Influence of laser photocoagulation on choroidal capillary cytoarchitecture

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Abstract

Aim—To identify if laser photocoagulation induces morphological changes specifically related to the choroidal capillary endothelial processes that protrude into Bruch’s membrane.

Methods—Two human eyes and one adult macaque monkey eye received retinal laser photocoagulation that was just suprathreshold, before enucleation or exenteration. They were examined by electron microscopy to determine the length of the endothelial processes emanating from the choroidal capillaries in the region around the laser burn. One human and two monkey untreated eyes were used for comparison.

Results—In human eyes, there was no increase in the number of processes 15 hours after laser treatment but at 5 days the processes were more numerous and longer within 400–500 µm of the burn than in the untreated half of the same eye. The processes were longer 9 days after photocoagulation in the monkey, when compared with untreated monkeys, and some breached the elastic lamina, a phenomenon not seen in the untreated eyes. Qualitative differences were also noted in the endothelial cell processes following photocoagulation. Neovascularisation was not observed.

Conclusions—Protrusion of choroidal endothelial cell processes into Bruch’s membrane is a normal anatomical feature but the number, length, and morphology of the processes change following mild photocoagulation. It is plausible that these processes may play a part in the clearance of debris from Bruch’s membrane, and represent an early stage of angiogenesis. If the latter is true prophylactic laser photocoagulation at just suprathreshold levels may carry a risk of inducing choroidal neovascularisation.

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There is considerable interest in the possible long term beneficial effects of laser photocoagulation in patients with high risk drusen as part of age related maculopathy (ARM). The mechanism whereby photocoagulation induces resolution of the drusen is unknown, but it must account for the observation that disappearance of drusen occurs over many months, and that the process begins in close proximity to the burn and then spreads outwards over an ever increasing area of the fundus (Fig 1).

It has been suggested by others that processes of choroidal pericytes penetrate into Bruch’s membrane and may contribute to the clearance of drusen as a result of observations in primates following experimental photocoagulation. Moreover, it was implied that they may have a physiological role in clearing debris from Bruch’s membrane. It is well established that choroidal capillaries also have cellular processes that penetrate the outer portion of Bruch’s membrane. They have been described in normal eyes of many species, but they have received little attention until recently. Their physiological significance is uncertain. Similar structural specialisations of the endothelial cell have been identified in other tissues, and various functions have been attributed to them, such as stabilising the thin fenestrated vascular wall of the renal vas a recti, sensing extracellular matrix in kidney through chemoreceptors and mechanoreceptors, and phagocytosis in trabecular meshwork and lung.

A variety of potential functions has been proposed for these endothelial processes in the choroid. It has been suggested that they may be important to metabolic exchange between choroid and retinal pigment epithelium, or contribute to debris in Bruch’s membrane. It is also possible that they may influence the clearance of debris from Bruch’s membrane in humans. The number and size of the processes do not correlate with age, nor any index of age change in the RPE, such as autofluorescence of the RPE or residual body content of the RPE, although there is a weak linear association with Bruch’s membrane thickness. It has also been suggested that choroidal endothelial processes may represent an early phase of choroidal neovascularisation, and they have been described as a low turnover vascular remodelling process, which occurs in response to preceding vascular degeneration in the choroid. It is well recognised that focal erosion of basement membrane around endothelial cells, with subsequent protrusion of cell processes through these defects is a sign of nascent angiogenesis in oncology studies. It is therefore possible that the same processes are potential precursors of neovascularisation if the appropriate extracellular conditions prevail.
Given that resolution of drusen and choroidal neovascularisation may occur following photocoagulation, and that the presence of choroidal capillary endothelial cell processes may contribute to the therapeutic effect of laser photocoagulation, we sought evidence that laser photocoagulation influences their density and/or morphology.

Materials and methods
Three female patients (patient 1, 58 years old; patient 2, 45 years old; patient 3, 61 years old) undergoing exenteration as part of their treatment for meibomian gland carcinoma, consented to donate their normal globes to research. The research was approved by the ethics committee of Moorfields Eye Hospital. Before surgery, the eye of patients 2 and 3 had just suprathreshold photocoagulation applied with an argon laser to the inferior macula as part of their treatment for meibomian gland carcinoma, consented to donate their normal globes to research. The research was approved by the ethics committee of Moorfields Eye Hospital. Before surgery, the eye of patients 2 and 3 had just suprathreshold photocoagulation applied with an argon laser to the inferior macula using variables similar to those used in current prophylactic studies of ARM. Patient 2 had seven 200 µm diameter burns, of 200 ms duration and 140 mW, applied 15 hours before surgery, and patient 3 had twenty 200 µm burns, of 200 ms duration and 200 mW, applied 5 days before surgery. All three eyes were fixed within 15 minutes of exenteration in mixed aldehydes (2.5% paraformaldehyde and 2% glutaraldehyde) in 100 ml sodium cacodylate buffer pH 7.4 and prepared for electron microscopy. The eyes that had been photocoagulated had segments taken from the superior macula, at least 4 mm from any laser burn, to compare with specimens taken from the inferior macula that incorporated the laser lesions.

Eyes from three adult macaque monkeys were enucleated after perfusion fixation in paraformaldehyde and then emersion fixed as described above. One of the monkeys had 12 suprathreshold photocoagulation burns (200–400 ms duration and 200–400 mW) applied to the right eye only with a Keeler indirect diode laser, 9 days before sacrifice. The untreated eyes served for comparison.

All globes were hemisected by a circumferential incision through the pars plana and the posterior pole was isolated. Sections of the maculas were obtained from the central macula in eyes that were not treated and from areas equidistant from the horizontal raphe in the superior and inferior macula in eyes with laser burns. Those specimens taken from the treated half always included a visible laser lesion.
Tissues were dehydrated in acetone and propylene oxide, post-fixed in osmic acid, and embedded in Araldite for electron microscopy. Ultrathin sections were stained with uranyl acetate and lead citrate. Sections were examined with an electron microscope (Jeol), and the number of processes on the internal aspect choroidal endothelial cell per 100 µm length of Bruch’s membrane were counted, measured, and photographed. A process was only counted if it clearly transgressed the basal lamina of the choriocapillaris. Untreated eyes had 10 continuous lengths of 100 µm of Bruch’s membrane counted. In the photocoagulated eyes the number of protrusions were counted in 100 µm intervals from the laser burn to establish if the number of protrusions varied with distance from the injury. Two separate sections of the laser burns were examined in eyes from patients and nine sections were examined in the treated monkey. Over 1000 µm of Bruch’s membrane were examined in each of the human eyes and the monkey to arrive at an average number of processes per 100 µm with each 100 µm from the laser centre. The longest processes were also recorded for each eye. The presence and location of long spacing collagen (LSC) and focal thickening in the basement membrane were noted.

Poisson statistics were used to look at differences between the number of processes found in the untreated sections compared with the treated maculas since the number of processes in untreated eyes appear to conform to a Poisson distribution.

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<th>Subject</th>
<th>Untreated</th>
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<tr>
<td></td>
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<td>M3</td>
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Figure 2 Electron micrograph of choriocapillaris and Bruch’s membrane in human eyes. (A) An untreated area of patient 3 (final magnification 48 000), and (B) an 71 year old man (final magnification 50 000). These show focal thickening of basal lamina (small arrow) around the base of a cell process (curved arrow) associated with long spacing collagen (LSC) (thick arrow) in the outer collagenous zone (OCZ). (C) 22 year old man (final magnification 100 000), (D) an 82 year old woman (final magnification 80 000); these show a magnified appearance of the processes (thick arrow).
Results (Table 1)

UNTREATED REGIONS OF HUMAN EYES

In patient 1 (P1), who did not receive photocoagulation, the average number of endothelial cell processes per 100 µm was 5.7 with a range of 3.0–8.3. The longest individual process was 1.0 µm.

In the superior half of the macula of patient 2 (P2) in which no photocoagulation had been undertaken there was an average of 6.5 processes per 100 µm (3.3–9.1). The longest individual process was 0.8 µm.

In the superior control half of the macula of patient 3 (P3) there was an average of 4.5 processes per 100 µm (0.7–8.3). The longest individual process was 0.7 µm.

No processes were observed to breach the elastic layer of Bruch’s membrane. The processes were often associated with focal thickening of the endothelial cell basement membranes. Each eye had large quantities of long spacing material in the outer collagenous layer of Bruch’s membrane, often in close juxtaposition to the endothelial processes and focal thickening in the basal lamina.

PHOTOCOAGULATED REGIONS OF HUMAN EYES

The laser lesion destroyed only focal areas of photoreceptors and RPE, leaving Bruch’s membrane intact and the choriocapillaris patent except in a few instances where it was absent at the site of the burn (Figs 3 and 4).

In patient 2, who had photocoagulation 15 hours before exenteration, the average number of processes per 100 µm ranged between 2.9–8.8 (Fig 5). The longest process was 0.5 µm at 200–300 µm from the laser burn but did not breach the elastic lamina of Bruch’s membrane. The number of processes was maximal at 100–400 µm from the laser burn (Fig 5). There was no statistically significant difference between the maximum number of processes seen after laser treatment and that seen in the control half of the same eye (p=0.38, two tailed test).

In patient 3, who had laser photocoagulation 5 days before exenteration, the average number of processes in each 100 µm from the laser burn ranged between 1.8–13.5 µm with the maximum number being between 400–500 µm from the burn and minimum number directly under the burn (Fig 6). The number of processes at 400–500 µm from the burn was statistically different from the average of 4.5 that was seen in the untreated half of the same eye (p=0.009, two tailed test). The longest process was 2.6 µm at 0–100 µm from the laser burn. Processes of this size were not seen in the untreated specimens, but none breached the elastic lamina of Bruch’s membrane. Neovascularisation was not seen.

As found in the untreated eyes, the processes were often associated with focal thickening of the endothelial cell basement membranes and large quantities of long spacing material in the outer collagenous layer of Bruch’s membrane, often in close juxtaposition to the endothelial processes and focal thickening in the basal lamina.

UNTREATED MONKEY EYES

Monkey 1 had no treatment and the total number of processes per 100 µm was 10.4 with a range of 9–12. The longest process was 0.8 µm.

Monkey 2 had no treatment and the total number of processes per 100 µm was 3.8 with a range of 2–5. The longest process was 0.7 µm.

Figure 3  Patient 2 treated 15 hours before exenteration. (A) Colour fundus photograph of the laser burns taken minutes after they were created. (B) Histology of the laser burn. The burn is localised to the photoreceptors (small arrow) and RPE (curved arrow), with Bruch’s membrane (thick arrow) still intact and most of the choriocapillaris surviving.

Figure 4  Patient 3 lasered 5 days before exenteration. (A) Colour fundus photograph of the laser burns taken minutes after they were created. (B) Histology of the laser burn. The burn is localised to the photoreceptors (small arrow) and RPE (curved arrow), with Bruch’s membrane (thick arrow) still intact and most of the choriocapillaris surviving.
Monkey 3 had laser photoagulation to the right eye, while the left eye had no treatment. The non-treated left eye had a total number of processes per 100 µm of 8.4. The longest process was 0.9 µm.

None of the control specimens had processes that breached the elastic lamina of Bruch’s membrane. Focal thickenings in the endothelial basement membranes were noted where the processes protruded into Bruch’s membrane, but there was no long spacing material in the outer collagenous layer as seen in the human eyes.

PHOTOCOAGULATED MONKEY EYE
Laser photocoagulation destroyed a focal area of photoreceptors and the RPE, leaving Bruch’s membrane intact and the choriocapillaris patent (Fig 7). The average number of processes ranged from 3.1–11.4 per 100 µm with a large variability in the numbers at each distance from the burn. The number of processes was greatest at a distance of 500–700 µm from the laser burn, although this was not significantly different from that seen in the control eye of this monkey (p=0.28). The longest process was 1.5 µm at 800–900 µm from the burn. Processes this size were not seen in the control eye of this monkey and 4% of processes at 500–600 and 800–900 µm from the burn breached the elastic lamina of Bruch’s membrane (Fig 8A, B). There was no evidence of neovascularisation.

Qualitative differences were also noted in the endothelial cell processes when compared with those in control sections. The cytoplasm of the endothelial cells contained more intracytoplasmic organelles than was seen in untreated specimens (Fig 8C, D). Many cellular processes were seen that did not breach the basal lamina, and therefore were not counted in our analysis. However, they appeared to be displacing the basal lamina from the plasma membrane of the endothelial cell and may represent an early phase of formation of cellular protrusion into Bruch’s membrane (Fig 8C, D).

Discussion
We have presented evidence that the cytoplasmic processes from the choroidal capillary endothelial cells increased in number and size in the human eye that was photoagulated 5 days before exenteration. In the monkey eye treated 9 days before sacrifice, we noted longer processes than seen in untreated specimens, that extended through the elastic lamina of Bruch’s membrane. The limitation of counting only those processes passing through the basal lamina of the capillary may have underestimated the influence of laser treatment on the endothelium since complex structures emanating from the endothelium were seen internal to the basal lamina, a phenomenon not seen in the untreated eyes. Changes have been noted previously in the choriocapillaris following photoagulation including prominent, expanded rough endoplasmic reticulum, and an increase in the number of lysosomes. We found similar evidence of morphological changes in the endothelial cells of the monkey,
consistent with increased activity. We did not see any cellular processes from the RPE traversing the full width of Bruch’s membrane as has occurred in rats following laser burns using high energy intensity.31

The potential relevance of these findings to the clearance of drusen following photocoagulation, and to neovascularisation, is unclear. It would be attractive if it were possible to demonstrate a long lived cellular response to photocoagulation to account for the time scale of resolution of drusen. We were unable to detect convincing evidence for phagocytosis of debris in Bruch’s membrane by these processes as has been found in other tissues.20 21 However, we cannot rule out the possibility that this exists as a normal function of endothelial cells of the choriocapillaris, and that this function is stimulated after photocoagulation. It is also possible that the endothelial cells may influence clearance of debris by modifying the biophysical properties of Bruch’s membrane. This would have been more convincing in humans if the processes had transgressed the elastic lamina as was seen in the monkey eye, since it is at this level that the principal barrier to fluid flow exists.32 The interval between photocoagulation and histological examination is short and had it been longer more persuasive evidence may have been found.

Endothelial processes such as those examined in this study have been interpreted as nascent neovascularisation since the initial step in new vessel formation is the egress of endothelial cell processes through the basement membrane sheath accompanied by production of proteases.22 33 34 The observed increase of endothelial activity following laser treatment would be in accord with the finding that choroidal neovascularisation occurs consistently after heavy laser photocoagulation at the primate macula,28 and follows laser treatment in humans.29 However, we saw no new vessel formation, implying that the difference may be one of degree determined by the intensity of photocoagulation. Consequently, choroidal neovascularisation may be a distortion of what is a normal feature of the choroid. A fine line must exist between the upregulation of process formation after photocoagulation and nascent angiogenesis. It is possible only to speculate on the circumstances that allow inward growth of formed blood vessels from the choroid, and several factors have been considered as being potentially relevant. Imbalance of inhibitory and stimulating diffusible factors, changes in the physicochemical properties of Bruch’s membrane, and the presence of macrophages may modulate this process.22 35 It is also possible that the basal lamina of endothelial cells may influence the effects of the release of growth factors. It is of interest therefore that quite marked focal thickenings of the endothelial cell basal lamina was noted to surround the base of these processes, especially in the young.18 We have postulated that the role of this focal thickening of basal lamina may be to act to distribute mechanical forces around the cytoplasmic peg that anchors the cell to the extracellular matrix. If focal thickening also
moderates the risk of neovascularisation, its absence in the elderly may serve to increase the risk of choroidal new vessel growth with age. It is hardly surprising that photocoagulation may alter the influences that regulate the generation of cellular processes intruding into Bruch’s membrane which is normal, and choroidal neovascularisation as part of disease. The results of the clinical trials of prolyphactic photocoagulation to patients with high risk drusen must be awaited before we can abandon concern over the potential induction of neovascularisation. Our observations suggest that the use of low rather than high energy settings may reduce the risk, but may not obviate it totally.

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