 eyes with pseudophakic bullous keratopathy (PBK) in which the mean corneal thickness decreased following CXL from 747 to 623 µ at 1 month FU. Six months after treatment, mean corneal thickness increased to 710 µ but remained less than the preoperative thickness. No changes in pain score were reported in this study.

Kozobolis et al reported on 2 eyes with PBK with FU of 2 months with decreased corneal edema and improvement in VA following CXL. Wollensak et al reported on 3 eyes with corneal edema as a result of BK with a mean decrease in corneal thickness of 94 ± 14 µ after 18 months. In cases where the cornea was not cicatrized there was improvement in VA. Krueger et al reported on 1 eye with corneal edema with a preoperative corneal thickness of 675 µ and VA of counting fingers (CF). Following CXL treatment corneal thickness was reduced to 550 µ and VA improved to 6/24, 6 months following treatment.

Barbosa et al reported on reduction in corneal edema and improvement in symptoms in 11 of 25 treated eyes (44%) after 6 months. The reduction in corneal edema was significant at 3 months following treatment. In 56% of eyes following CXL, the epithelium was not removed and riboflavin was injected directly into the corneal stroma.
there was recurrence of bullae after 6 months. Gadelha et al\textsuperscript{11} reported on two eyes with PBK in which pain was reduced following treatment. However, there was no significant reduction in corneal thickness or improvement in VA. Hafezi et al\textsuperscript{12} reported on CXL treatment for fluctuations in VA as a result of Fuch’s endothelial dystrophy (FED). The authors report improvement in VA and VA fluctuations, and decrease in corneal edema in a FU of 3 months.

**CASE REPORT**

A 28-year-old male was presented to our medical center in February 2010. After reviewing his medical records we revealed that he was first presented to his ophthalmologist in October 2008 with uncorrected visual acuity (UCVA) of 6/6 in his right eye and best spectacle-corrected visual acuity (BSCVA) of 6/7 with refractive correction of –1.25 to 0.50×165. In December 2009, he had a further visit to his ophthalmologist. He was noted to have UCVA of 6/6 in his right eye and BSCVA 6/9 with refractive correction of – 0.75 to 2.00×165. Slit-lamp examination was unremarkable at that stage but he was suspected to have keratoconus. Two weeks later, he presented again with further deterioration in his subjective VA. On examination, his UCVA was 6/30 and BSCVA of 6/15. On slit-lamp examination, he was suspected to have corneal edema. Endothelial cell count (ECC) was 3,106 in his right eye and 2,538 in his left eye with an edematous cornea and a normal appearing cell morphology. He was diagnosed with keratoconus and diffuse corneal edema (not hydrops). He was treated with PredForte 1% (prednisolone acetate) and Cosopt (Dorzolamide Hydrochloride-Timolol Maleate) and because of further decrease in VA and his unresponsiveness to medical treatment he was listed for PKP.

In February 2010, he presented to our medical center with UCVA of CF from 1.5 meters and BSCVA of 6/60-2 with correction of –6.00 × 165. Slit-lamp examination of his left eye revealed quite eye with normal appearing conjunctivae, diffuse corneal edema with no Descemet rupture, Descemet membrane opacities or keratic precipitates (KP). He had a normal anterior chamber (AC), clear lens and normal fundus examination including optic disk. His right eye slit-lamp examination was unremarkable with a normal fundus. Central corneal thickness (CCT) was 658 µ in his left eye and 572 µ in his right eye. Corneal topography of the left eye was suggestive of keratoconus with a normal topography of the right eye (Figs 1 to 3). He had normal ocular response analyzer (ORA) signals in the right eye and abnormal in the left eye with an IOPcc of 11.1 and 20.6 mm Hg respectively (Fig. 4) Xalacom (Latanoprost 0.05% + Timolol 0.5%) and PredForte 1% (prednisone) were prescribed till the next FU a week later. No improvement was noted in the subsequent FU a week later. We discussed the situation with the patient and made him aware that PKP can always be done at any stage, but we suggested CXL.

At that stage he underwent CXL to his left eye with view of reducing his corneal edema. We used the surgical technique recommended by Theo Seiler and his group. Surgical procedure was performed by one surgeon under topical anesthesia. Standard prepping and draping were performed. The eye was washed with poldine 4%. A 9 mm radius central corneal epithelium was removed. The patient was seated and riboflavin 0.1% combined with dextran 20% was instilled every 5 minutes for 30 minutes. The exposure of the cornea to UVA was done only after the appearance of a strong yellow flare in AC. Speculum was inserted and...
(CL) was applied until full epithelialization. Vigamox (moxifloxacin 0.5%) antibiotics were prescribed until full epithelialization which occurred after 3 days. A PredForte 6 time daily for 2 weeks was used and was tapered gradually. He was asked to continue his Xalacom. Valtrix (valacyclovir) 1 gm 3 times daily for 1 week was prescribed.

Five days postoperatively (PO) his BSCVA was CF at 1 meter with a CCT of 682 μ. Corneal topography with computerized corneal topography KR7000P (Topcon) was not recordable.

Three weeks PO his UCVA was 6/60 and BSCVA of 6/18+ with –1.00 to 2.50×165 correction. CCT of 572 μ in the right eye and 590 μ in the left eye with an IOPcc of 10.5 and 18.8 mm Hg respectively. Five weeks PO his CCT was 577 μ in the left eye and a bullae appeared in the inferiortemporal cornea with a CCT of 751 μ over the bullae. Three months PO his UCVA was 6/48 and BSCVA of 6/15– with –2.5×160 correction and CCT of 568 μ in the left eye.

At 4.5 months UCVA was 6/60 and BSCVA was 6/12– with +0.25 to 2.75×160 correction. He was noted to have haze +1 (Fig. 5). His corneal topography of K minimum of 43.37, K maximum of 47.12 and K average of 45.12 (Fig. 6).

The cornea and the UCVA, BSCVA, CCT, IOP and refraction were stable and the treatment (eyedrops) were discontinued.

Here is his last FU in a November 2011: BSCVA 6/9 partial, refraction plano = –2.5×160 and IOPCC 12.9 K minimum 43.62, K maximum 47.87, K average 45.62, CCT left eye 590 μ, right eye 564 μ (Figs 9 and 10).

Fig. 3: Ocular response analyzer (ORA). Upper: Demonstrating regular pattern with corneal resistance factor (CRF) and corneal hysteresis (CH) within normal limits in the right eye. Lower: Demonstrating irregular pattern with corneal resistance factor and corneal hysteresis lower than normal but as expected in keratoconic eye and edematous cornea.

Fig. 4: Normal orbscan of the right eye.

Fig. 5: Postoperative corneal haze.
DISCUSSION

Among the methods of CXL for treating corneal edema mentioned in the literature review. The method described by Hafezi et al seems to us the most suitable for treating corneal edema as the treatment is supposed to cover the major part of the stroma and not only the anterior part as in the classical methods. As we know, CXL affects the first 300 µ of the cornea and in an edematous cornea the treated stoma may constitute a negligible part of the anterior ‘real anterior stoma’ because of the huge edema. We performed the treatment according to the classical method because we were not aware of the method described by Hafezi et al which was published in May 2010 and our treatment was
performed in February 2010. We believe that the method may have an impact on efficacy of the treatment.

Another factor which may affect the efficacy of the treatment is the cause of the BK. The etiologies reported in the literature for corneal edema comprise more than one pathology (PBK, FED, absolute glaucoma and decompensated corneal graft) although the end result was corneal decompensation. Most cases suffered from PBK. CXL treatment aims at treating the end result of the pathologic process and not the cause. The surgical techniques were different although most reported cases were treated in the traditional way. Does the cause have any impact on the surgical outcome? Do causes, such as absolute glaucoma (in which the IOP is continuing to decompensate the endothelium) or PBK (in which the cause of the decompensation is trauma to the endothelium) have different outcomes?

Moreover, reports that show improvement in corneal edema at the last FU are short-term FU and others reports vanishing of the treatment effect during the FU. Hence, one may deduce that the CXL treatment is effective only as a short-term treatment till a PKP or DASEK are performed.

We present a retrospective case of a young male suffering from rapid deterioration of vision as a result of new onset keratoconus followed by a severe diffuse corneal edema (not hydrops), with no tearing of Descemet membrane and no drastic decrease of ECC. He is an adopted son and we could not detect his family for evaluation of their corneas. His vision was finger counting only, no local or general cause of the edema was found, medical treatment failed and he was candidate for PKP.

Differential diagnosis in our case is:
1. Iridocorneal endothelial syndrome (ICE) but we are missing iris and pupillary changes as well as glaucoma.
2. Endothelitis which is associated with cytomegalovirus, Herpes and EBV. Missing KP's and iritis. No active viral infection could be detected by blood tests.
3. Posterior polymorphous endothelial dystrophy, however, he is missing Descemet membrane changes.

CXL was offered for two purposes:
1. Arresting the progression of KC.
2. Reducing corneal edema in the hope that his endothelial dysfunction is temporary. In case that the edema recurs inspite of CXL he can undergo DASEK and not PKP because his KC is not a severe and CXL is supposed to arrest it in this nonadvanced stage.

CONCLUSION
CXL is effective method in reducing corneal edema and its symptoms and improving VA in patients suffering from endothelial decompensation. No long-term results are reported. In our case the FU is 20 months. Further studies are needed to evaluate the long-term efficacy. In the meantime, this treatment can be offered as a temporary treatment till permanent solution is offered.

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