The Effect of Therapeutic Low-Frequency Ultrasound applied to Myofascial Trigger Points: A Pilot Pre-Post Design Study

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Abstract

Ultrasound at a frequency of 1 or 3 MHz is frequently used to treat various musculoskeletal conditions, but research on ultrasound operating at 38-50 kHz frequencies (US-KHz) is lacking. Study aimed to evaluate the short-term effect of US-KHz on pain pressure threshold (PPT), ankle dorsiflexion range of motion (ROM), and motor performance (the Side Hop Test) in subjects with a myofascial trigger point (TrP) in the calf muscle. US-KHz was applied to the area of the palpable TrP in the calf muscle for 10 minutes (continuous pulse, transducer head size 19.6 cm², power 0.75-1.25 w/cm²) in twenty volunteers (18-45 years old). Significant improvements (p<0.001) were noted in the Side Hop Test 24 hours after the intervention. ROM improved significantly after 5 minutes with changes maintained 24 hours later. No change was found in the PPT. This pilot study presents preliminary evidence of the efficacy of US-KHz in treating TrPs.

Keywords: Therapeutic Ultrasound; Kilo-Herz Frequency; Trigger Points; Feasibility Study; Calf Muscles.

1. Introduction

Myofascial pain is a common pain syndrome of the skeletal muscles, frequently accompanied by myofascial trigger points (TrPs). These TrPs are sensitive spots found within the muscle and are generally identified by local muscle palpation [1, 2]. Although there is much debate amongst researchers and clinicians as to the diagnostic criteria of TrPs, a recent international Delphi panel concurred that a diagnosis is rendered when two of the following three criteria are met: a hypersensitive spot within a muscle which may include in addition to a sharp pain, sensations such as a dull ache, or tingling; a taut muscle band; and a referred sensation extending to distant sites [2, 3]. TrPs are usually classified as either active or latent. The local and referred symptoms of active TrPs are spontaneous and increase when manipulated. In contrast, while latent TrPs are not associated with spontaneous discomfort, local and referred symptoms may be provoked by mechanical stimulation, such as finger pressure exerted over the latent TrPs [4]. Similar to active TrPs, latent TrPs may restrict the range of motion (ROM) and may lead to muscle dysfunction, i.e., muscle weakness, muscle imbalance, or altered motor unit recruitment patterns [5, 6].

The etiology of myofascial pain and TrPs is not as yet fully understood. Treatment is usually symptomatic with the intent to reduce pain and improve ROM and muscle strength [4]. Diverse invasive and non-invasive modalities have been reported with conflicting results. These modalities include local injections, dry needling, deep tissue massage, stretching spray techniques, laser therapy, electrotherapy, and superficial and deep heat [7-9]. Therapeutic ultrasound (US) is a commonly used modality applied to treat myofascial pain [10]. The US transforms electrical energy into

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mechanical oscillation via piezoelectric crystals [11]. The oscillation of particles within the biological tissues, due to the longitudinal wave created by the US, results in an increase in deep tissue temperature and mechanical modifications at the cellular level [12], thereby, possibly modulating pain, increasing tissue elasticity and ROM, improving blood flow, reducing muscle spasms, and enhancing the tissue healing process.

The efficacy of using US to treat TrPs has been investigated in several studies culminating with conflicting results. A recent systematic review and meta-analysis of randomized controlled studies identified ten studies that assessed the therapeutic effects of US on myofascial pain [13]. The review indicated that US treatment significantly reduces pain intensity with no concurrent significant effect on ROM. The authors concluded that due to the high risk of bias and the significant heterogeneity of the studies in terms of treatment protocols and dosage, the results of the meta-analysis did not clearly support the use of US as an effective method to treat myofascial pain. The amount of US energy delivered and absorbed by the biological tissues depends on a variety of dosage parameters, i.e., pulse frequency, intensity, and duration of application [14-16]. Typical frequencies of conventional US range between 1-3 MHz, with the higher frequencies associated with a lower penetration power [17]. It has been suggested that much lower frequencies (associated with longer wavelengths) would provide a more divergent field shape. Furthermore, reducing the frequency would result in less attenuation of the sonic energy as it travels inward. There would also be a more uniform distribution of the absorbed energy in the direction of wave propagation, thus, expecting greater thermal and athermal effects in the deeper tissues [18]. Due to these properties, there is a growing interest in applying US at kHz frequencies.

Despite the theoretical advantages of low-frequency US, only a few studies have investigated its efficacy in clinical settings. Several studies have focused on soft tissue healing. A systematic review and meta-analysis of eight randomized controlled trials revealed that low-frequency US at both low and high intensities may accelerate wound healing in patients with venous stasis and diabetic foot ulcers [19]. Fewer research studies relating to musculoskeletal impairments have been conducted, resulting in conflicting outcomes. Thus, in a prospective randomized trial comparing the efficacy of US at 3 MHz with that of 45KHz in the treatment of acute ankle sprains, it was determined that the group treated with a lower US frequency demonstrated greater immediate improvements in gait parameters, remaining statistically significant three days following treatment [18]. In contrast, Meakins and Watson (2006) [19] found that low-frequency US is as effective as hot packs in increasing functional ankle mobility. In light of these conflicting results, the goal of this pilot study was to examine the immediate and short-term effects of kHz-frequency US on pain pressure threshold (PPT), ankle dorsiflexion ROM, and motor performance (Single Leg Hop Test) in subjects with latent TrPs located in the calf muscle.

2. Materials and Methods

2.1. Design

A pre-post design pilot interventional study.

2.2. Setting

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2.3. Sample

The study sample included 20 healthy students attending the Ben-Gurion University of the Negev, between the ages of 18-45. The trial was advertised through an ad posted on the university's bulletin board and social networks. Inclusion criteria consisted of the presence of at least one TrP located in the calf muscle of one leg, which was identified by manual palpation by a tester. If more than one TrP was found, the one with a lower PPT was used. Exclusion criteria included a background of neurological diseases, active systemic diseases affecting sensation or perception of pain (i.e., diabetes, peripheral vascular disease, cancer, rheumatoid arthritis, osteoarthritis), a fracture or ankle sprain during the six months prior to the study, pregnancy, local infection or inflammation, and contraindications to US (malignant tumors, active bleeding or open wound in the calf area, and systemic vascular disease).

2.4. Intervention

Each participant underwent a single US application of a 38-kHz frequency (SIRIO, BAC Technology, Italy), head size 19.6 cm², in a continuous current according to the manufacturer's instructions. A standard water-based US gel was used. The maximum power of the device was 2.5 W/cm². The intervention was performed by a physical therapy student trained in operating the device. The participant was informed prior to treatment that he would not feel any abnormal sensations such as strong heat or pain but might feel a buzzing in his ears and a low-medium heat sensation in the placement area. The treatment procedure was performed on the area of the marked point (a dot was drawn on the skin with a marker by the examiner at the initial examination, thereby, accurately mapping the TrP). The procedure lasted 10
minutes, commencing at a 30% maximum intensity, subsequently increased to 40% after 2 minutes of treatment. If the participant was still feeling comfortable, intensity was increased to 50%. After an additional 2 minutes, if the patient reaffirmed that he/she still felt comfortable, the therapist raised the intensity again. If after increasing the intensity, the patient reported discomfort or pain, the therapist lowered the intensity back to the previous intensity. At any stage throughout the treatment, the patient could ask to lower the intensity if he/she felt uncomfortable or in pain. We chose this regimen because in our preliminary stage when we tested different parameters of US (we aimed to apply the US at 50% of the maximum intensity), some subjects reported that 50% felt too hot.

2.5. Outcome Measures

All outcome measures were documented by a single researcher (a 4th-year physical therapy student), who had undergone intensive specific training in evaluation techniques. Demographics were collected using a questionnaire and included sex, weight, height, smoking, and background diseases. Body mass index (BMI) was calculated as weight (kg)/height (m)²).

2.5.1. Weight-Bearing Lunge Test (WBLT)

This test was used to assess ankle dorsiflexion ROM. The WBLT has been used to detect ROM deficits in individuals with chronic ankle instability [20] and to track the progress of ROM improvement during the rehabilitation [21]. Hall and Docherty (2017) [22] found that WBLT highly correlates with measurements performed using a 2D motion capture system and determined that WBLT is a valid and reliable tool to assess dorsiflexion ROM. Furthermore, Bennell et al. found that the WBLT demonstrated excellent inter and intra-rater reliability [23]. Another study found that the WBLT exhibited high test-retest reliability (ICC=0.93-0.99) [24].

We assessed the dorsiflexion ROM angle between the anterior border of the tibia and the vertical line using a digital inclinometer. The examiner used a non-permanent felt-tipped pen to mark a dot on the anterior border of the tibia, 15cm below the tibial tuberosity of both legs. The test procedure was demonstrated to the subjects together with standardized instructions before the commencement of the procedure. The subject then positioned the tested foot so that the heel line and the big toe were aligned on white masking tape. With both feet stationary, a controlled forward lunge was performed, thereby, flexing the knee as the subject attempted to reach maximal dorsiflexion. Pronation or supination of the foot was not controlled. During the tests, subjects were allowed to hold onto the wall for balance. At the maximum lunge point, the examiner placed the inclinometer on the tibial mark and recorded the achieved angle. Two additional measurements were taken after the subject stood and resumed a comfortable position. The best result was recorded by the examiner.

2.5.2. Side Hop Test

This test assessed functional performance ability and dynamic impairments. The gold standard for muscle strength evaluation is the isokinetic muscle torque test [25], however, its validity has been questioned due to the low correlation between the isokinetic muscle torque and functional performance [26]. It has been suggested to use dynamic muscle power tests to evaluate muscle function [27] given that the ability to produce high forces during high velocities is one of the most important factors in sports performances [28]. In addition to power tests, hop tests have been highly recommended to evaluate sport-specific performances in healthy athletes and after various interventions [29, 30]. A battery of hop tests evaluating hop performances has been previously described and analyzed for reliability [31].

There are three tests with a high capacity to discriminate hop performance: the vertical jump, the hop for distance, and the side hop. For the present study, the Side Hop Test was employed. Reliability was conducted for this test and determined to be good with an ICC of 0.84 [27, 32]. Docherty et al. method was used in this study [33]. All subjects were instructed to hop laterally on the right limb a 30cm distance and then return to the starting location. Subjects were told that the goal was to complete ten repetitions as quickly as possible. If a subject fell, or put the contralateral foot down, or did not completely clear the 30cm distance while hopping, the trial was regarded as unacceptable and the subject had to repeat the task. Subjects were familiarized with the task by completing 3-4 repetitions at partial speed. Following a rest period, they completed their first trial, rested for at least 60 seconds, then completed the second. The examiner recorded the time with a handheld stopwatch to the nearest hundredth of a second. The best trial (shortest) was used for analysis.

2.5.3. Pain Pressure Threshold

PPT of the identified TrP assessed the latent TrP pressure sensitivity using a mechanical algometer. The disc was placed perpendicularly on the TrP. The examiner then increased the intensity of pressure until the participant reported initial feelings of pain. The score was determined by averaging three consecutive measurements. The algometer was found to be valid and reliable in repeated measurements (Interrater, Intrarater) in normal muscles in healthy people with a minimal clinically important difference of 14.71-19.61 N/cm² [34]. Chesterton et al. [35] found high reliability when three repeated measures were averaged: ICC = 0.91 (CI 0.82; 0.97 95%) with a minimal clinically important difference
of 17.39 N/cm². All outcome measure evaluations were performed prior to the intervention, five minutes after the intervention, and ~24 hours after the intervention.

2.6. Research Procedure

The subjects chosen in accordance with the exclusion and inclusion criteria were informed of the experimental procedure. After signing an informed consent form, the examiner palpated the participant’s calf muscles while the subject lay on his/her stomach with a slightly bent knee. If a TrP was detected the examiner marked the TrP as described above. After completing the demographic questionnaire, the three tests were performed under the examiner’s guidance in the following order: PPT, ankle dorsiflexion ROM, and Side Hop Test. The subject was asked to lay prone, and the intervention was performed according to the above-detailed protocol. Five minutes after the intervention ended, PPT and ankle dorsiflexion ROM evaluations were repeated. Approximately 24 hours after the intervention, the patient was asked to return and perform an additional set of all three tests.

2.7. Statistical Analyses

All statistical analyses were performed using the SPSS statistical package (version 23 for Windows). Significance levels were set at P<0.05. Descriptive statistics characterized the sample. The comparisons between the background, 5-minute post-intervention, and 24-hour post-intervention measurements were calculated by the repeated measurements ANOVA using a posthoc analysis. A Pearson correlation analysis assessed the association between changes in the outcome measures (delta scores=24-hour post-intervention measurement minus the baseline measurement).

3. Results

The study included 20 subjects (19 females, one male) ranging in age from 24-32, the mean age of 26.65±2.78. The mean BMI was 21.90±2.84 kg/m²; most (85%) reported routinely exercising. Additional demographic characteristics are presented in Table 1.

Table 1. Descriptive statistics (N=20)

| Variables               | Mean±SD   |
|-------------------------|-----------|
| Age (years)             | 26.65±2.78|
| Weight (kg)             | 60.30±7.97|
| Height (m)              | 1.66±0.06 |
| BMI (kg/m²)             | 21.90±2.84|
| Sex (females)           | 19 (95%)  |
| Smoking                 | 1 (5%)    |
| Regular physical activity| 17 (85%) |

SD: standard deviation; BMI: body mass index

The main results are summarized in Table 2.

Table 2. Comparison (results of repeated measurements ANOVA) of measurements before and after the intervention

| Variable          | Before Mean±SD | 5 min. after Mean±SD | 24 hours after Mean±SD | Comparison     |
|-------------------|----------------|-----------------------|------------------------|----------------|
| PPT (kg/cm²)      | 0.22±0.552     | 0.44±0.712            | 0.32±0.782             | F=0.817, p=0.439|
| WBLT (○)          | 137.30±7.41    | 139.50±7.69           | 140.10±7.55            | F=8.767, p<0.001*|
| Side Hop Test (sec)| 28.54±22.34    | 21.47±16.17           |                        | F=15.027, p<0.001|

SD: standard deviation; PPT: pressure pain threshold; WBLT: weight-bearing lunge test;
Statistically significant (p<0.05) differences are marked in bold.

* In a pairwise comparison, significant differences were found between the 1st and 2nd measurements, and the 1st and 3rd measurements, but not between the 2nd and 3rd measurements.

The PPT showed some increase after the US application between the 1st (performed before placement) and the 2nd test (five minutes after placement), and between the 1st and 3rd test (24 hours after placement), although, this change was not statistically insignificant (P = 0.439).
In the ankle dorsiflexion ROM tested by the WBLT, an average improvement of ~2° between the 1st and 2nd test and an improvement of ~1° between the 2nd and 3rd test was observed. The differences between the 1st and 2nd test and between the 1st and 3rd test were statistically significant (p<0.001), however, no significant difference was found between the 2nd and 3rd test. Regarding motor performance tested by the Side Hop Test, a statistically significant difference was found between the 1st and 2nd tests (p<0.001). The mean difference between the 1st and 2nd test was 7.07 seconds (that is higher than a minimally detectable change of 2.8-5.7 seconds [36]). The Pearson correlation analysis assessed the relationship between changes in the measurement results; the only significant correlation (r = -0.443, p = 0.05) was found between changes in the PPT (five minutes after treatment) and changes in the Side Hop Test. Subjects who further improved in the PPT during the 2nd test also improved in motor performance in the 2nd test (24 hours after placement).

4. Discussion

We determined that a single application of US-KHz directed at latent TrPs located in the calf muscles of healthy subjects results in an immediate increase in the ROM lasting 24 hours after the intervention. We also discovered that the US-KHz led to improvements in the Side Hop Test 24 hours following the intervention, however, no effect on PPT was found. To the best of our knowledge, only three previously published studies have examined the effect of US-KHz applied to musculoskeletal structures in vivo. The earliest study, a randomized placebo-controlled work published in 1996 compared the immediate effects of a single treatment with low-frequency US (45-KHz) to treatment with US at a high frequency (3- MHz), following an acute ankle sprain. Significant greater improvements in gait parameters following the lower frequency US were observed. The authors attributed this to a greater analgesic effect [18]. In contrast, Basso and Pike’s [37] double-blinded study of subjects with a post-Colles fracture, found no difference in wrist ROM between patients treated with 46.39 US-KHz and sham treatments. However, it appears that the subjects were exposed to only one treatment and were examined two and eight weeks later; Thus, many confounding variables could have affected the results. The treatment and assessment protocol of the present study has more in common with the more recently published study by Meakins and Watson’s treatment protocol was based on Ward and Robertson [38] who applied US-KHz to non-living pig tissue demonstrating a 4-50C increase in superficial tissue temperature following 5-6 minutes of treatment. When examining the ROM of the human ankle, the authors increased the duration time by 4 minutes to compensate for the circulation’s cooling effect and the moving treatment head. As increased tissue temperature has been shown to enhance connective tissue extensibility, as well as altering nerve conduction velocity, and increasing blood flow [11, 39], the authors considered the increase in ankle ROM as the result of enhanced tissue extensibility due to increased tissue temperature.

Dosage parameters in the present study were based on the manufacturer’s recommendations (SIRIO, BAC Technology, Italy). While these parameters were not identical to those presented by Meakins and Watson, they were quite similar, i.e., treatment duration was 14 minutes in the present study, vs. 10 minutes in their study. Intensity ranged between 0.75-1.25 W/cm² in the present study vs. 0.94 W/cm² in their study. Hence, it appears that the same mechanism, i.e., increased tissue temperature, was responsible for the positive treatment effect on ankle ROM observed here as well. As latent TrP causes muscle dysfunction [6, 39], the thermal effect of the US-KHz might also explain the demonstrated improvement in the Side Hop Test [6]. Previous studies have shown positive effects of US-MHz frequencies on latent and active TrP in terms of ROM, PPT, pain relief, disability, and quality of life [11,40-44], however, no studies have as yet compared the effects of low frequency (KHz) and high frequency (MHz) US on TrPs. Based on the promising preliminary results of the current pilot feasibility study, it is essential to expand these results in follow-up larger controlled studies which will include comparisons between the effectiveness of US-KHz with the ‘traditional’ US-MHz in the treatment of TrPs.

In previous studies investigating MHZ frequency US, a significant beneficial effect on PPT indices of participants with soft tissue injury was observed [11, 40-42]. The gap found in these indices between the MHZ frequency studies and this study may lie in the deep penetration potential of the KHz frequency, which may be 20 times greater than the MHZ frequency. We, therefore, feel that a different and more subtle treatment protocol should be adopted due to the great power of the device.

We observed that some people were more sensitive than others and reported discomfort or pain during the intervention. For these individuals, the intervention continued until the end of the allotted time, but the US intensity was lowered to a level that was comfortable for the participant. Our impression was that people with greater sensitivity to treatment showed less improvement in terms of pain pressure index in post-treatment tests. It may be worthwhile to perform a sensitivity test before applying the treatment as part of the research protocol in order to monitor people with
skin sensitivity. Further studies should concentrate on the applied muscle, as previous studies have suggested that certain muscles are more easily influenced by superficial heating, resulting in an immediate change in muscle length and flexibility, for example, shoulder external rotator muscles were more affected by superficial heating than the triceps surae [39].

This study has several major limitations. Firstly, there was no control group; all subjects underwent the same treatment in the same way. This study design does not allow for controlling potential biases, i.e., a placebo effect or spontaneous change in outcomes (time effect). Moreover, a pre-post-designed study does not allow the blinding of tester or subjects. However, as mentioned before, this is a pilot, feasibility study performed as a preliminary stage to ascertain if US-KHz causes any change in subjects’ outcomes, in addition to serving as a basis for a full-scale clinical trial.

5. Conclusion

The current results support the potential application of US-KHz therapy as an effective treatment modality for TrPs, by affecting elasticity and muscle flexibility (represented in this study by ROM) and improving muscle performance of the muscle. If this efficacy will be confirmed in a further full-scale clinical trial, US-KHz can be a therapeutic alternative to invasive treatments such as dry needling in TrP conditions and can be added to the physical therapist's toolbox. US-KHz is a relatively new modality and only a few studies have examined its effect, therefore, high-quality, especially randomized controlled trials, are essential to establish its efficacy and treatment protocols. Further studies should also examine the effectiveness of this treatment on active TrPs as well as on TrPs in other areas of the body.

6. Declarations

6.1. Author Contributions

Conceptualization, M.E.G and L.K.; methodology, M.E.G and L.K.; formal analysis, M.E.G and L.K.; data curation, A.D and L.K.; writing— M.E.G A. D and L.K.; writing—review and editing, M.E.G and L.K.; visualization, M.E.G and L.K.; supervision, L.K.; project administration, L.K. All authors have read and agreed to the published version of the manuscript.

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6.4. Ethical Approval

The study was approved by the Ethical Review Board of the Ben Gurion University of the Negev, Israel. All subjects received a detailed explanation of the study goals and provided written consent before participation.

6.5. Data Availability Statement

The data presented in this study are available on request from the corresponding author.

6.6. Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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