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ORIGINAL RESEARCH

# Effects of Wearable Transcutaneous Electrical Nerve Stimulation on Fibromyalgia: A Randomized Controlled Trial

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Correspondence: Shai N Gozani NeuroMetrix, Inc., 4b Gill St., Woburn, MA, 01801, USA Tel +1 781 314 2789 Fax +1 781 890 1556 Email gozani@neurometrix.com **Purpose:** Fibromyalgia is a chronic condition characterized by widespread pain and interference with daily activities. The aim of this study is to assess the benefit of transcutaneous electrical nerve stimulation (TENS) for persons diagnosed with fibromyalgia.

**Patients and Methods:** Adults meeting diagnostic criteria for fibromyalgia were randomized in a double-blind trial to receive either an active (n=62) or sham (n=57) wearable TENS device for 3-months. Subjects were classified as having lower or higher pain sensitivity by Quantitative Sensory Testing (QST). Patient Global Impression of Change (PGIC, primary outcome) and secondary efficacy measures including Fibromyalgia Impact Questionnaire (FIQR), Brief Pain Inventory (BPI) and painDETECT questionnaire (PDQ) were assessed at baseline, 6-weeks and 3-months. Treatment effects were determined by a mixed model for repeated measures (MMRM) analysis of the intention-to-treat (ITT) population (N=119). A pre-specified subgroup analysis of pain sensitivity was conducted using an interaction term in the model.

**Results:** No differences were found between active and sham treatment on PGIC scores at 3-months (0.34, 95% CI [-0.37, 1.04], p=0.351) in the ITT population. However, in subjects with higher pain sensitivity (n=60), PGIC was significantly greater for active treatment compared to sham (1.19, 95% CI [0.24, 2.13], p=0.014). FIQR total score (-7.47, 95% CI [-12.46, -2.48], p=0.003), FIQR pain item (-0.62, 95% CI [-1.17, -0.06], p=0.029), BPI Interference (-0.70, 95% CI [-1.30, -0.11], p=0.021) and PDQ (-1.69, 95% CI [-3.20, -0.18], p=0.028) exhibited significant improvements for active treatment compared to sham in the ITT population. Analgesics use was stable and comparable in both groups.

**Conclusion:** This study demonstrated modest treatment effects of reduced disease impact, pain and functional impairment from wearable TENS in individuals with fibromyalgia. Subjects with higher pain sensitivity exhibited larger treatment effects than those with lower pain sensitivity. Wearable TENS may be a safe treatment option for people with fibromyalgia.

#### Clinicaltrials.gov Registration: NCT03714425.

**Keywords:** fibromyalgia, wearable, transcutaneous electrical nerve stimulation, neuromodulation, non-pharmacological treatment, clinical trial

### Introduction

Fibromyalgia is a chronic condition characterized by widespread pain and tenderness. Individuals with fibromyalgia also experience sleep disturbances, fatigue, cognitive impairment, and mood disorders.<sup>1</sup> The prevalence is estimated at 2-6%of the US adult population and is more common in women than men.<sup>2</sup> The specific

Journal of Pain Research 2021:14 2265-2282

© 2021 Jamison et al. This work is published and licensed by Dove Medical Press Limited. The full terms of this license are available at https://www.dovepress.com/terms work you hereby accept the Terms. Non-commercial uses of the work are permitted without any further permission from Dove Medical Press Limited, provided the work is properly attributed. For permission for commercial use of this work, please see paragraphs A2 and 5 of our Terms (https://www.dovepress.com/terms.php). cause of fibromyalgia is unknown; however, pathological changes in the central nervous system leading to pain hypersensitivity are likely involved.<sup>3</sup> Fibromyalgia diagnostic criteria have evolved over the past 30 years and may include characteristic symptoms, tenderness on physical exam and medical history.<sup>1,4,5</sup> However, diagnosis remains challenging, with many patients suffering for years before identification of the disease.<sup>6</sup> There is increasing interest in developing objective and quantitative biomarkers for the condition.<sup>7</sup> As an example, many individuals with fibromyalgia exhibit low pressure pain thresholds throughout the body and physiological signs of central sensitization and diminished endogenous pain inhibition when assessed by Quantitative Sensory Testing (QST).<sup>3</sup>

Few treatments have been shown to be effective in managing fibromyalgia. Three drugs (pregabalin, duloxetine, and milnacipran) have been approved by the Food and Drug Administration (FDA) for management of fibromyalgia but these pharmacological agents are often associated with side effects<sup>8–10</sup> and poor adherence.<sup>11,12</sup> Non-pharmacological treatments such as patient education and physical exercise are recommended as first-line therapy, potentially followed by alternative approaches such as cognitive behavioral therapy and acupuncture.<sup>13</sup> Although these non-pharmacological interventions are safe, their efficacy has not been conclusively demonstrated. In general, fibromyalgia treatments are associated with small to moderate effect sizes.<sup>14–16</sup>

Transcutaneous Electrical Nerve Stimulation (TENS) is believed to activate descending pain inhibition and reduce central excitability.<sup>17,18</sup> Theses mechanisms align with pathological changes in pain processing that are thought to underlie fibromyalgia. TENS is usually applied to the site of pain, however its analgesic effects can be widespread,<sup>19,20</sup> potentially making it effective for multisite pain.<sup>21,22</sup> For these reasons, TENS has been suggested as a non-pharmacological option for patients with fibromyalgia.<sup>20,23</sup> However, the efficacy of TENS in reducing fibromyalgia pain and symptoms is uncertain.<sup>24</sup> As a general matter, randomized controlled trials of TENS have been criticized for small sample sizes, poor controls, the absence of compliance monitoring, and inadequate stimulation intensity.<sup>25,26</sup> A recent trial of daily TENS use for 1-month addressed these deficiencies and demonstrated significant treatment effects in fibromyalgia patients compared to sham TENS and no TENS controls.<sup>23</sup>

Wearable TENS is an emerging category of noninvasive neuromodulation where the device is designed for placement directly on the body, typically at a fixed location such as the upper calf or arm. These devices may incorporate automated stimulation control and sensors to adaptively modulate stimulation and track objective outcomes.<sup>27</sup> Some wearable devices are designed for extended wear, including during sleep, which addresses a key limitation of traditional TENS.<sup>28</sup> Wearable TENS devices have been evaluated in chronic lower extremity pain,<sup>29</sup> chronic low back pain,<sup>30</sup> chemotherapy induced peripheral neuropathy (CIPN),<sup>31</sup> migraine,<sup>32</sup> and essential tremor<sup>33</sup> with generally encouraging results. Wearable TENS has not been evaluated for management of fibromyalgia.

The present RCT compared 3-months of at-home treatment with an active or sham wearable TENS device located on the upper calf in individuals with fibromyalgia. The upper calf location enables stimulation of sensory dermatomes S2 through L4 with a circumferential electrode. These dermatomes are typically targeted when treating lower extremity and low back pain,<sup>34</sup> which are common in fibromyalgia.<sup>4,35</sup> The primary hypothesis was that active treatment would produce greater improvements in pain, somatic symptoms, and functional impairment compared to sham treatment. A second hypothesis was that subjects with the greatest relative baseline pain sensitivity by QST would exhibit the largest treatment effects. This hypothesis was based on the proposition that analgesic treatments that target sensitized central neurons, such as TENS,<sup>17,18</sup> may be most effective in chronic pain characterized by hyperalgesia and central sensitization. Modelbased statistical analyses were employed to maximize power to detect treatment effects in the presence of low to moderate effect sizes characteristic of fibromyalgia.

## Materials and Methods Study Design and Subject Selection

The protocol was registered at ClinicalTrials.gov (NCT03714425) prior to initiation of the study. This single-site, parallel-group trial was conducted at a tertiary academic hospital between February 2019 and June 2020. Following a screening process, eligible individuals were scheduled for a baseline visit. All subjects signed a written informed consent and were randomized to either an active or sham device. All participants were administered a brief stimulation trial with the device and if they disliked the sensation they could immediately withdraw from the study. No subject chose to withdraw. QST was performed at this initial visit. Following the baseline visit, subjects were called weekly to monitor changes in analgesic use and adverse events. This study conforms to the Consolidated Standards of Reporting Trials (CONSORT) guidelines,<sup>36</sup> including a recently published pain-specific supplement.<sup>37</sup> This study protocol and all amendments were approved by the Human Research Committee (Institutional Review Board) of Mass General Brigham (Massachusetts General Hospital and Brigham and Women's Hospital, Boston, MA, USA). This study followed the principles outlined in the World Medical Association Declaration of Helsinki – Ethical Principles or Medical Research Involving Human Subjects.

The subject inclusion criteria were age 21 or older; able to speak and understand English; own a smartphone that can run the mobile app associated with the wearable TENS device; meet American College of Rheumatology 2010 diagnostic criteria for fibromyalgia;<sup>1</sup> physician diagnosis of fibromyalgia in the medical record; and average pain intensity  $\geq 4$  on an 11-point numerical rating scale (NRS). Exclusion criteria included a diagnosis of cancer or other malignant disease; acute osteomyelitis or acute bone disease; present or past psychiatric diagnosis (Diagnostic and Statistical Manual of Mental Disorders: DSM-5) that was judged by the principal investigator to interfere with study participation (no subjects were excluded on this basis in the present study); pregnancy; clinically unstable systemic illness that could interfere with treatment; pain condition requiring urgent surgery; active substance use disorder that could interfere with study participation; or an implanted cardiac pacemaker, defibrillator, or other implanted electronic device. Subjects were asked to continue their pre-study analgesic medications with selfreported changes tracked through a weekly phone interview.

## Randomization and Blinding

Subjects were randomized to an active or sham device with equal allocation. Active and sham devices were physically identical; only differing in whether they were loaded with standard software or modified software that implemented a sham stimulation protocol. A total of 120 devices were randomized by the manufacturer (www. <u>graphpad.com/quickcalcs/randomize1</u>) and provided to the trial site. The manufacturer had no interaction with the study subjects. The study coordinators and investigators could not determine whether a device was an active or sham device based on any markings or physical characteristics and did not discuss the stimulation experience with subjects. Subjects were told that two types of TENS were being evaluated, a "low intensity" device and a "high intensity" device. Blinding effectiveness for both subjects and study coordinators was assessed at the end of the study. Subjects in the sham treatment group were offered a device with standard software after completing the study.

## **TENS** Intervention

The active treatment was a commercially available wearable TENS device (Quell, NeuroMetrix, Inc., Woburn, MA, USA) that is placed on the lower extremity of either leg, typically the upper calf.<sup>27,29</sup> The device is comprised of a one-channel electrical stimulator, a stretchable band to secure the stimulator to the leg, an electrode array and a smartphone app. The electrode array consists of 4 hydrogel pads that provide a total stimulation surface area of  $60 \text{ cm}^2$ . The stimulator generates bipolar, current-regulated pulses with a duration of 290 microseconds and alternating leading phase polarity. Stimulation frequency is random with a uniform distribution between a 60 and 100 Hz. The stimulator communicates with a smartphone application through Bluetooth<sup>®</sup>. The mobile application serves as a remote control for stimulator functions, displays device status, and tracks utilization. It is linked with a cloud database for storage of deidentified utilization data.

Prior to first use, the device calibrates to the user's sensation threshold using an algorithm based on ascending and descending methods of limits. Subsequent stimulation is controlled automatically, although the user can also manually decrease or increase intensity. The initial stimulation level is 1.8 times the sensation threshold. This intensity is generally perceived as "strong but comfortable" by most individuals,<sup>38</sup> which is the target sensation for effective conventional TENS.<sup>26,39</sup> Each therapy session is 60 minutes, with sessions automatically starting every other hour as long as the device is on the body, including overnight. The active device provided 60-minutes of continuous stimulation during each therapy session. The sham device provided three 2-minute periods of stimulation during each session (at 0, 28, and 58 minutes) for a total of 6-minutes of stimulation. Sham TENS based on transient stimulation has been validated in healthy controls<sup>40</sup> and used in earlier osteoarthritis<sup>41</sup> and fibromyalgia RCTs.<sup>20,23</sup> The device placement on the upper calf and usage instructions were identical for the two devices. Subjects were instructed to maintain a strong but comfortable stimulation intensity<sup>26</sup> and to use their device for at least two 1-hour therapy sessions each day over the course of the study.

## Quantitative Sensory Testing

The multimodal QST procedures included mechanical and cold stimuli.<sup>30,42</sup> Responses to punctate mechanical stimuli were measured using a standard set of weighted probes (Touch-Test Sensory Evaluator; North Coast Medical, Inc.). Singular taps were performed on the metacarpophalangeal joint of the middle finger of the non-dominant hand.<sup>43</sup> The lowest-force stimulator that produced a sensation of discomfort at 10 out of 100 was used to assess temporal summation of pain that occurred with rapid administration of 10 identical stimuli at 1-second intervals. Participants rated the painfulness of the first, fifth, and tenth stimulus. Mechanical temporal summation was defined as the increase in pain from the first to the tenth stimulus. A Somedic pressure algometer (Somedic SenseLab AB) was used to measure pain pressure thresholds (PPT) at the trapezius muscle and thumb joint. Each site was assessed twice on the left and right sides. Mechanical pressure was applied using a 0.5-cm<sup>2</sup> probe covered with a 1-mm polypropylene pressure-transducing material. Pressure was increased at a steady rate of 30 kPA/s until the subject indicated that the stimulus was painful. Cuff algometry was used to assess responses to sustained mechanical pressure. A Hokanson rapid cuff inflator (D. E. Hokanson, Inc.) was used to inflate a standard blood pressure cuff around the gastrocnemius muscle of the dominant leg until the subject indicated a pain level of 40 out of 100. This pressure was maintained for 2 minutes, with the subject rating their pain at 30 second intervals.

Responses to noxious cold were evaluated using a repeated cold pressor task (CPT), which involved immersion of the right hand in a circulating water bath (Neslab RTE 17, Thermo Electron Corp.) maintained at 4 °C. Participants underwent a series of CPTs, with the first 2 consisting of serial immersions of the dominant hand for 15 seconds, with 2 minutes between immersions. Once the subject removed their hand, pain ratings were asked at 0, 15, 30, and 60 seconds. If the subject was not able to remain in the water for the full 15 seconds, they were able to remove it early with the same assessment intervals. Conditioned Pain Modulation (CPM) was measured by assessing PPT at the trapezius during the water bath immersions. The final CPT involved an immersion of the dominant hand until the participant reached maximum pain tolerance (or 3 minutes). Pain was assessed at 15 second intervals while submerged and as soon as the hand was removed from the water. The participants rated the intensity of cold pain on a 0–100 scale.

## Efficacy Measures

The primary outcome measure was the Patient Global Impression of Change (PGIC), which represents the subject's overall belief about the efficacy of treatment on a 7-point categorical verbal rating scale.<sup>44</sup> The scale ranges from (1) "no change or condition has gotten worse" to (7) "a great deal better and a considerable improvement that has made all the difference." There were 7 pre-specified secondary efficacy measures. Disease impact and health related quality-of-life was assessed with the Fibromyalgia Impact Questionnaire (FIQR).<sup>45</sup> The FIQR pain intensity and sleep quality items, which are components of the FIQR total score, were also separately analyzed because of their particular importance to individuals with fibromyalgia.46-48 Pain severity and pain interference with function were evaluated with the Brief Pain Inventory (BPI).49 Neuropathic pain was assessed with the 7-item painDETECT questionnaire (PDQ).<sup>50,51</sup> Pain-related disability was evaluated with the Pain Disability Index (PDI).<sup>52</sup> Psychological outcomes included the Hospital Anxiety and Depression Scale (HADS)<sup>53</sup> and the Pain Catastrophizing Scale (PCS).<sup>54</sup> These outcome measures have been widely used in prior fibromyalgia treatment trials and their psychometric properties have been validated in US patients with chronic pain.55 The psychometric characteristics of several instruments have been specifically evaluated in patients with fibromyalgia.45,56,57 All instruments were delivered in conventional paper format consistent with their original formulation and validation. All efficacy measures except for PGIC were taken at baseline, 6-weeks, and 3-months. PGIC was assessed at 6-weeks and 3-months. The 6-week outcome assessments were mailed to the study subjects, completed at home, and returned via mail. The 3-month efficacy assessments were intended to be completed by subjects at the 3-month in-person visit. Due to COVID-19 restrictions on clinic visits beginning March 2020, 27 subjects were asked to complete the 3-month outcome assessments via mail.

## Statistical Analyses

The sample size calculation was designed to confirm the hypothesis that the active device would be associated with

a greater PGIC score compared to the sham device at 3-months of 0.6 points with a standard deviation (SD) of 1.0.<sup>58–60</sup> The calculation assumed 85% power and a twosided Type I error rate of 0.05. The estimated sample size of 100 was increased to a target recruitment of 115 to account for 15% drop-out.<sup>30</sup> The primary analysis of treatment effects was conducted in the intention-to-treat (ITT) population, which included all 119 randomized subjects. In addition, a pre-specified subgroup analysis was carried out. The study protocol predicted that subjects with higher pain sensitivity based on QST measures would demonstrate the greatest treatment effects. Pain sensitivity was classified using the first principal component of the base-line QST data.<sup>61</sup>

The mean PGIC score at 3-months and the mean baseline to 3-month change scores for the secondary efficacy measures were compared between the active and sham treatment groups by a mixed model for repeated measures (MMRM) analysis. In the presence of missing data, a MMRM analysis typically has more power than a twosample t-test<sup>62</sup> or ANCOVA model.<sup>63</sup> The ITT model included fixed effects for treatment, visit (baseline, 6-weeks, 3-months), and a treatment-by-visit interaction. Covariates included the baseline value (except for PGIC), baseline pain severity (BPI average pain item) and BMI along with the three corresponding by-visit interactions. The subgroup model included all the parameters in the ITT model and an interaction between treatment and pain sensitivity. This model was first used to test for treatment heterogeneity by a significant interaction term at a twosided p-value less than 0.15.<sup>64,65</sup> The model was then used to estimate the marginal effects of treatment for subjects with lower and higher pain sensitivity.

A value of 1 ("no change") was assigned as the 3-month PGIC score if there was no 6-week or 3-month data for a subject. The outcome vector of the secondary efficacy measures included a change score of zero for the initial (baseline) visit to account for subjects with no treatment data.<sup>66,67</sup> Missing covariates were addressed with the missing-indicator approach.<sup>68</sup> Correlations among measurements taken on the same subject were first modeled with an unstructured covariance assumption. If the model failed to converge then a first-order autoregressive covariance structure was used. Marginal effects of treatment were determined at 3-months along with corresponding two-sided p-values. In the subgroup model, the marginal effects were further conditioned on pain sensitivity. Comparisons were deemed significant if the two-sided

p-value was less than 0.05. Adjustments for multiple comparisons of secondary efficacy measures were not performed to preserve Type II error rates at the risk of an elevated family-wise Type I error rate.<sup>69</sup> Multiplicity corrections (eg, Bonferroni) generally assume that outcomes are independent and will overcompensate for correlated measures leading to increased Type II errors. In the present study, there were moderate correlations among many of the efficacy measures.

A MMRM analysis uses all available data at each visit and implicitly imputes missing data under a missing at random (MAR) assumption. A sensitivity analysis was performed using reference-based multiple imputation (MI) with the jump-to-reference method.<sup>70,71</sup> The MI model included the same covariates as the primary MMRM analysis. The imputed datasets were analyzed with the original MMRM model and combined using Rubin's rule.<sup>70</sup> The impact of missing data was assessed by comparing the resulting estimates with the primary MMRM estimates.

Responder rates were compared between treatment groups using logistic regression. Missing outcomes due to study withdrawal were treated as non-responders. For subjects that completed the study, 6-week data were used if 3-month data were not available. The dependent variable in the model was a binary variable indicating whether the subject was a responder or non-responder. The model included treatment assignment as an independent variable and baseline pain severity (BPI average pain item) and BMI as covariates. An interaction term between treatment and baseline pain sensitivity was added for subgroup analyses of responder rates.

# Results

## Subjects

Of 170 individuals screened for the study, 119 met the inclusion/exclusion criteria and were randomized to an active (62) or a sham (57) device for 3-months (Figure 1). Among these subjects, 16 (7 active, 9 sham) withdrew and were lost to follow-up and 3 (3 active, 0 sham) withdrew but completed the 3-month assessments. The remaining 100 subjects completed 3-months of treatment, however 4 (1 active, 3 sham) did not return the 3-month assessments. A comparison of demographic data between the two groups is presented in Table 1. The active group exhibited non-significant but numerically higher efficacy measures at baseline (Table 2). The baseline characteristics were similar

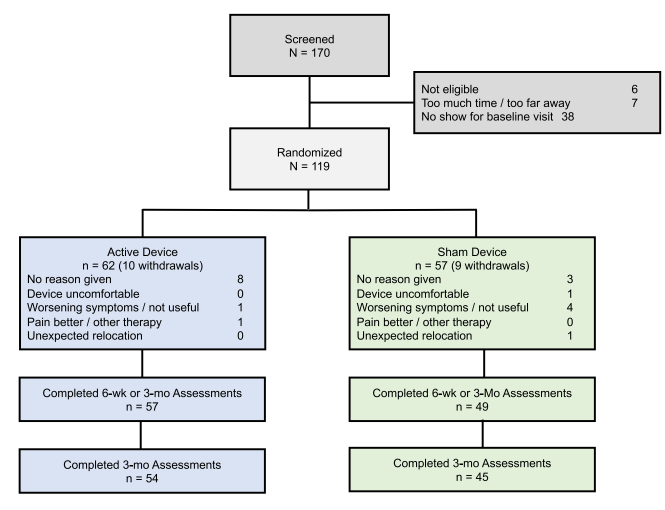


Figure I CONSORT diagram with intention-to-treat.

to other trials of pharmacological and non-pharmacological interventions for fibromyalgia.<sup>23,58</sup> Half of the subjects (47.1%) were taking over-the-counter analgesics, 31.9% were prescribed neuroleptics, 27.7% were prescribed an antidepressant, and 21.8% were taking an opioid, including tramadol. There was no discernable change in analgesic use over the course of the study and there was no significant difference between the treatment groups for any week (Supplemental Table 1). Most subjects were compliant with the target of 2 or more TENS sessions per day and utilized adequate stimulation intensity (Supplemental Table 2). Participants averaged 3.5 (SD 2.4) sessions per day and used their device on an average of 68.9 (SD 27.1) days and 23.7 (SD 23.4) nights during the study. No differences in TENS utilization were found between groups. Both the active and sham stimulation intensities (defined relative to sensation threshold) were comparable to values reported for successful long-term users of TENS.<sup>38</sup>

### **Baseline Pain Sensitivity**

Principal component analysis was applied to the correlation matrix of the baseline QST data (Table 3) to identify the prominent pain patterns in the study population. The loading weights for the first principal component, which accounted for 40% of the total variance, are shown in Figure 2. The component had negative loadings for mechanical pain thresholds, positive loadings for pain responses to punctate stimuli and cold stimuli, and a positive loading for temporal summation, which can be interpreted as a composite index of pain sensitivity. Subjects were classified as lower (< median) or higher ( $\geq$ median) pain sensitivity using this principal component. This yielded a lower pain sensitivity subgroup with 59 subjects and a higher pain sensitivity subgroup with 60 subjects. The proportions of subjects with higher pain sensitivity were 48.4% in the active group and 52.6% in the sham group, p=0.643. The proportions of higher pain

Characteristic	All Subjects (N=119)	Treatment		
	(14-117)	Sham (n=57)	Active (n=62)	
Age (years)	50.4 (13.5)	48.3 (13.1)	52.3 (13.8)	
Female (%)	(93.3)	53 (93.0)	58 (93.5)	
BMI (kg/m <sup>2</sup> )	27.5 (6.2)	27.0 (5.4)	28.0 (6.9)	
Education (years)	15.7 (2.9)	15.8 (2.7)	15.6 (3.0)	
Race (%)				
Caucasian	95 (79.8)	47 (82.5)	48 (77.4)	
African American	10 (8.4)	5 (8.8)	5 (8.1)	
Asian	I (0.8)	0 (0.0)	I (I.6)	
American Indian/Alaska Native	2 (1.7)	I (1.8)	l (l.6)	
Other	(9.2)	4 (7.0)	7 (11.3)	
Marital Status (%), n=116 responded				
Single	46 (39.7)	25 (46.3)	21 (33.9)	
Married	52 (44.8)	21 (38.9)	31 (50.0)	
Widowed	6 (5.1)	3 (5.6)	3 (4.8)	
Divorced	12 (10.3)	5 (9.3)	7 (11.3)	
Working Status (%)				
Full Time	33 (27.7)	15 (26.3)	18 (29.0)	
Part Time	26 (21.9)	15 (26.3)	(17.7)	
Not Working	59 (49.6)	27 (47.4)	32 (51.6)	
Medical Leave	I (0.8)	0 (0.0)	I (I.6)	
Compensation Status (%), n=115 responded				
None	71 (61.7)	38 (69.1)	33 (55.0)	
Workers Compensation	I (0.9)	I (I.8)	0 (0.0)	
Social Security Disability	31 (30.0)	12 (21.8)	19 (31.7)	
Retirement	5 (4.4)	2 (3.6)	3 (5.0)	
Unemployment	2 (1.7)	I (I.8)	l (l.7)	
Other	5 (4.4)	I (I.8)	4 (6.7)	
Tobacco Use (%), n=117 responded	13 (11.1)	6 (10.5)	7 (11.3)	

Note: Reported as mean (SD) or count (percentage).

Abbreviations: SD, standard deviation; BMI, body mass index.

sensitivity were similar in subjects that completed the study (50.0%) and those that withdrew early (52.6%), p=0.833.

## Patient Global Impression of Change

The mean PGIC score at 3-months was 3.58 (95% CI [3.09, 4.07]) in the active group and 3.24 (95% CI [2.74, 3.75]) in the sham group (Figure 3A). The mean difference of 0.34 (95% CI [-0.37, 1.04], p=0.351) was not significant and corresponded to an effect size of 0.17 (95% CI [-0.19, 0.53]). The prespecified subgroup analysis

demonstrated a significant interaction between treatment and pain sensitivity (p=0.011). In the higher pain sensitivity subgroup, the mean PGIC score at 3-months was 4.05 (95% CI [3.37, 4.73]) for active treatment and 2.86 (95% CI [2.19, 3.53]) for sham treatment (Figure 3A). The mean difference of 1.19 (95% CI [0.24, 2.13], p=0.014) was significant and corresponded to an effect size of 0.63 (95% CI [0.11, 1.15]). The difference between active treatment and sham treatment in subjects with lower pain sensitivity was not significant (mean difference -0.55, 95% CI [-1.52, 0.42], p=0.268).

Efficacy Measure	All Subjects	Treatment		
	(N=119)	Sham (n=57)	Active (n=62)	
FIQR Total Score	57.4 (17.2), 105	52.8 (17.3), 48	61.3 (16.3), 57	
FIQR Pain Item	6.5 (1.9), 118	6.2 (1.8), 57	6.8 (2.0), 61	
FIQR Sleep Item	7.4 (2.5), 119	7.0 (2.5), 57	7.8 (2.4), 62	
BPI Severity	5.6 (1.6), 118	5.5 (1.5), 57	5.8 (1.6), 61	
BPI Interference	5.7 (2.2), 119	5.4 (2.2), 57	5.9 (2.2), 62	
PDQ	16.5 (6.9), 115	15.6 (6.4), 55	17.3 (7.2), 60	
PDI	37.1 (15.7), 115	34.6 (15.5), 55	39.4 (15.6), 60	
HADS	17.2 (7.5), 117	15.8 (7.6), 56	18.5 (7.3), 61	
PCS	19.8. (12.9), 118	17.8 (12.3), 56	21.5 (13.2), 62	

Notes: Reported as mean (SD), sample size; sample size less than maximum if missing efficacy assessments at baseline due to missing individual items within questionnaires. Abbreviations: SD, standard error; Cl, confidence interval; FIQR, Fibromyalgia Impact Questionnaire Revised; BPI, Brief Pain Inventory; PDQ, painDETECT questionnaire; PDI, Pain Disability Index; HADS, Hospital Anxiety and Disability Scale; PCS, Pain Catastrophizing Scale.

# Secondary Efficacy Measures

After 3-months of active treatment, all secondary efficacy measures exhibited significant within-group improvement compared to their pre-treatment baseline (Table 4). Fibromyalgia disease impact (FIQR total score) decreased by 13.49 points (95% CI [-16.80, -10.18]) in the active group compared to a decrease of 6.02 points (95% CI [-9.57, -2.47]) in the sham group, which represented a significant difference of -7.47 points (95% CI [-12.46,

-2.48], p=0.003) and corresponded to an effect size of 0.55 (95% CI [0.18, 0.92]). Pain intensity (FIQR pain item) decreased by 1.83 points (95% CI [-2.20, -1.46]) for active treatment compared to a decrease of 1.21 points (95% CI [-1.61, -0.82]) for sham (Figure 3B). The difference was significant (mean -0.62, 95% CI [-1.17, -0.06], p=0.029) and represented an effect size of 0.41 (95% CI [0.04, 0.77]). Differences between the active and sham treatments primarily emerged between the 6-week

Table 3	Comparison	of Quantitative	Sensory Tests	at Baseline
Table 5	Comparison	Of Quantitative	Jensory resta	at Daschine

Measurement	All Subjects (N=119)	Treatment		
		Sham (n=57)	Active (n=62)	
Mechanical Pain Thresholds				
PPT Thumb Joint (kPa)	211 (95)	205 (88)	217 (104)	
PPT Trapezius Muscle (kPa)	254 (151)	249 (150)	259 (152)	
Cuff Pressure Gastrocnemius (mmHg)*	131 (66)	131 (67)	132 (65)	
Mechanical Pain Responses				
Punctate Stimulus, 1st of 10 <sup>†</sup>	12.1 (12.9)	12.4 (14.6)	11.8 (11.2)	
Punctate Stimulus, 5th of 10 <sup>†</sup>	20.6 (16.1)	19.9 (16.8)	21.2 (15.6)	
Punctate Stimulus, 10th of $10^{\dagger}$	26.1 (19.6)	25.8 (20.1)	26.3 (19.2)	
Cold Pain Responses				
CPT at 15 sec <sup>†</sup>	74.5 (21.6)	73.4 (20.1)	75.4 (23.0)	
CPT Aftersensation at 30 $sec^{\dagger}$	27.3 (24.0)	30.2 (23.0)	24.7 (24.8)	
Central Pain Processing				
Temporal Summation <sup>†</sup>	14.0 (14.8)	13.4 (13.9)	14.5 (15.7)	
Conditioned Pain Modulation (%)	51.9 (40.0)	52.5 (41.3)	51.4 (39.0)	

**Notes**: Reported as mean (SD); \*median imputation for 3 subjects without data; <sup>†</sup>pain rating scale from 0 to 100. **Abbreviations**: PPT, pressure pain threshold; CPT, cold pressure test.

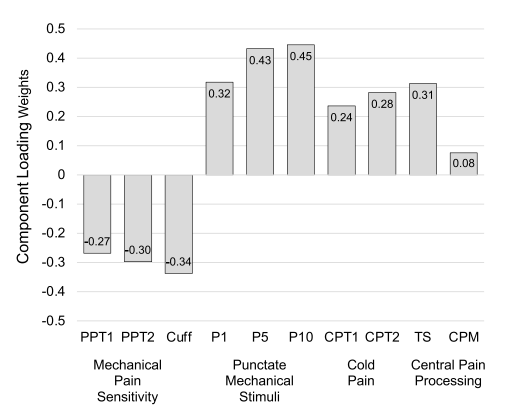


Figure 2 Loading weights for the first QST principal component. PPT1, PPT at trapezius muscle. PPT2, PPT at thumb joint. Cuff, cuff pressure at gastrocnemius, P1, 1st of 10 punctate stimuli delivered once per second rated on 0–100 pain scale. P5, 5th of 10 punctate stimuli. P10, last of 10 punctate stimuli. CPT1, cold pressor test at 15 seconds following hand immersion into cold water bath (pain 0–100). CPT2, 30-second after sensation following 15 seconds of hand immersion in cold water bath (pain 0–100). The first component can be interpreted as an index of pain sensitivity.

Abbreviations: TS, temporal summation; CPM, conditioned pain modulation; QST, Quantitative Sensory Testing; PPT, pressure pain threshold.

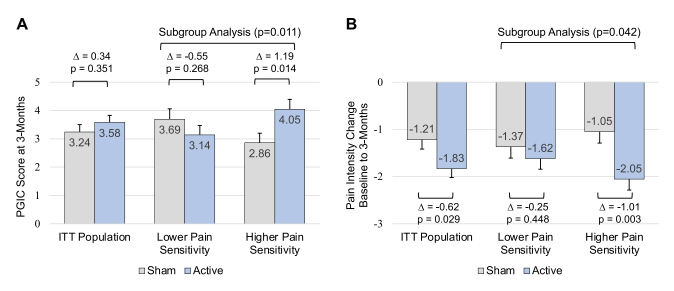


Figure 3 Comparisons of PGIC scores at 3-months (**A**). Comparisons of baseline to 3-month change scores in pain intensity (FIQR pain item) (**B**). Error bars indicate SE. Δ, treatment effect (Active - Sham). Treatment comparisons based on MMRM analyses of ITT population and of the lower and higher pain sensitivity subgroups. Subgroup analysis p-value is for the treatment by pain sensitivity interaction term in the subgroup MMRM model. **Abbreviations:** PGIC, Patient Global Impression of Change; FIQR, Fibromyalgia Impact Questionnaire Revised; ITT, intention-to-treat; MMRM, mixed model for repeated measures; SE, standard error.

#### Table 4 Mean Changes in Efficacy Measures from Baseline to 3-Months Using a MMRM Analysis

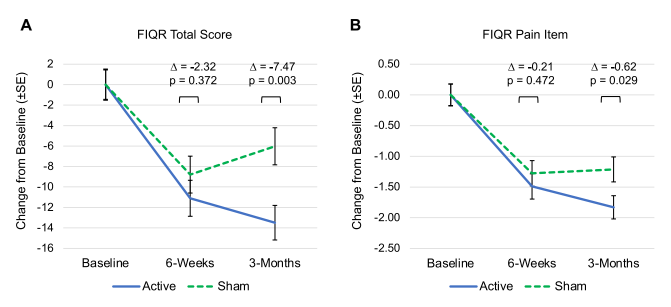
Measure	N	Mean Change	SE	Treatment Comparison (Active - Sham)		
				Difference	95% CI	p-value
FIQR Total Score						
Sham	57	-6.02 <sup>†</sup>	1.81			
Active	62	-13.49 <sup>‡</sup>	1.69	-7.47	-12.46, -2.48	0.003
FIQR Pain Item						
Sham	57	-I.2I <sup>‡</sup>	0.20			
Active	62	-1.83 <sup>‡</sup>	0.19	-0.62	-1.17, -0.06	0.029
FIQR Sleep Quality Item						
Sham	57	-0.55	0.31			
Active	62	-1.59 <sup>‡</sup>	0.28	-1.04	-1.87, -0.20	0.015
BPI Severity						
Sham	57	-0.89 <sup>‡</sup>	0.17			
Active	62	-1.19 <sup>‡</sup>	0.16	-0.40	-0.87, 0.08	0.102
BPI Interference						
Sham	57	-I.I4 <sup>‡</sup>	0.22			
Active	62	− <b>I.84</b> ‡	0.20	-0.70	-1.30, -0.11	0.021
PDQ						
Sham	57	0.06	0.56			
Active	62	-1.63 <sup>†</sup>	0.51	-I.69	-3.20, -0.18	0.028
PDI						
Sham	57	- <b>3.43</b> <sup>†</sup>	1.25			
Active	62	-6.07 <sup>‡</sup>	1.11	-2.64	-5.99, 0.70	0.121
HADS						
Sham	57	-I.0I <sup>†</sup>	0.51			
Active	62	-1.82 <sup>‡</sup>	0.49	-0.81	-2.22, 0.60	0.259
PCS						
Sham	57	-3.53 <sup>‡</sup>	0.86			
Active	62	-3.39 <sup>‡</sup>	0.78	0.15	-2.17, 2.46	0.902

**Notes**: <sup>†</sup>Significant within group improvement at p<0.05; <sup>‡</sup>Significant within group improvement at p<0.001.

Abbreviations: MMRM, mixed-model for repeated measures; SE, standard error; CI, confidence interval; FIQR, Fibromyalgia Impact Questionnaire Revised; BPI, Brief Pain Inventory; PDQ, painDETECT questionnaire; PDI, Pain Disability Index; HADS, Hospital Anxiety and Disability Scale; PCS, Pain Catastrophizing Scale.

and 3-month assessments (Figure 4). There were also significant improvements in pain interference with function (BPI interference subscale), neuropathic symptoms (PDQ) and sleep quality (FIQR sleep item) for active treatment compared to sham (Table 4). The treatment effects estimated by the sensitivity analysis were generally smaller but consistent with those in the original MMRM analysis (Supplemental Table 3).

There was a significant interaction between treatment and pain sensitivity for the baseline to 3-month change in pain intensity (p=0.042) but not for FIQR total score (p=0.245). In the higher pain sensitivity subgroup, the mean change in pain intensity was -2.05 (95% CI [-2.51, -1.60]) for active treatment and -1.05 (95% CI [-1.52, -0.57]) for sham treatment (Figure 3B). The mean difference of -1.01 (95% CI [-1.67, -0.35], p=0.003) was significant and corresponded to an effect size of 0.78 (95% CI [0.25, 1.30]). No group differences were found for lower pain sensitivity (mean -0.25, 95% CI [-0.91, 0.40], p=0.448). Treatment heterogeneity was also detected for the BPI severity subscale (p=0.106) and for PCS (p=0.111).



**Figure 4** Change in FIQR total score (**A**) and FIQR pain item (**B**) from baseline to 6-weeks and 3-months. Error bars indicate SE.  $\Delta$ , treatment effect (Active - Sham). Both treatment arms exhibit improvement from baseline to 6-weeks, however the group difference small and not significant. Between 6-weeks and 3-months, the active treatment arm continues to improve while the sham arm regresses or stays flat, leading to a significant group difference at the study endpoint. **Abbreviations:** FIQR, Fibromyalgia Impact Questionnaire Revised; SE, standard error.

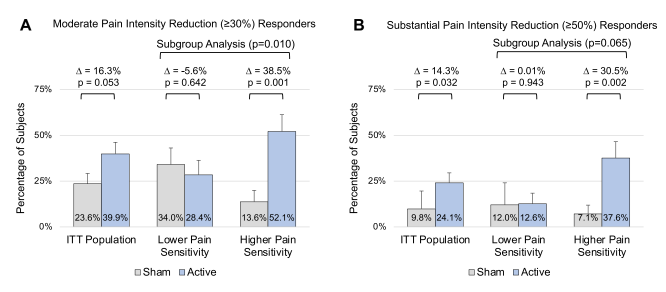


Figure 5 Comparison of pain intensity reduction responder rates, based on FIQR pain item, in the ITT population and for the lower and higher pain sensitivity subgroups. Responder rates for a moderate ( $\geq$ 30%) reduction in pain intensity (**A**). Responder rates for a substantial ( $\geq$ 50%) reduction in pain intensity (**B**). Error bars indicate SE.  $\Delta$ , treatment effect (Active - Sham). Responder rates based on logistic regression analyses. Subgroup analysis p-value is for the treatment-by-pain sensitivity interaction term in the subgroup logistic regression model.

Abbreviations: FIQR, Fibromyalgia Impact Questionnaire Revised; ITT, intention-to-treat; SE, standard error.

## **Responder Analysis**

Responder rates after 3-months of treatment based on the PGIC score and changes in pain intensity (FIQR pain item) were examined. A PGIC responder was defined as a score  $\geq 5$  ("moderately better" symptoms, functional abilities and overall health). Pain intensity responders were defined at the conventional levels of  $\geq 30\%$  reduction

and  $\geq$  50% reduction.<sup>46,72,73</sup> The treatment-by-pain sensitivity interaction term was significant for all responder definitions. For PGIC, the active group had a numerically greater responder rate than sham in the ITT population that was not significant (42.5% vs 34.5%, difference = 8.0%, p=0.372). The treatment difference in the higher sensitivity subgroup was significant (58.0% vs 30.2%, difference = 27.8%, p=0.024). In the lower pain sensitivity subgroup, active treatment had a smaller responder rate (28.2% vs 38.9%, difference = -10.8%, p=0.386). For pain intensity, active treatment had a significantly greater responder rate compared to sham in both the ITT population and in the higher sensitivity subgroup at both the 30% (p=0.053 for ITT) and 50% levels (Figure 5). In all instances, the differences in responder rates between active and sham were larger in the subset of subjects with higher pain sensitivity than in the overall ITT population.

## Blinding

Blinding was assessed by asking the coordinators and subjects whether a low intensity or high intensity device was used. The coordinator identified the correct treatment in 54.7% (95% CI [45.2, 64.2]) of the 103 subjects that completed the study (n=100) or withdrew but provided the 3-month assessment (n=3). The treatment was correctly identified 63.8% (95% CI [51.4, 76.2]) of the time for the active device and 43.8% (95% CI [29.7%, 57.8%]) of the time for the sham device. Of the 99 subjects that completed the 3-month assessment, 86 answered the blinding question. In these subjects, 50.0% (95% CI [39.4, 60.6]) named the correct treatment. Subjects in the active group correctly identified their treatment 17.4% (95% CI [6.4, 28.3]) of the time and subjects in the sham group correctly recognized their treatment 87.5% (95% CI [77.3, 97.7]) of the time. Among all subjects, 84.9% (95% CI [77.3, 92.5]) believed they received a low intensity device.

## Safety

A total of 12 (5 active, 7 sham) adverse events were reported (<u>Supplemental Table 4</u>). They included rash at the site of the device, numbness and tingling, and muscle cramping. Six (3 active, 3 sham) were determined to be related to TENS use, 3 (1 active, 2 sham) were deemed possibly related to TENS use, and 3 (1 active, 2 sham) were judged to be unrelated to TENS use by the principal investigator. The 9 events that were definitely or possibly related to TENS use were minor and self-limited.

## Discussion

This double blind, randomized, sham-controlled trial of wearable TENS did not meet its primary endpoint of a significant group difference in the 3-month PGIC score. The difference of 0.34 points was similar to milnacipran (0.47),<sup>58</sup> duloxetine (0.35-0.87),<sup>59,60</sup> and pregabalin (0.19-0.51),<sup>74-76</sup> which are FDA approved drugs widely

used for management of fibromyalgia. The current study was powered for an effect size of 0.6 and may have been underpowered to detect the small observed PGIC difference. The interpretation of global assessments in persons with fibromyalgia is challenging, likely reflecting the complex and heterogeneous nature of the disease.<sup>77</sup> Another potential complication with PGIC is recency bias (ie, favoring recent events over older ones).<sup>78</sup> As a result, therapeutic interventions should be evaluated by multiple outcomes that include measures anchored to the pretreatment baseline.<sup>77,79</sup> For this reason, FIQR may be a more appropriate primary endpoint for future fibromyal-gia trials evaluating wearable TENS.

Although the primary endpoint was not met, comparisons between the active and sham groups suggest that wearable TENS has specific treatment effects in individuals with fibromyalgia. Overall disease impact decreased in the active group as measured by changes in the Fibromyalgia Impact Questionnaire (FIQR) total score. FIQR is a comprehensive health-related quality-of-life assessment of fibromyalgia and is a core outcome in clinical trials and practice.<sup>80</sup> The items comprising the FIQR high everyday relevance to people with have fibromyalgia.<sup>35,81</sup> The within-group improvement of 13.5 points for active treatment exceeded the minimal clinically important difference of 8 points, which was derived for the earlier FIQ instrument.<sup>82</sup> Pain interference with function decreased in the active group as assessed by the BPI interference subscale. Individuals with chronic pain rank an increased ability to function and improved sleep to be important treatment objectives.<sup>83</sup> IMMPACT recommendations include BPI Interference as a core outcome measure for clinical trials of chronic pain interventions.<sup>55</sup> Neuropathic pain symptoms as assessed by the painDETECT questionnaire decreased. Neuropathic pain is a distressing symptom reported by many individuals with fibromyalgia.<sup>84,85</sup> In many study participants, the specific clinical benefits of active treatment were superimposed on concurrent use of analgesic medications. It is also possible that analgesic use limited the TENS associated reduction in pain intensity, as has been reported for combination pregabalin and duloxetine therapy.<sup>86</sup>

In addition to the findings in the ITT population, the interaction between treatment and pain sensitivity was significant in an MMRM analysis of PGIC, indicating treatment heterogeneity. In subjects with higher pain sensitivity, those receiving active treatment had a 1.2 point greater PGIC score compared to subjects receiving sham

treatment. This difference was significant and clinically meaningful.<sup>72,82,87</sup> This finding is consistent with the hypothesis that TENS is most effective in sensitized pain pathways.<sup>17,18,20,41</sup> It also follows from the observation that placebo pain relief is less effective in individuals with hyperalgesia.<sup>88</sup> The goal of personalized medicine is to identify treatments that are most clinically beneficial and least harmful for each patient based on their individual genetic, physiological, and psychological characteristics.<sup>89</sup> In clinical practice, it may be useful to identify fibromyalgia patients with higher pain sensitivity as optimal candidates for wearable TENS. However, traditional QST is impractical outside of specialized laboratories. As an alternative, it may be possible to predict pain sensitivity using simplified QST methods or from self-reported clinical variables.

Responder analyses demonstrated that over 40% of subjects in the active treatment group experienced clinically meaningful benefits on the individual outcome measures. Forty-three percent (43%) reported that their symptoms, functional abilities, and overall health were at least "moderately better" based on PGIC and 40% experienced a moderate reduction in pain intensity. The responder rates for active treatment exceeded sham treatment by 8 to 16% in the ITT population. The responder rate differences in the higher pain sensitivity subgroup were larger, ranging from 28 to 38%. These differences support a specific treatment effect in individuals using the active device, which may be particularly strong in those with hyperalgesia and central sensitization. The responder rate differences in the lower pain sensitivity subgroup were small. The absolute responder rates and treatment group differences in the ITT population were similar to pregabalin<sup>73,90</sup> and milnacipran.<sup>58</sup>

Most sham-controlled trials of non-invasive electrical stimulation utilize inactive devices as controls, which are difficult to blind.<sup>91,92</sup> In this study, the sham device provided intermittent stimulation totaling 6 minutes during each 1-hour therapy session. This translated to an average of 23 (SD 15) minutes per day for subjects using the sham device. This duration of stimulation is comparable to "active" devices in earlier RCTs that have been criticized for possibly under dosing patients.<sup>25</sup> Given the noticeable stimulation by the sham device, it is unlikely that subjects randomized to the sham group would conclude that they were receiving an inactive placebo. When asked at the end of the study whether they had received a low or high intensity device, nearly all subjects (85%) indicated low

intensity, regardless of the assigned treatment. Therefore, it is unlikely that differential efficacy expectations biased the study results. Fibromyalgia is associated with an enhanced sensitivity to all sensory stimuli.<sup>93</sup> It is possible that many participants assumed their device was low intensity because of a belief that high intensity would cause them discomfort.

The sham response in this study was not surprising as fibromyalgia is associated with substantial placebo effects.<sup>94</sup> Moreover, unlike drug trials that utilize inert placebos, the sham was not passive in order to protect blinding. It is possible that the 6-minutes of stimulation per session and the average of 23 minutes per day had a direct impact on pain perception by modulating pain pathways and through decreased sympathetic activity.<sup>95</sup> It is noteworthy that in contrast to placebo, no-treatment controls in fibromyalgia studies exhibit limited or no improvement.<sup>23,94</sup> Therefore, the within-group improvements reported for active treatment may represent clinical benefits in real-world use.

There have been few high-quality sham-controlled RCTs that examined the efficacy of TENS in fibromyalgia.<sup>24</sup> In a recently published RCT, Dailey and colleagues examined the benefits of TENS applied simultaneously to the lower and upper back for two hours a day for 4-weeks in 301 individuals with fibromyalgia.<sup>23</sup> Their study demonstrated reduced movement evoked and resting pain, decreased fatigue, less pain interference with function and decreased overall disease impact (FIQR total score) relative to both sham TENS and no-TENS controls. Interestingly, the Dailey study showed a significant difference in PGIC responder rates between active and sham treatments that is similar to the higher pain sensitivity subgroup in the present study. Although there were methodological differences between the Dailey RCT and the present RCT, both demonstrated that TENS reduces fibromyalgia associated pain, symptoms, and functional impairment.

The utility of wearable TENS has been examined in additional chronic pain states. The clinical benefits vary with the population characteristics, which is expected of a complex intervention such as TENS.<sup>96</sup> Jamison and colleagues compared wearable TENS against treatment as usual in 68 individuals with chronic low back pain in a RCT.<sup>30</sup> In that study, subjects in the active treatment arm reported significant reductions in pain intensity (BPI severity subscale), pain interference (BPI interference subscale) and pain catastrophizing (PCS) compared to the

control arm after 3-months. However, significant differences were not found for pain-related disability (PDI) and psychological burden (HADS). The positive treatment effects were larger than in the present study, which might be attributable to a comparison to an unblinded control rather than a blinded sham device.<sup>23</sup> Gewandter and colleagues evaluated wearable TENS in 26 patients with CIPN in a 6-week open-label pilot study.<sup>31</sup> Significant improvements were reported for a composite CIPN instrument (EORTC-CIPN20<sup>97</sup>), overall pain quality (Short-Form McGill Pain Questionnaire-2<sup>98</sup>), and pain intensity, tingling, numbness and cramping (latter four outcomes based on 11-point NRS daily dairy). Interestingly, an improvement in large fiber sensation (Utah Early Neuropathy<sup>99</sup>) was also reported. Several observational studies of real-world registry data in heterogenous chronic pain have also been reported.<sup>21,28,29</sup> These studies suggested that wearable TENS was comparatively more effective at reducing pain interference with function than pain intensity.

There are several factors that support generalizability of the treatment effects found in the present study. First, the ITT analyses included all randomized subjects and thus accounted for treatment dropouts that inevitably occur in practice. Second, subjects received limited training by the study coordinators and used their device as recommended by the manufacturer for regular use. Third, utilization levels observed in this study were similar to unsupervised real-world use of the device for chronic lower extremity and low back pain.<sup>29</sup>

## Study Limitations

This study has several limitations that should be considered when interpreting the results. There were only 8 male subjects. This makes it difficult to generalize the findings to men, which may account for at least 10% of fibromyalgia cases.<sup>35,100</sup> Although fibromyalgia was diagnosed by accepted criteria and confirmed by a physician's diagnosis in the medical record, the study did not incorporate an in-person medical history and physical examination. A physical examination is not always necessary to accurately diagnose fibromyalgia,<sup>1</sup> however a concurrent assessment of fibromyalgia may have influenced the makeup of the study population.<sup>101</sup> Data on analgesic use at baseline and changes during the study were collected by self-report, which is subject to recall bias and the ability and willingness of the subjects to provide accurate account. Within these constraints, there was no evidence of group differences at baseline or over the course of the study. Despite being encouraged to not change their fibromyalgia treatments during the study, some subjects may have pursued other nonpharmacological approaches (eg, physical therapy). It is difficult to know how other treatments or environmental factors such as weather<sup>102</sup> might have affected the study. Outside treatments were found to be evenly divided between the active and sham groups, so one group did not have an advantage in receiving more pain-related treatments compared with the other. Finally, treatment effects may be slightly larger in single-center compared to multi-center RCTs.<sup>103</sup>

# Conclusions

Fibromyalgia is a common chronic pain condition that reduces quality of life and is challenging to treat. Nonpharmacological interventions are recommended as firstline therapy.<sup>13</sup> There are three FDA approved drugs, and others off-label including are used opioids. Pharmacological agents are associated with substantial side effects<sup>8,9</sup> and poor adherence.<sup>11,12</sup> This study demonstrated clinical benefits of at-home wearable TENS use in individuals with fibromyalgia over a 3-month period. These included a reduction in pain and somatic symptoms, a decrease in functional impairment, and less overall disease impact. These treatment benefits were obtained with few adverse events that were minor and self-limited. Wearable TENS represents a safe and effective treatment option for people with fibromyalgia.

## **Abbreviations**

BMI, body mass index; BPI, Brief Pain Inventory; CI, confidence interval; CIPN, chemotherapy induced peripheral neuropathy; CPM, conditioned pain modulation; CPT, cold pressor test; FDA, Food and Drug Administration; FIQR, Fibromyalgia Impact Questionnaire Revised; HADS, Hospital Anxiety and Depression Scale; ITT, intention to treat; MAR, missing at random; MI, multiple imputation; MMRM, mixed model for repeated measures; NRS, numerical rating scale; PCS, Pain Catastrophizing Scale; PDI, Pain Disability Index; PDQ, painDETECT questionnaire; PGIC, Patient Global Impression of Change; PPT, pressure pain threshold; QST, Quantitative Sensory Testing; RCT, randomized controlled trial; SD, standard deviation; SE. standard error; TENS. Transcutaneous Electrical Nerve Stimulation.

## **Data Sharing Statement**

De-identified data will be made available to qualified academic researchers by reasonable request to the corresponding author (gozani@neurometrix.com).

## **Ethics Statement**

This study protocol and all amendments were approved by the Human Research Committee (Institutional Review Board) of Mass General Brigham (Massachusetts General Hospital and Brigham and Women's Hospital, Boston, MA, USA). This study followed the principles outlined in the World Medical Association Declaration of Helsinki – Ethical Principles or Medical Research Involving Human Subjects.

## Funding

NeuroMertrix, Inc. provided partial funding for this this trial.

## Disclosure

S. Gozani is an employee and shareholder of NeuroMetrix, Inc. He holds multiple patents related to the Quell device. CJ Gilligan reports Sponsored Research from Mainstay Medical and Sollis, personal fees from Medtronic, personal fees from Abbott, personal fees from Saluda, personal fees from Persica, outside the submitted work. The remaining authors report no potential conflicts of interest for this work.

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