

Therapeutic Effectiveness of AxioBionics Wearable Therapy Pain Management System in Patients with Chronic Lower Back Pain

Philip Muccio¹, Josh Schueller¹, Miriam van Emde Boas², Norm Howe³, Edward Dabrowski⁴ and David Durrant⁵

¹Axiobionics, LLC, Michigan, USA. ²University of Michigan Medical School, Department of Radiology, Nuclear Medicine, Functional Neuroimaging, Cognitive and Mobility Laboratory, Michigan, USA. ³Validation & Compliance Institute, LLC, Michigan, USA. ⁴Oakland University Beaumont School of Medicine, Michigan, USA. ⁵Chicago Neuroscience Institute, Illinois, USA.

Clinical Medicine Insights: Arthritis and Musculoskeletal Disorders Volume 14: 1–11 © The Author(s) 2021 Article reuse guidelines: sagepub.com/journals-permissions DOI: 10.1177/1179544121993778

\$SAGE

ABSTRACT: Chronic lower back pain is one of the most common medical conditions leading to a significant decrease in quality of life. This study retrospectively analyzed whether the AxioBionics Wearable Therapy Pain Management (WTPM) System, a customized and wearable electrical stimulation device, alleviated chronic lower back pain, and improved muscular function. This study assessed self-reported pain levels using the visual analog scale before and during the use of the AxioBionics WTPM System when performing normal activities such as sitting, standing, and walking (n = 69). Results showed that both at-rest and activity-related pain were significantly reduced during treatment with the AxioBionics WTPM System (% reduction in pain: 64% and 60%, respectively; P < .05). Thus, this study suggests that the AxioBionics WTPM System is efficacious in treating chronic lower back pain even when other therapies have failed to sufficiently decrease reported pain levels.

KEYWORDS: Lower back pain, AxioBionics WTPM system, wearable therapy pain management system

RECEIVED: September 19, 2020, ACCEPTED: January 19, 2021.

TYPE: Review

FUNDING: The author(s) disclosed receipt of the following financial support for the research, authorship, and/or publication of this article: Funding for this research project was provided by Axiobionics, LLC.

DECLARATION OF CONFLICTING INTERESTS: The author(s) declared the following potential conflicts of interest with respect to the research, authorship, and/or publication of

this article: Philip Muccio is an owner of Axiobionics, LLC
Josh Schueller is employed by Axiobionics, LLC
Norm Howe is a paid consultant to Axiobionos, LLC
Miriam van Emde Boas did not receive remuneration for her contribution to the paper
Edward Dabrowski did not receive remuneration for his contribution to the paper
David Durrant did not receive remuneration for his contribution to the paper

CORRESPONDING AUTHOR: Philip Muccio, Axiobionics, LLC, 6111 Jackson Rd., Suite 200, Ann Arbor, MI 48103, USA. Email: philipmuccio@axiobionics.com

Introduction

Pain not only affects individuals physically but can have detrimental emotional, mental, and financial effects as well. Approximately 11% of Americans aged 20 years and older experience chronic pain that lasts more than 1 year. This figure increases to 17.7% when including adults who have experienced pain lasting for at least a month. According to a survey, 13% of the US workforce has experienced a loss in productivity, either by decreased productive working hours or time off from a job, and the annual estimated cost of this productivity loss was \$61 billion.

Despite the numerous forms of pain management available, pain relief can be elusive for many individuals. In a survey of more than 2600 Americans with chronic, severe, non-cancer-related pain conducted in 1998 47% reported that they had changed their doctor, and 22% had done so 3 times or more. The most common reason for this change was the intense, remaining pain (42%).^{4,5}

Low back pain (LBP) is defined as pain, muscle tension, or stiffness localized below the costal margin and above the inferior gluteal folds, with or without leg pain (sciatica).⁶ Chronic LBP is a common neurological ailment, second only to headaches, and the most prevalent cause of disability that affects work productivity.¹ Approximately 28% of the patients surveyed with LBP reported a limitation of activity due to their condition.¹ They also had co-morbid manifestations such as loss of sleep,

increased muscle stiffness, and reduced spine mobility, often leading to reduced daily energy and mental focus, depression, and job loss.^{3,7-9} Total annual health care expenditures attributed to back pain are estimated at \$26.1 billion.²

Standard therapies for chronic LBP include administration of pain medications (eg, opiates and non-steroidal anti-inflammatory drugs), physical therapy (PT), chiropractic care, transcutaneous electrical nerve stimulation (TENS), and epidural steroid injections. These treatments can effectively mitigate pain in patients with LBP; however, in those who do not fully respond to them, a combination of treatments may be needed to optimize results.

Pain medications have played a major role in managing chronic LBP for many years; however, according to Chou et al, ¹⁰ there is no medication superior to others because there are complex cost/benefit trade-offs to consider. This conclusion is consistent with another study, suggesting that opioids are not superior to non-opioid medication regarding chronic back pain alleviation. ¹¹ In a recent systematic review and meta-analysis, Busse et al ¹² concluded that opioids have a statistically significant impact, but a small improvement in pain and physical functioning compared to placebo. In another study, the authors concluded that there was no evidence supporting opioid therapy for moderate to severe chronic back pain. Additionally, opioids induce unwanted side effects such as sedation, dizziness, nausea, vomiting, constipation, physical dependence, tolerance, and respiratory depression. ¹³ In general, both short and long-term pain

medication regimens can potentially cause undesirable side effects and health risks, thus contributing to a decrease in quality of life and poor response in pain reduction, both at rest and during physical activity. To further complicate the matter, the use of opioid therapy has contributed to a significant public health crisis in the United States, where more than 63% of the 52 404 drug-related overdose deaths in 2015 were opioid-related. 14,15

PT also has limitations. Patients undergoing various PT-related treatments may not be relieved of pain either after each session or at the end of a multi-month program. Fritz et al16 found no improvement in pain intensity at a 4-week, 3-month, or 1-year follow-up when comparing patients receiving conventional care and those undergoing PT. In a study comparing the McKenzie and Back School methods for treating LBP, Garcia et al¹⁷ reported a 19% and 25% reduction in pain, respectively, while at a 6-month follow-up, there was an average pain intensity of 5.2 between the 2 methods. In a systematic review of PT techniques for managing LBP, the authors concluded that the treatment ineffectively managed pain.¹⁸ In a retrospective study of 4597 patients with LBP, Eleswarapu et al¹⁹ concluded that a substantial percentage of patients did not meet minimal clinically important difference (MCID) for pain and function, following LBP treatment with PT. Fritz also concluded that PT did not result in MCID among adults; however, this study cohort presented recentonset LBP.16

Considering the lack of sufficient pain relief and drawbacks associated with medication and PT, TENS has been considered among the choices for LBP management, especially considering the low risk imposed on patients. However, its efficacy has been questioned by the Department of Health and Human Services due to the lack of supporting scientific evidence.²⁰ In our experience, TENS can fail in pain alleviation for a variety of reasons, including electrode placement errors, electrode fixation issues, low power output, and lack of professional instruction. In a recent systematic review, the benefits of TENS in patients with LBP was found inconclusive because the quality of the studies was low, and adequate parameters and timing of assessment were not uniformly used or reported.²¹ This study highlighted the importance of defining methods, electrical parameters, and quality of research protocols to better understand TENS.

TENS attempts to control pain by blocking the pain pathway proposed in the Gate Control Theory. ²² TENS is designed to stimulate only the cutaneous afferent nerves, without penetrating deeper into the body to stimulate and relax muscle spasms. This may be 1 reason some patients experience spasm relief while others do not. In a study of the direct and indirect benefits of TENS, Gladwell et al, ²³ found that while 89% of subjects with musculoskeletal pain had pain relief from TENS, only 71% had relief of muscle spasms. An additional drawback of TENS is that large electrodes cannot be used to cover a large area of pain as it could significantly reduce the current output

and distribution, thereby conferring limited pain relief. In addition to the deficiencies in its underlying principles, standard TENS technology, which uses adhesive skin electrodes, can fail to lower pain due to various issues. Patients with LBP commonly experience difficulty in placing the electrodes on the lower back due to the inability to twist their body or see the affected area. Even if patients succeed in placing the electrodes, there is no guarantee that the electrodes will be placed accurately or that they will stay affixed throughout the day. Sweat and body hair can interfere with electrode adherence, conductance, and performance. Furthermore, patients can experience contact dermatitis due to the chemicals present in the electrodes, which, for example, are added for adherence of electrodes to the skin.²⁴ A high prevalence of skin irritation has been noted in 40% after the use of $TENS^{25}$ and which can lead to rejection and failure of the therapy.

Although TENS can be useful in managing pain when being on the body, it loses its effect when removed, and there is no long-term, residual carry-over benefit. A meta-analysis of patients with chronic LBP showed that TENS was significantly different from placebo/control (P<.02) and that TENS/interferential stimulation intervention was better than placebo/control during therapy (P=.02), but not immediately after therapy (P=.08), or 1-3 months after (P=.99). The study results imply that keeping TENS and its electrodes affixed to the body with an active stimulus is the best way to achieve pain relief.

The output signal generated by TENS is much lower than that of neuromuscular electrical stimulation (NMES). There is growing evidence that NMES on trunk muscles in patients with LBP may increase core muscle strength, reduce pain, and improve function when compared to conventional care. The total charge generated is calculated by multiplying the amplitude and pulse width. TENS, with a maximum amplitude of 60 mA and 150 µs pulse width, produces a total charge of 9 μColoumbs. In contrast, NMES, with a maximum amplitude of 100 mA and a pulse width of 300 µs, produces a total charge of $30\,\mu\text{Coloumbs}$ and therefore is 3.3 times stronger than TENS. To stimulate muscle, an NMES signal must penetrate deeper than subcutaneous tissue to depolarize motor neurons and must be of sufficient current intensity to reach back muscles such as the multifidus muscle.²⁷ A deeply penetrating electrical signal is more effective at reducing pain than TENS in a comparative study of varying electrical stimulation devices.²⁸ Interestingly, there is scientific evidence for combining TENS and NMES into a treatment modality for pain. Moore and Shurman²⁹ assessed the use of TENS, NMES, and TENS-NMES and concluded that their combination was superior to TENS or NMES alone at reducing chronic back pain.

There is a growing body of evidence suggesting that NMES may be beneficial as a modality to strengthen muscles as well as to facilitate a reduction in chronic LBP.³⁰ In one study, electrodes were positioned over the abdominal and lower back paraspinal muscles using intensities sufficient to produce muscle

contraction. The results revealed clinically and statistically significant improvements in both muscle groups' performance, and the results were associated with significant improvements in self-reported pain levels, suggesting that NMES plays an important role in chronic LBP rehabilitation. Another study showed that electrical stimulation significantly increased (P<.05) the endurance of the back muscles compared with the control and exercise groups. The study showed that electrical stimulation could be a valuable treatment in the early care of patients with LBP in maintaining and increasing strength and endurance of back muscles when a more active exercise program is too painful to perform.

Volitional exercise may not provide enough tension through the atrophied type II fibers to induce hypertrophy of these fibers even when performed at near maximum intensities.³³ In a Cochrane review investigating physical activity and exercise, exercise did not consistently cause any change (positive or negative) in self-reported pain scores.³⁴ However, NMES has been shown to recruit type II muscle fibers and induce an increase in muscle strength,^{29,35-37} and this may have a beneficial effect on chronic LBP and function.^{33,38}

Not all studies managed to show benefit from the use of NMES in chronic LBP. In a study by Guo et al, ³⁹ there was no statistical difference between the NMES and control groups regarding pain reduction or disability score after a 4-week treatment period. However, the authors indicated that the study had limitations that should be overcome to produce a more confident conclusion in the future. NMES dose may need to be greater than that implemented in this study to effectuate positive changes in LBP.

An alternative to conventional care is needed to improve pain modulation and reduce the potential risks for patients. Thus, we investigated the effectiveness of the AxioBionics Wearable Therapy Pain Management (WTPM) System in the treatment of chronic LBP. This system employs both NMES and TENS technologies, which are integrated into a wearable system that automatically aligns electrodes on the body and keeps them affixed to it. It is hypothesized that the WTPM System would reduce at-rest and activity-based LBP

The WTPM System employed both TENS and NMES to maximize the therapeutic result. The TENS modality (low-level, sensory stimuli) was employed to alleviate pain and the NMES (moderate-level, supra-sensory, supra-motor threshold) to help reduce muscle spasms and atrophy^{27,33} and enhance the pain alleviation effect over time. It has long been acknowledged that conditioning of the trunk muscles is effective in the management of acute and chronic LBP.²⁷ Rather than relying solely on TENS, the WTPM System sought to improve patient outcomes by combining these 2 modalities into a wearable system. The WTPM System assured proper placement and fixation of electrodes to ensure that electrodes were aligned over areas of pain and motor points. To our knowledge, this is the

first study to delineate the effects of an electrical stimulation system on at-rest and activity-based pain.

Methods

Human Subjects Research: This study has been determined by the Institutional Review Board to be Exempt according to FDA 21 CFR 56.104 and 45CFR46.104(b)(4): (4) Secondary Research Uses of Data or Specimens on 06/16/2020. IRB Review provided by Gretchen Parker, PhD, RAC, CIP IRB Chair, Pearl IRB 29 East McCarty Street, Suite 100 Indianapolis, IN 46225.

Patients: Data from all patients who were treated with AxioBionics WTPM System were reviewed for inclusion in this retrospective analysis.

Inclusion Criteria: Sixty-nine adults (27 women and 42 men; age, 27–90 years) from 8 mid-western cities in the United States with at least 1 year of axial LBP alone or axial LBP with radicular pain to the buttocks or thigh, unilaterally or bilaterally (10-point visual analog scale [VAS] score, 4-10) were included in this study.

Exclusion Criteria: Patients with a history of pain for less than 1 year were excluded from treatment with the WTPM System. Additionally, patients were excluded if they had a cardiac condition, that is, requiring implantation of a demand cardiac pacemaker or defibrillator, atrial or ventricular fibrillation, congestive heart failure, recent myocardial infarction (<3 months), or hemodynamically significant valvular dysfunction. Moreover, pregnant women (safety of muscle stimulation during pregnancy is unknown) and patients with epilepsy, dementia, severe receptive or global aphasia that confounds testing and training (operationally defined as unable to follow 2-point commands that leads to an unlikely ability to understand the protocols and procedures), and active cancer were excluded from treatment.

Procedure: VAS was used to measure the AxioBionics WTPM System's at-rest and activity-based LBP alleviation capability. At-rest was defined as lying down or remaining immobile. Activity-based was defined as any position requiring muscle activity or body movement. All patients underwent a baseline pain assessment both at-rest and during activity. At VAS assessment, each patient had to choose a pain level that best described their average pain at rest and during activity: sitting (longer than 15 minutes), standing, and walking. Prior treatments were recorded to investigate whether there was a difference in outcome between patients having received a given therapy or combination of therapies prior to the use of the AxioBionics WTPM System. These included pain medications, PT, epidural steroid injections, manipulations, surgery, and TENS.

System Evaluation and Fitting: The AxioBionics WTPM System incorporates electrodes in a custom-fitted, lycra-spandex garment and combines NMES and TENS stimuli. The stimulator used in the WTPM System was a Focus 2 channel NMES/TENS device $(300\,\mu seconds\ pulse\ width,\ 50\,Hz,$



Figure 1. The method used to stimulate the area of pain during the NMES/TENS trial period.

10 seconds on, 10 seconds off, synchronous mode, manufacturer: EMPI, St. Paul, MN) Patients initially underwent an evaluation of their medical history, assessment of pain distribution, initial electrode set-up on the body (without garment), and a trial of NMES/TENS in the clinic, and instructions on appropriate intensity levels were given to produce mild to moderate muscle contraction in 1 or more muscles that included the lower back paraspinal muscles, gluteus maximus, and hamstrings, unilaterally or bilaterally. Guidance on stimulus level was intended to evoke both sensory fibers for pain control and alpha motor neurons for relaxation of muscle spasms and blood flow in the area. The patients were asked to use the NMES/TENS device with conventional 2" × 4" adhesive electrodes at home for 4weeks and advised to place the electrodes on or around the area of pain that provided the best pain relief (Figure 1). A wearing schedule was provided that started with 1 hour the first day and then increased by 1 hour each subsequent day until the patient was able to wear the stimulus as much as could be tolerated or needed to control pain. No limitations on wear time were given. Patients were encouraged to wear the system during the day and when sleeping if they had sleep disturbance from pain. Patients were free to choose their wear schedule.

After the 4-week home trial, the patients returned to AxioBionics for follow-up, and the results of the NMES/TENS trial were documented. The test garments were fabricated from the measurements of the patients and individually tailored for a secure fit. The fitting process led to the customization of the WTPM Systems that captured the array of 4 electrodes that optimized pain relief for each patient. Each patient received a WTPM System configured as a belt, a brief, belt/sleeve, or shorts (Figures 2 and 3). For those who purchased the WTPM System following the 4-weeks, additional

data were collected for another 4 weeks. No further data collection was performed on patients who did not purchase the system after the 4-week trial.

Data Analyses: All statistical analyses were performed using SPSS Statistics 17.0 (IBM Corp., Armonk, N.Y., USA). Descriptive statistical analysis using means, standard deviations, confidence intervals, and proportions were employed to describe the baseline characteristics of the combined sample of all patients. A P-value < .05 was used in all statistical analyses to indicate statistical significance. To evaluate the effectiveness of AxioBionics WTPM System to decrease at-rest and activity-related pain, t-tests were used to determine the significance of the decrease in pain on VAS. Univariate analysis of variance (ANOVA) was used to determine changes in pain levels between the groups (pain medications, PT, epidural steroid injections, surgery, TENS, manipulations) and among all combinations of patient categories.

Results

Table 1 shows the distribution of past treatments that patients underwent before treatment with the WTPM System.

Tables 2 and 3 show the change in pain levels before and during treatment with the WTPMS at-rest and with activity, respectively and whether the treatment met minimal important difference (MID).

Table 4 shows the percentage of patients who achieved 100% reduction in pain during the trial period, when fitted with the WTPMS and the combination.

Table 5 shows a comparison of the increase in pain related to activity with standard care and with the WTPMS.

A 1-way univariate analysis of variance test was performed to assess the changes in pain between all combinations of groups (Table 6). Combinations not listed had insufficient sample size for statistical calculations. Significant differences are not present among the listed categories or the combinations of categories.

Figures 4a and b show the mean pain scores for all patients at baseline with the WTPM treatment and changes in pain scores. Both at-rest and activity-related pain significantly reduced during treatment with the AxioBionics WTPM System (P-values < .05). Activity-related pain reduced more than at-rest pain (P< .05) when assessed using a 10-point VAS score; however, baseline activity-related pain was significantly greater than baseline at-rest pain (P< .05). In terms of percent reduction, the change in pain scores was more with at-rest pain than with activity-related pain. Both at-rest and activity-related pain categories showed improvement in the VAS score from the trial period to that with the use of the WTPM System. Overall, 32% of patients in the at-rest category achieved 100% pain relief, whereas only 8.7% of patients in the activity-related pain category did so.

Figure 5 shows the number of patients who were stratified into each of the 3 categories of pain (low, medium, and high)

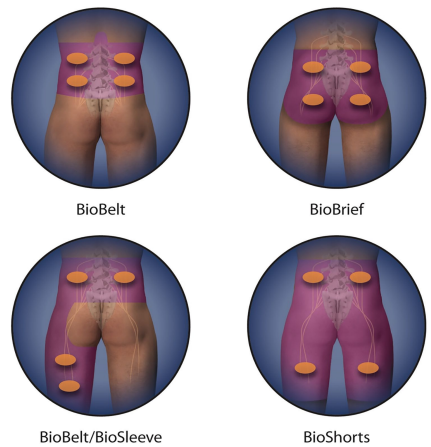


Figure 2. Various configurations of the WTPM system.



Figure 3. Actual depiction of the belt version of the WTPM system used to treat LBP.

based on their VAS scores. Although the bulk of the patients belonged to the high pain group before treatment, most of them were reassigned to the low or medium pain groups during the WTPM treatment.

Additional analyses were performed on the data of the study subgroups. Among the 56 patients whose data of at-rest pain (sitting) were recorded before the use of the WTPM System, all patients had difficulty in sitting for varying periods, ranging from 1 minute to 8 hours (Figure 6). Forty-nine of these patients reported improvement in their sitting ability using the WTPM System. The minimum improvement in sitting time was 50%. Figure 3 shows the number of patients in each range.

Standing ability was also limited before the use of the WTPM System in all assessed patients (n=49); times are shown in Figure 4. During the use of the WTPM System, 97.7% of the patients (n=44) reported standing improvement. The minimum improvement in the amount of time the patients could stand was 40% (n=20); the periods are shown in Figure 7.

Overall, 96% of the patients who were investigated (n=50) reported interruptions in their sleep. All 29 patients who were asked reported sleep improvements. All patients reported

Table 1. Involvement in past treatments (N=69).

	PAIN MEDICATIONS	PHYSICAL THERAPY	EPIDURAL STEROID INJECTIONS	SURGERY	TENS	MANIPULATION
N	66	62	36	27	21	14
%	96	90	52	39	30	20

Forty-eight of the 69 patients (70%) underwent the 4-week trial period and subsequently were fitted with the WTPM System, whereas 21 patients (30%) only underwent the 4-week trial

Abbreviations: N, number of patients; TENS, transcutaneous electrical nerve stimulation.

Table 2. Change in pain levels (at-rest).

N	CATEGORY OF TX	PAIN LEVEL BEFORE TX (SD)	PAIN LEVEL WITH TX (SD)	NUMERICAL CHANGE IN PAIN (SD)	% CHANGE IN PAIN	MINIMAL IMPORTANT DIFFERENCE (MID) ACHIEVED?
21	Trial only	6.4 (1.7)	2.6 (1.9)	3.9 (1.5)	60%*	Yes*
48	Fitted	6.4 (1.7)	2.1 (2.1)	4.4 (1.7)	68%*	Yes*
69	Combined	6.4 (1.7)	2.3 (2.0)	4.1 (1.7)	64%*	Yes*

Abbreviations: Tx, treatment; N, number of patients.

*P < .001 The P values were determined by Mann-Whitney U Test (AKA Wilcoxon rank-sum test). MID values determined by Wikström and Edelstam²⁶ and Jensen et al⁴⁰ achieved at 50% and 33% reduction in VAS Pain Scores, respectively.

Table 3. Change in pain levels (during activity).

N	CATEGORY OF TX	PAIN LEVEL BEFORE TX (SD)	PAIN LEVEL WITH TX (SD)	NUMERICAL CHANGE IN PAIN (SD)	% CHANGE IN PAIN	MINIMAL IMPORTANT DIFFERENCE (MID) ACHIEVED?
21	Trial only	9.0 (1.1)	3.8 (1.7)	5.2 (1.7)	58%*	Yes*
48	Fitted	8.8 (1.2)	3.3 (2.2)	5.4 (2.0)	62%*	Yes*
69	Combined	8.9 (1.2)	3.5 (2.1)	5.3 (1.9)	60%*	Yes*

Abbreviations: Tx, treatment; N, number of patients.

*P < .001 The P values were determined by Mann-Whitney U Test (AKA Wilcoxon rank-sum test). MID values determined by Wikström and Edelstam²⁶ and Jensen et al⁴⁰ achieved at 50% and 33% reduction in VAS Pain Scores, respectively.

Table 4. Percent of patients who achieved 100% pain relief.

N	CATEGORY OF TX	N ACHIEVING 100% RELIEF (AT-REST)	% ACHIEVING 100% RELIEF (AT-REST)	N ACHIEVING 100% RELIEF (ACTIVITY)	% ACHIEVING 100% RELIEF (ACTIVITY)
21	Trial period	5	24%	0	0%
48	Fitted	17	35%	6	13%
69	Combination	22	32%	6	8.7%

Abbreviations: N, number of patients; Tx, treatment.

Table 5. Comparison of the increase in activity-related pain.

	AVG PAIN WITH STANDARD CARE (SD)	AVG PAIN WITH WTPM SYSTEM (SD)
At-rest	6.4 (1.7)	2.3 (2.0)
Activity	8.9 (1.2)	3.5 (2.2)
Numerical Change	2.5	1.2

Abbreviations: WTPM, Wearable Therapy® Pain Management; Avg, average.

walking and concentration improvements, as well as decreased muscle spasm and stiffness; however, there were no consistent measures used that provided more quantitative results across the patient population. A graphical representation of the WTPM System pathway is provided in Figure 8.

Discussion

This study retrospectively investigated the data of all patients who were treated with the AxioBionics WTPM System for chronic LBP and elucidated the impact of the WTPM system on both at-rest and active-movement pain. Data collected from 69

Table 6. One-way univariate analysis of variance test results for the changes in pain between all combinations of groups.

CATEGORY	AT-REST CHANGE	ACTIVE CHANGE
	P-VALUE	P-VALUE
Corrected model	.880	.337
Age (young ≤55 years, old >55 years)	.894	.971
PainMeds	.346	.536
PhysTherapy	.661	.515
Manipulation	.708	.817
Injections	.514	.179
Surgery	.808	.322
TENS	.272	.680
AgeCategory * physTherapy	.429	1.000
AgeCategory * manipulation	.966	.535
AgeCategory * injections	.637	.160
AgeCategory * surgery	.412	.183
AgeCategory * TENS	.693	.787
PhysTherapy*injections	.203	.700
PhysTherapy*TENS	.932	.912
Manipulation*injections	.366	.957
Manipulation*surgery	.645	.674
Manipulation*TENS	.583	.692
Surgery*TENS	.719	.651
AgeCategory*manipulation*TENS	.670	.875
AgeCategory*injections*surgery	.920	.974
AgeCategory*surgery*TENS	.507	.159
Manipulation*injections*surgery	.578	.841
Manipulation*injections*TENS	.665	.659

Abbreviations: AgeCategory, the category of the age; PainMeds, pain medications; PhysTherapy, physical therapy; TENS, transcutaneous electrical nerve stimulation.

patients, diagnosed only with LBP, were analyzed to determine the therapeutic effectiveness of the AxioBionics WTPM System to relieve LBP. The results showed that the WTPM System significantly decreased both at-rest and activity-related pain; given that these patients came to AxioBionics after undergoing conventional therapies and treatments, the outcome of this study suggested that the WTPM system was more effective at alleviating LBP in patients who did not respond well to conventional treatments. Our results suggest that high-level active pain can be adequately controlled with the WTPM System, perhaps better than common treatments for chronic LBP.

Our results showed a significant 2.5 level VAS increase in patient movement-related pain relative to the at-rest category in the standard treatment grouping, whereas, in the patients using the WTPM System, there was only a minor increase of 1.2. The fact that the WTPM System lowered both at-rest and activity-based pain is a substantial finding and may be the reason for the concomitant increase in the ability to sit and stand for a longer period while wearing the WTPM System during activities of daily living. It also suggests the importance of keeping electrical stimulation placed on the body with the stimulus on, to ensure that activity-related pain is held in check. We can postulate from these results that TENS/ NMES is more effective at pain relief when operating on the body than when it is removed, and that it can play a significant role in helping patients remain active for a longer period. Resende et al²¹ showed that pain relief from TENS abates when the stimulus is removed, whereas, our results indicate that it should remain on the body as much as possible to achieve the optimum relief either at-rest or when physically active. Although this study did not compare the efficacy of adhesive electrodes and the WTPM System, keeping electrodes firmly affixed to the body with a garment system may significantly help patients achieve longer wear time without losing electrode contact. Future studies should investigate the electrode interface to elucidate the effectiveness of these 2 systems.

Another significant finding of this study was the number of patients who reported no pain when wearing the WTPM System when at-rest (32%) and when active (8.7%), and while no one reported having no pain in the active group during the WTPM System trial, 13% of those who were fit with the WTPM System reported having no pain despite an activity increase.

Pain relief significance achieved with the WTPM System for LBP

Two researchers have derived standards for meaningful pain reduction that we can use as measures. Wickström and Edelstam studied women with endometriosis and found that, to achieve a minimally important difference, a 50% pain reduction had to be achieved. Jensen, et. al. studied postoperative pain and concluded that to meet clinical significance, a 33% pain reduction had to be achieved. In this study, both at-rest and activity-related LBP reductions averaged at 64% and 60%, respectively, which is well above these 2 standards.

As can be seen from these results, pain can fluctuate from a lower to a higher value when patients become more active; therefore, it is important to know if treatment for pain can impact both values and whether that reduction will lead to an improvement in function and work capability. Our results show a concomitant increase in function (sitting and standing) when pain is lowered and held in check as movement increases.

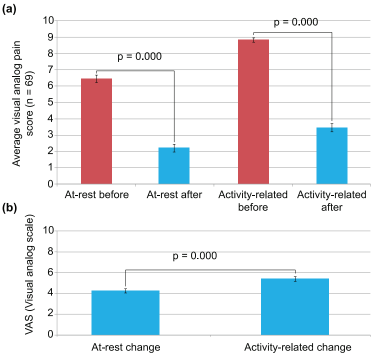


Figure 4. (a) AxioBionics Wearable Therapy Pain Management System significantly decreases both at-rest and activity-related pain, and (b) change in activity-related pain was significantly greater than that in at-rest pain with the use of the AxioBionics Wearable Therapy Pain Management System.

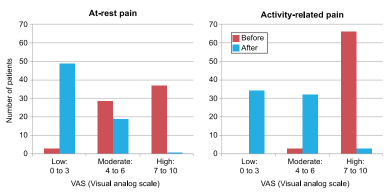


Figure 5. Separation of the visual analog scale into 3 categories. At-rest and activity-related pain are shown by the number of patients before and during treatment with the Wearable Therapy Pain Management System.

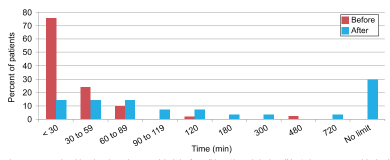


Figure 6. Percentages of patients categorized by the time they could sit before (N=50) and during (N=27) treatment with the Wearable Therapy Pain Management System.

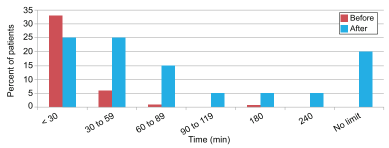


Figure 7. Percentages of patients categorized by the time they could stand before (N=41) and during treatment (N=20) with the Wearable Therapy Pain Management System. The minimum increase in standing time was 40%.

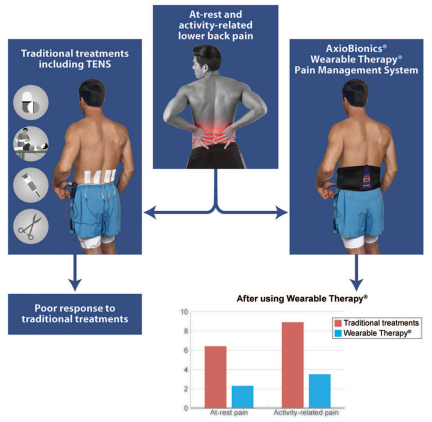


Figure 8. Graphical representation of the WTPM system pathway.

It is generally understood that not all treatments for chronic LBP will result in a favorable outcome. This study managed to show that, even when pain relief is inadequate from standard clinical options, such as pain medications, PT, and TENS, the WTPM System is a reasonable and efficacious alternative solution. The results also suggest that the WTPM System is equally effective at alleviating LBP in patients with and without a pain medication regimen.

Following, we present a brief discussion regarding possible mechanisms of action regarding pain improvement, following this study's supporting evidence of its null hypothesis (reduced LBP). We present 7 mechanisms that could contribute to pain reduction; (1) gate control theory; (2) muscle spasm reduction; (3) endogenous opiates; (4) muscle strengthening from increasing physical activity; (5) muscle strengthening derived from stimulated muscle; (6) improved spinal posture; and (7) increase in intra-abdominal pressure from the WTPM garment. Furthermore, longer sitting time (>7 hours/day) further increases the risk of LBP.⁴¹ Lessening sitting time, as well as sedentarism is, therefore, a reasonable pain reduction objective.

It is beyond the scope of this study to differentiate the contribution of each mechanism; however, there may be an

additive effect. For example, conventional TENS may not be equally efficacious with this new system. Furthermore, all actions invoke pain relief by naturally-occurring means and not by pharmaceutical or invasive procedures; therefore, the potential for patient harm is significantly reduced.

Although the use of the AxioBionics WTPM System showed pain improvement during sitting, standing, walking, concentration, muscle stiffness, and sleep, the type of recorded data was not consistent within each activity group. This limited the number of quantitative analyses that could be performed in this retrospective study. Therefore, future studies should include consistent and quantitative measures to elucidate the effectiveness of the AxioBionics WTPM System in LBP treatment. Finally, the sample sizes of some subpopulations in this study were not large enough to be evaluated for differences in the effectiveness of the WTPM System. Therefore, additional studies should include larger sample populations to determine whether combinations of different treatments affect the outcome of the WTPM System regarding LBP. Although this was a retrospective analysis and did not include a sham or control device, future studies would benefit from inclusion of a sham TENS/NMES WTPMS to control for the placebo effect. It should also include patientspecific pain medications and dosages to elucidate the medications' pain-relieving effects compared to the WTPM System. Future studies should also include prospective RCTs to establish unbiased evidence of the effectiveness of this unique approach.

Conclusion

The WTPM System has the potential to provide substantial pain alleviation in patients with chronic LBP when conventional treatments fail to deliver adequate relief. It eliminates the problems that have affected surface electrode use for TENS/NMES over the years by eliminating wrong electrode placement, electrode placement difficulty, electrode to skin surface integrity, and electrode failure. Unlike many conventional treatments for LBP, the WTPM System can be used independently, non-invasively, non-pharmacologically, and for any desired period.

Acknowledgements

We wish to thank David Durrant, D.C., Ph.D. for his science advisory and technical support.

ORCID iD

Philip Muccio https://orcid.org/0000-0002-3058-9640

REFERENCES

- National Center for Health Statistics (US). Health, United States, 2006: With Chartbook on Trends in the Health of Americans. National Center for Health Statistics (US); 2006.
- Luo X, Pietrobon R, Sun SX, Liu GG, Hey L. Estimates and patterns of direct health care expenditures among individuals with back pain in the United States.

- Spine (Phila Pa 1976). 2004;29:79-86.
- Stewart WF, Ricci JA, Chee E, Morganstein D, Lipton R. Lost productive time and cost due to common pain conditions in the US workforce. *JAMA*. 2003;290:2443-2454.
- Institute of Medicine (US) Committee on Advancing Pain Research, Care, and Education. Relieving Pain in America: A Blueprint for Transforming Prevention, Care, Education, and Research. (presentation). 2020. Updated 2020. Accessed July 4, 2020. https://www.ncbi.nlm.nih.gov/books/NBK91497/pdf/Bookshelf_ NBK91497.pdf
- Institute of Medicine; Board on Health Sciences Policy; Committee on Advancing Pain Research, Care, and Education. Relieving Pain in America: A Blueprint for Transforming Prevention, Care, Education, and Research. The National Academies Press (US); 2011.
- Ramasamy A, Martin ML, Blum SI, et al. Assessment of patient-reported outcome instruments to assess chronic low back pain. *Pain Med.* 2017;18: 1098-1110.
- Hardt J, Jacobsen C, Goldberg J, Nickel R, Buchwald D. Prevalence of chronic pain in a representative sample in the United States. Pain Med. 2008;9:803-812.
- Kelly GA, Blake C, Power CK, O'keeffe D, Fullen BM. The association between chronic low back pain and sleep: a systematic review. Clin J Pain. 2011;27: 169-181.
- Heffner KL, France CR, Trost Z, Ng HM, Pigeon WR. Chronic low back pain, sleep disturbance, and interleukin-6. Clin J Pain. 2011;27:35-41.
- Chou R, Huffman LH; American Pain Society; American College of Physicians. Medications for acute and chronic low back pain: a review of the evidence for an American Pain Society/American College of Physicians Clinical Practice Guideline. Ann Intern Med. 2007;147:50 5-514.
- Krebs EE, Gravely A, Nugent S, et al. Effect of opioid vs nonopioid medications on pain-related function in patients with chronic back pain or hip or knee osteoarthritis pain: the SPACE randomized clinical trial. *JAMA*. 2018;319: 872-882.
- 12. Busse JW, Wang L, Kamaleldin M, et al. Opioids for chronic noncancer pain: a systematic review and meta-analysis. *JAMA*. 2018;320:2448-2460.
- Benyamin R, Trescot AM, Datta S, et al. Opioid complications and side effects. Pain Physician. 2008;11:S105-S120.
- Rudd RA, Aleshire N, Zibbell JE, Matthew Gladden R. Increases in drug and opioid overdose deaths—United States, 2000–2014. MMWR Morb Mortal Wkly Rep. 2016;16:1323-1382.
- Vadivelu N, Kai AM, Kodumudi V, Sramcik J, Kaye AD. The opioid crisis: a comprehensive overview. Curr Pain Headache Rep. 2018;22:16. doi:10.1007/ s11916-018-0670-z
- Fritz JM, Magel JS, McFadden M, et al. Early physical therapy vs usual care in patients with recent-onset low back pain: a randomized clinical trial. *JAMA*. 2015;314:1459-1467.
- Garcia AN, Costa LD, da Silva TM, et al. Effectiveness of back school versus McKenzie exercises in patients with chronic nonspecific low back pain: a randomized controlled trial. Phys Ther. 2013;93:729-747.
- Cuenca-Martinez F, Cortés-Amador S, Espí-López GV. Effectiveness of classic physical therapy proposals for chronic non-specific low back pain: a literature review. Phys Ther Res. 2018;21:16-22.
- Eleswarapu AS, Divi SN, Dirschl DR, Mok JM, Stout C, Lee MJ. How effective is physical therapy for common low back pain diagnoses? a multivariate analysis of 4597 patients. Spine (Phila Pa 1976). 2016;41:1325-1329.
- Centers for Medicare & Medicaid Services. Decision Memo for Transcutaneous Electrical Nerve Stimulation for Chronic Low Back Pain (CAG-00429N). 2020. Updated 2020. Accessed May 27, 2020. https://www.cms.gov/medicarecoverage-database/details/nca-decision-memo.aspx?NCAId=256
- Resende L, Merriwether E, Rampazo ÉP, et al. Meta-analysis of transcutaneous electrical nerve stimulation for relief of spinal pain. Eur J Pain. 2018;22: 663-678.
- Melzack R, Wall PD. Pain mechanisms: a new theory. Surv Anesthesiol. 1967;11:89-90.
- Gladwell PW, Badlan K, Cramp F, Palmer S. Direct and indirect benefits reported by user of transcutaneous electrical nerve stimulation for chronic musculoskeletal pain: qualitiative exploration using patient interviews. *Phys Ther*. 2015;95:1518-1528.
- Weber-Muller F, Reichert-Penetrat S, Schmutz JL, Barbaud A. Contact dermatitis from polyacrylate in TENS electrode. *Ann Dermatol Venereol*. 2004;131:478-480. [Article in French].
- Green T, Pye RJ. Skin reactions to TENS equipment used for pain relief. Br J Dermatol. 1991;125:35.
- Wickström K, Edelstam G. Minimal clinically important difference for pain on the VAS scale and the relation to quality of life in women with endometriosis. Sex Reprod Healths. 2017;13:35-40.
- Sions JM, Crippen DC, Hicks GE, Alroumi AM, Manal TJ, Pohlig RT.
 Exploring neuromuscular electrical stimulation intensity effects on multifidus muscle activity in adults with chronic low back pain: an ultrasound

- $imaging-informed\ investigation.\ {\it Clin\ Med\ Insights\ Arthritis\ Musculoskelet\ Disord.}$ 2019;12:1179544119849570.
- Rajfur J, Pasternok M, Rajfur K, et al. Efficacy of selected electrical therapies on chronic low back pain: a comparative clinical pilot study. *Med Sci Monit*. 2017;23:85-100.
- Moore SR, Shurman J. Combined neuromuscular electrical stimulation and transcutaneous electrical nerve stimulation for treatment of chronic back pain: a double-blind, repeated measures comparison. *Arch Phys Med Rebabil.* 1997;78:55-60.
- Wigerstad-Lossing I, Grimby G, Jonsson T, Morelli B, Peterson L, Renström P. Effects of electrical muscle stimulation combined with voluntary contractions after knee ligament surgery. Med Sci Sports Exerc. 1988;20:93-98.
- Coghlan S, Crowe L, McCarthypersson U, Minogue C, Caulfield B. Neuromuscular electrical stimulation training results in enhanced activation of spinal stabilizing muscles during spinal loading and improvements in pain ratings. Conf Proc IEEE Eng Med Biol Soc. 2011;2011:7622-7625.
- Kahanovitz N, Nordin M, Verderame R, et al. Normal trunk muscle strength and endurance in women and the effect of exercises and electrical stimulation. Part 2: comparative analysis of electrical stimulation and exercises to increase trunk muscle strength and endurance. Spine (Phila Pa 1976). 1987;12:112-118.
- Hicks GE, Sions JM, Velasco TO, Manal TJ. Trunk muscle training augmented with neuromuscular electrical stimulation appears to improve function in older adults with chronic low back pain: a randomized preliminary trial. Clin J Pain. 2016;32:898-906.

- Geneen LJ, Moore RA, Clarke C, Martin D, Colvin LA, Smith BH. Physical activity and exercise for chronic pain in adults: an overview of Cochrane Reviews. Cochrane Database Syst Rev. 2017;1:CD011279.
- Cabric M, Appell HJ, Resic A. Fine structural changes in electrostimulated human skeletal muscle. Evidence for predominant effects on fast muscle fibres. Eur J Appl Physiol Occup Physiol. 1988;57:1-5.
- Gregory CM, Bickel CS. Recruitment patterns in human skeletal muscle during electrical stimulation. *Phys Ther.* 2005;85:358-364.
- Sinacore DR, Delitto A, King DS, Rose SJ. Type II fiber activation with electrical stimulation: a preliminary report. Phys Ther. 1990;70:416-422.
- Glaser JA, Baltz MA, Nietert PJ, Bensen CV. Electrical muscle stimulation as an adjunct to exercise therapy in the treatment of nonacute low back pain: a randomized trial. J Pain. 2001;2:295-300.
- Guo P, Wang JW, Tong A. Therapeutic effectiveness of neuromuscular electrical stimulation for treating patients with chronic low back pain. *Medicine (Balti-more)*. 2018;97:e13197.
- Jensen MP, Chen C, Brugger A. Interpretation of visual analog scale ratings and change scores: a reanalysis of two clinical trials of postoperative pain. J Pain. 2003;4:407-414.
- Park SM, Kim HJ, Jeong H, et al. Longer sitting time and low physical activity are closely associated with chronic low back pain in population over 50 years of age: a cross-sectional study using the Sixth Korea National Health and Nutrition Examination Survey. Spine J. 2018;18:2051-2058.