A Fish Scale–Derived Collagen Matrix as Artificial Cornea in Rats: Properties and Potential

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Submitted: February 1, 2013
Accepted: March 31, 2013

PURPOSE. A fish scale–derived collagen matrix (FSCM) is proposed as an alternative for human donor corneal tissue. Light scatter and light transmission of the FSCM were measured and compared with human cornea, and its short-term biocompatibility was tested in a rat model.

METHODS. Light scatter was determined with a straylight measuring device, whereas light transmission was measured using a broadband absorption spectrometer. For evaluation of the biocompatibility, three approaches were used: the FSCM was implanted as an anterior lamellar keratoplasty (ALK), placed in an interlamellar corneal pocket (IL), and placed subconjunctivally (SC). Transparency, neovascularization, and epithelial damage were followed for 21 days. Morphology and cellular infiltration were assessed histologically.

RESULTS. The amount of scattered light was comparable to that seen in early cataract and the percentage of light transmission was similar to the transmission through the human cornea. Implantation of the FSCM as an ALK led to mild haziness only, not obscuring the pupil, despite the development of neovascularization around the sutures; IL placement led to a moderate haze, partly obscuring the pupil, and to (partial) melting of the anterior corneal lamella. The SC group exhibited local swelling and induration, which decreased over time. Histology showed a chronic inflammation varying from mild and moderate in the ALK and IL group, to severe in the SC group.

CONCLUSIONS. In spite of technical difficulties, it was feasible to use the FSCM for ALK, whereas IL placement led to melting of the anterior lamella. Further studies are necessary for better understanding of its immunogenicity. The light scatter and transmission data show that the first version of this FSCM is comparable to human cornea tissue in this respect.

Keywords: keratoprosthesis, artificial cornea, biocompatibility, transparency

Corneal disease is a major cause of blindness worldwide, second only to cataract. Globally, more than 10 million individuals are bilaterally blind due to corneal pathology and even more unilaterally.1 Corneal transplantation is presently the only option to restore vision in these patients. Full-thickness corneal transplantation is one of the most successful forms of tissue transplantation and has been performed since 1905. One year after transplantation, the success rate is excellent in low-risk cases (avascular corneas). However, at longer follow-up, the overall acceptance of the grafts declines, particularly in high-risk cases.2–5 The worldwide demand for human donor corneas, however, exceeds the world’s supply, especially in developing countries.6 Artificial corneal substitutes have emerged to counter this shortage and overcome the disadvantages of human donor corneas, including immune rejection. These corneal substitutes range from completely synthetic prostheses, which primarily aim to restore the cornea’s refractive function,7,8 and tissue-engineered cell-based constructs,9 to hydrogels and scaffolds that facilitate the regeneration of the host tissue.10–12 Although much progress has been realized, keratoprostheses have not reached widespread use.13 At the moment, they are too expensive and complex for routine use in developing countries, where the need for implants is highest. Alternatively, biological explants are being developed and tissue-engineered corneal epithelial cell sheets have been successfully transplanted in patients.14–15 As corneal pathology often extends beyond the epithelium and affects the corneal stroma as well, replacement of the corneal stroma is usually necessary. To replace the corneal stroma, a cell-based construct or polymer scaffold of sufficient thickness is needed.

Ideally, such a scaffold for corneal regeneration allows reepithelialization, endothelialization, and repopulation with interstitial cells and nerves, preferably all of the patient’s own origin. However, human corneal endothelium hardly regenerates. For many corneal opacities without endothelial involvement, procedures that preserve the recipient’s endothelium are
used, such as anterior lamellar keratoplasty (ALK) and deep ALK (DALK). These anterior keratoplasties result in much better graft survival than the full-thickness grafts in penetrating keratoplasty (PK).\textsuperscript{16–18} The anterior lamellar approach is therefore a logical starting point for scaffold-based corneal regeneration. In 2010, this approach was studied with a biosynthetic implant in a phase I clinical trial with 10 patients. Corneal regeneration with restored vision and sensitivity was found after 24 months of follow-up,\textsuperscript{19} proving potential of this concept. The biosynthetic implant was synthesized from human recombined collagen type I, which is an elaborate and rather expensive procedure, and despite the claimed success in this study, no new trials have started so far. Collagen scaffolds that already exist in nature may reduce the cost of fabrication and promise a sufficient resource for clinical transplantation even in developing countries. A decellularized porcine corneal matrix as a xenographic scaffold for corneal regeneration has been studied for several years by several research groups, as it closely resembles the human corneal stromal organization.\textsuperscript{12,20,21} Indeed, such porcine matrices may offer a relative inexpensive and widely available alternative to human donor corneas. Results from clinical trials are not yet available, but a first phase I clinical trial is currently running.\textsuperscript{22} Collagen matrices that are even more widely available, easier to harvest, and at lower expense, therefore definitely offer an interesting alternative, and this is the topic of our study.

Here we present the results of a matrix made from naturally occurring collagen type I and obtained from scales of the tilapia fish. Tilapia (Oreochromis mossambicus) are farmed for consumption under controlled circumstances\textsuperscript{23} and the specific size of the fish can be selected on harvesting. Their scales were rinsed with 70% ethanol and stored until use in decellularize the fish scales. The decellularized and decalcified process as developed by Courtman et al.\textsuperscript{24} were cleaned in distilled water and cellular components were removed using a four-step detergent and enzymatic extraction process as developed by Courtman et al.\textsuperscript{24} Acetic acid was used to increase pore size and porosity and nitric acid to decellularize and decalcified scales were rinsed with 70% ethanol and stored until use in sterilized PBS at 4°C.

**Methods**

**Fish Scale–Derived Artificial Cornea**

A ±250-\(\mu\)m-thick decellularized and decalcified fish scale–derived extracellular matrix (ECM), consisting of collagen type I, was used for implantation (7-mm diameter, 0.2–0.3 mm thickness, P09011001; Aeon Astron Europe, Leiden, The Netherlands) (Fig. 1).\textsuperscript{25} In short, fresh tilapia scales were cleaned in distilled water and cellular components were removed using a four-step detergent and enzymatic extraction process as developed by Courtman et al.\textsuperscript{24} Acetic acid was used to increase pore size and porosity and nitric acid to decellularize the fish scales. The decellularized and decalcified scales were rinsed with 70% ethanol and stored until use in sterilized PBS at 4°C.

**Top Pattern, Light Scatter, and Transmission**

Phase-contrast images (Axio Observer.A1; Carl Zeiss AG, Jena, Germany) were taken of the prepared fish scale–derived collagen scaffold and used to create a composition photograph in Adobe Photoshop (CS3 Extended, version 10.0; Adobe Systems Incorporated, San Jose, CA) to visualize the whole top surface. Scanning electron microscope images were taken of the cutting edge of the prepared FSCM and of the micro pattern on the top surface.

To measure forward light scatter, we chose a similar approach as is used clinically to assess the functional effect of forward light scatter in patients.\textsuperscript{29} This way, the ex vivo results can directly be compared with clinical in vivo results. We applied a psychophysical technique known as “compensation comparison” and the outcome value is the straylight parameter “s.”\textsuperscript{30} This technique is implemented in a commercial instrument (C-Quant; Oculus GmbH, Wetzlar, Germany) for clinical use, measuring the light scatter between 5 and 10 degrees, which has proved representative for the total amount of straylight (light scatter ≥1). We used this instrument, with a slight adaption, to assess forward light scatter from physical samples.\textsuperscript{31} In short, two stimuli of the compensation comparison method, straylight flicker and comparison flicker, are presented to and compared by the subject simultaneously. To exclude the influence of light scatter provoked by the observer’s eye and to measure only the light scatter caused by the matrix, the straylight flicker source itself was shielded in such a way that it illuminated only the tested sample.\textsuperscript{31} Three FSCMs (P11251101; 8.0 ± 0.8 mm [diameter], 0.25–0.35 mm [thickness]) were put on a test glass, one drop of PBS was used to prevent dehydration, and a cover glass was put on top. A black, opaque disk, with a central hole of 6.5-mm diameter, was mounted over the test glass, leaving only the FSCM visible. The test glass, with the FSCM and black opaque disk, was put in the ocular of the C-Quant device to measure the light scatter. Three measurements per FSCM were performed. A holder containing PBS without an FSCM acted as a control experiment.

To measure light transmission, three FSCMs (P11251101; 8.0 ± 0.8 mm [diameter], 0.25–0.35 mm [thickness]), placed in a sample holder and kept hydrated with PBS, were studied by a broadband absorption spectrometer, covering the visible and near infrared spectrum with a spectral resolution of 0.55 nm. Details are available from Bouwman et al.\textsuperscript{32} The setup comprised a light source (LOT Oriel Xe-arc; 300-\(\mu\)m filament; LOT-QuantumDesign GmbH, Darmstadt, Germany) and spectrometer (Andor Shamrock SR-303i; Andor Technology PLC, Belfast, UK). The light source emitted white light with a homogeneous intensity pattern in the 200- to 760-nm region, fully covering the wavelength domain relevant to the human eye. This light was guided via a diaphragm through the center of the FSCM, which was placed 15 cm from the 10-\(\mu\)m wide inlet of the spectrometer. The sensor of the spectrometer measured light only in the horizontal plane. The maximum angle at which light could enter the spectrometer was 0.004 degrees. The spectrometer measured the direct light transmission, as straylight is defined as light being scattered at 1 or more degrees. All measurements were taken relative to
atmospheric air and normalized for background light. Normalization and measurement of transmission values for empty space were performed directly before each individual measurement. A holder containing only PBS was used to correct for light absorption or scatter caused by the holder itself. The light transmission of the visible spectrum was measured in steps of 0.56-nm wavelength and compared with the total light transmission of the human cornea, using the formula of van den Berg and Tan.33

Suturing

Two human donor eyes, obtained from the Euro Cornea Bank (Beverwijk, The Netherlands), were used to test the sutureability of the scaffolds with nylon 10/0 sutures (nr. 8065 198001; Alcon B.V., Gorinchem, The Netherlands). In short, two FSCMs of 6-mm diameter were sutured into human corneas, 1 day postmortem, by placing 12 interrupted sutures in one case and continuous suture in the other, including knotting and burial of the knots. Severity of tearing due to the suturing was observed.

Animals

Eighteen male Fischer 344/DuCrI albino rats (Charles River Laboratory, L’Arbresle Cedex, France), each weighing between 280 and 336 g, were used for ocular implantation with permission of the Animal Ethics Committee of the Leiden University Medical Center. All animals were treated in accordance with the ARVO Statement for Use of Animals in Ophthalmic and Vision Research. They were given 1 week to acclimatize.

Implantation

The rats were divided into three groups of six animals, and in order to determine biocompatibility, scaffolds were placed in the anterior eye. In the first group, the FSCM, with the pattern on top, was placed as an ALK. In the second group, the FSCM was placed in an intralamellar pocket (IL), and in the last group it was inserted subconjunctivally (SC). The rats were anesthetized with isoflurane, with the addition of a topical drop of oxybuprocain and a subconjunctival injection of 20 µL bupivacain. Prior to implantation, the FSCM was soaked for 5 minutes in antibiotics (polyspectran containing gramicidine, oxybuprocain and a subconjunctival injection of 20 µL bupivacain). Prior to implantation, the FSCM was soaked for 5 minutes in antibiotics (polyspectran containing gramicidine, oxybuprocain and a subconjunctival injection of 20 µL bupivacain).

ALK was performed by trephination of the right eye using a 3-mm trephine. The anterior tissue was removed using an air bubble, lamellar dissector, and scissors. The FSCM was cut to a diameter of 5.2 mm, inserted, and attached with eight nylon 10/0 sutures (nr. 8065 198001; Alcon B.V.). The surface of the FSCM was carefully leveled to line up with the native epithelium.

For the IL group, a nonpenetrating incision was made with a 15° stab knife and a pocket was created using an air bubble and a lamellar dissector. An FSCM with a diameter of 2.7 mm was inserted and the pocket was closed with two 10/0 nylon sutures.

SC implantation was performed by making an incision with a 15° stab knife. A blunt spatula was used to create a subconjunctival pocket, into which an FSCM with a diameter of 2.7 mm was inserted; the pocket was closed with two 10/0 nylon sutures.

Dehydration of the eye during surgery was prevented by regular wetting of the eye with special eye washings (balanced salt solution, 907553; Pharmacy LUMC, Leiden, The Netherlands). Directly after surgery, 1% chloramphenicol ointment (Chloramphenicol-POS 1%, 10 mg/g; Ursapharm Benelux B.V., Helmond, The Netherlands) was applied. In all cases, the unoperated left eye served as a control and was kept hydrated with gel (Vidisca Carbogel, carbomer 2 mg/g; Tramedico B.V., Weesp, The Netherlands) to prevent dryness during surgery.

In Vivo Observation

The animals were observed in the 3 weeks following implantation. Ocular drops with corticosteroids and antibiotics (Tobradex, containing dexamethasone and tobramycin; Alcon Cusi SA, Barcelona, Spain) were applied in the ALK and IL groups, once daily during the first week and every other day during the second week. All drops were stopped at the 2-week follow-up. The SC group did not receive any drops postoperatively.

At 2, 7, 13, and 21 days after implantation, all animals were examined with a microscope to judge neovascularization, transparency, and clinical signs of inflammation, such as conjunctival redness or purulent secretion. Corneal neovascularization was numerically scored from 0 to 5, with 0 = no vessels, 1 = growth of vessels at the limbus, 2 = vessels reaching the sutures/FSCM, 3 = vessels present underneath the FSCM, 4 = vessels entering the FSCM, and 5 = vessels present throughout the whole FSCM. Transparency of the cornea was assessed using a grading scale as previously used by Hackett et al.,34 with 0 = transparent, 1 = a mild haze not obscuring the pupil, 2 = a moderate haze partially obscuring the pupil, and 3 = an opaque area totally obscuring the pupil.

Histopathological Evaluation

After euthanizing the 16 rats with carbon dioxide, the eyes were enucleated and fixated with Davidson solution (composed of glacial acetic acid, ethyl alcohol, and buffered formalin) and then dehydrated and embedded in paraffin. Sections were cut on a microtome (Leica RM2165; Leica Microsystems GmbH, Wetzlar, Germany) at 5 µm and stained with hematoxylin and eosin (HE) for histological examination. All eyes were analyzed for the organization of the epithelium, stroma, and endothelium, and scored semiquantitatively for infiltration of immune cells and ingrowth of corneal cells into the FSCM. The immune infiltrate was characterized based on morphology.

Statistical Analysis

All statistical analyses were performed using a statistical software program (SPSS for Microsoft Windows, version 17.0.2; IBM SPSS Statistics, IBM Corporation, Chicago, IL). The χ² trend test was used for assessing differences between the groups regarding corneal neovascularization and opacification, and leukocyte infiltration. Statistical significance was assumed for resulting P values less than 0.05.

RESULTS

Top Pattern, Light Scatter, and Light Transmission

The whole top surface of the FSCM was visualized with a composed phase-contrast image. The micro pattern on top of the FSCM differed per area, with roughly one quarter having a spider-web appearance with micro ridges and channels, two quarters exhibiting circular running ridges and channels, but without intersecting lines, and the last quarter consisting of spikes (Fig. 2a). SEM images of the FSCM confirmed these findings (Figs. 2b–d). The micro ridges were wavelike in shape, ±5.7 µm high, and ±3.4 µm wide. The channels in between were indeed ±30 µm wide (Fig. 3).35
The amount of straylight was measured for three FSCMs using the compensation comparison method. Log values for forward light scatter caused by the holding device were beyond the lowest measurable value for the C-Quant device (log [s] < 0.40) and were considered to be negligible. The mean light scatter of the three FSCMs was log (s) = 1.62. All results are listed in Table 1.

Direct light transmission of the visible spectrum was measured on another three FSCMs. The mean direct light transmission of the three measured FSCMs, corrected for reduction of light transmission caused by the PBS-filled holder, amounted to 89%, 97%, and 85% and showed corresponding curved graphs (Fig. 4a). The three measurements were grouped and compared with the total light-transmission curve of the human cornea (Fig. 4b). The human corneal light-transmission values were within the SEM of the FSCM light-transmission values. The mean direct light transmission of the FSCM was 90%, whereas the total light transmission of the human cornea was 91.3%.

**Suturing**

Suturing tests with the two FSCMs revealed in both methods no tearing at the suture points and the knots could be buried. On bringing in the suture needle, confined cracking occurred. Micro-shearing at the suture points of the FSCM (and not of the human cornea) was observed on tightening some of the knots.

**Short-Term Biocompatibility**

To determine the scaffold’s biocompatibility, FSCMs were placed in the cornea as ALKs, placed intracorneally (intralamellar pockets) and placed subconjunctivally.

One week following transplantation, all six rats that underwent ALK had a clear and quiet cornea, with minimal neovascularization in maximally one quadrant. After 2 weeks, the scaffolds remained in place, while neovascularization was reaching the FSCM. In five cases, a mild haze at the edge of the FSCM, not obscuring the pupil, was observed, and in the sixth animal, the haze partially obscured the pupil. Sutures started to loosen slightly in all cases. At the 3-week end point, the neovascularization extended underneath the FSCM with vessels aimed at the sutures, but not penetrating the FSCM. The haze increased slightly, not obscuring the pupil. Gradual loosening of the sutures resulted in loss of the FSCM in two cases at day 18 and 21, as the rats were allowed to move freely and touch their eyes. These cases were excluded from further analysis.

During follow-up, the eyes were stained with fluorescein to analyze epithelialization. In the ALK group, the total area of the FSCM stained positively, indicating the absence of rec epithelialization of the FSCM surface after 3 weeks.

Implantation in an intralamellar pocket was performed in six rats. The results were less uniform than found for the ALK implants. After 1 week, two cases showed no haze, two cases exhibited a mild haze that did not obscure the pupil, and the remaining two rats showed a moderate haze, partially obscuring the pupil. In four cases, blood vessels approached the sutures located near the limbus. All sutures were removed after 1 week. After 2 weeks, superficial vessels reached the FSCM. The amount of haze remained stable, while some melting of the anterior lamella in front of the FSCM was noticed in five animals. Two animals were killed at 13 and 19 days postimplantation due to too much weight loss. When the 3-week end point was reached, blood vessels reached the FSCM in the remaining four animals, with some vessels starting to invade the over- or underlying stroma. Vessels did not invade the matrix and were less dense than in the ALK group.

After 3 weeks, the corneal haziness had slightly increased in three animals, ranging from a mild haze to an opaque area totally obscuring the pupil. In one case, however, the neovascularization had diminished and remained present only in the limbal area and a moderate corneal haze was observed.

**Table 1. Forward Light Scatter Results in Log(s) With Oculus (C-Quant)**

<table>
<thead>
<tr>
<th>Measurements</th>
<th>First</th>
<th>Second</th>
<th>Third</th>
<th>Mean</th>
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<td>1.53</td>
<td>1.46</td>
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<tr>
<td>FSCM 2</td>
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<td>1.69</td>
<td>1.74</td>
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<tr>
<td>Mean</td>
<td>1.62</td>
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**Figure 2.** Composition of several phase-contrast images showing the distribution of the micro pattern with the spokes and interconnecting channels in one quadrant (a-1), the spikes in the opposite quarter (a-2), and the circular running lines in the area (a-3) between those two quadrants. Detailed SEM-images depict the transitional area between the circular lines and spikes (b), between the lines and spokes (d), and a cross section showing the multiple layers (c).

**Figure 3.** Cross section of the FSCM showing the dimensions of the circular running ridges on the surface.
Corneal fluorescein staining revealed melted areas of the anterior lamella of variable size in all four remaining cases. The six rats receiving an SC implant showed slight SC swelling postoperatively at the implantation site. The swelling disappeared after 2 weeks in four rats and lasted until the end point in two rats. Local conjunctival hyperemia was present at a low degree until the end point. Sutures were not removed, as within 1 week the sutures were overgrown by the conjunctiva and removal without damaging the surrounding area proved to be difficult.

Overall
Except for the two cases mentioned, all implanted animals were in good health, kept their weight, and showed no symptoms of any ocular infection or noticeable irritation. The FSCM flattened the recipient rat cornea due to a curvature disassociation. End point neovascularization and opacification (Figs. 5a, 5b) did not differ significantly (P = 0.46 and P = 0.19, respectively) between the ALK and IL groups. Typical cases of each implantation model are depicted in Figure 6. An overview of in vivo and histological characteristics is given in Table 2.

Histopathological Evaluation
To visualize the clinically observed changes with histology, sections of the four scaffold-containing corneas in the ALK group were cut, stained with HE, and compared with the control lateral eye (Figs. 7a, 7b). HE staining showed mild corneal edema with blood vessels and infiltrating leukocytes in the stroma around the sutures and at a lower degree under the FSCM (Figs. 7c, 7d). A few leukocytes were seen within the FSCM. The leukocytes showed a typically chronic inflammatory reaction with mainly macrophages and lymphocytes and some neutrophils. Epithelial downgrowth occurred in all cases at the border of the FSCM. At the location of the sutures and at the border of the FSCM, the epithelium showed hyperplasia and metaplasia.

HE staining of the four cases of the IL group showed infiltration of immune cells, which consisted of the typical mixture with macrophages, lymphocytes, and some neutrophils. A few leukocytes were found within the FSCM (Fig. 7c). Local edema, metaplasia, and necrotic cells of the epithelium overlying the FSCM were observed in three of the four cases (Figs. 7c, 7f). However, one of the implants had no leukocytes or blood vessels infiltrating the stroma (Fig. 7f) and the other only a few (not shown). This corresponded with the in vivo observations (Figs. 6f–j). Infiltration of immune cells was not significantly different between the ALK and IL groups (P = 0.57) (Fig. 8).

The SC implants were completely surrounded by leukocytes in all six cases, with several leukocytes invading the FSCM. The leukocyte infiltration was limited to the adjacent surrounding tissue, again representing chronic inflammation. The SC group showed significantly more infiltration than the ALK group (P = 0.04) and the IL group (P = 0.02) (Fig. 8). The tissue surrounding the implant showed edema (Figs. 7g, 7h). Compared with nonoperated control eyes, more vessels were present.

DISCUSSION

Pattern and Light Scatter and Transmission

The composed phase-contrast image of the FSCM shows that the pattern that contains the micro channels covers only a quarter and not the full surface. This is in accordance with the natural pattern of the so-called ctenoid fish scales. Nevertheless, this nonhomogeneity of the pattern does not prevent corneal stromal cells from populating the scaffold’s surface, and is perhaps even stimulating cell spread. The circular running ridges had a height and width much larger than the visible wavelengths, thereby provoking light scatter. This is in
accordance with the observations that the pattern was clearly visible under the phase-contrast microscope. The latter visualizes differences in refraction, which may cause light scattering.

The FSCM had a forward light scattering of log(s) = 1.62, comparable to the amount of scattering caused by early cataract. The direct light transmission was 90%. Even though this value reflects the direct transmission and not the total light transmission, which includes forward-scattered light as well, it is very close to the total light transmission of 91% found for the human cornea. The light-transmission curves of the FSCM and human cornea are also very similar in shape, indicating corresponding values of Rayleigh scatter (i.e., light scatter caused by particles much smaller than the wavelength of light such as molecules). The displacement along the y-axis of the light-transmission curves for the different FSCMs can be attributed to a wavelength-independent scatter, plausibly caused by the micro pattern that roughens the surface of the FSCM with ridges larger in size than the wavelength of visible light. Although special care was taken for optimum position, it could not be prohibited that each FSCM was slightly differently positioned relative to the spectrometer and therefore caused different amounts of scatter. Removal of the micro pattern from the FSCM surface will improve light transmission and especially light-scatter values, which in turn will result in better visual acuity when implanted.

**Suturing**

The suturing test demonstrated that the FSCM can be sutured successfully into the cornea, when handled with care. The scaffold was brisker and had less mechanical strength than the human cornea (or rat cornea), indicated by the confined cracking and shearing. Increasing the elasticity and strength likely will improve the ease of handling and this may be needed to make the FSCM applicable for full-thickness transplantation in humans.

**In Vivo Biocompatibility**

The rat keratoplasty model has been used to study corneal transplantation for more than 25 years. Our findings of this first in vivo study with the FSCM correspond to the results seen with allogeneic corneal transplantations in rats. All corneal implants showed good acceptance up to 1 week postoperatively (first column: [a, f, k, p]) and up to 7 days (second column: [b, g, l, q]). Neovascularization and opacification were visible at 13 days (third column: [c, h, m, r]) and gradually increased until day 21 (fourth column: [d, i, n, s]). Neovascularization is seen even more clearly postmortem (last column: [e, j, o, t]).
both groups, vessels failed to invade the FSCM, perhaps due to obscuring pupil. Increased until it reached the FSCM and sutures, but vessels failed to penetrate the FSCM. The opacity gradually increased to a mild haze (stage 1) in the ALK group and a moderate haze (stage 2) in the IL group. Total opacification (stage 3) of the FSCM, indicative for a complete rejection, was observed in only one animal of the IL group. It is important to note that in humans with a nonvascularized cornea, a non-HLA–matched allogeneic corneal transplantation has a high acceptance rate, whereas in rats, this situation leads to rejection after 5 to 15 days. A rejected graft is recognized by complete corneal atrophy and opacification and infiltrating T cells on histology. The main local abnormalities that occurred in the corneal epithelium are largely explained by mechanical irritation, which leads to corneal damage and subsequent leukocyte infiltration.

The infiltration of leukocytes was mild to moderate and consisted predominantly of macrophages, lymphocytes, and some neutrophils, indicative of a chronic immune reaction. The leukocytic infiltration was primarily aimed at the sutures in the ALK group and surrounded the scaffold in the SC group. The SC group showed the presence of fibroblasts aligned around the scaffold, similar to a mild foreign body reaction. It is unlikely that this infiltration and swelling is caused by pathogens present on the FSCM prior to implantation, as FSCMs from the same batch were tested and found to be negative for bacteria and fungi. Furthermore, the collagen of the FSCM or residues of chemicals used for decellularization and decalcification could have provoked the immune response. A strong argument to support the notion that the inflammation was due to surgical trauma and mechanical irritation is the complete absence of inflammation in one case. The inflammation had been due to a heterologous protein, one would have expected an immune response. The fibril density of the FSCM or by yet to be identified antiangiogenic properties.

The amount of neovascularization was comparable to that seen in allogeneic corneal transplantations in rats and humans with a nonvascularized cornea, a non-HLA–matched allogeneic corneal transplantation has a high acceptance rate, whereas in rats, this situation leads to rejection after 5 to 15 days. A rejected graft is recognized by complete corneal atrophy and opacification and infiltrating T cells on histology. The main local abnormalities that occurred in the corneal epithelium are largely explained by mechanical irritation, which leads to corneal damage and subsequent leukocyte infiltration.

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<table>
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<th>Follow-Up, d</th>
<th>FSCM Lost</th>
<th>Corneal Vessel Score, d 21</th>
<th>Opacity Score, d 21</th>
<th>Excluded on Histology</th>
<th>Corneal Epith. Atrophy</th>
<th>Corneal Epith. Hyperplasty</th>
<th>Corneal Epith. Metaplasia</th>
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Other collagen matrices used for corneal regeneration, such as the biosynthetic artificial cornea, are already being used in clinical studies, following a decade of preclinical research. The novelty and advantage of the FSCMs are their wide availability, low cost, and simple manufacturing process. However, we are at an early phase in our research and working on overcoming technical challenges so as to develop a useful prototype.

**Conclusion**

This systemic study of the physical and biomedical short-term effects of the FSCM as corneal replacement, demonstrates its potential for future use. The availability of an easy obtainable and biocompatible FSCM can have a high impact on decreasing the shortage of donor corneas and make lifelong use of...
Immunosuppressive medication redundant. The potential of this FSCM is demonstrated for the first time by an adequate light transmission, reasonable light-scattering values, and ability to be used in keratoplasty. Future studies with a curved and thinner FSCM are necessary to prevent mechanical irritation and increase the understanding of its immunogenicity. Long-term in vivo studies and studies on modified FSCMs, for instance with the top pattern removed, are needed to develop and optimize this readily available collagen matrix as an artificial cornea.

Acknowledgments

The authors thank Jos J.M. Onderwater, BSc, and A. Mieke Mommaas-Kienhuis, PhD, of the Department of Molecular Cell Biology, LUMC, for their much appreciated work and help on the scanning electron microscopy. We thank Sarah J. Sparks, MSc, of the Department of Dermatology, LUMC, for all the additional sectioning and staining and Steven H. Cuylle, MSc, of the Leiden Observatory for the light transmission experiments. Supported by a grant from Agentschap.nl. Agentschap.nl had no involvement in study design, collection, analysis or interpretation of data, nor in the writing and decision to submit the paper. Aecon Astron Europe B.V. provided the fish scale-derived matrices.

Disclosure: T.H. van Essen, Aecon Astron Europe B.V. (F); C.C. Lin, Body Organ Biomedical Corporation (E); A.K. Hussain, None; S. Maas, None; H.J. Lai, Aecon Astron Europe B.V. (E, I, S); P. H. Linnartz, None; T.J.T.P. van den Berg, None; D.C.F. Salvatori, None; G.P.M. Luyten, None; M.J. Jager, None

References


