Mechanisms of pain relief by vibration and movement

Ryusuke Kakigi, Horoshi Shibasaki

Abstract
Mechanisms of pain relief induced by vibration and movement were investigated. A CO₂ laser beam, which is useful for pure nociceptive stimulation, was used for recording pain-related somatosensory evoked potentials (pain SEPs) and for measuring pain threshold and reaction time (RT). Concurrently applied vibratory stimuli to and active movements of the fingers significantly reduced and prolonged pain SEPs, increased pain threshold, and prolonged RT, indicating that an increase in the inhibitory mechanisms of painful feeling was induced by the concurrently adopted sensory inputs mediated by large myelinated fibres. In contrast, continuous cooling enhanced pain SEPs and decreased pain threshold, probably due to the spatial summation of two kinds of nociceptive impulses mediated by the same pathways. The results of this investigation throw light on the mechanisms of the alleviation of pain by vibration and movement.

One of the main hypotheses for the gate control theory reported by Melzack and Wall¹ ² is that afferent signals which are mediated by large myelinated fibres inhibit small pain fibres presynaptically in the dorsal horn of the spinal cord. This hypothesis is supported by the analgesic effect of transcutaneous electrical nerve stimulation (TENS) of the peripheral nerve.³ ⁴ There has been no method of evaluating its effect objectively, and quantitatively, however, owing to the lack of a method of applying pure thermal or painful stimuli while recording the responses from the CNS in humans.⁵ Golding et al⁶ and Nardone and Schieppati⁷ reported the effects of TENS on waveforms of electrically-stimulated somatosensory evoked potentials (SEPs) for elucidating analgesic effects by TENS, but electric stimuli are not appropriate for the objective described above.

We studied SEPs induced by painful but tolerable CO₂ laser beam (pain SEPs),⁸ ¹³ and proved that ascending sensory signals induced by the beam are mediated by A delta fibres and the spinothalamic or spinoreticular tracts. Pain SEP findings significantly correlate with an impairment of pain-temperature sensation. We analysed the effects of various interference stimuli such as vibration, movement, touch, and cooling on the pain SEPs, pain threshold, and reaction time (RT) to the painful stimuli induced by CO₂ laser stimulation. We aimed to confirm, by using objective methods, the analgesic effect by afferent signals mediated by large fibres and to elucidate its underlying mechanisms.

Subjects and methods
Fifteen normal volunteers, 11 women and four men, were studied. Their ages ranged from 21 to 36 years with the mean age of 25. Their height ranged from 150 to 174 cm (mean 158). No medication was given for sedation, and subjects were kept awake. Each subject gave informed consent. A special CO₂ laser stimulator for recording SEPs was made by Nippon Infrared Industries (Kawasaki, Japan). Its maximum power was 12.5 W, and the stimulus intensity could be continuously changed. The laser wavelength was 10.6 μm, the diameter of the irradiation beam was about 2 mm, and the stimulus duration was 20 msec. We adopted an intensity of approximately 18 mJ/mm² which elicited sharp pain that all subjects described as tolerable, “like a pin-prick”. To avoid habituation irradiation points were moved slightly for each stimulus. The laser beam was applied to the part of the dorsum of hand innervated by the radial nerve once every 3 seconds. Subjects’ eyes were protected by swimming goggles.

Silver disc electrodes (1 cm diameter) were attached to the scalp with collodion and filled with electrode jelly based on the international 10–20 system. An impedance was maintained at less than 3 kΩ. Three exploring electrodes were placed at Cz, C₁', and C₂' (2 cm behind C₁ and C₂, respectively). The latter two sites corresponded to the hand sensory area of each hemisphere. Linked ear lobes (A₁ + A₂) were used as the reference. The amplifier frequency response was 0.5–30 Hz (-3dB). The analysis time was 1 second, and the sampling rate was 1976 ms. Peak latency and amplitude were measured for each recognisable component by a computer cursor. Amplitude was measured from the preceding peak of the opposite polarity. Relative positivity at grid 1 resulted in a downward deflection. Interfering stimuli were continuously applied to the fingers of the same hand throughout each recording session of pain SEPs as follows. (1) Tactile stimulation: the dorsum of the 2nd and 3rd fingers was continuously stroked by the experimenter with a soft wad of tissue paper. (2) Vibration: vibratory stimulus with a frequency of approximately 500 Hz was applied to the dorsum of the 2nd and 3rd fingers with a battery powered...
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Results

PAIN SEP FINDINGS

A small negative and a large positive potential, termed N1 and P1, respectively, were identified in the “control” and each “interference” trial in all 15 subjects. As the responses recorded at the Cz electrode were much larger than those at C3′ or C4′ (figure 1) the former were analysed. The mean peak latencies of N1 and P1 in the “control” were 205-6 and 302-5 ms, respectively, and their amplitudes were 2-23 μV and 8-07 μV, respectively (table 1). Amplitude of both N1 and P1 recorded at C4′ was not significantly different from that at C3′.

Peak latencies of N1 and P1 were significantly prolonged in “movement” and “vibration” interference (table 1 and figure 2). Amplitude of both N1 and P1 was decreased by “movement” and “vibration” interference, and a change of P1 in both conditions was significantly large (p < 0-01; table 1 and figure 2). Amplitude change by “tactile” interference was not consistent, amplitude of both N1 and P1 was increased by the “cooling” interference, particularly P1 (p < 0-01; table 1 and figure 2). Wave form changes were also identified in each interference condition recorded at the C3′ and C4′ electrode, but their degrees were much smaller than those at the Cz. For example, the percentage of the amplitude decreased by the “movement” interference was 18-9%, 40-3%, and 20-0% at the C3′, Cz, and C4′ electrode, respectively.

PAIN THRESHOLD

The mean (SD) of pain threshold in “control” was 13-6 (0-5) mJ/mm². Pain thresholds in “movement” and “vibration” interference conditions were the same with or higher than that of “control” in all subjects, and their changes were significant, particularly for “vibration” interference (table 2). In contrast, pain threshold in the “cooling” interference condition was the same as or smaller than that of “control” in all subjects, and the difference between them was significant (table 2).

REACTION TIME (RT)

The mean (SD) RT to the CO₂ laser stimulus in the “control” condition was 334-7 (27-2) ms. It was increased in all “interference” conditions except “cooling”, and the change in “movement” and “vibration” interference was significant (table 3). The mean (SD) RT to electrical stimulation applied to the median nerve at the wrist was 170-0 (30-2) ms. The difference between the peak latency of N1 and RT (N1–RT) was also measured; that in “movement” and “vibration” interference conditions was significantly longer than that in the “control” condition (table 3). N1–RT in the “tactile” and “cooling” interference conditions was shorter than that in the “control” condition, but the changes were not significant. As the peak latency of P1 was longer than RT in several subjects, P1–RT was not measured.

Figure 1 Averaged wave forms of pain SEPs recorded at C3′, C4′, and Cz electrode after CO₂ laser stimulation to left hand with no interference (control wave form) in normal subject. Linked ear lobes were used as reference. Traces on top are superimposition of eight recordings at Cz electrode.
Discussion
A CO₂ laser beam applied in an appropriate condition seems a good method for analysing pain mechanisms because it provides a pure thermal and painful stimulation without causing mechanical distortion of the skin and it is possible to trigger other instruments with no time lag. A few laboratories have also studied pain SEPs by using CO₂ laser in normal subjects and reported similar wave forms to those of ours.¹⁷⁻¹⁹

As both N1 and P1 are maximal around the vertex and symmetrically distributed, as reported in our previous reports,¹⁸⁻¹⁹ and as in this study, the interference effects at Cz are much larger than those at C₁ or C₂; the generator source of N1 and P1 may be in the thalamus or both parietal lobes rather than the primary sensory cortex. The second sensory cortex (S II) and the cingulate gyrus should also be considered as the main sites for pain perception. The former was proposed by Hari and her colleagues by results of magnetoencephalogram after painful dental stimulation.²¹ The latter was proposed by Talbot et al.²² by analysing positron emission tomography during noxious thermal stimulation.

We reconfirmed the presence of pain relief by vibration and movement, which is consistent with the gate control theory reported by Melzack and Wall.¹ ² Unfortunately, it is impossible to record clear pain SEPs from the peripheral nerve or the spinal cord probably due to the small S/N ratio.²³ Therefore, the site where the interference effects are mainly caused and its underlying mechanisms have to be estimated from other information. It is, of course, possible that the interference occurred in the dorsal horn of the spinal cord as proposed by Melzack and Wall.²⁻² In that case, the innocuous vibratory stimuli or movements reduce the excitability of laser-activated dorsal horn neurons, thus reducing the number of activated spinthalamic tract neurons and simultaneously reducing the synchronicity of the active volley reaching the cerebral hemisphere. This would both reduce the amplitude and increase the latency of N1 and P1 and elevate the pain threshold.

It is also possible that the interference occurred in the cerebral hemisphere. If RT was represented by a loop in which the evoked potential latency represented the afferent limb and the difference between the evoked potential latency and RT represented the efferent limb the significant increase in the time interval (N1–RT) by movement and vibration interference suggest that the interference effects were caused in the cerebral hemisphere. The more pronounced degradation of the second potential (P1) compared with that of

Table 1 Latencies and amplitudes of pain SEPs recorded at Cz without (control) and with various interference

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>Tactile</th>
<th>Movement</th>
<th>Vibration</th>
<th>Cooling</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean (SD) peak latency (msec)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N1</td>
<td>205.6 (24.6)</td>
<td>216.8 (31.3)</td>
<td>224.4** (30.8)</td>
<td>232.1** (25.7)</td>
<td>212.3 (23.1)</td>
</tr>
<tr>
<td>P1</td>
<td>302.5 (18.6)</td>
<td>310.3 (19.0)</td>
<td>317.7* (34.4)</td>
<td>336.5** (28.9)</td>
<td>312.5 (18.6)</td>
</tr>
<tr>
<td>Mean (SD) amplitude (μV)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N1</td>
<td>2.23 (0.92)</td>
<td>2.09 (1.03)</td>
<td>1.87 (1.11)</td>
<td>1.89 (1.54)</td>
<td>2.47 (1.48)</td>
</tr>
<tr>
<td>P1</td>
<td>8.07 (2.74)</td>
<td>9.18 (4.01)</td>
<td>4.82** (2.33)</td>
<td>6.28* (2.80)</td>
<td>10.50** (4.53)</td>
</tr>
</tbody>
</table>

Significance of difference between control and each interference trial calculated by paired r test (*p < 0.02; **p < 0.01).
the primary potential (N1) also seems to indicate the cerebral hemisphere as the responsible site for the interference effects. Several hypotheses may account for this particular phenomenon. If some neurons in the thalamus or the cerebral cortex receive inputs not only from nociceptors through small fibres but also from mechanoreceptors through large fibres, interactions are expected to take place. Mechanisms underlying the inhibition of electric SEPs by “movement”, “vibration”, or “tactile” interference, so-called “gating”, 1.4-16 23-30 are generally explained by this hypothesis. In addition, we propose two hypotheses in relation to the cognitive process as a higher function of the CNS. The first is that humans cannot completely differentiate painful stimulation from sensory stimulation of other modalities which are applied concurrently, and tend to neglect unpleasant sensations. The second is that painful sensation is attenuated by an attention to ascending signals of other modalities such as vibration. The effects of “tactile” interference were fairly small compared with those of active movement or vibration interference, suggesting that the signals mediated by small fibres are not attenuated as much as those mediated by cutaneous sensory fibres as by movement or vibration. 

A prominent tactile interference effect on electric SEPs 1.4-16 has to be explained by a different mechanism underlying the interference between them. It is difficult to elucidate the underlying mechanisms of the contrary effect by the cooling interference. Continuous stimulation of the nociceptors by cooling during the superimposed laser stimulation might increase the excitability of the appropriate dorsal horn neurons and then that particular phenomenon might have the effects of increasing the synchronicity of the volley and the number of spinothalamic tract fibres producing an increased amplitude of N1 and P1 and a reduction of the pain threshold. The fact that painful sensation caused by injury in cold weather is felt more than that in warm weather might be compatible with this particular finding.

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### Table 2 Pain threshold (mV/mm²) in “control” and each “interference” condition

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>Tactile</th>
<th>Movement</th>
<th>Vibration</th>
<th>Cooling</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean (SD)</td>
<td>13.6 (0.5)</td>
<td>13.9 (0.6)</td>
<td>14.5** (0.5)</td>
<td>14.9 (0.5)</td>
<td>15.1** (0.4)</td>
</tr>
</tbody>
</table>

Significance of difference between control and each interference trial calculated by paired t test (*p < 0.02, **p < 0.01).

### Table 3 Reaction time (RT) (ms) to CO2 laser stimulation and the time interval between N1 of pain SEP and RT in “control” and each “interference” condition

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>Tactile</th>
<th>Movement</th>
<th>Vibration</th>
<th>Cooling</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean (SD)</td>
<td>334.7 (27.2)</td>
<td>341.6 (29.8)</td>
<td>372.0** (37.7)</td>
<td>398.7** (57.3)</td>
<td>323.0 (33.4)</td>
</tr>
<tr>
<td>Mean (SD) N1-reaction time</td>
<td>129.1 (30.2)</td>
<td>124.8 (41.2)</td>
<td>147.6* (30.5)</td>
<td>166.7** (58.7)</td>
<td>111.7 (25.7)</td>
</tr>
</tbody>
</table>

Significance of difference between control and each interference trial calculated by paired t test (*p < 0.02, **p < 0.01).

Di Difference between peak latency of N1 and reaction time.
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