Visual evoked potential latency and contrast sensitivity in patients with posterior chamber intraocular lens implants

J W Howe,† K W Mitchell,‡ M Mahabaleswara, and M N Abdel-Khalek

From the †University Department of Ophthalmology and the ‡Regional Department of Medical Physics, Royal Victoria Infirmary, Newcastle upon Tyne NE1 4LP

SUMMARY An electrophysiological investigation of visual evoked potential (VEP) latency and contrast sensitivity was performed in a group of 13 patients who had undergone extracapsular cataract surgery with posterior chamber lens implantation. In spite of good postoperative visual acuity, abnormalities were detected in nine of the group (69%). This study suggests that, even with successfully implanted lenses, there may be a reduction in visual function which could be the result of altered transmission through the plastic lenticulus or fibrosis of the posterior lens capsule, and/or subtle changes in retinal architecture, not observed ophthalmoscopically.

The optical disadvantages of spectacle lenses in the correction of aphakia make intraocular lenses (IOLs) an attractive alternative. Although intraoperative and postoperative complications have been widely reported, improvements in both lens design and surgical technique have reduced these problems, so that lens implantation is increasingly widely practised. In the USA alone it has been estimated that the implantation rate exceeds 500,000 annually,1 and that posterior chamber lenses account for nearly two-thirds of these, the remainder largely comprising anterior chamber implants. There is also an increasing trend towards using IOLs in the UK, a recent survey suggesting that 45% of cataract operations involved the use of lens implants.2

The success of an implanted lens can be judged on the basis of low complication rate and by assessment of eventual visual acuity. On the basis of an acuity of 6/12 or better a success rate of over 90% has been reported with posterior chamber implants3,4 and 87% in patients who have undergone intracapsular lens extraction with insertion of an anterior chamber implant.5 Visual acuity may, however, be normal in the presence of a variety of disorders in which overall contrast sensitivity is reduced.6 This is the type of visual impairment in which some patients report ‘fogginess’ or ‘haziness’ of vision despite achieving normal standards of visual acuity. In a recent report Hess et al.7 have detected abnormalities in contrast sensitivity, measured psychophysically, in patients with iris clip lenses inserted after intracapsular cataract extraction. As an alternative to psychophysical means, contrast sensitivity function can be estimated electrophysiologically by recording visual evoked potentials (VEPs).8 These are known to be influenced not only by retinal and optic nerve pathology but also by factors affecting the quality of the visual image on the retina, such as opacities of the media, refraction, and pupil size (for review, see Halliday9). On theoretical grounds alone it would appear that abnormalities in the VEP are likely to be seen in patients with IOLs.

We believe there have been no reports so far on electrophysiological changes in patients who have undergone intraocular lens implantation. We report here the results of a bimodal approach which has determined VEP latency and contrast sensitivity in 13 patients with good visual acuity following extracapsular surgery and posterior chamber IOL insertion.

Patients and methods

Patients Thirteen patients (age 57–70 years, mean 65·4 years) who had undergone successful extracapsular
Cataract extraction with insertion of a posterior chamber IOL were examined (Table 1). Each had a corrected visual acuity of 6/12 or better in the pseudophakic eye, with no clinical evidence of retinal or optic nerve pathology and a normal intraocular pressure (21 mmHg or less). In 10 of these patients there was a variable degree of cataract in the contralateral eye, but in the remaining three the cataract had been unilateral and the visual acuity in the fellow eye 6/6 or better. Results from these patients were compared with a normal, age matched population from our database (mean age 62.6, SD 9.3, n=55).

**Electrophysiology**

A checkerboard stimulus was produced by a video pattern generator on a high quality TV monitor. In the first part of the test luminance modulation of the pattern was selected to give the pattern reversal mode of stimulation. The checksize of the stimulus was 50' and the visual field subtended, 17°x 14°. The overall luminance of the TV screen—both with and without pattern—was maintained constant at 10 Cd m⁻². Pattern contrast (defined as \( \frac{L_{\text{max}} - L_{\text{min}}}{L_{\text{max}} + L_{\text{min}}} \) where \( L_{\text{max}} \) and \( L_{\text{min}} \) were the luminance of the bright and dark checks respectively) was adjusted to be 95% with a reversal rate of 2 per second.

The second part of the test, to measure contrast sensitivity, was performed with the onset-offset mode of stimulation. The rationale for the selection of this type of pattern modulation is fully discussed elsewhere. The pattern was present for 40 ms every 500 ms (note: the presentation rate—as with pattern reversal mode—was locked to the 50 Hz monitor frame rate). Using a check size of 19' for patients with

![Figure 1](http://bjo.bmj.com/)

**Fig. 1** VEPs from patient No. 7 (LE pseudophakic RE normal). Pattern reversal responses are shown in column (a) and onset-offset responses at different contrast levels in column (b).
a VA of 6/9 or better and 25' for those with an acuity of 6/12, we made recordings at (usually) five contrast levels between 5% and 80%. These were selected in a pseudorandom sequence to try to eliminate any possible order effects. The patients were given short periods of rest between each contrast measurement so as to minimise fatigue and any concomitant increase in response variability.

Silver/silver chloride disc electrodes were attached to the scalp with collodion, in the following positions: active—Oz; reference—Cz; earth—Pz.

A Medelec electrophysiological recording unit was used to amplify, average, and store the evoked potentials. The amplifier bandwidth was 0-8–80 Hz, and either 64 or 128 epochs of 300 ms duration were averaged depending on the size of the response. Two averages were obtained at each contrast level to check for consistency, and quantitative analysis was performed on the average of these two. A peak-peak amplitude measure was adopted: in the case of the pattern reversal VEPs it was that between P100 and N150, in those to onset-offset it was between components C1 and CII. A permanent record of the responses was made on an X-Y plotter. Amplitude data from the traces were then fed into a MINC PDP-11 computer for storage and further analysis. Monocular stimulation was adopted in all investigations, the subject being instructed to maintain fixation and focus on a small LED marker attached to the centre of the screen. This was checked during the investigation by closed circuit TV monitoring. Before the start of the test the subject was preadapted to the luminance of the blank screen for 5 minutes. This was the only source of illumination in an otherwise darkened room. The test was concluded with the measurement of the subject’s pupils under experimental conditions.

Results

In Fig. 1 the responses from a patient with a left intraocular implant and visual acuities of 6/6 in each eye are illustrated. The pattern reversal responses indicated a delay in the main P100 component in the pseudophakic eye as compared with the RE. The onset-offset responses clearly showed reduced amplitude at all contrast levels in the LE. When the amplitude of these responses was plotted against log contrast (Fig. 2), significant differences were made between the two eyes were observed, not only in amplitude but also in the electrophysiologically estimated contrast threshold (arrowed).

In Table 1 data concerning all 13 patients in the study are presented. P100 latency was significantly increased (at the 1% confidence level) in four of the group, and increased contrast threshold in eight. It is

![Fig. 2. Amplitudes of the onset-offset responses from patient No. 7. (Fig. 1) plotted against log-contrast. Closed symbols (●) = normal RE. Open symbols (○) = pseudophakic LE.](http://bjo.bmj.com/)

<table>
<thead>
<tr>
<th>Serial no.</th>
<th>Age</th>
<th>VA</th>
<th>P100 Latency (ms)</th>
<th>Contrast threshold (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>60 (M)</td>
<td>6/6</td>
<td>134-0*</td>
<td>3-3</td>
</tr>
<tr>
<td>2</td>
<td>61 (M)</td>
<td>6/9</td>
<td>115-0</td>
<td>5-0</td>
</tr>
<tr>
<td>3</td>
<td>69 (M)</td>
<td>6/6</td>
<td>122-0</td>
<td>15-0*</td>
</tr>
<tr>
<td>4</td>
<td>64 (F)</td>
<td>6/6</td>
<td>125-0</td>
<td>3-9</td>
</tr>
<tr>
<td>5</td>
<td>57 (F)</td>
<td>6/9</td>
<td>114-0</td>
<td>7-7*</td>
</tr>
<tr>
<td>6</td>
<td>70 (M)</td>
<td>6/12</td>
<td>102-0</td>
<td>10-9*</td>
</tr>
<tr>
<td>7</td>
<td>65 (F)</td>
<td>6/6</td>
<td>138-0*</td>
<td>13-8*</td>
</tr>
<tr>
<td>8</td>
<td>66 (F)</td>
<td>6/6</td>
<td>113-0</td>
<td>8-6*</td>
</tr>
<tr>
<td>9</td>
<td>70 (F)</td>
<td>6/9</td>
<td>111-0</td>
<td>4-6</td>
</tr>
<tr>
<td>10</td>
<td>61 (M)</td>
<td>6/9</td>
<td>132-0*</td>
<td>5-2*</td>
</tr>
<tr>
<td>11</td>
<td>68 (M)</td>
<td>6/6</td>
<td>110-0</td>
<td>3-9</td>
</tr>
<tr>
<td>12</td>
<td>69 (M)</td>
<td>6/12</td>
<td>129-0*</td>
<td>11-9*</td>
</tr>
<tr>
<td>13</td>
<td>70 (M)</td>
<td>6/6</td>
<td>105-0</td>
<td>6-1*</td>
</tr>
</tbody>
</table>

*Significant at 1% level.

Normal P100 latency=113-0 (SD 6-2) ms (99% CL=128-0 ms) (n=55). Normal contrast threshold=2-1 (SD 1-3)% (99% CL=5-1%) (n=49).
Table 2  Interocular comparison in those patients with pseudophakia and a normal contralateral eye

<table>
<thead>
<tr>
<th>Serial no.</th>
<th>VA</th>
<th>Diagnosis</th>
<th>P100 Latency (ms)</th>
<th>Contrast Threshold (%)</th>
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</thead>
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<tr>
<td></td>
<td>RE</td>
<td>L.E.</td>
<td>RE</td>
<td>L.E.</td>
</tr>
<tr>
<td>3</td>
<td>6/9</td>
<td>6/6</td>
<td>Normal</td>
<td>Pseudophakia</td>
</tr>
<tr>
<td>7</td>
<td>6/6</td>
<td>6/6</td>
<td>Normal</td>
<td>Pseudophakia</td>
</tr>
<tr>
<td>10</td>
<td>6/9</td>
<td>6/5</td>
<td>Pseudophakia</td>
<td>Normal</td>
</tr>
</tbody>
</table>

*Significant at 1% level.
Normal interocular latency difference Δt=2-0 (SD 1-9) ms (99% CL=6-3 ms) (n=55).
Normal interocular threshold difference Δt=0-8 (SD 0-8)% (99% CL=2-7%) (n=49).

significant in all three patients, and again higher values were observed in the operated eye.

Discussion

Before it is possible to attach pathological significance to a delayed and/or attenuated VEP or depressed contrast sensitivity (that is, increased threshold) several confounding causes must be eliminated. To this end special attention was given to the selection of patients with good central acuity and no evidence of any retinal or optic nerve disorder as judged by careful ophthalmoscopy, perimetry, and fluorescein angiography. In all patients the corneae were clear on slit-lamp examination, and no significant vitreous opacities were observed. The posterior capsule was intact in all patients. The size of the pupil can also be a source of error, as miosis reduces retinal illumination, which in turn increases VEP latency. Pupillary diameter was directly measured and was not significantly different from the normal range; it was never less than 3 mm in any of the eyes investigated, thus being excluded as a possible causal factor. At the average luminance of the stimulator the eyes were adapted to a low photopic level. The pupils were therefore partially dilated under these conditions, and this would theoretically increase the amount of distorted light due to spherical aberration, which can be an important consideration in IOL design.10 This blurring could produce a reduction in contrast, though we were unable to detect any correlation between pupil size and contrast sensitivity.

Visual acuity was measured on a conventional Snellen chart, viewed under photopic conditions (approximately 10 times the average luminance of the TV stimulus), and therefore the patient’s pupils would be more constricted, reducing the influence of peripheral distortion. From this reasoning we would speculate that, if the contrast sensitivity experiment was performed under photopic conditions (for example, 100 Cd m⁻²), the observed differences in contrast sensitivity in pseudophakia may be attenuated or even disappear, and conversely, performed at reduced luminance (for example, 1 Cd m⁻²), the difference may be enhanced.

The delayed pattern reversal VEPs in 31% of the patients and reduced contrast sensitivity in 61% were probably due to poorer light transmission through the optic media, in particular the intact posterior capsule or the implant itself. Although the posterior capsule fibrosis did not appear to be clinically significant, it could still explain some of our findings by virtue of its position on or near the nodal point of the eye, since pathological evidence frequently shows fibrosis and membrane formation.12 However, our data do not preclude the possibility that some of these changes may have resulted from subtle alterations in retinal architecture resulting from mild postoperative cystoid macular oedema, which was not ophthalmoscopically apparent.

The reason for the difference in detection rate in the two VEP measures was most likely due to differences in the stimulus. The pattern reversal stimulus was very coarse, utilising very large checks at high contrast. Experiment has confirmed that this form of pattern modulation, in conjunction with checks of subtense >20', elicits VEPs which tend to respond to the luminance of the individual checks as well as the spatial contrast between them. This luminance modulation effect would tend to overcome to some degree any reduction in spatial contrast of the retinal image produced by opacification within the media. The onset-offset mode of stimulation, however, produces VEPs which have much purer components to spatial contrast, and when contrast sensitivity was investigated by this approach the detection rate of these latent contrast changes was much enhanced. The checksize used in this part of the experiment (19' or 25') determined contrast sensitivity in the ‘mid-range’ of the total contrast sensitivity function (which is approximately equal to the maxima found in the function when it is obtained psychophysically to sinusoidal grating, that is, at 2 cycles degree⁻¹). The data therefore do not allow us to make predictions on whether overall contrast sensitivity was diminished, though the fact that all patients had a good standard of visual acuity suggests
that sensitivity to small, high contrast elements near the resolution limit was unaffected.

In conclusion, our data show that it cannot be assumed that a good standard of central acuity measured postoperatively in pseudophakic patients, necessarily implies 'normal vision'. We would submit that the role of contrast sensitivity testing as a means of assessing the efficacy of an IOL implant, either by psychophysical methods or by using evoked potentials as we have done, has merit as a useful adjunct in the clinical management of these patients.

References


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