Multicenter, randomized, double-blind, controlled trial of transcutaneous electrical nerve stimulation for pancreatic cancer related pain

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Abstract

Background: Up to 80% of patients with pancreatic cancer experience abdominal and back pain. Although pharmacologic medications provide some relief, many report inadequate analgesia and adverse effects. Transcutaneous electrical nerve stimulation (TENS) is a non-invasive physical modality and had been widely applied for pain relieving, yet no study has investigated the effectiveness of TENS for pain in pancreatic cancer.

Methods: Eligible patients were randomly assigned in a 1:1 ratio to TENS group or control group. The primary outcome was percentage change of numerous rating scale (NRS) after treatment. Secondary outcomes included percentage change of analgesic medication consumption and effect on constipation and poor appetite.

Results: One hundred seventy-one patients were recruited (84 to control group and 87 to TENS group). NRS in TENS group has been largely decreased 77.9% right after treatment and 27.1% in 2 hours, before applying any analgesic medication, while that in control group was slightly downregulated right after treatment but gave a trend to increase at 1, 2, and 3 hours. When comparing both groups, pain was significantly well controlled without analgesic medication supplement in TENS group at 0 hour (difference in mean percent change in NRS=134.0 [95% CI, 130.0–142.7], \( P < .01 \)) after treatment, and this analgesic effect last to 3 weeks after treatment cycle (difference in mean percent change in NRS=22.5 [95% CI, 17.6–27.3], \( P < .01 \)) without increase of analgesic medication consumption.

Conclusions: TENS reduces pain without increase analgesic medication consumption in patients with pancreatic cancer pain. It provides an alternative therapy for pain in pancreatic cancer.

Clinical Trial Registration: This study was registered at ClinicalTrials.gov, identifier NCT03331055.

Abbreviations: NRS = numerous rating scale, TENS = transcutaneous electrical nerve stimulation.

Keywords: pain, pancreatic cancer, transcutaneous electrical nerve stimulation

1. Introduction

Pain in pancreatic cancer has come up to 80%, with 50% to 70% suffering from severe pain.\cite{1} The experience of pain can either positively or negatively influence patient outcomes.\cite{2} Conventionally, non-steroidal anti-inflammatory agents and/or opioid analgesics are used for alleviating pain according to the pain...
management strategy recommended by World Health Organiza-
tion. However, there are still many patients suffered from 
refractory pain, which presents a big challenge to the physicians.
In addition, serious drug-related side effects bring more agony to 
many patients that can markedly reduce their quality of life.
Celiac plexus neurolysis, in which the celiac plexus is chemically 
ablated, has been widely performed as an alternative treatment 
for alleviating cancer-associated pain, but would finally leads to 
an intractable pain.

Transcutaneous electrical nerve stimulation (TENS) is a non-
invasive and easy operated modality. It is applied by transcuta-
neous (over the skin) electrical stimulation and is primarily used 
for pain control in a wide range of acute and chronic pain 
conditions. TENS units typically use adhesive electrodes 
applied to the skin surface to apply pulsed electrical stimulation 
that can be modified in terms of frequency (stimulation rate), 
tensity, and duration. It has been successfully applied in clinical 
treatment of various types of pain including: neuropathic pain, bone pain, postoperative pain, etc. Although its algentic 
mechanism is still unclear, electrical nerve stimulation results in 
consistent improvement of mechanical and thermal hyper-
algesia with reduction in the mechanism is still unclear, electrical nerve stimulation results in 
neurons, increased inhibitory input to the pain pathways at 
an intractable pain.

In addition, serious drug-related side effects bring more agony to 
refractory pain, which presents a big challenge to the physicians.

We conducted prospective, randomized, and sham-controlled 
trial to evaluate the efficacy and safety TENS for cancer related 
pain in pancreatic cancer patients.

2. Patients and methods

2.1. Patients

The protocol was approved by relevant ethics committees and 
institutional academical review boards of Fuda Cancer Hospital 
(2017-TCM-01) and was registered at ClinicalTrials.gov 
(NCT03331055). All participants signed consent forms when 
recruited. Patients were recruited in 4 sites from March 2016 
through March 2018. Key inclusion criteria were an age of 18 to 
70 years, primary or metastatic pancreatic cancer or liver cancer 
with cancer related visceral pain, no neuromytic celiac plexus 
block was done in the past 1 month, with anticipatory survival of 
more than 3 months, and normal lung and heart function. 
Exclusion criteria included who could not tolerate of maintaining 
30min of side position without movement, who has been 
recruited in other clinical trial for pain relieving, who underwent 
radiotherapy or local radiative seeds implantation in the past 1 
month, who imaging diagnosed with encephalic tumor or 
metastasis, who with cardiac pacemaker or metal stand, who 
with risk in portal or other embolism, who with not well-
controlled hypertension or diabetes.

2.2. Trial design

This randomized, single-blind, sham-controlled trial consisted of 
a screening visit, a 1-day pre-interventional analysis of numerous 
rating scale (NRS) and analgesic medication consumption 
baseline, a 1-week intervention duration, and a 4-weeks 
observation. On the basis of the screening visit and information 
collected in pre-interventional analysis process, patients were 
enrolled or were excluded if they were not eligible.

Eligible patients were randomly assigned in a 1:1 ratio of TENS 
group or control group. Randomization was performed by means 
of random number table, with stratification according to NRS of 
3 to 6 and >6. Points on 1.5 cm away from middle line of T8 to 
T12 vertebra (belong to acupoints of B3, BL18, BL19, BL20, and 
BL21 in traditional Chinese meridian theory system), RN12, and 
pain point on abdomen. On back, acupoints at the same level were 
attached a pair of electrical poles, while acupoints on 
abdomen were stimulated with another pair of pole. TENS group 
was applied electrical stimulation in 2/100Hz for 30minutes.

Tense was various and managed at the maximum comfortable 
critical points according to individual difference. Sham group 
was administered patches at the same acupoints which also 
attached electrical lines to electro-therapeutic apparatus but 
without electrical stimulation. Interventions were administered 
twice a day. Patients were unaware of the trial-group assign-
ments.

2.3. Primary and secondary outcomes

The primary study outcome was pain relief for each patient 
before painkiller applied, at 0, 1, 2, and 3 hours after each 
intervention, and at 1, 2, 3, and 4 weeks after treatment cycle, 
calculated as percent change in NRS at 0, 1, 2, 3 hours and 1, 2, 3, 
4 weeks, as compared with the patient’s baseline NRS. Secondary 
outcomes were percentage change in morphine use (expressed as 
the decrease of morphine equivalent consumption compared with 
baseline).

2.4. Complications

Complications were defined as any unplanned event considered 
related to TENS that required additional treatment after the 
procedure, and were recorded and classified in accordance with 
the Common Terminology Criteria of Adverse Events v4.0.

2.5. Sample size calculation

Estimations based on trial of acupuncture for cervical cancer 
pain of 64 patients with changing NRS from baseline of 
4.2 in acupuncture group and –2.2 in control group (P < .01), 
indicated that a sample of 168 patients (equally divided into 
control and treatment groups) would suffice to achieve 90% 
statistical power for detecting a significant greater decrease of 
NRS in TENS group than in control group at a two-sided 
significance level of 0.05. For these calculations, patients 
undergoing TENS were assumed to themselves experience a 
decrease of 2.2 in pain scores (compared with their baseline pain 
scores) due to a placebo effect. Aggregate attrition rate at the 2-
week mark (including both mortality and loss to follow-up) was 
assumed to be approximately 20%, therefore the total number of 
participants were 210.

2.6. Statistical analysis

Data were analyzed on an “intent to treat” basis conducted by 
using GraphPad Prism (San Diego, CA, version 6.0). Mean 
differences were expressed as coefficients in appropriate linear 
models and estimated using generalized estimating equations; this 
allowed the use of data from differing time points in the 
estimation procedure. CIs for these mean differences comparing 
outcomes between treatment groups were constructed using 
unpaired t test by assuming nonparametric test. Significant 
differences were indicated by P < .05 or P < .01.
3. Results

3.1. Patients recruitment

Between March 2016 through March 2018, a total of 254 patients were referred for this trial (Fig. 1). Eighty-three patients did not meet study entry criteria or refused to participate. One hundred seventy-one patients were randomly assigned, with 84 patients assigned to control group and 87 assigned to TENS group. Patients in both groups were comparable for all cogent variables (Table 1). All patients had locoregional disease. Patients who received TENS or sham TENS showed no evidence of early or late complications. Twenty-one patients were lost to follow-up (12 in the TENS group [mean baseline pain score 5.6] and 9 in the control group [baseline pain score 6.1]). All patients were included in the intention-to-treat analysis.

3.2. Primary outcome: percentage change in NRS

NRS in control group were tended to increase compared with baseline at 0 hour (mean percent change in NRS = −25.3 [95% CI, −26.7 to −24.0]), 1 hour (mean percent change in NRS = 26.2 [95% CI, 23.7 to 28.3]), 2 hours (mean percent change in NRS = 98.7 [95% CI, 95.1 to 102.3]), and 3 hours (mean percent change in NRS = 128.1 [95% CI, 124.6 to 131.7]) before taken analgesic medication after intervention, and was invariant at 1 week (mean percent change in NRS = −34.2 [95% CI, −34.2 to −25.9]), 3 weeks (mean percent change in NRS = −21.5 [95% CI, −26.2 to −16.8]), and 4 weeks (mean percent change in NRS = −131.7 [95% CI, 130.0 to 142.7]) after treatment cycle. When comparing both groups, pain was significantly well controlled without analgesic medication supplement in TENS group at 0 hour (difference in mean percent change in NRS = −30.0 [95% CI, 50.0 to 51.4], P < .01; Fig. 2A), 1 hour (difference in mean percent change in NRS = 63.0 [95% CI, 55.4 to 66.7], P < .01; Fig. 2A), 2 hours (difference in mean percent change in NRS = 116.7 [95% CI, 113.0 to 121.4], P < .01; Fig. 2A), and 3 hours (difference in mean percent change in NRS = 134.0 [95% CI, 130.0 to 142.7], P < .01; Fig. 2A) after intervention, and this analgesic effect last to 1 week (difference in mean percent change in NRS = 35.2 [95% CI, 32.1 to 38.3], P < .01; Fig. 2B), 2 weeks (difference in mean percent change in NRS = 30.6 [95% CI, 26.4 to 34.9], P < .01; Fig. 2B), and 3 weeks (difference in mean percent change in NRS = 22.5 [95% CI, 17.6 to 27.3], P < .01; Fig. 2B) after treatment cycle.

3.3. Secondary outcome: percentage change in morphine consumption

In the control group, morphine use (analgesic medication consumption in morphine-equivalent units) increased compared with baseline at 1 week (mean percent change in morphine consumption = 7.8 [95% CI, 5.2 to 10.5]), 2 weeks (mean percent change in morphine consumption = 8.1 [95% CI, 5.4 to 10.8]), 3 weeks (mean percent change in morphine consumption = 10.8 [95% CI, 8.3 to 13.3]), and 4 weeks (mean percent change in morphine consumption = 13.3 [95% CI, 10.8 to 15.8]) after treatment cycle. In contrast, in TENS group, morphine use (analgesic medication consumption in morphine-equivalent units) was invariant at 1 week (mean percent change in morphine consumption = −1.0 [95% CI, −0.5 to 2.4]), and 3 weeks (mean percent change in morphine consumption = −0.5 [95% CI, −0.2 to 2.3]) after treatment cycle.

Table 1: Baseline characteristics of all randomly assigned patients.

| Characteristic                               | Control   | TENS      |
|----------------------------------------------|-----------|-----------|
| No. of patients                             | 84        | 87        |
| Male sex                                    |           |           |
| %                                           | 57.1      | 49.4      |
| Age, yr                                     | 58.9      | 51.2      |
| SD                                          | 9.9       | 10.7      |
| Pain history, wk                            | 9.4       | 8.7       |
| SD                                          | 8.1       | 6.2       |
| Narcotic consumption, morphine-equivalent units | 44.8  | 39.7       |
| Mean                                         | 56.1      | 64.2      |
| Abdominal pain intensity, numeric rating scale | 5.4      | 5.8       |
| Mean                                         | 1.8       | 1.7       |
| Constipation                                | 26        | 33        |
| %                                           | 31.0      | 37.9      |
| Poor appetite                                | 48        | 46        |
| %                                           | 57.1%     | 52.9%     |

SD = standard deviation; TENS = transcutaneous electrical nerve stimulation.
weeks (mean percent change in morphine consumption $= 8.5$ [95% CI, 6.0–11.1]), and 4 weeks (mean percent change in morphine consumption $= 9.1$ [95% CI, 6.5–11.6]). In TENS group, morphine use also increased at 1 week (mean percent change in morphine consumption $= 6.6$ [95% CI, 5.4–8.4]), 2 weeks (mean percent change in morphine consumption $= 7.9$ [95% CI, 6.2–9.3]), 3 weeks (mean percent change in morphine consumption $= 7.3$ [95% CI, 5.9–9.5]), and 4 weeks (mean percent change in morphine consumption $= 7.5$ [95% CI, 5.9–9.1]). However, the differences between control group and TENS group were not significant at 1 week (difference in mean percent change in morphine consumption $= 2.2$ [95% CI, 1.0–5.4], $P < .20$; Fig. 3), 2 weeks (difference in mean percent change in morphine consumption $= 1.9$ [95% CI, 1.0–5.1], $P < .29$; Fig. 3), or 4 weeks (difference in mean percent change in morphine consumption $= 1.8$ [95% CI, 1.0 to 6.3], $P < .30$; Fig. 3), or 4 weeks (difference in mean percent change in morphine consumption $= 2.3$ [95% CI, 1.0 to 5.6], $P < .17$; Fig. 3).

No patient suffered constipation and poor appetite in control group was improved after treatment. Comparatively, in TENS group, all of 33 patients complained with constipation in TENS group told they had different degrees of improvements, and 42 out of 46 patients with poor appetite gain better appetite during and after treatment.

3.4. Safety

No side effect has occurred in any patient.

4. Discussion

TENS shows its benefits in reducing pancreatic cancer pain, which effect lasted till 3 weeks after treatment. More promising result is that TENS downregulated the NRS before analgesic medication had been applied within 2 hours after treatment procedure.

TENS has been widely studied and used for patients suffering cancer related pain with showing the potential to improve quality of life within specific types of cancer.[18] However, not all studies on TENS for cancer pain have the consistent results cause the sample size, study design, stimulation site, electric frequency, intensity, cycle frequency, method of administration, and outcome measures are various.[7,8] Hence in this study, we manage the methodological quality through methods of sample calculation by assuming result basing on a trial of acupuncture for cervical cancer pain, double-blind management, sham group design, etc. Besides these, core elements that would affect the results are mode of TENS, treatment frequency, and duration.

For electric frequency, even though some researches had gained negative results in pain relieving with TENS by using low electrical frequency or long treatment intervals (e.g., twice a week),[19–21] TENS has been used with varying success in analgesic treatment. Results from studies investigating the morphine-sparing effects of electrostimulation manifested that alternating-frequency (2/100Hz), high-intensity (9–12mA) stimulation of acupoints has >50% morphine-sparing effect in patients after lower abdominal gynecologic surgery.[22,23] Han and colleagues conducted a series of animal studies that have shown that low frequency TENS-induced anti-hyperalgesia (decreased sensitivity to pain) is mediated by enkephalin, b-endorphin, and endomorphin through d-opioid and m-opioid receptors, while high frequency TENS enhances dynorphin through k-opioid receptors.[24] But interestingly, it has been found that neither high nor low frequency alone could downregulate pain.[25] Similarly again, analgesic effect of alternating-
frequency TENS was proved in this study with showing significant downregulation after intervention without taking painkiller, and maintaining a lower NRS after treatment cycle in 4 weeks without increase of analgesic medication consumption compared with control group.

Moreover, intensity is also close related to the “pain relieving duration.” It had been proved stronger intensity can help to reach a longer duration of staying at a higher pain threshold, and this dose–effect relationship would contribute to a better pain management results. As it has been suggested the strongest comfortable intensity is normally work for pain in organs, this study conduct strongest but comfortable intensity for each individual rather than implement a fix intensity, which might cause a vast various of sense to different person, might be an important factor help reaching an obvious decrease right after treatment process and maintain the effect of pain relieving for 2 hours without applying analgesic medication.

For cycle frequency, it seems that it has been underestimated by many studies. However, from the results of this study, a peak of pain-relieving effect by 77.9% of downregulation of NRS right after treatment procedure, decrease to 27.1% in 2 hours after treatment before analgesic medication have to be applied to. Hence, we assume that 2 hours should be a maximum of rest duration to maintain the pain-relieving effects by TENS only. However, implement the TENS treatment for every 2 hours for patients, which mean 6 times a day, could be with very low patient compliance.

One week of continuous treatment proved to prolong the analgesic effect. When finishing a week cycle of treatment, average level of NRS seems to be maintained at a lower level without increase of analgesic medication consumption, of a 35.2% downregulation of NRS at 1 week and 22.5% at 3 weeks.

Despite other traditional standard therapy, such as intraoperative CPN, was proved to decrease 10% pain scores in pancreatic cancer patients at 1 month, TENS could benefit patients without invasion and too much additional economic burden.

In this initial research for intractable pancreatic cancer pain, we found that most patients’ pain was not well controlled due to various reasons, such as adverse function of analgesic medication of constipation and lost appetite. Without increase of analgesic consumption, pain score was significantly reduced after TENS treatment cycle.

However, even during the treatment period, the pain was not disappeared or maintain at a low level for all day long. Average 3 hours of downregulation effect on NRS might call for a higher cycle frequency to achieve a more stable analgesic effect. Moreover, a higher cycle frequency may also reduce the use of painkiller. Therefore, further research on exploring more therapeutic characteristics of TENS and setting up an optimal application model in analgesia treatment is needed. For this, we have designed a portable and wearable acupuncture electrical stimulation device (China patent: ZL201820740038.6), and will carry out further research in the future.

5. Conclusion

In final, we conclude TENS is safe and has significant effect on relieving pain in pancreatic cancer patients. In addition, it has many advantages such as non-invasive, seldom side effects, easy-operation, cheap, and with good patient compliance. Alternative frequency, maximum comfortable intensity, twice or more time treatment daily gave a model of TENS in pancreatic cancer pain treatment. TENS obviously may play a much more important role in dealing with cancer related pain in the future.

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