SHORT REPORT

Sensory deficits of a nerve root lesion can be objectively documented by somatosensory evoked potentials elicited by painful infrared laser stimulations: a case study

J Lorenz, H C Hansen, K Kunze, B Bromm

Abstract

Somatosensory evoked potentials (SEPs) in response to painful laser stimuli were measured in a patient with a unilateral sensory deficit due to radiculopathy at cervical levels C7 and C8. Laser evoked potentials (LEPs) were compared with SEPs using standard electrical stimulation of median and ulnar nerves at the wrist and mechanical stimulation of the fingertips by means of a mechanical stimulator. Early and late ulnar and median nerve SEPs were normal. Mechanical stimulation resulted in w shaped early SEPs from all five fingertips with some degree of abnormality at the fourth and fifth digits of the affected hand. Late LEPs were completely absent for stimulations at affected dermatomes and normal in the unaffected control dermatomes. The border between skin areas with normal or absent LEPs was very sharp and fitted the dermatomes of intact C6 and damaged C7 and C8 nerve roots. It is suggested that pain dermatomes are narrower than tactile dermatomes because thin fibres of the nociceptive system, activated by laser stimuli, probably do not overlap between adjacent spinal segments to the same extent as thick fibres of the mechanoreceptive system, activated by standard electrical or mechanical stimulation.

(J Neurol Neurosurg Psychiatry 1996;61:107–110)

Keywords: radiculopathy; laser evoked potentials; somatosensory evoked potentials; mechanical stimulation; electrical stimulation

Verification and topodiagnosis of nerve root affections usually cause no problem if sensory and motor signs indicate the same segment and the motor axonal damage can be proved by EMG. However, evaluation of the sensory deficit completely relies on the patient’s report so that in cases of isolated dorsal root affection or incongruent motor and sensory signs there is a need for objective methods regarding the sensory component. Standard electrical somatosensory evoked potentials (SEPs) can fail to document the sensory deficit.1-3 We present a patient with a unilateral sensory deficit in C7 and C8 dermatomes. Objective verification of the border between areas of normal and affected skin sensitivity was achieved by means of late SEPs in response to painful radiant heat stimuli delivered by a CO2 laser. This infrared laser selectively activates superficial Aδ and C fibres of the nociceptive system and has been previously reported as an appropriate tool to examine the functional integrity of peripheral small fibres and anterolateral tracts.4-6

Patient and methods

MEDICAL HISTORY AND CLINICAL FINDINGS

This 65 year old man complained about cervical pain and brachialgia in his right arm that had started some five years previously. For seven months he had noted numbness without paraesthesia of the ulnar portion of the forearm and the hand and loss of grip strength. Particularly, the fifth digit was reported to be

Institute of Physiology
J Lorenz
B Bromm

Neurological Department
tor, Hamburg, Germany.
H C Hansen
K Kunze

Correspondence to:
Professor Dr B Bromm,
Institute of Physiology,
University Hospital
Eppendorf, Martinistraße 52,
20246 Hamburg, Germany.

Received 25 September 1995
and in revised form
24 January 1996
Accepted 16 February 1996

Sensory testing results of affected (C7/C8) and non-affected (C6) dermatomes

<table>
<thead>
<tr>
<th>Sensory quality and function method</th>
<th>Affected dermatome (C7/C8)</th>
<th>Non-affected dermatome (C6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mechanosensitivity:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pressure threshold (PT) v Frey filaments</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Vibration threshold (VT) 128 Hz tuning fork; 8/8 scaled</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Light touch</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Cotton wool stabs</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Joint position sense passive finger/toe movements</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>M score (%)</td>
<td>25</td>
<td>100</td>
</tr>
<tr>
<td>Temperature sensitivity:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Warm sense</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Warm sense</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Gold sense</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Cold sense</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Gold sense</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Cold sense</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>T score (%)</td>
<td>0</td>
<td>100</td>
</tr>
<tr>
<td>Pain sensitivity:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mechanical pain pulling a hair</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Mechanical pain pin prick</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Sharp blunt discrimination</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Safety pin</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Heat pain threshold (HPT) CO2 laser</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>P score (%)</td>
<td>0</td>
<td>100</td>
</tr>
</tbody>
</table>

0 = not perceived; 1 = disturbed; 2 = normal. (PT) 0 = > 275 g; 1 = 0-8 g-275 g; 2 = < 0-8 g. (VT) 0 = 1/8; 1 = 1/8-4/8; 2 = > 4/8. (HPT) 0 = > 30 W; 1 = 18 W-30 W; 2 = < 18 W.
NEUROPHYSIOLOGICAL ASSESSMENT

Laser evoked potentials (LEPs) were elicited as published previously. In summary, cutaneous radiant heat stimuli delivered by a CO$_2$ laser were applied to the dorsal skin of affected and non-affected dermatomes. Laser intensities of 20 W (20 ms duration; 5 mm beam diameter) were used which is above normal pain threshold (10-0 (SD 2-3) W). Electrical nerve stimuli were also randomly interspersed between laser stimuli which served to evaluate late SEPs. Two blocks of 60 stimulations at both control and affected dermatomes were repeated. An EEG was recorded within 0-1 and 30 Hz over Cz and averaged according to stimulus modality (electrical vs laser) and stimulated dermatome (affected vs non-affected).

Conventional early cortical SEPs were recorded using electrical stimulation (3 Hz-repetition rate, 0-3 ms duration, constant current) of median and ulnar nerves at the wrist of the affected hand. Stimulus intensity was adjusted to the sum of sensory and motor threshold. The EEG was recorded within 10 and 1000 Hz at electrode positions over the somatosensory projection area against Fz reference. A total of 256 artefact free poststimulus epochs of 100 ms duration were digitised at 3000 Hz and averaged on line. The N20 component was compared with normal height corrected values from our laboratory.

Early cortical SEPs were additionally elicited at all five finger tips of the affected hand by a mechanical stimulator (Somedic TS 120) which had a 10 mm diameter stimulating probe with nine blunt tips, each with a diameter of 2 mm. The patient's stimulated finger rested on the probe, which was displaced by 500 µm with a slope of 1000 µm/ms at a stimulus rate of 2 Hz and a duration of 2 ms. This stimulation did not cause a movement of the finger. Previous experiments with the mechanical stimulator on normal subjects in our laboratory (unpublished data) have proved this setting appropriate to reliably elicit w shaped primary cortical responses consisting of N28-P35, likely to correspond to N20 and P27 of

“dead”. Neurological examination disclosed painful immobility of the cervical spine with spontaneous head tilt towards the right. Strength examination disclosed moderate paresis (MRC grade 4) of right wrist extensors and interossei without atrophy. Tendon reflexes of triceps muscles were slightly reduced on both sides. Strength and tendon reflexes of other muscles were normal.

The sensory deficit was determined according to a standard protocol (see table) that tested for mechanosensitivity (M score), temperature sensitivity (T score), and pain sensitivity (P score). Four subtests of each sensory quality were scored on a three point scale: normal (2 points), disturbed (1 point), and absent (0 points). The sum scores were normalised to percentage of maximum value. Within affected C7 and C8 dermatomes there was a complete loss of temperature and pain sensitivity and a 25% residual mechanosensitivity due to the patient’s ability to perceive vibration at 2/8 when stimulated with an 8/8 scaled 128 Hz tuning fork and to perceive pressure of calibrated von Frey nylon monofilaments at 164 g. Normal sensitivity was present in the non-affected C6 dermatome. The clinical findings were regarded as consistent with radiculopathy in right C7 and C8 vertebrae due to spondylosis.

Figure 1  Computed axial tomography shows hypodense disc material protruding into the vertebral canal and right neuroforamen. The scan corresponds to the level between C7 and Th1 vertebrae.

Figure 2  Laser evoked potentials are absent after stimulation of affected C7/C8 dermatomes (shaded area) and normal after stimulation of non-affected C6 dermatome.
Laser evoked potentials in radiculopathy

Figure 3  Early somatosensory evoked potentials (SEPs) after electrical stimulation of right median (top left) and ulnar nerves (top right) and after mechanical stimulation with pressure pulses of all five digits at the fingertips of the right hand. The shaded area indicates the extent of the sensory deficit within C7/C8 dermatomes. Electrical SEPs are normal. Mechanical SEPs yield poor configuration and latency delay of P50 at the fourth and fifth digits respectively.

Results

Needle EMG with semiquantitative motor unit potential (MUP) evaluation disclosed high amplitude MUPs (up to 4 mV) in the right extensor carpi radialis, extensor digitorum, extensor carpi ulnaris, interossei dorsalis I, and paraspinals C7/C8, but not in biceps and brachioradialis. Fibrillation potentials were absent in tested muscles. These results were considered to be consistent with a radiculopathy involving right C7 and C8 roots. Radiological studies of the cervical spine showed spondylosis between vertebrae C6/C7 and C7/Th1. Computed tomography disclosed narrowing in the right intervertebral nerve root canal at level CV 6/7 (fig 1). The lower adjacent segment was partly obscured by bone artifacts. An MRT was not available. Anatomical findings were consistent with a cervical spondylosis and suggested compression of the right nerve root C7.

The shaded area in fig 2 shows the extent of sensory deficit at the hand dorsum. Laser evoked potentials were absent for stimuli applied to the shaded skin areas, whereas
normal LEPs were elicited after stimulation of the non-shaded area. Thus LEPs confirmed clinical analgesia and thermanaesthesia and indicated very sharply the border between dermatomes of affected and non-affected cervical roots.

Figure 3 gives the results of early SEPs after electrical stimulation of median and ulnar nerves at the wrist and after mechanical stimulation of all five digits of the affected hand in this patient. Early electrical SEPs were well elicited from median and ulnar nerves as typical w shaped configurations. Latencies of N22 (median nerve) and N24 (ulnar nerve) were normal according to our standards. Late components of median and ulnar SEPs (not shown) were also normal. Mechanical stimulation of the digits also resulted in w shaped waveforms with correspondent negative amplitudes and positivities about 6 to 8 ms later and of smaller amplitudes when compared with components after direct electrical nerve stimulation at the wrist. This difference can be explained by the fact that mechanical finger pulses take additional receptor activation time, longer peripheral conduction time, and do not synchronise fast conducting α fibres. Comparison of the waveforms of all five fingers showed some degree of abnormality in configuration and latency of the P50 component after stimulation of digit four and—less significant—of digit five.

Discussion
Radiculopathies exhibit characteristic clinical signs such as muscle weakness, irradiating pain, and sensory deficits along myotomal and dermatomal innervation zones of the affected spinal roots. Normal nerve conduction velocity often allows differentiation from peripheral postganglionic nerve disease. Cortical or spinal SEPs after electrical nerve stimulation have been reported as useful in the evaluation of radiculopathies by some authors, but were considered of poor utility by others. Electrical dermatomal stimulation has been proposed, but again, doubt about its utility was raised both in lumbosacral and cervical radiculopathies. We also found normal early and late electrical SEPs from stimulation of mixed nerve trunks of the affected hand. We expected this result because median and ulnar nerve SEPs could be sufficiently conducted via intact C6 and Th1 roots in our case. Mechanical stimulation showed only minor abnormality in deficient dermatomes as indicated by poor configuration and delay of P50 particularly after stimulation of digit four compared with the intact thumb. By contrast, absence of LEPs after painful stimuli applied to affected dermatomes suggests considerable less intersegmental overlap of small diameter nociceptive afferents than large diameter mechanoreceptive afferents. This finding fits clinical experience of a narrow analgesic stripe as a typical feature of sensory loss in monoradiculopathy.

Foerster, as early as 1936, described greater caudo-oral extension of tactile dermatomes than pain dermatomes determined after surgical dissection of dorsal roots in patients treated for extreme spasticity. He also reported a case in which dissection of C4 and C5 dorsal roots only resulted in clinical described thermanaesthesia and analgesia at the shoulder whereas tactile sensitivity remained intact. Inoue and Buchthal confirmed the existence of overlap of dermatomes between two to three spinal segments by recording spinal nerve potentials from electrodes inserted near cervical roots after electrical stimulation of various proximal and distal mixed nerves and sensory fibres in the upper limbs of healthy subjects. This case study supports Foerster by providing evidence that thin fibres of the pain and temperature system exhibit less intersegmental overlap than thick fibres of the tactile system. Although not considered of major importance for routine diagnostics in nerve root lesions, recording of LEPs may supplement standard examinations when objective documentation of the sensory disturbance is needed.

This work was supported by grants of the Deutsche Forschungsgemeinschaft. We gratefully acknowledge W Eickhoff for the EMG study.