Effects of eccentric exercise in patients with subacromial impingement syndrome: a systematic review and meta-analysis

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Abstract

Background: Subacromial impingement syndrome is a common problem in primary healthcare. It often include tendinopathy. While exercise therapy is effective for this condition, it is not clear which type of exercise is the most effective. Eccentric exercises has proven effective for treating similar tendinopathies in the lower extremities. The aim of this systematic review was therefore to investigate the effects of eccentric exercise on pain and function in patients with subacromial impingement syndrome compared with other exercise regimens or interventions. A secondary aim was to describe the included components of the various eccentric exercise regimens that have been studied.

Methods: Systematic searches of PubMed, Cochrane Library and PEDro by two independent authors. Included studies were assessed using the PEDro scale for quality and the Cochrane scale for clinical relevance by two independent authors. Data were combined in meta-analyses. GRADE was applied to assess the certainty of evidence.

Results: Sixty-eight records were identified. Seven studies (eight articles) were included, six were meta-analysed (n = 281). Included studies were of moderate quality (median PEDro score 7, range 5–8). Post-treatment pain was significantly lower after eccentric exercise compared with other exercise: MD -12.3 (95% CI −17.8 to −6.8, I² = 7%, p < 0.001), but this difference was not clinically important. Eccentric exercise provided no significant post-treatment improvement in function compared with other exercise: SMD -0.10 (95% CI −0.79 to 0.58, I² = 85%, p = 0.76). Painful eccentric exercise showed no significant difference compared to pain-free eccentric exercise. Eccentric training regimes showed both similarities and diversity. Intervention duration of 6–8 weeks was almost as effective as 12 weeks.

Conclusions: Evidence of low certainty suggests that eccentric exercise may provide a small but likely not clinically important reduction in pain compared with other types of exercise in patients with subacromial impingement syndrome. It is uncertain whether eccentric exercise improves function more than other types of exercise (very low certainty of evidence). Methodological limitations of existing studies make these findings susceptible to change in the future.

Trial registration: PROSPERO CRD42019126917, date of registration: 29-03-2019.

Keywords: Subacromial impingement syndrome, shoulder impingement syndrome, Subacromial pain syndrome, Eccentric exercise, Eccentric training
Background
Subacromial impingement syndrome is a common healthcare problem, especially in adult populations. Prevalence is estimated to between seven and 26% of the general population [1], and almost half of all shoulder-related pain in patients seeking primary health care is related to subacromial impingement syndrome [2]. A thorough understanding of the treatment of this condition is therefore of importance for physiotherapists and other healthcare personnel.

Subacromial impingement syndrome is a condition where the subacromial space, the area directly below the acromion process and above the shoulder joint, has narrowed. This can happen for several reasons, traditionally categorised as either primary or secondary causes. Primary causes are structural changes or morphological pre-conditions of the acromion process [3, 4]. Secondary causes often depend on multiple factors such as rotator cuff syndrome (strains or tears to the muscles and/or tendons of the muscles making up the rotator cuff), tendinopathy (inflammation or degeneration) in one or more of the same muscle tendons (infra-spinatus, supraspinatus or subscapularis) and/or inflammation of the subacromial bursa [5, 6]. The long head/tendon of the biceps brachii can also be involved [6].

Eccentric exercises are exercises performed only during the elongation phase of muscle activation (i.e. the lowering or slowing-down phase of a limb) and normally at a high intensity. A possible hypothesis for their benefits is that they could potentially reverse painful neovascularization within damaged tendons, which has been shown in a study on eccentric exercise and Achilles tendinopathy [7]. Eccentric exercises have also been shown to decrease swelling of the Achilles tendon [8].

Due to their high intensity, and the fact that collagen growth in tendons tends to peak 24 to 72 h after training [9], enough time for recovery seems vital for effective rehabilitation with eccentric exercises. It could therefore be expected that in eccentric exercise regimens, not only how the exercise is performed (eccentric versus concentric phase) and its intensity, but also its frequency and duration will be of importance.

It also has been proposed that injuries to tendons and other soft tissue are the most common causes of long-term shoulder pain in general. Histological examinations of tendon injuries of the supraspinatus in patients with subacromial impingement syndrome have shown changes to the tendon resembling those in patients with similar injuries of the patellar and Achilles tendon [10]. Patellar and Achilles tendinopathy are two types of tendon injuries where high intensity eccentric exercise has been shown effective, not only in decreasing pain but also in stimulating tissue regeneration and restoring function [11–13]. This makes it relevant to investigate whether eccentric exercises could be equally effective in treating patients with subacromial impingement syndrome.

Earlier studies have shown that exercise, in general, is effective in treating subacromial impingement syndrome [14], at least as effective as corticosteroid injections for treating pain [15] and equally long-term effective as surgery [16]. Eccentric exercise in particular also has shown promising results in three uncontrolled studies [17–19].

It is not clear at present which type of exercise/training is the most effective for subacromial impingement syndrome, and whether this differs depending on involved structures and underlying mechanisms [14, 16, 20, 21]. Previous reviews [22–24] on eccentric exercise and subacromial impingement syndrome have only had access to a limited set of data, up to two randomised controlled trials (RCTs) [25, 26] and one or more of the uncontrolled studies mentioned above [17–19]. A new review including recently published studies and incorporating a meta-analysis, not previously performed, would therefore be able to generate new knowledge, especially since this is a relatively new field of research [22]. Therefore, the aim of this systematic review was to investigate the effects of eccentric exercise on pain and function in patients with subacromial impingement syndrome compared with other exercise regimens or interventions. A secondary aim was to describe the included components of the various eccentric exercise regimens that have been studied.

Methods
Protocol and registration
A protocol for this systematic review was registered in PROSPERO (PROSPERO 29-03-2019: CRD42019126917). The review was conducted and reported according to the Preferred Reporting Items for Systematic Review and Meta-Analysis statement [27].

Eligibility criteria
Participants: adult men and women, i.e. individuals from 18 years of age and upwards, with shoulder pain and diagnosed with subacromial impingement syndrome, defined as shoulder pain and a positive Neer’s impingement test, Hawkins-Kennedy impingement test, Empty can test (Jobe’s impingement test), Painful arc sign and/or positive response to a subacromial corticosteroid injection. Included individuals should be part of the general population and not only part of a specific subgroup, e.g. swimmers or tennis players, to allow for greater clinical relevance and applicability of the findings to the general population.

Intervention: eccentric exercise. The exercise interventions needed to be described in sufficient detail so that both exercise and execution could be clearly identified and so that the eccentric loading could be easily compared with any control, whether resistance training, other forms of eccentric exercise (e.g. pain versus no pain), or any other intervention. Eccentric exercises had to be the primary treatment method.
Comparisons: Other types of exercise (e.g. resistance, mobility, aerobic); other interventions (e.g. massage, mobilization/manipulation, acupuncture, TENS, corticosteroid injections); other types of eccentric exercise (where exercising according to different ratings of perceived pain have been compared).

Outcome measures: Pain, e.g. measured by visual analogue scale (VAS) or numerical pain rating scale (NPRS); function, e.g. measured by the disabilities of the arm, shoulder and hand (DASH) questionnaire or the Constant-Murley score; main components of the various eccentric exercise regimens.

Exclusion criteria were any uncontrolled study designs and shoulder pain due to fractures, dislocations or medical conditions (e.g. osteoarthritis, rheumatoid arthritis). Only RCTs qualified for inclusion because we aimed to assess intervention effects. Identification of any controlled but not randomised studies was reported. No limitation to population size was set.

**Literature search**

We conducted literature searches in the databases PubMed, Cochrane Library and the Physiotherapy Evidence Database (PEDro) in March 2019. We developed a search strategy for PubMed and subsequently adapted it to the other databases (Appendix 1). The search strategy combined search terms with medical subject headings and comprised a combination of the term "subacromial impingement", or synonyms thereof, with the term "eccentric exercise", or synonyms thereof. We did not apply any restrictions to language or date of publication. We performed backward and forward citation searches of included studies and identified previous reviews for additional potentially relevant studies, and searched ClinicalTrials.gov for ongoing studies. We also consulted content experts in the field.

**Study selection**

Two authors (RL and SB) screened the identified titles and abstracts independently for relevance (according to the inclusion/exclusion criteria) and assessed full text articles for inclusion. The authors were not blinded to trial identifiers such as authors’ and journals’ names.

**Data collection process**

Two authors (RL and SB) performed data extraction independently. Extracted data included study aim, population demographics (including age and sex), intervention components, control group intervention components, outcome measures and outcome data. When needed, we solicited and obtained additional data from trial investigators.

**Risk of bias assessment**

We assessed risk of bias in the included studies in duplicate (RL and SB or LN), using the PEDro scale for quality [28]. This instrument has been shown to have acceptably high reliability and validity [29, 30]. We also assessed clinical relevance using the Cochrane scale for clinical relevance [31]. Any disagreements were resolved by discussion among all authors until consensus was reached. To assess agreement among the reviewers, percentage agreement and Cohen’s kappa with 95% confidence intervals (CI) was calculated. The results of the risk of bias assessment was used, together with other criteria, in the overall assessment of the certainty of the evidence.

**Data synthesis and analysis**

When possible, we combined the data in meta-analyses for investigation of the aggregated post-treatment and intermediate to long-term effects. Because one study [32] compared painful versus pain-free eccentric exercise, we excluded it from the meta-analysis. When articles reported several different measures for pain, we extracted and tabulated all measures, but we chose one for the meta-analyses. “Worst pain” or “pain during activity” was chosen because we considered this measure to best correspond to “pain” in the other included studies. We also considered this to be the outcome most important to the patient.

We analysed pain by calculating the MD with corresponding 95% CI. To be able to present the aggregated effect, we rescaled the NPRS data from 0 to 10 to 0–100. When median and percentiles were reported instead of mean and SD, we approximated the missing mean and SD with the corresponding median and percentile range and imputed these values [33]. We assumed a normal distribution of pain scores on the VAS or the NPRS, and we considered a 15 mm difference on the VAS as representative of a minimal important difference (MID) in pain [34].

For function, five different instruments were used in the seven studies. Due to the many instruments, we calculated the aggregated effect using standardised mean difference (SMD) with 95% CI. Because the scales went in different directions, mean values were multiplied by –1 when necessary.

Statistical heterogeneity was assessed with the $\chi^2$ and $I^2$ statistics. Because heterogeneity was present ($I^2 > 30\%$), we used random-effect models. When possible, missing data were handled by imputing values from previous time-points, applying the “last observation carried forward” principle. We analysed outcomes post-treatment in two subgroups which were not pre-planned: six to 8 weeks and 12 weeks. Intermediate and long-term outcomes were analysed when reported, as time points closest to 1 year. We explored whether excluding high risk of bias trials would affect the result. We performed the meta-analyses in Revman 5.3 [35].

**Assessment of certainty of evidence**

To assess confidence in the combined estimates of effect, we applied the GRADE (Grading of Recommendations Assessment, Development and Evaluation) approach using the
following criteria: risk of bias, consistency, directness, precision, and reporting bias [36]. Because all included studies were RCTs, we initially assigned a high certainty level, but rated down one or more levels to moderate, low or very low if we detected issues with risk of bias, precision, consistency or directness. Publication bias was not assessed due to the small number of studies, but was not considered likely.

Results
Search results
The search procedure yielded 68 records, of which 51 unique articles remained after removing duplicates. After screening titles and abstracts for relevance, and when necessary assessing the full text articles, 43 articles could be excluded according to inclusion/exclusion criteria. No controlled studies that were not also randomised were identified. Scrutiny of reference lists identified one potentially relevant study, but on closer inspection it did not fulfil the inclusion criteria. In total, we included seven studies reported in eight articles [25, 26, 32, 37–41]. The search process and results are illustrated in Fig. 1. Of the included studies, six were included in the meta-analyses [25, 26, 37–41].

Characteristics of included studies
The seven included studies were conducted between 2012 and 2018. Two were conducted in the United States and five in Europe. Of the seven studies, six [25, 26, 37–40] compared eccentric exercise with other exercise. One [32] compared painful eccentric exercise (above the pain threshold: VAS < 40–50 mm) with pain-free eccentric exercise (VAS = 0 mm). For one of the studies, long-term results were reported in a separate article [41].

Risk of bias
Included studies had a PEDro score of between five and eight (out of 10), with a median value of seven (Table 1). Agreement among the reviewers was excellent, 95% (Cohen’s kappa 0.89; 95% CI 0.78 to 0.99). None of the studies had been able to blind patients nor therapists, but two studies [26, 38] had blinded the assessors. Three studies [26, 37, 40] did not use intention-to-treat analysis. In one study [25] groups differed at baseline and one study [40] did not publish measures of variability. On the Cochrane scale for clinical relevance [31] all studies were assigned
Table 1  Characteristics and risk of bias of included studies

| Author, year, country | Participants | Intervention (details in Table 2) | Control | Outcome measures | Follow-up period | PEDro score for quality (details in 2) | Cochrane score for clinical relevance (details in Appendix 3) |
|-----------------------|--------------|-----------------------------------|---------|-----------------|-----------------|----------------------------------------|-----------------------------------------------------------|
| Bateman et al. 2014, United Kingdom [38] | n = 11, not responded to previous conservative treatment, on waiting list for operation, image verified rotator cuff tendinopathy or positive response to subacromial corticosteroid injection. Intervention 4 individuals, mean age 52 years; control_1 3 individuals, mean age 53 years; control_2 4 individuals, mean age 55 years. | Eccentric exercise; Intensity: pain reproduction | Resistance exercise (concentric phase only); Intensity matched experiment group | Pain: VAS; Function: Oxford Shoulder Score | 8 weeks (post-treatment) | 5/10 | 4/5 |
| Blume et al. 2015, United States [35] | n = 34, shoulder pain, positive Neer’s, Hawkins or cross-body adduction test. Intervention 10 women 8 men, mean age 50.1 years (SD 16.9); control 10 women 6 men, mean age 48.6 years (SD 14.6). | Eccentric exercise; Intensity: 70–80% of 1RM | Resistance exercise (concentric phase only); Intensity matched experiment group | Pain: NPRS; Function: DASH | 8 weeks (post-treatment) | 8/10 | 5/5 |
| Chaconas et al. 2017, United States [34] | n = 46, shoulder pain at least 3 months, 3 or more positive: Neer’s, Hawkins, empty can test, painful arc sign, shoulder external rotation pain, palpation pain supra/infraspinatus. Intervention 10 women 15 men, mean age 43.4 years (SD 17.9); control 9 women 12 men, mean age 48.4 years (SD 16.9). | Eccentric exercise; Intensity: 65% of 1RM | Resistance exercise; Intensity lower than experiment group | Pain: NPRS; Function: WORC | 6 weeks (post-treatment) & 6 months (intermediate term follow-up) | 6/10 | 5/5 |
| Dejaco et al. 2017, Netherlands [37] | n = 36, subacromial pain at least 3 months, 2 of 3 positive: Neer’s, Hawkins, empty can test. Intervention 10 women 10 men, mean age 50.2 years (SD 10.8); control 7 women 9 men, mean age 48.6 years (SD 12.3). | Eccentric exercise; Intensity: pain reproduction | Resistance exercise; Intensity matched experiment group | Pain: VAS; Function: Constant-Murley Score | 12 weeks (post-treatment) & 6 months (intermediate term follow-up) | 7/10 | 5/5 |
| Hallgren et al. 2014, Sweden [50] | Reporting on the same study participant, intervention and control as Hallgren et al 2012 | | | Pain: VAS; Function: Constant-Murley Score | 1 year (long-term follow-up) | 7/10 | 5/5 |
| Holmgren et al. 2012, Sweden [26] | n = 97, diagnosed with primary SIS, shoulder pain at least 6 months, not responded to previous conservative treatment, on waiting list for operation, positive Neer’s injection test, 3 of 4 positive: Neer’s, Hawkins, empty can test, Patte’s manoeuvre. Intervention 14 women 37 men, mean age 52 years (SD 9); control 22 women 24 men, mean age 52 years (SD 8). | Eccentric exercise; Intensity: pain reproduction; Corticosteroid injection | Mobility exercise; Intensity lower than experiment group; Corticosteroid injection | Pain: VAS; Function: Constant-Murley Score | 12 weeks (post-treatment) | 7/10 | 5/5 |
| Maenhout et al. 2013, Belgium [25] | n = 61, shoulder pain at least 3 months, painful arc sign, palpation pain supra/infraspinatus, 2 of 3 positive: Neer’s, Hawkins, empty can test and 2 of 4 painful resistance tests. Intervention 16 women 15 men, mean age 40.2 years (SD 12.9); control 20 women 10 men, mean age 39.4 years (SD 13.1). | Eccentric exercise; Intensity: pain reproduction | Resistance exercise; Intensity matched experiment group | Pain and function: SPADI | 12 weeks (post-treatment) | 6/10 | 5/5 |
| Vallés-Carrascosa et al. 2018, | n = 22, diagnosed with SIS, positive painful arc sign. Intervention 8 women 3 men, mean age 57 years; control | Eccentric exercise, pain up to 40–50 mm | Eccentric exercise, pain free execution | Pain: VAS; Function: Constant-Murley Score | 4 weeks (post-treatment) | 7/10 | 5/5 |
scores of five out of five, except one [40], who lost one point due to the effect size being smaller than the MID for main outcomes. Complete PEDro scores and Cochrane relevance scores are available in Appendix 2 and 3.

**Participants**

The seven included studies involved a total of 303 participants with subacromial impingement syndrome; 140 women and 156 men (one study [40] did not specify sex of the participants). Mean age of participants ranged from 39 to 60 years. Upon enrolment, pain duration varied from 3 months [25, 37, 39] to 6 months [26], with three studies [32, 38, 40] not specifying any timeframe. Pain intensity ranged from 12 mm VAS (“best pain”, rescaled from NPRS) to 70 mm VAS (“worst pain”). Disability levels at enrolment were mild to moderate. Detailed characteristics of included studies are presented in Table 1.

**Interventions**

Exercise regimens in the included studies showed both similarities and differences. All interventions focused on one or both of the following exercises: shoulder external rotation with shoulder in neutral position and shoulder abduction in the scapular plane with thumb pointing up (full can exercise). Elastic exercise bands and/or dumbbells were used in all studies. Duration of the intervention varied between four and 12 weeks and frequency varied between two times per week to two times per day. Exercise intensity was specified in two studies at 65% and 70–80% of 1RM (one repetition maximum) respectively [37, 38]. All other studies used pain reproduction to specify exercise intensity, i.e. allowing or encouraging pain during exercise that matched the pain normally felt by the participant in everyday activities, as long as it did not exceed 50 mm on VAS.

Comparators in all included studies consisted of other exercise interventions (one study [40] also included a second control group who did not receive any intervention, these participants are not included in the meta-analyses). In five studies [25, 37–40] the comparator was resistance exercise (concentric or concentric/eccentric). In one study [32] it was pain-free eccentric exercise (the experimental group performed painful eccentric exercise). In all aforementioned studies frequency, intensity and type of training were matched to the eccentric exercise protocol of the experimental group, except one study [37], in which the intensity was markedly lower. In the last study [26], the control group performed mobility exercises. In five of the studies [26, 32, 37–39], resistance training was complemented with stretching for both experimental and control group. In one of these [26], both experimental and control group also received a corticosteroid injection as well as advice on ergonomics and posture. Details on exercise regimens are presented in Table 2.

**Outcome measures**

Pain and function were measured in all included studies. In four studies, as well as the one-year follow up [26, 32, 39–41], pain was measured using a 100 mm VAS [34]. In two studies [37, 38], a 10-point NPRS [42] was used. In one study [25] pain was measured with the Shoulder Pain and Disability Index (SPADI [43]) pain subscale.

For function, five different instruments were used in the seven studies: the Oxford Shoulder Score (OSS [44]), the Disability of the Arm, Shoulder and Hand (DASH [43]), the Western Ontario Rotator Cuff (WORC) index [45], the Constant-Murley (CM) Score [46], and the SPADI total score [43]. All measures have been validated.

**Summary of findings**

Summary of findings for the main comparisons are presented in Table 3 and described below for each outcome.

**Effects of eccentric exercise on pain**

Effects on all measures of pain are presented in Appendix 4. Although treatment lengths in the studies varied between four and 12 weeks, six of the included studies [25, 26, 37–40] measured post-treatment pain and were considered to be of sufficient clinical homogeneity to be combined. Intermediate to long-term effects (6–12 months follow-up) were measured in three studies [37, 39, 41]. Six studies with 281 participants were included in the meta-analysis of post-treatment effects on pain. In the three studies [37, 38, 40] with treatment lengths of six to 8 weeks, eccentric exercise provided no significant reduction in pain compared with other exercise
Table 2 Summary of eccentric training regimens

| Study                   | Duration | Frequency | Intensity | Sets/ reps | Equipment                        | Exercise(s)                                                                 |
|-------------------------|----------|-----------|-----------|------------|----------------------------------|-----------------------------------------------------------------------------|
| Bateman et al. (2014)   | 8 weeks  | 2 times/day | Same intensity for all participants, increased for all after 4 weeks, pain during execution allowed but not specified | 3 × 15 | Elastic exercise band; yellow and red | Full can with elastic band, only eccentric phase                           |
| Blume et al. (2015)    | 8 weeks  | 2 times/week | Approx. 70–80% of 1RM, no increase of pain during execution | 3 × 12 | Dumbbells                        | Full can; sidelying ER; sidelying IR; supine protraction; sidelying horizontal abduction; sidelying abduction; prone shoulder extension; all with dumbbells, only eccentric phase + Pectoralis minor and posterior shoulder stretch, thoracic spine extension, pain-free AROM in flexion and abduction (accessory exercises done daily) |
| Chaconas et al. (2017) | 6 weeks  | 1 time/day | Approx. 65% of 1RM (15–18 RM), no increase of pain during execution | 3 × 15 | TheraBand; green, blue, black, silver, gold | ER with elastic band, only eccentric phase + Scapular retraction with elastic band, cross body posterior shoulder stretch |
| Dejaco et al. (2017)   | 12 weeks | 2 times/day | Pain reproduction, but not over 5 of 10 (NPRS) during execution | 3 × 8–15 | Duraband Servofit + dumbbell 1 kg (when needed) | ER with elastic band, only eccentric phase; empty can with/without dumbbell, only eccentric phase |
| Larsson et al. BMC Musculoskeletal Disorders (2018) | 8 weeks  | 2 times/week | Same intensity for all participants, increased for all after 4 weeks, pain during execution allowed but not specified | 3 × 15 | Elastic exercise band; yellow and red | Full can with elastic band, only eccentric phase |
| Hallgren et al. (2014) | Same intervention as in Holmgren et al (2012), long-term results | | | | | |
| Holmgren et al. (2012) | 12 weeks | 2 times/day | Pain reproduction, but not over 5 of 10 (VAS/NPRS) during execution | 3 × 15 | Dumbbells + elastic exercise band | Full can with dumbbell, only eccentric phase; ER with elastic band, only eccentric phase + 3 scapular stabilization exercises; posterior shoulder stretch |
| Maenhout et al. (2013) | 12 weeks | 2 times/day | Pain reproduction, but not over 5 of 10 (VAS/NPRS) during execution | 3 × 15 | Dumbbells + TheraBand: different colors | Full can with dumbbell, only eccentric phase + ER/IR with elastic band, no increase of pain during execution (3 × 10, once daily) |
| Vallés-Carrascosa et al. (2018) | 4 weeks (painful eccentric exercise) | 5 times/week | Pain reproduction, but not over 40–50 mm VAS during execution | 3 × 10 | Dumbbells + elastic exercise band | Full can with dumbbell, only eccentric phase + ER/IR with elastic band; 2 scapular stabilization exercises; trapezius-stretch |
|同一研究的不同颜色 | 4 weeks (pain-free eccentric exercise) | 5 times/week | No pain during execution, VAS = 0 mm | 3 × 10 | Dumbbells + elastic exercise band | Full can with dumbbell, only eccentric phase + ER/IR with elastic band; 2 scapular stabilization exercises; trapezius-stretch |

Full can/empty can = shoulder abduction in scapular plane (scaption) with a thumbs up/thumbs down position; ER/IR external/internal rotation with neutral shoulder and elbow flexed 90°; RM Repetition Maximum; VAS Visual Analogue Scale; NPRS Numerical Pain Rating Scale

regimens: MD -13.5 (95% CI = 28.5 to 1.4, I² = 55%, p = 0.08). In the three studies [25, 26, 39] with treatment length of 12 weeks, eccentric exercise provided significant reduction in pain compared with other exercise regimens: MD -11.9 (95% CI = -18.2 to -5.5, I² = 0%, p < 0.001). In total, post-treatment pain was significantly lower after eccentric exercise compared with other exercise regimens: combined MD -12.3 (95% CI = -17.8 to -6.8, I² = 7%, p < 0.001) (Fig. 2). However, because the MD did not surpass the MID of 15 mm on VAS [34], the effect was not considered clinically relevant.

Three studies [37, 39, 41] with 167 participants were included in the meta-analysis of intermediate to long-term effects on pain, assessed at six or 12 months. Eccentric exercise did not provide an overall significant intermediate to long-term reduction in pain compared with other exercise regimens: MD -4.9 (95% CI = -15.4 to 5.6, I² = 50%, p = 0.36) (Fig. 3).

Certainty of evidence was assessed as low, for both short-term and intermediate to long-term effects on pain. We downgraded one level for risk of bias because all studies are at unclear risk of bias, mainly due to lack of blinding. We downgraded one level for imprecision because the 95% CI included both no clinically relevant effect (MID less than 15 mm on VAS) and clinically relevant effect (MID greater than 15 mm on VAS).

The study that compared painful versus pain-free eccentric exercise [32] only measured effects immediately after the four-week long intervention, and showed no significant difference in pain between experimental and control group. According to the PEDro scale it was assessed as having a low risk of bias.
Table 3 Summary of findings for the comparison eccentric exercise versus control exercise for subacromial impingement syndrome

| Outcomes, time frame | Absolute effect estimates (95% CI) | No. of participants (studies) | Certainty in effect estimates (GRADE) | Conclusion |
|----------------------|-----------------------------------|-------------------------------|--------------------------------------|------------|
| Pain: post-treatment (6–12 weeks) Measured by VAS or NPRS, converted to VAS 0–100 mm (lower better) MID: 15 mm | Mean post-treatment pain ranged across control groups from 15.0 to 63.9 mm Mean post-treatment pain in the experimental group was 12.3 mm lower (17.8 lower to 6.8 lower) | 281 (6 studies) | Low\(^a, b\) | Eccentric exercise may provide a small but likely not important reduction in pain post-treatment compared with other types of exercise. |
| Pain: intermediate to long-term (6–12 months) Measured by VAS or NPRS, converted to VAS 0–100 mm (lower better) MID: 15 mm | Mean intermediate/long-term pain ranged across control groups from 18.0 to 52.1 mm Mean intermediate/long-term pain in the experimental group was 4.9 mm lower (15.4 lower to 5.6 higher) | 167 (3 studies) | Low\(^a, b\) | Eccentric exercise may result in little or no important difference in pain compared with other types of exercise. |
| Function: post-treatment (6–12 weeks) Multiple scales of various range | Standardised mean post-treatment function in the experimental group was 0.10 SMD units better (0.79 better to 0.58 worse) | 281 (6 studies) | Very low\(^a, b, c\) | It is uncertain whether eccentric exercise improves function more than other types of exercise post-treatment follow-up. |
| Function: intermediate to long-term (6–12 months) Multiple scales of various range | Standardised mean intermediate/long-term function in the experimental group was 0.28 SMD units worse (0.67 better to 1.24 worse) | 167 (3 studies) | Very low\(^a, b, c\) | It is uncertain whether eccentric exercise improves function more than other types of exercise at intermediate/long-term follow-up. |

GRADE Working Group grades of evidence

- **High certainty**: We are very confident that the true effect lies close to that of the estimate of the effect
- **Moderate certainty**: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different
- **Low certainty**: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect
- **Very low certainty**: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of the effect

\(^a\)Downgraded one level due to serious risk of bias (mainly due to lack of blinding)

\(^b\)Downgraded one level due to serious imprecision (high heterogeneity in magnitude and direction of effect across studies, wide CIs, small study sizes)

\(^c\)Downgraded one level due to clear inconsistency of results

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**Fig. 2** Forest plot of post-treatment effects on pain, sub-grouped by treatment length of six to eight weeks and treatment length of 12 weeks. Dashed vertical line denotes minimal important difference (VAS 15 mm). IV = inverse-variance, VAS = visual analogue scale.
No studies reported any negative effects of (increased) pain during exercise, nor any other side effects.

Effects of eccentric exercise on function
Effects on function are presented in Appendix 5. Six studies with 281 participants were included in the meta-analysis of post-treatment effects on function; three studies [37, 38, 40] with treatment lengths of six to 8 weeks and three studies [25, 26, 39] with treatment length of 12 weeks. The meta-analysis shows considerable variation in effects. Eccentric exercise provided no significant post-treatment improvement in function compared with other exercise regimens, regardless of whether the intervention lasted six to 8 weeks or 12 weeks: SMD -0.10 (95% CI -0.79 to 0.58, \(I^2 = 85\%), p = 0.76\) (Fig. 4).

Three studies [37, 39, 41] with 167 participants were included in the meta-analysis of intermediate to long-term effects on function, assessed at six or 12 months. The meta-analysis showed considerable variation in effects. Eccentric exercise provided no significant intermediate or long-term improvement in function compared with other exercise regimens: SMD 0.28 (95% CI -0.67 to 1.24, \(I^2 = 87\%), p = 0.56\) (Fig. 5).

Certainty of evidence for short-term and intermediate to long-term effects on function was assessed as very low. We downgraded one level for risk of bias because all studies are at unclear risk of bias, mainly due to lack of blinding. We downgraded one level for imprecision because the 95% CI favoured both eccentric and control exercise. We also downgraded one level for inconsistency because of the variance across studies in both direction and size of the effect.

Discussion
Based on six trials of low to moderate risk of bias, including a total of 281 patients with subacromial impingement syndrome and comparing eccentric exercise to other types of exercise, the findings of this review and meta-analysis suggest that eccentric exercise provides slightly better effect on pain but not on function compared with other exercise. However, while the meta-analysis for pain showed that eccentric exercise reduced pain significantly more than other exercise post-treatment, this difference was not greater than the MID, and the difference was not sustained
at intermediate to long-term follow-ups. The meta-analysis for function showed no difference between eccentric exercise and other exercise. The certainty of evidence in these findings is low to very low, due to study limitations, imprecision and, for function, inconsistency across the studies. Based on one trial of low risk of bias, including 22 participants, painful eccentric exercise did not yield any significant difference compared with pain-free eccentric exercise.

Of the individual studies, Holmgren et al. [26] showed that eccentric exercise was better than non-loading mobility exercises for both pain and function and Chaconas et al. [37] showed a statistically significant and clinically important between-group difference in pain and function between strength training groups, favouring eccentric exercise. The remaining four RCTs [25, 38–40] did not find any such significant difference between groups, although a trend can be discerned across the studies in favour of eccentric exercise.

Our findings lend further support to earlier reviews that have investigated various types of exercise for subacromial impingement syndrome [14, 22–24]. In their review of eccentric exercise for subacromial impingement and lateral epicondylalgia, Ortega-Castillo & Medina-Porqueres [22] concluded that eccentric exercise may reduce pain and improve strength in upper limb tendinopathies, but that it was questionable whether it was more effective than other exercise. The certainty of evidence in these findings is low to very low, due to study limitations, imprecision and, for function, inconsistency across the studies. Based on one trial of low risk of bias, including 22 participants, painful eccentric exercise did not yield any significant difference compared with pain-free eccentric exercise.

The aggregated difference in pain reduction was similar for the six to 8 week interventions and those of 12 weeks, which may indicate that the shorter treatment length is sufficient to address pain. The fact that significance was reached for the longer intervention period but not for the shorter is likely due to the low power of those trials. None of the subgroups showed improvement greater than the minimally important difference (15 mm VAS [34]). In the 12-week study by Maenhout et al. [25], the greatest improvement was reported after the first 6 weeks, further supporting that interventions of six to 8 weeks could be sufficient to obtain measurable effects.

Although most studies focused on short-term effects of eccentric exercise, two of them reported a six-month follow-up [37, 39] and one was a one-year follow-up [41]. All found that the MD in pain between eccentric exercise and control was smaller at the follow-up, and the non-difference in function remained. Although it is likely that both experimental and control groups improved spontaneously over time, the findings suggest that after 1 year it does not matter which type of exercise you perform.

Chaconas et al. [37] have suggested that the purpose of eccentric exercise is to use training intensities that are so high that the exercises cannot be performed in the concentric phase. This is possible because individuals are on average 20% stronger during eccentric muscle contractions than during concentric contractions [47, 48]. It also has been reported that the experience of pain is less, and the reversion of pain faster, during eccentric exercises than during concentric ones [49]. Previous research also has shown that maximum intensity eccentric knee extensions (but not maximum intensity concentric extensions) may increase satellite cell activation necessary for muscle repair [50], and eccentric bench-press at 90% of 1RM may double growth hormone release, also associated with muscle repair and hypertrophy, compared with both concentric and eccentric bench-presses at 70% of 1RM [51]. However, in none of the included studies in this review were the eccentric training loads so high that they could not also be performed concentrically. In the study by Blume et al. [38], eccentric exercises were even used to calculate the intensity of the eccentric exercises, without adjusting for difference in strength. Only two of the seven included studies clearly defined training intensities at all; Blume et al. [38] used an intensity of 70–80% of 1RM for both experimental and control group, and Chaconas et al. [37] used 65% of 1RM for the eccentric exercise group. The remaining study protocols used pain reproduction to define the training intensity. Intensity of the control group exercises were relatively well

![Fig. 5 Forest plot of intermediate to long-term effects on function. Because the scales used in Dejaco 2017 and Hallgren 2014 go in the opposite direction (i.e. higher is better) than the one used in the other study, the mean values were multiplied by −1. As a rule of thumb, a standardised mean difference of 0.2 represents a small difference, 0.5 a moderate, and 0.8 a large difference. IV = inverse-variance](image-url)
matched to that of the experiment group, with two exceptions [26, 37]; these were also the two studies that showed the biggest between-group differences. It might therefore be speculated, that what matters most is not the type of exercise performed, but at what intensity.

As of yet, no studies have been published in which the effects of heavy (> 85% of 1RM) eccentric training of the rotator cuff have been investigated, but one study looking at heavy resistance training (concentric/eccentric) in patients with rotator cuff tendinopathies did not find any difference between higher (85% of 1RM) and lower (50% of 1RM) intensity resistance training [52]. Future research that aims to compare eccentric and other resistance exercise in patients with subacromial impingement syndrome should therefore focus on, and clearly define, heavy (80–90% of 1RM) eccentric exercise. Intensity should be individualised and defined as percentage of 1RM rather than aiming for pain reproduction or pain up to a certain level, since this review shows that a certain experience of pain does not indicate a better outcome than other levels of pain, or no pain at all [25, 32].

Of the seven included studies, aiming to investigate eccentric exercise for tendinopathy, only three [25, 37, 40] actually controlled whether the shoulder pain originated from a muscle tendon. This included pain on palpation of the supra- or infraspinatus tendon as a possible or required inclusion criterion, or ultrasound or MRI (magnetic resonance imaging) verified rotator cuff tendinopathy. It is thus not possible to know the proportion of participants in the different studies in whom the pain originated from a muscle tendon, and the proportion in whom structural changes to the acromion process or bursitis were the primary pathogeneses. Future studies in this field therefore need to specifically control for and verify tendinopathies in order to investigate whether there is a correlation between pain from one or more muscle tendons of the rotator cuff (or the long head of the biceps brachii) and the effects of eccentric exercise, compared with patients without a clear tendon involvement. It is likely that optimal exercise strategy will vary depending on affected tendon or other primary pathogenesis.

This review is based on small to medium-sized trials, of which the smallest only had 11 participants. A major limitation in all studies is the lack of blinding of both participants and therapists, a common problem in physiotherapy trials. Most studies also failed to blind the outcome assessors. Other limitations are the large variation in the exercise protocols and generally poor definition of training intensity, leading to high heterogeneity and precluding any conclusions to be drawn about the most effective exercise regimen. For example, it is possible that the results in the studies by Chaconas et al. [37] and Holmgren et al. [26] came from the greater training intensities rather than the specific type of exercise. It could therefore be argued that these studies should have been excluded from the meta-analyses. We instead chose to include them, partly because the aim of this review was to compare eccentric exercise to any intervention, regardless of what type, but also because high intensity loading can be seen as the hallmark of eccentric training and not a confounding factor. The high heterogeneity among the trials also became apparent when we pooled the outcome data for function. Besides the clinical heterogeneity in exercise protocols, there was also a wide range of outcome measures used to evaluate function, further contributing to the high heterogeneity. It could be questioned whether pooling the data for function was appropriate, but we determined that it was useful both for illustrating the heterogeneity and for assessing whether there was sufficient inconsistency among studies to downgrade the certainty of evidence.

A potential limitation with our review is that only three databases were searched, and no grey literature was searched. However, we chose the most relevant databases and had no reason to believe that any studies not published and indexed in either of the searched databases would exist; hence, we limited our search strategy to those databases. We did not apply a minimal important difference on the effects on function because we used SMD in our calculations to the heterogeneity in outcome measures. However, the difference between groups was not significant at any of the time points. Strengths of the review are the rigorous methodology, that we appraised evidence from randomised controlled trials only, that we performed meta-analyses, and that we assessed the body of evidence using GRADE.

In conclusion, evidence of low certainty suggests that eccentric exercise may provide a small but likely not clinically important reduction in pain post-treatment compared with other types of exercise in patients with subacromial impingement syndrome. At intermediate to long-term follow-up, eccentric exercise may result in little or no important difference in pain compared with other types of exercise. It is uncertain whether eccentric exercise improves function more than other types of exercise post-treatment and at intermediate to long-term follow-up in patients with subacromial impingement syndrome (very low certainty of evidence). Pain during exercise does not seem to provide greater improvement in pain or function compared with pain-free exercise. On the other hand, no negative effects have been observed. Eccentric training regimens have shown both similarities and diversity. Intervention durations of six to 8 weeks have been shown to be similar in effectiveness as an intervention duration of 12 weeks. For frequency and intensity no conclusions can be made, but it seems that exercise at higher intensities might yield better results. When it comes to exercise type, only shoulder external rotation with neutral shoulder and the full can exercise have been studied. It is likely that the optimal exercise will vary depending on underlying tendinopathy. Further research as outlined above is necessary.
## Appendix 1

### Search strategies and search results

**Database:** PubMed  
**Date:** 6 March 2019.  
**No. of results:** 37

| Search Query | Items found |
|--------------|-------------|
| #6 Search ((("shoulder impingement" OR "subacromial impingement" OR "subacromial pain" OR "rotator cuff") OR subacromial impingement syndrome OR rotator cuff) AND ("eccentric exercise" OR "eccentric exercises" OR "eccentric training" OR "eccentric strengthening" OR "eccentric strength training") OR subacromial impingement syndrome OR rotator cuff) | 37 |
| #5 Search "eccentric exercise" OR "eccentric exercises" OR "eccentric training" OR "eccentric strengthening" OR "eccentric strength training" | 1981 |
| #4 Search ((("shoulder impingement" OR "subacromial impingement" OR "subacromial pain" OR "rotator cuff") OR subacromial impingement syndrome OR rotator cuff) | 12594 |
| #3 Search rotator cuff | 5788 |
| #2 Search subacromial impingement syndrome | 1654 |
| #1 Search "shoulder impingement" OR "subacromial impingement" OR "subacromial pain" OR "rotator cuff" | 11637 |

**Database:** The Cochrane Library.  
**Date:** 6 March 2019.  
**No. of results:** 27  
Cochrane Reviews [2].  
Cochrane Protocols (0).  
Trials [25].  
Editorials (0).  
Special Collections (0).  
Clinical Answers (0).  
Other Reviews (0).

| ID | Search | Hits |
|----|--------|------|
| #1 | "shoulder impingement" OR "subacromial impingement" OR "subacromial pain" OR "rotator cuff":[tiab,kw] (Word variations have been searched) | 1425 |
| #2 | MeSH descriptor: [Subacromial impingement syndrome] explode all trees | 280 |
| #3 | MeSH descriptor: [Rotator Cuff] explode all trees | 318 |
| #4 | #1 OR #2 OR #3 | 1425 |
| #5 | 
| #6 | #4 AND #5 | 694 |

**Database:** PEDro.  
**Date:** 6 March 2019.  
**No. of results:** 4.

| Search | Records found |
|--------|---------------|
| Abstract and title: (shoulder impingement" OR "subacromial impingement" OR "subacromial pain" OR "rotator cuff") AND eccentric, | 4 |

### Reference lists

A comprehensive search of reference lists brought 1 new records.
### Table 4 PEDro scores for the included studies

| Study                      | Eligibility criteria and source | Random allocation | Concealed allocation | Groups similar at baseline | Participant blinding | Therapist blinding | Assessor blinding | < 15% dropouts | Intention-to-treat analysis | Between-group difference reported | Point estimates and variability reported | Total score (0 to 10) |
|----------------------------|---------------------------------|-------------------|----------------------|----------------------------|----------------------|---------------------|-------------------|----------------|--------------------------|--------------------------------------|------------------------------------------|------------------|
| Bateman et al. (2014) [38] | Yes                             | Yes               | Yes                  | No                         | No                   | No                  | Yes               | No             | Yes                      | No                                   | No                                      | 5                |
| Blume et al. (2015) [35]   | Yes                             | Yes               | Yes                  | No                         | No                   | Yes                 | Yes               | Yes            | Yes                      | Yes                                   | Yes                                     | 8                |
| Chaconas et al. (2017) [34]| Yes                             | Yes               | No                   | Yes                        | No                   | Yes                 | Yes               | No             | Yes                      | Yes                                   | Yes                                     | 6                |
| Dejaco et al. (2017) [37]  | Yes                             | Yes               | Yes                  | No                         | No                   | No                  | Yes               | Yes            | Yes                      | Yes                                   | Yes                                     | 7                |
| Hallgren et al. (2014) [50]| Yes                             | Yes               | Yes                  | No                         | No                   | Yes                 | Yes               | No             | Yes                      | Yes                                   | Yes                                     | 7                |
| Holmgren et al. (2012) [26]| Yes                             | Yes               | Yes                  | No                         | No                   | Yes                 | Yes               | No             | Yes                      | Yes                                   | Yes                                     | 7                |
| Maenhout et al. (2013) [25]| Yes                             | Yes               | Yes                  | No                         | No                   | No                  | Yes               | Yes            | Yes                      | Yes                                   | Yes                                     | 6                |
| Vallés-Carrascosa et al. (2018) [32]| Yes | Yes | Yes | No | No | No | Yes | Yes | Yes | Yes | Yes | Yes | 7 |
## Appendix 3

### Table 5 Cochrane scale for clinical relevance

| Study                        | Are the patients described in detail so that you can decide whether they are comparable to those that you see in your practice? | Are the interventions and treatment settings described well enough so that you can provide the same for your patients? | Were all clinically relevant outcomes measured and reported? | Is the size of the effect clinically important? | Are the likely treatment benefits worth the potential harms? | Total score (0 to 5) |
|------------------------------|----------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------|-----------------------------------------------|-------------------------------------------------------------|--------------------|
| Bateman et al. (2014) [38]    | Yes                                                                                                                         | Yes                                                                                                          | No                                                          | Yes                                           | Yes                                           | 4                  |
| Blume et al. (2015) [35]     | Yes                                                                                                                         | Yes                                                                                                          | Yes                                                         | Yes                                           | Yes                                           | 5                  |
| Chaconas et al. (2017) [34]  | Yes                                                                                                                         | Yes                                                                                                          | Yes                                                         | Yes                                           | Yes                                           | 5                  |
| Dejaco et al. (2017) [37]    | Yes                                                                                                                         | Yes                                                                                                          | Yes                                                         | Yes                                           | Yes                                           | 5                  |
| Hallgren et al. (2014) [50]  | Yes                                                                                                                         | Yes                                                                                                          | Yes                                                         | Yes                                           | Yes                                           | 5                  |
| Holmgren et al. (2012) [26]  | Yes                                                                                                                         | Yes                                                                                                          | Yes                                                         | Yes                                           | Yes                                           | 5                  |
| Maenhout et al. (2013) [25]  | Yes                                                                                                                         | Yes                                                                                                          | Yes                                                         | Yes                                           | Yes                                           | 5                  |
| Vallés-Carrascosa et al. (2018) [32] | Yes                                                                                                                        | Yes                                                                                                          | Yes                                                         | Yes                                           | Yes                                           | 5                  |
## Table 6: Outcome table for pain

| Study                      | Number of participants | Withdrawals/dropouts | Intervention                                      | Control                                      | Comments                                                                                     |
|----------------------------|------------------------|----------------------|---------------------------------------------------|----------------------------------------------|---------------------------------------------------------------------------------------------|
| Bateman et al. (2014) [38] | I: 4                   | C1: 1                | Eccentric exercise (painful eccentric exercise)   | VAS                                          | VAS                                                                                         |
|                            |                        | C1: 0                | Post-treatment (8 weeks): 43.5 (SD 26.5)          | Baseline: 53.0 (SD 14.2)                      | Baseline: 42.3 (SD 6.9)                                                                    |
|                            |                        | C2: 4                | Δ −9.5                                            | Post-treatment (8 weeks): 51.7 (SD 17.6)      | Δ 9.4                                                                                       |
|                            |                        |                      | Between-group difference in change: Δ 18.9; in favour of eccentric exercise (ns) |                                                                                             |                                                                                             |
| Blume et al. (2015) [35]  | I: 18                  | C: 0                 | NPRS                                              | NPRS                                         |                                                                                             |
|                            |                        | C: 4                 | Baseline: 2.0 (SD 1.8)                            | Baseline: 2.1 (SD 1.6)                        |                                                                                             |
|                            |                        |                      | Post-treatment (8 weeks): 10 (SD 1.3)             | Post-treatment (8 weeks): 1.5 (SD 2.2)        |                                                                                             |
|                            |                        |                      | Δ −1.0                                            | Δ −0.6                                       |                                                                                             |
|                            |                        |                      | Between-group difference in change: Δ 0.4; in favour of eccentric exercise (ns) |                                                                                             |                                                                                             |
| Chaconas et al. (2017) [34] | I: 25                  | C: 0                 | NPRS                                              | NPRS                                         |                                                                                             |
|                            |                        | C: 2                 | Baseline: 1.64 (SD 1.71)                          | Baseline: 1.22 (SD 1.28)                      |                                                                                             |
|                            |                        |                      | Post-treatment (6 weeks): 1.00 (SD 1.47)          | Post-treatment (6 weeks): 1.13 (SD 1.25)      |                                                                                             |
|                            |                        |                      | Δ −0.64                                           | Δ −0.09                                      |                                                                                             |
|                            |                        |                      | At 6 months: 0.54 (SD 1.18)                       | At 6 months: 1.00 (SD 1.30)                   |                                                                                             |
|                            |                        |                      | Δ −1.10                                           | Δ −0.22                                      |                                                                                             |
|                            |                        |                      | Average Pain: 3.72 (SD 2.03)                      | Average Pain: 3.30 (SD 1.61)                  |                                                                                             |
|                            |                        |                      | Post-treatment (6 weeks): 1.40 (SD 1.68)          | Post-treatment (6 weeks): 2.70 (SD 2.03)      |                                                                                             |
|                            |                        |                      | Δ −2.32                                           | Δ −0.60                                      |                                                                                             |
|                            |                        |                      | At 6 months: 1.04 (SD 1.62)                       | At 6 months: 2.14 (SD 1.83)                   |                                                                                             |
|                            |                        |                      | Δ −2.68                                           | Δ −1.16                                      |                                                                                             |
|                            |                        |                      | Worst pain: 3.88 (SD 2.40)                        | Worst pain: 7.00 (SD 2.09)                    |                                                                                             |
|                            |                        |                      | Δ −3.12                                           | Δ −6.39                                      |                                                                                             |
|                            |                        |                      | At 6 months: 3.32 (SD 2.91)                       | At 6 months: 6.39 (SD 2.59)                   |                                                                                             |
|                            |                        |                      | Δ −3.68                                           | Δ −0.61                                      |                                                                                             |
|                            |                        |                      | Between-group difference in change at 6 weeks (best pain): Δ 0.55; in favour of eccentric exercise (ns) |                                                                                             |                                                                                             |
| Study                          | Number of participants | Withdrawals/dropouts | Intervention                                    | Control                                      | Comments                                                                 |
|-------------------------------|------------------------|----------------------|------------------------------------------------|----------------------------------------------|---------------------------------------------------------------------------|
| Dejaco et al. (2017) [37]     | I: 20                  | C: 16                | Eccentric exercise (painful eccentric exercise) | Other exercise types (non-painful eccentric exercise) | VAS Baseline: 390 (SD 185) Post-treatment (12 weeks): 94 (SD 135) Δ − 29.6 At 6 months: 19.1 (SD 24.5) Δ − 19.9 Between-group difference in change at 12 weeks: Δ 6.5; in favour of eccentric exercise (ns) Between-group difference in change at 6 months: Δ 4.4; in favour of eccentric exercise (ns) |
| Hallgren et al. (2014) [50]   | I: 51                  | C: 46                |                                                | VAS Baseline: 42.0 (SD 27.0) Post-treatment (12 weeks): 18.9 (SD 15.8) Δ − 23.1 At 6 months: 19.8 (SD 18.5) Δ − 22.3 | VAS Pain at rest: Baseline: 15 (SD 19) At 1 year: 2 (SD 6) Δ = 13 Pain during activity: Baseline: 61 (SD 22) At 1 year: 18 (SD 23) Δ = −43 Long-term follow-up of the study by Holmgren et al. 2012. NB: 12 patients in the intervention group and 29 in the control group underwent surgery but were still included in 1-year follow-up. |
| Study                          | Number of participants | Withdrawals/dropouts | Intervention                          | Control                          | Comments                                                                 |
|-------------------------------|------------------------|----------------------|---------------------------------------|----------------------------------|--------------------------------------------------------------------------|
|                               |                        |                      | Pain at night:                         | Pain at night:                   |                                                                           |
|                               |                        |                      | Baseline: 46 (SD 28)                   | Baseline: 40 (SD 30)              |                                                                           |
|                               |                        |                      | At 1 year: 12 (SD 19)                 | At 1 year: 14 (SD 21)            |                                                                           |
|                               |                        |                      | △ − 34                                | △ − 26                           |                                                                           |
|                               |                        |                      |                                       |                                  |                                                                          |
|                               |                        |                      | Between-group difference in change (pain at rest): | Between-group difference in change (pain at night): |
|                               |                        |                      | △ 3; in favour of control (ns)         | △ 8; in favour of eccentric exercise (ns) |
|                               |                        |                      |                                       |                                  |                                                                          |
| Holmgren et al. (2012) [26]   | I: 51                  | I: 0                 | VAS                                    | VAS                              | * Holmgren reports an erroneous value for within group change for VAS at rest for the control group (−5). |
|                               | C: 46                  | C: 0                 | Pain at rest:                          |                                  |                                                                          |
|                               |                        |                      | Baseline: 15 (SD 19)                  |                                   |                                                                          |
|                               |                        |                      | Post-treatment (12 weeks):             |                                  |                                                                          |
|                               |                        |                      | 10 (SD 14)                            |                                   |                                                                          |
|                               |                        |                      | △ − 5                                 |                                   |                                                                          |
|                               |                        |                      | Pain during activity:                 |                                  |                                                                          |
|                               |                        |                      | Baseline: 61 (SD 22)                  |                                   |                                                                          |
|                               |                        |                      | Post-treatment (12 weeks):             |                                  |                                                                          |
|                               |                        |                      | 25 (SD 26)                            |                                   |                                                                          |
|                               |                        |                      | △ − 36                                |                                   |                                                                          |
|                               |                        |                      | Pain at night:                         |                                  |                                                                          |
|                               |                        |                      | Baseline: 46 (SD 28)                  |                                   |                                                                          |
|                               |                        |                      | Post-treatment (12 weeks):             |                                  |                                                                          |
|                               |                        |                      | 15 (SD 22)                            |                                   |                                                                          |
|                               |                        |                      | △ − 31                                |                                   |                                                                          |
|                               |                        |                      |                                       |                                  |                                                                          |
|                               |                        |                      | Between-group difference in change (pain at rest): | Between-group difference in change (pain at night): |
|                               |                        |                      | △ 5; in favour of eccentric exercise (ns) | △ 8; in favour of eccentric exercise (ns) |
|                               |                        |                      |                                       |                                  |                                                                          |
| Maenhout et al. (2013) [25]   | I: 31                  | I: 4                 | SPADI Pain subscale                    | SPADI Pain subscale              | Pain was measured using the SPADI. Data were provided via e-mail so that we could extract data for the pain subscale. Scale ranges from 0 to 50, higher score indicates worse pain. |
|                               | C: 30                  | C: 5                 | Baseline: 25.8 (SD 10.4)               | Baseline: 28.4 (SD 12.4)          |                                                                          |
|                               |                        |                      | Post-treatment (12 weeks):             |                                   |                                                                          |
|                               |                        |                      | 11.0 (SD 9.5)                         |                                   |                                                                          |
|                               |                        |                      | △ − 18; in favour of eccentric exercise (95% CI − 30.9 to − 7.2) | |                                                                          |
| Study | Number of participants | Withdrawals/dropouts | Intervention | Control | Comments |
|-------|------------------------|----------------------|--------------|---------|----------|
|       |                        |                      | Eccentric exercise (painful eccentric exercise) | Other exercise types (non-painful eccentric exercise) |          |
| Δ - 14.8 | Δ - 17.4 | Between-group difference in change: Δ 2.6; in favour of control (ns) | | | |
| Vallés-Carrascosa et al. (2018) [32] | I: 11 C: 11 | I: 0 C: 0 | VAS Baseline: 37.0 (32.0; 79.0) Post-treatment (4 weeks): 12.0 (3.0; 30.0) Δ - 25.0 | VAS Baseline: 55.0 (48.0; 68.0) Post-treatment (4 weeks): 28.0 (18.0; 370) Δ - 27.0 | Median (1st; 3rd quartile) |
|       |           |                      | Between-group difference in change: Δ 2.0; in favour of non-painful eccentric exercise (ns) | | | |

*RCT* randomized controlled trial, VAS Visual Analogue Scale, NPRS Numerical Pain Rating Scale, SPADI Shoulder Pain and Disability Index, ns not significant, CI confidence interval
### Appendix 5

#### Table 7 Outcome table for function

| Study                  | Number of participants | Withdrawals/dropouts | Intervention Eccentric exercise (painful eccentric exercise) | Control Other exercise types (non-painful eccentric exercise) | Comments |
|------------------------|------------------------|----------------------|--------------------------------------------------------------|---------------------------------------------------------------|----------|
| Bateman et al. (2014)  | I: 4                   | C1: 1                | Oxford Shoulder Score Baseline: 340 (SD 5.3) Post-treatment (8 weeks): 313 (SD 63) Δ = 27 | Oxford Shoulder Score Baseline: 240 (SD 3.7) Post-treatment (8 weeks): 24.3 (SD 42) Δ = 0.3 | Control group 2 (C2) excluded because no treatment was given. Function measured with the 12-item Oxford Shoulder Score. Range 12–60, where 12 indicates no pain and normal function and higher is worse. MID 6 points [43]. |
|                        | I: 1                   | C1: 1                |                                                              |                                                              |          |
|                        | C2: 0                  |                      |                                                              |                                                              |          |
|                        |                        |                      | Oxford Shoulder Score Baseline: 340 (SD 5.3) Post-treatment (8 weeks): 313 (SD 63) Δ = 27 | Oxford Shoulder Score Baseline: 240 (SD 3.7) Post-treatment (8 weeks): 24.3 (SD 42) Δ = 0.3 | Control group 2 (C2) excluded because no treatment was given. Function measured with the 12-item Oxford Shoulder Score. Range 12–60, where 12 indicates no pain and normal function and higher is worse. MID 6 points [43]. |
|                        |                        |                      |                                                              |                                                              |          |
| Blume et al. (2015)    | I: 18                  | C: 20                | DASH Baseline: 250 (SD 106) Post-treatment (8 weeks): 121 (SD 11.7) Δ = 129 | DASH Baseline: 21.2 (SD 6.5) Post-treatment (8 weeks): 9.3 (SD 7.1) Δ = 11.9 | Function measured with the 30-item DASH questionnaire. Range 0–150, higher is worse. MID = 10.5 points [40]. |
|                        | I: 0                   | C: 4                 |                                                              |                                                              |          |
|                        |                        |                      |                                                              |                                                              |          |
| Chaconas et al. (2017) | I: 25                  | C: 21                | WORC Baseline: 666 (SD 160) Post-treatment (6 weeks): 876 (SD 145) Δ = 21.0 | WORC Baseline: 65.4 (SD 14.1) Post-treatment (6 weeks): 70.6 (SD 16.8) Δ = 5.2 | Function measured with WORC. Total range 0–2100 (aggregate score of 6 total sub-domains), here reported as scores out of 100 i.e. a percentage of normal score, where higher is better. MID = 275 points [39]. |
|                        | I: 0                   | C: 2                 | At 6 months: 92.7 (SD 9.0) Δ = 26.1 | At 6 months: 77.2 (SD 15.9) Δ = 11.8 |          |
|                        |                        |                      |                                                              |                                                              |          |
| Dejaco et al. (2017)   | I: 20                  | C: 16                | Constant-Murley Score Baseline: 72.5 (SD 17.7) Post-treatment (12 weeks): 87.3 (SD 16.2) Δ = 148 | Constant-Murley Score Baseline: 78.9 (SD 8.5) Post-treatment (12 weeks): 87.6 (SD 78) Δ = 8.7 | Function measured with Constant-Murley Score. Four subscales (pain, ADL, ROM, strength). Total range 0–100, where 100 indicates excellent shoulder function. MID = 17 points [41]. |
|                        | I: 1                   | C: 1                 | At 6 months: 86.9 (SD 16.8) Δ = 144 | At 6 months: 888 (SD 8.1) Δ = 9.9 |          |
|                        |                        |                      |                                                              |                                                              |          |
Table 7 Outcome table for function (Continued)

| Study                      | Number of participants | Withdrawals/dropouts | Intervention (painful eccentric exercise) | Control (Other exercise types (non-painful eccentric exercise)) | Comments                                                                                     |
|----------------------------|------------------------|----------------------|-------------------------------------------|-----------------------------------------------------------------|--------------------------------------------------------------------------------------------|
| Hallgren et al. (2014) [50] | I: 51                  | I: 1                 | Constant-Murley Score                     | Constant-Murley Score                                          | Long-term follow-up of the study by Holmgren et al. 2012. Baseline values from the Holmgren et al. paper. |
|                            | C: 46                  | C: 1                 | Baseline: 48.5 (SD 15.0)                   | Baseline: 43.5 (SD 15.0)                                        |                                                                                             |
|                            |                        |                      | 1-year follow up: 83.0 (SD 140)            | 1-year follow up: 760 (SD 180)                                  |                                                                                             |
|                            |                        |                      | \( \Delta = 34.5 \)                       | \( \Delta = 32.5 \)                                            |                                                                                             |
| Holmgren et al. (2012) [26] | I: 51                  | I: 0                 | Constant-Murley Score                     | Constant-Murley Score                                          |                                                                                             |
|                            | C: 46                  | C: 0                 | Baseline: 48.5 (SD 15.0)                   | Baseline: 43.5 (SD 15.0)                                        |                                                                                             |
|                            |                        |                      | 12 weeks: 72.5 (SD 19.0)                  | 12 weeks: 52.5 (SD 230)                                        |                                                                                             |
|                            |                        |                      | \( \Delta = 24 \)                         | \( \Delta = 9 \)                                               |                                                                                             |
| Maenhout et al. (2013) [25] | I: 31                  | I: 4                 | SPADI                                      | SPADI                                                          | Measured with SPADI. A 13-item scale (5 items on pain, 8 on disability). Range 0–130, higher is worse. Pain and disability subscales not reported separately. MID = 18 points [40]. |
|                            | C: 30                  | C: 5                 | Baseline: 420 (SD 11.0)                   | Baseline: 44.3 (SD 11.5)                                       |                                                                                             |
|                            |                        |                      | Post-treatment (12 weeks): 17.0 (SD 11.4) | Post-treatment (12 weeks): 14.5 (SD 11.7)                      |                                                                                             |
|                            |                        |                      | \( \Delta = 25.7 \)                       | \( \Delta = 27.0 \)                                            |                                                                                             |
| Vallés-Carrascosa et al. (2018) [32] | I: 11               | I: 0                 | Constant-Murley Score                     | Constant-Murley Score                                          | Median (1st;3rd quartile)                                                                     |
|                            | C: 11                  | C: 0                 | Baseline: 35.0 (22.0; 47.0)               | Baseline: 36.0 (22.0; 45.0)                                    |                                                                                             |
|                            |                        |                      | Post-treatment (4 weeks): 59.0 (50.0; 68.0) | Post-treatment (4 weeks): 65.0 (55.0; 690)                    |                                                                                             |
|                            |                        |                      | \( \Delta = 24.0 \)                       | \( \Delta = 29.0 \)                                            |                                                                                             |

DASH Disabilities of the Arm, Shoulder and Hand questionnaire, WORC Western Ontario Rotator Cuff index, SPADI Shoulder Pain and Disability Index, MID minimal important differences, ns not significant.
Abbreviations
CI: Confidence interval; CM score: Constant-Murley score; DASH: Disabilities of the arm, shoulder and hand; IV: Inverse-variance; MD: Mean difference; MID: Minimal important difference; MRI: Magnetic resonance imaging; NPRS: Numerical pain rating scale; OSS: Oxford Shoulder Score; RCTs: Randomised controlled trials; RMR: Repetition maximum; SMD: Standardised mean difference; SPADI: Shoulder Pain and Disability Index; VAS: Visual analogue scale; WORC: Western Ontario Rotator Cuff index

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Authors’ contributions
RL and SB identified and screened titles and abstracts for relevance, performed data extraction and assessed risk of bias of included studies. RL was a major contributor in writing the manuscript. SB was responsible for analysis of data. LN assessed risk of bias of included studies. All authors read and approved the final manuscript.

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The datasets used and/or analysed during the current study are available from the original source articles or from the corresponding author on request.

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The authors declare that they have no competing interests.

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