Occult Fifth Nerve Dysfunction In Multiple Sclerosis

ANDREW EISEN, DONALD PATY, SHERRILL PURVES and MAUREEN HOIRCH

SUMMARY: Somatosensory evoked potentials (SEPs) were recorded following trigeminal nerve stimulation in 25 normal subjects. Mucosal stimulation of the lip resulted in a reduced stimulus artefact. The three initial peaks, N13, P19, and N30, measured respectively 12.8 ± 0.9 ST, 19.3 ± 1.4 ST, and 28.6 ± 1.7 ST. Blink reflex studies were also performed in most of these subjects. In 41.4% of 29 patients with established or suspected multiple sclerosis, the trigeminal SEP was abnormal. Additional use of the blink reflex raised the overall incidence of trigeminal nerve dysfunction to 51.7%. None of the patients had clinical evidence of fifth nerve involvement either historically or on examination. Four of seven patients with progressive spinal MS and two patients whose only deficit was that of optic neuritis, had abnormal trigeminal SEPs. It is concluded that occult involvement of the pontine fifth nerve structures occurs frequently in MS despite the rarity of corresponding clinical findings. The trigeminal SEP is a useful additional neurophysiological method.

INTRODUCTION

The value of evoked potentials (EPs) in the diagnosis of multiple sclerosis (MS) is well established (McDonald and Halliday, 1977; Poser, 1980; Chiappa, 1980; McDonald, 1980). These physiological tests, however, add little in patients with definite MS. Their real value lies in detecting subclinical second lesions in patients in whom the diagnosis is suspect.

In this study we describe the value of the trigeminal somatosensory evoked potential in revealing evidence of fifth nerve dysfunction in patients with established or suspected MS in whom facial nerve sensation was normal clinically.

METHODS

Subjects and patients

There were 25 control subjects. They either had no medical disease, being recruited from among the medical and paramedical staff of the department, or were patients referred to the laboratory for diseases such as carpal tunnel or cubital tunnel syndromes, considered to be irrelevant in terms of trigeminal evoked SEPs. None gave histories of systemic disease, alcoholism, or treatment with medication likely to affect peripheral or central nervous system conduction. The procedure was carefully explained to the control and patient groups. The mean age of the control group was 42.8 years, with a range of 19 - 65 years. Fifteen were female. Twenty-nine patients referred to the laboratory as having established or suspected MS were similarly studied. Their mean age was 41.4 years (range 19 - 66 years). Nineteen were female. Twenty-nine patients referred to the laboratory as having established or suspected MS were similarly studied. Their mean age was 41.4 years (range 19 - 66 years). Nineteen were female. The patients were classified according to the criteria of McDonald and Halliday (1977) as having definite MS - 6, probable MS (two lesions and
one episode or two episodes and one lesion) - 8; progressive spinal - 7; and suspected (one typical lesion) - 7. None of the patients had evidence of trigeminal dysfunction either by history or upon clinical examination.

**Stimulation and recording**

Electrophysiological testing was performed with the subjects supine. The room was semi-dark and air-conditioned, having an ambient temperature of between 20 and 22°C. Subjects were encouraged to relax and doze off if they so wished.

The stimulus was a square-wave current pulse of 0.2 S⁻³ duration applied at a rate of 5 Hz to each corner of the mouth sequentially. The cathode was so positioned as to be in contact with the mucosal surface of the lip. Stimulus intensity was adjusted to 2.5 times threshold, usually less than 7.5 mA. This was well tolerated. Scalp needle electrodes were used to record the evoked response. One electrode was placed either at C5 or C6 (International 10-20 system) contralateral to the side that was stimulated. A referential electrode was positioned at Fpz. A ground electrode was placed around the upper extremity ipsilateral to the side stimulated. Using this recording montage, little difficulty was encountered with the stimulus artefact, which could not be improved upon by placing the ground around the neck or head.

The input from the recording electrodes was led through differential amplifiers having a band pass of 0.5 Hz - 0.2 kHz. The time analyzed was 50 S⁻³ and 1024 epochs were averaged. After an interval of two or three minutes, a further 1024 epochs were added to the initial averaged response to enhance time-locked events.

Latency measurements were made by use of a cursor on the oscilloscope. Peaks were labelled according to the International nomenclature recommended by Donchin et al (1977). Thus, peaks having latencies of 20 S⁻³ or less were labelled to the nearest millisecond, whilst those with mean values greater than 20 S⁻³ were labelled to the nearest 5 S⁻³. A component was also identified by its polarity. N = negative, P = positive, and using the cephalic referential system described above, surface negativity was upward defllecting.

Primary and secondary components of the blink reflex (Kugelberg, 1952; Kimura et al, 1970; Shahani and Young, 1972) were recorded from the orbicularis oculi muscles bilaterally. A small scalp needle electrode was used to record the responses against a reference electrode positioned at Fpz, as used when recording the SEPs. Stimuli were delivered at a rate of not more than once every two seconds to the supraorbital notch. Stimulus intensity was adjusted so that the largest responses were obtained with the least discomfort. Five trials were superimposed and responses having the shortest latencies were measured.

In this study, latencies of the various peaks of the SEP and blink reflex components were considered abnormal if greater than three standard deviations above the normal mean.

**RESULTS**

Normal data for the trigeminal SEPs and blink reflex responses are summarized in Tables 1 and 2 respectively. Figures 1 and 2 are representative normal examples of these. Twelve of the 29 patients (41.4%) had abnormalities (prolonged latencies or absence) of their trigeminal SEPs. This occurred bilaterally on five occasions. Abnormalities were recorded in 3 of the 6 definite cases of MS, 2 of the 8 probable cases, 4 of the 7 progressive spinal cases, and 3 of the 8 patients with suspected MS. The patients had neither symptoms nor signs indicative of trigeminal nerve involvement. Figure 3 is an example of a patient with progressive spinal MS whose trigeminal SEPs were abnormal bilaterally, whereas a unilateral abnormality is exemplified in Figure 4 recorded from a patient with definite MS.

Eighteen of the 29 patients had blink reflex measurements. Five (27.8%) demonstrated abnormalities indicative of trigeminal nerve dysfunction. Thus, both ipsi- and contra-lateral secondary responses were absent or difficult to obtain when the involved side was stimulated, but both were of normal latency when the uninvolved side was stimulated (Kimura, 1973, 1975). In addition, the primary component was delayed or absent in these patients. There were abnormalities of the trigeminal SEPs and/or the blink reflex in 15 (51.7%) of the patients.

**Figure 1 — The SEPs shown were recorded following trigeminal nerve stimulation of the left corner of the mouth, the cathode being placed over the mucosal membrane. The top tracing is an average of the blink reflex in 15 (51.7%) of the patients.**
TABLE 1
Latencies and Side to Side Differences of SEP Peaks Following Trigeminal Nerve Stimulation

<table>
<thead>
<tr>
<th>Peak</th>
<th>N13</th>
<th>P19</th>
<th>N30</th>
</tr>
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<tbody>
<tr>
<td>Latency (S⁻³)</td>
<td>12.8 ± 0.9</td>
<td>19.3 ± 1.4</td>
<td>28.6 ± 1.7</td>
</tr>
<tr>
<td>(range)</td>
<td>(11.6 - 14.8)</td>
<td>(16.6 - 21.5)</td>
<td>(24.7 - 32.6)</td>
</tr>
<tr>
<td>Upper Limit (S⁻³)</td>
<td>15.5</td>
<td>23.6</td>
<td>33.6</td>
</tr>
<tr>
<td>Side to Side difference (S⁻³)</td>
<td>0.6 ± 0.5</td>
<td>0.6 ± 0.4</td>
<td>1.2 ± 0.8</td>
</tr>
<tr>
<td>(range)</td>
<td>(0.1 - 1.5)</td>
<td>(0 - 1.6)</td>
<td>(0 - 2.9)</td>
</tr>
<tr>
<td>Upper Limit (S⁻³)</td>
<td>2.1</td>
<td>1.9</td>
<td>3.7</td>
</tr>
</tbody>
</table>

TABLE 2
Normal Blink Reflex Studies

<table>
<thead>
<tr>
<th>Component</th>
<th>Latency (S⁻³)</th>
<th>Upper Limit (S⁻³)</th>
<th>Side to Side difference (S⁻³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary</td>
<td>10.0 ± 0.7</td>
<td>12.1 (8.9 - 11.4)</td>
<td>0.4 ± 0.3 (0 - 0.9)</td>
</tr>
<tr>
<td>Ipsilateral Secondary</td>
<td>33.6 ± 2.1</td>
<td>39.8 (29.4 - 36.1)</td>
<td>0.5 ± 0.4 (0 - 1.5)</td>
</tr>
<tr>
<td>Contralateral Secondary</td>
<td>33.9 ± 2.1</td>
<td>40.4 (29.9 - 37.6)</td>
<td>1.2 ± 0.8 (0.1 - 2.3)</td>
</tr>
</tbody>
</table>

Difference between ipsilateral and contralateral secondary responses (N = 40) measures 0.9 ± 0.8 S⁻³. Range 0 - 2.5 S⁻³, upper limit 3.3 S⁻³.

DISCUSSION:
Use of trigeminal nerve stimulation to evoke somatosensory potentials (SEPs) has rarely and only recently been described (Stöhr and Petrusch, 1979; Bennett and Jannetta, 1980; Stöhr et al, 1981). Difficulty can arise because of the large stimulus artefact resulting from the proximity of stimulating and recording electrodes. This problem can be considerably minimized by stimulating mucosal membrane rather than skin, which has a higher impedance (Bennett and Jannetta, 1980). When this is done the first negative peak, N13, is easily recognizable in normal subjects. Accurate positioning of the recording electrode over C5 or C6 (International 10-20 system), which approximately overlie the cortical representation of the face area, is critical because the waveform, especially N13, is rapidly degraded if the recording electrodes lie much outside these scalp positions. No attempt was made to measure the amplitude of the trigeminal SEPs as, for example, was done by Stöhr et al (1981). In our experience this is too variable in normals to be valuable, although a greater than 50% side to side difference in amplitude would certainly be relevant. The neural generators of the three peaks of the trigeminal SEP that were measured are not known. Presumably, N13 corresponds to N20 of SEPs recorded following upper extremity stimulation and P40 recorded with lower extremity stimulation (Eisen and Odusote, 1980), which represent arrival of impulses at the primary somatosensory cortex (Allison et al, 1979).

None of the patients in the present series had clinical evidence of trigeminal nerve dysfunction either by history or on examination. Nevertheless, half of those with definite MS showed abnormalities of their trigeminal SEPs. This incidence of abnormal...
evoked potentials in patients with clinically definite MS but in whom there were no corresponding clinical findings matches the incidence of abnormal visual evoked potentials (VEPs) in definite MS patients without visual symptoms or signs (McDonald, 1980). However, the incidence of abnormal brainstem auditory potentials (BAEPs) found in definite MS without brainstem involvement is considerably lower (Purves, 1981). The role of VEPs in uncovering occult lesions of the optic nerves in multiple sclerosis is now well established (Poser, 1980). They have been of particular value, for example, in patients with progressive spinal MS. In this subgroup of multiple sclerosis about 40% of patients have been shown to have abnormalities of VEPs (Halliday et al., 1973; Asselmann et al., 1975; Hennerici et al., 1977; Matthews et al., 1977; Paty et al., 1979). Abnormalities of the VEP in progressive spinal MS clearly indicates involvement outside the spinal cord and confirms the presence of disseminated plaques. All of the patients in this study with progressive spinal MS had abnormalities of SEPs evoked by lower extremity cutaneous nerve stimulation (Eisen and Elleker, 1980). Of much greater interest, however, was the finding that nearly 60% of this small group had abnormalities of their trigeminal SEP, indicating a lesion rostral to the spinal cord.

Just short of 40% of our suspects also had abnormal trigeminal SEPs. This in itself is not too helpful, since it mirrors the incidence of abnormal SEPs evoked by more conventional means in MS suspects. More pertinent was the finding that in two of these patients the only clinical abnormality had been one of optic neuritis. This combination of optic neuritis and an abnormal trigeminal SEP would again point to dissemination of lesions.

In a large series of patients, Kimura (1975) demonstrated abnormality of the blink reflex in 78% of patients with MS who had clinical evidence of pontine involvement. This incidence of abnormality dropped to 40% when there were no such clinical features. In the present much smaller series, just short of 30% of the patients had abnormalities of their blink reflexes, which were specifically indicative of trigeminal nerve involvement. These were all clinically asymptomatic. Using the trigeminal SEP and blink reflex in combination, occult involvement of trigeminal pathways was documented in half the patients. This is a slightly higher yield than obtained with VEPs in patients with possible or probable MS who had no history of optic neuritis (Chiappa, 1980). Although transitory perioral numbness

Figure 3 — In this patient who had progressive spinal MS, no convincingly recordable trigeminal SEPs could be obtained from either side. The patient had no clinical evidence of abnormal facial sensation.

Figure 4 — The top tracing of this patient with definite multiple sclerosis was recorded following left sided trigeminal stimulation. It is a normal response. The bottom tracing was recorded following stimulation of the contralateral side of the mouth and no measurable SEP was obtained.
and tingling are not uncommon in multiple sclerosis, isolated objective facial sensory deficit is rare apart from trigeminal neuralgia (McAlpine et al, 1965; Poser et al, 1980). This discrepancy between the apparent frequency of occult lesions and clinical findings should not cause concern. It is well known that the extent of lesions in multiple sclerosis is frequently greater than would be anticipated from clinical observations (Ghatak et al, 1974; Wisniewski et al, 1976). In chronic plaques there have been recent suggestions to indicate that relatively normal function can be maintained through the spread of sodium channels into the demyelinated internodal region so that impulse conduction can continue to be supported (Ritchie and Rogart, 1977; Waxman, 1977). Spread of sodium channels in this manner can be considered analogous to the spread of acetylcholine receptors in denervated muscle.

Because the trigeminal SEP appears to be useful in revealing occult brainstem plaques relatively frequently, its use could be advantageously combined with that of SEPs evoked by upper and/or lower extremity nerve stimulation. In this way it becomes possible to detect plaques in anatomically different parts of the somesthetic pathways, allowing confirmation of the diagnosis of multiple sclerosis.

ACKNOWLEDGEMENT

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REFERENCES


