Study of the blood-aqueous barrier in choroidal melanoma

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Abstract

Aims—Aqueous flare was used to determine the frequency and amount of blood-aqueous barrier breakdown and correlate it with tumour variables.

Methods—Aqueous flare was analysed prospectively by laser flare photometry in 139 consecutive patients seen in the oncology unit for choroidal melanoma. Both eyes of patients were examined with a laser flare cell meter in a standard fashion.

Results—Mean flare difference between healthy and tumour eyes was 3.01 (SD 2.5) photons per millisecond (ph/ms) in 32 cases of small melanomas (p<0.0001), 10.74 (13.9) ph/ms in 92 cases of medium and large melanomas (p<0.0001), and 19.23 (11.8) ph/ms in 15 cases of very large melanomas (p<0.0001). This mean differential flare was significantly higher in medium and large than in small melanomas (p<0.002) and in very large melanomas than in medium and large melanomas (p<0.028). A difference of ≥7 ph/ms between affected and healthy eyes was noted in 70 of 139 melanomas (50.4%). It was found in 3/32 small melanomas (9.4%), in 53/92 medium and large melanomas (57.6%), and in 14/15 very large melanomas (93.3%).

Conclusion—Multiple linear regression analysis showed that flare was most strongly correlated with tumour volume (r=0.43; p<0.0001) and tumour height (r=0.41; p<0.0008).

Laser flare photometry (Kowa laser flare cell meter FC-1000, Kowa Electronics and Optics, Tokyo, Japan) represents an enormous gain of sensitivity over classic slit-lamp evaluation of aqueous flare enabling an objective, quantitative, and non-invasive evaluation that is sufficiently accurate to measure precisely subclinical flare. 5 We used laser flare photometry to evaluate anterior chamber (aqueous) flare in patients with choroidal melanomas in order to determine the frequency and amount of blood-aqueous barrier breakdown and correlate it with different tumour variables.

Patients and methods

A total of 155 consecutive patients referred to the oncology unit during a period of 10 months with the diagnosis of choroidal melanoma were examined by laser flare photometry. The diagnosis of choroidal melanoma was made on the basis of standard clinical examination and paraclinical investigations in all patients. Depending on their size choroidal melanomas were subdivided into three groups, the demographics of which are detailed in Table 1:

(1) small melanomas (SM) with a diameter of less than 10 mm and a height of less than 3 mm;
(2) medium and large melanomas (MLM) with a diameter of 10–20 mm and a height of 3–10 mm;
(3) very large melanomas (VLM) with a diameter >20 mm and/or a height >10 mm.

Fourteen eyes with choroidal melanoma complicated by a neovascular glaucoma, an invasion of the optic nerve, a total retinal detachment, or a diffuse transcleral exteriorisation were excluded from the study. Two patients with a previous contralateral emulcation who had a melanoma in the remaining eye were also excluded. A total of 139 patients were included in the study. Laser flare photometry was performed in both eyes 30 minutes after instillation of dilating drops by the same technician, who did not know which eye the tumour was in. Measurement is based on the backscattering of light produced in the anterior chamber by a helium-neon laser beam of constant power. Backscattered photons are detected with a photomultiplier and analysed with a computer. 6 Flare intensity is proportional to the protein content of aqueous humour, itself reflecting blood-aqueous barrier disruption. Moreover, flare intensity (scattered light) is proportional to the size of molecules present in the aqueous. 6 Anterior chamber proteins are mainly composed of albumin but also of a proportion of proteins of higher molecular weight such as globulins. 7 As the
exact composition of inflammatory aqueous humour is variable, the standard flare units now routinely used are photon counts per millisecond (ph/ms), rather than traditionally reported bovine albumin equivalent values expressed in milligrams per millilitre.\(^8\) Presently a flare increase of 5–7 ph/ms is empirically considered to be clinically relevant.\(^4,9\) In the particle measurement mode, the laser flare cell meter counts cells or other particles in a volume of 0.075 mm\(^3\) and the units used are number of cells per 0.075 mm\(^3\). The instrument is programmed to count only particles of a size between 9–12 \(\mu\)m, ideally counting only inflammatory cells.\(^3\)

Flare difference between the affected (tumour) eye and the healthy eye was statistically analysed using Student’s paired \(t\) test. Flare values (absolute mean flare and mean differential flare between affected (tumour) and healthy eye) between groups were statistically analysed, using Student’s unpaired \(t\) test, applying Bonferroni’s correction when multiple comparisons were made. The proportion of clinically relevant flare increases (>5 ph/ms and >7 ph/ms) between groups was analysed using the \(\chi^2\) test.

All medium and large melanomas (n=92) were treated by proton beam irradiation. For this more homogeneous group computerised tumour variables were available and were correlated with the flare value. For this purpose the 92 patients of this group were subdivided into three subgroups according to flare value: flare <10 ph/ms, flare from 10 to 20 ph/ms, and flare >20 ph/ms.

The tumour variables studied were: (1) tumour height (H=mm), determined by B scan real time ultrasonography performed by two different examiners; (2) the largest tumour diameter (LTD=mm) determined by a computerised mapping of the tumour serving for the proton beam treatment plan; (3) the tumour volume (V=mm\(^3\)), calculated by a mathematical formula using the largest and smallest tumour diameter and height; (4) the location of the anterior margin of the tumour with reference to the equator of the globe determined with the same computerised mapping and expressed in negative (−) mm when posterior to the equator and in positive (+) mm when anterior to the equator; (5) the ciliary body invasion determined preoperatively by transillumination and by indirect binocular ophtalmoscopy (percentage of + cases); (6) Bruch’s membrane rupture, estimated clinically by indirect ophtalmoscopy, and/or by fluorescein angiography, and/or by ultrasonography (percentage of + cases); (7) retinal detachment estimated clinically by indirect binocular ophtalmoscopy (0–1–2–3 quadrants); and (8) the delay from diagnosis to flare determination (months). Flare was correlated with these eight tumour variables using a multiple linear regression model.

### Results

In all three studied groups (small, medium and large, and very large tumours) the absolute flare values were significantly higher in the tumorous eye than in the healthy eye (Table 2). There was also a significant linear progression from the small to the medium-large group and from the medium–large to the very large group both for mean absolute flare and mean differential flare (Table 2). The proportion of cases with a flare difference between paired eyes of 5 ph/ms or more and 7 ph/ms or more also significantly increased from one group to the other (Table 2). No significant difference of anterior chamber cells was found between paired eyes in any of the three groups.

For the group of medium–large melanomas treated by proton beam irradiation, the eight clinical and computer generated variables were compared in the three flare subgroups (flare <10 10–20; flare >20) using paired Student’s \(t\) test.
Discussion

Breakdown of the blood-aqueous barrier resulting in an influx of proteins (flare) can be caused by inflammation or by any other as yet unexplained non-inflammatory insult to the blood-aqueous barrier such as in diabetes.\textsuperscript{10} Blood-aqueous barrier disruption associated with choroidal melanomas was only rarely reported even in large series. In a group of 450 melanomas examined with the slit-lamp, Fraser et al reported an elevated flare in the anterior chamber in 4-9% of the cases.\textsuperscript{2} Using laser flare photometry we found that anterior chamber flare was elevated much more often in choroidal melanoma than previously suspected by slit-lamp examination. Flare was significantly higher in the tumorous eye when compared with the healthy fellow eye in groups of melanomas of all sizes. An empirically determined increase of flare of 5–7 ph/ms or more is presently considered clinically relevant.\textsuperscript{1,9} An increase of ≥5 ph/ms occurred in 62-6% and an increase of ≥7 ph/ms occurred in 50-4% of all cases and reached 93.3% in the group of very large melanomas, indicating that the tumour was causing a distinct breakdown of the blood-aqueous barrier in a significant number of cases. The amount of flare was linearly correlated with the size of the melanoma, being highest in the group of very large melanomas.

Multiple linear regression coefficients showed that the amount of flare was related to the size of the tumour, as flare value was clearly correlated both with tumour volume and height. The location of the tumour and whether it invaded the ciliary body or not, Bruch’s membrane rupture, the associated retinal detachment, and the delay from diagnosis to flare determination did not influence the amount of flare, indicating that, except for size (volume and height), no other clinical variable influenced the amount of blood-aqueous barrier breakdown. The reason why melanomas cause blood-aqueous breakdown is not known and can only be hypothesised. Recent experimental studies demonstrated the proinflammatory properties of melanin or of a melanin associated protein. In a mouse uveitis model where pretreated animals were challenged in the anterior chamber with horse serum or conalbumin, melanin was shown to increase inflammation.\textsuperscript{11} Binding of melanin to serum components such as antibodies was thought to be a possible explanation for its proinflammatory activity.\textsuperscript{11} Broekhuysen et al and other groups recently reported that a protein fraction extracted from the retinal pigment epithelium and from the choroid acted as an autoantigen and produced an autoimmune anterior uveitis in rats, called experimental autoimmune anterior uveitis (EAAU).\textsuperscript{12,13} EAAU could also be elicited with an extract from the iris and ciliary body.\textsuperscript{14} If a melanin associated protein is responsible for blood-aqueous breakdown, it seems logical that tumours of great size containing a larger amount of protein associated with melanogenesis would have an increased tendency to produce a higher degree of blood-aqueous

\[ \text{Flare} = 6.80 + 1.3513 \times H \]

\text{Regression coefficient: 0.41}

\[ \text{Flare} = 9.20 + 0.0063 \times V \]

\text{Regression coefficient: 0.43}

Figure 1 Regression line of flare values (photons/ms) in correlation with tumour height (mm) (A) and tumour volume (mm$^3$) (B).

\( t \) test (Table 3). There was a significant, linear progression from subgroup 1 to subgroup 2 and from subgroup 2 to subgroup 3 only for tumour volume and tumour height, indicating that flare increase seemed to be related to tumour volume and height (Table 3). This was confirmed when these variables were analysed by multiple regression analysis. Flare increase showed a significant correlation with tumour volume \((r=0.43, p<0.0001; \text{Fig} 1A)\) and tumour height \((r=0.41, p<0.0008; \text{Fig} 1B)\), but not to largest tumour diameter \((r=0.27, p=\text{NS})\), location of the anterior margin of the tumour \((r=0.32, p=\text{NS})\), ciliary body invasion \((r=0.23, p=\text{NS})\), associated retinal detachment \((r=0.22, p=\text{NS})\), Bruch’s membrane rupture \((r=0.26, p=\text{NS})\), or to delay from diagnosis to flare determination \((r=0.05, p=\text{NS})\).
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barrier breakdown. Other factors that possibly might influence blood-aqueous barrier breakdown are alterations of retinal vascular permeability overlying melanomas, lymphocytic infiltration, and amount of necrosis of the tumour.15 16

The most likely hypothesis, however, is that choroidal melanomas cause blood-aqueous barrier disruption in a similar fashion to diabetes. It was shown by laser flare photometry that the amount of diabetes associated blood-aqueous barrier disruption was directly related to the degree of diabetic retinopathy.10 17 These changes in blood-ocular barrier permeability are thought to be caused by diffusible angiogenic factors generated by hypoxic tissue. One of these factors, vascular endothelial growth factor (VEGF), that has been recently purified controls both angiogenesis and vascular permeability and was found to be significantly elevated in vitreous of patients with proliferative diabetic retinopathy.18 19 'Tumours also produce similar 'tumour angiogenic factors' causing blood vessels from adjacent normal tissue to grow towards and into the tumour providing the tumour with oxygen and other nutrients. As in diabetic retinopathy where flare is proportional to the extent of ischaemic areas, flare was directly related to the extent of the tumour (melanoma size). Another similarity to proliferative diabetic retinopathy is the neovascular glaucoma that can develop after irradiation of large choroidal melanomas despite limited irradiation of the adjacent retina obtained with the proton beam. This could be the result of massive release of diffusible angiogenic factors from hypoxic tumour tissue and/or radiation retinopathy as in proliferative diabetic retinopathy.20 21

Finally, it is worth stressing that laser flare photometry should not be considered as a sensitive diagnostic help in the examination of choroidal melanomas as suggested earlier in a study on 11 melanomas.22 Taking the lower value (5 ph/ms) of what is considered a clinically relevant flare rise, more than one third of melanomas (37.4%) had an increase of less than 5 ph/ms and in the group of small melanomas this proportion was as high as 72%.