Three-dimensional structure of the mammalian limbal stem cell niche

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1. Introduction

The limbal stem cell niche, situated at the anatomic border of the conjunctiva and the cornea (Levis et al., 2013; Eberwein and Reinhard, 2014, Rama et al., 2010), provides a microenvironment that contributes to the development and maintenance of the limbal epithelium. The existence of the limbal stem cell (LSC) niche is accepted, precise knowledge of its threedimensional (3D) architecture remains incomplete. The LSC niche was explored on freshly excised and organ-cultured corneoscleral rims from human donors and organ-cultured corneoscleral rims from organ-cultured corneas from human donors (n = 47), pigs (n = 15) and mice (n = 27) with full-field optical coherence microscopy (FFOCM). Limbal crypt features were detected in 90% of organ-cultured human corneoscleral rims, extending between the palisades of Vogt as radially oriented rectangular (74% of eyes) and/or rounded (23% of eyes) forms, often branching off to, or becoming interconnected by, sub-scleral radially or circumferentially oriented crypts (in 56% of eyes). Mean crypt volume represented 16% of sampled limbal volume on the vertical axis and 8% on the horizontal axis. In pigs, palisades were finer and crypts wider with relatively uniform distribution around the eye, and radial orientation, connecting to numerous narrow criss-crossing invaginations beneath the scleral surface. In mice, only a circumferential limbal trough was detected. Mean crypt volume represented 13% of sampled limbal volume in humans and 9% in pigs. FFOCM combined with fluorescence microscopy, showed presence of p63+ cells and cytokeratin-3+ cells in the limbal crypts. To assess colony forming efficiency (CFE), limbal epithelial cells were cultured at low density with full-field optical coherence tomography (FFOCM). Limbal epithelial cells were cultured at low density with mitomycin-arrested 3T3 feeders. CFE increased with limbal crypt volume and was not significantly decreased in organ-cultured cornea, despite degradation of the epithelial roof, suggesting that stem cells remain protected at the base of crypts during organ culture. CFE in human samples was significantly greater than in pig, and CFE in mouse was zero. Crypt architecture in the three species appears associated with eye exposure to light. LSC density increased with percentage limbal volume occupied by crypts.
are necessary to repopulate the regenerating corneal epithelium (Schlotzer-Schrehardt et al., 2007).

The existence of the limbal niche is accepted, but precise knowledge of its 3D architecture remains elusive (Dziasko et al., 2014). Exploration of the architecture of the corneal limbus and the limbal crypts has been carried out using different techniques, with different structures found (Goldberg and Bron, 1982; Townsend, 1991; Dua et al., 2005; Shortt et al., 2007; Shanmuganathan et al., 2007; Miri et al., 2012; Lathrop et al., 2012; Molvær et al., 2013). Both Goldberg and Bron (1982) and Townsend (1991) examined the corneal limbus with a slit lamp and described the palisades of Vogt as a series of radially oriented fibrovascular ridges, concentrated along the upper and lower corneoscleral limbus, separated by interpalisade epithelial rete ridges. They found high variability of the palisade zone from one individual to another and within the same eye, along with high variability of the shape of palisades and interpalisade epithelial crypts, including radially oriented rectangular and/or circular or oval forms, and branching or interconnection of the palisades to produce a trabecular pattern. Indeed, the high variability led Goldberg and Bron (1982) to conclude that the limbal palisade pattern is as individual as a fingerprint. Limbal crypt structure has been explored using histology (Yu et al., 2005; Shanmuganathan et al., 2007; Nunile et al., 2013; Molvær et al., 2013), confocal fluorescence microscopy (Shortt et al., 2007; Yeung et al., 2008), electron microscopy (Dua et al., 2005; Shanmuganathan et al., 2007; Shortt et al., 2007) in vivo confocal reflectance microscopy (Shortt et al., 2007; Miri et al., 2012; Deng et al., 2012) and optical coherence tomography (OCT) (Bizheva et al., 2011; Lathrop et al., 2012). Shortt et al. (2007) documented the portion of the limbus located toward the cornea using confocal and electron microscopy, and described the presence of both limbal crypts (LCs) between the palisades of Vogt and focal stromal projections (FSPs) at the corneal edge of the limbus, extending in a fingerlike pattern from the palisades and hence appearing circular/oval in the en face imaging plane of confocal microscopy. Dua et al. (2005) and Shanmuganathan et al. (2007) used histology to document the existence of larger, less frequent crypt structures that they termed limbal epithelial crypts (LECs) that contained radial, circumferential and oblique interconnecting components and descended from the epithelium to beneath the scleral surface. Confocal reflectance microscopy, using for example the HRT II with the Rostock module (Heidelberg, Germany) can perform non-invasive imaging of the limbal region, in ex vivo or in vivo tissues (Miri et al., 2012), but its small field size (300–400 μm × 300–400 μm) prevents viewing of the whole of the limbal region in depth in a single acquisition, and cross-sectional information is not available. In addition, histology and electron microscopy are invasive and require fixation, staining, and slicing of tissue. OCT imaging, along with confocal microscopy, of the 3D structure of the limbus in fixed human corneoscleral rims (Lathrop et al., 2012) revealed a combination of the structures described in the literature including the variety of palisade and interpalisade patterns with their intra and inter-individual variability, along with structures that may correspond to LCs, LECs and FSPs. The current study sought to assess stem cell presence in relation to morphology by exploring the 3D structure of the limbal crypts using FFOCM, a technique that allows non-invasive micrometer resolution exploration of the 3D volume in unfixed, unsliced ex vivo tissue without staining.

OCT is an imaging modality analogous to ultrasound but using light reflected from structures inside tissue (Huang et al., 1991). Conventional OCT is widely used in ophthalmology to image both the anterior and posterior segment. Axial resolution is typically on the order of 5 μm, and lateral resolution on the order of 10–20 μm. FFOCM (Beaurepaire et al., 1998; Dubois et al., 2002) is a variant of conventional OCT in which two dimensional (2D) en face images are captured on a camera and three-dimensional (3D) data sets may be obtained by scanning in the depth direction. This configuration and the use of a white light source allow for higher axial and transverse resolution than conventional OCT, on the order of 1 μm. No contrast agents are required as contrast is entirely endogenous. FFOCM can therefore perform micrometer resolution 3D imaging non-invasively in fresh or fixed ex vivo biological tissue samples. FFOCM has been used to image ophthalmic tissues including cornea (Ghouali et al., 2014; Akiba et al., 2007), lens and retina (Grieve et al., 2004), as well as skin (Dalimier and Salomon 2012), brain (Assayag et al., 2013a), breast (Assayag et al., 2013b) and gastrointestinal tissues (Grieve et al., 2014).

Here FFOCM was applied to imaging of the limbal region of the corneoscleral rim of humans, pigs and mice, in order to further understanding of the 3D architecture of the limbal stem cell niche in unfixed tissues in these three species.

2. Materials and methods

2.1. Tissues

This study was carried out according to the tenets of the Declaration of Helsinki and followed international ethical requirements for human tissues, and was approved by our local institutional review board for human experimentation. Forty seven human corneoscleral rims were imaged, of which 21 were from males, 18 females, 8 sex unknown, with an average donor age of 71 (range 40–93). Of the 47 corneoscleral rims imaged, 43 were organ-cultured and 4 fresh, of which 9 (7 organ-cultured and 2 fresh) were cultured for colony forming assays (CFA).

The forty three human cadaveric corneoscleral rims from donors originated from the Établissement Français de Sang (EFS) cornea bank at Saint-Antoine Hospital, Paris, and were retained after the central corneas had been used for keratoplasty procedures on patients (8-mm trephination of the donor tissue). The donor corneas were preserved in CorneaMax (EuroBio, France) medium for a maximum of 35 days at 31 °C, in accordance with European Eye Banking regulations. They were then placed in Corneajet (EuroBio, France) medium containing Dextran for deturgescence 48 h prior to transplantation, and transferred to the Quinze-Vingts Ophthalmology Hospital where grafts were performed. The corneoscleral rims were retained in Corneajet (EuroBio, France) medium and imaged within 24 h. Following imaging, they were transferred to the cell culture laboratory.

The four fresh cadaver donor corneas were obtained from the Surgery School of the Assistance Publique – Hôpitaux de Paris (AP-HP). These were imaged <24 h post-mortem, then transferred to the cell culture laboratory.

All animal manipulation was performed in accordance with the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research, and was approved by our local institutional review board for animal experimentation.

Fifteen pig eyes (albino English – pigmented Belgian cross-breed) were obtained from a private research facility. Eyes were removed within one hour after sacrifice of the animal and stored in phosphate buffered solution (PBS). On arrival at the FFOCM imaging location (<4 h post-mortem) they were transferred to DMEM medium and dissected for imaging. The corneoscleral button was removed and relief incisions made in the sclera in order to flatten the cornea for imaging. Of the fifteen pig eyes imaged, all were imaged fresh and one placed in organ-culture for 31 days in a flatmount position, and 3(2 fresh, plus the one organ-cultured) were cultured for CFA.
Twenty-seven mouse eyes (C57 BL/6, Janvier labs, France) were obtained from the animal facility of the Vision Institute. Of the 27 mouse eyes imaged, 21 were fresh, unfixed tissue prepared for the current study, while six were recuperated from an unrelated study and had been fixed using 4% paraformaldehyde (PFA) and mounted on microscope slides. Mice were deeply anesthetized by pentobarbital and sacrificed by cervical dislocation. After sacrifice, the eyes were immediately removed and dissected to flatten mount the cornea for imaging. Of the fresh mouse eyes 8 were organ-cultured for 31 days in a flatmount position, imaged, and then also cultured for CFA. Two of the fresh mouse eyes were organ-cultured for 31 days in a flatmount position, imaged, and then also cultured for CFA.

To resume, 47 human (4 fresh, 43 organ-cultured), 15 pig (15 fresh, of which 1 subsequently organ-cultured) and 27 mouse (22 fresh, of which 2 subsequently organ-cultured, 6 fixed) were cultured for CFA. Of the 9 human, 3 pig and 10 mouse samples were cultured for CFA.

### 2.2. FFOCM

The FFOCM instrument used is a commercial system (LI-Tech, France). It has been described previously (Ghouali et al., 2014). Illumination is provided by a halogen source, whose short coherence length leads to an axial resolution of 1 μm. The full field is illuminated and images are captured on a complementary metal oxide semiconductor (CMOS) camera. The interferometer arms hold a matched pair of microscope objectives in the Linnik configuration. 10× water immersion objectives with 0.3 numerical aperture lead to a lateral resolution of 1.6 μm. Penetration depth depends on tissue content and transparency and ranges from approximately 200 μm in sciera to 1000 μm in cornea. Wide field images can be displayed by automated mosaicing to image samples up to 25 mm in diameter, or stacks of images in depth can be captured. A 2 × 2 mm² stitched field is captured in less than five minutes, and a 200 μm × 1 mm × 1 mm 3D stack is also captured in less than five minutes.

### 2.3. Image acquisition

For all species, wide field en face FFOCM images of the whole of each corneoscleral rim were captured along with 3D stacks of images at four equally spaced sampling points around each corneoscleral rim. Each of these stacks could be navigated in multiplanar, video or 3D mode in order to view the crypt structure. A photograph of each corneoscleral rim was also saved.

The wide field stitched image of each corneoscleral rim was captured close to the surface at a depth where the palisades of Vogt were most clearly visible (Fig. 1A). The four wide field image stacks (of depth × width × length of 150 μm × 2 mm × 4 mm in human, 200 μm × 2 mm × 4 mm in pig, and 150 μm × 800 μm × 800 μm in mouse) were then captured at top, bottom, left and right regions of the corneoscleral rim (Fig. 1A–E), to provide the most complete coverage possible of the limbal region in a reasonable timeframe that yielded manageable data sets.

An incision was made in the sciera to mark 0° for the imaging data, and on transfer to the cell culture laboratory, the corneoscleral rim was cut into eight 45° segments for human and pig, numbered one to eight in a clockwise fashion. Each image data volume straddled two of the segments, so providing a sampling of crypt volume in each cultured explant. The protocol of donor cornea collection for the French Eye Banks and the Surgery School does not include recording information on the orientation of the cornea. In order to determine vertical (superior/inferior) and horizontal (temporal/nasal) axes, the long and short axes of the anterior surface of each cornea were measured on the photographs. Crypt distribution around the 360° circumference of the cornea could therefore be related to location on the vertical (superior/inferior) or horizontal (temporal/nasal) axes.

In mouse, each whole cornea was divided into four explants, each correlated to an image stack. Eye orientation was not recorded.

### 2.4. Image analysis

#### 2.4.1. Qualitative assessment of crypt-richness of entire corneoscleral rim

The wide-field FFOCM images (Fig. 1A) at a single depth typically 30 μm below the surface in organ-cultured human samples, 50 μm in fresh human samples (where intact, thick epithelium meant the palisades of Vogt were located deeper than in organ-cultured samples), 70 μm in pigs and 10 μm in mouse were assessed to qualitatively determine overall visibility of limbal crypt structures and the palisades of Vogt in each corneoscleral rim, and also to identify potential crypt-rich areas within each corneoscleral rim. The wide-field image depth tended to lie toward the surface of the palisades of Vogt and the interpalisade epithelial crypts, where the typical limbal palisade patterns were most easily recognized, while sub-scleral crypts tended to be deeper. Morphological crypt features including shape (rectangular or rounded) and orientations (radial or circumferential) were tallied for all samples to give a percentage of overall number of samples with a particular feature visible.

#### 2.4.2. Limbal crypt volume

Image processing software (Fiji, National Institutes of Health, USA) was used to perform thresholding and area quantification on slices averaged in 10 μm depth bundles in each 3D image stack. Volumes were expressed as the percentage of each stack occupied by crypts:

\[
%V = 100 \times \left( \sum \text{SA} \times 10 \right)/v
\]

where \( \text{SA} = \) surface area occupied by crypts in each 10 μm en face bundle, over volume \( v = \) depth × width × length of 150 μm × 2 mm × 4 mm in human, 200 μm × 2 mm × 4 mm in pig, and 150 μm × 800 μm × 800 μm in mouse.

#### 2.5. Fluorescent labeling of epithelial and stem cells

As the contrast mechanism in FFOCM is intrinsic, due to changes in refractive index, fluorescent labels are not detected in FFOCM. In order to demonstrate the presence of stem cells and epithelial cells in the limbal crypt architecture imaged by FFOCM, we labeled and imaged specimens using fluorescence microscopy. We used both an Olympus laser scanning microscope FV-1200 (Olympus France, Rungis) for confocal fluorescence imaging of labeled specimens, with overlay of the fluorescence and FFOCM images in post-processing, and a prototype multimodal optical microscope that allows simultaneous co-located FFOCM and fluorescence imaging. Compared to previous fluorescence-FFOCM setups (Harms et al., 2012; Aukorius et al., 2012), in the set-up used here, fluorescence is acquired simultaneously with the FFOCM measurements, meaning image capture is faster.

Briefly, this set-up operated in a similar fashion to conventional FFOCM but with the addition of a blue light emitting diode (LED) (Thorlabs, M470L2, 650 mW) centered at 470 nm blocked by an excitation filter (λ = 500 nm, Thorlabs, FES0500) to excite fluorophores. The two illumination beams are combined and focused onto the sample, and in the detection path, FFOCM and fluorescence signals are separated by a Single Edge Dichroic Beamsplitter
(BrightLine, $\lambda = 593$ nm, Di02-R594-22x27), and captured by two cameras, a CMOS from PhotonFocus for the FFOCM image and a scientific CMOS from PCO.edge to capture the fluorescence image, with additional filters placed in front of the detection cameras to ensure independence of the two paths. As the optical path into the tissue is parallel for the FFOCM and fluorescence channels, an
were not contiguous. The epithelial colonies were identified with a whole mount immunofluorescence assay. Briefly, after fixation with 4% PFA for 1 h, the cells were permeabilized and blocked with 0.5% triton X-100 and 0.2% gelatin in PBS for one night at room temperature followed by incubation with the anti cytokeratin 3 antibody (1:100) (monoclonal antibody Clone AE-5; Dako Trappes, France) and anti p63-α polyclonal antibody (1:50) (cell signaling technology, France) for 24 h at room temperature, and then with the secondary antibody (Alexa Fluor 488, 1:600) for 2 h at room temperature, and nuclei were counterstained with DAPI. One segment was incubated with phosphate-buffered solution (PBS) instead of the primary antibody to serve as a negative control.

Fluorescence-FFOCM images were taken of all three segments using the multimodal system. As the fluorescence-FFOCM set-up used allows imaging in only the green wavelength band, the Olympus confocal fluorescence microscope was preferred when we wished to label several cell types at once.

2.6. Colony forming assays

After FFOCM analysis, 9 human corneoscleral rims (2 fresh, 7 organ-cultured) were divided into 8 explants per rim, 3 pig corneoscleral rims (2 fresh, 1 organ-cultured) into 8 explants per rim, and 10 mouse whole corneas (8 fresh, 2 organ-cultured) into 4 explants per cornea. According to the protocol described in Ghoubay-Benallaoua et al., 2013, explants were digested using 1 mg/ml collagenase A (Roche Diagnostics, Mannheim, Germany) in cholera toxin-free Green medium at 37 °C for 18 h. Limbal epithelial cells were collected from the incubated tissues and dissociated into single cells by pipetting. The clonal growth ability of cultured limbal epithelial cells was evaluated by determining CFE. Swiss albino murine 3T3 fibroblasts (ATCC, Molsheim, France) were treated with 4 μg/ml mitomycin C for 2 h and then trypsinized and plated at a density of $2 \times 10^4/cm^2$ onto six-well culture plates. Enzymatically dissociated cells were seeded at low density (1000 cells/well) in six-well culture plates on 3T3 fibroblasts feeder layers (Nakamura et al., 2004). Cultures were incubated at 37 °C under 5% CO2. They were observed three times a week by inverted microscopy though had a greater quantity of intact morphological features though had a greater quantity of intact cellular details (Fig. 2, Supplementary movie 3). Indeed, under sufficient fluorescence-FFOCM images could be related to location on the vertical (superior/inferior) or horizontal (temporal/nasal) axes by measuring the short (vertical) and long (horizontal) axes in each cornea (Fig. 1A–E).

Crypts were detected on two opposing sides in 31% of eyes, on two neighboring sides in 18% of eyes, on 3 or 4 sides in 10% of eyes and on only one side in 41%. Average angular spread of crypts was followed to their epithelial outlet (Fig. 1 P, Q, R, Supplementary movie 2). Sub-scleral caverns could usually be followed to their epithelial outlet (Fig. 1 P–R). On occasions where these sub-scleral caverns appeared to be isolated in surrounding sclera, their path to the epithelium may have passed outside of the imaged zone.

Supplementary video related to this article can be found at http://dx.doi.org/10.1016/j.exer.2015.08.003.

3.1.1. Description of crypt morphology

3.1.1.1. Organ-cultured corneas. Limbal crypts were observed in 90% of the organ-cultured human corneoscleral rims. Morphological description tallies indicated that crypt pattern including shape and orientation was highly variable from one individual to another and from one region of the limbus to another in the same subject. In en face views, rectangular shaped, radially oriented interpalisade crypts were seen in 74% of eyes (Fig. 1F–J), rounded interpalisade crypts in 23% of eyes (Fig. 1K, L), with interconnecting circumferential crypts in 56% of eyes (Fig. 1I, J). Multiple crypt forms and orientations were seen in the same eye in 75% of eyes. Between the palisades of Vogt, crypts rose to the surface at the cornea, corresponding to LCs (interpalisades). Palisades often terminated in FSPs (Fig. 1M, N). On moving outward to the sclera, crypts then invaginated beneath the scleral surface (Fig. 1O–R) corresponding to LECs. In cross-sectional views, crypts appeared toward the corneal edge as simple valleys between the palisades of Vogt (seen in 79% of eyes), which on moving toward the sclera became enlarged with a narrower epithelial outlet (seen in 41% of eyes), and then invaginated to a cavern beneath a scleral roof at the scleral side of the limbus (seen in 40% of eyes) (Fig. 1H, J, P, Q, R, Supplementary movie 2). Sub-scleral caverns could usually be followed to their epithelial outlet (Fig. 1 P–R). On occasions where these sub-scleral caverns appeared to be isolated in surrounding sclera, their path to the epithelium may have passed outside of the imaged zone.

3.1.1.2. Fresh corneas. Little difference was observed in the crypt morphology between fresh and organ cultured specimens, aside from epithelial degradation. Fresh cadaver corneas had similar morphological features though had a greater quantity of intact epithelium, so maintaining a thicker epithelial roof with visible cellular details (Fig. 2, Supplementary movie 3). Indeed, under superficial and wing cells layers, small, highly scattering basal cells with large nuclei to cytoplasmic ratio could be identified clinging to palisade walls and in sub-scleral caverns.

3.1.2. Analysis of crypt distribution, area and volume around the limbus

Crypt distribution around the 360° circumference of the cornea could be related to location on the vertical (superior/inferior) or horizontal (temporal/nasal) axes by measuring the short (vertical) and long (horizontal) axes in each cornea (Fig. 1A–E).

Crypts were detected on two opposing sides in 31% of eyes, on two neighboring sides in 18% of eyes, on 3 or 4 sides in 10% of eyes and on only one side in 41%. Average angular spread of crypt
clusters was 50\(^\circ\) (range 15\(^\circ\)–120\(^\circ\)) with an average of 2 clusters per cornea (range 1–4) for all corneas with visible crypts. Average crypt depth in humans was 50 \(\mu\)m (range 35 \(\mu\)m–120 \(\mu\)m) and width 50 \(\mu\)m (range 15 \(\mu\)m–150 \(\mu\)m).

Limbal crypt volume per quadrant represented on average 13.0\% (range 0–62\%) of the limbal volume sampled (2 \(\text{mm} \times 4 \text{ mm} \times 150 \text{ \(\mu\)m}). This figure was 15.8\% for the vertical location and 8.1\% for the horizontal location (\(p = 0.007\)). The 95\%-confidence interval of the limbal crypt volume per cornea was 0.0\%–29.9\%.

\(V\) was significantly greater (\(p < 0.0001\)) in fresh (39.8\%, range 23\%–56\%) than in organ-cultured tissue.

No correlation of \(V\) with age or age group (in 10 year groups) was observed in this study (\(p = 0.29\)).

### 3.1.3. Fluorescent labeling of cells

Fluorescent labeling confirmed presence of p63\(^{+}\) stem cells and epithelial cells with corneal differentiation (i.e., cytokeratin 3 \(\pm\) cells) in the crypt architecture (Fig. 3). These are seen to be coincident with the crypt structures viewed with FFOCM (Fig. 3A–D). Controls were negative (Fig. 3E, F). Imaging with confocal fluorescence microscopy allowed merged viewing of stem cells with other cell populations, showing that both stem and epithelial cells are present in human limbal crypts (Fig. 3G, H), though not colocalized.

### 3.1.4. Correlation between morphology and clone production

Colonies grown from dissociated limbal epithelial cells were associated with areas visibly rich in crypts (Fig. 4). The CFE(\%) significantly increased with limbal crypt volume (\(V\)), per segment (\(r_e = 0.42, p = 0.001\)) and per cornea (\(r_e = 0.82, p = 0.02\); Fig. 4B). It should be noted that large \(V\) over the sampled area does not necessarily mean large crypts; large \(V\) can equally mean numerous small crypts.

The average CFE(\%) was 1.8\% overall, of which 2.4\% was on the vertical axis and 1.4\% on the horizontal axis (\(p = 0.17\)).

In fresh cadaver corneas, CFE(\%) was 2.1\% overall, which was not significantly greater than in organ-cultured cornea (\(p = 0.67\); Fig. 4D).

### 3.2. Pig

In pig eyes (Fig. 5B), the palisade zone was located further toward the scleral than the corneal side of the limbus. Palisades were fine and interpalisade crypts wide and oval, in a uniform radial orientation, connecting to numerous narrow invaginations beneath the sclera in a trabecular pattern. Distribution was relatively uniform and consistent around the 360\(^\circ\) of the eye as well as having similar depth (50 \(\mu\)m), width (90 \(\mu\)m), shape (oval) and orientation (radial) in all eyes. The limbal zone was 1.5 mm wide.

Limbal crypt volume represented on average 9.5\% (range 3–21\%) of the limbal volume sampled (200 \(\mu\)m \(\times\) 2 mm \(\times\) 4 mm). This figure was 9.2\% for the vertical location and 9.8\% for the horizontal location (\(p = 0.67\)).

Fluorescent labeling revealed p63\(^{+}\) cells contained within crypts in pig (Fig. 3I, J). CFE(\%) in fresh pig samples was 0.59\% overall, of which 0.45\% was on the vertical axis and 0.70\% on the horizontal axis (\(p = 0.17\)).

Organ-cultured pig cornea, after 31 days in organ culture conditions, showed epithelial degradation including detachment and undulations that increased surface scattering from the tissue, thereby decreasing penetration depth of the FFOCM images. The limbal crypt structure was faintly visible beneath the damaged epithelium and the palisade/interpalisade pattern appeared intact, though quantification was not possible due to the degraded image quality. CFE(\%) in organ-cultured pig cornea was zero. This was not significantly different from CFE(\%) in fresh pig cornea (Fig. 4D).

### 3.3. Mouse

In fresh mouse cornea, the palisades of Vogt were not observed and the limbal region contained only an epithelial trough circumferentially surrounding the cornea (Fig. 5C). \(V\) was therefore evaluated at zero for all mouse samples. Architecture of the limbus appeared unchanged after 31 days in organ culture, though the epithelium had thinned and surface cells were misshapen.

Fluorescent labeling with DAPI and p63\(^{+}\) using confocal fluorescence microscopy showed clumps of p63\(^{+}\) cells present in the limbal trough, along with fewer isolated p63\(^{+}\) cells distributed throughout the central cornea (Fig. 3K, L, M). CFA was unsuccessful in both fresh and organ-cultured mouse cornea, yielding a CFE(\%) of zero for all samples.

### 3.4. Interspecies variability

Morphology of the limbal stem cell niche differed between species (Fig. 5).

CFE(\%) according to tissue characteristics is shown in Fig. 4D. It was significantly higher in human tissues (either fresh or organ-cultured) than in pig and mouse tissues (\(p < 0.01\)). Differences between pig and mouse tissues were not significant (\(p > 0.2\)). For all 3 species there were no significant differences in CFE(\%) between fresh and organ-cultured tissues (\(p > 0.3\)).

### 4. Discussion

#### 4.1. Crypt architecture

Crypt structures were detected in 90\% of the human samples examined. This is in agreement with the reported value of visibility of the palisades of Vogt in 80–90\% of subjects (Townsend, 1991; Patel et al., 2006; Dua et al., 2006). Visibility of palisades has been shown to be lower in non-pigmented individuals (Miri et al., 2012) and to decrease with age (Goldberg and Bron, 1982). The corneoscleral rims we obtained had an average age of 71 (range...
Fluorescent labeling of stem cells. A, C, E: FFOCM images of crypts and stem cells (in green, D) in the crypt architecture. Controls (E, F) showed only a uniform noise background in the fluorescence channel (F). G, H: human limbus imaged with a confocal fluorescence microscope with an X10 objective (G) and zoom (H) showing p63+ cells (green), cytokeratin 3+ (red) and DAPI (blue) staining, where the sclera lies to the left and the cornea to the right of the image. p63+ cells are visible in the base of a few crypts, while cytokeratin 3+ cells are not colocalized with p63+ cells, and are present in greatest number in higher imaging planes toward the sample surface and toward the cornea. I, J: porcine cornea imaged with a confocal microscope with an X10 objective (I) and a X20 objective (J) showing p63+ cells (green), and DAPI (blue) staining, where the sclera lies to the left and the cornea to the right of the image. p63+ cells are visible within crypts. K: portion of a flat mounted mouse cornea at 5×, marked with p63+ and DAPI, showing location of central cornea (yellow star) and sclera. L: zoom on limbal region at 20× shows clumps of bright p63+ cells, and M: at 20× some p63+ cells distributed throughout central cornea.

Fig. 3. Fluorescent labeling of stem cells. A, C, E: FFOCM images of crypts (260 μm × 260 μm field size) from the fluorescence-FFOCM setup with the fluorescence channel switched off. B, D, F: With the simultaneous fluorescence imaging, the fluorescence-FFOCM combined images show presence of epithelial cells (in green, B) and stem cells (in green, D) in the crypt architecture. Controls (E, F) showed only a uniform noise background in the fluorescence channel (F). G, H: human limbus imaged with a confocal fluorescence microscope with an X10 objective (G) and zoom (H) showing p63+ (green), cytokeratin 3+ (red) and DAPI (blue) staining, where the sclera lies to the left and the cornea to the right of the image. p63+ cells are visible in the base of a few crypts, while cytokeratin 3+ cells are not colocalized with p63+ cells, and are present in greatest number in higher imaging planes toward the sample surface and toward the cornea. I, J: porcine cornea imaged with a confocal microscope with an X10 objective (I) and a X20 objective (J) showing p63+ cells (green), and DAPI (blue) staining, where the sclera lies to the left and the cornea to the right of the image. p63+ cells are visible within crypts. K: portion of a flat mounted mouse cornea at 5×, marked with p63+ and DAPI, showing location of central cornea (yellow star) and sclera. L: zoom on limbal region at 20× shows clumps of bright p63+ cells, and M: at 20× some p63+ cells distributed throughout central cornea.

3D navigation of our data sets to the depths at which this contrast inversion was observed, we suggest that the dark regions below palisades are caused by shadowing from the palisades on the surface, and at this depth we are in fact viewing only sclera. We identified interlinking dark structures towards the conjunctiva, at depths between 0 μm and 120 μm, as crypts rather than palisades.

While Goldberg and Bron (1982) and Townsend (1991) used slit lamp examination to describe the morphology of the limbal region as containing a variety of palisade and interpalisade patterns and forms, more precise microscopic morphology described by Shortt et al. (2007) included the presence of both LCs between the palisades of Vogt and FSPs at the corneal edge of the limbus, while Dua et al. (2005) and Shanmuganathan et al. (2007) describe the existence of larger, less frequent crypt structures that they term LECs that contained radial, circumferential and oblique components and descended beneath the scleral surface. In the current study, all of the features previously described were seen. FSPs were less frequently detected as distinct features than in previous studies (Shortt et al., 2007; Miri et al., 2012), possibly due to epithelial degradation in organ-cultured cornea making them appear simply as terminations of the palisades. Miri et al., 2012 used laser scanning confocal microscopy to image the limbal zone in vivo, in depth in 46 subjects. They describe a similar crypt architecture to that described here on the surface, however in depth, they suggest that palisades and interpalisade crypts (or epithelial reticulum) inverse their contrast, so that the structures which interlink the network beneath the conjunctival surface are identified as the palisades.

The precise non-invasive 3D imaging performed using FFOCM provides information on the morphology of the limbal crypt network on unfixed, unstained, whole tissue samples. This morphological information could be valuable in the design of artificial molds for stem cell culture for artificial limbus fabrication (Levis et al., 2013). Our quantitative assessment of the limbal crypts and although information on each individual's racial origin was not supplied with the donor corneas, racial origin of donors is predominantly European-Caucasian (non-pigmented) in the French Eye Bank network. This may explain why palisades were not detected in 10% of samples imaged.

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showed that stroma represented the most important component of the niche volume with the epithelial component representing less than 20%.

The data on pig and mouse crypt architecture indicate the differences between species. Radially oriented crypts extending beneath the scleral roof were found in humans and pigs but not in mice. This architecture provides protection from visible light and ultraviolet irradiation. Interestingly, the former species have a diurnal activity whereas the latter has a nocturnal activity. Ultraviolet (UV) B radiation induces damage to the human corneal...
epithelium, such as epithelial sloughing, decreased mitotic rate, and decreased Langerhans cell density (Borderie et al., 1996; Haaskjold et al., 1993). Interestingly, the porcine limbus has been shown to share features with the human limbus such as high CFE achieved with epithelial cells harvested from crypt regions and expression of the putative stem cell markers p63-a and integrin β1 (Notara et al., 2011). The absence of regional variation of crypt volume in pigs as compared to humans could be explained by the different orientation of the eyes with respect to the face. The exposure of pig eyes due to their position on the side of the head and the lack of brow may mean that there is less variation in UV and visible radiation over the 360° of the eye. In the nocturnal mouse, protection from light has less importance, and this may be the reason that the stem cells are distributed in a limbal trough close to the epithelial surface as well as distributed throughout the central cornea (Majo et al., 2008). Unsuccessful CFA in mouse by the method proposed for human and porcine samples is consistent with the literature (Majo et al., 2008).

4.2. Fresh versus organ-cultured samples

Organ culture of human donor corneas is known to induce epithelial sloughing. After the deswelling period in dextran-supplemented medium, the corneal epithelium is reduced to two cell layers (Borderie et al., 1995). The remaining epithelial cells appear flattened. They feature normal endoplasmic reticula, glycogen, mitochondria and nuclei, presence of hemidesmosomes on the basal cell membrane of the basal epithelial cells and desmosomes in the epithelium. The epithelial basement membrane is normal (Borderie et al., 1997). In the present study, 3% was reduced in organ-cultured tissue but CFE was preserved. Reduction in crypt volume in organ-cultured corneas may be explained by the reduced thickness of the limbal epithelium in organ-cultured tissue, while the limbal epithelial progenitors, located in the bottom of crypts, are preserved, thereby maintaining CFE.

4.3. Correlation of fluorescent labeling and FFOCM images

Use of an FFOCM system coupled with a fluorescence channel, that allows coincident and simultaneous OCT and fluorescence images to be acquired, successfully demonstrated presence of stem cells and epithelial cells within the crypt structures detected by FFOCM. Limitations of the multimodal fluorescence-FFOCM system used included lack of depth sectioning and use of a single color wavelength band for excitation and detection, plus low sensitivity in the fluorescence channel, so that the fluorescent signal was weak or absent in comparison to images obtained with the Olympus confocal fluorescence microscope. For these reasons, we display images from both fluorescence-FFOCM and confocal fluorescence microscopy. Determining the depth of the stem cells could be achieved by adding sectioning capability to the fluorescence-FFOCM system, and merged images using multiple markers at different emission wavelengths could be achieved by adding multiple illumination paths. A system including these upgrades is currently being developed.

4.4. Targeted biopsy

Imaging using FFOCM can provide a quick assessment of whether a corneoscleral rim is rich in crypts, and therefore a good candidate for assessing clone production in culture. This assessment step could improve efficiency in the culture of artificial limbus, by determining optimum samples for CFA. This type of targeted biopsy was previously proposed by Shortt et al., 2007 using confocal reflectance microscopy. However we propose that FFOCM can provide such information more quickly and easily. Screening of donor corneas in tissue banks to choose the best candidates for limbal graft could also be performed with FFOCM. Proposed future developments of FFOCM technology include use of a camera allowing far more rapid image acquisition, so improving the efficiency of the technique for ex vivo imaging, but also potentially permitting in vivo imaging in the future (Grieve et al., 2005).