



Using TENS for pain control: the state of the evidence

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Practice points

- High frequency (HF) and low frequency (LF) transcutaneous electrical nerve stimulation (TENS) activate different opioid receptors. Both applications have been shown to provide analgesia specifically when applied at a strong, nonpainful intensity. HF TENS may be more effective for people taking opioids.
- Effective analgesia for chronic pain conditions may be limited by the development of tolerance to TENS if repeated application of either LF or HF TENS at the same frequency and intensity is used daily (i.e., same dose). Strategies to prolong analgesia may include varying these parameters.
- Application of TENS electrodes at acupoint sites may increase analgesia.
- Targeting the use of TENS during movement or activity may be most beneficial.
- Systematic reviews suggest that TENS, when applied at adequate intensities, is effective for postoperative pain, osteoarthritis, painful diabetic neuropathy and some acute pain conditions.
- Emerging evidence suggests TENS may be helpful for people with fibromyalgia and spinal cord injury.
- TENS may be effective in restoration of central pain modulation, a measure of central inhibition.
- A clearer picture of TENS effectiveness will emerge as trials with attention to optimal dosing and appropriate outcome measures increase in numbers.

SUMMARY: Transcutaneous electrical nerve stimulation (TENS) is a nonpharmacological intervention that activates a complex neuronal network to reduce pain by activating descending inhibitory systems in the central nervous system to reduce hyperalgesia. The evidence for TENS efficacy is conflicting and requires not only description but also critique. Population-specific systemic reviews and meta-analyses are emerging, indicating both HF and LF TENS being shown to provide analgesia, specifically when applied at a strong, nonpainful intensity. The purpose of this article is to provide a critical review of the latest basic science and clinical evidence for TENS. Additional research is necessary to determine if TENS has effects specific to mechanical stimuli and/or beyond reduction of pain and will improve activity levels, function and quality of life.

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Background

Transcutaneous electrical nerve stimulation (TENS) is an inexpensive nonpharmacological intervention used in the treatment of acute and chronic pain conditions. These small battery-powered devices deliver alternating current via cutaneous electrodes positioned near the painful area. The parameters of pulse frequency, and pulse intensity are adjustable and linked to TENS efficacy. This article will provide a critical review of the latest basic science and clinical evidence for TENS. We will summarize mechanisms of action, factors that influence TENS efficacy, and describe and critique the use of TENS for pain control in a variety of patient populations. Findings of systematic reviews of TENS for pain management in the last 7 years will be presented. We will also highlight advances from Randomized Controlled Trials (RCT) published in the last 5–7 years, which are not included in the systematic reviews. This article offers a concise review of the basic science mechanisms for TENS as well as an up to date critique of current clinical research for TENS.

Mechanisms of TENS reduction on analgesia

TENS activates a complex neuronal network to result in a reduction in pain. At frequencies and intensities used clinically, TENS activates large diameter afferent fibers [1,2]. This afferent input is sent to the central nervous system to activate descending inhibitory systems to reduce hyperalgesia. Specifically, blockade of neuronal activity in the periaqueductal gray (PAG), rostral ventromedial medulla (RVM) and spinal cord inhibit the analgesic effects of TENS showing that TENS analgesia is maintained through these pathways [3–5]. In parallel, studies in people with fibromyalgia show that TENS can restore central pain modulation, a measure of central inhibition [6]. Therefore, TENS reduces hyperalgesia through both peripheral and central mechanisms.

• Neurotransmitters & receptors that mediate TENS analgesia

HF TENS increases the concentration of β -endorphins in the bloodstream and cerebrospinal fluid, and methionine-enkephalin in the cerebrospinal fluid, in human subjects [7,8]. The analgesia produced reduction in hyperalgesia by HF TENS is prevented by blockade of opioid receptors in the RVM or spinal cord, or

synaptic transmission in the ventrolateral PAG [4–5,9]. This opioid-mediated analgesia produced by HF TENS has been confirmed in human subjects [10]. Furthermore, the reduction in hyperalgesia produced by HF TENS is prevented by blockade of muscarinic receptors (M1 and M3) and GABA_A receptors in the spinal cord [11,12]. However, blockade of serotonin or noradrenergic receptors in the spinal cord has no effect on the reversal of hyperalgesia produced by HF TENS [13]. Thus, HF TENS produces analgesia by activating endogenous inhibitory mechanisms in the central nervous system involving opioid GABA, and muscarinic receptors.

The reduction in hyperalgesia by LF TENS is prevented by blockade of μ opioid receptors in the spinal cord or the RVM or spinal cord, and by synaptic transmission in the ventrolateral PAG [4,5,9]. Further, the reduction in hyperalgesia by LF TENS is prevented by blockade of GABA_A, serotonin 5-HT_{2A} and 5-HT₃, and muscarinic M1 and M3 receptors in the spinal cord [11–13], and is associated with increased release of serotonin [14]. This opioid mediated effect of LF TENS has been confirmed in human subjects [15]. In addition, LF TENS does not produce analgesia in opioid tolerant people and animals but HF TENS does [16,17]. Thus, LF TENS uses classical descending inhibitory pathways involving the PAG-RVM pathway activating opioid, GABA, serotonin and muscarinic receptors to reduce dorsal horn neuron activity and the consequent pain.

• Reduction in central excitability

In animals without tissue injury, both LF and HF TENS reduce dorsal horn neuron activity [18–22]. In animals with peripheral inflammation or neuropathic pain, enhanced activity of dorsal horn neurons (i.e., central sensitization) to both noxious and innocuous stimuli is reduced by both HF and LF TENS [23–26]. In parallel, there is a reduction in both primary and secondary hyperalgesia by both LF and HF TENS [23,25–31]. Furthermore, in people with fibromyalgia and osteoarthritis, there is a reduction in pressure pain thresholds not only at the site of stimulation, but also at sites outside the area of application [6,32], implicating a reduction in central excitability.

HF TENS also reduces central neuron sensitization [24], and release of the excitatory neurotransmitters glutamate and substance P in the spinal cord dorsal horn in animals with

inflammation [33,34]. The reduction in glutamate is prevented by blockade of δ -opioid receptors. Thus, one consequence of activation of inhibitory pathways by TENS is to reduce excitation and consequent neuron sensitization in the spinal cord.

• Peripheral mechanisms of TENS

Both HF and LF TENS have effects at the site of stimulation. HF TENS reduces substance P, which is increased in dorsal root ganglia neurons in animals after tissue injury [33]. Blockade of peripheral opioid receptors prevents the analgesia produced by LF, but not HF TENS [35,36]. Thus, TENS may also alter excitability of peripheral nociceptors to reduce afferent input to the central nervous system.

In α -2a adrenergic knockout mice, analgesia by LF and HF TENS does not occur [37]. Blockade of peripheral, but not spinal or supraspinal, α -2 receptors prevents the analgesia produced by TENS [37] suggesting a role for peripheral α -2a-adrenergic receptors in analgesia produced by TENS. Further, the reduction in cold allodynia by LF TENS is reduced by administration of systemic phentolamine to block α -adrenergic receptors [25]. This adrenergic effect may alter the autonomic system. There are increases in blood flow with LF TENS at intensities that produce motor contractions; greater than 25% above motor threshold [38–42]. Thus, some of the analgesic effects of TENS are mediated through peripheral adrenergic receptors.

Factors that directly affect TENS efficacy

The factors affecting TENS efficacy include the population and the outcome assessed, timing of the outcome measures, negative interaction of opioid use and the parameters of the TENS dose. Three important factors for TENS efficacy are tolerance to repeated TENS, intensity of the stimulation and electrode placement. A recent article by Sluka *et al.* [43] provides an extensive review of variables that can affect the clinical use of TENS.

• Tolerance to repeated TENS

Repeated application of either LF or HF TENS at the same frequency, intensity and pulse duration daily (i.e., same dose), produces analgesic tolerance in animals [17] and humans [44]. The analgesic tolerance by LF TENS results in cross-tolerance at μ -opioid receptors in the spinal cord,

and the analgesic tolerance by HF TENS results in cross-tolerance at δ -opioid receptors in the spinal cord in animals [17]. Prevention of analgesic tolerance occurs with pharmacological modulation of pathways involved in opioid tolerance. Specifically blockade of NMDA-glutamate receptors or CCK receptors in the spinal cord prevents analgesic tolerance to both LF and HF TENS [45,46]. Analgesic tolerance can also be prevented by modulating between LF and HF TENS within a treatment session [47], or by increasing intensity of TENS daily [48]. Thus, animal studies suggest TENS tolerance can be delayed with pharmacological methods as well as with non-pharmacological modulation of TENS parameters.

• Intensity of TENS established as a critical factor in efficacy

The intensity of stimulation utilized is critical with TENS application. Using the strongest intensity that remains comfortable produces hypoalgesia in healthy subjects; lower intensities are ineffective [49–56]. In addition to activation of greater numbers of sensory afferents, higher pulse amplitudes are proposed to activate deeper tissue afferents allowing for greater analgesia [2]. High intensity TENS decreases post-operative opioid requirements and negative opioid-side effects [57,58]. Even as researchers demonstrate the importance of intensity in TENS delivery, TENS systematic reviews continue to include studies with wide ranging intensity settings. In fact, as outlined below, application of TENS at inadequate intensities is one of the primary factors attributed to conflicting reports of TENS efficacy. Therefore, clinicians should strive to apply TENS at the maximally tolerated intensity for each individual person.

• Electrode site placement

The intersection of acupuncture and TENS is receiving increasing attention in research. Numerous studies have examined both electroacupuncture and traditional TENS pad electrodes applied over acupuncture sites [59–67]. Clinically, application of TENS at these acupoints reduces pain and may be more effective than when applied over non-acupoint sites when measuring pain and pain thresholds to heat and pressure in normal subjects [59–63], as well as in patient populations [64–67] when compared with sham TENS. In post-operative hysterectomy subjects, TENS at acupoint sites reduced opioid

intake, nausea and dizziness when compared with TENS at non-acupoint sites [64].

Evidence of TENS for pain management

• Systematic reviews/meta-analyses

In the last 7 years, there have been a number of systematic reviews/meta-analyses that have examined efficacy of TENS for pain reduction in people with neck pain [68], postoperative pain [69], cancer pain [70,71], labor pain [72], acute pain [73], low back pain [74,75] and osteoarthritis pain [58,76]. There have also been systematic reviews on the methodology of TENS [77,78]. As a whole, these reviews are conflicting with some showing efficacy and some showing no efficacy for the use of TENS. The challenge is often a lack of high quality studies or a lack of consistency between high-quality studies included in the systematic reviews with respect to clinical population homogeneity, dose of TENS (i.e., location of TENS electrodes, frequency and intensity of TENS stimulation, and frequency and duration of TENS delivery), description of blinding and the influence of analgesic medication.

Table 1 represents a summary of these systematic reviews. Below we address the evidence on postoperative pain, acute non-postoperative pain, low back pain, osteoarthritis pain and painful diabetic neuropathy as examples.

Postoperative pain

There have been reviews of TENS efficacy in the last 7 years on management of postoperative pain which present differing results. A systematic review shows inconclusive results, [86] and a subsequent review shows positive effects [87]. The review by Bjordal and colleagues grouped trials into those with adequate TENS parameters (adequate frequency: 1-8 Hz for LF -TENS or 25-150 Hz for HF TENS; adequate intensity: strong sub noxious, maximal tolerable, or ≥ 15 mA) and those that did not meet these criteria. They show that those with adequate TENS parameters ($n = 11$) showed a 36% reduction in analgesic intake compared with those with inadequate TENS parameters ($n = 10$) that showed a 4% reduction. In contrast, the Cochrane review [86] did not consider dosing. Additionally, TENS has been found to reduce movement (walking and vital capacity maneuvers), but not resting, pain postoperatively [88]. Since the above systematic reviews focused on TENS for resting or overall pain, this factor may have also contributed to the conflicting results.

Acute nonpostoperative pain

A Cochrane review addressing acute pain (i.e., pain less than 12 weeks duration associated with procedures such as cervical laser, venipuncture, sigmoidoscopy screen, postpartum uterine contraction and rib fractures) in adults used a minimum stimulation intensity of 'strong but comfortable' as an inclusion factor. However, with 12 studies included, the authors were unable to make any conclusions due to insufficient evidence [73]. Four studies were included in a separate meta-analysis of RCTs where TENS was utilized in a pre-hospital setting for acute pain, (defined as moderate to severe) with TENS delivered by emergency service personnel. All studies found TENS lead to a clinically significant reduction in pain severity as compared with placebo TENS [89]. This review only included studies where TENS was used short term in ambulance responses. These studies were excluded from the Cochrane review of TENS and acute pain [73] due to low stimulation intensity. Thus, short-term use of TENS in ambulance responses the required intensity may be less than that required for chronic or other types of acute conditions. Recent randomized controlled trials for TENS show significant reductions in postpartum pain [90] and pain during wound-care procedures [91]. Interestingly, the mechanical triggers of wound-care procedures are similar to movement pain, supporting the effect of TENS for pain caused by mechanical stimulation, such as muscle movement, pressure, or force.

Low back pain

Systematic reviews [74,80] and a meta-analysis [75] have examined the efficacy of TENS for low back pain with conflicting results from not recommended [80], inconclusive [74], and effective [75]. All analyses used different inclusion and exclusion criteria, all examined effects on pain at rest, several used a mixed patient population, and none used dosing or timing of outcome, or examined potential interactions with pharmacological agents.

For example, the systematic review by Dubinsky and Miyasaki [80] was based on only two studies with differing patient populations - one for chronic, non-specific low back pain [92] and the other for low back pain in people with multiple sclerosis. The pain of MS is related to direct injury and permanent damage to the central nervous system [93]; while chronic musculoskeletal pain is generally due to modifiable

Table 1. TENS systematic reviews 2007–2013.							
Year	Topic	Author	Review type	Studies (n)	Subjects (n)	Results	Ref.
2013	Neck pain	Kroeling	Systematic review	20	1239	Update on 2009 and 2005 systematic review. Authors were unable to determine effect of TENS in neck pain due to limited quality of evidence, but suggest active TENS may be more effective than placebo TENS. Limited number of studies with standardization and description of treatment characteristics.	[68]
2012	Thoracic surgery	Sbruzzi	Meta-analysis	11	545	Use of random effects models to assess TENS effect s/p thoracic surgery. In thoracotomy and sternotomy, TENS associated with pharmacological analgesia improved pain relief compared with placebo TENS. With sternotomy, TENS was more effective than pharmacological analgesia for pain relief. No change in pulmonary function.	[69]
2012	Cancer pain adults	Hurlow	Systematic review	3	176	Update of 2008 Robb article in cancer pain. Addition of one RCT suggesting TENS may improve bone pain on movement in a cancer population. Results remained inconclusive due to a limited number of RCTs for review.	[71]
2011	Methodological Quality TENS and pain	Bennett	Systematic review	38	2268	Review of three Cochrane systematic reviews: acute pain, chronic pain and cancer pain. Authors identified sources of potential bias related to study design including less than optimal dosing of TENS, outcome assessment and timing as well as blinding and application of TENS. Proposal of criteria for future studies to enhance fidelity.	[77]
2011	Pain in labor	Bedwell	Systematic review	14	1256	Update to 2009 Dowswell article. Limited evidence that TENS reduces pain in labor. TENS does not appear to have effect on other outcomes for mothers and infants.	[72]
2010	Phantom limb pain	Mulvey	Cochrane systematic review	0	0	No RCTs have been completed to examine decreased pain in amputees. Further investigation is needed.	[79]
cLBP: Chronic low back pain; DPN: Diabetic peripheral neuropathy; IFC: Interferential current; LBP: Low back pain; OA: Osteoarthritis; QOL: Quality of life; RA: Rheumatoid arthritis; RCT: Randomized controlled trials; TENS: Transcutaneous electrical nerve stimulation.							

Table 1. TENS systematic reviews 2007–2013 (cont.).

Year	Topic	Author	Review type	Studies (n)	Subjects (n)	Results	Ref.
2008	TENS dose response for chronic pain	Claydon	Review of systematic reviews	6	Hand RA: 78 cLBP: 175 Knee OA: 294 Chronic musculoskeletal pain: 984 Chronic pain: 1227 Total: 2758	Two of six reviews of TENS and chronic pain reported high intensity TENS applications were more effective compared with placebo than low intensity. Reviewed confounding variables of inadequate design, low statistical power and different TENS protocols – single treatment versus repeated treatments of TENS.	[78]
2010	Neurological disorders (LBP and DPN)	Dubinsky	Systematic review	11		Inconsistent evidence for pain reduction in cLBP; probable evidence for pain reduction with diabetic peripheral neuropathy.	[80]
2010	Diabetic Peripheral Neuropathy	Jin	Meta-analysis	3	78	Pain reduction significantly greater than placebo following 4–6 weeks of treatment. Reduced hyperalgesia and numbness and increased QOL also significantly improved with active TENS.	[81]
2010	Hand RA	Brouseau	Cochrane systematic review	3	78	Update of 2003 review; Acupuncture like TENS has benefit for reducing pain intensity and increasing grip over placebo while conventional TENS no benefit compared with placebo.	[82]
2009	Pain in labor	Dowswell	Cochrane systematic review	17	1466	Limited evidence that TENS reduces pain in labor. Little difference between TENS groups and control groups. Those women receiving TENS to acupuncture points were less likely to report severe pain.	[83]
2009	Non-specific low back pain	Machado	Meta-analysis	4	178	Random effects statistical model demonstrated moderate effect for TENS in acute and chronic LBP	[75]
2009/2011	Acute pain	Walsh	Cochrane systematic review	12	919	Insufficient evidence to draw any conclusions about the effectiveness of TENS for the treatment of acute pain in adults.	[73]
2008	Chronic low back pain	Khadilkar	Cochrane systematic review	4	585	Conflicting evidence about TENS benefit in reducing back pain intensity. Acupuncture like TENS responded similar to conventional TENS, two of four studies lacked adequate stimulation intensity.	[74]
cLBP: Chronic low back pain; DPN: Diabetic peripheral neuropathy; IFC: Interferential current; LBP: Low back pain; OA: Osteoarthritis; QOL: Quality of life; RA: Rheumatoid arthritis; RCT: Randomized controlled trials; TENS: Transcutaneous electrical nerve stimulation.							

Table 1. TENS systematic reviews 2007–2013 (cont.).

Year	Topic	Author	Review type	Studies (n)	Subjects (n)	Results	Ref.
2008	Cancer pain adults	Robb	Cochrane systematic review	2	64	Due to small number of subjects and studies, there is insufficient evidence to determine the effectiveness of TENS and cancer pain. One RCT no significant difference in TENS and placebo TENS; one RCT no significant difference in acupuncture-like TENS and sham TENS.	[70]
2008	OA of knee	Rutjes	Cochrane systematic review	18	813	Mixed review of trials for TENS, IFC and pulsed electrical stimulation. Inconclusive for the results of TENS for pain and function of the knee due to small trials and inadequate design and power.	[76]
2008	Chronic pain	Nnoahm	Cochrane systematic review	25	1281	13/22 inactive control studies demonstrate a positive analgesic outcome for active TENS treatments. For multiple treatment comparison studies 8/15 were in favor of active TENS. 7/9 active controlled studies found no difference in analgesic efficacy between high frequency TENS and low frequency TENS.	[84]
2007	Knee OA	Bjorndal	Systematic review and meta-analysis	11	425	Seven of 11 studies had optimal TENS dosing and demonstrated clinically relevant pain relief compared with placebo control. These studies included IFC, electro acupuncture and low level laser therapy.	[58]
2007	Chronic low back pain; DPN; Diabetic peripheral neuropathy; IFC: Interferential current; LBP: Low back pain; OA: Osteoarthritis; QOL: Quality of life; RA: Rheumatoid arthritis; RCT: Randomized controlled trials; TENS: Transcutaneous electrical nerve stimulation.	Johnson	Meta-analysis	38	1227	With resting pain as main outcome measure, the overall random effects meta-analysis model showed a decrease in pain with electrical nerve stimulation.	[85]

‘plastic’ changes in both the peripheral and central pain pathways (sensitization) [94–96]. Machado [75] used people with non-specific low back pain with positive results – however, they combined acute and chronic low back pain, which likely have different underlying mechanisms.

None of the reviews considered adequate dosing of TENS and there were studies included in each review that did not describe TENS parameters or used inadequate doses. For example, the study by Deyo and colleagues [92], comparing TENS with and without exercise to placebo TENS with and without exercise in people with chronic low back pain, was included in two systematic reviews [74,80] and is rated as a well-designed clinical trial using appropriate blinding, randomization and good description of withdrawal and dropouts. However there are significant weaknesses in the application of TENS, some of which have been discovered since the trial was conducted 23 years ago. Intensity was applied by having subjects set the amplitude to a pre-designated setting on the machine which corresponded to 15 mA as obtained from the manufacturer. Patient response to stimulation was not stated. In our preliminary data, application of TENS to the spine that results in a strong but comfortable intensity requires at least 30 mA and, thus, the amplitude used was likely below an effective dose. Thus, it is not clear if TENS is effective for low back pain. Future studies should design clinical trials with adequate dosing and appropriate outcome measures. Future systematic reviews need to use patient populations with similar pain physiology and adequate use of TENS parameters as inclusion criteria.

Osteoarthritis pain

Similar to the reviews of acute pain and low back pain, a recent Cochrane systematic review showed that TENS was not effective for knee osteoarthritis (OA) pain [97], and is in direct contrast to a prior systematic review by the same group that concluded TENS was effective for knee OA pain [98] and a meta-analysis that showed a significant reduction in knee OA pain with TENS [58]. Intensities in the included studies varied widely. For example in the recent Cochrane review [97], 12 included trials used adequate intensities, five trials used inadequate intensities (HF-TENS at sensory threshold or below [99–103] and two trials did not report TENS intensity [104,105]. To address dosing, Bjordal and colleagues performed

a systematic review on TENS for osteoarthritis pain and show that when given at adequate intensities and frequencies TENS produces a clinically significant reduction in pain when compared with studies of inadequate dosing [58]. Therefore TENS works for OA pain if used at adequate intensities. A recent randomized controlled trial applied TENS in people with knee OA as an adjunct to primary care and showed no added benefit. However, parameters were not standardized and, and participants were allowed to self-select from eight different TENS protocols in the 6 week trial making interpretation of findings challenging [106].

Diabetic peripheral neuropathy (DPN)

In people with painful DPN, TENS may also provide benefit. A meta-analysis including three RCTs (n = 78) reported reduction of pain that was significantly greater than placebo TENS following 4–6 weeks of treatment [81]. In addition, secondary outcomes of overall improvement in DPN symptoms (hyperalgesia, numbness, and quality of life) were significantly greater for active TENS groups when compared with placebo [107–109]. Therefore, there is support for the use of TENS in reducing pain and improving quality of life in people with painful DPN.

• TENS interventions: emerging evidence from recent clinical trials

Fibromyalgia (FM)

Recent evidence suggests that TENS can be effective for people with fibromyalgia. Although there have been several randomized controlled trials [6,110–113], no systematic reviews have been published and the quality of these studies and the intervention have varied significantly. Two trials compared TENS to a placebo and used an adequate dose. Dailey *et al.* [6] showed a one-time session of TENS (using a maximum tolerable intensity) significantly decreased movement pain and hyperalgesia. No changes were observed in resting pain [6]. Lauretti *et al.* [111] showed TENS using a strong intensity (60 mA) at two sites and at one site produced a significant decrease in pain at rest compared with placebo when applied over a seven day period. Two additional studies show reductions in pain with strong but comfortable intensity HF TENS compared with warmth therapy and to a no TENS group [110,112]. Thus, when used at a strong but comfortable sensation, TENS may be effective for both resting and movement pain in people with fibromyalgia.

Neuropathic pain

TENS may offer relief to people with neuropathic pain and complex regional pain syndrome. A crossover design trial investigating neuropathic pain in people with spinal cord injury, [114] found a favorable effect of both LF and HF TENS (LF TENS 38%; HF TENS 29%) on a global relief scale and 25% of subjects requested a unit for further treatment. However, this study did not compare against a placebo or control group, intensity was not reported, and there were a low number of study participants ($n = 24$). A more recent study reports LF TENS provided significant reduction in pain when compared with placebo TENS in people with spinal cord injury. Here the parameters of 4 Hz and 200 μ s were applied at sites below the level of injury at a set intensity of 50 mA [115]. Thus, LF TENS may be most effective for pain in people with spinal cord injury.

Other pain conditions

A recent randomized controlled trial of TENS as an adjunct treatment in the management of lateral epicondylalgia concludes that TENS does not provide additional benefit when used as an adjunct to primary care (education and therapeutic exercise) [116]. In review, while an appropriate intensity was used, the intervention was not monitored for dosing and low adherence was reported. Further, outcome measures were assessed through questionnaires and not necessarily while wearing the TENS device. Additional TENS reports are favorable for relief of chronic pelvic pain syndrome [117] and pain associated with latent upper trapezius trigger points [118]. Overall, the evidence suggests, TENS may be useful for a variety of pain conditions.

Summary & conclusion

Because no single profession holds all the keys to successful management of pain, further investigation is warranted to ensure optimal use of this safe, noninvasive, inexpensive and patient friendly intervention. The advantages of

obtaining pain relief without the negative side effects of many pharmaceutical interventions is welcomed and desired by certain patients. Both HF and LF TENS been shown to provide analgesia specifically when applied at a strong, non-painful intensity and HF TENS may be more effective for people taking opioids. Effective analgesia for chronic pain conditions may be limited by the development of tolerance to TENS if repeated application of either HF or LF TENS at the same frequency, intensity and pulse duration is used daily. Application of TENS electrodes at acupoint sites may increase analgesia and targeting the use of TENS during movement or required activity may provide the most benefit.

Experiments investigating the concept of TENS responders will enable clinicians to select this modality for the correct population. Additional investigation in the area of TENS tolerance is necessary to determine methods to decrease tolerance and to establish if a wash out period is required to determine when tolerance would no longer be a factor in the application of TENS in patient care. Although parameter selection is becoming clearer, investigating the parameters of electrode site selection, daily treatment duration, and long-term usage will further clarify appropriate dosing so that TENS may be given in the most effective manner. Further, examining a variety of outcomes, beyond resting pain, will determine if TENS has effects specific to mechanical stimuli and/or beyond reduction of pain and will improve activity levels, function and quality of life.

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