Localized Retinal Electrophysiological and Fundus Autofluorescence Imaging Abnormalities in Maternal Inherited Diabetes and Deafness

Caren Bellmann,1,2 Magella M. Neveu,1 Hendrik P. N. Scholl,1,2 Chris R. Hogg,1 Pamela P. Rath,1 Sbaron Jenkins,2 Alan C. Bird,1,2 and Graham E. Holder1

PURPOSE. To investigate retinal function in patients with maternally inherited diabetes and deafness (MIDD) and to correlate the findings with fundus autofluorescence (FAF) imaging.

METHODS. FAF was imaged in five patients (age range, 49–60 years) confirmed to have the mitochondrial DNA nucleotide A3243G point mutation. Retinal function was measured by full-field (Ganzfeld) electroretinography (ERG) and pattern ERG, incorporating the International Society for Clinical Electrophysiology of Vision (ISCEV) standards. Multifocal ERG (mfERG) was also performed. For analysis of the mfERG data, five regional ring groups of equal eccentricity were formed. For each ring, the peak amplitude (defined as the difference between P1 and N1) and the implicit time of P1 were determined and compared with normative values.

RESULTS. Visual acuity in the patients was between 20/20 and 20/40 (Early Treatment Diabetic Retinopathy Study [ETDRS] chart). Irregular increased FAF signals were observed adjacent to and between areas of atrophy of the retinal pigment epithelium (RPE). Ganzfeld ERGs were within normal limits in three patients. Pattern ERG was abnormal in five eyes of three patients. mfERG peak amplitude abnormalities were particularly present in rings 2 and 3 and were consistent with the distribution of FAF abnormalities. In all but one eye, no implicit times changes were present.

CONCLUSIONS. Significant mfERG abnormalities with normal Ganzfeld ERG are consistent with nonuniform damage to the central retina in MIDD, in keeping with the FAF findings. Reduced peak amplitudes with normal implicit times in the mfERG suggest localized loss of function and may indicate damage to the cone photoreceptor outer segments or cone photoreceptor loss in MIDD. (Invest Ophthalmol Vis Sci. 2004;45:2355–2360) DOI:10.1167/iovs.03-1090

From 1Moorfields Eye Hospital, London, United Kingdom; and the 2Institute of Ophthalmology, London, United Kingdom.

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Corresponding author: Graham E. Holder, Department of Electrophysiology, Moorfields Eye Hospital, City Road, London EC1V 2PD, UK; graham.holder@moorfields.nhs.uk.

FAF Imaging

FAF imaging was recorded with a confocal scanning laser ophthalmoscope (HRA; Heidelberg Engineering, Dossenheim, Germany). As described previously,15–17 an argon blue laser with a wavelength of 488 nm was used for excitation. Emission was recorded at higher than 500 nm. Before examination, the pupil of the study eye was dilated with phenylephrine 2.5% and tropicamide 1%. Recordings were performed with best image focus and after correction of refractive errors.
Electrophysiological Recordings

Assessment included Ganzfeld electroretinography (ERG) and PERG recorded incorporating the protocols recommended by the International Society for Clinical Electrophysiology of Vision (ISCEV).18,19

The mfERG was recorded with a commercial system (RETIscan System; Roland Consult, Wiesbaden, Germany). The stimulus used in the mfERG consisted of 61 hexagons, covering in total a visual field of 56.9°, at a viewing distance of 33 cm. The stimulus-distortion factor was set to 4 to compensate for differences in cone density across the retina, from center to periphery. Patients were asked to fixate on the center of a large cross in the central hexagon. Before examination, eyes were kept light adapted for at least 10 minutes. Pupils were dilated with phenylephrine 2.5% and tropicamide 1%. Average pupil diameter was 8 mm. Refractive errors were corrected. The mfERG was recorded binocularly in three patients and monocularly in one patient who had strabismus. One patient declined the mfERG.

All mfERG responses were recorded using corneal gold foil electrodes. The reference and ground electrodes were attached to the ipsilateral outer canthus and forehead, respectively. The surface electrode impedance was less than 5 kΩ. The signals were amplified and band-pass filtered (5–300 Hz; −3dB). Eight trials were recorded for each mfERG session, and the average recording time for each trial was 50 to 120 seconds. The total duration of a recording session was 20 minutes or less. These techniques are similar to those used in previous studies.20,21

The mfERG recordings were compared with FAF imaging results by superimposing the 61 trace arrays onto FAF images with external image analysis software (Photoshop 6.0; Adobe Systems, Mountain View, CA). In addition to qualitative analysis, further analyses were performed with the system (RETIscan; Roland) software. In particular, first-order kernels were considered for mfERG evaluation, because they correlate with the function of the outer retina.15

To reduce the complexity of the local 61 mfERG responses, five regional ring groups of equal eccentricity were formed. The ring groups were defined as follows: ring 1, from 0° to 2.1° eccentricity; ring 2, from 1.4° to 6.7° eccentricity; ring 3, from 5.7° to 12.0° eccentricity; ring 4, from 9.5° to 19.8° eccentricity; ring 5, from 15.1° to 28.5° eccentricity. For each ring group, the peak amplitude, defined as the difference between N1 and P1, was calculated, and the implicit time of P1 was determined. For analysis of the mfERG temporal data, the analysis was restricted to the implicit time of the positive deflection (P1). Results of peak amplitude and implicit time were compared with normative values ranging between the 5th and the 95th percentiles (Fig. 1).

RESULTS

Four female patients and one male patient were studied (age range, 49–60 years). All patients had significant neurosensory hearing loss. None showed signs of diabetic retinopathy, and all had stable glucose levels at the time of examination. The clinical data, together with Ganzfeld ERG and PERG are described in Table 1. mfERG results are demonstrated in Figure 1. In normal subjects, the amplitudes of each of the 61 local mfERG responses are almost equal, because of the stimulus scaling factor with increasing hexagon size toward the periphery, compensating for lower cone density (Fig. 2).

Patient 1

Visual acuity was 20/20 in both eyes. Fundus findings showed a symmetrical bilateral retinal involvement with irregular...
patches of the retinal pigment epithelium atrophy at the posterior pole. The fovea was normal. FAF imaging revealed a decreased FAF signal in areas of RPE atrophy. An irregular increased FAF signal was observed adjacent to and between the areas of RPE atrophy.

Ganzfeld ERGs showed mildly subnormal rod-specific responses as well as subnormal maximal ERG a- and b-wave amplitudes from both eyes. The 30-Hz flicker ERG showed delayed implicit times but no amplitude abnormalities. The PERG amplitude was below the normal range indicating macular involvement. Although the mfERG showed reduced peak amplitudes with normal implicit times in all five ring groups (Fig. 1), mfERG changes were particularly present in areas of FAF abnormalities and without any substantial reduction in amplitude of the response to the central hexagon (Fig. 3A).

Patient 2

Visual acuity in the right eye was 20/40 and in the left eye 20/32. Extensive areas of RPE atrophy, sparing the fovea, were present in both eyes on ophthalmoscopy. The right eye was more severely affected, leaving only a small island centrally. A band of increased FAF adjacent to the RPE atrophy was observed in both eyes (Fig. 3B). Scotopic and photopic Ganzfeld ERGs were bilaterally normal. The PERG was undetectable from the right eye, indicating severe maculopathy and reduced from the left eye. mfERG evaluation revealed reduced peak amplitudes in ring groups 2 and 3 from both eyes and additionally in ring group 1 on the right corresponding to the areas of RPE atrophy (Fig. 1A). Implicit times were within the normal range in all ring groups.

### Table 1. Visual Acuity and Electrophysiological Findings

<table>
<thead>
<tr>
<th>No.</th>
<th>Age</th>
<th>Eye</th>
<th>VA</th>
<th>Ganzfeld ERG (30 Hz) Amplitude</th>
<th>Implicit time</th>
<th>Rod-Specific b-wave Amplitude</th>
<th>Implicit Time</th>
<th>Pattern ERG Amplitude (P 50)</th>
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<tbody>
<tr>
<td>1</td>
<td>52</td>
<td>RE</td>
<td>20/20</td>
<td>n</td>
<td>a</td>
<td>a</td>
<td>n</td>
<td>a</td>
</tr>
<tr>
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<td>55</td>
<td>RE</td>
<td>20/40</td>
<td>n</td>
<td>n</td>
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<td>4</td>
<td>60</td>
<td>RE</td>
<td>20/25</td>
<td>n</td>
<td>n</td>
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<tr>
<td>5</td>
<td>56</td>
<td>RE</td>
<td>20/25</td>
<td>a</td>
<td>a</td>
<td>n</td>
<td>n</td>
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</tr>
</tbody>
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n, normal; a, abnormal.

![Figure 2. mfERG superimposed onto a FAF image in a normal subject (left eye). Amplitudes of the 61 local ERG responses are almost equal, because of the stimulus scaling with increasing hexagon size toward the periphery, compensating for lower cone density.](image-url)
groups on the left eye and mildly delayed in the second ring group on the right (Fig. 1B).

**Patient 3**

Visual acuity was 20/20 in both eyes. Widespread areas of perimacular RPE atrophy were distributed in both eyes (Fig. 3C). The FAF signal was extinguished in areas of RPE atrophy, whereas an irregular increased FAF signal was found adjacent to and between the areas of atrophy. In comparison to patient 1 the fundus changes were located at greater eccentricity (Fig. 3A).

The Ganzfeld ERG was normal. Although the PERG was within the normal range in each eye, the mfERG showed reduced central peak amplitudes. In addition, peak amplitudes were reduced or borderline in ring group 3, whereas those in ring groups 2, 4, and 5 were normal. The implicit times were all normal (Fig. 1).

The overlay of the mfERG onto the FAF image is shown in Figure 3C. In comparison to areas with normal FAF background, the peak amplitudes were reduced both in areas of increased and decreased FAF. No distinction could be made between areas of RPE atrophy and areas with increased FAF.

**Patient 4**

The visual acuity measured in both eyes was 20/25. Areas of small RPE atrophy and pigmentary changes were seen on clinical examination bilaterally. The location and appearance of the fundus changes were similar to those described in patient 3, although the areas of RPE atrophy were more restricted. The FAF pattern was similar to that described before showing an irregular, increased FAF signal adjacent to and between the areas of RPE atrophy forming a ring-shaped pattern dystrophy at the posterior pole.

Ganzfeld ERG and PERG were normal. mfERG peak amplitudes were reduced only in ring group 2 from the right eye. The implicit times were normal in all rings (Fig. 1).

On visual inspection, the central hexagon showed no significant reduction, whereas reduced amplitudes were found in...
areas of increased and decreased FAF signals on both eyes (Fig. 3D).

**Patient 5**

The visual acuity in each eye was 20/25. Clinical examination revealed the typical changes of MIDD with perimacular RPE atrophy and pigmentary changes. Drusen-like changes were present in the fovea. FAF images were similar to those in patient 3. A decreased FAF signal in areas of RPE atrophy and an increased FAF signal adjacent to these areas were observed. The rod-specific Ganzfeld ERG was normal, whereas the 30-Hz cone flicker ERG showed delayed implicit times from both eyes and reduced peak amplitudes on the right eye. The PERG was normal in the right eye and abnormal in the left (Fig. 4).

The patient declined mERG examination.

**DISCUSSION**

Clinical examination and FAF imaging in MIDD revealed a typical pattern with areas of irregular increased FAF adjacent to and between the areas of RPE atrophy (Rath PP, et al. IOVS 2002;43:ARVO EAbstract 4345) (Figs. 3, 4). The electrophysiological data revealed nonuniform central retinal dysfunction consistent with abnormalities suggested by FAF imaging. Only two patients demonstrated Ganzfeld ERG abnormalities suggestive of generalized retinal dysfunction, consistent with previous reports describing normal electrophysiology in most patients.10–13,22 The Ganzfeld ERG, however, is a mass response and is thus unaffected by localized dysfunction such as a pure maculopathy.

The PERG and the mERG are measures of central retinal function. A marked reduction in the amplitude of the PERG P50 component suggests dysfunction anterior to the ganglion cells in the visual pathway.14 The PERG within the normal range in some of the patients is in keeping with localized retinal dysfunction surrounding the macular and sparing the fovea, as revealed both in fundus examination and FAF imaging. In other patients, there was sufficient retinal dysfunction to cause absolute abnormalities in the PERG.

The mERG provides spatial information about the central retinal cones and their postreceptor cells not readily available in the Ganzfeld ERG.15 The mERG has been used for investigation for a variety of macular diseases.15,25–26 The present mERG analysis in MIDD reveals reduction of peak amplitudes whereas the implicit times were normal in all but one eye. These findings are compatible with loss of cone photoreceptor outer segments as hypothesized by Hood.15 One patient demonstrated a normal PERG, although the central hexagon of the mERG showed reduced peak amplitudes. The mERG may occasionally reveal subtle early changes in macular function that are not enough to produce an abnormal PERG.

In addition to the ring group analysis, we compared the mERG results with FAF findings. With the 61-hexagon stimulus, mERG responses were recorded over the optic nerve head. This is not an artifact. The hexagons do not fall completely on the optic disc, and light-scattering may contribute to the almost normal responses over the optic nerve head.27 Qualitative analysis revealed no substantial peak amplitude reduction in the central element. Amplitude changes in the mERG superimposed onto the FAF image corresponded well with the area of abnormality seen with FAF imaging (Fig. 3).

It is thought that an increased FAF signal is the result of a high metabolic turnover of photoreceptor outer segments leading to lipofuscin accumulation in RPE cells.28 Both the electrophysiological and imaging data suggest restricted photoreceptor damage in MIDD rather than generalized retinal dysfunction. In histologic study of patients with A3243G mutation and Kearns-Sayre or MELAS syndrome, investigators reported atrophic photoreceptor outer segments and abnormal mitochondria in the inner retinal segments. Ultrastructural changes were present in RPE cells together with enlarged mitochondria.9 It was postulated that photoreceptor outer segments are stressed both from the RPE and from photoreceptor inner segments, which contain 90% of the retina’s mitochondria.9,29

Although the FAF pattern in MIDD is specific, the changes have similarities with the pattern described in patients with geographic atrophy due to age-related macular degeneration.16 It is known that mitochondrial alterations are not only described in maternal inherited diseases but also appear in any aging postmitotic cell,30 so that there may be a common pathogenic pathway in the development of atrophy of the RPE. Mitochondria are involved in a number of metabolic processes including the generation of chemical energy in form of adenosine triphosphate (ATP).31 They are the main source of reactive oxygen species formation and important control centers for apoptosis.30–32 Thus, it is no surprise that mitochondrial DNA shows a high mutation rate and that mutations accumulate with age.33 Furthermore, defective mitochondria may be not properly autophagocytosed. Their components may undergo further oxidative modification within the lysosomes, resulting in the formation of additional undegradable material, such as lipofuscin in RPE cells, and progressively less mitochondrial recycling. Consequently, compensatory mechanisms may fail with time, followed by dysfunction and cell death, particularly in relation to postmitotic tissues with high energy demand, such as photoreceptors and RPE cells.

Age-dependent somatic selection favors the persistence of mitochondria carrying the mutation in many mitochondrial diseases and symptoms may appear only later in life. This is in
keeping with the slow progression and localized damage in our patients and in patients with other mitochondrial diseases.34
Rods photoreceptor dysfunction may be predicted in patients with MIDD, as much of the generated energy in rods supports the ionic pumps that keep the cell in a response-ready state (i.e., for photoreceptor outer segment disc turnover and for the phototransduction cascade).35 If rods were the first location susceptible to the effects of the A3243G mutation, it would explain the striking location of retinal changes, as histologic data describe a maximum rod density at 4 to 6 mm from the fovea.36 Nevertheless, global rod system ERG abnormalities in the Ganzfeld ERG were observed in only one patient.

The results clearly demonstrate a nonuniform retinal damage and suggest damage to the cone photoreceptor outer segments in MIDD. The results support further the hypothesis that both photoreceptor outer segments and RPE cells are involved in the pathogenesis of MIDD, consistent with histologic data. However, whether the two cell layers are equally susceptible to the effects of the A3243G mutation or the photoreceptor damage is a consequence of the disease principally involving the RPE is yet to be ascertained.

Acknowledgments

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References

15. Hoed DC. Assessing retinal function with the multifocal tech-

407–416.