Design of functional biomaterials as substrates for corneal endothelium tissue engineering

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Abstract
Corneal endothelium defects are one of the leading causes of blindness worldwide. The actual treatment is transplantation, which requires the use of human cadaveric donors, but it faces several problems, such as global shortage of donors. Therefore, new alternatives are being developed and, among them, cell therapy has gained interest in the last years due to its promising results in tissue regeneration. Nevertheless, the direct administration of cells may sometimes have limited success due to the immune response, hence requiring the combination with extracellular mimicking materials. In this review, we present different methods to obtain corneal endothelial cells from diverse cell sources such as pluripotent or multipotent stem cells. Moreover, we discuss different substrates in order to allow a correct implantation as a cell sheet and to promote an enhanced cell behavior. For this reason, natural or synthetic matrices that mimic the native environment have been developed. These matrices have been optimized in terms of their physicochemical properties, such as stiffness, topography, composition and transparency. To further enhance the matrices properties, these can be tuned by incorporating certain molecules that can be delivered in a sustained manner in order to enhance biological behavior. Finally, we elucidate future directions for corneal endothelial regeneration, such as 3D printing, in order to obtain patient-specific substrates.

Keywords: corneal endothelium; tissue engineering; cell therapy; biomaterials

Introduction
Diseases affecting the cornea are the fifth leading cause of blindness affecting more than 10 million people in the world [1, 2]. Nowadays, corneal transplant is the gold standard treatment for these diseases. In fact, it is the most frequent type of transplant performed worldwide [3].

An organ transplant is the replacement of a dysfunctional tissue or organ with a healthy one that can be from the same patient (autograft), from a human donor (allograft) or an animal (xenograft). Corneal transplant, named as keratoplasty, can be performed using two different techniques, mainly replacing the full cornea, known as penetrating keratoplasty (PK), or only replacing the damaged cornea layers, known as partial thickness transplants [4]. PK replaces the five layers of the cornea, whereas partial thickness transplants conserve the healthy and functional tissue from the cornea, presenting mainly two types of transplants, being the deep anterior lamellar keratoplasty (DALK) and the endothelial keratoplasty (EK). DALK replaces the anterior layers (epithelium, Bowman’s membrane and stroma), whereas EK selectively replaces the posterior layers (endothelium, Descemet’s membrane and, in some cases, part of the stroma; Fig. 1). Over the last decade, the number of PK has been decreasing and partial thickness transplants are the most selected option. In fact, according to the annual Eye Bank Association of America (EBAA) report from 2021, EK was the most performed keratoplasty, representing more than 60% of all corneal transplants [5]. The top indication for corneal transplant is related to Fuch’s dystrophy, a disease that affects the corneal endothelium, which represents 39% of all transplants performed [3].

Despite of the great advances in the clinical scenario, corneal transplantation still faces several unmet challenges, such as donor shortage and rejection, that limit the overall viability and urges the seek of alternative solutions [3, 6]. Taking into account that the most damaged and vulnerable tissue is the endothelium, there is a need to find alternative solutions such as the development of an in vitro tissue construct for corneal endothelium regeneration.

The main difference of the corneal endothelium with other tissues or cell layers in the human organs is that corneal endothelial cells (CEC), upon external aggression or aging, do not present regenerative capacity. This is mainly related to their non-proliferative state, leading to a decrease in functionality, reducing the transparency and allowing the formation of edema. Subsequently, this edema can result in a pathological condition
named as bullous keratopathy, requiring a surgical treatment in order to recover the corneal dysfunction [7].

Current alternatives are based on stem cell therapies that can be combined with synthetic matrices. Cell therapies have gained interest as there are several routes that may allow the production of CEC. Once these CEC are produced, these need to either assemble among them forming a cell sheet that can be directly implanted, or to be combined with extracellular matrices (ECM) materials for its subsequent implantation. In the latter case, these cells need to be in direct contact with a substrate similar to the Descemet’s membrane (DM) and attach onto it. Hence, in order to replicate the native scenario, biomaterials are being designed with several assets similar to the DM that will allow a better interaction and will, at the same time, allow direct implantation of a construct composed of biomaterials and cells.

Here, we aim at describing the current alternatives to corneal endothelial transplantation, taking special consideration in the design of biomaterials that can provide a support for CEC. In this sense, biomaterials can have two pivotal roles, allowing on the one hand, the production of cell sheets to allow direct implantation in a scaffold-free system or the combination of cells with adequate biomaterials. In both cases, biomaterial design will play a key role in providing the stimuli required for cells to produce the desired morphologies and functionalities.

**Required properties for corneal endothelium regeneration**

Native tissues present several conditions that allow the different functionalities to work properly. The different functionalities refer to the main characteristics and task each tissue should perform. For instance, bone is a tissue that provides structural support to the body, hence, having a damaged bone, may limit the structural support of the body. In the case of the cornea, and more specifically, the corneal endothelium is mainly related with a proper visualization, hence having alterations in the corneal endothelium may limit the visibility as well as induce visual impairment or even blindness. Nevertheless, whenever there is damage and the cells are found in abnormal conditions, normal functionalities are lost (Fig. 2). Abnormal conditions generally take place either due to an increase in cell size or to a change in cell morphology that may ultimately reduce the properties required for the proper functionality (Fig. 2).

CEC are of great relevance and, hence, these need to be carefully examined to evaluate the need of transplantation. Eventually, these CEC, in the case of deceasing, can be a potential cell source to be transplanted to a patient requiring corneal endothelium replacement. For this purpose, CEC quality has been deeply studied, showing that age significantly affects cell quality and proper donors need to be carefully selected. In this sense, abnormal cell morphology (Fig. 3) may lead to non-functional corneal endothelium which limits vision capacity [9].

In order to maintain the properties as similar as possible to native tissue, we initially provide an overview of the different aspects related to the corneal endothelium in order to take this into account when designing materials for its replacement. There are several parameters that define normal behavior of the corneal endothelium, which are listed below (Fig. 4).

**Transparency** refers to the ability of light to go through the corneal endothelium. CEC play an essential role in the maintenance of corneal transparency through its barrier and the Na+/K+ -ATPase pump functions that regulate corneal hydration [10]. The normal function of corneal endothelium is achieved with an optimum cell density, considered as 2500–3000 cells/mm² [11]. An abnormal pump function results in an excess of fluid in the corneal stroma due to fluid absorption from the aqueous humor. This excess of fluid in the cornea changes the composition of the proteins, which ends in corneal edema and, subsequently, in a decreased level of corneal transparency [10].

**Permeability** is the ability to allow the flow of nutritional elements through the CEC. As cornea is an avascular tissue, the corneal endothelium represents a nutritional gateway between the anterior chamber and the cornea. Most of the nutrients required for the cornea, which are allocated in the aqueous humor, are transferred into the cornea through CEC pumps [12]. In order to maintain this nutritional function, a minimum cell density of 400–500 cells/mm² is required [13].

**Mechanical stiffness** refers to the stability, flexibility and rigidity of the layer that will allow to properly maintain the shape of the layer. The different layers of the cornea have shown values of elastic modulus between 50 and 100 kPa in humans [14]. Nevertheless, the corneal endothelium has shown significantly lower levels, ranging between 5 and 10 kPa in rabbits [15]. Hence, the mechanical stability of the corneal endothelium is not strictly relevant, as the structural mechanical properties seem to be based on the other layers of the cornea. The main component of the DM is collagen, so using collagen as a substrate while providing a stiffness that is comparable to the native environment of CEC seems as a promising approach. In fact, Gutermuth et al. [16] evaluated that seeding cells onto

**Figure 1.** Anatomy of the eye and cornea. The cornea is the outer layer of the eye and contains different layers: epithelium, Bowman’s membrane, stroma, Descemet’s membrane and endothelium. Current transplant are DALK replaces posterior layers, EK replaces anterior layers, and PK replaces the entire cornea. DALK, deep anterior lamellar keratoplasty, EK, endothelial keratoplasty; PK, penetrating keratoplasty.
smooth collagen enhanced its functionality compared to a silicone substrate.

Topography refers to the presence of surfaces with determined roughness that are able to provide physical cues to the attached cells. Topography is among the leading physical cues to guide stem cell fate [17]. Different morphologies have been proposed, which can be prepared by replica molding or by stereolithography [18]. A recent study analyzed the microtopography of a DM using 3D confocal microscopy [16]. This consisted of flat hexagonal pits with an irregular morphology, which ends in the formation of the characteristic polygonal or hexagonal morphology of CEC [11]. Their width ranged from 10 to 20 μm and their maximal height was 1 μm [16]. The results showed that mimicking the native structure of DM allowed enhanced CEC functionality. Interestingly, the topography of the DM plays a key role as well in determining the fate of Fuchs’ endothelial dystrophy, which is based on the presence of microtopography on the DM [19]. Previous findings showed that when culturing cells on different topographical features, CEC tended to form monolayers on lower height and density substrates, whereas there seemed to be a reduction in the capacity to form CEC monolayer in the densely packed substrates. Hence, topographical features can as well be used to predict the ability of CEC to form the needed continuous monolayer [19].

Composition refers to the vast combination of proteins and polysaccharides present in the cornea that are able to provide the chemical cues and mechanical stability of the matrix. In general, basement membranes are rich in type IV collagen (COL IV), but more specifically, the DM also contains type VIII collagen (COL VIII) that perform a hexagonal pattern [20, 21]. Furthermore, the DM also contains laminin, nidogen, entactin and some glycoproteins and proteoglycans [22].

Thickness (although related with stiffness and transparency): DM thickness increases through age, its values vary from 3 μm at birth and arriving to 10 μm in adults [23]. Normally, when performing DMEK, the graft is usually between 10 and 15 μm.

Manipulation in surgery: Graft orientation is very important, as CEC have to be in contact with the humor aqueous to be functional. The ’up-side-down’ graft placement should be avoided [22].

Curvature: The internal part of the cornea has a circular diameter of 11.7 mm and a sagittal height of 2.18 mm to the center of the cornea [23]. Interestingly, a corneal map has been depicted in order to understand how the curvature affects the healing of the

Figure 2. Corneal morphology and functionality in normal and abnormal conditions. Cells present an hexagonal shape in normal conditions whereas its shape and size its modified in abnormal conditions (A). Light can enter through and arrive to the inner parts of the eye in normal conditions but light is not able to arrive in abnormal conditions (B). Adapted with permission from Ref. [8].
This may provide relevant information on how to design biomaterials with specific curvatures in order to adjust the tension produced by the curvature [24, 25]. For instance, an in vitro model using a radius of 8.4 mm and an angle of 100° allowed stimulating cell alignment and consequently enhance the human corneal tissue equivalents [26].
Sources of therapeutic cells

In order to provide a functional corneal endothelium, the presence of cells is necessary. As previously described, these cells have limited availability from donors and for this reason, it is of great importance to find alternative sources, which should be from the autologous origin where possible, that may allow replicating similar cell structures. In this sense, corneal endothelial regeneration using cell therapies is based on the isolation and expansion of human CEC (HCEC) or the differentiation of stem cells into CEC.

CEC culture conditions

It has been reported that the modulation of phosphatidylinositol 3-kinase (PI3K)/Akt and Smad2 signaling pathways promotes CEC proliferation and wound healing [27]. Hence, previous research has evaluated the effect of different molecules in cell culture medium as potential modulators of the mentioned pathways.

On the other hand, epidermal growth factor (EGF), basic fibroblasts growth factor (bFGF) and transforming growth factor beta 2 (TGF-β2), which are present in the endothelium and the aqueous humor, have been described to enhance in vitro cell migration, proliferation and corneal wound healing [27]. This can be explained as EGF and bFGF activate PI3K/Akt pathway, which permits CEC to enter in the proliferative S-phase of the cell cycle [28–32]. Differently, TGF-β2 induces the suppression of the cell proliferation. Despite of this suppression of the proliferative status, TGF-β2 is able to enhance cell migration and corneal endothelial wound healing through the Smad2 pathway.

On the other hand, it has been reported that insulin growth factor 1 (IGF-1), heresulgin beta and activin A permits an upregulation of PI3K/Akt and Smad2 signaling pathways, which promote CEC proliferation, migration and physiological functions. In this sense, these factors were shown to successfully recover the corneal thickness in a rabbit model [33, 34].

Moreover, it has been demonstrated that the Wnt/β-catenin pathway is also related to CEC proliferation. Specifically, through the expression of WNT10B, which activates the Cyclin D1 that is implicated in HCEC proliferation [35]. Therefore, the incorporation of molecules that activate this pathway, such as IL-1β or a GSK3β inhibitor, are also used to culture HCEC [35].

Although culture components are indispensable for the correct formation and expansion of CEC, the culture methodology is also a very important factor to consider. Cells can be cultured in adherent or in suspension conditions. Most cell types are cultured as a cell monolayer in adherent culture, using treated culture plates to enhance its adhesion and proliferation. Cells can also be cultured in suspension conditions, which forms a 3D cell culture system that mimics the physiological tissue environment and permits a higher cell-to-cell interaction. It has been demonstrated that the formation of 3D aggregates in suspension culture of stem cells resembles an early-embryogenic process, known as gastrulation, which permits the formation of a more primitive stage of differentiation and, subsequently, higher differentiation potential [36].

The main cell source for the formation of large number of HCEC is primary cells or differentiated stem cells. Primary cells are isolated from human cadaveric donors and expanded in vitro. These cultured cells are able to proliferate with signaling molecules, which can then be expanded to reach a sufficient cell number, allowing direct CEC transplantation [37–39]. However, the limited availability of this cell source limits its clinical use. For this reason, human stem cells from diverse origins have been evaluated, such as pluripotent stem cells (PSC), corneal stroma, skin, umbilical cord, bone marrow, adipose tissue and dental pulp. In the following sections, we will briefly describe how these cells from other sources can be used to obtain CEC.

CEC differentiation

Pluripotent stem cells

Different protocols have been used for the formation of CEC from PSC such as embryonic stem cells (ESC) or induced Pluripotent Stem Cells (iPSC). For instance, Chen et al. [40] generated CEC-like cells from ESC using two different conditioned culture medium. This process was regulated by the TFG-β2 signaling pathway. Their results showed that CEC-like cells presented the typical hexagonal morphology and were able to express CEC-specific markers such as N-cadherin, Na⁺/K⁺-ATPase, ZO-1 and vimentin at protein and RNA level [40]. Other study generated CEC-like cells in two phases [41]. First, ESC was cocultured with corneal stromal cells in a culture medium with EGF and bFGF and generated pericocular mesenchymal precursors (POMPs). Second, POMPs were differentiated into CEC-like cells using a conditioned medium. Their results showed that cells expressed CEC markers N-cadherin, FoxC1 and Pitx2, which are crucial in ocular development. Furthermore, CEC-like cells were transplanted and could restore corneal transparency in rabbit corneal endothelium dysfunction models [41]. Moreover, a two-step protocol has also been reported to derive PSC into CEC [42, 43]. For this purpose, Smad inhibitors or small molecules such as a GSK3β inhibitor were used in an initial step, which induced cells to differentiate into neural crest, which is a precursor state similar to the natural formation of CEC. Then, cells were cultured in presence of FGF-2, among others, and were able to generate CEC-like cells. Differentiated cells were similar to native CEC, with a hexagonal morphology and expressing typical markers as ZO-1, Na⁺/K⁺-ATPase, COL8A1 and COL8A2, among others [42, 43]. A recent study was able to generate corneal endothelial cell substitute cells from iPSC (named as CECSi cells) using a direct differentiation protocol [44]. Their differentiation medium contained IGF-1, among others, and iPSC were able to differentiate into CECSi cells in 11–14 days. These CECSi cells exhibited a hexagonal morphology and an increased expression of CEC markers such as Na⁺/K⁺-ATPase, ATP1A1, ZO-1, N-cadherin and Pitx2. Moreover, differentiated cells with ROCK inhibitor were transplanted into a monkey corneal edema model and a recovery of the edema was showed [44].

Bone marrow-derived endothelial progenitor cells

Human limb fetal bone marrow has also been used as a source for the generation of CEC-like cells [45]. For this purpose, bone marrow-derived endothelial progenitor cells were transdifferentiated into CEC using a coculture system with fetal CEC with CEC conditioned medium. Transdifferentiated cells expressed aquaporin 1 (AQP1), a CEC marker, and formed polygonal cells. Transplanted cells of a cat model with a corneal endothelium defect recovered corneal transparency after 28 days [45].

Corneal stroma precursors

Another study evaluated the use of stem cells from the corneal stroma, known as Corneal-precursors, to differentiate into CEC. Previous work demonstrated that cell differentiation was achieved based on the upregulation of Pitx2, which occurred in the presence of retinoic acid and a GSK3β inhibitor [46]. Differentiated cells were transplanted into rabbit corneas,
allowing the maintenance of its transparency and thickness for 8 days [46].

Adipose-derived stem cells

A protocol has also been described for obtaining CEC-like cells from autologous adipose-derived stem cells (ADSC). This was performed using an initial step of cell reprogramming and the coculture of ADSC with rabbit CEC. Their results showed that CEC-like cells were positive for CEC markers CD31, AQP-1 and ZO-1 [47].

Skin-derived precursors

The dermis contains a population of autologous stem cells that can be easily isolated through a minor surgery. Inagaki et al. [48] demonstrated that skin-derived precursors (SKP) from facial skin had the ability to differentiate into CEC-like cells. The differentiation was performed by the activation of Smad2 and Wnt/β-catenin pathway by supplementing the culture medium with, among others, a GSK3β inhibitor and with TGF-β2. Differentiated cells presented a polygonal morphology and were positive for specific CEC markers ZO-1 and Na+/K+-ATPase. Moreover, pump function in CEC-like cells was higher compared to SKP and 3T3 cells. In vivo analysis in a rabbit model of bullous keratopathy revealed that differentiated cells could maintain corneal transparency and thickness [48].

Umbilical cord stem cells

Umbilical cord stem cells (UCSC) represent a new allogenic alternative for regenerative medicine and tissue engineering. UCSC cultured with a GSK3β inhibitor showed the ability to differentiate into CEC-like cells. These cells presented typical characteristics like polygonal shape and the expression of CEC markers (ZO-1, Na+/K+-ATPase and Pitx2). Furthermore, CEC-like cells transplanted into a rabbit model of bullous keratopathy revealed that differentiated cells could maintain corneal transparency and thickness [49].

Dental pulp stem cells

Dental pulp tissue is gaining interest due to the regenerative ability of the dental pulp stem cells (DPSC). DPSC can be easily isolated from third molars and represent an autologous cell source. A two-step protocol showed the ability to differentiate DPSC into CEC-like cells. This process was performed through the activation of the PI3K/Akt pathway and the differentiated cells presented the characteristic polygonal-like shape together with an increased expression of typical markers (ZO-1, Na+/K+-ATPase and Pitx2). Furthermore, CEC-like cells transplanted into a rabbit model of bullous keratopathy were able to restore corneal transparency and thickness [49].

Materials to promote in vitro CEC tissue engineering

Materials

While the cells previously described are mandatory for the proper reconstruction of the corneal endothelium, biomaterials may provide an adequate environment for cells to perform their function. In general, the number of combinations and types of materials is high, which provides an elevated number of biomaterials that can be used for tissue regeneration and substitution. Depending on the targeted tissue, the type of materials as well as the properties need to be tuned. Not only there are different types of biomaterials available, but a combination of different biomaterials is as well possible, providing an unlimited number of possibilities for tissue engineering. Furthermore, their properties can be further enhanced by incorporating different nano- and micro-size morphological features in the materials as well as relevant chemical cues, such as growth factors or molecules that can be released at the desired time point.

Due to the inherent properties previously described for the corneal endothelium, the main types of materials that can be used are described below.

Polymers

These materials have been extensively used in tissue engineering due to their excellent properties. They can be chemically functionalized and present unique characteristics as biodegradation, biocompatibility and intrinsic biological domains. They can be classified depending on their origin, a synthetic or natural source [51].

Synthetic polymers

These polymers are made of diverse monomers of different lengths. There is almost an unlimited number of synthetic polymers that can be designed depending on the needs. For this reason, they offer the required mechanical and physicochemical properties as tensile strength, stiffness and biodegradation. Normally, synthetic polymers are cheap as they can be easily fabricated in large quantities. Some forms of poly(lactic acid) (PLA) and poly(lactic-co-glycolic acid) (PLGA) have been used in approved products for specific applications by the US Food and Drug Administration (FDA), and therefore could be promising materials to be used in the future as substrates for corneal tissue engineering [52–54].

Natural polymers

These materials are present in organism membranes and in ECM and include proteins and polysaccharides. Due to their natural origin, they offer great advantages like biocompatibility and binding domains. These domains help in cell attachment and differentiation as they recognize the natural substrate [55]. For corneal regeneration, collagen is the most important protein as it is the principal component of the cornea. All in all, diverse natural scaffolds have been used in this field [55].

Comparing synthetic and natural polymers, there are two main aspects that distinguish both of them. On the one hand, from a material point of view, the composition and properties of the synthetic polymers can be controlled to a much higher extent compared to the natural polymers. Natural polymers may suffer from batch-to-batch variations, which may alter the final properties of the product. On the other hand, regarding the biological response, synthetic polymers are generally more inert, although strategies can be used to modify or functionalize the surface,
whereas natural polymers have the needed amino acids and proteins required for proper cell attachment and cell guidance, being the possible adverse effect the immune response.

**Composites**
The union of a minimum of two distinct materials forms composites. This combination results in an enhancement of physical or biological properties. For example, collagen and chitosan together have been determined to enhance the optical transparency and mechanical strength in membranes for corneal tissue engineering [56].

**Overall morphology**
Besides the biomaterial composition, different morphologies of the final biomaterial determine the interaction with the surrounding tissue and the function to be achieved.

**Scaffolds (porous and fibrous)**
Three-dimensional scaffolds are the most commonly used structures for tissue engineering. There are several requirements for an ideal scaffold for tissue engineering, which are mainly to provide similar mechanical properties to those of the native tissue, to degrade while the new tissue is being formed and to present sufficient porosity allowing cell infiltration. These scaffolds can be fabricated by several methods, including freeze-drying, solvent casting, solid freeform fabrication or phase separation, among others [51]. A special type of preparation is electrospinning, which allows forming thin layers of nanofibrous materials, mimicking the fibrillar ECM structure [57]. The latter method is of special interest in cornea regeneration as the volume required for CEC culture is relatively low, as only a thin membrane is needed.

**Hydrogels**
Hydrogels are polymeric materials that are able to entrap high amounts of water within the polymeric network. The main advantage of hydrogels is that they possess injectability, which is the ability of the materials to be injected into the site of defect [58]. Based on this latter property, the risk of infection in surgery is reduced due to the limited invasiveness of the technique. Furthermore, these materials are prepared at low temperature, which are then able to gelify after injection upon contact with body temperature. This ability further allows hydrogels to encapsulate and deliver biologically active molecules as well as cells [58, 59]. The most commonly used hydrogels are based on polysaccharides and other natural proteins. Their degradation rates are generally fast but can be strategically modified to tune their degradation rates to match with the formation of new tissue.

Figure 5. Schematic representation of CEC production depending on the initial pluripotent status of cell source. On top, ESC differentiated into CEC; in the middle, somatic cells (e.g. fibroblasts), can be reprogrammed into iPSC, which can be differentiated directly into CEC; at the bottom, MSC can be differentiated directly into CEC or using a two-step protocol. CEC, corneal endothelial cells; DPSC, dental pulp stem cells; ESC, embryonic stem cells; iPSC, induced pluripotent stem cells; MSC, mesenchymal stem cells; NCSC, neural crest stem cells.
Microparticles
Microparticles (MP) are spherical-like particles in the range of tens of microns up to several hundreds of microns that combined with an adequate hydrogel, can be injected in the desired site. These MP are generally used either as microscaffolds onto which cells are able to grow to create a tissue or as systems to allocate molecules and allow the delivery of biologically relevant molecules [60].

Nanoparticles
In a similar way to MP, nanoparticles (NP) present unique properties based on their nanosizes. Their range of sizes down to the nano level confers them high-specific surface areas with electrostatic charge that allows their penetration into cells [61]. These are generally synthesized by chemical routes and their size ranges between 20 and 100 nm. In the case of cornea, these particles are generally made of polymeric and ceramic materials.

Biomaterials for producing scaffoldless CEC sheets
Substituting the corneal endothelium consists merely on substituting the thin layer of CEC found in the cornea. With this in mind, one plausible strategy has been the culture of cells in such environment that will enhance the cell-to-cell contact and force the formation of a single-cell layer. This single-cell layer can have the required properties in terms of handling and manipulation for surgeon to implant, as well as the biological requirements needed, as the cells will be maintained alive and hence allowing to perform the previously mentioned functions. For these cell layers to be adequately formed and to allow the proper phenotypic expression of the different markers as well as to easily manipulate the obtained cell sheet, biomaterials have appeared as candidate to enhance the cell-to-cell contact and contribute to the cell sheet formation. In this sense, most of the biomaterials used for this purpose are based on thermosensitive materials that, upon change of temperature, have a transition in the surface properties, transitioning from a hydrophilic material to a rather hydrophobic material that induces cell detachment [62]. Other methods to allow the detachment of cells are based on chemical methods, such as trypsinizing treatment, but their use may limit the maintenance of the cell sheet as well as may limit the phenotypic markers required.

One of the most commonly used biomaterials for cell sheet production is the material that can be thermally stimulated for the detachment of adherent cells. In this sense, poly(N-isopropylacrylamide) (PNiPAAm) has been considered as an excellent material for this purpose, as it allows the transition from hydrophilic to hydrophobic based on the temperature (Fig. 6) [62, 63]. Taking this material as the gold standard for cell sheet production, several works have been proposed with two main purposes: to establish a different temperature to allow the detachment of the cell sheet or to functionalize the surface of the thermoresponsive material to guide cells better into the desired phenotype. Previous works have reported that the temperature at which the transition takes place can be modulated by the introduction of certain copolymers or specific domains that can alter the transition temperature [64]. In this sense, in previous works, PNiPAAm was paired with poly(vinyl methyl ether)-based polymer, allowing not only the change of temperature switch but also provided thickness, stiffness and swelling behavior. CEC were seeded on the surface followed by the detachment of the cell sheet, showing specific CEC markers such as ZO-1 and Na⁺/K⁺-ATPase functionalities before and after the transfer [64, 65]. The combination with other polymers has also been shown to be possible, such as poly(N-N'-dimethylacrylamide)) [65].

Collagen has generally been used as a biomaterial to allow the culture of CEC on top while allowing its further extraction as a cell layer [66]. The combination with other polymers such as diethyleneeglycol methacrylate allowed tuning the temperature for detachment at around 32°C [67]. These cell sheets can be then implanted in rabbit corneas that have previously been denuded of the endothelium, showing that the clarity was gradually increased in the cell sheet group compared to the control groups [68]. In some cases, temperature could decrease up to 20°C and still allowed the detachment of the cell layer with no adverse behavior [69]. An important milestone of these thermo-responsive substrates is to culture them under serum-free conditions to allow its clinical translation [70, 71]. In this sense, thermo-responsive polymers were used with different ECM coatings to allow cell attachment. Interestingly, the type of CEC used had a significant effect on cell behavior, showing that immortalized heterogeneous CEC were less prone to attachment than an immortalized clonally grown CEC line (HCEC-B4G12). It was shown that this cell line was able to produce high levels of vinculin, which allowed having a strong adherence on the substrate even in the absence of serum and without previous conditioning with ECM molecules [70, 71].

Nanotechnology has as well emerged as a promising tool to allow the implantation of cells into the site of defect without the need of a scaffold. In this sense, human umbilical cord blood endothelial progenitor cells were labeled with CD34 immunomagnetic NP. These NP can be strongly attracted by a magnet. The combination of cells labeled with NP allowed to recapitulate the cells with NP and repair the corneal endothelium defects by simply placing the magnet in the anterior chamber [72]. Positive results have been obtained in terms of allowing substituting the endothelium of rabbits, maintaining good levels of intraocular pressure without any signs of corneal edema [73].

Substrates for CEC implantation
Besides the previously described cell sheet production, CEC implantation has been ameliorated with the use of several matrices that can provide a proper environment for cell attachment, proliferation and differentiation. Furthermore, these biomaterials themselves are also able to elicit a positive response in the integration and regeneration of the surrounding tissue. For this purpose, different matrixes with different complexities have been used (Fig. 7). On the one side, decellularized matrixes, which take advantage of a decellularized DM that is able to maintain the native architecture of the CEC environment. On the other side, the use of synthetic matrixes based on biomaterials, like polyvinylidene fluoride (PVDF) or polyvinyl alcohol (PVA), has been increasingly used in CEC implantation [74, 75]. Initially, most matrixes were designed as partially inert allowing only the placement of cells, considered as ECM mimicking matrixes, although their designs have evolved to present higher functionality, giving rise to the advanced mimicking ECM biomaterials. In this section, different examples of the current material used will be described.

Decellularized matrixes
Commonly used materials in tissue engineering are the decellularized tissues. These types of tissues take advantage of tissues from cadaveric origin or from other animal species, submitting
the tissues to several processes to remove the cellular content while maintaining the architecture of the native structure. These have shown adequate results for various tissues, demonstrating that these can be used as proper scaffolds for cell culture, allowing cell proliferation and maintaining the desired phenotype [76]. The corneal endothelium has also allowed such success, taking advantage of animal corneas, mainly from bovine origin. The posterior lamellae of the cornea can be decellularized by established protocols involving several processing using enzymes as well as detergents. CEC can then be seeded on the DM allowing the formation of a CEC polygonal monolayer, showing positive staining for typical CEC markers [77]. These structures can then be tuned depending on the needs and on different methodologies to improve CEC behavior. In this sense, a bilayered structure was built using an acellular porcine cornea matrix, which already contained good optical clarity, enough toughness for surgical procedures needed, as well as the compatibility needed for tissue engineering [78]. For this purpose, a full thickness cornea substitute...
was fabricated combining limbal epithelial cell-like cells with CEC. Cells were cultured on each side of the scaffold, creating 3–4 layers of the epithelium and a uniform monolayer of CEC, showing similar densities to native tissues. Immunofluorescence showed that CEC were able to express N-Cadherin, ZO-1 and Na⁺/K⁺-ATPase, which was then correlated with a gradual increase in transparency after an 8-week follow-up postimplantation in rabbits [78]. Similarly, other work used the human anterior lens capsule extracted during cataract surgery, which was then exposed to enzyme digestion and allowed the seeding of CEC on top of it [79]. The results showed that CEC grew in confluence similar to that of native tissue, showing strong positive staining for ZO-1, Connexin-43 and Na⁺/K⁺-ATPase. Hence, this was considered as another plausible scaffold for the ex vivo expansion of CEC which will ultimately allow maintaining intact the barrier function and the ionic pump functions [79].

The use of xenogeneic substrates is a plausible approach for corneal endothelium regeneration, although their long-term effects and the presence of remaining cellular residues from the original donor bring controversy on their real clinical use [80]. In order to avoid the use of xenografts, allografts have as well been proposed as substrates for decellularization. In this sense, taking advantage of the discarded corneas of eye donors, these could be digested by enzymatic digestion, allowing the culture of CEC for 14 days. Results showed that cells were functional having a new source of high-quality corneal tissue for transplantation [80]. The main benefit is the presence of ECM-based proteins that are able to induce the binding of CEC to a proper substrate. It was previously described that the use of COL I, COL IV and fibronectin (FN) allowed enhanced adhesion, whereas laminin did not seem to enhance in such manner the cell adhesion and proliferation. For these reasons, providing environments similar to those of native tissues, either using native decellularized tissues or combinations of different biomaterials is essential for a proper regeneration [81]. Collagen has been found as one of the molecules that has enhanced CEC behavior and should hence be considered as a substrate for CEC proliferation and differentiation [82]. Due to the great similarity of other membranes, previous studies also used amniotic membranes as possible carriers for CEC culture and transplant showing positive results with similar results to that of the control [38, 83].

**ECM mimicking biomaterials**

Initially, materials based on common polymeric materials were used to see the effect of the material composition on CEC behavior. These type of polymers do not fully resemble the natural ECM, although can be easily functionalized on the surface to endow cues that might be positive for CEC culture. For instance, polymers such as Poly (DL-lactide-co-glycolide) (PDLGA), which is a combination of PLA and glycolic acid, have been previously shown to have beneficial effect on CEC [79]. Interestingly, as the amount of PLA content increased in the samples, the biocompatibility of the membranes decreased, which was probably related with a higher degradation rate and hence a higher acidic pH in the culture media which eventually reduced cell viability [73]. In general, previous studies have tested different synthetic polymers with different degree of hydrophobic properties. In this sense, PVA, poly(ethylene-co-vinyl alcohol), tissue culture polystyrene (TCPs) and PVDF were tested using CEC [73, 74]. It was found that relatively low hydrophobic polymers, such as PVA, inhibited CEC attachment, whereas the more hydrophobic polymers allowed its attachment. Lower hydrophobic polymers showed the tendency to have rather fibroblastic cell morphologies after 3 weeks of culture, whereas the most hydrophobic substrate, PVDF, showed the maintenance of typical hexagonal shape, together with higher expression of CEC-related markers. Interestingly, it was hypothesized that the degree of hydrophobicity found in PVDF enhanced cell-material interaction, stimulating cells to deposit COL IV, establishing a favorable microenvironment for CEC to proliferate and differentiate [74].

Different combinations of synthetic polymers with natural polymers have been performed. For instance, biodegradable poly-caprolactone (PCL) substrates and natural polymeric materials such as chitosan were tested. Interestingly, CEC could not attach and proliferate on chitosan, whereas some degree of attachment was found on PCL [84, 85]. For this purpose, combinations of blends were prepared to analyze the effect of combining a natural polymer with a synthetic one [84]. Interestingly, despite PCL has little sites for cell attachment, as the PCL content was increased, the degree of cell adhesion and proliferation increased. Not only did the proliferation increase, but so did the markers of functionality, mainly N-Cadherins and ZO-1, combined with an hexagonal morphology [85]. The enhanced behavior was ascribed to several factors related with the biomaterials, mainly the composition and the topography. In this sense, the blend presented a higher topography than the chitosan by itself as well as the presence of the carbon-oxygen (C=O) bond of PCL, which resulted in higher protein expression of specific markers in the membrane compared to the TCPS, providing a favorable environment in terms of ECM for the culture of CEC [84, 86, 87].

Another promising synthetic polymer that has shown excellent transparency is poly(ethylene glycol) (PEG). This polymer can be manipulated to obtain 50 μm thin films with similar tensile strengths to that of human corneal tissue and with transparency higher than 98% [88, 89]. Furthermore, these types of substrates allowed the proliferation of CEC obtaining a 100% confluent layer after 7 days, allowing its implantation into a sheep corneal model (Fig 8). Despite the results showed positive results, further biological activity of the biodegradable systems should be sought in order to allow a better cell-material interaction [88, 89]. For this purpose, other combinations of synthetic polymers with natural polymers have combined PLA and cross-linkable gelatin [90]. For this combination, a spin coating technique was used, where droplets of the samples are placed on a rotating device which is then spined to allow centrifugal forces to create a thin coating. For this purpose, poly(d, L-lactide-block-acrylic acid) and different types of gelatin were used to form multilayered structures to allow the culture of CEC. The technique allowed the formation of ultrathin layers (lower than 1 μm) with high transparency (over 90%) and even showing similar behavior to control in term of in vitro behaviors. Hence, this methodology shows a plausible substitute for natural corneas [90]. Although synthetic polymers and composites are having promising results, in vivo studies are still required.

Rather than using synthetic grafts, some researchers have taken advantage of natural polymers such as collagen. Collagen has been shown to have adequate behavior when CEC were cultured and these were subsequently transplanted [39]. Other natural polymers such as hyaluronic acid have as well been used for corneal endothelium as a strategy to mimic the native tissue due to their presence in natural endothelium [91]. Nevertheless, collagen has been generally combined with other polymers to allow maintaining stability, since their degradation is generally fast. In this sense, the formation of interpenetrating networks with other polymers such as chitosan or sodium hyaluronate have been previously used, showing good transparency and excellent behavior.
of CEC seeded on top of the substrates. Similarly, chitosan has been combined as well with gelatin and chondroitin sulfate [92–94]. Despite its degradation can be observed after 5 months of implantation, the results are promising to continue exploring these blends of materials [95, 96]. Interestingly, despite results combining gelatin and chitosan have shown adequate results, blending gelatin with chitin has shown enhanced properties, presenting milder immune response although with a faster degradation, having full degradation in <8 weeks [97]. Chitosan was previously cross-linked with glutaraldehyde to increase the degradation rates and the mechanical properties [98]. For this purpose, collagen vitrigel, which is mainly a collagen sheet with transparent properties, was obtained using porcine atelocollagen and UV irradiation, allowing the culture of CEC on the surface. The vitrigel was given a spherical morphology to mimic to a higher extent the native tissue, showing that their consistency was better compared to the use of flat sheets once implanted in an animal model [99]. The vitrigel membrane together with CEC were transplanted into a rabbit model, showing that after 14 days, the CEC completely covered the grafts after this period of time, expressing high levels of ZO-1 and Na⁺/K⁺-ATPase. These results were significantly better than the control groups in the absence of transplanted CEC [100]. Similar to vitrigel, other strategies based on collagen have prepared membranes by compressing collagen

Figure 8. Procedure performed on live sheep corneas (a) before and (b) after insertion of the poly(ethylene glycol) hydrogel film (PHF) into anterior chamber. Images from (c–h) show evolution of transparency of control and implanted PHF, together with H&E staining of (i) control and (j) PHF after 28 d implantation. Adapted with permission from Ref. [89].
hydrogel until a thin layer is formed, showing as well excellent characteristic of the seeded cells on top. Nevertheless, the transparency of the membrane still remains a further issue to assess [101]. In order to overcome issues with transparency, gelatin, which is denatured collagen, can be easily processed, which upon mixing with heparin, allows an excellent tuning. In this sense, the mechanical properties could be increased by adjusting the heparin ratio as well as the crosslinking of the gelatin or using dehydrothermal processing, as well as changing the molecular weight and the properties of the gelatin [102–105]. Similarly, the mechanical properties of the compressed collagen can as well be improved by combining the sheets with PLGA aligned nanofibers [106]. Furthermore, collagen has been shown to be still distant from the gold standard, which is the use of human anterior lens capsules. These membranes were compared with bioengineered collagen sheets of different thickness, mainly 20 and 100 μm. The results showed that the collagen membranes behaved in an adequate manner but were still distant from the optimum control. The membranes were either too soft for the proper placement into the site of defect or too rigid to allow introducing the membranes within the keratoplasty. Nevertheless, the culture of CEC in vitro behaved in an excellent manner, showing proper morphology and adequate phenotype expression [107]. Hence, for further clinical applications, these materials still require improving the mechanical properties for optimal implantation.

Advanced CEC mimicking biomaterials

**Molecule eluting biomaterials**

Cells are able to communicate among them through the release of molecules that are sent as signaling cues. These molecules are able to elicit changes in the phenotypes of the receiving cells, affecting their phenotype and ultimately activating regeneration pathways. Taking into account that most of the materials used in CEC tissue engineering are based on water-soluble polymers, the previously mentioned molecules can be entrapped within these matrices to allow a sustained release over time and simulate the desired phenotypes and regeneration pathways. The excellent water adsorption capacity of gelatin and the fine-tuning when combined with heparin allowed the fabrication of membranes that encapsulated growth factors, as bFGF that was shown to combine with heparin, allowing the fabrication of membranes water adsorption capacity of gelatin and the fine-tuning when desired phenotypes and regeneration pathways. The excellent previously mentioned molecules can be entrapped within these CEC tissue engineering are based on water-soluble polymers, the effecting their phenotype and ultimately activating regeneration able to elicit changes in the phenotypes of the receiving cells, affecting their phenotype and ultimately activating regeneration pathways. Taking into account that most of the materials used in CEC tissue engineering are based on water-soluble polymers, the previously mentioned molecules can be entrapped within these matrices to allow a sustained release over time and simulate the desired phenotypes and regeneration pathways. The excellent water adsorption capacity of gelatin and the fine-tuning when combined with heparin allowed the fabrication of membranes that encapsulated growth factors, as bFGF that was shown to increase the proliferative capacity of cells, and were able to release them in a sustained manner. Most importantly, cells cultured on the membranes presented adequate surface markers for CEC and had enough rigidity for manipulation and implantation [104]. Hence, this provides evidence on the use of drug-eluting biomaterial implanted in the cornea rather than the direct placement of cell sheets or cell-loaded scaffolds [104]. Related with the release of molecules, dorzolamide hydrochloride was coated on the surface of PCL-based materials to release it once implanted in rabbit eyes, showing a decrease on the intraocular pressure [108, 109]. Furthermore, these materials were shown to be biocompatible with CEC, although no understating of cellular markers was given. Optimization of the membrane should be performed in order to obtain near-order kinetics rather than burst release. This could be mitigated by actually incorporating the drugs within the scaffolds rather than performing coatings [110]. Similar study was performed using gelatin cross-linked with a carbodiimide, which depending on the degree of crosslinking, allowed the release of the loaded drug within the initial 8 h [110]. Once again, the soaking of the drug within the surface reduces the sustained delivery and hence strategies to provide a prolonged release are needed. In a more sophisticated system, microspheres made of biodegradable PLGA incorporating ROCK inhibitor Y-27632 were prepared using gelatin in a double emulsion, which were injected into the anterior chamber in rabbit eyes [111]. ROCK inhibitor has been proposed as a treatment in clinical scenario, although patients still require the use of immunosuppressive treatments during a long period of time to avoid any immune response [112]. The release of the ROCK inhibitor allowed the enhancement of CEC proliferation in vitro and the release rate could be controlled by tuning the amount of lactic and glycolic ratio. Despite of the positive results in vitro, the release was shown to be complete after 10 days at most, which limits further outcomes of the described microspheres [111, 112]. Hence, other encapsulation strategies that allow a longer release must be sought as well as strategies to increase the sustained release of drugs.

**Surface modification**

Similar to drug-eluting materials, materials that are able to modulate the cell-material interaction as well as the tissue-material interaction by modifying the surface properties have as well gained significant attention in endothelium regeneration. In this sense, two commonly used strategies are based on the exposure of functionalized surfaces for cells to interact with, or the use of substrates with topographical cues. Some strategies have involved sophisticated technologies for performing topographies or the surface functionalization, whereas others are based on simple material mixtures. In the latter one, some strategies have used combinations of synthetic and natural polymers, such as blends of glycerol and silk fibroin. Glycerol may have great potential due to the transparent properties, but lacks of enough mechanical stability as well as limited biological interactions. In a similar manner, silk fibroin by itself lacks of enough transparency although has excellent biocompatibility and mechanical properties [113]. Nevertheless, during the preparation process cracks may appear, which have been shown to decrease in the presence of glycerol [114]. Hence, their combination has shown a rather uniform structure, with a reduced thickness and a rougher surface. The results demonstrated that CEC were shown to proliferate better on surfaces that had intermediate roughness with the presence of 1% glycerol in silk fibroin. The need of slight roughness for enhancing the behavior of cells is based on the improvement of protein adsorption on the surface of the substrate which inherently enhanced CEC interaction with the substrate [115].

Other strategies have coated silk fibroin with certain molecules such as COL IV, FN and chondroitin sulfate-laminin mixture, showing that the optimum results for fibroin treatment were collagen [82]. Agarose has as well been functionalized with different peptides to enhance the biological properties, showing that in general, gelatin-derived signals showed the best behavior of cultured CEC [116]. Similar studies were performed with Poly llysine as base substrate, which could as well be modified with synthetic peptides, such as RGD, to promote CEC behavior. Furthermore, their combination with natural-based proteins showed enhanced cell attachment and enhanced functionality (Fig 9) [117]. In a similar way, beta-carotenoids have as well been placed on the surface of silk fibroin to enhance the proliferation and enhancement of phenotypic response, as it is known that beta-carotenoids are able to protect patients suffering from eye diseases, due to its anti-oxidant properties [118]. The results showed an enhancement compared to the pristine silk film, mainly related to the proliferation and to the expression of CEC markers [118]. In an attempt to further increase the properties of silk, lysophosphatidic acid (LPA) was incorporated within the structure of silk fibroin. LPA is an endogenous
glycerophospholipid signaling molecule, which was previously shown to stimulate growth of different cell types as well as enhancing several biological functionalities. The results showed that the physicochemical properties were generally unaltered. Contrary to this, the biological properties of the composite containing LPA compared to pristine silk had enhanced biological functionalities, showing higher biocompatibility and higher specific markers expression [119].

In a similar manner, other natural polymers, such as gelatin, have been combined with synthetic ones to achieve enhanced properties. Nevertheless, not only is the composition important but also the morphology of the matrix is of even more relevance. The morphology of the substrates can be controlled at the macro or at the micron level. At the macroscopic level, it is relatively affordable and cost-effective to produce material sheets with optimized curvatures to mimic the curvature at which natural endothelium can be found [120]. For this purpose, a curved gelatin hydrogel sheet was prepared by hydrothermal method allowing the culture of CEC as well as its implantation in monkeys with bullous keratopathy. Interestingly, the sheets showed similar permeability to water and protein as controls, mainly atelocollagen and vitrigel sheets. An importance of the curved sheets is that there was no wrinkling of the structure after implantation, whereas the flat sheets produced wrinkles which inhibited proper cell function [120]. Moreover, patterned structures at the micro and nano level that can be created during the processing of the materials is an interesting approach to control the CEC fate [121]. For instance, gelatin methacrylate has been used to create patterned structures, showing that rather than the composition, the presence of gratings with different heights and widths had a direct effect on CEC behavior. More specifically, the gratings with 1 μm pillars with square and hexagonal morphology were optimum [121]. Despite the presence of macro structural features are interesting to modulate and conform CEC in a

Figure 9. Representative images of adhesion of CEC on poly-ε-lysine hydrogel uncoated or combined with the different proteins. Symbol * denotes significantly different P < 0.05 compared to poly-ε-lysine hydrogel only (A–H). The images were quantified to clearly observe the differences in cell adhesion at 24 h (I). Scale bar 100 μm. Adapted with permission from Ref. [117].
hexagonal manner, the morphology of the matrices at the nano level may as well have profound effect on CEC behavior. For instance, the matrices of synthetic polymers with high transparency, adhesion and signaling, the CEC fate and proliferation could be controlled in such a way. In this sense, several micro well and pits were fabricated showing that 1 μm pillars could enhance proliferation as well as ZO-1 expression, showing in general that topography is mandatory for the culture of CEC.

Since the topographical effect is inherently linked with a proper adhesion onto the substrate, topographical cues were combined with chemical cues, providing microwells and micropillars with different coatings on PDMS substrates [125, 126]. Interestingly, the results showed that not only the topographical cues are important but also the chemical cues, showing that optimum results were obtained when micropillar coated with FN and collagen mix were placed, observing highest CEC gene expression (Fig. 10). Nevertheless, further studies are needed to assess the trend using different patterns, as the results were not completely conclusive [127]. Furthermore, these studies have been performed in non-implantable materials, such as Polydimethylsiloxane (PDMS), so efforts should strive in obtaining similar results in implantable materials, such as collagen, gelatin or synthetic polymers. That is why a more applied technology using titanium as substrate, which is the same titanium used in the keratoprosthesis, showed that there was no effect on the range of topographies used, having a maximum Ra of 1.15 μm [128].

Even the use of short peptides that are able to self-assemble in aqueous media have been proposed for the encapsulation of CEC [129]. Actually, one of the main limitations of self-assembling peptides is the limited manipulation ability, which make them less viable for clinical applications. The short peptides based on Isoleucine-Valine-Lysine-Cysteine (IVKC) allowed the encapsulation and survival of CEC.

Conclusions and future perspectives

Current strategies to develop new materials from the successfully CEC transplant have been explored. The technologies previously described address two main strategies, mainly the use of cell sheets, where biomaterials are used to allow the initial attachment and proliferation to subsequently detach the cell sheet, whereas another strategy takes into account the use of biomaterials as implantable substrates for tissue engineering. All these technologies are still in an emergent status in the field of corneal endothelium, but is a promising strategy and valuable in other tissues, reaching clinical scenarios and showing promising results, even having established several companies in the field. In this sense, a significant number of companies have appeared in the last decade involving the use of biomaterials and their combinations for the regeneration of tissues [130]. In this sense, skin has for instance been tissue engineered for the production of this synthetic graft for patients with burns and similar situations, providing, in combination with autologous cells in some cases, a great combination for the faster regeneration of burned patients [131, 132]. In a similar way, cartilage tissue regeneration has been established as well combining collagen-based biomaterials with chondrocyte-like cells, which have been acquired from different cell sources, having mesenchymal stem cells (MSC) as a main source [133, 134]. As an final example, nerve regeneration, which is an extremely complex tissue, has proven the use of different cell guiding biomaterials with specific microstructures combined with different type of growth factors, such as neural growth factor, together with cells has provided optimistic results [135–137].

Cell sheets are generally based on different methodologies to control the stem cell fate or to control the ability of CEC to maintain the adequate phenotype, mainly using chemical substrates that can modulate the stem cell fate. The control over the stem cell fate has generally depended on the use of genes as well as chemical substrates that may limit further applications in real clinical scenarios. The use of cells extracted from the same patient that are cultured on substrates that allow the fabrication of cell sheets has been shown as an excellent approach for autologous transplantation. The role of stem cells and how to obtain autologous cells from the patients should be the direction in which research should be striving. These substrates have to be meticulously designed to allow controlling the appropriate detachment of cells in order to avoid affecting the cell sheet and the properties of the non-functional cells.

In a similar way, biomaterials need to be enhanced as the ideal partner for tissue engineering of the cornea. These biomaterials need to be designed and systematically studied in terms of topography, stiffness, curvatures, chemical composition or surface charge among others, which will provide physicochemical elements necessary to control the phenotype of the cultured CEC. Despite some works have shown their potential, results have been sparse and not placed together to understand the needed properties of fully functional scaffolds. Other scaffolds substrates for the adequate regeneration of the corneal endothelium such as nanofibers or NP, as well the use of drug delivery systems should be enhanced. Nanotechnology has arisen as a methodology to carefully allow directing cell behavior, which combined with drug delivery systems that elute specific amounts of signaling molecules in short periods of time can have great benefit. The eluted molecules may modulate an antibacterial response, reduce an inflammatory response and even enhance the cell response of the CEC present. Finally, new technologies such as 3D printing should be taken into account to further produce biomaterials that can be patient specific.

In this sense, rather than the use of substrates onto which cells can be cultured, injectable hydrogels have gained great interest in the last years. These can be controlled and tuned according to its chemistry and can be cross-linked in several
ways, mainly physical stimuli, magnetic stimuli or chemical stimuli. Injectable hydrogels offer a non-invasive system to allow the in situ placement of materials, which can be combined with cells or drugs to enhance the tissue regeneration. In this sense, several approaches have been made for this purpose. The use of hydrogels opens the route of cell encapsulation using hydrogels to construct patient-specific biomaterials with biological functionality using 3D printing [138]. For instance, chitosan, hydroxypropyl chitosan and sodium alginate were combined to allow the encapsulation of CEC which were then placed on a DM [139]. The CEC were able to survive as well as maintained a normal morphology, which opens a new opportunity for corneal endothelium reconstruction by in situ hydrogel formation [140]. Similarly, oxidized dextran cross-linked with adipic acid dihydrazide hydrogels were used as injectable systems. Interestingly, the time of gelation as well as their degradation properties could be controlled by tuning the degree of dextran oxidation and the concentration of both components. Results showed that cell adhesion and proliferation could easily take place, although low oxidized dextrans showed confluence in just 24 h of culture [140]. Furthermore, the release products from the dextrans did not show cytotoxicity in indirect cultures, rendering excellent biocompatibility.

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