Abstract

Objective: To evaluate early in vivo corneal wound healing findings after Descemet's Stripping Automated Endothelial Keratoplasty (DSAEK) by using in vivo confocal microscopy. Method: A total of 15 eyes of 15 patients were included. In vivo confocal microscopy (Confoscan 4, Fortune Technologies, Italy) was performed from 4 to 7 weeks after DSAEK. Measurements were scanned from the corneal endothelium to the corneal surface with a Navis® software (NIDEK, Multi-Instrument Diagnostic System, Japan). Results: Donor-receptor interface was found in an average of 114 ± 12.4 microns. Corneal stromal folds were observed from 111.1 ± 3.5 microns from the endothelium to 286 ± 94 microns (mean 175 ± 90.5 microns of the corneal stroma). Keratocites were activated in the donor tissue from 12 ± 1.4 microns from the endothelium to 105 ± 38.2 microns (mean 93 ± 36.9 microns of the corneal stroma). Conclusions: Donor keratocites are activated up to 7 weeks after DSAEK. Several corneal folds are present in proximity to the donor-receptor interface after DSAEK. Further evaluation of these findings is justified to determine its clinical significance.


Resumen

Objetivo: Evaluar los hallazgos cicatriciales tempranos en la córnea de pacientes operados de DSAEK (Descemet’s Stripping Automated Endothelial Keratoplasty) mediante microscopía confocal in vivo. Método: Se incluyeron 15 ojos de 15 pacientes. Se realizó microscopía confocal (Confoscan 4, Fortune Technologies, Italy) entre las 4 y las 7 semanas después de la cirugía de DSAEK. Las mediciones se realizaron desde el endotelio al epitelio corneal utilizando el programa Navis® software (NIDEK, Multi-Instrument Diagnostic System, Japan). Resultados: La interface donador-receptor se encontró a 114 ± 12.4 micras en promedio. Se comenzaron a observar pliegues en el estroma a 111.1 ± 3.5 micras desde el endotelio corneal hasta 286 ± 94 micras (promedio de 175 ± 90.5 micras del estroma corneal). Se comenzó a observar activación de queratocitos en el tejido donador a 12 ± 1.4 micras desde el endotelio hasta 105 ± 38.2 micras (promedio de 93 ± 36.9 micras del estroma corneal). Conclusiones: Los queratocitos del tejido donador se encontraron activados hasta 7 semanas después de la cirugía de DSAEK. En la proximidad de la interface donador-receptor se encontraron grandes pliegues en el estroma después de cirugía de DSAEK. Se requieren futuras evaluaciones para determinar la significancia clínica de estos hallazgos.


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Introduction

Endothelial transplantation using the Descemet’s Stripping Automated Endothelial Keratoplasty (DSAEK) technique has demonstrated to be a good surgical alternative to solve corneal pathologies where the corneal endothelium is damaged but the corneal stroma is intact, with a donor graft being required for the resolution of the problem\(^\text{1,2}\). Furthermore, it has been reported to be an option in the treatment of patients who experience failure after penetrating keratoplasty\(^\text{3}\).

The purpose of the present study is to evaluate early corneal scarring findings in DSAEK-operated patients using confocal microscopy in vivo.

Method

Fifteen eyes from 15 patients underwent corneal endothelium transplantation using the DSAEK technique\(^\text{4}\). All endothelial grafts were obtained on request from the eye bank, each with a thickness of 100 microns. Preoperative diagnoses of the patients undergoing DSAEK surgery were nine bullous keratopathies secondary to pseudophakia and six Fuchs dystrophies, also pseudophakic. Average age was 72.1 ± 4.9 years (range, 63 to 76 years). Seven patients were males and eight were females.

Confocal microscopy in vivo examination was carried out on all patient corneas, depending on corneal transparency, between 4 and 7 weeks after the DSAEK procedure using a confocal slit microscope (Confiscan 4; Fortune Technologies, Vigonza, Italy). On each confocal microscopy examination, a sequence of digital images was obtained, which consisted of four consecutive corneal thickness total depth scans, with a scan being equivalent to obtaining images from the endothelium to the epithelium and back to the endothelium, i.e., from the posterior to the anterior view, and then again to the posterior view, in order to enable moving along the central corneal thickness Z-axis. The Z-ring device (Confoscan, Fortune Technologies, Italy) was used, which maintains contact with the cornea in order to obtain reliable thickness measurements without the presence of anteroposterior movement of the eyeball.

Images were automatically captured in a computer hard drive. Measurements were made from the endothelium to the corneal epithelium using the Navis program, v. 3.5.0. (NIDEK, Multi-Instrument Diagnostic System, Japan).

Results

Corneal transparency was observed between 4 and 7 weeks after the DSAEK surgery in all operated corneas. Visual capacity was 20/50 or more in all patients 7 weeks after the procedure.

Donor-receptor interface was found due to the presence of hyper-reflective particles at 114 ± 12.4 microns (from endothelium to epithelium) on average (Fig. 1). Stromal folds began to be observed at 111.1 ± 3.5 microns from the corneal endothelium to up to 286 ± 94 microns towards the corneal epithelium (with an average of 175 ± 90.5-micron stromal thickness, with the presence of folds) (Fig. 2). Donor tissue keratocyte activation began to be observed at 12 ± 1.4 microns from the endothelium to up to 105 ± 38.2 microns towards the corneal epithelium (at an average of 93 ± 36.9-micron stroma thickness, with keratocyte activation) (Fig. 3).

Discussion

Endothelial transplantation is a technique that is taking an important place among the surgical resources for the treatment of corneal pathologies where the only damaged structure is the corneal endothelium and replacement of the entire cornea is therefore not necessary, as in the case of penetrating keratoplasty\(^\text{1,2,4}\).

The DSAEK technique has demonstrated its efficacy even in the management of primary failures in penetrating keratoplasties\(^\text{5}\). Furthermore, favorable results have been reported in terms of visual acuity and postoperative satisfaction of patients undergoing endothelial transplantation with the DSAEK technique\(^\text{5,6}\).

In the present study, early scarring changes were assessed in vivo in corneal tissue after DSAEK surgery. Confocal microscopy was carried out until there was adequate transparency on each one of the operated corneas, in order for the images of their different structures to be adequately evaluable, as well as to enable thickness measurements to be reliably performed without the presence of corneal edema.

Previous confocal microscopy studies in patients operated on using DSAEK report the presence of “fog” in the donor-recipient interface, as well as the presence of particles in said interface, similar to those found in refractive surgery\(^\text{6,8}\). Prasher et al.\(^\text{9}\) also reported the presence of particles in the donor-recipient interface 6 months after surgery, as well keratocyte activation in the anterior stroma, in contrast with the present study, where keratocyte activation was found in the
donor stroma, with the difference that we carried out the confocal microscopy procedure at between 4 and 7 weeks postoperatively. Keratocyte activation on confocal microscopy has been reported in the postoperative period of all techniques that elicit a scarring reaction of the corneal stroma, as in refractive surgery in all its varieties. Keratocyte activation has also been reported in the case of physical procedures, such as cross-linking, The corneal stroma depth at which keratocytes are activated, as well as their permanence, varies depending on the different procedures.

The presence of particles in the interface has been a constant confocal microscopy finding in all surgical techniques where an interface is generated in the corneal stroma, as in the case of Laser Assisted In Situ Keratomileusis (LASIK) surgery, regardless of the use of microkeratome, or femtosecond laser, as well as in deep lamellar keratoplasty.

On the other hand, corneal stroma folds have also been frequently reported with confocal microscopy studies in the postoperative period of corneal surgeries involving the presence of an interface in the corneal stroma, such as LASIK surgery and deep lamellar keratoplasty.

Conclusions

The present study focused completely on early scarring findings on confocal microscopy in patients after endothelial transplantation using the DSAEK technique, which showed both early keratocyte activity and the presence of particles in the donor-receptor interface, as well as folds in the corneal stroma. Clinical results were not the purpose of this study.

Ethical responsibilities

Protection of people and animals. The authors declare that no experiments have been conducted in humans or animals for this research.

Confidentiality of data. The authors declare having followed the protocols of their work center on the publication of patient data.

Right to privacy and informed consent. The authors declare that no patient data appear in this article.

Conflicts of interest

No author has any type of conflict of interest in this article.

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