Sham surgery versus labral repair or biceps tenodesis for type II SLAP lesions of the shoulder: a three-armed randomised clinical trial

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ABSTRACT

Background Labral repair and biceps tenodesis are routine operations for superior labrum anterior posterior (SLAP) lesion of the shoulder, but evidence of their efficacy is lacking. We evaluated the effect of labral repair, biceps tenodesis and sham surgery on SLAP lesions.

Methods A double-blind, sham-controlled trial was conducted with 118 surgical candidates (mean age 40 years), with patient history, clinical symptoms and MRI arthrography indicating an isolated type II SLAP lesion. Patients were randomly assigned to either labral repair (n=40), biceps tenodesis (n=39) or sham surgery (n=39) if arthroscopy revealed an isolated SLAP II lesion. Primary outcomes at 6 and 24 months were clinical Rowe score ranging from 0 to 100 (best possible) and Western Ontario Shoulder Instability Index (WOSI) ranging from 0 (best possible) to 2100. Secondary outcomes were Oxford Instability Shoulder Score, change in main symptoms, EuroQol (EQ-5D and EQ-VAS), patient satisfaction and complications.

Results There were no significant between-group differences at any follow-up in any outcome. Between-group differences in Rowe scores at 2 years were: biceps tenodesis versus labral repair: 1.0 (95% CI −5.4 to 7.4), p=0.76; biceps tenodesis versus sham surgery: 1.6 (95% CI −5.0 to 8.1), p=0.64; and labral repair versus sham surgery: 0.6 (95% CI −5.9 to 7.0), p=0.86. Similar results—no differences between groups—were found for WOSI scores. Postoperative stiffness occurred in five patients after labral repair and in four patients after tenodesis.

Conclusion Neither labral repair nor biceps tenodesis had any significant clinical benefit over sham surgery for patients with SLAP II lesions in the population studied.

Trial registration number ClinicalTrials.gov identifier: NCT00586742

INTRODUCTION

The superior glenoid labrum of the shoulder is a common site of injury and degeneration, which can cause both pain and disability. Snyder et al1 used the term SLAP (superior labrum anterior posterior) to describe a combined detachment of the long head of biceps tendon and the superior labrum from the glenoid rim (figure 1). The diagnosis is controversial; the validity of clinical tests,2 and even arthroscopy and MRI arthrography is questioned.3-6 In addition, a high prevalence of SLAP lesions has been reported in a middle-aged asymptomatic population.7

Labral repair is the most common procedure to treat labral tears, but because of high rates of complications and poor outcomes,10-17 it has been suggested that indications should be narrowed.10-17 Recent reports show a decrease in the number of labral tears performed and a decrease in the age of the patients undergoing these operations.18 19 Release of the biceps tendon (tenodesis or tenotomy) is increasingly used as an alternative13 18-22 to SLAP repairs in select patients, but the evidence for it is weak.13 20-22 One non-controlled study reported better results after tenodesis compared with labral repair,13 but no randomised trials have compared these procedures. Very little evidence about non-operative treatment for type II SLAP lesions was available when the present study was designed. Recent non-controlled studies suggest that non-operative treatment including graded exercise therapy is beneficial, and the postoperative rehabilitation in the present study followed those rehabilitation principles.21-25

Our aim was to conduct a randomised, double-blind, sham-controlled trial to assess the clinical effectiveness of labral repair and biceps tenodesis in patients with a type II SLAP lesion.

MATERIALS AND METHODS

Design
This blinded, three-armed randomised, sham-controlled study with a 2-year follow-up was conducted at Lovisenberg Diaconal Hospital, Oslo, Norway, from January 2008 to January 2014. The study protocol has been published previously.26 The patients, the treating physiotherapists/manual therapists and the persons collecting and analysing the data were blinded to the study group assignments. The first author was not blinded to group assignment, but had no role in the follow-up of the patients.

Participants
We enrolled patients 18–60 years of age who had shoulder pain (>3 months) that was unresponsive to conventional non-operative treatment (physiotherapy, non-steroidal anti-inflammatory drugs and/or corticosteroid injections) and had a history, clinical findings and MRI arthrography indicating a SLAP lesion. Patients were informed that final study inclusion and randomisation would be performed during arthroscopy if an isolated type II SLAP lesion was confirmed. Exclusion criteria included previous shoulder surgery, SLAP lesions with concomitant...
labral cysts, clinical and radiological signs of arthritis of the acromioclavicular (AC) or glenohumeral joint, and tears of the rotator cuff or the long head of the biceps tendon. Detailed criteria are described in the published study protocol.

**Diagnostic arthroscopy**

Arthroscopic examination of the shoulder was performed in all patients under general anaesthesia, with the use of standard posterior and anterior portals. The orthopaedic surgeon (CPS) evaluated both the subacromial space and the glenohumeral joint and noted the intra-articular findings (figure 1), and a video was created for each patient.

**Randomisation**

Participants who fulfilled the inclusion criteria and consented to take part in the trial were randomised to receive either (1) labral repair, (2) biceps tenodesis or (3) sham surgery. An independent statistician used the permuted block method of randomisation. Concealed allocation was organised by an independent secretary who distributed sealed opaque-numbered envelopes to the head nurse in the operating room. The nurse opened the envelope when a patient was confirmed to be eligible by arthroscopic diagnostic evaluation. Following confirmation of an isolated type II SLAP lesion, the patient was included in the randomisation procedure.

**Interventions**

A detailed description of each intervention is provided in the published protocol. A single experienced shoulder surgeon performed all arthroscopic evaluations and treatments.

**Operative and postoperative procedures**

During the labral repair, the superior glenoid rim was debrided with a motorised shaver, followed by a percutaneous placement of a drill guide and anchors through the myotendinous

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**Figure 1** CONSORT (Consolidated Standards of Reporting Trials) flow chart. AC, acromioclavicular; SLAP, superior labrum anterior posterior.
junction of the supraspinatus. All anchors were placed posterior to the biceps root, and single circular sutures were used. The biceps tenodesis was done with a mini-open technique. Under arthroscopic vision, a spinal needle was placed as laterally as possible and with a 90° angle central in the biceps tendon and a tenotomy was performed at the biceps insertion. With less than 2 cm skin incision (to mimic the scars in the other two groups), the spinal needle was followed down to the biceps pulley, the pulley was split and the biceps tendon was lifted out. The groove was debrided, a double-loaded metal anchor was placed in the inferior part of the groove, and the tendon secured with two sutures, each passing twice through the tendon. The proximal part of the tendon was excised and the wound closed.

For the sham surgery, standard diagnostic shoulder arthroscopy was performed. In addition, a 5 mm skin incision was made to mimic a labral repair. The patient was kept in the operation theatre for the amount of time required to perform an actual arthroscopic index shoulder surgery.

All operative procedures were recorded on video. For all patients, a sling was placed before they left the operating room, and standard postoperative care and instructions were provided.

Postoperative rehabilitation

Patients in all three groups had a standardised, but individually adjusted rehabilitation. Elbow, wrist, and finger mobilisation and gentle pendulum exercises were conducted, starting on the first day. A sling was used for 3 weeks. Exercises to improve the scapula-humeral rhythm, coordination and mobility were performed using sling exercise therapy.20 Gradual biceps loading was started at 12 weeks. Exercises to improve functional stability and scapula muscles were progressively emphasised after 6 weeks. Sports and job-specific rehabilitation were provided on an individual basis, usually starting 3 months postoperatively. Rehabilitation was continued for 3–6 months and included 12–16 sessions with a therapist and 20 self-administered exercises.

Outcome measures

The two primary outcome measures were the Rowe score30 and the Western Ontario Shoulder Instability Index (WOSI)31 at 6 and 24 months after surgery. The 1988 version of the Rowe score and the WOSI were validated for use in the present study.32–34 As recommended by Rowe, patient satisfaction was assessed separately using a self-reported question with response alternatives of poor, fair, good and excellent.35

Secondary outcomes were: WOSI and Rowe at 3 and 12 months, the Oxford Instability Shoulder Score (OISS),36 the EuroQol (EQ-5D, EQ-VAS) for generic health-related quality of life37 and change of the main symptom (pain) from baseline to all follow-ups.38 The EuroQol and OISS are validated for use in patients with SLAP lesions.33–34

A single-blinded observer assessed all participants and administered questionnaires at baseline and all postoperative follow-ups. Patient characteristics were assessed at baseline and included self-reported manual labour (yes/no); physical activity (1=competitive sports/2= frequent physical activity (three to four times a week)/3=occasional or none); dominant shoulder (yes/no); apprehension test (positive/negative); Crank test (positive/negative); and O’Brien test (positive/negative). Blinding was evaluated by a question asking the patients if they thought they were operated or had sham surgery. Adverse events were classified as serious if the patient needed prolonged inpatient hospital care, hospitalisation or death, and considerable if the patient had painful shoulder stiffness (capsulitis) with external rotation and forward flexion <30° and abduction <90°.39

Sample size and statistical analysis

The study was designed to detect a clinically important detectable difference of 10 points in mean Rowe score. To detect this difference among treatment groups (SD=15, α=0.05, β=0.80, one-way analysis of variance), the study required 36 patients in each group. Assuming 10% dropout, we planned to include 40 patients in each group. In a later study,36 we found the minimal clinically important change for Rowe score to be 17. At 6 months there were no crossovers and the results were analysed and interpreted blindly, as recommended by Järvinen et al.40

Continuous variables are presented as mean values with 95% CI and categorical values as numbers and percentages. We used a mixed models approach to analyse the effect of the interventions on primary and secondary variables at 6 and 24 months. For each model, we included a core set of adjustment variables as covariates: age, sex, time, frequency of physical activity, manual labour and dominant arm. In addition, the baseline measurement was included as a covariate to eliminate regression towards the mean for the estimated changes over time of the different treatments and for pairwise comparisons. The final model was tested for confounding with the covariates not included in the final main effects model. Confounding

## Table 1 Baseline characteristics of the patients according to study group

| Characteristic | Biceps tenodesis | Sham surgery | Labral repair |
|---------------|-----------------|--------------|--------------|
| Age (years), median (range) | 40 (18 to 64) | 40 (23 to 60) | 42 (22 to 57) |
| Female sex, n (%) | 15 (38.5) | 17 (42.5) | 15 (37.5) |
| Body mass index | 27.6 (5.6) | 26.1 (3.8) | 26.4 (4.3) |
| University education, n (%) | 18 (48.6) | 23 (59.0) | 22 (56.4) |
| Manual labour, n (%) | 19 (48.7) | 11 (28.2) | 17 (42.5) |
| Physical activity, n (%) | 18 (47.2) | 21 (56.4) | 28 (70.0) |
| No training | 22 (55.0) | 17 (42.5) | 12 (30.0) |
| Frequent physical activity | 17 (42.5) | 17 (42.5) | 27 (67.5) |
| Competitive | 1 (2.5) | 5 (12.5) | 1 (2.5) |
| Taking analgesics daily or weekly, n (%) | 7 (17.9) | 5 (12.5) | 6 (15.0) |
| Duration of pain, month median (IQR) | 24 (59) | 26 (61) | 24 (47) |
| Dominant shoulder involved, n (%) | 28 (71.8) | 33 (84.6) | 28 (70.0) |
| Positive O’Brien test, n (%) | 37 (94.9) | 33 (86.8) | 36 (90.0) |

*Plus-minus values are means±SD.
†The body mass index is the weight in kilograms divided by the square of the height in metres.
‡Manual labour was reported if the patient was engaged in daily manual work.
§Physical activity was reported if the patient had no regular training, frequent if the patient was training two times or more in a week on a regular basis, and competitive if the patient was competing on a national level.
¶The O’Brien test is positive if the patient has pain when resisting force from an examiner, with the arm in 90° of flexion, 20°adduction and supinated, and less pain with the arm otherwise in the same position but with pronation.
was defined as a change in the estimate of at least 25% when a term was added to a model. Finally, we tested for interactions between covariates and all outcomes at the 0.05 level only. If a significant difference was found, we applied Tukey’s test for multiple comparisons. The assumptions were explored using Cook’s distance, covariance ratio and trace statistics to assess the validity of the model. To ascertain the robustness of findings, we performed a bootstrap regression analysis with 1000 replications, adjusting for the baseline values and the variables mentioned above. Likewise, we performed multiple imputation using the Markov chain Monte Carlo method. The analysis was performed with the missing data as is and with the imputed values. For categorical variables, we applied the mixed models with a multinomial distribution. For sensitivity analysis, we performed per protocol analysis comparing results in those who adhered to the protocol. Post hoc analyses were used to compare primary outcome scores at 6 months in those who crossed over from sham surgery with scores in a similar number of patients with the most inferior scores of patients randomised to biceps tenodesis and labral repair. The analyses were performed in Statistical Analysis System (SAS, V.9.4) and R V.3.1.1.

RESULTS

Patient characteristics

Of the 445 patients eligible for inclusion, 262 were excluded before arthroscopy. Of the 183 patients who had arthroscopy, 65 were excluded peroperatively (figure 1). A total of 118 patients were randomised; 39 were assigned to sham surgery, 39 to biceps tenodesis and 40 to labral repair. The mean age was 40 years, ranging from 18 to 64. One patient aged 64 years was mistakenly included. The baseline characteristics are described in table 1.

Four patients were lost to follow-up at 2 years; two in the sham group and one in each of the other two groups.

Primary outcomes

There was a significant improvement from baseline to 6 and 24 months for all three study groups (figure 2 and table 2). Age, sex, trauma, manual work, frequency of physical activity, duration of pain and dominant shoulder were not significantly associated with primary outcomes at follow-up. The only
significant covariates were the baseline Rowe (p<0.001) and WOSI (p<0.0002) scores.

There were no significant between-group differences in the change from baseline to 6 or 24 months in any primary outcome, neither for the adjusted nor for the unadjusted results (table 3). Unadjusted results were marginally different and imputation did not materially change between-group differences.

### Secondary and other outcomes

All three groups improved on the secondary outcomes from baseline to 6 and 24 months (table 2), and there were no significant differences among the groups (table 3). In the sham group, 31/37 (84%) patients reported excellent or good results at 1-year follow-up, in the labral repair and biceps tenodesis groups the numbers were 29/35 (83%) and 34/38 (89%), respectively. The first 25 patients did not receive the blinding question. After sham surgery, 19/26 (73%) patients believed they were repaired, in comparison to 31/32 (97%) after biceps tenodesis and 31/35 (89%) after labral repair.

There were no serious adverse events, infections or nerve injuries. Ten patients experienced prolonged postoperative stiffness (capsulitis); five after labral repair, four after biceps tenodesis and 31/35 (89%) after labral repair.

### Per protocol analysis

There were no significant differences in the changes between groups for any primary or secondary outcomes in the per protocol analysis. At 6 and 24 months, respectively, the mean

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**Table 2** Primary and secondary outcomes at baseline and at 3, 6, 12 and 24 months after arthroscopy *

| Outcome                                      | Biceps tenodesis (n=39) | Sham surgery (n=39) | Labral repair (n=40) |
|----------------------------------------------|--------------------------|---------------------|----------------------|
| Primary outcome, mean (95% CI)              |                          |                     |                      |
| Rowe score (0 to 100)†                       |                          |                     |                      |
| Baseline                                     | 60.3 (57.0 to 64.0)      | 63.2 (59.8 to 66.7) | 62.7 (59.6 to 65.8)  |
| 3 Months                                     | 62.4 (57.1 to 67.7)      | 68.9 (63.5 to 74.3) | 63.8 (58.6 to 69.0)  |
| 6 Months                                     | 76.0 (70.7 to 81.1)      | 76.3 (71.1 to 81.6) | 76.1 (71.0 to 81.3)  |
| 12 Months                                    | 83.3 (78.4 to 88.1)      | 81.4 (76.4 to 86.5) | 83.5 (78.6 to 88.4)  |
| 24 Months                                    | 86.8 (82.2 to 91.4)      | 85.3 (80.6 to 89.9) | 85.8 (81.3 to 90.4)  |
| WOSI (0 to 2100)‡                           |                          |                     |                      |
| Baseline                                     | 1155 (1026 to 1283)      | 1062 (940 to 1183)  | 1044 (941 to 1146)   |
| 3 Months                                     | 1115 (829 to 1401)       | 833 (528 to 1138)   | 788 (506 to 1070)    |
| 6 Months                                     | 689 (548 to 829)         | 560 (420 to 701)    | 552 (414 to 691)     |
| 12 Months                                    | 490 (353 to 627)         | 475 (335 to 614)    | 429 (294 to 565)     |
| 24 Months                                    | 436 (313 to 559)         | 458 (334 to 582)    | 340 (219 to 461)     |
| Secondary outcome, mean (95% CI)            |                          |                     |                      |
| OISS (12 to 60)§                            |                          |                     |                      |
| Baseline                                     | 38.6 (36.1 to 41.1)      | 36.9 (34.3 to 39.6) | 36.7 (34.8 to 38.6)  |
| 3 Months                                     | 37.8 (35.3 to 40.4)      | 36.0 (33.4 to 38.6) | 36.0 (33.6 to 38.5)  |
| 6 Months                                     | 29.7 (27.0 to 32.4)      | 28.6 (25.9 to 31.3) | 27.4 (24.7 to 30.6)  |
| 12 Months                                    | 24.1 (21.1 to 27.1)      | 25.9 (22.8 to 28.9) | 23.3 (20.3 to 26.2)  |
| 24 Months                                    | 21.7 (18.5 to 24.8)      | 23.3 (20.1 to 26.5) | 22.7 (19.6 to 25.8)  |
| EQ-SD (0.59 to 1.00)¶                         |                          |                     |                      |
| Baseline                                     | 0.6 (0.5 to 0.7)         | 0.7 (0.6 to 0.7)    | 0.7 (0.6 to 0.7)     |
| 3 Months                                     | 0.6 (0.6 to 0.7)         | 0.7 (0.7 to 0.8)    | 0.6 (0.7 to 0.8)     |
| 6 Months                                     | 0.7 (0.7 to 0.8)         | 0.8 (0.8 to 0.9)    | 0.8 (0.7 to 0.9)     |
| 12 Months                                    | 0.8 (0.8 to 0.9)         | 0.8 (0.7 to 0.9)    | 0.8 (0.8 to 0.9)     |
| 24 Months                                    | 0.8 (0.8 to 0.9)         | 0.9 (0.8 to 0.9)    | 0.9 (0.8 to 0.9)     |
| EQ-VAS (0 to 100)¶                           |                          |                     |                      |
| Baseline                                     | 67.9 (63.3 to 73.4)      | 66.7 (62.2 to 73.1) | 68.4 (62.9 to 73.8)  |
| 3 Months                                     | 67.1 (60.7 to 73.5)      | 67.7 (61.6 to 74.3) | 72.9 (66.6 to 79.3)  |
| 6 Months                                     | 73.9 (68.3 to 79.4)      | 76.9 (71.3 to 82.4) | 81.3 (75.8 to 86.7)  |
| 12 Months                                    | 78.3 (72.5 to 84.1)      | 79.4 (73.5 to 85.3) | 79.2 (73.5 to 85.0)  |
| 24 Months                                    | 79.6 (74.0 to 85.2)      | 76.8 (71.2 to 82.5) | 81.7 (76.1 to 87.2)  |
| Change of main complaint (−9 to +9)          |                          |                     |                      |
| 3 Months                                     | 3.0 (1.5 to 4.5)         | 2.3 (0.8 to 3.9)    | 3.2 (1.7 to 4.6)     |
| 6 Months                                     | 3.8 (2.2 to 5.4)         | 2.8 (1.2 to 4.4)    | 3.7 (2.1 to 5.3)     |
| 12 Months                                    | 5.3 (3.9 to 6.6)         | 4.5 (3.1 to 5.9)    | 5.5 (4.2 to 7.0)     |
| 24 Months                                    | 5.7 (4.6 to 6.9)         | 5.5 (4.3 to 6.7)    | 5.3 (4.3 to 6.9)     |

* Unadjusted values are given.
† Higher scores indicating better shoulder function.
‡ Western Ontario Shoulder Instability Index, lower scores indicating better shoulder function.
§ Oxford Instability Shoulder Score, lower scores indicating better shoulder function.
¶ EuroQol (EQ-5D and EQ-VAS), higher scores indicating better health-related quality of life.
OISS, Oxford Instability Shoulder Score; WOSI, Western Ontario Shoulder Instability Index.

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The values are expressed as mean (95% CI).
* Adjusted for age, sex, baseline score, time, manual labour, physical activity and dominant shoulder. Between-group differences were not significant for any outcome. Time and baseline score significantly predicted outcome.
† Higher Rowe scores indicating better shoulder function.
‡ Western Ontario Shoulder Instability Index, lower score indicating better shoulder function.
§ Oxford Instability Shoulder Score, lower score indicating better shoulder function.
¶ EuroQol (EQ-5D and EQ-VAS), higher scores indicating better health-related quality of life.
§ Oxford Instability Shoulder Score; WOSI, Western Ontario Shoulder Instability Index.

** DISCUSSION**

In this three-armed randomised, double-blind, sham-controlled trial comparing labral repair, biceps tenodesis and sham surgery for symptomatic, isolated SLAP II lesions of the shoulder, we found a significant improvement in objective and subjective shoulder scores for all three groups, but no significant group differences at either the 6 or 24 months of follow-up. As there is no previous sham-controlled trial comparing labral repair, biceps tenodesis and sham surgery, we discuss the results of the present sham surgical trial in the context of sham surgical trials in knee patients with knee pain.74,42

**Clinically relevant outcome measures**

The choice of outcome measures is an important issue. The scores used in this study were validated for use on patients in the crossover group had a Rowe score over 80 and 7 of them rated the shoulder as good or excellent.
with a SLAP lesion and capture the most important aspects of shoulder function. Both the clinician-based and the patient-based scores showed comparable results and differences between groups were much smaller than the different scores are able to detect.

Potential limitations
Issues related to sample size, blinding, the population studied and external validity are possible limitations of the present study. This is a small study, but the CIs of the observed between-group differences indicate that the study was adequately powered to detect clinically relevant differences between the three groups. For Rowe scores, the CIs did not include 10, which the trial was designed to detect. That only 73% of patients in the sham group believed that they had been repaired may have biased the results. Unfortunately, data did not provide enough information to calculate a blinded index. The primary follow-up time of 6 months was selected mainly for ethical reasons. Given that the patients were referred for surgery and that we hypothesised better results after surgical repair compared with sham surgery, it was our opinion that it would be unethical to maintain patient blinding much beyond 6 months. Therefore, we advised patients that it would take time to recover from surgery and informed them that it was our aim to maintain the blinding as long as possible. However, if they were not satisfied by 6 months, their group assignment could be unblinded.

Our results at 2 years confirm the 6-month results with no statistically significant differences between the three groups. In addition, the comparison of crossovers with a similar number of patients with inferior results in the biceps tenodesis and labral repair groups suggests that the threshold for reoperation was lower in the sham group. However, it cannot be ruled out that the higher rate of reoperation in the sham group was related to unblinding rather than to differences in treatment failure. Even though there was no significant difference in mean Rowe score in the crossover group after reoperation, it is worth noting that 8 of the 14 patients had a Rowe score over 80.

We assessed 445 patients and included 118 according to strict inclusion criteria. This introduces a risk of selection bias but improves the internal validity of the study in order to assess isolated SLAP II lesions. Both athletes and patients with work-related injuries were included. Results were adjusted for frequency of physical activity and manual labour, but do not apply to a specific population. Future studies should compare non-operative and operative treatments in athletes, including return to sports in addition to the validated outcomes applied in the present study.

The choice of method regarding the biceps tenodesis may be questioned. Surgeons currently use several methods, both arthroscopic and open, with a variety of fixation devices. The point of fixation of the biceps tendon may be in the bicipital groove, subcoracoid humeral fixation or transferred to the conjoint tendon. We fixed the biceps with a mini-open technique using a suture anchor low in the bicipital groove. Systematic reviews and randomised studies comparing different tenodesis techniques have not shown any differences in outcome.

Strengths of this study
The study has several strengths: randomisation; inclusion of a sham surgery control; blinded assessments, analysis and interpretation of results; validated outcome measures; no crossovers at 6 months; and minimal loss to follow-up. Strict inclusion criteria were applied to increase the internal validity for evaluating the efficacy of the surgical procedures. The use of locally available physiotherapists and manual therapists for the post-operative exercise programme of these patients enhances the external validity of the study.

The physiotherapy provided to all groups (the sole treatment in the sham group) followed current guidelines. However, the study does not provide information about the effectiveness of physiotherapy with supervised exercises without diagnostic arthroscopy. Ideally, the study should have included an arm with patients not receiving any treatment (natural course), or an arm with physiotherapy without placebo surgery. We discussed this and decided that the inclusion of a fourth group would be too difficult to implement. Although physiotherapy might have contributed to the results in all groups, the impact of placebo, the natural course and regression to the mean should not be underestimated.

Clinical implications
The results of our trial extend previous reports of possible overtreatment of SLAP lesions and indicate a need to narrow indications. Patient age is debated and authors advocate that SLAP repair should be reserved for the young and active patient. There were no significant differences in function, patient satisfaction or complications by age in this study, but the groups are too small to perform subgroup analysis and identify factors associated with failures. The present study does not support either labral repair or biceps tenodesis for type II SLAP lesions in this population, as we found no significant difference between treatments in any outcome. Considering the lack of high-quality trials in this field, the results of this study should be interpreted with caution.

Based on this study, we believe future patients should be informed about the long recovery and possible complications after surgery, and that non-operative treatment has a good probability of success. Future studies should evaluate non-operative treatments and predictors of success.

We conclude that at 6 and 24 months of follow-up, there were no significant differences between labral repair, biceps tenodesis and sham surgery for patients with a type II SLAP lesion in the population studied.

What are the findings?
Labral repair, biceps tenodesis and sham surgery for patients with an isolated SLAP II lesion all led to significant improvement in both objective and subjective scores. There was no significant difference among (1) labral repair, (2) biceps tenodesis and (3) sham surgery in the population studied.

How might it impact on clinical practice in the future?
Patients should be told that non-operative treatment has a good probability of success, but further studies are needed to establish what treatment is the best for the young active patient.

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Contributors
CPS, OR and JIB conceived and designed the study. ØS and PM participated in setting up of the study. ØS participated in patient recruitment and data collection. PM conducted the analysis and, together with JIB, ØS, OR and CPS, interpretation. CPS and JIB drafted the first version of the manuscript. All authors helped revise the manuscript and gave their final approval of the submitted version.
Competing interests None declared.

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