Objective perimetry using the multifocal visual evoked potential in central visual pathway lesions

A I Klistorner, S L Graham, J Grigg, C Balachandran

Aims: To examine the ability of the multifocal pattern visual evoked potential (mVEP) to detect field loss in neurological lesions affecting the visual pathway from the chiasm to the cortex.

Method: The mVEPs recorded in the clinic were retrospectively reviewed for any cases involving central neurological lesions. Recordings had been performed with the AccuMap V1.3 objective perimeter, which used an array of four bipolar occipital electrodes to provide four differently oriented channels for simultaneous recording. 19 patients with hemianopias were identified. Of these there were 10 homonymous hemianopias with hemifield type loss, two bitemporal hemianopias, and seven homonymous hemianopias with quadrantanopic distribution. A comparison with subjective field results and CT/MRI findings was done to determine the relation between the two methods of visual field mapping and any relation with the anatomical location of the lesion and the mVEP results.

Results: In all hemianopic type cases (12) the defect was demonstrated on the mVEP and showed good correspondence in location of the scotoma (nine homonymous and two bitemporal). The topographic distribution was similar but not identical to subjective testing. Of the seven quadrantanopic type hemianopias, only four were found to have corresponding mVEP losses in the same areas. In the three cases where the mVEP was normal, the type of quadrantanopia had features consistent with an extra-striate lesion being very congruous, complete, and respecting the horizontal meridian.

Conclusions: The mVEP can detect field loss from cortical lesions, but not in some cases of homonymous quadrantanopia, where the lesion may have been in the extra-striate cortex. This supports the concept that the mVEP is generated in V1 striate cortex and that it may be able to distinguish striate from extra-striate lesions. It implies caution should be used when interpreting “functional” loss using the mVEP if the visual field pattern is quadratic.

The visual evoked potential (VEP) has traditionally been an important method for assessing the state of visual pathways. It has been most useful in the diagnosis of conditions such as optic neuritis and other optic neuropathies. There have been several reports of VEP abnormalities in cortical lesions, which demonstrated reductions in amplitudes when the affected part of the visual field was stimulated. Sequential quadrantic or hemifield stimulation was used in these studies. 1-4

The clinical findings from these studies were limited by the fact that the conventional pattern VEP is predominantly generated by cortical elements receiving projections from the central retina and it has been estimated that the central 2° of visual field contributes 65% of the response. 7-9 Therefore, the conventional VEP had limited ability to reflect field loss in non-central areas. Recent developments in multifocal stimulation techniques have provided a new method for assessing visual function, and expanded the scope of the VEP to enable assessment of multiple sites in the field out to 30° eccentricity. 10-11 Results published using multifocal techniques have demonstrated a good correspondence between visual field thresholds and the amplitude of the local multifocal VEP (mVEP) response. Scotomas can be identified, and field defects in glaucomatous optic neuropathy and optic neuritis have been well defined. 12-17

There are a few reports of mVEPs in cortical lesions, which have suggested that hemianopic and quadrant defects can be identified and topographically represented with the mVEP responses. 18 19 This has led to the notion that normal mVEP response is obtained from stimulation of the “blind” area of the field that the visual loss is “functional” rather than pathological. 19 If true then this would have implications for the investigation of suspected malingerers and cases of medicolegal compensation.

We have investigated a sample of confirmed visual pathway or cortical lesions with varying types of field defects to determine if the mVEP does represent the subjective losses seen on perimetric testing. Based on previous conventional VEP studies, which were thought to be a good differentiating tool for functional versus organic field loss, and limited mVEP research, it had been anticipated that all types of lesions would be detected. However, since the mVEP is thought to be derived from the striate V1 cortex 20-22 (although exact origins are yet to be confirmed), the possibility that extra-striate lesions might remain undetected still remains.

Methods

Subjects

A retrospective review was performed of all subjects tested in the clinic with mVEP. Nineteen subjects were identified who had various forms of homonymous or bitemporal field loss from lesions of the posterior visual pathway (pituitary to visual cortex). All subjects had subjective visual fields (Humphrey visual field Sita standard 24-2), and 17/19 had some form of neuroimaging (computed tomography (CT) or magnetic resonance imaging (MRI)) available. Three of the subjects had been tested twice on the mVEP.

Abbreviations: CVA, cerebrovascular accident; FDT, frequency doubled threshold; FIR, finite impulse response; HVF, Humphrey visual field; mVEP, multifocal pattern visual evoked potential; VEP, visual evoked potential
Informed consent was obtained from all subjects before mVEP testing. The study had ethics approval and followed the principles of the Declaration of Helsinki.

Stimulation and recording

The AccuMap system (ObjectiVision, Sydney, Australia) has been described in detail in previous publications. Briefly, a checkerboard pattern undergoes pseudo-random binary exchange at each of 56 sites in the visual field. Each input (stimulation site) is modulated in time according to a different sequence. The technique permits computation of the resulting signal by cross correlation of the response evoked by the sequence stimulation, with the sequence itself. Further runs then use different sequences for the same stimulation site to reduce cross contamination.

The visual stimulus is generated on a computer screen (22" Hitachi high resolution display, video frame rate 75 Hz). Fifty six close packed segments in a dartboard configuration are used. The segments are cortically scaled with eccentricity to stimulate approximately equal areas of cortical (striate) surface (see fig 1). The scaling therefore produces a signal of similar amplitude from each stimulated segment. Each segment contains a checkerboard pattern (16 checks), with the size of individual checks being proportional to the size of the segment and therefore also dependent on eccentricity. Luminance of the white check is 146 cd/m² and luminance of the black check 1.1 cd/m², producing Michelson contrast of 99%. Background luminance of the screen is maintained at a mean level of 73.5 cd/m². A dim room light is always on.

Subjects are seated comfortably in a chair and asked to fixate on the fixation number at the centre of the stimulus pattern. The distance to the screen is 30 cm, corresponding to a total subtense of the stimulus of 52° with an additional nasal step. All subjects are optimally refracted for near and the pupils are not dilated. All recordings are collected using monocular stimulation. The signal is amplified 100 000 times and band pass filtered between 1 Hz and 30 Hz, with later digital filtering down to 20 Hz, using finite impulse response (FIR) band pass filter. The data sampling rate is 502 Hz. Raw data are scanned in real time and runs contaminated by a ratio.

In all those cases there was clear topographic correspondence between subjective visual field and mVEP. However, the distribution of the field defect did not always show as clear a vertical demarcation as the subjective test, and in some cases did not fully extend to the periphery. Despite this, it was always identifiable as a hemifield defect, and was not obscured by surrounding noise. When average quadratic amplitude of mVEP from subjectively normal quadrants was compared with amplitude derived from quadrants which were affected on Humphrey visual field (HVF) (at least 75% of test points p < 0.05) the difference was highly statistically significant (p<0.0001, Student’s t test) (fig 2).

Examples showing subjective fields and corresponding mVEP are presented in figures 3–5, with one hemianopic type and two quadratic type hemianopias. The hemianopic defect is well defined on both subjective and objective tests (fig 3). The quadratic field defect in figure 4 just crossed the horizontal midline in the left eye making it incongruous, although in the mVEP it respects the horizontal. Figure 5 shows an incomplete left homonymous quadratic type defect, with an additional small defect in the superonasal field of the left eye which was caused by an old retinal lesion.

However, in three cases (all congruous quadratic type) the amplitude of mVEP was normal across the whole field despite clear abnormality seen on HVF (see example in fig 6). On examining quadratic mVEP amplitudes there was no statistical difference (p = 0.46, Student’s t test) between the mVEP amplitudes of the affected and unaffected quadrants.

Therefore, in all of the hemianopic type cases (12/12) but only 4/7 quadratic type lesions, corresponding mVEP losses were found in the same areas of the visual field. The topography of the field loss in the corresponding 15 cases showed very similar distributions, as seen in the figures 3–5.

![Figure 1](http://bjo.bmj.com/)  
**Figure 1** Stimulus pattern for mVEP. Note central check sizes and stimulus areas are much smaller than peripheral (cortical scaling of stimulus). External ring expands up to 26° of eccentricity with an additional nasal step up to 32°.
However, the vertical midline demarcation was not as clear in the mVEP as it was on subjective testing, with some abnormal points seen in several cases adjacent to the midline in the normal hemifield.

To determine if artefactual signals could be generated in a quadrant with no signal input we occluded a quadrant of the screen and recorded the mVEP in a normal subject (fig 7). This clearly showed loss of signal in that quadrant, so we do not believe the normal signals in the three cases with quadrantopic type defects with normal mVEPs are generated by cross contamination or artefact. It is important to realise that occluding a segment of the screen does not simulate a real scotoma, it merely confirms that the software is not generating artificial signals by cross contamination.

Table 1: Summary of clinical, mVEP, and CT/MRI findings of all patients

<table>
<thead>
<tr>
<th>Patient No</th>
<th>Age</th>
<th>Sex</th>
<th>Diagnosis</th>
<th>CT or MRI confirmed</th>
<th>Field defect</th>
<th>MVEP defect</th>
<th>Repeat MVEP defect</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>56</td>
<td>M</td>
<td>L occipital infarct</td>
<td>Yes</td>
<td>R hemianopia incomplete</td>
<td>Yes –</td>
<td>–</td>
</tr>
<tr>
<td>2</td>
<td>80</td>
<td>F</td>
<td>L occipital infarct</td>
<td>Yes</td>
<td>R complete hemianopia</td>
<td>Yes –</td>
<td>–</td>
</tr>
<tr>
<td>3</td>
<td>53</td>
<td>M</td>
<td>L occipital infarct</td>
<td>Yes</td>
<td>R hemianopia congruous</td>
<td>Yes –</td>
<td>–</td>
</tr>
<tr>
<td>4</td>
<td>65</td>
<td>M</td>
<td>Pituitary tumour</td>
<td>Yes</td>
<td>R hemianopia, incomplete, macular sparing</td>
<td>Yes –</td>
<td>–</td>
</tr>
<tr>
<td>5</td>
<td>43</td>
<td>M</td>
<td>Embolic CVA, partial recovery</td>
<td>No lesion on CT</td>
<td>L incomplete congruous</td>
<td>Yes –</td>
<td>–</td>
</tr>
<tr>
<td>6</td>
<td>35</td>
<td>M</td>
<td>Blunt head injury</td>
<td>Not available</td>
<td>L hemianopia, complete, congruous</td>
<td>Yes –</td>
<td>Yes –</td>
</tr>
<tr>
<td>7</td>
<td>80</td>
<td>F</td>
<td>R medial occipital lobe infarct</td>
<td>Yes</td>
<td>L incomplete hemianopia, congruous</td>
<td>Yes –</td>
<td>–</td>
</tr>
<tr>
<td>8</td>
<td>29</td>
<td>F</td>
<td>Craniohypophysealgia with chiasmal compression</td>
<td>Yes</td>
<td>Bitemporal hemianopia</td>
<td>Yes –</td>
<td>–</td>
</tr>
<tr>
<td>9</td>
<td>74</td>
<td>M</td>
<td>L occipital infarct</td>
<td>Yes</td>
<td>R hemianopia incomplete</td>
<td>Yes –</td>
<td>–</td>
</tr>
<tr>
<td>10</td>
<td>57</td>
<td>M</td>
<td>Massive R middle cerebral CVA/ craniotomy decompression</td>
<td>Yes</td>
<td>L hemianopia, R complete</td>
<td>field loss and optic atrophy</td>
<td>–</td>
</tr>
</tbody>
</table>

R: right; L: left; S: superior; I: inferior; CVA: cerebrovascular accident; CT: computed tomography scan; MRI: magnetic resonance imaging.
Repeat mVEP defect shows result if second test was done (3 cases only).

DISCUSSION

The use of multichannel multifocal VEP recording provides a new means of assessing the integrity of the visual pathway through to the visual cortex. There have been many previous studies on the ability of the conventional VEP to detect field loss and hemianopias. While it can be achieved, the topographical relation is difficult to define without the new dimension that multifocal techniques provide. Thus, two recent reports by Ringger (ARVO abstract no 248, 2004) and Wall indicated that hemianopic lesions can be detected by mVEP, but that subcortical lesions are more readily identified than cortical.

In this study all of the 12 hemianopic cases with hemifield type defects were found to have corresponding mVEP losses in the same areas. These hemianopic cases showed definite occipital lesions in eight cases, two more were from chiasmal compression, one from clipping of a cerebral aneurysm more than 15 years earlier and one from old head trauma from a motor vehicle accident. In all these cases the hemianopia was demonstrated on the mVEP, even if the topographic correspondence was not exact.

However, only four out of seven hemianopias with quadrantopic type defects showed corresponding defects on mVEP. In three quadrantopia cases the mVEP was normal across the whole field. A more detailed examination of the HVF in these cases was undertaken, and revealed a noticeable difference between the two groups. While in the four cases where the mVEP was affected the subjective scotoma did not quite reach horizontal meridian (three cases) or was slightly over it (one case), in the remaining three cases...

![Figure 2](image.jpg)

**Figure 2** Average quadranotopic amplitude of mVEP from normal quadrants on Humphrey visual field (HVF) Sita 24-2 compared with mVEP amplitude derived from quadrants which were affected on HVF (at least 75% of test points p<0.05). Data from the 10 cases where mVEP was abnormal (p<0.0001).
with a normal mVEP, the HVF defect respected horizontal meridian.

The mVEP is thought to be mostly derived from the striate cortex (area V1).14–22 Thus, recent studies based on principal component analysis have confirmed that the electrical dipole that generates the mVEP response follows the shape of V1 folding.22 We therefore would expect that any lesions before or including V1 would lead to reductions in signal. Alternatively, the presence of a mVEP should indicate that there is no abnormality present at least up to the level of neurons of the striate cortex. However, the possibility remains that focal cortical lesions involving extra-striate areas (which are also retinotopically organised) could produce subjective visual field loss but retain a normal amplitude mVEP signal.

While there is no information available about the topography of extra-striate V2/V3 areas in humans, animal experiments indicate that V2 surrounds V1 with projections of upper and lower hemifields being separated by the striate cortex, while V3 is wrapped around V2, being therefore with its upper and lower hemifields separated even further.24–26 Based on anatomical topography it has been suggested, and clinically confirmed in a few cases, that lesions on the border between V2/V3 or V1/V2 will manifest quadrantanopic type scotomas, which would exactly respect the horizontal meridian.27–30 Furthermore, it was proposed that for localised damage in V1 to produce a scotoma strictly respecting the horizontal meridian, the lesion must transect the striate cortex exactly along the base of the calcarine fissure all the way through to incorporate central and peripheral parts of the field.30 This is very unlikely for both vascular and neoplastic type lesions. The idea that lesions in extra-striate areas can produce dense perimetric scotomas has been questioned by Merigan and co-workers, however.31 Based on animal studies using chemical deactivation of neurons in area V2, they suggested that a quadrantic type scotoma is

Figure 3  Subject with hemianopia from head trauma (patient 6). Shows HVF grey scale and pattern deviation plots (Sita 24-2) for right and left eyes (A) and corresponding mVEP trace arrays and probability plots (B) from AccuMap. There is good correspondence between HVF and mVEP defects.

Figure 4  Subject with right superior quadrantic field defect (subject 13). The defect just crosses the horizontal meridian in the left eye making it not completely congruous.
more likely to be caused by damage to fibre tracts leading from V1 to extra-striate areas, rather than neuronal damage in extra-striate areas themselves. While this matter still remains controversial, it is accepted that a quadrantanopic type scotoma, which strictly respects the horizontal meridian can be caused by damage beyond the neurons of V1.

As was indicated, in the three cases of quadrantanopic type scotoma, which were not manifested on mVEP the defects strictly respected the horizontal as well as the vertical midline. This was consistent with a lesion beyond the striate cortex. The CT or MRI showed small localised occipital pole infarcts in two cases, and no detectable lesion in the third. Standard neuroimaging is not accurate enough to confirm an exclusively extra-striate lesion, but the small occipital infarcts were not inconsistent. A recent study using functional MRI has demonstrated in a human with a congruous quadrantic defect the loss of MRI signal in the corresponding extra-striate area, but preservation of the MRI signal in V1. In the four other cases of quadrantanopic type hemianopia where the mVEP was reduced, the field defects were more characteristic of lesions at or before the striate cortex. They had less congruous defects that did not respect the horizontal meridian. In these cases the mVEP was abnormal in the quadrants involved.

The observation that in many cases the mVEP was not as clearly respecting the vertical midline is important in terms of applying the mVEP technique to investigating neurological lesions. This may be because the stimulus extends right up to the midline in the mVEP, whereas in the Humphrey visual field the test points are 3° either side of the vertical. Shifts of fixation therefore can have a greater impact. A similar problem was reported in frequency doubled threshold (FDT)
perimetry\(^{11}\) and has been adjusted for in the latest version of the FDT system.

A further reason for lack of exact correspondence between mVEP and subjective fields is that in some areas the background noise (including alpha rhythm) is being incorrectly recognised as a true signal, so the probability plot is not flagging the area as abnormal. Improvements in signal recognition will help reduce this problem.

In conclusion, in the investigation of optic pathway or cortical lesions the mVEP can be used to confirm hemianopic defects, in lesions up to and involving the V1 striate cortex neurons. We propose that the three cases with quadrantanopic type defects with preserved mVEP (which showed somatoplasia respecting horizontal meridian) may have been lesions in extra-striate areas or the tracts leading from V1, as they had field defects typical for this. The findings are consistent with the notion that most of the mVEP signal appears to derive from V1. It also means that preservation of amplitude implies an intact pathway to V1, which helps localise the lesion. However, a quadrantic subjective field loss respecting horizontal meridian should not be judged as “functional” or maligning just because the mVEP is preserved, since it could be of extra-striate origin.

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**Authors’ affiliations**

A I Klistorner, S L Graham, J Grigg, C Balachandran, Save Sight Institute, Department of Ophthalmology, Sydney University, Sydney, Australia

Proprietary interest: AIL and SLG consultants for ObjectVision.

AIL is Sydney Medical Foundation research fellow.

**REFERENCES**


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