Assessment of Central Retinal Function in Patients with Advanced Retinitis Pigmentosa

Christina Gerth, Tom Wright, Elise Héon, and Carol A. Westall

PURPOSE. To assess central retinal function in patients with advanced retinitis pigmentosa (RP) using the multifocal (mf)ERG and static perimetry.

METHODS. Patients with RP; a nonrecordable, full-field (ff)ERG; and visual acuity (VA) of ≤ 1.0 logMAR were included. All patients underwent mfERG testing (103 hexagons, and 2.67 and 5.33 cd · s · m⁻² flash intensities) and static perimetry (103 corresponding areas) in the better eye. First-order kernel mfERGs were analyzed for total noise, signal-to-noise ratio, response amplitude, and implicit time. The number of areas with recordable mfERG responses were counted and compared with visual field (VF) sensitivity.

RESULTS. Twenty-nine patients aged 16 to 68 years with a VA of 0.02 to 1.0 logMAR and a kinetic VF of 10° to 60° in diameter were included. mfERGs were successfully performed in 22 of 29 patients. Responses were detected in at least one stimulated area in 22 of 22 patients, with an overall response detection of 9.8% in all stimulated areas and no difference between flash intensities. All responses were diminished severely in response density P1-N1, with normal P1 implicit time in 50% of the recordings. No predictive factors for recordable mfERG responses were identified. VF results were recorded reliably in 27 of 29 patients, with a 40% response detection rate.

CONCLUSIONS. mfERG responses were recordable in at least one area in all successfully tested patients with advanced RP. Response detection and performance was significantly higher for static perimetry. Static perimetry may be a more sensitive primary outcome measure of central vision function than the mfERG in patients with advanced RP and nonrecordable ffERGs. (Invest Ophthalmol Vis Sci. 2007;48:1312–1318) DOI: 10.1167/iovs.06-0630

Retinitis pigmentosa (RP) refers to a group of hereditary retinal degenerations with a prevalence of approximately 1 in 4000.¹ Characteristic symptoms include nyctalopia, impaired dark adaptation, and a progressive visual field loss that often leads to legal blindness.²⁻⁵ Advanced RP is characterized by a nonrecordable full-field (ff)ERG and a severely constricted visual field of less than 20°, but often well-preserved central vision.⁴ It has been shown that static visual field (VF) testing is a sensitive tool to quantify this preserved central VF in this patient population.⁵⁻⁸ The VF test, however, has limitations: (1) as a behavioral method, the response may undergo signal modulation and processing within the retinal and cortical visual pathway⁹; (2) the test is subjective in nature and relies on the patient’s cooperation and experience with the test. Electrophysiological tests, in contrast, are objective measures of retinal function. The origin of the ffERG (reviewed in Ref. 10) and multifocal (mf)ERG¹¹ responses has been correlated to distinct retinal layers and can indicate the point of cellular dysfunction. The ffERG is often nonrecordable above noise in patients with advanced diseases,²⁻¹² except when using specific signal acquisition and filtering techniques.¹³⁻¹⁵ The mfERG technique developed by Sutter and Tran¹⁶,¹⁷ permits a localized measurement and mapping of the retinal response. The cone-mediated mfERG with localized retinal stimulation may allow quantification of the remaining cone-mediated function, despite an advanced stage of disease.¹⁸⁻²³ This may be particularly useful in characterizing better advanced stages of disease. Several studies involving patients with RP and Usher syndrome with either recordable or nonrecordable ffERG responses suggest that the mfERG allows a quantification of the central cone-mediated function.¹⁸⁻²⁴ Validation of arising treatment options will require reliable outcome measures to quantify retinal function.²⁵ The objectives of our study were (1) to assess the central retinal responses in patients with advanced RP and a nonrecordable ffERG (International Society for Electrophysiology of Vision [ISCEV] standard),²⁶ by using the cone-mediated mfERG, and (2) to compare the mfERG sensitivity with static perimetry in those patients.

In the population tested, we identified mfERG responses in less than 10% of stimulated areas in 76% of the patients tested. Recordable mfERGs exhibited a greater reduction in amplitude than delay in implicit time. No predictive factors for an acceptable signal-to-noise response were identified. The static visual field test was more sensitive in detecting remaining central retinal responses than was the mfERG.

MATERIALS AND METHODS

Subjects

Participants were recruited from the Ocular Genetics Clinic at The Hospital for Sick Children. Written informed consent was obtained from all participants or their guardians in accordance with the tenets of the Declaration of Helsinki. The project was approved by the Research Ethics Board of the Hospital for Sick Children, Toronto. Inclusion criteria were diagnosis of syndromic or nonsyndromic RP with non-recordable ffERG (ISCEV standard), visual acuity (VA) in the better eye of 1.0 logMAR or better, an interpretable Goldmann visual field (test target III₄₄) and the ability to perform the proposed tests. Exclusion criteria were significant media opacities; cystoid macula edema (visualized by ophthalmoscopy and/or optical coherence tomography); or the presence of a maculopathy, glaucoma, nystagmus, myopia greater than −6.00 DS (which could interfere with mfERG responses)²⁷

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or any systemic (e.g., diabetes) or neurologic diseases that could affect vision or the capacity to perform the tests.

Vision Function Assessment
Best-corrected distance VA was assessed on a logMAR (logarithm of the minimum angle of resolution) scale by using the backlit Early Treatment Diabetic Retinopathy Study charts (ETDRS).28 Contrast sensitivity (CS) was measured using the Pelli-Robson Contrast Sensitivity charts at a 1-meter distance.29 Slit-lamp examination and intracocular pressure measurement were performed in all patients before testing. Goldmann visual fields were repeated if they had not been performed within the previous 6 months. The index eye was (1) the eye with the better VA, (2) the eye with the better CS in patients with equal VA in both eyes, or (3) a randomly chosen eye in patients with equal VA and CS in both eyes.

Multifocal ERG
Stimulation and primary analysis were performed using a stimulus-camera-refractor unit (VERIS, ver. 5.1; Electrodiagnostic Imaging [EDI] San Mateo, CA). After full pupil dilation, mfERGs were recorded from the index eye with a bipolar Burian-Allen contact lens electrode with a built-in infrared illuminator (Hansen Ophthalmic Development Laboratory, Coralville, IA, and EDI) placed on the corneal surface of the eye. Refractive errors were corrected by using the built-in refractor unit. The stimulus consisted of 103 scaled hexagons extending 20° in radius flashed in a pseudorandom pattern at intervals of 13.3 ms (m-sequence-length, 2^15 − 1) on a dark background (<1 cd · m^−2), as illustrated in Figure 1A.

A pilot study was undertaken in a similar patient group that indicated that a maximum flash intensity of 5.33 cd · s · m^−2 (400 cd · m^−2/75 Hz), as used by Hood et al.,19 might result in a higher number of retinal responses than a lower flash intensity (lower luminance). To confirm this, each patient was tested twice, first at a maximum flash intensity of 2.67 cd · s · m^−2 (200 cd · m^−2/75 Hz) and consecutively at 5.33 cd · s · m^−2 at a high contrast of 99% and adjusted surround of 50%. Signals were sampled at 1200 Hz (i.e., 0.83 ms between samples). The data were acquired at a gain of 50,000 over a frequency range of 10 to 300 Hz (preamplifier model ICP 511; Grass Telefactor, West Warwick, RI). Stable fixation was achieved displaying a black fixation cross 4° in diameter with a pen size of 20%, which covers less than 40% of the central hexagon and less than 10% of the adjacent hexagons. Stimulus luminance was calibrated with the auto-calibrator (EDI).

The total duration of the m-sequence used was 7 minutes 17 seconds and was divided into 16 segments of 27.29 seconds each. Steady fixation was ensured by viewing the fundus with a built-in infrared fundus camera. The criteria for repeating a segment were movements and/or saccades exceeding 3° or the size of the optic disc. Such movements usually did not exceed two to four times during each recording. Recorded segments with physiological saccades with immediate refixation were not excluded.

The protocol followed the recommended guidelines of the International Society for Electrophysiology of Vision (ISCEV) for basic mfERG, with the addition of the variable luminance.30

Central Visual Fields
Static central visual fields were assessed with the Humphrey Visual Field Analyzer (HFA II; Carl Zeiss Meditec, Inc., Dublin, CA) using a customized program (Humphrey Visual Field; HVF).19,24 The perimeter display was customized to match the mfERG test pattern. Thresholds were measured at 103 locations with a size 3, white test stimulus centered in the middle of each hexagonal area and the SITA protocol. The background illumination was 10 cd · m^−2. Full refractive correction was used. Eye and head tracking was monitored. Each patient was given 1-minute breaks every 3 to 4 minutes of testing to ensure alertness.

![Figure 1](image-url)  ![Figure 2](image-url)
Two iterations of artifact rejection, but no spatial averaging was applied to the raw mfERG data. mfERGs were analyzed for noise to exclude poor-quality recording as follows. The root mean square (RMS) of a higher-order kernel without a signal correlation was measured as a total noise estimate and compared with an already established control dataset \((n/H11005)150\). The control noise estimate approximately follows a normal distribution, as shown in Figure 2. mfERG data with a noise estimate larger than the 95th percentile of the control dataset were excluded from further analysis.

Two different analyses, the signal-to-noise ratio (SNR) analysis and the stretch-fit analysis described by Hood and Li,31 were applied to identify the more powerful method able to extract the expected small responses. (1) SNR analysis: First-order kernel responses were extracted from the epoch lengths 0 to 60 ms (signal-S), 60 to 120 ms (noise1), and 120 to 180 ms (noise2). (The late-epoch window of the first-order kernel response was a more stringent noise estimator than the fourth-order kernel, when applied to the SNR analysis in our patient cohort.) Waveform analysis was performed with custom software written in R.32 Responses with a low SNR may be caused either by high noise or low signal. It is very likely that RP-associated cone-mediated dysfunction will cause a low signal. Therefore, the noise ratio noise1/noise2 was calculated as an indicator for noise distribution for each individual recording. The SNR for each

| TABLE 1. Summary of Visual Function Test Results |

<table>
<thead>
<tr>
<th>Patient</th>
<th>VA (LogMAR)</th>
<th>CS (Log Units)</th>
<th>GVF (III%)</th>
<th>Diameter (°)</th>
<th>HVF Counts</th>
<th>mfERG Counts 200/400</th>
<th>mfERG: RD P1-N1 200/400 (nV/deg²)</th>
<th>mfERG: IT P1 200/400 (ms)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ADRP 1/m/27</td>
<td>0.02</td>
<td>1.5</td>
<td>87</td>
<td>8/5</td>
<td>14.4/3</td>
<td>25/26.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ADRP 2/t/57</td>
<td>0.22</td>
<td>1.05</td>
<td>33</td>
<td>No fix</td>
<td>NA</td>
<td>NA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ADRP 3/t/54</td>
<td>0.4</td>
<td>1.05</td>
<td>39</td>
<td>12/noise</td>
<td>3.3/NA</td>
<td>43.3/NA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ARRP 4/m/16</td>
<td>0.3</td>
<td>1.25</td>
<td>24</td>
<td>5/5</td>
<td>2.6/5.7</td>
<td>31.6/33.3†</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ARRP 5/t/50</td>
<td>0.64</td>
<td>0.45</td>
<td>26</td>
<td>No fix</td>
<td>NA</td>
<td>NA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>XLRP 6/t/21</td>
<td>0.1</td>
<td>1.20</td>
<td>88</td>
<td>6/10</td>
<td>10.3/3.3</td>
<td>22.5/26.6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>XLRP 7/t/49</td>
<td>0.28</td>
<td>1.05</td>
<td>37</td>
<td>8/4</td>
<td>1.4/2.8</td>
<td>31.6/25</td>
<td></td>
<td></td>
</tr>
<tr>
<td>XLRP 8/t/23</td>
<td>0.4</td>
<td>1.25</td>
<td>58</td>
<td>21/23</td>
<td>3.3/2.5</td>
<td>23.3/29.1†</td>
<td></td>
<td></td>
</tr>
<tr>
<td>sRP 9/t/41</td>
<td>0.05</td>
<td>1.5</td>
<td>42</td>
<td>19/5</td>
<td>3.1/25.7</td>
<td>26.6/27.5†</td>
<td></td>
<td></td>
</tr>
<tr>
<td>sRP 10/t/49</td>
<td>0.1</td>
<td>1.2</td>
<td>20</td>
<td>Noise/9</td>
<td>NA/2.1</td>
<td>NA/30.8†</td>
<td></td>
<td></td>
</tr>
<tr>
<td>sRP 11/t/37</td>
<td>0.1</td>
<td>1.20</td>
<td>60</td>
<td>6/13</td>
<td>3.3/5/5</td>
<td>26.6/29.1†</td>
<td></td>
<td></td>
</tr>
<tr>
<td>sRP 12/t/46</td>
<td>0.1</td>
<td>1.45</td>
<td>48</td>
<td>8/13</td>
<td>2.7/1.5</td>
<td>26.6/24.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>sRP 13/m/27</td>
<td>0.1</td>
<td>1.35</td>
<td>45</td>
<td>UR</td>
<td>11/53</td>
<td>15.5/13.6</td>
<td>50.0/27.5†</td>
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</tr>
<tr>
<td>sRP 14/m/45</td>
<td>0.6</td>
<td>0.3</td>
<td>15</td>
<td>42</td>
<td>Noise/11</td>
<td>NA/5.0</td>
<td>NA/26.7†</td>
<td></td>
</tr>
<tr>
<td>sRP 15/t/37</td>
<td>0.4</td>
<td>1.25</td>
<td>20</td>
<td>32</td>
<td>2.4/4.8</td>
<td>25.8/30.8†</td>
<td></td>
<td></td>
</tr>
<tr>
<td>sRP 16/t/40</td>
<td>0.4</td>
<td>1.1</td>
<td>20</td>
<td>65</td>
<td>Noise</td>
<td>NA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>sRP 17/t/34</td>
<td>0.42</td>
<td>1.65</td>
<td>35*</td>
<td>103</td>
<td>Noise</td>
<td>NA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>sRP 18/t/68</td>
<td>0.5</td>
<td>0.6</td>
<td>10</td>
<td>7</td>
<td>11/4</td>
<td>2.6/2.4</td>
<td>28.3/27.5</td>
<td></td>
</tr>
<tr>
<td>sRP 19/t/48</td>
<td>0.52</td>
<td>0.75</td>
<td>40</td>
<td>69</td>
<td>9/17</td>
<td>5.7/5.2</td>
<td>25.8/24.9</td>
<td></td>
</tr>
<tr>
<td>sRP 20/m/50</td>
<td>0.64</td>
<td>0.6</td>
<td>10</td>
<td>6</td>
<td>No fix</td>
<td>NA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>sRP 21/m/35</td>
<td>1.0</td>
<td>0.15</td>
<td>20*</td>
<td>UR</td>
<td>3/9</td>
<td>1.8/9.1</td>
<td>36.6/27.7</td>
<td></td>
</tr>
<tr>
<td>CHM 22/m/47</td>
<td>0.3</td>
<td>1.35</td>
<td>20*</td>
<td>35</td>
<td>Noise/6</td>
<td>NA/1.8</td>
<td>NA/25.8</td>
<td></td>
</tr>
<tr>
<td>USH 1 23/m/48</td>
<td>0.1</td>
<td>1.65</td>
<td>15</td>
<td>17</td>
<td>11/9</td>
<td>3.8/5.7</td>
<td>21.7/23.3</td>
<td></td>
</tr>
<tr>
<td>USH 1 24/m/35</td>
<td>0.26</td>
<td>1.3</td>
<td>20</td>
<td>36</td>
<td>No fix</td>
<td>NA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>USH 2 25/m/45</td>
<td>0.16</td>
<td>1.5</td>
<td>25</td>
<td>46</td>
<td>10/14</td>
<td>7.5/4.9</td>
<td>26.7/24.2</td>
<td></td>
</tr>
<tr>
<td>USH 2 26/t/41</td>
<td>0.1</td>
<td>1.55</td>
<td>10*</td>
<td>18</td>
<td>6/5</td>
<td>4.6/3.3</td>
<td>26.6/24.1</td>
<td></td>
</tr>
<tr>
<td>USH 2 27/t/43</td>
<td>0.1</td>
<td>1.55</td>
<td>20</td>
<td>54</td>
<td>6/10</td>
<td>6.7/5.7</td>
<td>301/24.1</td>
<td></td>
</tr>
<tr>
<td>USH 2 28/t/47</td>
<td>0.3</td>
<td>1.45</td>
<td>35</td>
<td>83</td>
<td>16/noise</td>
<td>8.2/NA</td>
<td>25.8/NA</td>
<td></td>
</tr>
<tr>
<td>USH 3 29/m/52</td>
<td>0.22</td>
<td>1.15</td>
<td>10</td>
<td>6</td>
<td>No fix</td>
<td>NA</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Counts refer to an area with measurable HVF threshold/mfERG responses. RD, response density; IT, implicit time; ADRP, autosomal-dominant RP; ARRP, autosomal-recessive RP; sRP, simplex RP; USH, Usher syndrome; CHM, choroideremia; NT, not tested; UR, unreliable; no fix, not able to see fixation target or test stimulus; NA, not applicable; VA, visual acuity; CS, contrast sensitivity; GVF, Goldman visual field; mfERG, multifocal electroretinogram (tested at 200 and 400 cd · m⁻²); HFV, Humphrey visual field.

† Implicit time delay >2 SD.

Response Analysis

Two iterations of artifact rejection, but no spatial averaging was applied to the raw mfERG data. mfERGs were analyzed for noise to exclude poor-quality recording as follows. The root mean square (RMS) of a higher-order kernel without a signal correlation was measured as a total noise estimate and compared with an already established control dataset \((n = 150)\). The control noise estimate approximately follows a normal distribution, as shown in Figure 2. mfERG data with a noise estimate larger than the 95th percentile of the control dataset were excluded from further analysis.

Two different analyses, the signal-to-noise ratio (SNR) analysis and the stretch-fit analysis described by Hood and Li,31 were applied to identify the more powerful method able to extract the expected small responses. (1) SNR analysis: First-order kernel responses were extracted from the epoch lengths 0 to 60 ms (signal-S), 60 to 120 ms (noise1), and 120 to 180 ms (noise2). (The late-epoch window of the first-order kernel response was a more stringent noise estimator than the fourth-order kernel, when applied to the SNR analysis in our patient cohort.) Waveform analysis was performed with custom software written in R.32 Responses with a low SNR may be caused either by high noise or low signal. It is very likely that RP-associated cone-mediated dysfunction will cause a low signal. Therefore, the noise ratio noise1/noise2 was calculated as an indicator for noise distribution for each individual recording. The SNR for each

| TABLE 2. Results of mfERG and HVF Performance and Response Count |

<table>
<thead>
<tr>
<th>Test</th>
<th>Patients n (%)</th>
<th>Patient Age (y)</th>
<th>Identified Response Number per Patient (max 103)</th>
</tr>
</thead>
<tbody>
<tr>
<td>mfERG successfully performed (200 cd · m⁻² and 400 cd · m⁻²)</td>
<td>24/29 (83%)</td>
<td>18–57 (mean 42)</td>
<td>5–103 (mean 41)</td>
</tr>
<tr>
<td>mfERGs within noise limit</td>
<td>19/24 (79%)</td>
<td>14–68 (mean 37)</td>
<td>3–21 (mean 9.7)</td>
</tr>
<tr>
<td>mfERGs exceeding noise limit:</td>
<td>20/24 (83%)</td>
<td>14–68 (mean 37)</td>
<td>3–33 (mean 11.0)</td>
</tr>
<tr>
<td>mfERG: RD P1-N1 200/400 (nV/deg²)</td>
<td>5/24 (21%)</td>
<td>34–49 (mean 43)</td>
<td>NA</td>
</tr>
<tr>
<td>mfERG: IT P1 200/400 (ms)</td>
<td>4/24 (17%)</td>
<td>34–54 (mean 44)</td>
<td>NA</td>
</tr>
<tr>
<td>HVF successfully performed</td>
<td>27/29 (93%)</td>
<td>14–68 (mean 40)</td>
<td>5–103 (mean 41)</td>
</tr>
</tbody>
</table>
individual mfERG was calculated by the quotient of S/noise. Responses were deemed detectable if the SNR was greater than the mean noise ratio + 1.96 SD. (More details are described in the Appendix.) (2) Stretch-fit analysis: Response templates from age-matched control subjects were created for each hexagon. Templates were stretched in both time and amplitude to obtain the best fit with a noise window (120–180 ms epoch) from the patient’s recording to build a distribution of template fits for each patient achieved randomly. Templates were then refitted to the signal components of each patient’s recording (0–60 ms epoch). Responses were considered detectable if hexagons achieved a fit greater than the mean noise fit + 1.96 SD.

Responses of each recording were counted, and response density P1-N1 and implicit time P1 were compared with age-matched control subjects (n = 26; ages, 13–51 years).

HVF tests with a fixation loss of less than 20% and false-positive or false-negative responses less than 33% were reliable and included in the analysis. HVF responses were counted for each patient. The number of detectable mfERG and HVF responses was compared, to identify which was the more sensitive test procedure in this patient cohort.

Clinical test results such as VA, CS, Goldmann visual field, age at testing or RP type were analyzed and correlated with the number of detectable hexagons using a one-way ANOVA for categorical variables and Spearman test.

RESULTS

General Findings

Twenty-nine patients (11 female and 18 male) aged 16 to 68 years (mean, 40) were included in the study. Diagnoses included RP (n = 21), choroideremia (n = 1), or Usher syndrome (n = 7). Heritance in patients with RP was autosomal dominant (3/29), autosomal recessive (2/29), or X-linked (3/29) or was categorized as simplex RP (13/29). Simplex RP refers to cases where only one member of the family (the patient tested) is affected and no other inheritance clues, such as mutations or carrier stage have been found in other family members.

Visual acuity ranged from 0.05 to 1.0 logMAR (mean, 0.3). Contrast sensitivity was reduced in 23 of 29 patients with a range of 1.65 to 0.15 log units (mean, 1.15). Kinetic visual fields were constricted from 10° to 60° in diameter (mean, 23°). Thirteen of 29 patients had been tested with the HVF or with a different static perimetry test at one time during the disease course, between 1.15 and 13 years before this study (mean, 3.6 years). Demographic data and vision function results are summarized in Table 1.

Multifocal ERG

Figure 1 shows a typical example of a patient recording compared with that of a control subject. Eighty-three percent of patients were able to perform the test. Five patients could not see the fixation target or even the test stimulus itself. Age, kinetic visual field size, VA, and CS were not different in those 5 patients compared with the other 24 patients.

Data from five recordings at 200 cd·m⁻² and four recordings at 400 cd·m⁻² in a total of seven patients were excluded from further analysis because of a large total noise estimate. Data from two of the seven patients exceeded the noise level for both intensities. The other five patients had a noise-acceptable mfERG for one of the stimulus intensities. The resultant 39 recordings from 22 of 29 patients (19 recordings at 200 cd·m⁻² and 20 at 400 cd·m⁻²), were analyzed for responses in each of the 103 stimulated areas.

Analyses Comparison

The SNR analysis and the stretch-fit analysis method were not significantly different in response identification (200 cd·m⁻²: P = 0.39; 400 cd·m⁻²: P = 0.27, paired t-test). The localization of identified mfERG responses identified by the two analysis methods did not correlate significantly (P = 0.25, McNe mar’s test). Averaged waveforms from identified responses were generated for each patient by each method. These waveforms were compared with template waveforms from the same hexagons in age-matched control subjects using a measure of correlation suggested by Woody. The SNR analysis showed a significantly better correlation with age-matched control tem-

![Image](http://iovs.arvojournals.org/pdfaccess.ashx?url=/data/journals/iovs/932941/)
TABLE 3. Correlation of mfERG Counts

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Correlation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>0.15</td>
</tr>
<tr>
<td>Diagnosis*</td>
<td>0.26</td>
</tr>
<tr>
<td>Visual acuity</td>
<td>-0.12</td>
</tr>
<tr>
<td>Contrast sensitivity</td>
<td>0.26</td>
</tr>
<tr>
<td>Goldmann visual field</td>
<td>0.50†</td>
</tr>
</tbody>
</table>

Data are Spearman r. Counts refer to an area with measurable mfERG responses.

* F statistic.

† P = 0.03, but considered as not significant, because applied Bonferroni adjustment.

plates (P = 0.04, paired t-test) than did the stretch-fit analysis in this patient population. We decided, therefore, that the SNR analysis was more suitable for this patient population and we applied it to each of the single mfERGs.

Response Identification

Responses were identified in 9.8% (395/4017) of the stimulated areas. There was no statistically significant sensitivity difference between the two stimulus intensities (P > 0.05, Wilcoxon rank sum test): 184 of 1957 signals (103 stimulated areas × 19 recordings) and 211 of 2060 signals (103 stimulated areas × 20 recordings) were identified as detectable responses for the 200 cd·m⁻² and 400 cd·m⁻² stimulus condition, respectively (Table 2).

Typical examples of the response analysis are shown in Figure 3. Identified mfERG responses are encircled in each of the trace plots. The number of identified responses range from 7 to 13 of the 103 areas stimulated in these four examples. The response average P1 implicit time was within normal limits in patients 1, 6, and 23 and delayed by 3.7 SD in patient 11. The response average P1 implicit time was within normal limits in 39% and 59% for the 200-cd·m⁻² and 400 cd·m⁻² stimulus condition, respectively. Nine responses were delayed >5 SD.

Central Visual Field Comparison with the mfERG

The test performance of the HVF was better than the mfERG. All but two patients were tested successfully with the HVF. Results from those two patients were excluded from the analysis because of the large number of false-positive and -negative responses, which were due to obvious reduced compliance during the test. Both of them had an mfERG recorded successfully. Visual field responses were detected in 40% of the tested areas (5–103; mean 41 areas) in 27 patients tested successfully.

Data from patients with both mfERG and HVF results were compared. The HVF test was significantly more sensitive than the mfERG in detecting localized responses (P < 0.001, paired t-test).

HVF sensitivity was significantly higher in areas with detectable mfERG responses than in areas without them (P = 0.003, 200 cd·m⁻²; P = 0.04, 400 cd·m⁻²; Welch two-sample t-test) as illustrated in Figure 5. A correlation between the mfERG and the HVF test was found in areas with measurable responses (P < 0.001; McNemar test). Therefore, mfERG responses were most likely found in areas with identified HVF responses.

DISCUSSION

Sensitive and reliable outcome measures are necessary to characterize the natural disease history and are fundamental in monitoring disease outcome. Electrophysiological tests, such as the scotopic and photopic full-field ERG, are objective and reliable measures to quantify overall retinal function. In patients with advanced RP, photoreceptor and inner retinal layer function may be too small to be measurable by the full-field ERG (ISCEV standard). We expected that the mfERG would be sensitive enough to quantify the cone-mediated function in most of the patients who had advanced RP and preserved central vision. Hence, we postulated that the mfERG

![Figure 4](http://iovs.arvojournals.org/pdfaccess.ashx?url=/data/journals/iovs/932941/ on 12/23/2017)
could be used as an outcome measure in further studies involving those patients with advanced retinal disease.

Previous studies assessing the applicability of the mfERG technique in patients with RP used different recording and analysis protocols, applied different inclusion criteria, and tested different RP subtypes than the present study.18–24 Signal-to-noise differentiation is crucial in the determination of detectable small-amplitude responses, which may not always be applied in the previous studies. Hood et al.19 successfully tested eight patients with RP, who had larger visual fields than those recorded in patients in the present study, and all but one had a recordable ffERG. They concluded that the total mfERG response depends on the relative size of the peripheral mfERG response. Seeliger et al.18 concluded that mfERG responses are nonrecordable in patients with RP with a Goldmann visual field of <10° and a photopic full-field 30-Hz flicker response of 1 μV or less. The present study shows that at least one response is recordable in all successfully tested patients using an objective signal-to-noise analysis paradigm, irrespective of the kinetic visual field size. We were able to identify responses even in patients with restricted kinetic visual fields smaller than 10° in radius. Severely diminished responses were associated with normal temporal characteristics in nearly 50% of the responses, which agrees with the findings of Hood et al.19 What is somewhat surprising is the finding that some patients lacked a central mfERG response despite a visual acuity of 0.4 logMAR or better and a normal HVF foveal threshold. Seiple et al.34 showed a similar example of a patient with RP, VA of 20/25, and preserved mfERG responses in the peripheral area without a central response. In such cases, the number of intact photoreceptors may be sufficient to resolve a small visual angle required for good VA. The mfERG, on the one hand, may not be sensitive enough to detect signals initiated by the same amount of photoreceptors. Static perimetry, on the other, is a threshold test with a higher sensitivity range due to cortical amplification.

The mfERG may not be recommended as a primary outcome measure in patients with advanced RP and nonrecordable ffERG. Three critical observations were made in this specific patient cohort: (1) Response identification was positive in only 10% of all areas tested; (2) 17% of the patients were not able to perform the mfERG, despite high motivation and cooperation. These patients did not differ in age, visual acuity, contrast sensitivity or visual field size from patients who were able to perform the test. A conventional screen presentation may facilitate localizing the stimulus by the patient, but at the expense of monitoring fundus fixation. This compromise, we believed, would not be acceptable; (3) test compliance and success rate were higher with static perimetry than with mfERG test in this specific patient cohort. Signal exceeded the expected noise level in 15% of the mfERG recordings, which may be caused by undetected fixation instability during recording. The static visual field test, in contrast, was more sensitive in detecting central retinal function in the same patients. There may be a bias due to previous experience with the test procedure in less than 50% of our patients. The long period since the previous static visual field in respect to changed visual perception due to disease progression makes it unlikely that this preexistent experience interferes with the results in our study. The heterogeneous and not yet identified genotype in our patient cohort may be a factor for response detection. However, those patients are all tested at an advanced stage of disease with the same inclusion criteria.

Small-amplitude mfERG responses are detectable in all successfully tested patients with advanced RP, using the described analysis method. However, static perimetry provides a better functional outcome measure of central visual function when compared with the mfERG in this advanced stage of disease. Both tests remain useful as a baseline measure in patients with advanced RP nonrecordable ffERGs. If the carefully performed mfERG analysis, including the signal-to-noise analysis fails to detect responses in the tested area, the central static perimetry would be a measure suitable for longitudinal studies of residual function using the patient’s individual reliability.34 The use of both tests in patients with recordable mfERG responses may be beneficial in assessing longitudinal dynamics of retinal dysfunction with disease progression or in quantifying outcome.

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References


**APPENDIX**

**Estimation of Recording Quality**

Processed data were exported from unfiltered trace plots of a high-order kernel (4-20; epoch 0–60 ms) from VERIS. The RMS (power estimate) was calculated as follows:

\[
\text{Noise} = \sqrt{\frac{1}{n} \sum_{b=1}^{b=105} x_i^2}
\]

where \(b\) is the hexagon number and \(i\) is the sample.

Waveforms were rejected when: noise > mean noise \(_{\text{controls}}\) + 2 SD noise \(_{\text{controls}}\) (i.e., when estimated noise exceeded the 95th percentile of our control dataset).

**Identification of Hexagons with Signals**

Processed data were exported from unfiltered trace plots of a first-order kernel (epoch 0–180 ms) from VERIS. For each hexagon, RMS \(_{\text{signal}}\) (epoch 0–60 ms) and RMS \(_{\text{noise}}\) (120–180 ms) were calculated.

\[
\text{RMS}_b = \sqrt{\frac{1}{n} \sum_{i=1}^{n} x_i^2}
\]

\[
\text{SNR}_b = \frac{\text{RMS}_{\text{signal}}}{\text{RMS}_{\text{noise}}}
\]

Hexagons were identified to contain a signal when

\[
\text{SNR}_{b=105} + 2 \cdot \sigma_{\text{SNR}_{b=105}} \leq \text{SNR}_b
\]