Phisical modalities on the functional performance in knee osteoarthritis: a sytematic review

Modalidades físicas no desempenho funcional na osteoartrite de joelho: uma revisão sistemática

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Abstract

Introduction: Despite recent advances in the treatment of osteoarthritis (OA), few studies have evaluated the longitudinal effect of physical modalities in functional capacity in patients with knee OA. Thereby, since the physical components and pain can affect the functional performance of daily activities, the effect of these treatment's form is still to be established. Objective: Evaluate the effectiveness of therapeutic ultrasound, electrical stimulation and phototherapy in the functional performance, in patients with knee osteoarthritis. Methods: Articles present in the PubMed, Lilacs, SciELO and PEDro's databases were evaluated. The used keywords were “pulsed ultrasound therapy”, “ultrasound therapy”, “electric stimulation” and “low level laser therapy” in combination with “knee osteoarthritis”. Were included in this presented review, randomized clinical studies using ultrasound, electrical and laser stimulation in subjects with knee osteoarthritis. To evaluate the methodological quality of the selected studies, was used the PEDro's scale. The dependent variables of the study were: pain, physical function, joint stiffness, life quality and functional performance.
Results: 268 studies were found, of these, 41 studies met eligibility criteria and were classified for analysis in full. The used methodology in the studies varied widely, however, in most cases there was improvement in functional performance of individuals with knee OA, with the use of physical modalities, for the pulsed ultrasound, continuous ultrasound, electrical stimulation and laser resources. Conclusion: The physical modalities used in the studies demonstrated improvement in functional performance of individuals with knee OA.

Keywords: Osteoarthritis. Treatment. Physical Therapy.

Introduction

Osteoarthritis (OA) is a chronic, progressive and degenerative osteo-articular disease characterized by arthralgia, stiffness and joint function limitation. The OA’s etiology involves biomechanical, biochemical and genetic’s factors that contribute to the instability between articular cartilage’s synthesis and destruction (1) may affect muscle performance (2).

The OA mainly affects the joints that support weight discharge and, among them, the knee joint is the most affected (1, 3).

The knee OA may affect the activities of daily life, reducing the strength, power and muscular endurance, providing decrease in proprioceptive acuity and body balance. These changes may affect the subjective perception of pain, stiffness and physical function and hinder the performance of functional activities, such as, walking, lifting and sit in a chair and up and down stairs (2).

Despite the availability of treatments provided for patients with knee osteoarthritis, the option is initially for non pharmacological interventions due to no deleterious side effects and are less aggressive for the patient compared to pharmacological and surgical treatments (4). The objective of non pharmacological treatments, including physical therapy, is the relief of the signs and symptoms of the disease and, if possible, the delay in progression, being that various treatments have been used for this purpose, including therapeutic ultrasound (5), electrical stimulation (5, 6) and the low power laser (7).

Despite recent advances in OA's treatment, few studies have evaluated the longitudinal effect of
Methods

Search strategies

The following electronic databases were searched from January to March 2014: PubMed, Lilacs, SciELO and PEDro. The keywords used were: “pulsed ultrasound therapy”, “ultrasound therapy”, “electric stimulation” and “low level laser therapy” in combination with “knee osteoarthritis”.

Two evaluators (LF and HC) independently selected the studies based on titles, excluding those which were not related to this review’s subject. After the selection, the evaluators reviewed the summaries of selected articles to identify those which met the inclusion and exclusion criteria of the study to then be analyzed in detail.

Eligibility criteria

The following inclusion criteria were used in this study:

1) Types of studies: only randomized controlled experiments involving application of physical modalities (ultrasound, electrical stimulation and laser) in patients with knee OA were selected.

2) Types of participants: studies involving patients diagnosed with knee OA were selected. The diagnosis was established based on valid instruments, such as the classification criteria of the American College of Rheumatology (8), radiographic or laboratory evidence or medical records. No other restriction on the duration of the disease and intensity were applied.

3) Types of interventions: studies comparing physical modalities (therapeutic ultrasound, electrical stimulation and low-level laser therapy), with groups without treatment (control) or placebo groups.

4) Types of result’s measures: the dependent variables of the study were: 1) pain, 2) physical function, or 3) joint stiffness. If available, data on life quality and functional performance, served as secondary outcome measures.

5) Score greater than or equal to 5 on the PEDro scale (9).

We selected randomized controlled experiments published in English, Portuguese or Spanish over the last 10 years.

Selection of studies

After removal of duplicate titles, summaries of all identified articles were analyzed by two reviewers. Full articles were then read in detail and the eligible ones were included in this systematic review.

Data collection

Two reviewers independently extracted data on study characteristics, such as participants, interventions, control conditions, co-interventions, outcome measures and results. Disagreements were analyzed by a third reviewer and resolved by discussion.

Endpoints

The study endpoints were defined with the use of tests to evaluate the physical and functional performance. The Western Ontario and McMaster Universities Osteoarthritis Index (WOMAC) (10) was used for the evaluation of the self-reported state of osteoarthritis and the Visual Analogue Scale for pain evaluation.

Methodological quality rating

The PEDro scale (9), which is based on the Delphi list (11), Portuguese translated in 2009, was used for methodological quality evaluation of the studies.
Were included in the study, articles higher than or equal to 5, since studies scored equal to or exceeding 5 (50%) are considered high quality, according to Moseley et al. (12).

**Results**

In the bibliographic research they were initially found 268 studies, of these, 148 were excluded because they are not in accordance with the proposed goal from the title or being duplicates, leaving 120 articles for the summary reading. 69 articles were excluded because they did not meet any of the study eligibility criteria, leaving 51 articles for full and detailed reading. Finally, 10 studies were excluded due to lower score than 5 on the PEDro scale. Were included 41 trials, reaching 2442 patients total (Figure 1). The sample size of the 41 studies ranged from 3 to 175 individuals with OA. To facilitate the visualization of the articles included in this review, more detailed results are described in Table 1.

The studies included in this review were divided to discuss the results according to each physical mode. Therefore, of these studies, 41, 4 referred to the pulsed therapeutic ultrasound, 7 to the continuous therapeutic ultrasound, 8 neuromuscular electrical stimulation, 12 to transcutaneous electrical nerve stimulation, 3 pulsed electrical stimulation and 7 to the low level laser therapy.

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**Figure 1** - Study selection results for keywords.
<table>
<thead>
<tr>
<th>Study</th>
<th>Sample</th>
<th>Groups</th>
<th>Program</th>
<th>Instruments</th>
<th>Endpoints Evaluated</th>
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<tbody>
<tr>
<td>Tascioglu et al. (2010)</td>
<td>82</td>
<td>CU x PU x placebo</td>
<td>CU: 1 MHz, 2 W/cm², 5 min. PU: 1 MHz, 2 W/cm², 1:4; 5 minutes. 10 sessions.</td>
<td>VAS, WOMAC, WT 20 minutes, goniometer</td>
<td>Pain, stiffness, physical function, ROM, functional performance.</td>
<td>All, ↑ pain, stiffness and physical function, PU ↑↑.</td>
<td>8</td>
</tr>
<tr>
<td>Mao-Hsiung Huang et al.</td>
<td>120</td>
<td>PE x CU + PE x PU + PE x CG</td>
<td>CU: 1 MHz e 1.5W/cm², 5 minutes per each spot, totaling 25 cm². PU: 1 MHz e 2.5W/cm², 1:4; 5 minutes per each spot, totaling 25 cm². 24 sessions.</td>
<td>VAS, Lequesne, WT 50 minutes, goniometer, dynamometry.</td>
<td>Pain, discomfort, physical function, MS, ROM, functional performance.</td>
<td>All, except o CG, ↑ pain, discomfort and muscle strenght, with PU ↑↑, CU e PU ↑ walking speed e ROM. PU ↑↑ walking speed.</td>
<td>5</td>
</tr>
<tr>
<td>Huang et al. (2005)</td>
<td>140</td>
<td>PE x PU + PE x PU + PE + HA x CG</td>
<td>PU: 1 MHz, 2.5 W/cm², 1:4 per 5 minutes for each treated spot, on total of 25 cm². 24 sessions.</td>
<td>VAS, Lequesne, WT 50 minutes, goniometer, dynamometry.</td>
<td>Pain, discomfort, physical function, ROM, MS, functional performance.</td>
<td>All, except CG, ↑ MS, pain and discomfort. Groups with PU ↑ROM and functional performance. PE+ PU +HA ↑ functional performance.</td>
<td>7</td>
</tr>
<tr>
<td>Cakir et al. (2013)</td>
<td>58</td>
<td>PE + CU x PE + PU x PE + placebo</td>
<td>CU: 1 MHz e 1 W/cm², 12 minutes. PU: 1 MHz, 1 W/cm², 1:4, 12 minutes 10 sessions.</td>
<td>VAS, WOMAC, WT 20 minutes.</td>
<td>Pain, stiffness, physical function, functional performance.</td>
<td>All ↑</td>
<td>7</td>
</tr>
<tr>
<td>Mascarin et al. (2012)</td>
<td>40</td>
<td>TENS x CU x PE</td>
<td>TENS: 100Hz e 50 µs, int till the sensory threshold, 20 minutes CU: 1MHz, 0.8 W/cm², 3 a 4 minutes 24 sessions.</td>
<td>VAS, WOMAC, goniometer, WT6.</td>
<td>Pain, physical function, functional performance, ROM.</td>
<td>PE e CU ↑functional performance. All ↑ pain e physical function.</td>
<td>6</td>
</tr>
<tr>
<td>Ozgonenel et al. (2008)</td>
<td>67</td>
<td>CU x placebo</td>
<td>CU: 1 MHz / 1 W/cm², 5 minutes 10 sessions.</td>
<td>VAS, WOMAC, WT 50 minutes.</td>
<td>Pain, physical function, stiffness, functional performance.</td>
<td>CU ↑</td>
<td>7</td>
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<tr>
<td>Luksurapan et al. (2013)</td>
<td>46</td>
<td>CU x PP</td>
<td>CU: 1 MHz e 1W/cm², 10 minutes. PP: 1 MHz e 1W/cm², 10 minutes + Piroxicam. 10 sessions.</td>
<td>VAS, WOMAC.</td>
<td>Pain, stiffness, physical function.</td>
<td>Both ↑, PP ↑↑, but without significant difference.</td>
<td>10</td>
</tr>
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<tr>
<td>Kulcu et al. (2009) (19)</td>
<td>45</td>
<td>PEF x CU x CG</td>
<td>PEF: 2 Hz, 100 Hz, 25 Hz, consecutively, 35 minutes. CU: 1 MHz, 1.5 W/cm², 10 minutes.</td>
<td>VAS, WOMAC.</td>
<td>Pain, stiffness, physical function.</td>
<td>PEF e CU ↑.</td>
<td>5</td>
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<tr>
<td>Özgüçlü et al. (2010) (20)</td>
<td>40</td>
<td>PEF + HC + CU + PE x HC + CU + PE</td>
<td>PEF: 50 Hz; 30-G; 90 seconds break, 30 minutes HC: 20 minutes CU: 1 MHz e 1.5 W/cm², 5 minutes 10 sessions.</td>
<td>VAS, WOMAC.</td>
<td>Pain, stiffness, physical function.</td>
<td>Both ↑</td>
<td>6</td>
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<tr>
<td>Ulus et al. (2012) (21)</td>
<td>42</td>
<td>CU + HC + IC + PE x Placebo + HC + IC + PE</td>
<td>CU: 1 MHz e 1W/cm², 10 minutes HC: 20 minutes IC: 10 minutes 15 sessions.</td>
<td>VAS, WOMAC, WT 50 minutes, Lequesne, HADS.</td>
<td>Pain, physical function, stiffness, functional performance, discomfort, psychological state.</td>
<td>Both ↑</td>
<td>7</td>
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<tr>
<td>Bennell et al. (2005) (22)</td>
<td>119</td>
<td>MT x Placebo</td>
<td>MT: 12 weeks of service + 12 weeks of self-management.</td>
<td>VAS, WOMAC, Likert scale, SF-36, AQol, dynamometry, step test for balance.</td>
<td>Pain, physical function, stiffness, patient’s global change, life quality, MS, body balance.</td>
<td>Both ↑ pain and patient’s global change.</td>
<td>8</td>
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<tr>
<td>Imoto et al. (2013) (6)</td>
<td>82</td>
<td>NMEE + PE x CG</td>
<td>NMEE: 50 Hz, 250µs, maximum tolerated intensity, TON: 10s, TOFF: 30s, 20 minutes</td>
<td>VAS, TUG, Lequesne, DAL scale.</td>
<td>Pain, physical function, functional performance, discomfort, DAL.</td>
<td>NMEE ↑ pain, physical function, discomfort and DAL.</td>
<td>7</td>
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<tr>
<td>Vaz et al. (2013) (23)</td>
<td>12</td>
<td>NMEE</td>
<td>NMEE: 80 Hz, 400ms, maximum intensity tolerated. 24 sessions.</td>
<td>WOMAC, ultrasonography, dynamometry.</td>
<td>Pain, stiffness, physical function, muscle structure, MS.</td>
<td>NMEE ↑ muscle thickness increase and fascicle length, MS, pain, stiffness and physical function.</td>
<td>-</td>
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<tr>
<td>Elboim-Gabyzon et al. (2013) (24)</td>
<td>50</td>
<td>NMEE + PE x PE</td>
<td>NMEE: 75 Hz, 250µs, maximum tolerated intensity, TON: 10s, TOFF: 50s, 45 minutes 12 sessions.</td>
<td>VAS, WOMAC, WT10m, TUG, U’nDT, myometry.</td>
<td>Pain, stiffness, physical function, functional performance, muscle performance.</td>
<td>NMEE + PE ↑ pain and muscle activation.</td>
<td>5</td>
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<tr>
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<tr>
<td>Durmus et al. (2007) (25)</td>
<td>50</td>
<td>NMEE x Biofeedback + PE</td>
<td>NMEE: 50 Hz, 200µs, visible contraction intensity, TON: 10s, TOFF: 10s, 20 minutes 20 sessions.</td>
<td>VAS, WOMAC, MR test, 10 MR, WT 50 minutes, U’nDT.</td>
<td>Pain, stiffness, physical function, functional performance, MS.</td>
<td>Both ↑</td>
<td>5</td>
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<tr>
<td>Gaines et al. (2004) (26)</td>
<td>38</td>
<td>NMEE x CG</td>
<td>NMEE: visible contraction intensity, TON: 10s, TOFF: 50s, 15 minutes.</td>
<td>Pain diary, McGill pain quiz, AIMS2-PS.</td>
<td>Pain.</td>
<td>NMEE ↑ pain only after 15 minutes.</td>
<td>5</td>
</tr>
<tr>
<td>Bruce-Brand et al. (2012)</td>
<td>41</td>
<td>PE x NMEE x CG</td>
<td>NMEE: 50 Hz, 100-400µs, maximum tolerated intensity, TON: 10s, TOFF: 50s, 20 minutes.</td>
<td>WOMAC, WT 25 minutes, GUSDT, U’nDT, SF-36, dynamometry, magnetic resonance.</td>
<td>Pain, stiffness, physical function, functional performance, life quality, muscle performance.</td>
<td>NMEE e PE ↑ functional performance and increased cross-sectional area of the quadriceps.</td>
<td>5</td>
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<tr>
<td>Burch et al. (2008) (28)</td>
<td>116</td>
<td>IC + standardized muscle stimulation x TENS</td>
<td>IC: 5.000Hz, 1 e 150Hz, mild tingling intensity, 15 minutes standardized muscle stimulation: 50Hz for 200ms each 1500ms, intensity between 3.39µs e 102.2µs, with average of 16.26 mA, 20 minutes TENS: 0.2 Hz, 300µs, 0.5mA of intensity, 35 minutes</td>
<td>VAS, WOMAC.</td>
<td>Pain, stiffness, physical function, life quality.</td>
<td>IC + standarized muscle stimulation ↑</td>
<td>5</td>
</tr>
<tr>
<td>Palmieri-Smith et al. (2010)</td>
<td>30</td>
<td>NMEE x CG</td>
<td>NMEE: 2.500Hz, 50 bursts/s, maximum tolerated intensity, TON: 10s, TOFF: 50s, 10 contractions. 12 sessions.</td>
<td>WOMAC, dynamometry, WT 12-19 minutes.</td>
<td>Pain, stiffness, physical function, MS and muscle activation, functional performance.</td>
<td>↔</td>
<td>7</td>
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<tr>
<td>Law et al. (2004) (30)</td>
<td>34</td>
<td>multiple frequencies TENS x placebo</td>
<td>TENS: 25-35mA of intensity, Frequency: 2Hz ou 100Hz or 2/100Hz alternated; 57µs or 200µs or 57/200µs, respectively, 40 minutes 10 sessions.</td>
<td>VAS, goniometer, TUG.</td>
<td>Pain, ROM, functional performance.</td>
<td>All, except placebo, ↑</td>
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Transcutaneous electrical nerve stimulation:
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<tbody>
<tr>
<td>Atamaz et al. (2012) (31)</td>
<td>175</td>
<td>TENS + PE x IC + PE x SWD + PE x placebos + PE</td>
<td>TENS: 80 Hz, 10-30mA intensity, IC: 100 Hz, 4KHz, tactile sensation intensity. SWD: 27.12MHz, 300W input and average of 3.2W. 15 sessions.</td>
<td>VAS, WOMAC, goniometer, WT15m, Nottingham Health Profile.</td>
<td>Pain, stiffness, physical function, ROM, functional performance, life quality.</td>
<td>TENS, IC, active SWD, lower intake of paracetamol.</td>
<td>9</td>
</tr>
<tr>
<td>Cetin et al. (2008) (32)</td>
<td>100</td>
<td>SWD + HC + PE x TENS + HC + PE x CU + HC + PE x HC + PE x PE</td>
<td>SWD: I: 27.12 MHz, 15 minutes TENS: 60-110 Hz, 60 µs, no contraction maximum intensity, 20 minutes. CU: 1 MHz, 1.5 W/cm², 10 minutes.</td>
<td>VAS, Lequesne, ISKO, WT 50 minutes, dynamometry.</td>
<td>Pain, discomfort, physical function, functional performance, MS.</td>
<td>All ↑ pain and incapacity (ISKOA). All, except isolated PE, ↑ pain, discomfort, physical function e MS. SWD e TENS ↑↑</td>
<td>6</td>
</tr>
<tr>
<td>Cheing et al. (2004) (33)</td>
<td>62</td>
<td>TENS x placebo x PE x TENS + PE</td>
<td>TENS: 80 Hz, 140µs, tingling intensity, 60 minutes, 20 sessions.</td>
<td>Dynamometry, spatiotemporal parameters of the march, goniometer.</td>
<td>Muscle performance, ROM.</td>
<td>TENS + PE shows a trend in the improvement of physical parameters, but showed no significant difference.</td>
<td>5</td>
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<tr>
<td>Pietrosimone et al. (2011) (34)</td>
<td>36</td>
<td>TENS + PE x placebo + PE x PE</td>
<td>TENS: 150 Hz, 150µs, strong sensorial intensity, for at least 8 hrs a day, 12 sessions.</td>
<td>WOMAC, dynamometry.</td>
<td>Pain, stiffness, physical function, muscle activation.</td>
<td>TENS + PE ↑ muscle activation. All ↑ pain, stiffness e physical function.</td>
<td>7</td>
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<tr>
<td>Kolen et al. (2012) (35)</td>
<td>74</td>
<td>TENS on different spots.</td>
<td>TENS: 80 Hz, 100µs, maximum tolerated intensity, 30-45 minutes.</td>
<td>VAS, WOMAC, WT6’, dynamometry, goniometer, HADS, Pain Anxiety Symptoms Scale, Pain Catastrophizing Scale, satisfaction quiz.</td>
<td>Pain, stiffness, physical function, functional performance, muscle performance, ROM.</td>
<td>TENS ↑ pain. ↑↑ when applied on low skin resistance spots.</td>
<td>7</td>
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<tr>
<td>Selfe et al. (2008) (36)</td>
<td>37</td>
<td>NIN x placebo.</td>
<td>TENS on low resistance spots, 20-30 minutes, 17 sessions.</td>
<td>VAS, WOMAC, SF-36.</td>
<td>Pain, stiffness, physical function, global evaluation, life quality.</td>
<td>NIN ↑ pain, without meaningful differences. NIN ↑ vitality (subscale SF-36) and global evaluation of the patient.</td>
<td>7</td>
</tr>
<tr>
<td>Vance et al. (2012) (37)</td>
<td>75</td>
<td>High frequency TENS x low frequency TENS x placebo</td>
<td>TENS: 100Hz or 4Hz, 100µs, 10% below the motor threshold intensity, 40-50 minutes, 1 session.</td>
<td>VAS, quantitative sensory testing, TUG.</td>
<td>Pain, functional performance.</td>
<td>All, except placebo, pressure pain. All ↑ rest pain TUG.</td>
<td>8</td>
</tr>
<tr>
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<tr>
<td>Pietrosimone et al. (2009) (38)</td>
<td>33</td>
<td>TENS x crioteraphy x CG</td>
<td>TENS: 150Hz, 150µs, sensory intensity, 45 minutes Crioteraphy: 2 bags of 1.5L of ice, one on the front and another on the back of the knee, 20 minutes 1 session.</td>
<td>Dynamometry.</td>
<td>Muscle activation.</td>
<td>All, except CG ↑ quadriceps activation immediately after application.</td>
<td>6</td>
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<tr>
<td>Chen et al. (2013) (39)</td>
<td>50</td>
<td>HA x TENS</td>
<td>TENS: 3 Hz and 20 Hz, 200µs, maximum tolerated intensity, 20 minutes 12 sessions. HA: 2.5ml, 5 shots.</td>
<td>VAS, Lequesne, goniometer, WT, pressure algometry.</td>
<td>Pain, physical function, discomfort, ROM, functional performance, global evaluation, DAL.</td>
<td>TENS ↑pain, discomfort and physical function.</td>
<td>6</td>
</tr>
<tr>
<td>Paker et al. (2006) (40)</td>
<td>60</td>
<td>HA x TENS</td>
<td>TENS: 150Hz, 20 minutes per 1 hour. HA: 3 shots of hylan GF20.</td>
<td>WOMAC, Lequesne, SF-36.</td>
<td>Pain, stiffness, physical function, discomfort, life quality.</td>
<td>Both ↑pain, physical function e stiffness. HA ↑↑ physical function.</td>
<td>6</td>
</tr>
<tr>
<td>Pietrosimone et al. (2010) (41)</td>
<td>36</td>
<td>TENS + PE x placebo + PE x PE</td>
<td>TENS: 150Hz, 150µs, Strong intensity without contraction for at least 8 hours a day.</td>
<td>WOMAC, dynamometry, three-dimensional march analysis at 15 meters.</td>
<td>Pain, stiffness, physical function, functional performance, muscle activation, MS.</td>
<td>All ↑ → knee flexion angle.</td>
<td>7</td>
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<tr>
<td>Fary et al. (2009) (42)</td>
<td>3</td>
<td>PES</td>
<td>PES: 100Hz, 2ms, subsensorial intensity, for at least 8 hours a day.</td>
<td>VAS, Likert scale, SF-36, global perceived effect scale, accelerometry.</td>
<td>Pain, physical function, global evaluation, life quality, exercise.</td>
<td>PES ↑ symptoms, supported by up to 16 weeks.</td>
<td>-</td>
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<tr>
<td>Garland et al. (2007) (43)</td>
<td>58</td>
<td>PES x placebo</td>
<td>PES: 100 Hz, subsensorial intensity, 6 hours or more.</td>
<td>VAS, WOMAC.</td>
<td>Pain, stiffness, physical function, global evaluation.</td>
<td>PES ↑</td>
<td>8</td>
</tr>
<tr>
<td>Fary et al. (2011) (44)</td>
<td>70</td>
<td>PES x placebo</td>
<td>PES: 100Hz, 4ms, 7 hours a day.</td>
<td>VAS, WOMAC, SF-36, health survey, Human Activity Profile, accelerometry, perceived global effect scale.</td>
<td>Pain, stiffness, physical function, life quality, exercise.</td>
<td>Both ↑</td>
<td>10</td>
</tr>
</tbody>
</table>

**Pulsed electrical stimulation:**

- Fary et al. (2009) (42)
- Garland et al. (2007) (43)
- Fary et al. (2011) (44)
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<th>Study</th>
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<tr>
<td>Kheshie et al.</td>
<td>53</td>
<td>High intensity AL + PE x low intensity AL + PE x placebo + PE</td>
<td>High intensity AL: Total power of 1250 J, 610 to 810 mJ/cm² application. Low intensity AL: Total power of 1250 J, 800 mW, 1 kHz, 50 J/cm².</td>
<td>VAS, WOMAC. Pain, physical function.</td>
<td>Both active ↑ High intensity AL ↑↑</td>
<td>7</td>
</tr>
<tr>
<td>Alfredo et al.</td>
<td>40</td>
<td>AL + PE x placebo + PE</td>
<td>AL: 904 nm, 700 Hz, 60 mW average, 20 mW maximum, 27 J, 3 J/point total dose, 50 s.</td>
<td>VAS, WOMAC, Lequesne, goniometer, dynamometry.</td>
<td>Pain, physical function.</td>
<td>AL ↑ pain, physical function e ROM.</td>
</tr>
<tr>
<td>Alghadir et al.</td>
<td>40</td>
<td>AL x placebo</td>
<td>AL: 850 nm, 50 mW, 6 J/poin, 60 s/poin, with 48 J/cm². 8 sessions.</td>
<td>VAS, WOMAC, WT 15 minutes.</td>
<td>Pain, physical function, functional performance.</td>
<td>AL ↑</td>
</tr>
<tr>
<td>Hegedus et al.</td>
<td>27</td>
<td>AL x placebo</td>
<td>AL: 830 nm, 50 mW, 6 J/poin, 48 J/cm² dose. LP: 0.5 mW. 8 sessions.</td>
<td>VAS, Ritchie index, goniometer, thermography.</td>
<td>Pain, ROM, local microcirculation, pressure sensitivity.</td>
<td>AL ↑</td>
</tr>
<tr>
<td>Al Rashoud et al.</td>
<td>49</td>
<td>AL on acupuncture points + PE x placebo</td>
<td>AL: 830 nm, 30 mW, 6 J total per session, 1.2 J/poin, 4 J/cm², 5 points, 40 s. 9 sessions.</td>
<td>VAS, SKFS.</td>
<td>Pain, physical function.</td>
<td>AL ↑</td>
</tr>
<tr>
<td>Yurtkuran et al.</td>
<td>52</td>
<td>AL on acupuncture points x placebo</td>
<td>AL: 904 nm, 4 mW, 0.48 J dose, 10 mW/cm² power density, 120 s/poin. 10 sessions.</td>
<td>VAS, WT50m, knee circumference, Medial Tenderness Score, Nottingham Health Profile, WOMAC.</td>
<td>Pain, functional performance, edema, knee sensibility, life quality, stiffness, physical function.</td>
<td>AL ↑ edema reduction.</td>
</tr>
<tr>
<td>Tascioglu et al. (2004)</td>
<td>60</td>
<td>3J AL x 1.5J AL x placebo</td>
<td>AL: 830 nm, 50 mW. 10 sessions.</td>
<td>VAS, WOMAC.</td>
<td>Pain, stiffness, physical function.</td>
<td>↔</td>
</tr>
</tbody>
</table>

Note: CU: continuous ultrasound; PU: pulsed ultrasound; ROM: range of motion; VAS: visual analog scale; WOMAC: Western Ontario and McMaster Universities Osteoarthritis; WT: walking test; PE: physical exercise; CG: control group; MS: muscle strength; HA: hyaluronic acid; SWD: shortwave diathermy; HC: hot compresses; TENS: Transcutaneous electrical nerve stimulation; ISKO: index of severity for knee osteoarthritis; WT6': walking test of 6 minutes; PP: piroxicam phonophoresis; PEF: pulsed electromagnetic field; HADS: Hospital Anxiety and Depression Scale; MT: multimodality therapy; SF-36: life quality quiz; NMEE: Neuromuscular Electrical Stimulation; DAL: daily life activities; TUG: timed get up and go; TO: timed up and down stairs test; MR: maximum resistance; AIMS2-PS: arthritis impact measurement scale 2-pain subscale; GUSDT: get up and sit down test; IC: intensityerferential current; NIN: Noninvasive intensityeractive neurostimulation; PES: pulsed electric stimulation; AL: active laser; SKFS: saudi knee function scale; TO: time on; TOFF: time off.

↑: Effective  ↔: Ineffective  ↑↑: Greater efficacy
Discussion

Therapeutic Ultrasound

Studies comparing pulsed and continuous ultrasound application effectiveness on pain, physical function stiffness, discomfort, range of motion (ROM), functional performance and muscle strength (FM), demonstrated that both application forms can be effective for patients with knee osteoarthritis. However, the increase in the variables analyzed was better in the group receiving the application of pulsed ultrasound (13, 14).

Huang et al. (15) checked the efficacy of exercise performed alone, pulsed ultrasound + exercise and pulsed ultrasound + exercise + hyaluronic acid in pain variables, discomfort and fibromyalgia. It was found that the groups with pulsed US were effective to increase the range of motion and functional performance and hyaluronic acid group was more effective in functional performance, discomfort and fibromyalgia. The parameters used in these studies (13-15) varied as: continuous US (1 MHz, 1.5 to 2 W/cm², 5 minutes) and pulsed US (1 MHz, 2 to 2.5 W/cm² working cycle 1:4, 5 minutes) with treatment time from 10 to 24 sessions.

Discordant results were obtained in the study of Cakir et al. (16), who reported improvement in pain, physical function, stiffness and functional performance, at the continuous US (1 MHz and 1 W/cm², 12 minutes), pulsed US (1 MHz, 1 W/cm² and a working cycle 1:4, 12 minutes) and placebo US groups, with no differences among the groups, after 10 therapy sessions. All groups performed home exercises and the authors suggest that these exercises masked the US effects.

Mascarin et al. (5) have proven the effectiveness of 24 sessions with transcutaneous electrical neurostimulation therapy (100Hz and 50µs, sensitive threshold intensity, 20 minutes), continuous US (1 MHz, 0.8 W/cm², 3-4 minutes) and exercise in pain and physical function. The presented results demonstrated efficacy only of continuous US and exercise in functional performance. Concordant to this study, Özgonenel et al. (17) observed the effectiveness of continued US (1 MHz and 1 W/cm², 5 to 10 minutes) in pain, physical function, stiffness and functional performance for 10 sessions compared to the placebo group. Luksurapan et al. (18) found no significant difference between the Piroxicam phonophoresis and continuous US, because both treatments showed improvements.

Kulcu et al. (19) verified the effectiveness of pulsed electromagnetic field therapy (PEMF) (frequency: 2 Hz, 100 Hz, 25 Hz, consecutively, 35 minutes/session) and continued US (1 MHz, 1.5 W/cm², 10 minutes) in pain, stiffness and physical function. Both therapies were effective for the analyzed variables. In contrast, Özgüçlü et al. (20) found no additional effects of pulsed electromagnetic field therapy (50 Hz, with an intensity of 30-G in a 90s interval, 30 minutes) to treatment with continuous US (1 MHz and 1.5 W/cm², 5 minutes), hot pack and exercise, with the same pain improvement, physical function and stiffness in treatments without pulsed electromagnetic field for 10 sessions.

Discordant results were described by Ulus et al. (21) who verified efficiency of both the continuous US (1 MHz and 1 W/cm² for 10 minutes) and the placebo US associated with interferential current (IF), hot packs and exercise, in pain, physical function, stiffness, functional performance, discomfort and psychological state, without difference among the groups, for 15 sessions. The authors infer that some positive US studies have low methodological quality and cite the use of other agents in the study may have masked the final effect and the study have a low sample size.

Bennell et al. (22) observed that both the multimodal therapy (taping, exercises, mobilization and massage) as the placebo US are effective in pain and patient global assessment, showing that only contact with the therapist can lead to positive changes in these patients.

Briefly, the 9 studies using ultrasound as a therapeutic strategy, 7 demonstrated positive effects on variables. There seems to be an indication that when the therapeutic ultrasound is isolated applied and compared to other therapies, show similar effectiveness to other therapeutic techniques. However, when comparing the different ultrasound application forms (continuous or pulsed) is suggested that the pulsed application mode appears to be more effective for improvement in functionality variables.

Neuromuscular electrical stimulation

Scientific evidences used in this review show that the neuromuscular electrical stimulation (NMES)
may be beneficial in patients with knee OA in variables such as pain (6, 23-26), physical function (6, 23, 25), functional performance (25, 27), discomfort (6), daily living activities (6) and stiffness (23, 25). Furthermore, NMES seems to compensate physiological declines occurring in the muscle of patients, acting both in structure as in function of the quadriceps (23) by means of cross-sectional area increases (23, 27) and muscular strength (23, 25) and also in the improvement of voluntary muscle activation when combined with an exercise program (24). The parameters used in these studies (6, 23-27) varied from: 50-80Hz, 100-400μs, intensity: visible muscle contraction until the maximum tolerated, time on: 10 seconds, off time: 10-50 seconds 15-45 minutes per session, for 12-36 sessions.

Burch et al. (28) reported the efficacy of standard muscle stimulation (50Hz for 200ms to 1500ms each, intensity between 3.39μs and 102.2μs with an average production of 16.26mA, 20 minutes) in pain, stiffness, physical function and quality when compared to low-intensity TENS (0.2 Hz, 300μs, 0.5mA, 35 minutes), after 8 weeks. The standard muscle stimulation refers to stimulation characteristics: triphasic stimulation patterned based on the normal activation time of the quadriceps and hamstrings, during a high-level run. Before the standardized muscle stimulation, the group received IR current (5.000Hz, pre-modulated between 1Hz and 150Hz, tingling intensity, 15 minutes).

Disagreeing with the presented studies, Palmieri-Smith et al. (29) didn’t find gains in strength or activation in the quadriceps muscle after application of NMES (2,500 Hz AC, 50 bursts per second, maximum tolerated intensity, time on: 10 seconds, time off: 50 seconds, with 10 electrically induced contractions, for 12 sessions). The authors believe that the lack of effect may be due to low dysfunction of the quadriceps presented in the voluntaries of this study or due to the dosing of the intervention.

From 7 studies using neuromuscular electrical stimulation, 6 reported increases in physical and functional role. In studies that have shown positive effects of the application of NMES on the parameters analyzed, the NMES average parameters were: frequency 50 Hz, pulse duration 200-250 microseconds and the maximum intensity tolerated by the patient. There seems to be an indication that the largest increases in muscle strength and functionality of the patients are inversely related to the deficit of muscle strength previous to treatment, in other words, the higher the deficit of muscle strength previous to treatment, the greater are the possibilities of improvement with use of NMES.

Transcutaneous electrical nerve stimulation

Law et al. (30) demonstrated the efficacy of different TENS parameters (2Hz, 100 Hz and 2/100Hz, 576 uS, 200μs and 576μs/200μs, respectively, comfortable intensity levels, 40 minutes) in pain, functional performance and ROM. The authors found that TENS was superior to independent placebo application of current application parameters. Atamaz et al. (31) assessed the TENS efficacy (80Hz, 10-30mA), IF current, SWD, as well as the placebos interventions for each resource associated with exercise, in pain, physical function, functional performance and life quality, for 15 sessions. The authors found that the proposed therapies were effective in the variables analyzed in the study, and that the intake of analgesics was lower during the active interventions compared with placebos. In contrast, Cetin et al. (32) studied the effect of short-wave diathermy (SWD) (27.12 MHz, field condenser technique, 15 minutes), transcutaneous electrical nerve stimulation (TENS) (60-110 Hz, 60 uS, maximum intensity without contraction 20 minutes), continuous US (1 MHz, 1.5 W / cm², 10 minutes) associated with a hot bag and exercise, compared with group of hot bag and exercise only or just exercise. The authors examined the effectiveness of SWD, TENS and continuous US interventions in pain, discomfort, physical function and MS. However, the best increments in the analyzed variables were obtained in SWD and TENS groups.

Studies (33, 34) have demonstrated the TENS effectiveness associated with physical exercise on muscle activation and a trend towards physical parameters improvement, when compared to isolated treatments. The parameters varied: 80-150Hz, 140-150μs, sensory intensity, 60 minutes and 8 hours a day, 12-20 sessions.

Kolen et al. (35) demonstrated that for the TENS application (80Hz, 100ms, maximum tolerated intensity, 30-45 minutes) to be effective, it must be applied in locations with lower electrical skin resistance to reduce pain. However, when functional is the therapeutic goal, TENS can be applied on the greatest pain spots or random locations.
Selfe et al. (36) studied the efficacy of adjuvant therapy of non-invasive interactive neurostimulation (NIN) in the “vitality” subscale of the SF-36 questionnaire and the overall patient evaluation, when compared to placebo, in 17 sessions for 20 to 30 minutes per session with progressive intensity. The NIN therapy refers to the application of TENS on acupuncture points, with low skin resistance. The NIN therapy also resulted in clinically significant reductions in pain, but without differences among groups, which may be explained by the small sample size and frequency of treatment have been less than ideal, since the frequency of weekly treatment decreased over study.

Only two studies (37, 38) analyzed the effectiveness of a single TENS session in knee OA. Vance et al. (37) observed efficacy of TENS (100 and 4Hz, 100µs and intensity of 10% below the motor threshold, 40-50 minutes) and placebo treatment in pain at rest and during walking, however, only TENS groups increased the pressure pain threshold. The authors concluded that the placebo effect would possibly be reduced with more treatment sessions. Pietrosimone et al. (38) found that both the TENS (150Hz, 150µs, sensory stimulation, 45 minutes) and the cryotherapy (2 bags of 50,721oz of ice for 20 minutes) has similar effects on quadriceps muscle activation immediately after the therapy application with no difference between the therapy groups.

Studies (39, 40) compared the effect of TENS and hyaluronic acid intra-articular injection in patients with knee OA. Chen et al. (39) observed greater efficiency of TENS (3-20Hz, 200µs, maximum intensity tolerated, 20 minutes) in pain, discomfort and physical function compared to the injection of hyaluronic acid in 12 sessions. Meanwhile, Paker et al. (40) observed efficacy of TENS (150 Hz, 20 minutes to 1 hour, 15 sessions) and hyaluronic acid injection in pain, stiffness and physical function, with greater effect of hyaluronic acid in physical function. The different results between studies may be due to the types of electrodes and hyaluronic acid products that differed in the studies (39).

Disagreeing with the studies presented, Pietrosimone et al. (41) found no effect of TENS (150Hz, 150µs, strong sensory intensity during each exercise session and a minimum of 8 hours a day) associated with exercise in angle and knee flexion peaks during pace. Moreover, the pain, stiffness, physical function, functional performance, muscle activation and MS have been improved in these patients after exercise regardless of the use of TENS. The absence of TENS effects can be attributed to insufficient sample size and the authors declare that the long treatment periods may be necessary to change these variables, since the quadriceps activation rate was higher in TENS group.

Briefly, TENS can be effective in reducing the pain perception and functionality in patients with knee osteoarthritis. For this, the electrodes application area can be a determining factor, because when the goal is to reduce pain, the electrodes must be applied in places of least resistance of the skin, but when the objective is to increase functionality, application of this feature in areas of greatest pain perception can be done. The use of TENS seems to be as or more effective than other analgesic therapies. From 11 articles included in this review, 10 articles evaluated the beneficial effects of TENS and 1 article didn’t notice increases in comparison with the placebo group. However it is believed that the results can be explained due to the sample size and/or due to the insufficient treatment period.

Three studies examined the effectiveness of pulsed electrical stimulation (PES) in the perception of pain and self-reported health status in patients with knee OA (42, 43, 44). Two studies (42, 43) verified efficiency of PES in pain, physical function, stiffness and self-reported health status of the patient, using the following parameters: 100Hz, 2ms, subsensorial intensity for more than 6 hours/day. Furthermore, Fary et al. (42) found that these changes could be sustained for at least 16 weeks. Discordant with previous studies, Fary et al. (44) found no increased effectiveness of PES (100Hz, 4ms, 7 hours a day for 26 weeks) compared to placebo in pain, physical function, stiffness, life quality and exercise. The authors note that the sample may not be representative of the OA population, due to its characteristics.

According to the studies inserted in this review, it appears that three studies have tested the use of pulsed electrical stimulation in patients with knee osteoarthritis. Two studies have shown beneficial effects on pain, patient’s global evaluation, physical function and joint stiffness. One study has shown no positive effects of pulsed electrical stimulation, but we believe the results can be explained because the sample may not be representative of the population of osteoarthritis.
Low-power laser therapy

Kheshie et al. (45) demonstrated the effectiveness of high intensity laser (pulsed YAG laser 1250J) in three phases: initial phase flow was adjusted to two applications of 710 and 810 mJ/cm², intermediate stage of 610 mJ/cm² and final phase, the same as first fluence for 45 minutes) and low intensity laser (BTL-5000 laser, As-Ga of 1250J, 50 J/cm², 830 nm, 1KHz, 800mW for 32.33 minutes) on pain and physical functioning in 12 sessions. The authors demonstrated that both therapies are effective, but the high intensity laser appears to have best results.

Alfredo et al. (7) observed the low-intensity laser efficiency (As-Ga, 27 J, being 3 J for 50 seconds per point, 904 nm, 700 Hz, 60 mW) in pain, physical function and ROM compared to placebo. Agreeing with this study, Alghadir et al. (46) verified the efficacy of 8 low-intensity laser sessions (As-Ga, 50mW, 850nm, 48 / cm² with 6 J / point) in pain, physical function and functional performance compared to placebo condition. Hegedus et al. (47) assessed the efficacy of low level laser (Ga-Al-As, 50 mW, 830 nm, 6 J per point, 48 J / cm² per session) in pain, ROM, pressure sensitivity and local microcirculation in 8 sessions compared the placebo condition.

Al Rashoud et al. (48) demonstrated the effectiveness of low level laser (Ga-As-Al, 1.2 J for 40 seconds per point, totaling 4 J / cm², 830nm, 30mW) applied on acupuncture points and associated with exercise for 9 sessions, in pain and physical function. Yurtkuran et al. (49) also evaluated the laser effectiveness (As-Ga 4 mW, 904nm and 0.48 J per session, 10mW / cm²) on acupuncture points and observed improvement only in the edema seen by knee circumference compared to the placebo group, in 10 sessions. The authors point out that the dose adopted in the study was lower than that recommended by the World Association of Laser Therapy, which may have influenced the results.

Disagreeing with the presented studies, Tascioglu et al. (50) found no efficacy in pain, physical function and stiffness in the different therapies with low intensity laser: Group 1: Laser for 2 minutes per point (Ga-Al-As, 50mW, 830nm, 3J per point, totaling 15J per session, a total of 10 minutes); Group 2: Laser for 1 minute per point (Ga-Al-As, 50mW, 830nm, 1.5J per session, totaling 5 minutes); and Group 3: placebo laser. The authors afirm that the results can be justified by the laser mode, dosages and wavelength adopted in the study that generated an ineffective treatment.

Of the 6 studies inserted in this review, 5 proved to be effective to improve pain and function in patients with knee osteoarthritis. From the analysis of the results presented, it is believed that the therapy with the low intensity laser can improve pain, range of motion and functional performance, especially compared to the placebo condition.

Study limitations

The studies used in this review demonstrate results variations according to each physical modality and should be carefully considered by professionals to choose the best physical modality for each individual. Furthermore, there are some limitations in this study, among which can be highlighted: 1) The inclusion of studies that evaluated the physical modalities associated with other forms of intervention, preventing conclude what the real effect of the isolated application mode is. Though, it is emphasized that such modalities are complementary and are not clinically used in isolation; 2) The difficulty of establishing a treatment protocol facing the results and divergent application methods, the lack of information in some studies, besides the scarcity of published articles.

Conclusion

The studies showed in this review demonstrated variations on the benefits of physical modalities with respect to the used parameters, frequency of treatment, and application sites. Although, it was found based on the last 10 years literature, using high-designed studies, these methods are effective for this population, improving their symptoms and signs. For more treatment effectiveness, these interventions should be adjusted depending on the goal that you want to achieve with each patient, with specific protocols for each clinical condition.

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Received in 08/31/2015

Recebido em 31/08/2015

Approved in 07/28/2016

Aprovado em 28/07/2016