Title Page Inter-individual differences in the responses to pain neuroscience education in adults with chronic musculoskeletal pain: A systematic review and meta-analysis of randomised controlled trials. Short title: Individual differences in response to pain neuroscience education Authors: James A Watson BSc (Hons) 1 J.A.Watson@tees.ac.uk Cormac G Ryan BSc, MSc, PhD₁ C.Ryan@tees.ac.uk Greg Atkinson BSc (Hons), PhD₁ Greg.Atkinson@tees.ac.uk Philip Williamson, BSc (Hons), MSc, PhD3 phil.williamson@york.ac.uk Dominic Ellington BSc (Hons) 2 D.Ellington@tees.ac.uk

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Highlights:

- Pain neuroscience education (PNE) is a pain management intervention.
- Little evidence of true individual differences in response to PNE for disability.
- Findings should be interpreted cautiously due to very wide prediction intervals.
 - Estimating individual differences should be applied to other pain interventions.

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Abstract

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Pain neuroscience education (PNE) is an approach used in the management of chronic musculoskeletal pain (CMP). Previous reviews on PNE and other pain interventions, have focussed on mean treatment effects, but in the context of "precision medicine", any inter-individual differences in treatment response are also important to quantify. If inter-individual differences are present, and predictors identified, PNE could be tailored to certain people for optimising effectiveness. Such heterogeneity can be quantified using recently-formulated approaches for comparing the response variance between the treatment and control groups. Therefore, we

conducted a systematic review and meta-analysis on the extracted standard deviations of baseline-to-follow up change to quantify the inter-individual variation in pain, disability and psychosocial outcomes in response to PNE. Electronic databases were searched between 01/01/2002 and 14/06/2018. The review included five randomised controlled trials (n=428) in which disability outcomes were reported. Using a random effects meta-analysis, the pooled SD (95% CI) for control groupadjusted response heterogeneity to PNE was 7.36 units /100 (95% CI: -3.93 to 11.12). The 95% prediction interval for this response heterogeneity SD was wide (-10.20 to 14.57 units /100). The control group-adjusted proportion of "responders" in the population who would be estimated to exceed a clinically important change of 10/100 ranged from 18-45%. Therefore, when baseline-to-follow up random variability in disability is taken into account (informed by the control arm), there is currently insufficient evidence for the notion of clinically important inter-individual differences in disability responses to PNE in people with CMP. The protocol was published on PROSPERO (CRD42017068436).

Perspective

We bring a novel method to pain science for calculating inter-individual differences in response to a treatment. This is conducted within the context of a systematic review and meta-analysis on PNE. We highlight how using erroneous methods for calculating inter-individual differences can drastically change conclusions when compared to appropriate methods.

Key words

Pain, neuroscience, education, Individual response variance

Introduction

Pain neuroscience education (PNE) is an educational approach used in the management of chronic pain. PNE aims to reconceptualise an individuals' understanding of their pain as less threatening to facilitate rehabilitation₂₃. Since its inception PNE has become increasingly popular in clinical practice₂₄. Our group recently published a mixed-methods systematic review and meta-analysis on the effectiveness of PNE for adults with chronic musculoskeletal pain (CMP)₃₉. Quantitatively we found no evidence to indicate that PNE results in clinically important changes over control for pain or disability. In contrast we found moderate quality evidence that PNE produces small clinically important changes over control for pain catastrophising and kinesiophobia. Qualitatively we found that achieving some degree of pain reconceptualisation following PNE can enhance peoples' ability to cope with their condition.

One question that arose during our previous research work was whether PNE may be effective for some types of people, implying that there may be some individual differences in response to PNE₃₉. The quantitative component of our review focused on the mean intervention/treatment effect. This focus on mean intervention effect whilst common in research on pain interventions_{5,15,30} could have obscured important inter-individual differences in response to PNE_{16,41}. Such response heterogeneity is particularly important within the context of precision medicine, an increasingly popular field which encompasses 'tailor-made' therapies based on the person's individual response to a given intervention₃₁. This individualised approach to medicine aims to improve the quality of care and reduce costs₃₃. The potential

importance of a tailored approach has been highlighted by some of our previous qualitative work on PNE. The relevance of PNE to the individual (i.e. how tailored the material is to that individual) appears to be an important factor in the success of PNE_{17,18,29,39}. Where PNE was reported to be relevant, people reported greater perceived benefit. The opposite was found where PNE was deemed not relevant_{17,18,29}.

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Some researchers₂₇ have attempted to complement the quantification of mean treatment effects with a quantification of how many people in each intervention group change above or below a pre-set threshold, termed sample responder counts. Crucially, this approach does not provide any information about response heterogeneity to a given intervention in the context of precision medicine. In fact, these responder counts lack statistical power and may merely reflect within-subject random variation between timepoints and/or group differences in mean change. Furthermore, the dichotomisation (responder or non-responder) also creates problems adjusting for baseline differences between study groups (comprehensive reviews are available 2,32). These sample responder counts tell us little about whether different people respond to different degrees to the same intervention, which is one of the fundamental questions in precision medicine. Should any interindividual differences be falsely identified using the above-mentioned methods, any follow-up analysis to explore potential moderators of the intervention effect to explain the individual differences in response are therefore unwarranted_{1,2}. Subsequent follow-up studies on the same participants is a waste of resources, and potentially unethical, if no true inter-individual differences in response exist to explain.

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Inter-individual differences in response can be quantified by comparing the SDs of the baseline-to-follow-up changes between the experimental and control groups_{1,4}. The difference between these SDs represents the SD for individual responses (SDir) which quantifies the individual variability in treatment response *per se*. The SD of the mean change score solely for the intervention group comprises treatment response variance *in addition to* the random variability in measurements between the baseline and follow-up timepoints. The SD of the changes in the control group represents this random variability in measurements between baseline and follow up – the random within-subjects variance component and measurement error.

Our qualitative analysis highlighted that PNE may be effective for some people but not for others implying that true inter-individual differences in response to PNE may exist which could be explored to facilitate appropriate targeting of PNE to those most likely to benefit³⁹. However, clinically relevant inter-individual response variation should first be conducted using appropriate methodology^{1,2,13,40,41} to confirm the presence of such inter-individual responses. If individual differences are observed, and predictors of individual response are identified, then PNE could be tailored to the individual optimising its effect⁴¹.

To date, there has been no investigation of 'true' individual response variation of the effect of PNE, or indeed any pain management intervention. Therefore, we aimed to conduct a systematic review and meta-analysis of the available research to quantify the 'true' inter-individual variation in pain, disability and psychosocial outcomes in response to PNE in adults with CMP.

Methods

The protocol for the systematic review was published on PROSPERO (CRD42017068436). The analysis of inter-individual differences is presented here in detail to ensure the background and rationale for this novel method within the field of pain is adequately reported. A detailed account of the full review-methods has been published elsewhere 39 but a brief summary is provided below.

Inclusion and Exclusion Criteria

Inclusion criteria

Studies including adults (≥18 years) who have CMP consistent with the British
Pain Society definition (chronic pain, that lasts beyond the time that tissue
healing would normally be expected to have occurred, often taken as ≥3
months)₃₅.

 RCTs that (i) compared the intervention with no treatment (true control) or usual care (ii) concomitant studies where PNE was delivered in addition to another intervention where that other intervention was received by both groups and (iii) head-to-head studies where PNE was compared to another active intervention.

• Studies reporting either pain and/or disability and/or psychosocial wellbeing.

The SD of the changes for the intervention and control groups must have been included within the publication, have been available from the author upon request, or could be calculated from other information given such as the

201 standard error. This is an additional criterion that was not included in the 202 registered protocol. 203 204 Exclusion criteria 205 206 Studies that included participants with non-musculoskeletal pain such as cancer pain, visceral pain or post stroke pain. 207 208 209 210 Search Strategy 211 Pre-identified keywords (Pain AND (Physiology OR Neurophysiology OR 212 Neuroscience OR Biology) AND Education) and index terms were searched across 213 214 all included databases (The Cochrane Library, AMED, CINAHL Complete, MEDLINE, PsycINFO, PEDro, Scopus, EMBASE, Education Resources Information 215 Centre (ERIC), Web of Science, clinicaltrials.gov, dissertations indexed with 216 217 ProQuest Dissertations and Theses Global and EThOS) from 2002-25 July 2017, and updated on 14 June 2018. 218 219 220 After removing duplicates, the title and abstracts were screened by two authors and disagreements were resolved through discussion or a 3rd reviewer. The full-text was 221 222 obtained for all records that could potentially fit the criteria. Upon reading the fulltexts those deemed not to meet the inclusion criteria were rejected. See 223 224 Supplementary Digital Content 1 for a list of excluded publications and reasons for 225 exclusion.

Deviation from protocol

In our previous review39 when the SD of change was not reported, and could not be obtained by contacting the authors, it was either calculated from other information given such as standard error, or estimated from the baseline and follow up SDs, according to methods described in the Cochrane handbook10. Where there was uncertainty regarding the validity of baseline, follow up and change score SDs from included studies we opted not to use this data to inform our calculations to estimate the SD of change scores. Instead, we used a robust data set of individuals with CMP where we were confident in the validity of the baseline, follow up and change score SDs. However, for the current review, given that to calculate the true inter-individual differences in response to an intervention the SD of the mean change score is of central importance1, it would be inappropriate to estimate the SD of the change or use a robust data set. Thus, an additional criterion for inclusion was created for the current review where the SD of the changes for the intervention and control groups must have been published in the article, available upon request by the author, or could be calculated from other information given, such as the standard error.

Assessment of methodological quality and data extraction

Articles selected for critical appraisal were independently assessed by two reviewers using the Cochrane tool for assessing risk of bias₉. Two reviewers independently extracted the data using JBI-SUMARI₃₆ including details about the interventions,

populations, study methods and outcomes of relevance to the review question/objectives. The Grades of Recommendation, Assessment, Development and Evaluation (GRADE) approach was used to rate the overall quality of quantitative evidence for each outcome. A summary of findings table created using GradePro is presented (Table 1 and 2).

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Meta-analysis

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To contextualise the results for individual response variance we conducted a random-effects meta-analysis for the mean difference in disability across the included studies using a restricted maximum likelihood (REML) model combined with the Knapp-Hartung method. This method uses quantiles of the t distribution to calculate a confidence interval for the average effect instead of the standard normal distribution in the more conventional methods₃₇. The Knapp-Hartung method has been shown to be superior to the DerSimonian-Laird method where there is a small number of studies (<20) and heterogeneity is present 11. We then extracted the standard deviation of the changes in disability for both control (C) and PNE (I) groups. The true individual response variance (intervention minus control) was then calculated by $\sqrt{(SD_{12}-SD_{C2})_{13}}$. The standard error (SE) for this variance was then calculated using the equation: $SE = \sqrt{[2(SD_1^4/DF_1 + SD_c^4/DF_c)]}$, where DF₁ and DF₂ are the degrees of freedom of the standard deviation in the PNE group and the control groups 13. A negative value for the individual response variance for the confidence intervals or prediction intervals implies greater variability in the changes in disability in the control versus PNE group.

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The individual response variances and their SEs were meta-analysed using an REML model combined with Knapp-Hartung method. It's important to highlight that the variances are unbiased, whereas the SD is not, and deriving a SE for the SD for individual responses is also problematic. Thus, we synthesised the individual response variances instead of the SDs for individual responses. The point estimate for the pooled individual response variance were derived together with a 95% CI to express its uncertainty. The point estimate and CIs were then square rooted to convert to an SD metric. If the lower limit was negative, the sign was ignored, the square root taken, and the sign re-applied. This approach is consistent with the 'no bound' option in SAS/STAT® software, which permits negative variances (SAS Institute Inc.).

Using the methods of Swinton et al.34 the proportion of responders in the population of interest within each included RCT was estimated. To estimate this, the observed mean change score and true individual response variance are needed for each RCT. Normal variance is assumed. The total area of any probability distribution is equal to one, thus the estimate of the proportion of response can be obtained by calculating the area of the derived normal distribution that lies beyond the minimally clinically important difference (MCID). An MCID of 10% was used in recent NICE guidelines for back and radicular pain25. The calculation estimating the proportion of response was performed via an online calculator28. The proportion of response was estimated for the intervention and control groups for all RCTs and has been used to demonstrate the difference in results, and thus conclusion that could be made if researchers erroneously ignored the control group data.

The tau statistic (τ) was used to quantify between-study heterogeneity – a SD that describes the typical variability of the mean effect between studies_{3,8}. A 95% prediction interval was calculated using the tau and the SE for the pooled mean effect to quantify the expected range of true effects in future similar studies₁₂. Stata (StataCorp. 2019. Stata Statistical Software: Release 16. College Ststion, TX: StataCorp LLC.) was used to conduct all statistical analysis.

Results

Following removal of duplicates, 12,136 publications were identified (Fig. 1). Fifty-seven full text articles were screened. Forty-nine articles were excluded at this stage. See document, supplementary digital content 1 for a list of excluded publications and reasons for exclusion. Thus, six publications reporting five RCTs were included_{6,19,20,21,26,38}. The included studies encompassed a total of 428 participants (I = 212, C = 216). Table 3 provides further details regarding the studies.

Methodological quality

Quality scores ranged from 1-6 out of 7 (Table 4). There was a high risk of performance bias due to lack of blinding of participants and personnel (Fig. 2 and 3 produced by using RevMan software (Review Manager. Version 5.3. Copenhagen: The Nordic Cochrane Centre. The Cochrane Collaboration, 2014).

326 327 Study outcomes 328 329 Jackson and Turner14 recommend only pooling data where the number of studies is 330 ≥5 to ensure adequate statistical precision. Disability was the only outcome measured consistently in all five included studies, thus our analysis focused solely 331 332 on this outcome. 333 334 The pooled mean group difference in pre/post changes in disability (intervention minus control) was -2.26 units /100 (95% CI: -6.49 to 1.97). See Fig. 4. Between 335 336 study heterogeneity in mean treatment effect was observed (τ = 2.49; 95% CI: 0.48 337 to 4.51). The prediction interval revealed that, were investigators to undertake a future trial, the 95% plausible range for mean disability change versus control would 338 be -11.56 to 7.04 units /100. 339 340 The pooled point estimate for the inter-individual variability in disability change in 341 342 response to PNE (SDIR) was 7.36 units /100 (95% CI: -3.93 to 11.12). Substantial 343 between-study heterogeneity was observed (τ = 6.55). The 95% prediction interval 344 for true inter-individual responses was -10.20 to 14.57. Appendix 1 provides a step by step guide for the calculations here. 345

Using the methods of Swinton et al.34 we estimated the proportion of responders in the population of interest within each included RCT (Table 5). The threshold reduction in disability for clinical relevance was set at -10/100, in keeping with recent NICE guidelines for back and radicular pain25. These proportions were adjusted for

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the apparent proportions exceeding this threshold in the comparator groups that were estimated to be due wholly to random variability in the pre to post measurements of disability. It can be seen that these proportions are generally lower than the proportion of participants who exceed the threshold in the intervention groups *per se*.

Discussion

We conducted a systematic review and meta-analysis of the literature in order to quantify the control-group adjusted inter-individual variation in pain, disability and psychosocial outcomes in response to PNE in adults with CMP. Several potential studies did not report the SD of the mean change, and this information could not to be obtained upon request meaning our analysis was restricted to disability.

The inter-individual difference in disability change in response to PNE, as indicated by our SDir of 7.36 /100 units, did not reach our criterion for clinical significance (10 /100 units). Therefore, there is insufficient evidence at present for the existence of inter-individual differences in people's response to PNE over and above random within-subjects variability between baseline and follow-up observations. Although this finding, seems at odds with previous qualitative study findings from our group_{17,18,29}, that qualitative work focused upon patient experience rather than attempting to objectively quantify inter-individual differences. Considering the upper 95% CI (11.12 /100 units) and wide 95% prediction interval -10.20 to 14.57 of the SDir, any inferences regarding "true" inter-individual responses are unclear. Given the small number of included studies, the wide prediction intervals are unsurprising and this

illustrates the importance of statistical power in any analysis of response heterogeneity_{1,2}.

Therefore, it is apparent that more high quality RCTs are needed that sufficiently report relevant data. We encourage researchers and reviewers of academic journals to ensure that the means and standard deviations of the change scores in all treatment groups are reported. This will provide the information required to include the study within meta-analyses of both individual responses and mean effect of treatment.

It is worth highlighting that the very common act of simply looking at the intervention group responses (Table 4) would have falsely led a researcher to think that substantial response heterogeneity was present. This may have led to follow-up analyses to explore potential moderators which may be unwarranted and a waste of resources. Furthermore, any follow-up studies on the same participants may be unethical if there are no true individual differences in response present to explain 1.

This is the first systematic review and meta-analysis to employ the method of calculating true inter-individual differences in response to an intervention within the pain sciences₃₄. Given the huge global burden of chronic pain, and the limited efficacy of current treatment options for matching peoples' individual responses to treatments, appropriate methodology needs to be applied across the pain field. This will hopefully lead to improved quality of care, reduced costs₃₃ and ultimately improve the quality of life of people with pain.

Limitations

Only five studies were eligible for this review which meant that we could only analyse disability data and the inter-individual differences in response to PNE for other outcomes are unknown. Six studies that were otherwise eligible, were excluded because they did not report the appropriate data needed to conduct an inter-individual differences meta-analysis and this data was not available upon email request. We have no reason to believe that authors would withhold this data and thus assume these studies are missing at random. Only studies published in English were eligible for inclusion as no facility for translation was available. Thus, important data from non-English studies may have been missed.

The nature of the comparison group will influence the calculation of the interindividual difference. In the case of usual care comparisons and other intervention comparisons, if these have inherent variability in response within them, beyond random variability (noise) of a true no intervention control, this may mask the degree of interindividual variability seen within the PNE (intervention of interest) group. Thus, this could have influenced the findings. Nevertheless, in the case of intervention vs usual care, if there are true individual differences in the responses to the novel component(s) of the intervention under study, then this should, in theory, manifest itself in a larger change variance in the intervention group vs the usual care group.

Conclusion

This is the first study to investigate "true" inter-individual differences in response within the field of pain. By this, we mean a quantification of response heterogeneity that takes into account the individual differences in baseline to follow-up change that can be observed in the comparator groups, and are attributable to random fluctuation in pain scores over time. Our findings provide little evidence at present of "true" variation in peoples' response to PNE regarding disability, but the evidence is very uncertain. Furthermore, given the wide 95% confidence and prediction intervals any inferences made regarding true individual variation in peoples' response to PNE are unclear. Moreover, given the small number of studies included in the analysis further work is warranted before firm conclusions can be drawn. Therefore, the data currently available does not allow us to clearly identify if individual differences in disability occur for people with CMP following PNE. We would recommend against studies exploring which factors may explain which people will benefit from PNE until such time as the existence of inter-individual differences has been confirmed using appropriate methodology and we would extend this recommendation to all pain interventions.

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569	analysis. <i>J Pain</i> 2019

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571	maximal oxygen uptake to exercise training: a critical review. Sports Med
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Table 1 Summary of findings:

PNE compared to control for treatment of adults with chronic musculoskeletal pain

Patient or population: treatment of adults with chronic musculoskeletal pain

Setting:

Intervention: PNE Comparison: control

Outcomes	Anticipated absolute effects* (95% CI)		Relative effect	Nº of	Certainty of the evidence	A
Outcomes	Risk with control	Risk with PNE	(95% CI)	participants (studies)	(GRADE)	Comments
Change in disability score in the short term. (ST Disability) assessed with: Validated measure of disability converted to percentage Scale from: 0 to 100 (worse)	The mean change in disability score in the short term. was -8.63 units	mean 2.26 units lower (6.49 lower to 1.97 higher)		428 (5 RCTs)	⊕⊖⊖⊖ VERY LOW a,b,c,d,e,f,g,h	PNE may reduce/have little to no effect on change in disability score in the short term. but the evidence is very uncertain.

^{*}The risk in the intervention group (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI).

CI: Confidence interval

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

Explanations

- a. A large proportion of the weight came from a study where there was concern over selection bias, performance bias, attrition bias, reporting bias and other bias. There was concern with most studies over performance bias which whilst normal of these types of studies may still impact the results.
- b. Some variation is size of the effect, however the difference between studies does not reach a clinically meaningful difference
- c. Good overlap of the confidence intervals.
- d. I-Squated above 50%
- e. Tau-Squared higher than point estimate.
- f. Sample of chronic musculoskeletal pain comparing PNE against control using an appropriate outcome measure.
- g. Has over 400 participants but imprecise due to prediction interval including null effect and clinically important benefit.
- h. A comprehensive search was conducted on electronic databases and trials registries. References lists and citing articles of included studies were searched to identify any further articles.

Table 1 Legend: Summary of findings, PNE compared to control for treatment of adults with chronic musculoskeletal pain

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Table 2 Summary of findings:

Do inter-individual differences in disability change in response to PNE exist in adults with chronic musculoskeletal pain?

Patient or population: treatment of adults with chronic musculoskeletal pain

Setting:

Intervention: PNE Comparison: control

Outcomes	Estimated absolute inter- individual difference in response (95% CI)	№ of participants (studies)	Certainty of the evidence (GRADE)	Comments
Inter-individual variability in disability change in the short term. SDIR assessed with: Validated measure of disability converted to percentage Scale from: 0 to 100 (worse)	mean 7.36 units (3.93 lower to 11.12 higher)	428 (5 RCTs)	⊕○○○ VERY LOW a,b,c,d,e,f,g	Little evidence of "true" variation in peoples' response to PNE for disability, but the evidence is very uncertain.

CI: Confidence interval

GRADE Working Group grades of evidence

High certainty: We are very confident that the true difference in response lies close to that of the estimate of the difference in response **Moderate certainty:** We are moderately confident in the difference in response estimate: The true difference in response is likely to be close to the estimate of the difference in response, but there is a possibility that it is substantially different

Low certainty: Our confidence in the difference in response estimate is limited: The true difference in response may be substantially different from the estimate of the difference in response

Very low certainty: We have very little confidence in the difference in response estimate: The true difference in response is likely to be substantially different from the estimate of difference in response

611 Explanations

- a. A large proportion of the weight came from a study where there was concern over selection bias, performance bias, attrition bias, reporting bias and other bias. There was concern with most studies over performance bias which whilst normal of these types of studies may still impact the results.
- b. Some variation in size of the effect, however the difference between studies does not reach a clinically meaningful difference
- c. Good overlap of the confidence intervals.
 - d. Tau-Squared higher than point estimate.
 - e. Sample of chronic musculoskeletal pain comparing PNE against control using an appropriate outcome measure.
- f. While the analysis includes over 400 participants this lack precision due to the very wide prediction interval including both a clinically important positive effect and clinically important negative effect.
- g. No evidence of publication bias. Sample sizes ranged from 62-120. A comprehensive search was conducted on electronic databases and trials registries. References lists and citing articles of included studies were searched to identify any further articles.
- Table 2 Legend: Summary of findings, Do inter-individual differences in disability
- change in response to PNE exist in adults with chronic musculoskeletal pain?

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Study	Methods	Sample	Participants	Intervention(s)	Duration of	Control	Authors conclusions/notes	Setting/country
		size			educational			
		(baseline)/			intervention			
		gender/						
		mean age						
		in years						
van	RCT	N = 105	Fibromyalgia	Written PNE + 1 phone	Unknown	Written Relaxation	Written PNE alone is not	Specialised
Ittersum et		7% M	diagnosed	call for		exercises + 1 phone	effective for changing the	centres for
al. 201338		46.7	according to The	motivation/questions +/-		call for	impact of the illness on daily	chronic pain
			American College	2x phone calls/emails		motivation/questions	life, pain catastrophising, or	and chronic
			of	for further		+/- 2x phone	illness perceptions in	fatigue.
			Rheumatology	clarification/questions		calls/emails for further	fibromyalgia patients.	Belgium.
			1990 criteria ₄₂ .			clarification/questions		
			18-65 years of age.		<i>y</i>			
			Baseline pain as					
			mean % = 71.5%					
			Duration of pain in					
			mean months =					
			unknown					
Gallagher,	RCT	N = 79	18-75 years of age	80-page booklet divided	Unknown	80-page booklet	Written material using	Unknown
McAuley		39% M	with pain that had	into 11 sections -		divided into 11	metaphors to explain key	Unknown
	1					I .	ı	ı

Moseley 20136 disrupt their activities of daily living for more than the previous 3 months. Baseline pain as mean % = 65%	and		43.5	been sufficient to	Metaphors and stories to		sections - Advice about	biological concepts increased	
living for more than the previous 3 months. Baseline pain as mean % = 65% Duration of pain in mean (SD) months = 28 (19.5) Pires, Cruz and Caeiro, RCT N = 62 Low back pain >3 2x 1.5h Group PNE. and Caeiro, 35% M months duration 12 sessions of aquatic exercise over 6 weeks.	Moseley			disrupt their	help understand the		managing pain (The	knowledge of pain biology and	
than the previous 3 months. Baseline pain as mean % = 65% Duration of pain in mean (SD) months = 28 (19.5) Pires, Cruz and Caeiro, RCT N = 62 Low back pain >3 2x 1.5h Group PNE. PNE 3h 12 sessions of aquatic exercise over 6 weeks. when compared to material that presented biopsychosocial advice for pain management. PNE is a clinically effective addition to aquatic exercise. clinic.	20136			activities of daily	biology of pain		back book and Manage	decreased catastrophic thought	
months. Baseline pain as mean % = 65% Duration of pain in mean (SD) months = 28 (19.5) Pires, Cruz and Caeiro, RCT N = 62 Low back pain >3 2x 1.5h Group PNE. The service of a quatic exercise over 6 weeks. The service of pain management advice for pain management. PNE 3h 12 sessions of aquatic exercise. PNE 3h 12 sessions of aquatic exercise.				living for more			your pain)	processes about pain and injury	
Baseline pain as mean % = 65% Duration of pain in mean (SD) months = 28 (19.5) Pires, Cruz RCT N = 62 Low back pain >3 2x 1.5h Group PNE. and Caeiro, and Caeiro				than the previous 3				when compared to material that	
Baseline pain as mean % = 65% Duration of pain in mean (SD) months = 28 (19.5) Pires, Cruz and Caeiro, RCT N = 62 Low back pain >3 Low back				months.				presented biopsychosocial	
mean % = 65% Duration of pain in mean (SD) months = 28 (19.5) Pires, Cruz RCT N = 62 Low back pain >3 2x 1.5h Group PNE. PNE 3h 12 sessions of aquatic exercise over 6 weeks. RCT N = 62 Low back pain >3 2x 1.5h Group PNE. PNE 3h 12 sessions of aquatic exercise. Clinic.								advice for pain management.	
Duration of pain in mean (SD) months = 28 (19.5) Pires, Cruz RCT N = 62 Low back pain >3 2x 1.5h Group PNE. PNE 3h 12 sessions of aquatic exercise. Outpatient and Caeiro, and Caeiro and Caeiro, and Caeiro and Caeiro, and Caeiro an				Baseline pain as					
mean (SD) months = 28 (19.5) Pires, Cruz RCT N = 62 Low back pain >3 2x 1.5h Group PNE. PNE 3h 12 sessions of aquatic exercise. PNE is a clinically effective addition to aquatic exercise. clinic.				mean % = 65%					
mean (SD) months = 28 (19.5) Pires, Cruz RCT N = 62 Low back pain >3 2x 1.5h Group PNE. PNE 3h 12 sessions of aquatic exercise. PNE is a clinically effective addition to aquatic exercise. clinic.									
Pires, Cruz RCT N = 62 Low back pain >3 2x 1.5h Group PNE. PNE 3h 12 sessions of aquatic PNE is a clinically effective Outpatient and Caeiro, 35% M months duration 12 sessions of aquatic exercise over 6 weeks. addition to aquatic exercise. clinic.				Duration of pain in					
Pires, Cruz RCT N = 62 Low back pain >3 2x 1.5h Group PNE. PNE 3h 12 sessions of aquatic exercise over 6 weeks. PNE is a clinically effective outpatient exercise over 6 weeks.				mean (SD) months					
and Caeiro, and Caeiro, months duration 12 sessions of aquatic exercise over 6 weeks. addition to aquatic exercise.				= 28 (19.5)					
	Pires, Cruz	RCT	N = 62	Low back pain >3	2x 1.5h Group PNE.	PNE 3h	12 sessions of aquatic	PNE is a clinically effective	Outpatient
	and Caeiro,		35% M	months duration	12 sessions of aquatic		exercise over 6 weeks.	addition to aquatic exercise.	clinic.
201526 51 +/- leg pain. 18-65 exercise over 6 weeks. Control 3h 30-50m each session. The addition of PNE resulted in Portugal	201526		51	+/- leg pain. 18-65	exercise over 6 weeks.	Control 3h	30-50m each session.	The addition of PNE resulted in	Portugal
years of age. 30-50m each session. statistically significant reduction				years of age.	30-50m each session.			statistically significant reduction	
in pain intensity at 3-month								in pain intensity at 3-month	
Baseline pain as follow up. No statistically				Baseline pain as				follow up. No statistically	
mean % = 42.9% significant differences were				mean % = 42.9%				significant differences were	
found for pain intensity at 6								found for pain intensity at 6	
Duration of pain in weeks follow up or functional				Duration of pain in				weeks follow up or functional	
mean (SD) months disability at either follow up.				mean (SD) months				disability at either follow up.	
= unknown				= unknown					
Louw et al. RCT N = 67 Patients with 0.5h individual PNE. PNE 0.5h Lumbar surgery alone Providing a single PNE session 7 Clinical site	Louw et al.	RCT	N = 67	Patients with	0.5h individual PNE.	PNE 0.5h	Lumbar surgery alone	Providing a single PNE session	7 Clinical sites
2014/16 _{19,20} 46% M lumbar + usual care to patients prior to lumbar in the US.	2014/1619,20		46% M	lumbar			+ usual care	to patients prior to lumbar	in the US.

		49.6	radiculopathy,	PNE booklet "your	Control 0		surgery results in significant	
			scheduled for	nerves are having back			reduction in healthcare costs 3-	
			lumbar surgery.	surgery" & Lumbar			years after LS.	
			18-65 years of	surgery + usual care				
			age.					
ļ								
			Baseline pain as					
			mean % = 48.4%					
			Duration of pain in					
			mean (SD) months					
			= 3 (7.5)					
Malfliet et	RCT	N = 120	Non-specific	3 PNE sessions	PNE 1.88h	3 biomedical education	PNE, and not neck/back school	University
al. 201821		39.2% M	chronic spinal pain	1. 0.5-1h group		sessions	education, is able to improve	hospitals in
		39.8	(neck and lower	(maximum of 6	Control	1. 0.5-1h group	kinesiophobia, beliefs regarding	Ghent and
			back) at least 3	patients).	1.88h	(maximum of 6	the negative impact of the	Brussels,
			days a week for at	Information booklet		patients).	illness on quality of life and	Belgium.
			least 3 months	provided at the		Information	functional capacity, and beliefs	
			since the first	end.		booklet provided	regarding the chronicity of pain	
			symptoms.	2. ~0.63h home-based		at the end.	and the time scale of illness	
				online e-learning		2. ~0.63h Home-	symptoms. However, none of	
			18-65 years of age	module containing		based online e-	the educational programs of this	
				3 explanatory		learning module	study were able to decrease the	
			Baseline pain as	videos and		containing 3	participants perceived disability	
			mean $\% = 50.65$	viueus allu		C	due to pain. Nevertheless, as	
						explanatory videos	kinesiophobia is generally	

	Duration of pain in	questions about		3.	0.5 Individual.	considered to be a strong
	mean (SD) months	pain.			Focus on patients'	predictor and mediator of
	= 82 (143.25)	0.5 Individual			personal needs	chronic pain, PNE is preferred
		education. Focus on			following	as the educational approach for
		patients' personal			difficulties with	people with non-specific
		needs following			session 2. Focus	chronic spinal pain.
		difficulties with			on the application	
		session 2. Focus on			of knowledge to	
		the application of	, (participants life.	
		knowledge to			por no parino irrei	
		participants life.				

Table 3 Legend: Randomized controlled trial, RCT. Male,

Table 3 Legend: Characteristics of included studies. PNE, Pain neuroscience education. SD, Standard deviation. RCT,

635 Randomised controlled trial

Table 4 Critical appraisal of quantitative studies

Study	Score /7	Percentage
Gallager 2013 ⁶	5	71%
Louw 2014/16 ^{19,20}	3	43%
Malfliet 2018 ²¹	6	86%
Pires 2015 ²⁶	3	43%
van Ittersum 2013 ³⁸	1	14%

638 Figure 4 Legend: Forest plot of PNE versus control in the short term; primary

outcome disability mean difference.

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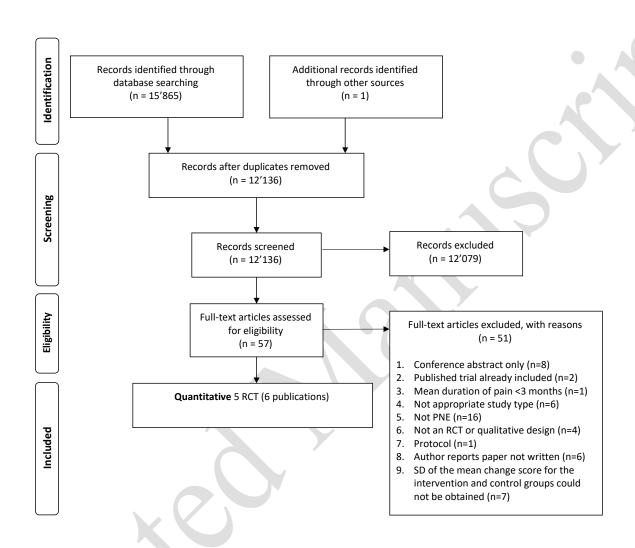
Table 5: Proportions of responders.

Study	Mean	SD	%	Mean	SD	%	Mean	SD for	0/0
	Change	(PNE)	responders	change	(Con)	Responders	treatment	true Ind	Responders
	(PNE)	, ,	(PNE)	(Con)		(Con)	effect	diffs	based on
	, ,						(PNE-		SDir ₃₄
							Con)		
van									
Ittersum et									0
al. 2013 ₃₈	0.7	4.2	0	0.3	2.9	0	0.4	3.0	
Pires, Cruz									29
and Caeiro,									
201526	-11.1	15.8	53	-7.7	10.6	41	-3.4	11.7	
Louw et al.									
2014/1619,20	-12.0	18.5	54	-11.1	13.8	53	-0.9	12.3	23
Malfliet et									
al. 201821	-1.1	13.8	26	1.6	11.2	15	-2.7	8.1	18
Gallagher,									
McAuley									
and									
Moseley		1							
20136	-36	17	94	-27.0	15.0	87	-9.0	8	45

Table 5 Legend: Proportions of responders. PNE, Pain neuroscience education.

644 Con, Control. SD, Standard deviation. SDir, Standard deviation for individual

645 responses.



647 Figure 1 Legend: PRISMA flow diagram of search and study selection process.

648 (Adapted from Moher et al.22).

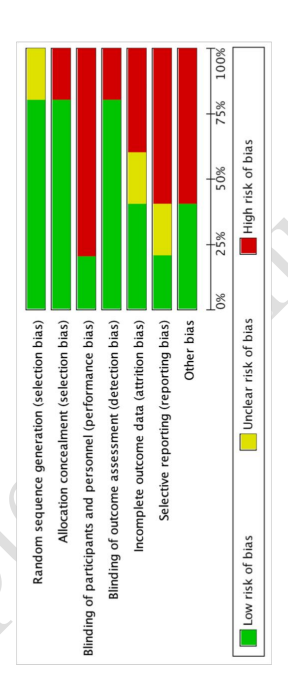
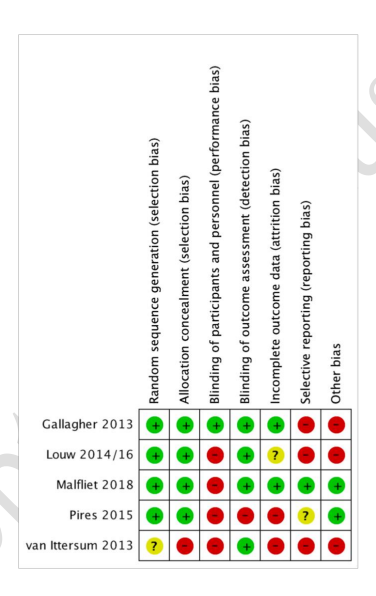


Figure 2 Legend: Risk of bias graph.



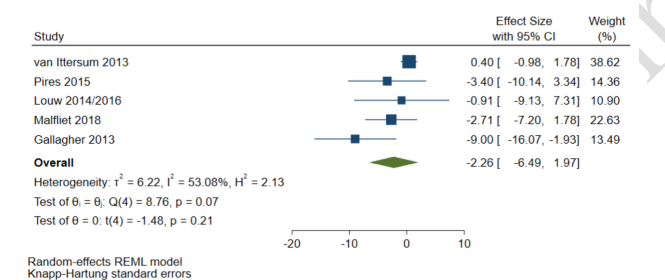


Figure 4 Legend: Forest plot of PNE versus control in the short term; mean

difference of disability between groups.

Step 1

We extracted the standard deviation (SD) of the changes in disability for both control (C) and PNE (I) groups.

Study	SDC	Mean I
van Ittersum	2.9	0.7
Gallagher	15	-36
Pires	10.6	-11.1
Louw	13.79	-12
Malfliet	11.15	-1.1

Step 2

The true individual response variance (intervention minus control) was then calculated by $V(SD_1^2-SD_C^2)$ (Hopkins, 2015).

Study	IR_Variance	SDI	SDC
van Ittersum	9.23	4.2	2.9
Gallagher	64	17	15
Pires	137.28	15.8	10.6
Louw	152.09	18.5	13.79
Malfliet	65.84	13.79	11.15

Step 3

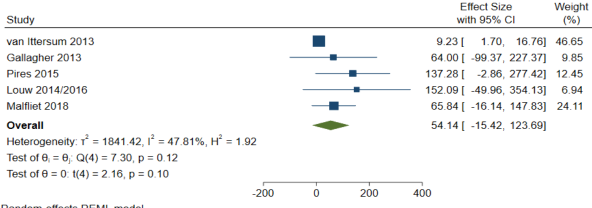
The standard error (SE) for this variance was then calculated using the equation: SE = $V[2(SD_14/DF_1 + SD_C4/DF_C)]$, where DF₁ and DF_C are the degrees of freedom of the standard deviation in the PNE group and the control groups (Hopkins, 2015).

Study	IR_Variance	SE	SDI	SDC	n I	n C
van Ittersum	9.23	3.83949378	4.2	2.9	53	52
Gallagher	64	83.3522758	17	15	40	39
Pires	137.28	71.5013373	15.8	10.6	30	32
Louw	152.09	103.087047	18.5	13.79	29	33
Malfliet	65.84	41.8303551	13.79	11.15	60	60

Step 4

The individual response variances and their SEs were meta-analysed using an REML model combined with Knapp-Hartung method. It's important to highlight that the variances are unbiased, whereas the SD is not, and deriving a SE for the SD for individual responses is also problematic. Thus, we synthesised the individual response variances instead of the SDs for individual responses. The point estimate for the pooled individual response variance were derived together with a 95% CI to express its uncertainty.

Forest plot of **Variance** Meta-analysis for estimating individual differences in response:



Random-effects REML model Knapp-Hartung standard errors

Step 5

The point estimate and CIs were then square rooted to convert to an SD metric. If the lower limit was negative, the sign was ignored, the square root taken, and the sign re-applied. This approach is consistent with the 'no bound' option in SAS/STAT® software, which permits negative variances (SAS Institute Inc. 2017. SAS/STAT 14.3 User's Guide. Cary, NC: SAS Institute Inc.).

			As SD with
			sign re-
	As variance	SD without sign	applied
Total point estimate	54.14		7.35798886
Lower Cl -	15.42	3.92683078	-3.9268308
Upper Cl	123.69		11.1216006

Steps to calculate the prediction interval for the inter-individual differences point estimate

PI = pooled estimate +/-
$$t_{(n-2)}x$$
 SQRT(SE² + tau^2)

Pooled Est 54.14

3.182 is the two-tailed t value for n-2 degrees of freedom = 3 degrees of freedom, and P=0.05. See: http://www.ttable.org/student-t-value-

3.182 calculator.html $t_{(n-2)} =$ SE= 25.0508232 $SE^2 =$ 627.543743 (From STATA) tau² = 1841.4235 $(SE^2 + tau^2) = 2468.96724$ $SQRT(SE^2 + tau^2) =$ 49.6887034 PI = Pooled est +/-SQRT(SE2 + tau2)t(n-2) Х PI = 54.14 +/-49.6887034 3.182 x 3.182 +/-PI = 158.109454

PI Upper = 212.249454 PI Lower = -103.96945

Square root the above values to convert from variance to SD to get to the PI for the SDir:

PI Upper = 14.5687836 PI Lower = -10.196541