Foveal electroretinograms and choroidal perfusion characteristics in fellow eyes of patients with unilateral neovascular age-related macular degeneration

Juancho F C Remulla, Alexander R Gaudio, Sumiko Miller, Michael A Sandberg

Abstract

Aims/Background—A prolonged choroidal filling phase on fluorescein angiography has been reported to be a common finding and associated with visual function abnormalities in patients with age-related macular degeneration (AMD). This investigation was carried out to determine whether this perfusion defect was related to the slowing of foveal cone electroretinogram (ERG) implicit time seen in patients with AMD.

Methods—Fluorescein angiograms and foveal cone ERGs were evaluated in the fellow eyes of 67 patients with unilateral neovascular AMD.

Results—Twenty-eight (42%) of the eyes had a choroidal perfusion defect. ERG implicit times averaged 1 ms slower (p=0.0167) and were more likely to be delayed (p=0.0078) in eyes with abnormal choroidal perfusion than in eyes with normal choroidal filling; significant relations were found also after controlling for age. ERG implicit time was also inversely related to ETDRS visual acuity and positively related to the extent of macular drusen; and the latter showed a borderline significant tendency to be more prevalent in eyes with prolonged choroidal perfusion. However, an association of a delayed ERG implicit time with prolonged choroidal filling remained after controlling for age, acuity, and the extent of drusen.

Conclusion—These findings further establish prolonged choroidal perfusion as a common finding in AMD and link it to retinal malfunction.


A prolonged choroidal filling phase on fluorescein angiography has been reported for 26% of 100 cases1 and 33% of 126 cases2 with age-related macular degeneration (AMD), and for 66% of 106 cases with the neovascular form of AMD.3 This perfusion abnormality is associated with patchy elevations of dark adapted threshold,4 an increased risk for developing geographic atrophy,5 and a slowed time course of cone and rod dark adaptation.6 It has been thought to reflect either an attenuated choriocapillaris, seen histologically,7,8 together with a diffusely thickened Bruch's membrane,9,10 or a thickened Bruch's membrane alone.2

We recently reported that fellow eyes of patients with unilateral neovascular AMD have on average foveal cone ERG amplitudes that are normal, but implicit (peak) times that are slower than normal.10 Since a defect in choroidal filling is apparently common in patients with AMD1–3 and is associated with visual dysfunction,4,6 we investigated whether this angiographic finding was related to the slowed foveal ERG seen in the fellow eyes of patients with the unilateral neovascular form.

Methods

We evaluated the fellow eyes of 67 patients (ages 61 to 89) with unilateral neovascular AMD. These patients were part of a larger group with unilateral neovascular AMD who are being followed prospectively with foveal cone ERGs and other tests of cone function to predict who will develop a choroidal neovascular membrane in the fellow eye; demographic characteristics and ocular findings of this larger group have been described previously.11 Eligibility criteria for the present report included a corrected Snellen visual acuity of 20/60 or better, sufficiently clear media to allow detailed evaluation of the fundus, macular drusen, and no sign of other retinal disease in the study eye.

These study eyes had a foveal cone ERG recorded and had angiograms of clear, readable quality throughout the transit. The angiograms were read by two of the authors (JCR and ARG), masked with respect to results of foveal cone electroretinography, to identify those angiograms with a prolonged choroidal filling phase; the latter was characterised by a non-uniform fluorescence extending over at least 5 disc diameters of the posterior pole persisting through the onset of the venous phase of the retinal circulation.1

Foveal cone ERGs were elicited with a 4° white stimulus flickering at 42 Hz presented by a hand held, dual beam stimulator ophthalmoscope (Maculoscope, Doran Instruments, Littleton, MA, USA) as previously described.10,12 Responses were monitored with a contact lens electrode, amplified, filtered, digitised, summed, and quantified by Fourier analysis with respect to amplitude and phase; phase was then converted to implicit time (that is, time interval from stimulus onset to the corresponding cornea positive response peak).
Possible confounding factors for the relation of foveal cone ERG implicit time to choroidal perfusion characteristics were also evaluated in the study eyes: we measured their visual acuity on an ETDRS chart,\(^\text{11}\) coded the presence of a lens opacity in non-pseudophakic eyes, and estimated their area of macular drusen from fundus photographs\(^\text{13}\) without reference to the ERG and angiographic results. Since the frequency distributions of drusen area and foveal cone ERG amplitude were positively skewed, both measures were converted to common logarithms to approximate better a normal distribution. This research followed the tenets of the Declaration of Helsinki and was approved by the internal review boards of the Massachusetts Eye and Ear Infirmary and Harvard Medical School.

**Results**

Angiographic evidence of prolonged choroidal filling was demonstrated in 28 of the 67 eyes (42%). Patients with prolonged choroidal filling were similar to those with normal choroidal filling with respect to mean age, mean ETDRS acuity in the study eye, and the prevalence of a lens opacity in the study eye (Table 1). On the other hand, patients with prolonged choroidal perfusion had a tendency to have more extensive macular drusen (Table 1).

The mean (SE) for foveal cone ERG implicit time was 38.3 (0.4) ms for patients with a prolonged choroidal filling phase and 37.0 (0.3) ms for those with normal choroidal filling; these means were significantly different (\(t\) test, \(p=0.0167\)). If we controlled for the positive relation between ERG implicit time and age by multiple regression,\(^\text{10}\) then mean implicit times for the two groups of patients remained significantly different (Table 2). If we divided the patients into those with a delayed ERG implicit time (that is, greater than 38 ms\(^\text{14}\)) and those with a normal implicit time, then the presence of a delayed implicit time was significantly related to the presence of prolonged choroidal filling by contingency table analysis (Table 3; \(p=0.0078\)) and by logistic regression controlling for age (\(p=0.0161\)). If a patient had a prolonged perfusion time, then the average odds were 3.9 to 1 that the patient also had a delayed implicit time. An example of a fluorescein angiogram with abnormal choroidal filling from a patient with a delayed ERG implicit time is illustrated in Figure 1.

We also found that a slower or delayed implicit time was inversely related to ETDRS acuity (0.0098 and 0.0229, respectively) and positively related to the extent of drusen (\(p=0.0026\) and \(p=0.0375\), respectively) when controlling for age; there was no relation of a slower or delayed implicit time to the presence

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**Table 1**  Patient age and study eye characteristics by category of choroidal filling

<table>
<thead>
<tr>
<th>Variable</th>
<th>Normal filling</th>
<th>Prolonged filling</th>
<th>(p) Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>72.3 (1.1) years*</td>
<td>74.3 (1.4) years*</td>
<td>0.28164</td>
</tr>
<tr>
<td>ETDRS acuity</td>
<td>38.8 (1.3) letters*</td>
<td>37.2 (1.6) letters*</td>
<td>0.44287</td>
</tr>
<tr>
<td>Lens opacity</td>
<td>58% of cases</td>
<td>56% of cases</td>
<td>0.88181</td>
</tr>
<tr>
<td>Extent of drusen</td>
<td>3.1 (0.6)%*</td>
<td>3.9 (0.9)%*</td>
<td>0.05099</td>
</tr>
</tbody>
</table>

*Mean (SE); the ETDRS acuities transposed to the Snellen scale were 20/27 for patients with normal choroidal filling and 20/29 for patients with prolonged choroidal filling.

**Table 2**  Multiple regression of foveal cone ERG implicit time on age and category of choroidal filling

<table>
<thead>
<tr>
<th>Term</th>
<th>Estimate</th>
<th>SE</th>
<th>(t) Ratio</th>
<th>(p) Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>0.11</td>
<td>0.03</td>
<td>3.32</td>
<td>0.0015</td>
</tr>
<tr>
<td>Choroidal filling*</td>
<td>1.02</td>
<td>0.47</td>
<td>2.17</td>
<td>0.0336</td>
</tr>
</tbody>
</table>

*Normal=0, prolonged=1.

**Table 3**  Distribution of cases by foveal ERG implicit time and category of choroidal filling

<table>
<thead>
<tr>
<th></th>
<th>Normal filling</th>
<th>Prolonged filling</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal implicit time*</td>
<td>28</td>
<td>11</td>
<td>39</td>
</tr>
<tr>
<td>Delayed implicit time</td>
<td>11</td>
<td>17</td>
<td>28</td>
</tr>
<tr>
<td>Total</td>
<td>39</td>
<td>28</td>
<td>67</td>
</tr>
</tbody>
</table>

*31 to 38 ms.

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**Figure 1**  Fluorescein angiogram from the fellow eye of a 66-year-old woman with unilateral neovascular age-related macular degeneration. The fluorescein angiogram shows persistence of a patchy choroidal fluorescence in the posterior pole at the onset of the venous phase. This eye had a foveal cone ERG implicit time of 39.4 ms.
We found that the percentage of fellow eyes of patients with unilateral neovascular AMD and a prolonged choroidal filling phase on fluorescein angiography (42%) fell within the range of previously reported values for eyes with AMD. By combining our results with those previously published, the prevalence of this defect becomes 166 of 399 cases (42%), further establishing it as a common angiographic sign in AMD.

Prolonged choroidal perfusion was associated with a foveal cone ERG implicit time that was slower and more likely to be delayed. This adds an objective measure of retinal malfunction to the previously described psychophysical abnormalities of impaired sensitivity in the dark and slowed recovery during dark adaptation for AMD eyes with this perfusion defect. This relation implicates an abnormality of the choriocapillaris and/or Bruch’s membrane, directly or indirectly, in slowing the foveal cone ERG in this disease.

We also noted that ERG implicit time was inversely related to ETDRS visual acuity and positively related to the extent of macular drusen in our patients. The second finding may help to account for the previous observation that foveal cone ERG implicit time was slower in fellow eyes with normal visual acuity of patients with unilateral neovascular AMD many of whom have extensive drusen, than in normal subjects, who have minimal (if any) drusen. Our patients with prolonged choroidal perfusion also showed a borderline tendency to have more extensive drusen. Nevertheless, prolonged choroidal perfusion remained a significant predictor of a delayed ERG implicit time even after controlling for acuity and extent of drusen. The association of ERG implicit time to prolonged choroidal perfusion also could not be explained by a difference in media clarity between the patients with and without the perfusion defect, because the two groups had comparable prevalences of lens opacity and because implicit time was unrelated to the presence of a lens opacity.

Foveal cone ERG amplitude was unrelated to the choroidal perfusion characteristics of the study eyes. A delayed cone ERG response with normal amplitude may be seen in the full field cone ERG of some patients with central retinal vein occlusion (Sandberg and Gaudio, unpublished observations) and in the foveal cone ERG of some patients with diabetic macular oedema. Conceivably, a vascular component may be similarly implicated in eyes with prolonged choroidal filling. It is possible that the angiographic finding of prolonged choroidal perfusion in patients with AMD is reflective of outer retinal ischaemia, resulting in foveal cone ERGs of normal amplitude but delayed implicit time.

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Discussion
We found that the percentage of fellow eyes of patients with unilateral neovascular AMD and a prolonged choroidal filling phase on fluorescein angiography (42%) fell within the range of previously reported values for eyes with AMD. By combining our results with those previously published, the prevalence of this

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History of ophtalmology

Discovering the uses of the ophthalmoscope

In the early years after Helmholtz invented the ophthalmoscope, the even greater task of discovering what was normal and what was pathological began. Not that the instrument was rapidly accepted – to quote the editor of the Medical Times, it was ‘ridiculed and denounced’ by many physicians as a dangerous toy! Jabez Hogg of the Westminster Ophthalmic Hospital worked hard to redeem this situation by publishing a book on its use. However, those attending the Ophthalmological Congress in Brussels in 1857 debated its utility at length.

Summaries of the normal and abnormal appearances of the retina began to appear in the literature. Opacities in the lens were noted, and Williams describes the ‘beautiful opaque streaks’ with enthusiasm. Floating bodies in the vitreous were seen ‘bounding and darting about’ and were considered common, although their cause was a mystery.

By 1865, funduscopy was confidently used in the diagnosis of renal disease. For example, ‘Mary Ann’ was a sewing machinist who had undergone much hardship. Her sight had begun to fail since Christmas 1864 when she had ‘sat all day in wet clothes’, and she presented to Mr Hulke. On ophthalmoscopy, her retinal veins were ‘turgid’, and grey-white patches were visible. Kidney disease was diagnosed, and confirmed by the finding of albumin and casts in the urine. Mr Hulke sent ‘Mary Ann’ away with some muriate of iron, and exhorted his colleagues to remember the use of fundoscopy in renal disease. Dr Hughlings Jackson agreed with this point, while stressing the association of fundal changes with cerebrovascular accidents. A pithy description of a patient with retinal haemorrhages who was ‘in a ripe condition for extensive cerebral apoplexy’, but ‘also ready to die in many other ways’ followed. It is hoped that this opinion was not communicated to the patient, or at least that his alleged slight deafness saved him from hearing it!

Later, in the early 1900s, more detailed investigation of ‘flame-shaped haemorrhages’ and arteriovenous nipping took place, and the connection with hypertension firmly established. Understandably, some misdirections occurred:

Bardsley differentiated the signs of arteriosclerosis from hypertension, and believed that the retinal signs could appear and disappear rapidly. Armed with a large supply of adrenaline and chloroform, a house officer he set out to prove this on his patients. After injecting enough adrenaline to secure a rise in blood pressure of at least 45 mm Hg – an effect which could be achieved in an ethics committee chairman merely by reading this account – the retina was examined. Bardsley reported that fullness of veins and AV nipping was seen – just as he had previously noted in cases of toxema, and that it disappeared when the blood pressure fell. Indeed, having tried this on all the patients, he felt that it should be possible to indicate the blood pressure merely from fundoscopy.

Others felt that even established arteriosclerosis was reversible. Crofton described a 60-year-old man with small, tortuous retinal arteries and severe AV nipping. Advised of this, the patient went on a sea voyage. On his return 2 months later, Crofton reported that all silver wiring was gone, and the AV nipping had considerably lessened.

Not all physicians believed in the prognostic implications of high blood pressure, although many prohibited the use of the eyes, feeling that the strain of vision might worsen the disease. Adams described a 64-year-old with an arterial pressure of 250 and a host of flame-shaped haemorrhages, who lived a further 4 years. ‘Why treat?’ insisted Adams, as hypertension is a sign to which the patient does not object.

Doubtless these physicians would be fascinated to hear about the developments that have occurred, although if Crofton did cure the retinal signs of hypertension, then it seems that some valuable knowledge has been lost.
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