Comparative electrophysiological responses in anisometropic and strabismic amblyopic children

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Purpose: To compare anisometropic hypermetropic amblyopic and strabismic amblyopic responses to pattern electoretinogram (PERG) and pattern visual evoked potential (PVEP).

Materials and methods: Fifty-six patients – 18 hypermetropic anisometropic amblyopic children (mean age 9.70±2.5 years), 19 strabismic amblyopic children (mean age 10.30±2.6 years) and 19 normal emetropic subjects (mean age 10.10±2.2 years) – were enrolled in this study. After routine ophthalmic examination, PERG and PVEP were recorded in response to checks reversed at the rate of two reversals/second stimulating macular area.

Results: The difference between hypermetropic anisometropic amblyopia and strabismus amblyopia with respect to P100/P50/N95 wave latencies (P=0.055/0.855/0.132) and P100/P50/N95 amplitudes (P=0.980/0.095/0.045) was not statistically significant. However, there was a significant statistical difference between strabismic amblyopia group and controls for P100/P50/N95 latencies (P=0.000/0.006/0.004).

Conclusion: Our findings indicated that despite clinical differences between anisometropic amblyopic and strabismic amblyopic patients, no differences were found in the responses of PVEP and PERG. The abnormal components of the PVEP and PERG in amblyopic subjects could reflect a retinal dysfunction in the visual pathway.

Keywords: amblyopia, strabismus, pattern electoretinogram, pattern visual evoked potential, anisometropia

Introduction

Amblyopia is a visual disorder defined as a decrease in visual acuity of one or both eyes that results from abnormal binocular interaction or pattern visual deprivation that occurs in the immature visual system of early childhood in eyes that appear typically normal.1,2 Amblyopia has a significant effect on psychosocial behavior, and therefore, early (before 4 years of age) visual testing is highly recommended for all children.3,4 Early strabismus and refractive errors are risk factors for unilateral amblyopia, whereas bilateral astigmatism and hyperopia are risk factors for bilateral amblyopia.5 Less common causes of visual deprivation such as eyelid ptosis and congenital cataracts may also result in amblyopia.6,7 Strabismus and anisometropia have different mechanisms of inducing amblyopia.8 Abnormal binocular interaction in strabismus generates suppression and amblyopia of the second diplopic image.9 On the other hand, unequal visual inputs that occur in anisometropia result in progressive suppression of the unfocused image of the ametropic eye that cannot harmonize with the image from the fellow eye.10 These two mechanisms may correspond to different anatomic locations for anisometropic and strabismic amblyopia. Although the anatomic basis of amblyopia seems to be located in the primary and secondary visual cortex,11 lateral geniculate nucleus dysfunction has been documented and retinal involvement may...
also be implicated. Abnormal macular structural images have been found using optical coherence tomography (OCT) in the amblyopic eye.\textsuperscript{12–16} These reports suggest that it is important to consider the relationship between possible structural anomalies and electrophysiological responses.\textsuperscript{17,18} Preganglionic-level retinal dysfunction may occur as suggested by reduction in the abnormal pattern electroretinogram (PERG) amplitude in the amblyopic eye.\textsuperscript{19} In this study, we propose to compare hypermetropic anisometric amblyopic and strabismic amblyopic responses to PERG and pattern visual evoked potential (PVEP).

**Materials and methods**

**Study design**

This is a cross-sectional study of 56 children. It followed the principles of Declaration of Helsinki for experimentation in humans and was approved by the Internal Ethical Committee Review Board of Antonio Pedro University Hospital of the Federal Fluminense University. Written informed consent was obtained from parents or guardians of all the volunteers. A complete comprehensive ocular examination including evaluation of best-corrected visual acuity, ocular motility and pupillary reflexes, slit lamp biomicroscopy, fundus examination and evaluation of cycloplegic refraction was conducted. Patients in the strabismic amblyopia group were esotropic and were treated for amblyopia before and after having undergone strabismic surgery by the same surgeon (LCSSL) and in the same hospital. Some minimal degree of misalignment and amblyopia persisted until the time of the study. Anisometropic patients were previously treated with corrective spectacles and occlusion. All subjects were assigned to one of the following three groups: 1) hypermetropic anisometropic amblyopic group (n=18), 2) strabismus-related amblyopic group (n=19) and 3) normal emetropic controls (n=19). Amblyopia was defined as a loss of the best-corrected visual acuity of greater than or equal to two lines on the visual acuity chart. Anisometropic amblyopia was defined as a refractive difference of (+)1.5 diopters with a spherical equivalent correction. Visuscopy was performed to determine the quality of fixation and exclude eccentric foveal fixation. Table 1 shows the group demographic and clinical data. Patients who had a history of intraocular surgery, neurological disease, retinal disease, glaucoma and nystagmus were excluded from the study.

**Electrophysiological exams**

To minimize diurnal variations, testing was conducted between 13:00 and 15:00 hours. Electrophysiologic tests were registered when the child was considered to be alert and cooperative. Pupil-modifying drugs and eye drops were not used before or during the examinations. PVEP and PERG were recorded at the Electrophysiological Laboratory of the Niterói Eye Hospital with a Nihon Kohden (Meb-7102K NeuroPack 2) electrophysiological apparatus in accordance with the PERG and PVEP International Society for Clinical Electrophysiology of Vision standards.\textsuperscript{20–22} The checkboard pattern stimulus was black and white with a check size of 16′ arc for a distance of 1 m. Monocular stimulation was performed after a 10-minute adaptation period in a semi-dark and acoustically isolated environment. Reverse checkerboard patterns (80% contrast and 110 cd/m\(^2\) mean luminance) were generated by a Cathodic Ray Tube (CRT) with a 75-Hz frame rate and two reversals per second. The standard PERG transient response was recorded using loop electrodes hooked to the eyelid into the lower fornix and in contact with the bulbar conjunctiva 5 mm below the corneal limbus. The electrodes were sterilized before and after each test.

Data were evaluated with Kruskal–Wallis and Mann–Whitney test for group analyses with SPSS 20. Bonferroni correction was applied to minimize multiple-comparison test interference. The correlation between visual acuity and PVEP P100 latency was determined using Pearson’s correlation coefficient. A P-value of <0.017 was considered statistically significant.

**Results**

A low positive correlation was found between visual acuity and PVEP P100 latency determined by Pearson’s correlation coefficient (\(r=0.45, P=0.001\)).

**Table 1 Demographics and clinical data**

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>Strabismic</th>
<th>Anisometric</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Amblyopic eye</td>
<td>Fellow eye</td>
<td>Amblyopic eye</td>
</tr>
<tr>
<td>Visual acuity (logMAR)</td>
<td>0.0</td>
<td>0.44±0.3</td>
<td>0.01±0.0</td>
</tr>
<tr>
<td>Refractive error* (SE)</td>
<td>0.25±0.3</td>
<td>0.62±0.6</td>
<td>0.66±0.6</td>
</tr>
<tr>
<td>Age (years)</td>
<td>10.10±2.2</td>
<td>10.30±2.6</td>
<td>9.70±2.5</td>
</tr>
<tr>
<td>Sex (male/female)</td>
<td>15/4</td>
<td>10/9</td>
<td>10/8</td>
</tr>
</tbody>
</table>

*Notes:* *None had cylinder >1.25. Data are presented as mean ± standard deviation.

*Abbreviation:* SE, spherical equivalent.
Table 2 Electrophysiological differences between controls, amblyopic eye and fellow eye

<table>
<thead>
<tr>
<th>Groups, N=56</th>
<th>Control, N=19</th>
<th>Strabismic, N=19</th>
<th>Anisometropic, N=18</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Amblyopic eye</td>
<td>Fellow eye</td>
<td>Amblyopic eye</td>
</tr>
<tr>
<td>P100-PVEP latency (ms)</td>
<td>105.28±14.9</td>
<td>130.68±21.9</td>
<td>110.37±14.8</td>
</tr>
<tr>
<td>P100-PVEP amplitude (μV)</td>
<td>12.81±7.3</td>
<td>8.96±6.2</td>
<td>11.88±7.9</td>
</tr>
<tr>
<td>P50-PERG latency (ms)</td>
<td>49.40±7.0</td>
<td>59.12±15</td>
<td>45.88±5.0</td>
</tr>
<tr>
<td>P50-PERG amplitude (ms)</td>
<td>2.28±1.4</td>
<td>2.09±1.6</td>
<td>3.11±1.8</td>
</tr>
<tr>
<td>N95-PERG latency (ms)</td>
<td>92.61±15.6</td>
<td>124.81±39.4</td>
<td>104.45±24.7</td>
</tr>
<tr>
<td>N95-PERG amplitude (μV)</td>
<td>4.02±1.7</td>
<td>3.94±1.4</td>
<td>5.14±1.9</td>
</tr>
</tbody>
</table>

Note: Data are presented as mean ± standard deviation.

Abbreviations: PVEP, pattern visual evoked potential; PERG, pattern electroretinogram.

Table 2 depicts the PVEP and PERG wave amplitudes and latencies of anisometropic amblyopic eyes, strabismic amblyopic eyes and controls. The data in the table show that there was a significant difference in the P100 (P=0.000) and P50/N95 (P=0.006/0.004) wave latencies between the strabismic amblyopic eye and controls. However, there was no significant difference in the P100/P50/N95 wave amplitudes (P=0.060/0.564/0.856) between the strabismic amblyopic eye and controls.

The comparison between both amblyopic group and controls was statistically significant for P100/P50/N95 latencies (P=0.001/0.008/0.007), but not for P100/P50 amplitudes (P=0.117/0.147/0.770). We found no significant difference between both anisometropic hypermetropic amblyopia and strabismus amblyopia with respect to P100/P50/N95 wave latencies (P=0.055/0.855/0.132) and P100/P50 amplitudes (P=0.980/0.095/0.045). The comparison between the P100/P50/N95 wave amplitudes (P=0.117/0.416/0.750) and latencies (P=0.543/0.416/0.876) of the fellow normal eyes and controls was also statistically significant.

Discussion

Our results are in accordance with previous papers that show a difference between the PVEP and PERG wave latencies of amblyopic eyes and controls, as well as no statistical difference in the P100/P50 wave amplitudes of controls and amblyopic eyes. The comparison between the P100/P50 wave amplitudes and latencies of the amblyopic fellow normal eyes and controls was also statistically significant notwithstanding a persistent treatment with topical atropine and full-or part-time occlusion. All patients included in our study were able to maintain fixation during examinations, as well as being collaborative despite their age, which is important for obtaining electrophysiological responses. We believe that clinical differences among the anisometropic hypermetropic amblyopic and strabismic amblyopic patients did not have an influence on the PVEP and PERG results.

The P50 and N95 components of the PERG may be affected in different diseases and could reflect a dysfunction of the retinal ganglion cell. The N95 wave component is more sensitive to stimulus contrast and special frequency, which are both more specific of ganglion cell function. On the other hand, the P50 wave component is poorly sensitive to stimulus contrast, but more sensitive to luminance, a more characteristic response of the overall internal retina. Therefore, PERG offers an assessment of the central retinal function and transcription of the retinal ganglion cell activity. Normal PERG wave amplitudes of amblyopic eyes and controls recorded in our sample may be a reflection of individually optimized alignment and stability. In our series, PVEP responses were obtained by high spatial frequencies (16-minute check) that specially stimulate the macular region. This region has a larger cortical representation, and abnormal PVEP responses represent the function of macular region in amblyopes. Small check (ie, 15-minute check) of probably active small axons prevalent in the parvocellular system and prolonged latencies of the PVEP could suggest deficiency in specific group of neurons.

Although our findings showed a low positive correlation between visual acuity and PVEP P100 latency, this weak correlation could not influence the results because the visual acuity was similar in both amblyopic groups.

It is known that the age of onset of amblyopia may influence the results of electrophysiological tests, and it is also of our knowledge that the age of amblyopia onset in cases of strabismus is often earlier than in anisometropic cases. For these reasons, during the enrollment phase of the study, many patients were excluded. In order to constitute more homogeneous groups without significant differences in age and visual acuity, we included patients with similar date of onset of amblyopia. As, in our study, the date of onset of amblyopia was recorded while interviewing the parents, we should consider that this can lead to misinformation about the exact time of onset of amblyopia.
Future studies of amblyopia and electrophysiological tests can be developed comparing other types of visual function exams, including spatial visual function with low contrast and contrast-sensitive function.

Conclusion

Our findings indicated that despite clinical differences between the anisometropic amblyopia and strabismic amblyopia, no differences were found in the responses of PVEP and PERG. On the other hand, the PVEP and PERG latencies were altered between the amblyopic group and the control group. The abnormal components of the PVEP and PERG in amblyopic subjects could reflect a retinal dysfunction in the visual pathway.

Acknowledgments

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Disclosure

The authors report no conflicts of interest in this work.

References
