Inhibition of microvascular endothelial cell proliferation by vitreous following retinal scatter photocoagulation

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Abstract

Laser photocoagulation of pig retina induced breakdown of the blood-retinal barrier, with the appearance of serum proteins in the vitreous as determined by sodium dodecyl sulphate/polyacrylamide gel electrophoresis (SDS-PAGE) and immunoblotting techniques. Vitreous from laserer eyes inhibited the proliferation of cultured retinal microvascular endothelial cells in comparison with vitreous from non-lasered control eyes, and the inhibitory effect in the lasered eyes persisted for at least seven days. Inhibition was specific for endothelial cells, since no effect was observed when retinal pericytes or Tenon's fibroblasts were the target cells. These results suggest that indirect scatter photocoagulation may induce regression of neovascularisation by causing breakdown of the blood-retinal barrier and thus releasing into the vitreous serum components which result in inhibition of retinal microvascular endothelial cell growth.

Preretinal neovascularisation is responsible for many of the blinding sequelae of diabetic retinopathy. Neovascularisation occurs in response to ischaemic damage to inner retinal tissue and is thought to be mediated via diffusible angiogenic factor(s). Panretinal photocoagulation induces regression of the new vessels and has become an established primary therapy for preretinal neovascularisation.

The exact mechanism of regression of neovascularisation is unknown, but a number of hypotheses have been advanced to explain this effect. They include a proposed reduction in angiogenic factor release from ischaemic retina owing to destruction of retinal tissue, to increased availability of oxygen through thinned outer retina, or to disruption of the outer blood-retinal barrier facilitating outward diffusion of the angiogenic factor into the choroid. Recently, on the basis of tissue culture experiments, it has been suggested that the retinal pigment epithelium (RPE) may produce a factor capable of inhibiting the proliferation of vascular endothelial cells. However, RPE cells can also produce retinal vascular mitogens.

In order to study the mechanisms underlying photocoagulation induced regression of neovascularisation we undertook scatter laser photocoagulation of pig retina and examined the properties of vitreous from lasered and un lasered eyes. Our initial study showed that vitreous from eyes which had received laser photocoagulation significantly inhibited the proliferation of retinal microvascular endothelial cells in vitro compared with vitreous from unlased eyes. These results suggested that an inhibitor of angiogenesis is released from the retina after photocoagulation and diffuses into the vitreous, where it acts directly on preretinal new vessels. In the present study, we have extended this work to an analysis of vitreous proteins, and our results indicate that the major change in vitreous content following laser therapy is an influx of serum proteins and other components, the net effect of which is growth inhibitory activity specifically directed towards endothelial cells.

Materials and methods

PHOTOCOAGULATION

Eight 1-year-old miniature pigs received scatter photocoagulation by an argon blue-green laser (HGM Model 5, Litechnica) according to the following protocol. After premedication with 10 mg/kg intramuscular ketamine 10 mg/kg methohexitone was administered intravenously to induce anaesthesia prior to intubation. Anaesthesia was maintained with spontaneous inhalation of a 30% O₂/70% N₂O mixture supplemented by 1-5% halothane. The animal's head was positioned on a slit-lamp and one thousand 500 μm laser burns at 300 mW and 100 ms duration were applied to the retina of the right eye through a Goldmann fundus contact lens. The left eye was not lasered and served as the control for each animal. Following photocoagulation the anaesthetic agents were discontinued, and the animals recovered.

PREPARATION OF VITREOUS

Five of the animals were killed four days after photocoagulation and three after seven days. In each instance the animals were anaesthetised as described earlier and both eyes were enucleated. A blood sample was also taken from some of the animals, and the serum was separated and stored. The animals were killed by intravenous potassium chloride. The globes were immediately dissected and the vitreous was removed and stored at −70°C. Before use the vitreous was thawed, liquefied by an ultrasonicator (MSE Soniprep 150), and finally filter sterilised through a 0·22 μm filter (Millipore, USA).

CELL CULTURE

Human eyes of variable age and with no known ophthalmic disease were obtained from Moorfields Hospital Eye Bank, London, after corneal

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buttons had been removed. Bovine eyes were collected from a local abattoir. Human RPE cells were isolated and cultured as previously described. Bovine retinal microvascular endothelial cells were grown in a 1:1 mixture of Dulbecco’s modified Eagles medium (DMEM) with 7.5% human platelet-poor plasma and pericyte conditioned medium. Bovine retinal pericytes and bovine Tenon’s fibroblasts were routinely grown in DMEM supplemented with 20% fetal calf serum.

Cultures between the second and fifth passage were used for all subsequent studies.

**Production of conditioned media**

Conditioned media were prepared as previously described. In brief, cultures of retinal pericytes and RPE cells were grown to confluence in 25 cm² flasks (Falcon) and washed twice with Dulbecco’s phosphate buffered saline without calcium and magnesium (PBSA). After they had been washed 8 ml of DMEM supplemented with 7.5% human platelet-poor plasma was added to each flask. After incubation for two days the conditioned medium was removed and stored at −20°C.

**Proliferation assays**

2×10⁴ retinal microvascular endothelial cells were seeded into each gelatinised well of 24-well plates (Flow) in DMEM supplemented with 7.5% human platelet-poor plasma. After a period of two days to allow for attachment and initial proliferation the medium was removed, and the cells were washed twice with plasma-free DMEM. Each well then received 1 ml of one of the following test media:

1. A 1:1 mixture of DMEM+7.5% plasma and pericyte conditioned medium to which was added 75 µl liquefied vitreous (either ‘lasered’ or ‘non-lasered’) or 75 µl PBSA as a control.
2. A 1:1 mixture of DMEM+7.5% plasma and RPE conditioned medium to which was added 75 µl liquefied vitreous (either ‘lasered’ or ‘non-lasered’) or 75 µl PBSA as a control.

After two days in test media the cell numbers were determined by means of a haemocytometer. Each experiment was performed on at least two separate occasions.

Representative vitreous samples were also examined for their effect on pericyte and fibroblast proliferation. 2×10⁴ cells were seeded in each well of uncoated 24-well plates as described above and incubated overnight to allow for attachment and proliferation. The test media (DMEM+7.5% plasma mixed 1:1 with pericyte conditioned media) plus 75 ml/l of either vitreous or saline were added and cell numbers were determined at two days in the case of fibroblasts and at five days for pericytes, since the latter grew comparatively slowly.

Aliquots of cells from each test were stained with trypan blue in order to determine a cytotoxic effect. Each experiment was performed on at least two separate occasions.

**Migration assays**

Migration assays were undertaken as described by Hackett and Campochiaro using 48-well modified Boyden chambers. Briefly the lower wells were filled with test media consisting of 75 ml/l of vitreous added to either DMEM with no serum or DMEM with 7.5% fetal calf serum. Each test medium type was added to eight wells. Control wells had the vitreous substituted with an equivalent volume of PBSA. The wells were then overlaid with a type 1 collagen (Ethicon) coated polycarbonate membrane with 10 µm pores and the chambers assembled. Upper wells were filled with a suspension of one of the cell types to be tested (endothelial cells, pericytes, RPE, or fibroblasts) in serum free medium at a density of 9×10⁴ cells/ml. The chambers were then incubated for six hours at 37°C. The filter was removed, fixed in methyl alcohol, and stained with modified Wright’s stain. The number of cells which had migrated through the polycarbonate filter were counted after the non-migrating cells had been wiped from the top surface. Cell nuclei in 10 high power fields were counted in each well.

**Gel electrophoresis**

Samples of vitreous (1 µl) were analysed by SDS-PAGE in an 8–25% gradient gel by the Pharmacia Phast System according to the manufacturer’s instructions. Samples were run under reducing and non-reducing conditions (with or without mercaptoethanol in the sample buffer).
In addition, native polyacrylamide gel electrophoresis of vitreous samples was performed. Gels were double stained with silver nitrate and Coomassie blue dye. Alternatively, proteins separated on the gels were transferred by diffusion blotting in the Pharmacia Phast System onto nitrocellulose membranes and detected with antisera against whole pig serum by the immunoperoxidase procedure according to the manufacturer’s instructions. Samples of pig serum and haemolysed pig serum were run on the same gels for comparison.

**STATISTICS**
All data from the proliferation and migration assays were expressed as the mean together with standard deviation of the mean. The statistical significance of vitreous related changes was examined by analysis of variance.

**Results**

**PROLIFERATION ASSAYS**
Vitreous from all laser treated eyes significantly inhibited the proliferation of retinal microvascular endothelial cells in the presence of both pericyte and RPE conditioned media when compared with controls containing PBSA (p<0.001) (Figs 1 and 2). In contrast, vitreous from unlasered eyes did not significantly affect endothelial cell proliferation (p>0.8). There was no apparent difference in the degree of inhibition between laser exposed samples from four- and seven-day post-treatment animals in either the presence of pericyte conditioned medium or RPE conditioned medium.

The inhibition of proliferation by vitreous from laser treated eyes appeared to be specific for retinal microvascular endothelial cells, since inhibition was not seen with either retinal pericytes or fibroblasts (Table I).

Trypan blue exclusion techniques did not demonstrate any cytotoxic effect by any of the vitreous samples on the cells during the test period.

**MIGRATION ASSAYS**
The vitreous from either lasered or non-lasered eyes showed no apparent effect on the migration of either endothelial cells, pericytes, RPE, or fibroblasts in comparison with appropriate controls (Table II).

**GEL ELECTROPHORESIS**
In analysis by gel electrophoresis samples of vitreous from untreated eyes showed a small number of protein bands, as predicted for normal vitreous proteins (Fig 3). Samples from laser treated eyes showed a large increase in protein concentration, whose electrophoretic pattern was closely similar to that of pig serum (Fig 3). Immunoblotting of the proteins with...
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Inhibitory effect of vitreous on microvascular endothelial cell proliferation was tested in vitro. The inhibitory effect was present at both four and seven days after laser treatment. These time periods were chosen because clinical regression of neovascularisation is usually evident during this period following panretinal photocoagulation. Owing to the limited quantity of vitreous available it was only possible to (a) use one concentration of vitreous (75 μl vitreous/1 of medium was chosen arbitrarily), and (b) test the effect of ‘lasered’ vitreous on the response of microvascular endothelial cells to two mitogenic stimuli. We chose pericyte conditioned medium because pericytes are (a) closely associated with retinal capillaries, (b) implicated in new vessel formation, and (c) capable of producing culture medium which is mitogenic for retinal microvascular endothelial cells. RPE conditioned medium was used because destruction of RPE cells after scatter photocoagulation results in regression of extra-retinal neovascularisation, and RPE cells produce retinal vascular mitogens in vitro. Since no significant differences were observed in the degree of inhibition in the presence of either conditioned media, it is likely that the ‘lasered’ vitreous acted via the same mechanism in both media.

The inhibitory effect of vitreous from laser treated eyes was specific for retinal capillary endothelial cells, since we observed no inhibitory effect on the proliferation of pericytes or fibroblasts in the presence of pericyte conditioned media. Since both pericytes and fibroblasts have previously been shown to respond to mitogenic factors in pericyte conditioned medium (Wong, unpublished) the results suggest that the mitogens which are responsible for endothelial cell proliferation are different from those for the other cell types studied.

Since the migration of retinal microvascular cells is as important as proliferation in new vessel formation, we examined the effect of lasered vitreous on the migration of retinal endothelial cells. Owing to the limited quantity of material available it was possible to examine the effect of vitreous only on serum induced migration. Thus, although vitreous had no effect on the migration of endothelial cells in the presence of serum, there may be an effect in the presence of conditioned media.

The nature of the laser induced inhibition of endothelial cell proliferation is obscure. Previously it has been suggested that RPE cells release an inhibitor of neovascularisation after laser injury, but RPE cells also release mitogens for endothelial cells, and clearly the role of RPE cells in this system is complex. Both xenon and laser photocoagulation cause damage to the outer blood-retinal barrier, and protein concentrations in the vitreous are known to be raised after laser treatment. While some of these proteins were acknowledged to have been derived from serum, it was also suggested by Campochoiaro and colleagues that a contribution was derived from damaged retinal cells, in particular the RPE. In the present study we have shown that the great majority of the proteins in the vitreous after laser therapy are derived from serum and that serum proteins persist in the vitreous for at least seven days following laser treatment.

Discussion

We have confirmed our previous observation that following panretinal photocoagulation a diffusible factor(s) which inhibits microvascular cell proliferation appears to accumulate in the vitreous. The inhibitory effect was present at both four and seven days after laser treatment. These time periods were chosen because clinical regression of neovascularisation is usually evident during this period following panretinal photocoagulation. Owing to the limited quantity of vitreous available it was only possible to (a) use one concentration of vitreous (75 μl vitreous/1 of medium was chosen arbitrarily), and (b) test the effect of ‘lasered’ vitreous on the response of microvascular endothelial cells to two mitogenic stimuli. We chose pericyte conditioned medium because pericytes are (a) closely associated with retinal capillaries, (b) implicated in new vessel formation, and (c) capable of producing culture medium which is mitogenic for retinal microvascular endothelial cells. RPE conditioned medium was used because destruction of RPE cells after scatter photocoagulation results in regression of extra-retinal neovascularisation, and RPE cells produce retinal vascular mitogens in vitro. Since no significant differences were observed in the degree of inhibition in the presence of either conditioned media, it is likely that the ‘lasered’ vitreous acted via the same mechanism in both media.

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Figure 4: A: Native PAGE of vitreous samples from 7 day laser treated eyes. Track (1) vitreous from laser treated animal; (2) control vitreous from untreated animal; (3) pig serum; (4) haemolysed pig serum. B: Corresponding nitrocellulose immunoblot of samples, stained with rabbit anti-whole pig serum, with goat anti-rabbit as the second antibody.

Figure 5: SDS-PAGE of vitreous samples from 4 day and 7 day laser treated eyes. Tracks 1, 3, 5, 7: control vitreous from untreated right eyes. Tracks 2, 4: vitreous samples from pig 1 and pig 2, 4 day laser treated eyes. Tracks 6, 8: vitreous samples from pig 3 and pig 4, 7 day laser treated eyes. Tracks 9, 10: molecular weight markers.
Serum is known to contain inhibitors of endothelial cell proliferation such as transforming growth factor β (derived from platelets) and other inhibitors such as plasminogen activator inhibitor-1, though at very low levels. However, for such a breakdown of the RPE blood-retinal barrier to cause regression of neovascularisation the effect must be in some way 'selective', since leakage of serum proteins into the vitreous is a characteristic of new vessels.

The consequence of this breakdown for retinal endothelial cells probably involves an interaction of extravasated serum components with local retinal and/or vitreous components, resulting in either loss of stimulatory activity or production of inhibitory activity. This effect may be quantitative or cumulative over a long time. Our attention should be directed towards identification and characterisation of these serum factors.

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