MAILBOX

Modified corneal collagen crosslinking reduces corneal oedema and diurnal visual fluctuations in Fuchs dystrophy

Crosslinking of corneal collagen with riboflavin and ultraviolet-A irradiation (CXL) induces crosslinks within and between collagen fibres. CXL increases corneal biomechanical and biochemical stability, and is currently used clinically to treat keratoconus. CXL also significantly reduces the stromal swelling capacity. We investigated whether a modified CXL treatment protocol would be beneficial in early Fuchs dystrophy with various degrees of corneal oedema and diurnal variations in visual acuity.

MATERIALS AND METHODS

CXL was performed as published previously, with the following modification: in cases where the stroma was thicker than 450 μm after abrasion and 30 min of instillation of isoosmolar riboflavin solution, glycerol 70% solution was applied every 5 s for 2 min, and the central corneal thickness (CCT) was measured using ultrasound pachymetry (Tomey GmbH, Erlangen, Germany). Glycerol 70% solution was administered repeatedly until the target corneal thickness of 370–430 μm was reached. During irradiation (UV-X, Peschke Meditrade, Cham, Switzerland), CCT was monitored by ultrasound pachymetry every 5 min, and glycerol 70% solution was applied, if necessary.

RESULTS

Three eyes in two patients were treated using this modified CXL protocol. A 50-year-old woman with Fuchs dystrophy and a history of 3 years of diurnal visual fluctuations was referred to us in March 2008. The preoperative best spectacle-corrected visual acuity (BSCVA) was 20/50. We performed modified CXL in the left eye.

At 1 month after CXL, Scheimpflug analysis (Pentacam, Oculus Instruments, Wetzlar, Germany) of CCT showed a reduction of more than 100 μm, and the corneal thickness spatial profile (CTSP) and percentage of increase in thickness (PIT) showed a regularisation of the ‘flattening’ typical for Fuchs dystrophy (figure 1). Accordingly, diurnal analysis of corneal thickness showed a distinct postoperative reduction in CCT at all time points measured (figure 2). One month after CXL, the patient reported a reduction in diurnal visual fluctuations, and we measured an increase in BSCVA to 20/32. The patient showed stable topographical and visual acuity at the 3-month follow-up.

DISCUSSION

Fuchs endothelial dystrophy is characterised by corneal swelling and oedema. In early stages of the disease, the endothelial proton pump is compromised, leading to diurnal CCT variations and fluctuations in visual acuity. Since CXL reduces the corneal swelling capacity, we investigated whether CXL would be beneficial in early Fuchs dystrophy.

We modified the standard CXL treatment protocol by reducing corneal thickness prior to irradiation. Because the standard parameters only treat the anterior 270–330 μm, we used glycerol 70% solution intraoperatively to dehydrate and thin the cornea prior to and during irradiation.

We saw a distinct reduction in CCT, an improvement in the corneal thickness spatial profile (CTSP) and an increase in BSCVA 1 month after treatment which remained stable at the 3-month follow-up.

![Figure 1](image-url) Scheimpflug analysis of corneal thickness before (A) and 1 month (B) after crosslinking (CXL). Central corneal thickness (CCT) is reduced by more than 100 μm at 1 month after CXL (arrows). Before CXL, analysis of the corneal thickness spatial profile and percentage increase in thickness shows a ‘flattening’ typical for Fuchs dystrophy (stars, left side of panel). Postoperatively, this ‘flattening’ is regularised (stars, right side of panel).
Endothelial cell counts were not performed in this case. However, the cornea showed no clinical signs of decompensation at 3 months after treatment.

Similarly, Ehlers and Hoedtard investigated whether oedema related to endothelial decompensation would diminish after CXL. They reported a distinct reduction in CCT after CXL which took months to occur. Re-treatments became necessary, probably because Ehlers and Hoedtard crosslinked a swollen cornea, whereas in our approach, the cornea was dehydrated with glycerol 70% solution, thus thinned prior to treatment. Wollensak et al. recently reported the use of CXL in patients with bullous keratopathy. In contrast to our modification of the CXL treatment protocol, they used glucose 40%, and they preoperatively dehydrated the cornea for 1 day instead of 30 min. Whereas the dehydrating properties of glucose solution and glycerol solution are virtually identical, we cannot determine whether corneal dehydration at 1 day prior to CXL is more beneficial than our approach. However, we believe that it is more feasible clinically to perform dehyderation immediately before the CXL procedure.

Patients with early Fuchs dystrophy and disturbing diurnal visual fluctuations represent a novel application for CXL. Although CXL may not prevent the outcome of the dystrophy, it may increase the patients’ visual comfort until keratoplasty becomes necessary.

Farhad Hafezi, Peter Dejica, Francois Majo

1 IROC, Institute for Refractive and Ophthalmic Surgery, Zurich, Switzerland
2 Private Practice, Schaffhausen, Switzerland
3 Hospital Ophthalmique Jules Goin, Lausanne, Switzerland

Correspondence to Professor Dr. Farhad Hafezi, IROC, Institute for Refractive and Ophthalmic Surgery, Stockerstrasse 37, Zurich 8002, Switzerland; farhad@hafezi.ch

Competing interests None.

Ethics approval Ethics approval was provided by the cantonal ethics committee of the Canton of Zurich.

Patient consent Obtained.

Provenance and peer review Not commissioned, externally peer reviewed.

Accepted 6 July 2009

doi:10.1136/bjo.2009.162479