

Descemet membrane endothelial keratoplasty: graft rejection, failure and survival Baydoun, L.

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General Introduction and Thesis Outline

GENERAL INTRODUCTION

Common indications for corneal transplantation include Fuchs Endothelial Corneal Dystrophy, bullous keratopathy, and failed transplants. These endothelial disorders used to be treated with a full thickness (penetrating) graft for more than a century, until the successful replacement of the host posterior stroma with the diseased Descemet membrane and the endothelial monolayer by a similar donor graft was introduced by Melles in 1998. In the following decade, this posterior lamellar (endothelial) keratoplasty approach underwent continuous anatomical refinement by eliminating the stroma from the graft and herewith achieving a selective replacement of only the diseased Descemet membrane with its endothelium. This approach was called Descemet Membrane Endothelial Keratoplasty (DMEK). DMEK convinces with excellent clinical outcomes and lower complication rates compared to earlier keratoplasty techniques.

This thesis investigates two important complications after DMEK that may have an impact on graft survival, namely allograft rejection and graft failure. By understanding reasons for failed or unsuccessful DMEK grafts, certain complications may be avoided, more quickly recognized, or even prevented in the future. To understand the subtle differences in corneal transplantation techniques, this introduction will provide an overview of the corneal anatomy and surgical approaches and their evolution.

THE HUMAN EYE

The eye is surrounded and shielded by the eyelids. From anterior to posterior, the eye consists of the cornea, conjunctiva and sclera, the anterior chamber with aqueous humor, iris and pupil, the crystalline lens, the vitreous and retina plus choroid, and the optic nerve (Figure 1). The cornea will be the focus of this thesis.

The cornea

The human cornea is the transparent, avascular, and usually invisible structure in the very front of the eye. In adults, the cornea's diameter measures about 11-12 mm, horizontally about 1-2 mm more than vertically, while the corneal thickness is about 500 micrometers (μ m) centrally and increases towards the periphery.¹⁻³ The cornea is a highly innervated tissue with a density of nerve endings that is about 300 to 400 times higher than in the skin.^{2,4} The innervation is required for tissue repair and for pain perception.² Already minor damage to the cornea or a

Figure 1. Cross-section of the human eye. Source: https://www.outlanderanatomy.com/a-real-eye-opener-the-eye-part.

very tiny foreign body on its surface can cause colossal pain and irritation. This can be a real nightmare, but in fact it is a blessing for the eye because ensuing permanent damage or inflammation of the cornea may cause irreversible visual impairment or even blindness. Besides, the cornea is the main refractive component in the eye's optical system that contributes to a sharp image and has a major responsibility in protecting the eye from infections.¹⁻³ Five layers can be distinguished within the cornea, of which three are cellular layers and two serve as interfaces. From the front to the back those layers are the epithelium, Bowman layer, stroma, Descemet membrane and endothelium (Figure 2).¹⁻³

The epithelium is about 40-50 μ m thick and, due to the junctional complexes between the epithelial cells, it forms a strong barrier to germs, toxins, dust, and other substances that may harm the inner eye. The epithelium comprises five to six cell sheets of three sorts of cell types: the outer multilayer of superficial cells which is covered by the tear film is followed by two to three layers of wing cells

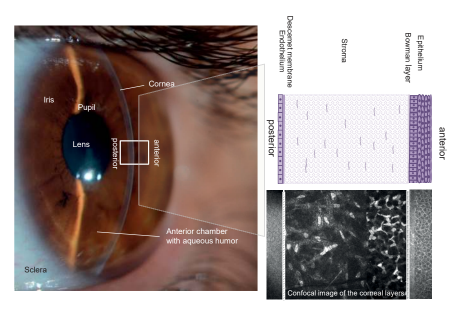


Figure 2. Slit lamp image of the author's anterior eye (left) and a cartoon image (right, middle) and confocal microscopy image (right, bottom) of the corneal layers from anterior to posterior (right to left).

and the deepest single columnar basal cell layer. The latter adheres posteriorly to the epithelial basement membrane.¹²

Directly adjacent to the epithelial basement membrane is the *Bowman layer*, named after William Bowman (1816-1892), who was an English ophthalmologist and anatomist. He discovered this 8-15 μ m thin membrane-like layer that becomes thinner with age and consists of interwoven collagen fibers. The collagen is secreted by stromal keratocytes, and the Bowman layer appears to merge with the stromal fibers, which could explain why it is histologically looked upon as the acellular superficial layer of the anterior stroma.¹² An anatomical characteristic of the Bowman layer that probably contributes to its stiffness and aids in sustaining the corneal shape is that, in contrast to the fibers of the stroma, those of the Bowman layer are smaller in diameter and randomly arranged.^{1-35,6}

The *stroma*, the major component of the cornea, makes up approximately 80-90% of the entire corneal thickness.^{1,3} Just like the Bowman layer, its anatomic structure plays an essential role in sustaining the characteristic corneal shape and rigidity.⁷ The stroma predominantly contains water and non-aqueous components, i.e. collagen fibers, proteoglycans and cells.^{1,2,8} The collagen fibers are

regularly organized in parallel bunches forming so-called fibrils with a characteristically small and uniform diameter.^{1,2,8,9} The interfibril spaces are relatively homogenous and constant, which is maintained by proteoglycans. Fibrils are united in lamellae which are surrounded by keratocytes that form a three-dimensional network and vary in density throughout the stroma. Keratocytes continuously process (digest and produce) stromal components. The central cornea comprises about 200 lamellae with a packing density decreasing from anterior to posterior.^{1,2,8-10} Because the lamellae show considerably more interweavements anteriorly than posteriorly, the posterior stroma swells more easily.^{7,8} In contrast to other collagenous tissues, the cornea is transparent and clear, allowing optimal optical performance. This clarity is achieved by the unique delicate arrangement and organization of the fibrils and their constant and balanced turnover, as well as the lack of blood and lymphatic vessels.^{2,8,9,11} In addition, corneal transparency strongly depends on the integrity of the epithelial and endothelial layers but also on the stable stromal water content of about 78%.²⁹ The deturgescence of the cornea is controlled by the endothelium and its basement membrane, the Descemet membrane.

The Descemet membrane, named after the French physician Jean Descemet (1732-1810), adheres anteriorly to the stroma and posteriorly to the corneal endothelial monolayer that secretes Descemet membrane. The latter is composed of a fine collagen meshwork that thickens after birth until adulthood from 3 μ m to about 10 μ m, respectively.¹² Histologically, three layers can be identified: a thin non-banded layer (0.3 μ m) adjacent to the posterior stroma, an anterior banded zone (2-4 μ m), and a posterior non-banded zone (>4 μ m), which thickens with age.^{12,13} The existence of a distinctive noncellular pre-Descemet stromal layer in the human cornea is controversial.^{14,15}

The endothelium is directly adjacent to the posterior side of Descemet membrane and lines the inner surface of the cornea that is in contact with the aqueous humor in the anterior chamber. The cells have mostly a hexagonal shape, are interlaced and contain various junctions to enable intercellular exchange of molecules and electrolytes.² When examined from posteriorly, this monolayer appears very 'relaxed' and 'peaceful' while resembling a honeycomb mosaic (Figure 3).¹² However, when looking more closely, the endothelial cells have a large nucleus and plenty of cytoplasmic organelles (e.g., mitochondria, Golgi apparatus, etc.) that imply a high metabolic activity.² This activity is essential to maintain corneal transparency to preserve vision. The cells accomplish this by providing a fluid pump and a leaky barrier that together form a so-called pump-

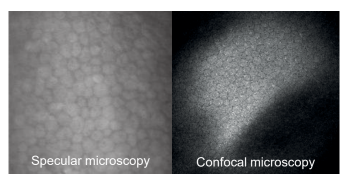


Figure 3. Specular (left) and confocal (right) microscopy images displaying the honeycomb mosaic of the corneal endothelial cell layer.

leak mechanism.^{9,6} Consequently, a relative corneal dehydration is realized by passive (energy-free) fluid diffusion along an osmotic gradient from the relatively hypotonic stroma towards the relatively hypertonic aqueous humor. This stream is promoted by active (energy-requiring) pumps in the endothelial cell walls that transport ions from the stroma to the aqueous humor to maintain the osmotic gradient. Simultaneously, the endothelium possesses a barrier function allowing a selective fluid/nutrients drift from the anterior chamber into the avascular stroma through leaky intercellular junctions.^{12,16-19} Given that the endothelial cells are healthy and sufficient, fluid inflow and outflow is in equilibrium. An imbalance due to endothelial dysfunction or major cell depletion caused, for example, by ocular surgery, inflammation, or trauma, can lead to corneal edema and thickening with subsequent loss of clarity.¹²

So far, the main treatment to restore corneal clarity in opaque corneas is corneal transplantation, also known as keratoplasty. Descemet Membrane Endothelial Keratoplasty (DMEK) is the most recent surgical innovation among endothelial keratoplasty techniques that selectively replaces the diseased endothelium with its Descemet membrane by a healthy donor endothelial Descemet sheet.

ADVANCES IN CORNEAL TRANSPLANTATION

Corneal grafting is the oldest, most frequently performed and most successful form of transplantation in medicine and was developed to restore corneal transparency in eyes with diseased and opaque corneas by replacing the affected tissue by a healthy donor graft. There is notable controversy in the literature about who first introduced the idea and concept of corneal transplantation. Fact is,

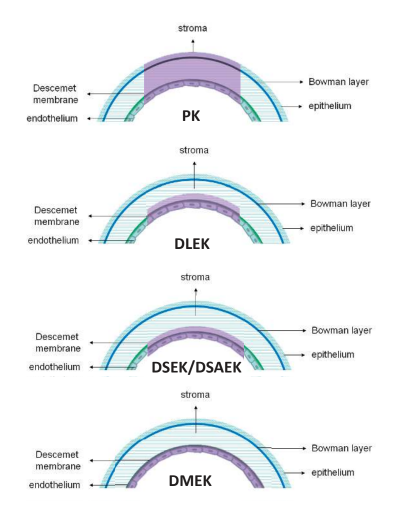


Figure 4. Evolution of Endothelial Keratoplasty.

PK = Penetrating Keratoplasty; DLEK = Deep Lamellar Endothelial Keratoplasty; DSEK/DSAEK = Descemet Stripping (Automated) Endothelial Keratoplasty; DMEK = Descemet Membrane Endothelial Keratoplasty.

that the early efforts can be traced back more than 200 years and, interestingly, the evolution of exchanging the entire corneal tissue (full-thickness or penetrating keratoplasty) or only selected anterior or posterior corneal layers (lamellar keratoplasty) took place side by side from the very beginning (Figure 4).²⁰

Penetrating keratoplasty

In 1789, the French surgeon Guillaume Pellier de Quengsy first suggested to replace corneal scars by glass framed in silver that was then attached to the sclera.²⁰⁻²² However, experiments in corneal transplantation did not begin until the early 19th century. In 1813, a German ophthalmologist from Braunschweig, Karl

CHAPTER 1

Himly, was the first to suggest that opaque animal corneas could be replaced by a healthy cornea of another animal.^{23,24} Later, his student Franz Reisinger initiated experimental animal full thickness corneal transplantations and in 1818 introduced the term 'keratoplasty'.^{20,25} Still, it required 70 more years until the first heterologous graft, a lamellar anterior graft, was successfully transplanted in 1888 by Arthur von Hippel, another German ophthalmologist, who already at that time believed that corneal transparency depended on the integrity of the endothelium and Descemet membrane.^{20,26}

In 1905, the Austrian ophthalmologist Eduard Zirm performed the first homologous penetrating procedure in a human.²⁷ Subsequently, many contemporary ophthalmologists furthered and refined the penetrating approach which appeared less demanding and back then resulted in better clinical outcomes compared to the lamellar approach.^{28,29} Undoubtedly, the advent of general anesthesia, microscopes, 'aseptic' surgical conditions, antibiotics and corticosteroids to treat allograft rejection, as well as the development and improvement of new instruments, trephines, suture material and donor tissue preparation and preservation played an important role in increasing the technique's popularity and success.

Penetrating keratoplasty (PK) remained for many decades the standard of care for a variety of corneal disorders affecting any corneal layer. However, although PK could, to some degree, provide acceptable functional improvement, there were obvious limitations, which included: 'open-sky surgery', suture-related delayed healing and inflammation, secondary ulcers, globe instability, corneal vascularization, higher risk of graft rejection, unpredictable refractive errors with unsatisfactory visual outcomes and graft failure due to endothelial cell decay.³⁰⁻⁴² Attempts in reducing complications and enhancing outcomes by applying laser-guided donor and recipient corneal trephination or secondary artificial lens implantation provided only some relief.⁴³⁻⁴⁶ In addition, strikingly, 30-50% of indications for PK affected only the corneal endothelium, which practically meant the unnecessary removal of also healthy anterior and central corneal tissue.⁴⁷

Posterior lamellar keratoplasty

In the 1950s, the ophthalmologists, José Barraquer from Spain, and Charles Tillet from the United States, were the first to introduce a technique for lamellar endothelial replacement by suturing the transplant underneath a manually dissected stromal flap.^{48,49} This was a promising concept; however, the technique was challenging and difficult to adopt. Later modifications by diverse scientists were not yet successful.^{50,51} In 1998, the Dutch ophthalmologist, Gerrit Melles, introduced a concept in which the adherence of the endothelial transplant to the recipient posterior stroma was achieved by injecting an air bubble into the anterior chamber underneath the graft while the patient remained in a supine position.^{52,53} Posterior Lamellar Keratoplasty (PLK) proved to be clinically successful and was the beginning of an exciting era of successional developments in new techniques (Figure 4).⁵⁴

Deep Lamellar Endothelial keratoplasty (DLEK)

In the primary PLK procedure, a 7-7.5mm recipient stromal disc with the diseased endothelium was excised and a similar-sized donor tissue of stroma and endothelium was inserted through a 9mm and later a 5mm sclero-corneal incision.^{54,55} The latter was technically possible because the graft was folded like a 'taco' and then unfolded inside the recipient's anterior chamber.⁵⁵ Melles` PLK technique was soon popularized in the United States by Mark Terry as 'Deep Lamellar Endothelial Keratoplasty' (DLEK).^{56,57} Although the preliminary results of the procedure were encouraging, the manual dissection of the recipient and donor lenticules were quite demanding which inhibited broad adoption.

Descemet Stripping (Automated) Endothelial Keratoplasty (DS(A)EK)

In 2002, Melles and colleagues modified the DLEK technique by facilitating the removal ('stripping') of the host's diseased corneal Descemet membrane and endothelium. This so-called 'descemetorhexis' was combined with the insertion of the same graft as used earlier in DLEK and was then redefined as 'Descemet-Stripping Endothelial Keratoplasty' (DSEK).⁵⁸⁻⁶⁰ To simplify donor tissue preparation, Mark Gorovoy popularized the use of a microkeratome and adjusted the nomenclature by an additional A: Descemet-stripping Automated Endothelial Keratoplasty' (DSAEK).⁶¹ Standardizing graft preparation was a decisive step in making this technique accessible to a broad range of surgeons because it enabled eye banks to provide pre-cut tissue and surgeons to prepare the tissue on their own. Subsequent adoption of endothelial keratoplasty could no longer be halted since, in contrast to its precursor PK, the lamellar techniques provided better and faster visual recovery with more predictable refractive errors due to the preserved ocular anterior surface. Furthermore, these novel techniques were associated with less complications and minimized the risk of intraoperative expulsive bleeding due to the 'closed-globe' surgery, of traumatic wound dehiscence due to the avoidance of large penetrating incisions, caused less wound

healing problems, less corneal vascularization due to the sutureless surgery, and a remarkably lower risk of allograft rejection.^{62,63}

After almost 100 years, in which PK had been the only available form of (unselective) corneal transplantation, PLK had its breakthrough and was implemented as the new 'gold standard' for the treatment of endothelial pathologies within 10 years of its creation.⁶⁴

Descemet Membrane Endothelial Keratoplasty (DMEK)

Still, also these innovative lamellar procedures showed functional limitations with undesired fluctuations in visual outcome and optical aberrations which seemed attributed to graft thickness and graft irregularity. The additional posterior stroma in the endothelial transplant could induce interface haze and scarring and the preparation of the graft both manually and with a microkeratome could result in an irregular graft thickness. The latter may produce posterior surface irregularities and aberrations that may be even higher than after PK.⁶⁵⁻⁷³

One might expect better visual outcomes after restoration of the human corneal anatomy. Consequently, Melles further refined endothelial keratoplasty by eliminating the stroma from the donor graft and hereby achieving a selective replacement of only the diseased Descemet membrane with its endothelium.⁷⁴⁻⁷⁵ In 2006, Melles performed the first surgery on a Dutch patient (Figure 5) with Fuchs Endothelial Corneal Dystrophy (FECD) using this new approach and named this innovation Descemet Membrane Endothelial Keratoplasty (DMEK).⁷⁶⁻⁷⁷



Figure 5. Photograph showing Dr. Gerrit Melles (left) and the first patient (right) who underwent a Descemet Membrane Endothelial Keratoplasty in August 2006 at the Melles Cornea Clinic, Rotterdam, The Netherlands in 2016. Patient's consent received for publication. Simultaneously, the Melles group developed a technique for DMEK graft preparation that preserved the anterior stroma for transplantation into another eye that suffered from stromal disease.^{78,79} Despite the convincing preliminary results of the thinner DMEK with exceptional visual acuity levels and a considerably lower incidence of allograft rejections, surgeons were hesitant to start with this new technique since preoperative graft preparation, intraoperative graft handling (unfolding, orientation), as well as postoperative graft dehiscence, i.e. the (in) complete DMEK graft detachment from the posterior stroma, were difficult steps and hurdles.⁸⁰⁻⁹¹ In the following years, technique standardization and refinement of preparation and surgery (Figure 6) steepened the surgeon's learning curve and lowered the threshold to offer DMEK to patients.⁹²⁻¹⁰² Aside from increased

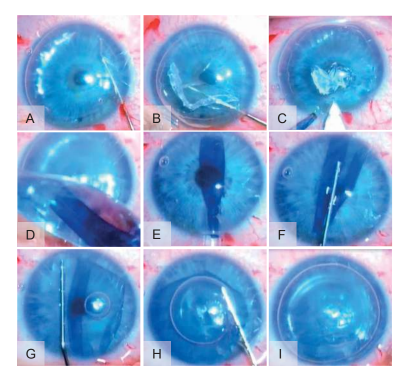


Figure 6. Steps of standardized 'no-touch' DMEK: (A, B) Scoring and stripping of the recipient's Descemet membrane from the posterior stroma, under an air-filled anterior chamber. (C) Creation of a 3.0mm limbal tunnel incision at the 12:00 o'clock position. (D, E) DMEK roll placed in an injector after staining with trypan blue solution before injection into the recipient anterior chamber. (F, G, H) Orientation of the graft (donor endothelial side down, donor Descemet membrane side up facing posterior stroma) and unfolding over the iris. (I) Injection of an air-bubble underneath the graft to position it onto the recipient posterior stroma. At the end of the surgery, the anterior chamber is left completely filled with air for about 30-60 minutes, followed by an air-liquid exchange, leaving a 30% to 50% air-bubble in the anterior chamber.

surgeon's experience, adoption was further encouraged by the decreasing detachment rates thanks to either air or gas tamponade.¹⁰³⁻¹⁰⁸

Since the introduction, DSEK/DSAEK as well as DMEK have evolved into generally established techniques for the treatment of endothelial disorders. Both techniques are still being modified: DSEK/DSAEK grafts are becoming thinner to reach DMEK outcomes in terms of visual acuity levels, whereas DMEK modifications aim at utilizing endothelial donor tissue more efficiently by transplanting different graft shapes or sizes.¹⁰⁸⁻¹¹⁹ Accordingly, in 2014 and 2016, respectively, the Melles group introduced 'Hemi-DMEK' and 'Quarter-DMEK' techniques in which from one donor cornea two semi-circular or four quarter-shaped DMEK grafts can be prepared that can potentially be transplanted into two or four recipients (Figure 7).¹¹²⁻¹¹⁹ This approach would then allow doubling and quadrupling the number of endothelial grafts recovered from the same donor pool. Although standard DMEK and modified DMEK techniques may be similar surgically, the indications for each surgery type may differ.

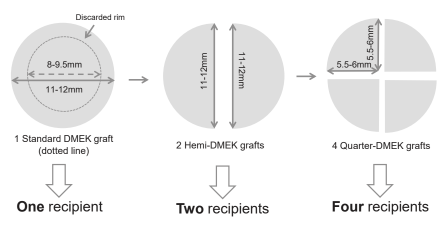


Figure 7. Different graft shapes of available DMEK techniques. (Left) standard DMEK: a central 8.5-9.5mm circular DMEK-graft is trephinated and transplanted into one recipient; the outer Descemet membrane rim is discarded. (Middle) Hemi-DMEK: the entire 11-12mm diameter Descemet membrane sheet is utilized to obtain two semi-circular grafts to be transplanted into two recipients. Each Hemi-DMEK graft has a similar surface area as a circular standard DMEK graft. (Right) Quarter DMEK: each Hemi-DMEK graft is further divided into two quarters, providing in total four Quarter-DMEK grafts to be transplanted into four recipients.

INDICATIONS FOR DMEK

Patients with visual impairment from endothelial dysfunction are the target group for endothelial keratoplasty. DMEK has gained broad popularity over the past years. Along with the rising numbers of procedures, also the surgical experience with this technique has expanded and with it the ease for surgeons to perform DMEK on more patients with diverse endothelial diseases.^{6,103,120,121}

Fuchs Endothelial Corneal Dystrophy (FECD)

FECD, the most common indication for DMEK, was first described in 1910 by the Austrian ophthalmologist, Ernst Fuchs, who initially termed it 'Dystrophia epithelialis corneae' since he assumed an epithelial pathology due to the morphologic changes in the corneal surface with reduced corneal sensitivity. Despite lacking a slit lamp, Fuchs assumed a defective endothelial layer, but since he could not judge the distinct anatomic changes, he suggested to redefine etiology and nomenclature later again if necessary.¹²² One typical characteristic of FECD is the presence of corneal 'guttae' which were first observed and termed by the Swiss ophthalmologist Alfred Vogt in 1921.¹²³ 'Guttae' result from focal thickening of the Descemet membrane forming excrescences due to accumulations of collagen that are produced by abnormal endothelial cells.² They are often scattered more horizontally and inferiorly over the posterior corneal surface than vertically and superiorly.^{124,125} Disease progression with continuous loss of endothelial cells ultimately results in corneal edema with subsequent fibrotic changes of stroma and epithelium. Patients complain of fluctuating or gradually deteriorating vision, glare (especially at night), but also of ocular pain from ruptured epithelial bullae due to uncovered nerve endings.^{2,126-128}

FECD is a slowly progressive, usually bilateral disease of the posterior corneal layers.² An early-onset and a late-onset phenotype can be distinguished. The early-onset form occurs in only 1% of FECD cases, is equally present in males and females and can manifest already in the first decade of life to progress further before the age of 50. The late-onset, more common form occurs predominantly in women and develops in the second or third decade, to become symptomatic in the fifth to sixth decade.^{2,127-129} Clinical progression of FECD is graded in different stages depending on the extension and confluence of guttae and the presence of corneal edema.^{2,127,130,131} While mild edema may not necessarily cause visual deterioration, the presence of guttae without edema can significantly induce light scattering and photophobia, as well as reduce contrast sensitivity that may be very bothersome for the patient.¹³²

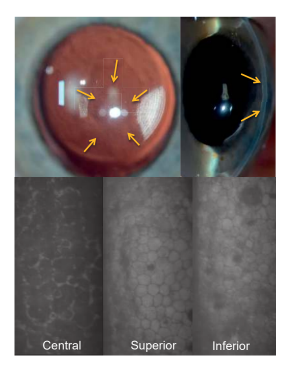


Figure 8. Slit lamp images (upper row) of a cornea in reflecting light (left) and with a slit from the side (right) showing centrally localized guttae (arrows). Specular microscopy images (lower row) show that in the same eye central endothelial cells are not visible whereas in the peripheral areas (superior, inferior) cells can be visualized with only few guttae (dark spots).

Recently, it has been suggested that in FECD eyes with only centrally localized guttae and no or only mild corneal edema (Figure 8), transplantation of a smaller Quarter-DMEK graft may be sufficient to enhance corneal clearance in the stripped area and diminish visual distortions.^{118,119} Removing only the affected central Descemet membrane and endothelium with a small central descemetorhexis may help to sustain viable peripheral recipient endothelial cells, potentially even eliminating the need for a transplant as in Descemetorhexis only (DSO).¹³³⁻¹³⁷

Bullous keratopathy

Bullous keratopathy stands for an entity of conditions that may show the same clinical picture but that differ in their underlying cause. These eyes are often characterized by pronounced and quickly progressing corneal edema that is associated with endothelial cell depletion. It often follows surgical trauma, as in 'pseudophakic bullous keratopathy' after (complicated) cataract surgery or after glaucoma (tube) surgery, but can also occur in eyes with congenital glaucoma, aphakia or rarely after ocular trauma.^{138,139} In contrast to patients with FECD, patients with bullous keratopathy often experience a faster non-fluctuating decrease in visual acuity, while visual distortions and pain are similar.

Failed transplants

Another indication for DMEK consists of eyes with corneal decompensation following corneal transplantation, so-called graft failure. Corneal edema results from the inability of the (remaining) graft endothelial cells to maintain corneal clarity. When irreversible and accompanied by major visual impairment, only repeat transplantation can restore vision. Graft failure can occur after any form of transplantation, i.e., PK, DSEK/DSAEK and DMEK. From long-term studies we know that certain eves are more prone to develop failure.¹⁴⁰⁻¹⁴² Risk factors for graft failure include an enhanced cell decline and low cell count early after keratoplasty, high risk recipient preoperative indications (e.g., herpetic eye disease), concomitant glaucoma, repeat grafting, prior rejection episodes, atopic ocular surface disease and corneal vascularization.^{38-42,140-145} In early keratoplasty studies, the term graft failure traditionally referred to endothelial failure, i.e., a non-functioning graft. More recent studies 'confusingly' also use this term to characterize any graft that requires repeat transplantation, like for unsatisfying refractive or visual outcomes after PK or DSEK/DSAEK, despite a clear graft with functioning endothelium.41,146

DMEK COMPLICATIONS

Complications can either occur intraoperatively and/or postoperatively after any surgery. With the novel PLK techniques, but especially with DMEK, the type, incidence and severity of complications has changed considerably. Still, prevention and improved management of complications will further enhance clinical outcomes and patient satisfaction. In the following part, DMEK complications that are relevant for this thesis will be described.

Graft dehiscence

Following the introduction of endothelial keratoplasty, graft detachment evolved as a new complication in the early postoperative period.⁸⁹⁻⁹¹ It describes the partial or complete non-adherence of the endothelial graft to the recipient posterior stroma. Compared to DSEK/DSAEK grafts, DMEK grafts detach more easily and tend to curl up, which often hinders spontaneous re-attachment.¹⁴⁷ Although it was initially assumed that complete apposition of the graft is required to achieve corneal clearance, we now know from clinical observations that corneas with smaller detachments, for example involving only graft edges (\leq 1/3 of the graft surface area), can clear without secondary intervention.¹⁴⁸⁻¹⁵¹ Spontaneous clearance was also observed in eyes with larger or complete detachments with

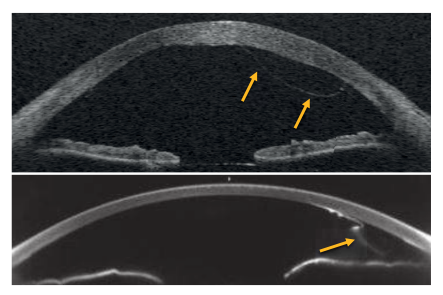


Figure 9. DMEK graft detachments (arrows) visible on Anterior Segment Optical Coherence Tomography (upper picture, notice the curling of the graft towards the stroma, indicating correct orientation) and on a Scheimpflug image (lower picture).

or without involvement of the optical axis (> 1/3 of the graft surface area).^{150,151} Because clearance would often take longer (> 3 months) and may not be longlasting, nowadays a re-injection of air/gas (so-called re-bubbling) or direct regrafting is recommended in larger detachments.^{147,152} Diverse causes for DMEK detachments have been identified, among which are: insufficient pressurization of the eye with wound leakage at the end of the surgery, residual Descemet remnants, eye rubbing and inward graft folds.¹⁵³ A preventable reason for DMEK detachment is an inverted graft.¹⁴⁷ After stripping the Descemet membrane from the posterior stroma, it usually forms a roll with the endothelium on the outer side. This knowledge is essential to decide on proper graft orientation during surgery.^{78,92} For identifying graft dehiscence, anterior segment Optical Coherence Tomography (OCT), that allows cross-sectional imaging of the anterior eye, has become indispensable (Figure 9).¹⁵⁴ Some surgeons use this tool also intraoperatively (intraoperative OCT) to determine and manage graft orientation and unfolding.^{102,155} In contrast to an edematous cornea, in reasonably clear corneas, detached graft areas can also be visualized on a Scheimpflug image (Figure 9).¹⁵⁴

Allograft rejection

The immune system is essential to fight (life)threatening pathogens and agents. We can differentiate two forms of immune reactions: a non-specific quick 'in-

nate (natural) immune system' that attracts immune cells to the required sites by excreting cytokines and that triggers the specific 'adaptive (acquired) immune system' by presenting specific antigens via the lymph nodes.² Although the adaptive immune system reacts initially with a delay, it can be more efficient because it is antigen-specific and develops a memory to react quicker in case of a second 'attack' by the same pathogen. The adaptive immune system utilizes two specific immune cells: B cells that direct the humoral (antibody) system and T cells, the cellular arm of the adaptive system; both are equipped with specific receptors. The B-cell receptor is an antibody that directly recognizes specific pathogens whereas the T-cell receptor can only recognize processed pathogens that are presented via HLA (human leukocyte antigen) molecules, which are cellsurface proteins on other immune cells. There are two classes of HLA antigens: 1. HLA Class I molecules, which present intracellular peptides to cytotoxic T cells (CD8 positive), and 2. HLA Class II molecules which present extracellular antigens (from outside a cell) to helper T cells (CD4 positive). Antigen presentation on HLA Class II molecules triggers the development of an immune response. HLA Class II molecules are expressed on a variety of immune cells, such as antigen presenting cells (APCs, e.g., dendritic cells, macrophages, monocytes), but also on B cells and activated T cells.²

The eye consists of vulnerable tissues with limited capabilities to regenerate: not only corneal endothelial cells but also retinal cells cannot proliferate. Consequently, an immune-mediated inflammation can lead to cell damage with functional loss or even blindness. To protect the eye, nature has provided it with a unique 'immune privilege'.^{156,157}

The Dutchman, Jacobus van Dooremaal, a student of F.C. Donders in Utrecht, already showed in 1873 that he could implant living tissue in the anterior chamber of the eye of dogs and rabbits without getting the normal severe immune response.¹⁵⁸

In 1948, the British zoologist Peter Medawar and colleagues were the first who discovered that the eye was an immune privilege site. Their conclusions were based on observations of prolonged survival of an allogeneic tissue graft that was placed in the anterior chamber of the eye.¹⁵⁹ Since this work was ground-breaking for tissue and organ transplantation, Medewar is often referred to as the 'father of transplantation'.¹⁶⁰

CHAPTER 1

To date, the immune privilege of the cornea and anterior chamber is explained by the following mechanisms: 1. anatomical, cellular, and molecular barriers, 2. eye-derived immunological tolerance, and 3. an active systemic ocular immuno-suppressive component.^{156,157,159,161} The blood-aqueous barrier and the avascular nature (lacking blood and lymphatic vessels) of the cornea hinder inflammatory cells from entering the eye and antigens to be presented to the immune-competent sites.^{162,163} Tolerance is achieved by different mechanisms, such as a relative absence of APCs and the moderate expression of HLA antigens in stroma and endothelium.¹⁶⁴ In addition, an active suppression is realized by the presence of immunosuppressive cytokines (e.g., transforming growth factor- β , α -melanocyte stimulating hormone, vasoactive intestinal peptide), and anterior chamber-associated immune deviation (ACAID), that specifically helps to suppress cellular delayed type hypersensitivity and antibody-medicated immune responses against antigens introduced into the anterior chamber.¹⁶⁵⁻¹⁶⁷

This immune privilege is thought to be the reason why corneal transplantation is among the most successful tissue transplantations even without tissue typing or systemic immunosuppression.¹⁶⁸ This relates only to low-risk keratoplasties.^{38,169} Some transplanted corneas still experience allograft rejection: a major risk factor is the extent of corneal vascularization and the state of the recipient bed.² Consequently, a breakdown in the immune privilege can be observed when a graft is placed in an inflamed eye, in cases with heavy recipient vascularization, after pre-sensitization by a previously rejected graft, in case of large eccentric grafts or application of pro-inflammatory medication.^{2,166,169-173}

Since the introduction of endothelial keratoplasty, rejection rates have decreased with each technical refinement, possibly due to an increasing experience of surgeons, leading to less surgical trauma, and by transplantation of less donor tissue.^{174,175} Until two years after surgery, rejection rates with PK, DSEK/DS(A)EK, and DMEK were 20%, 12%, and 1%, respectively, in a study with similar indications for grafting, comparable patient demographics and the same topical corticosteroid treatment.⁸⁵ This low rejection rate, but also the occurrence of milder rejection forms, were observed by several scientific groups.^{84,85,175-179} With DMEK, HLA matching is practically not performed anymore and rejection rates did not increase after lowering the steroid potency one month after surgery but did increase when steroids were discontinued in the second year after DMEK.^{178,179} Recently, the cumulative rejection rate 10 years post-DMEK was 4%.¹⁸⁰ Risk factors for rejection in DMEK are still unclear; however, in a study on DSEK, Afro-American race was found to be a significant risk factor for rejection.¹⁸¹ Interestingly, mor-

phologic endothelial cell changes were observed in eyes that later developed DMEK rejection.¹⁸² Although graft failure in eyes following DMEK-rejection in the first five years after surgery was rare,¹⁷⁵ it would be useful to avoid any allograft rejection and subsequent endothelial cell decline by recognizing eyes at risk of developing rejection.

Endothelial cell decay and graft failure

Bowman was the first to describe the appearance of endothelial cells under a microscope. In vivo visualization, however, only became possible in 1920, when Vogt described this mosaic of hexagonal cells visible in the specular reflection of the illuminator in a slit lamp biomicroscope.¹⁹ Nowadays, specular microscopy and meanwhile also confocal microscopy are standard tools to visualize and evaluate the corneal endothelial cells and assess their approximate number (Figure 3).^{183,184} At birth, the endothelial cell density measures around 6000 cells/mm² and decreases to 3500 cells/mm² within only five years and further to 2700 cells/ mm² before adolescence.^{16,185,186} While the cell decrease is accelerated in childhood, after the age of 18 years, the average rate of decrease stabilizes at 0.6% per year.^{185,186} Since endothelial cells have no mitotic activity and cannot regenerate, the damage or loss of cells cannot be compensated.¹ Instead, residual neighboring cells migrate and spread out to carpet the denuded space which results in deformation (loss of hexagonality = pleomorphism) and enlargement (polymegathism) of the cells as part of the endothelial wound healing process.^{12,16-19}

Endothelial cell density decrease can increase after ocular surgery and most importantly in post-keratoplasty corneas.¹⁸⁷⁻¹⁸⁹ Following DMEK, endothelial cell density decreases by approximately 25-40% in the early phase, and after the 6-month follow-up, by 7-9% yearly.^{190,191} Endothelial cell density is regularly evaluated after keratoplasty because its decrease is assumed to have impact on the survival of a graft. And although corneal clarity can be maintained with a density as low as 500 cells/mm², when a threshold is reached, the cornea may decompensate.¹⁸⁷

When the cornea fails to clear directly after keratoplasty, this is referred to as 'primary or early failure' mainly due to impaired donor quality but sometimes also due to difficult surgery. In recent DMEK reports, the term primary failure is also used in eyes that required re-grafting for DMEK detachment or upside-down graft positioning. However, in these cases, corneal edema derives rather from a technical failure, i.e., the graft is not positioned/oriented properly, than from endothelial dysfunction. If the graft clears after surgery but fails later, it is referred to

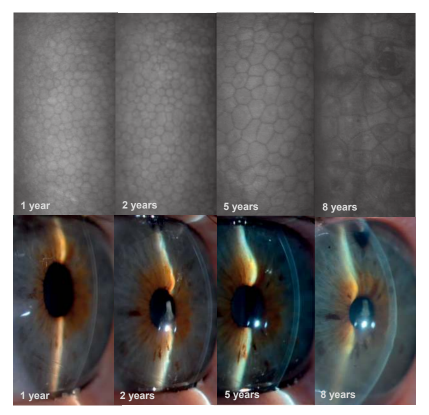


Figure 10. Specular Microscopy (upper row) and slit lamp images (lower row) of an eye eventually developing graft failure at 8 years after Descemet Membrane Endothelial Keratoplasty.

as 'secondary or later failure'. This can, for example, occur after rejection episodes, but also due to comorbidities like glaucoma or the primary indication for surgery (Figure 10). The number of failed grafts in a cohort tells us something about the survival rate of a graft. After PK, survival rates may vary from 75% to 95% at 3 and 5 years and from 50% to 80% at 10 years postoperatively, depending on indication and complications. After DSAEK and DSEK, 3- and 5-year survival rates of respectively 87% to 97% and 93% were reported. Early results after DMEK suggest that survival rates may at least be similar to earlier keratoplasty techniques.¹⁹²⁻¹⁹⁸

AIM AND THESIS OUTLINE

This thesis investigates two complications after DMEK that may have an impact on graft survival, namely graft failure and allograft rejection. Owing to the reduced complication rates in DMEK, we hypothesized that DMEK survival is promising and that outcomes of repeat DMEK to manage 'failed' grafts are acceptable. We further hypothesized that allograft rejection is a rare complication that appears to be more subtle after DMEK and that corneal changes before rejection may manifest themselves by other diagnostic approaches. To test our assumptions, we investigated the following objectives:

The first four chapters focus on graft longevity, graft failure, and repeat DMEK. In **Chapter 2**, the 10-year clinical outcome of the first ever DMEK patient is evaluated. **Chapter 3** depicts the longer-term endothelial survival after successful DMEK surgery in a large DMEK cohort. By identifying eyes with endothelial graft failure, possible reasons, and risk factors for DMEK graft failure are described. **Chapter 4** illustrates the problem of the diverging use of the term graft failure not only for endothelial failure but also for any DMEK graft that needs replacement without considering newly evolved complications such as graft detachment. In **Chapter 5**, the feasibility, and reasons for repeat DMEK along with the clinical outcomes and complications of repeat surgery up to 12 months are investigated.

The next four chapters focus on inflammation and allograft rejection. In **Chapter 6**, we assess whether aqueous flare can be used as a measure of subclinical inflammation in eyes following DMEK. In **Chapter 7**, we identify eyes with allograft rejection in a large consecutive DMEK series and determine the relevance of Scheimpflug imaging as a diagnostic tool for detection of corneal changes preceding allograft rejection. In **Chapter 8**, we correlate the occurrence of corneal changes seen on Scheimpflug images and specular microscopy in eyes before rejection manifests. In addition, we assess the increase in pachymetry and the decrease of the endothelial cell density.

In **Chapter 9,** we analyse the feasibility and outcome of transplanting a DMEK modification in which four smaller Quarter-DMEK grafts, potentially less-antigenic grafts, from the same donor are transplanted into four recipients.

REFERENCES

- DelMonte DW, Kim T. Anatomy and physiology of the cornea. J Cataract Refract Surg 2011; 37:588-98
- Cornea Textbook. Fundamentals, Diagnosis and Management. Volume 1. Krachemer JH, Mannis MJ, Holland EJ. 3rd Edition. Elvesier Mosby 2011
- 3. Massoudi D, Malecaze F, Galiacy SD. Collagens and proteoglycans of the cornea: importance in transparency and visual disorders. *Cell Tissue Res* 2016;363:337-49
- Müller LJ, Marfurt CF, Kruse F, Tervo TM. Corneal nerves: structure, contents, and function. Exp Eye Res 2003;76:521-42
- 5. Germundsson J, Karanis G, Fagerholm P, Lagali N. Age-related thinning of Bowman's layer in the human cornea in vivo. *Invest Ophthalmol Vis Sci* 2013;54:6143-9
- 6. Marshall J. The 2014 Bowman Lecture Bowman's and Bruch's: a tale of two membranes during the laser revolution. *Eye (Lond)* 2015;29:46-64
- 7. Müller LJ, Pels E, Vrensen GF. The specific architecture of the anterior stroma accounts for maintenance of corneal curvature. *Br J Ophthalmol* 2001;85:437-43
- 8. Meek KM, Knupp C. Corneal structure, and transparency. Prog Retin Eye Res 2015;49:1-16
- 9. Maurice DM. The cornea and sclera, In: Davidson H (Ed.). The Eye. New York Academic 1984
- Bergmanson JP, Horne J, Doughty MJ, Garcia M, Gondo M. Assessment of the number of lamellae in the central region of the normal human corneal stroma at the resolution of the transmission electron microscope. *Eye Contact Lens* 2005;31:281-7
- 11. Streilein JW. New thoughts on the immunology of corneal transplantation. Eye 2003;17: 943-8
- 12. Johnson DH, Bourne WM, Campbell RJ. The ultrastructure of Descemet's membrane. I. Changes with age in normal corneas. *Arch Ophthalmol* 1982;100:1942-7
- Bourne WM, Johnson DH, Campbell RJ. The ultrastructure of Descemet's membrane. III. Fuchs' dystrophy. Arch Ophthalmol 1982;100:1952-5
- 14. Dua HS, Faraj LA, Said DG, Gray T, Lowe J. Human corneal anatomy redefined: a novel pre-Descemet layer (Dua's layer). *Ophthalmology* 2013;120:1778-85
- 15. Schlötzer-Schrehardt U, Bachmann BO, Tourtas T, et al. Ultrastructure of the posterior corneal stroma. *Ophthalmology* 2015;122:693-9
- 16. Bourne WM. Biology of the corneal endothelium in health and disease. Eye 2003;17:912-8
- Fischbarg J, Maurice DM. An update on corneal hydration control. *Exp Eye Res* 2004;78: 537-41
- 18. Mergler S, Pleyer U. The human corneal endothelium: New insights into electrophysiology and ion channels. *Progress in Retinal and Eye Research* 2007;26:359-78
- 19. Bourne WM. Corneal endothelium-past, present, and future. *Eye Contact Lens* 2010;36: 310-4
- 20. Frank Joseph Goes. The Eye in History. Jaypee Brothers Medical Publishers. First Edition 2013
- 21. De Quengsy G (1789) Précis ou cours d'opérations sur la chirurgie des yeux: puisé dans le sein de la pratique, & enrichi de figures en taille-douce. Didot, Paris
- 22. Chirila TV, Hicks CR. The origins of the artificial cornea: Pellier de Quengsy and his contribution to the modern concept of keratoprosthesis. *Gesnerus* 1999;56:96-106
- 23. Moffatt SL, Cartwright VA, Stumpf TH. Centennial review of corneal transplantation. *Clin Experiment Ophthalmol* 2005;33:642-57

- 24. Crawford AZ, Patel DV, McGhee CNJ. A brief history of corneal transplantation: From ancient to modern. *Oman J Ophthalmol* 2013;6:12-7
- 25. Reisinger F. Die Keratoplastik: ein Versuch zur Erweiterung der Augenheilkunst. *Bayerische* Ann Chir Augenheilk 1824;207-15
- 26. Von Hippel A. Eine neue Methode der Hornhauttransplantation. Albrecht von Graefes Arch Ophthalmol 1888;34(I):108-30
- 27. Zirm E. Eine erfolgreiche totale Keratoplastik. Albrecht von Graefes Arch Ophthalmol 1906;54:580-93
- 28. Filatov VP. Transplantation of the cornea. Arch Ophthalmol 1935;13,321-47
- 29. Castroviejo R. Keratoplasty Am J Ophthalmol 1941;24:1-20
- 30. Tan DT, Dart JK, Holland EJ, Kinoshita S. Corneal transplantation. Lancet 2012;379:1749-61
- 31. Olson RJ. Complications associated with running 11-0 nylon suture in penetrating keratoplasty. Ophthalmic Surg 1982;13:558-61
- 32. Christo CG, van Rooij J, Geerards AJ, Remeijer L, Beekhuis WH. Suture-related complications following keratoplasty: a 5-year retrospective study. *Cornea* 2001;20:816-19
- 33. Rohrbach JM, Weidle EG, Steuhl KP, Meilinger S, Pleyer U. Traumatic wound dehiscence after penetrating keratoplasty. *Acta Ophthalmol Scand* 1996;74:501-5
- 34. Elder MJ, Stack RR. Globe rupture following penetrating keratoplasty. How often, why, and what can we do to prevent it? *Cornea* 2004;23:776-80
- 35. Nagra PK, Hammersmith KM, Rapuano CJ, Laibson PR, Cohen Ej. Wound dehiscence after penetrating keratoplasty. *Cornea* 2006;25:132-5
- Williams KA, Muehlberg SM, Lewis RF, Coster DJ. How successful is corneal transplantation? A report from the Australian Corneal Graft Register. *Eye* 1995;9:219-27
- 37. Felipe AF, Hammersmith KM, Nottage JM, et al. Indications, visual outcome, and ectasia in clear corneal transplants 20 years old or more. *Cornea* 2013;32:602-7
- Niziol LM, Musch DC, Gillespie BW, Marcotte LM, Sugar A. Long-term outcomes in patients who received a corneal graft for keratoconus between 1980 and 1986. Am J Ophthalmol 2013;155:213-9
- 39. Ing JJ, Ing HH, Nelson LR, Hodge DO, Bourne WM. Ten-year postoperative results of penetrating keratoplasty. *Ophthalmology* 1998;105:1855-65
- 40. Price MO, Thompson RW Jr, Price FW Jr. Risk factors for various causes of failure in initial corneal grafts. *Arch Ophthalmol* 2003;121:1087-92
- 41. Thompson RW Jr, Price MO, Bowers PJ, Price FW Jr. Long-term graft survival after penetrating keratoplasty. *Ophthalmology* 2003;110:1396-402
- 42. Inoue K, Amano S, Oshika T, Tsuru T. Risk factors for corneal graft failure and rejection in penetrating keratoplasty. *Acta Ophthalmol Scand* 2001;79:251-5
- 43. Bahar I, Kaiserman I, McAllum P, Rootman D. Femtosecond laser-assisted penetrating keratoplasty: stability evaluation of different wound configurations. *Cornea* 2008;27:209-11
- 44. Farid M, Kim M, Steinert RF. Results of penetrating keratoplasty performed with a femtosecond laser zigzag incision initial report. *Ophthalmology* 2007;114:2208-12
- 45. Price FW Jr, Price MO. Femtosecond laser shaped penetrating keratoplasty: one-year results utilizing a top-hat configuration. *Am J Ophthalmol* 2008;145:210-4
- 46. Srinivasan S, Ting DS, Lyall DA. Implantation of a customized toric intraocular lens for correction of post-keratoplasty astigmatism. *Eye (Lond)* 2013;27:531-7
- 47. Cursiefen C, Küchle M, Naumann GO. Changing indications for penetrating keratoplasty: histopathology of 1,250 corneal buttons. *Cornea* 1998;17:468-70

- 48. Barraquer JI. Lamellar keratoplasty. (Special techniques). Ann Ophthalmol 1972;4:437-69
- 49. Tillett CW. Posterior lamellar keratoplasty. *Am J Ophthalmol* 1956;41:530-3
- 50. Jones DT, Culbertson WW. 1998. Endothelial lamellar keratoplasty (ELK). *Investig Ophthalmol Vis Sci* 39:876 (Abstr.)
- 51. Culbertson WW. Endothelial replacement: flap approach. Ophthalmol Clin North Am 2003;16:113-8
- 52. Melles GRJ, Eggink FAGJ, Lander F, et al. A surgical technique for posterior lamellar keratoplasty. *Cornea* 1998;17:618-26
- 53. Melles GRJ, Lander F, Beekhuis WH, Remeijer L, Binder PS. Posterior lamellar keratoplasty for a case of pseudophakic bullous keratopathy. *Am J Ophthalmol* 1999;127:340-1
- Melles GRJ, Lander F, van Dooren BTH, Pels E, Beekhuis WH. Preliminary clinical results of posterior lamellar keratoplasty through a sclerocorneal pocket incision. *Ophthalmology* 2000;107:1850-7
- 55. Melles GR, Lander F, Nieuwendaal C. Sutureless, posterior lamellar keratoplasty: A case report of a modified technique. *Cornea* 2002;21:325-7
- 56. Terry MA, Ousley PJ. Deep lamellar endothelial keratoplasty in the first United States patients: early clinical results. *Cornea* 2001;20:239-43
- 57. Terry MA, Ousley PJ. Deep lamellar endothelial keratoplasty visual acuity, astigmatism, and endothelial survival in a large prospective series. *Ophthalmology* 2005;112:1541-8
- 58. Melles GR, Wijdh RH, Nieuwendaal CP. A technique to excise the Descemet membrane from a recipient cornea (descemetorhexis). *Cornea* 2004;23:286-8
- 59. Gorovoy M, Price, FW. New technique transforms corneal transplantation. *Cataract Refract* Surg Today 2005;11:55-8
- 60. Price FW Jr, Price MO. Descemet's stripping with endothelial keratoplasty in 50 eyes: a refractive neutral corneal transplant. *J Refract Surg* 2005;21:339-45
- 61. Gorovoy MS. Descemet-stripping automated endothelial keratoplasty. *Cornea* 2006;25:886-9
- 62. Chen ES, Terry MA, Shamie N, Hoar KL, Friend DJ. Precut tissue in Descemet's stripping automated endothelial keratoplasty donor characteristics and early postoperative complications. *Ophthalmology* 2008;115:497-502
- 63. Price MO, Gorovoy M, Benetz BA, et al. Descemet's stripping automated endothelial keratoplasty outcomes compared with penetrating keratoplasty from the Cornea Donor Study. *Ophthalmology* 2010;117:438-44
- 64. Eye Bank Association of America. 2015 Eye Banking Statistical Report. Washington, D.C., United States, 2016. Available at http://restoresight.org/wp-content/ uploads/2016/03/2015-Statistical-Report.pdf
- 65. Patel SV, Baratz KH, Hodge DO, Maguire LJ, McLaren JW. The effect of corneal light scatter on vision after Descemet stripping with endothelial keratoplasty. *Arch Ophthalmol* 2009;127:153-60
- 66. Price MO, Price FW Jr. Endothelial keratoplasty a review. *Clin Experiment Ophthalmol* 2010;38:128-40
- 67. McLaren JW, Patel SV. Modeling the effect of forward scatter and aberrations on visual acuity after endothelial keratoplasty. *Invest Ophthalmol Vis Sci* 2012;53:5545-51
- 68. Anshu A, Price MO, Tan DTH, Price FW Jr. Endothelial keratoplasty: a revolution in evolution? *Surv Ophthalmol* 2012;57:236-52

- 69. Dirisamer M, Parker J, Naveiras M, et al. Identifying causes for poor visual outcome after DSEK/DSAEK following secondary DMEK in the same eye. *Acta Ophthalmol* 2013;91:131-9
- Heinzelmann S, Bohringer D, Maier PC, Reinhard T. Correlation between visual acuity and interface reflectivity measured by pentacam following DSAEK. Acta Ophthalmol 2014;92:1-4
- Seery LS, Nau CB, McLaren JW, et al. Graft thickness, graft folds, and aberrations after Descemet stripping endothelial keratoplasty for Fuchs dystrophy. Am J Ophthalmol 2011;152:910-6
- 72. Chamberlain W, Omid N, Lin A, et al. Comparison of corneal surface higher-order aberrations after endothelial keratoplasty, femtosecond laser-assisted keratoplasty, and conventional penetrating keratoplasty. *Cornea* 2011;31:6-13
- Koh S, Maeda N, Nakagawa T, et al. Characteristic higher order aberrations of the anterior and posterior corneal surfaces in 3 corneal transplantation techniques. *Am J Ophthalmol* 2012;153:284-90
- Melles GRJ, Rietveld FJR, Pels E, Beekhuis WH, Binder PS. Transplantation of Descemet's membrane carrying viable endothelium through a small scleral incision. ARVO Abstract 1998
- 75. Melles GR, Lander F, Rietveld FJ. Transplantation of Descemet's membrane carrying viable endothelium through a small scleral incision. *Cornea* 2002;21:415-8
- 76. Melles GR. Posterior lamellar keratoplasty: DLEK to DSEK to DMEK. Cornea 2006;25:879-81
- 77. Melles GR, Ong TS, Ververs B, van der Wees J. Descemet membrane endothelial keratoplasty (DMEK). *Cornea* 2006;25:987-90
- 78. Lie JT, Birbal R, Ham L, van der Wees J, Melles GRJ. Donor tissue preparation for Descemet membrane endothelial keratoplasty. *J Cataract Refract Surg* 2008;34:1578-83
- 79. Lie JT, Groeneveld EA, Ham L, van der Wees J, Melles G.R.J. More efficient use of donor corneal tissue with Descemet membrane endothelial keratoplasty (DMEK). *Br J Ophthalmol* 2010;94:1265-6
- Melles GR, Ong TS, Ververs B, van der Wees J. Preliminary clinical results of Descemet membrane endothelial keratoplasty. Am J Ophthalmol 2008;145:222-7
- 81. Ham L, Dapena I, van Luijk C, van der Wees J, Melles GR. Descemet membrane endothelial keratoplasty (DMEK) for Fuchs endothelial dystrophy: review of the first 50 consecutive cases. *Eye (Lond)* 2009;23:1990-8
- 82. Ham L, Balachandran C, Verschoor CA, van der Wees J, Melles GR. Visual rehabilitation rate after isolated descemet membrane transplantation: descemet membrane endothelial keratoplasty. *Arch Ophthalmol* 2009;127:252-5
- 83. Dapena I, Ham L, Droutsas K, et al. Learning curve in Descemet's membrane endothelial keratoplasty: First series of 135 consecutive cases, *Ophthalmology* 2011;118:2147-54
- 84. Dapena I, Ham L, Netuková M, van der Wees J, Melles GR. Incidence of early allograft rejection after Descemet membrane endothelial keratoplasty. *Cornea* 2011;30:1341-5
- 85. Anshu A, Price MO, Price FW Jr. Risk of corneal transplant rejection significantly reduced with Descemet's membrane endothelial keratoplasty. *Ophthalmology* 2012;119:536-40
- Tourtas T, Laaser K, Bachmann BO, Cursiefen C, Kruse FE. Descemet membrane endothelial keratoplasty versus Descemet stripping automated endothelial keratoplasty. Am J Ophthalmol 2012;153:1082-90
- 87. Neff KD, Biber JM, Holland EJ. Comparison of central corneal graft thickness to visual acuity outcomes in endothelial keratoplasty. *Cornea* 2011;30:388-91

- 88. Dapena I, Ham L, Melles GRJ. Endothelial keratoplasty: DSEK/DSAEK or DMEK the thinner the better? *Curr Opin Ophthalmol* 2009;20:299-307
- 89. Dapena I, Moutsouris K, Ham L, Melles GR, Graft detachment rate, *Ophthalmology* 2010;117:847
- 90. Maier A-KB, Gundlach E, Pilger D, et al. Rate and localization of graft detachment in Descemet membrane endothelial keratoplasty. *Cornea* 2016;35:308-12
- 91. Cursiefen C. Descemet membrane endothelial keratoplasty: the taming of the shrew. JAMA Ophthalmol 2013;131:88-9
- 92. Groeneveld-van Beek EA, Lie JT, van der Wees J, Bruinsma M, Melles GR. Standardized 'no-touch' donor tissue preparation for DALK and DMEK: harvesting undamaged anterior and posterior transplants from the same donor cornea. *Acta Ophthalmol* 2013;91:145-50
- Kruse FE, Laaser K, Cursiefen C, et al. A stepwise approach to donor preparation and insertion increases safety and outcome of descemet membrane endothelial keratoplasty. *Cornea* 2011;30:580-7
- 94. Dapena I, Moutsouris K, Droutsas K, et al. Standardized "no-touch" technique for Descemet membrane endothelial keratoplasty. *Arch Ophthalmol* 2011;129:88-94
- 95. Liarakos VS, Dapena I, Ham L, van Dijk K, Melles GR. Intraocular graft unfolding techniques in Descemet membrane endothelial keratoplasty. *JAMA Ophthalmol* 2013;131:29-35
- 96. Bachmann BO, Laaser K, Cursiefen C, Kruse FE. A method to confirm correct orientation of descemet membrane during descemet membrane endothelial keratoplasty. *Am J Ophthalmol* 2010;149:922-5
- 97. McCauley MB, Price FW Jr., Price MO. 2009. Descemet membrane automated endothelial keratoplasty: hybrid technique combining DSAEK stability with DMEK visual results. J *Cataract Refract. Surg* 35:1659-64
- 98. Studeny P, Farkas A, Vokrojova M, Liskova P, Jirsova K. Descemet membrane endothelial keratoplasty with a stromal rim (DMEK-S). *Br J Ophthalmol* 2010;94:909-14
- 99. Da Reitz Pereira C, Guerra FP, Price FW Jr, Price MO. Descemet's membrane automated endothelial keratoplasty (DMAEK): visual outcomes and visual quality. *Br J Ophthalmol* 2011;95:951-4
- 100. Burkhart ZN, Feng MT, Price MO, Price FW. Handheld slit beam techniques to facilitate DMEK and DALK. Cornea 2013;32:722-4
- Veldman BP, Dye PK, Holiman JD, et al. The S-stamp in Descemet membrane endothelial keratoplasty safely eliminates upside-down graft implantation. *Ophthalmology* 2016;123:161-4
- Saad A, Guilbert E, Grise-Dulac A, Sabatier P, Gatinel D. Intraoperative OCT-assisted DMEK: 14 consecutive cases. Cornea 2015; 34:802-7
- Rodríguez-Calvo-de-Mora M, Quilendrino R, Ham L, et al. Clinical outcome of 500 consecutive cases undergoing Descemet's membrane endothelial keratoplasty. Ophthalmology 2015;122:464-70
- 104. Tourtas T, Schlomberg J, Wessel JM, et al. Graft adhesion in descemet membrane endothelial keratoplasty dependent on size of removal of host's descemet membrane. JAMA Ophthalmol 2014;132:155-61
- 105. Price MO, Price FW Jr. Descemet's membrane endothelial keratoplasty surgery: update on the evidence and hurdles to acceptance. *Curr Opin Ophthalmol* 2013;24:329-35

- 106. Güell JL, Morral M, Gris O, Elies D, Manero F. Comparison of Sulfur Hexafluoride 20% versus Air Tamponade in Descemet Membrane Endothelial Keratoplasty. Ophthalmology 2015;122:1757-64
- Terry MA, Straiko MD, Veldman PB, et al. Standardized DMEK Technique: Reducing Complications Using Prestripped Tissue, Novel Glass Injector, and Sulfur Hexafluoride (SF6) Gas. Cornea 2015;34:845-52
- 108. Schaub F, Enders P, Snijders K, et al. One-year outcome after Descemet membrane endothelial keratoplasty (DMEK) comparing sulfur hexafluoride (SF6) 20% versus 100% air for anterior chamber tamponade. Br J Ophthalmol 2017;101:902-8
- 109. Busin M, Madi S, Santorum P, Scorcia V, Beltz J. Ultrathin descemet's stripping automated endothelial keratoplasty with the microkeratome double-pass technique: two-year outcomes. Ophthalmology 2013;120:1186-94
- 110. Cheung AY, Hou JH, Bedard P, et al. Technique for Preparing Ultrathin and Nanothin Descemet Stripping Automated Endothelial Keratoplasty Tissue. *Cornea* 2018;37:661-6
- 111. Chamberlain W, Lin CC, Austin A, et al. Descemet Endothelial Thickness Comparison Trial: A Randomized Trial Comparing Ultrathin Descemet Stripping Automated Endothelial Keratoplasty with Descemet Membrane Endothelial Keratoplasty. Ophthalmology 2019;126:19-26
- 112. Lam FC, Baydoun L, Dirisamer M, et al. Hemi-Descemet membrane endothelial keratoplasty transplantation: a potential method for increasing the pool of endothelial graft tissue. JAMA Ophthalmol 2014;132:1469-73
- Lam FC, Baydoun L, Satué M, et al. One year outcome of hemi-Descemet membrane endothelial keratoplasty. Graefes Arch Clin Exp Ophthalmol 2015;253:1955-8
- 114. Lie JT, Lam FC, Groeneveld-van Beek EA, Van der Wees J, Melles GRJ. Graft preparation for hemi-Descemet membrane endothelial keratoplasty (hemi-DMEK). Br J Ophthalmol 2016;100:420-4
- 115. Gerber-Hollbach N, Parker J, Baydoun L, et al. Preliminary outcome of hemi-Descemet membrane endothelial keratoplasty for Fuchs endothelial dystrophy. Br J Ophthalmol 2016;100:1564-8
- 116. Müller TM, Baydoun L, Melles GR. 3-Year update on the first case series of hemi-Descemet membrane endothelial keratoplasty. *Graefes Arch Clin Exp Ophthalmol* 2017;255:213-5
- 117. Birbal RS, Hsien S, Zygoura V, et al. Outcomes of Hemi-Descemet Membrane Endothelial Keratoplasty for Fuchs Endothelial Corneal Dystrophy. *Cornea* 2018;37:854-8
- 118. Müller TM, Lavy I, Baydoun L, et al. Case report of Quarter-Descemet membrane endothelial keratoplasty for Fuchs endothelial dystrophy. *Cornea* 2017;36:104-7
- Zygoura V, Baydoun L, Ham L, et al. Quarter-Descemet membrane endothelial keratoplasty (Quarter-DMEK) for Fuchs endothelial corneal dystrophy: 6 months clinical outcome. Br J Ophthalmol 2018;102:1425-30
- 120. Oellerich S, Baydoun L, Peraza-Nieves J, et al. Multicenter Study of 6-Month Clinical Outcomes After Descemet Membrane Endothelial Keratoplasty. *Cornea* 2017;36:1467-76
- 121. Price MO, Gupta P, Lass J, Price FW Jr. EK (DLEK, DSEK, DMEK): New Frontier in Cornea Surgery. *Annu Rev Vis Sci* 2017;15;3:69–90
- 122. Fuchs E. Dystrophia epithelialis corneae. Graefe's Arch Ophthalmol 1910;76:478-508
- 123. Vogt A: Die Sichtbarkeit des lebenden Hornhautendothels. Ein Beitrag zur Methodik der Spaltlampenmikroskopie. *Graefe's Arch Ophthalmol* 1920; 101:123-44

- 124. Adamis AP, Filatov V, Tripathi BJ, Tripathi RC: Fuchs' endothelial dystrophy of the cornea. Surv Ophthalmol 1993;38:149-68
- 125. Fujimoto H, Maeda N, Soma T, et al. Quantitative regional differences in corneal endothelial abnormalities in the central and peripheral zones in Fuchs' endothelial corneal dystrophy. *Invest Ophthalmol Vis Sci* 2014;55:5090-8
- 126. Wilson SE, Bourne WM. Fuchs' dystrophy. Cornea 1988;7:2-18. Review
- 127. Krachmer JH, Purcell JJ Jr, Young CW, Bucher KD. Corneal endothelial dystrophy: A study of 64 families. *Arch Ophthalmol* 1978;96:2036-9
- 128. Vedana G, Villarreal G Jr, Jun AS. Fuchs endothelial corneal dystrophy: current perspectives. *Clinical Ophthalmology* 2016:10 321-30
- 129. Weiss JS, Møller HU, Aldave AJ, et al. IC3D classification of corneal dystrophies edition 2. Cornea 2015;34:117-59
- 130. Louttit MD, Kopplin LJ, Igo RP Jr, et al. FECD Genetics Multi-Center Study Group. A multicenter study to map genes for Fuchs endothelial corneal dystrophy: baseline characteristics and heritability. *Cornea* 2012 ;31:26-35
- 131. Eghrari AO, Garrett BS, Mumtaz AA, et al. Retroillumination Photography Analysis Enhances Clinical Definition of Severe Fuchs Corneal Dystrophy. *Cornea* 2015;34:1623-6
- Watanabe S, Oie Y, Fujimoto H, et al. Relationship between Corneal Guttae and Quality of Vision in Patients with Mild Fuchs' Endothelial Corneal Dystrophy. *Ophthalmology* 2015;122:2103-9
- 133. Moloney G, Chan UT, Hamilton A, et al. Descemetorhexis for Fuchs' dystrophy. Can J Ophthalmol 2015;50:68-72
- 134. Borkar DS, Veldman P, Colby KA. Treatment of Fuchs Endothelial Dystrophy by Descemet Stripping Without Endothelial Keratoplasty. *Cornea* 2016;35:1267-73
- 135. Moloney G, Petsoglou C, Ball M, et al. Descemetorhexis Without Grafting for Fuchs Endothelial Dystrophy-Supplementation With Topical Ripasudil. *Cornea* 2017;36:642-8
- 136. Garcerant D, Hirnschall N, Toalster N, et al. Descemet's stripping without endothelialkeratoplasty. Review *Curr Opin Ophthalmol* 2019;30:275-85
- 137. Artaechevarria Artieda J, Wells M, Devasahayam RN, Moloney G. 5-Year Outcomes of Descemet Stripping Only in Fuchs Dystrophy. Cornea 2020;39:1048-51
- 138. Morishige N, Sonoda KH. Bullous keratopathy as a progressive disease: evidence from clinical and laboratory imaging studies. *Cornea* 2013;32:77-83
- 139. Gonçalves ED, Paris Fdos S, Gomes JÁ, Kanecadan L, Campos M. Bullous keratopathy. Ophthalmology 2011;118:2303
- 140. Patel SV, Diehl NN, Hodge DO, Bourne WM. Donor risk factors for graft failure in a twentyyear study of penetrating keratoplasty. *Arch Ophthalmol* 2010;128: 418-25
- Sugar A, Tanner JP, Dontchev M, et al.. Recipient risk factors for graft failure in the Cornea Donor Study. Ophthalmology 2009;116:1023-8
- 142. Yu AL, Kaiser M, Schaumberger M, et al. Perioperative and postoperative risk factors for corneal graft failure. *Clin Ophthalmol* 2014;8:1641-7
- 143. Patel SV. Graft survival and endothelial outcomes in the new era of endothelial keratoplasty. *Exp Eye Res* 2012;95:40-7
- 144. Benetz B, Lass JH, Gal RL, Sugar A, et al. Cornea Donor Study Investigator Group. Endothelial morphometric measures to predict endothelial graft failure after penetrating keratoplasty. JAMA Ophthalmol 2013;131:601-8

- 145. Lass JH, Sugar A, Benetz BA, et al. Cornea Donor Study Investigator Group. Endothelial cell density to predict endothelial graft failure after penetrating keratoplasty. *Arch Ophthalmol* 2010;128:63-9
- 146. Letko E, Price DA, Lindoso EM, Price MO, Price FW Jr. Secondary graft failure and repeat endothelial keratoplasty after Descemet's stripping automated endothelial keratoplasty. Ophthalmology 2011;118:310-4
- Gerber-Hollbach N, Baydoun L, López EF, Frank LE, Dapena I, Liarakos VS, Schaal SC, Ham L, Oellerich S, Melles GRJ. Clinical Outcome of Rebubbling for Graft Detachment After Descemet Membrane Endothelial Keratoplasty. *Cornea* 2017;36:771-6
- 148. Romaniv N, Price MO, Price FW, Mamalis N. Donor Descemet membrane detachment after endothelial keratoplasty. Cornea 2006;25:943-7
- 149. Srinivasan S, Rootman DS. Slit-lamp technique of draining interface fluid following Descemet's stripping endothelial keratoplasty. *Br J Ophthalmol* 2007;91:1202-5
- 150. Balachandran C, Ham L, Verschoor CA, et al. Spontaneous corneal clearance despite graft detachment in descemet membrane endothelial keratoplasty. Am J Ophthalmol 2009;148:227-34
- 151. Dirisamer M, Dapena I, Ham L, et al. Patterns of corneal endothelialization and corneal clearance after descemet membrane endothelial keratoplasty for Fuchs endothelial dystrophy. Am J Ophthalmol 2011;152:543-55
- 152. Feng MT, Srinivas SP, Miller JM, Price FW. Air reinjection and endothelial cell density in Descemet membrane endothelial keratoplasty: five-year follow-up. *J Cataract Refract Surg* 2014;40:1116-21
- 153. Müller TM, Verdijk RM, Lavy I, et al. Histopathologic Features of Descemet Membrane Endothelial Keratoplasty Graft Remnants, Folds, and Detachments. *Ophthalmology* 2016;123:2489-97
- 154. Moutsouris K, Dapena I, Ham L, et al. Optical coherence tomography, Scheimpflug imaging, and slit-lamp biomicroscopy in the early detection of graft detachment after Descemet membrane endothelial keratoplasty. *Cornea* 2011;30:1369-75
- 155. Steven P, Le Blanc C, Velten K, et al. Optimizing descemet membrane endothelial keratoplasty using intraoperative optical coherence tomography. *JAMA Ophthalmol* 2013;131:1135-42
- 156. Hori J. Mechanisms of immune privilege in the anterior segment of the eye: what we learn from corneal transplantation. *J Ocul Biol Dis Infor* 2008;1:94-100
- 157. Niederkorn JY. See no evil, hear no evil, do no evil: the lessons of immune privilege. *Nat Rev Immuno* 2006;7:354-9
- 158. J.C. van Dooremaal, Over de gevolgen van het invoeren van levende weefsels en doode voorwerpen in het oog. Thesis, Utrecht, 1873
- 159. Medawar PB. Immunity to homologous grafted skin; the fate of skin homografts transplanted to the brain, to subcutaneous tissue, and to the anterior chamber of the eye. Br J Exp Pathol 1948;29:58-69
- 160. Starzl TE. Peter Brian Medawar: father of transplantation. J Am Coll Surg 1995;180:332-6
- 161. Streilein JW. Ocular immune privilege: therapeutic opportunities from an experiment of nature. *Nat Rev Immuno* 2003;3:879-89
- 162. Maurice DM & Mishima S (1984): Ocular pharmacokinetics. In: Sears M (ed.) Pharmacology of the Eye. Berlin: Springer-Verlag 19-116

- 163. Collin HB (1966): Endothelial cell-lined lymphatics in the vascularized rabbit cornea. *Invest Ophthalmol* 5:337-54
- 164. Whitsett CF & Stulting RD. The distribution of HLA antigens on human corneal tissue. Invest Ophthalmol Vis Sci 1984;25:519-24
- 165. D'Orazio TJ & Niederkorn JY. A novel role for TGF-b and IL-10 in the induction of immune privilege. *J Immunol* 1998;160:2089-98
- 166. Pleyer U, Schlickeiser S. The taming of the shrew? The immunology of corneal transplantation. Acta Ophthalmol 2009;87:488-97
- 167. Streilein JW. Peripheral tolerance induction: lessons from immune privileged sites and tissues. *Transplant Proc* 1996,28:2066-70
- Streilein JW, Yamada J, Dana MR, Ksander BR. Anterior chamber-associated immune deviation, ocular immune privilege, and orthotopic corneal allografts. *Transplant Proc* 1999;31:1472-5
- Qazi Y, Hamrah P. Corneal Allograft Rejection: Immunopathogenesis to Therapeutics. J Clin Cell Immunol 2013;2013(Suppl 9):006
- 170. Lapp T, Heinzelmann S, Shanab WA, Reinhard T, Boehringer D. Graft decentering in DSAEK: a risk factor for immune reactions? *Eye (Lond)* 2016;30:1147-9
- 171. Bachmann B, Taylor RS, Cursiefen C. Corneal neovascularization as a risk factor for graft failure and rejection after keratoplasty: an evidence-based meta-analysis. *Ophthalmology* 2010;117:1300-5
- 172. van Klink E, Jager MJ. Effect of interferon-alpha on MHC expression and influx of inflammatory cells on the rat eye. *Doc Ophthalmol* 1996-1997;92:17-28
- 173. Niederkorn JY. High risk corneal allografts and and why they lose their immune privilege. *Curr Opin Allergy Clin Immunol* 2010;10:493-7
- 174. Allan BD, Terry MA, Price FW Jr, et al. Corneal transplant rejection rate and severity after endothelial keratoplasty. *Cornea* 2007;26:1039-42
- 175. Price DA, Kelley M, Price FW Jr, Price MO. Five-Year Graft Survival of Descemet Membrane Endothelial Keratoplasty (EK) versus Descemet Stripping EK and the Effect of Donor Sex Matching. Ophthalmology 2018;125:1508-14
- 176. Schlögl A, Tourtas T, Kruse FE, Weller JM. Long-term Clinical Outcome After Descemet Membrane Endothelial Keratoplasty. *Am J Ophthalmol* 2016;169:218-26
- 177. Hos D, Tuac O, Schaub F, et al. Incidence and Clinical Course of Immune Reactions after DMEK: Retrospective Analysis of 1000 Consecutive Eyes. *Ophthalmology* 2017;124:512-8
- 178. Böhringer D, Schwartzkopff J, Maier PC, Reinhard T. HLA matching in keratoplasty: lessons learned from lamellar techniques. *Klin Monbl Augenheilkd* 2013;230:490-3
- 179. Price MO, Scanameo A, Feng MT, Price FW Jr. Descemet's Membrane Endothelial Keratoplasty: Risk of Immunologic Rejection Episodes after Discontinuing Topical Corticosteroids. *Ophthalmology* 2016;123:1232-6
- Vasiliauskaitė I, Oellerich S, Ham L, et al. Descemet Membrane Endothelial Keratoplasty: Ten-Year Graft Survival and Clinical Outcomes. Am J Ophthalmol 2020;217:114-20
- Price MO, Jordan CS, Moore G, Price FW Jr. Graft rejection episodes after Descemet stripping with endothelial keratoplasty: part two: the statistical analysis of probability and risk factors. Br J Ophthalmol 2009;93:391-5
- 182. Monnereau C, Bruinsma M, Ham L, et al. Endothelial cell changes as an indicator for upcoming allograft rejection following Descemet Membrane Endothelial Keratoplasty. Am J Ophthalmol 2014;158:485-95

- McCarey B, Edelhauser HF, Lynn MJ. Review of corneal endothelial specular microscopy for FDA clinical trials of refractive procedures, surgical devices and new intraocular drugs and solutions. *Cornea* 2008;27:1-16
- 184. Niederer RL, McGhee CNJ. Clinical in vivo confocal microscopy of the human cornea in health and disease. *Prog Retin Eye Res* 2010;29:30-58
- 185. Bourne WM, Nelson LR, Hodge DO. Central corneal endothelial cell changes over a tenyear period. *Invest Ophthalmol Vis Sci* 1997;38:779-82
- 186. Nucci P, Brancato R, Mets MB, Shevell SK. Normal endothelial cell density range in childhood. Arch Ophthalmol 1990;108:247-8
- 187. Bourne WM. Cellular changes in transplanted human corneas. Cornea 2001;20:560-9
- 188. Bourne WM. Morphologic and functional evaluation of the endothelium of transplanted human corneas. *Trans Am Ophthalmol Soc* 1983;81:403-50
- 189. Armitage WJ, Dick AD, Bourne WM. Predicting endothelial cell loss and long-term corneal graft survival. *Invest Ophthalmol Vis Sci* 2003;44:3326-31
- 190. Ham L, Dapena I, Liarakos VS, et al. Midterm Results of Descemet Membrane Endothelial Keratoplasty: 4 to 7 Years Clinical Outcome. *Am J Ophthalmol* 2016;171:113-21
- Baydoun L, Tong CM, Tse WW, et al. Endothelial cell density after Descemet membrane endothelial keratoplasty: 1 to 5-year follow-up. Am J Ophthalmol 2012;154:762-3
- 192. Ang M, Mehta JS, Lim F, et al. Endothelial cell loss and graft survival after Descemet's stripping automated endothelial keratoplasty and penetrating keratoplasty. *Ophthalmology* 2012;119:2239-44
- 193. Price MO, Gorovoy M, Price FW Jr., et al. Descemet's stripping automated endothelial keratoplasty. Three-year graft and endothelial cell survival compared with penetrating keratoplasty. Ophthalmology 2013;120:246-51
- 194. Price MO, Fairchild KM, Price DA, Price FW. Descemet's stripping endothelial keratoplasty. Five-year graft survival and endothelial cell loss. Ophthalmology 2011;118:725-9
- 195. Borderie VM, Boëlle P-Y, Touzeau O, et al. Predicted Long-term outcome of corneal transplantation. *Ophthalmology* 2009;116:2354-60
- 196. Lass JH, Benetz BA, Gal RL, et al. Writing Committee for the Donor Study Research Group. Donor age and factors related to endothelial cell loss 10 years after penetrating keratoplasty. Specular microscopy ancillary study. Ophthalmology 2013;120:2428-35
- Sugar A, Gal RL, Kollman C, et al. Writing Committee for the Donor Study Research Group.
 Factors associated with corneal graft survival. JAMA Ophthalmol 2015;133:246-54
- 198. Price MO, Giebel AW, Fairchild KM, Price FW Jr. Descemet's membrane endothelial keratoplasty (DMEK): Prospective multicenter study of visual and refractive outcomes and endothelial survival. *Ophthalmology* 2009;116:2361-8

