Plant Recombinant Human Collagen Type I Hydrogels for Corneal Regeneration

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Abstract

Purpose To determine feasibility of plant-derived recombinant human collagen type I (RHCI) for use in corneal regenerative implants

Methods RHCI was crosslinked with 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide (EDC) and N-hydroxysuccinimide (NHS) to form hydrogels. Application of shear force to liquid crystalline RHCI aligned the collagen fibrils. Both aligned and random hydrogels were evaluated for mechanical and optical properties, as well as in vitro biocompatibility. Further evaluation was performed in vivo by subcutaneous implantation in rats and corneal implantation in Göttingen minipigs.

Results Spontaneous crosslinking of randomly aligned RHCI (rRHCI) formed robust, transparent hydrogels that were sufficient for implantation. Aligning the RHCI (aRHCI) resulted in thicker collagen fibrils forming an opaque hydrogel with insufficient transverse mechanical strength for surgical manipulation. rRHCI showed minimal inflammation when implanted subcutaneously in rats. The corneal implants in minipigs showed that rRHCI hydrogels promoted regeneration of corneal epithelium, stroma, and nerves; some myofibroblasts were seen in the regenerated neo-corneas.

Conclusion Plant-derived RHCI was used to fabricate a hydrogel that is transparent, mechanically stable, and biocompatible when grafted as corneal implants in minipigs. Plant-derived collagen is determined to be a safe alternative to allografts, animal collagens, or yeast-produced recombinant human collagen for tissue engineering applications. The main advantage is that unlike donor corneas or yeast-produced collagen, the RHCI supply is potentially unlimited due to the high yields of this production method.

Lay Summary A severe shortage of human-donor corneas for transplantation has led scientists to develop synthetic alternatives. Here, recombinant human collagen type I made of tobacco plants through genetic engineering was tested for use in making corneal implants. We made strong, transparent hydrogels that were tested by implanting subcutaneously in rats and in the corneas
of minipigs. We showed that the plant collagen was biocompatible and was able to stably regenerate the corneas of minipigs comparable to yeast-produced recombinant collagen that we previously tested in clinical trials. The advantage of the plant collagen is that the supply is potentially limitless.

**Keywords** Plant collagen · Cornea regeneration · Limbal stem cells · Tissue engineering · Recombinant human collagen type I

**Introduction**

The human cornea is the transparent front of the eye that focuses incoming light onto the retina for vision. Its cellular layers comprise an outermost epithelium, a middle stroma composed of collagenous extracellular matrix (ECM) with interconnected keratocytes, and an innermost endothelium. It is heavily innervated, but avascular. Damage or disease leading to permanent transparency loss or surface distortion can result in corneal blindness. Historically, the only widely used treatment for restoring vision is human donor cornea (HDC) transplantation. However, a serious donor shortage has left an estimated 12.7 million patients worldwide waiting for corneal transplants [1]. Moreover, the outcome of conventional corneal transplantation is limited by immune rejection, ocular infection, and premature graft failure. Artificial corneas in the form of prostheses have been introduced in clinics, most notably the Boston KPro, AlphaCor, and osteo-odontokeratoprosthesis (OOKP) [2–4]. All three prostheses are associated with potentially serious side effects and are regarded as a last resort treatment for patients with severe pathology or previous conventional graft failures [5].

Given the shortcomings of conventional corneal transplantation and insufficient performance of prostheses, various research groups have focused on the development of a range of corneal replacements [6, 7].

In 2010, Fagerholm et al. published the 2-year results of a clinical trial describing the first-in-human in situ tissue regeneration of the cornea. Tissue regeneration was achieved by implanting cell-free, bio-responsive recombinant human collagen type III (RHCIII) hydrogels [8]. RHCIII was selected as a homogenous, non-xenogeneic protein that minimizes the risk of allergic reactions to animal source collagens, or zoonotic disease transmission [9]. The results of the RHCIII hydrogels are promising, but the human cornea is primarily composed of type I collagen. Here, we examine the biocompatibility and efficacy of recombinant human collagen type I (RHCI) collagen hydrogels as implants in minipig corneas, compared to syngeneic grafts.

The RHCI implants designed for this study are the first to use plant-derived type I collagen. A prior comparison of RHCI and RHCIII has been made, but both collagens were produced in *Pichia pastoris*. Protein production in yeast is a fermentation-based process that requires the additional expression of two enzymes: prolyl 4-hydroxylase (P4H) and pepsin. P4H is the enzyme that catalyzes the formation of 4-hydroxyproline amino acids from proline [10]. The 4-hydroxyproline amino acids are needed for stable triple helix formation. The recombinant pepsin is needed to cleave the telopeptides from the full-length protein before use [9]. This process is inherently limited by the fermentation batch size.

Plant-derived RHCI is generated from genetically modified tobacco plants, which allows the production of theoretically unlimited quantities of protein and supports agricultural development. The two genes encoding RHCI were co-expressed in tobacco plants with P4H and lysyl hydroxylase 3 (LH3) enzymes that were responsible for key post-translational modifications of the RHC [11]. Tobacco plants co-expressing all five vacuole-targeted proteins generated intact RHCI pro-collagen with stable triple helical structures. Tobacco plant–derived RHCI was shown to have equivalent biofunctionality to human tissue–derived collagen in other applications [11–13]. We recently reported that RHCI hydrogels were fully biocompatible with corneal cells and served as excellent substrates for corneal limbal epithelial stem cells, showing comparable efficacy to human amniotic membrane, the current gold standard for their culture [14].

Previous studies of collagen implants did not include attempts to align the collagen fibrils, although there have been numerous reports of attempts to align collagen fibrils for use as corneal implants, citing the need for transparency and mechanical strength [15, 16]. Here, we examined the properties of RHCI with aligned and random fibrils in vitro and their biocompatibility in vivo in a rat model. The RHCI hydrogel showing the most optimal properties for use as corneal implants was tested in a minipig model, where its efficacy was compared to that of allografting, simulating donor cornea transplantation.

**Materials and Methods**

The study followed the tenets of the Declaration of Helsinki and was approved by the Antwerp University Hospital Ethical Committee (EC: 14/30/319) for use of human amniotic membrane. Animal experiments in rats and Gottingen minipigs were performed in accordance with the Association for Research in Vision and Ophthalmology Statement for the Use of Animals in Ophthalmic and Vision Research, and the Swedish Animal Welfare Ordinance and the Animal Welfare Act, after ethical approval was acquired from the local Linköping ethical committee (Linköpings Djurförsöksetiska Nämnd).
Materials

All inorganic salts, enzymes, basic chemicals, and reagents were purchased from Sigma-Aldrich (St. Louis, MO, USA), unless stated otherwise. Plant-derived RHCI was provided by Collplant (Ness Ziona, Israel).

Recombinant Human Collagen Hydrogels

RHCI Hydrogels

To fabricate RHCI hydrogels with random fibrillogenesis (rRHCI), we use the protocol described in Haagdorens et al. [14]. Briefly, a 30% v/v solution of tobacco-derived RHCI in 10mM HCl was prepared by diluting with 100% ethanol. Fibrillogenesis was initiated by addition of 160 mM sodium phosphate buffer that was adjusted to pH 7.5 with 100 mM NaOH. The buffer was added at a ratio of 1:10 v/v to the original collagen-HCl volume. The solution was stirred for 2 h at 25°C; after which, water-diluted EDC and NHS were added to attain a 50 mM 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide (EDC) and 100 mM N-hydroxsuccinimide (NHS) concentration. The collagen solution was then stirred for 24 h at 4°C on a magnetic stirrer at 200 rpm. The solution was then transferred into a 50-mL tube for 6 cycles of washing to remove all excess EDC/NHS. Each cycle consisted of centrifuging the mixture at full-speed (5000 rpm, 10 min), discarding the supernatant and resuspending the collagen in 40 mL distilled water (DW). During the final wash cycle, the RHCI suspension was dispensed into a rectangular Teflon mold (4.5 × 4.5 cm) for in vitro characterization or into curved corneal-shaped and corneal-sized molds (10 mm diameter, 500-μm thick) as implants. The collagen was then air-dried under a sterile hood and immersed in 100% ethanol to promote detachment from the mold. The gels were stored in 100% ethanol until further use. Gels were rehydrated by five 2-h soaks in phosphate-buffered saline (PBS).

RHCI Hydrogels with Aligned Collagen Fibrils

For production of aligned RHCI membranes (aRHCI), shear force was applied to the collagen solution prior to fibrillogenesis and chemical crosslinking. Shear force was effected using a motorized automatic film applicator, Elcometer 4340 (Elcometer Ltd, Manchester, UK) and an Elcometer 3570 “Doctor Blade” that served as the orthogonal force. After ethanol dilution, the acidic RHCI was cast onto a glass plate that was mounted in the Elcometer. The glass plate had previously been surface-coated with a hydrophobic siloxane solution. After collagen casting, shear force was applied at a constant speed of 0.02 m/s and a thickness of 50 μm. After the RHCI had spread, the glass plate was immediately lifted from the Elcometer and immersed in a bath of fibrillogenesis buffer. This was performed very carefully in order not to disrupt the collagen membrane. Membranes were then crosslinked for 24 h in fibrillogenesis buffer and 50 mM EDC and 100 mM NHS. Crosslinked membranes were then rinsed four times in DW and peeled from the glass plate. Hydrogels were kept in 100% ethanol until further use.

Human Amniotic Membrane

Human amniotic membrane (HAM), prepared following our previous protocol, was used as a benchmark [14]. Written informed consent was collected from the donors, women undergoing planned cesarean section. Briefly, HAM was washed in an antibiotic cocktail and then flattened onto sterilized nitrocellulose filter paper, epithelial side up. The paper-supported HAM was then cut into 5cm × 5cm pieces and cryopreserved at −80°C in 50% Dulbecco’s Modified Eagle Medium (DMEM), 50% glycerol. The membrane was quarantined for 4 weeks during which time sterility and serology testing were performed to confirm safety. Prior to use, HAM was thawed, epithelial cells removed, and the cell-free membrane was washed. For use, HAM is oriented spongy layer up, after which excess water is mechanically expelled using cell scrapers. For stabilization, HAM was mounted “spongy layer” down in a CellCrown or interlockable ring [17].

Physical and Mechanical Characterization

To measure water content, samples were DW equilibrated for 24 h. Water content of samples was determined by weighing membranes that were blotted dry to remove excess surface liquid to obtain the wet weight (W0). These samples were then dried in a drying oven (60°C) for 24 h to obtain the dry weight (W). Equilibrated water content of hydrogels (Wt%) was obtained according to the following equation: \( W_t\% = (W_0 - W) / W_0 \times 100\% \).

The refractive index (RI) of samples was measured of fully hydrated membranes using an Abbe refractometer (Model C10, VEE GEE Scientific Inc., Kirkland, WA, USA) at 21°C with DW as the calibration agent.

To test membrane light transmission, 6-mm acellular discs of the sample were punched out and placed in a glass bottom 96-well plate. Using the previously published protocol [18], light absorbance was measured at room temperature for specific wavelengths of the visual spectrum (405, 450, 490, 530, and 630 nm) with the VICTOR3 microplate reader (PerkinElmer). Percentage of light transmittance was calculated with the following equation: transmission (\( % \)) = \( 10^{2 - \epsilon} \) - absorbance). Wells mounted with 100 μL of trypan blue or DW served as negative and positive controls, respectively. Transmittance values were normalized to demineralized water.
Oscillatory rheology was performed on a Discovery Hybrid Rheometer (DHR2; TA Instruments, Sollentuna, Sweden) using 8-mm diameter, parallel-plate, stainless steel geometry. Frequency sweeps were performed at a constant shear strain amplitude of 0.267% at 25 °C.

Tensile testing of the membrane samples was performed on a vertical stress-strain testing device (Instron 3345, Canton, MA, USA) fitted with a 100 N load cell. Testing was performed at a 2 mm/min rate. Membrane samples where cut into 4 × 0.5-cm-long strips and fixed at both ends to a 1-mm-thick polypropylene frame. The ultimate tensile strength (UTS) was calculated based on the area of the sample (sample thickness measured with optical coherence tomography; RTVue 100–2, Optovue, California, USA).

Hydrogel Stability and Susceptibility to Enzymes and Microbial Attack

The stability of the hydrogels was tested against in vitro degradation by collagenase enzyme [19]. In brief, samples (n = 3) equilibrated in Tris–HCl buffer (0.1 M, pH 7.4) were incubated in 5 mL 5 U/mL type I collagenase dissolved in Tris–HCl at 37°C. The undigested mass was weighed at time 0 (W₀) and every 60 min (Wₜ) until the sample was completely digested. At every interval, surface water was blotted away, and samples were weighed three times independently using an ultra-microbalance (SE2, Sartorius, Göttingen, Germany). The percentage of mass remaining after digestion was calculated according to the following equation: Residual mass (%) = (Wₜ/W₀) × 100%. As a control, HAM was incubated in Tris–HCl buffer only and weighed at the same time points.

For determination of attack by common bacteria causing blindness, such as Gram-negative Pseudomonas aeruginosa, fully hydrated samples equilibrated in PBS were cut into 1 cm² pieces. Hydrogels were mounted in a 24 well-plate and fixed using a CellCrown. As previously described by Dravida et al. [20], a dilution of 10⁻¹ of two P. aeruginosa strains (ATCC 15442 and ATCC 9027) was made in Tryptic Soy Broth (TSB). Using a 22 gauge needle, 100 μL solution was injected into the scaffold (n = 3 for each ATCC strain). The inoculated scaffolds were incubated in an oven (37°C) for 24 h. The samples were homogenized in PBS using a tissue grinder. The homogenized extracts were diluted in the range of 10⁻⁸ and plated on agar (TSA). The number of colony-forming units (CFU)/mL was determined 24 h later. To have sufficiently thick amnion to inject the solution, mechanical scraping of the spongy layer was not performed for this experiment. Injection of 100 μL bacterial solution in 10 mL TSB and 10 mL PBS served as negative and positive controls respectively.

Ultrastructural Characterization

Atomic Force Microscopy

Membrane samples of 10 × 10 mm² were cut out and placed onto a rigid support carrier. The carrier, a glass slide that had functional aldehyde surface groups (developed in-house), was then immersed in PBS to allow full rehydration of the sample prior to atomic force microscopy (AFM) imaging. HAM samples received additional gentle mechanical scraping with cell scrapers to remove excess spongy layer, thus exposing the underlying membrane and reducing height variation. While imaging, samples were kept hydrated. AFM imaging was carried out in PBS by a NanoWizard 3 (JPK Instruments, Germany) system using NSC35 probes (MicroMasch, Germany) and the force curve acquisition-based Quantitative Imaging™ (QI) mode. Topography images were generated with the setpoint height channel in the QI-mode, setpoint force 5–10 of nN. Fibril diameters were estimated from height cross-sections extracted from the AFM images. Full-width at half-maximum was measured where the image resolution was sufficient. In other cases (e.g., rRHCI characterized by very fine fibrils), a trained operator’s estimates were used.

Electron Microscopy

Samples of biomaterials were fixed and prepared for scanning electron microscopy (SEM) and transmission electron microscopy (TEM) as we previously described in Haagdorens et al. [14]. Gold sputter-coated samples were imaged on a SEM 515 Microscope (Philips, Eindhoven, The Netherlands) for surface features [14]. Ultra-thin sections were cut from osmicated samples embedded in EMbed 812 (Electron Microscopy Sciences, Hatfield, PA). Ultrathin sections were stained with lead citrate and examined using a Tecnai G2 Spirit Bio Twin Microscope (FEI, Eindhoven, The Netherlands) at 120 kV.

In vivo Biocompatibility Testing in Rats

Random RHCI hydrogels, 11 mm in diameter and 1-mm thick, were implanted subcutaneously in rats (following ISO standard 10993-6:2007). Denuded HAM was also implanted as a comparison for RHCI. To achieve samples of sufficient thickness, 7 layers of HAM were stacked upon each other and sealed with fibrin glue (Tissuel, Baxter, Deerfield, IL, USA). Clotted fibrin sealant molded into 1-cm³ blocks were implanted as a control. In total, 5 collagen hydrogels, 4 HAM samples, and 3 fibrin clots were implanted subcutaneously. Two sham surgeries were performed to ensure that the skin healing process does not affect the outcome of biocompatibility testing. Postoperatively, extended assessment of health conditions of animals was performed based on an observation-al grading scheme (not shown).
At 3 months post-implantation, the animals were euthanized, and the implants and surrounding tissues were collected. Each rRHCI samples was subdivided into quarters for analysis. Quarters one and two were fixed in formalin, then embedded in paraffin or O.C.T. compound, respectively for histology and immunohistochemistry. One quarter was fixed in glutaraldehyde for EM. The final quarter was not analyzed. Recovered HAM, fibrin, and sham samples were similarly processed for histology and immunohistochemistry.

Non-consecutive paraffin sections, cut at 7-μm thick, were stained with hematoxylin and eosin (H&E) for histological evaluation. Based on H&E staining, two slides, each containing three tissue sections of the same sample, were used to assess biocompatibility as per Table S1 (adapted from ISO10993-6:2007 Table E3). According to this grading table, the test sample was considered a non-irritant (0 up to 2.9), slight irritant (3.0 up to 8.9), moderate irritant (9.0 up to 15.0), or severe irritant (> 15). Grading was performed double-blinded. In brief, each H&E section was graded at five different sites, two at each respective end, and one in the middle of the sample. If remnants of the implant could be identified, the area of grading was at the junction of the sample and surrounding tissue.

Cryosections, 7-μm thick, were stained with antibodies to identify plasma cells (anti-syndecan-1), granulocytes (anti-myeloperoxidase), macrophages (anti-CD68), T-lymphocytes (anti-CD3), blood vessels (anti-vWF), and activated myofibroblasts (anti-αSMA) as previously described (Table S2) [38]. Immunofluorescent (IF) images were recorded using an inverted LSM-800 Zeiss confocal microscope (LSM800, Carl Zeiss Microscopy, Germany), and the percentage of positive staining area per view was calculated for each sample using the ImageJ software (National Institutes of Health, Bethesda, MD, USA). In each stained sample, the analysis was performed for three IF images at standard locations (left end, middle, and right end of the sample). A Kruskal-Wallis nonparametric test was used with a Dunn’s multiple comparisons post hoc test. Statistical analysis was performed in GraphPad Prism 8.4.3. \( P \leq 0.05 \) was considered significant.

**In vivo Efficacy Testing as Corneal Implants in Minipigs**

To test in vivo safety and efficacy of corneal regeneration, the implants were evaluated in Göttingen minipigs, which are genetically uniform. Four animals were used in each treatment group. The left eyes of the pigs underwent surgical treatment, while the right eyes served as unoperated controls. All animals were subjected to anterior keratoplasty. The central section of corneal epithelium and stroma was removed to a depth of 500μm using a 6.5-mm surgical trephine and a scalpel. Solid rRHCI gels were grafted into the wound bed, covered with HAM, and secured in place using interrupted sutures. In the control group, cornea excised from the other pigs in this study was used as implants, being secured with interrupted sutures. These were syngeneic grafts rather than conventional allografts, as they are from genetically identical donors and recipients. The syngeneic grafts were secured using interrupted sutures. All operated eyes received corticosteroids and antibiotics in the form of topical dexamethasone/tobramycin eye drops (Tobrasone, Alcon, Sweden). The dexamethasone/tobramycin was applied 3 times daily for the first 5 postoperative days. The pigs were monitored daily for any eye-related complications for the entire study duration. Clinical exams were performed pre-operation, and then at 6 weeks, and 3, 6, 9, and 12 months post-operation according to protocols as we described in McTiernan et al. [21]. Briefly, all operated and control corneas were examined by slit lamp biomicroscopy (for haze, redness), anterior segment optical coherence tomography (for corneal changes including thickness), Schirmer’s tear test (for tear production), and aesthesiometry (for touch sensitivity). At the endpoint of 12 months post-operation, all animals were euthanized according to animal facility guidelines, and tissue samples of both operated and contralateral eyes were collected.

Samples from implanted and control corneas were processed for TEM as described in McTiernan et al. [14, 21, 22]. In brief, samples were prepared for TEM by fixing a quarter of each cornea in 2.5% glutaraldehyde solution in 0.1 M sodium cacodylate buffer (pH 7.4). After fixing, the quarters were cut into 1-mm wide strips. The strips were postfixed in 1% OsO4 solution for 2 h, dehydrated through an ethanol series and embedded in EMBed 812 (Electron Microscopy Sciences, Hatfield, Pennsylvania). Ultrathin sections stained with lead citrate were imaged on a Tecnai G2 Spirit Bio Twin Microscope (FEI, Eindhoven, The Netherlands) at 120 kV.

In vivo confocal microscopy examinations were performed before surgery and at 3, 6, 9, and 12 months post-surgery. Images taken were used for nerve counts as described in McTiernan et al. [21]. When nerves where observed, they were defined as bright, slender, straight, or branched structures and were sufficiently different from the background intensity level. NeuronJ, a nerve tracing and analysis software, and FIJI were used together for measuring the total length of the observed nerves [23]. Corneal nerve densities were based on averaging the highest nerve density for each treatment group and time point from individual images.

**Statistics**

For all tests of hydrogels, a minimum of three samples was used in evaluations. For the in vivo subcutaneous studies, 5 RHCI hydrogels, 4 HAM samples, and 3 fibrin clots were implanted. The additional RHCI and HAM samples were implanted just in case animals died from unrelated causes, as these samples were more difficult to obtain. As all animals...
remained healthy and survived, so, all samples were analyzed for potential adverse immune cell infiltration or activation by differential cell counts followed by a Kruskal-Wallis nonparametric test as described above.

The unit of analysis for the in vivo study in minipigs was the eye (RHCI n=4, syngeneic graft n=4, unoperated n=8). Clinical variables over time (Fig. 3a–f) were analyzed with a mixed effects model with matched stacking, no assumption of sphericity, and \( \alpha = 0.05 \) using a post hoc Tukey’s multiple comparison test. Statistical analysis and graphing were performed using GraphPad Prism v8.4.3 (GraphPad Software LLC, San Diego, CA, USA).

Results

Physical and Mechanical Properties of RHCI Hydrogels

The properties of the aligned and random RHCI hydrogels, compared to HAM, are summarized in Fig. 1. Both types of collagen hydrogel and control HAM displayed a RI that approximates that of the human cornea (1.37–1.38). The rRHCI hydrogels allowed for light transmittance within the visual spectrum of over 91% (Fig. S1) while aRHCI hydrogels showed only 37% light transmission, appearing opaque (Fig. S1, 1a). Figure 1a also shows that the aRHCI hydrogels were prone to splitting along the direction of the collagen fibrils. HAM, which served as a control, showed light transmission of 66%. The opacity of the aRHCI hydrogels precluded its use for corneal implants, while it was tested for toxicity and biocompatibility as subcutaneous implants in rats, it was not tested as corneal implants in vivo. The rRHCI hydrogels used for corneal implantation, however, was robust, pliable, and easy to manipulate when molded into corneal-shaped and corneal-sized implants (Fig. S2).

Both aligned and random hydrogels, as well as HAM, showed a predominantly elastic, rather than viscous behavior with storage moduli much higher than loss moduli. However, due to the very low thickness of HAM, reliable data from oscillatory rheology was not possible to obtain and therefore, eliminated from further comparisons. The G’ of rRHCI gels was found to be 2.9 ± 0.03 kPa, whereas that of aRHCI hydrogels was found to be 0.383 ± 0.03 kPa indicating a 10-fold higher stiffness for the rRHCI (Fig. S3). The aRHCI hydrogels have slightly higher water content compared to their random counterparts. Moreover, the G’’ of the rRHCI hydrogels was found to be 0.158 ± 0.036 kPa resulting in a tan\( \delta \) of 0.05± 0.01. Stress strain analysis shows that both RHCI hydrogels display stress and strain values that are inferior to that of HAM membrane. These results reveal that HAM displayed roughly three times greater energy absorption compared to random RHCI hydrogels. Young’s modulus indicates that the latter are much stiffer than the HAM. The aRHCI hydrogels were inherently too weak to allow stress-strain analysis.

Collagenase degradation assays showed that aligned and random RHCI hydrogel degradation was complete by 8 h and 16 h respectively (Fig. S4). The degradation of HAM was slower, with complete degradation occurring after 24 h. Inoculation with \( P. \) aeruginosa showed that RHCI hydrogels, irrespective of fibril alignment or not, were 50- to 100-fold more resistant to microbial contamination for both strains of \( Pseudomonas \) (Table S3) compared to HAM. We recently showed in Haagdorens et al. that RHCI hydrogels were biocompatible with human corneal epithelial cells and supported their growth [14].

Ultrastructure

Surface scanning by AFM (Fig. 1d–f) confirmed the random arrangement of fibrils in rRHCI hydrogels, and the unidirectional alignment of the RHCI fibrils in the shear-mediated preparation of aRHCI. It was noted that the collagen fibrils in the aRHCI hydrogels are thicker (93.8 ± 30.6 nm in diameter) than the fibrils in the rRHCI hydrogels (23.6 ± 10.4 nm in diameter). This showed that the fibrils had assembled into distinct thicker bundles of fibrils or collagen fibers. Fibrils in the random hydrogel were very fine, but also matted in areas. HAM fibrils were more similar to aRHCI in thickness (112.6 ± 61.7 nm in diameter).

SEM (Fig. 1g–i) confirmed the unidirectional collagen orientation in the surface of aHCl hydrogels. The rRHCI hydrogels showed a smooth surface. SEM imaging of HAM revealed complete removal of amniotic epithelium and exposure of the underlying basement membrane. At the amniotic basement membrane, remaining cellular debris and extracellular matrix (ECM) could be seen between collagen fibers.
TEM of rRHCI hydrogels confirmed the presence of fine collagen fibrils distributed randomly throughout the sample (Fig. 1k) as revealed by AFM. In the aRHCI, the fibrils formed thick collagen bundles (Fig 1j) as seen in the cross-sectional views. They showed unidirectional alignment in contrast to the interpenetrating thin collagen fibrils.
In vivo Biocompatibility in Rats

After subcutaneous implantation of the rRHCI hydrogels in rats, the animals did not exhibit any general or implantation-site adverse symptoms. Upon recovery of the implants, there were no clear macroscopic signs of implant degeneration, and the RHCI hydrogels could easily be distinguished from the surrounding tissues. Based on grading of the H&E-stained samples (Fig. 5a–f), RHCI hydrogels were ruled as being slightly irritant, with an average grading score of 7.33, well comparable to controls with scores of 7.63 for fibrin, 9.75 for HAM and 7.93 for sham (Table S4).

Transmission electron microscopy images of implanted hydrogels showed isotropic collagen fibers interspersed with capillaries and small blood vessels, along with fibroblasts (Fig. 2b–e). Microscopically, some degree of disintegration was evident, and the RHCI hydrogels were mainly infiltrated by lymphocytes, fibroblasts, and macrophages with some ingrowth of capillary vessels (Fig. 2f). Even though granulocytes were not identified in the H&E sections, some cells stained positive for anti-myeloperoxidase, a granulocyte marker, in the IF pictures (Fig. 2f). Statistical analysis of the prevalence of infiltrating cells showed statistically significant \( P \leq 0.05 \) elevated levels of T-lymphocytes, von Willebrand factor (vWF) positive cells, macrophages, and granulocytes compared to HAM, but no significant difference compared to other groups (Fig. 2g). The plasma cell staining was higher in the RHCI group compared to both HAM and fibrin glue. No significant differences among groups were found in the myofibroblast counts.

In vivo Performance in Minipigs

Results for the clinical observations of syngeneic grafts and their unoperated contralateral untreated corneas were previously reported in McTiernan et al. as the same four animals served as controls for the two different studies [21]. Clinical results are summarized in Fig. 3. At 6 weeks post-implantation, the RHCI implants were completely covered with regenerated epithelium, as noted from clinical observation and slit lamp. The corneal thickness in the RHCI group was close to unoperated ranges at 3 months, and indistinguishable from unoperated by 6 months onwards (Fig. 3a). Even though there looks to be a trend towards a thicker cornea in the RHCI group compared to the syngeneic graft, this difference was statistically non-significant.

There is a statistically significant amount of haze in the RHCI group (Fig. 3c) that decreased with time and was scored at zero in two out of the four animals at the 12-month point. At the 12-month point, the difference between groups is not statistically significant (Tukey’s multiple comparisons \( \alpha = 0.05 \)).

The subepithelial nerve plexus recovered fast, and at 12 months, nerve plexus morphology in the RHCI group resembled that of unoperated animals as seen in IVCM in Fig. 4. Nerve density of the corneal stroma, as noted based on IVCM, was close to normal ranges in both groups at 3 months onwards (Fig. 3e). One animal in the RHCI group had haze in the regenerated cornea that did not allow for imaging to obtain nerve counts. The missing datapoints were taken into account by using a mixed model rather than ANOVA for this comparison. All the pigs regained touch response in the operated eye prior to 12 months. Statistical analysis could not be performed due to uneven state of sedation of the animals during measurement.

Intraocular pressure was indistinguishable from unoperated eyes at all timepoints in both the RHCI group and the syngeneic grafts (Fig. 3b). The normal intraocular pressure (IOP) shows that there is no weakening of the cornea capable of causing long-term distortions or leakage, nor was the implant too rigid to cause increased pressure.

H&E sections showed that the stratified corneal epithelium and stroma have regenerated in the neo-corneas of all four pigs, to resemble those in the syngeneic grafts and untreated, healthy controls (Fig. 5g–l). Immunohistochemical staining showed full regeneration of a mucin-positive tear film and fully differentiated epithelial cells (Fig. 5). Antibodies against smooth muscle actin (α-SMA) showed the presence of positive cells in RHCI, most likely myofibroblasts, but not the syngeneic or control untreated corneas (Fig. 5y–ad).

Discussion

Plant-derived RHCI was successfully fabricated into aligned and random hydrogels. The RHCI was aligned to mimic the highly aligned, lamellar structure of collagen in the cornea. Unfortunately, the aRHCI hydrogels were opaque. Despite the same net weight of RHCI in the random and aligned gels, the aRHCI gels were thicker as the collagen assembled into thicker fibrils with variable spacing in between fibers. They also demonstrated extreme transverse mechanical weakness. The optical and mechanical deficits of the hydrogels in comparison to cornea is most likely due to the structural deficits of RHCI-only fibrils and a lack of perpendicular structural elements (for review, see España et al.) [24]. In the cornea, collagen protofibril formation is regulated through the incorporation of collagen V to create uniform protofibrils for subsequent fibrillar assembly [24]. In the absence of collagen V, collagen I produces thicker, branching, heterogeneous fibers like the ones observed by AFM and TEM in this study. These fibers are known to be incompatible with corneal transparency. Collagen fiber spacing also contributes to the aRHCI structural deficits. The spacing of corneal collagen fibrils is regulated by small leucine-rich proteoglycans, preventing the compaction of the collagen fibrils seen in the aRHCI [24]. The high aRHCI fibril density also contributes to the limitations...
in light transmission through the hydrogels. Finally, the collagen fibrils in each lamellar layer of the cornea are aligned in different directions, providing uniform mechanical strength throughout the entire cornea [24]. The single layer, shear force alignment method used here was insufficient to produce the structural complexity required to fully mimic the human cornea and will undergo future modifications, in order to improve the transparency of the material.
By contrast, the rRHCI hydrogels were transparent and mechanically stronger in all directions. When compared to previously tested collagen hydrogels, the transparency of random RHCI hydrogels was equivalent to that of porcine...
collagen type I, yeast-sourced RHCI, RHCIII, and collagen-mimicking hydrogels [25–29]. Moreover, rRHCI hydrogels outperformed HAM and rat tail collagen type I gels in terms of light transmission and refractive index [30]. In this study, RHCI hydrogels degraded faster in collagenase than HAM. The data on HAM degradation times should however be interpreted carefully; in this study, HAM degraded much slower than earlier reported degradation times of 6–13 h [18, 20]. The variability in degradation speeds of HAM underscores the inconsistent state of donated HAM and the need for well-standardized carrier materials in ocular tissue engineering. Aligned RHCI degraded too quickly for future clinical purposes without further modifications to its chemical formulation [31].

The difference in degradation time between both types of RHCI hydrogel may be attributed to the difference in collagen fibril density. One would assume, it would take longer to degrade the densely packed fibers in the aRHCI; however, without the interpenetrating fibrils holding the aligned fibers together, the hydrogel disintegrates into isolated fibers. Given the dense fibrillar network in the rRHCI hydrogels, these hydrogels degraded slower than aRHCI. Collagenase resistance of rRHCI was comparable to reported resistance of pure RHCIII, porcine collagen type I, and rat tail collagen type I hydrogels, with 8 h, 12–20 h, and 8 h degradation time respectively [19, 20, 25, 32].

As demonstrated by Griffith et al., collagen hydrogels have a tendency of being more resistant to microbial contamination when compared to HAM or the human cornea [20, 33]. As we inoculated membranes with P. aeruginosa, a bacterium responsible for causing severe corneal ulcers, results confirmed...
the relative microbial resistance of RHCI compared to amniotic membrane. Microbial contamination of grafts is a serious and widespread problem in ocular surgery because it often causes devastating infections and loss of implanted material [20]. With a 100-fold resistance to microbial infection, RHCI has a clear advantage over HAM in a clinical setting, where substrates are manipulated repeatedly in clean rooms prior to transport to the operating theater.

By implanting RHCI hydrogels subcutaneously in rats, we demonstrated that they were biocompatible in mammals and elicit minimal inflammation. Furthermore, the recovered hydrogel implants had initiated partial degradation. As we and other groups previously reported intact hydrogel retrieval after 90 days of implantation, this might indicate that additional crosslinking or incorporation of interpenetrating phospholipids to RHCI might improve the stability of the implant [19, 25, 27, 28, 34]. There was formation of new collagen fibers within the implants. This suggests that infiltrating fibroblasts had begun to remodel the implant. Similar observations have been made in corneal implants that were implanted in pigs, where host keratocytes remodeled the implants’ fibrillar structure [19, 35]. By computer analysis of immunofluorescent images, we were able to quantify leukocytosis and neovascularization of retrieved implants and compare results to the isoproterenol-based H&E grading system. Interestingly, no granulocytes could be identified in H&E sections, whereas

![Fig. 5](image.png)

**Fig. 5** Columns 1–4 shows each of the individual animals in the RHCI group. Column 5 shows representative images from the syngeneic graft. Column 6 shows representative images from unoperated controls. a–f Eyes of all RHCI grafted pigs and controls, showing haze in a and b. g–l Hematoxylin and eosin stained sections show morphologies of regenerated neo-corneas are comparable to those of the controls. Mucin staining (m–r) shows that the animals in the RHCI group have intact tear films. Cytokeratin 12(Ck12) staining (s–x) shows differentiated corneal epithelial cells in all samples. y–ad α-Smooth muscle actin (α-SMA), a myofibroblast marker shows the presence of activated fibroblasts in RHCI corneas, particularly in the ones with haze.
anti-myeloperoxidase, a marker for granulocytes, showed to
be focally positive by immunohistochemistry, showing that
H&E histopathological examination should be supplemented
by immunohistochemistry using known markers for target
cells. In 2016, Van Essen et al. published their results on
automated analysis of IF area staining for fish scale-derived
collagen matrices that were implanted subcutaneously in rats
for 11 weeks [36]. With reported area of stain of <1% for
macrophages and T-lymphocytes, results are very similar to
the values we found. Moreover, area of stain for granulocytes
appears to be lower for RHCI hydrogels compared to reported
values for fish scale matrices. The presence of α-SMA posi-
tive cells that are myofibroblast-like in the RHCI corneas are
in keeping with the haze that was noted. They also correspond
to the results from the subcutaneous study showing that RHCI
was a mild irritant. It is possible that there may be traces of
plant material that contributed to the irritation and activation
of stromal cells.

Although there are collagens that are derived from bovine and
porcine skin which are widely available and offer a potentially
unlimited supply source, an important consideration for the use
of xeno-derived collagen is the potential risk for allergic reactions
to xenogeneic biomaterials [37] and zoonotic pathogen transmis-
sion [38] if tight processing controls are not adhered to. Overall,
plant-derived RHCI was largely biocompatible and immune
compatible. RHCI hydrogels promoted regeneration of corneal
tissue and nerves when grafted into the corneas of minipigs.
Finally, application of RHCI should not solely be restricted to
ocular regeneration, as collagen type I is abundantly present
throughout the body. Aligned RHCI, with its unique collagen
alignment, might find its way in musculoskeletal tissue engineer-
ing, whereas random hydrogels might be explored in cutaneous
or cardiac regeneration [39–41].

Conclusions

RHCI hydrogels made from random collagen fibrils were bio-
compatible and promoted corneal regeneration in minipigs. This
shows that the plant-derived collagen serves as a viable source of
collagen for tissue engineering and regenerative medicine appli-
cations, such as for promoting in situ tissue regeneration of the
cornea after further optimization and testing.

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Data Availability Data collected and analyzed for this study are provided
in the Electronic Supplemental Material. Other data generated during the
current study are available from the corresponding authors on reasonable
request. RHCI available from CollPlant Ltd.

Declarations

Ethics Approval The Declaration of Helsinki guidelines were applied
and were approved by the Antwerp University Hospital Ethical
Committee (EC: 14/30/319) for use of human amniotic membrane.
Animal experiments in rats and Göttingen minipigs were performed in
accordance with the Association for Research in Vision and
Ophthalmology Statement for the Use of Animals in Ophthalmic and
Vision Research and the Swedish Animal Welfare Ordinance and the
Animal Welfare Act, after ethical approval was acquired from the local
Linkoping ethical committee (Linköpings Djurförsökssetiskt Nämnd).

Competing Interests Dr. Oded Shoseyov is a founder and chief scientist
at CollPlant which produces and markets tobacco-derived RHCI. Other
authors have no competing interests.

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