Effectiveness of high-intensity laser therapy in the treatment of musculoskeletal disorders
A systematic review and meta-analysis of randomized controlled trials

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Abstract
Background: Although high-intensity laser therapy (HILT) has been used for the management of musculoskeletal disorders (MSD), studies examining the effectiveness of HILT have been limited. We investigated the effectiveness of HILT in MSD using a systematic review and meta-analysis.

Methods: We searched the ovid MEDLINE, ovid Embase, Cochrane CENTRAL library, and Web of Science until January, 2018. Relevant studies concerning the effectiveness of HILT in patients with MSD were included. Both placebo and active controls were considered as comparators and only randomized controlled trial (RCT) design studies were included. Risk of bias (ROB) was used for the quality assessment of the RCT. For continuous variables, a meta-analysis was conducted using an inverse variance random effects model. The mean difference (MD) for visual analog scale pain and standardized mean difference (SMD) for disability were applied.

Results: Twelve studies were selected for this systematic review. In 11 studies, comprising 736 patients, pain was significantly improved by HILT compared with a control group (MD: −1.01; 95% confidence interval [CI]: −1.28 to −0.74). From the analysis of 688 patients from 10 studies, the pooled standardized mean difference (SMD) of HILT showed a significant improvement in disability scores compared with those in the control group (SMD, −1.09; 95% CI: −1.77, −0.41). In subgroup analysis by treatment regions, the mean difference (MD) in neck pain was the highest at −1.02 (95% CI: −1.45, −0.59) than in controls, followed by back pain (MD, −0.91; 95% CI: −1.24, −0.59).

Conclusions: The results of this study show that HILT treatment for back and neck pain significantly improved pain and disability scores compared with controls. The ROB of the included studies was moderate; however, significant heterogeneity existed. Thus, additional well-designed studies involving larger samples with long-term follow-up are needed to further assess each laser application, treatment region, and comparator.

Abbreviations: CI = confidence interval, CMS = Constant Murley Scale, DASH = Disabilities of the Arm Shoulder and Hand questionnaire, GDP = gross domestic product, HILT = high-intensity laser therapy, HILT = high-intensity laser therapy, LLLT = low-level laser therapy, MD = mean difference, MSD = musculoskeletal disorders, NDI = Neck Disability Index, NSAIDs = nonsteroidal anti-inflammatory drugs, ODI = Oswestry Disability Index, PRTEE = Patient-related Tennis Elbow Evaluation, RCT = randomized controlled trial, ROB = risk of bias, SD = standard deviation, SMD = standardized mean difference, SPADI = Shoulder Pain and Disability Index, VAS = visual analog scale, WMD = weighted mean difference.

Keywords: high-intensity laser therapy, musculoskeletal disorder, pain, systematic review meta-analysis
1. Introduction
Musculoskeletal disorders (MSD) comprise the most common conditions worldwide. One in 2 people have a MSD in the United States and the estimated number of individuals affected reached about 126.6 million in 2012. In particular, individuals with back and neck pain total approximately 75.7 million. The cost of MSD was estimated to be approximately $213 billion in 2011, which is 1.4% of the gross domestic product (GDP) in the United States.

MSD derives from a soft-tissue injury or pain in the musculoskeletal system including muscles, nerves, tendons, joints, and cartilage in the upper and lower limb, neck, and lower back. Various causes of MSD pain can be attributed to result from damage of muscle tissue, trauma, postural strain, repetitive movements, overuse, and prolonged immobilization. The main purpose of MSD treatment is to improve pain and physical disabilities. Treatment for MSD pain includes nonsteroidal anti-inflammatory drugs (NSAID), analgesics, corticosteroid injections, as well as acupuncture or acupressure. In addition, self-management and education, exercise, manual therapy, and psychosocial therapy have also been considered nonpharmacological treatments. Recently, laser therapy including low-level laser therapy (LLLT) and high-intensity laser therapy (HILT) have been used for the management of MSD. Laser therapy is a noninvasive treatment having a low incidence of adverse effects.

In a systematic review, LLLT applied for pain treatment in MSD showed significant differences compared to control groups. HILT can stimulate joints more deeply and treat a wider area than LLLT; thus, the application of HILT for MSD may improve pain and function when compared to LLLT. However, studies regarding the effectiveness for HILT have been limited. Thus, it is necessary to investigate the effectiveness of HILT in MSD patients. The aim of the present study was to perform a systematic review and meta-analysis regarding the effectiveness of HILT in treatment of MSD.

2. Methods
2.1. Literature search
We searched the core databases including ovid MEDLINE, ovid Embase, the Cochrane CENTRAL library, and Web of Science up to January 17, 2018. The MeSH terms and text words included in the search strategy are shown in Supplementary Table 1, http://links.lww.com/MD/C600. To maximize the sensitivity of the included articles, an only intervention-related term was used for searching articles. The search terms included “high-intensity laser therapy” and “HILT.” In the Web of Science database, the term “pain” was also applied to the search strategy.

2.2. Study selection
We included studies conducted to examine the effectiveness of HILT in patients with MSD. Treatment regions for MSD included the back, neck, shoulder, arm, or hands. Placebo or active comparators such as ultrasound, brace, transcutaneous electrical nerve stimulation (TENS), transaction therapy, medical treatment bandage, or exercise were considered as comparators. The randomized controlled trial (RCT) design study was the only design included and studies written in both English and Korean were included. Two reviewers independently selected the studies based on the inclusion criteria. If disagreement occurred, a final decision for inclusion was confirmed by a third reviewer.

2.3. Quality assessment
The risk of bias (ROB) tool developed by Cochrane group was used for quality assessment. We estimated 3 levels (low, unclear, and high) for 7 items including random sequence generation, allocation concealment, blinding of participants, blinding of outcome assessment, incomplete outcome data, selective reporting, and funding source. Two independent reviewers assessed the ROB and a consensus was reached with another reviewer in cases of inconsistency.

2.4. Data extraction
Items for data extraction were prespecified by the authors. Patient characteristics such as diagnosis, mean age, and standard deviation (SD), and percentage of males in the included studies were extracted. Interventions and sample size, evaluation time, and country conducting the study were also summarized. Pain and disability were extracted as outcome measures. For HILT application methods, the type of laser, wavelength, output power, energy density, application time, number of total sessions and sessions per week, application site, and the process of application were listed.

2.5. Statistical analysis
For continuous variables, a meta-analysis was conducted using an inverse variance random effects model. The mean difference (MD) for VAS pain and standardized mean difference (SMD) for disability were applied. VAS pain was reported using the same unit; however, disability measures were reported using diverse units due to the different instruments used. For 3 arm studies, splitting of the shared group was applied according to the Cochrane group guidelines. For example, in studies examining HILT versus Placebo versus active comparator, the HILT group was included in both the HILT versus Placebo and HILT versus active comparator groups. Thus, the number of individuals in the HILT was divided, but the same mean and SD were applied in both groups. This method was applied for continuous outcomes. For disability, a higher score presented higher disability except for one study. Thus, the mean of the study with an opposite value was multiplied by −1 in order to align the direction.

The pooled estimate of the subgroup according to treatment region was analyzed. The placebo comparator was used in subgroup analysis by treatment region or follow-up period, if the study included a 3-arm intervention. The subgroup analysis according to comparators (placebo or active controls) was also estimated. Heterogeneity was assessed using forest plot visually and Higgins $I^2$ with $P$-value. In order to identify publication biases, contour funnel plots were used. The Review Manager 5.3 program was used for all analysis.

3. Results
3.1. Literature search
From the initial literature search, 626 articles were identified and 225 articles were included for title or abstract screening after removing duplicates (Fig. 1). Following the full text review of 107 articles, 94 were excluded because the studies did not focus on HILT, no patients with MSD were included, no outcomes of interest were reported, the study design was ineligible, or papers were not original articles. Ultimately, 12 studies were included in this systematic review.
3.2. General characteristics of the included studies

The included studies consisted of patients having back, neck, shoulder, arm, and hand pain (Table 1). Comparators were varied and included placebo, exercise, ultrasound, transcutaneous electrical nerve stimulation (TENS), brace, transaction therapy, and bandages. The mean age of the subjects included in the studies ranged from 32 to 58 years old and the percentage of males ranged from 0% to 100%. Trials were conducted in Egypt, Saudi Arabia, Turkey, Italy, Poland, China, and South Korea.

Pain was assessed using a visual analog scale (VAS). A 10-cm VAS was used to measure pain and possible scores ranged from 0 (no pain) to 10 (worst pain). Thus, higher VAS scores indicated higher pain. A functional activity assessment was used according to the treatment regions. For low-back pain, disability was measured as the Oswestry Disability Index (ODI) and the Neck Disability Index (NDI) was used for estimating neck disability. The Shoulder Pain and Disability Index (SPADI) and the Constant Murley Scale (CMS) for shoulder pain, and the Patient-related Tennis Elbow Evaluation (PRTEE), and the Disabilities of the Arm Shoulder and Hand questionnaire (DASH) for arm/hand pain were used as disability measures. The outcomes were evaluated after 2 weeks to 12 weeks of treatment. The characteristics used for HILT in the included studies are summarized in Table 2.

3.3. Quality assessment

The results of quality assessment are presented in Fig. 2. In random sequence generation, selective reporting, and funding source, the percentage of low-quality studies was over 75% and also more than 50% indicated blinding of participants of outcome assessment, as well as incomplete outcome data. The percentage of low risk for allocation concealment was below 50% as for most of the included studies it was neither conducted nor reported.

3.4. Pain

In 11 studies including 736 patients, HILT significantly improved pain compared to the control group (MD: −1.01; 95% CI: −1.28, −0.74) (Fig. 3). There was no apparent systematic bias in the contour funnel plot. Although an asymmetry was detected, the missing values were both in significant and non-significant areas (Fig. 4A). In subgroup analysis by treatment regions, the MDs for the neck were the highest at −1.02 (95% CI: −1.45, −0.58) compared to the control group, followed by the back (MD: −0.91; 95% CI −1.24, −0.59) and the arms/hands subgroups (MD: −0.82; 95% CI: −1.43, −0.21). There was no significant difference for the shoulder pain subgroup between the HILT and control groups. The heterogeneity in the neck and shoulder pain subgroups was significant ($I^2 = 73\%$; $P = .02$ for the neck and $I^2 = 88\%$, $P = .004$ in the shoulder subgroups); however, there was no significant heterogeneity in the back pain subgroup ($I^2 = 0\%$; $P = .88$) or the arm/hand subgroup ($I^2 = 0\%$; $P = .42$). In particular, the MD of HILT for pain was −1.03 (95% CI: −1.28, −0.77) and −0.82 (−1.39, −0.26) for the placebo and active control groups, respectively (Table 3). Heterogeneity was
| Study                         | Diagnosis                                      | Sample size and intervention | Mean age (SD) | Male | Evaluation          | Country   | Measures                                                                 |
|-------------------------------|-----------------------------------------------|------------------------------|---------------|------|---------------------|-----------|--------------------------------------------------------------------------|
| Alayat et al 2014[13]         | Chronic low back pain                         | HILT + EX 28, EX 24, HILT 20 | 32.81 (4.48)  | 100% | 4 weeks             | Egypt     | Pain assessment                                                          |
|                               |                                               |                              |               |      |                     |           | **VAS**                                                                  |
|                               |                                               |                              |               |      |                     |           | **MDQ2:** consisting of 10 items. Each item was scored from 0 to 5, with higher scores representing greater disability. |
| Alayat et al 2016[14]         | Chronic neck pain                             | HILT + EX 30, EX 30          | 35.47 (4.18)  | NA   | 6 weeks (after treatment) | Saudi Arabia | Pain assessment                                                          |
|                               |                                               |                              |               |      |                     |           | **VAS**                                                                  |
|                               |                                               |                              |               |      |                     |           | **NDI:** cervical-related disabilities and a valid and reliable measurement tool for patients with neck pain. Measured by 5 questions for a maximum total score of 50. |
| Casale et al 2013[15]         | Symptomatic Carpal tunnel syndrome            | HILT 10, TENS 10             | 57.3 (12.9)   | 50%  | 3 weeks (after treatment) | Italy     | Pain assessment                                                          |
|                               |                                               |                              |               |      |                     |           | **VAS**                                                                  |
| Chen et al 2018[16]           | Lumbar disc protrusion                        | HILT + SDS 32, SDS 31        | 39.27 (12.58), SDS 43.00 (12.19) | 65.1% | 2 weeks             | China     | Pain assessment                                                          |
|                               |                                               |                              |               |      |                     |           | **VAS** (lumbosacral portion pain, lower limb radiation pain)            |
|                               |                                               |                              |               |      |                     |           | **ODI:** composed of 10 items, including pain degree, personal life and self-help ability, carrying, walking, sitting, standing, sleep, sexual life, social life, and travel. Evaluated by total scores within 0–50 points with higher score indicating more serious dysfunction. |
| Dundar et al 2015[17]         | Myofascial pain syndrome of the trapezius     | HILT + EX 38, EX 37          | 40.2 (12.9),  38.4 (12.1) | 0%   | 4 weeks             | Turkey    | Pain assessment                                                          |
|                               |                                               |                              |               |      |                     |           | **VAS** (pain at rest, movement, and night)                              |
|                               |                                               |                              |               |      |                     |           | **MDQ:** measure of changes in functional disability. Each of 10 items is scored from 0 to 5. The maximum score is 50. The obtained score can be multiplied by 2 to produce a percentage score. |
| Dundar et al 2015[18]         | Lateral epicondylitis                         | HILT 30, PL 31, Brace 30    | 32.6 (10.9), PL 33.4 (11.2), Brace 33.6 (9.8) | 54.8% | 4 weeks             | Turkey    | Pain assessment                                                          |
|                               |                                               |                              |               |      |                     |           | **VAS** (pain at rest and under strain)                                  |
|                               |                                               |                              |               |      |                     |           | **PRTEE:** measures the changes in functional disability. 15-item questionnaire designed for patients with lateral epicondylitis. Each of 15 items has response option (0 = no difficulty, 10 = unable to perform). The overall score ranging from 0 (best) to 100 (worst). |
| Study | Diagnosis | Sample size and intervention | Mean age (SD) | Male | Evaluation | Country | Measures |
|-------|-----------|-----------------------------|---------------|------|------------|---------|----------|
| Fiore et al 2011[19] | Low back pain | HILT 15, US 15 | 51.2 (6.0) | 63.3% | 3 weeks (post-treatment) | Italy | Pain assessment | VAS | Functional activity assessment | OLBPDQ measures the changes in functional disability. |
| Halafaj et al 2017[20] | Cervical spondylosis | HILT 86, Traction therapy 88 | 45.5 | 34.9% | 4 weeks, 12 weeks | Poland | Pain assessment: | VAS | Functional activity assessment | ADI: cervical-related disabilities and a valid and reliable measurement tool-Polish version for patients with neck pain. Composed of 10 parts, including pain intensity, personal care and lifting, reading, headaches, concentration, work, driving, sleeping, and recreation. Measures 6 for a maximum total score of 60 (converted into percentages; 0% to 100%). |
| Kim et al 2015[21] | Frozen shoulder | HILT+EX 33, EX 33 | HILT+EX 57.5 (8.7), EX 55.6 (7.9) | 81.8% | 3 weeks, 8 weeks | South Korea | Pain assessment | VAS |
| Pekyavas et al 2016[22] | SAIS | HILT+MT+KT+EX 19, MT+KT+EX 16, KT +EX 20, EX 15 | 47.1 (13.8) | NA | 15 days (post-treatment) | Turkey | Functional activity assessment | SPADI: measure shoulder pathology-related pain and disabilities. Evaluated by total scores within 0–100 points for pain, 0–50 points for function/disability. Higher scores indicated greater pain and disability. | VAS |
| Salli et al 2016[23] | Lateral epicondylitis | HILT 31, Bandage 34 | 46.5 (8.1) | 27.7% | 6 weeks | Turkey | Pain assessment | VAS | Functional activity assessment | DASH: composed of 30 items, assessing the ability to perform upper extremity activities. Each item was scored from 0 to 5. Higher scores represent greater disability. |
| Santamato et al 2009[24] | SAIS | HILT 35, US 35 | 54.1 (9.0) | 40.0% | 2 weeks (after treatment) | Italy | Pain assessment | VAS | Functional activity assessment | CMS: Evaluated by total scores within 0–100 points (35 points for pain and function, 65 points for ROM and strength including activities of daily living-work, recreation, sleep, and ability to work). Higher scores indicate greater pain and disability. |

CMS = Constant Murley Scale, DASH = Disabilities of the Arm Shoulder and Hand questionnaire, EX = exercise, HILT = high-intensity laser therapy, MODQ = Modified Oswestry Disability Questionnaire, NDI = Neck Disability Index, ODI = Oswestry Disability Index, OLBPDQ = Oswestry Low Back Pain Disability Questionnaire, PL = placebo, PRTEE = Patient-related tennis elbow evaluation, SAIS = subacromial impingement syndrome, SD = standard deviation, SPADI = shoulder pain and disability index, TBNS = transcutaneous electrical nerve stimulation, VAS = visual analog scale.
| Study                  | Type of laser/wavelength | Output power/energy density | Total dose (energy)/session | Application time/No. of total sessions/No. of sessions/week | Application sites                                      | Method of application                                                                 |
|-----------------------|--------------------------|-----------------------------|-----------------------------|---------------------------------------------------------------|-------------------------------------------------------|----------------------------------------------------------------------------------------|
| Alayat et al 2014[13] | Pulsed Nd:YAG laser/1064 nm | 3 kW/510–1780 mJ/cm         | 3000 J                      | 15 min/12/3                                                  | Lower-back area of L1–L5 and S1                       | Initial phase: the laser fluency was set to 3 successive subphases of 610, 710, and 810 mJ/cm², for a total of 1400 J with fast manual scanning.  
Intermediate phase: the handpiece toward 8 paravertebral points from L1 to S3, with 25 J, a fluency of 610 mJ/cm², and a time of 14 s at each point, for a total of 200 J.  
Final phase: the same as the initial phase, except that slow manual scanning was used. |
| Alayat et al 2016[14] | Pulsed Nd:YAG laser/1064 nm | 3 kW/510–1780 mJ/cm         | 2250 J                      | 15 min/12/2                                                  | Posterior neck on the paraspinal area, upper back, interscapular area, trapezius, sternocleidomastoid muscles, and posterior and lateral shoulder areas. |
| Casale et al 2013[15] | NIR 830–1064 nm          | 25 W/250 J/cm               | 100 sec/15/7                | Median nerve at the wrist                                    |                                                       | Treated with combined 830–1064 nm laser (radiating dose: 250 J/cm²) delivered to the skin overlying the course of the median nerve at the wrist for 100 s at 25 W (18 W [1064 nm] + 7 W [830 nm]) via a fiber-optic probe with a spot size of ~1 cm².  
The probe was moved along the course of the nerve for 10 cm and it was maintained in an upright position using a cast in order that the incident light remained perpendicular to the skin, to avoid any thermally induced sensations. |
| Chen et al 2018[16]   | Gallium-arsenide diode laser (BTL-6000 laser)/1064 nm | 12W/150 J/cm²              | 7500 J                      | 10 min/10/5                                                  | Rib inferior margin and posterior superior iliac spine | Initial phase: performed with fast manual scanning with a total energy of 3000 J.  
Intermediate phase: key treated areas comprised pain points between rib inferior margin and posterior superior iliac spine, utilizing 1500 J total energy.  
Final phase: same as the initial phase except that slow manual scanning was employed. |
| Dundar et al 2015[17] | Pulsed Nd:YAG laser/1064 nm | 3 kW/360–1780 mJ/cm         | 1060 J                      | 15 min/15/7                                                  | Bilateral trapezius muscles                           | First phase: fast manual scanning (100 cm² per 30 s), both the transverse and longitudinal directions over the bilateral the trapezius muscles, total energy dose of 500 J, 3 subphase of 360 mJ/cm² (166.7 J), 410 mJ/cm² (166.8 J), and 510 mJ/cm² (166.5 J), for a total of 500 J.  
Second phase: the handpiece with spacers fixed vertically at 90° to the trigger points, bilaterally on 3 trigger points (total of 6 points) over the trapezius muscle with 10 J, a fluency of 610 mJ/cm², and a time of 6 s at each point, for a total of 60 J. |

(continued)
| Study               | Type of laser/ wavelength | Output power/ energy density | Total dose (energy)/ session | Application time/No. of total sessions/ No. of sessions/week | Application sites                                                                 | Method of application                                                                 |
|--------------------|---------------------------|----------------------------|------------------------------|---------------------------------------------------------------|----------------------------------------------------------------------------------|-------------------------------------------------------------------------------------|
| Dundar et al 2015  | Pulsed Nd:YAG laser/ 1064 nm | 10.5 W/ 360-1780 mJ/cm²     | 1275 J                       | 15 min/15/7                                                  | Soft tissues near the lateral epicondyle, and extensor muscles extending over forearm from lateral epicondyle | First phase: fast manual scanning (100 cm² per 30 s), CET, soft tissues near the lateral epicondyle, and extensor muscles extending over forearm from lateral epicondyle, 3 subphases of 510 mJ/cm² (208 J), 810 mJ/cm² (208 J), and 970 mJ/cm² (209 J), for a total of 625 J.  
Second phase: CET near the lateral epicondyle, 360 mJ/cm² (5 J), 510 mJ/cm² (9 J), and 610 mJ/cm² (10 J), a time of 6 s at time, for a total of 25 J.  
Third phase: slow manual scanning (100 cm² per 60 s), same area treated in first phase up to a total energy dose of 625 J.  
Starting initial phase: fast manual scanning (100 cm² per 30 s), muscular contracture on the lumbar and dorsal muscles, latissimus dorsi, obliquus externus and magus gluteus, a total of 1200 J.  
Intermediate phase: total of 200 J  
Final phase: slow manual scanning (100 cm² per 60 s), same area treated in first phase until a total energy dose of 1200 J.  
The treatment was started at a distance of 3–5 cm, directly above the transverse processes of each of the cervical vertebrae, from the C4 to Th4.  
The handpiece was moved contact-free in a continuous spiral motion, slightly inward of each of the spinal segments. Within 3.5 min of treatment, 195 J of energy was provided.  
Biostimulation application was carried out paraspinally in a continuous wave mode, using radiation power density P≈300 mW, and continuous handpiece motion parallel to muscle fibers, from C4 to Th4.  
The average energy density was Ed≈50 J/cm² at a wavelength of 980 nm and procedure duration of 6.5 minutes with provided energy of 1250 J.  
Initial phase: rapid manual scanning (100 cm²/30 s) of the anterior joint line and posterior joint line of the shoulder with one shot of 850 mJ at a frequency of 30 Hz. The scanning was performed parallel to the joint line, with the patient’s arm internally rotated on the posterior scan and externally rotated on the anterior scan. The total energy dose administered during this phase was 4000 J.  
Intermediate phase (a fixed scan phase): with one-shot emission of 350 mJ at a frequency of 20–25 Hz. The total delivered energy was 4000 J. |
| Fiore et al 2011   | Pulsed Nd:YAG laser/ 1064 nm | 6 W/760 mJ/cm²              | 2600 J                       | 10 min/15/5                                                  | Muscular contracture on the lumbar and dorsal muscles, latissimus dorsi, obliquus externus and magus gluteus, a total of 1200 J. |
| Halefaj et al 2017 | BLT-6000/980 nm            | 7 W/50 J/cm²                | 1250 J                       | 10 min/10/5                                                  | Cervical vertebrae (C4–Th4).                                                                                                   | The treatment was started at a distance of 3–5 cm, directly above the transverse processes of each of the cervical vertebrae, from the C4 to Th4.  
The handpiece was moved contact-free in a continuous spiral motion, slightly inward of each of the spinal segments. Within 3.5 min of treatment, 195 J of energy was provided.  
Biostimulation application was carried out paraspinally in a continuous wave mode, using radiation power density P≈300 mW, and continuous handpiece motion parallel to muscle fibers, from C4 to Th4.  
The average energy density was Ed≈50 J/cm² at a wavelength of 980 nm and procedure duration of 6.5 minutes with provided energy of 1250 J. |
| Kim et al 2015     | Nd:YAG laser/ 1064 nm      | 8000 W/350–850 mJ/cm²        | 4000 J                       | 15 min/9/3                                                   | Shoulder anterior joint line and posterior joint line.                                                                          |

(continued)
Table 2 (continued).

| Study | Type of laser/wavelength | Output power/energy density | Total dose (energy)/session | Application time/No. of total sessions/ No. of sessions/week | Application sites | Method of application |
|-------|--------------------------|-----------------------------|-----------------------------|---------------------------------------------------------------|-------------------|-----------------------|
| Pekýavas et al 2016[20] | Nd:YAG laser/1064 nm | 3 kW 510–710 J/cm² | 2050 J | 30 min/15/3 | Upper trapezius, supraspinatus, and deltoid muscles and anteriorly for the pectoralis minor muscle of the shoulder | The hand piece was applied vertically perpendicular to the shoulder joint for 5 s. In each scan, the scanning points included both the anterior and posterior joint line of the shoulder. Initial phase: fast manual scanning of the same areas treated in the initial phase and the deltoid area until a total energy dose of 2000 J. Intermediate phase: Application of the handpiece with fixed spacers vertically to 90° on the trigger points including 4 phases for each point until a pain reduction of 70%–80% was achieved in 6 s. In this phase, the mean energy dose was 50 J. Final phase: Slow manual scanning (100 cm²/60 s) of the same areas treated in the initial phase until a total energy dose of 1000 J was achieved. |
| Salli et al 2016[21] | NR | 6 W/150 J/cm² | Approximately 600–900 J | 75 sec for 4 sessions, 750 sec for 6 sessions/10/5 | Elbows with lateral epicondylitis | First 4 sessions: 75 s at a dose of 4 W 6 J/cm² targeting the most painful areas in circular motion from the center to the outside. Subsequent 6 sessions: 30 s at a dose of 6 W 100–150 J/cm² at the pain-inflicting region in linear motion. Intermediate phase: Application of the handpiece with fixed spacers vertically to 90° on the trigger points until a pain reduction of 70%–80% was achieved. In this phase, the mean energy dose was 50 J. Final phase: Slow manual scanning (100 cm²/60 s) of the same areas treated in the initial phase until a total energy dose of 1000 J was achieved. |
| Santamato et al 2009[19] | Pulsed Nd:YAG laser/1064 nm | 1 kW/760 mJ/cm² | 2050 J | 10 min/10/5 | Upper trapezius, supraspinatus, and deltoid muscles and anteriorly for the pectoralis minor muscle of the shoulder | The hand piece was applied vertically perpendicular to the shoulder joint for 5 s. In each scan, the scanning points included both the anterior and posterior joint line of the shoulder. Intermediate phase: Application of the handpiece with fixed spacers vertically to 90° on the trigger points until a pain reduction of 70%–80% was achieved. In this phase, the mean energy dose was 50 J. Final phase: Slow manual scanning (100 cm²/60 s) of the same areas treated in the initial phase until a total energy dose of 1000 J was achieved. |

CET = common extensor tendon, HILT = high-intensity laser therapy, Nd:YAG = neodymium-doped yttrium aluminum garnet, NR = not reported.
Figure 2. Quality assessment of included studies using risk of bias assessment (A) ROB graph and (B) ROB summary. +: low ROB; --: high ROB; ?: unclear ROB. ROB = risk of bias.

Figure 3. Mean difference in visual analog scale pain between high-intensity laser therapy and comparator.
not significant compared to placebo ($I^2=8\%$; $P=.37$), while there was significant heterogeneity compared to the active control group ($I^2=78\%$; $P<.001$). According to the follow-up periods, the pooled pain effect did not show any significant differences.

### 3.5. Disabilities

A total of 688 patients from 10 studies indicated that the pooled SMD of HILT significantly improved disability compared with the control group (SMD: $-1.09$; 95% CI: $-1.77$, $-0.41$) (Fig. 5). We could not detect any obvious publication bias by the contour funnel plot despite asymmetry (Fig. 4B). Both significant and nonsignificant data were missing. In the subgroup analysis for each treatment region, a similar trend in SMD was observed in both the disability scores and pain scores; however, the SMD in the shoulder pain subgroup significantly improved and there was no significant difference in the arm/hand subgroup. The disability following HILT significantly improved compared to the placebo group (SMD: $-0.96$; 95% CI: $-1.28$, $-0.64$), however, there was no significant improvement between the HILT and the active control groups (SMD: $-1.06$; 95% CI: $-2.50$, 0.37) (Table 3). In comparison with the control group, there was no significant heterogeneity ($I^2=47\%$; $P=.09$), while a significant heterogeneity was observed in comparison with the active control group ($I^2=97\%$; $P<.001$). The SMD for disability stratified according to the follow-up period showed improvement with a longer follow-up period, however, the CIs overlapped.

### 4. Discussion

Application of HILT for pain and functional ability of MSD showed significant improvement compared to controls. The studies included in this meta-analysis generally had moderate ROB.

We were unable to find any previous reviews evaluating the effectiveness of HILT for the management of MSD, but a systematic review for LLLT in pain was available. The systematic review included 1462 patients with MSD from 8 studies.$^{10}$ The pain in the LLLT group significantly decreased compared to the control group following treatment (MD: $-0.85$; 95% CI: $-1.22$, $-0.48$), a result, which was similar to our study showing that pain in the HILT-exposed group significantly decreased compared to controls (MD: $-1.01$; 95% CI: $-1.28$, $-0.74$).

Huang et al.$^{25}$ conducted a systematic review for LLLT application in chronic back pain. The weighted mean difference (WMD) of LLLT in pain was $-13.57$ (95% CI: $-17.42$, $-9.72$)

![Figure 4. Contour funnel plot of included studies (A) pain (B) disability.](image)

| Subgroup     | Studies, n | Patients, n | Random effects, MD/SMD [95% CI] | Effect, $P$-value | $I^2$, % | Heterogeneity, $P$-value |
|--------------|------------|-------------|---------------------------------|-------------------|---------|-------------------------|
| **VAS pain** |            |             |                                 |                   |         |                         |
| Comparators  |            |             |                                 |                   |         |                         |
| Placebo      | 6          | 362         | $-1.03$ [$-1.28$, $-0.77$]      | $<.001$           | 8       | $.37$                   |
| Active control | 6         | 404         | $-0.82$ [$-1.39$, $-0.26$]      | $<.001$           | 78      | $<.001$                 |
| Follow-up period |        |             |                                 |                   |         |                         |
| $\leq 4$ weeks | 5         | 246         | $-1.24$ [$-1.65$, $-0.83$]      | $<.001$           | 47      | $.11$                   |
| $>4$ weeks   | 6          | 490         | $-0.89$ [$-1.19$, $-0.60$]      | $<.001$           | 38      | $.16$                   |
| **Disability** |          |             |                                 |                   |         |                         |
| Comparators  |            |             |                                 |                   |         |                         |
| Placebo      | 6          | 334         | $-0.90$ [$-1.28$, $-0.64$]      | $<.001$           | 47      | $.09$                   |
| Active comparator | 5       | 384         | $-1.06$ [$-2.50$, 0.37]         | $<.001$           | 97      | $<.001$                 |
| Follow-up period |        |             |                                 |                   |         |                         |
| $\leq 4$ weeks | 4         | 198         | $-0.81$ [$-1.26$, $-0.36$]      | $<.001$           | 56      | $.08$                   |
| $>4$ weeks   | 6          | 490         | $-1.26$ [$-2.31$, $-0.21$]      | $.02$             | 96      | $<.001$                 |
compared to placebo and there was no significant difference in the disability score using the ODI (SMD: −0.38; 95% CI: −1.14, 0.39). In our study, exposure to HILT in patients with back pain resulted in a significant improvement in both pain and disability scores compared to the control groups (MD: −0.91; 95% CI: −1.24, −0.59 for pain and SMD: −1.21; 95% CI: −1.38, −0.85).

In the study by Chow et al., LLLT applied to patients with neck pain significantly improved pain and disability scores compared to placebo (WMD: 19.86; 95% CI: 10.04–29.68 for pain and SMD: 1.38; 95% CI: 0.39–2.37 for the disability score).[26] These results were similar to the results of our study. The pain and disability scores in the HILT group significantly improved compared to the control group (MD: −1.02; 95% CI: −1.45, −0.58 for pain and SMD: −1.92; 95% CI: −3.61, −0.23).

HILT has been known to reduce heat accumulation in tissues and to have photothermal and photochemical effects in deep tissues for limited periods.[11] These properties favor treatment of deep tissues and structures by increasing cell metabolism, vascular permeability, and blood flow.[13,24,27] The pain control effect achieved by HILT might be attributed to multiple mechanisms. In the central nervous system, the secretion of endogenous opioids such as β-endorphins is increased by laser therapy and these could centrally inhibit pain sensations.[23] In the peripheral nervous system, substance P sensitizes pain-transmitting neurons and leads to hyperalgesia; however, laser therapy has been reported to decrease the secretion of substance P by peripheral receptors.[14,29] Laser therapy might increase the latency and decrease the conduction velocity of sensory nerves by inhibiting A- and C-fiber transmission; these in turn may decrease the transmission of pain signals.[30] In tissues, laser therapy may also reduce the release of histamine and bradykinin in injured tissues and increase the pain threshold.[31,32] These multiple actions of laser therapy may represent the underlying mechanisms involved in the control the pain in MSD. In addition, a decrease in pain sensation has a significant effect on the increase of range of motion and the quality of life of the patient.[22] Thus, functional ability in patients with MSD could also be improved.

The present study has several strengths. First, to the best of our knowledge, this study is the first systematic review examining the effectiveness of HILT for MSD. Thus, our results present the best available evidence for pain and disability recovery following HILT in patients with MSD including back, neck, shoulder, arm, and hand pain. Second, the ROB of the included studies was not high on quality assessment. Therefore, the results of this systematic review may be considered reliable because of the moderate quality of the included studies. Third, the effectiveness of HILT was present not only for MSD overall but also for different treatment regions, thus allowing a comprehensive evaluation of the effects of HILT.

The study has some limitations that should be considered while interpreting the results. First, some heterogeneity identified in the pain and disability scores, thus we performed the meta-analysis using a random effects model. Subgroup analyses were conducted, which allowed a decrease in the heterogeneity of some subgroups as in case of comparing placebo with treatment of the back pain. Second, there may be a possibility of a publication bias. The results from the contoured funnel plot produced an asymmetry. However, the contoured funnel plot
showed the missing data in both significant and not significant regions and the asymmetry of the contoured funnel plot could not completely explain the publication bias. Third, the detailed information relative to the application of the laser could not be presented because of the diversity of lasers, such as type of laser or application method, used in the included studies. Finally, all of the included studies were RCTs with short-term follow-up to elucidate the clinical effectiveness of HILT for pain management. However, the present study is meaningful as it presents the effectiveness of HILT for treatment of MSD.

5. Conclusions

The results of this study showed that HILT treatment for back and neck pain significantly improved pain and functional disability compared to controls. The ROB of the included studies was moderate; however, significant heterogeneity existed. Thus, in the future, larger, well-designed studies are warranted to assess different laser applications, treatment regions, and comparators.

Author contributions

Contributions: Conception and design: Hyun-Ju Seo and Hyun Jin Song; Literature search: Hyun-Ju Seo, Hyun Jin Song, Youngjin Lee, and Sung Kyu Kim; Quality assessment: Hyun-Ju Seo, Hyun Jin Song, Youngjin Lee, and Sung Kyu Kim; Data extraction: Hyun-Ju Seo, Hyun Jin Song, Youngjin Lee, and Sung Kyu Kim; Analysis and interpretation of data: Hyun-Ju Seo and Hyun Jin Song; Drafting of article: Hyun Jin Song; Obtaining of funding: Hyun-Ju Seo.

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