Thermal sensitivity is not changed by acute pain or afferent stimulation

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SUMMARY The effect of conditioning stimulation on thermal sensitivity and clinical pain was studied in 40 patients and six healthy subjects. Thresholds regarding cold, warm and heat pain perception did not differ significantly between the painful and non-painful skin areas in patients or between patients and healthy subjects before stimulation. The patients received either 100 Hz TENS, 2 Hz TENS, 100 Hz vibration, or placebo. No significant changes in thermal sensitivity were observed during and after conditioning stimulation in any of the test groups, although 24/40 (60%) of the patients reported reduction of their clinical pain intensity. The results indicate that (a) thermal sensitivity is not influenced by the presence of clinical pain, (b) the effects of stimulation on thermal sensitivity (thresholds) and clinical pain are not closely related, (c) central inhibitory effects of TENS and vibration are crucial for their pain relieving capacity.

Several studies have been performed to elucidate the pain reducing effect of transcutaneous electrical nerve stimulation (TENS) in experimentally induced pain in healthy volunteers as well as in pain patients. Few studies have actually compared the effect of afferent stimulation on experimental and clinical pain in the same patient (cf. ref. 1). Such a comparison seems important since studies on experimental pain in healthy subjects and pain patients have yielded some conflicting results (cf. ref. 1).

In the present study we have compared the effect of different types of stimulation, that is, low and high frequency TENS as well as vibration and placebo, on both experimental and clinical pain. Earlier reports on TENS2 and dorsal column stimulation (DCS)3 have dealt with long-standing chronic pain. We have concentrated our efforts on patients with acute pain of short duration.

First, we have observed to what extent on-going clinical pain influences temperature and pain perception, intra- and extrasegmentally with regards to the origin of the pain. This might be of interest regarding the development of sensory abnormalities due to interaction between activity in different sets of nerve fibres subserving various sensory modalities.4

Secondly, we have tried to find out if there is a correlation between the effect of afferent stimulation on experimental versus clinical pain. If such a correlation exists, it could be used in a predictive manner in pain treatment with methods such as TENS.

Material and methods

The study was carried out on 40 patients, 23 males and 17 females aged 20–58 years and on six pain-free subjects, three males and three females, aged 29–47 years.

Clinical pain

Patients were admitted to an emergency clinic for dental and oral surgery due to acute pain from teeth and/or surrounding tissues. The pain was due to pulp inflammation, apical periodontitis, pericoronitis or postoperative pain following operative removal of an impacted tooth. In all cases pain was ipsilaterally perceived corresponding to the trigeminal area innervating the area of the inflammatory lesion. The affected tissue was innervated either by the maxillary (13 cases) or the mandibular (27 cases) division.

The patients had suffered pain for 1–4 days. No patient had taken any analgesics within at least 10 hours before experimental procedures. All patients reported constant pain, that is not varying more than ±10% of its intensity over the hour.

All patients were examined, told their diagnosis and asked if they would take part in the experiments. If they agreed to participate, they were informed about the experimental pro-
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cedures in general, and that they could stop the procedures at any moment they desired. They were also told that they would get conventional dental treatment following the test session.

The subjects were informed that they might experience pain alleviation, no change or pain aggravation during stimulation. Care was taken to avoid suggestion.

Subjects were assigned to one of five groups: (1) vibration 100 Hz, eight patients, (2) placebo vibration, five patients, (3) 100 Hz TENS, 11 patients, (4) 2 Hz TENS, 11 patients, and (5) placebo TENS, 5 patients.

All patients rated their pain intensity initially using a graded five level verbal scale: light, light-moderate, moderate, moderate-severe and severe pain; and a visual analogue scale (VAS) both initially and following stimulation. The VAS consisted of a 10 cm horizontal line on a card. The words “no pain” and “worst pain ever” were placed on the left and right extreme ends of the line, respectively. Patients were instructed to mark the line at a point representing their pain.

During the experimental session each patient continuously rated his/her pain intensity using a graphic rating scale (GRS) consisting of a lever attached to a potentiometer controlling the position of a pen on a chart-recorder out of sight of the patient. The patients were instructed to move the lever from zero position (indicating pain intensity before starting stimulation) to one side when pain was reduced (endpoint = 10, meaning no pain) and to the opposite side if pain increased in intensity (endpoint = 10, meaning worst pain ever). Evaluation of the obtained pain reduction during stimulation was performed using the GRS for continuous registration. The values obtained with the VAS and GRS were consistent.

Thermal sensibility

All patients, regardless of the kind of conditioning stimulation used, were tested with respect to warm-cold, and thermal heat pain perception.

Thermal stimulation was delivered to the skin using a feed-back controlled thermode, consisting of four Peltier elements with an interposed thermocouple, stimulus surface 10 x 10 mm. The output current from the thermocouple measuring the temperature of the skin at the stimulus/probe interface was amplified and fed to the control unit for the Peltier elements. The side of the Peltier elements not facing the skin was cooled/heated by circulating water through a small chamber attached to this side with thermally conducting epoxi. The stimulating surface of the Peltier elements could be heated or cooled depending on the direction of current through the elements, with a temperature change which was linear over the temperature range 20–60°C. Measurements were made with a temperature change of 0–8°C/s. All measurements were started from an adapting temperature of 34–35°C, the same in all tests for each individual. Measurements were made from recordings on a chart-recorder of the output from the thermocouple.

Warm-cold perception Warm and cold thresholds were measured using the technique described by Fruhstorfer et al. The thermode was applied to the skin and the patient was instructed to press the button of a hand-held switch at the first sensation of warmth; this reversed the current to the Peltier elements shifting the thermode temperature in the cooling direction. The patient was also instructed to press the button of the switch at the first sensation of cold thereby shifting the thermode temperature in the warming direction. This procedure was repeated several times until stable values were reached. Determination of warm and cold thresholds was made from the last three to five measurements and the means were taken.

Pain threshold The probe temperature was continuously increased until the subject reported the stimulus as painful, at which instant the current to the Peltier elements was reversed by the subject, using the hand-held switch, returning the thermode to start temperature. Pain thresholds were measured twice and the mean was taken.

Conditioning stimulation

Mechanical vibratory stimulation The vibrator (Bruel & Kjaer, 4806) was driven by sinusoidal pulses at 100 Hz. Stimulation amplitude was 400–800 μm. The disc-shaped probe, diameter 3 cm, was applied at right angles to the skin in an attempt to exert pressure on the underlying bone.

TENS (2 Hz and 100 Hz) The stimulator produced monophasic square wave pulses of 0.2 ms duration at 100 Hz or a 71 Hz pulse train (duration 84 ms) at 2 Hz. A pair of conducting rubber electrodes, each measuring 3 x 3 cm, was positioned on the skin overlying the painful area. The most distally placed electrode was always connected to the anode and the proximal electrode to the cathode. With 100 Hz TENS the stimulus intensity was set to give a tingling sensation and with 2 Hz TENS prominent muscular contractions. The TENS stimulation was never reported as painful.

Placebo (vibration and TENS) Placebo stimulation was accomplished by applying the vibrator probe or TENS electrodes in contact with the skin of the painful area as for active stimulation but without transmitting any actual vibratory or electrical stimulation. The general procedure was the same as for the patients receiving vibration or TENS except that the patients were informed that some people might not experience the stimulation.

General experimental procedure

Before starting conditioning stimulation the thresholds for warmth, cold and pain were assessed: (A) on the skin within the painful area distal to the vibrator probe and the TENS electrodes respectively (the distance between thermode and the distal electrode/probe varied between 2–4 cm); (B) on a corresponding contralateral area; (C) on the dorsal aspect of the hand, ipsilateral to the painful side, in order to study extrasegmental effects.

Warm-cold thresholds and heat pain threshold measurements were made at separate spots, and the interval between successive stimulations at a certain spot was in general 15 min in order to avoid skin sensitisation or suppression. The same areas were used for each test before, during, and after conditioning stimulation. Measurements were always made in the order of warm-cold and pain threshold. Next, conditioning stimulation (CS) was started. Fifteen minutes later, during CS, the same measurements were repeated. After a total time of 30 min CS was terminated. Fifteen min later measurements were repeated.
Control group
In order to compare the influence of CS on perception of experimental stimulation in patients having clinical pain with that of pain-free subjects, six healthy normal volunteers were tested with the same types of CS and experimental parameters as those described above.

Statistical analysis
Analysis of temperature thresholds before, during, and after stimulation was made with a general analysis of variance, ANOVA. The ANOVA was made with conditioning stimulation, skin areas, and time as fixed factors. To approximately fulfill the assumption for the ANOVA, threshold values were transformed to logarithms before the analysis. Comparison of the number of patients reporting pain reduction using the different methods were made using the chi-square test. The significance level \( p < 0.05 \) was considered significant in the statistical analysis.

Results

**Temperature and pain perception** Mean temperature thresholds for perception of cold, warmth, and heat pain in patients and normal subjects at different skin areas before receiving conditioning stimulation are seen in fig 1. The mean values of the thermal thresholds for eliciting cold, warmth, and pain sensations were clearly separated in both painful and non-painful areas in patients as well as in the normal subjects. Mean difference limens (the warm-cold threshold difference) were in patients 6.7–6.9°C and in the normal subjects 4.8–5.5°C. No significant differences were found regarding painful versus non-painful skin areas of patients, in patients versus normal subjects, or between normal subjects in any of the measured variables. Thermal stimulation did not produce any dysesthetic sensations.

**Effects of afferent stimulation on temperature perception and on heat pain thresholds** During and after afferent stimulation small threshold changes (less than 3°C) were observed, both in patients and normal subjects, as illustrated in fig 2. No significant effects were observed with the different types of conditioning stimulation either in patients or in normal subjects. For comparison with all data before, during, and after afferent stimulation the data were therefore pooled, disregarding the type of conditioning stimulation used. This did not change the results of the statistical analysis in any significant way. In fact, the group mean values usually varied 0.1–0.3°C, and were always less than 0.8°C during and after stimulation as compared to initial measurements.

Fig 1 Average cool, warm and heat pain thresholds for all patients (left side diagrams) and pain-free subjects (right side diagrams) in different skin areas. Vertical bars represent 95% confidence limits. \( I \) = skin area innervated by trigeminal branch carrying the pain (patients) or corresponding skin area in subjects. \( C \) = contralaterally located skin area, corresponding to painful area. \( H \) = dorsal aspect of the hand, ipsilateral to the painful side.

Fig 2 Mean values regarding cold, warm and heat pain thresholds in the painful area (ipsilateral area) in patients (solid line) or a corresponding area in normal subjects (broken line). Vertical bars represent 95% confidence limits. Measurements before and after conditioning stimulation represents mean values of all patients \((N = 40)\) and normal subjects \((N = 6)\). Values during stimulation refer to effect of \( V \) = vibration; 2 = 2 Hz TENS; 100 = 100 Hz TENS; \( P \) = placebo.
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![Graph showing number of patients with different degrees of pain](image)

Fig 3 (A) Number of patients with different degrees of pain (L = light pain, M = moderate pain, S = severe pain) who received 100 Hz vibration (V), 100 Hz TENS (100), 2 Hz TENS (2), placebo-TENS or vibration (P). Patients who experienced pain reduction exceeding 50% represented by hatched areas. (B) Effects of conditioning stimulation on subjective pain intensity (a) pain increase; (b) no change in pain intensity; (c) pain reduction less than 50%; (d) pain reduction exceeding 50%.

No correlation was found between a change in clinical pain intensity and a change in cold, warm and heat pain thresholds during or after conditioning stimulation.

Clinical pain The distribution of patients having different pain intensities prior to stimulation, as rated on the verbal scale, is given in fig 3A showing a roughly even distribution.

The effect of TENS, vibration, and placebo on subjective pain intensity is seen in fig 3B. Of all patients tested, three reported complete pain relief (one patient receiving vibration and two patients receiving 100 Hz TENS). No type of active stimulation was superior to the others concerning number of patients reporting pain reduction. Placebo was significantly less effective than active stimulation.

Discussion

Thermal sensitivity before conditioning stimulation The cold and heat pain thresholds both in patients and in pain-free subjects were in the same range as reported by others, although warm thresholds tended to be somewhat higher than those previously described. This difference in findings might be due to the difference in stimulating area of the thermode, in this study 10 x 10 mm as compared with 25 x 25 mm used by others, since thermal thresholds depend on spatial summation.

The findings that thresholds did not differ in various skin areas in patients or subjects, or between patients and subjects indicate that the ongoing acute pain did not influence thermal sensitivity significantly. Lindblom and Meyerson did not find any relationship between chronic pain and perception of mechanically induced cutaneous pain, except in one patient with an abnormally low pain threshold in a hyperaesthetic area. Ischaemic pain has, however, been reported to elevate dental pain thresholds and decrease (non significant) thermal sensitivity in healthy subjects.

Thermal sensitivity during and after conditioning stimulation The present finding that TENS and vibration did not change thermal sensitivity, even in the painful and stimulated area, is in agreement with reports on the effect of DCS and intracerebral stimulation on experimental pain in chronic pain patients. A comparison between those former and the present results indicates that the duration of pain does not seem to be a crucial factor in determining the degree of interaction between pain and thermal sensitivity or the influence of TENS and vibration on activity in those systems. It seems that there is a rather secure transmission of the thermally induced activity from the skin. However, in dysesthetic skin areas a parallel increase has been reported between mechanical pain thresholds and reduction of chronic pain.

The finding that conditioning stimulation did not affect skin heat pain thresholds but did reduce clinical pain is interesting since in both areas activity in small-diameter nerve fibres transmitted to similar second order neurons is thought to be responsible for the perceived sensations (cf. ref. 17). Possible explanations for this discrepancy could be the following.

First, the peripheral site of origin might be of importance. The temperature stimulation was given to the skin whereas the clinical pain originated more deeply and was probably mediated by activity in deep somatic afferents. The present results could reflect a difference in susceptibility to central modulation induced by TENS and vibration on input from small diameter cutaneous and deep somatic afferents, as proposed by Woolf. Thus, TENS and vibration have been shown to increase the tooth pain threshold, and high-frequency TENS to reduce ischaemic pain in normal subjects but negative findings have been reported using cutaneously applied mechanical and thermal stimuli.

Secondly, a difference in impulse pattern in the fibres activated by the thermal stimulation as com-
pared with the one set up by the pathological process might also be of importance. The thermal stimulation with a rather rapid change in temperature would be expected to give a more synchronous activation of afferent nerve fibres than that seen during the constant clinical pain. In a previous study two types of painful stimulation, probably creating different afferent temporal and spatial input patterns, were differently susceptible to TENS and vibration, that is, acute orofacial pain due to pathology but not the pain induced by operative procedures was reduced by TENS and vibration.

It has been suggested that at least part of the pain-reducing effect induced by TENS is caused by a peripheral blockade or fatigue of pain transmitting fibres although this has been rejected by Janko and Trontelj. The present finding that stimulation had no effect on the thermal thresholds but diminished clinical pain would argue against such peripheral effects of TENS considering local anatomy. No reports exist demonstrating nociceptive afferents with peripheral divergences innervating both the tooth pulp, the surrounding tissues as well as cutaneous areas of the face. Such afferents would be a prerequisite to induce direct peripheral effects by TENS.

The present data also argue against distraction by the afferent stimulation as an alternative explanation for its pain reducing effect. The paraesthesia induced by the conditioning stimulation disappeared shortly after termination of stimulation, whereas pain reduction persisted for a period of 15 min to 5 hours.

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