Research Article

Effects of Rhythmic Sensory Stimulation on Ehlers–Danlos Syndrome: A Pilot Study

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Ehlers–Danlos syndrome (EDS) is a connective tissue disorder characterized by joint hypermobility and skin extensibility and is often accompanied by chronic pain. Rhythmic sensory stimulation (RSS) can be defined as the stimulation of the senses in a periodic manner within a range of low frequencies. Music plus sound delivered through a vibroacoustic device is a form of RSS and has demonstrated utility in managing pain. In this current study, we conducted an open-label pilot study of 15 patients with hypermobile EDS using RSS as the intervention. Posttreatment improvements were seen in 11 of the 15 patients (73%), whereas 3 of the 15 patients (20%) experienced worse outcomes. Of the 14 patients that completed the experiment, 6 participants (43%) were classified as “responders” to the device while 8 participants (57%) were classified as “nonresponders.” Responders demonstrated significant improvements in pain interference (51.5 ± 16 preintervention vs. 43.5 ± 16.4 postintervention BPI score) and depression symptoms (34.0 ± 15.9 preintervention vs. 26.8 ± 12.1 postintervention CESD score). Poststudy interviews confirm the improvements of pain interference, mood, and bowel symptoms. Furthermore, analysis of medical conditions within the responder group indicates that the presence of depression, anxiety, irritable bowel syndrome, and fibromyalgia may indicate a greater likelihood for patients to benefit with vibroacoustic applications. These results indicate a possible potential for RSS, delivered using a vibroacoustic device, in managing pain-related symptoms. Further research is necessary to elucidate the exact mechanism behind the physiological benefits of RSS.

1. Introduction

Ehlers–Danlos syndrome (hEDS) is a congenital, heterogeneous group of connective tissue disorders characterized by joint hypermobility and skin extensibility [1]. Many forms of hEDS are caused by genetic mutations coding for various types of fibrillar collagen leading to weak muscles, ligaments, tendons, and organs [2]. EDS is now classified under 13 different subtypes [3], with hypermobile EDS (hEDS) being the most common [1, 4]. Patients with hEDS may develop chronic pain that occurs due to multiple contributions from joint wear and tear, musculoskeletal stiffness, neurogenic pain, and central nervous system dysfunction [5–7]. hEDS patients with chronic pain may also experience fatigue, anxiety, depression, and reduced quality of life [7–10]. There is currently no curative treatment for hEDS, and the most common form of treatment is using analgesics (e.g., opioids) [2, 5, 7, 11]. Given the complications of pain medications such as tolerance [12], substance abuse [13], and persistent pain, despite medication [2], integrating complementary strategies to manage pain could be part of a comprehensive multidisciplinary treatment approach.

Several theories have been proposed to explain the mechanism(s) of chronic pain. Melzack and Wall's gate
control theory (GCT) involves a bottom-up approach which posits that stimulation of peripheral nerves by nonnoxious stimuli can impede the transmission of painful stimuli [14]. Melzack later developed the neuromatrix (NM) theory of pain, a top-down model in which a collection of central nervous system components determines whether afferent noxious stimuli can be translated into pain perception [15]. Under the NM theory, sensory, cognitive, and affective aspects influence pain perception.

Rhythmic sensory stimulation (RSS) is defined as the stimulation of the senses in a periodic manner within a range of low frequencies. Music plus low-frequency sound delivered through a vibroacoustic device is a form of RSS and has demonstrated utility in managing pain [16, 17]. This intervention is based on two forms of rhythmic sensation: (1) sound captured by the auditory system and (2) vibrations captured by the somatosensory system. Although the mechanisms underpinning RSS for pain management are not well understood, theoretical grounds for utilizing RSS for treating pain in hEDS include positive cognitive/affective effects from music on brain function [18–21], neural entrainment of rhythmic stimuli [22–24], and sensory inhibition of pain via vibration [14, 25].

Although research using RSS has been conducted within an array of conditions, little research has focused on chronic pain in hEDS. In a previous single-case study of a patient with hEDS [26], we found an improvement in pain interference scores after using RSS via a vibroacoustic device for four weeks. In addition, the study found an improvement in bowel function, musculoskeletal tension, and energy levels. In the current study, we conducted an open-label pilot study of 15 patients with hEDS using RSS as the intervention. Based on a previous single-subject case report of an hEDS patient using the RSS device [26], it was anticipated that this device could improve pain interference scores as well as mood symptoms in a larger sample of hEDS patients. A washout period of two weeks followed the postintervention stage.

2. Materials and Methods

2.1. Participants. A total of 15 participants entered the study, and 14 completed the study. The inclusion criteria comprised the following: (1) a clinical diagnosis of hEDS by the clinical staff at Mount Sinai Hospital’s Wasser Pain Management Centre and Medical Genetics Department or a completed and signed referral form by health care practitioners indicating diagnosis; (2) joint hypermobility; (3) confirmed joint dislocations; (4) a Beighton score of 5 or more out of 9; (5) ongoing chronic pain of at least 6 months; (6) the ability to read and write English adequately; (7) self-reported satisfactory bilateral hearing; (8) the ability to operate the supplied RSS device, the Sound Oasis VTS1000, and to pick up and return the device at the Wasser Pain Management Centre.

The exclusion criteria were as follows: (1) serious comorbid illnesses, such as malignancy and vertebral fracture; (2) active inflammatory conditions, such as spinal infection and ankylosing spondylitis; (3) history of pathological fracture; (4) pregnancy; (5) coccydynia; (6) metabolic bone disease; (7) bleeding or clotting disorder; (8) hypotension; (9) active psychiatric disorders, such as bipolar disorder, schizophrenia, severe/psychotic depression, or anxiety disorder (at the discretion of clinicians/health care practitioners instead); (10) the use of pacemakers, implanted defibrillators, or neurostimulators.

2.2. Design and Procedure. A one-group pre-post treatment design was implemented. There were three assessments. The first assessment served as a baseline before study initiation and utilized four self-reported measures, including the Brief Pain Inventory, Short Form (BPI-SF) [27]; the Profile of Mood States, Short Form (POMS-SF) [28]; the Short Form 36 Health Survey (SF-36) [29]; and the Center for Epidemiologic Studies Depression Scale (CES-D) [30]. Following 4 weeks of treatment, the second assessment was performed with the same four questionnaires plus a Patient Global Impression of Change (PGI-C) and a 30-minute semistructured interview. After the two-week washout period, the final assessment included the same five questionnaires and a 30-minute semistructured interview. The primary endpoints were changes in scores from baseline to the end of active treatment in the BPI-SF, POMS-SF, SF-36, CES-D, and PGI-C. Permission was obtained to use each questionnaire. A short summary of each of these measures are as follows:

(1) BPI-SF: the brief pain inventory assesses the severity of pain and the impact of pain on daily functions. Although first made for cancer patients, it is reliable and valid for assessing noncancer-related pain [31]. Scores for questions are rated out of 10, with greater scores indicating worse conditions. Relief is measured as a percentage, with a greater percentage indicating better relief from pain.

(2) POMS-SF: the profile of mood states is used to measure psychological distress. A five-point scale ranging from “not at all” to “extremely” is administered, and a number from 0 to 4 is associated with this scale in a direction depending on the type of questions. Greater scores are associated with worse conditions. Reliability and validity for the POMS have been conducted [32].

(3) SF-36: the short form healthy survey measures the quality of health and is commonly used in health economics to calculate cost-effectiveness of a health treatment. The SF-26 consists of eight scaled scores that are weighted sums of a section of questions. Each scale is converted into a 0–100 point value in which a lower score indicates more disability and a higher score indicates less disability. Reliability and validity for the SF-36 have been conducted [33].

(4) CES-D: this is a brief self-report to measure depressive symptoms. The questionnaire consists of 20 questions, scores from 0 to 3, totaling 0–60 points. Higher scores indicate greater depressive symptoms.
Reliability and validity for the CES-D have been conducted [34].

(5) PGI-C: PGI-C consists of one item that measures the change in patient health. The range is 0–7 with a score of 1–3 indicating improvement, 4 being no improvement, and 5–7 being worse after the treatment. Reliability and validity for the PGI-C have been conducted [35].

Secondary endpoints included changes in scores from time of treatment termination to the final assessment following the washout period. Qualitative analysis was conducted on the responses from the semistructured interviews to explore patients' experience and perceived effectiveness of the RSS system for managing pain and mood. Open-ended questions from semistructured interviews included the following questions:

(1) Can you please tell me about your experience with using the RSS system?
(2) Have you started any new treatments to help manage your pain since you began this study 4 weeks ago? (probe: type of treatment, duration of treatment, and effectiveness of treatment)
(3) Have you changed your medication regime since you began this study 4 weeks ago? (probe: reduced or increased dose, added medication, removed medication, and switched medications)
(4) If you have changed your medication regime, why did you do so? (probe: change in prescription, side effects, difference in pain, mood, or sleep levels)
(5) Did you experience any negative side effects using the RSS stimulation system? (probe: if yes, what were the most bothersome side effects)
(6) Did you notice any positive effects/changes that you feel are a result of using the RSS system? (probe: improvement in pain levels or sleep, changes in behaviour, attitude, coping methods, emotions, and relationships)
(7) What did you dislike about using the RSS stimulation system, if anything? (probe: inconvenience, frequency or duration of use, vibrations, music, and comfort of chair)
(8) Would you choose to use the RSS system in the future? (probe: if yes, how often)
(9) On a scale of 0–10 with 0 representing not helpful at all and 10 representing extremely helpful, how helpful do you feel the RSS system was in helping you manage the severity of your pain?
(10) On a scale of 0–10 with 0 representing not helpful at all and 10 representing extremely helpful, how helpful do you feel the RSS system was in improving your mood?
(11) On a scale of 0–10 with 0 representing not helpful at all and 10 representing extremely helpful, how helpful do you feel the RSS system was in helping you manage daily activity levels?

To analyze and present this qualitative data, all answers were compiled and categorized into common phrases represented by 1–E2 words, counted, and then presented in bar graphs as a percentage of the total number of participant responses for each question of interest.

During this 4-week study, patients were given an RSS device (Sound Oasis VTS1000) which they self-administered daily for 30 minutes, five days per week. The patients were instructed to select a time of day to use the RSS device; however, it was recommended that participants use the RSS device in the morning and use it consistently at the same time. The patients were given a diary in order to record their usage and the presence or absence of side effects.

Participants were instructed to place the device on a bed or chair, turn the device on, and choose the “Energize therapy session” program that is supplied on this commercially available device. “Energize” comprises of three tracks (41 Hz to 73 Hz with 41 Hz dominant; 36 Hz to 61 Hz with 41 Hz dominant; and 36 Hz to 65 Hz with 55 Hz dominant). The music uses the bass, guitar, piano, and digital keyboard and features mono and binaural high alpha and beta entrainment. A somatosensory vibration generated by an audible low-frequency sound wave can be felt from the shoulders to the low-back, and relaxing but not “ambient” instrumental music can be heard prompting the participant to begin timing their 30-minute session. Patients were instructed to turn the low-frequency sound driven vibration intensity to 15 and the volume level of the music to 1 or 2. However, participants were informed that if discomfort was experienced due to these settings, they were to adjust the vibration and volume level to alleviate the discomfort. Participants were asked to note any deviations to the recommended procedure in the diary.

All patients consented to the study after being provided a clear description of the study and their rights of participation, withdrawal, and confidentiality. The study was conducted in accordance with the principles set out in the Declaration of Helsinki [36] and approved by a Toronto Academic Hospital Research Ethics Board.

2.3. Data Analysis. Descriptive statistics including percentages, means, and standard deviations were used to report demographic data such as age, gender, pain location, and pain duration. Data were analyzed by comparing baseline, posttreatment, and washout measurements using repeated measures ANOVA analysis. Interview answers were compiled, and common keywords or phrases were derived from each answer, counted, and then presented as a percentage of total participants. Outcomes were analyzed using SPSS for Windows, version 22, with 5% as the significance threshold.

3. Results

3.1. Patient Demographic Characteristics. A total of 15 patients with hEDS were recruited for participation, of which 14 completed the intervention and poststudy interviews. The average age was 35.8 years (SD = 13.2 years), ranging from 23
to 59 years. All patients were female and reported the presence of a concomitant condition such as depression, anxiety, or chronic pain conditions. More than half of the patients were either married or in a relationship. The demographic and clinical characteristics have been described in Table 1.

### Table 1: Patient demographics.

<table>
<thead>
<tr>
<th>hEDS patients (n = 15)</th>
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<tbody>
<tr>
<td>Age</td>
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<td>Medical conditions</td>
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<td>Headache</td>
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<td>Spontaneous dislocations/subluxations</td>
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<td>Chronic back pain</td>
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<tr>
<td>Depression</td>
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<td>Irritable bowel syndrome</td>
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<td>Flat feet, fallen arches, neuromas, and plantar fasciitis</td>
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<td>Anxiety</td>
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<td>Chronic pelvic pain</td>
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<td>Insomnia</td>
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<td>Chronic facial pain</td>
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<td>Fibromyalgia</td>
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<td>Dysautonomia</td>
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<td>Mast cell activation syndrome</td>
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<td>Thyroid condition</td>
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<td>Cracked, weak, or crowded teeth</td>
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<td>Raynaud’s syndrome</td>
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<td>Chiari malformation</td>
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<td>Other*</td>
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*Other includes the number of participants that reported at least one medical condition not on the original list of medical conditions. There are no reported cases of the following medical conditions: complex regional pain syndrome, substance abuse or dependence, diabetes, radiographically confirmed tethered cord, spontaneous cerebrospinal fluid leaks, or previous neurosurgery.

3.2. Initial Findings. Posttreatment improvements in both pain interference and depression were seen in 11 of the 15 patients (73%) (Figures 1(a) and 1(c)). During the washout period, 4 of these patients (36%) reported an increase in pain interference, whereas 6 of these patients (55%) reported continued improvements. Depression symptoms generally became worse during the postwashout period with only 2 patients (18%) reporting improvements in mood. Average scores for all patients showed improvements in pain severity, pain interference, and depression from pre-post treatment but did not reach significance ($p < 0.05$, Table 2). A closer look at the data of each participant reveals that 3 of the 15 hEDS patients (20%) experienced worse outcomes in pain interference and depression symptoms after treatment, which improved after stopping the treatment (Figures 1(c) and 1(d)).

Qualitative analysis of poststudy interviews indicated that 57% of patients reported the device to have a positive effect on their quality of life, with 21% reporting a neutral effect and 21% reporting a negative effect (Figure 2(a)). Open-ended questions suggested effects of the device on pain, muscle relaxation, sleep, and mood. Although patients were notified that they could adjust the intensity of vibrations to a comfortable level, some patients reported that the vibrations were too strong (Figures 2(c) and 2(d)). It is possible that they did not follow the RSS device adjustment instructions.

3.3. Clinically Meaningful Change in Pain and Depression. The analysis of individual data points along with poststudy interviews indicated that the experience of the RSS device varied among hEDS patients, with a small subset of patients reporting their symptoms worsening. Therefore, we were interested in determining the minimal clinically important difference (MCID) to report how many patients experienced a meaningful change after treatment. Following the procedure outlined in a review on MCID, the PGI-C scale was used to determine the MCID (a score of “3” or better) [37]. Of the 14 patients that completed the study, 6 participants (43%) were classified as “responders” to the device while 8 participants (57%) were classified as “nonresponders.” Mean age of the responders was found to be significantly different from the mean age of nonresponders (46.67 ± 13.2 years old for responders vs. 29.63 ± 9.1 for nonresponders, $p < 0.05$). Data analysis on responders indicated a significant improvement in pain interference and depression symptoms but not pain severity from pretreatment to posttreatment, with $p < 0.05$ (Table 3). Analysis of individual data points
indicated that half of the responders continued improving in pain interference during the washout period (Figure 3(b)), whereas only 33% of these responders continued improving in depression symptoms during washout (Figure 3(c)).

3.4. Examining Medical Conditions in Responders and Nonresponders. To study the difference between responders and nonresponders in more detail, an analysis of the prevalence of concurrent medical conditions was conducted and this yielded a potentially important observation (Figure 4).

Medical conditions with at least a 10 percent greater prevalence among responders than nonresponders were considered as possible candidates for screening for nonresponders. Responders had a high prevalence of depression, anxiety, insomnia, irritable bowel syndrome, and fibromyalgia.

4. Discussion

The present study investigated the effects of rhythmic sensory stimulation on pain and mood symptoms of hEDS.
patients. Our findings revealed that 43% of hEDS reported benefit from using RSS in their symptoms. The proportion of hEDS responders to this RSS treatment is similar to the proportion of fibromyalgia responders (40%) in a previous study that used a similar protocol [38]. The mean age of responders was significantly higher than the mean age for nonresponders. It is possible that older patients have a greater disease severity and more complications and are thus more likely to be responders. To study the difference between responders and nonresponders in more detail, an analysis of the prevalence of concurrent medical conditions was conducted and yielded a potentially important observation (Figure 4). Responders had a high prevalence of depression, anxiety, insomnia, irritable bowel syndrome, and fibromyalgia. These findings are consistent with observations from our previous case study. Therefore, these concurrent medical conditions could potentially serve as a selection tool to determine likely candidates among hEDS patients that may benefit from using this device.

Consistent with the initial case report, we found that pain interference is significantly improved (i.e., reduced) in responders without significant changes in pain severity.
Furthermore, poststudy interviews reveal that one of the common positive benefits of the device was muscle relaxation. This suggests that the vibrotactile sensation produced by the RSS device could have improved mobility-related pain and functionality, which has been demonstrated in previous studies using low-frequency stimulation [39–41]. Mechanical stimulation of the Pacinian corpuscle, a subcutaneous mechanoreceptor sensitive to vibrations of 60 Hz to 600 Hz, is thought to produce this kind of effect [16]. Furthermore, low-frequency vibrations have been shown to produce other physiological responses that include changes in breathing rate, metabolic rate, blood pressure, fatigue, headache, nausea, and tension [42–45].

We also found significant improvements in depression symptoms in responders after using the intervention. Improvements seen in mood symptoms may be due to improvements seen in pain interference, as this increases the types of activities that patients can engage in and thus improves quality of life. However, the cognitive and affective effects of music also play a major role, increasing dopamine and serotonin levels in the brain and improving mood symptoms more directly [18, 19]. A previous study using the same intervention on patients with major depressive disorder found an improvement in depressive symptoms such as sleep quality and anhedonia after music-based RSS for five weeks [46]. The effects of music therapy on improving depression and anxiety are well documented [47].

An additional consideration is the evidence for aberrant neural oscillations that have been associated with the sensation of chronic pain [48, 49], depression [50], and poor sleep quality [51, 52]. In examining the role of rhythmic oscillatory coherence, clinical and animal studies suggest that neurogenic pain is linked to consistent abnormal activity between the thalamus and cortical areas that manifest as abnormal thalamocortical rhythms [53–58]. This phenomenon, termed thalamocortical dysrhythmia, forms the basis for a model of neurogenic pain, which is prevalent in 68% of hEDS patients [59]. Any stimuli that can normalize these aberrant neural oscillations are purported to potentially reverse or reduce the associated symptoms of chronic pain. RSS has been shown to have such an effect of synchronizing neural oscillations, and this phenomenon has been termed as entrainment in the

![Figure 3: Analysis of effects of the intervention on pain interference and mood for responders. (a) Line graph depicting BPI-severity scores in patient responders at all time points in response to the intervention. (b) BPI-interference scores in patient responders in response to the intervention. (c) CES-D scores in patient responders in response to the intervention. *p < 0.05.](image)
neuroscience literature [23, 24]. The entrainment of brain rhythms to reduce chronic pain may be one of the mechanisms responsible for the benefits observed following use of RSS and must be investigated further.

5. Conclusion

Auditory and vibrotactile RSS can be an additional tool to reduce hEDS symptoms including pain, mood, sleep, and potentially bowel movements. hEDS patients that report these particular symptoms may be more likely to benefit from 40 Hz of auditory and vibrotactile, while a small percentage of patients may find it discomforting. Further research is necessary to elucidate the mechanisms underpinning the physical and psychological effects of RSS.

Data Availability

The data used to support the findings of this study are included within the article.

Conflicts of Interest

V. V., A. M., D. P., L. P., H. F., M. E., and A. G. declare that there are no conflicts of interest. L. B. has served as a paid scientific consultant to Somerset Group and Sound Oasis and receives limited royalties from the Somerset Group for the Sonic Aid series and from Sound Oasis on the sales of the Vibroacoustic Therapy System VTS-1000 device.

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