Type VII Collagen in the Human Accommodation System: Expression in Ciliary Body, Zonules, and Lens Capsule

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Purpose. To investigate intraocular expression of COL7A1 and its protein product type VII collagen, particularly at the accommodation system.

Methods. Eyes from 26 human adult donors were used. COL7A1 expression was analyzed in ex vivo ciliary epithelium by microarray. Type VII collagen distribution was examined by Western blot analysis, immunohistochemistry, and immuno-electron microscopy.

Results. COL7A1 is expressed by pigmented and nonpigmented ciliary epithelia. Type VII collagen is distributed particularly at the strained parts of the accommodation system. Type VII collagen was associated with various basement membranes and with ciliary zonules. Anchoring fibrils were not visualized.

Conclusions. Type VII collagen distribution at strained areas suggests a supporting role in tissue integrity.

Keywords: collagen type VII, inner limiting membrane, basement membrane, accommodation, lens capsule

Type VII collagen (Col VII) is renowned as the major component of anchoring fibrils.1 It is essential for epithelium-to-stroma anchorage in skin, mucosa, and cornea.2,3 Although predominantly expressed in skin, Col VII is estimated to comprise a mere 0.001% of the total skin collagen content.4 Yet, the relevance of such levels of Col VII expression is clinically evidenced in severe dystrophic epidermolysis bullosa, in which patients lack functional Col VII (i.e., anchoring fibrils). Their skin blisters readily at small amounts of friction, damaging their epithelial basement membranes at each event. Such repetitive wounding is accompanied by severe and extensive scar formation, mutilating deformations, and recurrent infections. Patients succumb to skin cancer or sepsis often before age 35.5,6 Extraocular manifestations are well documented, and encompass mainly corneal and conjunctival (both surface ectoderm) erosions accompanied by scar formation, symblepharon, pannus, and so forth.6 Intraocular manifestations are mentioned only in rare case reports, where no clinical or histological abnormalities are found,7 or are limited to lens (also surface ectoderm) sclerosis.7–9 Interestingly, however, retinal COL7A1 gene expression was recently established (FANTOM5 consortium10). Moreover, its protein product was demonstrated at the vitreoretinal junction11,12 and inner layers of the normal retina,12,13 although no anchoring fibrils were visualized. Anchoring fibrils, however, are reported to be far more numerous at basement membrane zones of mechanically strained tissues.1,4,11,14,15 To investigate the characteristics of intraocular Col VII further, we explored the accommodation system. We observed COL7A1 expression at the pigmented and nonpigmented ciliary epithelia, and Col VII protein at the ciliary body and zonules.

Materials and Methods

Samples

Eyes were provided by the Euro Cornea Bank, Beverwijk (http://www.eurotissuebank.nl/corneabank/, in the public domain), The Netherlands. In The Netherlands, the usage of donor material is provided for by the Organ Donation Act (Wet op de orgaandonatie [WOD]). In accordance with this law, donors provide written informed consent for donation, with an opt-out for the usage of leftover material for related scientific research. Specific requirements for the use in scientific research of leftover material originating from corneal grafting have been described in an additional document formulated by the Ministry of Health, Welfare, and Sport, and the BIS Foundation (Eurotransplant; Leiden, July 21, 1995; 6714.h.t). The current research was carried out in accordance with all requirements stated in the WOD and the relevant documents. Approval of the local medical ethics committee was not required, because the data were analyzed anonymously. Skin tissue was obtained from a cosmetic surgery procedure, after written informed consent of the patient and approval by the institutional review board.
We analyzed eyes from a total of 26 human donors (age 35–83 years, mean age 63.7 years), which were without known ophthalmic pathology (Table 1). In general, corneas had been removed for transplantation purposes. All eyes were fixed, frozen, or processed within 48 hours postmortem.

Embedding

Eleven eyes were embedded in paraffin or resin as described previously.12 For light microscopy, the anterior parts of four eyes were positioned in 2% agarose before paraffin embedding, and sectioned at 3 to 4 μm (Leica RM2265; Microtome, Heidelberg, Germany). For electron microscopy, the anterior parts of three eyes were washed, dehydrated, pre-infiltrated, and embedded in T8100 resin (Technovit 8100; Heraeus Kulzer, Wehrheim, Germany). Areas of interest were sawn out, trimmed and sectioned at 100-nm thickness (Ultracut type 701; Reichert-Jung, Vienna, Austria). Three eyes and a skin sample were not fixed, but mounted in optimal cutting temperature compound (Tissue-Tek; Sakura Finetek Europe, Alphen aan den Rijn, The Netherlands), snap-frozen in isopentane propane in liquid nitrogen, and cut in 7- to 10-μm thick sections (CM3050 S Cryostat; Leica Microsystems, Wetzlar, Germany).

Immunohistochemistry

Paraffin, T8100, and cryosamples were processed as described previously.12 For light microscopy, the anterior parts of four paraffin-embedded eyes were deparaffinized with xylene followed by ethanol rehydration steps. After washing, a protease K (IHC Select kit; Chemicon/Millipore, Billerica, MA, USA) dilution of 1:5 in PBS was added for 15 minutes to unmask epitopes. After washing, endogenous peroxidases were blocked (0.3% H2O2/PBS for 30 minutes). Primary anti-Col VII antibodies (Supplementary Table S1) were allowed to incubate (1:100 in PBS for 1 hour). These comprised two monoclonal antibodies, designated mAb(12) (LH7.2; Abcam, Cambridge, UK) and mAb(14) (clone 32; EMD Millipore, Billerica, MA, USA) and two polyclonal antibodies, designated pAb(16) (anti-tissue type Col VII; Calbiochem, San Diego, CA, USA) and pAb(72) (recombinant LH7.2, a kind gift from Alex Nyström, PhD, University of Freiburg, Freiburg im Breisgau, Germany). After thorough washing in PBS, sections were incubated in corresponding horseradish conjugated secondary antibodies (rabbit-anti-mouse and goat-anti-rabbit; DAKO, Glostrup, Denmark) (1:500 in PBS for 1 hour). Sections were washed, then stained using 3-amino-9-ethylcarbazole (AEC Staining Kit; Sigma-Aldrich, St. Louis, MO, USA) and counterstained with hematoxylin. For signal enhancement, an avidin/biotin labeling kit was used (Vectastain Elite ABC kit; Vector Labs, Burlingame, CA, USA), according to the manufacturer’s instructions. An HC DMR microscope (Leica) was used to analyze the samples. All proceedings were performed at room temperature. In negative control sections, incubation with primary antibodies was omitted.

Immuno-Electron Microscopy

Sections from T8100 embedded eyes were mounted on 150-mesh, 0.6% formvar-coated nickel grids. For antigen retrieval, samples were incubated in 0.05% trypsin (Gibco, Paisley, Scotland) in 0.1 M Tris-buffer (pH 7.8) containing 0.1% CaCl2, for 15 minutes at 37°C. The sections were washed, then stained using 3-amino-9-ethylcarbazole (AEC Staining Kit; Sigma-Aldrich, St. Louis, MO, USA) and counterstained with hematoxylin. For signal enhancement, an avidin/biotin labeling kit was used (Vectastain Elite ABC kit; Vector Labs, Burlingame, CA, USA), according to the manufacturer’s instructions. An HC DMR microscope (Leica) was used to analyze the samples. All proceedings were performed at room temperature. In negative control sections, incubation with primary antibodies was omitted.

**Table 1. Donor Characteristics**

<table>
<thead>
<tr>
<th>Assay</th>
<th>Embedding</th>
<th>Age</th>
<th>Sex</th>
<th>Cause of Death</th>
<th>Comorbidity</th>
<th>Fixative</th>
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<tr>
<td>IHC</td>
<td>Paraffin</td>
<td>83</td>
<td>Female</td>
<td>Cerebrovascular accident</td>
<td>Polymyalgia rheumatica</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>65</td>
<td>Male</td>
<td>Cardial</td>
<td>Bladder cancer</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>64</td>
<td>Male</td>
<td>Cardial</td>
<td></td>
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<tr>
<td></td>
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<td>78</td>
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<td>Cryo</td>
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<td>67</td>
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<tr>
<td></td>
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<td>68</td>
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<td>Malignancy</td>
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<tr>
<td></td>
<td></td>
<td>35</td>
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<td>N/A (control, abdominal skin)</td>
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<td>Aortic dissection</td>
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<td>Cerebrovascular accident</td>
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<td></td>
<td>73</td>
<td>Male</td>
<td>Cardial</td>
<td></td>
<td>None</td>
</tr>
</tbody>
</table>

PF, paraformaldehyde.
* Skin sample only.

We analyzed eyes from a total of 26 human donors (age 35–83 years, mean age 63.7 years), which were without known ophthalmic pathology (Table 1). In general, corneas had been removed for transplantation purposes. All eyes were fixed, frozen, or processed within 48 hours postmortem.
Sections were then incubated in primary antibody pAb(16) (1:100) in a 1% BSA-c/PBS buffer, initially for 2 hours at 37°C, then at room temperature overnight. Afterward, the sections were washed, and incubated with gold-labeled anti-rabbit IgG (goat-anti-rabbit, Gold Colloid 6 nm, 1:200; Aurion) for 1 hour. The samples were washed in ultrapure water, fixed in 2% glutaraldehyde for 2 minutes, silver-enhanced according to kit protocol (R-Gent SE-LM Silver enhancement kit; Aurion) for 10 minutes at room temperature, and washed again with ultrapure water. Then, they were contrasted by applying methylcellulose-uranyl acetate (9:1) at 4°C for 15 minutes, after which the solution was removed. The samples were allowed to dry for at least 15 minutes, and were analyzed in a CM100 BioTwin transmission electron microscope (TEM) (Philips, Eindhoven, The Netherlands) using immuno-electron microscopy (iTEM) software (ResAlta Research Technologies, Golden, CA, USA) and a Morada 11-MP TEM camera (Olympus Soft Imaging Solutions GmbH, Münster, Germany).

**Antibody Epitope Mapping**

The antibodies mAb(12), mAb(70) (US Biological, Swampscott, MA, USA), and pAb(72) target Col VII at its amino terminus (NC-1), whereas mAb(14) targets the collagenous domain toward the amino-terminal end. To assess the target epitopes of pAb(16), this antibody was mapped by a third party commercial laboratory (PEPperPRINT, Heidelberg, Germany) (Supplementary Fig. S1).

**Western Blotting**

Seven bulbi (three pairs pooled for pAb[16], one bulbus for mAb[14]) were microscopically dissected in cooled 90% glycerol (diluted in 1% EDTA/PBS). Their anterior segment was isolated by cutting through the pars plana of the ciliary body circumferentially. The vitreous was dissected close to the anterior hyaloid, the iris was bluntly removed by forceps. The anterior segment was stretched and fixed on a clear silicone layer by insect pins. The zonules and lens capsule were contrasted with MembraneBlue-Dual dye (DORC, Zuidland, The Netherlands). The zonules were cut closely to the inner limiting membrane (ILM), minimizing contamination by ILM and ciliary epithelium. After opening of the capsular bag and removing the lens, visible lens fibers were flushed out. Traces of iris pigment that stuck to the anterior lens capsule were removed with Q-tips. Then, the pars plicata of the ciliary body was isolated from remaining retinal and choroidal fragments. Excess fluids were removed by centrifugation (85 g for 2 minutes). Tissues were homogenated by turraxing them in 0.5% Triton X-100 in Tris-buffered saline. Some samples were
subjected to bacterial type VII collagenase (high purity grade, 30 units for 1 hour at 37°C; Sigma, City, State, Country) or pepsin (vortexing 5 minutes at room temperature, 4 mg/mL pepsin in 0.5 M acetic acid at one-fourth sample volume; Sigma) digestion before sample buffer incubation. Sample buffer, SDS-PAGE, and Western blotting were performed as described earlier.12 Briefly, nitrocellulose membranes were blocked with 2% fat-free milk and incubated overnight in mAb(14) or pAb(16) (Supplementary Table S1). The membranes were incubated in corresponding secondary antibodies (goat-anti-mouse or goat-anti-rabbit; Jackson ImmunoResearch, West Grove, PA, USA), washed, incubated in tertiary alkaline phosphatase-conjugated antibodies (rabbit-anti-goat-AP; Jackson ImmunoResearch) for 1 hour each, and developed in NBT/BCIP (BioRad, Hercules, CA, USA). All steps were performed at RT. In controls, the primary antibody was omitted.

**Gene Expression**

To assess COL7A1 expression in pigmented epithelium (PE) and nonpigmented epithelium (NPE), gene expression level data were provided by Janssen et al.16 Data were derived from their total dataset, including the unpublished remainder of their selection of top 10% relevant PE and NPE genes. In short, PE and NPE cells from seven snap-frozen donor samples were separately collected through laser dissection microscopy. Then, RNA isolation, amplification, labeling, and hybridization against 44k Agilent microarrays was performed. Gene expression data were analyzed with R and the knowledge database Ingenuity.16

**RESULTS**

**Immunohistochemistry (IHC) and iTEM: Polyclonal Antibody**

By light microscopy, pAb(16) labeling was observed at the ciliary zonules, stromal fibroblasts, and the basement membranes of PE and blood vessels. The zonules showed anti-Col VII labeling along their entire span (Figs. 1, 2). Through that labeling, zonules could be traced to fibrillar structures at the NPE layer (Figs. 1A, 2A). Labeling of the PE basement membrane was sharply delineated from pars plana (Fig. 1A)
to pars plicata. However, at the bases of the ciliary processes (Fig. 1B), the area in-between the PE basement membrane and that of nearby blood vessels labeled broadly (i.e., Figs. 1B, 2B). Such labeling around blood vessels diminished rapidly toward the tips of the ciliary processes, where the sharp delineation returned (Fig. 1C). The lens and lens capsule remained unlabeled (Fig. 1D). At higher magnification, however, small intracapsular structures were seen at the antero- and post-equatorial capsule, where they corresponded to zonules (Figs. 2C, 2D). They disappeared toward the anterior and posterior poles. The basal lamina of the posterior iris PE was slightly labeled, posteriorly more so than anteriorly. The iris stroma showed almost no labeling. Lens fibers showed some faint background labeling (on rare occasion), and were considered “unlabeled” (Fig. 1D).

By iTEM, such IHC labeling was confirmed. The zonules, including their NPE origin and capsular insertions, were labeled, as was the basement membrane of PE. No anchoring fibrils could be distinguished (Figs. 3, 4). The basement membrane of the PE labeled moderately throughout its entire thickness (Figs. 3E, 5G).

At the capsular surface, some approximating zonules extended protruding fibrils, perpendicular to the zonular course and lens capsule surface (Fig. 4E). Here, the assembled zonules ran parallel to the capsular surface, thus forming the zonular lamella. At areas without such perpendicular fibrils, a lucid unlabeled plane was seen between the zonules and lens capsule (Fig. 4D; Supplementary Fig. S2). Intracapsular densities (or linear densities) were readily distinguishable from the lens epithelial surface (Supplementary Fig. S2B), no actual intracellular labeling was seen. The intracapsular densities had round or elongated (linear) shapes. Their fibrillar aspect and labeling were less outspoken than that of zonules (Figs. 4C, 4D). The round densities had lucent envelopes, which were less distinct in the elongated variant (Figs. 4C, 4D; Supplementary Fig. S2B). At the (antero-) equatorial lens capsule zonular fibrils mingled with the capsule at certain points (Fig. 4E), but otherwise paralleled a lucent superficial capsular plane (Supplementary Figs. S2A, S2B). No connections between zonules and intracapsular densities could be visualized convincingly; although on occasion faint intermediary fibrillar shapes could be discerned (Supplementary Fig. S2C). The NPE and PE cells labeled negligibly, as did the collagenous stroma underneath the PE. In both IHC and iTEM, the vitreous cortex, the lens fibers, and the lens epithelium...
remained unlabeled. Negative controls (IHC/iTEM) had no significant labeling or background. Results are summarized in Table 2.

Antibody Epitope Mapping

Seven epitopes could be identified on the pAb(16) antibody with specific binding affinity for Col VII. The three epitopes with the best signal-to-noise ratio (E-value), had affinity for the collagenous triple helical domain of Col VII. Of the four remaining epitopes, two epitopes would target NC-1 peptide sequences in the fibronectin (FN3) domain and two in the von Willebrand factor (vWFA) domain (Supplementary Fig. S1).

IHC and iTEM: Confirmation Antibodies

To confirm our immunohistochemical results obtained by pAb(16) in ocular tissues, a comparison with other validated antibodies was made in cryosections of skin (Supplementary Fig. S3) and ciliary body (Fig. 5). All antibodies labeled the dermal-epidermal basement membrane zone specifically. In cryosections of ocular tissue, the labeling of the pAb(16) antibody was partially reproduced by the other antibodies. The zonular fragments that survived cryosectioning were stained

Table 2. Summary of pAb(16) Labeling Intensities, Semi-Quantitative Col VII Determination

<table>
<thead>
<tr>
<th>Wb</th>
<th>IHC</th>
<th>iTEM</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Stroma (fibroblasts)</td>
<td>++</td>
</tr>
<tr>
<td></td>
<td>Vascular BM</td>
<td>++</td>
</tr>
<tr>
<td></td>
<td>BM</td>
<td>+++</td>
</tr>
<tr>
<td></td>
<td>NPE BM (ILM)</td>
<td>+++</td>
</tr>
<tr>
<td></td>
<td>Intercellular zonular origins</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>Zonules/lens capsule complex</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>Zonules</td>
<td>+++</td>
</tr>
<tr>
<td></td>
<td>Lens capsule</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Zonular membrane</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>Linear densities</td>
<td>++</td>
</tr>
<tr>
<td></td>
<td>Lens</td>
<td>-</td>
</tr>
</tbody>
</table>

BM, basement membrane; -, no signals; +, weak; ++, moderate; ++++, intense.
faintly, and only by the pAb(16) antibody. Intracapsular densities were not discernible. Avidin-biotin signal enhancement often resulted in basement membrane labeling. Evaluation by iTEM could not be performed, because none of the comparison antibodies proved able to label positive control sections of skin in T8100 adequately.

**Western Blotting**

Col VII could be demonstrated in accommodation system lysates by mono- and polyclonal antibodies (Fig. 6). Although Col VII was not easily extracted from cornea control tissue, a limited collagenase digestion of the crude lysate showed a 290-kDa signal. Tissue type Col VII could be detected in zonules/lens capsule and ciliary body lysates, although a brief pepsin treatment of the crude lysate was needed for mAb(14) detection.

**Gene Expression**

COL7A1 is equally expressed in PE and NPE at “low” levels (compared with the total dataset), as indicated by a “log2” conversion.
transformed absolute expression level”13-16 (Table 3). These data can be found in the Gene Expression Omnibus database (GSE37957).

**DISCUSSION**

This study establishes Col VII expression in accommodation system tissues. Col VII is mainly immunolocalized at the zonules, and around the blood vessels at the bases of ciliary processes. COL7A1 mRNA is expressed in human PE and NPE cells, at low levels.

The zonules are “woven” by the NPE during organogenesis,18 when fine zonular fibers originate at the intercellular spaces of the NPE,19 aggregate into zonules, and then transverse the ILM. With aging, the ILM is infolded into those intercellular spaces,16 which corresponds to the “invaginations” we currently describe. Upon their lens capsule penetration, the zonules resolve into broadly fanning microfibrils,20 which anchor to “poorly defined structures” near the epithelial cells.21,22 This anchoring fashion withstands the repetitive mechanical strain of accommodation.23 By Col VII labeling, we underline that the zonules penetrate the lens capsule, transverse intracapsularly, and represent the “linear densities” (Supplementary Fig. S2A) of the anterior and equatorial lens capsule.17 Col VII might support the microfibrillar interaction at either zonular terminus.

Zonules were labeled by pAb(16) in cryo and paraffin slides, but not by the other antibodies. Because monoclonal antibodies were able to detect Col VII by Western blot, Col VII epitopes might be unavailable when embedded within the zonular matrix, or due to steric hindrance of the antibodies. Because pAb(16) has several binding sites for Col VII, such restrictions may apply to a lesser extent, thus possibly explaining different staining patterns. To assess the origin of zonular Col VII, we obtained microarray data from ex vivo NPE and PE cells. Their COL7A1 mRNA expression profiles, combined with the data from Western blots, add to our IHC results. Our hypothesis of a possible role of Col VII in the accommodation system is therefore supported. The low levels of COL7A1 expression at detectable Col VII protein amounts suggest a low turnover (e.g., in comparison with skin)23.

The demonstration of Col VII at ciliary stromal cells, PE, and blood vessels might relate to the repetitive mechanical strain these tissues are subjected to. The contact points between PE and the blood vessels at the ciliary base, for example, are intensely labeled. The ciliary epithelia need a rich vasculature for their active secretory functions. Unaided, such vascular accommodation might diminish the structural tenacity of the processes. Interestingly, accommodation system histopathology has not been reported in Col VII-deficient patients.7,8

By Western blot, the intense zonular IHC labeling appears to be due to the presence of NC-1 globuli. Although isolation of the relative minute quantities of Col VII from any extracellular matrix can be complicated due to its firm embedding therein, mAb(1-i) was able to detect a full-length Col VII (290-kDa) signal in zonular/lens capsule lysates. However, none of the mAbs was able to target zonules in sections successfully, in contrast to ciliary body and cornea. Thus, the IHC/IEM results obtained with pAb(16) could not be completely reproduced with other antibodies. We have tried to validate pAb(16) in various ways. The pAb(16) is frequently used in Col VII investigations, and does not show significant cross-reactivity to Col VII-deficient human12 or mouse tissues24 or keratocytes.25

Interestingly, convincing anchoring fibrils were not visualized, although some looped shapes of thin fibrils were situated between anti-Col VII-labeled structures (Supplementary Fig. S2C). Still, the function of Col VII might be established indirectly, through determination of interacting proteins. Proteomic studies have detected Col VII in human ciliary body,11 ILM,11,26 retinal blood vessel,11 and bovine zonule (Col VII fragment C9JBL3_HUMAN27) samples, but the established dermal Col VII interactors (i.e., type I and IV collagen, laminin 332) were detected incompletely (ciliary body,26,29 zonule,27 lens capsule11). Interactions of Col VII and fibrillin, the main component of zonules and part of a tissue mechanosensing complex,30 have not been documented.

To address the potential functions of intraocular Col VII further, the discrepancy between Col VII labeling in the absence of anchoring fibrils would be supported by identifying the functional ocular Col VII isoforms and possibly through thorough ophthalmological examination of patients with recessive dystrophic epimysolysis bulbosa.

**Acknowledgments**

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**References**


