Conditioned medium from mixed retinal pigmented epithelium and Müller cell cultures reduces in vitro permeability of retinal vascular endothelial cells

M Tretiach, M C Madigan, M C Gillies

Aim: To investigate the in vitro effect of laser photoagulation on blood-retinal barrier permeability. Methods: Retinal capillary endothelial cells were exposed to supernatants from long term co-cultured cells that were argon laser treated. Endothelial cell permeability was analysed by (1) measurement of transendothelial electrical resistance and (2) equilibration of [3H] inulin and [14C] albumin across the cell monolayer. Results: Laser photoagulation of various retinal cells and control ECV304 cells in the lower chamber did not appreciably improve permeability of the endothelial cell monolayer compared with that of unlayered cells. However, medium that was conditioned by mixed retinal pigmented epithelium and Müller cells significantly reduced both inulin (43.2% (SD 6.5%) equilibration in mixed cultures v 59.8% (SD 7.0%) control cells, p<0.05) and albumin (15.1% (SD 3.8%) v 31.1% (SD 6.7%), p<0.05) permeability of the endothelial cell monolayers. A fourfold increase in transendothelial electrical resistance was also seen. Conclusions: These results are consistent with the hypothesis that interaction of Müller cells with retinal pigmented epithelium induced by laser treatment results in secretion of soluble factor(s), which reduces permeability of retinal vascular endothelium. Identification of these factor(s) may have implications for the clinical treatment of macular oedema secondary to diabetic retinopathy and other diseases.

The blood-retinal barrier (BRB) exists at the level of the retinal capillary endothelium ("inner" BRB) and the retinal pigmented epithelium (RPE) ("outer" BRB). Macular oedema secondary to breakdown of the inner BRB is the most common cause of vision impairment in diabetic retinopathy. Retinal laser therapy reduces the risk of blindness in eyes with diabetic macular oedema. However, laser photoagulation is generally administered late in the course of the disease when vision loss is imminent, may not always work, and is inherently destructive. Understanding how retinal laser treatment affects a leaking BRB is important for developing better treatments for macular oedema.

Changes in retinal morphology after laser have been well described in rats, rabbits, monkeys, and humans. Although some laser energy may directly affect the retinal vessels, it is generally accepted that the major site of absorption is the RPE and choroid. The laser affected areas of the photoreceptor outer segments and RPE exhibit signs of necrosis including cell disruption, vacuolisation, and condensation of cytoplasmic proteins within a few hours after treatment to an extent that is commensurate with the intensity of the burn. Within days, RPE cells migrate across Bruch’s membrane to fill the lesion with subsequent scar formation. Müller cells and astrocytes replace the damaged outer nuclear layer of the retina, interdigitating with the migrated RPE cells. Müller cells undergo widespread and long lasting changes after photoagulation including increased expression of glial fibrillary acidic protein (GFAP) associated with hypertrophy, migration, and scar tissue formation.

We propose that laser photoagulation stimulates cells to produce soluble factor(s) that can restore a leaky BRB. In this study, we tested the hypothesis that supernatants from RPE, Müller cells, pericytes, and control ECV304 cells can reduce the in vitro permeability of a retinal capillary endothelial cell (RCEC) monolayer.

Materials and Methods

Cell isolation and culture

A mixed cell population of postmortem activated RPE and migrating Müller cells was isolated from 24-48 hour postmortem bovine eyes using a modification of Edwards’ method and based upon previous observations on the characteristics of postmortem Müller cells. A coronal section of the globe was made and the cornea, lens, and vitreous tissues were removed as one piece. The retina was carefully dislodged, and the remaining traces of retinal tissue were removed from the optic nerve with a sterile scalpel blade. Two eyecups were filled with 0.25% trypsin ethylenediamine tetra-acetic acid (TE) (ThermoTrace Biosciences, Melbourne, Australia) and incubated at 37°C for 15 minutes. The enzyme mixture was pipetted to gently loosen adherent cells, and the eyecups containing TE were incubated for a further 15 minutes. Dissociated cells were removed and transferred into Dulbecco’s modified Eagle’s medium (DMEM) (ThermoTrace Biosciences) containing 20% heat inactivated fetal bovine serum (FBS) (ThermoTrace Biosciences), 2 mM glutamine, 100 U/ml penicillin, and 100 μg/ml streptomycin (ThermoTrace Biosciences). Cells were centrifuged at 163 g for 6 minutes. The cells were resuspended in fresh medium and seeded into a 25 cm² flask (Nunc A/S, Roskilde, Denmark). Eyecups were filled with fresh TE and incubated for a further 30 minutes. The product of the second eyecup digestion was centrifuged and incubated as above. The first digest of the eyecup produced a mixed population of cells, whereas the second digest...
provided a high yield of RPE cells. Cells were incubated undisturbed for 1–2 weeks at 37°C (95% relative humidity, 5% CO2). At the first or second passage, cells were seeded (15 000 cells per well) into the lower chamber of 24 well Costar Transwell plates (Corning Inc, Acton, MA, USA) with DMEM containing 10% FBS. Medium was replaced 24 hours after seeding, and then twice weekly for 3 months as previously described for long term RPE cultures.19

Other cell types were cultured in 24 well plates as described above. Pericytes were isolated by enzyme digestion of bovine retinas;19 Müller cells were isolated from bovine retinas following the method of Tretiach et al,20 and bovine RCEC were isolated using an enzyme digestion technique followed by filtration to collect the microvesSEL fragments.21 A human bladder carcinoma derived epithelial cell line (EVC304, European Collection of Cell Cultures, Salisbury, UK) was included as an epithelial cell control.22 23 Cultures were photographed with Kodak Ektachrome T160 (Kodak, Rochester, NY, USA) film using a Zeiss Telaval 31 inverted microscope (Carl Zeiss, North Ryde, NSW, Australia).

**Immunohistochemistry**

Cells from primary cultures were routinely immunostained with a panel of antibodies (table 1). Antibodies, except antiretinaldehyde binding protein (CRALBP), were visualised using either Alexa 488 or 568 conjugated secondary antibodies. For CRALBP localisation, early passage Müller cells and RPE were grown to confluence on glass coverslips, incubated overnight at 4˚C with the primary antibody or 2% NDS on the negative controls. Coverslips were rinsed in PBS, incubated in biotinylated antirabbit Ig antibody (1:50) (Amersham Pharmacia, Baulkham Hills, NSW, Australia) for 1 hour, then ExtraAvidin peroxidase (1:200) (Sigma-Aldrich P/L, Castle Hill, NSW, Australia) for 45 minutes. Bound antibody was detected with Vector red chromagen (Vector Laboratories, Burlingame, CA, USA). Coverslips were dehydrated through a series of alcohols and xylene and mounted with DePeX. Images were captured with Leica DC Viewer Computer Software (Version 3) (Leica Microsystems Ltd, Heerbrugg, Switzerland) using a Leitz Diaplan light microscope (Leitz Messtechnik GmbH, Wetzlar, Germany).

### Table 1 Antibodies used for immunohistochemistry

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Dilution</th>
<th>Cell type</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polyclonal antihuman</td>
<td>1:50</td>
<td>RPE</td>
<td>Zymed Labs Inc, San Francisco, CA, USA</td>
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<tr>
<td>ZO-1</td>
<td></td>
<td></td>
<td>Prof J Saari, University of Washington, USA</td>
</tr>
<tr>
<td>Polyclonal anticow</td>
<td>1:100</td>
<td>Müller</td>
<td>Dako, Sydney, NSW, Australia</td>
</tr>
<tr>
<td>CRALBP</td>
<td></td>
<td>RPE</td>
<td>Dako, Australia</td>
</tr>
<tr>
<td>Monoclonal antiratine</td>
<td>1:100</td>
<td>Müller</td>
<td>Dako, Australia</td>
</tr>
<tr>
<td>vitamin (V9)</td>
<td></td>
<td>RPE</td>
<td>Dako, Australia</td>
</tr>
<tr>
<td>Polyclonal antihuman</td>
<td>1:50</td>
<td>RCEC</td>
<td>Dako, Australia</td>
</tr>
<tr>
<td>vWF</td>
<td></td>
<td></td>
<td>Dako, Australia</td>
</tr>
<tr>
<td>Monoclonal antiratine</td>
<td>1:50</td>
<td>Pericytes</td>
<td>Dako, Australia</td>
</tr>
<tr>
<td>αSMA (1A4)</td>
<td></td>
<td></td>
<td>Dako, Australia</td>
</tr>
<tr>
<td>Polyvalinal anticow</td>
<td>1:500</td>
<td>Astrocytes</td>
<td>Dako, Australia</td>
</tr>
<tr>
<td>GFAP</td>
<td></td>
<td>RCEC</td>
<td>Dako, Australia</td>
</tr>
<tr>
<td>Monoclonal antihuman</td>
<td>1:10</td>
<td>Neural cells</td>
<td>International, Soco, Maine, USA</td>
</tr>
<tr>
<td>NCAM</td>
<td></td>
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</table>

CRALBP, cellular retinaldehyde binding protein; NCAM, neural cell adhesion molecule; GFAP, glial fibrillary acidic protein; αSMA, alpha smooth muscle actin; vWF, von Willebrand’s Factor; ZO-1, zonula occludens.

**Conditioned medium and argon laser studies**

Second passage RCEC (5000 cells per well) were seeded onto coated 0.4 μm pore size polycarbonate filter (0.33 cm²) inserts of two chamber Transwell plates.21 Inserts were placed into wells containing the long term cultured cells (described above) to study the effects of conditioned medium on endothelial cell (EC) permeability. The conditioned medium and control cell groups (7–8 wells per group) are described in table 2. Long term cultured cells were adapted in EC medium22 for 24 hours before co-culturing with EC. Seven days later, cells in the lower chamber were lasered as follows. Filter inserts were removed from the 24 well Transwell plate and returned to the incubator in humidified dishes. Medium was decanted from cells in the lower wells and the plates were held sideways in situ on the chin rest of a Coherent argon blue-green laser. Four wells per group received 70 shots per well with 200 μm spot size, pulse duration 0.1 s, 150 mW. The lowest dose that caused a visible reaction in the RPE monolayer on phase contrast microscopy had been established previously in dose response studies (not shown). After laser treatment, fresh medium was added to cells in the lower chamber. The Transwell filter inserts containing EC were replaced into medium that was now conditioned by the “lasered” and “unlasered” wells.

**Permeability studies**

Transendothelial electrical resistance (TEER) was measured using a Millipore ERS resistance meter (Millipore, NSW, Australia) as previously described.21 TEER was recorded from day 3 after the filter inserts containing EC had been added to the cell groups providing conditioned medium, and then every second day until day 7, when these groups were lasered. TEER was measured every 12 hours thereafter. We prospectively determined that comparisons between groups should be carried out when the control EC reached peak resistance. Presumably this situation best reflects the in vivo BRB. The mean resistance of the different groups was calculated by subtracting the average reading from the “no cell” wells (group 7) multiplied by the area of the Transwell filter (0.33 cm²). Results were expressed in ohms cm². The experiment was repeated three times. Permeability of radiolabelled macromolecular tracers across the EC barrier was determined as follows. Within one hour of laser treatment, a mixture of radiolabelled tracers—[methoxy-3H] inulin (NEN Life Science Products Inc, Boston, MA, USA) and [methyl-14C] methylated (bovine serum) albumin (NEN Life Science Products Inc)—was added to medium in the upper Transwell chamber. The concentration of tracer was predetermined to provide a sufficient number of counts (5000–50 000 dpm) in the final volume added to the luminal chamber (total count). Medium was removed from the lower chamber at 24, 36, and 48 hours. A Tricarb 2100TR Packard liquid scintillation β counter (Packard Instrument Co, Meriden, CT, USA) using the full spectrum DPM technique

### Table 2 Conditioned medium and control cell groups used in the barrier assay

<table>
<thead>
<tr>
<th>Group number</th>
<th>Description</th>
<th>Total number of wells</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Mixed RPE &amp; Müller cells</td>
<td>8</td>
</tr>
<tr>
<td>2</td>
<td>Mixed pericyte &amp; Müller cells</td>
<td>8</td>
</tr>
<tr>
<td>3</td>
<td>RPE cells alone</td>
<td>8</td>
</tr>
<tr>
<td>4</td>
<td>Müller cells alone</td>
<td>8</td>
</tr>
<tr>
<td>5</td>
<td>EVC304 cells</td>
<td>8</td>
</tr>
<tr>
<td>6</td>
<td>RCEC alone</td>
<td>7</td>
</tr>
<tr>
<td>7</td>
<td>Coated filters only</td>
<td>2</td>
</tr>
</tbody>
</table>

*Control groups.
(Operation Manual, Packard Instrument Co) was used to calculate radioactivity of the individual radionuclides. A ratio of the radioactivity from each of the lower wells to the total count was determined for each time point and expressed as the percentage equilibration.

Statistics
As we anticipated that the in vitro EC barrier enhancing effect occurred after a delayed response to photocoagulation, we compared the conditioned medium groups with the control EC at the later timepoints: between 24–48 hours after lasering. Results were calculated as follows: (1) mean (standard deviation) electrical resistance (ohms cm²) for the TEER, and (2) equilibration of tracers was expressed as mean (SD) per cent (%) for the permeability studies. Repeated measures ANOVA followed by linear contrast was used to analyse the data. Time was treated as the within subject factor and group as the between subject factor. The level of significance was set at p = 0.05.

RESULTS
Cell morphology and immunohistochemistry
All primary cells grew to confluence within one week after seeding into culture flasks. Cells cultured from the first digest of the eyecups contained a mixed population of two distinct cell types (fig 1A). Retinal pigmented epithelium and Müller cells were identified in these cultures, where RPE displayed intense pigmentation and characteristic cobblestone morphology and Müller cells exhibited long, delicate radial fibre structures and distinctive varicosities around the cell body. Cells cultured from the second digest consisted of >90% RPE (fig 1B) as determined with specific antibody labelling (fig 1C). Müller cells isolated from retinal tissue were identified by anti-CRALBP (fig 1D) and antivimentin (fig 1C). Müller cell culture showing growth with contact inhibition (G) (×250 magnification).

Long term cultures
With phase microscopy, long term mixed RPE and Müller cell cultures (P1-2) grown in 24 well plates appeared as flat, uniform sheets with foci of darkly pigmented areas (fig 1E). RPE cells alone appeared to grow irregularly with areas of multilayering (fig 1F). Mixed pericytes and Müller cells grew without extensive contact in the long term cultures albeit without detectable pigmentation (fig 1G). Many pericytes became detached from the lower well surface by completion of the experiment, when the upper Transwells were removed. The predominant population remaining on the lower well surface were Müller cells. ECV304 cell derived Müller cells remained as a stable monolayer with characteristic cobblestone morphology (fig 1H) for up to 4 months in culture. Lasering of the long term cultured cells did not appear to change cell morphology dramatically.

Permeability results
Transendothelial electrical resistance of control EC reached 9.0 Ω cm² on day 5 and remained fairly constant thereafter, peaking at 11.7 Ω cm² on day 8 (thick line in fig 2A). By day 9, TEER of EC grown in medium conditioned by mixed RPE and Müller cell cultures (fig 2A) was fourfold that of the control EC and the other conditioned medium groups (fig 2B). Conditioned medium from mixed RPE and Müller cells that were laser treated did not significantly affect TEER of the EC barrier (not shown).

As there was no difference in results between the lasered and unlased cells in each group, we combined the outputs from both groups to improve the statistical power. Groups in which there was an obvious effect on EC permeability were analysed using repeated measures ANOVA as described above. The difference between EC exposed to supernatants from the mixed RPE and Müller cell group (43.2% (SD 6.5%) equilibration) and control EC (59.8% (SD 7.0%) equilibration) was significant for inulin (p<0.05) (fig 2C) and albumin leakage (15.1% (SD 3.8%) vs 31.1% (SD 6.7%), p<0.05) (fig 2D). Conditioned medium from other groups had no discernible effect on the overlying EC.

DISCUSSION
In this study, we set out to examine whether supernatants from a variety of lasered cells could reduce the permeability of an RCEC monolayer using a two chamber in vitro assay. Although laser treatment of cells in the lower chamber did not have any effect on EC permeability, we observed that conditioned medium from mixed RPE and Müller cells significantly reduced monolayer permeability, consistent with the hypothesis that barrier enhancing factors are released from cells within the laser scar.
Various theories about the mechanism(s) of retinal laser therapy have been proposed. The therapeutic benefit of retinal laser appears to be an indirect effect related to a secondary tissue response, rather than to the immediate burn. Incompetent retinal vessels regain potency when the outer blood-retinal barrier is repaired after laser therapy. These observations suggest that the laser scar that forms after photocoagulation may be a source of factors that restore leaking retinal vessels. RPE and Müller cells react to tissue destruction by assembling at the site in the immediate and early phases, suggesting that they play an important role in repair of the outer blood-retinal barrier. Although the response of RPE cells to injury appears to be rapid, that of the Müller cells is temporarily delayed. Patterns of growth factor expression following photocoagulation in normal pig retinas have been studied to understand how RPE and Müller cells might play a contributory role in retinal wound healing. RPE may orchestrate the initial response(s) via TGF-β which is a chemoattractant for inflammatory cells and promotes matrix deposition—as well as PDGF, EGF, TGF-α, and FGF—to promote the proliferation of RPE and other cells.

Thereafter the reparative effects appear to be mediated by a combination of autocrine and paracrine signals from the major cell types, including RPE and Müller cells. We used long term cell cultures to control for artefacts that might be mistaken as a “laser induced” response, as we have previously found that short term cultured cells appear to have an activated phenotype (unpublished). Other in vitro studies have used short term RPE cultures to investigate the effect of conditioned medium on EC proliferation. We believe that our study is the first to investigate the effect of diffusible substances from RPE cells on EC permeability. The short versus long term (present study) culture conditions and variations in cell confluency may elicit different factors from RPE. Endothelial cell responses to RPE grown under different culture conditions are likely to be variable.

Inulin is a low molecular weight molecule (MW 5000–5500). Its leakage across EC monolayers reflects TEER and is an accepted measure of paracellular permeability. In the present study, the larger albumin molecule (MW 69 000) was more successfully retarded by the EC barrier (see fig 2C and D). We have observed that in vitro EC may grow unreliably,
forming multilayers that do not always achieve complete barrier formation. Nevertheless, the significant degree to which conditioned medium from the mixed RPE and Müller cells contributed to decreased permeability of the above mentioned macromolecules, provides further evidence of the in vitro plasticity of the RCEC monolayer.

It is well established that the scar formed after laser therapy is comprised predominately of RPE and Müller cells. In this study we found that only the supernatants from the laser induced scar may contribute to tightening leaky retinal blood vessels. Further work is necessary to identify factor(s) that may contribute to the barrier tightening effect.

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