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Effects of Whole-body Electromyostimulation on knee pain and physical function in adults with symptomatic knee osteoarthritis: a randomized controlled trial

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Abstract

In a randomized, controlled study, whole-body electromyostimulation (WB-EMS) was investigated as a promising alternative to conventional strength training for the treatment of knee osteoarthritis (OA). 72 overweight participants with symptomatic knee OA were randomly assigned to WB-EMS (n=36) or a usual care control group (CG, n=36). For seven months, the WB-EMS group received three times per fortnight a WB-EMS training, while the CG was prescribed 6x physiotherapeutic treatments. The primary outcome, change in the pain subscale of the Knee injury and Osteoarthritis Outcome Score (KOOS), significantly improved in favour of the WB-EMS group, with a mean increase of 16.7 points versus 7.0 points in the CG (absolute difference between groups 9.0 points, 95%CI 2.9 to 15.1, p=0.004). Secondary outcomes, including the other KOOS subscales (symptoms, function in daily living, function in sports/recreational activities and quality of life), 7-day pain diary, isometric muscle strength and lower limb function (30s sit-to-stand test), were also in favour of WB-EMS. With few dropouts and no reported adverse events, WB-EMS had a participation rate of 88% ± 10%. Overall, WB-EMS was found to be effective in relieving knee pain symptoms and improving physical function in individuals with symptomatic knee OA compared to usual care treatment.
Introduction

Knee osteoarthritis (OA) is a leading cause of global disability [1]. The individual burden and socioeconomic impact of knee OA is profound and is expected to increase in the coming decades [2-4]. With no cure for OA currently, clinical guidelines emphasize treatments that relieve symptoms of the disease and improve function, such as exercise, weight loss (for those overweight) and education [5-7].

Various exercise programs, such as resistance and endurance training, have a positive effect on pain and function in knee OA [8]. In a recent systematic review, resistance training was effective in reducing pain and/or improving function in daily living in 11 out of 12 studies (with a moderate to large effect size) [9]. However, despite the high level of evidence regarding the benefits of physical activity and exercise for knee OA, the majority of individuals with knee OA do not meet recommendations for physical activity [10].

In individuals with knee OA, a vicious cycle of pain, avoidance of physical activity, reduced muscle strength and further functional limitations has been proposed [11]. As such, there can be barriers for participation in resistance training to improve strength [12]. In contrast to conventional resistance exercise, Whole-body Electromyostimulation (WB-EMS) is an approach characterized by intense activation of muscles via an adjustable impulse delivered via surface electrodes with low voluntary effort. This approach may be an attractive alternative for individuals with knee OA who may have an inability to sufficiently voluntarily contract muscles to facilitate muscle strength gains and associated symptomatic relief. In previous studies, WB-EMS has shown positive effects on muscle strength, muscular morphology and fat mass in healthy, sarcopenic and/or functionally impaired participants [13-19].

The majority of existing EMS studies in individuals with knee OA concentrated on the effects of local EMS. A systematic review by de Oliveira et al. [20] showed moderate evidence in favour of neuromuscular electrical stimulation (NMES) alone or in combination with exercise for isometric quadriceps strengthening. A recent meta-analysis by Carvalho et al. [21] reported insufficient evidence on the effects of NMES combined with exercise compared to exercise alone on patient-reported outcomes (e.g. pain). Due to the lack of comparability between studies (methodological differences, e.g. study design, training protocol, type of stimulation), the evidence for NMES in individuals with knee OA remains limited.

WB-EMS could have some advantages compared to local EMS. WB exercise increases overall physical performance and may also exhibit positive systemic anti-inflammatory effects by activating large muscle groups [22,23].

The aim of this study was to compare the effects of a 7-months WB-EMS application to a usual care control group (CG) in overweight individuals with symptomatic knee OA. Our primary hypothesis was that WB-EMS will result in significantly greater reductions in knee pain compared to the usual care CG. We further hypothesized that, compared to the CG, WB-EMS will result in significantly greater improvements in self-reported function in daily living, recreational activities and quality of life, quadriceps strength and physical function.
Method

Study design

The EMSOAT (Whole-Body Electromyostimulation for the Treatment of knee OA) study is a parallel-group (1:1 allocation) superiority randomized controlled trial (RCT) conducted at the Institute of Medical Physics (IMP), Friedrich-Alexander University of Erlangen-Nürnberg (FAU), and the Department of Radiology, University Hospital Erlangen Germany. The RCT was approved by the FAU ethics committee (Nr. 352_20 B) and all participants provided written informed consent prior to enrolment. The project fully complies with the Helsinki Declaration [24] and was prospectively registered at clinicaltrials.gov, NCT05672264, on 05/01/2023.

Participants

Participants were recruited between March and June 2022 in the metropolitan area of Erlangen-Nürnberg, Germany. As in previous studies, we recruited potential participants by reports and expert interviews on knee OA and corresponding study calls in local newspapers and social media. The call listed the key study eligibility criteria, contact person and an email address. Furthermore, we contacted eight medical practices (practitioners with qualification in sports medicine and orthopaedists) via letter and provided information flyers for their patients.

Inclusion criteria were (1) men or women 40-70 years of age, with (2) overweight (BMI>25 kg/m²), (3) confirmed femorotibial OA equivalent to Kellgren-Lawrence grades (KL) 2 and 3 [25] (see explanation below), (4) knee pain for at least 3 months, (5) pain in the last 30 days at least on 50% of the days and (6) an average pain intensity > 2.5 [26] on a scale 0-10 (NRS).

Exclusion criteria were: (1) Any WB-EMS training or more than 60min of resistance exercise training per week in the last year, (2) glucocorticoid or opioid medication, (3) trauma to the knee joint within the last 3 months, (4) intra-articular knee injection in the last 3 months, (5) conditions and diseases (and corresponding medication) with relevant impact on study outcomes (i.e. other rheumatic diseases e.g. rheumatoid arthritis, fibromyalgia, serious cardiovascular diseases), (6) conditions or diseases that are contraindications for WB-EMS (e.g. electric implants, epilepsy, cardiac pacemakers [27]) and (7) absence ≥4 weeks during the intervention period.

As radiographs could not be obtained for study purposes only [28], potential participants were asked to provide externally acquired anterior-posterior radiographs of their index (more painful) knee when available. These were assessed by an experienced musculoskeletal radiologist (FWR) and those with KL 2 or KL3 were included [25]. Participants without externally acquired radiographs or radiographs older than 2 years were screened by MRI and those with full-thickness cartilage damage at both the femur and tibia in at least one compartment (grades 3.2 or 3.3 in at least one central femoral and one subregion of the anterior, central and posterior tibial subregions on the MOAKS (MRI Osteoarthritis Knee Score) [29] scale) were excluded. Also, those with no or only focal cartilage damage (maximum of 1.0 or 1.1. in the 10 femorotibial subregions of the MOAKS instrument) were excluded. Using these MRI definitions, the likelihood of including KL 0 and 1 knees or knees with end stage structural OA (KL4) was minimized [30].

If both knees of a single participant were eligible, we defined the side that caused more pain as the “index limb” (affected knee).
**Intervention**

**WB-EMS application**
WB-EMS was applied using a system with medical device approval (miha bodytec®, Type II, Gersthofen, Germany) that enables simultaneous stimulation of up to 10 main muscle groups (thighs and upper arms, hip/bottom, abdomen, chest, lower back, upper back, latissimus dorsi and two free options) with an overall area of stimulation of about 2600 cm². The system allows intensities to be chosen for each region. We established a consistently supervised, video-guided WB-EMS program 3 times per fortnight (e.g. every Monday or Tuesday and every second Thursday or Friday) for 6 months (from August 2022 to January 2023) plus one month of conditioning (July 2022; see below). All participants started the intervention at the same time. We used an impulse protocol that was applied in research [14,15,17,18,31-33] and most commercial settings in order to allow transferability of our approach. Bipolar electric current with a frequency of 85Hz, an impulse-width of 350 µs and a rectangular impulse pattern was used for 20 minutes in an interval approach with 6 sec of EMS stimulation and 4 sec of rest. Participants completed two sets with 6-8 repetitions of seven exercises (e.g. light dynamic squatting with knee angles ≥ 120° and arm curls) in a standing position (Figure 1). Of importance, we designed low-intensity movements/exercises to keep the effect of the voluntary movements itself as low as possible.

The intensity of the EMS was regulated based on the rate of perceived exertion (RPE) scale. We applied a perceived exertion rate to generate and maintain a sufficient but tolerable intensity of the EMS.
application. Before the 6 months of WB-EMS training, we implemented 4 weeks of conditioning with lower impulse intensity and shorter sessions (July 2022). We started with 12 minutes in the first session and increased time by 2 minutes per session. After conditioning, participants were encouraged to exercise at an EMS-induced RPE of “6-7” (i.e. “hard+ to very hard”) on the Borg CR10 Scale [34]. Impulse intensity was individually adapted for each body region in close interaction with the participant. During the session, instructors slightly increased (impulse) intensity every 2-3 min in close cooperation with the participants to maintain the prescribed RPE (“6-7”) during the session. From mid-September 2022, all participants used a second pair of circular electrodes for the thighs, to adequately stimulate the thighs and maintain the intensity. All training sessions took place in the Institute of Medical Physics. We applied a personal training setting with one licensed and experienced instructor responsible for two participants. Instructors monitored compliance with the prescribed exercise intensity and recorded attendance rate accurately. In case of non-participation, participants reported absence by email or telephone. Possible adverse events were recorded on a weekly basis during the entire course of the study. Further, the international guideline of safe and effective WB-EMS application was strictly respected [35].

Control intervention (referral to physiotherapy)

The participants received a prescription for 6 physiotherapy treatment sessions (20 min each) with the recommendation have those within the first three months at a frequency of 1x/week. Physiotherapy treatment was carried out individually in the sense of "usual care" in a diagnosis-orientated manner. The specific content was at the decision of the treating physiotherapists containing techniques and exercises for reducing pain and detonisation of muscle tissue, increasing mobility of the knee joint and strengthening leg muscles. It was recommended that the therapy be carried out in one of three cooperating practices. However, participants were free to take the prescription to another practice of their choice. All practices were informed about the study and the aims of the study in a letter accompanying the prescription.

Education (both groups)

Both groups were invited to participate in a training program for self-management of OA [36]. Six units (60 min each) were offered over a period of 12 weeks. Before each of the 6 sessions, an invitation with a brief information was sent via email to the participants of both groups. The 6 sessions were led by different experts, each of them was blinded to the group allocation. The aim of the program was education, information and counselling to improve quality of life and mobility. Self-management, personal responsibility and coping strategies of the participants to cope with bio-psycho-social (stress) factors was promoted and supported. Overall, we intended to reduce fear and avoidance behaviour.

Outcomes

Primary outcome

- Changes in the pain subscale of the Knee injury and Osteoarthritis Outcome Score (KOOS-Pain) from baseline to 7-month follow-up (FU)

Secondary outcomes

- Changes in the other four subscales of the KOOS over 7 months covering (a) symptoms, (b) function in daily living, (c) function in sports/ recreational activities and (d) quality of life.
- Changes in knee pain intensity over 7 months as determined by a 7-day knee pain protocol applying the numerical rating scale (NRS) [37,38].
- Changes in maximum strength of the hip/leg extensors ("leg press") over 7 months
- Changes in objective lower-limb function (30s sit-to-stand test) over 7 months

**Exploratory outcomes**
- Changes of total body-fat content and lean body mass over 7 months as determined by a direct segmental multi-frequency bioelectrical impedance analysis (DSM-BIA)
- Changes in pain medication use as determined by 7-day knee pain protocol over 7 months

**Outcome measures**
Participants were requested to refrain from intense physical activity and exercise 48 hours before the assessments. Baseline and FU assessments were consistently performed by the same research assistant using the identically calibrated devices, in exactly the same setting and at about the same time of the day (±90 min).

**Knee pain diary and questionnaire**
Knee pain and self-reported functional status was determined using the KOOS questionnaire [39,40] which comprises five subscales (dimensions): pain, other symptoms, activities of daily living (ADL), sports and recreation function (Sport/Rec) and knee-related quality of life (QoL). Each of these dimensions is scored separately, using a Likert scale with five possible answers ranging from 0 (no problems) to 4 (extreme problems). According to a formula, described in detail by Roos [39,40], scores are transformed to a 0–100 scale, with zero representing extreme knee problems and 100 representing no knee problems.

In addition to the KOOS subscale pain, the intensity of knee pain was monitored using a numerical rating scale from 0 (no pain) to 10 (worst possible pain) [37,38] conducted over 7 days, before and during the last week of the intervention. We provided standardized logs and requested the participants to rate their highest daily knee pain intensity every evening. The average 7-day pain intensity at baseline and FU was included in the analysis. Additionally, participants were asked to record pain medication daily in their logs. Average numbers of days using analgesics during the 7-day periods of monitoring were included in the analysis.

Lastly, we asked all participants in a baseline questionnaire for demographic parameters, diseases, medication and confounding lifestyle factors (physical activity, exercise and nutrition). The follow-up questionnaire specially addressed changes of this parameters in order to detect factors that may confound our results.

**Functional testing**
Maximum isokinetic hip-/leg-extension strength was tested using a linear isokinetic leg press (CON-TREX LP, Physiomed, Laipersdorf, Germany). Maximum strength was measured unilateral on the index limb (affected knee). Participants were sitting in a slightly supine (seatback 55°) position, fixed by hip and chest straps. Using the standard velocity of 0.5 m/s, range of motion was within 30° to 90° knee angle. After briefing and one familiarization trial with low effort, participants were requested to conduct two sets of five repetitions each with maximum voluntary effort ("push as strongly as possible") separated by 60 s of rest. The highest force value of the two trials was included in the analysis. The present protocol has been applied in prior studies (e.g. [15,16,41,42]).

In order to determine the physical function of the lower extremities (objective lower-limb function), the 30-second sit-to-stand test ("Chair Rise Test") was used, which is a recommended performance-based test in individuals with knee OA [43]. With arms folded across their chests, participants were
instructed to complete as many sit-to-stand movements as possible from a chair within 30s. Knees and hips had to be extended in the standing position, while the buttocks had to touch the seat in the lower position. Following a demonstration by the tester, a practice trial of one repetition was given to check proper form, followed by the 30s test trial. We did not adjust the seat height for lower extremity length. The same standard chair was used for all assessments [44,45].

**Anthropometry**

**Body mass and composition** was determined through direct-segmental, multi-frequency Bio-impedance-Analysis (DSM-BIA; InBody 770, Seoul, Korea). This device measures impedance of the trunk, arms and legs separately using an eight-point tactile electrode system that applies six frequencies between 1 and 1000 kHz.

**Sample size calculation**

The sample size analysis was based on the primary endpoint of KOOS-Pain. Since there is a lack of data on the effect of WB-EMS in OA, the power analysis was based on the effects of conventional strength training on pain in knee OA. In the meta-analysis by Goh et al. [46], a sub-analysis (89 studies; n = 7184) on the effect of strength training compared to "usual care" showed an SMD of 0.73 (0.49 - 0.98). With a power of 80% and an α-level of 5%, a two-sided t-test results in a required number of cases of n = 31/group. Since the meta-analysis of Goh et al. included predominately passive control groups, while our study implemented a usual care control group (6 physiotherapeutic sessions), we designed our sample size analysis more conservatively by increasing the number of cases by 15% which is equivalent to assuming an SMD of 0.67. Correspondingly, we aimed to include 36 subjects per group (WB-EMS: n=36, CG: n=36).

**Randomization and blinding**

Using two strata for pain intensity (NRS, assessed as inclusion criteria), the 72 eligible participants were allocated to the study groups based on drawing small opaque capsules placed in a bowl. In detail, 36 capsules of WB-EMS and 36 capsules of CG were put in the bowl, prepared by a researcher not involved in the trial. Thus, neither participants nor researcher knew the allocation beforehand (allocation concealment). After the randomization procedure, the principal investigator (SK) registered participants and instructed them in detail about study specifications.

Our blinding strategy focused on research assistants who assessed the outcome parameters and were kept unaware of the participants’ group status (WB-EMS or CG) and were not allowed to ask, either.

**Statistical analysis**

Intention to treat (ITT) analyses were applied. Multiple imputation (ITT) was performed using R statistics software (R Development Core Team Vienna, Austria [47]) in combination with Amelia II [48]. We used the full data set for multiple imputations, with imputation repeated 100 times. Over imputation diagnostic plots (“observed versus imputed values”) were checked by Amelia II. For pooling, the results R package mice [49] was used. Additionally, we applied per protocol (PP) analyses for all participants with complete datasets (baseline and 7-months assessment), independent of their compliance, for all the primary and secondary study outcomes. The results of PP and ITT analyses were similar and identical with respect to significances. Assumptions, such as normal distribution, were checked graphically (qq-plots, residual plots). The changes over time within groups were analysed by paired t-tests. The group differences at follow-up (“effects”) were determined by ANCOVA, adjusting
for baseline data using the group as covariate. Categorical variables were addressed using the Chi-Square test. Differences in use of pain medication (yes vs no) were determined by a two-way Analysis of Deviance (logistic regression) using the likelihood-ratio-test. All tests were 2-tailed and significance accepted at p <0.05. According to the suggestion of Li et al. [50], we did not adjust secondary outcomes for multiplicity. Standardized Mean Difference (SMD) according to Cohen (Cohen’s d) [51] was also calculated to indicate the size of the effect for primary and secondary outcome variables. SMDs ≥0.2, 0.5 and 0.8 represent small, medium and large effect sizes.

**Results**

A total of 440 women and men responded by email or telephone. After sending detailed study information via email, potential participants were further assessed for eligibility by phone calls. Of the remaining 113 participants, 12 were unwilling to be randomly assigned to the groups, 6 were unwilling to attend MRI and 23 declined to participate for other reasons. Finally, 72 participants could be included in the study. Participant flow through the study is displayed in Figure 2.

![Figure 2. Study flow diagram (according to CONSORT, Consolidated Standards of Reporting Trial)](image)

Table 1 lists the baseline data for the two groups. Of the 72 subjects randomized, 4 subjects were lost to FU for reasons unrelated to the study (CG: n=1; WB-EMS: n=3) (Fig. 2). Two participants of the WB-EMS group quit the intervention. One of these persons quit the trial after 11 weeks of training because
of orthopaedic problems unrelated to the exercise program. The second person quit after 5 months of training because of personal reasons. Please add Table 1 about here.

On average, participants attended 88% ± 10% of WB-EMS sessions (3 times per fortnight) over the period of 7 months (including condition). In most cases, the reason given for the absence was illness, whereby three participants had longer periods (4-8 weeks) of inactivity due to viral infections. No adverse or unintended effects or injuries were observed during the WB-EMS sessions, and no participant reported any WB-EMS-related discomfort during or after WB-EMS application. More than 90% of the participants in the CG have redeemed the prescription with the 6 physiotherapy treatments. The participation rate regarding the self-management program was around 50%. Both groups participated equally.

Table 2 displays the results of primary and secondary outcomes. KOOS-Pain scores improved significantly more in the WB-EMS group compared with the CG (mean difference (MD) 9.0 points, 95% CI 2.9 to 15.1, p=0.004). In detail, the score improved by 12.5% in CG (p=0.003) and by 30.7% in the WB-EMS (p<0.001). Thus, we confirmed our primary hypothesis that 7 months of WB-EMS application positively changes knee OA pain as assessed by KOOS-Pain subscale more than control. Please add Table 2 about here.

All secondary outcomes (other KOOS subscales, NRS, sit-to-stand test, muscle strength) also improved significantly more in the WB-EMS group compared to the control group at 7-month FU (Table 2). More in detail, in KOOS-Symptoms score there was a net benefit in favour of the WB-EMS group of 14.7% (MD 8.6 points, 95% CI 2.8 to 14.4). The result for KOOS-ADL score was similar: WB-EMS improved the score by 16.2% compared to CG (MD 10.8 points, 95% CI 5.3 to 16.3). The fourth and fifth KOOS dimensions Sport/REC and QoL also changed more favourably in the WB-EMS. The Sport/REC score was 49.2% (MD 11.5 points, 95% CI 3.3 to 19.6) and the QoL score was 33.9% (MD 9.5 points, 95% CI 3.1 to 16.0) higher in the WB-EMS than in the CG.

In parallel, the average knee pain intensity (NRS), which was recorded via 7-day diary, decreased significantly in WB-EMS by 25.3% compared to the CG (MD -1.04, 95% CI -1.75 to -0.33). The number of “sit-to-stands” in 30s (Chair Rise) developed in favour of the WB-EMS compared to the CG (MD 3.9 reps, 95% CI 2.0 to 5.8). In line with the changes in sit-to-stand test, there was a significant between-group difference for change in maximum isokinetic hip/leg extensor strength (MD 79.0 N, 95% CI 6.9 to 151.2) favouring WB-EMS group.

Table 3 displays the results of the exploratory outcomes. In contrast to the results described above, the WB-EMS program did not lead to a significant change or between-group differences in body weight. With respect to body composition, lean body mass remained stable in WB-EMS, whereas it significantly decreased (p=0.02) in the CG. The difference between the groups was non-significant (p=0.09). CG significantly gained fat mass (Tab. 3), whereas the increase in fat mass in WB-EMS group was not significant. Again, the between group difference were not significant (Tab. 3). Please add Table 3 about here.

No significant between-group differences with respect to physical activity (p=0.106), exercise or diet were reported. The weekly intake of analgesics, assessed via 7-day protocol, tendentially increased in the CG (BL: 0.81±2.47; FU: 1.36±2.85) and decreased in the WB-EMS (BL: 0.64±1.33; FU: 0.32±1.36). The intergroup difference was borderline non-significant (p=0.059). Of note, the number of subjects...
who took oral analgesics, as determined via the 7-day protocol, was 8 in CG and 9 in WB-EMS at baseline. At FU 10 participants in CG and 2 in WB-EMS used oral analgesics. After 7 month of intervention a significant reduction of number of participants taking analgesics in the WB-EMS compared to CG was observed (p= 0.033).

Discussion

In the present study, we examined whether a 7-month WB-EMS training program improves knee pain and function in individuals with symptomatic knee OA. In summary, our findings demonstrated that WB-EMS was highly effective in alleviating pain (KOOS) as primary outcome and improving the other four KOOS scores. Along with the enhancement of the KOOS scores, WB-EMS was more effective in improving pain intensity (NRS), objective lower-limb function (30s sit-to-stand) and maximum strength of hip-/leg extensors compared to a usual care approach.

To our knowledge, only one other study investigated the effect of WB-EMS in individuals with knee OA [22]. However, the pilot study of Park et al. included individuals with early knee OA (KL 1-2) and pain was not an inclusion criterion. Accordingly, the baseline KOOS-Pain score in their study was on average 18 points higher compared to our study. The study of Park [22] also pursued a fundamentally different approach: they examined the effectiveness of isometric strength exercise superimposed by WB-EMS compared to the exercises alone and a passive control. Worth mentioning, the isometric exercises alone showed an effect on maximum knee extension strength and the KOOS scores symptoms, ADL, Sports/Rec and QoL compared to passive control. However, the WB-EMS application led to additional effects. The KOOS scores for pain, symptoms and ADL were significant higher in the combined WB-EMS group compared to exercise alone [22].

We pursued a low-threshold approach in which the muscles are activated predominantly via EMS while performing light and less strenuous movements. This method might be attractive especially for the large target group of people who are not willing or able (e.g. because of pain) to perform intensive and strenuous strength training exercises. Following our philosophy of low barriers, the training frequency was 3 sessions per fortnight, compared to 3 sessions per week in Park’s study.

All other studies that have investigated the effect of EMS – mostly the term neuromuscular electrical stimulation (NMES) is used in literature – in knee OA have only used a local stimulation. The results of two recent meta-analysis on the effect of local EMS in individuals with knee OA indicate an increase in quadriceps muscle strength [52], but no significant reduction in pain [21,52].

It has to be noted that WB-EMS is not comparable with local EMS. The difference is not just that WB-EMS stimulates all major muscle groups at the same time. By using cuff electrodes, agonists and antagonists (e.g. quadriceps and hamstrings) are activated simultaneously over a large area. In most of the local EMS studies, the quadriceps muscle was stimulated in isolation with adhesive electrodes. This approach appears suboptimal, considering the importance of the hamstring muscles and intermuscular and proprioceptive coordination for the stability of the knee joint [53]. Strengthening the hamstring muscles in addition to strengthening the quadriceps muscles has even been shown to be beneficial for pain symptoms, mobility and function in knee OA [54]. In our study, we combined WB-EMS with dynamic functional movements because it leads to more pronounced effects on muscle mass and function than static, passive WB-EMS [55]. In the majority of studies on local EMS, the muscles are stimulated statically without movement or passively without movement and without voluntary activation of the muscles.

We focussed on overweight participants, because overweight/obesity is a strong risk factor for the development and progression of knee OA [3,56,57]. Study results suggest that not only the higher
mechanical stress is associated with obesity, but in particular the visceral fat with its pro-inflammatory
effect plays a role in the development and progression of OA [58]. In this context, it should be
mentioned that our WB-EMS program did not result in any significant intergroup differences in weight,
muscle mass and fat mass, even though an increase in fat mass and a decrease in LBM was recorded
within the CG. From this perspective, the effects of our WB-EMS training program on body composition
are rather small. Our WB-EMS approach was time-efficient and required only 30 minutes of training
per week. The low training volume was probably not sufficient to induce major body composition
changes. However, study results suggest that muscle activity is associated with the secretion of anti-
inflammatory substances, which could be one mechanism of pain reduction [23,59]. There is some
evidence of positive effects of WB-EMS application on inflammatory biomarkers in elderly women with
early knee OA [22].

The pain-relieving effect of WB-EMS could take place via different pathways. Another pathway could
be an improvement in knee joint stability and mechanics through an increase in muscle strength as we
observed in the study. Finally, the EMS current, which is a TENS current, may have contributed to the
effect [60].

Our project has various strengths. Great emphasis was placed on the safety aspect. This refers to an
individual dosage and a slow progressive increase in intensity to ensure safety and adaptation of the
muscles. To achieve that, we conducted 1 month of conditioning with an initial lower intensity (i.e.
current intensity) and a shorter application duration to prepare the participants well for the WB-EMS
training. The aim of this method was to avoid high levels of creatine kinase (CK) after initial applications
[61]. Moreover, we wanted to ensure that the training sessions set over threshold stimuli for the whole
period of 6 months. After the initial phase, an RPE target of “6-7” on the Borg CR10 was used. Lastly,
the training was carried out by qualified trainers with a supervision ratio of 1:2 (trainer:participant) to
ensure a high level of safety through optimal assistance and monitoring.

We observed a high attendance rate (88%). Further it indicated that our exercise protocol was not only
effective but obviously attractive, even in this cohort with a low affinity to conventional resistance
training. The high attractiveness was confirmed by the low drop-out rate, as there were only 3
dropouts in the WB-EMS group (all were unrelated to the program). No participant showed intolerance
to electrical stimulation and no EMS related side effects were reported.

Apart from its effectiveness and safety, high importance was attached to generalizability and
transferability. We included a representative cohort of individuals with knee OA and we applied a WB-
EMS protocol used in the majority of commercial settings. This ensures a good transferability of the
results and enables the findings to be applied more broadly using existing structures of commercial
providers.

In order to rule out the possibility of the use of pain medication distorting the results, we recorded the
medications as part of the pain diary. It was notable that the number of participants taking pain
medication significantly decreased in the WB-EMS group and the amount of medication taken
decreased tendentially, which excludes the possibility that the medication distorted the study results.

Some limitations of our trial should be noted. One limitation is that it was not blinded at participant
level. To be blinded, the CG would have had to receive the identical intervention as the training group,
with the difference that the WB-EMS devices would have provided electrical stimuli only below
motorical threshold. However, since low-threshold electrical stimuli, applied as transcutaneous
electrical nerve stimulation (TENS), showed pain-relieving effects in individuals with knee OA [60], we
did not use a blinded study design with low-intensity TENS, but pragmatically implemented a usual
care CG. In this context, it should be mentioned once again that the exercises performed during WB-
EMS were designed in such a way that they should not lead to muscular adaptations. However, it
cannot be ruled out that the dynamic movements without electrical stimulation also had a pain-
relieving effect. Our design does not allow us to separate the possible effects of WB-EMS and the
movements. Another limitation is that OA was not uniformly defined radiologically as an inclusion criterion using the Kellgren-Lawrence score. Since, for reasons of time and economy, no application was made to the Federal Office for Radiation Protection for the production of X-ray images, we examined existing X-ray images and, if not available or too old, MRI images were taken. However, with this procedure, the likelihood of including KL 0 and 1 knees or knees with end stage structural OA (KL4) was minimized [30].

According to various international guidelines [6,7,62], targeted physical training is a critical component of the treatment of knee OA. In summary, we could show that 3 times per fortnight of WB-EMS positively effects knee pain and function in individuals with knee OA. The effects in our study were at least as pronounced as those in studies in which conventional strength training was used [46]. Due to its time efficiency, low weight-bearing joint load and low subjective effort, WB-EMS has the potential to reach the large target group of individuals with knee OA who are not receptive to physical training. However, WB-EMS is an exclusive and more expensive form of training compared to conventional training, which in turn restricts the target group.

Data availability

Data relative to this work will be available upon reasonable request to the corresponding author.


16. Kemmler, W. *et al.* Whole-body electromyostimulation to fight sarcopenic obesity in community-dwelling older women at risk. Resultsof the randomized controlled FORMOsA-


31. Micke, F. *et al.* Similar Pain Intensity Reductions and Trunk Strength Improvements following Whole-Body Electromyostimulation vs. Whole-Body Vibration vs. Conventional Back-


47. R: A Language and Environment for Statistical Computing. v. 4.3.0 (R Foundation for Statistical Computing, Vienna, Austria., 2023).


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Author contributions

All authors made substantial contributions to the study conception and design, data acquisition, or data analysis and interpretation; drafting the article or revising it critically for important intellectual content; providing final approval of the manuscript for submission. The specific contributions of the authors were as follows: Study conception and design: S.K., S.v.S. and W.K.; Data analysis and interpretation: M.K., S.K., S.v.S. and W.K.; Drafting the article: S.K. and S.v.S.; Critically editing and revising the article: S.K., S.v.S., W.K., F.R., A.G., A.M., M.K. and M.U. All authors reviewed and approved the final version of the manuscript.

Additional information

Competing interests: The author(s) declare no competing interests.
Figure and table legend

Figure 1. WB-EMS training session (Written informed consent was obtained from the participants to publish this picture)

Figure 2. Study flow diagram (according to CONSORT, Consolidated Standards of Reporting Trial)

Table 1. Baseline characteristics of the study participants

Table 2. Baseline data and changes of primary and secondary outcomes in the WB-EMS and CG.

Table 3. Baseline data and changes of exploratory outcomes in the WB-EMS and CG.
**Table 1. Baseline characteristics of the study participants**

<table>
<thead>
<tr>
<th>Variable</th>
<th>CG (n=36)</th>
<th>WB-EMS (n=36)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>57.9 ± 7.0</td>
<td>58.3 ± 7.2</td>
</tr>
<tr>
<td>Gender (women/men) [n]</td>
<td>24 / 12</td>
<td>22 / 14</td>
</tr>
<tr>
<td>Body mass index (BMI) [kg/m²]</td>
<td>29.3 ± 3.6</td>
<td>31.1 ± 4.6</td>
</tr>
<tr>
<td>Body height [cm]</td>
<td>174.3 ± 9.0</td>
<td>173.2 ± 9.9</td>
</tr>
<tr>
<td>Body mass [kg]</td>
<td>89.5 ± 15.1</td>
<td>93.2 ± 15.1</td>
</tr>
<tr>
<td>Lean body mass (LBM) [kg]</td>
<td>58.1 ± 11.8</td>
<td>60.2 ± 12.5</td>
</tr>
<tr>
<td>Total body fat [%]</td>
<td>35.0 ± 7.7</td>
<td>35.2 ± 9.2</td>
</tr>
<tr>
<td>Physical activity [Score] ¹</td>
<td>3.70 ± 1.11</td>
<td>3.58 ± 1.28</td>
</tr>
<tr>
<td>No exercise [n] ²</td>
<td>12 (33%)</td>
<td>13 (36%)</td>
</tr>
<tr>
<td>Knee pain intensity [NRS] ³</td>
<td>4.07 ± 1.61</td>
<td>4.05 ± 1.45</td>
</tr>
</tbody>
</table>

All values are expressed as mean value ± standard deviation.

CG, control group; NRS, numeric rating scale (0-10); WB-EMS, whole-body electromyostimulation group.

¹ self-rated physical activity (“very low” (1) to “very high” (7), assessed by questionnaire
² assessed by questionnaire
³ average knee pain intensity, assessed by 7-day protocol
Table 2. Baseline data and changes of primary and secondary outcomes in the WB-EMS and CG.

<table>
<thead>
<tr>
<th>Outcome</th>
<th>CG (n=36)</th>
<th>WB-EMS (n=36)</th>
<th>Difference</th>
<th>SMD</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>MV ± SD</td>
<td>MV ± SD</td>
<td>MV (95% CI)</td>
<td>d^1</td>
<td></td>
</tr>
<tr>
<td><strong>KOOS Pain</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>56.1 ± 12.9</td>
<td>54.4 ± 12.4</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FU</td>
<td>63.1 ± 15.1</td>
<td>71.1 ± 13.9</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Changes</td>
<td>7.0 ± 13.6**</td>
<td>16.7 ± 13.9***</td>
<td>9.0 (2.9 to 15.1)</td>
<td>0.65</td>
<td>.004</td>
</tr>
<tr>
<td><strong>KOOS Symptoms</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>57.5 ± 15.4</td>
<td>57.7 ± 14.5</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FU</td>
<td>61.7 ± 15.3</td>
<td>70.3 ± 13.4</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Changes</td>
<td>4.1 ± 13.8 ns</td>
<td>12.6 ± 14.1***</td>
<td>8.6 (2.8 to 14.4)</td>
<td>0.62</td>
<td>.004</td>
</tr>
<tr>
<td><strong>KOOS ADL</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>64.6 ± 13.6</td>
<td>65.1 ± 13.9</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FU</td>
<td>68.0 ± 13.2</td>
<td>79.1 ± 12.6</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Changes</td>
<td>3.4 ± 13.7 ns</td>
<td>14.0 ± 13.9***</td>
<td>10.8 (5.3 to 16.3)</td>
<td>0.78</td>
<td>&lt;.001</td>
</tr>
<tr>
<td><strong>KOOS Sports/REC</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>33.1 ± 21.1</td>
<td>28.8 ± 20.8</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FU</td>
<td>41.4 ± 22.5</td>
<td>50.2 ± 19.2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Changes</td>
<td>8.3 ± 18.7*</td>
<td>21.4 ± 19.1***</td>
<td>11.5 (3.3 to 19.6)</td>
<td>0.61</td>
<td>.007</td>
</tr>
<tr>
<td><strong>KOOS QoL</strong></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>33.3 ± 16.5</td>
<td>31.4 ± 13.2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FU</td>
<td>39.1 ± 18.5</td>
<td>47.4 ± 13.6</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Changes</td>
<td>5.7 ± 14.3*</td>
<td>16.0 ± 14.7***</td>
<td>9.5 (3.1 to 16.0)</td>
<td>0.66</td>
<td>.004</td>
</tr>
<tr>
<td><strong>Knee pain intensity (NRS)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>4.07 ± 1.60</td>
<td>4.05 ± 1.45</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FU</td>
<td>3.31 ± 1.87</td>
<td>2.26 ± 1.29</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Changes</td>
<td>-0.76 ± 1.73*</td>
<td>-1.78 ± 1.75***</td>
<td>-1.04 (-1.75 to -0.33)</td>
<td>0.60</td>
<td>.005</td>
</tr>
<tr>
<td><strong>Maximum isokinetic Hip/Leg Extensor Strength [N]</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>749.2 ± 224.8</td>
<td>798.5 ± 230.5</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FU</td>
<td>778.5 ± 235.6</td>
<td>903.4 ± 278.9</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Changes</td>
<td>29.3 ± 151.3 ns</td>
<td>104.9 ± 152.6***</td>
<td>79.0 (6.9 to 151.2)</td>
<td>0.52</td>
<td>.03</td>
</tr>
<tr>
<td><strong>Sit-to-stand test (Chair Rise) [n]</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>17.7 ± 6.6</td>
<td>18.7 ± 5.9</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FU</td>
<td>18.2 ± 7.53</td>
<td>23.0 ± 5.74</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Changes</td>
<td>0.53 ± 4.06 ns</td>
<td>4.30 ± 4.07***</td>
<td>3.9 (2.0 to 5.8)</td>
<td>0.96</td>
<td>&lt;.001</td>
</tr>
</tbody>
</table>

All values are expressed as mean value (MV) ± standard deviation (SD).

CG, control group; CI, confidence interval; FU, 7-months follow-up; KOOS, Knee injury and Osteoarthritis Outcome Score (0-100, 0=extreme problems, 100=no problems); NRS, numeric rating scale (0-10, 0=no pain, 10=worst possible pain); SMD, standardized mean difference; WB-EMS, whole-body electromyostimulation group.

^1 Small effect; ^2 measured unilateral (knee of interest)

*p<0.05; **p<0.01; ***p<0.001; ns non-significant (changes within groups)
### Table 3. Baseline data and changes of exploratory outcomes in the WB-EMS and CG.

<table>
<thead>
<tr>
<th></th>
<th>CG (n=36)</th>
<th>WB-EMS (n=36)</th>
<th>Difference</th>
<th>SMD</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>MV ± SD</td>
<td>MV ± SD</td>
<td>MV (95% CI)</td>
<td>d¹</td>
<td></td>
</tr>
<tr>
<td><strong>Body fat content [%]</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>35.0 ± 7.7</td>
<td>35.2 ± 9.2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FU</td>
<td>36.2 ± 8.1</td>
<td>35.6 ± 9.1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Changes</td>
<td>1.21 ± 1.95***</td>
<td>0.42 ± 2.02 ns</td>
<td>-0.79 (-1.73 to 0.15)</td>
<td>0.40</td>
<td>.098</td>
</tr>
<tr>
<td><strong>Lean body mass [kg]</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>58.1 ± 11.8</td>
<td>60.2 ± 12.5</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FU</td>
<td>57.4 ± 11.7</td>
<td>60.1 ± 11.8</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Changes</td>
<td>-0.62 ± 1.58*</td>
<td>-0.08 ± 1.62 ns</td>
<td>0.62 (-0.10 to 1.35)</td>
<td>0.39</td>
<td>.09</td>
</tr>
<tr>
<td><strong>Pain medication [weekly dose]</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>0.81±2.47</td>
<td>0.64±1.33</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FU</td>
<td>1.36 ± 2.85</td>
<td>0.32 ± 1.40</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Changes</td>
<td>0.56 ± 2.38 ns</td>
<td>-0.31 ± 2.43 ns</td>
<td>-0.98 (-1.97 to 0.04)</td>
<td>0.41</td>
<td>.059</td>
</tr>
</tbody>
</table>

All values are expressed as mean value (MV) ± standard deviation (SD).

CG, control group; CI, confidence interval; FU, 7-months follow-up; SMD, standardized mean difference; WB-EMS, whole-body electromyostimulation group.

¹ d ≥ 0.2 small effect; d ≥ 0.5: moderate effect; d ≥ 0.8: high effect

* p<0.05; ** p<0.01; *** p<0.001; ns non-significant (changes within groups)