Word counts: Text: 2698, Abstract: 300

### Measuring lumbar reposition accuracy in patients with unspecific low back pain – Systematic Review and Meta-analysis

Rausch Osthoff, Anne-Kathrin MScPT<sup>a</sup>; Ernst, Markus MScPT<sup>a</sup>; Rast, Fabian MSc<sup>a</sup>; Mauz, Danica MSc<sup>a</sup>, Graf, Eveline PhD<sup>a</sup>; Kool, Jan PhD<sup>a</sup>; Bauer, Christoph MSc<sup>ab</sup>

<sup>a</sup>Zurich University of Applied Sciences (ZHAW), Department of Physiotherapy, Winterthur, Switzerland <sup>b</sup>University of Tampere, Faculty of Medicine, Tampere, Finland

**Conflict of interest statement:** None of the authors has a conflict of interest related to the content of the manuscript.

Funding: None of the authors received any funding.

Correspondence: Christoph Bauer Zurich University of Applied Sciences Department of Physiotherapy Technikumstrasse 71 8401 Winterthur Switzerland P.O. Box Phone: +41 58 934 64 49 Fax : +41 58 935 64 49 E-mail: bauc@zhaw.ch

### Abstract:

Study Design. Systematic review and meta-analysis.

**Objective.** To evaluate if patients with nonspecific chronic low back pain (NSCLBP) show a greater lumbar reposition error (RE) than healthy controls.

**Summary of Background Data.** Studies on lumbar RE in patients with NSCLBP present conflicting results.

**Methods.** A systematic review and meta-analysis of the available literature were performed to evaluate differences in RE between NSCLBP patients and healthy controls. Data on absolute (AE), constant (CE) and variable error (VE) were extracted and effect sizes (ES) were calculated. For the CE flexion pattern and active extension pattern, subgroups of patients with NSCLBP were analyzed. Results of homogeneous studies were pooled. Measurement protocols and study outcomes were compared. The quality of reporting and the authors' appraisal of risk of bias were investigated.

**Results.** The original search revealed 178 records of which 13 fulfilled the inclusion criteria. The majority of studies showed that patients with NSCLBP produced a significantly larger AE (ES 0.81 [CI .13-1.49]) and VE (ES 0.57 [CI 0.05-1.09]) compared to controls. CE is direction- specific in flexion and active extension pattern subgroups of patients with NSCLBP (ES 0.39 [CI -1.09-0.3] and ES 0.18 [CI -.3-0.65], respectively). The quality of reporting and the authors' appraisal of risk of bias varied considerably. The applied test procedures and instrumentation varied between the studies, which hampered the comparability of studies.

**Conclusions.** Whilst patients appeared to produce a larger lumbar RE compared to healthy controls, study limitations render firm conclusions unsafe. Future studies should pay closer attention to power, precision and reliability of the measurement approach, definition of outcome measures and patient selection. We recommend a large, well powered, prospective randomised control study which uses a standardized measurement approach and definitions for AE, CE, and VE to address the hypothesis that proprioception may be impaired with CLBP.

**Keywords:** Low back pain, proprioception, spine, posture, review, meta-analysis, lumbar reposition error, lumbosacral region, lumbar spine, motor control, movement control

### Key Points:

- Patients with NSCLBP tend to produce a larger lumbar RE compared to healthy controls.
- The applied test procedures and instrumentation varied between studies.
- We recommend a standardized measurement approach and the use of standardized and accurate definitions for lumbar reposition error to be used in future studies.

### Mini Abstract:

A systematic review and meta-analysis were performed to investigate differences in lumbar reposition error (RE) between patients with non-specific chronic low back pain (NSCLBP) and controls. Patients with NSCLBP produce greater RE compared to controls. We recommend standardized measurement approaches and definitions for RE to be used in future studies.

Low back pain (LBP) affects up to 84% of people in industrialized countries<sup>(1)</sup>. In 2005, the total direct costs of LBP in Switzerland amounted to €2.6 billion<sup>(2)</sup>. Evidence recommends the use of a prognostic sub-classification including cognitive, physical and lifestyle factors for all chronic LBP (CLBP) patients who do not display underlying red flag disorders; specific pathoanatomical disorders or pain disorders driven from the forebrain with a dominance of non-organic factors <sup>(3,4,5,6,7)</sup>. The physical factor of this classification system includes a large subgroup of patients with mal-adaptive movement or control disorders<sup>(3,4,5,6)</sup>. Movement and control disorders are interpreted as mal-adaptive primary physical compensations, after an initial painful episode, which drive the CLBP state<sup>(3)</sup>. They presumably lead to a proprioceptive deficit, due to stress on local muscle spindles and joint receptors in the painful area resulting from stress to a joint caused by an individual's maladaptive movement<sup>(3)</sup>. Proprioceptive deficits may lead to altered central sensory-motorcontrol mechanisms and disrupted body schema. Subsequently abnormal joint and tissue loading during daily activities and postures may affect local proprioceptors and maintain this vicious circle<sup>(7,8,9,10,11,12,13)</sup>. Reposition error (RE) is regarded as a measure reflecting proprioception deficits in the lower spine and typically involves participants trying to reproduce a specific target body position<sup>(14,15,16)</sup>.

RE can be expressed as absolute error (AE), constant error (CE), or variable error (VE). AE represents the error magnitude and is defined as the absolute difference between the target lumbar angle and actual lumbar angle. CE represents the error magnitude direction such that CE indicates bias towards a particular direction where negative CE typically represents a bias in the undershooting direction. VE describes the variability of the subjects' performance equivalent to the standard deviation of RE. High VE values reflect high variability in repositioning<sup>(17)</sup>.

Using lumbar RE as an outcome measure several studies have investigated deficits in proprioception in patients with LBP<sup>(11,12,14,15,16,17,18,19,20,21,22,23,24,25)</sup>. In these tests, patients are asked to reproduce a specific (e.g., neutral) lumbar position after performing an active or passive movement. Some studies reported an increased lumbar RE of patients with LBP compared to a healthy population<sup>(12,14,15,16,18,21,22,23)</sup>. Classifying patients with nonspecific CLBP (NSCLBP) based on movement and control impairments<sup>(3)</sup> revealed direction-specific differences in lumbar RE between flexion pattern (FP) and active extension pattern (AEP) subgroups of NSCLBP patients<sup>(14,16)</sup>. A recent RCT showed that these lumbar spine position sense deficits were treatable with a classification guided postural intervention<sup>(26)</sup>. However, other studies have shown no differences between patients with LBP and healthy controls when testing for lumbar position sense<sup>(17,19,21)</sup>, even after they were sub-grouped according to a McKenzie classification system or ICD-10 codes<sup>(17)</sup>.

As it is discussed controversial if proprioception is altered in patients with NSCLBP that display physical factors a meta-analysis of the earlier results is advisable and a systematic review may contribute to a better understanding of this issue.

Measurement procedures for assessing RE and findings vary among studies in patients with LBP and healthy controls. Therefore, the aim of this systematic review and meta-analysis was to evaluate if patients with NSCLBP produce a greater lumbar RE. Thus, a statistical pooling of homogeneous study results was performed. Furthermore, design and measurement methods of RE studies were compared to state recommendations for further research.

### MATERIALS AND METHODS

### Data Sources and Searches

Study identification commenced by electronic searching, using the MEDLINE (through Pubmed), CINAHL, and Cochrane Library, on articles published between January 1, 1990 and September 30, 2013. Search terms used were low back pain, proprioception, position sense, kinesthesis, reposition, and repositioning. Both Medical Subject Headings terms and free text words were entered. A combination of these terms was used to extract a comprehensive list of articles, from which the titles and abstracts were screened for eligibility. An additional search for grey literature on issue-specific databases<sup>(27,28,29)</sup>, citation tracking, and key author searches was conducted.

### Eligibility Criteria

The following criteria were applied to determine the eligibility of each study for inclusion in the meta-analysis:

- patients with NSCLBP and healthy controls,
- at least one measure reflecting RE (AE, CE, VE),
- published in English or German

Two reviewers independently evaluated records for eligibility. Disagreement was resolved by discussion and consensus. To avoid duplication in pooling, data were included only once if they were reported in previously published work.

### Quality Assessment

Two reviewers independently analysed the quality of the included studies as recommended by the PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analyses) statement for reporting systematic reviews and meta-analyses of studies that evaluate health care interventions: explanation and elaboration<sup>(30,31)</sup>. Accordingly, the STROBE (Strengthening the Reporting of Observational Studies in Epidemiology) statement was used to analyze both the quality of reporting and the author's 'appraisal of risk of bias'<sup>(32,33)</sup>. Discrepancies were solved by consensus. Results were summarized in tabular form to enable a sensitivity analysis based on quality criteria.

### Data Analysis

Two reviewers independently extracted information of each study including the setting of the study, characteristics of patients, inclusion and exclusion criteria, instrumentation, test protocol, and outcomes (tasks and variables). Those data were presented narratively in tabular form. Data on reliability and measurement error of the test protocols were extracted and presented in tabular form.

Descriptive data for continuous variables were expressed as mean and standard deviation (SD). The Cochrane collaboration's Revman 5.2.7 software was used for a pooled data analysis. Data were reported as AE, CE, or VE. Effect sizes of single studies were expressed as Hedges g or Cohens r, if the original data was non-normally distributed, with 95% confidence intervals. Those studies describing results reflecting AE, CE, and VE evaluated with neutral-slumped-neutral sitting were used for meta-analysis using a random effects model, subgrouped for adults and adolescents. Neutral-slumped-neutral was chosen as pooling criteria because six studies used this setup. All other setups were used once. Additionally, CE was analyzed independently for FP and AEP subgroups of NSCLBP, as CE is direction specific in these subgroups<sup>(14,15,16)</sup>. As the definition of undershooting into a flexed position and overshooting into an extended position varied between the studies, we applied a common definition and changed the sign of study results in one study<sup>(16)</sup> according to this definition. Undershooting into a flexed position was given a negative sign while overshooting into an extended position was given a positive sign. To assess heterogeneity, the Q-statistic and its p value were calculated.  $l^2$  was calculated as a mass of between-study heterogeneity (for each set of effect sizes) according to Borenstein<sup>(34)</sup>. The meta-analyses were first performed including all studies fulfilling the above criteria. As a sensitivity analysis, the metaanalysis were then repeated by excluding studies with poor quality of reporting and studies appearing as outliers to assess their influence on the meta-analysis.

### RESULTS

The search revealed 178 records; 31 of them were screened in full-text (Figure 1). Eighteen studies were excluded due to study design (e.g., interventional studies, no healthy control group), outcome variables (no AE, CE, VE), or the character of included subjects (no NSCLBP). A total of 13 studies<sup>(11,12,14,15,16,17,18,19,20,21,22,23,24,25)</sup> fulfilled the inclusion criteria (Table 1). Four out of thirteen of the included studies did not provide sufficient data on reposition error (mean, SD)<sup>(17,20,21,22)</sup>. Upon contacting the corresponding authors, we did not

receive this information from them. The overall loss of subjects was 148 patients with NSCLBP and 86 controls.

Table 2 summarizes the applied test procedures and instrumentation, which varied largely between the studies. Table 3 shows the reported variables and calculated effect sizes. The majority of the studies showed that NSCLBP patients produced a significantly larger AE and VE compared to controls. The quality of reporting and the authors' appraisal of risk of bias (STROBE) varied considerably. Some studies do not present information on risk of bias and attempts to reduce bias (Table 4). Reporting on reliability and measurement error was inconsistent with studies not reporting either or referring to measurement error and reliability of the measurement device (Table 5) <sup>(12, 15, 18, 19)</sup>.

Six studies were included in the meta-analysis as they shared the same measurement protocol (neutral-slumped-neutral in sitting) (Figure 2). The studies were subgrouped, according to the age of the participants, into adults<sup>(12,15,16,24,25)</sup> and adolescents<sup>(14)</sup>.

The overall effect size of 0.81 [CI 0.13-1.49] illustrates that patients with LBP produce a larger AE than healthy controls. The overall heterogeneity of study effects was considerable ( $I^2$ =83%, p<.05); it was no longer restricted to studies with poor quality of reporting but to all studies included in the meta-analysis. Heterogeneity did not change when single studies were excluded from the meta-analysis.

Two studies were included in a meta-analysis on VE (Figure 3). The overall effect size for VE of 0.68 [CI 0.01-1.36] illustrates that patients with NSCLBP have a higher deviation of the reposition error than healthy controls. The heterogeneity of study effects was substantial and significant ( $l^2=75\%$ , p<.046).

Three studies were included in a meta-analysis of CE (Figures 4 and 5). Again, the studies were subgrouped, according to the age of participants, into  $adults^{(15,16)}$  and  $adolescents^{(14)}$  and further for FP and AEP. The overall effect size for CE for FP 0.39 [CI -1.09-0.3] indicates that FP NSCLBP patients undershoot into flexion compared to healthy controls. The overall effect size for CE for AEP 0.18 [CI -0.3-0.65] indicates that AEP NSCLBP patients overshoot into extension compared to healthy controls. However, the results are not significant. The adolescent sample in the study by Astfalck and colleagues showed a reverse pattern<sup>(14)</sup>. The heterogeneity of study effects for the FP was considerably (I<sup>2</sup>=75%, p<.05). Removing the study of Astfalck and colleagues<sup>(14)</sup> lowered the heterogeneity considerably (I<sup>2</sup>=26%, p=.24). The heterogeneity of study effects for the AEP subgroup was neglectible (I<sup>2</sup>=36%, p=.21)

### DISCUSSION

The results of this study indicate that lumbar reposition sense is impaired in patients with NSCLBP compared to healthy controls. In the majority of the studies, patients with NSCLBP produced a greater AE and VE than healthy controls. Additionally, patients with FP NSCLBP tend to undershoot into flexion while patients with AEP NSCLBP overshoot into extension. Recent studies tend to report RE for FP and AEP subgroups of NSCLBP patients based on a better and improved understanding of NSCLBP. These studies showed that the direction of RE differs between subgroups. AE and CE tend to show larger effect sizes than VE.

The meta-analysis is based on data of neutral-slumped-neutral sitting<sup>(12,14,15,16)</sup> because these studies used a comparable measurement procedure and patient criteria. The meta-analysis showed similar findings for adults and adolescents regarding AE and VE.

However study limitations render firm conclusions unsafe. The quality of reporting and the authors' appraisal of risk of bias, in some studies, were limited. Some studies recruited only small samples<sup>(12,15,18,20,21,22,23,24,25)</sup>.

In some studies the inclusion and exclusion criteria were imprecise which however did not affect the studies of the meta-analysis<sup>(11,17,20)</sup>.

It is hypothesised that reduced proprioception is present in the group of CLBP disorders where patients present movement or control impairments<sup>(3)</sup>. Shortcomings in former studies to screen for this specific group and exclude patients with underlying red flag disorders, specific pathoanatomical disorders and pain disorders with a dominance of non-organic factors may have added to the inconsistency of the findings<sup>(17,19,20)</sup>. Only five studies reported attempts to minimize selection bias by using matching criteria<sup>(12,14,15,17,23)</sup>.

However within the meta-analysis, studies which included NSCLBP patients with dominant physical factors were included.

The measurement approach varied considerably among studies. Different testing positions, number of repetitions, movement instructions and measurement systems make it difficult to compare findings. Some studies used a warm up phase, practice trials, or demonstrations<sup>(11,12,18)</sup> while others did not<sup>(16,21)</sup>.

The most frequently used test position was sitting<sup>(11,12,15,16,17)</sup> The test positions can influence the results of lumbar position sense testing as proprioceptive input may differ depending on which segment of the spine moves (proximal or distal segment) and on the loading of the spine (unloaded vs. loaded). As lumbar RE appears direction specific in FP and AEP NSCLBP populations, the tested movement direction might influence the outcome<sup>(14,16,26)</sup>. Measurement systems varied and the scale and accuracy of these systems may differ and affect the measurement outcome when measuring small angular differences. The placement of devices/markers varied considerably with some studies assessing the total lumbar spine<sup>(12,16,17,21,22,24,25)</sup> while others assessed the lower part of the lumbar spine<sup>(14,15,18)</sup> or larger areas<sup>(21,23)</sup>. The number of repetitions varied between studies and ranged from 3 to  $10^{(14,17)}$ . The number of repetitions influences the stability of the results.

Several studies reported only one specific aspect of RE, usually AE, which limited the information that could be extracted from these studies<sup>(18,19,21,23,24,25)</sup>. The definitions of AE, CE, and VE were described rather vaguely in some studies<sup>(16,18,20,23)</sup>. This hampers comparability, as it is not clear if the same mathematical definition was used for the same type of error.

### Recommendations for future research

Future studies, using sufficiently large, matched sample sizes should use adequate screening and diagnostic instruments including the O'Sullivan classification system<sup>(35)</sup>, imagining techniques, response to facet-joint injection and questionnaires such as the STarT Back screening tool<sup>(36)</sup>, the Orebro questionnaire<sup>(37)</sup> or the Fear-avoidance beliefs questionnaire (FABQ)<sup>(38)</sup>. Collaboration between allied health and medical professions is required to elucidate the veracity of their hypotheses and for precise patient and control selection.

For future studies we recommend a test position and movement directions that are reported as an aggravating factor by the tested population, such as flexion and extension in sitting for CLBP patients with physical factors<sup>(12,15,16)</sup>. We further recommend an analysis of criterion validity and between-day reliability of both measurement error and reliability of the measurement device and approach, a standardized and validated placement of the devices and defining the adequate number of repetitions through a D-study<sup>(39,40)</sup>.

We recommend that authors present exact formulas for AE, CE, and VE and suggest the following definitions, with E being the expected error (E) which is equivalent to the mean error in finite populations:

AE is the mean absolute difference between the starting  $(\Theta)$  and final position (X).

$$AE = E[|X - \Theta|]$$

CE is the mean signed difference between  $\Theta$  and X.

$$CE = E[X - \Theta]$$

VE is the square root of the error variance.

$$VE = \sqrt{Var([X - \Theta])}$$

We recommend continuing to evaluate various aspects of error (AE, CE, and VE). Other aspects of RE are hardly mentioned in this review. Movement time or velocity<sup>(20)</sup>, learning

 phase, mean-squared RE, and the relevance of visual or verbal feedback need to be investigated. Further prospective randomized controlled studies (RCT) are needed to assess if improvements in movement control are associated with improvements in proprioception. The association of lumbar RE errors to other movement dysfunctions and other dimensions of LBP should be assessed. In summary only a large, well powered, prospective RCT with a standardized measurement approach can address the hypothesis that proprioception is impaired in CLBP patients with physical factors and treatable through a classification guided intervention.

### Limitations of this study

It has been discussed that using a funnel plot should assess publication bias when 10 or more studies can be pooled. As only six studies were included in the meta-analysis, a funnel plot would have been inconclusive regarding publication bias<sup>(41)</sup>. We considered a factor analysis of elements in the study design that would determine if a study found differences between NSCLBP patients and controls. However, due to the limited number of studies and the great variety in study designs, this was not possible. Therefore, we focused to choose the presented qualitative appraisal of methodological differences and their effect on the study design.

### Clinical implication

Clinical measures of RE are being used to assess proprioceptive deficits. The studies included in this review and meta-analysis strengthens the assumption that patients with NSCLBP produce greater RE than healthy controls and, therefore, have proprioceptive deficits compared to healthy controls. So far, only one study has investigated the responsiveness of RE to treatment. This study has shown an improvement in pain and RE after a classification guided intervention<sup>(3,26)</sup>. Until conclusions can be drawn from larger studies we propose clinical interpretation of RE with caution.

### CONCLUSION

Whilst patients appeared to produce a larger lumbar RE compared to healthy controls, study limitations render firm conclusions unsafe. Future studies should pay closer attention to power, precision and reliability of the measurement approach, definition of outcome measures and patient selection. We recommend a large, well powered, prospective randomised control study which uses a standardized measurement approach and definitions for AE, CE, and VE to address the hypothesis that proprioception may be impaired with CLBP.

### REFERENCES

- 1. Walker BF, The prevalence of low back pain: A systematic review of the literature from 1966 to 1998. *J Spinal Disord*, 2000. 13(3): p. 205-17.
- 2. Wieser S, Horisberger B, Schmidhauser S, et al., Cost of low back pain in switzerland in 2005. *Eur J Health Econ*, 2011. 12(5): p. 455-67.
- 3. O'Sullivan P, Diagnosis and classification of chronic low back pain disorders: Maladaptive movement and motor control impairments as underlying mechanism. *Man Ther*, 2005. 10(4): p. 242-55.
- 4. Waddell G, The back pain revolution. second ed. 2004: Churchill Livingstone.
- 5. Hill JC, Foster NEHay EM, Cognitive behavioural therapy shown to be an effective and low cost treatment for subacute and chronic low-back pain, improving pain and disability scores in a pragmatic rct. *Evidence Based Medicine*, 2010. 15(4): p. 118-9.
- 6. Vlaeyen JW Linton SJ, Fear-avoidance model of chronic musculoskeletal pain: 12 years on. *Pain*, 2012. 153(6): p. 1144-7.
- 7. Adams MA, Burton KBogduk N, The biomechanics of back pain. 2006: Churchill Livingstone Elsevier.
- 8. Richardson C, Hodges PW, Hides J, et al., Therapeutic exercise for lumbopelvic stabilization: A motor control approach for the treatment and prevention of low back pain. 2004: Churchill Livingstone.
- 9. Holm S, Indahl ASolomonow M, Sensorimotor control of the spine. *J Electromyogr Kinesiol*, 2002. 12(3): p. 219-34.
- 10. Bray H Moseley GL, Disrupted working body schema of the trunk in people with back pain. *Br J Sports Med*, 2011. 45(3): p. 168-73.
- 11. Brumagne S, Cordo P, Lysens R, et al., The role of paraspinal muscle spindles in lumbosacral position sense in individuals with and without low back pain. *Spine (Phila Pa 1976)*, 2000. 25(8): p. 989-94.
- 12. O'Sullivan PB, Burnett A, Floyd AN, et al., Lumbar repositioning deficit in a specific low back pain population. *Spine (Phila Pa 1976)*, 2003. 28(10): p. 1074-9.
- 13. Luomajoki H Moseley GL, Tactile acuity and lumbopelvic motor control in patients with back pain and healthy controls. *Br J Sports Med*, 2011. 45(5): p. 437-40.
- 14. Astfalck RG, O'Sullivan PB, Smith AJ, et al., Lumbar spine repositioning sense in adolescents with and without non-specific chronic low back pain an analysis based on sub-classification and spinal regions. *Man Ther*, 2013. 18(5): p. 410-7.
- 15. O'Sullivan K, Verschueren S, Van Hoof W, et al., Lumbar repositioning error in sitting: Healthy controls versus people with sitting-related non-specific chronic low back pain (flexion pattern). *Man Ther*, 2013.
- 16. Sheeran L, Sparkes V, Caterson B, et al., Spinal position sense and trunk muscle activity during sitting and standing in nonspecific chronic low back pain: Classification analysis. *Spine (Phila Pa 1976)*, 2012. 37(8): p. E486-95.
- 17. Asell M, Sjolander P, Kerschbaumer H, et al., Are lumbar repositioning errors larger among patients with chronic low back pain compared with asymptomatic subjects? *Arch Phys Med Rehabil*, 2006. 87(9): p. 1170-6.
- 18. Georgy EE, Lumbar repositioning accuracy as a measure of proprioception in patients with back dysfunction and healthy controls. *Asian Spine J*, 2011. 5(4): p. 201-7.
- 19. Koumantakis GA, Winstanley JOldham JA, Thoracolumbar proprioception in individuals with and without low back pain: Intratester reliability, clinical applicability, and validity. *J Orthop Sports Phys Ther*, 2002. 32(7): p. 327-35.
- 20. Descarreaux M, Blouin JSTeasdale N, Repositioning accuracy and movement parameters in low back pain subjects and healthy control subjects. *Eur Spine J*, 2005. 14(2): p. 185-91.
- 21. Newcomer K, Laskowski ER, Yu B, et al., Repositioning error in low back pain. Comparing trunk repositioning error in subjects with chronic low back pain and control subjects. *Spine (Phila Pa 1976)*, 2000. 25(2): p. 245-50.

- 22. Newcomer KL, Laskowski ER, Yu B, et al., Differences in repositioning error among patients with low back pain compared with control subjects. *Spine (Phila Pa 1976)*, 2000. 25(19): p. 2488-93.
- 23. Gill KP Callaghan MJ, The measurement of lumbar proprioception in individuals with and without low back pain. *Spine (Phila Pa 1976)*, 1998. 23(3): p. 371-7.
- 24. Lam SS, Jull GTreleaven J, Lumbar spine kinesthesia in patients with low back pain. *J Orthop Sports Phys Ther*, 1999. 29(5): p. 294-9.
- 25. Maffey-Ward L, Jull GWellington L, Toward a clinical test of lumbar spine kinesthesia. *J Orthop Sports Phys Ther*, 1996. 24(6): p. 354-8.
- 26. Sheeran L, van Deursen R, Caterson B, et al., Classification-guided versus generalized postural intervention in subgroups of non-specific chronic low back pain: A randomized study. *Spine (Phila Pa 1976)*, 2013.
- 27. (NDLTD) EWGotNDLoTaD, *Dart europe e-thesis portal*. http://www.darteurope.eu/basic-search.php.
- 28. Swiss National Libary, Swiss national libary. http://www.nb.admin.ch/?lang=en.
- 29. Europe L, Open grey literature europe. http://www.opengrey.eu/.
- 30. Liberati A, Altman DG, Tetzlaff J, et al., The prisma statement for reporting systematic reviews and meta-analyses of studies that evaluate health care interventions: Explanation and elaboration. *J Clin Epidemiol*, 2009. 62(10): p. e1-34.
- 31. Moher D, Liberati A, Tetzlaff J, et al., Preferred reporting items for systematic reviews and meta-analyses: The prisma statement. *J Clin Epidemiol*, 2009. 62(10): p. 1006-12.
- 32. Vandenbroucke JP, von Elm E, Altman DG, et al., Strengthening the reporting of observational studies in epidemiology (strobe): Explanation and elaboration. *Ann Intern Med*, 2007. 147(8): p. W163-94.
- 33. von Elm E, Altman DG, Egger M, et al., The strengthening the reporting of observational studies in epidemiology (strobe) statement: Guidelines for reporting observational studies. *Ann Intern Med*, 2007. 147(8): p. 573-7.
- 34. Borenstein M, Hedges LV, Higgins JPT, et al., Introduction to meta-analysis. 2011, Chichester, West Sussex, UK: John Wiley & Sons Ldt.
- 35. Dankaerts W O'Sullivan P, The validity of o'sullivan's classification system (cs) for a sub-group of ns-clbp with motor control impairment (mci): Overview of a series of studies and review of the literature. *Manual Therapy*, 2011. 16(1): p. 9-14.
- 36. Hill JC, Dunn KM, Lewis M, et al., A primary care back pain screening tool: Identifying patient subgroups for initial treatment. *Arthritis Rheum*, 2008. 59(5): p. 632-41.
- 37. Linton SJ Boersma K, Early identification of patients at risk of developing a persistent back problem: The predictive validity of the orebro musculoskeletal pain questionnaire. *Clin J Pain*, 2003. 19(2): p. 80-6.
- 38. Waddell G, Newton M, Henderson I, et al., A fear-avoidance beliefs questionnaire (fabq) and the role of fear-avoidance beliefs in chronic low back pain and disability. *Pain*, 1993. 52(2): p. 157-68.
- 39. Ernst M, Rast F, Bauer C, et al., Determination of thoracic and lumbar spinal processes by their percentage position between c7 and the psis level. *BMC Research Notes*, 2013. 6(1): p. 58.
- 40. Larivière C, Mecheri H, Shahvarpour A, et al., Criterion validity and between-day reliability of an inertial-sensor-based trunk postural stability test during unstable sitting. *Journal of Electromyography and Kinesiology*, 2013. 23(4): p. 899-907.
- 41. Higgins JPT Green S. Cochrane handbook for systematic reviews of interventions. Version 5.1.0. 2011 [cited 2014 7th February 2014].
- 42. O'Sullivan K, O'Sullivan L, Campbell A, et al., Towards monitoring lumbo-pelvic posture in real-life situations: Concurrent validity of a novel posture monitor and a traditional laboratory-based motion analysis system. *Manual Therapy*, 2012. 17(1): p. 77-83.
- 43. Sheeran L, Sparkes V, Busse M, et al., Preliminary study: Reliability of the spinal wheel. A novel device to measure spinal postures applied to sitting and standing. *European Spine Journal*, 2010. 19(6): p. 995-1003.

Origin		NS(	NSCLBP	Crit	Criteria		-	Healthy controls	()
Study	⊆	m/f	Age (y)	Inclusion	Exclusion	۲	m/f	Age (y)	matching
				NSCLBP >3months,18-65y,	Previous back surgery, neurologic				
O'Sullivan				increasing symptoms during	symptoms, ear/visual disturbance,				Age,
2013 <sup>(15)</sup>	15	10/5	31.3 ±10.3	prolonged sitting, reduced	red flags, pregnancy/<6months	15	10/5	32.1 ±9.2	gender,
0 07				symptoms during standing and	post-partum				BMI
				walking					
				NSCLBP >3months, 14-16y, MBI<	Specific diagnosis, previous back				Age,
				28kg/m, mechanically induced pain	surgery, neurologic symptoms,				gender,
Actfolck				in area between T12 to gluteal folds,	pelvic or abdominal pain, lower limb				pubertal
2012 <sup>(14)</sup>	28	14/14	15.7 ±0.5	no peripheral pain referral,	surgery/current injury, pregnancy/<6	28	14/14	15.4 ±0.5	stage,
6102				moderate ongoing LBP (NRS >3,	months post-partum, not English				socio-
				most days of the week)	speaking, inability to assume test				economic
					posture				status
				LBP >3months, mechanical basis of	Red flags, yellow flags,				
				disorder, motor control impairment	pregnancy/breastfeeding, revious				
Cheeron				(flexion/active extension pattern)	back surgery,				
опсенан, 2012 <sup>(16)</sup>	06	31/59	35±10.9		ear/vestibular/neurologic	35	13/22	36.0 ±10.3	ı
7 0 7					dysfunction affecting balance, not				
					able to sit or stand up from a stool				
					unaided				
				LBP >3months, mechanical	Previous inner ear infection or				
				dysfunction, NRS >5, lumbar ROM	vestibular disorder with balance				
Georgy,	ע ע	ç	101+61	of at least 50% of normal range	disturbance, history of head trauma	и 7	ç	38 5 ±5 0	
2011 <sup>(18)</sup>	2				with residual neurological deficits,	2		0.00	
					metabolic diseases,				
					pregnancy/breastfeeding, spinal				

## Table 1: Study design and subjects.

					surgery, severe back pain				
				LBP >6months	Diabetes, psychiatric diagnoses,				
Asell,	0	15/17	38 47		neurologic and rheumatic disorders,	, ,	16/1E	36 +0	Age,
2006 (17)	36	40,41	17 00		dizziness, vestibular disease,			00 H 30	gender
					surgery last 3 months				
				NSLBP >6months	Spondylolosthesis, Spondylolysis,				
					ankylosis spondylitis, osteoarthritis,				
					inflamatory arthritis, nerve root				
Descarreaux,	4	11/5	4 4 4		compression, trunk neuromuscular	ц Т	0/6	38 7	I
2005 <sup>(20)</sup>	2		+ -		disease, scoliosis (>15°), previous	2	310	7.00	I
					spinal surgery, malignant tumour,				
					hypertension,				
					pregnancy/breastfeeding				
				Recurrent LBP >3months, diagnosis	Neurologic involvement, recent back				
O'Shillivan				of lumbar segmental instability	surgery, pain preventing the test,				Age,
	15	6/9	38.8 ±12	flexion pattern	recent motor control rehabilitation,	15	6/9	38.2 ±10.9	height,
0007					ear/visual disturbance, severe soft				weight
					tissue tightness around hip/trunk				
				Recurrent mechanical NSCLBP with	trunk or lower limb pathology,				
Koumontokie				at least 2 episodes within the past	deformity, or condition that may				
2002 <sup>(19)</sup>	62	30/32	38.2 ±10.7	year with pain duration of less than	affect motor control	18	8/10	24.6 ±4.0	ı
2002				half the days in the past year, still					
				working, no neurological condition					
				Mechanical NSCLBP	Recent history of inner ear infection				
Brinsdae					with associated balance or				
	23	7/16	21.8 ±2.1		coordination problem, history of	21	6/15	22.3 ±3.8	I
0007					cerebranl trauma with unresolved				
					neurosensory symptoms, vestibular				
							1		

					disorder, previous spinal surgery, specific balance or stabilization				
					training in the last 6 months, pain				
					medication				
				NSCLBP (mechanical, nonradicular)	Severe pain preventing cooperation				
				≥3months, ROM of at least 50% of	with the study, pregnancy and				
				normal value	lactation, previous back surgery,				
Newcomer,		0110	V 117 C UC		current lower extremity problems,		0110	0 1 1 1 1 0 0	
2000 <sup>(21)</sup>	70	0/ 17	4.11 E C.80		radiculopathy, vertebral	N Z	<u>c</u>	09.1 H I I.O	
					compression fracture, neurologic				
					deficit, symptoms of vertigo or				
					dizziness				
				Pain between L1 and the gluteal	Severe pain preventing cooperation				
				folds ≥6months, average pain level	with the study, pregnancy and				
				of 5 of 10 in the preceding week,	lactation, previous back surgery,				
				ROM of at least 50% of normal	scoliotic curvature greater than 15°,				
2000 <sup>(22)</sup>	20	9/11	44.2 ±10.6	value	neurologic or current lower	20	9/11	39.8 ±12.7	
0007					extremity problems, lumbar				
					radiculopathy, vertebral				
					compression fracture, symptoms of				
					vertigo or dizziness				
1 am 1000 <sup>(24)</sup> &				Mechanical back pain ≥3months	Back pain from a				
Moffor Mord	ĊĊ	0/ 7 7	31.00		nonmusculoskeletal pathology,	0	5/5	сс	
малеу-үүаги, 1006 <sup>(25)</sup>	70				neurologic involvement, previous	2	0	04	
000					surgery in back/abdomen/chest				
Ē				Chronic mechanical low back pain	Neurologic deficit, psychological				
1008 <sup>(23)</sup>	20	7/13	43.3	>12months	component, further medical	20	7/13	32.9	gender
000					problems, nerve root pain				

Table 1: Study design and subjects. NSCLBP=non-specific chronic low back pain, LBP=low back pain, n=number of patients, m/f =male/female, BMI=body mass index, y=years, ROM=range of motion

Table 2: test procedure and instrumentation

Study	Movement task <sup>a</sup>	Measurement details	EO/EC	Instrument (I), Sensor position (SP)
O'Sullivan, 2013 <sup>(15)</sup>	P: Sitting, warming up by performing max trunk flex/ex, 1 practice trial IP: Sitting (90°hips, knees, ankles), arms supinated on thighs, neutral lumbo-pelvic spinal posture, (maintained 5 s) M: Slumped position (maintained 5 s) TP: Initial position (maintained 5 s)	n: 3 rest (s): ? feedback <sup>b</sup> : undergarments, shorts feedback <sup>c</sup> : no		l: "Body Guard" (Sels Instruments, Belgium) SP: L3, S2
Astfalck, 2013 <sup>(14)</sup>	P: Sitting, 3x ROM, 2 practice trials IP: Sitting (90° hips and knees), arms supinated on tighs, mid-range sitting posture position (maintained 5 s) M: Slumped position (maintained 5 s) TP: Initial position	n: 3 rest (s): ? feedback <sup>b</sup> : undergarments, shorts feedback <sup>c</sup> : no	EC	I : "Fastrak" (Polhemus Navigation Sciences Division, Vermont, USA) SP : L3, S2
Sheeran, 2012 <sup>(16)</sup>	P: Sitting/standing, 3x ROM IP: 1) Sitting, arms loose on thigh; 2) Standing, feed shoulder width apart, neutral lumbar and thoracic mid-range position (maintained 5 s) M: 1) Relaxed usual sitting (maintained 5 s); 2) Relaxed usual standing TP: Initial position	n: 4 rest (s): ? feedback <sup>b</sup> : loose clothing feedback <sup>c</sup> : no	С Ш	l: Vicon 512 (Vicon Motion Systems Ltd, Oxford, UK) SP: T12, S1
Georgy, 2011 <sup>(18)</sup>	P: Sitting, stabilized by straps, 3 practical trials IP: Sitting, passively moved to 30° of lumbar flexion (maintained 10 s) M: Upright neutral sitting TP: 30° lumbar flexion (maintained 3 s)	n : 3 rest (s) : 10 feedback <sup>b</sup> : ? feedback <sup>c</sup> : no		I: Biodex System 3 Pro Isokinetic Dynamometer (Biodex Medical Inc., Shirley, New York, USA) SP: Axis of actuator arm with L5/S1
Asell, 2006 <sup>(17)</sup>	P: Sitting, 2x sit-to-stand, 2x ROM, 6 practical trials (3 verbally, 3 pre- recorded instructions) IP: Sitting, hips and knees at 90°, guarded to the target position (maintained 2 s)	n: 10 rest (s): 3 feedback <sup>b</sup> : undergarments, hair in a bun, boldered armpits. No drinking	EC	l : "Fastrak" (Polhemus Navigation Sciences Division, Vermont, USA) SP : T7, S2, midpoint between

	M: Lumbar flexion until auditory signal (90% of max flex S2)	or eating 2h prior to testing		those 2 segments
	TP: 1/3 of the way towards maximal extension from the subjects normal	feedback <sup>c</sup> : no		
	sitting position, verbal signal by subject			
	P: Standing, Max ROM, learning phase with visual accuracy feedback			-
Descarreaux,	till 5 consecutive trunk positioning within 10% margin	n: 10 (a 5 s)		I: Loredan (Loredan Biomedical,
2005 (20)	IP: Neutral (0° flex or ex), pelvis and legs immobilised	rest (s): ?	·	West Sacramento, USA)
0	M: Flexion (15°, 30°, 60°), Extension (15°), randomised	feedback <sup>c</sup> : no		SP: ?
	TP: Flexion (15°, 30°, 60°), Extension (15°), randomised			
	P: Sitting, 3 x ROM	در 2		l · "Fastrak" (Polhemus
O'Sullivan	IP: Sitting (90°hips, knees, ankles), arms relaxed on thighs, neutral	rest (s): 2		Navidation Sciences Division
2003 <sup>(12)</sup>	spine posture (maintained 5 s)	feedback <sup>b</sup> . Undergarments shorts	C E	Vermont 11SA)
0007	M: Full lumbar flexion (maintained 5 s)			
	TP: Initial position	ICCODACK . ILO		or . I Iz, Lz, L4, Jz
	P: Standing, practicing with visual feedback	n : 3 within 30s		I: Lumbar Motion Monitor (LMM,
Koumantakis,	IP: Standing, hip leaning against a bench	foothool <sup>b</sup> , loono alothing	Ċ	Chattecx Corp., Chattanooga,
2002 <sup>(19)</sup>	M : Flexion, rotation, side-flexion	herden vouse cloumig.	) L	TN, USA)
	TP : 20° Flexion,15° rotation, 15° side-flexion	feedback <sup>c</sup> : no		SP:?
	P: Standing, 10 x pelvic tilt to warm up, ROM pelvic tilt	n: 5		
Brumagne,	IP: criterion position varying around neutral (maintained 5 s)	rest (s): ?		I: electrogoniometer
2000 <sup>(11)</sup>	M: Anterior pelvic tilt	feedback": shorts	ı	SP: ?
	TP: Criterion position	feedback*: no		
	P: Standing			
	IP: Standing, feet at shoulder's width apart and arms at side,1) neutral;			I : "Fastrak" (Polhemus
Newcomer,	2) 50% max ROM of flexion, extension, rotation, side-flexion	2	EO/EC	Navigation Sciences Division,
2000 <sup>(21)</sup>	M: 1) flexion, extension, rotation, side-flexion; 2) to neutral			Vermont, USA)
	TP: 1) neutral position (5 s to move to desired position, maintained 2 s)			SP : L1, S1
	2) 50% of max ROM of Flexion, extension, rotation, lateral-flexion (5 s			
		_		

	to move to desired position, maintained 2 s)			
	P: Standing, feet shoulder-width apart, arms at side, lower extremity			
	and pelvic immobilized, ROM			I : "Fastrak" (Polhemus
Newcomer,	IP: Standing, feet shoulder-width apart, arms at side, lower extremity	ç	C L	Navigation Sciences Division,
2000 <sup>(22)</sup>	and pelvic immobilized, neutral	••	2	Vermont, USA)
	M: Flexion, extension, side-flexion (5 s to move to desired position)			SP : T1, S1
	TP: 30%, 60%, 90% of max ROM (maintained for 2 s)			
	P: Cycling (5 minutes), ROM, 5 practice trials	n: 3		I : "Fastrak" (Polhemus
Moffow Mord	IP: Sitting with hips and knees 90°, neutral upright posture	rest (s): 15	C L	Navigation Science Division,
1006 <sup>(25)</sup>	M: Full lumbar flexion (maintained 3 s)	feedback: shorts, undergarments,	2	Vermont, USA)
0661	TP: Initial position	no drinking or eating 2h prior testing		SP : T10, S2
	P: 10 practical trails with visual feedback from screen			I: Lumbar Motion Monitor (LMM,
Ē	IP: 1) Standing: arms crossed; 2) Four-point-kneeling: 90° of hips,	n: 10 within 30 s		Chattecx Corp., Chattanooga,
1000 <sup>(23)</sup>	knees, shoulders	rest (s): ?	EC	TN, USA)
0661	M: Lumbar flexion	feedback <sup>b</sup> : loose clothing		SP : Harness, inferior binding
	TP: lumbar flexion 20°			posts level of T7
Table 2: test pro	Table 2: test procedure and instrumentation. <sup>a</sup> P=Preparation, IP= Initial Position, M=Movement, TP= Target Position, <sup>b</sup> sensory feedback (clothing, organs), <sup>c</sup> acustic or verbal	ement, TP= Target Position, <sup>b</sup> sensory feedback (clothing,	eedback (	clothing, organs), <sup>c</sup> acustic or verbal

feedback during measurements. S=seconds, EO/CE=eyes open/eyes closed, C=cervical, T=thoracic, S=sacral, max=maximal, ROM= Range of Motion, n= number of

trials

Absolute error						COLIFICOIS			Effect size		
Absolute error	position	Direction									
			mean	SD	۲	mean	SD	<u>ح</u>	Hedges g/ Cohens r*	95%CI LL	95%CI UL
O'Sullivan, 2013 <sup>(13)</sup>	sitting	flexion	11.5	6.4	15	5.1	3.6	15	1.20	0.41	1.99
Astfalck, 2013 <sup>(14)</sup>	sitting	flexion	4.1	2.3	28	3.1	1.3	28	0.53	-0.01	1.06
Sheeran, 2012 <sup>(16)</sup>	sitting	flexion	7.7	4.1	06	1.8	ø.	35	1.67	1.23	2.11
Georgy, 2011 <sup>(18)</sup>	sitting	extension	7.5	3.3	15	2.8	<u>б</u>	15	1.88	1.04	2.72
O'Sullivan, 2003 (12)	sitting	flexion	1.7	ø.	15	1.1	9.	15	0.83	0.08	1.58
Lam/Maffey <sup>(24, 25)</sup>	sitting	flexion	2.3	6.	20	2.6	1.2	10	-0.29	-1.05	0.47
Gill 1998 <sup>(23)</sup>	standing	flexion	6.7	5.0	20	4.5	3.4	20	.26	-0.12	1.20
Sheeran, 2012 <sup>(16)</sup>	standing	flexion	6.3	3.7	06	1.9	1.3	35	1.67	1.23	2.11
Koumantakis 2002 (19)	standing	flexion	5.5	3.5	62	3.7	1.8	18	0.55	0.03	1.08
Brumagne, 2000 <sup>(11)</sup>	standing	extension	4.3	~	23	1.6	9.	21	3.18	2.30	4.06
<b>Constant error</b>											
O'Sullivan, 2013 <sup>(15)</sup>	sitting	flexion	-6.9	11.5	15	2.6	5.0	15	-1.04	-1.79	-0.30
Astfalck, 2013 <sup>(14)</sup>	sitting	flexion	-	4.2	28	8 <sup>.</sup> -	2.6	28	0.20	-0.32	0.72
Sheeran, 2012 <sup>(16)</sup>	sitting	flexion	<u>ю</u>	7.7	06	2	1.1	35	0.11	-0.28	0.49
Brumagne, 2000 <sup>(11)</sup>	standing	extension	-2.5	2.5	23	9	1.0	21	-0.96	-1.58	-0.35
Sheeran, 2012 <sup>(16)</sup>	standing	flexion	-1.9	5.2	06	5	0.9	35	-0.31	-0.70	0.08
Variable error											
O'Sullivan, 2013 <sup>(15)</sup>	sitting	flexion	4.3	2.4	15	3.6	2.9	15	0.25*	-0.44	0.95
Astfalck, 2013 <sup>(14)</sup>	sitting	flexion	3.4	2.1	28	2.8	1.6	28	0.32	-0.21	0.84
Sheeran, 2012 <sup>(16)</sup>	sitting	flexion	4.2	2.6	06	1.9	~	35	1.01	0.60	1.42
Koumantakis 2002	standing	flexion	2.2	1.6	62	1.7	1.0	18	0.33	-0.19	0.86

Table 3: outcomes and effect size measures

Brumagne, 2000 ( <sup>III)</sup>	standing	extension	3.3	1.4	23	1.7	0.7	21	1.40	0.75	2.05
Sheeran, 2012 <sup>(16)</sup>	standing	flexion	4.2	2.4	06	1.8	1.2	35	1.12	0.70	1.53
Table 3: outcomes and effect size mean	nd effect size	measures * indicates that data was non-normally distributed and Cohens r was calculated as effect size measu	s that data	was noi	n-normal	ly distribu	ited and	Cohens I	- was calcula	ated as effect si	ze measure.

ent based on STROBE <sup>(32, 33)</sup>	
Table 4. Study quality assessme	NC+11-dv

Sheeran <sup>(16)</sup>	٦	0		-	-		0	0	-	-	0	-	-	-	-	٢	-	0
O'Sullivan <sup>(12)</sup>	-	-		۲	1		٢	0	٦	٦	0	1	1	0	0	٢	0	0
<sup>(35)</sup> nevillu2'O	0	٦		٢	1		0	0	٢	٢	1	1	٢	0	0	٢	0	0
Newcromer <sup>(22)</sup>	0	0		1	1		0	0	٦	0	1	1	0	0	1	٢	1	0
Newcomer <sup>(21)</sup>	0	0		1	0		0	0	٦	0	0	1	0	0	1	٢	1	0
Maffey-Ward <sup>(25)</sup>	0	٦		٢	1		0	0	٢	0	0	1	٢	0	0	٢	0	0
רשש <sub>(גל)</sub>	1	1		1	1		0	0	1	0	1	1	1	0	0	1	0	0
<sup>(01)</sup> si≯lanemuoX	٢	-		-	-		0	0	-	0	-	-	-	0	-	-	0	0
Cill ( <sub>S3)</sub>	٢	0		1	1		0	0	1	0	0	1	1	0	0	0	0	0
Georgy <sup>(18)</sup>	١	٦		1	1		1	0	٦	0	0	1	٦	0	0	٢	1	0
Descarreaux <sup>(20)</sup>	٢	0		1	1		0	0	1	0	1	1	0	0	1	1	1	0
Brumagne <sup>(11)</sup>	-	-		۲	1		0	0	٦	0	0	1	1	0	0	٢	0	0
Astfalck <sup>(14)</sup>	0	-		-	1		0	-	-	-	٢	0	٢	۲	-	-	-	0
<sup>(\t)</sup> IIəɛA	١	0		٢	1		٦	0	٢	٢	1	1	٢	0	٢	٢	1	0
Study	1a	1b		2	3		4	5	6a	6b	7	8	9	10	11	12a	12b	12c
STROBE <sup>(33)</sup>	Title & Abstract		Introduction	Background	Objectives	Methods	Study Design	Setting	Participants		Variables	Data Sources	Bias	Study Size	Quantitative variables	Statistical Methods		

<b></b>	[		[			[		[	[	[	r			[	[	[	[	r
0	0		~	0	0	~	0	~	0	0	0		~	~	0	~	~	-
0	0		0	0	0	~	0	0	0	0	0		0	~	~	~	~	0
0	0		0	0	0	~	0	~	~	0	0		0	~	~	~	~	-
0	0		0	0	0	1	0	0	0	0	0		1	1	1	~	1	0
0	0		0	0	0	-	0	0	-	0	0		٢	-	0	0	-	-
0	0		0	0	0	0	0	~	0	0	0		0	0	0	~	-	0
0	0		0	0	0	0	0	~	0	0	0		0	-	0	~	-	0
0	0		0	0	0	-	0	~	-	0	0		٢	0	0	~	-	~
0	0		0	0	0	0	0	٦	0	0	0		-	-	0	٦	-	~
0	0		0	0	0	Ļ	0	~	0	0	0		٢	Ļ	-	~	-	0
0	0		-	0	0	-	0	٦	0	0	0		-	-	0	٦	-	~
0	0		0	0	0	-	0	٦	0	0	0		0	-	0	٦	-	0
-	0		-	-	-	-	0	-	-	0	0		-	-	-	-	-	~
0	0		0	0	0	-	0	0	0	0	0		-	-	-	-	-	~
12d	12e		13a	13b	13c	14a	14b	15	16a	16b	16c		17	18	19	20	21	22
		Results	Participants			Descriptive Data		Outcome Data	Main Results			Discussion	Other Analysis	Key Result	Limitation	Interpretation y	Generalizabilit	Other Information (Funding)

### Ľ

Table 5. Reliability and Measurement Error			
Author	Reliability	Measurement error	Conclusion
Koumantakis 2002 <sup>(19)</sup>	NSCLBP: all RE-tests ICC= 0.24 to 0.64	NSCLBP: SEM= 0.45° to 1.34° (large)	Low ICC and high SEM The reliability is low in patients with LBP
	AE for flexion and rotation: ICC= 0.76 to HC: SEM= 0.45° to 3.90° 0.80	HC: SEM= 0.45° to 3.90°	-
	Other RE-tests: ICC = 0.2 to 0.69		
Asell 2006 (17)	Only tested in HC and with a slightly		Reliability is acceptable
	modified of the sitting pelvic test		
	VE: ICC= 0.75		
	CE: ICC =0.86		
Descarreaux 2005 <sup>(20)</sup>	Not specified	Not specified	
Astfalck 2013 <sup>(14)</sup>	Refer to Maffey-Ward 1996 & Lam 1999.		This task has previously been shown to
	(24, 25)		have good reliability in adults both with and
			without LBP (24, 20)
Newcomer 2000a <sup>(21)</sup>		SEMean= 0.48°	

Newcomer 2000b (22)		SEMean= 0.27°	SEMean decreased compared to the
			previous study
Lam 1999 <sup>(24)</sup>	No difference in error magnitude between	No difference in error magnitude	Suggest that either the study group did not
	days	between days	have kinaesthetic deficits associated with
			their condition or that the repositioning test
			in the sitting position lacks sensitivity
Georgy, 2011 <sup>(18)</sup>	Not specified	Not specified	
O'Sullivan 2003 <sup>(12)</sup>	Reliability is only indicated for the	Measurement error is only indicated for	Reliability and Measurement Error are not
	measurement device.	the measurement device.	specified for the testing protocol.
O'Sullivan 2013 <sup>(15)</sup>	ICC > 0.80 for the measurement device (42)	error for the	This device has been shown to have very
		measurement device <sup>(42)</sup>	good reliability and measurement error for
			the measurement of lumbo-pelvic posture.
Sheeran et al., 2012 <sup>(16)</sup>	Reliability is only indicated for the		
	measurement device (spinal wheel ICC=		
	0.95-0.98) (43)		
Table 5. Reliability and Measurement Erro	5. Reliability and Measurement Error NSCI BP= Nonspecific Chronic Low Back Pain RE=Reposition Error AF=Absolute Error CF=Constant Error VE=Variable Error	Pain RF=Renosition Frror AF=Absolute F	rror CF=Constant Frror VF=Variable Frror

Iable 5. Reliability and Measurement Error. NSCLBP= Nonspecific Chronic Low Back Pain, KE=Reposition Error, AE=Bosolute Error, CE=Constant Error, VE=Variable Error, HC= Healthy Controls, ICC= Intraclass Correlation Coefficient, SEM= Standard Error of the Measurement, SEMean= Standard Error of the Mean; LBP= Low Back Pain

Reposition error in LBP

### Figure legends

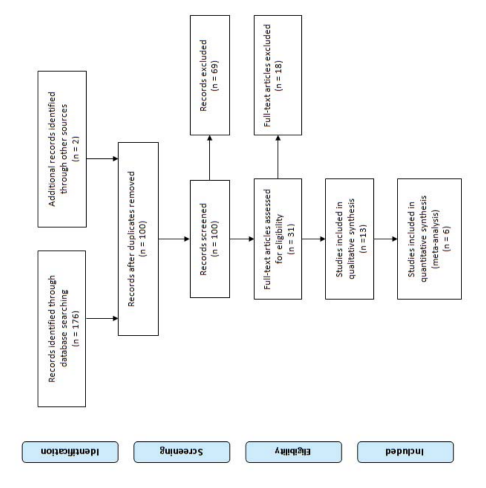
Figure 1. Flow chart according to PRISMA.

*Figure 2.* Forrest Plot showing the results of the meta-analysis of Absolute Error (AE) subgrouped for adults and adolescents. The overall effect size of 0.81 [Cl 0.13-1.49] picture that patients with unspecific low back pain (LBP) have a larger absolute error than healthy controls.

*Figure 3.* Forrest plot showing the results of the meta-analysis of Variable Error (VE) subgrouped for adults and adolescents. The overall mean difference of 0.57 [CI 0.05-1.09] illustrate that patients with unspecific low back pain (LBP) have a higher deviation of reposition error than healthy controls.

*Figure 4 and 5.* Forrest Plots showing the results of a meta-analysis on constant error (CE) subgrouped for adults and adolescents. The overall mean difference CE for FP is -0.39 [CI - 1.09-0.3] indicates that FP NSCLBP patients undershoot into flexion,. The overall mean difference CE for AEP is 0.18 [CI -.3-0.65] indicates that AEP NSCLBP patients overshoot into extension.

### PRISMA 2009 Flow Diagram



From: Moher D, Liberati A, Tetzlaff J, Altman DG, The PRISMA Group (2009). Preterned Reporting items for Systematic Reviews and Meta-Analyses: The PRISMA Statement. PLoSS Med 6(6): e1000097. doi:10.1371/journal.pmed1000097

Analyses: The PRISMA Statement. PLOS Med 6(6): e1000097. doi:10.1371/journal.pmed100 For more information, visit <u>www.prisma-statement.org</u>.

## Figure 2 (AE)

	NSC	<b>n</b>		ů :	Control			Std. Mean Difference	Std. Mean Difference
Study or Subgroup 2.1.1 Adults	Mean SU		lotal	Mean	R	lota	weight	lotal Mean SD lotal weight IV, Kandom, 95% CI	IV, Kandom, 95% CI
Sheeran 2012	7.7	1.1	90	1.8	0.8	35	22.3%	1.67 [1.23, 2.11]	ŧ
0´Sullivan 2003	1.7 0.8	8.(	15	1.1	0.6	15	19.0%	0.83 [0.08, 1.58]	ł
Lam/Maffey-Ward 1999/1996	2.3 0.9	6.0	20	2.6	1.2	10	18.8%	-0.29 [-1.05, 0.47]	ŧ
0 Sullivan 2013 Subtotal (95% CI)	11.5 6.4	4.0	15 140	5.1	3.6	15 75	18.5% <b>78.6</b> %	1.20 [0.41, 1.99] 0.88 [0.02, 1.73]	∮∳
Heterogeneity: Tau <sup>2</sup> = 0.64; Chi <sup>2</sup> = 19.64, df = 3 (P = 0.0002); l <sup>2</sup> = 85% Test for overall effect: Z = 2.01 (P = 0.04)	= 19.64 P = 0.04	, df	= 3 (P :	= 0.00(	02); l <sup>i</sup>	<sup>2</sup> = 859	9		•
2.1.2 Adolescents									
Astfalck 2013 Subtotal (95% CI)	4.1 2.3	č.	28 28	3.1	1.3	28 28	21.4% <b>21.4</b> %	0.53 [-0.01, 1.06] 0.53 [-0.01, 1.06]	+ ♦
Heterogeneity: Not applicable Test for overall effect: Z = 1.94 (P = 0.05)	(P = 0.05)	-							
Total (95% CI)			168			103	103 100.0%	0.81 [0.13, 1.49]	•
Heterogeneity: Tau <sup>2</sup> = 0.48; Chi <sup>2</sup> = 23.13, df = 4 (P = 0.0001); l <sup>2</sup> = 83% Test for overall effect: Z = 2.34 (P = 0.02) Test for subgroup differences: Chi <sup>2</sup> = 0.47, df = 1 (P = 0.49), l <sup>2</sup> = 0%	r = 23.13 (P = 0.02) hi <sup>2</sup> = 0.47	, df	= 4 (P : = 1 (P	= 0.00( = 0.49	01); ľ	² = 83% = 0%	9	Τ.	Control NSCLBP

## Figure 3 (VE)

	NSC	NSCLBP		ű	Control		•.	Std. Mean Difference	Std. Mean Difference	Difference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Mean SD Total Mean SD Total Weight	IV, Random, 95% CI	IV, Random, 95% CI	m, 95% CI	
3.1.1 Adults											
Sheeran 2012	4.2 2.6	2.6	6	1.9	-	35	53.0%	1.01 [0.60, 1.42]		+	
Subtotal (95% CI)			06			35	53.0%	1.01 [0.60, 1.42]		•	
Heterogeneity: Not applicable	plicable										
Test for overall effect: $Z = 4.80 (P < 0.00001)$	Z= 4.80	Ē	0.000.0	Ê							
3.1.2 Adolescents											
Astfalck 2013	3.4 2.1	2.1	28	2.8	2.8 1.6	28	47.0%	0.32 [-0.21, 0.84]	1	<b>1</b>	
Subtotal (95% CI)			28			28	47.0%	0.32 [-0.21, 0.84]	•	•	
Heterogeneity: Not applicable	plicable										
Test for overall effect: Z= 1.18 (P = 0.24)	Z= 1.18	Е.	0.24)								
Total (95% CI)			118			63	63 100.0%	0.68 [0.01, 1.36]		♦	
Heterogeneity: Tau <sup>2</sup> = 0.18; Chi <sup>2</sup> = 4.08, df = 1 (P = 0.04); P = 75%	: 0.18; Ch	1 1 1 1	.08, df	= 1 (P	0.0	t); I <sup>2</sup> = 7	.5%				Ţ
Test for overall effect: Z = 1.98 (P = 0.05)	Z= 1.98	<u>–</u> –	0.05)					-	-4 -2 Control	U Z	4
Test for subgroup differences: $Chi^2 = 4.08$ , df = 1 (P = 0.04), P = 75.5%	erences:	Chi≊	= 4.08,	df= 1	Ē	0.04), F	= 75.5%				

## Figure 4 (CE FP)

Difference Std. Mean Difference	om, 95% CI IV, Random, 95% CI		.81, -0.27]					0.32 [-0.31, 0.95]			-0.39 [-1.09, 0.30]	<u>-4 -2 0 2 4</u>	NICOLDD
Std. Mean Difference	Mean SD Total Weight IV, Random, 95% CI		29.0% -1.04 [-1.81, -0.27]										
	Weight		29.0%	38.2%	<b>07.7%</b> = 26%			32.8% <b>32.8</b> %			78 100.0%	= 75%	
_	Total		15	35	24): l <sup>2</sup>			28 28			78	.02); I <sup>2</sup>	
Control	SD		5	-0.2 1.1	P = 0			-0.8 2.6				P = 0.	
ŭ			2.6	-0.2	f = 1 (			-0.8				f = 2 (	
	Total		15	51	35 d	0.005)		15 15		0.32)	81	7.95, d	1.11.0
NSCLBP	SD		11.5	-3 6.9	hi <sup>2</sup> = 1	0 (P =		0.3 4.5		= d) 6		$hi^2 = 7$	
NS	Mean		-6.9 11.5	°-	0.04: 0	Z = 2.8		0.3	plicable	Z = 0.9		= 0.28; C	T.T = 7
	Study or Subgroup Mean SD Total	4.1.1 Adults	O'Sullivan 2013 FP	Sheeran 2012 FP	Subtotal (95% Ct) $00^{-1}$ $00^{-1}$ $00^{-1}$ $00^{-1}$ $00^{-1}$ $00^{-1}$ $00^{-1}$ $00^{-1}$ $00^{-1}$ $00^{-1}$ $00^{-1}$	Test for overall effect: $Z = 2.80$ (P = 0.005)	4.1.2 Adolescents	Astfalck 2013 FP Subtotal (95% CI)	Heterogeneity: Not applicable	Test for overall effect: $Z = 0.99$ (P = 0.32)	Total (95% CI)	Heterogeneity: Tau <sup>2</sup> = 0.28; Chi <sup>2</sup> = 7.95, df = 2 (P = 0.02); l <sup>2</sup> = 75% Tast for sourcell officier $7 = 1.11$ (P = 0.27)	ובאר וחו האבומוו בווברוי

# Figure 5 (CE AEP)

ference Std. Mean Difference	, 95% CI IV, Random, 95% CI		8, 0.84]	10' 0'04]				5, 0.51]	'5, 0.51] ◆			0, 0.65]		NSCI RP undershoot NSCI RP overshoot	Number of the second of the se
Std. Mean Difference	Mean SD Total Weight IV, Random, 95% CI		0.38 [-0.08, 0.84]	n-n_1 oc-n				-0.12 [-0.75, 0.51]	-0.12 [-0.7			0.18 [-0.30, 0.65]	I		
0,	Weight		59.6%	0.0.60				40.4%	40.4%			63 100.0%	= 36%		
	Total		35	2				28	28			63	.21); l <sup>2</sup>		
Control	SD		1.1					4.2					(P = 0		
ŭ	Mean		-0.2 1.1					-0.1 4.2					lf = 1		
	Total		30 80	5		: 0.11)		15	15		: 0.71)	54	1.56, (	: 0.47)	
NSCLBP	S		2 7.9			0 (P =		3.9			7 (P =		hi² =	2 (P =	r
NS	Mean SD Total		2		plicable	: Z = 1.6		-0.6 3.9		plicable	: Z = 0.3		= 0.04; C	Z = 0.7	,
	Study or Subgroup	5.1.1 Adults	Sheeran 2012 AEP	Subtotal (23% CI)	Heterogeneity: Not applicable	Test for overall effect: $Z = 1.60$ (P = 0.11)	5.1.2 Adolescents	Astfalck 2013 AEP	Subtotal (95% CI)	Heterogeneity: Not applicable	Test for overall effect: $Z = 0.37 (P = 0.71)$	Total (95% CI)	Heterogeneity: Tau <sup>2</sup> = 0.04; Chi <sup>2</sup> = 1.56, df = 1 (P = 0.21); $I^2 = 36\%$	Test for overall effect: $Z = 0.72$ (P = 0.47)	