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REVIEW

Effects of low-level laser therapy on pain in patients with musculoskeletal disorders: a systematic review and meta-analysis

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ABSTRACT

INTRODUCTION: This meta-analysis investigated the effectiveness of low-level laser therapy (LLLT) on pain in adult patients with musculoskeletal disorders.

EVIDENCE ACQUISITION: A systematic literature search was conducted in the Medline and PEDro databases. Two researchers independently screened titles and abstracts of the retrieved studies for eligibility. Quality assessment of the eligible studies was conducted using the PEDro rating scale. Studies that scored \geq 4 were included. A random-effects model was used for this meta-analysis. Subgroup meta-analyses were conducted to evaluate the influence of the adherence of the applied LLLT to the World Association of Laser Therapy (WALT) guidelines, the anatomical site under investigation and the study design on the overall weighted mean effect size. Meta regression was used to assess the possible influence of the study quality on the individual study effect sizes.

EVIDENCE SYNTHESIS: Eighteen studies allowing for 21 head-to-head comparisons (totaling N.=1462 participants) were included. The pooled raw mean difference (*D*) in pain between LLLT and the control groups was -0.85 (95% CI: -1.22 to -0.48). There was high (I^2 =85.6%) and significant between study heterogeneity (Cochran's Q = 139.2; df=20; P<0.001). The subgroup meta-analysis of the comparisons not following the WALT guidelines revealed a D=-0.68 (95% CI: -1.09 to -0.27). In this group, heterogeneity decreased to I^2 =72.6% (Q=51.2; df=14; P<0.001). In the WALT subgroup D equaled -1.52 (95% CI: -2.34 to -0.70). This between groups difference was clinically relevant although statistically not significant (Q=3.24; df=1; P=0.072).

CONCLUSIONS: This meta-analysis presents evidence that LLLT is an effective treatment modality to reduce pain in adult patients with musculoskeletal disorders. Adherence to WALT dosage recommendations seems to enhance treatment effectiveness.

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Introduction

In musculoskeletal rehabilitation, low-level laser therapy (LLLT) is frequently used as an adjunct in the management of pain in patients with musculoskeletal disorders.^{1, 2}

LLLT refers to a non-invasive, phototherapy or pho-

tobiomodulation that uses photons at a non-thermal irradiance to stimulate biological activity and has been classified as a safe, non-invasive treatment modality.³

Indeed, several possible mechanisms have been attributed to LLLT such as: increased endogenous opioid neurotransmitter production,⁴ raised threshold to thermal pain and enhanced local blood circulation,^{5, 6} in-



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Group by	Study name		Statistics for each study					Difference in means and 55% Ci					
WOLT		Difference in means	Standard error	Variance	Lower limit	Upper limit	Z/Value	p-Value					
no-WALT	Abrisham et al. (2011)	-0.900	0.177	0.081	-1.248	-0.554	-5.091	0.000	1	I -•	- 1		1
no-WALT	Al Restoud et al. (2013)	-0.600	D,632	0,400	-1,839	0,639	-0.949	0,343	- 1				
no-WALT	Alfredo et al. (2012)	-0.210	0.953	0.908	-2.077	1,657	-0.220	0.826	- 1		-	_ 1	
no-WALT	Ay et al. (2010 acute)	0.700	0.453	0,205	-0.187	1,587	1.547	0,122	- 1		-		
TJAW-on	Av et al. (2010 chronic)	0.000	0.455	0,207	-0.893	0.883	0.000	1,000	- 1		_	-	
no-WALT	Dogen et al. (2010)	-0.870	0.492	0.242	-1.834	0.094	-1.768	0.077	- 1				
no-WALT	Jiang et al. (2011 mild CTS)	-2.2890	D, 327	0,107	-2,900	-1,620	-6,921	0,000				1	
no-WOALT	Jiang et al. (2011 moderate C	TS) -1.100	0.515	0.265	-2,170	-0.150	-2.252	0.024				1	
no-WALT	Rheahle et al. (2014)	-0.980	0.275	0,075	-1,498	-0,422	-3,494	0,000	- 1	_ I →	- 1		
no-WALT	Meireles et al. (2010)	0.990	0.602	0,363	-0,191	2,171	1,644	0,100	- 1		-		
no-WALT	Santos et al. (2012)	-1.000	0.585	0.342	-2.147	0.147	-1.709	0.087	- 1		-		
no-WALT	Tascioglu et al. (2012 - 1.5.J/p	oint) -0.750	D.589	0,347	-1,905	0,405	-1,273	0,203	- 1		-		
no-WALT	Tascioglu et al. (2012 - 3J/poir	nt) -0.500	0.725	0.525	-1,920	0.920	-0.690	0,490	- 1			-	
no-WALT	Valione et al. (2014)	-1,400	0,276	0,078	-1,940	-0,860	-5,080	0,000	- 1			1	
no-WALT	Yeldan et al. (2009)	-0.050	0,503	0,253	-1,005	0,935	-0.099	0,921	- 1	- 1	-	-	
no-WALT		-0.680	0.210	0.044	-1.091	-0.268	-3.235	0.001	- 1	I ◄			
WALT	Erranet et al. (2010)	-0.850	D, 36T	0,135	-1,588	-0,131	-2,317	0,021	- 1		Ě-I	1	
WALT	Fusekul et al. (2014)	-0.330	1,301	1,683	-2.880	2,220	-0.254	0.800	- 1			<u> </u>	
WALT	Kittei et al. (2010)	-2.200	D,874	0,784	-3,913	-0,487	-2,517	0,012		_	-		
WALT	Konstantinovic et al. (2010a)	-0.610	0,257	0,066	-1,113	-0,107	-2,375	0,018		_ I -		1	
WALT	Konstantinovic et al. (2010b)	-1,428	0,067	0,004	-1,558	-1,298	-21,454	0.000			_	1	
WALT	Maliaropoulos et al. (2013)	-3,181	0,252	0,064	-3,675	-2,687	-12,617	0,000	_ i –∎			1	
WALT		-1.520	0.417	0,174	-2,338	-0.702	-3,642	0.000	1		- 1	1	1
Overall		-0.850	D, 188	0,035	-1,217	-0,482	-4,527	0,000				1	
									-4,00	-2,00	0,00	2,00	4,00
										Pavona LULT		Favora Control	

Figure 1.—Forest plot of the 18 trials (21 head-to-head comparisons) evaluating the effects of LLLT on pain versus control in patients with musculoskeletal disorders and subgroup analysis of adherence to WALT guidelines.

creased oxygen consumption by accelerating the redox reaction rate of the electron respiratory chain of mitochondria,⁷ increased adenosine triphosphate (ATP) production at the cellular level,⁸⁻¹⁰ increased production of anti-inflammatory cytokines.¹¹⁻¹³

Although LLLT is used in a variety of clinical settings, controversial results on its effectiveness in the treatment of pain in patients with musculoskeletal disorders have been reported.¹⁴⁻¹⁷

These conflicting results can be explained by the following facts: 1) the underlying cellular photobiostimulating mechanisms of LLLT are not well understood with as a consequence a largely empirical use and 2) the complexity of the appropriated parameter selection before each treatment session.^{3, 18} Therefore, an essential factor for the effective administration of LLLT is the certainty of optimal dosing to reach a sufficient volume of pathological target tissue.¹⁹ Although the World Association of Laser Therapy (WALT) introduced evidence based dosage recommendations for optimal administration of LLLT in the treatment of musculoskeletal pain, there are still RCT studies published without applying the WALT recommendations in their treatment protocol.^{14-17, 20-27} This can lead to low treatment efficacy (Figure 1).^{17, 24}

Evidence acquisition

This study was performed following the guidelines on the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) statement. Inclusion and exclusion criteria were set *a priori*. Eligible for inclusion were clinical trials, RCTs, reviews, meta-analyses, practice guidelines, studies on adult humans, published during the past five years in the English or German language. Only studies comparing LLLT *versus* a sham/placebo LLLT or studies comparing usual therapy + LLLT *versus* usual therapy were selected. Studies on the use of LLLT in the context of mandibular joint disorders were excluded. VAS was used to quantify pain in all studies.^{28, 29}

Outcomes

Within the context of evidence based practice this systematic review and meta-analysis aimed to answer the following questions:

— Is LLLT effective in treatment of pain in patients with musculoskeletal disorders?

— What is the effect of implementing the WALT dosage recommendations on the overall effect size?

— Is the pain relieving effect of LLLT affected by the anatomical site of the lesion?

— Does the study design or methodological study quality influence the individual effect size?

Data sources and search strategies

An electronic search was conducted in the MED-LINE (PubMed) and PEDro (Physiotherapy Evidence Database) databases with a latest update on 11.11.2015. Based on the PICO acronym, the following search algorithm was developed to evaluate the effects of LLLT in patients with musculoskeletal problems:

((((("musculoskeletal diseases" [MeSH Terms] AND "low-level light therapy" [MeSH Terms] OR ("lowlevel light therapy" [MeSH Terms] OR ("low-level" [All Fields] AND "light" [All Fields] AND "therapy" [All Fields]) OR "low-level light therapy" [All Fields] OR "Illt" [All Fields])) OR (Low-power [All Fields] AND ("lasers" [MeSH Terms] OR "lasers" [All Fields] OR "laser" [All Fields]))) OR (Low-intensity [All Fields] AND ("lasers" [MeSH Terms] OR "lasers" [All Fields] OR "laser" [All Fields]))) OR (low-laser [All Fields] AND ("therapy" [Subheading] OR "therapy" [All Fields] OR "therapeutics" [MeSH Terms] OR "therapeutics" [All Fields]))) AND ("placebos" [MeSH Terms] OR "placebos" [All Fields] OR "placebo" [All Fields])) NOT ("temporomandibular joint" [MeSH Terms] OR ("temporomandibular" [All Fields] AND "joint" [All Fields]) OR "temporomandibular joint" [All Fields] OR "tmj" [All Fields]) AND (Clinical Trial [ptyp] AND hasabstract[text] AND "2011/07/01"[PDat]: "2016/06/28" [PDat] AND "humans" [MeSH Terms]).

Manual searching and searching conference books of abstracts was not conducted. Pain was the outcome of interest in this study. In case of incomplete data reporting, the corresponding author of a study was contacted to obtain the missing data. A trial would be excluded from the meta-analysis if authors did not react to the request.

Study selection

Two researchers (AB and RC) independently screened titles and abstracts of the retrieved studies for their eligibility. Agreement was achieved by consensus. The reference lists of interpretation of the results. The 95% confidence intervals [95% CI] for the individual study effect sizes as well as the overall weighted mean were calculated.

Mixed effects subgroup analyses were conducted to evaluate the influence of covariates, such as the adherence of the applied LLLT to the WALT dosage guidelines, anatomical site under investigation and the study design. Meta regression was used to assess the possible influence of the study quality on the individual study effect sizes.

Statistical analysis

The Cochran's *Q* statistic and its corresponding P value were calculated to test the hypothesis that there was no heterogeneity across the individual effect sizes. *I*² was calculated to assess the degree of heterogeneity across studies. Higgins' suggested bench marking values were applied for the interpretation of the observed heterogeneity. Publication bias was assessed using visual analysis of the funnel plot and by formal testing for funnel plot asymmetry using the "trim and fill" and the "fail 'n safe" algorithms. For all analyses, P values less than 0.05 were considered significant. All calculations and plots were conducted using the CMA-2 software (Comprehensive Meta-Analysis 2nd version, Biostat, Englewood, NJ, USA).

Evidence synthesis

Study characteristics

Our search resulted in the identification of 124 potentially relevant studies. Three studies were suggested by experts and added in the further processing. After removing duplicates, the initial search yielded 94 articles which were screened on title, abstract and full-text. A total of 19 studies fulfilled the *a priori* set inclusion criteria (Figure 2).^{14-17, 20-26, 30-35} From the total of N.=1462 participants, N.=768 were in the LLLT group and N.=694 in the control group. Gender distribution was reported in 19 comparisons (overall females: N.=848; males: N.=528) while this information could not be revealed from one study.¹⁴

In five of the 19 studies, the reviewers independently agreed on all the items of the inclusion and exclusion criteria. One study ³⁶ showed important methodological limitations (PEDro score =2) and, therefore, was excluded from the further analysis.

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Figure 2.—Flow chart of the study selection process.

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Thus, 18 studies with a PEDro score ranging from 5 to 10 remained for the quantitative analysis. Three studies showed to be more-armed studies.^{14, 17, 26} The arms were included as separate head-to-head comparisons, totaling the number of comparisons in the meta-analysis to 21 (Table I).^{14-17, 20-27, 30-35}

Comparison 1: what is the effect of low-level laser therapy on pain compared to control in patients with musculoskeletal disorders?

All 21 comparisons analyzed the effect of LLLT on pain in patients with musculoskeletal disorders (Table I). The results were extracted from the studies and were analyzed using the random-effects model because of the expected high heterogeneity between studies. The overall weighted raw mean difference (*D*) in pain between LLLT and the control groups was 0.85 [95% CI: -1.22 to -0.48] (P<0.001). Heterogeneity analysis showed high (I^2 = 85.6%) and significant between study heterogeneity (Cochran's *Q*=139.2; df=20; P<0.001).

Author/year	Diagnosis	N.	Gender distribution	Exp./contr.	Intervention
Abrisham et al. (2011) ²⁰	Subacromial syndrome	80	30 males/50 females	40/40	LLLT and exercise vs. placebo LLLT and exercise
Al Rashoud <i>et al.</i> (2013) ²¹	Osteoarthritis knee	49	31 males/18 females	26/23	LLLT and exercise vs. placebo LLLT and exercise
Alfredo <i>et al.</i> (2011) ²²	Osteoarthritis knee	40	9 males/31 females	20/20	LLLT and exercise vs. placebo LLLT and exercise
Ay et al. (2010) 17 (acute)	Acute low back pain	40	14 males/26 females	20/20	LLLT and hot-pack vs. placebo LLLT and hot-pack
Ay et al. (2010) 17 (chronic)	Chronic low back pain	40	20 males/20 females	20/20	LLLT and hot-pack vs. placebo LLLT and hot-pack
Dogan <i>et al.</i> (2010) ¹⁶	Subacromial impingement	52	19 males/33 females	30/22	LLLT and cold-pack and exercise vs. placebo LLLT and cold-pack and exercise
Emanet et al. (2010) 30	Lateral epicondylitis	46	13 males/33 females	23 /23	LLLT and exercise vs. placebo LLLT and exercise
Fusakul <i>et al.</i> (2014) ³¹	Carpal tunnel	112	58 males/54 females	56/56	LLLT and neutral wrist splint <i>vs.</i> placebo LLLT and neutral wrist splints
Jiang <i>et al.</i> (2011) ¹⁴ (moderate CTS)	Carpal tunnel syndrome	33	NM	18/15	LLLT vs. placebo LLLT
Kheshie <i>et al.</i> (2014) ²³	Osteoarthritis knee	53	53 males/0 females	38 /15	High-intensity laser therapy and exercise vs. LLLT and exercise vs. placebo LLLT and exercise
Kiritsi et al. (2010) ³²	Plantar fasciitis	25	10 males/15 females	15/10	LLLT vs. placebo LLLT
Konstantinovic <i>et al</i> $(2010)^{33}$	Acute neck pain	60	25 males/35 females	30/30	LLLT vs. placebo LLLT
Konstantinovic <i>et al.</i> $(2010)^{34}$	Low back pain with radiculopathy	546	231 males/315 females	182/182	LLLT and NSAID vs. NSAID vs.placebo LLLT and NSAID
Malliaropoulos et al. (2013) ³⁵	Meniscal pathology	64	20 males/44 females	32/32	LLLT vs. placebo LLLT
Meireles <i>et al.</i> (2010) ²⁴	Rheumatoid arthritis	78	2 males/76 females	41/37	LLLT and NSAID vs. placebo LLLT and NSAID
Santos et al. (2012) 25	Episiotomy	52	0 males/52 females	26/26	LLLT vs. placebo LLLT
Tascioglu <i>et al.</i> (2012) ²⁶ (1.5 J/ point) and (3.0 J/point)	Carpal tunnel syndrome	60	14 males/46 females	40/20	LLLT 1.5 J vs. LLLT 3.0 J vs. placebo LLLT
Vallone et al. (2014) ¹⁵	Nonspecific chronic low back pain	100	43 males/57 females	50/50	LLLT and exercise vs. exercise
Yeldan <i>et al.</i> (2009) ²⁷	Subacromial impingement	60	13 males/47 females	34/26	LLLT and cold-pack and exercise <i>vs.</i> placebo LLLT and cold-pack and exercise

NSAID: nonsteroidal anti-inflammatory drugs; LLLT: low-level laser therapy; VAS: visual analog scale.

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Despite the observed inconsistency in the effect size of LLLT on pain, the present meta-analysis presents good evidence for the use of LLLT in the treatment of pain in adult patients with musculoskeletal disorders. From the 21 head-to-head comparisons, 17 favored LLLT while four comparisons (extracted from three studies) reported no beneficial effects of LLLT on pain (Figure 1).

Figure 3 depicts the funnel plot of standard error by D. The classic "fail-safe N" algorithm showed that 1179 non-significant studies would be needed to increase the P value above the set alpha level of 0.05, indicating that there was but very low risk for publication bias.

Comparison 2: does implementing the WALT dosage recommendations affects the overall effect size?

Six of the analyzed studies followed the 2005 published WALT guidelines for the LLLT intervention.¹⁹ To test if adherence to WALT guidelines had an effect on the overall weighted raw mean difference a subgroup meta-analysis was conducted. Subgroup meta-analysis



Figure 3.—Funnel plot of the included studies.

showed no significant relationship between the positive pain relieving effects and the use of WALT treatment dosage recommendations. Interestingly, only six studies (Table I) implemented the WALT dosage recommendations whilst a large variety in reported dose and

Outcome parameter	PEDro Score	WALT dosage recommendations
VAS for pain, ROM	9/10	No
VAS for pain, ROM	6/10	No
VAS for pain, ROM, muscle strength, Lequesne for functionality, WOMAC questionnaire for Activity	8/10	No
VAS and Likert scale for pain, ROM, Roland Disability Questionnaire and Modified Oswestry Disability Questionnaire for function	8/10	No
VAS and Likert scale for pain, ROM Roland Disability Questionnaire and Modified Oswestry Disability Questionnaire for function	7/10	No
VAS for pain, ROM, Shoulder Pain and Disability Index for functional status	9/10	No
VAS for pain, tenderness (pressure algometry), Painless grip strength (dynamometry)	6/10	Yes
VAS for pain, Symptom Severity Scale (SSS), Functional Status Scale (FSS), pinch strength, grip strength	8/10	Yes
VAS for pain, Boston Questionnaire scale for discomfort symptoms of CTS, Phalen's maneuver and Tinel's sign test for neurological signs of CTS, NCS	7/10	No
VAS for pain, WOMAC Scale for knee joint function	7/10	No
VAS for pain, ultrasonography for plantar fascia thickness	7/10	Yes
VAS for pain, neck disability index for neck mobility, SF-12 questionnaire health survey	10/10	Yes
VAS for pain, modified Schober Test for lumbar mobility, Oswestry disability scale for daily activities, SF-12 questionnaire health survey	10/10	Yes
VAS for pain, Lysholm Knee Scoring System for knee function, pain and swelling	9/10	Yes
VAS for pain, HAQ (Health Assessment Questionnaire) and DASH questionnaire (Disabilities of the arm shoulder and hand)	10/10	No
VAS for pain, REEDA Scale for healing process	8/10	No
VAS for pain, SSS, FSS, grip strength, nerve conduction studies, Ultrasonography evaluation	7/10	No
VAS for pain	5/10	No
VAS for pain, DASH questionnaire, Shoulder Disability Questionnaire (SDQ), Dynamo-metry for muscle strength, ROM	7/10	No

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beam parameter was used. The subgroup meta-analysis of the 15 head-to-head comparisons described in the studies which did not follow the WALT guidelines revealed a mean change in VAS of D=-0.68 [95% CI: -1.09 to -0.27]. In this group, heterogeneity decreased to I^2 =72.6% (Q=51.2; df=14; P<0.001). In the WALT subgroup, the mean change in VAS equaled D=-1.52 [95% CI: -2.34 to -0.70]. Under random-effects conditions, the between groups difference was statistically not significant at the 5% level (Q=3.24; df=1; P=0.072).

Comparison 3: is the pain relieving effect of LLLT affected by the anatomical site of the lesion?

In the 21 head-to-head comparisons included in the 18 studies, the effect of LLLT on pain in patients with musculoskeletal disorders was investigated at nine different anatomical sites: back (k=4), elbow (k=1), foot (k=1), hand (k=1), knee (k=4), neck (k=1), perineal (k=1), shoulder (k=3), wrist (k=5). To test if LLLT had different effects on pain at the different anatomical sites another subgroup meta-analysis was conducted. For the subgroups including more than one study per anatomical site, LLLT had the strongest effect on pain in patients with knee disorders with D=-1.34 [95% CI: -2.88 to 0.20], followed by wrist disorders with D=-1.22[95% CI: -2.05 to -0.39], should disorders with D=-0.76 [95% CI: -1.19 to -0.33] and back disorders with D=-0.63 [95% CI: -1.48 to 0.23]. Under random-effects conditions, the between groups difference was statistically not significant at the 5% level (Q=13.51; df=8; P=0.096).

Comparison 4: does the methodological study quality influence the individual effect size?

A subgroup meta-analysis comparing RCT versus CT studies was conducted. The RCT studies yielded an overall weighted raw mean difference of D=-0.82 [95% CI: -1.23 to -0.40] while the overall weighted effect size in the CT subgroup was D=-1.45 [95% CI: -2.40 to -0.51]. Again, the between groups difference was statistically not significant at the 5% level (Q=1.45; df=1; P=0.228).

To test for an eventual effect of the study quality on the effect size, individual studies effect-sizes were meta-regressed over their PEDro score which yielded a slope estimate of -0.086 [95% CI: -0.16 to -0.01].

Discussion

This systematic review and meta-analysis of 21 head-to-head comparisons extracted from 18 studies (totaling N.=1462 participants) was conducted to assess the available clinical evidence for the use of LLLT in the treatment of pain in adult patients with musculoskeletal disorders. The secondary objectives were to determine if the study outcome was affected by the adherence to the WALT dosage recommendations, if the pain relieving effect of LLLT was related to the anatomical site of the affected structure, and finally if the observed effect size was influenced by study design or study quality.

In the included studies a large variety in reported dose and beam parameter was used, this observed heterogeneity is in line with the findings of Jenkins *et al.* who stated that LLLT effectiveness studies frequently lack in accurate and complete reporting of technical and treatment parameters and that there is a need for more standardized reporting of these parameters.³⁷ Standardized reporting of beam and treatment parameters and the adherence to the evidence based WALT guidelines will significantly enhance the reproducibility and the body of knowledge on clinical application of LLLT.

Although the between group difference of the effects of adherence to the WALT guidelines did not reach statistical significance, this difference seems to be of important clinical relevance. Several authors have investigated the clinical effectiveness of VAS score reduction by defining the minimum clinically important difference (MCID) on the VAS pain score for a treatment intervention. Todd et al. stated that a VAS reduction of 13 mm was perceived as clinically relevant in patients with acute trauma pain, while Gallagher et al. concluded an MCID of 16 mm to be of clinical relevance in patients with acute abdominal pain.^{38, 39} In the present meta-analysis, a clinical relevant difference of 15.2 mm was found in the LLLT interventions following WALT guidelines. The absence of between groups significance could be the result of the low number of included studies and study subjects.

The studies investigating the effect of LLLT treatment on pain in adult patients with musculoskeletal disorders showed a high variety of anatomical treatment sites. The present meta-analysis suggests that the beneficial effects of LLLT on pain seem to be independent from the anatomical lesion site as the analysis of the between

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group difference reached no statistical significance.

To see if the overall weighted mean effect was affected by the study type, a subgroup meta-analysis comparing RCT *versus* CT studies was conducted, yielding no significant difference between the two study types. Despite the methodological flaws in reporting of technical and treatment parameters, the methodological quality spectrum of the included studies ranged from PEDro score 5 to 10 which can be interpreted as moderate to good methodological quality. The regression of the PE-Dro score on the study effect size reached no significance indicating that the conflicting evidence regarding the effectiveness of LLLT in the treatment of pain in patients with musculoskeletal disorders can only be partially explained by the methodological quality of the studies.

This meta-analysis suggests that remaining strictly to WALT guidelines during treatment may affect the clinical pain relieving outcome. Hence, therapists applying LLLT for the pain relief treatment of patients with musculoskeletal disorders, should prefer the use of evidence based treatment strategies and WALT dosage recommendations to optimize treatment effect. Future studies evaluating the effect of LLLT in the treatment of patients with musculoskeletal disorders should be conducted using standardized beam and treatment parameters to enhance reproducibility and the body of knowledge on the clinical application of LLLT.

Strengths and limitations of the study

A strength of the present study is the systematic review of the literature yielding an important number of clinical trials and randomized clinical trials of moderate to high methodological quality, all assessing pain on the same scale. This allowed for a quantitative analysis by pooling the individual study effect sizes expressed in their original units (*i.e.* mm on VAS) facilitating the interpretation of the results for the clinician. Furthermore, an analysis of the influence of covariates such as adherence to the WALT dosage recommendations and anatomical sites on the overall weighted effect size was conducted, providing information with important clinical relevance.

Limitations that may hamper the outcome of this study should be mentioned also. In the fast technical

developing field of LLLT, the authors choose to provide an actual status of the evidences for LLLT including only studies of the last five years. We acknowledge that this is another limitation of this study. Beside Medline only one specific physiotherapy database (PEDro) was searched while a gray literature search was omitted. Despite this limitation, the meta-analysis showed but very low risk for publication bias.

Conclusions

Based on the results of this study, LLLT appears to be an effective treatment modality to achieve pain relief in adult patients with musculoskeletal disorders. Therapists applying LLLT should follow the WALT dosage recommendations to yield clinically significant better pain relieving effects when treating patients with musculoskeletal disorders. Although the included studies showed a high heterogeneity in anatomical treatments sites, the beneficial effect of LLLT on pain seem to be unaffected by the anatomical site of the lesion.

References

- 1. Jamtvedt G, Dahm KT, Christie A, Moe RH, Haavardsholm E, Holm I, *et al.* Physical therapy interventions for patients with osteoarthritis of the knee: an overview of systematic reviews. Phys Ther 2008;88:123-36.
- 2. Watson T. The role of electrotherapy in contemporary physiotherapy practice. Man Ther 2000;5:132-41.
- 3. Chung H, Dai T, Sharma SK, Huang YY, Carroll JD, Hamblin MR. The nuts and bolts of low-level laser (light) therapy. Ann Biomed Eng 2012;40:516-33.
- Hagiwara S, Iwasaka H, Hasegawa A, Noguchi T. Pre-Irradiation of blood by gallium aluminum arsenide (830 nm) low-level laser enhances peripheral endogenous opioid analgesia in rats. Anesth Analg 2008;107:1058-63.
- Schindl A, Schindl M, Schon H, Knobler R, Havelec L, Schindl L. Low-intensity laser irradiation improves skin circulation in patients with diabetic microangiopathy. Diabetes Care 1998;21:580-4.
 Podogrodzki J, Lebiedowski M, Szalecki M, Kepa I, Syczewska M,
- Podogrodzki J, Lebiedowski M, Szalecki M, Kepa I, Syczewska M, Jozwiak S. [Impact of low level laser therapy on skin blood flow]. Dev Period Med 2016;20:40-6.
- 7. Yu W, Naim JO, McGowan M, Ippolito K, Lanzafame RJ. Photomodulation of oxidative metabolism and electron chain enzymes in rat liver mitochondria. Photochem Photobiol 1997;66:866-71.
- Benedicenti S, Pepe IM, Angiero F, Benedicenti A. Intracellular ATP level increases in lymphocytes irradiated with infrared laser light of wavelength 904 nm. Photomed Laser Surg 2008;26:451-3.
- 9. Karu T. Photobiology of low-power laser effects. Health Phys 1989;56:691-704.
- Ferraresi C, de Sousa MV, Huang YY, Bagnato VS, Parizotto NA, Hamblin MR. Time response of increases in ATP and muscle resistance to fatigue after low-level laser (light) therapy (LLLT) in mice. Lasers Med Sci 2015;30:1259-67.

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- 11. Bjordal JM, Lopes-Martins RA, Iversen VV. A randomised, placebo controlled trial of low level laser therapy for activated Achilles tendinitis with microdialysis measurement of peritendinous prostaglandin E2 concentrations. Br J Sports Med 2006;40:76-80; discussion 76-80
- 12. Mizutani K, Musya Y, Wakae K, Kobayashi T, Tobe M, Taira K, et *al.* A clinical study on serum prostaglandin E2 with low-level laser therapy. Photomed Laser Surg 2004;22:537-9.
- 13. Houreld NN, Sekhejane PR, Abrahamse H. Irradiation at 830 nm stimulates nitric oxide production and inhibits pro-inflammatory cytokines in diabetic wounded fibroblast cells. Lasers Surg Med 2010;42:494-502
- 14. Jiang JC, Wu JH. Low- level laser therapy treatment relieves pain and neurological symptoms in patients with carpal tunnel syndrome. J Phys Ther Sci 2011;23:661-5.
- 15. Vallone F, Benedicenti S, Sorrenti E, Schiavetti I, Angiero F. Effect of diode laser in the treatment of patients with nonspecific chronic low back pain: a randomized controlled trial. Photomed Laser Surg 2014;32:490-4
- 16. Dogan SK, Ay S, Evcik D. The effectiveness of low laser therapy in subacromial impingement syndrome: a randomized placebo controlled double-blind prospective study. Clinics (Sao Paulo) 2010:65:1019-22
- 17. Ay S, Dogan SK, Evcik D. Is low-level laser therapy effective in acute or chronic low back pain? Clin Rheumatol 2010;29:905-10.
- 18 Chipchase L. Is there a future for electrophysical agents in musculoskeletal physiotherapy? Man Ther 2012;17:265-6.
- 19. Bjordal JM. Low level laser therapy (LLLT) and World Association for Laser Therapy (WALT) dosage recommendations. Photomed Laser Surg 2012;30:61-2.
- 20 Abrisham SM, Kermani-Alghoraishi M, Ghahramani R, Jabbari L Jomeh H, Zare M. Additive effects of low-level laser therapy with exercise on subacromial syndrome: a randomised, double-blind, controlled trial. Clin Rheumatol 2011;30:1341-6.
- Al Rashoud AS, Abboud RJ, Wang W, Wigderowitz C. Efficacy of low-level laser therapy applied at acupuncture points in knee osteoar-21 thritis: a randomised double-blind comparative trial. Physiotherapy 2014;100:242-8.
- 22. Alfredo PP, Bjordal JM, Dreyer SH, Meneses SR, Zaguetti G, Ovanessian V, et al. Efficacy of low level laser therapy associated with exercises in knee osteoarthritis: a randomized double-blind study. Clin Rehabil 2012;26:523-33
- Kheshie AR, Alayat MS, Ali MM. High-intensity versus low-level 23 laser therapy in the treatment of patients with knee osteoarthritis: a randomized controlled trial. Lasers Med Sci 2014;29:1371-6.
- Meireles SM, Jones A, Jennings F, Suda AL, Parizotto NA, Natour J. Assessment of the effectiveness of low-level laser therapy on the hands of patients with rheumatoid arthritis: a randomized doubleblind controlled trial. Clin Rheumatol 2010;29:501-9
- 25. Santos Jde O, de Oliveira SM, da Silva FM, Nobre MR, Osava RH,

Riesco ML. Low-level laser therapy for pain relief after episiotomy: a double-blind randomised clinical trial. J Clin Nurs 2012;21:3513-22

- Tascioglu F, Degirmenci NA, Ozkan S, Mehmetoglu O. Low-level 26 laser in the treatment of carpal tunnel syndrome: clinical, electrophysiological, and ultrasonographical evaluation. Rheumatol Int 2012;32:409-15.
- Yeldan I. Cetin E. Ozdincler AR. The effectiveness of low-level laser 27 berapy on shoulder function in subacromial impingement syndrome. Disabil Rehabil 2009;31:935-40.
- Bijur PE, Silver W, Gallagher EJ. Reliability of the visual analog 28 scale for measurement of acute pain. Acad Emerg Med 2001;8:1153-
- Breivik H, Borchgrevink PC, Allen SM, Rosseland LA, Romundstad 29 L, Hals EK, et al. Assessment of pain. Br J Anaesth 2008;101:17-24
- 30 Emanet SK, Altan LI, Yurtkuran M. Investigation of the effect of GaAs laser therapy on lateral epicondylitis. Photomed Laser Surg 2010.28.397-403
- Fusakul Y, Aranyavalai T, Saensri P, Thiengwittayaporn S. Low-31. level laser therapy with a wrist splint to treat carpal tunnel syn-drome: a double-blinded randomized controlled trial. Lasers Med Sci 2014.29.1279-87
- 32. Kiritsi O, Tsitas K, Malliaropoulos N, Mikroulis G. Ultrasonographic evaluation of plantar fasciitis after low-level laser therapy: results of a double-blind, randomized, placebo-controlled trial. Lasers Med Sci 2010;25:275-81
- Konstantinovic LM, Cutovic MR, Milovanovic AN, Jovic SJ, Dragin 33. AS, Letic M, et al. Low-level laser therapy for acute neck pain with radiculopathy: a double-blind placebo-controlled randomized study. Pain Med 2010-11-1169-78
- Konstantinovic LM, Kanjuh ZM, Milovanovic AN, Cutovic MR, 34 Djurovic AG, Savic VG, et al. Acute low back pain with radiculopathy: a double-blind, randomized, placebo-controlled study. Photomed Laser Surg 2010;28:553-60.
- Malliaropoulos N, Kiritsi O, Tsitas K, Christodoulou D, Akritidou 35. A, Del Buono A, et al. Low-level laser therapy in meniscal pathology: a double-blinded placebo-controlled trial. Lasers Med Sci 2013;28:1183-8.
- 36. Alves Mde P, de Araújo GC. Low-Level Laser Therapy after Carpal Tunnel Release. Rev Bras Ortop 2011;46:697-701.
- Jenkins PA, Carroll JD. How to report low-level laser therapy (LLLT)/ 37 photomedicine dose and beam parameters in clinical and laboratory studies. Photomed Laser Surg 2011;29:785-7
- 38 Todd KH, Funk JP. The minimum clinically important difference in physician-assigned visual analog pain scores. Acad Emerg Med 1996:3:142-6.
- 39. Gallagher EJ, Bijur PE, Latimer C, Silver W. Reliability and validity of a visual analog scale for acute abdominal pain in the ED. Am J Emerg Med 2002;20:287-90.

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