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# Manual therapy and exercise for rotator cuff disease (Review)

Page MJ, Green S, McBain B, Surace SJ, Deitch J, Lyttle N, Mrocki MA, Buchbinder R

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#### [Intervention Review]

# Manual therapy and exercise for rotator cuff disease

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### ABSTRACT

#### Background

Management of rotator cuff disease often includes manual therapy and exercise, usually delivered together as components of a physical therapy intervention. This review is one of a series of reviews that form an update of the Cochrane review, 'Physiotherapy interventions for shoulder pain'.

#### Objectives

To synthesise available evidence regarding the benefits and harms of manual therapy and exercise, alone or in combination, for the treatment of people with rotator cuff disease.

#### Search methods

We searched the Cochrane Central Register of Controlled Trials (CENTRAL; 2015, Issue 3), Ovid MEDLINE (January 1966 to March 2015), Ovid EMBASE (January 1980 to March 2015), CINAHL Plus (EBSCO, January 1937 to March 2015), ClinicalTrials.gov and the WHO ICTRP clinical trials registries up to March 2015, unrestricted by language, and reviewed the reference lists of review articles and retrieved trials, to identify potentially relevant trials.

#### **Selection criteria**

We included randomised and quasi-randomised trials, including adults with rotator cuff disease, and comparing any manual therapy or exercise intervention with placebo, no intervention, a different type of manual therapy or exercise or any other intervention (e.g. glucocorticoid injection). Interventions included mobilisation, manipulation and supervised or home exercises. Trials investigating the primary or add-on effect of manual therapy and exercise were the main comparisons of interest. Main outcomes of interest were overall pain, function, pain on motion, patient-reported global assessment of treatment success, quality of life and the number of participants experiencing adverse events.

#### Data collection and analysis

Two review authors independently selected trials for inclusion, extracted the data, performed a risk of bias assessment and assessed the quality of the body of evidence for the main outcomes using the GRADE approach.



#### **Main results**

We included 60 trials (3620 participants), although only 10 addressed the main comparisons of interest. Overall risk of bias was low in three, unclear in 14 and high in 43 trials. We were unable to perform any meta-analyses because of clinical heterogeneity or incomplete outcome reporting. One trial compared manual therapy and exercise with placebo (inactive ultrasound therapy) in 120 participants with chronic rotator cuff disease (high quality evidence). At 22 weeks, the mean change in overall pain with placebo was 17.3 points on a 100-point scale, and 24.8 points with manual therapy and exercise (adjusted mean difference (MD) 6.8 points, 95% confidence interval (Cl) -0.70 to 14.30 points; absolute risk difference 7%, 1% fewer to 14% more). Mean change in function with placebo was 15.6 points on a 100-point scale, and 22.4 points with manual therapy and exercise (adjusted MD 7.1 points, 95% Cl 0.30 to 13.90 points; absolute risk difference 7%, 1% fewer to 14% more). Fifty-seven per cent (31/54) of participants reported treatment success with manual therapy and exercise compared with 41% (24/58) of participants receiving placebo (risk ratio (RR) 1.39, 95% Cl 0.94 to 2.03; absolute risk difference 16% (2% fewer to 34% more). Thirty-one per cent (17/55) of participants reported adverse events with manual therapy and exercise compared with 8% (5/61) of participants receiving placebo (RR 3.77, 95% Cl 1.49 to 9.54; absolute risk difference 23% (9% to 37% more). However adverse events were mild (short-term pain following treatment).

Five trials (low quality evidence) found no important differences between manual therapy and exercise compared with glucocorticoid injection with respect to overall pain, function, active shoulder abduction and quality of life from four weeks up to 12 months. However, global treatment success was more common up to 11 weeks in people receiving glucocorticoid injection (low quality evidence). One trial (low quality evidence) showed no important differences between manual therapy and exercise and arthroscopic subacromial decompression with respect to overall pain, function, active range of motion and strength at six and 12 months, or global treatment success at four to eight years. One trial (low quality evidence) found that manual therapy and exercise may not be as effective as acupuncture plus dietary counselling and Phlogenzym supplement with respect to overall pain, function, active shoulder abduction and quality life at 12 weeks. We are uncertain whether manual therapy and exercise improves function more than oral non-steroidal anti-inflammatory drugs (NSAID), or whether combining manual therapy and exercise with glucocorticoid injection provides additional benefit in function over glucocorticoid injection alone, because of the very low quality evidence in these two trials.

Fifty-two trials investigated effects of manual therapy alone or exercise alone, and the evidence was mostly very low quality. There was little or no difference in patient-important outcomes between manual therapy alone and placebo, no treatment, therapeutic ultrasound and kinesiotaping, although manual therapy alone was less effective than glucocorticoid injection. Exercise alone led to less improvement in overall pain, but not function, when compared with surgical repair for rotator cuff tear. There was little or no difference in patient-important outcomes between exercise alone and placebo, radial extracorporeal shockwave treatment, glucocorticoid injection, arthroscopic subacromial decompression and functional brace. Further, manual therapy or exercise provided few or no additional benefits when combined with other physical therapy interventions, and one type of manual therapy or exercise was rarely more effective than another.

#### **Authors' conclusions**

Despite identifying 60 eligible trials, only one trial compared a combination of manual therapy and exercise reflective of common current practice to placebo. We judged it to be of high quality and found no clinically important differences between groups in any outcome. Effects of manual therapy and exercise may be similar to those of glucocorticoid injection and arthroscopic subacromial decompression, but this is based on low quality evidence. Adverse events associated with manual therapy and exercise are relatively more frequent than placebo but mild in nature. Novel combinations of manual therapy and exercise should be compared with a realistic placebo in future trials. Further trials of manual therapy alone or exercise alone for rotator cuff disease should be based upon a strong rationale and consideration of whether or not they would alter the conclusions of this review.

# PLAIN LANGUAGE SUMMARY

#### Manual therapy and exercise for rotator cuff disease

#### Background

Rotator cuff disease is a common cause of shoulder pain. People with rotator cuff disease often describe their pain as being worse at night and exacerbated by movement in specific directions including overhead activity. It is often associated with loss of function and some people describe weakness.

Manual therapy comprises movement of the joints and other structures by a healthcare professional (e.g. physiotherapist). Exercise includes any purposeful movement of a joint, muscle contraction or prescribed activity. The aims of both treatments are to relieve pain, increase strength and joint range, and improve function.

#### **Study characteristics**

This summary of an updated Cochrane review presents what we know from research about the benefits and harms of manual therapy and exercise compared with placebo, no intervention or any other intervention in people with rotator cuff disease. After searching for all relevant studies published up to March 2015, we included 60 trials (3620 participants), however only 10 looked at manual therapy



and exercise in combination. Among the included participants, 52% were women, average age was 51 years and average duration of the condition was 11 months. The average duration of manual therapy and exercise interventions was six weeks.

# Key results: one trial of manual therapy and exercise compared with placebo (inactive ultrasound therapy) for 10 weeks in people with chronic rotator cuff disease

#### Overall pain (higher scores mean more improvement in pain reduction)

People who had manual therapy and exercise had improvements in pain that were little or no different to people who had placebo. Improvement in pain was 6.8 points more (ranging from 0.7 points less to 14.3 points more) at 22 weeks (7% absolute improvement).

People who had manual therapy and exercise rated their change in pain score as 24.8 points on a scale of 0 to 100 points.

People who had placebo rated their change in pain score as 17.3 points on a scale of 0 to 100 points.

#### Function (higher scores mean more improvement in function)

People who had manual therapy and exercise improved slightly more than people who had placebo. Improvement in function was 7.1 points more (ranging from 0.3 to 13.9 points more) at 22 weeks (7% absolute improvement).

People who had manual therapy and exercise rated their change in function as 22.4 points on a scale of 0 to 100 points.

People who had placebo rated their change in function as 15.6 points on a scale of 0 to 100 points.

#### Treatment success

16 more people out of 100 rated their treatment as successful with manual therapy and exercise compared with placebo, 16% absolute improvement (ranging from 2% less to 34% more improvement).

Fifty-seven out of 100 people reported treatment success with manual therapy and exercise.

Forty-one out of 100 people reported treatment success with placebo.

#### Side effects

23 more people out of 100 people had minor side effects such as temporary pain after treatment with manual therapy and exercise compared with placebo.

Thirty-one out of 100 people reported side effects with manual therapy and exercise.

Eight out of 100 people reported side effects with placebo.

### **Quality of the evidence**

High quality evidence from one trial suggested that manual therapy and exercise improved function only slightly more than placebo at 22 weeks, was little or no different to placebo in terms of other patient-important outcomes (e.g. overall pain), and was associated with relatively more frequent but mild adverse events.

Low quality evidence suggested that there may be little or no difference in overall pain and function when manual therapy and exercise is compared with glucocorticoid injection, there may be little or no difference in overall pain and function when manual therapy and exercise is compared with arthroscopic subacromial decompression, and people who receive acupuncture plus dietary counselling and Phlogenzym supplement may have less pain and better function than people receiving manual therapy and exercise.

We are uncertain whether firstly, manual therapy and exercise improves function more than oral non-steroidal anti-inflammatory drugs (NSAID), and secondly, combining manual therapy and exercise with glucocorticoid injection provides additional improvement in function over glucocorticoid injection alone, because the quality of the evidence was very low.

# SUMMARY OF FINDINGS

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# Summary of findings for the main comparison. Manual therapy and exercise compared to placebo for rotator cuff disease

# Manual therapy and exercise compared to placebo for rotator cuff disease

Patient or population: rotator cuff disease

Settings: Public hospital physiotherapy units and private physiotherapy practices, Australia

Intervention: soft tissue massage, glenohumeral joint mobilisation, thoracic spine mobilisation, cervical spine mobilisation, scapular retraining, postural taping and supervised exercises in 10 sessions over 10 weeks along with home exercises for 22 weeks

**Comparison:** inactive ultrasound therapy and application of an inert gel in 10 sessions over 10 weeks

Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of Partici- pants	Quality of the evidence	Comments
	Assumed risk	Corresponding risk		(studies)	(GRADE)	
	Placebo	manual therapy and ex- ercise				
Overall pain Assessed with SPADI pain score Scale from 0-100 (higher score denotes less pain) Follow-up: 22 weeks	The mean im- provement in overall pain score in the control group was <b>17.3</b> <sup>1</sup>	The mean improvement in overall pain score in the intervention group was <b>6.8 points higher</b> (-0.7 lower to 14.3 higher)	-	120 (1 RCT)	⊕⊕⊕⊕ HIGH	Absolute risk difference 7% (1% fewer to 14% more); relative per- centage change 14% (1% fewer to 30% more) NNTB not applicable
Function Assessed with SPADI to- tal score Scale from 0-100 (high- er score denotes greater function) Follow-up: 22 weeks	The mean im- provement in function score in the control group was <b>15.6</b> <sup>1</sup>	The mean improvement in function score in the in- tervention group was <b>7.1</b> <b>points higher</b> (0.3 higher to 13.9 higher)	-	120 (1 RCT)	⊕⊕⊕⊕ HIGH	Absolute risk difference 7% (1% to 14% more); relative percentage change 16% (1% to 32% more) NNTB 6 (3 to 103)
Pain on motion Assessed with VAS Scale from 0-10 (higher score denotes less pain)	The mean im- provement in pain on motion score in the con- trol group was <b>1.6</b> <sup>1</sup>	The mean improvement in pain on motion score in the intervention group was <b>0.9 points higher</b> (-0.03 lower to 1.7 higher)	-	120 (1 RCT)	⊕⊕⊕⊕ HIGH	Absolute risk difference 9% (1% to 17% more); relative percent- age change 18% (1% fewer to 35% more) NNTB not applicable

Global assessment of treatment success - Follow-up: 22 weeks	· · ·		<b>RR 1.39</b> (0.94 to 2.03)	112 (1 RCT)	⊕⊕⊕⊕ HIGH	Absolute risk difference 16% (2% fewer to 34% more); relative per-
	<b>414 per 1000</b> <sup>2</sup> <b>575 per 1000</b> (393 to 840)		(0.54 to 2.05)		mon	centage change 39% (6% fewer to 103% more)
						NNTB not applicable
<b>Quality of life</b> Assessed with AQoL	The mean im- provement in quality of life	The mean improvement in quality of life score in the intervention group was	-	120 (1 RCT)	⊕⊕⊕⊕ HIGH	Absolute risk difference 5% (3% to 7% more); relative percentage change 10% (5% to 14% more)
Scale from -0.4 to 1 higher score denotes higher quality of life)	score in the con- trol group was 0 <sup>1</sup>	<b>0.07 points higher</b> (0.04 higher to 0.1 higher)				NNTB not applicable
Follow-up: 22 weeks						
<b>Adverse events</b> Follow-up: 11 weeks	Study population		<b>RR 3.77</b> (1.49 to 9.54)	116 (1 RCT)	⊕⊕⊕⊕ HIGH	Absolute risk difference 23% (9% to 37% more); relative percentage
		<b>809 per 1000</b> 122 to 782)				change 277% (49% to 854% more) NNTH 5 (26 to 2)
						Adverse events were mild, includ- ing short-term pain during or after treatment in the clinic, short-term pain after home exercises, or mild irritation with taping.
		n control group risk across stu roup and the <b>relative effect</b> o			e corresponding ri	<b>sk</b> (and its 95% confidence interval) is
	confident that the tru moderately confiden	e effect lies close to that of th t in the effect estimate: The tr ate is limited: The true effect r	ue effect is likely t may be substantia	o be close to the		

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# Summary of findings 2. Manual therapy and exercise compared to glucocorticoid injection for rotator cuff disease



### Manual therapy and exercise compared to glucocorticoid injection for rotator cuff disease

Patient or population: rotator cuff disease

Settings: Military hospital-based outpatient clinic, USA; Primary care (general practitioner), UK

Intervention: Either joint and soft tissue mobilisation, manual stretches and supervised and home exercises twice a week for three weeks or active and passive mobilisa-

tion, home exercises and therapeutic ultrasound once a week for six weeks

**Comparison:** glucocorticoid injection

Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of Partici- pants	Quality of the evidence	Comments	
	Assumed risk	Corresponding risk		(studies)	(GRADE)		
	Glucocorticoid injection	manual therapy and exercise					
<b>Overall pain</b> Assessed with NRS Scale from 0-10 (lower score denotes less pain) Follow-up: 1 month	The mean over- all pain score in the control group was <b>1.7</b> <sup>1</sup>	The mean overall pain score in the intervention group was <b>0.1 points</b> <b>lower</b> (0.92 lower to 0.72 higher)	-	88 (1 RCT)	⊕⊕©© LOW <sup>2</sup>	Absolute risk difference 1% (9% fewer to 7% more); relative per- centage change 3% (28% fewer to 22% more) NNTB not applicable	
Function Assessed with SPADI total score Scale from 0-100 (low- er score denotes greater function) Follow-up: 1 month	The mean func- tion score in the control group was <b>23.2</b> <sup>1</sup>	The mean function score in the intervention group was <b>1 point lower</b> (8.77 lower to 6.77 high- er)	-	88 (1 RCT)	⊕⊕©© Low <sup>2</sup>	Absolute risk difference 1% (9% fewer to 7% more); relative per- centage change 2% (19% fewer to 15% more) NNTB not applicable	
Pain on motion	See Comments column	See Comments column	-	-	-	Outcome not measured	
Global assessment of treatment success	Study population		<b>RR 0.33</b> (0.14 to 0.79)	198 (1 RCT)	⊕⊕⊝© LOW 2,4	Absolute risk difference 12% (21% to 3% fewer); relative percentage	
Follow-up: 6 weeks	<b>184 per 1000</b> <sup>3</sup>	<b>61 per 1000</b> (26 to 145)	(0.14 (0 0.15)	(1101)		change 67% (86% to 21% fewer) NNTB 9 (7 to 26)	

Quality of life Assessed with Global Rat- ing of Change Scale Scale from -7 to 7 (higher score denotes higher qual- ity of life) Follow-up: 1 month	The mean quali- ty of life score in the control group was <b>3</b> <sup>1</sup>	The mean quality of life score in the intervention group was <b>no different</b> (1.37 lower to 1.37 high- er)	-	88 (1 RCT)	⊕⊕⊙© LOW <sup>2</sup>	Absolute risk difference 0% (10% fewer to 10% more); relative per- centage change 0% (46% fewer to 46% more) NNTB not applicable
Adverse events	Study population		not estimable	94 (1 RCT)	⊕⊕⊝⊝ LOW <sup>2</sup>	"Other than transient pain from the CSI [injection], there were no oth-
Follow-up: 12 months	<b>0 per 1000</b> <sup>5</sup>	<b>0 per 1000</b> (0 to 0)		(I KCI)	LOW 2	er adverse events reported by pa- tients in either group."
*The basis for the <b>assumed</b> based on the assumed risk in <b>CI:</b> Confidence interval					corresponding ris	<b>sk</b> (and its 95% confidence interval) is
GRADE Working Group grade High quality: We are very co Moderate quality: We are n stantially different. Low quality: Our confidence Very low quality: We have v	onfident that the true noderately confident e in the effect estima	in the effect estimate: The tr te is limited: The true effect i	rue effect is likely to may be substantia	o be close to the lly different from	the estimate of th	
This table summarises data fr <sup>1</sup> Mean score glucocorticoid in <sup>2</sup> Downgraded (-2) for risk of b <sup>3</sup> Risk in glucocorticoid injectio <sup>4</sup> Downgraded (-1) for indirect <sup>5</sup> Risk in glucocorticoid injectio	jection group in Rho ias. Participants coul on group in Hay 2003 ness. Only 75% of pa	n 2014 used as assumed cont d not be blinded (risk of perf used as assumed risk rticipants had rotator cuff dis	ormance bias and		)	
Summary of findings 3.	Manual therapy ar	nd exercise compared to	arthroscopic su	bacromial dec	ompression for	rotator cuff disease
Manual therapy and exerci	se compared to arth	nroscopic subacromial deco	ompression for ro	tator cuff disea	se	
Patient or population: rota Settings: Hospital, Ringkjoe Intervention: 12 weeks of n Comparison: arthroscopic s	bing County, Denma nanual therapy (soft 1	tissue treatment) plus superv	vised exercises (sta	bilising and stre	ngthening)	

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Outcomes			Relative effect - (95% CI)	No of Partici- pants	Quality of the evidence	Comments	
	Assumed risk Corresponding risk		- (55% CI)	(studies)	(GRADE)		
	Arthroscopic sub- acromial decom- pression	Manual therapy and ex- ercise					
<b>Overall pain</b> Assessed with Con- stant-Murley pain score Scale from 0-15 (higher score denotes less pain) Follow-up: 6 months	The mean improve- ment in overall pain score in the control group was <b>3.8</b> <sup>1</sup>	The mean improvement in overall pain score in the intervention group was <b>0.1 lower</b> (1.68 lower to 1.48 higher)	-	84 (1 RCT)	⊕⊕⊙© LOW <sup>2</sup>	Absolute risk difference 1% (11% fewer to 10% more); rel- ative percentage change 2% (40% fewer to 35% more) NNTB not applicable	
Function Assessed with Con- stant-Murley total score Scale from 0-100 (high- er score denotes greater function) Follow-up: 6 months	The mean improve- ment in function score in the control group was <b>19.9</b> <sup>1</sup>	The mean improvement in function score in the in- tervention group was <b>1.4</b> <b>points higher</b> (7.63 lower to 10.43 higher)	-	84 (1 RCT)	⊕⊕⊙© LOW <sup>2</sup>	Absolute risk difference 1% (79 fewer to 10% more); relative percentage change 4% (23% fewer to 31% more) NNTB not applicable	
Pain on motion	See Comments col- umn	See Comments column	-	-	-	Outcome not measured	
Global assessment of treatment success	Study population		<b>RR 1.14</b> (0.82 to 1.61)	79 (1 RCT)		Absolute risk difference 9% (13% fewer to 30% more); rel-	
Follow-up: 4-8 years	<b>590 per 1000</b> <sup>3</sup>	<b>673 per 1000</b> (484 to 950)	- (0.02 to 1.01)		LOW <sup>2</sup>	(13% fewer to 30% more); rel- ative percentage change 14% (18% fewer to 61% more)	
Quality of life	See Comments col- umn	See Comments column	-	-	-	Outcome not measured	
Adverse events	See Comments col- umn	See Comments column	-	-	-	Outcome not measured	

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GRADE Working Group grades of evidence

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

Moderate quality: 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 quality: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect.

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

This table summarises data from the Haahr 2005 trial.

<sup>1</sup>Mean score in arthroscopic subacromial decompression group in Haahr 2005 used as assumed control group risk

<sup>2</sup>Downgraded (-2) for risk of bias. Participants were not blinded (risk of performance and detection bias)

<sup>3</sup>Risk in arthroscopic subacromial decompression group in Haahr 2005 used as assumed risk



### BACKGROUND

### **Description of the condition**

This review is one in a series of reviews aiming to determine the evidence for efficacy of common interventions for shoulder pain. This series of reviews form the update of an earlier Cochrane review of physical therapy for shoulder disorders (Green 2003). Since our original review, many new clinical trials studying a diverse range of interventions have been performed. To improve usability of the review, we have subdivided the reviews by type of shoulder disorder, as patients within different diagnostic groupings may respond variably to different interventions. This review focuses on manual therapy and exercise interventions alone or in combination for rotator cuff disease. A separate review of electrotherapy modalities for rotator cuff disease is underway. Reviews of manual therapy and exercise for adhesive capsulitis (frozen shoulder) (Page 2014a) and electrotherapy modalities for adhesive capsulitis (Page 2014b) were published in 2014.

Shoulder pain is common, with a point prevalence ranging from 7% to 26% in the general population (Luime 2004). Although not life-threatening, it impacts on the performance of tasks essential to daily living (such as dressing, personal hygiene, eating and work), and often results in substantial utilisation of healthcare resources (Largacha 2006; Mroz 2014; Van der Heijden 1999; Virta 2012). The most common cause of shoulder pain in primary care is disorders of the rotator cuff (Linsell 2006; Ostör 2005), which comprises the supraspinatus, infraspinatus, subscapularis and teres minor muscles. These muscles facilitate both movement and dynamic stabilisation of the shoulder joint (Whittle 2015).

Numerous diagnostic labels have been used in the literature to describe disorders of the rotator cuff (for example, subacromial impingement syndrome, rotator cuff tendinopathy or tendinitis, partial or full rotator cuff tear, calcific tendinitis and subacromial bursitis) but the terms are not standardised (Schellingerhout 2008). The term 'rotator cuff disease' was proposed as an umbrella term to classify disorders of the rotator cuff regardless of the cause of disorder (e.g. degeneration or acute injury) and specific anatomical location (Buchbinder 1996; Whittle 2015).

People with rotator cuff disease often describe their shoulder pain as being worse at night and exacerbated by overhead activity, and some describe weakness or loss of function; however, there are few data regarding the diagnostic accuracy of individual symptoms in rotator cuff disease without tears (Whittle 2015). In addition to history-taking and clinical evaluation, the use of physical examination manoeuvres has been recommended for the diagnosis of rotator cuff disease. A systematic review of diagnostic test accuracy studies found that a positive painful arc test result and a positive external rotation resistance test result were the most accurate findings for detecting rotator cuff disease, whereas the presence of a positive lag test result (external or internal rotation) was most accurate for diagnosis of a full-thickness rotator cuff tear (Hermans 2013).

Rotator cuff disease has been found to increase in prevalence with age (Yamamoto 2010) and in those participating in occupational or sporting activities (e.g. swimming, tennis) that require repetitive overhead use of the arms (Edmonds 2014; Walker 2012). The condition is often self-limiting (Reilingh 2008; Whittle 2015), though 14% of patients, particularly the elderly, have been found to

continue consulting their GP for shoulder pain beyond two years after initial presentation (Linsell 2006).

#### **Description of the intervention**

Manual therapy and exercise, usually delivered together as components of a physical therapy intervention, are commonly used in the management of rotator cuff disease (Whittle 2015). Manual therapy includes any clinician-applied movement of the joints and other structures, for example mobilisation (of which several types exist, e.g. Kaltenborn 1976; Maitland 1977) or manipulation. Exercise includes any purposeful movement of a joint, muscle contraction or prescribed activity, which may be performed under the supervision of a clinician or unsupervised at home. Commonly prescribed exercises include range of motion (ROM), stretching, stabilising and strengthening (Dewhurst 2010).

Manual therapy and exercise are delivered by various clinicians, including physiotherapists, physical therapists, chiropractors, and osteopaths. The aims of both types of interventions are to improve function, promote healing, increase joint range, strengthen weakened muscles and correct imbalance in the stabilising function of the rotator cuff (Brantingham 2011; Kelly 2010; Kuhn 2009). In practice, people with rotator cuff disease seldom receive a single intervention in isolation (i.e. manual therapy alone or exercise alone) (Dziedzic 1999; Glazier 1998; Kooijman 2013; Roberts 2014). Often, electrotherapy modalities (e.g. therapeutic ultrasound, laser therapy) are also delivered as part of a multimodal physical therapy intervention (Kooijman 2013; Struyf 2012), and manual therapy and exercise may also be used in conjunction with other interventions such as non-steroidal anti-inflammatory drugs (NSAIDs) or glucocorticoid injection, or both.

#### How the intervention might work

Manual therapy and exercise interventions are hypothesised to produce a number of beneficial physiological and biomechanical effects. Manual therapy is employed to reduce pain by stimulating peripheral mechanoreceptors and inhibiting nociceptors, and to increase joint mobility by enhancing exchange between synovial fluid and cartilage matrix (Bialosky 2009). Exercise aims to improve muscle function and range of motion by restoring shoulder mobility, proprioception and stability (Kay 2012).

When delivered together, it is unclear whether the effects of manual therapy with exercise represent the effects of manual therapy, the effects of exercise, or an interaction between the two. It has been suggested that the short-term analgesic effects of manual therapy may allow people with other musculoskeletal conditions (e.g. neck pain) to perform exercises designed to produce long-term changes in muscle function and range of motion (Miller 2010; Miller 2014). A similar mechanism of action may occur in people with rotator cuff disease.

#### Why it is important to do this review

The previous version of this review (Green 2003) included four trials investigating the efficacy of manual therapy or exercise (or both) for rotator cuff disease (Bang 2000; Brox 1993; Conroy 1998; Winters 1997), and concluded that firstly, exercise alone was more effective than placebo and secondly, mobilisation was an effective add-on to exercise for people with this condition. However, it was unclear whether manual therapy alone, or manual therapy and exercise, were effective. Many new trials have been published since



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the 2003 review (as summarised in recent systematic reviews, including Brantingham 2011, Braun 2013, Gebremariam 2014, Hanratty 2012, Littlewood 2012 and Van den Dolder 2014). To best inform current practice, an up-to-date review which incorporates the most recently available evidence is needed.

# OBJECTIVES

To synthesise available evidence regarding the benefits and harms of manual therapy and exercise, alone or in combination, for the treatment of people with rotator cuff disease.

# METHODS

# Criteria for considering studies for this review

#### **Types of studies**

We included randomised controlled trials (RCTs) of any design (e.g. parallel, cross-over, factorial) and controlled clinical trials using a quasi-randomised method of allocation, such as by alternation or date of birth. Reports of trials were eligible regardless of the language or date of publication.

### **Types of participants**

We included trials that recruited adults (> 16 years of age) with rotator cuff disease, as defined by the authors (e.g. using terminology such as subacromial impingement syndrome, rotator cuff tendonitis or tendinopathy, supraspinatus, infraspinatus or subscapularis tendonitis, subacromial bursitis, or rotator cuff tears), for any duration.

We also included trials with participants with unspecified shoulder pain provided that the inclusion/exclusion criteria were compatible with a diagnosis of rotator cuff disease. If trials included participants with either rotator cuff disease or adhesive capsulitis, we attempted to retrieve the data for rotator cuff disease participants from the trialists; if unsuccessful, we included the trial only if more than 75% of participants had rotator cuff disease.

We excluded trials that included any participants with a history of significant trauma or systemic inflammatory conditions such as rheumatoid arthritis, osteoarthritis, hemiplegic shoulders, or pain in the shoulder region as part of a complex myofascial neck/ shoulder/arm pain condition.

#### **Types of interventions**

We included trials comparing any manual therapy or exercise intervention to placebo, no treatment, a different type of manual therapy or exercise, or another active intervention (e.g. glucocorticoid injection). Trials evaluating the primary or add-on effects of manual therapy and exercise, manual therapy alone, and exercise alone were eligible.

Eligible manual therapy interventions included mobilisation, manipulation and massage. Eligible exercise interventions included supervised or home exercises, which could be land-based or water-based, but had to comprise tailored shoulder exercises rather than just general activity, for example, swimming.

We excluded trials primarily evaluating the effect of electrotherapy modalities such as therapeutic ultrasound, laser therapy, transcutaneous electrical nerve stimulation (TENS), pulsed electromagnetic field therapy, interferential current, phonophoresis, iontophoresis, or short wave diathermy. Electrotherapy modalities for rotator cuff disease have been analysed in a separate Cochrane review.

### Types of outcome measures

We did not consider outcomes as part of the eligibility criteria.

#### Main outcomes

- Overall pain (mean or mean change measured by visual analogue scale (VAS), numerical or categorical rating scale).
- Function. Where trialists reported outcome data for more than one function scale, we extracted data on the scale that was highest on the following *a priori* defined list:
  - \* Shoulder Pain and Disability Index (SPADI) (Roach 1991);
  - \* Croft Shoulder Disability Questionnaire (Croft 1994);
  - \* Constant-Murley Score (Constant 1987);
  - \* any other shoulder-specific function scale.
- Pain on motion measured by VAS, numerical or categorical rating scale.
- Global assessment of treatment success as defined by the trialists (e.g. proportion of participants with significant overall improvement).
- Quality of life as measured by generic measures (such as components of the Short Form-36 (SF-36)) or disease-specific tools).
- Number of participants experiencing an adverse event in the trial (however defined by the authors).

#### Other outcomes

- Night pain measured by VAS, numerical or categorical rating scale.
- Pain with resisted movement measured by VAS, numerical or categorical rating scale.
- Range of motion (ROM) (e.g. flexion, abduction, external rotation and internal rotation (measured in degrees or other e.g. handbehind-back distance in centimetres)). Where trialists reported outcome data for both active and passive ROM measures, we extracted the data on active ROM only.
- Strength.
- Work disability.
- Surgery (e.g. surgical decompression, rotator cuff repair).

We extracted efficacy outcome measures (e.g. function or overall pain) at the following time points:

- up to three weeks;
- longer than three and up to six weeks (this was the main time point);
- longer than six weeks and up to six months, and;
- longer than six months.

If data were available in a trial at multiple time points within each of the above periods (e.g. at four, five, and six weeks), we only extracted data at the latest possible time point of each period.

We extracted adverse events reported at all time points.

We collated the main results of the review into 'Summary of findings' (SoF) tables which provide key information concerning the quality of evidence and the magnitude and precision of the effect of the interventions. We included the main outcomes (see above) in the SoF tables, with results at, or nearest, the main time point (six weeks) presented.

### Search methods for identification of studies

#### **Electronic searches**

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We searched the Cochrane Central Register of Controlled Trials (CENTRAL; *The Cochrane Library* 2015, Issue 3), Ovid MEDLINE (January 1966 to March 2015), Ovid EMBASE (January 1980 to March 2015), and CINAHL Plus (EBSCO, January 1937 to March 2015). The complete search strategies are presented in Appendix 1. Note that the search terms used also included clinical terms relevant to adhesive capsulitis and electrotherapy interventions as the current review and Cochrane reviews of electrotherapy modalities for rotator cuff disease, manual therapy and exercise for adhesive capsulitis, and electrotherapy modalities for adhesive capsulitis, were conducted simultaneously.

#### Searching other resources

We searched for ongoing trials and protocols of published trials in the clinical trials registry that is maintained by the US National Institute of Health (http://clinicaltrials.gov) and the Clinical Trial Registry at the International Clinical Trials Registry Platform of the World Health Organization (http://www.who.int/ictrp/en/). We also reviewed the reference lists of the included trials and any relevant review articles retrieved from the electronic searches, to identify any other potentially relevant trials.

#### Data collection and analysis

#### **Selection of studies**

Two review authors (MJP and BM) independently selected trials for possible inclusion against a predetermined checklist of inclusion criteria (see Criteria for considering studies for this review). We screened titles and abstracts and initially categorised studies into the following groups.

- Possibly relevant: trials that met the inclusion criteria and trials from which it was not possible to determine whether they met the criteria either from their title or abstract.
- Excluded: those clearly not meeting the inclusion criteria.

If a title or abstract suggested that the trial was eligible for inclusion, or we could not tell, we obtained a full-text version of the article and two review authors (MJP and BM) independently assessed it to determine whether it met the inclusion criteria. The review authors resolved discrepancies through discussion or adjudication by a third author (SG or RB).

#### Data extraction and management

Pairs of review authors (MJP, BM, SS, JD, NL and MM) independently extracted data using a standard data extraction form developed for this review. The authors resolved any discrepancies through discussion or adjudication by a third author (SG or RB), until consensus was reached. We pilot tested the data extraction form and modified it accordingly before use. In addition to items for assessing risk of bias and numerical outcome data, we also recorded the following characteristics.

- Trial characteristics, including type (e.g. parallel or cross-over), country, source of funding, and trial-registration status (with registration number recorded if available).
- Participant characteristics, including age, sex, duration of symptoms, and inclusion/exclusion criteria.
- Intervention characteristics, including type of manual therapy or exercise, duration of treatment, use of co-interventions.
- Outcomes reported, including the measurement instrument used and timing of outcome assessment.

One author (MJP) compiled all comparisons and entered outcome data into Review Manager (RevMan) 5.3 (RevMan 2014).

For a particular systematic review outcome there may be multiple results available in the trial reports (e.g. from multiple scales, time points and analyses). To prevent selective inclusion of data based on the results (Page 2013), we used the following *a priori*-defined decision rules to select data from trials.

- Where trialists reported analysis of covariance- (ANCOVA) adjusted mean differences along with either final values and change from baseline values for the same continuous outcome, we extracted ANCOVA-adjusted mean differences.
- Where trialists reported final values and change from baseline values for the same continuous outcomes, we extracted final values.
- Where trialists reported data analysed based on the intentionto-treat (ITT) sample and another sample (e.g. per-protocol, astreated), we extracted ITT-analysed data;
- For cross-over RCTs, we extracted data from the first period only.

#### Assessment of risk of bias in included studies

Pairs of review authors (MJP, BM, SS, JD, NL and MM) independently assessed the risk of bias in included trials using The Cochrane Collaboration's tool for assessing risk of bias, as described in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). We assessed the following domains.

- Random sequence generation.
- Allocation concealment.
- Blinding of participants and personnel.
- Blinding of outcome assessment (assessed separately for selfreported and objectively assessed outcomes).
- Incomplete outcome data.
- Selective reporting.
- Other sources of bias (for example, baseline imbalance).

We rated each item as being at 'low risk', 'unclear risk' or 'high risk' of bias. We classified the overall risk of bias as low if all domains were at low risk of bias, as high if at least one domain was at high risk of bias, or as unclear if at least one domain was at unclear risk of bias and no domain was at high risk. We assessed the selective reporting domain for all trials, and documented it in the risk of bias tables, but did not consider it in the overall risk of bias judgement if the only types of selective reporting identified were non- or partial reporting of outcomes. Non- or partial reporting of outcomes biases the results of meta-analyses that cannot include the relevant

Manual therapy and exercise for rotator cuff disease (Review)

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data, not the results of trials, and is therefore considered under the Assessment of reporting biases section (Kirkham 2010). We resolved any discrepancies through discussion or adjudication by a third author (SG or RB).

#### Measures of treatment effect

We used the Cochrane Collaboration statistical software, Review Manager 5.3, (RevMan 2014) to perform data analysis. We expressed dichotomous outcomes as risk ratios (RRs) with 95% confidence intervals (CIs) and continuous outcomes as mean differences (MDs) with 95% CIs if different trials used the same measurement instrument to measure the same outcome. Alternatively, we analysed continuous outcomes using the standardised mean difference (SMD) when trials measured the same outcome but employed different measurement instruments. To enhance interpretability of dichotomous outcomes, we calculated risk differences and number needed to treat for an additional beneficial outcome (NNTB) or the number needed to treat for an additional harmful outcome (NNTH).

### Unit of analysis issues

The unit of analysis was the participant. No trials included participants with bilateral shoulder pain.

#### Dealing with missing data

When required, we contacted trialists via email (twice, separated by three weeks) to retrieve missing information about trial design, outcome data, or attrition rates such as drop-outs, losses to followup and post-randomisation exclusions in the included trials. For continuous outcomes with no standard deviation (SD) reported, we calculated SDs from standard errors (SEs), 95% CIs or P values. If no measures of variation were reported and SDs could not be calculated, we planned to impute SDs from other trials in the same meta-analysis, using the median of the other SDs available (Ebrahim 2013). We have reported in the tables of Characteristics of included studies where outcome data were imputed.

#### Assessment of heterogeneity

We assessed clinical heterogeneity by determining whether the characteristics of participants, interventions, outcome measures and timing of outcome measurement were similar across trials. We assessed statistical heterogeneity using the Chi<sup>2</sup> statistic and the I<sup>2</sup> statistic (Higgins 2002). We interpreted the I<sup>2</sup> statistic using the following as an approximate guide:

- 0% to 40% may not be important heterogeneity;
- 30% to 60% may represent moderate heterogeneity;
- 50% to 90% may represent substantial heterogeneity;
- 75% to 100% may represent considerable heterogeneity (Deeks 2011).

#### Assessment of reporting biases

To assess small study effects, we planned to generate funnel plots for meta-analyses including at least 10 trials of varying size. If asymmetry in the funnel plot was detected, we planned to review the characteristics of the trials to assess whether the asymmetry was likely due to publication bias or other factors such as methodological or clinical heterogeneity of the trials (Sterne 2011). To assess outcome reporting bias (non- or partial reporting of a pre-specified outcome, which prevents the inclusion of data

in a meta-analysis), we compared the outcomes specified in trial protocols with the outcomes reported in the corresponding trial publications; if trial protocols were unavailable, we compared the outcomes reported in the methods and results sections of the trial publications (Dwan 2011; Kirkham 2010).

#### **Data synthesis**

For this review update, we identified a large number of trials, which studied a diverse range of interventions. To define the most clinically important questions to be answered in the review, after completing data extraction, one review author (MJP) sent the list of all possible trial comparisons to both of the original primary authors of this review (SG and RB). After reviewing the list of possible trial comparisons, both of these review authors discussed and drafted a list of clinically important review questions and categorised each trial comparison under the most appropriate review question. This process was conducted iteratively until all trial comparisons were allocated to a single review question, and was conducted without knowledge of the results of any outcomes. We defined the following review questions.

- Is manual therapy and exercise (with or without electrotherapy) more effective than placebo, no intervention, or another active intervention (e.g. glucocorticoid injection, oral non-steroidal anti-inflammatory drug (NSAID), arthroscopic subacromial decompression)?
- Is manual therapy and exercise delivered in addition to another active intervention more effective than the other active intervention alone?
- Is manual therapy alone more effective than placebo, no intervention, or another active intervention?
- Is manual therapy delivered in addition to another active intervention more effective than the other active intervention alone?
- Are supervised or home exercises alone more effective than placebo, no intervention, or another active intervention?
- Are supervised or home exercises delivered in addition to another active intervention more effective than the other active intervention alone?
- Is one type of manual therapy or exercise more effective than another (i.e. one type of manual therapy versus another type of manual therapy, or one type of exercise versus another type of exercise)?

We considered the first two to be the main questions of the review, as a multi-modal intervention comprising manual therapy and exercise is most reflective of current clinical practice (Klintberg 2015; Kooijman 2013; Roberts 2014; Struyf 2012).

We planned to pool results of trials with similar characteristics (participants, interventions, outcome measures and timing of outcome measurement) to provide estimates of benefit and harm. We planned to synthesise effect estimates using a random-effects meta-analysis model based on the assumption that clinical and methodological heterogeneity was likely to exist and to have an impact on the results. Where we could not pool data, we presented effect estimates and 95% CIs of each trial in tables and summarised the results in the text.

#### Subgroup analysis and investigation of heterogeneity

We did not undertake any subgroup analyses.



#### Sensitivity analysis

We planned to perform a sensitivity analysis to investigate the robustness of the treatment effect (of main outcomes) to allocation concealment and participant blinding, by removing the trials that reported inadequate or unclear allocation concealment and lack of participant blinding from the meta-analysis to see if this changed the overall treatment effect.

#### Summary of findings tables

We presented the results of the most important comparisons of the review in 'Summary of findings' tables, which summarise the quality of evidence, the magnitude of effect of the interventions examined and the sum of available data on outcomes, as recommended by Cochrane (Schünemann 2011a). The 'Summary of findings' tables include an overall grading of the evidence related to each of the main outcomes, using the GRADE (Grades of Recommendation, Assessment, Development and Evaluation Working Group) approach (Schünemann 2011b).

In the Comments column of the 'Summary of findings' table, we report the absolute per cent difference, the relative per cent change from baseline and the number needed to treat for an additional beneficial outcome (NNTB) (the NNTB is provided only when the outcome shows a statistically significant difference).

For dichotomous outcomes (global assessment of treatment success, adverse events), the absolute risk difference was calculated using the risk difference statistic in RevMan (RevMan 2014), and the result expressed as a percentage; the relative per cent change was calculated as the risk ratio -1 and was expressed as a percentage. For continuous outcomes (overall pain, function, pain on motion, quality of life), the absolute risk difference was calculated as the improvement in the intervention group minus the improvement in the control group, expressed in the original units (i.e. mean difference from RevMan divided by units in the original scale), expressed as a percentage. The relative per cent change is calculated as the absolute change (or mean difference) divided by the baseline mean of the control group, expressed as a percentage.

In addition to the absolute and relative magnitude of effect provided in the 'Summary of findings' table, for dichotomous outcomes we calculated the NNTB or the number needed to treat for an additional harmful effect (NNTH) from the control group event rate, and the risk ratio (RR) using the Visual Rx NNT calculator (Cates 2004). For continuous outcomes of function and overall pain, we calculated the NNTB using Wells calculator software, which is available at the Cochrane Musculoskeletal (CMS) editorial office (http://musculoskeletal.cochrane.org). We assumed a minimal clinically important difference (MCID) of 1.5 points on a 10-point scale (or 15 points on a 100-point scale) for pain (Hawker 2011), and 10 points on a 100-point scale for function or disability (for example SPADI, Constant-Murley, Disabilities of the Arm, Shoulder and Hand (DASH)) (Angst 2011; Roy 2009; Roy 2010) for input into the calculator.

# RESULTS

### **Description of studies**

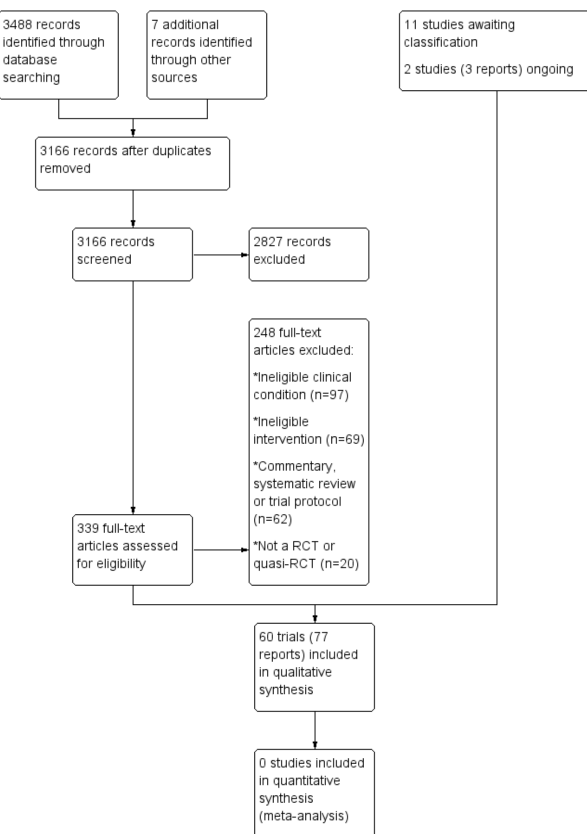
#### **Results of the search**

The search conducted up to March 2015 resulted in 3488 records across the four databases. Seven additional records were identified from screening reference lists of previously published systematic reviews and included trials. After removal of duplicates, we screened the titles and abstracts of 3166 unique records. We screened 339 full-text articles and identified 60 trials (77 reports) which were included in the review (Ainsworth 2009; Al Dajah 2014; Atkinson 2008; Bae 2011; Bang 2000; Bansal 2011; Barbosa 2008; Barra 2011; Barra Lopez 2013; Baskurt 2011; Beaudreuil 2011; Bennell 2010; Bialoszewski 2011; Blume 2014; Brox 1993; Celik 2009; Citaker 2005; Clews 1987; Cloke 2008; Conroy 1998; Cook 2014; Dickens 2005; Djordjevic 2012; Engebretsen 2009; Ginn 2005; Giombini 2006; Haahr 2005; Haik 2014; Hay 2003; Heredia-Rizo 2013; Holmgren 2012; Janse van Rensburg 2012; Kachingwe 2008; Kardouni 2014; Kassolik 2013; Kaya 2014; Kromer 2013; Littlewood 2014; Lombardi 2008; Ludewig 2003; Maenhout 2013; Martins 2012; Marzetti 2014; McClatchie 2009; Moosmayer 2014; Munday 2007; Osteras 2008; Rhon 2014; Senbursa 2007; Senbursa 2011; Struyf 2013; Subasi 2012; Surenkok 2009; Szczurko 2009; Teys 2008; Van den Dolder 2003; Walther 2004; Wang 2006; Winters 1997; Yiasemides 2011).

Eleven additional trials are awaiting classification. Six require translation (Acosta 2009; Bicer 2005; Just 2009; Leblebici 2007; Werner 2002; Wiener 2005) and five are only available as a conference abstract (Bube 2010; Ellegaard 2013; Ginn 2009; Pribicevic 2006; Wies 2008); see table of Characteristics of studies awaiting classification). Two ongoing trials (Roddy 2014; Van den Dolder 2010) were identified in clinical trials registries (see table of Characteristics of ongoing studies). A flow diagram of the study selection process is presented in Figure 1.



# Figure 1. Study flow diagram.





#### **Included studies**

A full description of all included trials is provided in the Characteristics of included studies table.

### Design

All trials except one were described as RCTs (Kassolik 2013 used a quasi-random method of allocation). All trials except two used a parallel-group design (McClatchie 2009 and Teys 2008 used a cross-over design). Forty-eight trials included two intervention arms (Ainsworth 2009; Al Dajah 2014; Atkinson 2008; Bae 2011; Bang 2000; Bansal 2011; Barbosa 2008; Barra 2011; Baskurt 2011; Beaudreuil 2011; Bennell 2010; Bialoszewski 2011; Blume 2014; Celik 2009; Citaker 2005; Conroy 1998; Cook 2014; Dickens 2005; Djordjevic 2012; Engebretsen 2009; Haahr 2005; Haik 2014; Hay 2003; Heredia-Rizo 2013; Holmgren 2012; Janse van Rensburg 2012; Kardouni 2014; Kassolik 2013; Kaya 2014; Kromer 2013; Littlewood 2014; Lombardi 2008; Ludewig 2003; Maenhout 2013; Martins 2012; Marzetti 2014; McClatchie 2009; Moosmayer 2014; Munday 2007; Osteras 2008; Rhon 2014; Senbursa 2007; Struyf 2013; Subasi 2012; Szczurko 2009; Van den Dolder 2003; Wang 2006; Yiasemides 2011), 10 included three arms (Barra Lopez 2013; Brox 1993; Clews 1987; Ginn 2005; Giombini 2006; Senbursa 2011; Surenkok 2009; Teys 2008; Walther 2004; Winters 1997) and two included four arms (Cloke 2008; Kachingwe 2008).

#### Participants

A total of 3620 participants were included in the 60 trials, and the number of participants per trial ranged from nine to 207. The median of the mean age of participants was 51 (interquartile range (IQR) 46 to 56) years, and the median of the mean duration of symptoms was 11 (IQR 5.5 to 25) months. Fifty-two per cent of the participants were women.

Diagnostic labels used by trialists included subacromial impingement syndrome (n = 36: Al Dajah 2014; Bae 2011; Bang 2000; Barra 2011; Barra Lopez 2013; Baskurt 2011; Beaudreuil 2011; Blume 2014; Brox 1993; Celik 2009; Citaker 2005; Conroy 1998; Cook 2014; Dickens 2005; Djordjevic 2012; Engebretsen 2009; Haahr 2005; Haik 2014; Heredia-Rizo 2013; Holmgren 2012; Janse van Rensburg 2012; Kachingwe 2008; Kardouni 2014; Kaya 2014; Kromer 2013; Lombardi 2008; Ludewig 2003; Maenhout 2013; Martins 2012; Munday 2007; Osteras 2008; Rhon 2014; Senbursa 2007; Struyf 2013; Subasi 2012; Walther 2004), rotator cuff tendinitis (n = 4: Atkinson 2008; Clews 1987; Littlewood 2014; Szczurko 2009), supraspinatus tendinitis (n = 3: Bansal 2011; Barbosa 2008; Giombini 2006), painful arc (n = 2: Cloke 2008; McClatchie 2009), rotator cuff tear (n = 2: Ainsworth 2009; Moosmayer 2014), chronic rotator cuff disease (n = 1: Bennell 2010), chronic rotator cuff injury (n = 1:Bialoszewski 2011) or a mixture of labels (i.e. some participants with impingement, others with tendinitis) (n = 3: Senbursa 2011; Surenkok 2009; Van den Dolder 2003). However, there were inconsistencies in the diagnostic criteria for (or definitions of) each of the conditions (see Characteristics of included studies).

Six (10%) trials (Ginn 2005; Kassolik 2013; Teys 2008; Wang 2006; Winters 1997; Yiasemides 2011) included participants with non-specific shoulder pain that was compatible with a diagnosis of rotator cuff disease. Two trials (Hay 2003; Surenkok 2009) included patients with either rotator cuff disease or adhesive capsulitis, but

participants with the latter condition comprised less than 25% of the sample.

Trials were conducted in USA (n = 9), Turkey (n = 8), UK (n = 7), Australia (n = 6), Brazil, Norway (n = 4 each), Spain (n = 3), Belgium, Canada, Italy, Poland, The Netherlands (n = 2 each), Denmark, France, Germany, India, Republic of Korea, Saudi Arabia, Serbia, South Africa and Sweden (n = 1 each).

#### **Interventions and Comparisons**

A detailed description of the interventions delivered in each trial is summarised in the Characteristics of included studies and a summary of the intervention components across trials is presented in Table 1. The median duration of the physical therapy interventions was six weeks (range one to 24), with a median of two treatment sessions delivered per week (range one to seven). The types of manual therapy and exercise delivered were very heterogeneous across the trials.

Manual therapy interventions included:

- joint mobilisation (glenohumeral or acromioclavicular joint) (n
   = 21: Bang 2000; Barbosa 2008; Bennell 2010; Bialoszewski 2011; Conroy 1998; Dickens 2005; Djordjevic 2012; Ginn 2005; Hay 2003; Kachingwe 2008; Kaya 2014; Kromer 2013; McClatchie 2009; Rhon 2014; Senbursa 2007; Senbursa 2011; Surenkok 2009; Struyf 2013; Teys 2008; Winters 1997; Yiasemides 2011);
- soft tissue mobilisation or massage of the shoulder (n = 9: Al Dajah 2014; Bennell 2010; Clews 1987; Haahr 2005; Heredia-Rizo 2013; Kaya 2014; Rhon 2014; Senbursa 2007; Van den Dolder 2003);
- spinal or neck mobilisation or manipulation (n = 5: Bennell 2010; Cook 2014; Haik 2014; Kardouni 2014; Kaya 2014);
- shoulder manipulation (n = 4: Atkinson 2008; Janse van Rensburg 2012; Munday 2007; Winters 1997);
- deep friction massage (n = 4: Bansal 2011; Bialoszewski 2011; Senbursa 2007; Senbursa 2011);
- proprioceptive neuromuscular facilitation stretching techniques (n = 4: Al Dajah 2014; Kaya 2014; Senbursa 2007; Senbursa 2011);
- diacutaneous fibrolysis (n = 2: Barra 2011; Barra Lopez 2013).

Two trials (Citaker 2005; Cloke 2008), did not report details about the type of manual therapy delivered.

Many different types of exercises were delivered in the trials. These included:

- strengthening exercises (n = 23: Ainsworth 2009; Bae 2011; Baskurt 2011; Bennell 2010; Bialoszewski 2011; Brox 1993; Celik 2009; Conroy 1998; Cook 2014; Dickens 2005; Djordjevic 2012; Haahr 2005; Janse van Rensburg 2012; Kachingwe 2008; Kaya 2014; Martins 2012; Marzetti 2014; Rhon 2014; Senbursa 2007; Senbursa 2011; Subasi 2012; Wang 2006; Yiasemides 2011);
- stretching exercises (n = 18: Ainsworth 2009; Baskurt 2011; Blume 2014; Celik 2009; Conroy 1998; Cook 2014; Giombini 2006; Holmgren 2012; Kromer 2013; Ludewig 2003; Martins 2012; Marzetti 2014; Senbursa 2007; Senbursa 2011; Subasi 2012; Walther 2004; Wang 2006; Yiasemides 2011);
- range of motion exercises (n = 13: Ainsworth 2009; Bialoszewski 2011; Brox 1993; Celik 2009; Conroy 1998; Cook 2014; Djordjevic

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2012; Ginn 2005; Kaya 2014; Rhon 2014; Senbursa 2011; Subasi 2012; Szczurko 2009);

- progressive resistance training (n = 7: Dickens 2005; Engebretsen 2009; Littlewood 2014; Lombardi 2008; Ludewig 2003; Marzetti 2014; Osteras 2008);
- Codman's pendulum exercises (n = 7: Bansal 2011; Citaker 2005; Giombini 2006; Heredia-Rizo 2013; Kaya 2014; Martins 2012; Marzetti 2014);
- eccentric training exercises (n = 6: Barbosa 2008; Blume 2014; Citaker 2005; Holmgren 2012; Maenhout 2013; Moosmayer 2014):
- postural exercises (n = 5: Ainsworth 2009; Bang 2000; Bennell 2010; Blume 2014; Kachingwe 2008);
- motor control exercises (n = 5: Bae 2011; Marzetti 2014; Moosmayer 2014; Struyf 2013; Yiasemides 2011);
- proprioceptive exercises (n = 4: Ainsworth 2009; Baskurt 2011; Heredia-Rizo 2013; Martins 2012);
- self-mobilisation techniques (n = 2: Bang 2000; Rhon 2014);
- dynamic humeral centring (Beaudreuil 2011).

Two trials (Barra Lopez 2013; Cloke 2008) did not report the specific type of exercises delivered. Exercises were performed:

- only under supervision in 20 trials (Ainsworth 2009; Bae 2011; Barbosa 2008; Barra Lopez 2013; Baskurt 2011; Bialoszewski 2011; Conroy 1998; Djordjevic 2012; Engebretsen 2009; Heredia-Rizo 2013; Janse van Rensburg 2012; Lombardi 2008; Martins 2012; Marzetti 2014; Moosmayer 2014; Osteras 2008; Senbursa 2011; Szczurko 2009; Walther 2004; Winters 1997);
- only at home in seven trials (Cook 2014; Littlewood 2014; Ludewig 2003; Senbursa 2007; Senbursa 2011; Walther 2004; Yiasemides 2011);
- or both under supervision and at home in 21 trials (Bang 2000; Bansal 2011; Beaudreuil 2011; Bennell 2010; Blume 2014; Brox 1993; Celik 2009; Citaker 2005; Dickens 2005; Ginn 2005; Giombini 2006; Haahr 2005; Hay 2003; Holmgren 2012; Kaya 2014; Kromer 2013; Maenhout 2013; Rhon 2014; Struyf 2013; Subasi 2012; Wang 2006).

Trialists investigated the primary or add-on effects of:

- manual therapy and exercise in 10 trials (Bennell 2010; Cloke 2008; Dickens 2005; Ginn 2005; Haahr 2005; Hay 2003; Kachingwe 2008; Rhon 2014; Szczurko 2009; Winters 1997);
- manual therapy alone in 29 trials (Al Dajah 2014; Atkinson 2008; Bang 2000; Bansal 2011; Barbosa 2008; Barra 2011; Barra Lopez 2013; Bialoszewski 2011; Citaker 2005; Clews 1987; Conroy 1998; Cook 2014; Haik 2014; Heredia-Rizo 2013; Janse van Rensburg 2012; Kachingwe 2008; Kardouni 2014; Kassolik 2013; Kaya 2014; Kromer 2013; McClatchie 2009; Munday 2007; Senbursa 2007; Senbursa 2011; Surenkok 2009; Teys 2008; Van den Dolder 2003; Winters 1997; Yiasemides 2011); or
- exercise alone in 26 trials (Ainsworth 2009; Bae 2011; Baskurt 2011; Beaudreuil 2011; Blume 2014; Brox 1993; Celik 2009; Djordjevic 2012; Engebretsen 2009; Ginn 2005; Giombini 2006; Holmgren 2012; Kachingwe 2008; Littlewood 2014; Lombardi 2008; Ludewig 2003; Maenhout 2013; Martins 2012; Marzetti 2014; Moosmayer 2014; Osteras 2008; Senbursa 2011; Struyf 2013; Subasi 2012; Walther 2004; Wang 2006).

Comparators were also diverse, including:

- placebo (Barra 2011; Bennell 2010; Brox 1993; Haik 2014; Kardouni 2014; McClatchie 2009; Munday 2007; Surenkok 2009; Teys 2008);
- no intervention (Dickens 2005; Kachingwe 2008; Lombardi 2008; Ludewig 2003; Surenkok 2009; Teys 2008; Van den Dolder 2003);
- glucocorticoid injection (Cloke 2008; Ginn 2005; Hay 2003; Rhon ٠ 2014; Winters 1997);
- surgery (Brox 1993; Haahr 2005; Moosmayer 2014);
- electrotherapy modalities (e.g. therapeutic ultrasound, microwave diathermy) (Al Dajah 2014; Bansal 2011; Giombini 2006);
- naturopathic care (Szczurko 2009);
- oral NSAID (Cloke 2008); •
- extracorporeal shock wave treatment (Engebretsen 2009); •
- kinesiotaping (Kaya 2014);
- a functional brace (Walther 2004).

Nineteen trials investigated whether there was benefit in adding manual therapy or exercise to another physical therapy intervention (Ainsworth 2009; Atkinson 2008; Bae 2011; Bang 2000; Barbosa 2008; Barra Lopez 2013; Baskurt 2011; Beaudreuil 2011; Bialoszewski 2011; Clews 1987; Conroy 1998; Cook 2014; Janse van Rensburg 2012; Kachingwe 2008; Kromer 2013; Maenhout 2013; Martins 2012; Senbursa 2011; Yiasemides 2011), and in 18 trials, one type of manual therapy or exercise was compared with another (Blume 2014; Celik 2009; Citaker 2005; Djordjevic 2012; Heredia-Rizo 2013; Holmgren 2012; Kachingwe 2008; Kassolik 2013; Littlewood 2014; Marzetti 2014; Osteras 2008; Senbursa 2007; Senbursa 2011; Struyf 2013; Subasi 2012; Walther 2004; Wang 2006; Winters 1997).

#### Outcomes

The outcomes measured in each trial are summarised in Table 2. Of the main outcomes, most trials included a measure of overall pain (n = 48) and function (n = 44), but fewer included measures of pain on motion (n = 16), global assessment of treatment success (n = 17), quality of life (n = 13) or adverse events (n = 17). Overall pain was most commonly measured using a zero to 10 or zero to 100 VAS, though several different descriptors for the maximum score on the scale (e.g. "worst imaginable pain", "severe pain", "intolerable pain") were noted. Function was most commonly measured using the SPADI or the Constant-Murley Score. Of the other outcomes, most trials included measures of range of motion (n = 38), but fewer included measures of night pain (n = 9), pain with resisted movement (n = 1), strength (n = 19), work disability (n = 7) or surgery (n = 2).

We contacted authors of three trials to retrieve missing data for unreported or partially reported outcomes (Cloke 2008; Dickens 2005; Kachingwe 2008), but received no responses.

#### **Excluded studies**

We excluded 248 full-text articles. Many of these were excluded because they were eligible for inclusion in one of the other three reviews in this series (i.e. focused on electrotherapy modalities for rotator cuff disease or adhesive capsulitis, or manual therapy or exercise for adhesive capsulitis). The reasons for exclusion were that the clinical condition was ineligible (n = 97), the intervention was ineligible (n = 69), the article was a commentary or systematic review (n = 62) or the study was not a RCT or quasi-RCT (n = 20).

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We have listed in the table of Characteristics of excluded studies 16 studies which required full-text screening by a third author (the full list of 248 excluded studies is available on request).

# **Risk of bias in included studies**

A summary of the risk of bias in included trials is presented in Figure 2 and Figure 3.

Figure 2. Risk of bias graph: review authors' judgements about each risk of bias item presented as percentages across all included studies. White areas mean that either subjective or objective outcomes were not measured in some of the trials, so an assessment of the risk of bias due to lack blinding of such outcomes was not applicable.

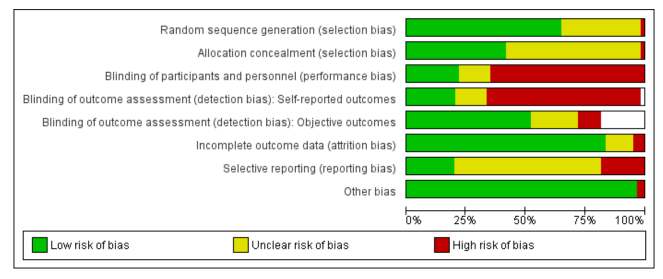


Figure 3. Risk of bias summary: review authors' judgements about each risk of bias item for each included study. Empty cells mean that either subjective or objective outcomes were not measured in the trial, so an assessment of the risk of bias due to lack blinding of such outcomes was not applicable.



Manual therapy and exercise for rotator cuff disease (Review)

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Figure 3. (Continued)

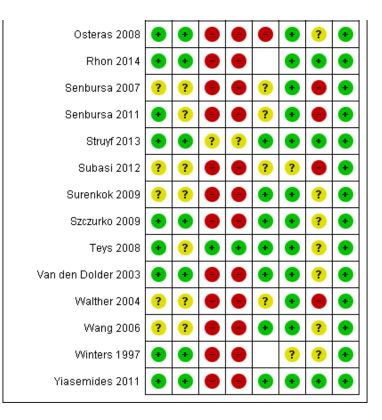
Citaker 2005	?	?	?	?	?	?	?	•
Clews 1987	?	?	•	•	•	•	?	•
Cloke 2008	?	?	•	•		•	•	•
Conroy 1998	?	?	•	•	•	•	?	•
Cook 2014	•	?	•	•		•	?	•
Dickens 2005	?	?	•	•	•	•	?	•
Djordjevic 2012	•	?	•		•	•	•	•
Engebretsen 2009	•	•	•	•	•	•	•	•
Ginn 2005	?	?	•	•	•	•	?	•
Giombini 2006	•	?	•	•	•	•	•	•
Haahr 2005	•	•	•		•	•	?	•
Haik 2014	•	?	•	•		•	?	•
Hay 2003	•	•			•	•	?	•
Heredia-Rizo 2013	?	•	••	?	•	•	?	•
Holmgren 2012	•	•	•	•	•	•	•	•
Janse van Rensburg 2012	•	•	•	•	•	•	?	•
Kachingwe 2008	•	?	•	•	÷	•	?	•
Kardouni 2014	•	•	•	•		•	?	•
Kassolik 2013	•	•	•	•	•	•	•	•
Kaya 2014	•	?		•		•	?	•
Kromer 2013	•	•	•	•		•	•	•
Littlewood 2014	•	•	•	•		•	•	•
Lombardi 2008	•	•	•	•	•	•	?	•
Ludewig 2003	•	?	•	•		•	?	•
Maenhout 2013	?	?	•	•	•	•	•	•
Martins 2012	?	?	?	?		•	•	•
Marzetti 2014	•	•	?	?	•	•	•	•
McClatchie 2009	•	?	•	•	•	•	?	•
Moosmayer 2014	•	•	•	•	•	•	?	•
Munday 2007	•	•	•	•	?	?	•	•
Osteras 2008	•	•	•	•	•	•	?	•
			-	-				- 1

Manual therapy and exercise for rotator cuff disease (Review)

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#### Figure 3. (Continued)



#### Allocation

We rated 39 (65%) trials at low risk of allocation bias because the method used to generate the random allocation sequence was adequate. We also rated 25 (42%) trials at low risk of allocation bias because the method used to conceal the allocation sequence was adequate. We rated one trial at high risk of allocation bias because participants were allocated to groups using a quasi-random sequence. In 20 (33%) trials the method of sequence generation was not reported and in 34 (57%) trials the method of allocation bias in these trials was therefore unclear.

#### Blinding

We rated 13 (22%) trials at low risk of performance bias because participants were successfully blinded. We rated eight (13%) trials at unclear risk of performance bias because participants received different types of manual therapy or exercise, but it is unclear whether they were provided with any information that would make them perceive the type of manual therapy or exercise they received as superior or inferior to the alternative type of manual therapy or exercise. We rated all 39 (65%) remaining trials at high risk of performance bias because participants were not blinded, which may have led them to deviate from the interventions as planned because of their beliefs about the intervention they received.

Self-reported outcomes were measured in all but one trial, and of these, we rated 12 (20%) at low risk of detection bias because it was clear that participants were blinded, eight (14%) at unclear risk of detection bias because it was unclear whether participants were blinded, and the 39 remaining trials (66%) at high risk of

detection bias for self-reported outcomes because participants were not blinded. Of 49 trials with outcome measures that were objectively rated (e.g. range of motion, strength), blinding of outcome assessors was reported in 31 (63%) and thus we rated these trials at low risk of detection bias for objective outcomes. In six (12%) trials there was no blinding of assessors of objective outcomes, so the risk of detection bias for objective outcomes was high. In 12 (24%) trials it was unclear whether such blinding was done, so the risk of detection bias for objective outcomes was unclear.

#### Incomplete outcome data

Fifty (83%) trials either had no dropouts, losses to follow-up or exclusions, or had a small amount of attrition that was deemed unlikely to bias the results. In three (5%) trials there was differential dropout across groups, with reasons that appeared to be related to the treatments received, and thus we rated these trials at high risk of attrition bias. In the remaining seven (12%) trials the quantity of or reasons for incomplete outcome data were not reported so the risk of attrition bias was unclear.

#### Selective reporting

We rated 12 (20%) trials at low risk of selective reporting bias because all outcomes specified in the trial registry entry or the trial protocol were fully reported in the trial publication. We rated 11 (18%) trials at high risk of selective reporting bias because data for at least one outcome that was listed in the trial registry entry or the methods section of the publication were not reported in the results section at all. We rated the remaining 37 (62%) trials at unclear risk of selective reporting bias for one of two reasons.

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Firstly, outcome data were completely reported for all outcomes specified in the methods section of the publication, but none of these trials was registered in a trials registry or had an available trial protocol, so it was unclear whether other outcomes were measured but not reported based on the nature of the results; or secondly, outcome data were incompletely reported (e.g. reporting means without measures of variation), but it was unclear whether data were incompletely reported based on the statistical significance or magnitude of the results.

#### Other potential sources of bias

In Engebretsen 2009 (supervised exercises versus radial extracorporeal shockwave treatment), there was an imbalance between groups in the number of additional treatments received outside of the trial setting, which was likely to bias the results in favour of the shockwave treatment group. In Kassolik 2013 (Swedish massage versus massage based on the tensegrity principle), there was baseline imbalance in range of motion, which may have biased results in favour of the group receiving massage based on the tensegrity principle. We rated both trials at high risk of other bias, and all other trials (97%) as free from other potential sources of bias.

# **Effects of interventions**

See: Summary of findings for the main comparison Manual therapy and exercise compared to placebo for rotator cuff disease; Summary of findings 2 Manual therapy and exercise compared to glucocorticoid injection for rotator cuff disease; Summary of findings 3 Manual therapy and exercise compared to arthroscopic subacromial decompression for rotator cuff disease

We were unable to perform any meta-analyses because of clinical heterogeneity or incomplete outcome reporting. Summary data and effect estimates (with 95% CIs) for all trials are presented in the Additional tables section. To enhance readability, we have reported in the following section the summary data, effect estimates and 95% CIs for main outcomes reported in trials addressing the two primary questions of the review (i.e. "Is manual therapy and exercise (with or without electrotherapy) more effective than placebo, no intervention or another active intervention?" and "Is manual therapy and exercise delivered in addition to another active intervention alone?"). For the remaining questions, the relevant data are presented in the Additional tables. If an outcome is not referred to within a sub-section or table, then no data for that outcome were available in the trial(s).

#### Is manual therapy and exercise (with or without electrotherapy) more effective than placebo, no intervention or another active intervention (e.g. glucocorticoid injection, oral non-steroidal anti-inflammatory drug (NSAID), arthroscopic subacromial decompression)?

In 10 trials, manual therapy and exercise was compared with either placebo (Bennell 2010), no intervention (Dickens 2005; Kachingwe 2008) or another active intervention (Cloke 2008; Ginn 2005; Haahr 2005; Hay 2003; Rhon 2014; Szczurko 2009; Winters 1997).

#### Manual therapy and exercise versus placebo

See Summary of findings for the main comparison. In one trial of 120 participants with chronic rotator cuff disease, judged at low risk of bias overall, a manual therapy and exercise package was

compared with placebo (Bennell 2010). The package comprised soft tissue massage, glenohumeral joint mobilisation, thoracic spine mobilisation, cervical spine mobilisation, scapular retraining and postural taping in 10 sessions over 10 weeks along with home exercises primarily focused upon strengthening the rotator cuff muscles for 22 weeks. The placebo consisted of inactive ultrasound therapy and application of an inert gel in 10 sessions over 10 weeks.

No data were available at our primary time point (three to six weeks). At 22 weeks, the mean change in overall pain with placebo was 17.3 points on a 100-point scale, and 24.8 points with manual therapy and exercise (adjusted mean difference (MD) 6.8 points, 95% confidence interval (CI) -0.70 to 14.30 points). Mean change in function with placebo was 15.6 points on a 100-point scale, and 22.4 points with manual therapy and exercise (adjusted MD 7.1 points, 95% CI 0.30 to 13.90 points). Mean change in pain on motion with placebo was 1.6 points on a 10-point scale, and 2.6 points with manual therapy and exercise (adjusted MD 0.9 points, 95% CI -0.03 to 1.70 points). Fifty-seven per cent (31/54) of participants reported treatment success with manual therapy and exercise compared with 41% (24/58) of participants receiving placebo (risk ratio (RR) 1.39, 95% CI 0.94 to 2.03). Mean change in quality of life with placebo was 0 points on a 1.4-point scale, and 0.07 points with manual therapy and exercise (adjusted MD 0.07 points, 95% CI 0.04 to 0.10 points). None of these differences were considered to be clinically important (Table 3). Thirty-one per cent (17/55) of participants reported adverse events with manual therapy and exercise compared with 8% (5/61) of participants receiving placebo (RR 3.77, 95% CI 1.49 to 9.54). However adverse events were mild and short-lived (short-term pain following treatment). We considered the evidence from this trial to be high quality.

#### Manual therapy and exercise versus no treatment

Two trials (89 participants), both at high risk of bias overall, compared manual therapy and exercise with no active intervention other than advice to maintain normal activities (Dickens 2005) or advice regarding posture and overhead activities (Kachingwe 2008). The physical therapy intervention in Dickens 2005 comprised mobilisation of the glenohumeral joint, acromioclavicular joint and thoracic and cervical spine, exercise therapy including attention to muscle imbalance, postural advice, strapping and occasionally electrotherapy, while in Kachingwe 2008, physical therapy comprised either glenohumeral mobilisation and supervised and home exercises or mobilisation with movement and supervised and home exercises.

In Dickens 2005, at six months the mean change in function with no treatment was 0.65 on a 100-point scale, and 20 points with manual therapy and exercise (MD 19.35, 73 participants) but the 95% CI was not estimable. No other outcomes were reported in this trial. Usable outcome data were not available in Kachingwe 2008, although the authors claimed that there were no statistically significant differences between groups in overall pain, function and active shoulder flexion at six weeks (Table 4). We downgraded by two points for high risk of performance and detection bias, and one point for imprecision, and so consider this evidence to be very low quality.

#### Manual therapy and exercise versus glucocorticoid injection

See Summary of findings 2. Five trials (507 participants), all at high risk of bias overall, compared manual therapy and exercise with glucocorticoid injection (Cloke 2008; Ginn 2005; Hay 2003; Rhon



2014; Winters 1997). The physical therapy interventions comprised: six sessions over 18 weeks of manual therapy and exercise (no details provided) (Cloke 2008); passive joint mobilisation and range of motion exercises twice a week for five weeks (Ginn 2005); active and passive mobilisation, home exercises and therapeutic ultrasound once a week for six weeks (Hay 2003); joint and soft tissue mobilisation, manual stretches and supervised and home exercises twice a week for three weeks (Rhon 2014); and massage and exercises once a week for six weeks (Winters 1997). The total number of glucocorticoid injections delivered varied across the trials, from a single injection at baseline only (Ginn 2005; Hay 2003), injection at six, 12 and 18 weeks (Cloke 2008), injection at baseline, one week later and two weeks later if necessary (Winters 1997), and as many as three injections administered one month apart during the one-year period (Rhon 2014). Due to the heterogeneity in interventions, comparators, diagnoses of shoulder pain, outcome measures, timing of outcome assessment and incomplete reporting of outcome data in some trials, we were unable to synthesise any data in meta-analyses.

Based on results of single trials, there was no clinically important difference between manual therapy and exercise and glucocorticoid injection with respect to overall pain at four weeks (mean 1.6 versus 1.7 on a 10-point scale, MD -0.10, 95% CI -0.92 to 0.72, 88 participants), 11 weeks (mean 11.5 versus 9.2 on a 28-point scale, MD 2.30, 95% CI 0.50 to 4.10, 82 participants), six months (mean 1.7 versus 2.2 on a 10-point scale, MD -0.50, 95% CI -1.32 to 0.32, 84 participants) or 12 months (mean 2.1 versus 2.5 on a 10point scale, MD -0.40, 95% CI -1.23 to 0.43, 94 participants). Further, there was no clinically important difference between groups in function at four weeks (mean 22.2 versus 23.2 on a 100-point scale, MD -1.00, 95% CI -8.77 to 6.77, 88 participants), five weeks (mean change 5.3 versus 5.2 on a 27-point scale, MD 0.10, 95% CI -1.62 to 1.82, 84 participants), six weeks (mean change 2.56 versus 3.03 on a 23-point scale, MD -0.47, 95% CI -2.11 to 1.17, 197 participants), 18 weeks (mean 27.73 versus 29.81 on a 48-point scale, MD -2.08, 95% CI-10.46 to 6.30, 49 participants), six months (mean 21.5 versus 22.2 on a 100-point scale, MD -0.70, 95% CI -8.52 to 7.12, 84 participants) or 12 months (mean 21.6 versus 23.1 on a 100-point scale, MD -1.50, 95% CI -9.07 to 6.07, 94 participants). Only one trial (Rhon 2014) measured adverse events, and found that apart from transient pain due to the injection, there were no other adverse events reported in either group (Table 5).

Three trials measured global treatment success (Ginn 2005; Hay 2003; Winters 1997). While Ginn 2005 found no difference between groups at five weeks (effect not estimable), Hay 2003 found 6% (6/100) of participants receiving manual therapy and exercise reported global treatment success at six weeks compared with 18% (18/98) of participants receiving glucocorticoid injection (RR 0.33, 95% CI 0.14 to 0.79; 198 participants). A similar effect was found in Winters 1997 at 11 weeks (51% [18/35] manual therapy and exercise versus 89% [42/47] glucocorticoid injection, RR 0.58, 95% CI 0.41 to 0.81; 82 participants). However, the difference between groups was less certain at six months in Hay 2003 (23% [23/99] manual therapy and exercise versus 18% [17/97] glucocorticoid injection, RR 1.33, 95% CI 0.76 to 2.32; 196 participants). Quality of life was measured in two trials (Hay 2003; Rhon 2014). In Hay 2003, scores were no different between groups at six weeks or six months (95% CIs not estimable). In Rhon 2014, quality of life scores were no different between groups at one month (mean 3 versus 3 on a -7 to +7 scale, MD 0.00, 95% CI -1.37 to 1.37, 88 participants), six months (mean 3 versus 3 on a -7 to +7 scale, MD 0.00, 95% CI -2.17 to 2.17, 84 participants) or 12 months (mean 3 versus 3 on a -7 to +7 scale, MD 0.00, 95% CI -1.38 to 1.38, 94 participants). Night pain scores at six weeks and six months in one trial (Hay 2003) were no different between groups (95% CIs not estimable). Active range of motion was reported in two trials (Ginn 2005; Hay 2003). In Ginn 2005, mean differences between groups in active shoulder abduction, flexion and hand-behind-back distance were very small and the precision could not be estimated due to incomplete reporting. In Hay 2003, fewer participants in the manual therapy and exercise group had impairment in active shoulder abduction and external rotation but these differences were not statistically significant (Table 5).

In summary, the overall impression from these five trials is that there were no clinically important differences between manual therapy and exercise and glucocorticoid injection with respect to overall pain, function, quality of life, night pain and active range of motion at both short- (four to six weeks) and long-term (six to 12 months). However, global treatment success was more common up to 11 weeks in participants receiving glucocorticoid injection. The lack of difference at long-term is not surprising given that glucocorticoid injections are short-acting interventions which only have evidence of benefit over placebo at short-term follow-up (Buchbinder 2003). A key limitation of these trials is the lack of participant blinding, which may have biased results in either direction if participants had different pre-conceived beliefs about the efficacy of physical therapy and glucocorticoid injection. Therefore, we downgraded by two points for high risk of performance and detection bias, and consider this evidence to be low quality.

#### Manual therapy and exercise versus NSAID

One trial (39 participants) at high risk of bias overall (Cloke 2008), compared six sessions over 18 weeks of manual therapy and exercise (no details provided) with regular NSAID or "simple analgesic intake" (dose and duration not reported). There was no clinically important difference between groups in function at 18 weeks (mean 27.73 versus 30.47 on a 48-point scale, MD -2.74, 95% CI -10.21 to 4.73) or 12 months (mean 28.94 versus 30.07 on a 48-point scale, MD -1.13, 95% CI not estimable, unclear number of participants) (Table 6). We downgraded by two points for high risk of performance and attrition bias, and one point for imprecision, and thus consider this evidence to be very low quality.

#### Manual therapy and exercise versus arthroscopic subacromial decompression

See Summary of findings 3. One trial (84 participants) at high risk of bias overall (Haahr 2005), compared 12 weeks of manual therapy (soft tissue treatment) and supervised exercises (stabilising and strengthening) with arthroscopic subacromial decompression. There was no clinically important difference between groups in overall pain at six months (mean change 3.7 versus 3.8 on a 15-point scale, MD -0.10, 95% CI -1.68 to 1.48, 84 participants), 12 months (mean change 3.7 versus 3.6 on a 15-point scale, MD 0.10, 95% CI -1.49 to 1.69, 84 participants) or four to eight years (mean change 3 versus 1.9 on a 10-point scale, MD 1.10, 95% CI -0.14 to 2.34, 79 participants). There was also no clinically important difference between groups in function at six months (mean change 21.3 versus 19.9 on a 100-point scale, MD 1.40, 95% CI -7.63 to 10.43, 84 participants), 12 months (mean change 23 versus 18.8 on a 100-point scale, MD 4.20, 95% CI -5.03 to 13.43, 84 participants) and four



to eight years (mean change 11.4 versus 9.1 on a 36-point scale, MD 2.30, 95% CI -2.06 to 6.66, 79 participants). With respect to the remaining outcomes, there was no clinically important difference between groups in global treatment success at four to eight years (68% [27/40] versus 59% [23/39], RR 1.14, 95% CI 0.82 to 1.61; 79 participants), active range of motion at six months (mean change 10.3 versus 9.6 on a 40-point scale, MD 0.70, 95% CI -3.83 to 5.23, 84 participants) and 12 months (mean change 11.6 versus 8.2 on a 40-point scale, MD 3.40, 95% CI -1.34 to 8.14, 84 participants), or strength at six months (mean change 2.7 versus 2.9 on a 25point scale, MD -0.20, 95% CI -2.50 to 2.10, 84 participants) and 12 months (mean change 3.2 versus 3.3 on a 25-point scale, MD -0.10, 95% CI -2.68 to 2.48, 84 participants) (Table 7). The lack of patient blinding may have influenced patients in both groups to pursue alternative interventions and influenced their responses to self-reported outcomes, which may have biased results in either direction. For this reason we downgraded by two points for high risk of performance and detection bias, and consider this evidence to be low quality.

# Manual therapy and exercise versus naturopathic care (dietary counselling, acupuncture, and Phlogenzym supplement)

One trial (85 participants) at high risk of bias overall (Szczurko 2009), compared naturopathic care comprising dietary counselling and acupuncture once per week and daily Phlogenzym supplement (recommended by some naturopaths for pain relief) for 12 weeks with manual therapy and supervised exercises (range of motion and strengthening) once per week and daily placebo tablet for 12 weeks. The authors observed clinically important differences favouring naturopathic care over manual therapy and exercise at 12 weeks with respect to overall pain (mean 2.75 versus 4.05 on a seven-point scale, MD 1.30, 95% CI 0.56 to 2.04) and function (mean 35.3 versus 56.24 on a 130-point scale, MD 20.94, 95% CI 6.40 to 35.48). Further, several domains of the SF-36 quality of life measure (physical functioning, role limitations due to physical health, bodily pain, and general health), and active shoulder abduction, flexion and extension were statistically significantly lower in the manual therapy and exercise group at 12 weeks (see Table 8). However, the lack of participant blinding may have biased results in either direction if participants had different pre-conceived beliefs about the efficacy of physical therapy and naturopathic care. Therefore, we downgraded by two points for high risk of performance and detection bias, and thus consider this evidence to be low quality.

#### Is manual therapy and exercise delivered in addition to another active intervention more effective than the other active intervention alone?

# Manual therapy and exercise and glucocorticoid injection versus glucocorticoid injection

One trial (47 participants) at high risk of bias overall (Cloke 2008), compared six sessions over 18 weeks of manual therapy and exercise (no details provided) along with glucocorticoid injection (single injection at six, 12 and 18 weeks) with glucocorticoid injection alone. There was no clinically important difference between groups in function at 18 weeks (mean 27.8 versus 29.81 on a 48-point scale, MD -2.01, 95% CI -13.09 to 9.07) or at 12 months (mean 23.79 versus 26.47 on a 48-point scale, MD -2.68, 95% CI not estimable) (Table 9). We downgraded by two points for high risk of performance and attrition bias, and one point for imprecision, and thus consider this evidence to be very low quality.

# Is manual therapy alone more effective than placebo, no intervention or another active intervention?

Twelve trials compared manual therapy alone with either placebo (Barra 2011; Haik 2014; Kardouni 2014; McClatchie 2009; Munday 2007; Surenkok 2009; Teys 2008), no intervention (Surenkok 2009; Teys 2008; Van den Dolder 2003), or another active intervention (Al Dajah 2014; Bansal 2011; Kaya 2014; Winters 1997).

#### Manual therapy alone versus placebo

The types of manual therapy investigated in the seven trials that compared it to placebo were diacutaneous fibrolysis (Barra 2011), thoracic spinal manipulative therapy (Haik 2014; Kardouni 2014), lateral cervical glide mobilisation (McClatchie 2009), chiropractic shoulder girdle adjustments (Munday 2007), scapular mobilisation (Surenkok 2009), and mobilisation with movement (Teys 2008). The placebo treatment consisted of sham mobilisation or manipulation in six trials (Barra 2011; Haik 2014; Kardouni 2014; McClatchie 2009; Surenkok 2009; Teys 2008) and detuned ultrasound in one trial (Munday 2007). Six trials were considered to be at unclear risk of bias overall due to unclear allocation concealment (Barra 2011; Haik 2014; McClatchie 2009; Surenkok 2009; Teys 2008) or unclear blinding of outcome assessment and attrition (Munday 2007), while one was rated at low risk of bias overall (Kardouni 2014).

Barra 2011 found that participants receiving one session of diacutaneous fibrolysis were more likely to report global treatment success (RR 2.14, 95% CI 1.06 to 4.34; 50 participants) and have greater improvement in active shoulder abduction, flexion and internal rotation immediately post-treatment (see Table 10). Across the trials there were no clinically important differences between groups in overall pain (Barra 2011, Kardouni 2014; McClatchie 2009, Munday 2007, Surenkok 2009), function (Kardouni 2014; Surenkok 2009), pain on motion (Haik 2014), or quality of life (Kardouni 2014). Three trials measured adverse events (Barra 2011; Munday 2007; Teys 2008); none were reported in Barra 2011 and Teys 2008 while in Munday 2007 there were no reports of serious adverse reactions to shoulder girdle adjustment (such as persistent severe stiffness or pain) although there were five reports of minor, temporary posttreatment soreness (Table 10). We downgraded by one point for unclear risk of allocation bias in most trials, and one point for imprecision in all trials, and thus consider this evidence to be low quality.

#### Manual therapy alone versus no treatment

We judged all three trials that compared manual therapy to no treatment to be at high risk of bias overall. Van den Dolder 2003 found that two weeks of soft tissue massage led to less overall pain (MD -22.00, 95% CI -41.19 to -2.81; 100-point scale, 29 participants), better function (MD 7.20, 95% CI 2.20 to 12.20; 30point scale, 29 participants), and more active shoulder abduction, flexion and internal rotation than no treatment. Further, one session of mobilisation with movement increased active shoulder elevation in Teys 2008. In contrast, Surenkok 2009 found that one session of scapular mobilisation led to no important differences in overall pain (MD -8.96, 95% CI -33.01 to 15.09; 100-point scale, 26 participants), function (MD 9.07, 95% CI -12.09 to 30.23; 100-point scale, 26 participants), pain on motion (MD -2.08, 95% CI -19.49 to 15.33; 100-point scale, 26 participants) and active shoulder abduction or flexion when compared with no treatment (Table 11). We downgraded by two points for high risk of performance and detection bias, one point for imprecision and one point for



inconsistency, and thus consider this evidence to be very low quality.

#### Manual therapy alone versus another active intervention

We judged the four trials that compared manual therapy to another active intervention to be at high risk of bias overall. Al Dajah 2014 found soft tissue mobilisation plus proprioceptive neuromuscular facilitation resulted in statistically significantly less overall pain (MD -1.43, 95% CI -1.97 to -0.89; 10-point scale, 30 participants) and greater external rotation immediately post-treatment compared with therapeutic ultrasound, though these differences were not clinically important. Bansal 2011 compared deep friction massage with therapeutic ultrasound, and found no clinically important differences between groups in overall pain (MD -0.7, 95% CI not estimable; 10-point scale, 40 participants) and active shoulder abduction at the end of 10 days' treatment. Kaya 2014 found no clinically important differences between manual therapy and kinesiotaping with respect to rest pain (MD -0.32, 95% CI -1.48 to 0.84; 10-point scale, 54 participants), function (MD -3.10, 95% CI -11.40 to 5.20; 100-point scale, 54 participants), and pain on motion (MD 1.19, 95% CI -0.02 to 2.40; 10-point scale, 54 participants) at 6 weeks, but night pain was higher in the manual therapy group (MD 1.91, 95% CI 0.47 to 3.35; 10-point scale, 54 participants). Winters 1997 found that participants receiving shoulder manipulation once a week for six weeks had overall pain that was statistically significantly higher (MD 3.40, 95% CI 1.34 to 5.46; 28-point scale, 79 participants) and were half as likely to have global treatment success (RR 0.49, 95% CI 0.33 to 0.73; 79 participants) at 11 weeks than participants receiving glucocorticoid injection (Table 12). We downgraded by two points for high risk of performance and detection bias, and one point for imprecision, and thus consider this evidence to be very low quality.

# Is manual therapy delivered in addition to another active intervention more effective than the other active intervention alone?

Thirteen trials examined whether there is benefit in adding manual therapy (either mobilisation, manipulation or massage) to another physical therapy intervention (either an exercise programme, an electrotherapy modality or multi-modal physical therapy) (Atkinson 2008; Bang 2000; Barbosa 2008; Barra Lopez 2013; Bialoszewski 2011; Clews 1987; Conroy 1998; Cook 2014; Janse van Rensburg 2012; Kachingwe 2008; Kromer 2013; Senbursa 2011; Yiasemides 2011). All except one trial were rated at high risk of bias overall due to lack of participant blinding; Conroy 1998 blinded participants but was at unclear risk of bias overall due to unclear allocation concealment. Due to the heterogeneous diagnoses of shoulder pain and content of interventions, we chose not to synthesise any data in meta-analyses.

For overall pain, seven out of nine trials reported mean differences favouring the group with manual therapy as an add-on, but the difference was clinically important in only three of these trials (Bang 2000; Bialoszewski 2011; Conroy 1998). For function, five out of eight trials had mean differences favouring the group with manual therapy as an add-on; however none of the differences in any trial were clinically important. Only four trials measured adverse events (Atkinson 2008; Cook 2014; Janse van Rensburg 2012; Kromer 2013) and none were reported by any participant. Pain on motion was measured in one trial (Bang 2000), where a clinically important difference favouring the group with manual therapy as an add-on was noted. There were only slight differences between groups in the number of participants with global treatment success in the three trials which measured this outcome (Barra Lopez 2013; Kromer 2013; Yiasemides 2011). Of seven trials measuring range of motion (Atkinson 2008; Barra Lopez 2013; Bialoszewski 2011; Conroy 1998; Janse van Rensburg 2012; Kachingwe 2008; Yiasemides 2011), only Barra Lopez 2013 and Bialoszewski 2011 found a statistically significant, albeit small, difference between groups on some measures (Table 13). The overall impression from these trials is that adding manual therapy to another physical therapy intervention infrequently conferred clinically important benefits over the other physical therapy intervention alone. We downgraded by two points for high risk of performance and detection bias, and one point for imprecision, and thus consider the evidence from these 13 trials to be very low quality.

# Are supervised or home exercises alone more effective than placebo, no intervention or another active intervention?

In nine trials, an exercise programme delivered alone was compared with either placebo (Brox 1993), no intervention (Kachingwe 2008; Lombardi 2008; Ludewig 2003) or another active intervention (Brox 1993; Engebretsen 2009; Ginn 2005; Giombini 2006; Moosmayer 2014; Walther 2004).

#### Exercises alone versus placebo

In Brox 1993, supervised and home exercises were compared with inactive (placebo) laser (each delivered twice a week for six weeks). The trial was judged to be at unclear risk of bias overall due to unclear allocation concealment. Mean differences favouring the exercises group were noted for overall pain (MD 10; 35-point scale, 80 participants) and function (MD 10; 30-point scale, 80 participants) at six months, although the data were incompletely reported so 95% CIs were not estimable. The authors stated that there were no statistically significant differences between groups in pain on motion, global treatment success, night pain, range of motion, or number of days on sick leave (Table 14). No participant in either group reported adverse events. We downgraded by one point for unclear risk of allocation bias, and one point for imprecision, and thus consider this evidence to be low quality.

#### Exercises alone versus no treatment

Benefits of exercise alone when compared with no treatment were observed in two trials (Lombardi 2008; Ludewig 2003), both at high risk of bias overall. Lombardi 2008 found participants receiving progressive resistance training exercises twice a week for eight weeks had less overall pain (MD -1.90, 95% CI -3.27 to -0.53; 10point scale, 60 participants), disability (MD -15.50, 95% CI -28.94 to -2.06; 100-point scale, 60 participants), and pain on motion (MD -1.90, 95% CI -3.05 to -0.75; 10-point scale, 60 participants) at two months; these differences were clinically important. Also, statistically significant differences favouring the exercise group were noted for active shoulder internal rotation and measures of strength (Table 15). However, active shoulder abduction, flexion and external rotation were not significantly different between groups, nor were quality-of-life scores on the SF-36. In Ludewig 2003, participants receiving a daily home exercise programme for 10 weeks had better function (MD 6.90, 95% CI 0.59 to 13.21; 83point scale, 62 participants) and less work-related pain (MD -1.30, 95% CI -2.10 to -0.50; 10-point scale, 62 participants) and workrelated disability (MD -1.20, 95% CI -2.00 to -0.40; 10-point scale, 62 participants) at 10 weeks; however, these effects were not clinically

important. Usable outcome data were not available in Kachingwe 2008, though the authors claimed that there were no statistically significant differences between groups in overall pain, function and active shoulder flexion at six weeks (Table 15). We downgraded by two points for high risk of performance and detection bias, one point for imprecision and one point for indirectness because the trial by Ludewig 2003 was restricted to construction workers. Therefore, this evidence was considered very low quality.

#### Exercises alone versus another active intervention

Other active interventions which have been compared with an exercise programme alone include tendon repair surgery for rotator cuff tear (Moosmayer 2014), radial extracorporeal shockwave treatment (Engebretsen 2009), microwave diathermy (Giombini 2006), therapeutic ultrasound (Giombini 2006), glucocorticoid injection (Ginn 2005), arthroscopic subacromial decompression (Brox 1993), and a functional brace (Walther 2004).

In Moosmayer 2014, mini-open or open tendon repair surgery was compared with supervised exercises (twice weekly for 12 weeks, with increasing intervals during the following six to 12 weeks). We judged the trial to be at high risk of bias overall. Overall pain was higher in the exercise group at six months (MD 1.60, 95% CI 0.90 to 2.30; 10-point scale, 103 participants), 12 months (MD 1.20, 95% CI 0.60 to 1.80; 10-point scale, 103 participants) and five years (MD 1.00, 95% CI 0.20 to 1.80; 10-point scale, 103 participants), although the difference was clinically important only at six months. The authors observed no clinically important difference between the supervised exercise group and surgery group in function at six months (MD -2.80, 95% CI -10.10 to 4.50; 100-point scale, 103 participants), 12 months (MD -8.50, 95% CI -15.00 to -1.90; 100-point scale, 103 participants) or five years (MD -6.50, 95% CI -13.60 to 0.70; 100-point scale, 103 participants). Active shoulder abduction was lower in the exercise group at 12 months (MD -16.80, 95% CI -32.40 to -1.20; 103 participants) and five years (MD -14.70, 95% CI -29.40 to -0.10; 103 participants), although differences in active shoulder flexion and strength were not clinically important. The lack of participant blinding may have influenced participants in both groups to pursue alternative interventions and influenced their responses to self-reported outcomes, which may have biased results in either direction. For this reason we downgraded by two points for high risk of performance and detection bias, and consider this evidence to be low quality.

In Engebretsen 2009, radial extracorporeal shockwave treatment (once weekly for four to six weeks) was compared with supervised exercises (twice weekly for up to 12 weeks). We judged the trial to be at high risk of bias overall. There was no clinically important difference between groups in overall pain at six weeks (MD -0.3, 95% CI -0.9 to 0.4; 9-point scale, 103 participants), 18 weeks (MD -0.2, 95% CI -0.7 to 0.3; 9-point scale, 103 participants) and 1 year (MD -0.5, 95% CI -1.22 to 0.22; 9-point scale, 94 participants), or in pain on motion at six weeks (MD -0.7, 95% CI -1.6 to 0.1; 9point scale, 103 participants), 18 weeks (MD -0.6, 95% CI -1.3 to 0.2; 9-point scale, 103 participants) and 1 year (MD -0.2, 95% CI -1.13 to 0.73; 9-point scale, 94 participants). The authors found that participants receiving supervised exercises had less disability at six weeks (MD -10, 95% CI -17.6 to -2.3; 100-point scale, 103 participants), 18 weeks (MD -8.4, 95% CI -16.5 to -0.6; 100-point scale, 103 participants), and one year (MD -3.9, 95% CI -14.04 to 6.24; 100-point scale, 94 participants), although none of these differences were considered to be clinically important. The number of participants working at 18 weeks was higher in the supervised exercise group (RR 1.46, 95% CI 1.07 to 1.99; 100 participants), but work disability occurred at a similar frequency in both groups at 1 year (RR 1.1, 95% CI 1.0 to 1.2; 91 participants). One participant in the exercise group and two in the shockwave group had the adverse event of aggravation of pain after treatment (RR 0.50, 95% CI 0.05 to 5.34; 100 participants). A limitation of the trial was that more participants receiving shockwave treatment sought additional care outside of the trial setting, which may have biased results against the exercise group. We downgraded by two points for high risk of performance and detection bias, and consider this evidence to be low quality.

In Giombini 2006, microwave diathermy (three times a week for four weeks) was compared with exercises (once a week supervised and daily at home for four weeks). We judged the trial to be at high risk of bias overall. The authors observed clinically important differences favouring microwave diathermy over exercises in terms of overall pain at four weeks (MD 2.90, 95% CI 2.45 to 3.35; 10-point scale, 25 participants) and 10 weeks (MD 3.70, 95% CI 3.08 to 4.32; 10-point scale, 25 participants), function at four weeks (MD -16.90, 95% CI -20.26 to -13.54; 100-point scale, 25 participants) and 10 weeks (MD -18.73, 95% CI -23.18 to -14.28; 100-point scale, 25 participants), and global treatment success (number of patients returning to sport) at 10 weeks (RR 0.42, 95% CI 0.19 to 0.95; 25 participants). No participant reported adverse events. We downgraded by two points for high risk of performance and detection bias, one point for imprecision and one point for indirectness because the trial was restricted to professional athletes, and thus consider this evidence to be very low quality.

All differences in outcomes between exercise and the remaining active interventions (therapeutic ultrasound (Giombini 2006), glucocorticoid injection (Ginn 2005), arthroscopic subacromial decompression (Brox 1993), and functional brace (Walther 2004)) were not clinically important or statistically significant (see Table 16). We downgraded the evidence from these four trials by two points for high risk of performance and detection bias, and one point for imprecision, and thus consider it to be very low quality.

#### Are supervised or home exercises delivered in addition to another active intervention more effective than the other active intervention alone?

Six trials investigated the effects of exercise as an add-on to another intervention (Ainsworth 2009; Bae 2011; Baskurt 2011; Beaudreuil 2011; Maenhout 2013; Martins 2012). The trials investigated the effects of adding a package of strengthening, stretching, and rangeof-motion exercises to ultrasound, glucocorticoid injection and advice (Ainsworth 2009), adding motor control and strengthening exercises to heat pack, TENS and ultrasound (Bae 2011), adding scapular stabilisation exercises to stretching and strengthening exercises (Baskurt 2011), adding Dynamic Humeral Centering to massage and exercise (Beaudreuil 2011), adding heavy load eccentric training to traditional rotator cuff training (Maenhout 2013), or adding proprioception exercises to stretching and strengthening exercises plus cryotherapy (Martins 2012). The overall risk of bias was low in one study (Beaudreuil 2011), unclear in two trials (Bae 2011; Baskurt 2011), and high in three trials (Ainsworth 2009; Maenhout 2013; Martins 2012).

The addition of exercises resulted in better function in Ainsworth 2009, Bae 2011, Beaudreuil 2011, and Maenhout 2013, but the



difference was only clinically important in Bae 2011. No clinically important differences between groups were observed for overall pain (Baskurt 2011; Beaudreuil 2011; Martins 2012), pain on motion (Baskurt 2011), global treatment success (Maenhout 2013), quality of life (Ainsworth 2009; Baskurt 2011; Martins 2012) or strength (Bae 2011; Baskurt 2011; Beaudreuil 2011; Maenhout 2013), although Bae 2011 found that the "exercise add-on" group had better active range of motion (Table 17). We downgraded the evidence from these six trials by two points for high or unclear risk of performance and detection bias in all but one trial, and one point for imprecision, and thus consider it to be very low quality.

# Is one type of manual therapy or exercise more effective than another?

Eighteen trials compared one type of manual therapy or exercise with another. Trials compared:

- eccentric progressive resistance exercises versus concentric progressive resistance exercises (Blume 2014)
- exercises below 90 degrees flexion versus exercises above 90 degrees flexion (Celik 2009)
- manual mobilisation versus proprioceptive neuromuscular facilitation (Citaker 2005)
- mobilisation with movement and taping versus supervised exercises (Djordjevic 2012)
- soft tissue techniques versus mobilisation, proprioceptive neuromuscular facilitation and exercise (Heredia-Rizo 2013)
- specific exercise programme targeting the rotator cuff and scapular stabilisers versus non-specific movement exercises for the neck and shoulder (Holmgren 2012)
- glenohumeral mobilisation versus mobilisation with movement (Kachingwe 2008)
- classic Swedish massage versus massage based on the tensegrity principle (Kassolik 2013)
- self-managed loaded exercise programme versus multi-modal physiotherapy (Littlewood 2014)
- neurocognitive therapeutic exercise versus traditional therapeutic exercise (Marzetti 2014)
- high-dose exercise programme versus low dose exercise programme (Osteras 2008)
- manual therapy programme versus self-training programme (Senbursa 2007)
- supervised exercises versus home exercises (Senbursa 2011)
- scapular-focused treatment versus stretching, muscle friction and eccentric rotator cuff training (Struyf 2013)
- water-based exercise programme versus land-based exercise programme (Subasi 2012)
- self-training centring and stretching exercises versus supervised stretching exercises (Walther 2004)
- customised exercises versus standardised exercises (Wang 2006)
- massage and supervised exercises versus manipulation (Winters 1997).

The overall risk of bias was low in one trial (Holmgren 2012), unclear in eight due to unclear allocation concealment, participant blinding or attrition (Blume 2014; Citaker 2005; Djordjevic 2012; Heredia-Rizo 2013; Kachingwe 2008; Marzetti 2014; Struyf 2013; Winters 1997), and high in nine due to lack of participant blinding or allocation concealment (Celik 2009; Kassolik 2013; Littlewood 2014; Osteras 2008; Senbursa 2007; Senbursa 2011; Subasi 2012; Walther 2004; Wang 2006)

One participant-blinded trial (Holmgren 2012) investigated the effects of a 12-week specific exercise programme targeting the rotator cuff and scapular stabilisers (strengthening eccentric exercises for the rotator cuff and concentric/eccentric exercises for the scapula stabilisers) compared with 12 weeks of nonspecific movement exercises for the neck and shoulder (without any external load). All participants received glucocorticoid injection prior to the exercise programme. For all outcomes assessed at three months, statistically significant differences favouring the specific exercise programme were found, most of which were clinically important: overall pain (MD -10.00, 95% CI -18.18 to -1.82; 100-point scale, 97 participants), function (MD 20.00, 95% CI 11.55 to 28.45; 100-point scale, 97 participants), pain on motion (MD -16.00, 95% CI -26.57 to -5.43; 100-point scale, 97 participants), global treatment success (RR 2.87, 95% CI 1.66 to 4.96; 97 participants), night pain (MD -12.00, 95% CI -21.87 to -2.13; 100-point scale, 97 participants), quality of life (MD 0.13, 95% CI 0.05 to 0.21; scale range from -0.59 to 1, 97 participants), and having surgery at some point between three months and one year follow-up (RR 0.37, 95% CI 0.22 to 0.64; 97 participants). We considered the evidence from this trial to be high quality.

Of the remaining 17 trials, 11 found no clinically important differences between groups on any outcome (Blume 2014; Celik 2009; Citaker 2005; Kachingwe 2008; Littlewood 2014; Marzetti 2014; Senbursa 2007; Senbursa 2011; Walther 2004; Wang 2006; Winters 1997). In the other six trials (Djordjevic 2012; Heredia-Rizo 2013; Kassolik 2013; Osteras 2008; Struyf 2013; Subasi 2012), some statistically significant differences in outcomes were noted (see Table 18). We downgraded the evidence from these 17 trials by two points for high or unclear risk of performance and detection bias, and one point for imprecision, and thus consider it to be very low quality.

# Subgroup and sensitivity analyses, and assessment of publication bias

Given the inability to conduct meta-analyses, no subgroup or sensitivity analyses were undertaken. Also, we were unable to generate funnel plots to assess small study effects. Despite the lack of funnel plots, we considered the risk of publication bias to be low because nearly all of the published studies reported statistically non-significant results for most outcomes. It is possible that some unpublished studies with non-significant results exist, but their inclusion in the review is unlikely to change our conclusions.

#### DISCUSSION

### Summary of main results

In this systematic review we have considered the results of 60 trials investigating the benefits and harms of manual therapy and exercise for rotator cuff disease. The combination of manual therapy and exercise (the most clinically relevant intervention (Klintberg 2015)) was examined in 10 trials (Bennell 2010; Cloke 2008; Dickens 2005; Ginn 2005; Haahr 2005; Hay 2003; Kachingwe 2008; Rhon 2014; Szczurko 2009; Winters 1997), but the variation in intervention content and comparators meant trials could not be pooled.



High quality evidence from one trial of 120 participants with chronic rotator cuff disease indicated no clinically important differences between manual therapy and exercise and placebo with respect to overall pain, function, pain on motion, global treatment success, quality of life and strength at 22 weeks, although manual therapy and exercise was associated with relatively more frequent but mild adverse events (short-term pain following treatment) (Bennell 2010). Very low quality evidence from two trials that compared manual therapy and exercise to no treatment was broadly consistent with these results (Dickens 2005; Kachingwe 2008).

Low quality evidence from five trials (Cloke 2008; Ginn 2005; Hay 2003; Rhon 2014; Winters 1997) revealed no clinically important differences between manual therapy and exercise and glucocorticoid injection with respect to overall pain, function, quality of life, night pain and active range of motion up to 12 months. However, global treatment success was more common up to 11 weeks in participants receiving glucocorticoid injection (based on low quality evidence). Low quality evidence from one trial showed no important differences between manual therapy and exercise and arthroscopic subacromial decompression with respect to overall pain, function, active range of motion and strength at six and 12 months, or global treatment success at four to eight years (Haahr 2005). Low quality evidence from one trial found that manual therapy and exercise may not be as effective as acupuncture plus dietary counselling and Phlogenzym supplement with respect to overall pain, function, active shoulder abduction and quality-of-life at 12 weeks in postal workers (Szczurko 2009). Very low quality evidence from one trial suggested that firstly, there was no important difference between manual therapy and exercise compared with oral NSAID, and secondly, no added benefit of manual therapy and exercise over glucocorticoid injection alone with respect to function at 18 weeks and 12 months (Cloke 2008).

### **Overall completeness and applicability of evidence**

Participants in the included trials were mostly representative of populations most affected by rotator cuff disease. Nearly all trials enrolled a community sample of people attending routine physical therapy care. Across the trials, the gender ratio was equal, and the median age was 51 (IQR 46 to 56) years. Thus, results are applicable to both male and female older adult populations, which is useful given that the incidence of rotator cuff disease increases with age (Linsell 2006; Yamamoto 2010). Further, trials were conducted in 21 different countries, including a range of high- and low- to middle-income countries. However, it is difficult to determine how representative participants in the included trials were with respect to duration of symptoms, as this characteristic was not reported in 31 (52%) trials.

Manual therapy and exercise are most often delivered together in physical therapy practice (Glazier 1998; Kooijman 2013; Roberts 2014; Struyf 2012), yet the effects of this multi-modal intervention were investigated in only 10 (17%) trials. Our review was dominated by trials investigating whether manual therapy or exercise provided benefit when added to another physical therapy intervention (e.g. manual therapy plus therapeutic ultrasound versus therapeutic ultrasound alone), or whether one type of manual therapy or exercise intervention was more effective than another.

In several trial reports, the components of the exercise programmes were incompletely described. For example, some trialists specified the type of exercise broadly (e.g. "range of movement exercises") without specifying the planes of movement addressed, the frequency of exercises (e.g. once or twice per day) or the setting in which they were undertaken (e.g. clinic or home). Incomplete descriptions such as these hinder replication of the trial, and limit reliable implementation of the exercise programme into clinical practice. A standardised and internationally agreed template for explicit reporting of exercise programmes has recently been developed (Slade 2014; Slade 2016). This guidance will hopefully improve the quality of reporting in future trials.

Another concerning issue is the variable choice of outcomes measured in the trials. Overall pain and function were measured commonly (80% and 73%, respectively), but these domains should be measured in all rotator cuff disease trials given that pain and functional limitations are the most common presenting symptoms of the condition (Whittle 2015). Further, adverse events were measured in less than a third of trials (28%). The proportions of trials measuring the other main outcomes of the review were relatively low: pain on motion (27%), global assessment of treatment success (28%), and quality of life (22%). Outcome measurement has certainly improved since the first version of our review (Green 1998), where function was measured in only 26% of trials (none with a validated disability index), and quality of life was measured in no trials. A core domain set and core outcome measurement set for rotator cuff disease trials would likely improve measurement of patient-important outcomes in future trials, and would facilitate efforts to synthesise the evidence in future (Buchbinder 2003; Page 2015). We are currently developing these core sets with the support of the Outcome Measures in Rheumatology (OMERACT) initiative, who approved a special interest group session on shoulder pain at the OMERACT 2016 meeting.

#### **Quality of the evidence**

Although we presented 'Summary of findings' tables only for trials addressing the primary questions of the review, we used the GRADE approach (Schünemann 2011b) to assess the quality of all the evidence examined. Most of the evidence was downgraded to low quality for a combination of two out of three reasons. Firstly, the risk of performance and detection bias for self-reported outcomes was high, secondly, evidence was based on small, single trials, leading to concerns about imprecise effect estimates, and thirdly, trialists examined a sample of people whose outcomes may not apply to the general population (e.g. construction workers, professional athletes). Regarding the first of these concerns, we rated very few trials (22% and 20%, respectively) at low risk of performance bias and detection bias for self-reported outcomes because participants and personnel were not blinded.

Blinding of participants and personnel is difficult to achieve in procedural trials, so performance bias and detection bias are often difficult to minimise. However, this is problematic because it is estimated that trials with unblinded assessment of subjective outcomes (such as function and pain) exaggerate treatment benefits by 22% on average (ratio of odds ratios 0.78, 95% credible interval 0.65 to 0.92) (Savovic 2012). For most comparisons and outcomes in our review, the true effects of the interventions may be different to the effect estimates observed because of this bias. On the other hand, Bennell 2010 found that they achieved a moderate to high degree of participant blinding with the use of inactive ultrasound therapy and application of an inert gel as a

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placebo control, increasing confidence that their results present an unbiased estimate of the true effect of manual therapy and exercise. This control intervention has also been found to be a realistic placebo for physical therapy in other trials performed by this group (Bennell 2014; Buchbinder 2007).

#### Potential biases in the review process

We searched CENTRAL, MEDLINE, EMBASE and CINAHL, but not PEDro, a database of randomised trials, systematic reviews and clinical practice guidelines in physiotherapy. A study comparing the indexing of 400 physiotherapy trials in eight bibliographic databases found that almost all were indexed in CENTRAL (95%), PEDro (92%) MEDLINE (89%) and EMBASE (88%), and only one of the 400 trials was uniquely indexed in PEDro (Michaleff 2011). Therefore, it is very unlikely that we missed relevant trials that would change the conclusions of our review. Two review authors independently assessed the trials for inclusion in this review, extracted data and assessed the risk of bias, and a third review author adjudicated when any discrepancy arose. Two of the review authors (SG and RB) are authors of one of the trials included in this review (Bennell 2010). To avoid bias, the paper was sent to an independent review author for eligibility assessment. Neither review author was involved in data extraction or assessment of risk of bias of this trial. Review questions of interest were defined with full knowledge of the possible comparisons that could be undertaken, but no knowledge of the results of any comparisons. To prevent selective inclusion of results (Page 2013), we used predefined decision rules to select data from trials when multiple measurement scales, time points and analyses were reported.

A limitation of the review process was that several trials addressing the main questions of the review did not present data for all measured outcomes, or presented outcome data incompletely, which prevented us from calculating effect estimates and 95% CIs; attempts to obtain unpublished data from trialists were unsuccessful. Another potential limitation was that we excluded four trials (Miller 2004; Mörl 2011; Seok-Hwa 2013; Tachibana 2012) which may have included participants with rotator cuff disease, but the eligibility criteria and participant characteristics were not reported in enough detail for us to determine this. Further, we excluded one trial (Ginn 1997) that included participants with shoulder pain due to either rotator cuff disease, adhesive capsulitis, osteoarthritis, biceps muscle tear or no specific diagnosis, and we were unable to obtain data on the rotator cuff disease subgroup, which comprised 65% of the sample. However, none of these trials addressed the two main questions of the review, so our overall conclusions remain unchanged. In addition, we did not search for grey literature (e.g. proceedings of specific conferences, theses or unpublished reports). However, we believe that had we identified unpublished studies with non-significant results, their inclusion in the review would be unlikely to change our conclusions since the majority of the evidence we considered had 'negative' findings.

# Agreements and disagreements with other studies or reviews

Following the earlier Cochrane review of physical therapy for shoulder disorders (Green 2003), there have been several systematic reviews of manual therapy (Brudvig 2011; Camarinos 2009; Ho 2009; Pribicevic 2010), exercise (Dewhurst 2010; Hanratty 2012; Kelly 2010; Kuhn 2009; Littlewood 2012; Marinko 2011), and manual therapy and exercise (Brantingham 2011; Braun 2010; Braun 2013; Gebremariam 2014; Kromer 2009; Nyberg 2010; Saltychev 2015; Van den Dolder 2014) for rotator cuff disease. All of these reviews have been narrower in scope than ours. Review authors either restricted their study eligibility criteria according to the diagnostic label used by trialists (e.g. focusing only on subacromial impingement syndrome or rotator cuff tendinopathy), or used broad participant eligibility criteria but focused on only one type of manual therapy (e.g. soft tissue massage). Therefore, to our knowledge ours is the most comprehensive review of manual therapy and exercise interventions for rotator cuff disease.

Our conclusions about the benefits and harms of manual therapy and exercise, manual therapy alone and exercise alone are consistent with nearly all other systematic reviews. In a few cases, other review authors (Dewhurst 2010; Kuhn 2009; Marinko 2011) had more favourable conclusions than ours. However, these discrepancies appear to be driven by less frequent consideration of the overall quality of evidence in these reviews (i.e. while study risk of bias was assessed, other domains of the GRADE approach (imprecision, inconsistency, indirectness and publication bias) were not).

#### AUTHORS' CONCLUSIONS

#### **Implications for practice**

Despite identifying 60 trials meeting the inclusion criteria for this review, only one trial compared a combination of manual therapy and exercise reflective of common current practice to placebo. Based upon high quality data from this trial, there was no clinically important benefit of manual therapy and exercise over placebo. Adverse events were relatively more frequent with manual therapy and exercise but mild in nature (short-term pain following treatment). Effects of manual therapy and exercise may be similar to those of glucocorticoid injection and arthroscopic subacromial decompression, but this is based on low quality evidence. Until further evidence confirms or refutes these results, practitioners should communicate the uncertainty of effect and consider other approaches or combinations of treatment.

#### Implications for research

Novel combinations of manual therapy and exercise should be compared with realistic placebo (e.g. use of inactive ultrasound therapy and application of an inert gel) in high quality randomised trials. Further trials of manual therapy alone or exercise alone for rotator cuff disease should be based upon a strong rationale and consideration of whether or not they would alter the conclusions of this review. The interventions should be described in enough detail to inform interpretation of findings and allow replication. Trials should use strategies designed to minimise the potential for bias, including adequate allocation concealment and blinding of participants by delivering a realistic physical therapy placebo. Development of a core set of outcomes for trials of rotator cuff disease and other shoulder disorders would facilitate our ability to synthesise the evidence in future.

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\* Indicates the major publication for the study

# CHARACTERISTICS OF STUDIES

### Characteristics of included studies [ordered by study ID]

Design: Parallel group RCT		
<b>Design:</b> Parallel group RCT		
Setting: One district general hospital and 9 local community hospitals, United Kingdom		
Intervention: Exercise, therapeutic ultrasound, glucocorticoid injection and advice		
Control: Therapeutic ultrasound, glucocorticoid injection and advice		
Source of funding: Not reported		
Diagnostic label used by trialists: Rotator cuff tear		
Criteria for defining the shoulder condition being treated		
<ul> <li>Radiological diagnosis of full thickness tear of &gt; 5 cm</li> </ul>		
Inclusion Criteria (not listed above)		
Aged 18 years and over		
Able to give informed consent		
Exclusion criteria (not listed above)		
Neurological abnormality affecting shoulder complex		
Involved in industrial claim or litigation		
Rotator cuff considered to be repairable		
Baseline characteristics		



Ainsworth 2009 (Continued)	Number randomised: 3	30; mean (range) age = 78.4 (65-96) years; male/female = 14/16; duration of symp-	
	toms: not reported		
	Control		
	Number randomised: 3 toms: not reported	30; mean (range) age = 78 (68-88) years; male/female = 15/15; duration of symp-	
Interventions	Intervention: exercise-based rehabilitation programme		
	<i>Components of intervention</i> : the programme was usually started with the participant lying in a supine position. The participant was taught to start with a flexed elbow and to raise the arm to a vertical position. The participant was then taught to control the arm with sways in a 20 degree arc before elevating and lowering the arm using a weight of approximately 0.75 kg. When the participant could carry out these activities supine, the head of the treatment couch was gradually inclined until they were able to perform the exercises in a sitting position. The participants also carried out stretching exercises to improve ranges of elevation, internal and external rotation, resistance band exercises into internal and external rotation, activities to improve proprioception, posture correction and adaptation of functional activities.		
	Frequency of administration: 6 treatment sessions (unclear how many sessions per week)		
	Control: therapeutic ultrasound, glucocorticoid injection and advice		
	<i>Components of intervention</i> : participants only received the treatment common to both groups (see be- low)		
	Both groups		
	<i>Components of intervention</i> : therapeutic ultrasound, glucocorticoid injection if needed for pain, and advice		
	Dose: not reported		
	Frequency of administration: 6 treatment sessions (unclear how many sessions per week)		
Outcomes	Outcomes assessed at 3, 6 and 12 months		
	<ul> <li>Function: Oxford Shoulder Score, scored from 0 (worst score) to 48 (best score)</li> <li>Quality of life: SF-36 (0-100 scores with higher scores denoting better quality of life)</li> <li>Passive shoulder external rotation and internal rotation using a goniometer</li> </ul>		
Notes	Conflicts of interest: the authors declared that they had no conflicts of interest		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Quote: "Patients were allocated a numbered envelope in the sequential order that they were recruited to the trial. Inside each envelope was the allocation which was generated from a random allocation table by an independent sta- tistician".	
		Comment: An adequate method was used to generate the allocation sequence	
Allocation concealment (selection bias)	Low risk	Comment: See quote above. An adequate method was used to conceal the al- location sequence	

Manual therapy and exercise for rotator cuff disease (Review)

Ainsworth 2009 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "The weakness in this study, as with many physiotherapy trials, was the lack of blinding. Efforts were made to reduce bias in this study but the lack of blinding is acknowledged to be a potential source of bias"
All outcomes		Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported some out-comes
Blinding of outcome as- sessment (detection bias) Objective outcomes	High risk	Comment: Assessors of objective outcomes (ROM) were not blind to treatment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Of the 60 patients recruited to the trial six were lost to follow up. Two patients withdrew from the trial after recruitment but before treatment had begun. One patient became too ill to proceed with the trial and one patient was widowed and no longer wished to engage with rehabilitation. One patient died of unrelated causes after completing treatment but before the 3 month assessment. Two further patients died, one before the 6 month assessment from kidney failure and the other before the 12 month assessment from blad- der cancer. One patient was lost to follow up before the 12 month assessment when he moved out of the area without leaving a forwarding address."
		Quote: "Although six patients in total were lost to follow up during the course of the trial and all of them belonged to the intervention group, none of them withdrew due to lack of confidence in their treatment. Three patients who had received the intervention treatment died from unrelated factors dur- ing the course of the trial. Data for all six patients were included in the analysis of baseline characteristics and up to the point where they dropped out of the trial. Analysis of data was on an intention-to-treat basis whereby patients were compared in the groups to which they were originally randomly assigned."
		Comment: The approach to dealing with missing participant data was likely to have minimised bias in the results
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were incompletely reported for ROM, but unclear if this was related to the nature of the results. Also, without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias identified

Al Dajah 2014
---------------

Methods Study design: Parallel group RCT		
	Setting: Physiotherapy outpatient department, Saudi Arabia	
	Intervention: Soft tissue mobilisation and proprioceptive neuromuscular facilitation	
	Control: Therapeutic ultrasound	
	Source of Funding: Not reported	
Participants	Diagnostic label used by trialists: Shoulder impingement syndrome	

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Al Dajah 2014 (Continued)

## Criteria for defining the shoulder condition being treated

- · Positive results in the Neer impingement test
- Negative results in the capsule stretch test
- Visual analogue scale (VAS ≥ 5)
- External rotation = 35 degrees ± 5 degrees
- Overhead reach of 155 ± 10 cm

### Inclusion Criteria (not listed above)

- Aged between 40 and 60 years
- No use of analgesics and anti-inflammatory drugs and muscle relaxants within 24 hours before the participation in the study

## **Exclusion Criteria (not listed above)**

- Open wounds
- Infection
- Acute injuries or fractures
- Recent surgeries
- Swelling
- Rheumatoid arthritis
- Reflex sympathetic syndrome
- Adhesive capsulitis

### Baseline characteristics: not reported

#### Interventions

# Intervention: soft tissue mobilisation (STM) and proprioceptive neuromuscular facilitation (PNF)

*Components of intervention:* the subjects were positioned with the humerus abducted to 45 degrees with elbow flexed to 90 degrees, and the humerus was externally rotated to a midrange position, typically about 20 degrees to 25 degrees of check, degrees used elsewhere external rotation. The subscapularis was palpated in the axilla to identify areas of myofascial mobility restrictions, taut bands, or trigger points. Identified restrictions were treated with STM utilising a combination of sustained manual pressure, and slow deep strokes to the subscapularis myofascia for 7 min. The STM was followed by contract-relax PNF for the subscapularis and other glenohumeral medial rotators, beginning in the same position used for the STM. The participants were instructed to perform maximal glenohumeral internal rotation against an opposing, isometric, manual resistance applied by the treating physical therapist for 7 seconds. Afterwards, the participant actively moved the humerus into full available external rotation. This position was maintained for 15 seconds. This 7-second internal rotation contraction against resistance followed by full active external rotation was repeated 5 times. Subjects were then instructed to actively move through the PNF flexion-abduction external-rotation diagonal pattern for 5 repetitions with manual facilitation

### Dose: 10 min

Frequency of administration: once

### **Control: therapeutic ultrasound**

*Components of intervention:* the arm was abducted to 45 degrees and the forearm was rested on the pillow for support. Ultrasound therapy was given to the subscapularis muscle insertion at the shoulder region

Dose: frequency - 3 MHz; intensity - 0.5 W/cm<sup>2</sup>; duration: 10 min

Frequency of administration: once

Outcomes

Outcomes assessed immediately after one treatment session (day 1):

• overall pain: VAS (scale units not reported but assumed 0 - 10)

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Al Dajah 2014 (Continued)

	ROM: external rotation using a goniometer (unclear if active or passive)		
Notes	Conflicts of interest: not reported		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Quote: "The subjects were assigned randomly into two groups by lot method"	
		Comment: An adequate method was used to generate the allocation sequence	
Allocation concealment (selection bias)	Unclear risk	Comment: There was no information on how the allocation sequence was con- cealed	
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention	
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants, who may have had different expectations about the benefits of the intervention they received, self-reported pain	
Blinding of outcome as- sessment (detection bias) Objective outcomes	Unclear risk	Comment: There was no information about whether assessors of objective outcomes were blinded	
Incomplete outcome data (attrition bias)	Low risk	Comment: There was no attrition because all participants were treated and as- sessed in a single session	

All outcomes		
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias identified

### Atkinson 2008

Methods	Study design: Parallel group RCT	
	Setting: University, South Africa	
	Intervention: Manipulation plus mobilisation	
	Control: Placebo laser treatment plus mobilisation	
	Source of Funding: Department of Chiropractic at Durban University of Technology	
Participants	Diagnostic label used by trialists: Rotator cuff tendinopathy	
	Criteria for defining the shoulder condition being treated	
	Diagnosis of rotator cuff tendinopathy by the researcher and confirmed by a specially trained physician and doctor of chiropractic and 3 of:	

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Atkinson 2008 (Continued)

- palpable tenderness along the anterior edge of the acromion
- a painful arc of abduction between 60 and 120 degrees
- a positive shoulder abduction (empty-can supraspinatus) stress test

### Inclusion Criteria (not listed above)

· Restricted motion/glide or shear of the acromioclavicular and/or glenohumeral joint

### **Exclusion Criteria (not listed above)**

- History of traumatic dislocation
- Instability indicating rupture
- A positive drop arm test indicating rupture
- Significant bony crepitus (a loose body, labral defect, advanced arthritis)
- Pain radiating distally below the elbow
- Shoulder surgery in the two previous years
- · Cardiac, pulmonary or systemic disease that refers pain to the shoulder found on exam
- If diagnosis required clarification of imaging studies
- Treatment for the shoulder within six weeks
- No joint acromioclavicular or glenohumeral joint dysfunction found per Shafer and Faye's description and the "PARTS" formula per Peterson and Bergmann

### **Baseline characteristics**

#### Intervention

Number randomised: 30; mean (range) age = 41.53 (18-63) years; male:female = 22:8; duration of symptoms: not reported

#### Control

Number randomised: 30; mean (range) age = 42.00 (20-76) years; male:female = 21:9; duration of symptoms: not reported

Interventions

### Intervention: manipulation

*Components of intervention:* high velocity, low-amplitude, gentle-impulse, shoulder adjustive thrust based on extensive motion palpation (completed at the 1st, 3rd and 6th visits) of the shoulder to detect restriction. The participant was positioned either sitting or lying

Dose: not reported

Frequency of administration: 6 sessions across a 2-week period

### Control: sham laser

*Components of intervention:* motion palpation (equivalent to grades 3 and 4 mobilisation) of the shoulder. Laser unit set to zero. The participant was seated in a comfortable position with the shoulder girdle exposed

### Dose: 5 min

Frequency of administration: 6 sessions across a 2-week period

### **Both groups**

Both groups received full-motion palpation of the shoulder prior to randomisation and before the 1st visit, to assess restriction. This equated to grades 3 and 4 mobilisations of the shoulder

Outcomes

Outcomes assessed at the 1st, 3rd and 6th visits in the 2-week period, except for ROM in the Manipulation group, which was measured at each of the 6 visits

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Atkinson 2008 (Continued)

- Pain using a numeric rating scale (NRS-101)
- ROM (shoulder flexion, extension, abduction, adduction, external rotation, horizontal abduction); unclear if active or passive
- Adverse events

Conflicts of interest: not reported

## Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "Randomization was accomplished by folding 60 sheets of paper, 30 marked Group 1, 30 marked Group 2, and mixing them together thoroughly to assure discontinuity. They were then placed in a box. At each subject randomization time point, the box was held to ensure all folded slips were completely obscured".
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Low risk	Quote: "Randomization was accomplished by folding 60 sheets of paper, 30 marked Group 1, 30 marked Group 2, and mixing them together thoroughly to assure discontinuity. They were then placed in a box. At each subject randomization time point, the box was held to ensure all folded slips were completely obscured".
		Comment: An adequate method was used to conceal the allocation sequence
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "Participants were informed they might be randomized into either group, treatment or placebo".
		Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants, who may have had different expectations about the benefits of the intervention they received, self-reported pain
Blinding of outcome as- sessment (detection bias) Objective outcomes	High risk	Quote: "This study would have been improved by a fully powered sample, the addition of a blind assessor."
		Comment: ROM and algometry were assessed by a non-blinded assessor
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Quote: "Five patients dropped out of Group 2 (placebo) and were replaced. Therefore out of a total of 35 [control] patients, 30 patients completed treat- ment. No patients dropped out of Group 1 and no patients complained or dropped out of the trial because of significant side effects such as persistent severe stiffness and/or pain"
		Comment: It is not clear when in the process and why these participants dropped out
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: The study appears to be free of other bias

Manual therapy and exercise for rotator cuff disease (Review)



Methods	Study design: Parallel group RCT			
	Setting: University, South Korea			
	<b>Intervention:</b> Motor control exercises and strengthening exercises plus hot packs plus TENS plus ultra sound			
	Control: Hot packs plus TENS plus ultrasound			
	Source of Funding: Supported by Sahmyook University			
Participants	Diagnostic label used by trialists: Impingement			
	Criteria for defining the shoulder condition being treated			
	At least one of:			
	<ul> <li>painful arc of movement during flexion or abduction</li> <li>positive Neer or Kennedy-Hawkins impingement signs</li> <li>pain on resisted lateral rotation or the Jobe test</li> </ul>			
	Inclusion Criteria (not listed above)			
	• None			
	Exclusion Criteria (not listed above)			
	<ul> <li>type III acromion</li> <li>calcification or fracture</li> <li>shoulder instability</li> <li>previous shoulder surgery</li> <li>cervicobrachialgia or shoulder pain during neck movement</li> </ul>			
	Baseline characteristics			
	Intervention			
	Number randomised: 17; mean age: 49.9 $\pm$ 5.5 years old; sex: F/M 11/6; duration of symptoms: not reported			
	Control			
	Number randomised: 18; mean age: 48.3 ± 4.3 years old; sex: F/M 12/6; duration of symptoms: not re- ported			
Interventions	Intervention: motor control and strengthening exercises			
	Components of intervention			
	<ul> <li>Motor control exercises: motor control training was performed to increase the mobility of the scapula against gravity during arm elevation. Shoulder control progressed following six-phase retraining exercises to control arm elevation in the frontal, sagittal and scapular planes. Movement training was performed under the supervision of a physiotherapist who gave feedback aimed at correcting the shoulder girdle movement. The retraining phases were graded according to the level of resistance applied to the shoulder during arm elevation (no resistance/passive movement, active assisted, active with or without external resistance) and the use of feedback during the movement. The phases ranged from no resistance with feedback to active movement with external resistance without feedback. Durine each retraining phase, the ROM was gradually increased as shoulder control improved until proper control was achieved for the full ROM in each vertical plane. When the subject was able to perform series of 10 repetitions with proper control, exercise series were added to reach 3 in total. The subject then moved up to the next phase. Once abduction over a range of 90 degrees was properly controlled humeral lateral rotation at 90 degrees of abduction was performed</li> </ul>			

Manual therapy and exercise for rotator cuff disease (Review)



Bae 2011 (Continued)				
(continued)	around the scapulo nal rotation and int low trapezius exerci the ROM, the repeti	cises: the strengthening exercise was performed to increase the muscle strength thoracic and scapulohumeral joints. The strengthening exercises included; exter- ternal rotation, scaption, chair press, push-up plus, press-ups, upright rows, and ise. The intensity of the exercises was assessed according to the movement plane, tions, the velocity and the resistance. A 10-min rest period was provided between and the strengthening exercise. All exercises were performed pain free		
	Dose: 30 min of exercis	e		
	<ul> <li>Control: the exercis joint</li> <li>Strength: 3 sets of 1</li> </ul>	e intensity was adjusted for the movement pattern and the pain in the shoulder 0 repetitions		
	Frequency of administration: 3 times per week for 4 weeks Control: hot packs plus TENS plus ultrasound			
	<i>Components of intervention</i> : participants only received the treatment common to both groups (see be- low) <b>Both groups</b>			
	<i>Components of interver</i> sound. No other detail	<i>ntion</i> : conservative physical therapy including applied hot packs, TENS and ultra- ed provided		
	Dose: 45 min per day (20 min of hot packs), (20 min of TENS), (5 min of therapeutic ultrasound)			
	Frequency of administration: 3 times per week for 4 weeks			
Outcomes	Outcomes assessed at 4 weeks			
	<ul> <li>Function: SPADI scale total score (0-100)</li> <li>Active ROM (flexion, extension, abduction, external rotation, internal rotation) using a goniometer</li> <li>Strength: concentric isokinetic evaluations (peak torque of external rotator, peak torque of interrotator, each at speed of 60 degrees/second and 180 degrees/second) using an isokinetic dynamo ter</li> </ul>			
Notes	Conflicts of interest: not reported			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence genera-	Unclear risk	Quote: "The thirty-five participants were randomly assigned to two groups"		
tion (selection bias)		Comment: No information about how the allocation sequence was generated was reported		
Allocation concealment (selection bias)	Unclear risk	Comment: No information about how the allocation sequence was concealed was reported		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Comment: Participants received different multimodal interventions, but it is unclear whether they were provided any information that would make them perceive the intervention they received as superior or inferior to the alterna- tive intervention		
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	Unclear risk	Comment: Participants self-reported pain and function, but it is unclear whether they were provided any information that would make them perceive the intervention they received as superior or inferior to the alternative inter- vention		

Manual therapy and exercise for rotator cuff disease (Review)

## Bae 2011 (Continued)

Blinding of outcome as- sessment (detection bias) Objective outcomes	Unclear risk	Comment: There was no information about whether assessors of strength and ROM were blind to treatment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: No attrition was reported and outcome data were reported as based on the total number of randomised participants
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: The study appears to be free of other bias

Methods	Study design: Parallel group RCT			
	Setting: Physical therapy clinic, US			
	Intervention: Manual physical therapy plus supervised flexibility and strengthening exercises			
	Control: Supervised flexibility and strengthening exercises			
	Source of Funding: Kaiser Foundation Research Institute, Northern California			
Participants	Diagnostic label used by trialists: Impingement			
	Criteria for defining the shoulder condition being treated			
	<ul> <li>Pain with: passive overpressure at full shoulder flexion with the scapula stabilised or passive internary rotation at 90 degrees shoulder flexion in the scapular plane and in progressive degrees of horizontary abduction; and one of</li> </ul>			
	<ul> <li>Pain on active shoulder abduction or pain on a resisted break test during abduction, internal rotatio or external rotation</li> </ul>			
	Inclusion Criteria (not listed above)			
	Between 18 and 65			
	• Be willing to remain on current level medication for study duration and for 2 weeks beforehand			
	Exclusion Criteria (not listed above)			
	Any other form of treatment for shoulder pain during study period			
	<ul> <li>Pending litigation on workman's compensation</li> </ul>			
	<ul> <li>History and exam suggestive of rotator cuff tear or adhesive capsulitis</li> </ul>			
	<ul> <li>History of shoulder subluxation, dislocation or fracture</li> </ul>			
	Cervical radiculitis or radiculopathy			
	History of cervical, shoulder or upper back surgery			
	History of systemic or neurological disease			
	Physical therapy or chiropractic treatment for the shoulder, neck or upper back in the past 12 months			
	<ul> <li>Insufficient English language skills to comprehend all explanations and respond to questions</li> </ul>			
	Baseline characteristics			
	Intervention			

Bang 2000 (Continued)	Number randomised: 28; mean age: 42 ± 10.1 years old; sex: F/M 10/18; duration of symptoms: 5.6 ± 3.7 months
	Control
	Number randomised: 24; mean age: 45 $\pm$ 8.4 years old; sex: F/M 12/12; duration of symptoms: 4.4 $\pm$ 2.8 months
Interventions	Intervention: manual physical therapy
	<i>Components of intervention</i> : techniques applied to movement limitations in the upper quarter that were relevant to the participant's problem. Mostly passive accessory or passive physiological joint mobilisation Maitland grades I-V were used. Typical initial treatment involved manual therapy techniques to enhance glenohumeral caudal glide in positions of flexion or abduction, and increase physiological flexion or internal rotation, but were adapted and progressed based on evaluation. Typical subsequent treatment involved manual therapy techniques to improve the combined physiological movements of hand behind back or shoulder quadrant, increase upper thoracic extension and side bend, or enhance extension, rotation, or side bend of the cervical spine. Techniques also included soft tissue massage and muscles stretching, and 1-2 extra home exercises designed to specifically support their mobilisation therapy (e.g. simple cervical and thoracic postural exercises such as chin tucks, and self-mobilisation such as caudal glides of the glenohumeral joint)
	Dose: 30 min in total
	Frequency of administration: twice weekly for 3 weeks
	Control: supervised flexibility and strengthening exercises
	<i>Components of intervention</i> : participants only received the treatment common to both groups (see be- low)
	Both Groups
	<i>Components of intervention</i> : standardised flexibility and strengthening exercise programme consist- ing of 2 passive stretching exercises (1 for the anterior musculature and 1 for the posterior capsule and musculature) and 6 strengthening exercises (flexion, scaption, rowing, horizontal extension-external rotation, seated press-up and elbow push-ups). Four of the 6 strengthening exercises were performed with Theratubing. Each stretch was held for 30 seconds, performed 3 times and with a 10-second rest between each exercise. Each Theratubing strengthening exercise was performed to a maximum of 10 repetitions for 3 sets with 60 seconds' rest between each exercise. The two other strengthening exercis- es were performed till fatigue or 25 reps
	Dose: 30 min in total
	<i>Frequency of administration</i> : manual therapy group = daily stretching exercises and 3-times weekly strengthening exercises; control group = twice weekly sessions for 3 weeks in-clinic and on other days, daily stretching exercises and 3-times weekly strengthening exercises at home
Outcomes	Outcomes assessed at 8 weeks (pain and function); isometric strength measured sometime between 4-8 weeks (not specified)
	<ul> <li>Pain: VAS from 0 (no pain) to 1000 mm (worst pain I can imagine)</li> <li>Function: modified Oswestry Low Back Disability Questionnaire scored from 0 to 45 with a higher score indicating better function</li> <li>Isometric strength measurements (for abduction, external rotation and internal rotation, with a composite measured), measured in Newtons using a dynamometer</li> </ul>
Notes	Conflicts of interest: not reported
Risk of bias	
Bias	Authors' judgement Support for judgement

Manual therapy and exercise for rotator cuff disease (Review)

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Bang 2000 (Continued)		
Random sequence genera- tion (selection bias)	Low risk	Quote: "On day 1, subjects signed the informed consent and were appointed to either the exercise group or the manual therapy group using the table of random numbers."
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Unclear risk	Comment: No information on how the allocation sequence was concealed was reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants, who may have had different expectations about the benefits of the intervention they received, self-reported pain and function
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "The testers were responsible for measurements of all dependent vari- ables and were blinded to group assignment for each subject."
		Comment: Outcome assessors of objective outcomes were likely blind to treat- ment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Two subjects did not complete the study. One subject from the manu- al therapy group was excluded from the study after the second visit due to in- juries sustained in a motor vehicle accident. The other subject, from the exer- cise group, elected to drop from the study after day 1 citing job related issues. All home exercise program compliance logs were returned and indicated that patients from both groups were fully compliant"
		Comment: The amount and reasons for attrition are unlikely to have affected the results
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: The study appears to be free of other bias

Bansal	2011
Dalisa	ZUII

Bansal 2011			
Methods	Design: Parallel group RCT Setting: University, India		
	Intervention: Deep friction massage plus Codman's exercises		
	Control: Therapeutic ultrasound plus Codman's exercises		
	Source of funding: Not reported		
Participants	Diagnostic label used by trialists: Supraspinatus tendinitis		
	Criteria for defining the shoulder condition being treated		
	Supraspinatus tendinitis defined by:		
	<ul> <li>point tenderness at greater tuberosity of humerus;</li> </ul>		

Manual therapy and exercise for rotator cuff disease (Review)



Bansal 2011 (Continued)

- positive empty can test;
- painful resisted abduction

# Inclusion Criteria (not listed above)

None

## Exclusion criteria (not listed above)

- History of trauma around shoulder
- Corticosteroid injections in the past
- Infective conditions
- Surgery around shoulder region
- Bony changes on radiological investigation

### **Baseline characteristics**

#### Intervention

Number randomised: 20; mean (SD) age = 30.90 (5.33) years; male:female = 12:8; duration of symptoms: not reported

## Control

Number randomised: 20; mean (SD) age = 30.35 (5.76) years; male:female = 9:11; duration of symptoms: not reported

Interventions	Intervention: deep friction massage		
	<i>Components of intervention:</i> deep friction massage to supraspinatus tendon in a transverse direction with the tip of the index finger, reinforced by middle finger. Participants were positioned half-lying with hand behind back (shoulder adduction and internal rotation)		
	Dose: 10-12 min for 10 sessions over 10 days		
	Frequency of administration: not explicitly reported, assumed daily for 10 days		
	Control: therapeutic ultrasound		
	<i>Components of intervention:</i> ultrasound applied to the supraspinatus tendon with the participants posi- tioned with hand behind back		
	<i>Dose:</i> intensity 0.6 w/cm <sup>2</sup> , frequency 1 MHz, pulse rate 4:1 for 6-8 min for 10 sessions over 10 days <i>Frequency of administration:</i> not explicitly reported, assumed daily for 10 days		
	Both groups		
	All participants were instructed in Codman's exercises consisting of pendulum or swinging motion of the arm in flexion, extension, horizontal abduction, adduction and circumduction. Dosage was not re- ported. Intensity (arc of motion) was increased as tolerated. Participants were also advised to avoid strenuous work involving the affected upper limb		
Outcomes	Outcomes assessed at 5 days and 10 days		
	<ul> <li>Pain using a visual analogue scale, ranging from 0 (no pain) to 10 (maximum pain)</li> <li>Active range of shoulder abduction measured using a goniometer with the participant in a seated position</li> </ul>		
Notes	Conflicts of interest: not reported		
Risk of bias			

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## Bansal 2011 (Continued)

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "The individuals were randomly divided into two groups"
		Comments: No information on how the allocation sequence was generated was reported
Allocation concealment (selection bias)	Unclear risk	Comment: No information on how the allocation sequence was concealed was reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants, who may have had different expectations about the benefits of the intervention they received, self-reported pain
Blinding of outcome as- sessment (detection bias) Objective outcomes	Unclear risk	Comment: No information was reported regarding the assessors of the objec- tive outcome, active range of shoulder abduction
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Comment: No dropouts, losses to follow-up or exclusions were reported, how- ever it is unclear whether the outcome data reported were based on the total number of randomly assigned participants
Selective reporting (re- porting bias)	Unclear risk	Comment: Only mean scores (no measures of variation) were reported for all outcomes. However, it is not clear whether data were incompletely reported based on the statistical significance or magnitude of the results. Also, without a trial protocol, it is unclear whether other outcomes were assessed but not re- ported based on the results
Other bias	Low risk	Comment: No other sources of bias were identified

## Barbosa 2008

Methods	Design: Parallel group RCT		
	Setting: Hospital (orthopaedics clinic), Brazil		
	Intervention: Mobilisation plus eccentric muscle training plus therapeutic ultrasound		
	Control: Eccentric muscle training plus therapeutic ultrasound		
	Source of Funding: Not reported		
Participants	Diagnostic label used by trialists: Supraspinatus tendinopathy		
	Criteria for defining the shoulder condition being treated		
	<ul> <li>Shoulder pain and/or dysfunction</li> <li>without a diagnosis of frozen shoulder</li> <li>demonstrated pain on palpation of the supraspinatus and/or biceps brachii muscle tendons</li> <li>positive in one or more special tests for detecting dysfunctions in the supraspinatus muscle tendon (like the Jobe test) and biceps brachii muscle tendon (like the Speed test and Yergason test)</li> </ul>		
	Inclusion Criteria (not listed above)		

Barbosa 2008 (Continued)	• Adults		
	Exclusion Criteria (not listed above)		
	<ul><li>Rupture of one or more of the rotator cuff tendons</li><li>Closed calcified tendinopathy diagnosed by imaging</li></ul>		
	Baseline characteristics:		
	Intervention		
	Number randomised: 7; mean age: 43.57 $\pm$ 7.49; sex: F/M 4/3; duration of symptoms: not reported		
	Control		
	Number randomised: 7; mean age: 48.71 ± 7.27; sex: F/M 5/2; duration of symptoms: not reported		
Interventions	Intervention: joint mobilisation		
	<i>Components of intervention</i> : front, back, lower longitudinal and lateral relaxations of the glenohumeral joint, anteroposterior movements of the acromioclavicular (squeeze) joint and anteroposterior, inferior or-superior and superior- inferior movements of the sternoclavicular joint.		
	<i>Dose</i> : 1 min of mobilisation for each movement (2-3 cycles per second), and 1 min of active free abduc- tion movement in the scapular plane, over the arc of movement without pain		
	Frequency of administration: 3 sessions per week for 4 weeks		
	Control: eccentric muscle training plus therapeutic ultrasound		
	<i>Components of intervention:</i> participants only received the treatment common to both groups (see be- low)		
	Both groups		
	Components of intervention		
	• Ultrasound: the Sonacel Dual therapeutic ultrasound equipment (Bioset) was applied by direct con- tact, using the contact medium of ultrasound transmission gel prepared within the Pharmacy Section of HCFMRP-USP, with continuous movement of the transducer		
	• Eccentric training exercises. the 'empty the can' movement (the participant performs abduction movements of the shoulder in the scapular plane, with medial rotation) when treating the supraspinatus muscle, or the "right curl" movement (the participant flexes his elbow, with the arm abducted beside the body) when treating biceps brachii dysfunctions. Movement resistance was offered manually, always by the same researcher and respecting the participant's pain limit		
	Dose		
	<ul> <li>Ultrasound: frequency of 3 MHz, with a SATA dosage of 1.0 W/cm<sup>2</sup> and a pulsed exit of 1:1 (50%). The US was applied for three min to the supraspinatus muscle tendon or for four min to the tendon of the long head of the biceps brachii muscle. The total emitted energy was 900 or 1600 J, resulting in emitted energy densities of 60 J/ cm<sup>2</sup></li> <li>Eccentric training exercises: 3 series of 20 repetitions</li> </ul>		
	Frequency of administration: 3 sessions per week for 4 weeks		
Outcomes	Outcomes assessed at 4 weeks		
	<ul> <li>Function: Constant score out of 100 with a higher score indicating less disability</li> <li>Bain on movement (scale not reported, and no outcome data reported)</li> </ul>		
	<ul> <li>Pain on movement (scale not reported, and no outcome data reported)</li> <li>ROM using goniometry (no outcome data reported)</li> </ul>		
Notes	Conflicts of interest: not reported		

Manual therapy and exercise for rotator cuff disease (Review)

# Barbosa 2008 (Continued)

**Risk of bias** 

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "They were randomly selected to participate in one of the treatment protocols (A or B)."
		Comment: No information on how the allocation sequence was generated was reported
Allocation concealment (selection bias)	Unclear risk	Comment: No information on how the allocation sequence was concealed was reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants, who may have had different expectations about the benefits of the intervention they received, self-reported pain and function
Blinding of outcome as- sessment (detection bias) Objective outcomes	Unclear risk	Comment: There was no information about whether assessors of objective outcomes were blind to treatment
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Comment: No dropouts, losses to follow-up or exclusions were reported, how- ever it is unclear whether the outcome data reported were based on the total number of randomly assigned participants
Selective reporting (re- porting bias)	High risk	Comment: No outcome data for pain on motion or ROM were reported (despite being listed as outcomes in the methods section of the review). Also, without a trial protocol, it is unclear whether other outcomes were assessed but not re- ported based on the results
Other bias	Low risk	Comment: The study appears to be free of other bias

# Barra 2011

Methods	Study design: Parallel group RCT	
	Setting: Primary health care centre in the Spanish National Health Service, Spain	
	Intervention: Diacutaneous fibrolysis (type of manual therapy)	
	Control: Placebo diacutaneous fibrolysis	
	<b>Source of Funding:</b> Jordi Gol Institute of Research in Primary Health Care funded translation of the manuscript into English. Funding of the trial not reported	
Participants	Diagnostic label used by trialists: Impingement	
	Criteria for defining the shoulder condition being treated	
	Referred to the physiotherapy unit with painful shoulder of periarticular origin	
	Inclusion Criteria (not listed above)	



Barra 2011 (Continued)

- Over 18 years
- Have had no previous diacutaneous fibrolysis

### **Exclusion Criteria (not listed above)**

- Adhesive capsulitis
- Damaged skin and/or cutaneous lesions in the shoulder area
- · Vascular abnormalities and a concomitant treatment with platelet anti-aggregant agents
- Previous shoulder surgery
- An acute (less than one week ago) inflammatory shoulder condition
- · People with a pending litigation or court claim

#### **Baseline characteristics**

#### Intervention

Number randomised: 25; mean age mean (SD):  $56.8 \pm 10.3$  years old; sex: F/M 16/9; duration of symptoms mean (SD):  $14.5 \pm 16.2$  months

#### Control

Number randomised: 25; mean age mean (SD):  $60.8 \pm 10.1$  years old; sex: F/M 13/12; duration of symptoms mean (SD):  $17.9 \pm 27.6$  months

Interventions

## Intervention: diacutaneous fibrolysis

*Components of intervention*: applied by means of a set of metallic hooks ending in a spatula with bevelled edges that allow a better distribution of the pressure on the skin. The aim is to release adherences between the different musculoskeletal structures, such as muscles, aponeurosis, tendons and others. Three consecutive steps are carried out. In the first one, or manual palpation step, the hand that is not holding the hook, the so-called palpatory hand, localises the intermuscular septum and remains in permanent control of the implementation of the technique. In the second step, or instrumental palpatory hand and both together (finger and hook) slide along the intermuscular septum, making short and brief movements perpendicular to the muscular fibres, allowing the detection of areas of movement restriction where the presence of adherences is presumed. The third step, the fibrolysis step, wherever a sensation of movement restriction is felt, a brief supplementary traction is carried out with the hook in order to tear the hypothetical adherent connective fibres. For this study, regardless of pain location, all the participants were treated with the same standardised protocol which involves the musculature of the scapula, the lateral region of the shoulder and arm, and the front part of the shoulder and chest. In the case of bilateral pain, this treatment was applied only to the most painful shoulder

Dose: 15 min

Frequency of administration: 1 session

#### Control: placebo diacutaneous fibrolysis

*Components of intervention*: the steps of manual and instrumental palpation occur as for the intervention group, but strictly at a superficial level. In the third step, instead of fibrolysis, a pinch of skin is held with the thumb of the palpatory hand and the tip of the spatula, so that the participant feels the hook distinctly but without any action taking place on the deep tissue levels

Dose: 15 min

Frequency of administration: 1 session

Outcomes Outcomes assessed immediately post-treatment (1 day)

• Pain: pain on internal rotation on a 100 mm VAS from 0 (no pain) to 100 (intolerable pain)

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Barra 2011 (Continued)

- Global assessment of treatment success: measured on a 5-point Likert scale consisting of "much better", "better", "no change", "worse" and "much worse". "Much better" or "better" were taken as a reported success
- Adverse events
- Active ROM (flexion, extension, abduction, external rotation and internal rotation) measured using a goniometer (except internal rotation which was measured in cm as hand behind back distance)

Notes

Conflicts of interest: the authors stated that they had no conflicts of interest

#### **Risk of bias**

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "The patients who consented to participate in this study were random- ly allocated to one of two groups: the intervention group or the placebo group. The randomization was stratified by centre. The DatInf RandList 1.2 software (DatInf GmbH, Tu¨bingen, Germany) was used."
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Unclear risk	Quote: "A physiotherapist was in charge of recruiting the participants, collect- ing the demographic data, measuring the initial variables and assigning a cor- relative number to each participant. Subsequently, the second physiothera- pist, who was the only person with access to the random allocation list gener- ated by the randomization software, implemented the technique according to the group assigned to the participant's number."
		Comment: No information on how the allocation sequence was concealed was reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Comment: When asked about their group allocation, 42/50 participants did not know and 5/8 of those who chose a group were correct. Four of these partici- pants were in the intervention group. The physiotherapist applying the inter- vention was not blinded, although this is unlikely to have affected outcomes given that no adjuvant treatment was provided
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	Low risk	Comment: Blinded participants self-reported pain and treatment success
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "After the participant had received the treatment, the first physiothera- pist, still blinded to the technique applied, took the final measurements of the variables"
		Comment: Assessor of objective outcomes was likely blind to treatment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: There was no loss to follow-up of participants
Selective reporting (re- porting bias)	Low risk	Comment: Outcome data fully reported for all outcomes specified in the Clini- calTrials.gov registry entry
Other bias	Low risk	Comment: The study appears to be free of other bias

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Barra Lopez 2013	
Methods	Study design: Parallel group RCT
	Setting: Primary healthcare centres, Spain
	<b>Intervention:</b> Diacutaneous fibrolysis plus standardised physiotherapy (exercise, electrotherapy and cryotherapy)
	Control 1: Sham diacutaneous fibrolysis plus standardised physiotherapy
	Control 2: Standardised physiotherapy alone
	Source of funding: Not reported
Participants	Diagnostic label used by trialists: Impingement
	Criteria for defining the shoulder condition being treated
	<ul> <li>Subacromial impingement syndrome diagnosed with the Neer impingement sign and Hawkins- Kennedy impingement test</li> </ul>
	Inclusion Criteria (not listed above)
	Aged 18 years and over
	Exclusion Criteria (not listed above)
	<ul> <li>Damaged skin</li> <li>Cutaneous lesions or vascular abnormalities in the shoulder area</li> <li>A concomitant treatment with platelet antiaggregant agents</li> <li>Acute inflammatory condition of the shoulder (&lt; 1 week)</li> <li>Previous shoulder surgery</li> <li>People with a pending litigation or court claim</li> </ul>
	Baseline characteristics
	Intervention
	Number randomised: 40; mean (SD) age = 56.2 (12) years; male:female = 15:25; mean (SD) duration of symptoms = 17.4 (24.9) months
	Control 1
	Number randomised: 40; mean (SD) age = 60 (10) years; male:female = 13:27; mean (SD) duration of symptoms = 24.2 (59.1) months
	Control 2
	Number randomised: 40; mean (SD) age = 59.1 (11.5); male:female = 17:23; mean (SD) duration of symp- toms = 14.7 (21.6)
Interventions	Intervention: diacutaneous fibrolysis
	<i>Components of intervention:</i> application of a metallic hook as deeply as possible following the inter- muscular septum between the muscles of the cervico-scapular (trapezius, rhomboideus major, rhom- boideus minor and levator scapulae) and shoulder region (infraspinatus, teres minor, teres major, tri- ceps brachii long head, deltoid, pectoralis major and biceps brachii long head tendon) in a centripetal direction towards the pain location
	Dose: not reported
	Frequency: 2 sessions per week for 3 weeks
	Control 1: sham diacutaneous fibrolysis

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<b>3arra Lopez 2013</b> (Continued)	rection as the interven ing place on the deep t	ntion: application of a metallic hook over the same muscles and in the same di- tion group, but only at a superficial level and without any mechanical action tak- issue layers. For the participant to feel the hook distinctly, a pinch of skin was f the palpatory hand and the tip of the hook
	Dose: not reported	
	Frequency: 2 sessions p	per week for 3 weeks
	Control 2: standardis	ed physiotherapy alone
	Components of interver	ntion: participants only received the treatment common to all groups (see below)
	All groups	
		<i>ntion:</i> all participants received a protocolised treatment of therapeutic exercises, py and cryotherapy. No other details provided
	Dose: not reported	
	Frequency: 5 sessions p	per week for 3 weeks
Outcomes	Outcomes assessed at	post-treatment (3 weeks) and 3 months
		ng a 100 mm visual analogue scale with "no pain" at the lower end and "intolerable
		Constant-Murley score (maximum possible score of 75 points, where a higher score
	<ul> <li>Active ROM (should versal two-armed get</li> </ul>	er flexion, extension, abduction, external rotation and internal rotation) using uni-
	Global assessment	of treatment success: participant's perception of the results using 5-point Likert "much worse" to "much better")
Notes	Conflicts of interest: r	not reported
	Trial registered in Clini	calTrials.gov (NCT01424579)
		o were suffering from bilateral subacromial impingement syndrome were provided treatment in both shoulders but, for the purpose of the study, only the most was evaluated
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "The allocation sequence was determined before the study using a computer generated randomisation list. It was stratified for each centre and treating physiotherapist (two in Cornellà, one in Ponteareas). As a result, the treating physiotherapist in Ponteareas has treated 10 patients per group and the two treating physiotherapists in Cornellà have each treated 15 patients per group."
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Unclear risk	Comment: No information on how the allocation sequence was concealed was reported
Blinding of participants and personnel (perfor-	High risk	Quote: "As Diacutaneous Fibrolysis is a manual technique, the therapist could not be blinded."

Quote: "When asked about the technique they thought they had received,

41% of the participants in the intervention group (sham n = 1, do not know n = 14) and 89% of the participants in the placebo group (real n = 21, do not know

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mance bias)

All outcomes



Barra Lopez 2013 (Continued)		n = 12) did not correctly determine the right response. Only four people (10%) in the placebo group correctly determined that they had received sham Diacu- taneous Fibrolysis treatment. As expected, all the participants in the control group answered "no additional technique" Comment: Participants in the diacutaneous fibrolysis (DF) and sham DF groups were blinded to whether they received DF or sham DF, but knew they did not receive the control condition. All participants in the control group were not blinded. Personnel delivering treatment were not blinded
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants, who may have had different expectations regarding the benefits of the intervention they received, self-rated pain and function
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "Assessment was carried out pre-treatment, post-treatment, and at a three-month follow-up by a different physiotherapist to the treating physio-therapist, and who was blinded to the group assignment."
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: Assessors of ROM were blind to group allocation Quote: "Analyses followed intention-to-treat principles. Missing observations due to lost to follow-up were completed with the last value observed from each subject" Quote: "Twenty-four participants were lost to follow-up. Seven participants (6%) dropped out during the treatment phase due to personal reasons unre- lated to the trial, and seventeen participants (14%) did not take the follow-up evaluation. It was not possible to contact thirteen of the participants for fol- low-up and four people were living in another city at that point in time." Comment: The number of participants lost to follow-up was similar across groups. The reasons for drop out for majority of those lost to follow-up is un- clear (they were unable to be contacted). Where reasons for drop-out were reported, it is unclear whether these were evenly distributed across groups. Analysis by ITT and missing data are unlikely to affect continuous outcomes
Selective reporting (re- porting bias)	Unclear risk	Comment: One outcome (participants' perception of results) is reported in the publication but was not listed on the registered trial information on the U.S. National Institutes of Health ClinicalTrials.gov website. However, it is unclear whether this outcome was introduced based on its results and the results of other outcomes
Other bias	Low risk	Comment: No other sources of bias identified

# Baskurt 2011

Saskult 2011				
Methods	Study design: Parallel group RCT			
	Setting: Orthopaedic physiotherapy unit, Turkey			
	Intervention: Scapular stabilisation exercises (PNF) plus stretching and strengthening exercises			
	Control: Stretching and strengthening exercises			
	Source of Funding: Not reported			
Participants	Diagnostic label used by trialists: Impingement			
	Criteria for defining the shoulder condition being treated			
	Pain on Neers, Hawkins or Jobe's test			
	<ul> <li>Diagnosis confirmed on radiography and ultrasound</li> </ul>			

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# Baskurt 2011 (Continued)

## Inclusion Criteria (not listed above)

• None

### **Exclusion Criteria (not listed above)**

- Inability to raise arm to 140 degrees of elevation
- Shoulder instability
- Adhesive capsulitis
- Cervical pathology
- Neurological deficit in the upper extremity
- Upper extremity surgery
- Musculoskeletal or cardiovascular pathologies limiting rehabilitation

### **Baseline characteristics**

## Intervention

Number randomised: 20; mean (SD) age: 51.50 (8.40) years old; male/female: not reported; mean (SD) duration of symptoms: 8.55 (10.78) months

Control

Number randomised: 20; mean (SD) age: 51.25 (11.55) years old; male/female: not reported; mean (SD) duration of symptoms: 11.60 (9.52) months

Interventions Intervention: scapular stabilisation exercises Components of intervention: scapular proprioceptive neuromuscular facilitation (PNF) exercises, scapular clock exercise, standing weight shift, double arm balancing, scapular depression, wall push up, wall slide exercises. Sessions were performed under physiotherapist supervision. When participants were able to do 3 sets of 10 repetitions without feeling substantial pain or fatigue, then the strongest elastic band was used Dose: 3 sets of each exercise Frequency of administration: 3 times per week for 6 weeks **Control: stretching and strengthening exercises** Components of intervention: participants only received the treatment common to both groups (see below) **Both groups** Components of intervention: flexibility exercises consisted of anterior, posterior and inferior capsule stretching, forward flexion ROM, abduction ROM and internal rotation stretching (with towel). Strengthening exercises consisted of subscapularis, infraspinatus, supraspinatus, and the anterior and posterior part of deltoid strengthening. Sessions were performed under physiotherapist supervision. When participants were able to do 3 sets of 10 repetitions without feeling substantial pain or fatigue, then the strongest elastic band was used. At the beginning of the study participants were educated on how to best use their injured shoulders (avoiding above the head work etc.) Dose: 3 sets of each exercise Frequency of administration: 3 times per week for 6 weeks Outcomes Outcomes assessed at 6 weeks • Pain: VAS scale 0-10 with a higher score indicating worse pain Pain on activity: VAS scale 0-10 with a higher score indicating worse pain ROM (flexion, abduction, internal rotation in 90 degrees abduction, external rotation in 90 degrees abduction) (unclear if active or passive)

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Baskurt 2011 (Continued)

• Strength (lower trapezium, middle trapezium, upper trapezium, serratus anterior, supraspinatus, subscapularis, infraspinatus, each measured in kg using a hand held dynamometer)

• Quality of life: WORC (0-2100) with a higher score indicating worse quality of life

Notes

Conflicts of interest: not reported

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "All patients meeting the criteria were separated into 2 groups accord- ing to simple random table."
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Unclear risk	Comment: No information on how the allocation sequence was concealed was reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Comment: Participants received different multimodal interventions, but it is unclear whether they were provided with any information that would make them perceive the intervention they received as superior or inferior to the al- ternative intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	Unclear risk	Comment: Participants self-reported pain, but it is unclear whether they were provided with any information that would make them perceive the interven- tion they received as superior or inferior to the alternative intervention
Blinding of outcome as- sessment (detection bias) Objective outcomes	Unclear risk	Comment: There was no information about whether assessors of objective outcomes were blind to treatment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: All randomised participants completed the study
Selective reporting (re- porting bias)	Unclear risk	Comment: Without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias were identified

Beaudreuil 2011	
Methods	Study design: Parallel group RCT
	Setting: Outpatient clinic, France
	Intervention: Dynamic Humeral Centering plus massage and home exercise
	Control: Non-specific mobilisation plus massage and exercise
	Source of Funding: Assistance Publique–Hôpitaux de Paris and the French Society of Rheumatology
Participants	Diagnostic label used by trialists: Impingement
	Criteria for defining the shoulder condition being treated
	At least 2 of Neer, Yocum and Hawkin's impingement tests

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# Beaudreuil 2011 (Continued)

## Inclusion Criteria (not listed above)

- Age over 30 years
- A Constant score of less than 80

## **Exclusion Criteria (not listed above)**

- Reduced passive ROM
- Interoposterior instability
- Tendinous calcification
- Corticosteroid injection within the previous 30 days
- Previous surgery
- Humeral fracture
- Inflammatory joint disease
- Neoplastic disorders

### **Baseline characteristics**

#### Intervention

Number randomised: 35; mean age:  $57.9 \pm 10.7$  years old; sex: F/M 21/14; mean duration of symptoms:  $35.7 \pm 81.6$  months

#### Control

Number randomised: 35; mean age: 59.4 ± 10.0 years old; sex: F/M 26/8; mean duration of symptoms: 20.9 ± 27.6 months

### Interventions

### Intervention: dynamic humeral centring

*Components of intervention*: the programme consisted of two parts:

- learning the lowering of the humeral head during passive abduction of the shoulder. It included muscular control of the scapula, perception of the passive lowering of the humeral head in the glenohumeral joint, active contraction of the pectoralis major and latissimus dorsi, perception of the lowering effect and co-contraction of these muscles during passive abduction of the shoulder
- actively lowering the humeral head by co-contraction of the pectoralis major and latissimus dorsi
  during active abduction of the shoulder. It was first performed with the elbow in a flexed position at
  90 degrees, from 0 degrees to 90 degrees of shoulder abduction. The active movement with the cocontracted pectoralis major and latissimus dorsi was then repeated with the elbow in an extended
  position, covering the entire range of shoulder abduction without and then with the participant holding a 0.5 kg weight

Home exercises consisted of 10 co-contractions of the pectoralis major and latissimus dorsi three times a day, the shoulder actively positioned in abduction and the elbow in the flexed or extended position according to the stage of progression

### Dose: see above

*Frequency of administration*: dynamic humeral centring - 15 sessions over 6 weeks (3 times a week for the first 3 weeks, and twice a week for the next 3 weeks); home exercises - daily for 6 weeks

#### **Control: non-specific mobilisation**

Components of intervention: the programme consisted of three stages

- Passive mobilisation of the shoulder with a painless ROM. Home exercises were at this stage 10 pendular movements of the shoulder 3 times a day
- Active mobilisation of the shoulder with a painless ROM. The home exercises were then 10 active anterior elevations of the shoulder in the lateral rotated position 3 times a day
- Active mobilisation of the shoulder performed with slight manual resistance applied by the physiotherapist along with the second part of the home exercises

	Active ROM: Constant sub-score from 0-40 points with 0 indicating the highest impairment
	<ul> <li>Function: Total Constant score from 0–100 with 0 indicating the highest impairment</li> <li>Pain: Constant sub-score for pain from 0–15 points with 0 indicating the highest impairment</li> <li>Strength: Constant sub-score from 0-25 points with 0 indicating the highest impairment</li> </ul>
Outcomes	Outcomes assessed at 3 months and 12 months
	<i>Frequency of administration</i> : 15 sessions over 6 weeks (3 times a week for the first 3 weeks, and twice a week for the next 3 weeks)
	Dose: 10 min for massage
	<i>Components of intervention</i> : each session began with massage of the neck and shoulder region with the participant lying on one side or sitting. Physiotherapists were allowed to adjust the intensity of the treatment according to the participant's capabilities. Participants also performed exercises at home depending on the intervention group
	Both groups
	<i>Frequency of administration</i> : manual therapy - 15 sessions over six weeks (3 times a week for the first 3 weeks, and twice a week for the next 3 weeks); home exercises - daily for 6 weeks
Beaudreuil 2011 (Continued)	Dose: see above

# Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "Patients were assigned in permuted blocks of six".
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Low risk	Quote: "Allocations were sealed in opaque and consecutively numbered en- velopes. Envelopes were opened by an independent investigator who was not involved in the eligibility assessment, outcome assessment or treatment. Allo- cation was revealed to the physiotherapist before the patients presented for treatment."
		Comment: An adequate method was used to conceal the allocation sequence
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Patients were informed that two treatment procedures were being evaluated, with no further information on the superiority of one treatment over the other. Patients were therefore blinded to the study hypothesis"
		Comment: Participants, but not personnel, were blind to treatment (though the latter is unlikely to have affected outcomes given that no adjuvant treat- ment was provided)
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	Low risk	Comment: Blinded participants self-reported pain and function
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "An assessor blinded to treatment assessed all outcomes."
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "A total of 34 patients underwent DHC and 35 control treatment. At 3 months, 90% of included patients were available for assessment, and at 12 months 70%."

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# Beaudreuil 2011 (Continued)

Other bias	Low risk	Comment: No other sources of bias were identified
Selective reporting (re- porting bias)	Low risk	Comment: Outcome data were fully reported for all outcomes specified in the ClinicalTrials.gov registry entry (NCT 01022775)
		Comment: The rate of dropout was relatively similar between groups, though no reasons for dropout were provided. However, analysis was by intention-to-treat, where missing data were imputed using the multiple imputation method

# Bennell 2010

Methods	Study design: Parallel group RCT			
	Setting: Public hospital physiotherapy units and private physiotherapy practices, Australia			
	<b>Intervention:</b> Physiotherapy (soft tissue massage, joint mobilisation, scapular retraining, postural taking, ing, supervised and home exercises)			
	Control: Placebo physiotherapy (sham ultrasound only)			
	Source of Funding: National Health and Medical Research Council			
Participants	Diagnostic label used by trialists: Chronic rotator cuff disease (encompassing several conditions)			
	Criteria for defining the shoulder condition being treated			
	<ul> <li>Severity of pain on movement greater than 3/10 on a 0-10 numerical rating scale</li> <li>Pain on active abduction or external rotation</li> <li>Positive quick test for shoulder impingement</li> </ul>			
	Inclusion Criteria (not listed above)			
	Aged 18 years or older			
	Exclusion Criteria (not listed above)			
	<ul> <li>Pain on rest greater than 7/10</li> <li>A suspected full rotator cuff tear</li> <li>Previous shoulder surgery</li> <li>Radiological evidence of osteoarthritis, calcification or previous fracture</li> <li>Systemic pathology including inflammatory joint disease or neoplastic disorders</li> <li>Greater than 50% restriction of ROM of passive movement in two or more shoulder planes</li> <li>Referred pain from the spine diagnosed by spinal clearing tests</li> <li>Symptoms of complex regional pain syndrome</li> <li>Active intervention in the previous three months</li> <li>Anti-inflammatory drugs in the previous two weeks</li> <li>Inability to understand written or spoken English</li> </ul>			
	Baseline characteristics			
	Intervention			
	Number randomised: 59; mean age: 59.3 $\pm$ 10.1 years old; sex: F/M 25/34; mean duration of symptoms 24 months			

Control

Bennell 2010 (Continued)	Number randomised: 61; mean age: 60.8 ± 12.4 years old; sex: F/M 31/30; mean duration of symptoms: 14 months
Interventions	Intervention: physiotherapy
	<i>Components of intervention</i> : directed at improving dynamic scapular control, strengthening scapular stabiliser and rotator cuff muscles, improving shoulder and thoracic posture, and increasing ROM of thoracic extension. The treatment was administered in a standardised way by a trained physiotherapist. Analgesia as required was permitted, and behaviour modification strategies such as education, goal setting, motivation and positive reinforcement were provided
	<ul> <li>Soft tissue massage: anterior and posterior shoulder tissues were massaged in supine and side-lyin positions respectively</li> </ul>
	<ul> <li>Glenohumeral joint mobilisation: anteroposterior and inferior joint glides in supine position with th shoulder at 45 degrees and 90 degrees respectively</li> </ul>
	<ul> <li>Thoracic spine mobilisation (T1-8): in prone position, using central posteroanterior technique</li> <li>Cervical spine mobilisation (C5-7): in prone position using central posteroanterior unilateral technique on both sides</li> </ul>
	<ul> <li>Scapular retraining: in side-lying position, therapist passively moves shoulder through range from elevation/protraction to retraction/depression, then assisted by participant, then independently by participant; isometric holds in retraction/depression</li> </ul>
	<ul> <li>Postural taping: taping of shoulders and scapula to encourage scapular retraction and depression an thoracic extension</li> </ul>
	<ul> <li>Home exercises: most exercises required the participant to incorporate their scapular retraining wit strengthening of the rotator cuff muscles. Some exercises reinforced and facilitated correct posture Resistance for specific exercises was provided by hand weights or elastic theraband. Exercises wer taught and performed during each treatment session and were otherwise self-administered at hom</li> </ul>
	Dose: total 30 - 45 min each session
	<ul> <li>Soft tissue massage: 6 min each position</li> <li>Glenohumeral joint mobilisation: 4 x 30 seconds each position</li> <li>Thoracic spine mobilisation (T1-8): Grade IV on each level for total 4 min</li> <li>Cervical spine mobilisation (C5-7): Grade IV on each level for total 4 min</li> <li>Scapular retraining: Weeks 1 and 2 only, 15 repetitions by 5 repetitions with 10-second holds</li> <li>Postural taping: continuously worn for 2 weeks and reapplied after 1 week</li> <li>Home exercises: twice daily in the first 2 weeks, once a day thereafter maintained for 12 weeks after the conclusion of the 10-week programme</li> </ul>
	<i>Frequency of administration</i> : in-clinic sessions were performed twice weekly for the first 2 weeks, once weekly for the next 4 weeks and then once fortnightly for the last 4 weeks, totaling 10 sessions over 10 weeks. After the 10-week programme, participants were instructed to maintain their daily home exercise programme for 12 weeks
	Control: sham ultrasound
	<i>Components of intervention</i> : sham ultrasound and light application of a non-therapeutic gel to the shoulder region
	Dose: 10 min for the ultrasound and 10 for the gel
	<i>Frequency of administration</i> : twice weekly for the first 2 weeks, once weekly for the next 4 weeks and then once fortnightly for the last 4 weeks, totaling 10 sessions over 10 weeks. After the 10-week pro- gramme, participants did not receive any intervention and were not instructed to do any home exercis es
	Co-interventions
	Use of analgesics and non-steroidal anti-inflammatory drugs was similar in the active and placebo

Use of analgesics and non-steroidal anti-inflammatory drugs was similar in the active and placebo groups over both the intervention period (analgesics: 11/55 (20%) active v 14/61 (23%) placebo; non-



## Bennell 2010 (Continued)

	steroidal anti-inflammatories: 12/55 (22%) v 13/61 (21%)) and the follow-up period (analgesics: 8/49 (16%) v 8/55 (15%); non-steroidal anti-inflammatories 6/49 (12%) v 8/55 (15%))
Outcomes	Outcomes assessed at 11 weeks and 22 weeks post-randomisation
	<ul> <li>Pain at rest: VAS (0-10) with a higher score indicating greater pain</li> <li>Drive an accuracy of the Control of the contro</li></ul>
	Pain on movement: VAS (0-10)     Date: SDADLasin sub-second state of the state
	• Pain: SPADI pain sub-score scaled from 0–100 with a higher score indicating more pain
	<ul> <li>Function: SPADI function subscore (0-100) with a higher score indicating more dysfunction</li> </ul>
	<ul> <li>Function: SPADI total score (0-100) with a higher score indicating more pain/dysfunction</li> </ul>
	<ul> <li>Quality of life: SF-36 physical component score and mental component score (each 0 to 100) with a higher score indicating better health</li> </ul>
	• Quality of life: AQoL (-0.04 to 1) with a higher score indicating better quality of life
	• Strength: isometric abduction, external rotation and internal rotation strength measured in kg with a Nicholas Manual Muscle tester
	<ul> <li>Proportion with successful treatment: % (defined by 5-point Likert scale, where 5 indicates "much better")</li> </ul>
	<ul> <li>Global rating of change overall: measured on a 5-point Likert scale from 1 ("much worse") to 5 ("much better"), with a score of 5 indicating a successful outcome</li> </ul>
	Adverse events: recorded in a log book
Notes	Conflicts of interest: authors declared that they have no financial or non-financial interests that may

be relevant to the submitted work

## Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "Participants had a baseline assessment and were randomised in per- muted blocks of six and eight, stratified by treating physiotherapist, to receive either active manual therapy and home exercise treatment or placebo treat- ment according to a computer generated table of random numbers created by the study biostatistician".
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Low risk	Quote: "Allocations were sealed in opaque and consecutively numbered envelopes kept in a central locked location. An independent administrator opened the envelopes in sequence and then revealed the group allocation to the relevant physiotherapist by facsimile just before the participant presented for treatment."
		Comment: An adequate method was used to conceal the allocation sequence
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Comment: While the personnel were not blinded to treatment allocation, when asked which treatment they were receiving participants' answers showed a moderate to high degree of blinding with a blinding index of 0.7 (1 indicates complete blinding, 0 indicates no blinding, and 0.5 would be expect- ed if participants were randomly guessing)
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	Low risk	Comment: Blinded participants self-reported pain and function
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "The same blinded assessor (EW) evaluated all participants at baseline, at 11 weeks (at the conclusion of the supervised active or placebo intervention), and at 22 weeks after randomisation."

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# Bennell 2010 (Continued)

Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: Attrition rates were low and similar across both groups, and data were analysed according to an intention-to-treat analysis, where missing data were replaced by the last score carried forward
Selective reporting (re- porting bias)	Low risk	Comment: Outcome data were fully reported for all outcomes specified in the study protocol
Other bias	Low risk	Comment: No other sources of bias were identified

# Bialoszewski 2011

Methods	Study design: Parallel group RCT
	Setting: University, Poland
	Intervention: Manual therapy plus TENS plus ultrasound plus exercise
	Control: TENS plus ultrasound plus exercise
	Source of Funding: Not reported
Participants	Diagnostic label used by trialists: Chronic rotator cuff injury
	Criteria for defining the shoulder condition being treated
	Confirmed diagnosis of rotator cuff injury without indications for surgical treatment
	Inclusion Criteria (not listed above)
	<ul> <li>No co-existing medical conditions</li> <li>No concomitant anti-inflammatory or analgesic medications</li> </ul>
	Exclusion Criteria (not listed above)
	• None
	Baseline characteristics
	Intervention
	Number randomised: 15; mean (range) age: 52.6 (38–61) years old; sex: F/M 5/10; mean duration of symptoms: 4.8 months
	Control
	Number randomised: 15; mean (range) age: 50 (38–60) years old; sex: F/M 7/8; mean duration of symp- toms: 4.4 months
Interventions	Intervention: manual Therapy
	<i>Components of intervention</i> : mobilisation of the glenohumeral joint and soft tissues using Kaltenborn's roll-glide techniques, Cyriax deep friction massage, Mulligan's mobilisation with movement and typical techniques of joint mobilisation in the anteroposterior direction
	Dose: not reported
	Frequency of administration: not reported
	Control: TENS plus ultrasound plus exercise



### Bialoszewski 2011 (Continued)

*Components of intervention:* participants only received the treatment common to both groups (see below)

## **Both groups**

Components of intervention

Standard rehabilitation involving TENS, therapeutic ultrasound and kinesiotherapy

- TENS: Triangular pulsed current was used. 6 x 6 cm rubber electrodes were placed at supraspinatus insertion region (cathode) and on the scapula (anode)
- Therapeutic ultrasound: 4 cm<sup>2</sup> ultrasound probe placed over the supraspinatus insertion region with an ultrasound gel served as a coupling substance
- Kinesiotherapy: standard passive and active exercises used to improve the ROM and restore muscle strength. The rotator cuff was initially strengthened in the painless ROM by performing active, passive and self-assisted exercises. Once the full ROM had been achieved, strengthening exercises were applied, ranging from flexion, abduction and external rotation to internal rotation adduction and extension

#### Dose

- TENS: frequency 100 Hz and width 1.0 ms, current set in accordance with participant's sensations for 20 min
- Therapeutic ultrasound: frequency of 1 MHz, maximum power of 10 W. Pulsed waves were used with a duty cycle of 20% and a frequency of 48 Hz. Power density of 0.5 W/cm<sup>2</sup> was used in the first session and increased by 0.1 W/cm<sup>2</sup> each session. The sessions lasted 4 to 9 min
- Kinesiotherapy: not reported

Frequency of administration: not reported

Outcomes Outcomes assessed at 4 time points, but it is not clear when these occurred (in terms of weeks) nor how long the intervention lasted

- Pain: VAS from 0–10 with a higher score indicating worse pain
- Active and passive ROM (flexion, abduction, internal rotation, external rotation) using a goniometer

## Notes Conflicts of interest: not reported

### **Risk of bias**

Bias	Authors' judgement	Support for judgement
Random sequence genera-	Unclear risk	Quote: "Patients were randomly assigned to two matched groups"
tion (selection bias)		Comment: No information about how the allocation sequence was generated was reported
Allocation concealment (selection bias)	Unclear risk	Comment: No information about how the allocation sequence was concealed was reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants, who may have had different expectations about the benefits of the intervention they received, self-reported pain

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# Bialoszewski 2011 (Continued)

Blinding of outcome as- sessment (detection bias) Objective outcomes	Unclear risk	Comment: There was no information on whether assessors of objective out- comes were blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: All randomised participants were analysed
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias were identified

Methods	<b>Study design:</b> Parallel group RCT <b>Setting:</b> Outpatient rehabilitation department, USA <b>Intervention 1:</b> Supervised eccentric progressive resistance exercises plus ice plus home exercises			
				Intervention 2: Supervised concentric progressive resistance exercises plus ice plus home exercises
	Source of Funding: Not reported			
Participants	Diagnostic label used by trialists: Subacromial impingement syndrome			
	Criteria for defining the shoulder condition being treated			
	<ul> <li>At least 1 positive impingement test (Neer, Hawkins-Kennedy, or coracoid or 'cross body' adductio impingement test) and one negative full-thickness rotator cuff tear test (infraspinatus test, drop arr test, or "empty can" test)</li> </ul>			
	Inclusion Criteria (not listed above)			
	<ul><li>Aged 18 years or older</li><li>Had not yet initiated physical or occupational therapy treatment</li></ul>			
	Exclusion Criteria (not listed above)			
	<ul> <li>History of shoulder, cervical, or thoracic surgery</li> <li>History of shoulder fracture, dislocation, labral tear, or full-thickness rotator cuff (RTC) tear</li> <li>Treatment for neoplasm in the last year</li> <li>Rheumatic disease</li> <li>Adhesive capsulitis</li> </ul>			
	<ul> <li>Shoulder pain currently rated greater than or equal to 7/10 on the NPRS</li> <li>Cardiac, neurological, or musculoskeletal condition that precludes ability to perform upper extremi resisted exercise</li> </ul>			
	<ul> <li>Pregnancy</li> <li>Inability to arrange transport to evaluation or treatment sessions or not planning to stay in the are long enough to complete study</li> </ul>			
	Baseline characteristics			
	<i>Intervention 1</i> Number randomised: 17 (17 completed); mean age: 48.8 ± 16.5 years old; sex: F/M 9/8; duration of symptoms: 28.2 ± 23.6 months			



Blume 2014 (Continued)			
	Intervention 2 Number randomised: 1 symptoms: 23 ± 27.8 mo	7 (13 completed); mean age: 47.2 $\pm$ 14.7 years old; sex: F/M 7/6; duration of onths	
Interventions	Intervention 1: eccent	ric progressive resistance exercises	
	and scapular muscles. E external rotation (ER) w duction, and prone sho and all were performed the lowering portion of	<i>tion:</i> supervised eccentric progressive resistance exercises for the rotator cuff Exercises included the seated 'full can', sidelying internal rotation (IR), sidelying with towel roll, supine protraction, sidelying horizontal abduction, sidelying ab- ulder extension. All exercises were performed using a dumbbell for resistance in the participant's pain-free AROM. The eccentric exercise group performed the exercises with the therapist repositioning the weight to the starting position the lifting portion of each exercise	
	Dose: 3 sets of 12 repeti	tions of each exercise; duration of session was 1 hour	
	Frequency of administro	ation: twice a week for 8 weeks	
	Intervention 2: concer	ntric progressive resistance exercises	
	and scapular muscles.	<i>tion:</i> supervised concentric progressive resistance exercises for the rotator cuff Same exercises as above, except this group performed the lifting portion of the apist repositioning the weight to the start position to avoid resistance in the low- rcise	
	Dose: 3 sets of 12 repeti	tions of each exercise; duration of session was 1 hour	
	Frequency of administration: twice a week for 8 weeks		
	Both groups - Ice and home exercises		
	<i>Components of intervention:</i> participants received ice treatment on the shoulder for 15 min at the end of each supervised clinic session. A home exercise programme (HEP) of stretching and postural correction exercises (pectoralis minor and posterior shoulder self-stretching and thoracic spine self-mobilisation into extension along with pain-free AROM in flexion and abduction standing in front of a mirror to monitor for excessive scapular elevation)		
	Dose: 3 sets of 12 repetitions of each exercise		
	Frequency of administrc weeks	<i>ation:</i> once daily on the days the participant was not exercising in the clinic, for 8	
Outcomes	Outcomes assessed at 5 weeks and 8 weeks		
	<ul><li>disability and a score</li><li>Active ROM: scaption</li></ul>	s of the Arm, Shoulder and Hand (DASH) 0-100, where a score of zero indicates no e of 100 indicates total disability n using an inclinometer	
	Isometric strength for	or shoulder external rotation and abduction using a hand held dynamometer	
Notes	Conflicts of interest: n	ot reported	
	Report only available as a PhD thesis		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Unclear risk	Comment: There was no information on how the allocation sequence was generated	

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Blume 2014 (Continued)		
Allocation concealment (selection bias)	Low risk	Quote: "The participants were randomly assigned to one of the two inter- vention groups using pre-prepared, sealed folders selected in numeric order which became their participant identification number"
		Comment: An adequate method was used to conceal the allocation sequence
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Comment: Participants were likely blinded, as they were unlikely to have no- ticed the difference between the two exercise programmes
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	Low risk	Comment: Blinded participants self-reported some outcomes
Blinding of outcome as- sessment (detection bias)	Low risk	Quote: "Outcomes were assessed by an investigator blinded to the treatment intervention group assignment"
Objective outcomes		Comment: Assessor of objective outcomes was blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Following group allocation, there were 17 participants in each group. One enrolled participant in the concentric group dropped out at three weeks due to an unrelated medical issue. Sixteen participants in the concentric group and seventeen participants in the eccentric group completed the fifth week assessments. Three participants in the concentric group withdrew after five weeks, one due to travelling out of state and two for financial and work con- flict reasons. As a result, a total of thirty participants completed the 8-week study and the final assessments, 13 in the concentric group and 17 in the ec- centric group".
		Comment: The attrition was unrelated to the interventions, and the amount was small, so is unlikely to have biased the results
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the thesis, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias identified

## Brox 1993

DI 0X 1999				
Methods	Study design: Parallel group RCT			
	<b>Setting:</b> Hospital orthopaedics department and hospital physical medicine and rehabilitation depart- ment, Norway			
	Intervention: Supervised exercise			
	Control 1: Arthroscopic subacromial decompression			
	Control 2: Placebo laser treatment			
	Source of Funding: Norwegian Research Council			
Participants	Diagnostic label used by trialists: Impingement			
	Criteria for defining the shoulder condition being treated			
	Pain in the shoulder			

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Brox 1993 (Continued)

- Dysfunction or pain on abduction
- Normal passive glenohumeral range of movement
- Pain during two of three isometric tests (abduction at 0 degrees, at 30 degrees and external rotation
- Positive results for impingement tests
- Reduced pain 15 min after injection of 6 ml 10 mg/ml lignocaine into the subacromial space

## Inclusion Criteria (not listed above)

- Aged 18 66
- pain had been resistant to outpatient physiotherapy and NSAIDs and steroids

## **Exclusion Criteria (not listed above)**

- · Arthritis of the acromioclavicular joint
- Cervical syndrome
- Rotator cuff rupture
- Glenohumeral instability
- Bilateral muscular pain with tenderness and severely decreased ability to relax the shoulder, neck or temporomandibular joint on examination
- Were reluctant to accept one or more treatment regimes in the study

### **Baseline characteristics**

#### Intervention

Number randomised: 50; mean age: 47 years old; sex: F/M 28/22; duration of symptoms: < 6 months (n = 6); 6 months to 1 year (n = 6); 1–3 years (n = 13); > 3 years (n = 25)

## Control 1

Number randomised: 45; mean age: 48 years old; sex: F/M 16/32; duration of symptoms: < 6 months (n = 8); 6 months to 1 year (n = 8); 1–3 years (n = 9); > 3 years (n = 20)

### Control 2

Number randomised: 30; mean age: 48 years old; sex: F/M 15/15; duration of symptoms: < 6 months (n = 5); 6 months to 1 year (n = 5); 1–3 years (n = 5); > 3 years (n = 14)

#### Interventions

### Intervention: supervised exercise

*Components of intervention*: supervised exercises to normalise dysfunctional neuromuscular patterns and to "increase the nutrition of the collagen in the rotator cuff". To eliminate gravitational forces and to start the exercises the arm was suspended in a sling fixed to the roof. Relaxed repetitive movements (first rotation, then flexion-extension, and finally abduction-adduction) were performed for about an hour in a daily training session. Resistance was gradually added to strengthen the short shoulder rotator and scapular stabilising muscles. Participants also received 3 lessons on the anatomy and function of the shoulder, pain management and ergonometrics

## Dose: 1 hour

*Frequency of administration*: participants were supervised twice weekly for 6 weeks. On the other days they followed the same exercise programme at home. The training continued for 3 to 6 months, with the supervision gradually being reduced

## Control 1: arthroscopic subacromial decompression

*Components of intervention*: the aim of the procedure was to make more space for the rotator cuff to reduce the risk of impingement. Standard treatment consisted of bursectomy and resection of the anterior and the lateral part of the acromion and the coracoacromial ligament. Participants received post-operative rehabilitation on day one and physiotherapy within the first week. Exercises prescribed by the surgeon were performed against low resistance and repeated many times. Unrestricted activities were advised after 4 to 6 weeks

Brox 1993 (Continued)			
	Dose: NA		
	Frequency of administration: 1 surgical procedure		
	Control 2: placebo laser		
	Components of intervention: a detuned soft laser delivered by the hospital physiotherapist		
	Dose: not reported		
	Frequency of administration: twice weekly for 6 weeks		
Outcomes	Outcomes assessed at 3 months and 6 months		
	<ul> <li>Function: Neer clinical testing of function (muscle strength, reaching ability and stability score) rang- ing from 0–30 points with a higher score indicating better function</li> </ul>		
	<ul> <li>Pain: Neer verbal rating pain score of pain the previous week, from 0–35 with a higher score indicating less pain</li> </ul>		
	<ul> <li>Active ROM: Neer ROM score from 0–25 with a higher score indicating better ROM</li> </ul>		
	<ul> <li>Global assessment of treatment success: number of participants with a good or an excellent Neer shoulder score (&gt; 80 points)</li> </ul>		
	Adverse events		
	<ul> <li>Pain on activity (during the previous week), on 9-point scale (1 = no pain, 9 = worst pain)</li> </ul>		
	<ul> <li>Pain at rest (during the previous week), on 9-point scale (1 = no pain, 9 = worst pain)</li> </ul>		
	<ul> <li>Night pain (during the previous week), on 9-point scale (1 = no pain, 9 = worst pain)</li> </ul>		
	Work disability: number of days sick leave		
Notes	Conflicts of interest: not reported		

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "Treatments were allocated by the method of random permuted blocks"
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Unclear risk	Comment: No information about how the allocation sequence was concealed was reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention. The comparison of supervised exercise versus arthroscopic sub- acromial decompression is of concern, while the comparison of supervised ex- ercise versus placebo is not
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants, who may have had different expectations about the benefits of the intervention they received, self-reported outcomes
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "Blind follow up measurements were carried out at three and six months after the first day of treatment. At follow up tests the patients wore a T-shirt to hide a possible scar from surgery. They were carefully told not to talk about their treatment."
		Comment: Outcome assessors of objective outcomes were probably blind to treatment

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Incomplete outcome data (attrition bias) All outcomes	High risk	Quote: "Unexpectedly, many patients dropped out from surgery. Four of them did not attend follow up examination at six months, which might have biased our results". The comparison of supervised exercise versus arthroscopic sub- acromial decompression is of concern; the comparison of supervised exercise versus placebo is not
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias were identified

Methods	Study design: Parallel group RCT				
	Setting: Marmara University Faculty of Medicine, Physical Medicine and Rehabilitation, Turkey				
	<b>Intervention:</b> Exercise below 90 degrees plus TENS plus pulsed therapeutic ultrasound plus cold pack plus NSAID				
	<b>Control:</b> Exercise above 90 degrees plus TENS plus pulsed therapeutic ultrasound plus cold pack plus NSAID				
	Source of funding: Not reported				
Participants	Diagnostic label used by trialists: Impingement				
	Criteria for defining the shoulder condition being treated				
	<ul> <li>Positive Neer impingement test, Hawkins signs, Jobe supraspinatus test</li> <li>30% less limitation in passive movement compared to that of the other side</li> <li>No degenerative arthritis or mesoacromion on imaging</li> <li>No pathologic symptoms except the edema in subacromial bursa on MRI</li> </ul>				
	Inclusion Criteria (not listed above)				
	Participants must not be playing any sports				
	Exclusion Criteria (not listed above)				
	<ul> <li>Receiving shoulder surgery</li> <li>Receiving other physical therapy or rehabilitation</li> <li>Receiving psychiatric treatment</li> </ul>				
	Baseline characteristics				
	Overall cohort of participants				
	Number randomised: 33; mean (range) age: 52 (34–70) years old; sex: F/M 23/7; duration of symptoms: not reported				
Interventions	Intervention: exercise below 90 degrees				
	<i>Components of intervention</i> : supervised shoulder flexion below 90 degrees, abduction, T-bar (wand) exercises containing internal-external rotation and extension, posterior capsule stretching and internal rotation exercises and rotator cuff strengthening exercises				
	Dose: 30 repetitions in the hospital and 30 repetitions twice more at home				

Manual therapy and exercise for rotator cuff disease (Review)

Celik 2009 (Continued)	Frequency of administr	ration: daily for 2 weeks		
	<i>Frequency of administration</i> : daily for 2 weeks <b>Control: exercise above 90 degrees</b>			
		-		
	<i>Components of intervention</i> : supervised exercises over 90 degrees, posterior and inferior capsule stretching exercises, rotator cuff strengthening and internal rotation exercises			
	Dose: 30 repetitions in	the hospital and 30 repetitions twice more at home		
	Frequency of administr	ration: daily for 2 weeks		
	Both groups			
	<i>Components of intervention</i> : TENS, pulsed therapeutic ultrasound, ice application and oral tenoxicam <i>Dose</i> :			
	<ul> <li>Ice: 15 min, applied at home after the exercises were performed</li> <li>TENS: 20 min</li> <li>Ultrasound: 1 w/cm<sup>2</sup> pulsed ultrasound for 4 min</li> <li>Oral tenoxicam: 20 mg once a day</li> <li>Frequency of administration: daily for 2 weeks</li> </ul>			
Outcomes	Outcomes assessed at 2 weeks and 16 weeks			
	Pain: VAS from 0 (no pain) to 10 (extremely severe pain)			
	<ul> <li>Global assessment of treatment success (participant satisfaction scale 0-4, higher score denotes higher satisfaction)</li> </ul>			
Notes	Conflicts of interest: not reported			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "Patients who agreed to participate were divided into two groups ran- domly"		
		Comment: No information about how the allocation sequence was generated was reported		
Allocation concealment (selection bias)	Unclear risk	Comment: No information about how the allocation sequence was concealed was reported		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Comment: Participants received slightly different types of exercises, but it is unclear whether they were provided with any information that would make them perceive the intervention they received as superior or inferior to the al- ternative intervention		
Blinding of outcome as- sessment (detection bias)	Unclear risk	Comment: Participants self-reported some outcomes but it is unclear whether they were provided with any information that would make them perceive the		

intervention they received as superior or inferior to the alternative interven-

Comment: The only objective outcome (physical therapist satisfaction) was

Quote: "First group had 17 patients, while the second group consisted of 16

patients. In the first group, one patient had difficulty in transportation to the

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Self-reported outcomes

Blinding of outcome as-

**Objective outcomes** 

(attrition bias)

sessment (detection bias)

Incomplete outcome data

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High risk

Low risk

tion

measured by an unblinded assessor



Celik 2009 (Continued)		
All outcomes		hospital, and another patient had indication of surgery while attending exer- cise program. For these reasons, two patients were left out of the study. In the second group, one patient left the exercise program due to pain. Patients in 2 groups of 15 people (23 females, 7 males; Age 52; range 34-70) completed the treatment."
		Comment: The amount of attrition was low and unlikely to have impacted on the results
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other source of bias was identified

Methods	Study design: Parallel group RCT			
	Setting: University rehabilitation department, Turkey			
	<b>Intervention:</b> Manual mobilisation plus hot pack plus theraband exercises plus home exercises (Cod- man)			
	<b>Control:</b> Proprioceptive neuromuscular facilitation (PNF) plus hot pack plus theraband exercises plus home exercises (Codman)			
	Source of funding: Not reported			
Participants	Diagnostic label used by trialists: Impingement			
	Criteria for defining the shoulder condition being treated			
	Clinically and radiographically confirmed shoulder impingement syndrome			
	Inclusion Criteria (not listed above)			
	• None			
	Exclusion Criteria (not listed above)			
	<ul> <li>No previous surgery, physical treatments or rehabilitation programmes for their condition before the study</li> </ul>			
	No previous local steroid injections			
	Baseline characteristics			
	Intervention			
	Number randomised: 20; mean (SD) age: $52.80 \pm 9.86$ years; sex: not reported; duration of symptoms: not reported			
	Control			
	Number randomised: 20; mean (SD) age: 55.50 ± 8.95 years; sex: not reported; duration of symptoms: not reported			
Interventions	Intervention: manual mobilisation			
	Components of intervention: no details provided			

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# Citaker 2005 (Continued)

#### Dose: not reported

*Frequency of administration*: 20 sessions (number of sessions per week and total number of sessions not reported)

### Control: proprioceptive neuromuscular facilitation

Components of intervention: no details provided

Dose: not reported

*Frequency of administration*: 20 sessions (number of sessions per week and total number of sessions not reported)

#### **Both groups**

*Components of intervention*: hot pack plus theraband exercises plus home exercises (Codman). The hot packs which were used for the treatment were Nonius<sup>™</sup>, with dimensions 30–50 cm. They were applied for 20 min. The elastic therabands used for the treatments (Theraband<sup>™</sup>) were 7.6 cm wide and 152 cm long. They were fixed to an object such as a doorknob. Six colour-coded bands were available; each provided increasing resistance from 0.5 to 2.7 kg with increments of 0.5 kg. Theraband exercises permit concentric and eccentric strengthening of the shoulder muscles. The exercises begun with the elbow flexed 90 degrees and the shoulder in the neutral position. The exercises were performed through an arc of 45 degrees in each of the 5 planes of motion. In addition, Codman pendulum exercises were utilised as a home programme in both groups

Dose: hot pack applied for 20 min

*Frequency of administration*: 20 clinic sessions (number of sessions per week and total number of sessions not reported) followed by 3 weeks of theraband exercises

Outcomes

Outcomes assessed at 3 weeks

- Function: University of California at Los Angeles (UCLA) shoulder rating scale total score (0-35, where higher scores denote better function)
- Night activity pain (0-10 VAS)
- Night rest pain (0-10 VAS)
- Day activity pain (0-10 VAS)
- Day rest pain (0-10 VAS)
- ROM (flexion, abduction, external rotation, internal rotation, hyperextension) using a goniometer (unclear if active or passive)

Notes

Conflicts of interest: not reported

#### **Risk of bias**

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "The patients were divided into two equal groups randomly." Comment: No information on how the allocation sequence was generated was reported
Allocation concealment (selection bias)	Unclear risk	Comment: No information on how the allocation sequence was concealed was reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Comment: Participants received slightly different multi-modal interventions, but it is unclear whether they were provided with any information that would make them perceive the intervention they received as superior or inferior to the alternative intervention

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## Citaker 2005 (Continued)

Blinding of outcome as- sessment (detection bias) Self-reported outcomes	Unclear risk	Comment: Participants self-reported some outcomes, but it is unclear whether they were provided with any information that would make them perceive the intervention they received as superior or inferior to the alternative interven- tion
Blinding of outcome as- sessment (detection bias) Objective outcomes	Unclear risk	Comment: There was no information about whether assessors of ROM were blinded
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Comment: No drop-outs, losses to follow-up or exclusions were reported, but it is unclear whether outcome data were based on an analysis of all 40 randomised participants, and it is unclear how many participants were ran- domised to each group
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data fully reported for all outcomes specified in the meth- ods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other source of bias identified

## **Clews 1987**

Methods	Study design: Parallel group RCT			
	Setting: Australian Institute of Sport, Australia			
	Intervention: Massage plus ice			
	Control 1: Therapeutic ultrasound plus ice			
	Control 2: Placebo ultrasound plus ice			
	Source of funding: Not reported			
Participants	Diagnostic label used by trialists: Rotator cuff tendinitis			
	Criteria for defining the shoulder condition being treated			
	<ul> <li>Unilateral shoulder pain</li> <li>Localised dull pain in the antero/lateral shoulder region with no radiation of symptoms</li> <li>Tenderness to palpation at least on the long head of biceps in the bicipital groove, the insertion of the supraspinatus tendon or the musculotendinous portion of the long head of biceps</li> <li>Pain on resisted shoulder abduction, flexion or resisted supination of the forearm</li> <li>A positive impingement sign</li> <li>Absence of cervical sign symptoms or signs pointing the problem being referred from the neck, including negative Elvey's test</li> <li>No treatment other than ice having been instituted for the injury</li> </ul>			
	Inclusion Criteria (not listed above)			
	• None			
	Exclusion Criteria (not listed above)			
	• None			
	Baseline characteristics			
	Overall cohort of participants			



## Clews 1987 (Continued)

Number randomised: 18 (6 per group); age (mean and SD, or range): not reported; sex: not reported; duration of symptoms: not reported

Interventions	Intervention: massage									
	Components of intervention: massage of several muscles delivered as follows									
	<ul> <li>Long head of biceps: with the participant lying supine with their arm held at the elbow and held away from the table, long strokes with firm pressure delivered by the thumb were applied to the long head of the biceps. This was then followed by deep transverse frictions through the musculotendinous junctions of the biceps</li> <li>Biceps tendon consisted of a series of very firm long strokes and short transverse strokes using the thumb and working slowly towards the tendon's origin. The arm was resting at the participants side or held away from the table</li> <li>Pectorals: the tips of the middle three fingers were held tightly together. The other hand was used to reinforce and apply pressure, and each stroke was applied from the sternum to the shoulder</li> <li>Supraspinatus: participant placed in prone position with the affected arm resting at their side on the table. Warm-up strokes were then applied to the muscle. Standing at the head of the participant, the thumbs were pressed firmly into the muscle and then rocked backwards and forwards while changing position every few seconds. Using the thumb and massaging from the acromioclavicular joint towards the superior angle of the scapula, long and deep strokes were also applied</li> </ul>									
						<ul> <li>Infraspinatus: with the participant's arm hanging over the side of the table, each stroke began at the vertebral border of the scapula and moved through to the muscle's insertion. One hand was used to reinforce the other, and pressure was applied through the points of the middle three fingers. The therapist then moved to the other side of the table and worked the muscle longitudinally by applying deep pressure with the thumb. This was followed by short, transverse strokes along the inferior border of the spine of the scapula</li> </ul>				
						Dose: 15 min				
	Frequency of administration: every day for 3 days									
	Control 1: therapeutic ultrasound									
	Components of intervention: pulsed ultrasound									
	Dose: 15 min at an intensity 0.8 w/cm <sup>2</sup>									
	Frequency of administration: every day for 3 days									
	Control 2: sham ultrasound									
	Components of intervention: sham ultrasound									
	Dose: 15 min									
	Frequency of administration: every day for 3 days									
	All groups									
	Components of intervention: ice packs applied to the affected shoulder, and NSAIDs									
	Dose: ice for 15 min twice daily and 1 tablet of Voltaren taken with meals									
	Frequency of administration: every day for 3 days									
Outcomes	Outcomes assessed at 3 days									
	<ul> <li>Pain: VAS scale on strength testing from 0-10 with a higher score indicating worse pain</li> <li>Strength (maximal isometric force production, measured in peak force)</li> </ul>									
Notes	Conflicts of interest: not reported									

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## Clews 1987 (Continued)

**Risk of bias** 

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "After the diagnosis had been made and inclusion in the study was con- firmed, each subject was randomly assigned to one of these three groups."
		Comment: No information about how the allocation sequence was generated was reported
Allocation concealment (selection bias)	Unclear risk	Comment: No information about how the allocation sequence was concealed was reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported some out-comes
Blinding of outcome as- sessment (detection bias)	Low risk	Quote: "One co-author did all the testing and was not aware of the subjects' group assignment"
Objective outcomes		Comment: Outcome assessor of objective outcomes was blind to treatment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: There was complete follow-up of all randomised participants in the study
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias were identified

### **Cloke 2008**

Methods	Study design: Parallel group RCT		
	Setting: Primary care, United Kingdom		
Intervention: Exercise and manual therapy package			
	Control 1: Injections of corticosteroid and local anaesthetic		
	<b>Control 2:</b> Combination of exercise and manual therapy package plus injections of corticosteroid and local anaesthetic		
	Control 3: Controlled medical treatment with regular NSAIDs		
	Source of funding: Chartered Society of Physiotherapy Charitable Trust		
Participants <b>Diagnostic label used by trialists:</b> Painful arc (synonymous with the diagnosis of subacromia pingement, subacromial bursitis, subdeltoid bursitis, rotator cuff tendinitis, supraspinatus ter and rotator cuff tendinopathy)			



Cloke 2008 (Continued)

Trusted evidence. Informed decisions. Better health.

Criteria for defining the shoulder condition being treated · Painful arc (pain in the subacromial region between 60 degrees and 120 degrees of active shoulder abduction against gravity) positive impingement test (Neer or Hawkin's) Inclusion Criteria (not listed above) Aged over 18 **Exclusion Criteria (not listed above)** · Pain originating from the neck and radiating to the shoulder • Systemic inflammatory arthritis Severe loss of ROM exceeding 50% lateral rotation or 30 degrees of elevation compared with the unaffected contralateral side consistent with primary frozen shoulder or severe secondary capsulitis Glenohumeral or acromioclavicular joint osteoarthritis as a primary pathology or presentationA clinically incompetent rotator cuff, that is, marked weakness of rotator cuff muscles (pseudoparalysis) Shoulder injection by a recognised technique • • Physiotherapy using a EMTP Chiropractic or osteopathic treatment within the previous three months Known sensitivity/allergic reaction to local anaesthetic agents or steroid carrier compounds **Baseline characteristics** Overall cohort of participants Number randomised: 112; mean (range) age: 54.5 (23 - 88); sex: F/M 64/48; duration of symptoms: not reported Interventions Intervention: exercise and manual therapy package (EMTP) Components of intervention: an exercise and manual therapy regime based on a literature review (described in Kibler W. Shoulder rehabilitation: principles and practice. Med Sci Sports Exerc 1998;30:S40-50) and the opinions of an expert group of physiotherapists. No other details provided Dose: not reported Frequency of administration: 6 sessions over 18 weeks Control 1: glucocorticoid and anaesthetic injection Components of intervention: a course of injections of methylprednisolone and lidocaine The injections were placed 1 cm inferior to the posterior corner of the acromion, directed upward toward the subacromial region Dose: 40 mg of methylprednisolone and 10 ml of 1% lidocaine Frequency of administration: 1 injection every 6 weeks for a maximum of 3 injections Control 2: combination of EMTP and glucocorticoid and anaesthetic injection See details of each above **Control 3: NSAIDs or analgesia** Components of intervention: regular NSAIDs or simple analgesia if unable to tolerate NSAIDs Dose: not reported

Frequency of administration: not reported

Outcomes Outcomes assessed at 18 weeks and one year

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Cloke 2008 (Continued)

- Function: Oxford shoulder score (12-60) with a higher score indicating worse disability
- Global assessment of treatment success: participant's perception of outcome (same, better, or worse) (no outcome data reported)
- Requiring surgery at 1 year (no usable outcome data reported as the number of participants in each group at 1 year was not reported)

Notes

## Conflicts of interest: not reported

Standard deviations for the Oxford Shoulder Score were not reported in numerical format so were estimated from Figures

## **Risk of bias**

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "Those who consented to enter the trial were randomized by closed envelope"
		Comment: No information about how the allocation sequence was generated was reported
Allocation concealment (selection bias)	Unclear risk	Quote: "Those who consented to enter the trial were randomized by closed envelope"
		Comment: It is not clear whether envelopes were sequentially numbered or opaque or how the sequence was generated, so it is unclear if the allocation sequence was successfully concealed
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported some out-comes
Incomplete outcome data (attrition bias) All outcomes	High risk	Quote: "After randomization, the control group had 27 patients, EMTP had 29, the injection group had 28, and both interventions had 29. Ninety patients (80%) completed the trial: control group, 20; EMTP, 22; injections, 26, both interventions, 22. Of the 22 patients who did not complete the trial, 1 was lost to follow-up, and 21 withdrew during the duration of the trial (control, 7; EMTP, 7; injection, 2; and both, 5), and (18.75% noncompletion rate). Those who withdrew were invited for follow-up in the standard outpatient clinic. Sixty-two patients returned the follow-up questionnaire at 1 year (55% of those randomized). By 1 year, 2 patients in the injection group and 1 each in the control and EMTP groups had gone on to surgery."
		Quote: "Analysis was on an intention-to-treat basis"
		Comment: Reasons for dropout were not reported and numbers of dropout were unbalanced between groups. The authors state that an intention-to-treat analysis was performed, though report outcome data based on the per-proto- col sample
Selective reporting (re- porting bias)	High risk	Comment: No measures of variation were reported for any outcomes (except for the Oxford Shoulder Score, where SDs and 95% CIs were reported in Fig- ure format). However, it is not clear whether data were incompletely report- ed based on the statistical significance or magnitude of the results. No usable outcome data were reported for global assessment of treatment success or

Manual therapy and exercise for rotator cuff disease (Review)



## Cloke 2008 (Continued)

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	requiring of surgery. Also, without a trial protocol, it is unclear whether othe outcomes were measured but not reported based on the results	:1
Other bias Low	Comment: No other sources of bias were identified	

Methods	Study design: Parallel group RCT				
	Setting: Primary care, USA				
	Intervention: Mobilisation plus comprehensive treatment (hot packs, AROM, stretching, strengthen- ing, soft tissue mobilisation and participant education)				
	Control: Comprehensive treatment alone				
	Source of funding: Not reported				
Participants	Diagnostic label used by trialists: Impingement				
	Criteria for defining the shoulder condition being treated				
	<ul> <li>Pain about the superolateral shoulder region and one or more of</li> <li>* active ROM deficits in humeral elevation</li> </ul>				
	* painful subacromial compression				
	<ul> <li>limited functional movement patterns in an elevated position</li> </ul>				
	Inclusion Criteria (not listed above)				
	Negative upper quadrant clearing exam to rule out cervical, elbow, wrist and hand involvement				
	Exclusion Criteria (not listed above)				
	Shoulder instability				
	primary scapulothoracic dysfunction				
	stage II or III adhesive capsulitis				
	<ul> <li>third degree musculo-tendinous tears</li> </ul>				
	<ul> <li>advanced acromioclavicular joint disease</li> </ul>				
	<ul> <li>advanced calcific tendonitis or bursitis</li> </ul>				
	<ul> <li>severe degenerative bony or ligamentous changes</li> </ul>				
	neurological involvement				
	unstable fracture of the humerus, scapula or clavicle				
	Baseline characteristics				
	Intervention				
	Number randomised: 7; age mean (SD): 55.0 $\pm$ 10.2 years old; sex: not reported; duration of symptoms: not reported				
	Control				
	Number randomised: 7; age mean (SD): 50.7 ± 16.5 years old; sex: not reported; duration of symptoms: not reported				
Interventions	Intervention: joint mobilisation				
	<i>Components of intervention</i> : the Foley method (Foley R, Janos S, Johnson R, Petersen C: Active and Passive Movement Testing of the Extremities, Spine, Pelvis, and Temporomandibular Joint, Petersen				

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Conroy 1998 (Continued)

C (ed), Chicago, IL: Northwestern University Medical School Programs in Physical Therapy, 1994) was used to deliver the mobilisation. Depending on the direction of restriction in capsular extensibility, inferior glide, posterior glide, anterior glide or long axis traction could be applied to the participant with oscillatory pressure. Stretch could also be applied in the case of muscle spasm

*Dose*: each indicated technique was administered 2-4 times (30 seconds each). As a result, the intervention group received a maximum of 15 min additional treatment compared with the control group

Frequency of administration: 3 times per week for 3 weeks

### **Control: comprehensive treatment alone**

*Components of intervention*: participants only received the treatment common to both groups (see below)

### **Both groups**

*Components of intervention*: hot packs, active ROM, physiologic stretching and muscle strengthening exercises, soft tissue mobilisation and participant education

- · Hot packs: placed on the posterior, anterior and superior aspects of the shoulder
- · Active ROM exercises: pain-free pendulum exercises and postural correction
- Physiologic stretching: cane-assisted flexion and external rotation, towel-assisted internal rotation and non-involved arm-assisted horizontal abduction
- Muscle strengthening exercises: chair press, internal and external rotation isometrics
- Advice: avoid increased pain with all exercises and daily activities and advised to position the upper extremity in a supported 40 50 degree scapular plane elevation position (loose packed position) when not using the extremity
- Soft tissue mobilisation: effleurage, friction, and kneading techniques, with the subject sitting with the arm supported in a relatively loose packed position. The friction technique was specifically applied to the supraspinatus, bicipital long head and the subscapularis tendons. Pressure was applied in a cephalic medial way towards the body

#### Dose

- Hot packs: 15 min
- Exercises: 45-60 min
- Soft tissue mobilisation: 10 min, with each technique performed for 1 min and repeated 3 times (the last min was effleurage)

Frequency of administration: 3 times per week for 3 weeks

Outcomes	Outcomes assessed at 3 weeks	
	<ul> <li>Function: overhead function measured using 3 tests: -         <ul> <li>reach behind head and touch the external occipital protuberance with the long finger, with the back and arm against the wall;</li> <li>reach across and around the upper body to the lowest cervical or thoracic spinous process that</li> </ul> </li> </ul>	
	they could reach with the long finger; and	
	<ul> <li>* using the long finger, touch a mark on the wall that required 135 degrees of shoulder flexion – each rated as number who answered "can do", "can do in spite of pain" or "cannot do"</li> </ul>	
	• Pain: worst pain in the last 24 hours on a VAS scale of 0 (no pain) to 100 mm (worst pain imaginable)	
	<ul> <li>Active ROM: flexion, abduction, elevation, external rotation and internal rotation, measured using a goniometer</li> </ul>	
Notes	Conflicts of interest: not reported	
Risk of bias		
Bias	Authors' judgement Support for judgement	
BIBS	Autnors' judgement Support for judgement	

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Conroy 1998 (Continued)		
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "Subjects were randomly assigned to either the experimental (mobili- sation) or the control (no mobilisation) group".
		Comment: No information about how the allocation sequence was generated was reported
Allocation concealment (selection bias)	Unclear risk	Comment: No information about how the allocation sequence was concealed was reported
Blinding of participants	Low risk	Quote: "Both the subject and examiner were blinded to group assignment."
and personnel (perfor- mance bias) All outcomes		Comment: Participants received different multimodal interventions that were unlikely to be distinguishable to participants. Thus, participants were likely blinded to the intervention they received
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	Low risk	Comment: Blinded participants self-reported pain
Blinding of outcome as-	Low risk	Quote: "Both the subject and examiner were blinded to group assignment."
sessment (detection bias) Objective outcomes		Comment: Assessors of objective outcomes were likely blinded to the intervention
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "One subject dropped from the control group due to lack of under- standing instructions".
		Comment: The amount of dropout was small and unlikely to have affected the results
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias were identified

## Cook 2014

Methods	Study design: Parallel group RCT			
	Setting: Outpatient clinical/academic centres (USA or South Africa)			
	<b>Intervention:</b> Neck manual therapy plus standard physiotherapy (shoulder manual therapy to the shoulder, self- and externally-applied stretching, isotonic strengthening, and restoration of normative movement)			
	Control: Standard physiotherapy			
	Source of funding: Not reported			
Participants	Diagnostic label used by trialists: Impingement			
	Criteria for defining the shoulder condition being treated			
	Report of pain or dysfunction with overhead activities			
	Demonstration of pain during active shoulder movements			
	Demonstration of a positive Neer/Hawkins-Kennedy test			
	Recent onset within last 12 months			

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Cook 2014 (Continued)

- Report of non-traumatic onset
- Demonstration of a painful arc of the arm from 60-120 degrees of flexion
- Baseline pain level more than or equal to 2/10 on an 11-point numeric scale

## Inclusion Criteria (not listed above)

• None

## **Exclusion Criteria (not listed above)**

- History of frozen shoulder
- Disorders of the acromioclavicular joint
- Degenerative arthritis of the glenohumeral joint
- Calcifying tendonitis
- Shoulder instability
- Post-traumatic disorders
- Shoulder surgery and/or elbow, hand and wrist
- Blatantly misdiagnosed cervical spine disorders

## **Baseline characteristics**

### Intervention

Number randomised: 38; mean (SD) age:  $51.1 \pm 12.9$  years old; sex: F/M 15/23; duration of symptoms:  $12.9 \pm 17.6$  weeks

#### Control

Number randomised: 36; mean (SD) age: 51.0 ± 15.5 years old; sex: F/M 22/14; duration of symptoms: 10.4 ± 10.6 weeks

#### Interventions

### Intervention: neck manual therapy

*Components of intervention*: the manual therapy interventions to the neck consisted of grade III posterior-anterior mobilisations, performed in prone for 30 repetitions for 3 sets. Since any comparable shoulder symptoms during mobilisation to the cervical spine was an exclusion criterion, and since none of the subjects exhibited active neck symptoms, the posterior-anterior mobilisation was performed to the stiffest or the participant's most painful segment. When no joint signs were present, the posterior-anterior was performed to either C5-C6, or C6-C7 at the same side of the neck as the shoulder impingement. Where both pain and stiffness were present at multiple levels the clinician was able to identify the targeted level for mobilisation

Dose: 30 repetitions for three sets

*Frequency of administration*: participant discharge, treatment length, and frequency of treatment were determined by the physiotherapists, although some participants terminated treatment themselves. Participants had a mean (SD) of 59.7 (70.2) days in care

### **Control: standard physiotherapy**

*Components of intervention*: participants only received the treatment common to both groups (see below)

## **Both groups**

*Components of intervention*: Kuhn's (2009) approach, which advocates the use of a modified treatment that is unique to each individual and is based on their hypothesised underlying dysfunctions/causes, was used. The treatment methods included manual therapy, self- and externally-applied stretching, isotonic strengthening, and restoration of normative movement. The clinical and home-treatment programmes were modified for all subjects in each phase regardless of presentation, and the dosage of the interventions was specific to the examination findings

Dose: not reported

Cook 2014 (Continued)	<i>Frequency of administration</i> : participant discharge, treatment length, and frequency of treatment were determined by the physiotherapists, although some participants terminated treatment themselves. Participants had a mean (SD) of 52 (29.6) days in care		
Outcomes	Outcomes assessed at 2 days and at discharge		
	• Function measured with QuickDASH: 11 questions associated with various activities of daily living (range from 1-5)		
	<ul> <li>Pain: measured on numerical pain rating scale for pain (NPRS, where 0 = no pain and 10 = worst possible pain)</li> </ul>		
	Adverse events		
	<ul> <li>Global assessment ported)</li> </ul>	of treatment success: self-reported rate of recovery (RoR) (no outcome data re-	
Notes	Conflicts of interest: not reported		
	Trial is registered in ClinicalTrials.gov (NCT01744002)		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Quote: "patients were randomised by roll of die into the shoulder treatment plus neck mobilisations or the shoulder treatment only groups".	

()		
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Unclear risk	Comment: No information on how the allocation sequence was concealed was reported
Blinding of participants	High risk	Quote: "The study was a randomised, single blinded, controlled trial."
and personnel (perfor- mance bias) All outcomes		Quote: "Physiotherapists were blinded to the collected self-report outcomes in the study."
		Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as-	High risk	Quote: "The study was a randomised, single blinded, controlled trial."
sessment (detection bias) Self-reported outcomes		Quote: "Physiotherapists were blinded to the collected self-report outcomes in the study."
		Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported all outcomes to this review (e.g. pain, function, patient-acceptable symptom state, self-re- port rate of recovery)
Incomplete outcome data (attrition bias)	Low risk	Quote: "Seventy-four (74) subjects were enrolled and of these, six did not re- turn for a required follow-up visit."
All outcomes		Comment: A CONSORT flow chart (Schultz 2010) shows that two participants were lost to follow-up from the shoulder plus neck treatment group and four were lost to follow-up from the shoulder treatment group. The reasons for all six losses are "did not return after initial visit". It is unclear what the reasons for not returning were, but regardless of the reasons, the small amount of attrition and relatively equal distribution of attrition is unlikely to bias the outcomes

comes

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Cook 2014 (Continued)		
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for three outcomes specified in the ClinicalTrials.gov registry entry (NCT01744002). However self-reported rate of recovery was listed as an outcome in the methods section (but not in the registry), and no outcome data for it were reported in the Results section. However it is unclear if this outcome was not reported based on the results
Other bias	Low risk	Comment: No other sources of bias identified

# Dickens 2005

Methods	Study design: Parallel group RCT				
	Setting: Orthopaedic department in a district general hospital, UK				
	Intervention: Physiotherapy (joint mobilisation, exercise therapy and/or electrotherapy)				
	Control: Maintain normal daily activities				
	<b>Source of funding:</b> Supported in part by the Physiotherapy Research Foundation, Project Reference No. PRF/99/2				
Participants	Diagnostic label used by trialists: Impingement				
	Criteria for defining the shoulder condition being treated				
	<ul> <li>Subacromial impingement on history, examination and radiographic findings</li> <li>Diagnostic local anaesthetic injection into the subacromial space and acromioclavicular joint</li> <li>Showed no improvement or had persisting pain, loss of function or a positive impingement test after three steroid injections into the subacromial space six weeks apart.</li> </ul>				
	Inclusion Criteria (not listed above)				
	• None				
	Exclusion Criteria (not listed above)				
	<ul> <li>Cervical radiculopathy</li> <li>Adhesive capsulitis</li> <li>Clinically obvious rotator cuff tear</li> <li>Had previously received a course of physiotherapy</li> <li>Grade III subacromial spur on their shoulder supraspinatus outlet radiograph</li> </ul>				
	Baseline characteristics				
	Intervention				
	Number randomised: 45; age: mean (range) 55 (27–68); sex: F/M 19/26; duration of symptoms: not re- ported				
	Control				
	Number randomised: 40; age: mean (range) 54 (26–73); sex: F/M 18/22; duration of symptoms: not re- ported				
Interventions	Intervention: physiotherapy				
	<i>Components of intervention</i> : personalised training programme, both in-hospital and at home. All partic- ipants received some or all of: acromioclavicular joint, thoracic, cervical spine and glenohumeral joint mobilisation, exercise therapy including attention to muscle imbalance, postural advice, strapping and, very occasionally, electrotherapy. The exercises were carried out once or twice per week in hospi-				

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## Dickens 2005 (Continued)

tal until participants felt comfortable performing their therapy programme without supervision. The aim of the physiotherapy was to reduce inflammation and pain, directly using electrotherapy modalities and indirectly by altering the movement patterns (strapping and exercises) to alter the stresses placed on the subacromial structures, to a level where the exercise programme could be undertaken. The need for joint mobilisations was decided upon at the physiotherapy assessment. The physiotherapist assessed the range of accessory movement available in each participant's glenohumeral (anteroposterior, longitudinal caudad), acromioclavicular (anteroposterior, longitudinal caudad), cervical (posterior-anterior) and thoracic spine joints (posterior-anterior and transverse) with passive accessory movements. Any joints that were found to have restricted movement were addressed with mobilisations into the direction of resistance and pain to help restore full pain-free range of movement. All participants were given exercises for the recruitment and strength of scapulothoracic muscles (especially lower trapezeius and serratus anterior). The exercise programme was progressed to involve strengthening of infraspinatus, subscapularis and teres minor relative to the supraspinatus and deltoid. The rotator cuff exercises were done with the use of resistance and participants were given Theraband for home use. The exercises started in neutral positions with isometric contractions and were progressed to inner range, through range, outer range and into functional positions. The resistance and speed of these exercises were altered and progressed

Dose: dependent on participant

*Frequency of administration*: twice a day for exercises. Overall duration of physical therapy programme not reported

### **Control: normal daily activities**

Components of intervention: advised to maintain their normal activities of daily living whilst waiting for surgery

Dose: NA

Frequency: NA

Outcomes

Outcomes assessed at 6 months

• Function: Constant score from 0-100 with a higher score indicating better function

Notes Conflicts of interest: not reported

#### **Risk of bias**

Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "Patients were randomised into two groups using unmarked envelopes in clinic to achieve simple randomisation. There were 100 envelopes, 50 of which contained the word 'control' and 50 of which contained the word 'phys- iotherapy'."	
		Comment: The random sequence may have been generated by shuffling of envelopes (given that the envelopes were unmarked), but this is not clear	
Allocation concealment (selection bias)	Unclear risk	Comment: No information on how the allocation sequence was concealed was reported. Envelopes were unmarked rather than sequentially numbered, which suggests that they were shuffled, though this is not clear	
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention	

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Dickens 2005 (Continued)		
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants, who may have had different expectations about the benefits of the intervention they received, self-reported some out-comes
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "All patients in both groups were re-examined and Constant scores were performed at 6 months by JLW. JLW was blinded to the group allocation of the patients."
		Comment: Assessor of objective outcomes was likely blind to treatment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Three patients in the physiotherapy group were unable to complete the physiotherapy programme for social reasons and were therefore excluded, leaving 42 patients".
		Quote: "In the control group, nine patients initially randomised not to under- go a physiotherapy programme refused to attend follow-up for repeat assess- ment. These patients felt that surgery was inevitable and that further assess- ment was not indicated. Although we have not been able to analyse the Con- stant score results on this group of nine patients, they all underwent surgery. These patients were included in the statistical analysis on an intention-to-treat basis."
		Comment: Participants who did not complete follow-up were analysed using a worst-case scenario intention-to-treat method where possible (i.e., it was assumed they showed no improvement). This is an appropriate imputation method given the reasons for missing data
Selective reporting (re- porting bias)	Unclear risk	Comment: Only mean and range for the Constant score in each group was re- ported (i.e. no SDs, SEs or 95% CIs). However, it is not clear whether data were incompletely reported based on the statistical significance or magnitude of the results. Also, without a trial protocol, it is unclear whether other outcomes were assessed but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias were identified

## Djordjevic 2012

Methods	Study design: Parallel group RCT	Study design: Parallel group RCT			
	Setting: Rehabilitation clinic, Serbia				
	Intervention: Mobilisation with movement and kinesiotaping				
	Control: Supervised exercise programme				
	Source of funding: No funding	Source of funding: No funding			
Participants	Diagnostic label used by trialists: Impingement				
	Criteria for defining the shoulder condition being treated				
	Rotator cuff lesion and/or shoulder impingement syndrome				
	Inclusion Criteria (not listed above)				
	• None				
	Exclusion Criteria (not listed above)				
	Shoulder girdle fractures and dislocation				
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- Shoulder surgery in the last 12 months
- Physician diagnosis of adhesive capsulitis, full thickness rotator cuff tear, cervicobrachial pain due to cervical spine pathology, neuromuscular disorders in upper extremities
- Use of corticosteroid and/or nonsteroidal anti-inflammatory therapy within 10 days before the first day of measuring ROM

#### **Baseline characteristics**

### Intervention

Number randomised: 10; mean (SD) age:  $51.8 \pm 5.3$  years; sex: F/M 6/4; mean (SD) duration of symptoms:  $4.7 \pm 0.6$  months

#### Control

Number randomised: 10; mean (SD) age: 54.1  $\pm$  6.8 years; sex: F/M 7/3; mean (SD) duration of symptoms: 4.8  $\pm$  0.9 months

#### Interventions

Intervention: mobilisation with movement (MWM) and kinesiotape

Components of intervention

- MWM: during the MWM treatment, the participant was seated, and the therapist was positioned on the
  opposite side of participant's painful shoulder. The therapist applied the thenar of one hand on the
  anterior aspect of the participant's humeral head and the other hand on his/her scapula. The hand
  on the humeral head performed a posterolateral glide, while the other hand stabilised the scapula.
  During this manoeuvre, the participant was encouraged to perform active shoulder movement to the
  point of the first onset of pain
- Kinesiotape: standard 5 cm black Kinesio Tex tape applied to the supraspinatus muscle, deltoid muscle, and glenohumeral joint. The first strip of tape was torn down just above the anchor point where the Y strip was formed. The anchor point of the strip was taped to the projection of insertion of the supraspinatus muscle on the greater tubercle, and then the whole strip was taped along the supraspinatus muscle along the spine of the scapula to the muscle's origin, with paper-off tension and with approximately 20% to 25% stretch. Deltoid muscle was taped using Y strip as well, which was applied from anchor site, 3 cm below deltoid insertion to its origin, with paper-off tension. The front tail of Y strip was taped along the anterior edge of the deltoid, and the back tail was applied along the posterior edge of the deltoid muscle. Finally, the glenohumeral joint was taped using an I strip, which was applied from a coracoid process following laterally, below the acromion, and around the posterior deltoid edge.

*Dose*: MWM - 10 repetitions, 3 sets daily, 30-second rest period between sets, in 10 sessions with 24hours' rest between sessions. Kinesiotape - applied after initial ROM measure, removed on day 5 and reapplied following ROM measures

Frequency of administration: daily for 10 days

#### **Control: supervised exercise**

*Components of intervention*: pendulum exercises and pain-limited, active ROM exercises of shoulder elevation, depression, flexion, abduction, rotations, and strengthening exercises. Strengthening exercises were isometric in nature, working on the external shoulder rotators, internal rotators, biceps, deltoid, and scapular stabilisers (rhomboids, trapezius, serratus anterior, latissimus dorsi, and pectoralis major). The participants were instructed to perform all the exercises to the first onset of pain

*Dose*: 10 repetitions in 1 set daily, 30-second rest periods between sets of different types of exercises; 10 sessions with 24 hours between sessions

Frequency of administration: daily for 10 days

Outcomes

Outcomes assessed at 5 days and 10 days

• Active ROM (abduction and flexion) measured using a goniometer

Manual therapy and exercise for rotator cuff disease (Review)



## Djordjevic 2012 (Continued)

Notes

Conflicts of interest: the authors stated that they had no conflicts of interest

**Risk of bias** 

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "Participants were randomly allocated to group 1 or group 2. To ensure balance between the 2 groups, we used a minimization process as a form of restricted randomization. Minimization was run by Minim version 1.5, a mini- mization program for allocating patients to treatments in clinical trials". Comment: An adequate method was used to generate the allocation se- quence
Allocation concealment (selection bias)	Unclear risk	Comment: No information on how the allocation sequence was concealed was reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The third author, a physiotherapist and certified MWM and KT prac- titioner with experience in orthopedic rehabilitation of more than 15 years, was responsible for both groups' treatments. This third author was blind to the group assignment and also to the ROM measured on days 0, 5, and 10. She was also instructed not to discuss with the subject if his/her treatment was any dif- ferent from the usual program applied to the painful shoulder." Comment: Participants received different multimodal interventions, but were not provided with any information that would lead them to believe one inter- vention was superior or inferior to the other
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "Outcome measures were measured by the second author, who also re- mained blind to the group assignment."
Objective outcomes		Comment: Only objective outcomes were measured (ROM) and these were measured by a blinded assessor
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "All subjects went through each phase of the study (flow diagram show- ing the progress of subjects at each stage of the clinical trial)" Comment: There were no drop-outs, exclusions or losses to follow-up (there- fore all randomised participants were analysed)
Selective reporting (re- porting bias)	Low risk	Comment: Outcome data were fully reported for all outcomes listed in the Aus- tralian and New Zealand Clinical Trials Registry entry
Other bias	Low risk	Comment: No other sources of bias identified

Engebretsen 2009	
Methods	Study design: Parallel group RCT
	Setting: Outpatient clinic of physical medicine and rehabilitation department in Oslo, Norway
	Intervention: Supervised exercises (1 session weekly for up to 12 weeks)
	<b>Control:</b> Radial extracorporeal shockwave treatment (1 session weekly for 4–6 weeks)
	Source of funding: Supported by Health Region East, Norway
Participants	Diagnostic label used by trialists: Impingement
	Criteria for defining the shoulder condition being treated

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## Engebretsen 2009 (Continued)

Subacromial shoulder pain using the following diagnostic criteria: dysfunction or pain on abduction, normal passive glenohumeral ROM, pain on two of three isometric tests (abduction at 0 degrees or 30 degrees, external or internal rotation), and a positive Kennedy-Hawkins sign

## Inclusion Criteria (not listed above)

- Age: 18-70 years old
- People with rotator cuff rupture were included if they fulfilled the above criteria

### **Exclusion Criteria (not listed above)**

- Bilateral shoulder pain
- · Previous surgery on the affected shoulder
- Instability
- Clinical signs of a cervical syndrome
- Rheumatoid arthritis
- Clinical and radiological signs of glenohumeral or acromioclavicular arthritis
- Inability to understand Norwegian
- Serious psychiatric disorder
- Use of anticoagulant drugs (except low dose aspirin)
- Pregnancy
- Previous experience of one of the study interventions
- Unwillingness to accept either of the interventions in the study

## **Baseline characteristics**

#### Intervention

Number randomised: 52; mean (SD) age: 49 (9.3) years; sex: F/M 26/26; duration of symptoms: 3-6 months: 19 (37); 6-12 months: 15 (29); 12-24 months: 8 (15); > 24 months: 10 (19)

### Control

Number randomised: 52; mean (SD) age: 47 (11.7) years; sex: F/M 26/26; duration of symptoms: 3-6 months: 15 (29); 6-12 months: 15 (29); 12-24 months: 6 (12); > 24 months: 16 (31)

## Interventions Intervention: supervised exercises

*Components of intervention*: the principle focus was on relearning of normal movement patterns, which could then be transferred to daily activities. The initial aim was to unload the stress on the rotator cuff and subacromial structures. During this phase, a mirror for awareness of posture, an elastic rubber band and a sling fixed to the ceiling were used. The participants received immediate feedback and correction (supervision) by the physiotherapist. Once dysfunctional neuromuscular patterns were normalised, endurance exercises were performed with gradually increasing resistance. Participants had an adjusted programme at home, which consisted of correction of alignment during daily living and simple low loaded exercises with a thin elastic cord to provide assistance and resistance to the movement. Simple advice was given

Dose: 45 min

Frequency of administration: 2 sessions weekly for up to 12 weeks

## Control: radial extracorporeal shockwave treatment

*Components of intervention*: 3 to 5 tender points were treated each time. Points were identified through a participant-oriented biofeedback process (insertion of supraspinatus tendon, dorsolaterally below the acromion, and a maximum of three trigger points in the rotator cuff muscles). Radial extracorporeal shockwave treatment uses low to medium energy shockwaves generated when a projectile is accelerated by compressed air and hits an applicator. These impulses are delivered into the tissue and spread as spherical 'radial' waves (rather than being focused). Participants were informed that the suggested



Engebretsen 2009 (Continued)			
	mechanism for pain relief was hyperstimulation analgesia and increased neurovascularisation that im- proves regeneration of tissue. Participants were advised to avoid activities that elicited pain		
	<i>Dose</i> : frequency: 12-8 Hz with 2000 pulses per session, with a pressure between 2.5 and 4.0 Bar, de- pending on what the participant tolerated without anaesthetic		
	Frequency of administration: 1 session weekly for 4-6 weeks		
Outcomes	Outcomes assessed at 6 weeks, 12 weeks, 18 weeks and 1 year		
	• Function: Shoulder Pain And Disability Index (SPADI); score: 0 -100, higher score indicating worse shoulder pain and disability		
	• Rest pain in the previous week, measured on a 9-point Likert-type scale, 1 indicates no pain and 9 indicates severe pain		
	<ul> <li>Pain during activity in the previous week, (9-point Likert-type scale, 1 indicates no pain and 9 indicates severe pain)</li> </ul>		
	Active ROM (no outcome data reported)		
	<ul> <li>Work disability (recorded as working full time or &lt; 50% or unemployed)</li> </ul>		
	Adverse events		
Notes	Conflicts of interest: the authors stated that they had no conflicts of interest		
	Trial registered in ClinicalTrials.gov (NCT00653081)		

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "A statistician not involved in data collection or analysis randomly allo- cated patients to treatment groups in blocks of four to six. Randomisation was stratified by sex."
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Low risk	Quote: "A person not involved in the treatments opened the sealed envelopes and assigned appointments according to treatment group."
		Comment: An adequate method was likely used to conceal the allocation se- quence
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "Participants and personnel could not be blinded for this trial."
		Given the nature of the interventions, participants were not blind to treat- ment, and may have had different expectations about the benefits of each in- tervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants, who may have had different expectations about the benefits of the intervention they received self-reported some out-comes
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "A blinded physiotherapist made the baseline and follow-up measure- ments. The patients were instructed not to discuss their treatment with the blinded physiotherapist."
		Comment: Assessor of objective outcomes was likely blinded to the interven- tion
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: 10 participants per group did not return for all follow-up measures, and while reasons for loss to follow-up were not reported, an intention-to- treat analysis was performed

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## Engebretsen 2009 (Continued)

Selective reporting (re- porting bias)	High risk	Comment: No outcome data were reported for active ROM despite this out- come being listed in the methods section of the trial report. Several outcomes not specified in the ClinicalTrials.gov registry entry were added to the publica- tion (e.g. function, active ROM, work status)
Other bias	High risk	Quote: "Thirteen patients in the radial extracorporeal shockwave group and three patients in the supervised exercise group received additional treatment (cortisone injections, chiropractic treatment, physical therapy/supervised ex- ercises) between 12 and 18 weeks (odds ratio 5.5, 95% confidence interval 1.3 to 26.4; P=0.014)."
		Quote: "In the follow-up period [up to 1 year], 10 participants in the SE group and 20 participants in the rESWT group had additional treatments (P = 0.024)".
		Comment: There was an imbalance between groups in the number of addi- tional treatments received outside of the trial setting, which is likely to have biased the results in favour of the radial extracorporeal shockwave group

Methods	Study design: Parallel group RCT			
Methous	Study design: Parallel group RC1			
	Setting: Metropolitan hospital, Australia			
	Intervention: Physical modalities, passive joint mobilisation and ROM exercises			
	Control 1: Glucocorticoid injection			
	Control 2: Exercise therapy			
	Source of funding: Cumberland Research Grants from The University of Sydney, Sydney, Australia			
Participants	Diagnostic label used by trialists: none specified			
	Criteria for defining the shoulder condition being treated			
	<ul> <li>Unilateral shoulder pain of local mechanical origin, defined as pain over the shoulder joint and/or proximal arm, which was exacerbated by active shoulder movements. Pain could be with or without stiffness. Pain for more than one month</li> </ul>			
	Inclusion Criteria (not listed above)			
	Over 18 years of age			
	Able to understand spoken English			
	Exclusion Criteria (not listed above)			
	Bilateral shoulder pain			
	Associated with instability			
	Due to an inflammatory or neoplastic disorder			
	<ul> <li>Referred from vertebral column structures - if pain was not reproduced by active shoulder mov ments, if it was reproduced by active neck movements or by palpation of the cervico-thoracic vert bral columns or if paraesthesia were present in affected upper limb</li> </ul>			
	Due to trauma within previous four weeks			
	Baseline characteristics			
	Intervention			

Ginn 2005 (Continued)	Number randomised: 42; mean (range) age: 57.4 (29-90) years sex: F/M 16/26; mean (SD) duration of symptoms: 7.4 (10.9) months				
	Control 1				
	Number randomised: 48; mean (range) age: 55.4 (29-87) years; sex: F/M 19/29; mean (SD) duration of symptoms: 7.4 (11.2) months				
	Control 2				
	Number randomised: 48; mean (range) age: 52.6 (22-83) years; sex: F/M 21/27; mean (SD) duration of symptoms: 7.3 (8.1) months				
Interventions	Intervention: physical modalities, passive joint mobilisation and ROM exercises				
	<i>Components of intervention</i> : combination of electrophysical modalities (interferential therapy, ul- trasound therapy, hot packs and ice packs), passive joint mobilisation at the sternoclavicular and acromioclavicular joints and ROM exercises (functional movements of the arm and could incorporate the use of aids to achieve additional range of movement). The aim of the exercise component was to increase the range of hand placement but excessive scapular movement was discouraged. There was no requirement that the ROM exercises be performed in a pain-free manner. Exercises were upgraded from active assisted to active to resisted active exercises using free weights or elastic resistance. The specific treatment for each of the subjects was individually determined by the treating physical thera- pist using data from the initial interview and musculoskeletal assessment and any additional informa- tion gathered				
	<i>Frequency of administration</i> : twice weekly attendance for application of passive joint mobilisation and electrophysical modality components, and daily adherence to prescribed exercise programme for 5 weeks Control 1: glucocorticoid injection				
	<i>Components of intervention</i> : single injection of 40 mg methylprednisolone acetate, administered in- to the sub-acromial space under local anaesthesia with lignocaine. Participant was encouraged to at- tempt to use their affected upper limb in a normal manner and to await contact from the investigators at the end of the 5-week treatment period to arrange a time for reassessment				
	Components of intervention: single injection given				
	Control 2: exercise therapy				
	<i>Components of intervention</i> : the target exercise treatment was directed toward the restoration of nor- mal shoulder muscle function in order to restore dynamic stability and muscle co-ordination at the shoulder region. This comprised stretches aimed at lengthening shortened shoulder muscles, exercise aimed at strengthening weakened shoulder muscles, including improving co-ordination between mus- cles, and motor retraining aimed at restoring scapulohumeral rhythm during the performance of uppe limb tasks. All exercises were to be pain-free and subjects in this treatment group were also advised to avoid/limit pain producing activities. Particular emphasis was placed on restoring the normal muscle force couple co-ordination and the dynamic stabilising function of shoulder muscles. The specific exer- cises for each of the subjects was individually determined by the treating physical therapist, using data from the initial interview and musculoskeletal assessment and any additional information gathered by the treating physical therapist. The exercise treatment was administered as a home-based, daily exer- cise programme with supervision by the physical therapist once per week, to correct and upgrade the intensity and complexity of the exercises				
	Frequency of administration: daily for 5 weeks				
Outcomes	Outcomes assessed at 5 weeks				
	<ul> <li>Pain after standardised reaching task measured on a 10 cm vertical visual analogue scale labelled "n pain" and "severe pain" at its extremes</li> </ul>				

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Ginn 2005 (Continued)

- Function measured using a 4-point scale of increasing difficulty ranging from 0 = "can perform with no shoulder pain" to 3 = "cannot perform because of shoulder pain". Functional limitation score = summation of scores for each item with maximum of 27 denoting worst function
- Global assessment of treatment success: perceived change in symptoms was measured by 3-point scale which included "getting better", "staying the same" and "getting worse"
- Active ROM: abduction and flexion measured using photographic method, hand-behind-back (HBB) ROM score determined by subtracting the affected side measurement of HBB from the unaffected side measurement of HBB (HBB = distance between T1 spinous process and the radial styloid process with a tape measure with the subject standing)
- Strength: Isometric abduction measured using a hand-held dynamometer

Conflicts	of interest: not reported
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## **Risk of bias**

Notes

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "Randomly allocated to 1 of the 3 treatment groups by 1 of the re- searchers, using intervention assignment schedules previously prepared sepa- rately for those subgroups."
		Comments: No information on how the allocation sequence was generated was reported
Allocation concealment (selection bias)	Unclear risk	Comments: No information on how the allocation sequence was concealed was reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants, who may have had different expectations about the benefits of the intervention they received, self-reported some out-comes
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "Two senior physical therapists otherwise not associated with the clin- ical trial, acted as assessors over the length of the study; the first for the initial 22 months and the second for the remaining 24 months. Subjects were specifi- cally requested not to discuss their treatment with the assessor to ensure she/ he remained unaware of the treatment group to which the subject had been allocated."
		Comment: Assessors of objective outcomes were likely blind to treatment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Eleven subjects were unavailable for reassessment at the end of the 5-week treatment period: 6 from the PR subgroup and 5 from the P subgroup. One subject from the injection group died during the treatment period; 1 sub- ject from the exercise group moved interstate; and 9 subjects, 2 from the injec- tion group, 4 from the exercise group and 3 from the MPM group, were unavail- able for unknown reasons"
		Comment: The amount and reasons for attrition are unlikely to have affected the results
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results

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#### Ginn 2005 (Continued)

Other bias

Low risk

# Giombini 2006 Methods Study design: Parallel group RCT Setting: Athletes who attended the Physiotherapy Department of the Sport Science Institute, Italy Intervention: Exercises Control 1: Microwave diathermy Control 2: Therapeutic ultrasound Source of funding: Not reported Participants Diagnostic label used by trialists: Supraspinatus tendinopathy Criteria for defining the shoulder condition being treated: Diagnosis of supraspinatus tendinopathy of the dominant shoulder based on following three criteria · Impingement with a positive Hawkins sign in internal rotation or impingement in 90 degrees of forward flexion with forced external rotation; Pain with supraspinatus muscle testing in the 'empty can' position; Ultrasonographic evidence of nonhomogenous signal intensity without a frank tear in the supraspinatus tendon Inclusion Criteria (not listed above) Gradual onset of pain Participant engaged in sport at county, regional, national or international level and training in chosen sport at least 3 times a week All participants were secondary referrals to the fellowship-trained sports physicians or orthopaedic surgeons with a special interest in sports traumatology or shoulder surgery from family practitioners or physical therapists, as well as tertiary referrals from other orthopaedic surgeons or sports physicians. All participants had undergone nonoperative management, including complete or modified rest from their sports, and several (3-8) 1-week cycles of nonsteroidal anti-inflammatory drugs **Exclusion Criteria (not listed above)** · Athletes without full passive ROM of the affected shoulder Supraspinatus tendinopathy after a single traumatic episode • Severe neck pain, frozen shoulder, calcific tendinopathy, degenerative joint disease of the acromioclavicular or glenohumeral joint Intra-articular or subacromial injections of corticosteroids Clinical or ultrasonographic diagnosis of a rotator cuff tear · Previous surgery in the affected or contralateral shoulder **Baseline characteristics** Intervention Number randomised: 11; mean (SD, range) age: 26.3 ± 6.2 years, range 20-38 years; sex: F/M 2/9; duration of symptoms: not reported

#### Control 1

Number randomised: 14; mean (SD, range) age: 25.3 ± 4.8 years, range 19-37 years; sex: F/M 2/12; duration of symptoms: not reported Giombini 2006 (Continued)

#### Control 2

Number randomised: 12; mean (SD, range) age:  $28.6 \pm 6.6$  years, range 19-43 years; sex: F/M 4/8; duration of symptoms: not reported

Interventions Intervention: exercises Components of intervention: supervised and home exercises, consisting of pendular swinging in the prone position in flexion and extension of the shoulder and passive glenohumeral stretching exercises to tolerance Frequency of administration: supervised exercises once a week for 4 weeks; home exercises 5 min per day, every day for 4 weeks **Control 1: microwave diathermy** Components of intervention: an ALBA Hyperthermia System was used which was equipped with a 433.92 MHz microwaves generator with a maximum output power of 100 W; a microstrip antenna applicator, with a curve shape specific for semicylindrical joint volumes of 20-30 cm in diameter and with a total radiating area of 240 cm<sup>2</sup> and an effective field size; and a pad of silicone 0.5 cm thick, filled with thermostatic deionized water that allows the greatest energy transfer to be achieved while preventing overheating of superficial tissues near the radiant source. A hydraulic thermoregulation and one or two skin temperature sensors were also used. The thermocouple was placed on the shoulder with the participant lying supine and the arm at 60 degrees of abduction and externally rotated. It was placed over the middle third of the joint line between the glenoid fossa and the humeral head. The thermocouple on the skin was perpendicular to the electromagnetic field Dose: 434 MHz; administered at a power between 50 and 70 W, a pilot temperature on the skin between 38 and 40 degrees centigrade, and a water pad temperature between 35 and 37 degrees centigrade according to the depth of the subcutaneous fat of each participant. Each session lasted 30 min Frequency of administration: 3 times a week for 4 weeks **Control 2: therapeutic ultrasound** Components of intervention: continuous ultrasound was administered with the participant in the same position as participants receiving hyperthermia and by slowly moving the transducer in a circular fashion along the area distal to the anterior border of the acromion and the inferior third of a line between the glenoid fossa and the humeral head. A gel couplant was used between the ultrasound transducer and the skin of the area undergoing treatment. A Level 730 device was used. It was equipped with an emission probe of 1 MHz frequency, a sound head with an effective radiating area of 10cm<sup>2</sup> and a maximum output power of 22 W. Dose: 1 MHz at an intensity of 2.0 w/cm<sup>2</sup>; each session lasted 15 min Frequency of administration: 3 times a week for 4 weeks Outcomes Outcomes assessed at 4 weeks and 10 weeks Function measured by Constant-Murley score (0-100) Rest pain measured on a 0-10 VAS Global assessment of treatment success: measured by number of participants who felt ready to return to sport at the end of the experimental period Night pain measured on a 0-10 VAS (no outcome data reported) • Pain on activity measured on a 0-10 VAS (no outcome data reported) • Pain with resisted movement measured on a 4-point scale (0 = no pain, 1 = slight pain but full strength, 2 = moderate pain and reduced strength; 3 = severe pain and inability to exert any strength against minimal manual resistance); measured with active resisted abduction in the neutral position, active abduction in external rotation and active resisted abduction in internal rotation (no usable outcome data reported) Adverse events

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## Giombini 2006 (Continued)

Notes

Conflicts of interest: the authors stated that they had no conflicts of interest

**Risk of bias** 

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "Subjects were randomised into 3 groups using a computer-generated list."
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Unclear risk	Comment: No information on how the allocation sequence was concealed was reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported some out-comes
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "The subjects were assessed by fully trained sports physicians who had never seen the patients and were unaware as to which intervention the pa- tients had been allocated."
		Comment: Assessor of objective outcome was likely blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: There was no loss to follow-up and all randomised participants were analysed
Selective reporting (re- porting bias)	High risk	Comment: Data for pain on resisted movement were reported in figure only as means with no error bars. No data for night pain, pain on movement, rest pain and painful arc were reported, despite being listed as outcomes in the meth- ods section of the trial report
Other bias	Low risk	Comment: No other sources of bias were identified

## Haahr 2005

Methods	Study design: Parallel group RCT		
	Setting: Herning Hospital, Ringkjoebing County, Denmark		
	<b>Intervention:</b> Exercises plus heat, cold packs or soft tissue treatment (i.e. not all participants received soft tissue treatment)		
	Control: Arthroscopic subacromial decompression		
Source of funding: Medical Research Unit of Ringkjoebing County, Denmark			
Participants	Diagnostic label used by trialists: Impingement		
	Criteria for defining the shoulder condition being treated		

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Haahr 2005 (Continued)

The presence of shoulder pain, pain on abduction of the shoulder with painful arc, positive impingement sign (Hawkin's sign), positive impingement test (relief of pain within 15 min after injection of local anaesthetic into the subacromial space)

### Inclusion Criteria (not listed above)

- Fulfilment of all diagnostic criteria
- Age between 18 and 55 years old
- Normal passive glenohumeral movement
- Previous treatment with rest, non-steroidal anti-inflammatory drugs, subacromial injection, and physiotherapy were allowed

#### **Exclusion Criteria (not listed above):**

- Impaired rotation in the glenohumeral joint
- History of acute trauma
- · Previous surgery or history of fracture in the proximity of the affected shoulder
- Known osteoarthritis in the acromioclavicular or glenohumeral joints
- Calcifications exceeding 2 cm in the rotator cuff tendons
- Signs of a rupture of the cuff
- · Cervical root syndromes

#### **Baseline characteristics**

#### Intervention

Number randomised: 45; mean (SD): 44.5 (1.2) years; sex: F/M 29/14; duration of symptoms: < 6 months: 3; 6-12 months: 10; > 1 year: 29

#### Control

Number randomised: 45; mean (SD) age: 44.3 (1.3) years; sex: F/M 29/12; duration of symptoms: < 6 months: 4; 6-12 months: 3; > 1 year: 34

Interventions

## Intervention: physiotherapy

*Components of intervention*: the treatments started with application of heat, cold packs, or soft tissue treatments. This was followed by active training of the periscapular muscles (rhomboid, serratus, trapezoid, levator scapulae, pectoralis minor muscles) and strengthening of the stabilising muscles of the shoulder joint (rotator cuff). This was done within the limits of pain

Dose: 60 min

*Frequency of administration*: total 19 sessions. First 2 weeks: 3 times weekly. Next three weeks: twice weekly. Last seven weeks: once weekly. After 12 weeks of the trial, participant encouraged to continue the programme 2 to 3 times per week at home

## Control: arthroscopic subacromial decompression

*Components of intervention*: investigation for stability of the shoulder joint under general anaesthetic followed by an arthroscopic examination of the glenohumeral joint, the rotator cuff and the subacromial bursa. The treatment consisted of bursectomy with partial resection of the antero-inferior part of the acromion and the coracoacromial ligament. Before discharge, the participant was instructed in performing light movements of the arm within the limits of pain. Stitches were removed by general practitioners after 10 days. At the same time, the participant was instructed by a physiotherapist to carry out increasingly active exercises, including exercises for strengthening the rotator cuff muscles. The team instructing the physiotherapy group was different from the group treating the surgery group. The surgeon then saw the participants after 6-8 weeks

Outcomes

Outcomes assessed at 3, 6 and 12 months, and 4-8 years

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Risk of bias	
Notes	Conflicts of interest: not reported
	Work disability (self-reported as currently working) at 4-8 years
	• Strength: Constant sub-score (range 0-25 where higher = more strength) at 3, 6 and 12 months
	<ul> <li>Active ROM: Constant sub-score (range 0-40 where higher = more ROM) at 3, 6 and 12 months</li> </ul>
	<ul> <li>Global assessment of treatment success ("recovered or improved" versus "unchanged" or "worse or much worse") at 4-8 years</li> </ul>
	<ul> <li>Overall pain: Constant sub-score measured on a VAS scale (range 0–15; 15 = no pain) at 3, 6 and 12 months; PRIM VAS (0-9; 0 = no pain) at 4-8 years</li> </ul>
	<ul> <li>Function: Constant total score (0 – 100 scale where a higher score = normal function) at 3, 6 and 12 months; PRIM score (0-36 where a higher score = worse function) at 4-8 years</li> </ul>

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "A computer program was used to generate a random sequence of allo- cation."
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Low risk	Quote: "The same specialist (SØ) carried out all the assessments, obtained in- formed consent for participation, and randomised the patients into one of two intervention groups by opening a sealed envelope containing the result of ran- domisation, which was unknown to SØ."
		Comment: An adequate method was used to conceal the allocation sequence
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported some out-comes
Blinding of outcome as- sessment (detection bias) Objective outcomes	High risk	Quote: "The same specialist (SØ) carried out all the assessments, obtained in- formed consent for participation, and randomised the patients into one of two intervention groups by opening a sealed envelope containing the result of ran- domisation, which was unknown to SØ."
		Quote: "Physiotherapists were not blinded to the treatment given when as- sessing the Constant score".
		Comment: Assessor of objective outcomes was not blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Ninety consecutive patients with subacromial impingement agreed to participate. Forty five cases were randomised to conservative treatment and 45 to surgical treatment. Among those assigned to conservative treatment, one withdrew from participation because of work problems and one failed to fill in the baseline questionnaire, leaving 43 cases in this group. In the surgery group, four cases dropped out before the start of the study (one because of work problems, one with a tumour in the humerus, one because his wife ad- vised against participation, and one for unknown reasons), leaving 41 cases in this group. Within the conservative treatment group, a further six participants were operated on within the 12 months of the study (five because of unsatis- factory improvement during exercises and in one case because a labral lesion was suspected). In the physiotherapy group 42 persons (93%) were followed

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Haahr 2005 (Continued)		
		for 12 months with the main outcome measure (Constant score). In the surgery group 40 persons (89%) had complete follow up data.
		Quote: "Seventy-nine (88%) answered the final questionnaire" [at 4-8 years' follow-up]
		Comment: The amount and reasons for drop-out are unlikely to have affected the results
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: no other sources of bias were identified

Methods	Study design: Parallel group RCT				
	Setting: University, orthopaedic clinics, and community public places, Spain				
	Intervention: Thoracic spine manipulation				
	Control: Sham manipulation				
	Source of Funding: Coordenacao de Aperfeicoamento de Pessoal de Nivel Superior				
Participants	Diagnostic label used by trialists: Shoulder impingement syndrome				
	Criteria for defining the shoulder condition being treated				
	<ul> <li>Present with at least three of the following findings: positive Neer impingement test, positive Hawkin impingement test, positive Jobe test, pain with passive or isometric resisted shoulder lateral rotation pain with active shoulder elevation, pain with palpation of rotator cuff tendons, and pain in the C5 of C6 dermatone region</li> </ul>				
	Able to reach at least 150 degrees of arm elevation				
	Inclusion Criteria (not listed above)				
	• None				
	Exclusion Criteria (not listed above)				
	<ul> <li>Red flags for spinal manipulation (e.g. fracture, osteoporosis, malignancy, infection, and active in flammatory process)</li> </ul>				
	Pregnancy				
	Systemic illnesses				
	<ul> <li>Physical therapy or manual therapy treatment within 6 months prior to the evaluation</li> <li>Signs of complete rotator cuff tear or acute inflammation</li> </ul>				
	<ul> <li>Cervicothoracic spine-related symptoms (positive cervical compression test and excessive kyphosi</li> </ul>				
	Scoliosis				
	Glenohumeral instability (positive apprehension, anterior drawer, or sulcus tests)				
	Previous upper extremity fracture or shoulder surgery				
	Baseline characteristics				
	<i>Intervention</i> Number randomised: 26 (25 completed); mean age: 33.8 ± 12.2 years old; sex: F/M 11/14; duration of symptoms: 49 ± 96 months				



Haik 2014 (Continued)	<i>Control</i> Number randomised: 2 symptoms: 42.6 ± 66 m	26 (25 completed); mean age: 29.7 ± 9.3 years old; sex: F/M 7/18; duration of nonths	
Interventions	Intervention: thoraci	c spine manipulation	
	<i>Components of intervention:</i> low-amplitude, high velocity thrust thoracic spine manipulation. The par- ticipant assumed a seated position and the physiotherapists performed a thrust technique, targeting the midthoracic spine		
	Dose: if no cavitation w	vas detected with the manipulation, the thrust was repeated up to 3 times	
	Frequency of administr	ration: once	
	Control: sham manip	ulation	
	<i>Components of intervention:</i> the participant assumed the same seated position and the physiotherapist held the participant in the same position as that of the thrust manipulation intervention. The physio-therapist applied the same forces as those of a thrust manipulation, while holding the position for a few seconds, without actually performing a thrust manipulation		
	Dose: as above		
	Frequency of administration: once		
Outcomes	<ul> <li>Outcomes assessed immediately post-intervention (day 1)</li> <li>Pain on motion (elevation and lowering of the arm): numerical pain rating sca (worst pain)</li> </ul>		
Notes	<b>Conflicts of interest:</b> authors stated they had "no affiliations with or financial involveme ganisation or entity with a direct financial interest in the subject matter or materials discuarticle"		
	Trialists also assessed	scapular kinematics but these were not included in our review outcomes	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Quote: "Using computer-generated lists, one for the impingement group and the other for the asymptomatic group, subjects were randomly assigned to 1 of 4 groups: a TSM impingement group (n = 25), a sham impingement group (n = 25), a TSM asymptomatic group (n = 24) and a sham asymptomatic group (n = 23)".	
		Comment: An adequate method was used to generate the allocation sequence	
Allocation concealment (selection bias)	Unclear risk	Comment: There was no information on how the allocation sequence was con- cealed	
Blinding of participants and personnel (perfor- mance bias)	Low risk	Quote: "The subjects were given only general information about the purpose of the study to control expectations and to conduct an effective sham intervention".	
All outcomes		Comment: Participants were blinded	

Blinding of outcome as-<br/>sessment (detection bias)Low riskComment: Blinded participants self-reported all outcomes of interest to the<br/>reviewSelf-reported outcomesSelf-reported outcomes

Manual therapy and exercise for rotator cuff disease (Review)

## Haik 2014 (Continued)

Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: One participant from each group was excluded post-randomisation because of a fault in the equipment used to assess scapular kinematics. How- ever, this small amount of attrition is unlikely to have biased the results
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias identified

## Hay 2003

Methods	Study design: Parallel group RCT		
	Setting: Primary care – general practitioner, UK		
	<b>Intervention:</b> Physiotherapy (all participants: advice and instructions on pain relief and active shoul- der exercises at home; dependent on participant: ultrasound and active and passive mobilisation) <b>Control:</b> Glucocorticoid injection		
			Source of funding: Arthritis Research Council
	Participants	Diagnostic label used by trialists: None specified	
Criteria for defining the shoulder condition being treated			
<ul> <li>New episode of unilateral shoulder pain diagnosed by a general practitioner. New episode defined as "pain in shoulder region, including the upper arm, elicited or exacerbated by active or passive shoul der movement, and no consultation for this pain in the affected shoulder in the previous 12 months</li> </ul>			
Inclusion Criteria (not listed above)			
Aged 18 and above			
Exclusion Criteria (not listed above)			
<ul> <li>History of inflammatory arthritis, polymyalgia rheumatica or gross structural or neurological abnormality of the shoulder</li> </ul>			
Contraindications to local steroid injection			
<ul> <li>History of examination leading to suspicion of potentially serious disease</li> </ul>			
Referred pain from neck or internal organs			
Clinical findings of ruptured cuff			
Previous fracture or surgery to shoulder, upper limbs, neck or thorax			
<ul><li>Previous physical therapy for shoulder pain within past 12 months</li><li>Pregnancy or breast feeding</li></ul>			
Baseline characteristics			
Intervention			
Number randomised: 103; mean (SD) age: 57.5 ± 13 years; sex: F/M 50/53; mean (range) duration of symptoms: 51 (21-120) days			
Control			
Number randomised: 104; mean (SD) age: 57.6 ± 14 years; sex F/M 60/44; mean (range) duration of symptoms: 58 (28-128) days			



Hay 2003 (Continued)

Interventions

### Intervention: physiotherapy

*Components of intervention*: the most frequently used modality at the assessment visit was a standardised education and advice leaflet for shoulder pain (85%) followed by a home exercise programme (79%) which was reinforced throughout the trial treatment course. The most frequently utilised modalities over the treatment period were ultrasound (42%), active mobilisations (41%) and passive mobilisations (41%)

#### Dose: 20 min

Frequency of administration: 8 individual physiotherapy sessions delivered within a 6-week period

### **Control: glucocorticoid injection**

*Components of intervention*: injection of methylprednisolone with lidocaine into the subacromial space, administered by GP according to standard technique: the tip of the acromium and head of the humerus were identified by palpation and the injection point (just behind the mid-line in the gap between the acromium and head of the humerus) was marked; the skin was cleaned and the needle inserted perpendicular to the skin pointing slightly upwards under the acromium and local steroid with lidocaine injected easily without resistance. Participants were advised to avoid overuse of the shoulder for 48 hours and told that they could make an appointment to return within 4 weeks if their symptoms persisted. If they did return, they were offered a second injection.

Dose: 40 mg of methylprednisolone with 4 ml 1% lidocaine (lignocaine)

*Frequency of administration*: 1 injection; if symptoms persisted participant could make appointment within 4 weeks of initial injection and have a second injection

Outcomes	Outcomes assessed at 6 weeks and 6 months		
	<ul> <li>Function: Croft Shoulder Disability Questionnaire (SDQ) scores range 0–23, 23 indicating severe dis- ability</li> </ul>		
	Day pain measured	on 10 cm VAS.	
	<ul> <li>Night pain measure</li> </ul>		
	<ul> <li>Global assessment of treatment success: participant's global assessment of change compared with baseline measured on 5-point scale of "complete recovery" to "much worse"</li> <li>Quality of life measured on EuroQol</li> </ul>		
<ul> <li>Active ROM: restricted active abduction - subjects not achieving maximum 180 degrestricted active external rotation - subjects with restriction of &gt; 50% compared wit arm; restricted passive external rotation - subjects with restriction &gt; 50% compared wit arm</li> </ul>			
Notes	<b>Conflicts of interest:</b> not reported 25% of participants had adhesive capsulitis		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Quote: "Treatment allocation was according to the study number. Numbers were issued in a predetermined random sequence, in blocks of 10 by general practice, generated by a random number table."	
		Comment: An adequate method was used to generate the allocation sequence	
Allocation concealment (selection bias)	Low risk	Quote: "The number corresponded with that on a sealed envelope issued to the patient by the nurse. Participants were instructed not to open the enve- lope until the nurse had left. The envelope contained information instructing the participant to either make an appointment with one of the trial physiother- apists or to return to their GP for a local steroid injection."	

Manual therapy and exercise for rotator cuff disease (Review)

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Hay 2003 (Continued)

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nay 2003 (Continuea)		Comment: An adequate method was used to conceal the allocation sequence
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported some out- comes
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "Outcome assessments were performed by the study nurse, who was unaware of the treatment allocation." Comment: Assessor of objective outcomes was likely blinded to the interven-
		tion
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "The completion rate of the trial at six months was 95% (196/207) with the following reasons for loss to follow up: five other medical complications, two personal problems, four could not be contacted/refused visit. Intention to treat analysis was used."
		Comment: The amount and reasons for dropout are unlikely to have affected the results
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias were identified

Methods	Study design: Parallel group RCT		
	Setting: Private physiotherapy practice, Spain		
	<b>Intervention:</b> Manual therapy based on soft tissue techniques in the cervical and upper thoracic re- gions plus infrared plus TENS plus ultrasound		
	<b>Control:</b> Mobilisation, proprioceptive neuromuscular facilitation, supervised exercises plus infrared plus TENS plus ultrasound		
	Source of funding: Not reported		
Participants	Diagnostic label used by trialists: Impingement		
	Criteria for defining the shoulder condition being treated		
	Negative response to cervical compression tests		
	<ul> <li>Positive results in two out of three specific tests which evaluate the compromise of the subacromia space (Neer's test, Jobe's test and Yergason's test)</li> </ul>		
	Inclusion Criteria (not listed above)		
	Older than 18 years of age		
	Exclusion Criteria (not listed above)		
	Diagnosed acromial malformations		



Heredia-Rizo 2013 (Continued)	<ul> <li>History of neurodegenerative disease of the central or peripheral nervous system</li> <li>Previous history of fracture and/or surgical interventions of the shoulder joint, the scapular or the spine at any level</li> <li>Osteitis, rheumatic or tumoral diseases at any joint or spinal level</li> <li>Shoulder infiltrations in the three months before the study</li> <li>Previous rotator cuff damage</li> </ul>		
	Baseline characteristics		
	Intervention		
	Number randomised: 11; mean (SD) age: 54 $\pm$ 9.75 years; sex F/M 4/7; duration of symptoms: not reported		
	Control		
	Number randomised: 11; mean (SD) age: 62 ± 10.96 years; sex: F/M 5/6; duration of symptoms: not re- ported		
Interventions	Intervention: manual therapy based on soft tissue techniques		
	<i>Components of intervention</i> : manual therapy was started with micro-mobilisations of the cervical struc- tures in all movement axes. Thus, a selective traction of each vertebra in the longitudinal axis was per- formed with additional lateral movements in the transverse axis of the restricted zones. Subsequent- ly relaxation manoeuvres were performed to fascial restrictions involving the cervical and scapulo- humeral region, especially of the trapezius, the sternocleidomastoid, levator scapulae, subscapularis and pectoral muscles. Finally, adhering to the principles of orthopaedic manual therapy described by Kaltenborn, a repositioning of the head of the humerus was conducted in different stages:		
	<ul> <li>traction of the head and the diaphysis of the humerus (caudal sliding), aimed towards separation of osseous surfaces;</li> </ul>		
	<ul> <li>dorsal sliding of the head of the humerus with scapular fixation;</li> <li>"combined" movement of both (dorsal and caudal sliding). Active exercises were not recommended, except for pendular movements using 1 kg of weight in the prone position</li> </ul>		
	Dose: 40 min		
	Frequency of administration: 5 days a week for 3 weeks		
	Control: mobilisation, proprioceptive neuromuscular facilitation, supervised exercises		
	Components of intervention: this comprised 20 min of:		
	<ul> <li>passive, active and active-assisted mobilisations of the shoulder without causing any pain;</li> <li>variations of proprioceptive neuromuscular facilitation (PNF) patterns.</li> </ul>		
	In addition they completed a daily programme of 20 min of supervised active exercises, such as pendu- lar movements using 1 kg of weight in prone, assisted active movements with a pulley, and propriocep- tive exercises with a ball in the horizontal plane		
	Dose: 40 min		
	Frequency of administration: 5 days a week for 3 weeks		
	Both groups		
	<i>Components of intervention</i> : in the first place, infrared was applied on the shoulder for 15 min (Infra 2000, EnrafNonius). Afterward, TENS was used with a frequency of 80 Hz, 150 ms for 30 min (Med 911, Enraf-Nonius). Lastly, an ultrasound device (Sonopuls 492, Enraf- Nonius) with a power of 1.5 W/cm <sup>2</sup> and a frequency of 3 MHz in pulsating mode was applied for 5 min.		
	Frequency of administration: 5 days a week for 3 weeks		

# Heredia-Rizo 2013 (Continued)

Outcomes

Outcomes assessed at 3 weeks

- Function: DASH questionnaire
- Active and passive ROM (flexion-extension, abduction-adduction, internal and external rotation) measured using a goniometer

Notes

Conflicts of interest: the authors stated that they had no conflicts of interest

**Risk of bias** 

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "After signing an informed consent according to the principles gath- ered in the Helsinki Declaration (2008 version), the patients were randomized to the Conventional (n = 11) or Experimental Group (n = 11) by means of sealed opaque envelopes."
		Comment: No information regarding how the allocation sequence was gener- ated was reported
Allocation concealment (selection bias)	Low risk	Quote: "After signing an informed consent according to the principles gath- ered in the Helsinki Declaration (2008 version), the patients were randomized to the Conventional (n = 11) or Experimental Group (n = 11) by means of sealed opaque envelopes."
		Comment: An adequate method was probably used to conceal the allocation sequence
Blinding of participants and personnel (perfor- mance bias)	Unclear risk	Quote: "A randomized and single blind (the practitioner carrying out the mea- surements remained unaware of the subject's group membership) clinical study was conducted."
All outcomes		Comment: Despite being described as assessor-blinded only, participants re- ceived slightly different types of manual therapy and exercise, but it is unclear whether they were provided with any information that would make them per- ceive the type of manual therapy and exercise they received as superior or in- ferior to the alternative type of manual therapy and exercise
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	Unclear risk	Comment: Participants self-reported function, but it is unclear whether they were provided with any information that would make them perceive the type of manual therapy and exercise they received as superior or inferior to the al- ternative type of manual therapy and exercise
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "A randomized and single blind (the practitioner carrying out the mea- surements remained unaware of the subject's group membership) clinical study was conducted."
		Comment: Outcome assessor of objective outcomes (ROM) was blind to treat- ment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: No drop-outs, losses to follow-up or exclusions were reported, and the number of participants randomised was reported as the number of participants analysed
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias identified

Manual therapy and exercise for rotator cuff disease (Review)



#### Holmgren 2012

Methods	Study design: Parallel group RCT				
	Setting: Department of orthopaedics, University hospital, Sweden				
	<b>Intervention:</b> Specific exercise programme plus subacromial corticosteroid injection, information about shoulder condition, ergonomic advice and advice on correction of posture, and manual therapy when required				
	<b>Control:</b> Non-specific exercises plus subacromial corticosteroid injection, information about shoulder condition, ergonomic advice and advice on correction of posture, and manual therapy when required				
	<b>Source of funding:</b> This research received no specific grant from any funding agency in the public, commercial, or not-for-profit sectors				
Participants	Diagnostic label used by trialists: Impingement				
	Criteria for defining the shoulder condition being treated				
	Primary subacromial impingement syndrome, diagnosed by an orthopaedic specialist and on waiting list for arthroscopic subacromial decompression. Shoulder condition defined by:				
	<ul> <li>typical history of pain located in proximal lateral aspect of the arm, especially with the arm raised;</li> <li>3 of the following: impingement sign according to Neer, impingement syndrome according to Hawkins-Kennedy, positive result on Jobes test, positive result on Patte's manoeuvre;</li> <li>positive Neer's impingement test (injection of 1 ml of 20 mg/mL triamcinolon mixed with 6 ml of 10 mg/mL mepivacain)</li> </ul>				
	Inclusion Criteria (not listed above)				
	<ul> <li>Lack of response to various conservative treatments (including exercise treatment) for at least 3 months</li> </ul>				
	Exclusion Criteria (not listed above)				
	<ul> <li>Significant loss of flexion, abduction or strength in rotation indicating a major cuff tear</li> <li>Radiologically verified malignancy</li> <li>Osteoarthritis of the glenohumeral joint</li> <li>Os acromiale decreasing the subacromial space</li> <li>Acromioclavicular arthritis</li> <li>Previous fractures in the shoulder complex or shoulder surgery on the affected side or both</li> <li>Clinically verified polyarthritis, rheumatoid arthritis, fibromyalgia</li> <li>Instability in any joint of the shoulder complex</li> <li>Frozen shoulder</li> <li>Symptoms from the cervical spine and pseudoparalysis</li> <li>Receipt of a glucocorticoid injection in the previous three months for the current problem</li> <li>Inability to understand written and spoken Swedish</li> </ul>				
	Baseline characteristics				
	Intervention				
	Number randomised: 50; mean (SD) age: 52 (9) years; sex: F/M 14/37; duration of symptoms: median 24 months, range: 6–120 months				
	Control				
	Number randomised: 52; mean (SD) age: 52 (8) years; sex F/M 22/24; duration of symptoms: median 12 months, range: 6-156 months				



## Holmgren 2012 (Continued)

Interventions

#### Intervention: specific exercise programme

*Components of intervention*: the programme consisted of 6 different exercises: 2 eccentric exercises for the rotator cuff (supraspinatus, infraspinatus, and teres minor), 3 concentric/eccentric exercises for the scapula stabilisers (middle and lower trapezius, rhomboideus, and serratus anterior), and a posterior shoulder stretch. The exercises were individually adjusted and progressed with increased external load by using weights and elastic rubber band at the physiotherapist visits once every other week during the whole rehabilitation period. The individual resistance for each participant was determined by using the pain monitoring model. The participant was not allowed to exceed 5 on this 0-10 scale when they performed the exercises however, they were recommended to feel some pain during loading. After completion of an exercise session, increased pain had to revert to levels before exercise before the next session otherwise the external load was decreased. Great emphasis was placed on teaching good posture (thoracic spine extension and retracted shoulder) and to maintain this position during the exercises

*Dose:* each strengthening exercise performed 15 times in 3 sets, twice daily for 8 weeks; posterior shoulder stretch performed 30-60 seconds 3 times daily for 8 weeks. From week 8–12 the exercises were repeated once a day. After 12 weeks when the specific exercise programme had finished, participants were encouraged to maintain the daily home exercises for 2 months

*Frequency of administration*: participants saw research physiotherapist once a week for the first 2 weeks and once every other week for the next 10 weeks (a total of 7 visits). The first visit lasted about 60 min, and the subsequent visits lasted about 30 min. In between these supervised sessions, participants performed home exercises once or twice a day for 12 weeks

### **Control: non-specific exercises**

*Components of intervention*: 6 unspecific movement exercises for the neck and shoulder without any external load (shoulder abduction in the frontal plane, shoulder retraction, shoulder elevation, neck retraction, stretch of upper trapezius and pectoralis major). The unspecific exercise programme was thought to have limited effect in people with subacromial impingement syndrome and therefore acted as a control. Participants did the same programme without any progression during the whole rehabilitation period

*Dose*: each movement exercise was repeated 10 times, and each strengthening exercise 3 times twice daily at home and once every other week at the physiotherapist visits

*Frequency of administration*: participants saw research physiotherapist once a week for the first 2 weeks and once every other week for the next 10 weeks (a total of 7 visits). The first visit lasted about 60 min, and the subsequent visits lasted about 30 min. In between these supervised sessions, participants performed home exercises once or twice a day for 12 weeks

#### **Both groups**

*Components of intervention:* subacromial glucocorticoid injection at the inclusion visit. When necessary, the physiotherapist performed manual treatment by stretching the posterior glenohumeral capsule and pectoralis minor during the visits. Thorough information about their shoulder condition, ergonomic advice and correction of their posture. Exercises introduced 2 weeks after the injection

OutcomesOutcomes assessed at 3 months and 12 months\*• Function: Constant-Murley shoulder assessment score – consists of objective (ROM and strength) and<br/>subjective (pain assessment, work load and strength) measurements; score 0–100; higher score indi-<br/>cates better function• Rest pain during the previous 24 hours measured on a visual analogue score (VAS); score: 0–100• Pain during activity measured on a VAS score 0–100• Night pain during the previous 24 hours measured on a VAS score 0–100• Quality of life: EuroQol instrument (EQ-5D)• Global assessment of treatment success: participant's global impression of change in symptoms be-<br/>cause of treatment measured on a 5-point Likert scale: worse, unchanged, small improvement, large

improvement or recovered



Holmgren 2012 (Continued)	Required surgery: whether the participant underwent surgery with the year after the 12-week exercise programme ended
Notes	Conflicts of interest: The authors stated that they had no conflicts to declare
	Trial registered in ClinicalTrials.gov (NCT01037673)
	*We only extracted 12-month outcome data for the 'required surgery' outcome, as 12-month data for all other outcomes were sub-grouped by whether participants underwent surgery post-exercise intervention or not

# Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "An independent physiotherapist prepared the random allocation se- quence beforehand. Equal numbers of the two treatment alternatives, 55 of each, were prepared and concealed in opaque envelopes. These were then mixed by hand and numbered. At the inclusion visit, the orthopaedic specialist (HB) coded the patients consecutively."
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Low risk	Quote: "The two treatment alternatives were prepared and concealed in opaque envelopes. Treatment allocation was performed at the first visit to the physiotherapist, within two weeks of the inclusion visit. The research physio- therapist received the envelope with the corresponding code revealing the as- signed treatment alternative out of a central locked location just before the participants presented for one of the two treatments: specific exercises (spe- cific exercise group) or unspecific exercises (control exercise group)."
		Comment: An adequate method was used to conceal the allocation sequence
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "The control exercise programme consisted of six unspecific movement exercises for the neck and shoulder without any external load (shoulder ab- duction in the frontal plane, shoulder retraction, shoulder elevation, neck re- traction, stretch of upper trapezius and pectoralis major. Each movement ex- ercise was repeated 10 times, and each stretching exercise three times twice daily at home and once every other week at the physiotherapist visits. The pa- tients did the same programme without any progression during the whole re- habilitation period. The unspecific exercise programme was thought to have a limited effect in patients with subacromial impingement syndrome and there- fore acted as a control".
		Comment: Participants (but not personnel) were blind to treatment (non-spe- cific movements were completed by the control group, who were likely un- aware that their movements were placebo exercises)
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	Low risk	Comment: Blinded participants self-reported some outcomes
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "The same orthopaedic specialist evaluated all primary and secondary outcome measures at the inclusion visit before patients started the exercises (baseline) and after three months when patients had completed their exercise programme. The specialist was blinded to the group assignment throughout the study."
		Comment: Assessor of objective outcomes was likely blind to treatment

Manual therapy and exercise for rotator cuff disease (Review)

Holmgren	2012	(Continued)
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Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "A total of 152 patients were eligible for inclusion; 102 patients met the inclusion criteria and provided written informed consent to participate. Three weeks after inclusion, five patients were excluded: two patients developed a frozen shoulder, diagnosed by the physiotherapist three weeks after inclusion, and three patients changed their minds about participating in the study and declined participation at the first physiotherapist visit because of lack of time. A total of 97 patients were compliant with the study protocol from baseline to the three month assessment and were included in the statistical analysis". Quote: "Ninety-five patients were assessed at 1-year follow-up. Two patients in each group did not attend the 1-year follow-up due to non-related disease" Comment: The amount and reasons for attrition are unlikely to have affected the results
Selective reporting (re- porting bias)	Low risk	Comment: Outcome data fully reported for all outcomes specified in the clini- cal trials registry
Other bias	Low risk	Comment: No other sources of bias were identified

anse van Rensburg 2			
Methods	Study design: Parallel group RCT		
	Setting: National Health Service physiotherapy department, UK		
	Intervention: Thoracic spinal manipulation plus mobilisation plus supervised exercises		
	Control: Mobilisation plus supervised exercises		
	Source of funding: Not reported		
Participants	Diagnostic label used by trialists: Impingement		
	Criteria for defining the shoulder condition being treated		
	<ul> <li>Clinical findings demonstrating subacromial impingement syndrome</li> <li>Shoulder pain of more than 1 week's duration</li> <li>Limitation of shoulder range of movement</li> <li>Pain produced during flexion and/or abduction of the shoulder</li> <li>Thoracic hypomobility detected on clinical examination</li> <li>Irrespective of diagnosis of subacromial impingement syndrome (SIS) by imaging, at least 2 of the following clinical signs must be present: positive painful arc, positive Neer's sign, positive Hawkin</li> </ul>		
	Kennedy sign Inclusion Criteria (not listed above)		
	<ul> <li>Men and women between 18 and 65 years of age</li> </ul>		
	Exclusion Criteria (not listed above)		
	<ul> <li>Reproduction of shoulder symptoms during active or passive cervical movements</li> <li>Presence of clinical signs of glenohumeral instability, e.g. positive sulcus, load and shift sign or a ca sular pattern</li> <li>History of severe trauma such as fracture, dislocation or cuff tear or shoulder surgery on the sympt matic shoulder</li> </ul>		
	<ul> <li>Possible serious pathology (previous cancer or tuberculosis, bone infections, significant unexplain weight loss, HIV (significant risk of concurrent serious pathology))</li> </ul>		



Janse van Rensburg 2012 (Continued)

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	Both groups	<i>ition</i> : physiotherapy comprising active or passive glenohumeral mobilisa-	
	low)	<i>ntion</i> : participants only received the treatment common to both groups (see be-	
		<b>plus supervised exercises</b>	
		ation: once a week for 6 weeks	
		to the shoulder	
	Components of interver nipulation was applied		
	Components of intervention: a specific high velocity low amplitude 'extension with rotation' thrust ma-		
Interventions	Intervention: thoracic spinal manipulation		
	symptoms: 5 (4-6) mon		
	Number randomised: 3	; mean (range) age: 60 (58-65) years; sex: F/M 1/2; mean (range) duration of	
	Control		
	symptoms: 9 (1-24) mo		
		; mean (range) age: 53 (41-60) years; sex: F/M 2/4; mean (range) duration of	
	Intervention		
	Baseline characteristi	cs	
	Refusal to participat	te in the study	
	Treatment from and	other practitioner for the same condition in the last 6 weeks	
		urbance/widespread motor changes, pregnancy)	
	bromatosis, non-me	echanical thoracic pain, bilateral paraesthesia in upper or lower limbs, sphincter	
		rns (oral steroids/chemotherapy medication, Ehlers–Danlos syndrome, neurofi-	
		sis, osteogenesis imperfecta, Paget's disease, significant thoracic trauma)	
	•		
		abnormal clotting, anticoagulant therapy, deep vein thrombosis/pulmonary em-	
	•		
	•	se (rheumatoid arthritis, ankylosing spondylitis, psoriatic arthritis)	
	•		
	•		
	•		
	•		
	•	abnormal clotting, anticoagulant therapy, deep vein thrombosis/pulmonary em-	
		abnormal clotting, anticoagulant therapy, deep vein thrombosis/pulmonary em-	
		aononnai ciotting, anticoagutant therapy, ueep vent thrombosis/puthollary em-	
		abilitina county, and cougarant and apy, acep vent an on bools/partionally ent	
	bolism)		
	bolism)		
	•		
	Other clinical conce	erns (oral steroids/chemotherapy medication, Ehlers–Danlos syndrome, neurofi-	
	bromatosis, non-me	echanical thoracic pain, bilateral paraesthesia in upper or lower limbs, sphincter	
	alterations, gait dist	urbance/widespread motor changes, pregnancy)	
	-		
	<ul> <li>Treatment from and</li> </ul>	ther practitioner for the same condition in the last 6 weeks	
	<ul> <li>Treatment from and</li> </ul>	ther practitioner for the same condition in the last 6 weeks	
		•	
	Refusal to participat	te in the study	
	Baseline characteristi	cs	
	Intervention		
	mervention		
	Control		
	Number randomised: 3	: mean (range) age: 60 (58-65) years: sex: E/M 1/2: mean (range) duration of	
Interventions	Intervention: thoracic spinal manipulation		
	Dose: 30 min		
	Frequency of administra	ation: once a week for 6 weeks	
	Control: mobilisation	plus supervised exercises	
	Control: mobilisation	plus supervised exercises	
		<i>tion</i> : participants only received the treatment common to both groups (see be-	
	Both groups		
	Components of interver	ation: physiotherapy comprising active or passive glenohumeral mobilisa-	
	tion; transverse friction	n massage to the rotator cuff tendons; exercises to stimulate the lower fibres of	
		otator cuff-strengthening. Exercises, and ergonomic and lifestyle advice was	
	provided in line with cu	irrent physiotherapy practice	
	Dose: 30 min		
	D030. 30 mm		
	Frequency of administre	ation: Once a week for 6 weeks	
Outcomes	Outcomes assessed at 6 weeks		
Outcomes	Outcomes assessed at 6 weeks		
	<ul> <li>ROM in flexion and a</li> </ul>	abduction using a goniometer (unclear if active or passive)	
	<ul> <li>Function using the I</li> </ul>	Disability of the Arm, Shoulder and Hand (DASH) score	
	-		
	<ul> <li>Adverse events</li> </ul>		
Notes	Conflicts of interest: r	not reported	
		·	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
DINJ	Autors judgement		
Random sequence genera-	Low risk	Quote: "The participants were randomized into the control or experimental	
tion (selection bias)	LOW HISK	group using unmarked envelopes. As it was anticipated that 20 subjects would	
		would using unmarked envelopes as it was anticipated that 70 subjects would	

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be recruited, there were 20 envelopes, 10 of which contained a sticker with the

Janse van Rensburg 2012 (Co	ontinued)	
		words 'control – no manipulation' and 10 which contained a sticker with the words 'experimental – manipulation'. The investigator had no involvement in the randomization. As participants arranged their treatment sessions at re- ception, the administration staff asked each patient to choose one of the un- marked envelopes.
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Low risk	Quote: "The participants were randomized into the control or experimental group using unmarked envelopes. As it was anticipated that 20 subjects would be recruited, there were 20 envelopes, 10 of which contained a sticker with the words 'control – no manipulation' and 10 which contained a sticker with the words 'experimental – manipulation'. The investigator had no involvement in the randomization. As participants arranged their treatment sessions at reception, the administration staff asked each patient to choose one of the unmarked envelopes. The administration staff opened the envelope and placed the sticker on the inside of the patient's treatment card so that the treating therapist would know which group the patient was assigned to and the patient card, for instance in the filing cabinet, she would not know which group the patient was in as there would be nothing to identify the patient to a particular treatment group"
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants, who may have had different expectations about the benefits of the intervention they received, self-reported function
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "Pre and post every treatment session, the investigator or second trained data collector (who did not undertake treatments and were therefore blinded to group allocation) measured the shoulder range of movement of each patient."
		Comment: Assessor of objective outcome was likely blinded to treatment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Thirty-five patients were assessed by the researcher over a 4-week period, nine of whom met the inclusion criteria and were re- cruited into the pilot study. One subject did not wish to continue with physio- therapy and dropped out after the first treatment session. This participant had been randomized to the control group" Comment: The number of drop-outs was low and unlikely to affect the results
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias identified

# Kachingwe 2008

Methods	Study design: Parallel group RCT
	Setting: University health centre, USA

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Kachingwe 2008 (Continued)	Intervention 1: Glenohumeral mobilisation plus supervised and home exercises
	Intervention 2: Mobilisation with movement technique plus supervised and home exercises
	Control 1: Supervised and home exercise only
	Control 2: Physician advice only
	<b>Source of funding:</b> Supported by the California State University, Northride Research, Scholarship and Creative Activity Award
Participants	Diagnostic label used by trialists: Impingement
	Criteria for defining the shoulder condition being treated
	• Primary shoulder impingement diagnosed by the referring physician and superolateral shoulder pain and 2 out of 4 specified objective signs and symptoms: a positive Neer impingement test, a positive Hawkins-Kennedy impingement test, painful limitation of active shoulder elevation (flexion, abduc- tion, scaption), and pain or limitation with the functional movement patterns of hand-behind-back or hand-behind-head.
	Inclusion Criteria (not listed above)
	18 years of age or older
	Exclusion Criteria (not listed above)
	<ul> <li>Physician diagnosis of adhesive capsulitis</li> <li>Grade III rotator cuff tear</li> <li>Calcific tendonitis confirmed by radiology</li> <li>Systemic or neurological disorder</li> <li>Cervical radiculopathy</li> <li>History of shoulder surgery</li> <li>Corticosteroid injection within the past month</li> <li>Physical therapy treatment for the shoulder within the past 3 months</li> </ul>
	Baseline characteristics
	Intervention 1
	Number randomised: 9; mean (SD) age: 43.4 ± 14.7 years; sex: F/M 5/4; mean (SD) duration of symp- toms: 19.2 ± 24.6 months
	Intervention 2
	Number randomised: 9; mean (SD) age: 48.9 $\pm$ 13.7 years; sex: F/M 4/5; mean (SD) duration of symptoms: mean: 22.6 $\pm$ 17.4 months
	Control 1
	Number randomised: 8; mean (SD) age: 47.3 $\pm$ 20.1 years; sex: F/M 4/4; mean (SD) duration of symptoms: mean: 32.5 $\pm$ 60.2 months
	Control 2
	Number randomised: 7; mean (SD) age: 45.6 $\pm$ 13 years; sex: F/M 3/4; mean (SD) duration of symptoms: mean: 70 $\pm$ 92.4 months
Interventions	Intervention 1: glenohumeral joint mobilisations (group 1)
	<i>Components of intervention</i> : joint mobilisation was administered based on assessment of glenohumer- al joint anterior, posterior and inferior glides and long-axis distraction passive accessory motions using a 0-6 accessory motion scale. For situations where there was reactivity within the capsular ROM, grade

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Kachingw	e 2008	(Continued)
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I-II mobilisation were applied. For situations where there was no reactivity but capsular hypomobility, grade III-IV accessory motions were applied

*Dose*: 3 sets of 30-second mobilisations (each mobilisation applied for 30 seconds at a rate of approximately 1 mobilisation every 1-2 seconds, followed by a 30-second rest).

Frequency of administration: once a week for 6 weeks

### Intervention 2: glenohumeral joint mobilisation with movement (group 2)

*Components of intervention*: Mulligan technique: involved the therapist applying a sustained posterior accessory glide to the glenohumeral joint while the subject simultaneously actively flexed the shoulder to the pain-free endpoint and applied a gentle overpressure force using the contralateral arm. Total abolition of pain during the technique was mandatory; if the participant started to experience pain during active motion, the therapist would investigate different force planes and/or grades of force until pain-free motion was sustained; if pain commenced during any repetition of any set, the technique was terminated

Frequency of administration: once a week for 6 weeks

#### Control 1: supervised and home exercises (groups 1, 2 and 3)

*Components of intervention*: supervised exercises including posterior capsule stretching, postural correction exercises, and an exercise programme focusing on rotator cuff strengthening and scapular stabilisation. Each session ended with subjects receiving a cold pack for 10-15 min to decrease potential inflammation and delayed muscle soreness. Participants were instructed to perform a home exercise programme mimicking the exercises performed in the clinic

Frequency of administration: supervised exercise - once a week for 6 weeks; home exercises - once per day

### Control 2: advice (group 4)

*Components of intervention*: participant education on postural awareness and limitation of overhead activities. Advice administered by the referring physician during their initial physical examination. The physician also provided the subject with a standard shoulder impingement home exercise programme without any input from the physical therapist.

Outcomes	Outcomes assessed at 6 weeks			
	<ul> <li>Function measured in function</li> </ul>	by the Shoulder Pain and Disability Index (SPADI); score: 0-130, 130 = worst deficit		
	• Pain: maximum pain intensity over preceding 24 hours measured by 0-10 VAS			
	Active ROM (flexion and scaption) using a goniometer			
Notes	Conflicts of interest: not reported			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence genera- tion (selection bias)	Low risk	Quote: "Patients were randomly assigned to one of four intervention groups according to the block randomisation method. E.g. subject #1 had an equal chance of drawing an envelope assigning them to A, B, C or D. If they drew 'A', then the card was removed so the next participant had an equal chance of		

		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Unclear risk	Comment: It is unclear if adequate safeguards were put in place to conceal the allocation sequence

drawing an envelope with B, C or D. etc."

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Kachingwe 2008 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "Each subject was informed of his/her protocol but remained blinded to other group assignments to avoid subject bias" Comment: Despite not knowing what other participants received, expecta- tions about the effectiveness of interventions received may have differed be- tween groups, particularly between those receiving exercise and mobilisation versus those receiving physician advice only
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants, who may have had different expectations about the benefits of the intervention they received, self-reported some out-comes
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "One physical therapist with 12 years of clinical experience performed the pre- and post-treatment assessment measurements. This assessor was blinded to group assignment and all intervention protocols." Comment: Assessor of objective outcomes was likely blinded to the interven- tion
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: No dropouts and outcome data reported as based on total number of randomised participants
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias were identified

Kardouni 2014	
Methods	Study design: Parallel group RCT
	Setting: Physical and occupational therapy offices and physician's clinics, USA
	Intervention: Thoracic spinal manipulative therapy
	Control: Sham manipulative therapy
	<b>Source of Funding:</b> Clinical and Translational Science Award No. UL1TR000058 from the National Cen- ter for Advancing Translational Sciences and the AD Williams' Fund of the Virginia Commonwealth Uni- versity
Participants	Diagnostic label used by trialists: Subacromial impingement syndrome
	Criteria for defining the shoulder condition being treated
	<ul> <li>Have 3 of the following 5 clinical signs of subacromial impingement syndrome: positive Hawkin's Test positive Neer Test; pain during active elevation &gt; 60 in the scapular or sagittal plane; positive Jobe Empty Can test for pain or weakness; pain or weakness with resisted shoulder external rotation wit the arm at the side</li> </ul>
	Inclusion Criteria (not listed above)
	<ul> <li>Pain for 6 weeks</li> <li>Typical daily shoulder pain 2/10 on an 11-point numeric pain rating scale (NPRS)</li> <li>18 to 60 years of age</li> </ul>
	Exclusion Criteria (not listed above)

Trusted evidence. Informed decisions. Better health.

Kardouni 2014 (Continued)				
	<ul> <li>A primary complain</li> <li>Signs of central nervices</li> <li>Signs of cervical nervices</li> <li>Contraindications to arthritis</li> <li>Adhesive capsulitis</li> <li>Instability of the shot</li> <li>Shoulder or arm pair</li> </ul> Baseline characteristic	o manipulative therapy such as osteoporosis, metastatic disease, or systemic oulder n with cervical rotation to the ipsilateral side, axial compression, or Spurling's Test <b>ics</b>		
	65.9 months <i>Control</i>	24; mean age: 31.1 ± 12.3 years old; sex: F/M 14/10; duration of symptoms: 40.2 ± 24 (21 completed); mean age: 31.2 ± 12.1 years old; sex: F/M 9/12; duration of months		
Interventions	Intervention: thoraci	c spinal manipulative therapy (SMT)		
	thrust was applied at the lower thoracic SMT, the	ntion: during administration of the thoracic SMT, a high-velocity, low-amplitude he end of available spinal motion after the participant exhaled. For the mid and e participants were prone, and the thrust was directed in the posterior to anteri- rvicothoracic junction SMT, participants were seated, and the thrust was an axial		
	Dose: twice at each of the 3 regions, for a total of 6 manoeuvres			
	Frequency of administration: once			
	Control: sham manip	ulative therapy		
		<i>ntion:</i> the therapist maintained manual contact through the ROM during exhala- ive thrust was delivered		
	Dose: twice at each of t	he 3 regions, for a total of 6 manoeuvres		
	Frequency of administr	ation: once		
Outcomes	Outcomes assessed im	mediately post-treatment (pain) and at 1 to 2 days (pain, function, QoL)		
	<ul> <li>Function: Penn Sho</li> <li>= no pain or function</li> </ul>	ulder Score (patient-rated shoulder function/disability questionnaire) 0-100 (100 nal loss)		
		ic pain rating scale, from 0 (no pain at all) to 10 (pain as bad as it can be) al Rating of Change, from -7 (a great deal worse) through to 0 (no change), to +7		
Notes		'The authors certify that they have no affiliations with or financial involvement in tity with a direct financial interest in this study."		
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence genera- tion (selection bias)	Low risk	Quote: "A randomization list for treatment group assignments of the partici- pants was computer generated with random blocking using nQuery Advisor software (Statistical Solutions, Saugus, MA)".		

Manual therapy and exercise for rotator cuff disease (Review)



Kardouni 2014 (Continued)		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Low risk	Quote: "Treatment assignments were placed into sequentially numbered pri- vacy envelopes to conceal treatment group allocation".
		Comment: An adequate method was used to conceal the allocation sequence
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Participants were blinded to treatment assignment, and were told pri- or to the start of testing they could receive an active or a placebo treatment. In an effort to help maintain blinding and prevent participants from knowing that they were receiving the active or placebo treatment, both treatment groups were assigned names representative of active treatments. Participants ran- domized to the thoracic SMT group were told that they were receiving "spinal manipulative therapy" while those randomized to the sham thoracic SMT group were told they would receive a "therapist-assisted range of motion" treatment."
		Comment: Participants were blinded
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	Low risk	Comment: Blinded participants self-reported all outcomes of interest to the review
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Forty-eight (n = 48) individuals with SIS were randomly assigned to re- ceive thoracic SMT (n = 24) or sham thoracic SMT (n = 24). Three participants were excluded from the final analysis (all in the sham thoracic SMT group) because it was discovered after testing that they had pain in both shoulders, leaving n = 45 for final analysis".
		Comment: The amount of attrition is small and reasons were unrelated to the intervention, so attrition is unlikely to have biased the results
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias identified

# Kassolik 2013

Methods	Study design: Parallel group RCT
	Setting: General practice, Poland
	Intervention: Classic (Swedish) massage
	Control: Massage using techniques based on the tensegrity principle
	Source of funding: University School of Physical Education in Wroclaw
Participants	Diagnostic label used by trialists: None specified
	Criteria for defining the shoulder condition being treated
	<ul> <li>Shoulder pain of any duration determined by provocation of participant's pain on active shoulder abduction, flexion or external rotation</li> </ul>
	Inclusion Criteria (not listed above)
	Adults referred to participating in physiotherapy by General Practitioner

Manual therapy and exercise for rotator cuff disease (Review)

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Kassolik 2013 (Continued)

Cassolik 2013 (Continued)	Able to provide write	ten informed consent	
	Exclusion Criteria (no	t listed above)	
	<ul> <li>Any previous surgica</li> <li>Systemic condition polymyalgia, rheum</li> </ul>	tion of the affected shoulder in the last 5 years al intervention on the affected shoulder in the last 5 years with a significant musculoskeletal component (inflammatory joint disease atic, neoplastic cancer) ers (cervical, or thoracic spine hernia)	
	Baseline characteristi	cs	
	Overall cohort of partici	ipants	
		5 (18 in one group and 17 in the other); mean (SD) age of women 53.9 ± 16 years; I3.6 ± 12.3 years; sex: F/M 19/11; duration of symptoms: not reported	
Interventions	Intervention: classic (	Swedish) massage	
	performed in a side rec (Swedish) were used - s	<i>ntion:</i> classic massage of the shoulder girdle and glenohumeral joint was umbent position. During the massage, typical classic massage techniques stroking with the palms (effleurage), friction with the palms, kneading (petris- ottement), and vibration	
		vas performed 7 to 8 times in particular body parts (frequency 60 to 70 moves ulse rate); percussion and vibration were performed for 1 min on average	
	Frequency of administration: 5 times a week for 2 weeks		
	Control: massage using techniques of the tensegrity principle		
	of classic massage but anatomical structures	ntion: the techniques used for this method were the same as in the methodology were aimed at additional areas. Before the massage, palpation of the selected was carried out. The purpose of the assessment was to determine which tissues ivity and which showed increased tension. Based on palpation results, the mas- was performed	
	Dose: 20 min		
	Frequency of administre	ation: 5 times a week for 2 weeks	
Outcomes	Outcomes assessed at 2	2 and 6 weeks	
	(SF-MPQ)	g the 10 cm VAS (numerical pain scale) of Short Form McGill Pain Questionnaire	
	Active ROM (flexion,	extension, abduction, external and internal rotation) using a goniometer	
Notes	Conflicts of interest: the authors stated that they had no conflicts of interest		
	Trial was registered in ClinicalTrials.gov (NCT01307826)		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	High risk	Quote: "Patients assigned with even numbers were included in the classic group and ones with odd numbers to the tensegrity group".	
		Comment: A quasi-random allocation sequence was used	

Manual therapy and exercise for rotator cuff disease (Review)



# Kassolik 2013 (Continued)

Allocation concealment (selection bias)	High risk	Comment: A quasi-random (i.e. predictable) allocation sequence was used, therefore the allocation sequence was not concealed
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Patients were blinded to which group they were in." Comment: Given the nature of the interventions (i.e. one versus another type of massage), it is unlikely that participants perceived the type of massage they received as superior or inferior to the alternative type of massage
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	Low risk	Quote: "Patients were blinded to which group they were in." Comment: Blinded participants self-reported pain
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "Range of motion measurements of the glenohumeral joint by the go- niometric method were conducted by a physiotherapistwho had no knowl- edge of which group the patient was assigned." Comment: Outcome assessor of objective outcomes was blind to treatment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: The CONSORT flow chart (Schultz 2010) shows that three and two participants, respectively, did not receive the allocated intervention for rea- sons unrelated to the intervention (e.g. unable to attend appointment because of work), which is unlikely to bias the results. No other drop-outs, losses to fol- low-up or exclusions occurred
Selective reporting (re- porting bias)	Low risk	Comment: Outcome data were fully reported for all outcomes specified in the ClinicalTrials.gov registry entry (NCT01307826)
Other bias	High risk	Quote: "Before the therapy, the groups differed in 7 of 10 measured motions (lower range in the group with massage based on the tensegrity rule) while af- ter the therapy, the difference appeared in only 1 case. Statistically significant changes in the group with massage based on the tensegrity rule in some of the motions may prove to be the result of worse ROM at the very beginning of the study in this group. Perhaps, in the opinion of the authors, in such a condition of glenohumeral joint, it is easier to achieve such results." Comment: There was baseline imbalance in ROM, which may have favoured the group receiving massage based on the tensegrity principle

Methods	Study design: Parallel group RCT		
	Setting: Outpatient physiotherapy clinic, Turkey		
	Intervention 1: Manual therapy and exercise plus cold pack		
	Intervention 2: Kinesiotaping plus exercise plus cold pack		
	Source of Funding: No funding		
Participants	Diagnostic label used by trialists: Subacromial impingement syndrome		
	Criteria for defining the shoulder condition being treated		
	<ul> <li>Diagnosis of subacromial impingement syndrome based on the Hawkins-Kennedy impingement sign the painful arc sign, and the infraspinatus muscle test</li> </ul>		
	Inclusion Criteria (not listed above)		

Kaya 2014 (Continued)

• Aged between 30 and 60 years

# Exclusion Criteria (not listed above)

- Cervical spine involvement
- Presence of a glenohumeral joint adhesive capsulitis, or instability
- History of previous shoulder surgery
- Having another physiotherapy treatment of this disorder in the past 6 weeks
- Steroid injection into or around the shoulder in the past 2 months
- Recurrent complaints or long history of complaints over a year
- Massive rotator cuff or labral tear (assessed via MRI)

### **Baseline characteristics**

Intervention 1

Number randomised: 30 (26 completed); mean age:  $47.15 \pm 9.44$  years old; sex: F/M 16/10; duration of symptoms: 6-28 weeks

#### Intervention 2

Number randomised: 30 (28 completed); mean age: 50.85 ± 5.17 years old; sex: F/M 17/11; duration of symptoms: 6-26 weeks

Interventions

# **Intervention 1: manual therapy**

*Components of intervention:* general mobilisation, including superoinferior gliding, rotations, and distractions to the scapula, were applied 3 to 5 times. Also, neuromuscular facilitation techniques for scapula motions at anterior elevation–posterior depression and posterior elevation–anterior depression planes were performed up to 5 to 6 repetitions. Glenohumeral joint mobilisation with long axis traction and posterior or inferior glide techniques to improve shoulder internal rotation limitations were applied according to the individual requirements of the participants. Soft tissue massage and joint mobilisation of the neck, thoracic region, and elbow areas, according to the involvement, and deep friction massage with specific ischaemic compression technique were applied to supraspinatus muscle

Dose: total duration 1.5 hours

Frequency of administration: once a week for 6 weeks

# **Intervention 2: Kinesiotaping**

*Components of intervention:* application of kinesiotaping to the tissue that was in need of help. According to the Wright test result and muscle strength tests, the affected weak muscle groups including supraspinatus, upper and lower trapezius, deltoideus, teres minor, and levator scapulae were identified. The muscle technique was applied to the specifically affected muscle with no tension on band with a Y shape. Then, a correction technique for the protracted shoulder and a ligament technique for the overall shoulder were applied

Dose: 24 hours per day

*Frequency of administration:* standard 2 inch (5 cm) Kinesio Tex tape was applied once per week for 6 weeks. Each taping was removed after 4 to 5 days in situ

# Both groups - Exercise plus cold pack

*Components of intervention:* supervised and home exercises, including strengthening, flexibility (ROM) and Codman's pendulum exercises. Flexibility exercises were composed of posterior capsule with "cross-body stretch", upper thoracic extension stretch, and active ROM stretching for glenohumeral joint for flexion and abduction. Strengthening exercises had 3 sets of 10 repetitions, using a 150 cm long precut section of Thera-Band. The participants began exercising using the no-latex yellow band at mild tension, and when able to perform 3 sets of 15 repetitions without significant pain or fatigue, they were progressed to the next colour-resistive band in the sequence: red, green, and blue. Phase 1 emphasised the strengthening of the rotator cuff with avoidance of excessive upper trapezius activity and



(aya 2014 (Continued)			
	was instructed to cont on wall and push-up pl	g. Shoulder elevation exercises were added in phase 2, and in phase 3, the subject inue the exercises from phase 2 in addition to the new exercises such as push-up lus with Thera-Band. Cold pack gel application on the shoulder was recommend- nes a day, especially before and after exercises	
	Dose: total duration 1.	5 hours	
	Frequency of administr	ation: once a week for 6 weeks (in clinic); daily at home	
Outcomes	Outcomes assessed at	6 weeks	
	<ul> <li>Function: Disabilitie more disability</li> </ul>	es of the Arm, Shoulder and Hand (DASH) 0-100, where a higher score indicates	
		p pain) to 10 (unbearable pain)	
	<ul> <li>Pain on motion: VAS, 0 (no pain) to 10 (unbearable pain)</li> <li>Night pain: VAS, 0 (no pain) to 10 (unbearable pain)</li> </ul>		
Notes	<b>Conflicts of interest:</b> "No funding sources or conflicts of interest were reported for this study."		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Quote: "Randomization was designed according to the random case sample in SPSS program (SPSS, Chicago, IL). The SPSS software randomly assigned par- ticipants to one of the groups".	
		Comment: An adequate method was used to generate the allocation sequence	
Allocation concealment (selection bias)	Unclear risk	Comment: There was no information on how the allocation sequence was con- cealed	
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention	
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants, who may have had different expectations about the benefits of the intervention they received, self-reported all out- comes of interest to the review	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Four participants of 30 from the manual therapy group (MT group) dis- continued the study because of personal reasons. Two participants of 30 from the kinesiotaping group (KT group) left the study; 1 of them had severe skin ir- ritation. The other did not like to use the tape throughout the study."	
		Quote: "The subset of per-protocol analysis is an "as-treated" analysis in which only participants adherent to the intervention were included from all random- ized participants by using baseline-post-intervention analysis."	
		Comment: The amount of attrition is small, and while related to the interven- tion for one participant, is unlikely to have biased the results	
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results	
Other bias	Low risk	Comment: No other sources of bias identified	

Manual therapy and exercise for rotator cuff disease (Review)



# Kromer 2013

Methods	Study design: Parallel group RCT		
	Setting: Referred from general practitioners or orthopaedic surgeons, Germany		
	Intervention: Individualised manual physiotherapy plus individually adapted exercises		
	Control: Individually adapted exercises alone		
	Source of funding: Not reported		
Participants	Diagnostic label used by trialists: Impingement		
	Criteria for defining the shoulder condition being treated		
	Symptoms for at least 4 weeks		
	<ul> <li>Region of complaint is the glenohumeral joint region of the proximal arm</li> </ul>		
	<ul> <li>Presence of one of the following signs indicating subacromial impingement syndrome: Neer impingement sign, Harkins-Kennedy impingement test, painful arc with active abduction of flexion</li> </ul>		
	<ul> <li>Pain during one of the following resistance tests: external rotation, internal rotation, abduction or flexion</li> </ul>		
	Inclusion Criteria (not listed above)		
	Age between 18 and 75 years		
	Exclusion Criteria (not listed above)		
	<ul> <li>Mean 24 h pain of 8.10 or more on a VNRS</li> <li>Primary scapulothoracic dysfunction due to paresis</li> <li>Diagnosed instability or previous history of dislocation</li> <li>Adhesive capsulitis (frozen shoulder)</li> <li>More than 1/3 restriction of elevation compared with unaffected side</li> <li>Substantial shoulder weakness or loss of active shoulder function</li> <li>Shoulder surgery in the last 12 months on the involved side</li> <li>Development of symptoms with active or passive cervical movements</li> <li>Neurological involvement with sensory and muscular deficit</li> <li>Inflammatory joint disease (e.g. Rheumatoid arthritis)</li> <li>Diabetes mellitus</li> <li>Psychotherapeutic drug intake</li> <li>Compensation claims</li> <li>Inability to understand written or spoken German</li> </ul>		
	Baseline characteristics		
	Intervention		
	Number randomised: 46; mean (SD) age: 50.1 $\pm$ 12.2 years; sex: F/M 22/24; mean (SD) duration of symptoms: 27.4 $\pm$ 28.4 weeks		
	Control		
	Number randomised: 44; mean (SD) age: 53.7 ± 9.9 years; sex: F/M 24/20; mean (SD) duration of symp- toms: 40.8 ± 53.4 weeks		
Interventions	Intervention: individualised manual physiotherapy		
	<i>Components of intervention</i> : painful and angular and/or translatory restricted peripheral joints were treated with manual glide techniques according to the concept of Kaltenborn. Comparable signs of the		

Manual therapy and exercise for rotator cuff disease (Review)



### Kromer 2013 (Continued)

spine segments were treated with posterior-anterior glides or coupled movements. Shortened muscles were stretched according to the description of Evjenth & Hamberg. Neural tissue was treated according to Butler. Treatment intensity was limited by pain of > 4/10. Subsequent treatment decisions were made with the help of an adapted clinical reassessment process based on the test-retest principle described by Maitland

*Dose*: initial duration of the glide techniques and the stretches was 20–30 seconds.Further dosage was based on reassessment results. Session duration was 20-30 min

*Frequency of administration*: 10 treatment sessions within 5 weeks, followed by exercise programme 3 times a week for 7 more weeks

Control: individually adapted exercises alone

*Components of intervention*: participants only received the treatment common to both groups (see below)

### **Both groups**

Components of intervention: core exercise programme - dynamic exercises started with 2 sets of 10 repetitions and with low resistance (yellow rubber band); shoulder and neck stretches were held for 10 seconds and repeated twice; Isometric scapular training positions were held for 10 seconds and repeated twice. If participants performed the core programme without problem, sets were increased from 2 to 3, repetitions (respectively seconds for the static exercises) were increased from 10 to 20, and in a last step, resistance was increased from the yellow to the red and to the green rubber band. Exercises from an 'additional programme' could be added if the participant could still perform the core programme without problems. Participants were instructed on how to perform each single exercise. They received a booklet with pictures and descriptions of the exercises and the individually defined dosage. Participants had to stop an exercise if they had pain of more than 3 out of 10 on a VNRS during the exercises or longer than approximately 30 seconds after they had stopped an exercise. Participants recorded performance and difficulties with the programme in their log books which enabled the therapist to check the 24-hour effect of the programme and to make adaptations. If the total load of the programme was too provocative, participants were allowed to split the programme into 2 parts performing them at different times during the day. For some exercises an alternative version could be used (e.g. exercises C6b instead of C6a). If an exercise could not be performed due to pain, it was left out for the next 2 training sessions and was replaced by exercises AP1 and AP2. Contact time for the control group was 15-20 min

*Frequency of administration*: participants performed the exercises twice a day for the first week, then once daily. Minimum exercises frequency during the week was 4, maximum 7. Thereafter both groups continued their exercise programme for 3 times a week for 7 more weeks

Outcomes	Outcomes assessed at 5 and 12 weeks and 1 year		
	<ul> <li>Function measured using the total SPADI score (1-100) where a higher score indicates greater impairment</li> <li>Pain during the past week measured using visual numeric rating scale (11 point scale: 0 = no pain)</li> <li>Global assessment of treatment success measured using the Patient Global Impression of Change (PGIC) scale. Participants rated as "slightly better" or "much better" were considered successes</li> </ul>		
	Work disability (days of sick leave)		
	Adverse events		
Notes	Conflicts of interest: not reported		
	Trial is registered in Current Controlled Trials (ISRCTN86900354)		
Risk of bias			
Bias	Authors' judgement Support for judgement		

Kromer 2013 (Continued)		
Random sequence genera- tion (selection bias)	Low risk	Quote: "Patients who fulfilled the eligibility criteria were asked to sign in- formed consent, they underwent baseline assessment and were subsequent- ly allocated to treatment groups in blocks of 6 using central randomization via the internet."
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Low risk	Quote: "To guarantee allocation concealment, therapists received the infor- mation about patient allocation immediately before the first treatment by the Department of Epidemiology, Maastricht University".
		Comment: Central randomisation (i.e. an adequate method) was used to con- ceal the allocation sequence
Blinding of participants and personnel (perfor- mance bias)	High risk	Quote: "Due to the nature of the intervention it was impossible to blind thera- pist and participants. However, we blinded therapists for the control group to all clinical information about their patients."
All outcomes		Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported some out- comes (e.g. pain, function, global assessment)
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Finally, 90 participants were randomly allocated, with 44 patients in the control group (IAEX) and 46 patients in the intervention group (IAEX + IMPT). At 5 weeks all patients were analysed with no loss to follow-up. At 12 weeks 2 patients in the intervention group discontinued treatment, 1 without giving a reason, the other reported that treatment took too much effort."
		Quote: "After 1 year data were available for 87 patients; 44 patients in the IMPT group and 43 in the IEP group".
		Comment: Only three participants dropped out, one for reasons relating to the intervention. This small dropout rate is unlikely to have had a substantial impact on the results
Selective reporting (re- porting bias)	Low risk	Comment: Outcome data were fully reported for all outcomes specified in the clinical trial registry entry (Current Controlled Trials ISRCTN86900354) and published trial protocol
Other bias	Low risk	Comment: No other sources of bias identified

# Littlewood 2014

Methods

Study design: Parallel group RCT

Setting: Private physiotherapy clinic, UK

Intervention: Self-managed loaded exercise

**Control:** Usual physiotherapy (might include advice, stretching, exercise, manual therapy, massage, strapping, acupuncture, electrotherapy, corticosteroid injection at the discretion of the treating physiotherapist)

Source of funding: International Mechanical Diagnosis and Research Foundation (IMDTRF)

Manual therapy and exercise for rotator cuff disease (Review)

# Littlewood 2014 (Continued)

Participants

# Diagnostic label used by trialists: Rotator cuff tendinitis

## Criteria for defining the shoulder condition being treated

- Primary complaint of shoulder pain with or without referral into upper limb for > 3 months
- No/minimal resting shoulder pain
- Range of shoulder movement largely preserved
- Shoulder pain provoked consistently with resisted muscle tests (abduction and lateral rotation)

## Inclusion Criteria (not listed above)

- Aged over 18 years
- Willing and able to participate

## **Exclusion Criteria (not listed above)**

- Shoulder surgery within the last 6 months
- Possible systemic pathology including inflammatory disorders
- Cervical repeated movement testing affects shoulder pain and/or range of movement

# **Baseline characteristics**

### Intervention

Number randomised: 12; mean (range) age: 62.6 (46-76) years; sex: F/M 7/5; mean (range) duration of symptoms: 29 (3-120) months

#### Control

Number randomised: 12; mean (range) age: 63.9 (44-79) years; sex: F/M 5/7; mean (range) duration of symptoms: 49 (3-168) months

Interventions Intervention: self-managed loaded exercise Components of intervention: the intervention was prescribed by the physiotherapist but completed by the participant independently. It involved exercising the affected shoulder against gravity, a resistive therapeutic band or hand weight over three sets of 10 to 15 repetitions completed twice per day. Exercise prescription was guided by symptomatic response requiring that pain was produced during exercise, but overall, symptoms were no worse upon cessation of that exercise. The exercise was prescribed and operationalised within a self-managed framework which included focus upon knowledge translation, exercise/skill acquisition, self-monitoring, goal setting, problem solving and pro-active follow-up Frequency of administration: mean number of treatment sessions was 3.9 (participants received a maximum of four funded sessions) **Control: usual physiotherapy** Components of intervention: might include a range of interventions including advice, stretching, exercise, manual therapy, massage, strapping, acupuncture, electrotherapy, glucocorticoid injection at the discretion of the treating physiotherapist Frequency of administration: mean number of treatment sessions was 7.6 (participants received a maximum of eight funded sessions) Outcomes Outcomes assessed at 3 months post-treatment Function: SPADI score (0-100) where a higher score indicates greater pain and disability Quality of life measured using SF-36 scores (0-100) where a higher score indicates a better quality of life. SF-scores included: physical functioning, role-physical, bodily pain, general health, vitality, social functioning, role emotion and mental health Notes Conflicts of interest: the authors stated that they had no conflicts of interest

Manual therapy and exercise for rotator cuff disease (Review)

# Littlewood 2014 (Continued)

# **Risk of bias**

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "A computer generated randomisation sequence was produced by SJW in blocks of two and four to ensure an equal number of participants were ran- domised to each group".
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Low risk	Quote: "The treating physiotherapists allocated participants to the self-man- aged exercise or usual physiotherapy treatment group by selecting the next consecutively numbered sealed opaque envelope, which concealed the group allocation."
		Comment: An adequate method was used to conceal the allocation sequence
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "Design: A single-centre pragmatic unblinded Parallel-group RCT." Quote: "Similar to other RCTs of physiotherapy interventions, this trial was un- blinded which introduces a potential source of bias. Although we initially pro- posed a double-blind study, i.e. patient and hence outcome assessor, this was regarded as unacceptable by the ethics committee."
		Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported some out- comes (e.g. pain, function, quality of life)
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "100% retention was attained with all participants completing the SPADI at three months."
		Comment: There were no dropouts, losses to follow-up or exclusions
Selective reporting (re- porting bias)	Low risk	Comment: Outcome data were fully reported for all outcomes specified in the published protocol of this trial
Other bias	Low risk	Quote: "The groups appeared well balanced at baseline except that the self- managed exercise group reported higher baseline shoulder pain and disability via the SPADI and the usual physiotherapy treatment group reported a longer mean duration of symptoms (49 versus 29 months). This estimate is influenced by one participant who reported duration of 168 months. When the influence of this outlier was removed the revised estimate of mean duration of symp- toms was 37 months for the usual physiotherapy group."
		Comment: There was some baseline imbalance in SPADI score and duration of symptoms, though it is unlikely to have had an impact on the results

# Lombardi 2008

Methods

Study design: Parallel group RCT

Setting: Outpatients attending clinics of the Federal University of São Paulo, Brazil

Intervention: Progressive resistance training programme for the musculature of the shoulder

Manual therapy and exercise for rotator cuff disease (Review)



Lombardi 2008 (Continued)	Control: Waiting list control			
	Source of funding: Not reported			
Participants	Diagnostic label used by trialists: Impingement			
	Criteria for defining the shoulder condition being treated			
	<ul> <li>A positive Neer test and Hawkin test for the diagnosis of shoulder impingement syndrome in the pre- vious 2 months and pain between 3 and 8 on the numeric pain scale in the arc of movement that pro- duces the greatest shoulder pain</li> </ul>			
	Inclusion Criteria (not listed above)			
	• None			
	Exclusion Criteria (not listed above)			
	<ul> <li>History of shoulder fractures or dislocation, cervical radiculopathy, degenerative joint disease of the glenohumeral joint</li> <li>History of surgery of on the shoulder, back or thorax</li> <li>History of inflammatory arthropathy</li> <li>Infiltration of the shoulder in the previous 3 months</li> <li>Undergoing any type of physical intervention</li> </ul>			
	Baseline characteristics			
	Intervention			
	Number randomised: 30; mean (SD) age: 56.3 ± 11.6 years; sex: F/M 21/9; mean (SD) duration of symp- toms: 13.7 ± 9.6 months			
	Control			
	Number randomised: 30; mean (SD) age: 54.8 ± 9.4 years; sex F/M 25/5; mean (SD) duration of symp- toms: 13.9 ± 9.3 months			
Interventions	Intervention: training programme			
	<i>Components of intervention</i> : progressive resistance training programme. The exercises were flexion, ex- tension, medial rotation and lateral rotation of the shoulder. Participants underwent a muscle strength assessment using a repetition maximum (RM) exercise in which participants performed 6 repetitions with the maximum bearable weight thereby determining the 6-repetition maximum (6 RM). Once the 6 RM load was determined, training was divided into: 2 series of 8 repetitions, the first series with 50% of the 6 RM and the second series with 70% of the 6 RM, respecting the participant's pain threshold. The exercise was interrupted if the participant felt pain and performed another movement. Between the first and second series, there was a resting period of 2 min. The speed of movement was 2 seconds for both the eccentric and concentric phases. The 6 RM load was re-evaluted every 2 weeks. Multipulley muscle-building equipment was used for the exercises. To strengthen the flexors of the shoulder, the participant was positioned with his or her back to the equipment and the elbow flexed at 90 degrees; the participant performed the flexion movement of the shoulder from 0–90 degrees. In the extensor strengthening exercise, the participant faced the equipment with the elbow flexed at 45 degrees and the shoulder at 60 degrees of flexion and 30 degrees of extension. In the strengthening of the medial and lateral rotators, the participant was positioned alongside the equipment with the elbow flexed at 90 degrees; for the medial rotation, the participant started at 45 degrees of lateral rotation and moved to 45 degrees of medial rotation; for the lateral rotation, the participant began the movement at 45 de- grees of medial rotation and moved to 30 degrees of lateral rotation			

Frequency of administration: twice a week for a period of 8 weeks

# Control: no training programme

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Lombardi 2008 (Continued)	<i>Components of intervention</i> : participants remained on a waiting list and were informed that they would receive physiotherapeutic treatment after 2 months had passed		
	Both groups		
	<i>Components of intervention:</i> 750 mg of acetaminophen every 8 hours when experiencing pain. In cas- es where the pain surpassed 7 on the visual pain scale, the participant could take 50 mg of diclofenac every 8 hours until the pain reached a 5 on the pain scale. This was done at the participant's discretion		
Outcomes	Outcomes assessed at 2 months		
	<ul> <li>Function: DASH questionnaire; DASH 2 (used for laborious function), DASH 3 (activities of daily living) and optional module DASH 2 (module for labourers); score 0-100; 0 = best state, 100 = worst state</li> </ul>		
	<ul> <li>Rest pain measured on a 10 cm VAS; 0 cm = no pain, 10 cm = unbearable pain</li> </ul>		
	<ul> <li>Pain on movement measured on a 10 cm VAS; 0 cm = no pain, 10 cm = unbearable pain</li> </ul>		
	<ul> <li>Active ROM measured for flexion, abduction, internal rotation with shoulder at 90 degrees abduction, external rotation with shoulder at 90 degrees abduction, external rotation with arm alongside body, and extension using a goniometer</li> </ul>		
	<ul> <li>Strength: isokinetic strength (peak torque (Nm) and total work (joules) at a velocity of 60 degrees/sec- ond) respectively measured for flexion, extension, abduction, adduction, internal rotation and exter- nal rotation, measured using an isokinetic dynamometer</li> </ul>		
	• Quality of life measured using Brazilian form of the SF-36 (0-100) where a higher score indicates a bet- ter quality of life. SF-scores included: physical functioning, role-physical, bodily pain, general health, vitality (0-100), social functioning, role emotion and mental health		
	<ul> <li>Global assessment of treatment success: degree of participant satisfaction measured using a Likert scale: much worse, a little worse, unchanged, a little better, much better</li> </ul>		

Notes

Conflicts of interest: not reported

**Risk of bias** 

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "A computer-generated randomization list was utilized to randomly allocate patients into experimental and control groups and a concealed ran- domization with an opaque sealed envelope was performed."
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Low risk	Quote: "A computer-generated randomization list was utilized to randomly allocate patients into experimental and control groups and a concealed ran- domization with an opaque sealed envelope was performed."
		Comment: An adequate method was used to conceal the allocation sequence
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported the SPADI
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "Evaluations were carried out at the beginning and end of the treat- ment program by the same blinded examiner for both groups".
		Comment: Assessor of objective outcomes was likely blinded to treatment

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Lombardi 2008 (Continued)

Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "In cases of interruption or abandonment of treatment, the data was analyzed as intent-to-treat." Quote: "Sixty patients were randomly assigned to the experimental and con- trol groups, with 30 patients in each group. Four patients from the control group failed to finish the study: 1 who started having difficulties appearing at the rehabilitation center but appeared for the final evaluation, and 3 who failed to return for the final evaluation, stating difficulties appearing at the evaluation locale. Data from the prior evaluation of the patients from the con- trol group were used for the intent-to-treat analysis." Comment: The amount and reasons for dropout are unlikely to have affected the results
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias were identified

Methods	Study design: Parallel group RCT
	Setting: Construction workers recruited through local unions
	Intervention: Home exercise programme of 5 shoulder stretching and strengthening exercises
	Control: No treatment
	<b>Source of funding:</b> Center to Protect Worker's Rights, the Public Health Service, and the University of Iowa, USA (grant # U60/CCU317202)
Participants	Diagnostic label used by trialists: Impingement
	Criteria for defining the shoulder condition being treated
	<ul> <li>Reported history of shoulder pain localised to the glenohumeral joint region excluding cervical ar periscapular pain, but including the common site of referred pain of the rotator cuff to the C5–6 de matome above the deltoid insertion</li> </ul>
	<ul> <li>Present with at least two positive shoulder impingement tests (Neer, Hawkins/Kennedy, Yocum, Job and/or Speeds tests) and pain reproduction during two of three additional categories of clinical test These categories included:</li> </ul>
	<ul> <li>a painful arc on active scapular plane abduction of the arm;</li> <li>tendemonstern election of the biener engettern enff tendemonsterne</li> </ul>
	<ul> <li>* tenderness to palpation of the biceps or rotator cuff tendons; and</li> <li>* pain with one or more resisted glenohumeral joint motions (flexion, abduction, internal rotatio or external rotation). Flexion and abduction were resisted at 90 degrees of elevation, and intern and external rotation were resisted both at the subject's side and at 90 degrees of abduction</li> </ul>
	Inclusion Criteria (not listed above)
	<ul> <li>Self-reported occupational exposure to overhead work for longer than one year</li> <li>Minimum of 130 degrees of active scapular plane abduction as measured goniometrically during clinical examination</li> </ul>
	Exclusion Criteria (not listed above)
	History of rotator cuff surgery
	<ul> <li>History of glenohumeral dislocation or other traumatic injury to the shoulder</li> </ul>



Ludewig 2003 (Continued)	<ul> <li>Only periscapular or cervical pain during arm elevation</li> <li>Shoulder symptoms reproduced by a cervical assessment</li> </ul>			
	Baseline characteristics			
	Intervention			
	Number randomised: 34; mean (SD) age: 48 (1.8) years; sex: men only; duration of symptoms: not re- ported			
	Control			
	Number randomised: 3 not reported	3; mean (SD) age: mean: 49.2 (1.8) years; sex: men only; duration of symptoms:		
Interventions	Intervention: home exercise programme			
<i>Components of intervention</i> : two stretches (pectoralis minor stretch and posterior shoulder st muscle relaxation exercise for the upper trapezius performed in front of a mirror, and progres sistance strengthening exercises for two muscle groups (serratus anterior muscle and humer nal rotation). Participants received instructions from a licensed therapist. Each subject receive ten/pictorial instructions for home reference and a daily exercise log to monitor compliance				
	daily); progressive resis to 3 sets of 15 repetitio	onds each repetition and 5 repetitions daily); muscle relaxation exercise (5 times stance strengthening exercises (3 sets of 10 repetitions the first week, progress ns the second week and three sets of 20 days the third week, after which, partici- increasing weight resistance and repeat the repetition sequence)		
	Frequency of administre	ation: daily for 10 weeks		
	Control: no intervention			
Outcomes	Outcomes assessed at	an average of 10 weeks (between 8-12 weeks)		
		Function measured by the Shoulder Rating Questionnaire (SRQ); score: 17–100, higher score indicat- ing greater shoulder function and fewer shoulder symptoms		
	<ul> <li>Pain: questionnaire answers compiled to produce a work-related pain score, ranging from 1-10, with higher scores indicating increased pain with work (derived from SPADI score)</li> </ul>			
	-	e ranging from 1-10, with higher scores indicating greater difficulty with work per-		
Notes	Conflicts of interest: r	not reported		
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence genera- tion (selection bias)	Low risk	Quote: "Randomisation was performed by an investigator blindly selecting one of two slips of paper indicating group assignment."		
		Comment: An adequate method was used to generate the allocation sequence		
Allocation concealment (selection bias)	Unclear risk	Comment: It is unclear if adequate safeguards were put in place to conceal the allocation sequence		

Quote: "Researchers were not blinded to group assignment, but were to base-

Comment: Given the nature of the interventions, participants were not blind to

treatment, and may have had different expectations about the benefits of each

line measurements at the time of follow up."

intervention

Manual therapy and exercise for rotator cuff disease (Review)

Blinding of participants

and personnel (perfor-

mance bias)

All outcomes

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High risk

Ludewig 2003 (Continued)		
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported all outcomes
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Ninety two per cent of subjects completed the study. Seven subjects were lost to follow up, four (11.8 %) in the exercise intervention group, and three control subjects (one symptomatic (3%) and two asymptomatic (8%)). One intervention subject withdrew after experiencing a new injury at work that interfered with continuation of the exercises. Another intervention subject was referred by his physician for additional outpatient physical therapy and subsequently withdrew from the study. A third intervention and one symptomatic control subject were not able to return for follow up for personal reasons (death in the family, custody dispute). The remaining three subjects either were no shows or were unable to be reached after multiple attempts at the time of post-test. Subjects lost to follow up were similar to the full sample with regard to demographic characteristics."
		Quote: "The initial analysis included all subjects from whom post-test da- ta were obtained, regardless of their level of compliance with the exercise programme. A secondary complete "intention to treat" analysis was also performed where all subjects initially enrolled were analysed. Missing post- test data were replaced with imputed values based on the average observed means from the two symptomatic groups."
		Comment: The amount and reasons for attrition are unlikely to have affected the results
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other source of bias were identified

Maen	hout	2013
mach		2020

Methods	Study design: Parallel group RCT			
	Setting: Shoulder surgery clinic, Belgium			
	Intervention: Heavy load eccentric training plus traditional rotator cuff training			
	Control: Traditional rotator cuff training			
	Source of funding: Not reported			
Participants	Diagnostic label used by trialists: Impingement			
	Criteria for defining the shoulder condition being treated			
	<ul> <li>Unilateral pain for at least 3 months in the anterolateral region of the shoulder</li> <li>Painful arc</li> <li>2 out 3 impingement tests positive</li> <li>2 out of 4 resistance tests painful</li> </ul>			
	Pain with palpation of the supraspinatus and/or infraspinatus tendon insertion			
	Inclusion Criteria (not listed above)			
	Age over 18 years old			



Maenhout 2013 (Continued)

## **Exclusion Criteria (not listed above)**

- Demonstration of partial or full ruptures of the rotator cuff by technician investigation
- History of shoulder surgery
- Shoulder fracture or dislocation
- Traumatic onset of pain
- Osteoarthritis
- Frozen shoulder
- Traumatic glenohumeral instability or shoulder nerve injuries
- Concomitant disorders (such as cervical pathology or systemic musculoskeletal disease)

### **Baseline characteristics**

### Intervention

Number randomised: 31; mean (SD) age: 40.2 (12.9) years; sex: F/M 16/15; duration of symptoms: not reported

Control

Number randomised: 30; mean (SD) age: 39.4 (13.1) years; sex: F/M 20/10; duration of symptoms: not reported

Interventions

# Intervention: heavy load eccentric training

*Components of intervention*: eccentric exercise consisted of full can (thumb up) abduction in the scapular plane, which was performed with a dumbbell weight. Participants were asked to perform the eccentric phase at a speed of 5 min/repetition. Starting position of the eccentric phase at full scapular abduction had to be pain free, and, if not, participants were advised to stretch out the arm at a slightly lower degree of scapular abduction

*Dose*: based on pain monitoring model. Whenever the pain was no longer present during the last set of repetitions, dumbbell weight was increased with 0.5 kg

Frequency of administration: 3 sets of 15 repetitions performed twice a day, at home for 12 weeks

### Control: traditional rotator cuff training

Participants only received the treatment common to both groups (see below)

### **Both groups**

*Components of intervention:* performed two traditional rotator cuff strengthening exercises at home that involved internal and external rotation resisted with an elastic band (Thera-Band). Participants were instructed to perform the exercises at a speed of 6 min'/repetition (2 min concentric phase, 2 min isometric phase and 2 min eccentric phase). The colour of the band was chosen so that the participant did not experience significantly more pain during the exercise than at rest. Load was increased by changing colour of the elastic band as soon as pain decreased. In addition, physiotherapy treatment sessions aimed at correcting performance of the exercises, increasing load and emphasising the importance of adherence to the home exercises were delivered. Treatments included glenohumeral mobilisation, scapulothoracic mobilisation, scapula setting and posture correction

*Frequency of administration*: exercises - 1 per day for 3 sets of 10 repetitions, performed at home for 12 weeks. Nine physiotherapy treatments delivered over 12 weeks

 Outcomes
 Outcomes assessed at 6 and 12 weeks

 • Function: SPADI questionnaire scaled 0-100 where a higher score indicates more pain and disability

 • Isometric strength measured with hand held dynamometer at 0, 45 and 90 degrees abduction, also measured at external and internal rotation (degrees)

 • Global assessment of treatment success: participant-rated improvement on a 5-point scale: "No change" equalled a score of 0, "better" was scored between 1 and 5 and "worse" between -1 and -5

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### Maenhout 2013 (Continued)

# • Adverse events (collected but not reported)

# Notes

Conflicts of interest: not reported

Trial was registered in ClinicalTrials.gov (NTC00782522)

**Risk of bias** 

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "Prior to the intervention, baseline outcome measurements were per- formed. Subsequently, patients were randomly allocated to the traditional ro- tator cuff strength training (TT) group or the TT combined with heavy load ec- centric training (TT + ET) group."
		Comment: No information on the method used to generate the allocation se- quence was reported (in either the registry entry or publication)
Allocation concealment (selection bias)	Unclear risk	Comment: No information on the method used to conceal the allocation se- quence was reported (in either the registry entry or publication)
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported some out- comes (e.g. pain, function, treatment success)
Blinding of outcome as- sessment (detection bias) Objective outcomes	High risk	Quote: "All tests were completed at the laboratory of the Department of Re- habilitation Science and Physiotherapy of Ghent University. This investigator could not be blinded to treatment group."
		Comment: The assessor of objective outcomes was not blinded and may have assessed outcomes for each group differently based on prior expectations of the benefits of each intervention
Incomplete outcome data (attrition bias)	High risk	Quote: "Intention to treat principle was respected, and all patients were in- cluded in analysis as randomized."
All outcomes		Comment: Trialists report that an intention-to-treat analysis was performed, but data for participant-reported treatment success were reported based on per-protocol analysis (and it is unclear what sample size other outcomes were based on in the analysis). A CONSORT flow chart (Schultz 2010) shows that more participants in the control group dropped out at week 12 due to "No im- provement". This attrition is likely to underestimate the difference between groups in participant-reported treatment success (i.e. bias in favour of the con- trol group). Other outcomes may also be affected if not based on an inten- tion-to-treat analysis
Selective reporting (re- porting bias)	High risk	Comment: Some outcomes specified in the registry entry (NTC00782522) were not reported in the publication (ROM, force reproduction, subacromial space). Also, patient-reported treatment success was not pre-specified in the registry entry
Other bias	Low risk	Comment: No other sources of bias identified

Manual therapy and exercise for rotator cuff disease (Review)

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Martins 2012	
Methods	Study design: Randomised, prospective and comparative trial
	Setting: Rehabilitation Centre, Brazil
	Intervention: Proprioception exercises plus stretching and strengthening exercises plus cryotherapy
	Control: Stretching and strengthening exercises plus cryotherapy
	Source of funding: Not reported
Participants	Diagnostic label used by trialists: Impingement
	Criteria for defining the shoulder condition being treated
	Diagnosis of rotator cuff disorder (impingement syndrome)
	Inclusion Criteria (not listed above)
	<ul> <li>Employment as a registered nurse, nurse technician, or nurse's aide at the Institution</li> <li>Availability and interest in participating in the study</li> </ul>
	Exclusion Criteria (not listed above)
	<ul> <li>Significant pain that would prevent performance of the physical therapy programme</li> <li>Medically diagnosed cognitive alterations</li> <li>Associated disability conditions</li> <li>Previous shoulder surgery or other shoulder complex disorders (adhesive capsulitis, degenerative alterations of the glenohumeral joint, tendinous calcification)</li> <li>Absence from more than 3 physical therapy sessions</li> <li>Use of medication or other treatment for shoulder pain during the physical therapy treatment</li> <li>Unwillingness to take part in the study</li> </ul> Baseline characteristics Intervention Number randomised: 9: age: 30 < 40 years (n = 0): 41 < 50 years (n = 5): > 50 years (n = 3): sey: F/M 7/1:
	Number randomised: 9; age: 30 ≤ 40 years (n = 0); 41 ≤ 50 years (n = 5); > 50 years (n = 3); sex: F/M 7/1; duration of symptoms: not reported <i>Control</i>
	Number randomised: 9; age: 30 ≤ 40 years (n = 2); 41 ≤ 50 years (n = 2); > 50 years (n = 4); sex: F/M 7/1; duration of symptoms: not reported
Interventions	Intervention: proprioception exercises
	<i>Components of intervention:</i> exercises with joint position, rhythmic stabilisation and repositioning of the members, unstable base, proprioceptive neuromuscular facilitation, and speed and accuracy
	Dose: resistance during strength exercises was increased every 3 sessions
	Frequency of administration: 2 sessions per week for 6 weeks
	Control: stretching and strengthening exercises plus cryotherapy
	<i>Components of intervention</i> : participants only received the treatment common to both groups (see be- low)
	Both groups
	<i>Components of intervention</i> : Codman's pendulum exercises of the shoulder, stretching of the cervical spine and shoulder muscles, exercises with a stick (to maintain or improve ROM), exercises to strength-



Martins 2012 (Continued)	en the muscles of the rotator cuff and scapular stabilisers, cryotherapy (ice pack for 20 min, performed at the end of the treatment session), and education regarding joint protection and posture <i>Frequency of administration</i> : 2 sessions per week for 6 weeks
Outcomes	<ul> <li>Outcomes assessed at 6 weeks</li> <li>Pain (Visual Numeric Scale ranging from "no pain" to "worst possible pain"; score range not reported)</li> <li>Quality of life measured using the Western Ontario Rotator Cuff Index (WORC). The total score ranges from 0-2100. Thus, 0 implies no reduction in quality of life and the worst score is 2100</li> <li>Work disability measured using the Occupational Stress Indicator, which allows the assessment of occupational satisfaction according to 22 psychosocial aspects by means of 6-point Likert scales, which vary from enormous dissatisfaction to enormous satisfaction. The sum of these measures provides an indicator of job satisfaction given by a global score that ranges from 22-123 points</li> </ul>
Notes	Conflicts of interest: not reported

Trial registration ClinicalTrials.gov NCT01465932

# Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "All subjects were then randomized and allocated to Group 1 (control) or Group 2 (experimental) by a person not participating in the application of data collection instruments and/or rehabilitation programs." Quote: "In order to promote greater homogeneity with the clinical status of the patients, subjects were initially subdivided according to the presence or absence of shoulder movement deficits (measurement of shoulder ROM with a goniometer) and the level of pain intensity, as shown by the Visual Numer- ic Scale (VNS)The randomization occurred so that the first subject of each subgroup was randomly assigned to one group and the second subject was as- signed to the other group, and so on." Comment: The specific method used to generate the allocation sequence is unclear
Allocation concealment (selection bias)	Unclear risk	Quote: "All subjects were then randomized and allocated to Group 1 (control) or Group 2 (experimental) by a person not participating in the application of data collection instruments and/or rehabilitation programs."
		Comment: The specific method used to conceal the allocation sequence was not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Comment: Participants received different multimodal interventions, but it is unclear whether they were provided with any information that would make them perceive the intervention they received as superior or inferior to the al- ternative intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	Unclear risk	Comment: Participants self-reported all outcomes, but it is unclear whether they were provided with any information that would make them perceive the intervention they received as superior or inferior to the alternative interven- tion
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: 9 participants were randomly allocated to each group, and one dropped out of each group, both for the same reason (lack of commitment to rehabilitation programme). This is unlikely to have biased the results
Selective reporting (re- porting bias)	High risk	Comment: Mean (SD) data for pain were reported for each group by subgroup only (mild, moderate, severe). No measures of variation were reported for the

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# Martins 2012 (Continued)

		health-related quality-of-life measure. Pain had not been pre-specified in the clinical trials registry entry
Other bias	Low risk	Comment: No other sources of bias identified

Methods	Study design: Parallel group RCT
	Setting: Outpatient clinic, Italy
	Intervention: Neurocognitive therapeutic exercise
	Control: Traditional therapeutic exercise
	Source of Funding: Not reported
Participants	Diagnostic label used by trialists: Shoulder impingement syndrome
	Criteria for defining the shoulder condition being treated
	<ul> <li>Diagnosis of Neer stage I shoulder impingement syndrome, degenerative rotator cuff tendinopath without tendon tears and/or subacromial bursitis, determined using four isometric tests (abduction at 0-30 degrees, external or internal rotation, positive Kennedy-Hawkins sign and positive Neer sign) X-ray on anteroposterior, axillary and outlet views, and MRI or echography of the affected shoulder</li> </ul>
	Inclusion Criteria (not listed above)
	<ul><li>Aged 18 years or older</li><li>Shoulder pain lasting for at least 3 months</li></ul>
	Exclusion Criteria (not listed above)
	<ul> <li>Inability or unwillingness to sign informed consent</li> <li>Rotator cuff and/or subscapularis tendon partial/full thickness tears</li> <li>Capsulolabral pathology responsive to surgical repair</li> <li>Congenital abnormalities of the acromion</li> <li>Previous surgery on the affected shoulder</li> <li>Inflammatory or neurological (systemic or local) diseases involving shoulder girdles</li> <li>Cognitive or psychiatric disorders</li> <li>Local tumour metastasis or application of radiotherapy</li> <li>Acute infections or osseous tuberculosis</li> </ul>
	Baseline characteristics
	<i>Intervention</i> Number randomised: 24; mean age: 61.6 ± 11.2 years old; sex: F/M 12/12; duration of symptoms: not re ported
	<i>Control</i> Number randomised: 24; mean age: 62.6 ± 13.9 years old; sex: F/M 15/9; duration of symptoms: not re- ported
Interventions	Intervention: neurocognitive therapeutic exercise (NCTE)
	<i>Components of intervention:</i> the neurocognitive protocol contained 10 exercises. The first 3 aimed at restoring shoulder fragmentation and counterbalance; the second set consisted of 4 exercises aimed a centring the humeral head in the glenoid fossa during active movements and introducing counterbal- ancing mechanism of the scapula during upper limb movements; the last 3 exercises aimed at recov-

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Marzetti 2014 (Continued)							
	(e.g. inclined table with higher cortical function	of movement of the affected shoulder. Exercises involving specific instruments h a board with 5 concentric circles) were taught to promote the stimulation of ns useful to select the most important proprioceptive information necessary to haviour and recover fine motor skills. The execution of the exercises was facilitat gery					
	Dose: duration 1 hour Frequency of administration: 3 times a week for 5 weeks Control: traditional therapeutic exercise (TTE)						
					<i>Components of intervention:</i> the traditional therapeutic exercise protocol contained mainly strength- ening exercises focused on the rotator cuff and scapular stabilising muscles, stretching exercises, Cod- man's pendulum exercises and exercises against elastic band resistance <i>Dose:</i> duration 1 hour		
	Outcomes	Outcomes assessed at 5, 12 and 24 weeks					
		<ul> <li>Function: Constant-Murley total score (0-100, higher = best result)</li> <li>Rest pain: VAS, 0 (no pain) to 10 (most severe pain)</li> <li>Pain on motion: VAS, 0 (no pain) to 10 (most severe pain)</li> <li>Adverse events</li> </ul>					
	Notes	Conflicts of interest: not reported					
	Trial registered in ClinicalTrials.gov (NCT01785745)						
Risk of bias							
Bias	Authors' judgement	Support for judgement					
Random sequence genera- tion (selection bias)	Low risk	Quote: "Patients were randomly assigned to either NCTE (group 1) or TTE (group 2) using a random sequence generator (www.random.org)"					
		Comment: An adequate method was used to generate the allocation sequence					
Allocation concealment (selection bias)	Low risk	Quote: "Allocation sequence was performed using closed envelopes, and the assignment code of each patient revealed to the researcher who performed the treatment only at the beginning of the therapeutic protocol".					
		Comment: An adequate method was used to conceal the allocation sequence					
Blinding of participants	Unclear risk	Quote: "Single-blind randomized, non-inferiority trial."					
and personnel (perfor- mance bias) All outcomes		Quote: "Outcome measures were determined by an assessor blinded to pa- tient allocation."					
		Comment: It can be inferred from the above quotes that the trialists consid-					

Comment: It can be inferred from the above quotes that the trialists considered participants to be unblinded. However, it is unclear whether participants were provided with any information that would make them perceive the type of exercise they received as superior or inferior to the alternative type of exercise

Blinding of outcome as-<br/>sessment (detection bias)Unclear riskComment: Participants self-reported some outcomes, but it is unclear whether<br/>they were provided with any information that would make them perceive the<br/>type of exercise they received as superior or inferior to the alternative type of<br/>exercise

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Marzetti 2014 (Continued)		
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "Outcome measures were determined by an assessor blinded to pa- tient allocation."
		Quote: "As a check on blindness, the assessor was asked to guess treatment allocation after the final outcome assessments were completed. The analysis of these guesses showed a correctness of approximately 30%, which is consid- ered not better than chance"
		Comment: Assessor of objective outcomes was blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Missing data at follow-up were managed by the Last Observation Car- ried Forward (LOCF) method. Analyses were performed according to the inten- tion-to-treat principle"
		Quote: "All of the participants completed the treatment protocol. Two par- ticipants in the NCTE group and three in the TTE group did not attend the fol- low-up visit at T2. Two participants in the TTE group did not attend the fol- low-up visit at T3."
		Quote: "reasons for lack of follow-up were not recorded. However, only a few participants were lost at follow-up (10.4%) and dropouts occurred to a similar extent in the two treatment groups, which did not substantially affect the results"
		Comment: No reasons for loss to follow-up were reported, but the amount is small and relatively balanced between groups, so is unlikely to have biased the results. All randomised participants were included in the analyses
Selective reporting (re- porting bias)	Low risk	Comment: Outcome data fully reported for all outcomes specified in the Clini- calTrials.gov registry entry
Other bias	Low risk	Comment: No other sources of bias identified

McClatchie 2009			
Methods	Study design: Cross-over RCT		
	Setting: Private orthopaedic practice, Canada		
	Intervention: Lateral cervical glide mobilisations		
	Control: Placebo mobilisation		
	<b>Source of funding:</b> Financial support provided by the Graduate Department of Rehabilitation Science, University of Toronto, Toronto, Canada		
Participants	Diagnostic label used by trialists: Painful arc		
	Criteria for defining the shoulder condition being treated		
	Generalised unilateral shoulder pain		
	Insidious onset of pain		
	Painful arc on shoulder abduction		
	Inclusion Criteria (not listed above)		
	Aged over 18		

McClatchie 2009 (Continued)	pain through "tradi	e been unresponsive to 2–4 recent physiotherapy sessions addressing shoulder tional" methods of movement patterns, strengthening and modalities such as ul-	
	trasound and cryotherapy Exclusion Criteria (not listed above)		
	<ul> <li>Symptoms of paraesthesia or neurological deficits</li> <li>Previous surgery or dislocation of the affected shoulder</li> </ul>		
	Clinically definitive	arthritis of the shoulder on X-ray	
	Had a cortisone inje	ection for the current episode of shoulder pain	
	Baseline characteristics		
	Overall cohort of partic	ipants	
		21 (7 mobilisation group, 14 placebo group in first period); mean (SD) age: 49.8 1 14/7; duration of symptoms: not reported	
Interventions	Intervention: lateral o	cervical glide mobilisations	
	the chair, head in a neu lap. The lateral aspect side of the participant' spinous process of C5,	ntion: participant was seated with the thoracic spine resting against the back of utral position, feet resting flat on the floor, and arms relaxed with hands in their of the spinous processes of C5, C6, and C7 was landmarked on the ipsilateral s painful shoulder. The examiner's thumb remained on the lateral aspect of the with the opposite hand placed on the participant's non-affected shoulder or ce as a lateral movement toward the non-painful side was applied with the mo-	
	Dose: 2 min each at C5, C6 and C7, with small amplitude end range movements (Grade IVþ)		
	Frequency of administration: once, within 4 days of the cross-over treatment		
	Control: placebo mobilisation:		
	<i>Components of intervention</i> : involved the examiner resting their hands in the same positions as the mo- bilisation technique, but without the application of external force		
	Dose: hands held at C5, C6 and C7 for 2 min each		
	Frequency of administration: once, within 4 days of the cross-over treatment		
Outcomes	Outcomes assessed between 1 and 4 days		
	<ul> <li>Pain: VAS score ranging from 0–10 cm with a higher score indicating worse pain</li> <li>Strength (abduction) measured in kilogram-force using manual muscle testing</li> </ul>		
	Note that active cervical spine ROM was also measured but was not extracted as we were only interest- ed in shoulder ROM		
Notes	Conflicts of interest: not reported		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera-	Low risk	Quote: "Subjects were randomized by a coin toss"	
tion (selection bias)		Comment: An adequate method was used to generate the allocation sequence	
Allocation concealment	Unclear risk Comment: No information on how the allocation sequence was concealed wa reported		

Manual therapy and exercise for rotator cuff disease (Review)

McClatchie 2009 (Conti	nued)
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Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Each subject recognized that the mobilization and placebo interven- tions were different from each other, however, no subject realized that the placebo intervention was not therapeutic." Comment: Participants were likely blind to the intervention they received at each session
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	Low risk	Comment: Blinded participants self-reported some outcomes
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "All outcome measures were assessed both before and after the inter- vention and conducted by the first investigator, who was blinded to which treatment intervention was received." Comment: Assessors of objective outcomes were likely blind to treatment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: There was no loss to follow-up. and all randomised participants were analysed
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias were identified

Moosmayer 2014			
Methods	Study design: Parallel group RCT		
	Setting: Recruited by general practitioners and referred to orthopaedic surgeon		
	Intervention: Physiotherapy comprising exercises only		
	Control: Surgery		
	Source of funding: Not reported		
Participants	Diagnostic label used by trialists: Full rotator cuff tear		
	Criteria for defining the shoulder condition being treated		
	<ul> <li>Symptomatic small (&lt; 1 cm) or medium-sized (1–3cm) tears of the rotator cuff. Demonstration of a full thickness tear by sonography or MRI, a tear size not more than 3 cm on short and long axis ultrasound scans and muscle atrophy on MRI not exceeding Stage 2</li> </ul>		
	Inclusion Criteria (not listed above)		
	<ul> <li>Lateral shoulder pain at rest or with exercise</li> <li>Painful arc</li> <li>Positive impingement signs</li> <li>Passive range of movement of at least 140 degrees for abduction and flexion</li> <li>Traumatic and atraumatic tears included</li> </ul>		
	Exclusion Criteria (not listed above):		
	• Age < 18 years		



Moosmayer 2014 (Continued)

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Continuea)	<ul> <li>Tears with an absolute indication for surgery such as those involving substantial parts of the supraspinatus tendon</li> <li>The presence of other local or systemic diseases affecting shoulder function</li> <li>Previous tendon surgery on the relevant shoulder</li> <li>Medical comorbidities</li> <li>Inability to comply with follow-up</li> </ul>
	Baseline characteristics
	Intervention
	Number randomised: 51; mean (range) age: 61 (46-75) years; sex: F/M 15/36; mean (SD) duration of symptoms: 9.8 (9.8) months
	Control
	Number randomised: 52; mean (range) age: 59 (44-75) years; sex: F/M 15/37; mean (SD) duration of symptoms: 12.3 (18.7) months
Interventions	Intervention: physiotherapy
	<i>Components of intervention</i> : supervised exercises only, with particular attention directed towards correction of upper quarter posture and restoration of scapulothoracic and glenohumeral muscular control and stability. Local glenohumeral control was addressed by exercises to centre the humeral head in the glenoid fossa. Isometric exercises and exercises against eccentric and concentric resistance for shoulder rotators were given. When local glenohumeral control was achieved, exercises were given with increasing loads and progressed from neutral to more challenging positions. During all exercises, scapular stability had to be maintained. Additional exercises were given for specific demands in work, sports and leisure activities. Twelve of 51 participants initially randomised to physiotherapy underwent tendon repair surgery during the 5-year follow-up, but were analysed in the group to which they were allocated.
	Dose: 40 min
	<i>Frequency of administration</i> : 20 sessions given on average twice weekly for 12 weeks and with increas- ing intervals during the following 6–12 weeks
	Control: surgery
	<i>Components of intervention</i> : tendon repair surgery was performed in a standard manner by mini-open (9 participants) or open (42 participants) tendon repair. All were performed in the deck-chair position under interscalene block regional anaesthesia and total intravenous anaesthesia without the use of inhalation agents, by one of three experienced orthopaedic surgeons. Following diagnostic arthroscopy and through a deltoid splitting approach, an anteroinferior acromioplasty was performed. With the arm at the side, the rotator cuff was mobilised until the tear was fully exposed. The footprint was prepared to bleeding bone and tendon repair performed with a combination of tendon-to-tendon and tendon-to-bone techniques by passing sutures through bone tunnels in the greater tuberosity. The deltoid was repaired to the acromion through drill holes. Tenodesis of the long head of biceps tendon was performed in 18 participants in whom arthroscopy had shown inflammation of a partial tear. Mini-open tendon repair differed from open repair by a shorter incision and arthroscopic acromioplasty. Post-operatively the arm was mobilised in a sling and passive range-of-movement exercises commenced. Active-assisted movements where initiated after 6 weeks, and supplemented by strengthening exercises 12 weeks after surgery
Outcomes	Outcomes assessed at 6 and 12 months, 2 years and 5 years
	<ul> <li>Function: Constant score scaled 0-100 with a higher score indicating less disability</li> <li>Active ROM: flexion and abduction measured in degrees using the Constant subscore</li> <li>Strength: Constant subscore shoulder strength (kg)</li> <li>Pain measured using the American Society of Shoulder and Elbow Surgeons (ASES) pain subscore (VAS 0-10 where a higher score indicates worse pain)</li> </ul>

# Moosmayer 2014 (Continued)

 Quality of life: Short Form 36 Health survey, score: 0–100; 100 indicating best possible health conditions (physical component score and mental component summary score)

Notes

**Conflicts of interest:** the authors state "Although none of the authors has received or will receive benefits for personal or professional use from a commercial party related directly or indirectly to the subject of this article, benefits have been or will be received but will be directed solely to a research fund, foundation, educational institution, or other nonprofit organisation with which one or more of the authors are associated."

#### **Risk of bias**

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "A computer-generated randomisation list (block length 20, ratio 1:1) was drawn up by our statistician."
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Low risk	Quote: "Sequentially numbered, sealed envelopes were used to assign treat- ment according to the participants' study number, given at baseline assess- ment. The randomisation sequence was concealed from the study's collabora- tors until treatment was assigned."
		Comment: An adequate method was used to conceal the allocation sequence
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported some out-comes
Blinding of outcome as-	Low risk	Quote: "Only the outcome assessor remained blinded throughout the study."
sessment (detection bias) Objective outcomes		Comment: Assessor of objective outcomes was likely blind to treatment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "In order to avoid interpretation bias and loss of power it was decid- ed to perform all outcome analyses 'as randomised' by following an inten- tion-to-treat principle. As a consequence, results from patients who withdrew (one from the surgery group) and those who changed treatment (nine from the physiotherapy group) were assessed as originally randomised. For the patient who withdrew from the surgery group, the baseline data carried forward gave him a result far below the mean of his group. The nine patients who changed treatment after failed physiotherapy had all performed at least 15 treatment sessions, and it seemed adequate to interpret their final score pre-operative- ly as the best estimate for the final result from physiotherapy. Following these patients without further treatment would have been unethical, and elimina- tion of their results from analysis would have led to an overestimation of the effect of physiotherapy."
		Quote: "Twelve nonoperative patients reported an insufficient treatment ef- fect. This subgroup had a mean increase of the Constant score of 1.8 points (range, 220 to 22 points) and underwent secondary surgery within the first two years (three patients during the first six months, six patients during the follow- ing six months, and three patients during the second year)."

Quote: "The five-year follow-up rate was 98%."



Moosmayer 2014 (Continued)		Comment: An appropriate method was used to deal with participants who did not remain in their allocated group. Twelve of 51 participants initially randomised to physiotherapy underwent tendon repair surgery, but were analysed in the group to which they were allocated.
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias were identified

# Munday 2007

Methods	Study design: Parallel group RCT			
	<b>Setting:</b> General population, South Africa (participants recruited through local advertising, flyers placed in general practitioners' and chiropractors' offices, and around universities, colleges, schools and businesses)			
	Intervention: Shoulder girdle adjustments (chiropractic)			
	Control: Detuned ultrasound (placebo)			
	Source of funding: Masters in Technology: Chiropractic (Durban University of Technology)			
Participants	Diagnostic label used by trialists: Impingement			
	Criteria for defining the shoulder condition being treated			
	<ul> <li>Shoulder impingement syndrome (SIS) with criteria of history of shoulder pain &gt; 6 weeks (stages 1-3) plus three of the following:         <ul> <li>palpable tenderness at the greater tuberosity;</li> <li>palpable tenderness at the anterior acromion;</li> <li>a painful arc of abduction between 60 and 120 degrees;</li> <li>positive impingement sign</li> </ul> </li> <li>SIS diagnosis was confirmed by a single, specially trained, on-duty clinician (a doctor of chiropractic)</li> <li>Inclusion Criteria (not listed above)</li> <ul> <li>&lt; 40 years of age (targeting SIS stage 1 or 2)</li> <li>No local or systemic pathology</li> <li>No shoulder treatment in the last 6 weeks</li> </ul> <li>Exclusion Criteria (not listed above)</li> </ul>			
	History of traumatic shoulder dislocation			
	Frequent, severe crepitus			
	<ul> <li>Weakness of internal rotation and abduction to resistance</li> </ul>			
	Pain radiating distally below the elbow			
	Shoulder surgery in the previous 2 years			
	Baseline characteristics			
	Overall			
	Sex: F/M 14/16			
	Intervention			

Munday 2007 (Continued)	Number randomised: 1 not reported	15; mean (range) age: 22 (16-38) years; sex: not reported; duration of symptoms:	
	Control		
	Number randomised: 1 not reported	15; mean (range) age: 23 (19-32) years; sex: not reported; duration of symptoms:	
Interventions	Intervention: should	er girdle adjustments	
	end feel or joint play w exposed. Adjustments	<i>ntion</i> : high velocity, low-amplitude manipulation in the direction of restricted ras performed. Participants sat in a comfortable position with the shoulder girdle to the acromioclavicular joint were most common, although adjustments to the ohumeral joints were made as well. The spine was not adjusted in this trial	
	Frequency of administration: 8 sessions over 3 weeks		
	Control: placebo (det	uned ultrasound)	
		<i>ntion</i> : participants were asked to sit in a comfortable position with the shoulder ceived detuned ultrasound	
	Dose: no frequency or	times. Duration 6 min	
	Frequency of administration: 8 sessions over 3 weeks		
Outcomes	Outcomes assessed at 3 and 7 weeks		
	<ul><li>Pain measured on a</li><li>ROM measured usir</li><li>Adverse events</li></ul>	n visual analogue scale ng a goniometer (unclear if active or passive, and no outcome data reported)	
Notes	Conflicts of interest: not reported		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Quote: "Randomisation was accomplished by utilising 30 folded sheets of paper (15 marked Group A, 15 marked Group B), thoroughly mixed together to assure discontinuity and then placed in a hat. At each subject randomisa- tion time point, the hat was held so that all folded slips were completely ob- scured."	
		Comment: An adequate method was used to generate the allocation sequence	
Allocation concealment (selection bias)	Low risk	Quote: "At each subject randomised time point, the hat was held so that all folded slips were completely obscured."	
		Comment: An adequate method was used to conceal the allocation sequence	
Blinding of participants and personnel (perfor- mance bias)	Low risk	Quote: "Patients were made aware that they might be randomized into either group (treatment or placebo) A or B. At the end of the trial, those in the placebo group were offered up to 8 free treatments".	
All outcomes		Comment: Patients were likely blinded to treatment (i.e. unaware that the ul- trasound machine was not switched on)	
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	Low risk	Comment: Blinded participants self-reported some outcomes	

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# Munday 2007 (Continued)

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Blinding of outcome as- sessment (detection bias) Objective outcomes	Unclear risk	Comment: There was no information about whether assessors of objective outcomes were blinded
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Quote: "Four patients were excluded due to either non-compliance with treat- ment protocols or exclusion factors. Additional patients were recruited and randomised to achieve a total of 30 patients (out of N=34) completing the trial. Intention to treat analysis was used."
		Comment: The flow of participants throughout the trial is unclear
Selective reporting (re- porting bias)	High risk	Comment: No outcome data were reported for ROM, which was specified as an outcome in the methods section
Other bias	Low risk	Comment: No other sources of bias were identified

Methods	Study design: Parallel group RCT		
	Setting: Primary care and orthopaedic surgeon clinics		
	Intervention: High dose medical exercise therapy		
	<b>Control:</b> Low dose medical exercise therapy		
	Source of funding: No funding		
Participants	Diagnostic label used by trialists: Impingement		
	Criteria for defining the shoulder condition being treated		
	<ul> <li>Unilateral primary shoulder impingement defined by positive subacromial impingement test (stan dardised procedure so that the interpretation was consistent from physician to physician; humeru abducted 90 degrees in the scapulae plan, maximal passive inward rotation should give subacromia pain); and elbow and thoracic spine function with no referred pain from this area to the shoulder</li> </ul>		
	Inclusion Criteria (not listed above)		
	<ul> <li>No previous shoulder surgery</li> <li>Normal neck (no neurological signs)</li> <li>No neurological diseases</li> <li>No history of shoulder dislocation, subluxation or fracture</li> <li>No vestibular or visual disturbances</li> <li>No chiropractic or physiotherapy treatment within the last 6 months prior to entering the study</li> </ul>		
	Exclusion Criteria (not listed above)		
	<ul> <li>Individuals with any cardiovascular, respiratory, systemic or metabolic condition limiting their abilit to participate in the study</li> <li>People showing signs of rotator cuff tears (drop arm test) or other disorders in the glenohumeral joir during the physical examination</li> </ul>		
	Baseline characteristics		
	Intervention		
	Number randomised: 31; mean (SD) age: 46.1 ± 11.2 years; sex: F/M: 66.9%/33.1%; mean (SD) duration of symptoms: 3.6 ± 5.1 years		

Osteras 2008 (Continued)	Control		
		80; mean (SD) age: 41.8 ± 14.5 years; sex: F/M 74%/26%; mean (SD) duration of ars	
Interventions	Intervention: high do	se exercise	
	<i>Components of intervention</i> : supervised high dose progressive resistance exercise therapy, comprising global aerobic exercises using a stationary bike, a treadmill, or a step machine, and semiglobal and local exercises using such medical exercise therapy equipment as wall pulley apparatus, lateral pulley apparatus, inclines board, angle bench, multiple purpose bench, shoulder rotator, dumbbells or barbells. The participants' exercise programme was graded in such a way that it was performed pain free, or close to pain free (a maximum of 3 on a 10-point VAS)		
	and local exercises par exercise programme (4 ter the last 4 exercises,	ormed eight exercises, each of 3 sets of 30 repetitions. Prior to the semiglobal ticipants warmed up for 15–20 min on an ergometer cycle. Half way through the exercises each of 3 sets of 30 repetitions) the participants cycled for 10 min. Afthe participants did another 10 min' stationary ergometer cycling. The intensies was moderate to high (i.e. a heart rate frequency of 70%–80% of the maximal	
	Frequency of administr	ation: 3 treatments a week for 12 weeks	
	Control: low dose exe	rcise	
	<i>Components of intervention</i> : same components as the high dose exercise group, but at a lower dose (see below)		
	<i>Dose</i> : participants started each treatment with 5–10 min on an ergometer cycle and then performed 5 semiglobal and local exercises using medical exercise therapy equipment performing 2 sets of 10 repetitions of each exercise. The intensity during the cycle exercises was moderate to high (i.e. a heart rate frequency of 70%–80% of the maximal heart rate). The crucial difference between the groups were time on the bike (35 min in the high dose group compared to 10 min in the low dose group), number of exercises (8 compared to 5), and number of repetitions (3 times 30 compared to 2 times 10 per exercise)		
	Frequency of administration: 3 treatments a week for 12 weeks		
Outcomes	Outcomes assessed at 3, 9 and 15 months		
	<ul> <li>Function: Shoulder rating Questionnaire (SRQ) scored 17-90 with higher score indicating better func- tion and fewer symptoms</li> </ul>		
	<ul> <li>Pain: VAS score 0-10 with higher score indication worse pain (cm)</li> <li>ROM (abduction, flexion) (unclear if active or passive)</li> </ul>		
	<ul> <li>Kow (abduction, nexton) (unclear in active of passive)</li> <li>Strength: isometric strength of abduction, flexion, external and internal rotation (recorded in Newtons)</li> </ul>		
Notes	Conflicts of interest: t	he authors stated that they had no conflicts of interest	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Quote: "The randomization procedure was concealed from the experimenters and treating physiotherapist. The envelopes containing numbers regarding HI versus LD were randomly drawn from a basket and kept in a locked place."	
		Comment: An adequate method was used to generate the allocation sequence	
Allocation concealment (selection bias)	Low risk	Quote: "The randomization procedure was concealed from the experimenters and treating physiotherapist. The envelopes containing numbers regarding HI versus LD were randomly drawn from a basket and kept in a locked place."	

Manual therapy and exercise for rotator cuff disease (Review)



Osteras 2008 (Continued)

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		Comment: An adequate method was used to conceal the allocation sequence
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions (high intensity versus low in- tensity exercise), participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported some out- comes
Blinding of outcome as- sessment (detection bias) Objective outcomes	High risk	Quote: "The outcome measurements were not obtained by a blinded asses- sor"
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "92% of patients completed the study. Five patients were lost to post- test; two in the HD group (3%) and three in the LD group (5%). In the HD group, one patient moved away from the city and was therefore unable to keep in touch, and the second HD subject withdrew after experiencing a new injury at work that interfered with the continuation of the exercise treatment. In the LD group, one subject was referred by his physician for additional outpatient ther- apy and therefore withdrew from the study. The two last patients in the LD in- terventions were not able to return for the post-test for personal reasons."
		the results
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias were identified

Rhon 2014		
Methods	Study design: Parallel group RCT	
	Setting: Military hospital-based outpatient clinic, USA	
	<b>Intervention:</b> Manual physical therapy (joint and soft tissue mobilisations, manual stretches, con- tract-relax techniques, supervised exercises) plus home exercises	
	Control: Glucocorticoid injection plus home exercises	
	<b>Source of funding:</b> Cardon Rehabilitation Products through the American Academy of Orthopaedic Manual Physical Therapists	
Participants	Diagnostic label used by trialists: Impingement	
	Criteria for defining the shoulder condition being treated	
	<ul> <li>Primary symptom of unilateral shoulder pain</li> <li>Meets diagnostic criteria for shoulder impingement. To be included in the study participants are required to have: pain with one of the 2 tests in category I, and pain with one test from either category II or category III, where pain is defined as reproduction of the usual pain that the subject experiences that makes up the nature of their complaint.</li> <li>* Category I: Impingement signs - passive overpressure at full shoulder flexion with the scapula stabilised; passive internal rotation at 90 degrees of shoulder flexion in the scapular plane and in progressive degrees of horizontal adduction.</li> </ul>	

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Rhon 2014 (Continued)

- \* Category II: active shoulder abduction.
- \* Category Ill: resisted break tests: abduction; internal rotation; external rotation

#### Inclusion Criteria (not listed above)

- Age 18 and older
- Read, write, and speak English
- Eligible for healthcare at a military medical treatment facility

### **Exclusion Criteria (not listed above)**

- History of shoulder dislocation, fracture, or adhesive capsulitis
- History of glucocorticoid injection or physical therapy for the shoulder pain in the past 3 months
- Baseline SPADI score less than 20%
- Reproduction of shoulder symptoms with cervical spine examination
- History of systemic or neurologic disease affecting the shoulder
- Positive rotator cuff lag sign or history of full-thickness rotator cuff tear
- Pending litigation
- Inability to attend physical therapy for 3 consecutive weeks

#### **Baseline characteristics**

#### Intervention

Number randomised: 52; mean (SD) age: 40  $\pm$  12 years; sex: F/M: 17/29; mean (SD) duration of symptoms: 4.9  $\pm$  4.4 months

#### Control

Number randomised: 52; mean (SD) age: 42  $\pm$  12 years; sex: F/M: 14/38; mean (SD) duration of symptoms: 6.5  $\pm$  13.9 months

Interventions	Intervention: manual physical therapy			
	<i>Components of intervention</i> : the manual physical therapy intervention consisted of a combination of joint and soft-tissue mobilisations; manual stretches; contract–relax techniques reinforcing exercises directed to the shoulder girdle or thoracic or cervical spine. Participants did not receive identical treatments, but the manual physical therapy techniques were matched to individual impairments identified on examination. Home exercises (including wand ROM exercises, scapular retraction, scapular protraction, thoracic self-mobilisation, butterfly stretch) were prescribed to reinforce clinic interventions			
	Dose: 30 min (manual physical therapy); 2-3 times per day (home exercises)			
	Frequency of administration: twice weekly over a 3-week period			
	Control: glucocorticoid injection			
	<i>Components of intervention</i> : injected 40 mg of triamcinolone acetonide to the subacromial space of the symptomatic shoulder. Participants also received printed instructions to perform a gentle gravity-assisted distraction and oscillatory pendulum exercise			
	<i>Dose</i> : the physician spent approximately 30 min with each subject explaining the rationale for the injection, relevant anatomy, performing the procedure and reviewing the pendulum exercises			
	<i>Frequency of administration</i> : as many as three total injections could be administered by the study physician (1 month apart) during the 1-year period			
Outcomes	Outcomes were assessed at baseline, 1 month, 3 months, 6 months, and 1 year			
	• Function using the SPADI (0-100)			



Rhon 2014 (Continued)	
	<ul> <li>Pain using a numeric pain rating scale (0-10, with 0 representing no pain and 10 representing worst pain imaginable)</li> </ul>
	<ul> <li>Quality of life using the Global Rating of Change scale (-7 to +7)</li> </ul>
	Adverse events

Notes

**Conflicts of interest:** the authors stated that they had no conflicts of interest

Trial registered in ClinicalTrials.gov (NCT01190891) and trial protocol published in BMJ Open

#### **Risk of bias**

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "The randomization schedule was computer-generated"
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Low risk	Quote: "The randomization schedule was computer-generated, with assign- ments placed in opaque, sequentially numbered envelopes by an off-site in- vestigator not involved with patient care or follow-up. Treatment allocation was revealed after collection of baseline outcomes"
		Comment: An adequate method was used to conceal the allocation sequence
Blinding of participants	High risk	Quote: "Patients and treating clinicians were not blind to the intervention."
and personnel (perfor- mance bias) All outcomes		Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias)	High risk	Quote: "The research assistant who collected outcome assessments at each time point was blind to group assignment."
Self-reported outcomes		Comment: Despite having a blinded research assistant record patients' re- sponses, unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported all outcomes
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Most patients (96%) returned for follow-up visits at 1 year."
		Quote: "The primary analyses of effectiveness included all available data from patients who received their assigned treatment (that is, the CSI or at least 1 session of MPT). We used a linear mixed-effects model, which is flexible in accommodating data assumed to be missing at random with data from 5 time points (0, 1, 3, 6, and 12 months) for the SPADI (primary outcome) and NPRS and 4 time points for GRC. The intervention (MPT or CSI) was the fixed effect with random effects for the repeated measures over time within a patient; the primary treatment comparison was the difference between groups from baseline to 1 year. For the sensitivity analysis to explore the effect of missing data, they were imputed for the 3 outcome variables at all follow-ups (20 imputations using MULTIPLE IMPUTATION-FULLY CONDITIONAL SPECIFICATION)"
		Quote: "We performed a sensitivity analysis with imputation for missing data and the results remained unchanged."
		Comment: Attrition was dealt with using an appropriate method
Selective reporting (re- porting bias)	Low risk	Comment: Outcome data were fully reported for all outcomes pre-specified in the trial protocol
Other bias	Low risk	Comment: No other sources of bias identified

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# Senbursa 2007

Methods	Study design: Parallel group RCT		
	Setting: Outpatient clinic of Physiotherapy Rehabilitation, Hacettepe University, Ankara, Turkey		
	<b>Intervention:</b> Manual therapy programme (12 clinic-run sessions of joint and soft tissue mobilisation, ice, stretching and strengthening exercise programmes and education 3 times per week)		
	<b>Control:</b> Self-training programme (active ROM, strengthening and stretching exercise programm times a week)		
	Source of funding: Not reported		
Participants	Diagnostic label used by trialists: Impingement		
	Criteria for defining the shoulder condition being treated		
	<ul> <li>Shoulder impingement syndrome of the shoulder with: shoulder pain with no major joint trauma marked loss of active and passive shoulder motion or painful ROM</li> </ul>		
	Inclusion Criteria (not listed above)		
	<ul> <li>Taken no treatment at another physiotherapy clinic in the last 2 years</li> <li>Magnetic resonance imaging as a reference standard</li> </ul>		
	Exclusion Criteria (not listed above)		
	<ul> <li>History of frozen shoulders</li> <li>Disorders of the acromioclavicular joint</li> <li>Degenerative arthritis of the glenohumeral joint</li> <li>Calcifying tendonitis</li> <li>Shoulder instability</li> <li>Post-traumatic disorders</li> <li>Shoulder surgery</li> <li>Elbow, hand, wrist or cervical spine disorders</li> </ul>		
	Baseline characteristics		
	Intervention		
	Number randomised: 15; mean (SD) age: 48.1 (7.5) years; sex: not reported; duration of symptoms: not reported		
	Control		
	Number randomised: 15; mean (SD) age: 49.5 (7.9) years; sex: not reported; duration of symptoms: not reported		
Interventions	Intervention: manual therapy		
	<i>Components of intervention</i> : joint and soft tissue mobilisation: deep friction massage on supraspinatus muscle tendon, radial nerve stretching, scapular mobilisation, glenohumeral joint mobilisation, proprioceptive neuromuscular facilitation techniques including rhythmic stabilisation and hold-relax. An education programme, ice application, stretching and strengthening exercise programme and home training were also performed. The stretching and strengthening exercise programme was supervised by a physiotherapist and a shoulder exercise brochure also provided instructions. Home training was self-delivered with an elastic Thera-band		
	Dose: not reported		
	Frequency of administration: 12 sessions in total, completed in sessions 3 times per week for 4 weeks		

Manual therapy and exercise for rotator cuff disease (Review)

Senbursa 2007 (Continued)	Control: self-training:		
	<i>Components of intervention</i> : an active ROM, stretching and strengthening exercise programme includ- ing rotator cuff muscles, rhomboids, levator scapulae and serratus anterior which was self-adminis- tered at home using an elastic band at home after being taught by a physiotherapist		
	<i>Dose</i> : 10–15 min		
	Frequency of administration: 7 times a week for 4 weeks		
Outcomes	Outcomes assessed at 3 months		
	<ul> <li>Function: Neer functional assessment questionnaire from 0–100, with a higher score indicating better function (no outcome data reported)</li> </ul>		
	Rest pain: VAS scores ranging from 0-10, with a higher score indicating worse pain		
	<ul> <li>Night pain: VAS scores ranging from 0-10, with a higher score indicating worse pain</li> </ul>		
	<ul> <li>Pain on motion: VAS scores ranging from 0-10, with a higher score indicating worse pain</li> </ul>		
	<ul> <li>ROM (flexion, abduction, internal rotation and external rotation) measured using a goniometer (un- clear if active or passive) (no usable outcome data reported)</li> </ul>		
	<ul> <li>Strength: flexion, abduction, internal rotation and external rotation measured using manual muscle testing (no outcome data reported)</li> </ul>		

Notes Conflicts of interest: not reported

**Risk of bias** 

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "The short-term clinical effectiveness of manual physical therapy com- pared with usual care was assessed in a randomized clinical trial"
		Comment: No information on how the allocation sequence was generated was reported
Allocation concealment (selection bias)	Unclear risk	Comment: No information on how the allocation sequence was concealed was reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions (one delivered by physiother- apist, other delivered at home), participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported some out-comes
Blinding of outcome as- sessment (detection bias) Objective outcomes	Unclear risk	Comment: There was no information about whether assessors of objective outcomes were blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: There was no attrition reported, and outcome data were reported as based on the number of participants randomised
Selective reporting (re- porting bias)	High risk	Comments: No outcome data for the Neer function tests scores and strength outcomes were reported, despite being listed as outcomes in the methods section of the trial report. Also, without a trial protocol it is unclear whether other outcomes were assessed but not reported based on the results

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#### Senbursa 2007 (Continued)

Other bias

Low risk

Methods	Study design: Parallel group RCT
	Setting: Hacettepe University, Physiotherapy and Rehabilitation Department, Turkey
	Intervention: Joint and soft tissue mobilization plus supervised exercise programme
	Control 1: Supervised exercise programme
	Control 2: Home-based exercise programme
	Source of funding: Not reported
Participants	Diagnostic label used by trialists: Supraspinatus tendinopathy or Impingement
	Criteria for defining the shoulder condition being treated
	<ul> <li>Stage 1 supraspinatus tear or subacromial impingement syndrome</li> <li>Diagnosed by clinical exam (Neer and Hawkins tests) and MRI</li> </ul>
	Inclusion Criteria (not listed above)
	• None
	Exclusion Criteria (not listed above)
	<ul> <li>Shoulder trauma</li> <li>Shoulder instability</li> <li>Frozen shoulder</li> <li>Acromioclavicular and glenohumeral joint problems</li> <li>Calcified tendonitis</li> <li>Shoulder surgery</li> <li>Disease of the hand, wrist or cervical region</li> <li>Physical therapy or rehab programme in the last two years</li> </ul>
	Baseline characteristics
	Intervention
	Number randomised: 30; mean (SD) age: 50.5 $\pm$ 10.6 years old; sex: not reported; duration of symptom not reported
	Control 1
	Number randomised: 25; mean (SD) age: 48.2 $\pm$ 7.9 years old; sex: not reported; duration of symptoms not reported
	Control 2
	Number randomised: 22; mean (SD) age: 48.0 ± 9.0 years old; sex: not reported; duration of symptoms not reported
nterventions	Intervention: joint and soft tissue mobilisation
	<i>Components of intervention</i> : deep friction massage on the supraspinatus muscle, radial nerve stretch- ing, scapular mobilisation, glenohumeral joint mobilisation, and proprioceptive neuromuscular facili- tation techniques

Senbursa 2011 (Continued)			
	Dose: not reported		
	Frequency of administration: 3 times per week for 12 weeks		
	Control 1: supervised exercises		
	<i>Components of intervention</i> : ROM, stretching and strengthening exercises for the rhomboid, levator scapulae, serratus anterior and rotator cuff muscles supervised and at home		
	Dose: 3 sets of 10 repet	itions	
	Frequency of administration: 3 times per week with a physio and self-administered daily for 12 weeks		
	Control 2: home exercises		
	<i>Components of intervention</i> : ROM, stretching and strengthening exercises for the rhomboid, levator scapulae, serratus anterior and rotator cuff muscles at home only		
	Dose: 3 sets of 10 repet	itions	
	Frequency of administr	ation: daily for 12 weeks	
Outcomes	Outcomes assessed at	4 and 12 weeks	
	<ul> <li>Function: Modified American Shoulder and Elbow Surgery (MASES) questionnaire (no usable outcome data; in Figure only with unlabelled errors bars)</li> <li>Rest pain: VAS scored from 0-10 with a higher score indicating worse pain</li> <li>Night pain: VAS scored from 0-10 with a higher score indicating worse pain</li> <li>Pain on motion: VAS scored from 0-10 with a higher score indicating worse pain</li> <li>ROM (no outcome data reported)</li> <li>Strength measured using Lovett's manual muscle test, scored on a scale from 0-5 (no outcome data reported)</li> </ul>		
Notes	Conflicts of interest: the authors declared that they had no conflicts of interest		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Quote: "The patients were randomly assigned by the SPSS software to one of three consecutive treatment groups".	
		Comment: An adequate method was used to generate the allocation sequence	
Allocation concealment (selection bias)	Unclear risk	Comment: No information on how the allocation sequence was concealed was reported	
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention	
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported some out- comes	
Blinding of outcome as- sessment (detection bias) Objective outcomes	Unclear risk	Comment: There was no information about whether assessors of objective outcomes were blinded	

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Senbursa 2011	(Continued)
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Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "All patients underwent rehabilitation for 12 weeks" Comment: There was no attrition reported, and outcome data were reported as based on the number of participants randomised
Selective reporting (re- porting bias)	High risk	Comment: No outcome data were reported for several outcomes (e.g. ROM and strength); the authors only report that there was no statistically significant difference between groups
Other bias	Low risk	Comment: No other sources of bias identified

# Struyf 2013

Methods	Study design: Parallel group RCT			
	Setting: Private medical clinics or private physiotherapy practices, Belgium			
	Intervention: Scapular-focused treatment (stretching and scapular motor control training)			
	Control: Stretching, muscle friction and eccentric rotator cuff training, plus therapeutic ultrasound			
	<b>Source of funding:</b> This study was financially supported by MSD Europe bvba, Nijverheidsstraat 18, Londerzeel, Belgium, and by research grant (G842) supplied by the Department of Health Sciences, Artesis University College Antwerp, Antwerp, Belgium. The material used in this study was provided by MSD Europe bvba			
Participants	Diagnostic label used by trialists: Impingement			
	Criteria for defining the shoulder condition being treated			
	<ul> <li>Shoulder impingement lasting at least 30 days, with a prescription for their impingement symptoms from the physician or orthopaedic surgeon</li> </ul>			
	Inclusion Criteria (not listed above)			
	<ul> <li>Informed consent</li> <li>Age 18 years or older</li> <li>Ability to complete questionnaires (no dementia, sufficient knowledge of the Dutch language)</li> </ul>			
	Exclusion Criteria (not listed above)			
	<ul> <li>Shoulder pain onset due to trauma</li> <li>A history of shoulder fractures or dislocation, cervical radiculopathy, degenerative joint disease of the shoulder, surgical interventions on the shoulder, inflammatory arthropathy</li> <li>Infiltration of the shoulder in the previous 3 weeks</li> <li>Non-steroidal anti-inflammatory drug use</li> <li>Participants undergoing shoulder treatment (including physical therapy) 1 year prior to the first assessment</li> </ul>			
	Baseline characteristics			
	Intervention			
	Number randomised: 10; mean (SD) age: 45.4 $\pm$ 15.1 years; sex: F/M 5/5; duration of symptoms: not reported			
	Control			



Struyf 2013 (Continued)	Number randomised: 1 ported	.2; mean (SD) age: 46.2 ± 13.5 years; sex: F/M 7/5; duration of symptoms: not re-	
Interventions	Intervention: scapula	r-focused treatment	
	la, including: passive m or tilting), home stretcl pectoralis minor muscl	ntion: manual mobilisations, stretching and motor control training of the scapu- nanual mobilisation (to improve passive scapular upward rotation and posteri- hing exercises for the levator scapulae and rhomboids muscles, stretching of the le length by the physiotherapist and scapular motor control training with em- rientation exercise (SOE) Scapular orientation exercises completed with 10 repe-	
	Dose: 30-min session		
	<i>Frequency of administration</i> : 9 sessions in total, delivered between 1 and 3 times per week (depending on practical issues of the participant)		
	Control		
	Components of intervention		
	cles (15 min) and st into the following ra pain threshold. Betw flexion, extension, n ticipant was asked t starting position	I and home eccentric muscle strength training programme of the rotator cuff mus- rength training performed with the use of an elastic band. Training was divided egimen: three series of 15 repetitions, once per day, respecting the participant's ween the different series, there was a resting period of 2 min. The exercises were nedial rotation, and lateral rotation of the shoulder. During each exercise, the par- o quickly move in the desired direction and consequently slowly returning to the	
	• Manual therapy: passive (multidirectional) glenohumeral mobilisation (5 min) and friction massage therapy (5 min)		
	• Therapeutic ultrasound: performed with intermittent pulsations (100 Hz) of a 3 cm <sup>2</sup> probe, 2 w/cm <sup>2</sup> for 5 min, focused on the subacromial region		
	Dose: 30 min session		
	<i>Frequency of administration</i> : 9 sessions in total, delivered between 1 and 3 times per week (depending on practical issues of the participant)		
Outcomes	Outcomes assessed at the end of 9 treatment sessions (4-8 weeks) and 3 months after the final treat- ment session		
	<ul> <li>Function using the Shoulder Disability Questionnaire (Dutch version). The score is calculated by the summation of all yes answers, divided by all answered questions (yes or no), and subsequently multiplied by 100. This results in a score between 0 (no disabilities) to 100 (severely disabled).</li> <li>Rest pain using a 100 mm VAS</li> </ul>		
	<ul> <li>Pain during activity using a 100 mm VAS</li> </ul>		
	<ul> <li>Strength: isometric elevation strength measured in the Jobe's test position (arm elevated to 90 de- grees in the plane of the scapula and internally rotated by pointing the thumb down) using a hand- held dynamometer</li> </ul>		
Notes	Conflicts of interest: the authors stated that they had no conflicts of interest		
	Trial registered (ISRCTN20736216)		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Low risk	Quote: "The patient took a form (with a letter A (n=23) or B (n=23)) indicating allocation to either groups from a closed envelop. A list with patient numbers and the group allocation that resulted from this randomization procedure was	

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Struyf 2013 (Continued)

Struy 2023 (continued)		stored in a sealed envelope. Only the therapist had direct access to the ran- domization list. In this way, patients were randomly allocated to either treat- ment group A or B." Comment: An adequate method was used to generate the allocation se- quence
Allocation concealment (selection bias)	Low risk	Comment: An adequate method was used to conceal the allocation sequence
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Comment: Participants received slightly different multi-modal interventions, but it is unclear whether they were provided with any information that would make them perceive the intervention they received as superior or inferior to the alternative intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	Unclear risk	Comment: Participants self-reported some outcomes, but it is unclear whether they were provided with any information that would make them perceive the intervention they received as superior or inferior to the alternative interven- tion
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "All assessments were performed by the same examiner blinded for group allocation. The order of the assessments (primary and secondary out- comes) was randomized to avoid order effects." Comment: A blinded assessors measured objective outcomes (e.g. muscle strength)
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: Two participants dropped out of the control group, one because of cervical pain, the other was unable to be contacted. No participants dropped out of the intervention group. Analysis was by intention-to-treat with last ob- servation carried forward. Missing data unlikely to affect the results
Selective reporting (re- porting bias)	Low risk	Comment: Outcome data were fully reported for all outcomes reported in the clinical trials registry
Other bias	Low risk	Comment: No other sources of bias identified

### Subasi 2012

Methods	Study design: Parallel group RCT			
	Setting: Outpatients			
	<b>Intervention:</b> Water-based exercise programme and surface heat and TENS and deep heat (ultra- sound)			
	Control: Land-based exercise programme and surface heat and TENS and deep heat (ultrasound)			
	Source of funding: Not reported			
Participants	Diagnostic label used by trialists: Impingement			
	<b>Criteria for defining the shoulder condition being treated:</b> Subacromial impingement syndrome de- fined by:			
	fined by:			

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Subasi 2012 (Continued)

#### Inclusion Criteria (not listed above)

- Outlet and axillary radiographs if required
- MRI if required

#### **Exclusion Criteria (not listed above)**

- Participants with non-shoulder-related pathologies that could lead to shoulder pain
- Infections and malignancies
- Shoulder instability
- Calcified tendinitis
- Calcified tendonitis and bursitis detected with conventional radiography
- A history of cervical, shoulder, or back surgery
- Corticosteroid injections or physical therapy due to a similar complaint involving the shoulder in the last 6 months
- Cervical radiculopathies
- Total rotator cuff tears
- Fractures or dislocations as a result of severe acute trauma
- Dementia or other psychiatric illnesses
- Adhesive capsulitis

#### **Baseline characteristics**

#### Intervention

Number randomised: 28 participants (35 shoulders); mean (SD) age: 58.3 ± 8.6 years old; sex: F/M 21/7; mean (SD) duration of symptoms: 8.9 ± 7.5 months

Control

Number randomised: 29 participants (35 shoulders); mean (SD) age:  $56.2 \pm 11.3$  years old; sex: F/M 15/14; mean (SD) duration of symptoms:  $10.0 \pm 13.2$  months

Interventions

#### Intervention: land-based exercise

*Components of intervention:* supervised land-based exercises. For the first 10 days, ROM and stretching exercises, and for the following 10 days, strengthening exercises were performed. After completion of 20 days' therapy, participants continued a home exercise programme twice daily

Dose: not reported

Frequency of administration: daily for the first 20 days and then twice daily for 3 months

#### **Control: water-based exercise**

*Components of intervention:* supervised water-based exercises. For the first 10 days, ROM and stretching exercises, and for the following 10 days, strengthening exercises in water by using dumbbells were performed. Water-based exercises took place in a therapy pool maintained at 28-30°C, which was 8 metres in width, 12 metres in length, and 1.4 metres at its deepest point. After completion of 20 days' therapy, participants continued a home exercise programme twice daily

Dose: Not reported

Frequency of administration: daily for the first 20 days and then twice daily for 3 months

#### **Both Groups**

Components of intervention

 Surface heat: heat packs filled with silica gel. Heated to 75°C and then administered wrapped in a towel



Subasi 2012 (Continued)

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Subasi 2012 (Continued)	<ul> <li>TENS: Sonopuls 492 machine using 4 carbon-silicone composite electrodes (2 x 2 cm in size). The electrodes were placed over the region of shoulder pain and operated using "the conventional method".</li> <li>Therapeutic ultrasound: Sonopuls 492 machine and Ultrasound Gel Therascanc. Continuous and circular motion to the anterior, posterior and lateral parts of the involved shoulder</li> </ul>			
	Dose:			
	-	n ncy for 60 microseconds with an amplitude under the motor threshold for 20 min und: 1.5 w/cm <sup>2</sup> and frequency 1 MHz for 8 min		
	Frequency of administr	ation:		
	<ul><li>Surface heat: not re</li><li>TENS: not reported</li><li>Therapeutic ultraso</li></ul>	ported und: 5 times a week for 20 sessions		
Outcomes	Outcomes assessed at	end of 20 days treatment and at 3 months post-treatment initiation (~8 weeks)		
	<ul> <li>Function: SPADI sco</li> <li>Quality of life: WORG of life and the worst</li> </ul>	n 0–10 with a higher score indicating more pain) re (0-100 with higher scores indicating more dysfunction) C score. The total score ranges from 0-2100 where 0 implies no reduction in quality c score is 2100 ROM (no outcome data reported)		
Notes	Conflicts of interest: the authors stated that they had no conflicts of interest			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "Of the patients, 28 were randomized into the land-based exercise group (LG) and 29 were randomized into the water-based exercise group (WG)."		
		Comment: No information on how the allocation sequence was generated was reported		
Allocation concealment (selection bias)	Unclear risk	Comment: No information on how the allocation sequence was concealed was reported		
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention		
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported some out-comes		
Blinding of outcome as- sessment (detection bias) Objective outcomes	Unclear risk	Comment: There was no information about whether assessors of objective outcomes were blind to treatment		
		Comment: The flow of participants through the trial was not described, and		

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## Subasi 2012 (Continued)

Selective reporting (re- porting bias)	High risk	Comment: The authors state in the methods section that specific tests (un- specified) and active and passive ROM were measured, but no data were re- ported for these outcomes. Also, without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias identified

### Surenkok 2009

Methods	Study design: Parallel group RCT				
	Setting: Sports physiotherapy clinic, Turkey				
	Intervention: Scapular mobilisation				
	Control 1: Sham mobilisation				
	Control 2: No treatment				
	Source of funding: Not reported				
Participants	Diagnostic label used by trialists: tendinitis or tenosynovitis				
	Criteria for defining the shoulder condition being treated				
	<ul> <li>Painful limitation of shoulder range of movement that had persisted for at least 4 weeks</li> <li>Inability to elevate the arm more than 100 degrees in the scapular plane because of pain over the anterior aspect of either shoulder</li> </ul>				
	Inclusion Criteria (not listed above)				
	• None				
	Exclusion Criteria (not listed above)				
	<ul> <li>Cervical symptoms (numbness or tingling in the upper extremity)</li> <li>A history of onset of symptoms because of a traumatic injury</li> <li>A history of shoulder surgery</li> </ul>				
	Baseline characteristics				
	Intervention				
	Number randomised: 13; mean (SD) age: 55.07 (13.36) years; sex: F/M 10/3; duration of symptoms: not reported				
	Control 1				
	Number randomised: 13; mean (SD) age: 54.30 (12.70) years; sex: F/M 2/11; duration of symptoms: not reported				
	Control 2				
	Number randomised: 13; mean (SD) age: 55.53 (17.15) years; sex: F/M 10/3; duration of symptoms: not reported				
Interventions	Intervention: scapular mobilisation				
	<i>Components of intervention</i> : application of superior and inferior gliding, rotations, and distraction to the scapula of the affected shoulder. The participants laid the affected forearm on their back. The therapist stood before the participant's affected shoulder, placing the index finger of one hand under the				

Manual therapy and exercise for rotator cuff disease (Review)



Surenkok 2009 (Continued)

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Surenkok 2009 (Continued)	moved superiorly and i ward and downward for	r, the other hand grasping the superior border of the scapula. The scapula was inferiorly for superior and inferior glide, and then the scapula was rotated up- or scapular rotation. Second, while the participant was in the same position the e ulnar fingers under the medial scapular border and distracted the scapula from		
	Dose: Sets of 10 repetit	ions with 30 seconds between each set		
	Frequency of administr	ation: once		
	Control 1: sham mobilisation			
	positioning. The therap	<i>ntion</i> : the sham condition replicated the treatment condition except for the hand bist placed one hand on the medial aspect of the scapula and the other hand on A simulated scapulothoracic movement, but with minimal pressure, was actually		
	Dose: sets of 10 repetit	ions with 30 seconds between each set		
	Frequency of administr	ation: once		
	Control 2: no treatme	nt		
		<i>ntion</i> : the participant was seated for the same length of time, but no manual con- pist and the participant took place		
	Dose: not applicable			
	Frequency: once			
Outcomes	Outcomes assessed immediately post-treatment (i.e. same day of treatment)			
	<ul><li> Rest pain: VAS from</li><li> Pain on motion: VAS</li></ul>	Shoulder Score from 0–100, with a higher score indicating better function 0 mm (no pain) to 100 mm (extreme pain) 6 from 0 mm (no pain) to 100 mm (extreme pain) uction and flexion using a goniometer		
Notes	Conflicts of interest: r	not reported		
	22 (56%) participants h sive capsulitis	ad tendinitis, 10 (26%) had tenosynovitis, and 7 (18%) of participants had adhe-		
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "Participants received 1 of 3 treatment conditions (SM, sham, or con- trol) in a randomized order known only by the treating therapist."		
		Comment: No information on how the allocation sequence was generated was reported		
Allocation concealment (selection bias)	Unclear risk	Comment: No information on how the allocation sequence was concealed was reported		
Blinding of participants and personnel (perfor- mance bias)	High risk	Quote: "The same physiotherapist applied all treatment conditions; a second physiotherapist took pre-outcome and post-outcome measurements. Both physiotherapists were blind to the group of each subject."		
All outcomes		Comment: Given the nature of the interventions, participants and physiother- apists could not be blind to treatment for the comparison of "mobilisation versus no treatment", and participants may have had different expectations		

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#### Surenkok 2009 (Continued)

Surenkok 2009 (continued)		about the benefits of each intervention. However, participants in the compari- son of "mobilisation versus sham mobilisation" were blind to treatment
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: In the comparison between mobilisation and no treatment, un- blinded participants who may have had different expectations about the ben- efits of the intervention they received self-reported some outcomes. However, in the comparison between mobilisation and sham mobilisation, blinded par- ticipants self-reported outcomes
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "The same physiotherapist applied all treatment conditions; a second physiotherapist took pre-outcome and post-outcome measurements." Comment: Assessor of objective outcomes was blind to treatment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: There were no dropouts, losses to follow-up or exclusions
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias were identified

#### Szczurko 2009

Study design: Parallel group RCT				
Setting: Postal employees who were members of the Canadian Union of Postal Workers, Canada				
Intervention: Physical exercise, hands on shoulder muscle and joint therapy and placebo tablets				
Control: Naturopathic care (dietary counselling, acupuncture, and Phlogenzym supplement)				
<b>Source of funding:</b> "Supported by The Canadian Union of Postal Workers and the Canada Post Corporation, Joint Benefits Committee. Mucos Pharma, Puhonice, Czech Republic and Heel Canada, Anjou, Quebec, Canada supplied the study drug"				
Diagnostic label used by trialists: Rotator cuff tendinitis				
Criteria for defining the shoulder condition being treated				
Symptoms consistent with rotator cuff tendonitis				
<ul> <li>Physical exam suggesting rotator cuff tendonitis (ROM limitations; Neer Impingement, Speeds, Apprehension, Subscapularis Lift tests)</li> </ul>				
Inclusion criteria (not listed above)				
Between 18 and 65 years old				
<ul> <li>Judged to be able to adhere to the given protocol</li> </ul>				
Pain in at least 1 shoulder for the previous 6 weeks				
Exclusion criteria (not listed above)				
<ul> <li>Allergies to the trial supplement</li> <li>Receiving corticosteroid injection therapy</li> </ul>				
Taking daily warfarin or antibiotics				
-				



Szczurko 2009 (Continued)	<ul> <li>Had a severe concurrent illness</li> <li>Pregnant or breastfeeding</li> </ul> Baseline characteristics			
	Intervention			
	Number randomised: 42; mean (SD) age = 50.9 (7.86) years; sex: F/M 25/17; duration of symptoms: not reported			
	Control			
	Number randomised: 4 reported	3; mean (SD) age = 50.7 (8.16) years; sex: F/M 25/18; duration of symptoms: not		
Interventions	Intervention: physica	l therapy		
	and active ROM muscle therapy for shoulder in	<i>ition:</i> standardised exercise programme consisting of passive, active assisted strengthening and joint therapy, reportedly consistent with standard physio- juries; hands-on shoulder muscle and joint therapy; placebo tablets consisting nce, matched to Phlogenzym in appearance, smell and taste		
	Dose: 30-min physical t	herapy consultations; 2 placebo tablets, 3 times per day		
	Frequency of administre	ation: once weekly for 12 weeks		
	Control: naturopathic care			
	<i>Components of intervention:</i> individualised dietary counselling with emphasis on reduction of alcohol intake and increase in consumption of fish, berries, fruits, vegetables, nuts and whole grains; standard-ised acupuncture treatment – needle insertion at LI15, SJ14, SI19, SI10-13 and BL41-46 plus up to 4 Ashi points of pain (needles were inserted and briefly stimulated using perpendicular thrusting technique); phlogenzym supplement (90 mg bromelain, 48 mg trypsin, 100 mg rutin)			
	<i>Dose &amp; Frequency of administration:</i> dietary counselling and acupuncture: one 30-min session per week for 12 weeks; supplement: 2 tablets, 3 times per day for 12 weeks			
Outcomes	Outcomes assessed at 4, 8 and 12 weeks post randomisation (only 12 week data reported)			
	<ul> <li>Function: Shoulder pain and disability index (SPADI) total score (0-130 score, where a higher score indicates worse pain and disability)</li> </ul>			
	<ul> <li>Overall pain: average degree of pain over a week using a VAS. Scores ranging from 0 (no pain at all) to 7 (severe pain)</li> </ul>			
	<ul> <li>Quality of life using the SF-36 questionnaire (8 domains with scores from 0-100, summarised into a physical function score and mental health score, where a higher score represents better health)</li> <li>Active ROM (shoulder flexion, extension, abduction, adduction, internal rotation, external rotation) using a goniometer/inclinometer</li> <li>Adverse events</li> </ul>			
Notes	Adverse events  Conflicts of interest: not reported			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence genera- tion (selection bias)	Low risk	Quote: "participants were randomized using age- and sex-matched comput- er randomization"		
		Comment: An adequate method was used to generate the allocation sequence		
Allocation concealment (selection bias)	Low risk	Quote: "Allocation concealment using central randomization was preserved up to the point of treatment"		

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Szczurko 2009 (Continued)

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		Comment: An adequate method was used to generate the allocation sequence
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "participants were blinded to allocation and supplements were de- livered using identical-looking tablets for all supplements and placebo, it was not possible to mask the interventions from the participants or the clinicians delivery care".
		Comment: Given the nature of the interventions, participants were not blind to treatment and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the interventions they received self-reported some out-comes
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "The flexion, extension, abduction, adduction, internal rotation, and external rotation of the affected shoulder were assessed by a coordinator blinded to treatment and using a goniometer/inclinometer"
		Comment: Assessor of objective outcomes was blinded to treatment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "In total, 89 patients were randomized and enrolled in the study. Four of the 89 patients decided to not start the study after reconsideration or be- came unreachable before the first treatment visit. None of these 4 participants withdrew with the knowledge of what type of treatment they would be receiv- ing. Of the 85 participants who started treatment, 17 (10 control, 7 active) did not complete the 12-week course of study: 1 participant broke her leg, 6 be- came unreachable, and 10 could not commit the time or lost interest. Of the 43 participants who started treatment in the NC group, 41 completed week 8 and 36 completed week 12. Of the 42 participants who started treatment in the PE group, 36 completed week 8 and 32 completed week 12."
		Comment: The amount and reasons for attrition are unlikely to have affected the results
Selective reporting (re- porting bias)	Unclear risk	Comment: No data for some measures of ROM (active internal rotation and ex- ternal rotation) were reported, but it is unclear if non-reporting was related to the results. Also, without a trial protocol, it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: No other sources of bias identified

Teys 2008	
Methods	Study design: Cross-over RCT
	Setting: General population in south-east Queensland, Australia
	Intervention: Postero-lateral glide (mobilisation with movement)
	Control 1: Sham postero-lateral glide
	Control 2: No treatment
	Source of funding: Not reported
Participants	Diagnostic label used by trialists: None specified
	Criteria for defining the shoulder condition being treated

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#### Teys 2008 (Continued)

- Painfully limited shoulder for more than 1 month but less than 1 year
- Inability to elevate the arm more than 100 degrees in the plane of the scapula because of the presence of pain over the anterior aspect of either shoulder

#### Inclusion Criteria (not listed above)

• None

#### **Exclusion Criteria (not listed above)**

- · Shoulder pain not deemed of musculoskeletal origin
- Medical treatment which would exclude the patient from physiotherapy treatment
- Active inflammatory disease
- Infection
- Cancer
- Neuromuscular disorders
- Fractures around the shoulder
- Evidence of cervical spine referral of pain to the shoulder

#### **Baseline characteristics**

Overall cohort of participants

Number randomised: 24; mean (SD) age: 46.1 (9.86) years old; sex: F/M 13/11; duration of symptoms: not reported

# Interventions Intervention: postero-lateral glide (Mulligans' mobilisation with movement) Components of intervention: participant was seated and the therapist stood beside the participant on the opposite side to the affected shoulder. One hand was placed over the scapula posteriorly while the thenar eminence of the other hand was placed over the anterior aspect of the head of the humerus. A posterior gliding force was applied to the humeral head. The participant was then asked to raise the affected arm in the plane of the scapula to the point of pain onset while the therapist sustained the gliding force to the humeral head, with care to avoid the sensitive coracoid process. The therapist endeavoured to maintain the glide at right angles to the plane of movement throughout the entire range Dose: 3 sets of 10 repetitions with a 30-second rest between each set Frequency of administration: once **Control 1 - Sham Mobilisation** Components of intervention: the therapist stood on the opposite side of the participant and placed one hand along the clavicle and sternum and the other on the posterior aspect of the humeral head of the affected shoulder. A simulated anterior glide was performed but with minimal pressure actually applied. The participant was asked to elevate the affected shoulder in the plane of the scapula through half of their available pain-free range to minimise the likelihood of pain provocation Dose: 3 sets of 10 repetitions with a 30-second rest between each set Frequency of administration: once **Control 2 - No treatment** Components of intervention: participant seated for the same length of time as in the other groups, but no manual contact made Dose: NA Frequency of administration: once

Outcomes Outcomes assessed immediately post-treatment (i.e. same day of treatment)



Teys 2008 (Continued)

- Active ROM: pain-free motion in the scapular plane measured with a universal goniometer (degrees)
- Adverse events

Notes	Conflicts of interest: not reported	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "The treatment allocation sequence was block randomized using the drawing of lots and concealed from the investigator who took the outcome measurements"
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Unclear risk	Comment: No information on how the allocation sequence was concealed was reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Participant blinding was facilitated by recruitment of people who had no experience of the manipulative therapy techniques applied to the shoulder and by careful instruction that did not refer to the study's aims of evaluation of a treatment technique. Subjects were informed that the study was investi- gating the effects of manual handling on shoulder pain. An exit questionnaire assessed the adequacy of patient blinding. Results of the exit questionnaire showed that three participants (12%) correctly guessed they had only received active treatment and none had correctly guessed that they had received either a sham or control."
		Comment: Participants were likely blinded to the intervention they received
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	Low risk	Comment: Blinded participants self-reported pressure pain threshold
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "The outcome measures were taken by an investigator skilled in their application and who remained blind to the allotted treatment condition."
		Comment: Assessor of ROM was blinded to treatment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "There was no loss to follow-up"
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: The study appears to be free of other sources of bias

#### Van den Dolder 2003

Methods	Study design: Parallel group RCT	
	Setting: Repatriation hospital in Sydney, Australia	
	Intervention: Soft tissue massage	
	Control: No treatment	
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Source of funding: Not reported

#### Van den Dolder 2003 (Continued)

Participants Diagnostic label used by trialists: Impingement, supraspinatus/rotator cuff tear, supraspinatus tendinitis, or rotator cuff tendinitis Criteria for defining the shoulder condition being treated Shoulder pain Inclusion Criteria (not listed above) • Able to understand spoken English • Aged between 18 and 80 **Exclusion Criteria (not listed above)** • Pain was due to trauma within the previous 4 weeks Pain was reproduced on combined cervical extension, ipsilateral rotation or side flexion with overpressure Pain was due to a neoplastic disorder · Pain was of an acute inflammatory nature There was no palpable tenderness over the posterior aspect of the shoulder or over the anterior portion of the deltoid muscle or the pectoralis major muscle **Baseline characteristics** Intervention Number randomised: 15; mean (SD) age: 63.1 (9.9) years old; sex: F/M 4/11; median (IQR) duration of symptoms: 26 (13-26) weeks No treatment Number randomised: 14; mean (SD) age: 65.9 (9.2) years old; sex: F/M 5/9; median (IQR) duration of symptoms: 30 (23-91) weeks Interventions Intervention: soft tissue massage Components of intervention: soft tissue massage of the shoulder performed as seen fit by the treating therapist. The areas focused on were the lateral border of the scapula, in full shoulder flexion; posterior deltoid, at end of range horizontal flexion; anterior deltoid, at end of range hand-behind-back; and pectoralis major, in the stretch position Dose: 15-20 min Frequency of administration: 6 treatments over 2 weeks **Control: no treatment** Components of intervention: stayed on a waiting list Frequency: 2 weeks Outcomes Outcomes assessed at 2 weeks Function: Patient Specific Functional Disability Measure (PSFDM) score rated 0-30 with a higher score indicating better function Pain: Short Form McGill Pain Questionnaire (SFMPQ) VAS pain score rated 0-100 (mm) with a higher score indicating greater pain in the last 24 hours Active ROM: flexion, abduction (both in degrees) and hand-behind-back (HBB) distance (cm) Notes Conflicts of interest: not reported

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#### Van den Dolder 2003 (Continued)

#### **Risk of bias**

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "Randomisation was performed by selection of a sealed envelope from a container of identical envelopes, inside which were instructions regarding which group the patient was to be allocated to."
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Low risk	Quote: "A third person, who arranged all necessary follow-up appointments, opened each envelope. This ensured concealment of allocation to both pa-tients and assessor"
		Comment: An adequate method was used to generate the allocation sequence
Blinding of participants	High risk	Quote: "Blinding of the patients to allocation was not possible".
and personnel (perfor- mance bias) All outcomes		Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported some out-comes
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "A third person, who arranged all necessary follow-up appointments, opened each envelope. This ensured concealment of allocation to both patients and assessor."
		Comment: Assessor of objective outcomes was likely blind to treatment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: There was no attrition and all outcome data were analysed based on the number of randomised participants
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: The study appears to be free of other sources of bias

Valther 2004		
Methods	Study design: Parallel group RCT	
	Setting: Not reported (Germany)	
	Intervention: Standardised self-training	
	<b>Control 1:</b> Conventional physiotherapy (stretching exercises)	
	Control 2: Functional brace	
	Source of funding: Not reported	
Participants	Diagnostic label used by trialists: Impingement	
	Criteria for defining the shoulder condition being treated	
	and a family state of the second state of the	

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Walther 2004 (Continued)

Subacromial impingement confirmed by:

- clinical examination
- radiographs of the shoulder in three planes
- ultrasound
- positive Neer test (subacromial injection of 10 ml pure bupivacaine)

#### Inclusion Criteria (not listed above)

• None

#### **Exclusion Criteria (not listed above)**

- Cervical radiculopathy
- Frozen shoulder
- Full-thickness tear of the rotator cuff
- Disorders of the acromioclavicular joint
- Degenerative arthritis of the glenohumeral joint
- Calcifying tendonitis
- Shoulder instability
- Post-traumatic disorders
- · Involvement in workers' compensation claims

#### **Baseline characteristics**

#### Intervention

Number randomised: 20; mean (range) age: 52.1 (40 – 66) years; sex: F/M 11/9; mean (range) duration of symptoms: 23 (3 – 72) months

Control 1

Number randomised: 20; mean (range) age: 51.5 (37 – 66) years; sex: F/M 9/11; mean (range) duration of symptoms: 32 (2 – 120) months

#### Control 2

Number randomised: 20; mean (range) age: 48.6 (25 – 61) years; sex: F/M 6/14; mean (range) duration of symptoms: 27 (5 – 60) months

Interventions

#### Intervention: standardised self training

*Components of intervention*: standardised self-training programme of centring and stretching exercises that affected the shoulder. Instructions for the exercise programme were printed with PhysioTools software. For most of the exercises, an elastic Thera-Band was used that was chosen according to the results of the initial force measurements. The self-training programme was taught to participants under the guidance of a physiotherapist for a maximum of 4 sessions

Dose: 10-15 min

Frequency of administration: 5 times per week for 12 weeks

#### **Control 1 - Conventional Physiotherapy**

*Components of intervention*: physiotherapy consisting of centring training for the rotator cuff. Stretching was added in case of any limitation of the ROM at the first examination

Dose: 10 sessions initially and further sessions prescribed by family doctor

Frequency of administration: 2–3 times per week for 12 weeks

#### **Control 2 - Functional brace**

Walther 2004 (Continued)	<i>Components of intervention</i> : participants were supplied with a functional shoulder brace (Coopercare Lastrap; Coopercare Inc, Toronto, Ontario, Canada). They were instructed on how to use the brace and told to use it as long as possible during the day and, if comfortable, also at night. The Coopercare Lastrap shoulder brace consists of a cotton sleeve and special Thermovibe pads. The presumed effect of the brace is the absorption of vibrations and the accumulation of heat. The brace is fixed with two elastic Velcro straps
	Dose: as long as possible during the day and at night if possible
	Frequency: Every day for 12 weeks
Outcomes	Outcomes assessed at 6 weeks and 12 weeks
	• Function: Constant total score from 0–100 with a higher score indicating better function (no usable outcome data reported)
	• Rest pain: VAS from 0 (pain free) to 100 (maximum pain) (no usable outcome data reported)
	• Night pain: VAS from 0 (pain free) to 100 (maximum pain) (no usable outcome data reported)
	• Pain on motion: VAS from 0 (pain free) to 100 (maximum pain) (no usable outcome data reported)
	Active ROM: Constant sub-score (0-40) (no usable outcome data reported)
	• Strength: Constant sub-score (0-25) (usable outcome data reported but not extracted as data selec- tively reported based on the statistical significance of the results)
	Work disability: number of months with inability to work
	Adverse events

Conflicts of interest: not reported

# Risk of bias

Notes

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "After informed consent was obtained, 60 consecutive patients with painful disabling impingement syndrome of the shoulder were randomized into three different conservative treatment groups"
		Comment: No information on how the allocation sequence was generated was reported
Allocation concealment (selection bias)	Unclear risk	Comment: No information on how the allocation sequence was concealed was reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported some out-comes
Blinding of outcome as- sessment (detection bias) Objective outcomes	Unclear risk	Comment: There was no information about whether assessors of objective outcomes were blind to treatment
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "None of the patients treated with physiotherapy or self-training dropped out of the therapy regimen. However, one of the patients treated with the brace complained about being bothered by the brace at work, especially while working overhead. Another patient had eczema of the skin develop un- derneath the pads. Both patients continued to wear the brace during the re- mainder of the 12-week therapy period."

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#### Walther 2004 (Continued)

		Comment: There was no attrition
Selective reporting (re- porting bias)	High risk	Comment: Measures of variation were only reported for the few outcomes that had statistically significant results
Other bias	Low risk	Comment: The study appears to be free of other bias

Methods	Study design: Parallel group RCT			
	Setting: Outpatient physical therapy clinic affiliated with an academic intuition, USA			
	Intervention: Customised exercises			
	Control: Standardised exercises			
	Sources of funding: Texas Physical Therapy Foundation			
Participants	<b>Diagnostic label used by trialists:</b> one of eight scapular or humeral syndromes - scapular downward rotation syndrome, scapular depression syndrome, scapular abduction syndrome, scapular winging syndrome, humeral anterior glide syndrome, humeral superior glide syndrome, humeral medial rotation syndrome, humeral hypomobility syndrome			
	Criteria for defining the shoulder condition being treated			
	• Shoulder pain for more than 10 days (classified as one of 8 scapular or humeral syndromes)			
	Inclusion Criteria (not listed above)			
	At least 21 years old			
	Exclusion Criteria (not listed above)			
	<ul> <li>Received physical therapy within the past 6 months</li> <li>Had concurrent neck or thoracic disorders</li> <li>Had systemic disease such as rheumatoid arthritis or diabetes mellitus</li> </ul>			
	Baseline characteristics			
	Intervention			
	Number randomised: 20; mean (SD) age: 39.3 ± 13.2 years; sex: F/M 9/6; mean (SD) duration of symp- toms: 49.6 ± 52.4 weeks			
	Control			
	Number randomised: 18; mean (SD) age: 49.9 ± 18.3 years; sex: F/M 6/9; mean (SD) duration of symp- toms: 41.6 ± 51.5 weeks			
nterventions	Intervention: customised exercises			
	<i>Components of intervention</i> : mainly supervised and home self-stretching and strengthening exercises es for scapular stabilisers, rotator cuff and scapulohumeral muscles. There were 6 exercises for each of the potential 8 categories that participants could be classified into based on the examination of the physical therapist. When subjects were able to complete the shoulder exercises without difficulty or without increasing symptoms, they were progressed to the next level of exercises. In general, strength ening exercises were progressed by increasing repetitions and resistance, while stretching exercises in creased by hold time. Resistance was provided by Thera-band or gravity in weaker muscles. When par- ticipants had difficulty or symptoms increased with the prescribed exercises, the exercise level was de			



Wang 2006 (Continued)

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vated the symptoms.

	valed the symptoms.	
	<i>Dose</i> : 5 repetitions of a with participant ability	5-second hold for each exercise. Resistance level or hold time was increased
	Frequency of administr for 8 weeks	<i>ation</i> : Supervised - once weekly for 8 weeks; Home - twice daily 5 times per week
	Control: standardised	l exercises
		<i>ntion</i> : same as above, except that 5 standardised strengthening exercises were xors, abductors, extensors, external rotators and internal rotators.
	<i>Dose</i> : 5 repetitions of a with participant ability	5-second hold for each exercise. Resistance level or hold time was increased
	<i>Frequency of administr</i> for 8 weeks	<i>ation</i> : supervised - once weekly for 8 weeks; home - twice daily 5 times per week
Outcomes	Outcomes assessed at	4 and 8 weeks
	• Pain: VAS scale rate	d 0-10 with a higher score indicating worse pain (cm)
		scale of shoulder function from 1–50, with a higher score indicating better function tion, internal rotation and external rotation) recorded using a standard universal es)
		strength of abductors, external rotator, internal rotator, middle trapezius and low- red using a handheld dynamometer (N-m)
Notes	Conflicts of interest:	not reported
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "Subjects were then randomly assigned to one of two exercise groups by selecting a number in a pre- prepared sealed envelope"
		Comment: No information on how the allocation sequence was generated pri- or to inserting numbers in envelopes
Allocation concealment (selection bias)	Unclear risk	Quote: "Subjects were then randomly assigned to one of two exercise groups by selecting a number in a pre- prepared sealed envelope"
		Comment: Not clear if envelopes were sequentially numbered or shuffled by participants
Blinding of participants and personnel (perfor-	High risk	Quote: "This is a single-blinded randomized clinical trial. The tester was blind- ed from the subject's treatment group."
mance bias) All outcomes		Comment: Participants and personnel were not blinded
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants self-reported pain and function
Blinding of outcome as- sessment (detection bias)	Low risk	Quote: "Four outcome measurements were obtained by a tester who was blinded to the assigned treatment group"
		binded to the assigned treatment group
Objective outcomes		Comment: Assessor of objective outcomes was likely blind to treatment

creased by decreasing the resistance, decreasing the repetitions, or stopping the exercises that aggra-

Manual therapy and exercise for rotator cuff disease (Review)

#### Wang 2006 (Continued)

Incomplete outcome data (attrition bias) All outcomes	Low risk	Comment: 5 and 3 participants dropped out of each respective group, but the reasons were balanced between groups, so attrition is unlikely to have affected the results
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: The study appears to be free of other bias

### Winters 1997

Methods	Study design: Parallel group RCT			
	Setting: General practices, The Netherlands			
	Intervention: Physiotherapy (exercise, massage)			
	Control 1: Manipulation			
	Control 2: Glucocorticoid injection			
	Source of funding: Ministry of Welfare, Health and Culture			
Participants	Diagnostic label used by trialists: None specified			
	Criteria for defining the shoulder condition being treated			
	Patients consulting their GP with shoulder complaints as follows:			
	<ul> <li>pain localised in the region of the deltoid muscle, acromioclavicular joint, superior part of the trape- zoid muscle and Scapula;</li> </ul>			
	<ul> <li>radiation of the pain in the arm could be present;</li> </ul>			
	The ROM of the upper arm or shoulder girdle could be limited			
	The participants were then divided into three diagnostic subgroups			
	<ul> <li>Synovial group: participants with pain or limited movement in one or several directions of the glenohumeral joint. These complaints originated from disorders of the subacromial structures, the acromioclavicular joint, the glenohumeral joint, or combination of these (the synovial structures)</li> <li>Shoulder girdle group: consisted of participants with pain and sometimes slightly limited range of active movement of the glenohumeral joint. These problems were not related to the synovial structures but, instead, probably originated from functional disorders of the cervical spine, upper thoracic spine, or the upper ribs (the shoulder girdle)</li> </ul>			
	• Combination group: consisted of participants with pain and sometimes slightly limited range of active or passive movement of the glenohumeral joint together with pain or limited range of movement of the cervical spine, upper thoracic spine, or upper ribs. Both the synovial structures and the structures of the cervical spine, upper thoracic spine, or upper ribs could have caused these complaints. This group was combined with the synovial group because a previous study had shown that the course of complaints of the combination group and the synovial group was the same			
	Inclusion Criteria (not listed above)			
	• None			
	Exclusion Criteria (not listed above)			
	Treatment for the condition in the past 6 months			

• Bilateral shoulder complaints



Winters 1997 (Continued)

- Presence of a specific rheumatic disorder (rheumatoid arthritis, etc.)
- Complaint caused by acute, severe trauma such as fracture, dislocation or cuff rupture
- Herniated cervical disc
- Presence of dementia or other psychiatric disorder
- Refusal to participate

#### **Baseline characteristics**

#### Shoulder Girdle subgroup

#### Intervention

Number randomised: 29; Mean (SD) age: 46.4 (11.2) years old; Sex: F/M 18/11; Median duration of symptoms: 4 weeks

#### Control 1

Number randomised: 29; Mean (SD) age: 43.9 (12.6) years old; Sex: F/M 15/14; Median duration of symptoms: 3 weeks

#### Synovial subgroup

#### Intervention

Number randomised: 35; Mean (SD) age: 53.1 (12.6) years old; Sex: F/M 14/21; Median duration of symptoms: 4 weeks

#### Control 1

Number randomised: 32; Mean (SD) age: 46.7 (12.1) years old; Sex: F/M 17/15; Median duration of symptoms: 9 weeks

Control 2

Number randomised: 47; Mean (SD) age: 53.5 (12.5) years old; Sex: F/M 32/15; Median duration of symptoms: 8 weeks

Interventions

#### Intervention: physical therapy

*Components of intervention*: regimes of "classic" physiotherapy, possibly including physical applications, massage and exercise therapies. No mobilisation techniques or manipulative techniques were allowed

Dose: not reported

Frequency of administration: twice per week (number of weeks in total not reported)

#### **Control 1: manipulation**

*Components of intervention*: mobilisation and manipulation of the cervical spine, upper thoracic spine, upper ribs, acromioclavicular joint and the glenohumeral joint

Dose: not reported

Frequency of administration: once a week for a maximum of 6 sessions

#### **Control 2: glucocorticoid Injections**

*Components of intervention*: injection of triamcinolone acetonide and lignocaine. Two out of 3 synovial structures (glenohumeral joint capsule, subacromial space and acromioclavicular joint) were injected. The intra-articular injection was given from the posterior side, the subacromial injection from the lateral side and the acromioclavicular injection perpendicularly from the upper side of the joint

*Dose*: 1 ml of 40 mg/ml triamcinolone acetonide in combination with 9 ml of 10 mg/ml lignocaine for 1–3 injections

# Winters 1997 (Continued)

	Frequency of administration: after randomisation, 1 week later and another 2 weeks later if needed	
Outcomes	Outcomes assessed at 11 weeks and 2-3 years (only for treatment success)	
	<ul> <li>Pain: measured using the Shoulder Pain Score, ranging from 7 (no pain) to 28 (severe pain)</li> <li>Global assessment of treatment success: cure defined as disappearance of shoulder complaints or a decrease of shoulder complaints to such an extent that they were no longer inconvenient, did not need treatment, or no longer interfered with normal working</li> </ul>	
Notes	Conflicts of interest: the authors stated that they had no conflicts of interest	
	Only data for the synovial group were included in the review	

# **Risk of bias**

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "The university's Department of Family Practice was in charge of the randomisation to treatment. For each diagnostic category, we had made a series of closed unnumbered envelopes which contained instructions of the treatment to be given. The participating general practitioners had to call a sec- retary and state the diagnostic category of each patient. The secretary in turn would draw an envelope to assign treatment"
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Low risk	Comment: An adequate method was used to conceal the allocation sequence
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment, and may have had different expectations about the benefits of each intervention. The comparison of physiotherapy versus manipulation is of less concern
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Comment: Unblinded participants who may have had different expectations about the benefits of the intervention they received self-reported all out- comes. The comparison of physiotherapy versus manipulation is of less con- cern
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Quote: "Drop out because of treatment failure was significantly higher in the physio therapy group (45% (13/29) of patients) than in the manipulation group (20% (6/29) patients)."
		Quote: "Drop out because of treatment failure was much lower in the injection group (17% (7/47)) than in the physio therapy group (51% (18/35)) and manip- ulation group (59% (19/32))."
		Comment: All withdrawals were recorded and their reasons published in the article. An intention to treat analysis was performed but it is not clear how missing data were imputed
Selective reporting (re- porting bias)	Unclear risk	Comment: Outcome data were fully reported for all outcomes reported in the methods section of the publication, but without a trial protocol it is unclear whether other outcomes were measured but not reported based on the results
Other bias	Low risk	Comment: The study appears to be free of other bias

Manual therapy and exercise for rotator cuff disease (Review)

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Yiasemides 2011	
Methods	Study design: Parallel group RCT
	Setting: Metropolitan teaching hospital, Australia
	Intervention: Passive shoulder mobilisation and exercise and advice
	Control: Exercise and advice alone
	Source of Funding: Partially funded by Musculoskeletal Physiotherapy Australia research Grant (2005)
Participants	Diagnostic label used by trialists: None specified
	Criteria for defining the shoulder condition being treated
	<ul> <li>Painful active flexion or abduction shoulder movements</li> <li>Minimal shoulder movement restriction</li> </ul>
	<ul> <li>Pain, tenderness, or restriction during passive accessory movements at the glenohumeral, acromio- clavicular, or sternoclavicular joint or during passive scapular movements</li> </ul>
	Greater than 1 month duration
	Inclusion Criteria (not listed above)
	• None
	Exclusion Criteria (not listed above)
	Less than 18 years of age
	Unable to understand spoken English
	Shoulder symptoms were reproduced during active cervical spine movements or during palpation of cervical or thoracic region joints
	<ul> <li>They reported paraesthesia in the affected upper limb</li> </ul>
	Passive shoulder region joint mobilization was contraindicated
	• Shoulder flexion or abduction ROM was less than 140 degrees, as determined from digital pho- tographs
	Shoulder pain was due to an inflammatory or neoplastic disorder
	• They had had surgery or trauma to the shoulder in the previous 4 weeks
	They reported a feeling of shoulder instability
	Baseline characteristics
	Intervention
	Number randomised: 47; mean (range) age = 62 (35-85) years; sex: F/M 27/20; mean (SD) duration of symptoms: 9.7 (12) months
	Control
	Number randomised: 51; mean (range) age = 58 (27-81) years; sex: F/M 24/27; mean (SD) duration of symptoms: 22 (38) months
Interventions	Intervention: passive shoulder region joint mobilisation
	<i>Components of intervention:</i> individually tailored low-velocity passive joint mobilisations to any of the shoulder region joints and passive mobilisation of the scapula. Either sustained or oscillatory tech- niques were utilised
	Dose: individually determined
	<i>Frequency of administration:</i> as above (minimum of 60% of all treatments involved passive shoulder mobilisation)
	Control: participants only received the treatment common to both groups (see below)

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Yiasemides 2011 (Continued)	Both groups:		
	<i>Components of intervention:</i> both groups received advice and an individually tailored exercise pro- gramme. Advice included avoidance of painful shoulder movements and maintenance of normal scapulohumeral rhythm. Exercises included stretching, strengthening and motor retraining		
	Dose: not reported		
	<i>Frequency of administration:</i> daily home exercise performance. Participants attended therapy sessions for exercise technique revision and progression 1-2 times per week for 4 weeks, with a maximum of 12 treatment sessions over 8 weeks		
Outcomes	Outcomes assessed 1 month, 3 months and 6 months		
	<ul> <li>Function: SPADI total score rated 0-100 with a higher score indicating worse function</li> <li>Pain: SPADI pain subscore rated 0-100 with a higher score indicating worse pain</li> <li>Global assessment of treatment success: 6-point Likert scale rated 0-5 where higher score indicates better recover. Scores of 4 ("greatly improved") and 5 ("fully recovered") were taken to indicate treatment success</li> <li>Active ROM: flexion and abduction measured with a photographic method</li> <li>Adverse events</li> </ul>		
Notes	Conflicts of interest: not reported		
	Trial registration: ACTRN: 12605000151639		

#### **Risk of bias**

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "Random allocation of participants was performed using a previously determined treatment assignment schedule with random numbers generated from the data analysis function in Microsoft Excel."
		Comment: An adequate method was used to generate the allocation sequence
Allocation concealment (selection bias)	Low risk	Quote: "To ensure concealment, the randomization procedure was carried out by a researcher (K.A.G.) not involved in participant recruitment, treatment, or assessment, and the treatment assignment schedule was stored in consecu- tively numbered, sealed opaque envelopes."
		Comment: An adequate method was used to conceal the allocation sequence
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: Given the nature of the interventions, participants were not blind to treatment and may have had different expectations regarding the benefits of the interventions
Blinding of outcome as- sessment (detection bias) Self-reported outcomes	High risk	Quote: "Primary outcome measurements of pain, functional impairment, and self-rated improvement were obtained from participants who were not blind-ed to treatment group allocation"
		Comment: Unblinded participants who may have had different expectations regarding the benefits of the interventions received self-reported some out-comes
Blinding of outcome as- sessment (detection bias) Objective outcomes	Low risk	Quote: "Secondary outcome measurements of painful AROM were obtained by a researcher (R.Y.) blinded to group allocation at the same time points." Comment: Outcome assessor of objective outcomes was blind to treatment

Manual therapy and exercise for rotator cuff disease (Review)

Yiasemides 2011	(Continued)
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Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "All analyses were conducted using an intention-to-treat approach. Missing data (lost to follow-up) were replaced with values obtained by impu- tation using regression models within each variable and group at all avail- able time points. For the 2 control group participants who were lost prior to re- assessment at 1 month after recruitment and, therefore, did not have a self- rated change in symptoms score, the average of the group was used for their missing scores." Comment: Amount of drop out and reasons were similar between groups
Selective reporting (re- porting bias)	Low risk	Comment: Outcome data were fully reported for all outcomes specified in the trial registry entry
Other bias	Low risk	Comment: No other sources of bias identified

AROM = active range of motion; PROM = passive range of motion; ROM = range of motion; VAS = visual analogue scale

## Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion	
Bergman 2004	Ineligible clinical condition: did not exclude people with pain radiating to the neck region or to the lower part of the arm	
Bron 2011	Ineligible clinical condition: participants had non-specific shoulder pain and myofascial trigger points	
Chen 2009	Ineligible clinical condition: mixture of shoulder disorders, where some participants had concomi- tant neck pain, and the inclusion/exclusion criteria was more compatible with a diagnosis of adhe- sive capsulitis	
Geraets 2005	Ineligible clinical condition: did not exclude patients with pain radiating to the neck region or to the lower part of the arm (>50% participants had a concomitant neck problem)	
Ginn 1997	Ineligible clinical condition: mixed shoulder disorders, where patients with rotator cuff disorders comprised only 65% of the sample (remaining patients had osteoarthritis, adhesive capsulitis, biceps muscle tear or no diagnosis)	
Hakguder 2011	Not a randomised or quasi-randomised trial	
Jinhwa 2012	Not a randomised or quasi-randomised trial	
Krischak 2013	Ineligible intervention: occupational therapy is ineligible, and both groups received exercise	
Merolla 2013	Not a randomised or quasi-randomised trial	
Miller 2004	Ineligible clinical condition: inclusion/exclusion criteria not compatible with a diagnosis of rotator cuff disease, and unclear if participants with adhesive capsulitis, a history of significant trauma or systemic inflammatory conditions such as rheumatoid arthritis, osteoarthritis, hemiplegic shoul- ders, or pain in the shoulder region as part of a complex myofascial neck/shoulder/arm pain condi- tion were excluded	
Molsberger 2010	Ineligible intervention: compared physical therapy plus NSAID to acupuncture or sham acupunc- ture. Cannot separate the effect of physical therapy from NSAID	
Muth 2012	Not a randomised or quasi-randomised trial	

Manual therapy and exercise for rotator cuff disease (Review)

Study	Reason for exclusion
Mörl 2011	Ineligible clinical condition: inclusion/exclusion criteria not compatible with a diagnosis of rotator cuff disease, and unclear if participants with adhesive capsulitis, a history of significant trauma or systemic inflammatory conditions such as rheumatoid arthritis, osteoarthritis, hemiplegic shoul- ders, or pain in the shoulder region as part of a complex myofascial neck/shoulder/arm pain condi- tion were excluded
Saggini 2010	Ineligible intervention: examined the effect of a proprioceptive Multi Joint System device, not man- ual therapy or exercise
Seok-Hwa 2013	Ineligible clinical condition: inclusion/exclusion criteria not compatible with a diagnosis of rotator cuff disease, and unclear if participants with adhesive capsulitis, a history of significant trauma or systemic inflammatory conditions such as rheumatoid arthritis, osteoarthritis, hemiplegic shoul- ders, or pain in the shoulder region as part of a complex myofascial neck/shoulder/arm pain condi- tion were excluded
Tachibana 2012	Ineligible clinical condition: inclusion/exclusion criteria not compatible with a diagnosis of rotator cuff disease, and unclear if participants with adhesive capsulitis, a history of significant trauma or systemic inflammatory conditions such as rheumatoid arthritis, osteoarthritis, hemiplegic shoul- ders, or pain in the shoulder region as part of a complex myofascial neck/shoulder/arm pain condi- tion were excluded

#### **Characteristics of studies awaiting assessment** [ordered by study ID]

Acosta 2009	
Methods	Requires translation
Participants	
Interventions	
Outcomes	
Notes	

Bicer 2005	
Methods	Requires translation
Participants	
Interventions	
Outcomes	
Notes	

Bube 2010	
Methods	

Available only as a conference abstract

Manual therapy and exercise for rotator cuff disease (Review)

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Participants	
Interventions	
Outcomes	
Notes	

## Ellegaard 2013

Methods	Available only as a conference abstract
Participants	
Interventions	
Outcomes	
Notes	

#### Ginn 2009

Methods	Available only as a conference abstract	
Participants		
Interventions		
Outcomes		
Notes		

Just 2009	
Methods	Requires translation
Participants	
Interventions	
Outcomes	
Notes	

#### Leblebici 2007

Methods

**Requires translation** 

Manual therapy and exercise for rotator cuff disease (Review)



#### Leblebici 2007 (Continued)

Participants	
Interventions	
Outcomes	
Notes	

#### **Pribicevic 2006**

Methods	Available only as a conference abstract
Participants	
Interventions	
Outcomes	
Notes	

#### Werner 2002

Methods	Requires translation
Participants	
Interventions	
Outcomes	
Notes	

Wiener 2005	
Methods	Requires translation
Participants	
Interventions	
Outcomes	
Notes	

#### Wies 2008

Methods

Available only as a conference abstract

Manual therapy and exercise for rotator cuff disease (Review)



#### Wies 2008 (Continued)

Participants	
Interventions	
Outcomes	
Notes	

## Characteristics of ongoing studies [ordered by study ID]

#### Roddy 2014

Trial name or title	The SUPPORT trial (SUbacromial imPingement syndrome and Pain: a randomised controlled trial Of exeRcise and injecTion)	
Methods	Factorial RCT	
Participants	Inclusion criteria	
	<ul> <li>18 years and over</li> <li>No history of significant shoulder trauma, for example, fracture, clinically-suspected full thickness cuff tear</li> <li>A clinical diagnosis of subacromial impingement syndrome (pain in deltoid insertion area, positive Neer and Hawkins-Kennedy tests, pain on shoulder abduction)</li> </ul>	
	Exclusion criteria	
	<ul> <li>Below 18 years old</li> <li>Those whose main complaint is due to neck problems, acromioclavicular pathology, or other primary shoulder disorders including adhesive capsulitis or clinically-suspected full thickness cuff tear</li> </ul>	
	<ul> <li>Potentially serious pathology (inflammatory arthritis, polymyalgia rheumatica, malignancy etc) or ipsilateral shoulder surgery/replacement</li> </ul>	
	<ul> <li>Those already on a surgical waiting list for shoulder surgery</li> </ul>	
	<ul> <li>Contra-indications to local corticosteroid injection (known blood coagulation disorders, warfarin therapy)</li> <li>Participation in a shoulder-focused exercise programme or shoulder injection in the last month</li> <li>Inability to provide informed consent, complete written questionnaires, or read instruction leaflets written in English</li> </ul>	
Interventions	<ul> <li>Ultrasound-guided subacromial glucocorticoid injection and a physiotherapist-led exercise pro- gramme</li> <li>Ultrasound-guided subacromial glucocorticoid injection and an advice and exercise leaflet</li> <li>Blind subacromial glucocorticoid injection and a physiotherapist-led exercise programme</li> <li>Blind subacromial glucocorticoid injection and an advice and exercise leaflet</li> </ul>	
Outcomes	Outcomes assessed at 6 weeks, 6 months and 12 months	
	<ul> <li>Overall pain (0-10 NRS)</li> <li>Global change</li> <li>SPADI</li> <li>Effect of shoulder disability on typical everyday activities</li> <li>Pain at night</li> <li>Quality of life (Euro-QoL and SF-12)</li> <li>Health care utilisation</li> </ul>	

Manual therapy and exercise for rotator cuff disease (Review)



Roddy 2014 (Continued)

• Effect of shoulder problem on work

	Adverse events
Starting date	01/03/2011
Contact information	Edward Roddy, Keele University, UK. Email: e.roddy@keele.ac.uk
Notes	Trial registration number: ISRCTN42399123

#### Van den Dolder 2010

Trial name or title	Is soft tissue massage an effective treatment for mechanical shoulder pain?	
Methods	Parallel group RCT	
Participants	Inclusion criteria	
	<ul> <li>Aged between 18 and 80 years</li> <li>Referred to physical therapy for management of shoulder pain</li> <li>Able to understand spoken English</li> </ul>	
	Exclusion criteria	
	<ul> <li>Shoulder pain due to trauma in the previous 4 weeks</li> <li>Shoulder pain reproduced with any neck movement</li> <li>Shoulder pain due to serious pathology (e.g. neoplasm, acute inflammatory condition, recent/un-united fracture)</li> <li>Worker's compensation claim relating to the shoulder pain</li> </ul>	
Interventions	<ul><li>Soft tissue massage plus exercise</li><li>Exercise alone</li></ul>	
Outcomes	<ul> <li>Outcomes assessed at 5 weeks and 4 months</li> <li>Overall pain (Short Form McGill Pain Questionnaire)</li> <li>Patient Specific Disability Measure</li> <li>SPADI</li> </ul>	
	<ul><li>Percentage improvement in Pain Questionnaire</li><li>ROM: flexion, abduction, hand-behind-back distance</li></ul>	
Starting date	22/06/2007	
Contact information	Paul van den Dolder, University of Sydney, Sydney, Australia. Email: pvan0651@mail.usyd.edu.au	
Notes	Trial registration number: ACTRN12607000336482	

## ADDITIONAL TABLES

Study ID	Manual therapy compo- nent(s)	Exercise component(s)	Duration of session (min- utes)	Number of sessions per week	Number of weeks treat- ment
Ainsworth 2009	None	The participant was taught to start with a flexed elbow and to raise the arm to a vertical posi- tion. The participant was then taught to control the arm with sways in a 20-degree arc before elevating and lowering the arm using a weight of approximate- ly 0.75 kg. When the participant could carry out these activities supine, the head of the treatment couch was gradually inclined until they were able to perform the exercises in a sitting posi- tion. The participants also carried out stretching exercises to im- prove ranges of elevation, inter- nal and external rotation, resis- tance band exercises into inter- nal and external rotation, activi- ties to improve proprioception, posture correction and adapta- tion of functional activities	Not reported	Not reported	Not reported (assumed to be 6 weeks)
Al Dajah 2014	Soft tissue mobilisation for the subscapularis for 7 minutes and 5 repeti- tions of the contract-relax proprio- ceptive neuromuscular facilitation (PNF) tech- nique for the shoulder in- ternal rotator muscles followed by 5 repetitions of a PNF-facilitated abduction and external rotation diagonal pattern	None	10	1	1
Atkinson 2008	Manipulation (high veloc- ity, low-amplitude, gen- tle-impulse, shoulder ad- justive thrust based on extensive motion palpa- tion)	None	Not reported	3	2
Bae 2011	None	Motor control and strengthening exercises	30	3	4
Bang 2000	Passive accessory or pas- sive physiological joint mobilisation Maitland grades I-V; soft tissue massage and muscle stretching	Home exercises: simple cervical and thoracic postural exercises such as chin tucks, and self-mo- bilisation such as caudal glides of the glenohumeral joint	30	2	3

## Table 1. Characteristics of manual therapy and exercise interventions

Manual therapy and exercise for rotator cuff disease (Review)

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Table 1. Characteristics of manual therapy and exercise interventions (Continued)

Bansal 2011	Deep friction massage to supraspinatus tendon in a transverse direction	Codman's exercises consisting of pendulum or swinging motion of the arm in flexion, extension, hor- izontal abduction, adduction and circumduction. Intensity (arc of motion) was increased as tolerat- ed	10	7	10 days
Barbosa 2008	Front, back, lower lon- gitudinal and lateral re- laxations of the gleno- humeral joint, antero- posterior movements of the acromioclavicu- lar (squeeze) joint and anteroposterior, inferi- or-superior and superior- inferior movements of the sternoclavicular joint	Eccentric training exercises: the 'empty the can' movement (the participant performs abduction movements of the shoulder in the scapular plane, with medi- al rotation) when treating the supraspinatus muscle, or the 'right curl' movement (the partic- ipant flexes his elbow, with the arm abducted beside the body) when treating biceps brachii dys- functions. Movement resistance was offered manually, always by the same researcher and respect- ing the participant's pain limit	Not reported	3	4
Barra 2011	Diacutaneous fibrolysis: application of a metal- lic hook as deeply as possible following the intermuscular septum between the muscles of the cervico-scapular and shoulder region in a centripetal direction to- wards the pain location	None	15	1	1
Barra Lopez 2013	Diacutaneous fibrolysis: application of a metal- lic hook as deeply as possible following the intermuscular septum between the muscles of the cervico-scapular and shoulder region in a centripetal direction to- wards the pain location	No details provided	Not reported	2	3
Baskurt 2011	None	Intervention group: scapular pro- prioceptive neuromuscular facil- itation (PNF) exercises, scapular clock exercise, standing weight shift, double arm balancing, scapular depression, wall push up, wall slide exercises, strength- ening and stretching exercises Control group: strengthening and stretching exercises	Not reported	3	6

Manual therapy and exercise for rotator cuff disease (Review)

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Table 1. Characteristics of manual therapy and exercise interventions (Continued)

Beaudreuil 2011	Passive mobilisation of the shoulder with a pain- less ROM	Dynamic humeral centring: learn- ing the lowering of the humeral head during passive abduction of the shoulder, and actively lower- ing the humeral head by co-con- traction of the pectoralis major and latissimus dorsi during active abduction of the shoulder. Home exercise: no details provid-	Not reported	3 x wk 1-3; 2 x wk 4-6	6
Bennell 2010	Soft tissue massage, glenohumeral joint mo- bilisation, thoracic spine mobilisation, cervi- cal spine mobilisation, scapular retraining, pos- tural taping	ed Most exercises required the par- ticipant to incorporate their scapular retraining with strength- ening of the rotator cuff mus- cles. Some exercises reinforced and facilitated correct posture. Resistance for specific exercises was provided by hand weights or elastic theraband. Exercises were taught and performed dur- ing each treatment session and were otherwise self-administered at home	30 to 45	2 x wk 1-2; 1 x wk 3-6; fort- nightly wk 7-10	10
Biasoszewski 2011	Mobilisation of the glenohumeral joint and soft tissues using Kaltenborn's roll-glide techniques, Cyriax deep transverse massage, Mul- ligan's mobilisation with movement and typical techniques of joint mo- bilisation in the antero- posterior direction	Standard passive and active ex- ercises used to improve the ROM and restore muscle strength. The rotator cuff was initially strength- ened in the painless ROM by performing active, passive and self-assisted exercises. Once the full ROM has been achieved, strengthening exercises were ap- plied, ranging from flexion, ab- duction and external rotation to internal rotation adduction and extension	Not reported	Not reported	Not reported
Blume 2014	None	Supervised exercises: eccentric or concentric exercises included the seated 'full can', sidelying in- ternal rotation (IR), sidelying ex- ternal rotation (ER) with towel roll, supine protraction, sidelying horizontal abduction, sidelying abduction, and prone shoulder extension. All exercises were per- formed using a dumbbell for re- sistance. Home exercises: stretching and postural correction exercises	60	2	8
Brox 1993	None	Supervised exercises: ROM and strengthening exercises	60	2	24
Celik 2009b	None	Intervention group: supervised shoulder flexion below 90 de-	Not reported	7	2

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able 1. Chara	acteristics of manual ther	apy and exercise interventions grees, abduction, T-bar (wand)	(Continued)		
		exercises containing internal-ex- ternal rotation and extension, posterior capsule stretching and internal rotation exercises and rotator cuff strengthening exer- cises			
		Control group: supervised shoul- der flexion exercises above 90 de- grees, posterior and inferior cap- sule stretching exercises, rotator cuff strengthening and internal rotation exercises			
Citaker 2005	Manual mobilisation (de- tails not provided)	Theraband exercises permit- ting concentric and eccentric strengthening of the shoulder muscles. The exercises began with the elbow flexed 90 degrees and the shoulder in the neutral position. The exercises were per- formed through an arc of 45 de- grees in each of the 5 planes of motion. In addition, Codman pendulum exercises were utilised as a home programme	Not reported	Not reported	Not reported
Clews 1987	Massage of the long head of biceps, biceps tendon, pectorals, supraspinatus and infraspinatus	None	15	3	1
Cloke 2008	Manual therapy (details not provided)	Exercises (details not provided)	Not reported	6 sessions over 18 weeks	18
Conroy 1998	Mobilisation: depend- ing on the direction of re- striction in capsular ex- tensibility, inferior glide, posterior glide, anterior glide or long axis traction could be applied to the participant with oscil- latory pressure. Stretch could also be applied in the case of muscle spasm.	<ul> <li>Active ROM exercises (pain-free pendulum exercises and postural correction)</li> <li>Physiologic stretching: cane-assisted flexion and external rotation, towel-assisted internal rotation and non-involved arm-assisted horizontal abduction</li> <li>Muscle strengthening exercises: chair press, internal and external rotation isometrics</li> </ul>	15	3	3
Cook 2014	Grade III posterior-ante- rior mobilisations to the neck	Self- and externally-applied stretching, isotonic strengthen- ing, and restoration of normative movement	Not reported	Varied per participant	Mean of 8 (varied per participant)
Dickens 2005	Acromioclavicular joint, thoracic, cervical spine and glenohumeral joint mobilisation. The phys- iotherapist assessed the range of accessory	Exercises for the recruitment and strength of scapulothoracic mus- cles (especially lower trapezius and serratus anterior). The exer- cise programme was progressed to involve strengthening of infra-	Dependent on participant	Not reported	Not reported

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Table 1. Char	racteristics of manual ther movement available in each participant's gleno- humeral (anteroposte- rior, longitudinal cau- dad), acromioclavicu- lar (anteroposterior, lon- gitudinal caudad), cer- vical (posterior–anteri- or) and thoracic spine joints (posterior–anteri- or and transverse) with passive accessory move- ments. Any joints that were found to have re- stricted movement were addressed with mobilisa- tions into the direction of resistance and pain to help restore full pain-free range of movement	<b>apy and exercise interventions</b> spinatus, subscapularis and teres minor relative to the supraspina- tus and deltoid. The rotator cuff exercises were done with the use of resistance and participants were given Theraband for home use. The exercises started in neu- tral positions with isometric con- tractions and were progressed to inner range, through range, out- er range and into functional posi- tions. The resistance and speed of these exercises were altered and progressed	(Continued)		
Djordjevic 2012	Mobilisation with move- ment (MWM): during the MWM treatment, the par- ticipant was seated, and the therapist was po- sitioned on the oppo- site side of participant's painful shoulder. The therapist applied the thenar of one hand on the anterior aspect of the participant's humer- al head and the other hand on his/her scapula. The hand on the humer- al head performed a pos- terolateral glide, while the other hand stabilised the scapula. During this manoeuvre, the partici- pant was encouraged to perform active shoulder movement to the point of the first onset of pain	Pendulum exercises and pain- limited, active ROM exercises of shoulder elevation, depres- sion, flexion, abduction, rota- tions, and strengthening exercis- es. Strengthening exercises were isometric in nature, working on the external shoulder rotators, in- ternal rotators, biceps, deltoid, and scapular stabilizers (rhom- boids, trapezius, serratus ante- rior, latissimus dorsi, and pec- toralis major)	Not reported	7	10 days
Engebretsen 2009	None	Supervised exercises: the initial aim was to unload the stress on the rotator cuff and subacromi- al structures. During this phase, a mirror for awareness of pos- ture, an elastic rubber band and a sling fixed to the ceiling were used. The participants received immediate feedback and correc- tion (supervision) by the phys- iotherapist. Once dysfunction- al neuromuscular patterns were normalised, endurance exercis- es were performed with gradu-	45	2	12

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able I. Chara	cteristics of manual ther	apy and exercise interventions ally increasing resistance. Par- ticipants had an adjusted pro- gramme at home, which con- sisted of correction of alignment during daily living and simple low loaded exercises with a thin elas- tic cord to provide assistance and resistance to the movement	(Continued)		
Ginn 2005	Passive joint mobilisa- tion at the sternoclavicu- lar and acromioclavicular joints	ROM exercises: exercises were upgraded from active assisted to active to resisted active exercises using free weights or elastic resis- tance	Not reported	2	5
Giombini 2006	None	Supervised and home exercis- es, consisting of pendular swing- ing in the prone position in flex- ion and extension of the shoul- der and passive glenohumeral stretching exercises to tolerance	Not reported	1	4
Haahr 2005	Soft tissue treatments (details not provided)	Supervised exercises: active training of the periscapular mus- cles (rhomboid, serratus, trape- zoid, levator scapulae, pectoralis minor muscles) and strengthen- ing of the stabilising muscles of the shoulder joint (rotator cuff). This was done within the limits of pain	60	3 x wk 1-2; 2 x wk 3-5; 1 x wk 6-12; 2-3 x wk 13-19	12
Haik 2014	Low-amplitude, high velocity thrust thoracic spine manipulation	None	3	1	1
Hay 2003	Active and passive mobil- isation	Home exercise programme	20	1 to 2	6
Heredia-Rizo 2013	Manual therapy based on soft tissue techniques: micro-mobilisations of the cervical structures in all movement axes, relax- ation manoeuvres per- formed to fascial restric- tions involving the cervi- cal and scapulohumer- al region, and a reposi- tioning of the head of the humerus as recommend- ed by Kaltenborn	Supervised exercises: pendular movements using 1 kg of weight in prone, assisted active move- ments with a pulley, and propri- oceptive exercises with a ball in the horizontal plane	40	5	3
Holmgren 2012	None	Supervised exercises: two eccen- tric exercises for the rotator cuff (supraspinatus, infraspinatus, and teres minor), three concen- tric/eccentric exercises for the scapula stabilisers (middle and lower trapezius, rhomboideus,	30	1 x wk 1-2; fortnightly wk 3-12	12

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		and serratus anterior), and a pos- terior shoulder stretch			
Janse van Rensburg 2012	Thoracic spinal manipu- lation: a specific high ve- locity low amplitude "ex- tension with rotation" thrust manipulation was applied to the shoulder	Exercises to stimulate the lower fibres of trapezius and specific ro- tator cuff-strengthening	30	1	6
Kachingwe 2008	Intervention group: glenohumeral joint mo- bilisation was adminis- tered based on assess- ment of glenohumeral joint anterior, posterior and inferior glides and long-axis distraction pas- sive accessory motions using a 0-6 accessory motion scale. For situa- tions where there was re- activity within the cap- sular ROM, grade I-II mo- bilisation were applied. For situations where there was no reactivity but capsular hypomobili- ty, grade III-IV accessory motions were applied. Control group: gleno- humeral joint mobili- sation with movement (Mulligan technique) in- volved the therapist ap- plying a sustained pos- terior accessory glide to the glenohumeral joint while the subject simul- taneously actively flexed the shoulder to the pain- free endpoint and ap- plied a gentle overpres- sure force using the con- tralateral arm	Supervised exercises including posterior capsule stretching, pos- tural correction exercises, and an exercise programme focus- ing on rotator cuff strengthen- ing and scapular stabilisation. Participants were instructed to perform a home exercise pro- gramme mimicking the exercises performed in the clinic	Not reported	1	6
Kardouni 2014	Thoracic spinal manip- ulation: a high velocity, low amplitude thrust ap- plied to the lower tho- racic spine, mid thoracic spine, and cervicotho- racic junction	None	Not reported	1	1
Kassolik 2013	Classic massage of the shoulder girdle and glenohumeral joint was performed in a side re- cumbent position. Dur-	None	Not reported	5	2

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Table 1. Char	acteristics of manual ther ing the massage, typi- cal classic massage tech- niques (Swedish) were used - stroking with the palms (effleurage), fric- tion with the palms, kneading (petrissage), percussion (tapotte- ment), and vibration	apy and exercise interventions	(Continued)		
Kaya 2014	Scapular mobilisation (superoinferior gliding, rotations, and distrac- tions to the scapula), neuromuscular facili- tation techniques for scapula motions at ante- rior elevation-posterior depression and posteri- or elevation-anterior de- pression planes, gleno- humeral joint mobilisa- tion with long axis trac- tion and posterior or in- ferior glide techniques to improve shoulder inter- nal rotation limitations, and soft tissue massage and joint mobilisation of the neck, thoracic region, and elbow areas	Supervised and home exercises, including strengthening, flexibil- ity (ROM) and Codman's pendu- lum exercises	90	1	6
Kromer 2013	Painful and angular and/ or translatory restricted peripheral joints were treated with manual glide techniques accord- ing to the concept of Kaltenborn. Compara- ble signs of the spine segments were treated with posterior-anterior glides or coupled move- ments. Shortened mus- cles were stretched ac- cording to the descrip- tion of Evjenth & Ham- berg. Neural tissue was treated according to But- ler	Core exercise programme - dy- namic exercises started with 2 sets of 10 repetitions and with low resistance (yellow rub- ber band); shoulder and neck stretches were held for 10 sec- onds and repeated twice; isomet- ric scapular training positions were held for 10 seconds and re- peated twice	20 to 30	2 x wk 1-5; 3 x wk 6-12	12
Littlewood 2014	Manual therapy or mas- sage (no details provid- ed)	Self-managed loaded exercise: involved exercising the affected shoulder against gravity, a resis- tive therapeutic band or hand weight over 3 sets of 10 to 15 rep- etitions completed twice per day. The exercise was prescribed and operationalised within a self- managed framework which in-	Not reported	1	8

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		cluded focus upon knowledge translation, exercise/skill-acqui- sition, self-monitoring, goal-set- ting, problem-solving and pro-ac- tive follow-up.			
Lombardi 2008	None	Progressive resistance training programme. The exercises were flexion, extension, medial rota- tion and lateral rotation of the shoulder	Not reported	2	8
Ludewig 2003	None	Home exercise programme in- volving: two stretches (pec- toralis minor stretch and poste- rior shoulder stretch), a muscle relaxation exercise for the upper trapezius performed in front of a mirror, and progressive resis- tance strengthening exercises for two muscle groups (serratus an- terior muscle and humeral exter- nal rotation)	Not reported	7	10
Maenhout 2012	None	Eccentric exercise consisted of full can (thumb up) abduction in the scapular plane, which was performed with a dumbbell weight	Not reported	7	12
Martins 2012	None	Intervention group only: proprio- ception exercises: exercises with joint position, rhythmic stabili- sation and repositioning of the members, unstable base, propri- oceptive neuromuscular facilita- tion, and speed and accuracy	Not reported	2	6
		Both groups: pendulum exercis- es of the shoulder, stretching of the cervical spine and shoulder muscles, exercises with a stick (to maintain or improve ROM), exer- cises to strengthen the muscles of the rotator cuff and scapular stabilisers			
Marzetti 2014	None	Supervised exercises: neurocog- nitive exercises (intervention group) or strengthening exercis- es focused on the rotator cuff and scapular stabilising muscles, stretching exercises, Codman's pendulum exercises and exercis- es against elastic band resistance (control group)	60	3	5
McClatchie 2009	Lateral cervical glide mobilisations: the lat- eral aspect of the spin-	None	Not reported	1	1

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	ous processes of C5, C6, and C7 was landmarked on the ipsilateral side of the patient's painful shoulder. The examin- er's thumb remained on the lateral aspect of the spinous process of C5, with the opposite hand placed on the patient's non-affected shoulder or head for counterbalance as a lateral movement toward the non-painful side was applied with the mobilising hand					
Moosmayer 2014	None	Supervised exercises only, with particular attention directed towards correction of upper quarter posture and restoration of scapulothoracic and gleno- humeral muscular control and stability. Local glenohumeral control was addressed by exercis- es to centre the humeral head in the glenoid fossa. Isometric ex- ercises and exercises against ec- centric and concentric resistance for shoulder rotators were given. When local glenohumeral con- trol was achieved, exercises were given with increasing loads and progressed from neutral to more challenging positions	40	2 x wk 1-12; > 2 x wk 12-24	24	
Munday 2007	Shoulder girdle adjust- ments: high-velocity, low-amplitude manip- ulation in the direction of restricted end feel or joint play was performed. Participants sat in a com- fortable position with the shoulder girdle ex- posed. Adjustments to the acromioclavicular joint were most com- mon, although adjust- ments to the ribs, scapu- la and glenohumeral joints were made as well. The spine was not adjust- ed in this trial	None	None	3	3	
Osteras 2008	None	Supervised progressive resis- tance exercise therapy, compris- ing global aerobic exercises us- ing a stationary bike, a treadmill, or a step machine, and semiglob-	40	3	12	

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		al and local exercises using such medical exercise therapy equip- ment as wall pulley apparatus, lateral pulley apparatus, inclines board, angle bench, multiple pur- pose bench, shoulder rotator, dumbbells or barbells			
Rhon 2014	Combination of joint and soft-tissue mobilisa- tions; manual stretches; and contract–relax tech- niques	Supervised exercises: reinforcing exercises directed to the shoul- der girdle or thoracic or cervi- cal spine. Home exercises: wand ROM exercises, scapular retrac- tion, scapular protraction, tho- racic self-mobilisation, butterfly stretch	30	2	3
Senbursa 2007	Joint and soft tissue mo- bilisation: deep friction massage on supraspina- tus muscle tendon, ra- dial nerve stretching, scapular mobilisation, glenohumeral joint mo- bilisation, proprioceptive neuromuscular facilita- tion techniques includ- ing rhythmic stabilisation and hold-relax	An active ROM, stretching and strengthening exercise pro- gramme including rotator cuff muscles, rhomboids, levator scapulae and serratus anterior which was self-administered us- ing an elastic band at home after being taught by a physiotherapist	Not reported	3	4
Senbursa 2011	Deep friction massage on the supraspinatus mus- cle, radial nerve stretch- ing, scapular mobilisa- tion, glenohumeral joint mobilisation, and propri- oceptive neuromuscular facilitation techniques	ROM, stretching and strengthen- ing exercises for the rhomboid, levator scapulae, serratus ante- rior and rotator cuff muscles su- pervised and at home	Not reported	3	12
Struyf 2013	Manual mobilisations, stretching and motor control training of the scapula, including: pas- sive manual mobilisa- tion (to improve passive scapular upward rotation and posterior tilting)	Home stretching exercises for the levator scapulae, stretching of the pectoralis minor muscle length by the physiotherapist and scapular motor control training with emphasis on a scapular ori- entation exercise	30	1 to 3	3 to 9
Subasi 2012	None	Intervention group: supervised land-based exercises. For the first 10 days, ROM and stretching ex- ercises, and for the following 10 days, strengthening exercises.	Not reported	7	3
		Control group: supervised wa- ter-based exercises. For the first 10 days, ROM and stretching ex- ercises, and for the following 10 days, strengthening exercises in water by using dumbbells			

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Surenkok 2009	Scapular mobilisation: application of superior and inferior gliding, rota- tions, and distraction to the scapula of the affect- ed shoulder	None	Not reported	1	1
Szczurko 2009	Hands-on shoulder mus- cle and joint therapy	Standardised exercise pro- gramme consisting of passive, active assisted and active ROM muscle strengthening and joint therapy	30	1	12
Teys 2008	Postero-lateral glide (Mulligans' mobilisation with movement)	None	Not reported	1	1
Van den Dold- er 2003	Soft tissue massage of the shoulder performed as seen fit by the treating therapist. The areas fo- cused on were the lateral border of the scapula in full shoulder flexion; pos- terior deltoid at end of range horizontal flexion; anterior deltoid at end of range hand-behind-back; and pectoralis major in the stretch position.	None	15 to 20	3	2
Walther 2004	None	Intervention: standardised self- training programme of centring and stretching exercises that af- fected the shoulder. For most of the exercises, an elastic Thera- Band was used that was chosen according to the results of the ini- tial force measurements Control: physiotherapy consist- ing of centring training for the ro- tator cuff. Stretching was added in case of any limitation of the ROM at the first examination	10 to 15	5	12
Wang 2006	None	Intervention group: customised supervised and home self- stretching and strengthening ex- ercises for scapular stabilisers, rotator cuff and scapulohumeral muscles Control group: standardised strengthening exercises - shoul- der flexors, abductors, extensors, external rotators and internal ro- tators	Not reported	1	8

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shoulder region joints and passive mobilisation of the scapula. Either sustained or oscillatory techniques were de-

livered

Table I. Char	acteristics of manual ther	apy and exercise interventions	(Continuea)		
Winters 1997	Intervention group 1: massage (details not pro- vided), with no mobilisa- tion techniques or ma- nipulative techniques al- lowed	Supervised exercises (details not provided)	Not reported	1	6
	Intervention group 2: mobilisation and ma- nipulation of the cer- vical spine, upper tho- racic spine, upper ribs, acromioclavicular joint and the glenohumeral joint				
Yiasemides 2011	Individually tailored low- velocity passive joint mo- bilisations to any of the	Supervised and home exercises including stretching, strengthen- ing and motor retraining	Individually determined	1 to 2	8

#### Table 1. Characteristics of manual therapy and exercise interventions (Continued)

#### Table 2. Outcome matrix

Study ID	Overall pain	Function	Pain on motion	Global as- sessment	Quality of life	Adverse events
Ainsworth 2009		Х			Х	
Al Dajah 2014	Х					
Atkinson 2008	Х					Х
Bae 2011		Х				
Bang 2000	Х	Х				
Bansal 2011	Х					
Barbosa 2008		Х	х			
Barra 2011	Х			Х		Х
Barra Lopez 2013	Х	Х		Х		
Baskurt 2011	Х		х		х	
Beaudreuil 2011	Х	Х				
Bennell 2010	Х	Х	х	Х	Х	Х

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## Table 2. Outcome matrix (Continued)

Biasoszewski 2011	Х					
Blume 2014		х				
Brox 1993	Х	х	х	х		Х
Celik 2009	Х			х		
Citaker 2005	Х	х	х			
Clews 1987	Х					
Cloke 2008		х		х		
Conroy 1998	Х	х				
Cook 2014	Х	Х		х		Х
Dickens 2005		Х		·		
Djordjevic 2012						
Engebretsen 2009	Х	Х	Х	·		Х
Ginn 2005	Х	Х		Х		
Giombini 2006	Х	Х	Х	х		Х
Haahr 2005	Х	Х				
Haik 2014	Х					
Hay 2003	Х	Х		Х	Х	
Heredia-Rizo 2013		Х				
Holmgren 2012	Х	Х	Х	Х	Х	
Janse van Rensburg 2012		Х				Х
Kachingwe 2008	Х	Х				
Kardouni 2014	Х	Х			Х	
Kassolik 2013	Х					
Kaya 2014	Х	Х	Х			
Kromer 2013	Х	Х		Х		
Littlewood 2014		Х			Х	
Lombardi 2008	Х	Х	Х	Х	Х	
Ludewig 2003	Х	Х				

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#### Table 2. Outcome matrix (Continued)

Rhon 2014	Х	Х			Х	х
Osteras 2008	Х	Х				
Senbursa 2007	Х	х	х			
Senbursa 2011	Х	Х	Х			
Struyf 2013	Х	Х	Х			
Subasi 2012	Х	Х			х	
Surenkok 2009	Х	Х	Х			
Szczurko 2009	Х	Х			Х	Х
Teys 2008						х
Van den Dolder 2003	Х	Х				
Walther 2004	Х	Х	Х			Х
Wang 2006	X	Х				
Winters 1997	Х			Х		
Yiasemides 2011	Х	х		Х		X
FREQUENCY	48	44	16	17	13	17

## Table 3. Manual therapy and exercise versus placebo

## Study ID: Bennell 2010

Intervention: soft tissue massage, joint mobilisation, scapular retraining and supervised and home exercises Control: sham ultrasound

OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTI- MATE	
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)	
Overall pain (SPADI pain score 0-100) change from baseline to 22 weeks	24.8	23.7	59	17.3	19.6	61	6.8 (-0.7, 14.3)*	
Function (SPADI total score 0-100) change from baseline to 22 weeks	22.4	22	59	15.6	17.8	61	7.1 (0.3, 13.9)*	
Pain on movement (VAS 0-10) change from baseline to 22 weeks	2.6	2.9	59	1.6	2.4	61	0.9 (-0.03, 1.7)*	
Quality of life (AQoL -0.4 to 1) change from baseline to 22 weeks	0.07	0.2	59	0	0.1	61	0.07 (0.04, 0.1)*	
Quality of life (SF-36 PCS 0-100) change from baseline to 22 weeks	10.8	25	59	4.7	22.3	61	6.3 (-2, 14.5)*	
Quality of life (SF-36 MCS 0-100) change from baseline to 22 weeks	-1	19.7	59	1.8	15.8	61	0.6 (-5.2, 6.4)*	
Strength: abduction (kg) change from base- line to 22 weeks	1.1	4.4	59	0.4	2.5	61	1.2 (0.1, 2.3)*	
Strength: external rotation (kg) change from baseline to 22 weeks	0.3	4.3	59	-0.1	1.9	61	0.9 (-0.1, 1.9)*	
Strength: internal rotation (kg) change from baseline to 22 weeks	1.3	3.4	59	0	2.7	61	1.5 (0.4, 2.5)*	
	Events	Total		Events	Total		Risk ratio (95% Cl)	
Total adverse events during 11-week inter- vention period	17	55		5	61		3.77 (1.49, 9.54)	

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Adverse events: short-term pain during or af- ter the treatment session (during 11-week in- tervention period)		55		5	61		0.67 (0.17, 2.66)
Adverse events: increased short-term pain with the home exercises (during 11-week in- tervention period)	12	55		0	61		27.68 (1.68, 456.77)
Adverse events: mild irritation to the tape used for postural taping (during 11-week in- tervention period)	2	55		0	61		5.54 (0.27, 112.84
Total adverse events during 11-week fol- low-up period (i.e. from 11-22 weeks) (note the only adverse event was increased short- term pain with the home exercises)	7	49		0	58		17.70 (1.04, 302.29)
Global assessment of treatment success (suc cessful outcome ("much better") compared with those reporting an unsuccessful out- come (either "slightly better", "no change",	- 31	54		24	58		1.39 (0.94, 2.03)
"slightly worse", or "much worse")) at 22 weeks							
	ersus no tre and home ex	eatment vercises	outcome)	CONTROL			EFFECT ESTIMATE
*ANCOVA adjusted mean differences presented <b>Table 4. Manual therapy and exercise v</b> <b>Study ID: Dickens 2005</b> Intervention: mobilisation and supervised Control: advice to maintain normal activiti	ersus no tre and home ex es	eatment vercises	n outcome)	CONTROL Mean	Range	n	EFFECT ESTIMATI Mean difference (95% CI)

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# Table 4. Manual therapy and exercise versus no treatment (Continued)Intervention: glenohumeral mobilisation plus supervised and home exercisesControl: advice to regarding posture and overhead activities

OUTCOME	INTERVEN	TION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (VAS 0-10 with higher scores denoting worse pain) % change from base- line to 6 weeks	44.2	38.6	9	14.4	119.8	7	Not estimable
Function (SPADI total score 0-130 with higher scores denoting worse function) % change from baseline to 6 weeks	56.7	29.8	9	34.2	58.9	7	Not estimable
Active range of flexion % change from base- line to 6 weeks	-15.9	116.6	9	42.6	15.8	7	Not estimable
Study ID: Kachingwe 2008 Intervention: mobilisation with movement Control: advice to regarding posture and o			kercises				
OUTCOME	INTERVEN			CONTROL			EFFECT ESTIMAT

OUTCOME	INTERVENT	TION		CONTROL		EFFECT ESTIMATE		
	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)	
Overall pain (VAS 0-10 with higher scores denoting worse pain) % change from base- line to 6 weeks	55.2	31.9	9	14.4	119.8	7	Not estimable	
Function (SPADI total score 0-130 with higher scores denoting worse function) % change from baseline to 6 weeks	55.5	20.1	9	34.2	58.9	7	Not estimable	
Active range of flexion % change from base- line to 6 weeks	46.7	31.9	9	42.6	15.8	7	Not estimable	

## Table 5. Manual therapy and exercise versus glucocorticoid injection

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## Table 5. Manual therapy and exercise versus glucocorticoid injection (Continued)Intervention: manual therapy and exercise (no details provided)

Control: glucocorticoid injection

OUTCOME	INTERVENT	TION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Function (Oxford Shoulder Score (12-60) with a higher score indicating worse disabil- ity) at 18 weeks	27.73	16	22	29.81	13.4	27	-2.08 (-10.46, 6.30)
Function (Oxford Shoulder Score (12-60) with a higher score indicating worse disabil- ity) at 12 months	28.94	NR	NR	26.47	NR	NR	2.47 (95% CI not es timable)
Study ID: Ginn 2005 Intervention: mobilisation, range of motior Control: glucocorticoid injection			modalities				
ОИТСОМЕ	INTERVENT	TION		CONTROL		EFFECT ESTIMATI	
	Mean	95% CI	n	Mean	95% CI	n	Mean difference (95% Cl)
Pain (VAS 0-10) change from baseline to 5 weeks	1	0 - 2.5	39	0.2	0 - 1.7	45	0.80 (-1.26, 2.86)
Function (categorical rating scale, 0-27 with higher scores denoting worse function) change from baseline to 5 weeks	5.3	4.1 - 6.5	39	5.2	3.9 - 6.5	45	0.10 (-1.62, 1.82)
Active range of abduction (degrees) change from baseline to 5 weeks (subgroup with decreased ROM and shoulder pain)	97	81 to 113	?	98	82 to 114	?	-1 (95% CI not es- timable)
Active range of abduction (degrees) change from baseline to 5 weeks (subgroup with full ROM despite shoulder pain)	30	8 to 52	?	28	13 to 44	?	2 (95% Cl not es- timable)
Active range of flexion (degrees) change from baseline to 5 weeks (subgroup with	104	92 to 116	?	111	102 to 120	?	-7 (95% CI not es- timable)

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## Table 5. Manual therapy and exercise versus glucocorticoid injection (Continued)

Active range of flexion (degrees) change from baseline to 5 weeks (subgroup with full ROM despite shoulder pain)	1	0 to 14	?	0	0 to 8	?	1 (95% CI not es- timable)
Active hand-behind-back distance change from baseline to 5 weeks	7.3	4.7 to 10	39	7.5	4.9 to 10.2	45	-0.20 (-3.77, 3.37)
Strength (isometric abduction force %) change from baseline to 5 weeks	60	46 to 75	39	66	55 to 76	45	-6.00 (-23.27, 11.27)
	% range	Total		% range	Total		Risk ratio (95% CI)
Global assessment of treatment success (% participants rated as "improved") at 5 weeks	33 to 85%	39		35 to 78%	45		Not estimable

### Study ID: Hay 2003

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Intervention: physiotherapy (all participants: advice and instructions on pain relief and active shoulder exercises at home; dependent on participant: ultrasound and active and passive mobilisation)

Control: glucocort	ticoid injection
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OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMAT
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Function (Croft SDQ, 0-23 with higher scores denoting worse disability) change from baseline to 6 weeks	2.56	5.4	99	3.03	6.3	98	-0.47 (-2.11, 1.17)
Function (Croft SDQ, 0-23 with higher scores denoting worse disability) change from baseline to 6 months	5.97	5.4	99	4.55	5.9	97	1.42 (-0.16, 3.00)
	Median	IQR	n	Median	IQR	n	Mean difference (95% Cl)
Night pain (VAS 0-10) at 6 weeks	2	1 to 4	99	3	0 to 6	98	Not estimable
Night pain (VAS 0-10) at 6 months	1	0 to 3	99	1	0 to 4	97	Not estimable
Quality of life (EuroQoL, scored from -1 to 1) at 6 weeks	0.76	0.66 - 0.8	99	0.76	0.59 - 0.8	98	Not estimable

Quality of life (EuroQoL, scored from -1 to 1) at 6 months	0.76	0.69 - 0.88	99	0.76	0.66 - 1	97	Not estimable
	Events	Total		Events	Total		Risk ratio (95
Global assessment of treatment success (completely recovered) at 6 weeks	6	100		18	98		0.33 (0.14, 0.7
Global assessment of treatment success (completely recovered) at 6 months	23	99		17	97		1.33 (0.76, 2.3
Active range of abduction (restriction of > 50% compared with non-involved arm) at 6 weeks	40	99		53	98		0.75 (0.55, 1.0
Active range of abduction (restriction of > 50% compared with non-involved arm) at 6 months	31	99		38	97		0.80 (0.55, 1.1
Active range of external rotation (restriction of > 50% compared with non-involved arm) at 6 weeks	8	99		12	98		0.66 (0.28, 1.5
Active range of external rotation (restriction of > 50% compared with non-involved arm) at 6 months	7	99		8	97		0.86 (0.32, 2.2

#### Study ID: Rhon 2014

Intervention: joint and soft tissue mobilisations, manual stretches, contract-relax techniques, supervised exercises and home exercises Control: glucocorticoid injection

OUTCOME	INTERVENT	TION		CONTROL		EFFECT ESTIMATE	
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (NPRS 0-10) at 1 month	1.6	1.93	42	1.7	2.02	46	-0.10 (-0.92, 0.72)
Overall pain (NPRS 0-10) at 6 months	1.7	1.85	39	2.2	2.00	45	-0.50 (-1.32, 0.32)
Overall pain (NPRS 0-10) at 12 months	2.1	2.02	46	2.5	2.07	48	-0.40 (-1.23, 0.43)

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	Events	Total		Events	Total		Risk ratio (95% C
er score denotes worse pain) at 11 weeks							
Overall pain (shoulder pain score 0-28, high-	11.5	4.4	35	9.2	3.7	47	2.30 (0.50, 4.10)
	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)
OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMAT
Study ID: Winters 1997 Intervention: exercise and massage Control: glucocorticoid injection							
Adverse events	"Other than	transient pain fr	om the CSI [injec	tion], there were no	o other adverse e	vents reported b	y patients in either group
Quality of life (Global Rating of Change scale, -7 to +7) at 12 months	3	3.37	46	3	3.44	48	0.00 (-1.38, 1.38)
Quality of life (Global Rating of Change scale, -7 to +7) at 6 months	3	6.17	39	3	3.33	45	0.00 (-2.17, 2.17)
Quality of life (Global Rating of Change scale, -7 to +7) at 1 month	3	3.21	42	3	3.37	46	0.00 (-1.37, 1.37)
Function (SPADI total score, 0-100 where higher scores denote worse function) at 12 months	21.6	18.86	46	23.1	18.60	48	-1.50 (-9.07, 6.07)
Function (SPADI total score, 0-100 where higher scores denote worse function) at 6 months	21.5	17.89	39	22.2	18.64	45	-0.70 (-8.52, 7.12)
Function (SPADI total score, 0-100 where higher scores denote worse function) at 1 month	22.2	18.61	42	23.2	18.52	46	-1.00 (-8.77, 6.77)

\*Outcome data extracted from Figure using DigitizeIt software

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### Table 6. Manual therapy and exercise versus NSAID

## Study ID: Cloke 2008

Intervention: manual therapy and exercise (no details provided) Control: NSAIDs

OUTCOME	TION		CONTROL		EFFECT ESTIMATE		
	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)
Function (Oxford Shoulder Score (12-60) with a higher score indicating worse dis- ability) at 18 weeks	27.73	16	22	30.47	7	17	-2.74 (-10.21, 4.73)
Function (Oxford Shoulder Score (12-60) with a higher score indicating worse dis- ability) at 12 months	28.94	NR	NR	30.07	NR	NR	-1.13 (95% CI not es- timable)

## Table 7. Manual therapy and exercise versus arthroscopic subacromial decompression

#### Study ID: Haahr 2005

Intervention: exercises plus heat, cold packs or soft tissue treatment (i.e. not all participants received soft tissue treatment) Control: arthroscopic subacromial decompression

OUTCOME	INTERVENTION			CONTROL	CONTROL		
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (Constant-Murley pain sub-score, 0-15 with higher = less pain) change from baseline to 6 months	3.7	3.57	43	3.8	3.80	41	-0.10 (-1.68, 1.48)
Overall pain (Constant-Murley pain sub-score, 0-15 with higher = less pain) change from baseline to 12 months	3.7	3.25	43	3.6	4.12	41	0.10 (-1.49, 1.69)
Overall pain (VAS 0-9, 0 = no pain) change from baseline to 4-8 years	3	2.50	40	1.9	3.08	39	1.10 (-0.14, 2.34)

Function (Constant-Murley total score, 0-100 with higher = better function) change from baseline to 6 months	21.3	19.17	43	19.9	22.81	41	1.40 (-7.63, 10.4
Function (Constant-Murley total score, 0-100 with higher = better function) change from baseline to 12 months	23	19.82	43	18.8	23.13	41	4.20 (-5.03, 13.4
Function (total PRIM score 0-36, higher = worse function) change from baseline to 4-8 years	11.4	8.44	40	9.1	11.11	39	2.30 (-2.06, 6.66
Active ROM (Constant-Murley ROM sub-score, 0-40 with higher scores denoting better ROM) change from baseline to 6 months	10.3	10.40	43	9.6	10.77	41	0.70 (-3.83, 5.23
Active ROM (Constant-Murley ROM sub-score, 0-40 with higher scores denoting better ROM) change from baseline to 12 months	11.6	10.72	43	8.2	11.41	41	3.40 (-1.34, 8.14
Strength (Constant-Murley force sub-score, 0-25 with higher scores denoting better strength) change from baseline to 6 months	2.7	3.57	43	2.9	6.65	41	-0.20 (-2.50, 2.10
Strength (Constant-Murley force sub-score, 0-25 with higher scores denoting better strength) change from baseline to 12 months	3.2	4.87	43	3.3	6.97	41	-0.10 (-2.68, 2.4
	Events	Total		Events	Total		Risk ratio (95% Cl)
Global assessment of treatment success (re- covered or improved) at 4-8 years	27	40		23	39		1.14 (0.82, 1.61)
Work disability (self-reporting as currently working) at 4 to 8 years	21	40		20	39		1.02 (0.67, 1.57)

#### Study ID: Szczurko 2009

Intervention: physical exercise, hands on shoulder muscle and joint therapy and placebo tablets

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Table 8.	Manual therapy and exercise versus naturopathic care (Continued)

Control: naturopathic care (dietary counselling, acupuncture, and Phlogenzym supplement)

OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMAT
	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)
Overall pain (VAS 0-7) at 12 weeks	4.05	1.69	42	2.75	1.77	43	1.30 (0.56, 2.04)
Function (SPADI total score 0-130, with higher scores denoting worse function) at 12 weeks	56.24	36.57	42	35.3	31.57	43	20.94 (6.40, 35.48
Quality of life (SF-36 Mental Component Score, 0-100 with higher scores better) at 12 weeks	50.05	10.4	42	50.08	10.97	43	-0.03 (-4.57, 4.51)
Quality of life (SF-36 physical functioning score, 0-100 with higher scores better) at 12 weeks	61.29	22.4	42	74.02	25.2	43	-12.73 (-22.86, -2.60)
Quality of life (SF-36 role-physical score, 0-100 with higher scores better) at 12 weeks	61.61	22.85	42	72.71	24.68	43	-11.10 (-21.21, -0.99)
Quality of life (SF-36 bodily pain score, 0-100 with higher scores better) at 12 weeks	47.81	19.67	42	59.6	19.7	43	-11.79 (-20.16, -3.42)
Quality of life (SF-36 general health score, 0-100 with higher scores better) at 12 weeks	56.6	24.99	42	70.44	19.08	43	-13.84 (-23.31, -4.37)
Quality of life (SF-36 vitality score, 0-100 with higher scores better) at 12 weeks	56.08	18.21	42	63.72	20.55	43	-7.64 (-15.89, 0.6
Quality of life (SF-36 social functioning score, 0-100 with higher scores better) at 12 weeks	74.13	23.54	42	77.44	24.72	43	-3.31 (-13.57, 6.9
Quality of life (SF-36 role-emotional score, 0-100 with higher scores better) at 12 weeks	74.12	27.24	42	80.08	23.63	43	-5.96 (-16.81, 4.89
Quality of life (SF-36 mental health score, 0-100 with higher scores better) at 12 weeks	71.81	18.83	42	74.51	18.83	43	-2.70 (-10.71, 5.3

## Table 8. Manual therapy and exercise versus naturopathic care (Continued)

Adverse events	5	42		2	43		2.56 (0.53, 12.47)
	Events	Total		Events	Total		Risk ratio (95% CI)
Active range of adduction (degrees) at 12 weeks	36.28	11.05	42	35.39	7.42	43	0.89 (-3.12, 4.90)
Active range of extension (degrees) at 12 weeks	35.44	10.26	42	42.39	11.18	43	-6.95 (-11.51, -2.39)
Active range of flexion (degrees) at 12 weeks	121.08	40.53	42	159.39	25.97	43	-38.31 (-52.82, -23.80)
Active range of abduction (degrees) at 12 weeks	105.36	45.05	42	148.63	34.73	43	-43.27 (-60.40, -26.14)

## Table 9. Manual therapy and exercise and glucocorticoid injection versus glucocorticoid injection

## Study ID: Cloke 2008

Intervention: exercise and manual therapy package plus glucocorticoid injection

**Control: glucocorticoid injection** 

Ουτςομε	INTERVENT	ΓΙΟΝ		CONTROL		EFFECT ESTIMATE	
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Function (Oxford Shoulder Score (12-60) with a higher score indicating worse dis- ability) at 18 weeks	27.8	22.5	20	29.81	13.4	27	-2.01 (-13.09, 9.07)
Function (Oxford Shoulder Score (12-60) with a higher score indicating worse dis- ability) at 12 months	23.79	NR	NR	26.47	NR	NR	-2.68 (95% Cl not es- timable)

## Table 10. Manual therapy alone versus placebo

Study ID: Barra 2011

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Intervention: diacutaneous fibrolysis

## Table 10. Manual therapy alone versus placebo (Continued) Control: placebo diacutaneous fibrolysis

OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (VAS 0-100) change from baseline to immediately post 1 treatment session (i.e. same day)	9.2	15.2	25	7.5	13.7	25	1.70 (-6.32, 9.72)
Active range of abduction (degrees) change from baseline to immediately post 1 treatment session (i.e. same day)	7.9	9.6	25	0.6	8.7	25	7.30 (2.22, 12.38)
Active range of flexion (degrees) change from baseline to immediately post 1 treat- ment session (i.e. same day)	9.5	10.9	25	-1.9	9	25	11.40 (5.86, 16.94)
Active range of extension (degrees) change from baseline to immediately post L treatment session (i.e. same day)	2.1	5.7	25	0.2	6.4	25	1.90 (-1.46, 5.26)
Active range of external rotation (degrees) change from baseline to immediately post L treatment session (i.e. same day)	0.8	9.6	25	0.2	5.2	25	0.60 (-3.68, 4.88)
Active range of internal rotation (hand be- nind back distance in cm) change from paseline to immediately post 1 treatment session (i.e. same day)	4.5	6.8	25	1.4	3.1	25	3.10 (0.17, 6.03)
	Events	Total		Events	Total		Risk ratio (95% Cl
Global assessment of treatment success ("some or a lot of improvement") immedi- ately post 1 treatment session (i.e. same day)	15	25		7	25		2.14 (1.06, 4.34)
Total adverse events	Zero events	in both groups					
Study ID: Haik 2014 Intervention: thoracic spine manipulation Control: sham manipulation	n						

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## Table 10. Manual therapy alone versus placebo (Continued)

OUTCOME	INTERVENTION			CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Pain on motion (VAS 0-10) immediately post 1 treatment session (i.e. same day)	2.4	2.7	25	2.2	2.3	25	0.20 (-1.19, 1.59)
Study ID: Kardouni 2014 Intervention: thoracic spinal manipulativ Control: sham manipulative therapy	e therapy						
OUTCOME	INTERVENTION			CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (VAS 0-10, higher = more pain) at 1-2 days	2.4	1.6	24	2	1.5	21	0.40 (-0.51, 1.31)
Function (Penn Shoulder Score 0-100, higher = better function) at 1-2 days	80.6	11.1	24	83	9.8	21	-2.40 (-8.51, 3.71)
Quality of life (Global Rating of Change, from -7 (a great deal worse) to +7 (a great deal better)) at 1-2 days	1.3	2	24	2	2.2	21	-0.70 (-1.94, 0.54)
Study ID: McClatchie 2009 Intervention: lateral cervical glide mobili Control: placebo mobilisation	sation						
Ουτςομε	INTERVENTION			CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
				3.2	2.5	14	-0.80 (-2.83, 1.23)
Overall pain (VAS 0-10) at 1-4 days	2.4	2.1	7	5.2	2.5	11	0.00 ( 2.00, 2.20)

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### Table 10. Manual therapy alone versus placebo (Continued)Intervention: shoulder girdle adjustments (chiropractic)Control: placebo ultrasound

OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)
Overall pain (VAS 0-100) at 3 weeks	16.73	18.19	15	24	11.72	15	-7.27 (-18.22, 3.68)
Overall pain (VAS 0-100) at 7 weeks	10.73	18.72	15	19.83	12.26	15	-9.10 (-20.42, 2.22)
	Events	Total		Events	Total		Risk ratio (95% CI)
Adverse events				us adverse reactions orts of minor, tempo			uch as persistent severe stif
Study ID: Surenkok 2009 Intervention: scapular mobilisation Control: sham mobilisation							
ОИТСОМЕ	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)
Overall pain (VAS 0-100) immediately post-treatment (day 1)	35.84	21.62	13	20.15	24.7	13	15.69 (-2.15, 33.53)
Function (Constant-Murley total score 0-100, with higher scores denoting bet- ter function) immediately post-treatment (day 1)	54.3	5.9	13	51.84	6.51	13	2.46 (-2.32, 7.24)
Pain on motion (VAS 0-100) immediately post-treatment (day 1)	61.3	23.45	13	42.76	32.84	13	18.54 (-3.40, 40.48)
Active range of abduction (degrees) imme- diately post-treatment (day 1)	149.07	38.21	13	132.3	36.17	13	16.77 (-11.83, 45.37
Active range of flexion (degrees) immedi-	167.3	15.89	13	154.38	16.78	13	12.92 (0.36, 25.48)

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Study ID: Teys 2008

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## Table 10. Manual therapy alone versus placebo (Continued)Intervention: mobilisation with movementControl: sham mobilisation

OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Active ROM (elevation, degrees) immedi- ately post-treatment (day 1)	NR	NR	NR	NR	NR	NR	9.9 (4.3, 15.6)
Adverse events	0 events in t	ooth groups					
able 11. Manual therapy alone versus Study ID: Surenkok 2009 Intervention: scapular mobilisation Control: no treatment	s no treatme	ent					
DUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (VAS 0-100) immediately post-treatment (day 1)	35.84	21.62	13	44.8	38.6	13	-8.96 (-33.01, 15.09)
Function (Constant-Murley total score 0-100, with higher scores denoting bet- ter function) immediately post-treatment (day 1)	54.3	5.9	13	45.23	38.48	13	9.07 (-12.09, 30.23)
Pain on motion (VAS 0-100) immediately post-treatment (day 1)	61.3	23.45	13	63.38	21.81	13	-2.08 (-19.49, 15.33)
Active range of abduction (degrees) imme- diately post-treatment (day 1)	149.07	38.21	13	144.07	29	13	5.00 (-21.08, 31.08)

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#### Table 11. Manual therapy alone versus no treatment (Continued)

#### Study ID: Teys 2008

Intervention: mobilisation with movement

#### **Control: no treatment**

OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Active ROM (elevation, degrees) immedi- ately post-treatment (day 1)	NR	NR	NR	NR	NR	NR	11.4 (2.3, 20.5)
Adverse events	Zero events	in both groups					
Study ID: Van den Dolder 2003 Intervention: soft tissue massage Control: wait-list control							
OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (VAS 0-100 on Short-Form McGill Pain Questionnaire) at 2 weeks	31.8	26.4	15	53.8	26.3	14	-22.00 (-41.19, -2.81
Function (Patient Specific Functional Dis- ability Measure (PSFDM), 0 – 30 with high- er score denoting better function) at 2 weeks	17.6	8	15	10.4	5.6	14	7.20 (2.20, 12.20)
Active range of abduction (degrees) at 2 weeks	135.6	24.1	15	91.2	28.6	14	44.40 (25.08, 63.72)
Active range of flexion (degrees) at 2 weeks	129.5	18.5	15	103.4	23.1	14	26.10 (10.80, 41.40)
Active hand behind back distance (cm) at 2 weeks	19.9	10.2	15	8.1	16.2	14	11.80 (1.87, 21.73)

#### Table 12. Manual therapy alone versus another active intervention

#### Study ID: Al Dajah 2014

Intervention: soft tissue mobilisation plus proprioceptive neuromuscular facilitation Control: therapeutic ultrasound

ОИТСОМЕ	INTERVENTION					EFFECT ESTIMATE	
	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)
Overall pain (VAS 0-10) immediately after one treatment session (day 1)	3.8	0.79	15	5.23	0.72	15	-1.43 (-1.97, -0.89)
External rotation (degrees, unclear if active or passive) immediately after one treatment session (day 1)	52.4	4.9	15	40.33	5.6	15	12.07 (8.30, 15.84)

#### Study ID: Bansal 2011

Intervention: deep friction massage technique plus Codman's exercises

Control: ultrasound therapy plus Codman's exercises

OUTCOME	INTERVENT	ION		CONTROL		EFFECT ESTIMATE	
	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)
Overall pain (VAS 0-10) at 10 days	1.4	NR	20	2.1	NR	20	-0.7 (95% Cl not es- timable)
Active range of abduction (degrees) at 10 days	107.15	NR	20	105.65	NR	20	1.5 (95% CI not estimable)

#### Study ID: Kaya 2014

Intervention: manual therapy and exercise plus cold pack

Control: kinesiotaping plus exercise plus cold pack

OUTCOME	UTCOME INTERVENTION						EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)
Rest pain (VAS 0-10) at 6 weeks	1.5	2.28	26	1.82	2.05	28	-0.32 (-1.48, 0.84)
Function (DASH 0-100, higher = worse function) at 6 weeks	35.61	15.66	26	38.71	15.41	28	-3.10 (-11.40, 5.20)
Pain on motion (VAS 0-10) at 6 weeks	5.11	2.68	26	3.92	1.71	28	1.19 (-0.02, 2.40)

Night pain (VAS 0-10) at 6 weeks	3.19	3.28	26	1.28	1.88	28	1.91 (0.47, 3.35)
Study ID: Winters 1997 Intervention: manipulation Control: glucocorticoid injection							
ОИТСОМЕ	INTERVENTION			CONTROL			EFFECT ESTIMATE
-	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)
Overall pain (shoulder pain score D-28, higher score denotes worse pain) at 11 weeks	12.6	5.1	32	9.2	3.7	47	3.40 (1.34, 5.46)
	Events	Total		Events	Total		Risk ratio (95% CI)
Global assessment of treatment suc-	14	32		42	47		0.49 (0.33, 0.73)
ble 13. Manual therapy alone as		nother physica	ll therapy inte	ervention versus	the other phy	sical therapy i	intervention
able 13. Manual therapy alone as Study ID: Atkinson 2008 Intervention: manipulation plus mob Control: sham laser plus mobilisation	ilisation		l therapy inte			sical therapy i	
cess ("cured") at 11 weeks able 13. Manual therapy alone as Study ID: Atkinson 2008 Intervention: manipulation plus mob Control: sham laser plus mobilisation OUTCOME	ilisation INTERVE	NTION		CONTRO	DL		EFFECT ESTIMATE
able 13. Manual therapy alone as Study ID: Atkinson 2008 Intervention: manipulation plus mob Control: sham laser plus mobilisation	ilisation		I therapy inte			sical therapy i	
able 13. Manual therapy alone as Study ID: Atkinson 2008 Intervention: manipulation plus mob Control: sham laser plus mobilisation OUTCOME	ilisation INTERVE	NTION		CONTRO	DL		EFFECT ESTIMATE Mean difference
able 13. Manual therapy alone as Study ID: Atkinson 2008 Intervention: manipulation plus mob Control: sham laser plus mobilisation DUTCOME Overall pain (NRS 0-100) at 2 weeks Range of abduction (degrees, unclear if	ilisation INTERVE Mean 23.2	NTION	n	CONTRO Mean	DL	n	EFFECT ESTIMATE Mean difference (95% Cl)
able 13. Manual therapy alone as Study ID: Atkinson 2008 Intervention: manipulation plus mob Control: sham laser plus mobilisation	ilisation INTERVEI Mean 23.2 ac- 157.2	NTION SD 15.5	n 30	CONTRO Mean 30.1	DL SD 17.4	n 30	EFFECT ESTIMATE Mean difference (95% CI) -6.90 (-15.24, 1.44)

Range of adduction (degrees, unclear if ac- tive or passive) at 2 weeks	66.2	12.2	30	64.5	12.6	30	1.70 (-4.58, 7.98)
Range of external rotation (degrees, unclear if active or passive) at 2 weeks	75.5	16.6	30	72.9	17.9	30	2.60 (-6.14, 11.34)
Range of internal rotation (degrees, unclear if active or passive) at 2 weeks	58.3	9.1	30	57.4	10.6	30	0.90 (-4.10, 5.90)
Total adverse events	Zero events	in both groups					
Study ID: Bang 2000 Intervention: manual physical therapy plus Control: supervised flexibility and strength			engthening exe	ercises			
OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (composite of five 0-100mm VAS pain scores, total score range 0-500) at 8 weeks	174.41	183.06	27	360.64	272.32	22	-186.23 (-319.33, -53.13)
Function (Functional Assessment Question- naire 0-45, with higher scores denoting bet- ter function) at 8 weeks	38.22	4.68	27	33.26	7.84	23	4.96 (1.30, 8.62)
Pain on motion (active abduction, 0-100mm VAS) at 8 weeks	16.82	21.02	27	37.54	29.01	23	-20.72 (-34.98, -6.46)
Resisted abduction pain (0-100mm VAS) at 8 weeks	22.7	26.27	27	32.64	29.45	23	-9.94 (-25.53, 5.65)
Resisted external rotation pain (0-100mm VAS) at 8 weeks	15.85	21.92	27	30.23	29.72	23	-14.38 (-29.07, 0.31
Resisted internal rotation pain (0-100mm VAS) at 8 weeks	21.04	27.97	27	33.5	27.57	23	-12.46 (-27.90, 2.98
Isometric abduction strength (Newtons) at	225.3	111.86	27	147.14	81.11	23	78.16 (24.50,

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Isometric external strength (Newtons) at 8 weeks	159.05	77.83	27	101.88	42.06	23	57.17 (23.15, 91.19
Isometric internal strength (Newtons) at 8 weeks	191.96	82.29	27	153.62	58.63	23	38.34 (-0.87, 77.55)
Study ID: Barbosa 2008 Intervention: mobilisation plus eccentric n Control: eccentric muscle training plus the			tic ultrasound				
OUTCOME	INTERVENT	ΓΙΟΝ		CONTROL			EFFECT ESTIMATI
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Function (Constant-Murley total score, 0-100) at end of 4 weeks treatment	84.43	6.97	14	74.14	5.18	14	10.29 (5.74, 14.84)
_							
Study ID: Barra Lopez 2013 Intervention: diacutaneous fibrolysis plus Control: placebo diacutaneous fibrolysis p OUTCOME		sed physiothera	ру	CONTROL			EFFECT ESTIMATI
Intervention: diacutaneous fibrolysis plus Control: placebo diacutaneous fibrolysis p	lus standardi	sed physiothera	py n	CONTROL Mean	SD	n	EFFECT ESTIMATI Mean difference (95% CI)
Intervention: diacutaneous fibrolysis plus Control: placebo diacutaneous fibrolysis p OUTCOME Overall pain (VAS 0-100) change from base-	IUS Standardi	sed physiothera			<b>SD</b> 28.9	<b>n</b> 40	Mean difference
Intervention: diacutaneous fibrolysis plus Control: placebo diacutaneous fibrolysis p	lus standardi INTERVENT Mean	sed physiothera	n	Mean			Mean difference (95% Cl)
Intervention: diacutaneous fibrolysis plus Control: placebo diacutaneous fibrolysis p OUTCOME Overall pain (VAS 0-100) change from base- line to 3 weeks Overall pain (VAS 0-100) change from base-	INTERVENT Mean 22.5	SD 19.3	<b>n</b> 40	<b>Mean</b> 18.9	28.9	40	Mean difference (95% CI) 3.60 (-7.17, 14.37)
Intervention: diacutaneous fibrolysis plus Control: placebo diacutaneous fibrolysis p OUTCOME Overall pain (VAS 0-100) change from base- line to 3 weeks Overall pain (VAS 0-100) change from base- line to 3 months Function (Constant-Murley score 0-100)	INTERVENT Mean 22.5 28.6	SP 19.3 24.1	<b>n</b> 40 40	Mean 18.9 27.1	28.9 32	40	Mean difference (95% CI) 3.60 (-7.17, 14.37) 1.50 (-10.91, 13.91

	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)
OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMAT
Study ID: Barra Lopez 2013 Intervention: diacutaneous fibrolysis plus s Control: standardised physiotherapy	standardised	physiotherapy					
Global assessment of treatment success self-reported as "better" or "much better") at 3 weeks	33	37		28	37		1.18 (0.95, 1.46)
	Events	Total		Events	Total		Risk ratio (95% C
Active range of internal rotation (hand be- hind back distance in cm) change from baseline to 3 months	3.5	8.3	40	2.7	7.8	40	0.80 (-2.73, 4.33)
Active range of internal rotation (hand be- hind back distance in cm) change from baseline to 3 weeks	2.7	5.9	40	2.2	5.1	40	0.50 (-1.92, 2.92)
Active range of external rotation (degrees) change from baseline to 3 months	6.1	11.4	40	5.1	12.4	40	1.00 (-4.22, 6.22)
Active range of external rotation (degrees) change from baseline to 3 weeks	5.9	9.5	40	4.7	12.2	40	1.20 (-3.59, 5.99)
Active range of extension (degrees) change from baseline to 3 months	6.3	9.5	40	4.4	8.2	40	1.90 (-1.99, 5.79)
Active range of extension (degrees) change from baseline to 3 weeks	5.7	9	40	3.3	8.3	40	2.40 (-1.39, 6.19)
Active range of flexion (degrees) change from baseline to 3 months	16.5	21.5	40	12.1	19.9	40	4.40 (-4.68, 13.48)
Active range of flexion (degrees) change from baseline to 3 weeks	12.3	17.5	40	8.3	18.3	40	4.00 (-3.85, 11.85)
from baseline to 3 months							

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Overall pain (VAS 0-100) change from base- line to 3 weeks	22.5	19.3	40	15.1	24.6	40	7.40 (-2.29, 17.09)
Overall pain (VAS 0-100) change from base- line to 3 months	28.6	24.1	40	22.7	26.6	40	5.90 (-5.22, 17.02)
Function (Constant-Murley score 0-100) change from baseline to 3 weeks	9.9	8.5	40	4.2	7.8	40	5.70 (2.12, 9.28)
Function (Constant-Murley score 0-100) change from baseline to 3 months	12.4	11.7	40	9.9	8.9	40	2.50 (-2.06, 7.06)
Active range of abduction (degrees) change from baseline to 3 weeks	12	21.2	40	4.5	22.5	40	7.50 (-2.08, 17.08)
Active range of abduction (degrees) change from baseline to 3 months	14.3	28.1	40	14.6	24.9	40	-0.30 (-11.94, 11.34
Active range of flexion (degrees) change from baseline to 3 weeks	12.3	17.5	40	2.5	19.4	40	9.80 (1.70, 17.90)
Active range of flexion (degrees) change from baseline to 3 months	16.5	21.5	40	14.8	21.1	40	1.70 (-7.64, 11.04)
Active range of extension (degrees) change from baseline to 3 weeks	5.7	9	40	-1.3	7.3	40	7.00 (3.41, 10.59)
Active range of extension (degrees) change from baseline to 3 months	6.3	9.5	40	0.6	8.3	40	5.70 (1.79, 9.61)
Active range of external rotation (degrees) change from baseline to 3 weeks	5.9	9.5	40	-0.6	9.2	40	6.50 (2.40, 10.60)
Active range of external rotation (degrees) change from baseline to 3 months	6.1	11.4	40	0	12	40	6.10 (0.97, 11.23)
Active range of internal rotation (hand be- hind back distance in cm) change from baseline to 3 weeks	2.7	5.9	40	0.9	4.9	40	1.80 (-0.58, 4.18)
Active range of internal rotation (hand be- hind back distance in cm) change from baseline to 3 months	3.5	8.3	40	2.7	6.6	40	0.80 (-2.49, 4.09)

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#### Table 13. Manual therapy alone as an add-on to another physical therapy intervention versus the other physical therapy intervention (Continued)

ualth		Events	Total	Events	Total	Risk ratio (95% CI)
erapy and	Global assessment of treatment success self-reported as "better" or "much better") at 3 weeks	33	37	26	38	1.30 (1.02, 1.66)

#### Study ID: Biasoszewski 2011

#### Intervention: Manual therapy plus TENS plus ultrasound plus exercise

#### Control: TENS plus ultrasound plus exercise

OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (VAS 0-10) change from base- line to end of 4 treatment sessions (exact timing unclear)	5.27	2.28	15	3.2	1.32	15	2.07 (0.74, 3.40)
Active range of abduction (degrees) at the end of 4 treatment sessions (exact timing unclear)	147	37.93	15	130	26.19	15	17.00 (-6.33, 40.33)
Active range of flexion (degrees) at the end of 4 treatment sessions (exact timing un- clear)	156.67	93.4	15	143	23.74	15	13.67 (-35.10, 62.44)
Active range of external rotation (degrees) at the end of 4 treatment sessions (exact timing unclear)	50.67	8.63	15	40.33	8.55	15	10.34 (4.19, 16.49)
Active range of internal rotation (degrees) at the end of 4 treatment sessions (exact timing unclear)	61.67	13.18	15	54.67	13.43	15	7.00 (-2.52, 16.52)
Study ID: Clews 1987 Intervention: massage plus ice Control: therapeutic ultrasound plus ice							
OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATI
	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)

Pain after strength test (VAS 0-10) at 3 days	2.8	1.2	6	3.2	1.2	6	-0.40 (-1.76, 0.96)
Strength (maximal isometric force produc- tion, measured in peak force) % change from baseline to 3 days	9.8	8.8	6	11	9.5	6	-1.20 (-11.56, 9.16)
Study ID: Clews 1987 Intervention: massage plus ice Control: sham ultrasound plus ice							
ОИТСОМЕ	INTERVENT	TION		CONTROL			EFFECT ESTIMAT
	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)
Pain after strength test (VAS 0-10 at strength testing) at 3 days	2.8	1.2	6	2.7	1.9	6	0.10 (-1.70, 1.90)
Strength (maximal isometric force produc-	9.8	8.8	6	-1.5	9	6	11.30 (1.23, 21.37)
tion, measured in peak force) % change from baseline to 3 days							
from baseline to 3 days Study ID: Conroy 1998 Intervention: mobilisation plus standardise Control: standardised physiotherapy				CONTROL			EFFECT ESTIMAT
from baseline to 3 days Study ID: Conroy 1998 Intervention: mobilisation plus standardise	INTERVENT	TION		CONTROL	50		EFFECT ESTIMATI
from baseline to 3 days Study ID: Conroy 1998 Intervention: mobilisation plus standardise Control: standardised physiotherapy			n	CONTROL Mean	SD	n	EFFECT ESTIMAT Mean difference (95% Cl)
from baseline to 3 days Study ID: Conroy 1998 Intervention: mobilisation plus standardise Control: standardised physiotherapy	INTERVENT	TION	<b>n</b> 7		<b>SD</b> 33.26	<b>n</b> 7	Mean difference
from baseline to 3 days Study ID: Conroy 1998 Intervention: mobilisation plus standardise Control: standardised physiotherapy OUTCOME	INTERVENT Mean	SD		Mean			Mean difference (95% Cl) -33.36 (-60.37,
from baseline to 3 days Study ID: Conroy 1998 Intervention: mobilisation plus standardise Control: standardised physiotherapy OUTCOME Overall pain (VAS 0-100) at 3 weeks Active range of abduction (degrees) at 3	INTERVENT Mean 12.5	<b>SD</b> 14.93	7	<b>Mean</b> 45.86	33.26	7	Mean difference (95% CI) -33.36 (-60.37, -6.35)

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#### Table 13. Manual therapy alone as an add-on to another physical therapy intervention versus the other physical therapy intervention (Continued)

	Events	Total		Events	Total		Risk ratio (95% CI
Function: number of participants who can reach to external occipital protuberance at 3 weeks	4	7		5	7		0.80 (0.36, 1.77)
Function: number of participants who can reach overhead 135 degrees at 3 weeks	5	7		5	7		1.00 (0.52, 1.94)
Function: number of participants who can reach to the spinous processes at 3 weeks	2	7		2	7		1.00 (0.19, 5.24)
Study ID: Cook 2014 Intervention: neck manual therapy plus sta Control: standardised physiotherapy	andardised ph	ysiotherapy					
OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (numerical rating scale 0-10) at discharge (mean of 56 (SD 55) days)	2.3	1.8	36	2.2	1.2	32	0.10 (-0.62, 0.82)
Function (QuickDASH Questionnaire, 1-5 where higher scores denote worse dysfunc- tion) at discharge (mean of 56 (SD 55) days)	13.6	10.5	36	13.6	6.6	32	0.00 (-4.12, 4.12)
Total adverse events	Zero events	in both groups					
Study ID: Janse van Rensburg 2012 Intervention: thoracic spinal manipulation Control: mobilisation plus exercises	plus mobilisa	tion plus exerc	ises				
OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Function (DASH, 0-100 with higher scores	11.92	6.48	6	20.35	12.37	2	-8.43 (-26.34, 9.48)

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Range of abduction (degrees, unclear if ac- tive or passive) at 6 weeks	142.17	12.73	6	130	14.14	2	12.17 (-9.92, 34.26)
Range of flexion (degrees, unclear if active or passive) at 6 weeks	142.33	7.31	6	135	0	2	7.33 (95% CI not es timable)
Adverse events	Zero events	in both groups					
Study ID: Kachingwe 2008 Intervention: glenohumeral mobilisation p Control: exercises	lus exercises						
OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (VAS 0-10 with higher scores denoting worse pain) percent change from baseline to 6 weeks	44.2	38.6	9	20.8	112.3	8	Not estimable
Function (SPADI total score 0-130 with high- er scores denoting worse function) percent change from baseline to 6 weeks	56.7	29.8	9	61.6	35.9	8	Not estimable
Active range of flexion percent change from baseline to 6 weeks	-15.9	116.6	9	27.6	41.7	8	Not estimable
Study ID: Kachingwe 2008 Intervention: mobilisation with movement Control: exercises	plus exercise	25					
OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (VAS 0-10 with higher scores denoting worse pain) percent change from baseline to 6 weeks	55.2	31.9	9	20.8	112.3	8	Not estimable

Function (SPADI total score 0-130 with high- er scores denoting worse function) percent change from baseline to 6 weeks	55.5	20.1	9	61.6	35.9	8	Not estimable
Active range of flexion percent change from baseline to 6 weeks	46.7	31.9	9	27.6	41.7	8	Not estimable
Study ID: Kromer 2013 Intervention: manual therapy and exercise Control: exercises	S						
ОИТСОМЕ	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (VAS 0-10) at 5 weeks	2.9	1.6	46	3.3	1.6	44	-0.40 (-1.06, 0.26)
Overall pain (VAS 0-10) at 12 weeks	2.3	1.8	46	2.3	1.8	44	0.00 (-0.74, 0.74)
Overall pain (SPADI pain sub-score 0-100) at 1 year	17.7	21.8	44	12.4	16.9	43	5.30 (-2.89, 13.49)
Function (SPADI total score, 0-100 with higher scores denoting worse function) at 5 weeks	23.5	17.5	46	26.8	17.8	44	-3.30 (-10.60, 4.00)
Function (SPADI total score, 0-100 with higher scores denoting worse function) at 12 weeks	16.1	17.2	46	19.8	19.5	44	-3.70 (-11.31, 3.91)
Function (SPADI total score, 0-100 with higher scores denoting worse function) at 1 year	15.3	20.3	44	10.2	15.2	43	5.10 (-2.42, 12.62)
	Events	Total		Events	Total		Risk ratio (95% CI
Global assessment of treatment success ("much better" on PGIC) at 5 weeks	22	46		20	44		1.05 (0.68, 1.64)
Global assessment of treatment success ("much better" on PGIC) at 12 weeks	"No differen	ce between grou	ups"				

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#### Table 13. Manual therapy alone as an add-on to another physical therapy intervention versus the other physical therapy intervention (Continued)

Global assessment of treatment success ("much better" on PGIC) at 1 year	"No differenc	e between groups"			
Work disability (on sick leave) between 0 and 5 weeks	1	40	5	38	0.19 (0.02, 1.55)
Work disability (on sick leave) at 6 and 12 weeks	1	38	3	35	0.31 (0.03, 2.82)
Work disability (on sick leave) at 12 weeks and 1 year	2	38	3	37	0.65 (0.11, 3.67)
Adverse events during 1 year trial period	"One patient der."	had a 12-point deterioratio	on and another patient a 3	8-point deterioration afte	r an accident involving the shoul-
Study ID: Senbursa 2011 Intervention: joint and soft tissue mobilisa Control: supervised exercises	tion plus super	vised exercises			
ОИТСОМЕ	INTERVENTI	N	CONTROL		EFFECT ESTIMATE
	% Events	Total	% Events	Total	Risk ratio (95% CI)
Rest pain (number of participants with no pain as measured on VAS 0-10) at 4 weeks	83%	Unclear	64%	Unclear	Not estimable
Rest pain (number of participants with no pain as measured on VAS 0-10) at 12 weeks	97%	Unclear	92%	Unclear	Not estimable
	47%	Unclear	36%	Unclear	Not estimable
Night pain (number of participants with no pain as measured on VAS 0-10) at 4 weeks					
	83%	Unclear	88%	Unclear	Not estimable
pain as measured on VAS 0-10) at 4 weeks Night pain (number of participants with no	83%	Unclear Unclear	88%	Unclear Unclear	Not estimable Not estimable

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#### Table 13. Manual therapy alone as an add-on to another physical therapy intervention versus the other physical therapy intervention (Continued)

#### Study ID: Senbursa 2011 Intervention: joint and soft tissue mobilisation plus supervised exercises Control: home exercises

CONTROL OUTCOME INTERVENTION **EFFECT ESTIMATE** Total % Events Total Risk ratio (95% CI) % Events Rest pain (number of participants with no 83% Unclear 82% Unclear Not estimable pain as measured on VAS 0-10) at 4 weeks Rest pain (number of participants with no 97% Unclear 91% Unclear Not estimable pain as measured on VAS 0-10) at 12 weeks Night pain (number of participants with no 47% Unclear 45% Unclear Not estimable pain as measured on VAS 0-10) at 4 weeks Night pain (number of participants with no 83% Unclear 82% Unclear Not estimable pain as measured on VAS 0-10) at 12 weeks Unclear Pain on motion (number of participants 23% Unclear 14% Not estimable with no pain as measured on VAS 0-10) at 4 weeks Pain on motion (number of participants 63% Unclear 41% Unclear Not estimable with no pain as measured on VAS 0-10) at 12 weeks Study ID: Yiasemides 2011 Intervention: passive mobilisation plus exercise and advice Control: exercise and advice

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OUTCOME	INTERVE	NTION		CONTRO	DL		EFFECT ESTI- MATE
Study ID: Brox 1993 Intervention: supervised exercises Control: placebo laser							
able 14. Exercises alone versus placebo	0						
Global assessment of treatment success (6- point Likert scale, 0 = much worse, 5 = fully recovered) at 6 months	4.6	1	47	4.8	0.7	51	-0.20 (-0.54, 0.14)
Global assessment of treatment success (6- point Likert scale, 0 = much worse, 5 = fully recovered) at 1 month	4.2	0.8	47	3.9	0.8	51	0.30 (-0.02, 0.62)
Active range of flexion painful arc (degrees) at 6 months	3	9	47	3	6	51	0.00 (-3.05, 3.05)
Active range of flexion painful arc (degrees) at 1 month	14	23	47	19	19	51	-5.00 (-13.39, 3.39
Active range of abduction painful arc (de- grees) at 6 months	7	15	47	6	11	51	1.00 (-4.24, 6.24)
Active range of abduction painful arc (de- grees) at 1 month	28	24	47	36	25	51	-8.00 (-17.70, 1.70
Function (SPADI disability sub-score, 0-100 with higher scores denoting worse function) at 6 months	13	18	47	12	16	51	1.00 (-5.76, 7.76)
Function (SPADI disability sub-score, 0-100 with higher scores denoting worse function) at 1 month	32	23	47	30	19	51	2.00 (-6.39, 10.39)
with higher scores denoting worse pain) at 6 months		20	47	18	20	51	0.00 (-7.93, 7.93)

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#### Table 14. Exercises alone versus placebo (Continued)

	Median	IQR	n	Median	IQR	n	Mean difference (95% CI)
Overall pain (Neer shoulder score pain score 0-35 with higher scores denoting less pain) at 6 months	25	NR	50	15	NR	30	10 (95% CI not es timable)
Function (Neer shoulder score function score 0-30 with higher scores denoting better func- tion) at 6 months	25	NR	50	15	NR	30	10 (95% CI not es timable)
Night pain (0-9, 1 = no pain, 9 = worst possible pain) at 6 months	3	NR	50	4	NR	30	-1 (95% CI not es- timable)
Pain on motion (0-9, 1 = no pain, 9 = worst possible pain) at 6 months	3	NR	50	6	NR	30	-3 (95% CI not es timable)
ROM (Neer shoulder score ROM score 0-25 with higher scores denoting better ROM) at 6 months	23	NR	50	19	NR	30	4 (95% CI not es- timable)
Work disability: days of sick leave	Reported as	not significantl	y different betwe	en groups			
Global assessment of treatment success (number of participants with a good or an ex- cellent Neer shoulder score (> 80 points))	Reported as	not significantl	y different betwe	en groups			
Total adverse events	Zero events	in both groups					
Table 15. Exercises alone versus no treatStudy ID: Kachingwe 2008Intervention: supervised exercisesControl: advice regarding posture and overh			usual care				
OUTCOME	INTERVENTION			CONTROL			EFFECT ESTI- MATE
	Mean	SD	n	Mean	SD	n	Mean difference



Overall pain (VAS 0-10 with higher scores de- noting worse pain) % change from baseline to 6 weeks	20.8	112.3	8	14.4	119.8	7	Not estimable
Function (SPADI total score 0-130 with high- er scores denoting worse function) % change from baseline to 6 weeks	61.6	35.9	8	34.2	58.9	7	Not estimable
Active range of flexion % change from base- line to 6 weeks	27.6	41.7	8	42.6	15.8	7	Not estimable
Study ID: Lombardi 2008 Intervention: progressive resistance training Control: wait-list control	exercises						
OUTCOME	INTERVEN	ΓΙΟΝ		CONTROL			EFFECT ESTI- MATE
	Mean	SD	n	Mean	SD	n	Mean differenc (95% Cl)
Function (DASH laborious function, 0-100 with higher scores denoting worse functional- ity) at 2 months	28.7	24.8	30	44.2	28.2	30	-15.50 (-28.94, -2.06)
Function (DASH activities of daily living, 0-100 with higher scores denoting worse functional- ity) at 2 months	33.2	18.7	30	43.4	22.8	30	-10.20 (-20.75, 0.35)
Overall pain (VAS 0-10) at 2 months	2.4	2.1	30	4.3	3.2	30	-1.90 (-3.27, -0.5
Pain on motion (VAS 0-10) at 2 months	5.2	2	30	7.1	2.5	30	-1.90 (-3.05, -0.7
Active range of abduction (degrees) at 2 months	136.9	28.5	30	127.2	31.6	30	9.70 (-5.53, 24.9
Active range of flexion (degrees) at 2 months	137.1	24.8	30	130.6	27.4	30	6.50 (-6.72, 19.7
Active range of internal rotation (degrees) at 2 months	45.3	13.3	30	35.6	15.7	30	9.70 (2.34, 17.06
Active range of external rotation (degrees) at	82.7	18	30	70.5	31.7	30	12.20 (-0.84, 25.24)

Active range of extension (degrees) at 2 months	54.5	8.8	30	46.9	12.2	30	7.60 (2.22, 12.98
Strength: Peak torque (Nm) at velocity of 60 degrees/second - flexion	29.2	11.46	30	21.97	13.46	30	7.23 (0.90, 13.56
Strength: Peak torque (Nm) at velocity of 60 degrees/second - extension	40.03	18.42	30	28.4	19.27	30	11.63 (2.09, 21.3
Strength: Peak torque (Nm) at velocity of 60 degrees/second - abduction	24.37	11.98	30	15.8	11.44	30	8.57 (2.64, 14.50
Strength: Peak torque (Nm) at velocity of 60 degrees/second - adduction	33.8	18.39	30	21.73	18.42	30	12.07 (2.76, 21.3
Strength: Peak torque (Nm) at velocity of 60 degrees/second - internal rotation	22.23	9.28	30	17.13	7.99	30	5.10 (0.72, 9.48)
Strength: Peak torque (Nm) at velocity of 60 degrees/second - external rotation	12	5.12	30	9.53	4.15	30	2.47 (0.11, 4.83)
Quality of life (SF-36 physical functioning score, 0-100 where a higher score indicates a better quality of life) at 2 months	64.3	19	30	62.8	22.3	30	1.50 (-8.98, 11.9
Quality of life (SF-36 role-physical score, 0-100 where a higher score indicates a better quality of life) at 2 months	36.7	41.4	30	30.8	39.8	30	5.90 (-14.65, 26.45)
Quality of life (SF-36 bodily pain score, 0-100 where a higher score indicates a better quali- ty of life) at 2 months	54.3	16	30	46.7	24.1	30	7.60 (-2.75, 17.9
Quality of life (SF-36 general health score, 0-100 where a higher score indicates a better quality of life) at 2 months	73.9	20.3	30	68.2	25.3	30	5.70 (-5.91, 17.3
Quality of life (SF-36 vitality score, 0-100 where a higher score indicates a better quali- ty of life) at 2 months	54.8	24.7	30	49.4	26.9	30	5.40 (-7.67, 18.4

Quality of life (SF-36 social functioning score, 0-100 where a higher score indicates a better quality of life) at 2 months	76.7	27.4	30	65.4	27.2	30	11.30 (-2.52, 25.12)
Quality of life (SF-36 role-emotional score, 0-100 where a higher score indicates a better quality of life) at 2 months	62.22	40.8	30	55.5	42.3	30	6.72 (-14.31, 27.75)
Quality of life (SF-36 mental health score, 0-100 where a higher score indicates a better quality of life) at 2 months	62.9	22	30	56.5	25.1	30	6.40 (-5.54, 18.34)
	Events	Total		Events	Total		Risk ratio (95% CI)
							<u> </u>
Global assessment of treatment success at 2 months				at the experimental p. This difference w			r of "much better" and "a n groups (P = 0.001)."
months Study ID: Ludewig 2003 Intervention: home exercise programme		' responses than					
months Study ID: Ludewig 2003 Intervention: home exercise programme Control: no treatment	little better	' responses than		p. This difference w			n groups (P = 0.001)." EFFECT ESTI-
months Study ID: Ludewig 2003 Intervention: home exercise programme Control: no treatment	little better	' responses than	the control grou	p. This difference w	as statistically sig	gnificant betwee	n groups (P = 0.001)." EFFECT ESTI- MATE Mean difference
months         Study ID: Ludewig 2003         Intervention: home exercise programme         Control: no treatment         OUTCOME         Function (Shoulder Rating Questionnaire, 17-100, with a higher score indicating better	little better	' responses than TON SD	n	p. This difference w CONTROL Mean	as statistically sig	n n	n groups (P = 0.001)." EFFECT ESTI- MATE Mean difference (95% Cl)

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#### Table 16. Exercises alone versus another active intervention

#### Study ID: Ginn 2005 Intervention: exercise therapy

Control: glucocorticoid injection

Ουτςομε	INTERVENTIO	DN		CONTROL			EFFECT ESTIMATE
	Mean	95% CI	n	Mean	95% CI	n	Mean difference (95% Cl)
Pain (VAS 0-10) change from baseline to 5 weeks	0.3	0 to 2.3	48	0.2	0 - 1.7	45	0.10 (-2.33, 2.53)
Function (categorical rating scale, 0-27 with higher scores denoting worse function) change from baseline to 5 weeks	4.6	3.5 to 5.6	48	5.2	3.9 - 6.5	45	-0.60 (-2.26, 1.06)
Active range of abduction (degrees) change from baseline to 5 weeks (subgroup with decreased ROM and shoulder pain)	116	100 to 132	?	98	82 to 114	?	18 (95% Cl not es- timable)
Active range of abduction (degrees) change from baseline to 5 weeks (subgroup with full ROM despite shoulder pain)	24	10 to 37	?	28	13 to 44	?	-4 (95% CI not es- timable)
Active range of flexion (degrees) change from baseline to 5 weeks (subgroup with decreased ROM and shoulder pain)	114	104 to 124	?	111	102 to 120	?	3 (95% CI not es- timable)
Active range of flexion (degrees) change from baseline to 5 weeks (subgroup with full ROM despite shoulder pain)	1	0 to 10	?	0	0 to 8	?	1 (95% CI not es- timable)
Active hand-behind-back distance change from baseline to 5 weeks	6.1	3.1 to 9.1	48	7.5	4.9 to 10.2	45	-1.40 (-5.26, 2.46)
Strength (isometric abduction force %) change from baseline to 5 weeks	70	58 to 82	48	66	55 to 76	45	4.00 (-11.85, 19.85
	% range	Total		% range	Total		Risk ratio (95% C
Global assessment of treatment success (% participants rated as "improved") at 5 weeks	33 to 77%	48		35 to 78%	45		Not estimable

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#### Table 16. Exercises alone versus another active intervention (Continued)

#### Study ID: Brox 1993

OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Median	IQR	n	Median	IQR	n	Mean difference (95% Cl)
Overall pain (Neer shoulder score pain score 0-35 with higher scores denoting less pain) at 6 months	25	NR	50	25	NR	45	0 (95% CI not es- timable)
Function (Neer shoulder score function score 0-30 with higher scores denoting bet- ter function) at 6 months	25	NR	50	28	NR	45	- 3 (95% Cl not es- timable)
Night pain (0-9, 1 = no pain, 9 = worst possi- ble pain) at 6 months	3	NR	50	2	NR	45	1 (95% CI not es- timable)
Pain on motion (0-9, 1 = no pain, 9 = worst possible pain) at 6 months	3	NR	50	3	NR	45	0 (95% CI not es- timable)
ROM (Neer shoulder score ROM score 0-25 with higher scores denoting better ROM) at 6 months	23	NR	50	22	NR	45	1 (95% CI not es- timable)
Work disability: days of sick leave	Reported as	not significantly	y different betwe	en groups			
Global assessment of treatment success (number of participants with a good or an excellent Neer shoulder score (> 80 points))	Reported as	not significantly	y different betwe	en groups			
Total adverse events	Zero events	in both groups					
Study ID: Moosmayer 2014 Intervention: supervised exercises Control: rotator cuff tear repair surgery							
OUTCOME	INTERVENTION			CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)

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Overall pain (VAS 0-10) at 6 months	2.7	2.2	51	1.1	1.3	52	1.6 (0.9, 2.3)*
Overall pain (VAS 0-10) at 12 months	1.6	1.6	51	0.5	1.2	52	1.2 (0.6, 1.8)*
Overall pain (VAS 0-10) at 5 years	1.6	1.6	51	0.6	1.4	52	1 (0.2, 1.8)*
Function (Constant-Murley total score, 0-100, higher = better function) at 6 months	63.9	20.2	51	65.6	16.3	52	-2.8 (-10.1, 4.5)
Function (Constant-Murley total score, 0-100, higher = better function) at 12 months	70.3	19.1	51	77.7	13.4	52	-8.5 (-15, -1.9)*
Function (Constant-Murley total score, 0-100, higher = better function) at 5 years	74.2	20.3	51	79.8	15	52	-6.5 (-13.6, 0.7)
Active range of abduction (degrees) at 6 months	135.4	47.9	51	135.4	41.7	52	-2.2 (-20.3, 15.8
Active range of abduction (degrees) at 12 months	143.8	43.9	51	158.4	33.7	52	-16.8 (-32.4, -1.
Active range of abduction (degrees) at 5 years	155.1	41.2	51	167.3	30.6	52	-14.7 (-29.4, -0.
Active range of flexion (degrees) at 6 months	146.6	46.3	51	147.3	34.5	52	-2.1 (-18.1, 13.9
Active range of flexion (degrees) at 12 months	155.6	38.4	51	166.1	27.5	52	-10.3 (-23.6, 3.1
Active range of flexion (degrees) at 5 years	163.5	35.4	51	170.6	27.9	52	-8.3 (-21, 4.4)*
Strength (Constant-Murley strength sub- score, kg) at 6 months	10.6	5.4	51	8	4.6	52	2.5 (0.7, 4.2)*
Strength (Constant-Murley strength sub- score, kg) at 12 months	11.9	5.1	51	11.1	4	52	0.8 (-0.9, 2.4)*
Strength (Constant-Murley strength sub- score, kg) at 5 years	11.4	5.4	51	12.1	4.7	52	-0.8 (-2.7, 1.1)*

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# Manual therapy and exercise for rotator cuff disease (Review) Copyright © 2016 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd. Table 16. Exercises alone versus another active intervention (Continued)

Intervention: supervised exercises

Control: radial extracorporeal shockwave treatment

OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (9-point Likert scale, 1 = no pain, 9 = severe pain) at 6 weeks	2.6	1.9	51	2.9	2.1	52	-0.3 (-0.9, 0.4)*
Overall pain (9-point Likert scale, 1 = no pain, 9 = severe pain) at 18 weeks	2.5	1.9	51	2.7	2	52	-0.2 (-0.7, 0.3)*
Overall pain (9-point Likert scale, 1 = no pain, 9 = severe pain) at 1 year	2.1	1.5	48	2.6	2	46	-0.5 (-1.22, 0.22)
Function (SPADI total score 0-100) at 6 weeks	25.8	21.5	51	33.5	23.3	52	-10 (-17.6, -2.3)*
Function (SPADI total score 0-100) at 18 weeks	24.5	25.6	51	29.2	25.9	52	-8.4 (-16.5, -0.6)*
Function (SPADI total score 0-100) at 1 year	24	23.4	48	27.9	26.6	46	-3.9 (-14.04, 6.24)
Pain on motion (9-point Likert scale, 1 = no pain, 9 = severe pain) at 6 weeks	3.9	2	51	4.6	2.4	52	-0.7 (-1.6, 0.1)*
Pain on motion (9-point Likert scale, 1 = no pain, 9 = severe pain) at 18 weeks	3.6	2.3	51	4.1	2.5	52	-0.6 (-1.3, 0.2)*
Pain on motion (9-point Likert scale, 1 = no pain, 9 = severe pain) at 1 year	3.5	2.2	48	3.7	2.4	46	-0.2 (-1.13, 0.73)
	Events	Total		Events	Total		Risk ratio (95% CI
Work disability (number of participants working) at 18 weeks	38	50		26	50		1.46 (1.07, 1.99)
Work disability (number of participants working < 50% or unemployed) at 1 year	38	45		30	46		1.1 (1.0, 1.2)*

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Table 16.	Exercises alone	versus another activ	ve intervention	(Continued)
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Adverse events (aggravation of pain after	1	50	2	50	0.50 (0.05, 5.34)
treatment) within 1 year study period					

#### Study ID: Giombini 2006 Intervention: supervised and home exercises Control: microwave diathermy

OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (VAS 0-10) at 4 weeks	5.3	0.65	11	2.4	0.46	14	2.90 (2.45, 3.35)
Overall pain (VAS 0-10) at 10 weeks	4.9	0.88	11	1.2	0.63	14	3.70 (3.08, 4.32)
Function (Constant-Murley total score, 0-100, higher = better function) at 4 weeks	61.2	4.28	11	78.1	4.23	14	-16.90 (-20.26, -13.54)
Function (Constant-Murley total score, 0-100, higher = better function) at 10 weeks	63.27	5.56	11	82	5.73	14	-18.73 (-23.18, -14.28)
	Events	Total		Events	Total		Risk ratio (95% CI
Global assessment of treatment success (ready to return to sport) at 4 weeks	4	11		11	14		0.46 (0.20, 1.06)
Global assessment of treatment success (ready to return to sport) at 10 weeks	4	11		12	14		0.42 (0.19, 0.95)
Adverse events	Zero events	in both groups					
Study ID: Giombini 2006 Intervention: supervised and home exercis Control: therapeutic ultrasound	es						
OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (VAS 0-10) at 4 weeks	5.3	0.65	11	5.8	0.96	12	-0.50 (-1.17, 0.17)

Overall pain (VAS 0-10) at 10 weeks	4.9	0.88	11	5.15	0.87	12	-0.25 (-0.97, 0.47
Function (Constant-Murley total score, 0-100, higher = better function) at 4 weeks	61.2	4.28	11	60	3.21	12	1.20 (-1.91, 4.31)
Function (Constant-Murley total score, 0-100, higher = better function) at 10 weeks	63.27	5.56	11	61.75	4.18	12	1.52 (-2.53, 5.57)
	Events	Total		Events	Total		Risk ratio (95%
Global assessment of treatment success (ready to return to sport) at 4 weeks	4	11		6	12		0.73 (0.28, 1.91)
Global assessment of treatment success (ready to return to sport) at 10 weeks	4	11		4	12		1.09 (0.36, 3.34)
Adverse events	Zero events	in both groups					
Study ID: Walther 2004 Intervention: standardised self-training of Control: functional brace			cises	CONTROL			FEFECT ESTIMA
Intervention: standardised self-training of	INTERVENT	TON		CONTROL			
Intervention: standardised self-training of Control: functional brace			n	CONTROL Mean	SD	n	
Intervention: standardised self-training of Control: functional brace	INTERVENT Mean	TON	n		SD	n	Mean difference
Intervention: standardised self-training of Control: functional brace OUTCOME Rest pain (VAS 0-100) at both 6 and 12	INTERVENT Mean "No signific	TION	<b>n</b> tween groups"		SD	n	EFFECT ESTIMA Mean difference (95% CI)
Intervention: standardised self-training of Control: functional brace OUTCOME Rest pain (VAS 0-100) at both 6 and 12 weeks Function (Constant-Murley total score) at	INTERVENT Mean "No signific "No signific	SD ant difference be	<b>n</b> tween groups" tween groups"		SD	n	Mean difference
Intervention: standardised self-training of Control: functional brace OUTCOME Rest pain (VAS 0-100) at both 6 and 12 weeks Function (Constant-Murley total score) at both 6 and 12 weeks Pain on motion (VAS 0-100) at both 6 and 12	INTERVENT Mean "No signific "No signific	TION SD ant difference be ant difference be	<b>n</b> tween groups" tween groups"		<b>SD</b>	<b>n</b>	Mean difference (95% CI)
Intervention: standardised self-training of Control: functional brace OUTCOME Rest pain (VAS 0-100) at both 6 and 12 weeks Function (Constant-Murley total score) at both 6 and 12 weeks Pain on motion (VAS 0-100) at both 6 and 12 weeks Strength (Constant-Murley strength sub-	INTERVENT Mean "No signific "No signific	SD ant difference be ant difference be ant difference be	n tween groups" tween groups" tween groups"	Mean			Mean difference

Number of adverse events	0	20		2	20		0.20 (0.01, 3.92)				
Adverse events	patients trea Another pati	None of the patients treated with physiotherapy or self-training dropped out of the therapy regimen. However, one of the patients treated with the brace complained about being bothered by the brace at work, especially while working overhead. Another patient had eczema of the skin develop underneath the pads. Both patients continued to wear the brace during the remainder of the 12-week therapy period"									
Study ID: Walther 2004 ntervention: supervised stretching exercis Control: functional brace	ses										
DUTCOME	INTERVENT	ION		OUTCOME		EFFECT ESTIMA- TRE					
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)				
Rest pain (VAS 0-100) at both 6 and 12 weeks	"No significa	ant difference be	etween groups"								
Function (Constant-Murley total score) at both 6 and 12 weeks	"No significa	ant difference be	etween groups"								
Pain on motion (VAS 0-100) at both 6 and 12 weeks	"No significa	ant difference be	etween groups"								
Strength (Constant-Murley strength sub- score 0-25) at 12 weeks	11.8	5.4	20	14.4	5.4	20	-2.60 (-5.95, 0.75)				
Work disability (number of months with in- ability to work)	1.6	NR	20	1.5	NR	20	0.1 (95% CI not es- timable)				
	Events	Total		Events	Total		Risk ratio (95% CI)				
Number of adverse events	0	20		2	20		0.20 (0.01, 3.92)				
Adverse events	patients trea Another pati	ated with the bra	ace complained abo of the skin develop	ut being bothere	d by the brace at	work, especially	nen. However, one of the while working overhead. wear the brace during the				

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#### Table 17. Exercises alone as an add-on to another physical therapy intervention versus the other physical therapy intervention

#### Study ID: Ainsworth 2009

Intervention: supervised exercises, ultrasound, glucocorticoid injection (if needed) and advice

Control: ultrasound, glucocorticoid injection (if needed) and advice

OUTCOME	INTERVEN	ΓΙΟΝ		CONTROL			EFFECT ESTI- MATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)
Function (Oxford Shoulder Score 0-48 with higher scores denoting better function) change from baseline to 6 months	9.42	6.23	26	4.43	7.23	30	4.99 (1.46, 8.52)
Function (Oxford Shoulder Score 0-48 with higher scores denoting better function) change from baseline to 12 months	8.96	5.3	24	6.27	7.93	30	2.69 (-0.85, 6.23)
Quality of life (SF-36 physical functioning score, 0-100 where higher = better) change from baseline to 6 months	4.42	22.33	26	-3	13.93	30	7.42 (-2.51, 17.35)
Quality of life (SF-36 physical functioning score, 0-100 where higher = better) change from baseline to 12 months	5.21	13.39	24	-3.17	17.19	30	8.38 (0.22, 16.54)
Quality of life (SF-36 role-physical score, 0-100 where higher = better) change from baseline to 6 months	2.89	49.16	24	14.17	42.89	30	-11.28 (-36.23, 13.67)
Quality of life (SF-36 role-physical score, 0-100 where higher = better) change from baseline to 12 months	-5.21	44.22	26	18.33	37.1	30	-23.54 (-45.11, -1.97)
Quality of life (SF-36 bodily pain score, 0-100 where higher = better) change from baseline to 6 months	3.65	22.64	26	4.58	28.01	30	-0.93 (-14.20, 12.34)
Quality of life (SF-36 bodily pain score, 0-100 where higher = better) change from baseline to 12 months	1.56	30.3	24	4.42	29.88	30	-2.86 (-19.02, 13.30)

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Quality of life (SF-36 general health score, 0-100 where higher = better) change from baseline to 6 months	5	15.36	26	4.33	17.75	30	0.67 (-8.00, 9.34)
Quality of life (SF-36 general health score, 0-100 where higher = better) change from baseline to 12 months	-4.79	15.77	24	-1.17	16.64	30	-3.62 (-12.30, 5.06)
Quality of life (SF-36 vitality score, 0-100 where higher = better) change from baseline to 6 months	2.5	15.64	26	0.67	19.38	30	1.83 (-7.35, 11.01)
Quality of life (SF-36 vitality score, 0-100 where higher = better) change from baseline to 12 months	-3.54	15.91	24	2.5	17.94	30	-6.04 (-15.08, 3.00)
Quality of life (SF-36 social functioning score, 0-100 where higher = better) change from baseline to 6 months	-0.48	26.81	26	0.83	28.03	30	-1.31 (-15.69, 13.07)
Quality of life (SF-36 social functioning score, 0-100 where higher = better) change from baseline to 12 months	-7.81	34.74	24	5	28.16	30	-12.81 (-29.98, 4.36)
Quality of life (SF-36 role-emotional score, 0-100 where higher = better) change from baseline to 6 months	6.41	47.44	26	0	41.98	30	6.41 (-17.22, 30.04)
Quality of life (SF-36 role-emotional score, 0-100 where higher = better) change from baseline to 12 months	-5.56	30.56	24	3.33	29.49	30	-8.89 (-25.04, 7.26)
Quality of life (SF-36 emotional wellbeing score, 0-100 where higher = better) change from baseline to 6 months	4.15	15.11	26	3.33	18.59	30	0.82 (-8.01, 9.65)
Quality of life (SF-36 emotional wellbeing score, 0-100 where higher = better) change from baseline to 12 months	-4.17	18.89	24	4.93	17.44	30	-9.10 (-18.90, 0.70)
Passive external rotation (degrees) change from baseline to 6 months	8.75	NR	26	-3.7	NR	30	12.45 (95% Cl not estimable)

Table 17. Exercises alone as an add-on to another physical therapy intervention versus the other physical therapy intervention (Continued)

#### Table 17. Exercises alone as an add-on to another physical therapy intervention versus the other physical therapy intervention (Continued)

Passive external rotation (degrees) change from baseline to 12 months	7.43	NR	24	4.4	NR	30	3.03 (95% CI not estimable)
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#### Study ID: Bae 2011

Intervention: motor control exercises and strengthening exercises plus hot packs plus TENS plus ultrasound Control: hot packs plus TENS plus ultrasound

OUTCOME	INTERVEN	ΓΙΟΝ		CONTROL		EFFECT ESTI- MATE	
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Function (SPADI total score 0-100 with higher scores denoting worse function) at 4 weeks	20.7	4.1	17	32.1	6	18	-11.40 (-14.79, -8.01)
Active range of abduction (degrees) at 4 weeks	129.1	19.7	17	106.6	10.4	18	22.50 (11.97, 33.03)
Active range of flexion (degrees) at 4 weeks	155.6	8.4	17	146.2	10.9	18	9.40 (2.97, 15.83)
Active range of extension (degrees) at 4 weeks	40.2	4.8	17	36.2	5.8	18	4.00 (0.48, 7.52)
Active range of external rotation (degrees) at 4 weeks	76.5	4.5	17	70.1	6.3	18	6.40 (2.79, 10.01)
Active range of internal rotation (degrees) at 4 weeks	47	8.5	17	43.7	7.7	18	3.30 (-2.08, 8.68)
Isokinteic strength: peak torque (Nm) of ex- ternal rotator 60 degrees/sec at 4 weeks	21.1	5.4	17	14.5	4.6	18	6.60 (3.27, 9.93)
Isokinteic strength: peak torque (Nm) of ex- ternal rotator 180 degrees/sec at 4 weeks	65.2	6.6	17	68.4	7.3	18	-3.20 (-7.81, 1.41)
Isokinteic strength: peak torque (Nm) of inter- nal rotator 60 degrees/sec at 4 weeks	24.7	4.5	17	22.8	5.7	18	1.90 (-1.49, 5.29)
Isokinteic strength: peak torque (Nm) of inter- nal rotator 180 degrees/sec at 4 weeks	26.1	6.3	17	21.9	7.5	18	4.20 (-0.38, 8.78)

Study ID: Baskurt 2011

Intervention: scapular stabilisation exercises (PNF) plus stretching and strengthening exercises

Table 17. Exercises alone as an add-on to another physical therapy intervention versus the other physical therapy intervention (Continu	ed)
Control: stretching and strengthening exercises	

Ουτςομε	INTERVENT	ION		CONTROL		EFFECT ESTI- MATE	
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (rest pain VAS 0-10) at 6 weeks	0.85	1.08	20	1.4	1.78	20	-0.55 (-1.46, 0.36
Pain on activity (VAS 0-10) at 6 weeks	3	1.55	20	3.2	2.11	20	-0.20 (-1.35, 0.95
Range of abduction (degrees, unclear if active or passive) at 6 weeks	179.75	11.11	20	177	13.4	20	2.75 (-4.88, 10.38
Range of flexion (degrees, unclear if active or passive) at 6 weeks	179.75	1.11	20	178.5	4.61	20	1.25 (-0.83, 3.33)
Range of internal rotation in 90 degrees ab- duction (degrees, unclear if active or passive) at 6 weeks	88.5	3.66	20	87.5	5.5	20	1.00 (-1.90, 3.90)
Range of external rotation in 90 degrees ab- duction (degrees, unclear if active or passive) at 6 weeks	87.5	4.13	20	84.5	10.9	20	3.00 (-2.11, 8.11)
Quality of life (WORC 0-2100, with higher scores denoting worse quality of life) at 6 weeks	82.61	10.33	20	70.82	19.7	20	11.79 (2.04, 21.5 <sup>4</sup>
Strength: lower trapezium (kg) at 6 weeks	10.96	1.2	20	9.22	1.24	20	1.74 (0.98, 2.50)
Strength: middle trapezium (kg) at 6 weeks	11.15	1.41	20	10.21	1.31	20	0.94 (0.10, 1.78)
Strength: upper trapezium (kg) at 6 weeks	12.19	1.28	20	11.24	1.59	20	0.95 (0.06, 1.84)
Strength: serratus anterior (kg) at 6 weeks	10.19	1.61	20	8.78	1.59	20	1.41 (0.42, 2.40)
Strength: supraspinatus (kg) at 6 weeks	11.64	1.25	20	10.79	1.55	20	0.85 (-0.02, 1.72)
Strength: subscapularis (kg) at 6 weeks	6.59	1.44	20	6.02	1.31	20	0.57 (-0.28, 1.42)
Strength: infraspinatus (kg) at 6 weeks	7.05	1.3	20	6.81	1.13	20	0.24 (-0.51, 0.99)

#### Table 17. Exercises alone as an add-on to another physical therapy intervention versus the other physical therapy intervention (Continued)

#### Study ID: Beaudreuil 2011

Intervention: Dynamic Humeral Centering plus massage and home exercise

Control: non-specific mobilisation plus massage and exercise

DUTCOME	INTERVEN	ΓΙΟΝ		CONTROL	CONTROL		
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (Constant-Murley pain sub-score 0-15, with higher scores denoting less pain) at 3 months	12.2	2.8	30	9.9	2.9	32	2.30 (0.88, 3.72)
Overall pain (Constant-Murley pain sub-score 0-15, with higher scores denoting less pain) at 12 months	13.1	2	22	10.8	3.7	26	2.30 (0.65, 3.95)
Function (Constant-Murley total score 0-100) at 3 months	63.8	16.9	30	54	19.8	32	9.80 (0.65, 18.95)
Function (Constant-Murley total score 0-100) at 12 months	68.9	17	22	62	21.1	26	6.90 (-3.88, 17.68
ROM (Constant-Murley ROM sub-score 0-40, with higher scores denoting better ROM) at 3 months	26.7	9.6	30	22.2	9.7	32	4.50 (-0.31, 9.31)
ROM (Constant-Murley ROM sub-score 0-40, with higher scores denoting better ROM) at 12 months	30.3	10	22	27.8	11.2	26	2.50 (-3.50, 8.50)
Strength (Constant-Murley strength sub- score 0-25, with higher scores denoting better strength) at 3 months	8.6	5	30	8	6.8	32	0.60 (-2.36, 3.56)
Strength (Constant-Murley strength sub- score 0-25, with higher scores denoting better strength) at 12 months	8	6	22	7.9	5.6	26	0.10 (-3.20, 3.40)

Table 17. Exercises alone as an add-on to another physical therapy intervention versus the other physical therapy intervention (Continued)

Ουτςομε	INTERVEN	ΓΙΟΝ		CONTROL			EFFECT ESTI- MATE	
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)	
Function (SPADI total score 0-100 with higher scores denoting worse function) at 6 weeks*	25.4	11.9	31	17.7	12	30	7.70 (1.70, 13.70)	
Function (SPADI total score 0-100 with higher scores denoting worse function) at 12 weeks*	17	11.4	31	14.5	11.7	30	2.50 (-3.30, 8.30)	
Isometric strength at 0° abduction (Newton) at 6 weeks*	150.8	27.6	31	142.7	27.5	30	8.10 (-5.73, 21.93	
Isometric strength at 0° abduction (Newton) at 12 weeks*	154.3	27.6	31	147.1	27.2	30	7.20 (-6.55, 20.9	
Isometric strength at 45° abduction (Newton) at 6 weeks*	79.7	12	31	81.7	12	30	-2.00 (-8.02, 4.02	
Isometric strength at 45° abduction (Newton) at 12 weeks*	81.6	12.2	31	83.5	11.8	30	-1.90 (-7.92, 4.12	
Isometric strength at 90° abduction (Newton) at 6 weeks*	74.8	12.3	31	72.5	12.3	30	2.30 (-3.87, 8.47)	
Isometric strength at 90° abduction (Newton) at 12 weeks*	78	12.5	31	70	12.2	30	8.00 (1.80, 14.20	
Isometric strength external rotation (Newton) at 6 weeks*	94.3	12.2	31	90.5	12.5	30	3.80 (-2.40, 10.00	
Isometric strength external rotation (Newton) at 12 weeks*	96	12.4	31	92.7	12.3	30	3.30 (-2.90, 9.50)	
Isometric strength internal rotation (Newton) at 6 weeks*	126.5	17.6	31	123.2	17.5	30	3.30 (-5.51, 12.1	
Isometric strength internal rotation (Newton) at 12 weeks*	129	17.9	31	125	17.2	30	4.00 (-4.81, 12.8	

#### Table 17. Exercises alone as an add-on to another physical therapy intervention versus the other physical therapy intervention (Continued)

	Events	Total		Events	Total		Risk ratio (95% CI)
Global assessment of treatment success ("very large improvement or "large improve- ment") at 6 weeks	14	30		10	27		1.26 (0.68, 2.35)
Global assessment of treatment success ("very large improvement or "large improve- ment") at 12 weeks	14	27		13	20		0.80 (0.49, 1.30)
Study ID: Martins 2012 Intervention: proprioception exercises plus s Control: standardised physiotherapy	tandardised	physiotherapy					
OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTI- MATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)
Quality of life (WORC, 0-2100, with higher score denoting worse QoL) at 6 weeks	0.01	NR	8	0.06	NR	8	-0.05 (95% CI no estimable)
Work disability (Occupational Stress Indica- tor, 22-123) at 6 weeks	90.2	20.8	8	87	18.2	8	3.20 (-15.95, 22.35)
	Events	Total		Events	Total		Risk ratio (95% Cl)
Overall pain (number participants with mild pain i.e. score < = 3 on VAS 0-10) at 6 weeks	6	8		5	8		1.20 (0.61, 2.34)
Overall pain (number participants with mod- erate pain i.e. score 4 < = 7 on VAS 0-10) at 6 weeks	2	8		3	8		0.67 (0.15, 2.98)
Overall pain (number participants with severe pain i.e. score > 7 on VAS 0-10) at 6 weeks	0	8		0	8		Not estimable

\*Mean scores in Maenhout 2013 adjusted for baseline score

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#### Table 18. One type of manual therapy or exercise versus another

#### Study ID: Blume 2014

Intervention: supervised eccentric progressive resistance exercises plus ice plus home exercises Control: supervised concentric progressive resistance exercises plus ice plus home exercises

OUTCOME	INTERVENT	ION		CONTROL	EFFECT ESTIMATE		
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Function (DASH 0-100 where higher = worse function) at 5 weeks	14.6	8.7	17	11.7	6.6	13	2.90 (-2.57, 8.37)
Function (DASH 0-100 where higher = worse function) at 8 weeks	11.5	11.8	17	8.4	6.7	13	3.10 (-3.59, 9.79)
Active ROM: scaption (degrees) at 5 weeks	143.4	24	17	141.9	25.7	13	1.50 (-16.54, 19.54)
Active ROM: scaption (degrees) at 8 weeks	145.7	23.6	17	143.7	31.3	13	2.00 (-18.38, 22.38)
Strength: abduction torque (lbs) at 5 weeks	239.3	132.8	17	193.9	110.6	13	45.40 (-41.78, 132.58)
Strength: abduction torque (lbs) at 8 weeks	281.5	165.6	17	259.4	138.4	13	22.10 (-86.79, 130.99)
Strength: external rotation torque (lbs) at 5 weeks	173.5	121.7	17	140.2	103.1	13	33.30 (-47.25, 113.85)
Strength: external rotation torque (lbs) at 8 weeks	203.5	121.7	17	186.1	122.5	13	17.40 (-70.81, 105.61)
Study ID: Celik 2009 Intervention: exercise below 90 degrees pl Control: exercise above 90 degrees standa			ру				
OUTCOME	INTERVENT	ION		CONTROL		EFFECT ESTIMATE	
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (VAS 0-10) at 2 weeks	2.9	1.7	15	4.1	1.6	15	-1.20 (-2.38, -0.02)
Overall pain (VAS 0-10) at 4 months	1.1	1	15	1.6	1.6	15	-0.50 (-1.45, 0.45)

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Global assessment of treatment success (patient satisfaction scale, 0-4, higher score nigher satisfaction) at 2 weeks	2.6	0.7	15	2.2	0.6	15	0.40 (-0.07, 0.87)
Global assessment of treatment success patient satisfaction scale, 0-4, higher score nigher satisfaction) at 2 weeks	3.3	0.4	15	3.1	0.1	15	0.20 (-0.01, 0.41)
Study ID: Citaker 2005 Intervention: manual mobilisation plus sta Control: proprioceptive neuromuscular fac			nysiotherapy				
DUTCOME	INTERVEN	ΓΙΟΝ		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Function (total UCLA score, 0-35, where nigher scores denote better function) at 3 weeks	33.22	2.95	20	29.97	4.6	20	3.25 (0.86, 5.64)
Day pain at rest (VAS 0-10) at 3 weeks	0.75	1.45	20	0.25	0.91	20	0.50 (-0.25, 1.25)
Night pain at rest (VAS 0-10) at 3 weeks	1.75	2.55	20	1.65	2.54	20	0.10 (-1.48, 1.68)
Day pain on motion (VAS 0-10) at 3 weeks	0.6	1.27	20	0.6	1.19	20	0.00 (-0.76, 0.76)
Night pain on motion (VAS 0-10) at 3 weeks	1.5	2.3	20	1.85	2.64	20	-0.35 (-1.88, 1.18)
Range of abduction (degrees, unclear if ac- ive or passive) at 3 weeks	170.5	21.52	20	174.75	9.8	20	-4.25 (-14.61, 6.11)
Range of flexion (degrees, unclear if active or passive) at 3 weeks	170.4	11.74	20	173	10.44	20	-2.60 (-9.49, 4.29)
Range of external rotation (degrees, unclear f active or passive) at 3 weeks	77.5	19.23	20	80.25	10.57	20	-2.75 (-12.37, 6.87)
Range of internal rotation (degrees, unclear f active or passive) at 3 weeks	85.5	13.11	20	85.25	9.1	20	0.25 (-6.74, 7.24)

Intervention: mobilisation with movement and kinesiotaping

# Table 18. One type of manual therapy or exercise versus another (Continued) Control: supervised exercise programme

ОИТСОМЕ	INTERVENTIO	N		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)
Active range of abduction (degrees) at 10 days	170	17.89	10	60.5	15.72	10	109.50 (94.74, 124.26)
Active range of flexion (degrees) at 10 days	166	20.59	10	86	21.89	10	80.00 (61.37, 98.63)

# Study ID: Heredia-Rizo 2013

# Intervention: soft tissue techniques

# Control: mobilisation, proprioceptive neuromuscular facilitation and exercises

OUTCOME	INTERVENT	ION		CONTROL	CONTROL			
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)	
Function (DASH, 0-100 with higher scores denoting worse functionality) at 3 weeks	34.69	11.21	11	38.61	20.8	11	-3.92 (-17.88, 10.04)	
Active range of abduction (degrees) at 3 weeks	142.5	25.3	11	115.55	20.68	11	26.95 (7.64, 46.26)	
Active range of flexion (degrees) at 3 weeks	153	20.97	11	140.01	16	11	12.99 (-2.60, 28.58)	
Active range of extension (degrees) at 3 weeks	65.5	12.79	11	51.66	11.72	11	13.84 (3.59, 24.09)	
Active range of external rotation (degrees) at 3 weeks	68	19.32	11	52.77	21.95	11	15.23 (-2.05, 32.51)	
Active range of internal rotation (degrees) at 3 weeks	85	8.49	11	73.33	16	11	11.67 (0.97, 22.37)	
Study ID: Holmgren 2012 Intervention: specific exercise programme Control: non-specific exercises								
OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE	

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 Table 18. One type of manual therapy or exercise versus another (Continued)

	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)
Overall pain (VAS 0-100) at 3 months	10	14	51	20	25	46	-10.00 (-18.18, -1.82)
Function (Constant-Murley total score, 0-100 with higher scores denoting better function) at 3 months	72.5	19	51	52.5	23	46	20.00 (11.55, 28.45
Pain on motion (VAS 0-100) at 3 months	25	26	51	41	27	46	-16.00 (-26.57, -5.43)
Night pain (VAS 0-100) at 3 months	15	22	51	27	27	46	-12.00 (-21.87, -2.13)
Quality of life (EuroQoL EQ-5D, -0.59 to 1 where lower scores denote worse QoL) at 3 months	0.82	0.14	51	0.69	0.24	46	0.13 (0.05, 0.21)
	Events	Total		Events	Total		Risk ratio (95% C
Global assessment of treatment success ("recovered" or "large improvement") at 3 months	35	51		11	46		2.87 (1.66, 4.96)
Had surgery between 3 months and 1 year	12	51		29	46		0.37 (0.22, 0.64)
Study ID: Kachingwe 2008 Intervention: glenohumeral mobilisation p Control: mobilisation with movement plus							
OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATI
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (VAS 0-10 with higher scores denoting worse pain) % change from base- line to 6 weeks	44.2	38.6	9	55.2	31.9	9	Not estimable

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Function (SPADI total score 0-130 with higher scores denoting worse function) % change from baseline to 6 weeks	56.7	29.8	9	55.5	20.1	9	Not estimable
Active range of flexion % change from base- line to 6 weeks	-15.9	116.6	9	46.7	31.9	9	Not estimable
Study ID: Kassolik 2013 Intervention: classic Swedish massage Control: massage based on the tensegrity p	principle						
ОИТСОМЕ	INTERVENT	TION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (VAS 0-10 in Short-Form McGill Pain Questionnaire) at 2 weeks	3.16	1.77	15	1.66	0.48	15	1.50 (0.57, 2.43)
Overall pain (VAS 0-10 in Short-Form McGill Pain Questionnaire) at 6 weeks	3.7	1.88	15	1.06	0.45	15	2.64 (1.66, 3.62)
Active range of abduction (degrees) at 2 weeks	145.8	18.6	15	136.2	11.9	15	9.60 (-1.57, 20.77)
Active range of abduction (degrees) at 6 weeks	144.4	17.6	15	136.6	10.6	15	7.80 (-2.60, 18.20)
Active range of flexion (degrees) at 2 weeks	156.2	20.9	15	143.3	13.9	15	12.90 (0.20, 25.60)
Active range of flexion (degrees) at 6 weeks	154.4	20.4	15	144.6	15	15	9.80 (-3.01, 22.61)
Active range of extension (degrees) at 2 weeks	36.1	5.4	15	29.6	8.3	15	6.50 (1.49, 11.51)
Active range of extension (degrees) at 6 weeks	33.6	5.6	15	29.6	7.9	15	4.00 (-0.90, 8.90)
Active range of external rotation (degrees) at 2 weeks	14.9	5.4	15	13.8	5.2	15	1.10 (-2.69, 4.89)
Active range of external rotation (degrees) at 6 weeks	13.8	5.4	15	13.6	5.5	15	0.20 (-3.70, 4.10)



Table 18. One type of manual therapy of	or exercise ver	sus another (Cor	ntinued)				
Active range of internal rotation (degrees) at 2 weeks	25.2	5.9	15	24.4	5.9	15	0.80 (-3.42, 5.02)
Active range of internal rotation (degrees) at 6 weeks	23.2	6.4	15	24.4	6.1	15	-1.20 (-5.67, 3.27)

#### Study ID: Littlewood 2014

Intervention: self-managed loaded exercise Control: usual physiotherapy (might include advice, stretching, exercise, manual therapy, massage, strapping, acupuncture, electrotherapy, corticosteroid injection at the discretion of the treating physiotherapist)

OUTCOME	INTERVENT	ION		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Function (SPADI total score, 0-100 with higher scores denoting worse function) at 3 months	20.9	19.2	12	20.7	20.3	12	0.20 (-15.61, 16.01)
Quality of life (SF-36 physical functioning score, 0-100 where a higher score indicates a better quality of life) at 3 months	78.2	17.7	12	73.3	29.3	12	4.90 (-14.47, 24.27)
Quality of life (SF-36 role-physical score, 0-100 where a higher score indicates a bet- ter quality of life) at 3 months	88.5	18	12	79.2	20	12	9.30 (-5.92, 24.52)
Quality of life (SF-36 bodily pain score, 0-100 where a higher score indicates a bet- ter quality of life) at 3 months	61.4	13.4	12	71.8	18.2	12	-10.40 (-23.19, 2.39)
Quality of life (SF-36 general health score, 0-100 where a higher score indicates a bet- ter quality of life) at 3 months	74.2	20.3	12	72.9	11.6	12	1.30 (-11.93, 14.53)
Quality of life (SF-36 vitality score, 0-100 where a higher score indicates a better quality of life) at 3 months	69.3	12.1	12	70.8	21.5	12	-1.50 (-15.46, 12.46)
Quality of life (SF-36 social functioning score, 0-100 where a higher score indicates a better quality of life) at 3 months	45.8	11.1	12	50	10.7	12	-4.20 (-12.92, 4.52)

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Table 18. One type of manual therapy o	or exercise v	ersus another	(Continued)				
Quality of life (SF-36 role-emotional score, 0-100 where a higher score indicates a bet- ter quality of life) at 3 months	95.8	10.4	12	97.2	7.4	12	-1.40 (-8.62, 5.82)
Quality of life (SF-36 mental health score, 0-100 where a higher score indicates a bet- ter quality of life) at 3 months	84.6	12.9	12	82.5	13.1	12	2.10 (-8.30, 12.50)
Study ID: Marzetti 2014 Intervention: neurocognitive therapeutic e Control: traditional therapeutic exercise	exercise						
OUTCOME	INTERVEN	ΓΙΟΝ		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Rest pain (VAS 0-10) at 5 weeks	0.95	NR	24	2.19	NR	24	-1.24 (-2.93, 0.46)
Rest pain (VAS 0-10) at 24 weeks	0.45	NR	24	2.05	NR	24	-1.59 (-3.29, 0.10)
Function (Constant-Murley total score 0-100, higher = better function) at 5 weeks	75.5	NR	24	74.57	NR	24	0.93 (-11.94, 13.8)
Function (Constant-Murley total score 0-100, higher = better function) at 24 weeks	83.27	NR	24	76.95	NR	24	6.32 (-6.55, 19.19)
Pain on motion (VAS 0-10) at 5 weeks	3.73	NR	24	4.1	NR	24	-0.37 (-2.35, 1.62)
Pain on motion (VAS 0-10) at 24 weeks	1.86	NR	24	3.33	NR	24	-1.47 (-3.46, 0.52)
Adverse events	Zero events	in both groups					
Study ID: Osteras 2008 Intervention: high dose medical exercise th Control: low dose medical exercise therapy							
OUTCOME	INTERVEN	ΓΙΟΝ		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (VAS 0-10) at 3 months	2.1	1.6	29	4.1	1.7	27	-2.00 (-2.87, -1.13)

Overall pain (VAS 0-10) at 15 months	1.2	1	26	4.2	2.1	23	-3.00 (-3.94, -2.06)
Function (Shoulder Rating Questionnaire, 17 - 90 with a higher score indicating better function) at 3 months	69.1	13.3	29	51.5	14.2	27	17.60 (10.38, 24.82)
Function (Shoulder Rating Questionnaire, 17 - 90 with a higher score indicating better function) at 15 months	79.1	7.6	26	54.7	18.7	23	24.40 (16.22, 32.58)
Range of abduction (degrees, unclear if ac- tive or passive) change from baseline to 3 months	42	42.0632	29	12	35.3905	27	30.00 (9.69, 50.31)
Range of abduction (degrees, unclear if ac- tive or passive) change from baseline to 9 months	49	40.4462	27	14	38.7616	25	35.00 (13.47, 56.53)
Range of flexion (degrees, unclear if ac- tive or passive) change from baseline to 3 months	31	63.0949	29	7	22.751	27	24.00 (-0.51, 48.51)
Range of flexion (degrees, unclear if ac- tive or passive) change from baseline to 9 months	34	22.751	27	8	16.9582	25	26.00 (15.14, 36.86)
Strength (isometric strength in abduction, Newtons) change from baseline to 3 months	34	26.2895	29	17	20.2231	27	17.00 (4.76, 29.24)
Strength (isometric strength in abduction, Newtons) change from baseline to 9 months	45	42.9741	27	14	46.0294	25	31.00 (6.74, 55.26)
Strength (isometric strength in flexion, New- tons) change from baseline to 3 months	33	34.1764	29	4	12.6395	27	29.00 (15.68, 42.32)
Strength (isometric strength in flexion, New- tons) change from baseline to 9 months	49	60.6694	27	28	62.9876	25	21.00 (-12.66, 54.66)
Strength (isometric strength in external ro- tation, Newtons) change from baseline to 3 months	28	52.579	29	9	30.3347	27	19.00 (-3.30, 41.30)



Strength (isometric strength in external ro- tation, Newtons) change from baseline to 9 months	36	63.1973	27	3	31.4938	25	33.00 (6.16, 59.84)
Strength (isometric strength in internal ro- tation, Newtons) change from baseline to 3 months	15	21.0316	29	13	20.2231	27	2.00 (-8.81, 12.81)
Strength (isometric strength in internal ro- tation, Newtons) change from baseline to 9 months	21	22.751	27	9	24.226	25	12.00 (-0.80, 24.80)
Study ID: Senbursa 2007 Intervention: manual therapy program Control: self-training program							
OUTCOME	INTERVENTIO	DN		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Rest pain (VAS 0-10) at 3 months	0.7	1.4	15	0.9	0.2	15	-0.20 (-0.92, 0.52)
Night pain (VAS 0-10) at 3 months	2.2	2.4	15	1.2	1.6	15	1.00 (-0.46, 2.46)
Pain on motion (VAS 0-10) at 3 months	3.1	2	15	2.5	1.5	15	0.60 (-0.67, 1.87)
Study ID: Senbursa 2007 Intervention: supervised exercises Control: home exercises							
OUTCOME	INTERVENTIO	ON		CONTROL			EFFECT ESTIMATE
	% Events	Total		% Events	Total		Risk ratio (95% CI
Rest pain (number of participants with no pain as measured on VAS 0-10) at 4 weeks	64%	Unclear		82%	Unclear		Not estimable
Rest pain (number of participants with no pain as measured on VAS 0-10) at 12 weeks	92%	Unclear		91%	Unclear		Not estimable
Night pain (number of participants with no pain as measured on VAS 0-10) at 4 weeks	36%	Unclear		45%	Unclear		Not estimable

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# Table 18. One type of manual therapy or exercise versus another (Continued)

Night pain (number of participants with no pain as measured on VAS 0-10) at 12 weeks	88%	Unclear		82%	Unclear		Not estimable
Pain on motion (number of participants with no pain as measured on VAS 0-10) at 4 weeks	16%	Unclear		14%	Unclear		Not estimable
Pain on motion (number of participants with no pain as measured on VAS 0-10) at 12 weeks	36%	Unclear		41%	Unclear		Not estimable
Study ID: Struyf 2013 Intervention: scapular-focused treatment Control: stretching, muscle friction and eco	centric rotato	or cuff training					
OUTCOME	INTERVEN	ΓΙΟΝ		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Rest pain (VAS 0-10) at 4-8 weeks	1.3	1.5	10	2.3	2.6	10	-1.00 (-2.86, 0.86)
Function (SDQ-NL, 0-100 with higher scores denoting worse function) at 4-8 weeks	35	14	10	48.7	11.3	10	-13.70 (-24.85, -2.55)
Pain on motion (VAS 0-10) at 4-8 weeks	3	1.9	10	5.1	2	10	-2.10 (-3.81, -0.39)
Strength (isometric elevation strength, Newtons) at 4-8 weeks	55.79	18.71	10	74.11	34.28	10	-18.32 (-42.53, 5.89)
Study ID: Subasi 2012 Intervention: water-based exercise progra Control: land-based exercise programme	mme						
OUTCOME	INTERVEN	ΓΙΟΝ		CONTROL			EFFECT ESTIMATE
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (VAS 0-10) at 4 weeks	3.2	1.4	29	3.7	1.4	28	-0.50 (-1.23, 0.23)

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Function (SPADI total score 0-100, with higher scores denoting worse function) at 4 weeks	16.7	12.6	29	20.1	10.5	28	-3.40 (-9.41, 2.61)
Function (SPADI total score 0-100, with higher scores denoting worse function) at 8 weeks	12	9.1	29	20.9	10.2	28	-8.90 (-13.92, -3.88
Quality of life (WORC, 0-2100, where 2100 is worst score) at 4 weeks	599.7	417.5	29	739.7	332.9	28	-140.00 (-335.69, 55.69)
Quality of life (WORC, 0-2100, where 2100 is worst score) at 8 weeks	475	269.9	29	733.1	331.6	28	-258.10 (-415.37, -100.83)
Study ID: Walther 2004 Intervention: standardised self-training of Control: supervised stretching exercises OUTCOME	centring and	_	ises	CONTROL			EFFECT ESTIMATI
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Rest pain (VAS 0-100) at both 6 and 12 weeks	"No significa	ant difference be	tween groups"				
	"No signific:	ant difference be	tween groups"				
Function (Constant-Murley total score) at both 6 and 12 weeks	5						
		ant difference be	tween groups"				
both 6 and 12 weeks Pain on motion (VAS 0-100) at both 6 and 12		ant difference be 4.6	tween groups" 20	11.8	5.4	20	-0.90 (-4.01, 2.21)
both 6 and 12 weeks Pain on motion (VAS 0-100) at both 6 and 12 weeks Strength (Constant-Murley strength sub-	"No significa			11.8	5.4 NR	20	-0.90 (-4.01, 2.21) -0.4 (95% CI not es timable)
both 6 and 12 weeks Pain on motion (VAS 0-100) at both 6 and 12 weeks Strength (Constant-Murley strength sub- score 0-25) at 12 weeks Work disability (number of months with in-	"No significa 10.9	4.6	20				-0.4 (95% Cl not es

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# Table 18. One type of manual therapy or exercise versus another (Continued)

#### Study ID: Wang 2006 Intervention: customised exercises Control: standardised exercises

Ουτςομε	INTERVENTION			CONTROL		EFFECT ESTIMATE	
	Mean	SD	n	Mean	SD	n	Mean difference (95% Cl)
Overall pain (VAS 0-100) at 4 weeks	20.1	14.5	15	23.1	18	15	-3.00 (-14.70, 8.70)
Overall pain (VAS 0-100) at 8 weeks	21.6	12.5	15	21.2	17.6	15	0.40 (-10.52, 11.32
Function (Flexi-SF Score, 1-50 with a higher score indicating better function) at 4 weeks	32.3	9.8	15	32.4	6.6	15	-0.10 (-6.08, 5.88)
Function (Flexi-SF Score, 1-50 with a higher score indicating better function) at 8 weeks	36.2	6.5	15	33.8	7	15	2.40 (-2.43, 7.23)
Active range of abduction (degrees) at 4 weeks	147.1	36.5	15	140.4	40.6	15	6.70 (-20.93, 34.33
Active range of abduction (degrees) at 8 weeks	149.3	32.5	15	143.4	37.4	15	5.90 (-19.17, 30.97
Active range of external rotation (degrees) at 4 weeks	81	22.1	15	71.9	27.3	15	9.10 (-8.67, 26.87)
Active range of external rotation (degrees) at 8 weeks	81.8	18.4	15	73.1	26.9	15	8.70 (-7.79, 25.19)
Active range of internal rotation (degrees) at 4 weeks	40.6	14.7	15	45.8	14.8	15	-5.20 (-15.76, 5.36
Active range of internal rotation (degrees) at 8 weeks	44.9	15.7	15	44.5	16.9	15	0.40 (-11.27, 12.07
Strength (isometric strength of abductors in N-m) at 4 weeks	48.3	20.9	15	36.3	20	15	12.00 (-2.64, 26.64
Strength (isometric strength of abductors in N-m) at 8 weeks	53.9	21.9	15	42.2	24.4	15	11.70 (-4.89, 28.29

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Global assessment of treatment success	18	35		14	32		1.18 (0.71, 1.95)
	Events	Total		Events	Total		Risk ratio (95% C
Overall pain (shoulder pain score 0-28, high- er score denotes worse pain) at 11 weeks	11.5	4.4	35	12.6	5.1	32	-1.10 (-3.39, 1.19)
	Mean	SD	n	Mean	SD	n	Mean difference (95% CI)
OUTCOME				CONTROL			EFFECT ESTIMAT
Study ID: Winters 1997 Intervention: exercise and massage Control: manipulation							
Strength (isometric strength of lower trapezius in N-m) at 8 weeks	37.2	20.3	15	29.4	18	15	7.80 (-5.93, 21.53)
Strength (isometric strength of lower trapezius in N-m) at 4 weeks	31.8	16.3	15	26	16.2	15	5.80 (-5.83, 17.43)
Strength (isometric strength of middle trapezius in N-m) at 8 weeks	41	19.4	15	30.8	17.9	15	10.20 (-3.16, 23.56
Strength (isometric strength of middle trapezius in N-m) at 4 weeks	34.8	15.5	15	28.1	17.9	15	6.70 (-5.28, 18.68)
Strength (isometric strength of internal ro- tators in N-m) at 8 weeks	37.5	15.7	15	28.4	14.6	15	9.10 (-1.75, 19.95
Strength (isometric strength of internal ro- tators in N-m) at 4 weeks	31.7	11.9	15	28	16.6	15	3.70 (-6.64, 14.04)
Strength (isometric strength of external ro- tators in N-m) at 8 weeks	36.3	15.1	15	29.3	15.9	15	7.00 (-4.10, 18.10)
Strength (isometric strength of external ro- tators in N-m) at 4 weeks	34.3	14	15	27.7	17.4	15	6.60 (-4.70, 17.90

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#### APPENDICES

### **Appendix 1. Search strategies**

Search strategy for CENTRAL:

- 1. MeSH descriptor: [Shoulder Pain] explode all trees
- 2. MeSH descriptor: [Shoulder Impingement Syndrome] explode all trees
- 3. MeSH descriptor: [Rotator Cuff] explode all trees
- 4. MeSH descriptor: [Bursitis] explode all trees
- 5. ((shoulder\* in All Text or rotator\* in All Text) and (bursitis in All Text or frozen in All Text or impinge\* in All Text or tendonitis in All Text or tendonitis in All Text or tendinopathy in All Text or pain\* in All Text))
- 6. "rotator cuff" in All Text
- 7. "adhesive capsulitis" in All Text
- 8. #1 or #2 or #3 or #4 or #5 or #6 or #7
- 9. MeSH descriptor: [Rehabilitation] explode all trees
- 10.MeSH descriptor: [Physical Therapy Modalities] explode all trees
- 11.MeSH descriptor: [Exercise Movement Techniques] explode all trees
- 12.MeSH descriptor: [Ultrasonography, Interventional] explode all trees
- 13.rehabilitat\* in All Text or physiotherapy\* in All Text or "physical therap\*" in All Text or "manual therap\*" in All Text or exercis\* in All Text
- 14. (ultrasound in All Text or ultrasonograph\* in All Text or this in All Text or tens in All Text or shockwave in All Text or electrotherap\* in All Text or mobili\* in All Text)
- 15.#9 or #10 or #11 or #12 or #13 or #14
- 16.#8 and #15

Search strategy for MEDLINE (Ovid):

- 1. shoulder pain/
- 2. shoulder impingement syndrome/
- 3. rotator cuff/
- 4. exp bursitis/
- 5. ((shoulder\$ or rotator cuff) adj5 (bursitis or frozen or impinge\$ or tendinitis or tendonitis or tendinopathy or pain\$)).mp.
- 6. rotator cuff.mp.
- 7. adhesive capsulitis.mp.
- 8. or/1-7
- 9. exp rehabilitation/
- 10.exp physical therapy techniques/
- 11.exp musculoskeletal manipulations/
- 12.exp exercise movement techniques/
- 13.exp ultrasonography, interventional/
- 14. (rehabilitat\$ or physiotherap\$ or physical therap\$ or manual therap\$ or exercis\$ or ultrasound or ultrasonograph\$ or TNS or TENS or shockwave or electrotherap\$ or mobili\$). mp.

15.or/9-14

- 16.clinical trial.pt
- 17.random\$.mp.
- 18.((single or double) adj (blind\$ or mask\$)).mp.

19.placebo\$.mp.

20.or/16-19

21.8 and 15 and 20

Search strategy for EMBASE (Ovid):

- 1. 'shoulder pain'/exp
- 2. 'shoulder impingement syndrome'/exp
- 3. 'rotator cuff'/exp

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- 4. 'bursitis'/exp
- 5. ((shoulder\* OR rotator\*) AND ('bursitis'/de OR frozen OR impinge\* OR 'tendonitis'/de OR 'tendinitis'/de OR 'tendinopathy'/de OR pain\*))
- 6. 'rotator cuff'
- 7. 'adhesive capsulitis'
- 8. #1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7
- 9. 'rehabilitation'/exp
- 10.'physiotherapy'/exp
- 11. 'kinesiotherapy'/exp
- 12.'endoscopic echography'/exp
- 13. rehabilitat\* OR physiotherapy\* OR 'physical therapy' OR 'manual therapy' OR kinesiotherap\* OR exercis\*
- 14. 'ultrasound'/de OR ultrasonograph\* OR 'transcutaneous nerve stimulation' OR 'transcutaneous electrical nerve stimulation' OR shockwave OR electrotherap\* OR mobili\*
- 15.#9 OR #10 OR #11 OR #12 OR #13 OR #14
- 16.'randomized controlled trial'/exp
- 17.#8 AND #15 AND #16

Search strategy for CINAHL Plus (EBSCO):

- S1 MH "shoulder pain"
- S2 MH "shoulder impingement syndrome"
- S3 MH "rotator cuff"
- S4 MH bursitis+
- S5 TX (shoulder\* N5 bursitis) or TX(shoulder\* N5 frozen) or TX(shoulder\* N5 impinge\*) or TX(shoulder\* N5 tend?nitis) or TX(shoulder\* N5 tendinopathy) or TX(shoulder\* N5 pain\*)
- S6 TX (rotator cuff N5 bursitis) or TX(rotator cuff N5 frozen) or TX(rotator cuff N5 impinge\*) or TX(rotator cuff N5 tend?nitis) or TX(rotator cuff N5 tend?nitis) or TX(rotator cuff N5 tend?nitis)
- S7 TX rotator cuff
- S8 TX adhesive capsulitis
- S9 S1 or S2 or S3 or S4 or S5 or S6 or S7 or S8
- S10 MH Rehabilitation+
- S11 MH physical therapy+
- S12 MH Manual Therapy+
- S13 MH Therapeutic Exercise+
- S14 MH Ultrasonography+
- S15 TX rehabilitat\* or physiotherapy\* or physical therap\* or manual therap\* or exercise\* or ultrasound or ultrasonograph\* or TNS or TENS or shockwave or electrotherapy\* or mobili\*
- S16 S10 or S11 or S12 or S13 or S14 or S15
- S17 PT clinical trial
- S18 TX random\*
- S19 TX(single blind\*) or TX(single mask\*)
- S20 TX(double blind\*) or TX(double mask\*)
- S21 placebo\*
- S22 S17 or S18 or S19 or S20 or S21
- S23 S9 and S16 and S22

#### WHAT'S NEW

Date	Event	Description
27 May 2016	New search has been performed	The original review, 'Physiotherapy interventions for shoulder pain' (Green 2003) was split into four reviews upon updating: this review, 'Manual therapy and exercise for rotator cuff dis- ease', 'Electrotherapy modalities for rotator cuff disease' (ongo- ing), 'Manual therapy and exercise for adhesive capsulitis (frozen

Manual therapy and exercise for rotator cuff disease (Review)

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Date	Event	Description
		shoulder)' (Page 2014a), and 'Electrotherapy modalities for adhe- sive capsulitis (frozen shoulder)' (Page 2014b). The review has al- so been broadened by including all randomised and quasi-ran- domised clinical trials regardless of whether outcome assess- ment was blinded.

#### HISTORY

Review first published: Issue 6, 2016

Date	Event	Description
1 May 2008	Amended	Converted to RM5. CMSG ID C067-R
24 February 2003	New citation required and conclusions have changed	Substantive amendment
24 February 2003	Amended	This review is based on the original review of 'Interventions for shoulder pain'. Please see published notes for further details.

#### CONTRIBUTIONS OF AUTHORS

MJP was responsible for writing the review, performing the searches, selecting trials, performing risk of bias assessment, data extraction, analysing the data and interpreting the results of the updated review. SG was responsible for performing the searches, selecting trials and performing the data extraction and quality assessment for the original review, defining the review comparisons and outcomes of interest of the original and updated review, analysing and interpreting the results, and contributing to writing both the original and updated review. BM was responsible for selecting trials, performing risk of bias assessment, data extraction and contributing to writing the manuscript for the updated review. SS, JD, NL and MM were responsible for performing risk of bias assessment, data extraction and quality assessment for the original review, defining the review comparisons and outcomes of interest of both the original and updated review. RB was responsible for performing the data extraction and quality assessment for the original review, defining the review comparisons and outcomes of interest of both the original and updated review. RB was responsible for performing the data extraction and quality assessment for the original review, defining the review comparisons and outcomes of interest of both the original and updated review, analysing and interpreting the review, defining the review comparisons and outcomes of interest of both the original and updated review, analysing and interpreting the results, and contributing to writing both the original and updated review.

#### DECLARATIONS OF INTEREST

SG and RB are authors of one of the trials included in this review (Bennell 2010). To avoid any bias, the paper was sent to an independent review author to assess whether it met the inclusion criteria for this review. Neither author was involved in the data extraction or risk of bias assessment of this trial. RB is Joint Co-ordinating Editor of Cochrane Musculoskeletal. To avoid bias, RB was excluded from the editorial and publication process for this review. SG is a practicing physiotherapist in part-time private physiotherapy practice (self-employed), and as such receives remuneration for the delivery of physiotherapy interventions. BM is a practicing physiotherapist in private physiotherapy practice and as such receives remuneration for the delivery of physiotherapy interventions.

#### SOURCES OF SUPPORT

#### **Internal sources**

- Australasian Cochrane Centre, School of Public Health and Preventive Medicine, Monash University, Melbourne, Australia.
- Department of Epidemiology and Preventive Medicine, Monash University, Melbourne, Australia.

#### **External sources**

• Australian National Health and Medical Research Council (NHMRC) Early Career Fellowship (1088535), Australia.

#### DIFFERENCES BETWEEN PROTOCOL AND REVIEW

The original review outcomes were pain, range of motion (active and passive), function/disability and quality of life, strength, return to work, participants' perception of overall effect, global preference, physicians' preference and adverse effects. The outcomes reported in

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this review have been modified from the original review to make them as consistent as possible with other Cochrane reviews on shoulder disorders and other chronic pain conditions. To improve succinctness of the review, we only included one measurement instrument per outcome domain. We assessed study risk of bias using The Cochrane Collaboration's 'Risk of bias' tool in this update of the review. We have included a 'Summary of findings' table.

#### NOTES

The original review, 'Physiotherapy interventions for shoulder pain' (Green 2003) was split into four reviews upon updating: this review, 'Manual therapy and exercise for rotator cuff disease', 'Electrotherapy modalities for rotator cuff disease' (ongoing), 'Manual therapy and exercise for adhesive capsulitis (frozen shoulder)' (Page 2014a), and 'Electrotherapy modalities for adhesive capsulitis (frozen shoulder)' (Page 2014a), and 'Electrotherapy modalities for adhesive capsulitis (frozen shoulder)' (Page 2014b). The review has also been broadened by including all randomised and quasi-randomised clinical trials regardless of whether outcome assessment was blinded.

#### INDEX TERMS

#### Medical Subject Headings (MeSH)

\*Musculoskeletal Manipulations; \*Rotator Cuff; Exercise Therapy [\*methods]; Muscular Diseases [\*therapy]; Randomized Controlled Trials as Topic; Shoulder Pain [etiology] [\*therapy]

#### **MeSH check words**

Adult; Humans; Male; Middle Aged