Estimation of the brain and spinal cord conduction time in man by means of the somatosensory evoked potentials and F and H responses

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SUMMARY Somatosensory evoked potential (SSEP) recordings from the scalp were performed in 17 healthy subjects. In seven of these SSEP was also recorded at the level of the second lumbar spine. In the other ten F and H responses and the corresponding M responses were studied. By means of the SSEP recordings at the level of the second lumbar spine and the F- and H-responses, the conduction time in the brain and spinal cord, that is central latency, was calculated and the following results were obtained: 160 ms with standard deviation (SD) ±1.1 ms (by means of SSEPs), 9.5±2.4 ms (by means of F response) and 13.1±1.5 ms (by means of H response). Of the three methods used the H response method seems to be the best for clinical purposes: it is easy to perform and statistically it is more stable than the F response recording; moreover the recording can be performed reliably even in persons with thick back muscles and subcutaneous fat, unlike the evoked potential procedure which only with difficulty shows detectable responses at the lumbosacral levels in such persons. Three patients are presented to illustrate the technique; in one of these the recording evoked potentials from the epidural space were recorded.

Electroneuromyography of the peripheral nervous system occupies an important position in clinical neurodiagnostic. However, for study at various levels of the spinal cord and brain it is less valuable. During the last few years somatosensory evoked potential (SSEP) recordings have improved the diagnostic approach to the spinal cord and brain: it is now possible to record scalp potentials evoked by stimulation of the tibial or peroneal nerves.1-4

In order to distinguish peripheral lesions from these at a higher level, that is in the spinal cord and brain, spinal SSEP recordings have now made it possible to differentiate conduction time at the root and peripheral nerve levels. Techniques available to perform the latter include surface recordings,5-7 and, with their better signal/noise ratio, epidural8-10 and subdural11-15 recordings. Dorfman4 has introduced a method to calculate the peripheral nerve and root conduction time by means of F response. In 1956 Dawson and Merton16 suggested that this F response was due to “recurrent” discharges from motoneurones activated antidromically. This suggestion has gained support from other investigators.17 18 However, it is known that the recognition of F response and the determination of latency to onset of F response may sometimes be difficult; it may vary in different persons and even in the same subject in different times of examination.19 In the proximal part of the sciatic nerve and roots the conduction velocity may be calculated also by means of Hoffman's (H) response,20 and the H response is easy to detect in clinical practice.

In this study three recording methods were used for the measurement of the conduction times at the peripheral levels (roots and peripheral nerve): (1) the somatosensory evoked potentials from the region of the second lumbar spine, (2) F response,
and (3) H response. From the measurements obtained, and from scalp somatosensory responses, the conduction time in the brain and spinal cord has been calculated.

Material and methods

Equipment During the recordings the subjects were supine and were instructed to minimise voluntary movements and swallowing. The recordings were carried out with routine EEG electrodes on the scalp and with chlorided silver cup electrodes of 9 mm diameter on the body. The signals were led first through a high impedance differential amplifier with bandpass of 20 Hz–0.5 kHz into an EMG-machine (Disa 14 C 12) and from there into a two-channelled signal analyser, HP-5481A, in which the analysis of both recordings was possible simultaneously. The resolution of sampling (bin-width) was 200 µs. Single stimuli were produced with the stimulator (Disa 3 K 62) at a rate of four per second, square wave pulses, duration 0.3 ms. The stimulation sites were popliteal fossa or ankle (peroneal or tibial nerve) or both. Stimulus intensity was adjusted in order to produce a slight twitching of the foot. The averaging of 512 responses was carried out from the scalp, recording site 2.5 cm behind the vertex with the reference electrode on the mid-forehead (Fz: 10–20 EEG system). The H responses were measured with the routine method applied for clinical purposes: the latency was measured to the beginning of the maximal H response during the stimulation of the tibial nerve from popliteal fossa, recorded from the surface of the soleus muscle; the reference electrode 20 mm distally from the proximal one. The latency measurement of F response was also performed to the beginning of the maximal F wave. In SSEP-recordings of the second lumbar spine the latency measurement was performed at the peak of response, since the peak was the clearest and most reliable indicator of latency.

In order to calculate the conduction time of the central nervous system in the brain and spinal cord, that is “central latency,” the measurements of conduction time of the peripheral nervous system were performed with three different methods: in seven subjects by means of evoked potentials from the level of the second lumbar spine with surface recordings, in ten by means of both H response and also F response.

Approach I In seven healthy subjects, aged 15 to 34 years, body length from 165 cm to 178 cm, stimulation of the peroneal nerve was performed separately from the popliteal fossa of both legs.

Recordings were taken with surface electrodes simultaneously from the scalp and also from the second lumbar spine. (Recordings from the inion were also carried out, but their signal/noise ratio was low and therefore they were not included in the present results.) Latency measurements were carried out at the onset of the scalp response (lat No), also at the peak P1 (lat P1), that is the positive peak of component I, and also the peak of negative deflection of the evoked response at the level of the second lumbar spine (lat L2). Figure 1 shows the typical configurations and the important landmarks of the responses in Approach I.

The conduction velocity of the sciatic trunk between the popliteal fossa and the second lumbar spine also was calculated. This conduction velocity (CVs) may be formulated:
(a) \( CVs = \frac{d}{\text{lat } L2} \) where \( d \) = distance between the popliteal fossa and the second lumbar spine (mm).

The conduction time at the level of brain and spinal cord, central latency (CLs), may be presented as the formula:
(b) \( CLs = \text{lat } No - \text{lat } L2 \).

Also the ratio of CLs and body length (B1) was
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calculated. This ratio (CLI<sub>s</sub>) is formulated as:

(c) CLI<sub>s</sub>=CL<sub>s</sub>/B1.

**Approach II** Ten volunteers, six males and four females aged 22 to 47 years, (body length from 154 to 199 cm), were studied by stimulation of the tibial nerve at the ankles of both legs. Recording from the scalp was taken as in Approach I: the latency was measured at the onset of the response (lat No) and at the positive peak P1. The latencies of F responses from both legs were measured at the beginning of the maximal F response (lat F) and at the one of the corresponding maximal muscle (M) response (lat M). The schematic representations of the scalp and F response are presented in fig 2.

The conduction time of the tibial nerve and the sciatic trunk (CT<sub>t</sub>) was calculated with the method presented by Dorfman, which may be formulated as:

(d) CT<sub>t</sub>=\frac{lat F-lat M-1}{2} (ms)

1 (ms)=the central delay of the spinal cord, proposed by Dorfman. Further, the conduction time at the level of brain and spinal cord, the central latency (CL<sub>r</sub>) may be presented as the formula:

(e) CL<sub>r</sub>=lat No - \left( \frac{lat F-lat M-1}{2} \right) (ms).

The value of, CL<sub>r</sub> was calculated in all the ten controls and was also compared to the body length (B1). This central latency index (CLI<sub>r</sub>) may be formulated as:

(f) CLI<sub>r</sub>=CL<sub>r</sub>/B1 (ms/m).

**Approach III** In the same controls as presented in Approach II, stimulation of the tibial nerve in the popliteal was performed separately on both legs. The recording site on the scalp was the same as in Approaches I and II. The latency was measured at the onset of the response, lat No, and at peak P1, lat P1. H responses from the soleus muscles on both legs were studied, and latencies were measured at the onset of the maximal H response (lat H) and the corresponding maximal M response (lat M). The schematic representations of the scalp and H responses are presented in fig 3. The method of averaging the conduction time (CT<sub>n</sub>) and the conduction velocity (CV<sub>n</sub>) of

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**Fig 2** Diagram of the scalp and F response, with the corresponding M response, induced by stimulation of tibial nerve at the ankle (Approach II). Upper tract: scalp response, latency measurements to the onset of response (No) and to the peak (P1) of the first positive deflection. No-P1-N1 is component I. Lower tract: F response with the corresponding M response; latency measurements were performed to the onset of F and M responses.

**Fig 3** Diagram of the scalp and H response with the corresponding M response induced by stimulation of tibial nerve in the popliteal fossa (Approach III). Upper tract: scalp response, latency measurements to the onset of response (No) and to the peak (P1) of the first positive deflection. Part of response, No-P1-N1, is component I. Lower tract: H response with the corresponding M response; latency measurements were performed to the onset of H and M responses.
the sciatic trunk (between popliteal fossa and spinal cord) has been formulated as follows:

\[(g) \quad CT_H = \frac{\text{lat } H - \text{lat } M - 1}{2} \text{ (ms)}\]

\[(h) \quad CV_H = \frac{80\% \times \text{body length (B1)}}{\text{lat } H - \text{lat } M - 1} \text{ (m/s)}\]

1 ms = the monosynaptic delay of the spinal cord, a time estimated from studies in animals and in man. The figure 80% is based on examinations of cadavers. The formula (h) expresses the average conduction velocity of the afferent and efferent neural pathways of the sciatic trunk.

The conduction time at the level of brain and spinal cord, that is the central latency (CLn), was also calculated in these 10 controls; it may be formulated as follows:

\[(i) \quad CL_n = \text{lat } No - \left( \frac{\text{lat } H - \text{lat } M - 1}{2} \right) \text{ (ms)}\]

Further, the CLn was compared to the body length of the controls and this value, the central latency index (CLH), may be formulated as:

\[(j) \quad CL_n = CL_n / B1 \text{ (ms/m)}\]

Student's *t* test for paired observations was used for statistical evaluation of the differences between CLf and CLn values in Approaches II and III.

**Patient studies** Case 1, a 30-year-old male, who had earlier been healthy but who had sustained acute trauma to the back about two months preceding the examination showed symptoms typical of radicular lesion of lumbosacral roots, especially on the right leg. Stimulation of the tibial nerve was performed separately on both legs from the popliteal fossa; 512 responses were recorded from the scalp and their averages taken. By means of H response and the corresponding M response the conduction velocity, CVH, was calculated for both sides. Thereafter stimulation of the tibial nerve from the popliteal fossa was performed and the evoked responses with surface electrodes were recorded from the second lumbar spine, the reference being on the forehead. 256 responses were averaged. Recordings of evoked responses were then carried out at the level of the third lumbar segment, between the third and fourth lumbar spine, with a platinum iridium wire electrode of 0.2 mm diameter placed in the epidural space using the technique presented by Shimoji et al. The reference electrode, also a platinum iridium wire, was inserted into subcutaneous tissue (depth about 0.5 cm) in the region of the right upper gluteal area. The stimulation of the tibial nerve was performed separately on both legs from the popliteal fossa, and 256 responses were averaged.

Case 2, a 38-year-old male with Hodgkin's disease who had suddenly suffered from acute paraplegia; myelography showed extradural metastasis at the level of the sixth to tenth thoracic segments, the CSF space being totally obliterated. Stimulation was performed from the tibial nerve at the level of popliteal fossa and recording was taken from the scalp. Conduction velocity, CVH, was calculated for both sides. Recordings from the scalp and the seventh cervical segment were also taken after stimulation of the median nerve from the wrist, separately in both hands (for details, see Siivola et al.)

Case 3, a 30-year-old male with multiple sclerosis (MS) had muscle weakness and sensory deficits mainly in the right limbs, especially on the right leg. Stimulation of the tibial nerve from the popliteal fossa separately on both sides was performed and recording taken from the scalp. The conduction velocity, CVH, was calculated by means of H response from both sides.

**Results**

The main results are presented in tables 1, 2, and 3.

### Table 1  The measured and calculated values of Approach I

<table>
<thead>
<tr>
<th>lat No (ms)</th>
<th>lat No/B1 (ms/m)</th>
<th>lat P1 (ms)</th>
<th>lat L2 (ms)</th>
<th>CVH (ms/m)</th>
<th>CLf (ms)</th>
<th>CLH (ms/m)</th>
<th>A-P1/N1 (μV)</th>
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<tr>
<td>Mean</td>
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<td>17.1</td>
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lat No = latency to the onset of scalp response
lat No/B1 = latency compared to the body length (B1) of controls
lat L2 = latency to the peak of evoked response at the level of the second lumbar spine
CVH = conduction velocity of the sciatic nerve and roots
CLf = central latency, conduction time at the level of the brain and spinal cord
CLH = CLf compared to the body length (B1) of controls
A-P1/N1 = amplitude between peaks P1 and N1 of scalp response
N = number of controls
Table 2  The measured and calculated values of Approach II

<table>
<thead>
<tr>
<th></th>
<th>lat No</th>
<th>lat No/Bl</th>
<th>lat P1</th>
<th>lat F</th>
<th>CT1</th>
<th>CL1</th>
<th>CLI1</th>
<th>A-P1/N1</th>
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<td></td>
<td>(ms)</td>
<td>(ms/m)</td>
<td>(ms)</td>
<td>(ms)</td>
<td>(ms)</td>
<td>(ms)</td>
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<tr>
<td>Mean</td>
<td>31.9</td>
<td>18.6</td>
<td>36.9</td>
<td>51.0</td>
<td>22.6</td>
<td>9.5</td>
<td>5.8</td>
<td>2.6</td>
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<tr>
<td>SD</td>
<td>2.2</td>
<td>1.2</td>
<td>2.4</td>
<td>6.8</td>
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</tbody>
</table>

lat No = latency to the onset of scalp response  
lat No/Bl = lat No compared to the body length of controls  
lat P1 = latency to the peak P1 of scalp response  
CT1 = conduction time of the tibial nerve and roots  
lat F = latency to the onset of the measured F-response  
CL1 = central latency conduction time at the level of the brain and spinal cord  
CLI1 = CL1 compared to the body length of controls  
A-P1/N1 = amplitude between peaks P1 and N1  
N = number of controls

Table 3  The measured and calculated values of Approach III

<table>
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<tr>
<th></th>
<th>lat No</th>
<th>lat No/Bl</th>
<th>lat P1</th>
<th>lat H</th>
<th>CT1</th>
<th>CV1</th>
<th>CL1</th>
<th>CLI1</th>
<th>A-P1/N1</th>
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<td>(ms)</td>
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<td>(μV)</td>
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<tr>
<td>Mean</td>
<td>25.0</td>
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<td>29.0</td>
<td>28.9</td>
<td>11.9</td>
<td>58.0</td>
<td>13.1</td>
<td>7.7</td>
<td>1.4</td>
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<tr>
<td>SD</td>
<td>2.1</td>
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<td>3.5</td>
<td>2.7</td>
<td>1.2</td>
<td>3.2</td>
<td>1.5</td>
<td>0.8</td>
<td>0.7</td>
</tr>
<tr>
<td>N</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>10</td>
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<td>10</td>
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</tr>
</tbody>
</table>

lat No = latency to the onset of scalp response  
lat No/Bl = lat No compared to the body length (Bl) of controls  
lat P1 = latency to the peak of P1 of scalp response  
CT1 = conduction time of the sciatic nerve and roots  
CV1 = conduction velocity of sciatic nerve and roots  
CL1 = central latency, conduction time at the level of the brain and spinal cord  
CLI1 = CL1 compared to the body length (Bl) of controls  
A-P1/N1 = amplitude between peaks P1 and N1 of scalp response  
lat H = latency to the onset of the maximal H response  
N = number of controls

Case 1, the conduction velocity of the sciatic nerve, CV_H, was 48 m/s on the right (the lesion side), a slightly abnormal value (differing more than 2 SD from the mean), and the corresponding CV_H value on the left was 55 m/s (normal). The responses from the scalp were not clearly distinguishable. The surface recordings from the second lumbar level induced with the stimulation of the tibial nerve (fig 4) were hardly detectable, but the recordings from the epidural space with platinum iridium wire electrode showed obvious responses (fig 5).

Case 2, the scalp recordings induced with the stimulation of the tibial nerve from the popliteal fossa showed no responses, but those produced by stimulation of the median nerves at the wrist showed obvious responses, as did recordings from the seventh cervical segment. Figure 6 shows the evoked responses at both levels induced by stimulation of the right median nerve. The H response was easy to detect on both sides and CV_H was normal on the right (55 m/s), but slightly decreased on the left (45 m/s).

Case 3, the H response was present on both sides and the CV_H values were normal. The latency at the onset of the scalp response of the right hemi-
Fig 5 Case 1, with radicular lesion at the lumbosacral level. Upper tract: recording of response from the epidural space with platinum-iridium wire electrode at the level of the third lumbar segment induced by stimulation of the right tibial nerve in the popliteal fossa (reference in subcutaneous tissue, about five cm from cathode). Lower tract: stimulation of the left tibial nerve in the popliteal fossa; recording as above (256 responses were averaged).

Fig 6 Case 2, with Hodgkin's disease. Upper tract: scalp response induced by stimulation of the right median nerve at the wrist. × shows the peak of the primary response of cortex peak-N20 (for details, see Siivola et al.19) Lower tract: response at the seventh cervical segment, recorded with surface electrodes between the sixth and seventh cervical spine, induced by stimulation of the right median nerve at the wrist. × shows the peak N13, the top of the negative deflection of the response.

Fig 7 Case 3, with MS. Upper tract: scalp response induced by stimulation of the right tibial nerve in the popliteal fossa. Lower tract: scalp response induced by stimulation of the left tibial nerve in the popliteal fossa.

Discussion

Somatosensory evoked potential recordings offer one method of measuring the conduction time of the central nervous system, brain and spinal cord. SSEPs, however, are much more difficult to record with surface electrodes from the thoracic or lumbar levels of the spinal cord than from the brachial plexus and cervical cord. F and H responses offer a non-invasive method of determining the conductivity of the lumbar roots and sciatic nerve.
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In Approach I the observed latency at the onset of scalp response (lat No, 29.2 ms, and that at the peak P1 (lat P1, 33.4 ms), are near reported values Larson et al.24, Giblin25 and Tsumoto et al.2 The latency values No and P1 were calculated from the scalp recordings because they originate from the component I of the scalp response (see figs 1–3) which, it has been suggested, results from postsynaptic activation of the thalamocortical neural pathways.2 Component I is the first cerebral manifestation in the scalp recording following the activation of the medial lemnisci tracts. The selection of the site of the recording electrode, about 2-5 cm behind from the vertex, is based on the investigations of Tsumoto et al., who found the component I to be maximal in this position. The conduction velocity of the sciatic nerve (CVs) in Approach I is quite near the corresponding conduction velocity of the sciatic nerve, CVs, in Approach III, but the central latency index (CLI) is much higher than the respective values in Approaches II and III (CLIF and CLIn). This difference may be due partially to the different subjects, but this is unlikely to be the only cause. More study is needed to explain this point.

In Approach II the latency at the onset of scalp response (31.9 ms) corresponds well with the results reported by Jones and Small5 and Dorfman.4 The differences are even slighter if the results are correlated to body lengths (B1) of controls (lat No/B1); the value of 18.6 msec/m in the present study compared with 17.6 ms/m obtained by Dorfman and Bosley.26 The observed lat P1 (36.8 ms) is about 3 ms less than reported by Jones,5 but that difference may be due to the differences in the body lengths of the controls. The latency of the F response (lat F) (51.0 ms), is quite similar to the value reported by Dorfman.4 When calculating the CLF values the scatter was large (SD 26% of the mean), while the distribution (expressed as SD) of CLIn in Approach III was only 11% of the mean. This is probably because H response is easy to produce, and its latency and amplitude are much more stable than those of F response, which in the present study changed very much even during the same experiment (see also Fra and Brignolio19). In Approach III lat No (25.0 ms) and lat P1 (29.0 ms) were in the same range as the findings reported by Delbeke et al.6 The conduction velocity of the sciatic nerve and roots (CVR) (58 m/s), corresponds well with the values reported by Vecchierini-Blineau and Guiheneuc.20 The central latency values (CLs) and the corresponding central latency indexes in Approaches II and III, (CLIF and CLIn) were, in theory, analogous values since they involved the same subjects. However, the values CLF and CLIn were statistically significantly different. This is due partially to the fact that the F response only tests the efferent neural pathways, but the H response estimates the average conduction time in both the afferent and efferent neural pathways, and it is known that conduction velocity in afferent proprioceptive paths is faster than in efferent motor pathways.19

Measurements of the evoked potentials at the lumbar levels using surface electrodes seems to be of limited value. In persons with relatively thick back muscles and subcutaneous fat, it is only with great difficulty that responses may be recorded with surface electrodes, and if a patient has difficulty in relaxing his back muscles the signal–noise ratio is low. Since the responses with skin electrodes are scarcely detectable, the differentiation of lesions at the brain, cervical and thoracic levels of the spinal cord is difficult with only stimulation of the peroneal or tibial nerve without invasive intrathecal or epidural recording.25 Dorfman4 has resolved this problem as follows: the conduction time at the thoracic level may be calculated as the difference between the latency of the scalp response for tibial nerve stimulation and conduction time in the cervical spinal cord calculated from the scalp and cervical responses of the median nerve. This method makes it possible to calculate indirectly the conduction velocity of the thoracic cord; however, the scatter of individual conduction velocities with this method is quite large (SD 18% of the mean). The method described in this paper is similar to that reported by Dorfman.4 In case 2 the conduction time of the cervical cord was evaluated from the scalp and cervical responses induced with the stimulation of the median nerve from the wrist; normal responses were found on the scalp and the seventh cervical segment, but the responses were lacking on the scalp after stimulation of the tibial nerve from popliteal fossa. These findings suggested a lesion site at the level of thoracic spinal cord. This was confirmed with myelography. In case 1, the responses of the skin recordings from the level of the second lumbar spine were scarcely detectable, but the recordings with platinum iridium wire electrodes from the epidural space at the level of the third lumbar segment showed clear asymmetry of the responses. The configuration of the response induced by stimulation of the left tibial nerve was similar to that reported by Shimoji et al., but the response induced with stimulation of the right tibial nerve was a flat dispersed potential. The
epidural recording technique is perhaps a good aid in cases where the skin recordings technique proves inadequate and it carries less risk of infection than the subdural technique.

Case 3, with MS showed scalp responses of prolonged conduction time (lat No) but in the periphery the CV values from both sides were normal. This suggests a decreased conduction velocity in the central nervous system, but normal conduction velocity in peripheral nerves, a typical finding in MS reported by many investigators.27

The present results suggest that the evaluation of the neural conduction time in the brain and spinal cord is more difficult than that of the peripheral nervous system. The measurement of the evoked potentials from the lumbar levels, the F response and H response, may overcome this problem. The H response method seems to be the best of these three; it is easy to perform even in persons with thick back muscles and subcutaneous fat; and statistically it is more stable than F response recording. In radicular lesions however, the recording of the evoked potentials with wire electrodes from the epidural space may offer a useful diagnostic tool for clinical practice.

References

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