The effects of transcranial electrical stimulation on opiate-induced analgesia in rats

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BACKGROUND AND OBJECTIVES: Recent experiments have shown that transcranial electrical stimulation significantly increases the potency and duration of the analgesic effects of opioids in humans and rats. In the present study, the influence of transcranial electrical stimulation (TCES) on the analgesic effect of remifentanil hydrochloride (HCl) in rats was determined.

METHODS: Experiments were performed on 80 albino male Wistar rats. Rats were randomly assigned to four groups: remifentanil HCl, remifentanil HCl and TCES, TCES, and control (n=20/group). Remifentanil HCl was injected on the 55th minute. Analgesia was assessed using the tail-flick latency test.

RESULTS: In the remifentanil HCl group, analgesia (10.85±1.04 s) was reached at the fifth minute, and the analgesia was high for the first 10 min. In the remifentanil HCl and TCES group, the latency time peaked (16.60±1.19 s) at the fifth minute. This peak was 150% higher than that for the remifentanil HCl group, and 251% higher than the control or TCES groups. Analgesia in the remifentanil HCl and TCES group was sustained for 20 min at a statistically higher rate than the other treatment groups (P<0.001).

CONCLUSIONS: TCES markedly increased the duration and analgesic potency of remifentanil HCl in rats. This effect appeared to be related to the release of enkephalins from brain structures, thus enhancing opioid analgesia.

Key Words: Analgesia; Remifentanil; Transcranial electrical stimulation

Les effets de l’électrostimulation crânienne sur une analgésie aux opiacés chez des rats

HISTORIQUE ET OBJECTIFS : Des expériences récentes démontrent que la stimulation électrique transcrânienne augmente considérablement la puissance et la durée des effets analgésiques des opiacés chez les humains et les rats. Dans la présente étude, l’influence d’une électrostimulation crânienne (ESC) sur l’effet analgésique de l’hydrochlorure de remifentanil (HCR) chez les rats a été déterminée.

MÉTHODOLOGIE : Des expériences ont été exécutées sur 80 rats Wistar mâles albinos. Les rats ont été divisés aléatoirement en quatre groupes : HCR, HCR et ESC, ESC et groupe témoin (n=20 par groupe). L’HCR a été injecté à la 55e minute. L’analgésie a été évaluée au moyen du test de latence de rétraction de la queue.

RÉSULTATS : Dans le groupe HCR, l’analgésie (10,85±1,04 s) a été obtenue à la 5e minute et a été élevée pendant les 10 premières minutes. Dans le groupe HCR et ESC, la période de latence a atteint un sommet (16,60±1,19 s) la 5e minute. Ce sommet était 150 % plus élevé que dans le groupe HCR, et 251 % plus élevé que dans les groupes témoin et ESC. L’analgésie dans le groupe HCR et ESC a été maintenue 20 minutes à un taux statistiquement plus élevé que dans les autres groupes de traitement (P<0.001).

CONCLUSIONS : L’ESC augmentait de manière considérable la durée et la puissance analgésique du HCR chez les rats. Cet effet semblait relié à la libération d’enképhalines dans les structures cérébrales, qui accroissait l’analgésie aux opiacés.
approximately 10 min to 20 min (7). Intraperitoneal remifentanil HCl reaches its peak effect in 5 min, and its duration of action is approximately 10±5 min in rats (8).

The aim of the present study was to determine the influence of TCES with Limoge’s current on the analgesic effect and duration of short-acting remifentanil HCl in rats.

METHODS

Animals

The present study was approved by the Committee for Research and Ethical Issues of Ankara University’s Veterinary Faculty (Ankara, Turkey). Experiments were performed on 80 albino male Wistar rats (Hifzisihha Institute, Ankara, Turkey) weighing 140 g to 180 g. Rats were housed on a 12 h:12 h alternating light/dark cycle with food and water ad libitum. The study rats had no prior exposure to electric stimulation or analgesics. A day before each experiment, the rats were placed in a transparent pliant box for 60 min to acclimate them to the experimental conditions. Experiments took place between 16:00 and 20:00. Rats were randomly assigned to four study groups, each consisting of 20 rats: the remifentanil HCl group (R), the remifentanil HCl and TCES group (R+TCES), the TCES group and the control group (C).

Electrode placement

Each rat was confined to a transparent pliant box designed to keep the rat’s head and tail free and out of the box. The box was perforated for injection into the abdomen. The rat was restrained to ease the conduct of the experiment. Venous needles were used to make the electrodes. The frontal electrode was placed on the cranium between the eyes on the metopic suture, and the two posterior electrodes were placed behind the mastoid bones on each side.

TCES

The stimulation was provided by a home-made generator calibrated with a digital oscilloscope (Multimeter, Hewlett Packard 973 A, USA). The electrical currents (Limoge’s current) consisted of low-frequency trains of high-frequency current (high-frequency intermittent bursts of bidirectional balanced currents: 166 kHz, low frequency; 4 ms at 100 Hz; current intensity: 100 mA). In each high-frequency cycle, the current was positive for 2 µs and negative for 4 µs (Figure 1). The frontal electrode was connected to the negative pole of the generator and received the negative impulse of weak intensity of 33 mA (long duration 4 µs). The two posterior electrodes were connected to the positive pole and received the positive impulse of 67 mA (short duration 2 µs) (3). The average intensity was zero, which completely eliminated electrode burns from electrolysis. A constant current circuit was added to maintain the same intensity whenever the electrodes interfaced impedance change. After the placement of the electrodes, the TCES and R+TCES groups were stimulated for 60 min. The stimulation was continued in on-and-off 5 min cycles until the end of the experiment. The electrodes were placed in all groups, but the R and control groups were not exposed to electrical stimulation.

Analgesia

Remifentanil HCl was dissolved in isotonic saline and injected intraperitoneally (1 µg/kg) with a 25 gauge needle (total volume of 1 mL) (Table 1). Groups R and R+TCES received remifentanil HCl on the 55th minute of stimulation, whereas the TCES and control groups were injected with only 1 mL of isotonic saline to equalize the painful stimulus effect of injection.

Measurement of analgesia

Analgesia was assessed using the wet tail-flick latency (TFL) test (9). The terminal third of the rat’s tail was immersed in a thermostat water bath (50°C), and the time required for initiating the retreat reflex time (seconds) was measured. For TFL measurements, the rats were disconnected from the stimulator for a short period. The time required for the retreat reflex was measured with a chronometer in the immersion TFL as previously described (10). The first tail movement imposed a time count interrupt. If the tail retreat reflex did not occur after 20 s, the tail was manually withdrawn from the water to avoid tissue damage.

The first TFL test was applied to all groups on the 60th minute of TCES, and then repeated every 5 min until the same values were measured in all groups.

To observe the effects of TCES on rat movements (head, muzzle and eye), rat reactions were monitored in the TCES group and compared with those of the control group.

TABLE 1

<table>
<thead>
<tr>
<th>Groups and experimental procedure</th>
<th>Remifentanil hydrochloride (R) or saline treatment</th>
<th>TFL</th>
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<tbody>
<tr>
<td>Group (n=20) TCES</td>
<td></td>
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<tr>
<td>R</td>
<td>– 1 µg/kg 5 min intervals</td>
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<tr>
<td>R+TCES</td>
<td>Limoge’s current (5 min) 1 µg/kg 5 min intervals</td>
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<tr>
<td>TCES</td>
<td>Limoge’s current (5 min) Isotonic saline 5 min intervals</td>
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<tr>
<td>Control*</td>
<td>– Isotonic saline 5 min intervals</td>
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</tbody>
</table>

*Drug and stimulation-free. TCES Transcranial electrical stimulation; TFL Tail-flick latency

Figure 1) Characteristics of transcranial electrical stimulation with Limoge’s current. (A) High frequency 166 kHz, 2 µs+, 4 µs and (B) intermittent low frequency 100 KHz, 4 ms

(3). Electrodes were connected to an electrical generator with thin and flexible cables.
Statistical analysis
Quantitative data was compared in terms of mean and SD of TFL. One-way ANOVA was used for the comparisons between and within the groups for time-dependent differences. The Dunnett test was done for comparisons with the control group. The Tamhane test was performed when comparing the period of non-homogeneous distribution (5 min to 15 min) between groups and the Takey HSD test on those with homogeneous distribution. All tests were performed using SPSS 11.5 (SPSS Inc, USA).

RESULTS
All results were expressed as mean ± SD. No major adverse effects were observed throughout the procedures.

The control TFL test value was 4.75±0.72 s at the fifth minute. The retreat reflex time in the TCES group was similar to the control TFL test result at fifth to 30th minute after the injection (Figure 2). TCES with Limoge's current did not modify the pain threshold during stimulation in the TCES group. Analgesia was increased by remifentanil HCl in the R group. In addition, remifentanil HCl-induced analgesia was increased by TCES.

In the R group, the retreat reflex time reached a peak value (10.85±1.04 s) at the fifth minute (60th minute of TCES), which was significantly longer than the TCES and control group values at the fifth and 10th minute (P<0.001). TFL test results at the 15th minute returned to control TFL values and were similar to the TCES and control group values.

The retreat reflex time reached a peak value (16.60±1.19 s) at the fifth minute (60th minute of TCES) in the R+TCES group; this value was five times greater than those of the TCES and control groups. In the R+TCES group, the fifth to 20th minute following remifentanil HCl injection, retreat reflex time was significantly longer than the other groups (P<0.001). TCES-induced analgesia with remifentanil HCl was observed for 20 min; however, a progressive decrease in analgesia with time was observed, and values reached the control TFL values by the 25th minute in the R+TCES group.

The TFL value of the R group was 131% higher than the TFL values of the control and TCES groups at the fifth minute (P<0.001). In the R+TCES group, this increase reached 251% compared with the control and TCES groups at the fifth minute (P<0.001). Furthermore, the analgesic potency for the R+TCES group increased by 150%, 162%, 124% and 93% compared with the R group at the fifth, 10th, 15th and 20th minute, respectively (P<0.001). The results suggested that remifentanil HCl-induced analgesia was potentiated by TCES. The analgesia in the R+TCES group decreased to control values by the 25th minute following remifentanil HCl administration.

Nonquantitative observations indicated that TCES did not affect the movements of the rats in the TCES group. The rat movements were similar in the TCES and control groups.

DISCUSSION
The R+TCES group presented a statistically significant nociceptive threshold above that of the R group. We found a positive correlation between TCES and analgesic effects.

Mathematical analysis of Limoge's current is based on the Fourier series. The presence of these electric fields in the brain may influence the endogenous opioid system (11). This is supported by the fact that TCES has been successfully used during opiate withdrawal in human heroin addicts without any adverse reactions (12). In addition, nonquantitative observations have indicated that TCES with Limoge's current does not affect the behaviour of drug-free rats (2,3,13,14). In the present study, TCES did not affect the movements of the rats in the TCES group, thus allowing for the administration of stimulation without side effects.

TCES produces a naloxone-reversible analgesia in drug-free rats that is not potent (13,14). This finding is supported by studies (12,15) that found that opiate withdrawal syndrome faded with TCES and that this effect could be reversed with naloxone. However, other studies (2,3) have shown that TCES does not modify the pain threshold in drug-free animals and humans, possibly due to the very low basal activity of central opiate systems. In our study, TCES alone did not modify the pain threshold in rats.

Stinus et al (3) observed that, in rats, morphine, fentanyl, alfentanil HCl and dextromoramide provided a 174%, 176%, 160% and 267% increase in TFL, respectively, compared with control values; these increases reached 306%, 336%, 215% and 392%, respectively, with the addition of TCES. In the present study, analgesia increased 131% in the R group compared with control TFL test results at the fifth minute. This increase reached 251% in the R+TCES group at the fifth minute with the addition of stimulation.

There are different perspectives concerning the starting time of stimulation to increase analgesic effects. The beginning of analgesia, the time required for maximal effect and the end of analgesia can differ due to the current used, the location of the stimulation, the technique used to measure analgesia and the duration of the stimulation. Stinus et al (3) reported that the analgesic effects of subcutaneous morphine were increased by starting TCES 3 h before injection. Different observers have successfully reached analgesia using TCES for less time. Analgesia has been shown to start on the 20th minute of low-frequency stimulation (10 Hz) in rats (13-16). In our study, we used TCES with high frequency and began the stimulation 5 min before drug administration. Analgesia with
TCES was high at the 60th minute (at the fifth minute of remifentanil HCl injection) and it remained high for 20 min in the R+TCES group. The analgesia lasted approximately two times longer when TCES was added to remifentanil HCl (compared with remifentanil HCl alone).

The mechanisms and neurobiological substrates of TCES remain unknown. TCES may alter the kinetics of the drug and change blood-brain barrier permeability, increasing the penetration of the drug into the brain. Perhaps it creates a new steady state in brain activity, which could be revealed by a pharmacological challenge. This phenomenon is likely centrally mediated and occurs via the enhancement of analgesia elicited by the release of endogenous opioids such as enkephalins and endorphins (1-6,16,17). Stimulation of the periaqueductal gray matter induces highly specific naloxone-reversible pain suppression that is associated with increased immunoreactivity for beta-endorphins in ventricular fluid (18,19). Enkephalinase inhibitors increase the analgesia caused by TCES. Opioid antagonists, such as naloxone, reverse this effect. It is believed that the endogenous opioids participate in this mechanism (2,3,12,15,17). On the other hand, TCES appears to cause an increase in the synthesis or release of serotonin, dopamine and noradrenaline in the midbrain, and dopamine and serotonin in the hypothalamus (20). Moreover, noradrenergic fibres (21), serotonergic fibres (22,23) or both (24) have been shown to play a key role in the medication of analgesics at the spinal level.

CONCLUSION

TCES markedly increases the duration and analgesic potency for remifentanil HCl in rats without affecting movements. This effect appears to be related to the release of enkephalins from brain structures, enhancing opioid analgesia.

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