Outcomes of Reverse Shoulder Arthroplasty Following Failed Superior Capsular Reconstruction

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Abstract

Background: History of prior rotator cuff repair (RCR) may adversely affect the outcomes of reverse total shoulder arthroplasty (RTSA), but there is no information regarding the influence of prior superior capsular reconstruction (SCR) surgery on the outcomes of RTSA. The purpose of this study is to evaluate the outcomes of RTSA following failed arthroscopic SCR.

Methods: All patients who underwent RTSA for failed SCR (SCR cohort) at our institution were identified from our institutional database. A comparative cohort of patients who had RTSA with a history of failed RCR (Control cohort) was also reviewed. Demographic information, 90-day complication rate, 90-day emergency room (ED) visits, length of stay (LOS), and outcome scores (patient-reported outcomes measurement information system physical function upper extremity [PROMIS PFUE], visual analog score [VAS], and range of motion [ROM]) were compared.

Results: From 2015 to 2020, 87 arthroscopic SCRs were performed at our institution and of these, 13 patients underwent RTSA at a mean time of 14.6 months (5.8–32.4) after SCR and were followed up for an average of 17.9 months (1.6–44.6). The average number of shoulder surgeries prior to RTSA was 2.8 (1–7), with the last surgery being SCR. During the same period, we identified 15 patients who underwent a RTSA after a failed RCR (Control cohort). The RTSA in the control cohort was performed on average at 12.8 months (1.5–39.5) following the last RCR, and patients were followed up for an average of 27.7 months (2.8–53.9). The average number of shoulder surgeries before the RTSA in the control cohort was 1.4 (1–3). Although the SCR cohort had significant improvements in pain scores and forward flexion (FF), there was
only a modest functional improvement with PROMIS scores and no meaningful improvement with external rotation (ER). Complications (23%) in the SCR cohort included one peri-prosthetic joint infection requiring two-stage revision, one acromion stress fracture, and one ulnar neuritis. Overall, compared to the SCR cohort, patients in the control cohort had better function (PROMIS PFUE), lower VAS score, and greater ROM (FF and ER) preoperatively and at last follow-up, but there were no differences in the LOS and 90-day ED visits, infection, and complication rate between the two cohorts.

Conclusions: RTSA after failed SCR improves pain and forward flexion but is associated with modest functional improvements and high complication rates. However, these findings will require confirmation in a larger cohort with longer follow-up.

Level of Evidence: Level III; Retrospective Cohort Comparison; Treatment Study

Keywords: Superior Capsular Reconstruction (SCR); Irreparable Rotator Cuff Tear; Rotator Cuff Tear; Reverse Total Shoulder Arthroplasty; Failed Superior Capsular Reconstruction; Failed Rotator Cuff Repair

Superior capsular reconstruction (SCR) is a graft reconstruction of the superior capsule of the glenohumeral joint and provides a static restraint to proximal migration of humeral head in a superior rotator cuff deficient shoulder. The static restraint function of SCR centers the humeral head on the glenoid during active elevation and allows the remaining, intact rotator cuff and deltoid to provide overhead elevation. Although the indications for SCR are continuously evolving, it is an attractive treatment option in younger patients without advanced glenohumeral
arthrosis (Hamada classification 1 and 2) and a painful shoulder due to an irreparable superior rotator cuff tear. Although this procedure was initially described with an autograft, allograft is more popular in the United States. However, even with the recent success of this operation, up to 55% of SCR surgeries have been reported to fail, re-tear, or have non-healing of the graft. Additionally, 5% of patients undergoing SCR require revision surgery, and 20% of patients undergoing this operation report being dissatisfied.

The initial indications for reverse total shoulder arthroplasty (RTSA) were primarily limited to the treatment of cuff tear arthropathy and good to excellent outcomes have been reported for this indication. The indications for RTSA have since then gradually expanded to include conditions other than cuff tear arthropathy, such as treatment of failed RCR or irreparable RC tears without arthritis, revision arthroplasty, and non-reconstructable proximal humerus fractures. With increased utilization of RTSA, risk factors have been identified that can adversely affect the outcomes of RTSA, and include a history of prior shoulder surgery, young age (<50), axillary nerve dysfunction, greater preoperative function, higher expectations and demands, and more recently reported, history of prior arthroscopic rotator cuff repairs (RCR).

Although reverse total shoulder arthroplasty (RTSA) is a salvage operation for failed SCR, to our knowledge, the outcomes of RTSA following failed SCR have not been reported previously. Therefore, the aim of this study was to determine the outcomes of patients undergoing RTSA with a history of an ipsilateral SCR. The null hypothesis of this study was that outcomes of RTSA after a failed ipsilateral arthroscopic SCR would be similar to the outcomes of RTSA with a history of failed ipsilateral arthroscopic RCR.
Methods

Study design

This was a retrospective case-control study (Level of Evidence: III).

Patient Identification and Study Design

An institutional review board approval was obtained for this retrospective case-control study. The SCR cohort was identified from our institutional database by screening for patients who had a history of SCR but subsequently underwent an RTSA due to persistent pain and/or compromised shoulder function. The control cohort was identified in a similar manner from our institutional database by screening for patients who underwent a RTSA for a failed rotator cuff tear (RCT) but did not have advanced radiographic changes (Hamada grade <3) prior to RTSA. The Hamada grade was determined in two ways. First, when available, data were extracted from patients’ medical records according to the preoperative notes of the operating surgeons. Additionally, all pre-RTSA radiographs were reviewed retrospectively by the first author during screening and analysis of cases. The control cohort was selected because patients with prior shoulder surgery, including arthroscopic RCR, have been shown to negatively affect the outcomes after RTSA. Since all the patients in the SCR cohort had a history of prior RCR, this control cohort of RTSA after failed RCR would more closely isolate the SCR procedure as an independent variable to assess its effect on outcomes after RTSA.

Data Collection

Baseline demographics, radiographic characteristics, clinical outcomes, complications, and 90-day emergency department (ED) visits and readmissions were collected and reported.
Baseline demographics included age at the time of RTSA, gender, arm dominance, number of prior rotator cuff repair surgeries, time to RTSA following the last shoulder surgery, and length of stay (LOS) were collected. Preoperative radiographic Hamada grading 8, clinical outcome measures including PROMIS physical function upper extremity score (PROMIS PFUE), Visual Analogue Score (VAS) for pain, and range of motion, [forward flexion (FF) and external rotation (ER)] were collected. 90-day ED visits and readmissions, and 90-day complications including superficial and deep infections, wound drainage, and hematomas, as well as other complications through the entirety of the follow-up were reported.

**Surgical Technique**

All patients were indicated for a RTSA following failure of non-surgical treatment for continued pain and/or shoulder dysfunction (pseudoparalysis) following their previous shoulder surgery (arthroscopic SCR or RCR). Once all the non-operative measures failed, patients underwent RTSA via a standard deltopectoral approach in the beach chair position. A subscapularis tenotomy was performed if subscapularis was present and intact and was repaired at the conclusion of the case. The humeral-sided anchors from the previous arthroscopic procedures were removed prior to reaming and broaching of the medullary canal (Figure 1). Some of the anchors, especially the lateral row anchors, can mechanically interfere with broaching and result in varus placement of the humeral component. The SCR allograft was removed during glenoid preparation. The glenoid-sided anchors were removed if they were in the way during reaming or implantation of the glenoid component. Depending on surgeon's preference, RTSA with onlay or inlay humeral stems as well as intermediate and lateralized glenosphere designs were utilized (Exactech Inc., Gainesville, FL, USA; DJO Surgical, Dallas,
Drains were used in all cases and removed on postoperative day #1. A standardized institutional postoperative rehabilitation protocol was used for these patients, which included immobilization in a sling postoperatively for up to 4 weeks. Passive shoulder range of motion (forward flexion as tolerated and external rotation to 20 degrees) and isometric deltoid strengthening exercises were started on the first postoperative day. Active assisted and active range of motion and isometric external rotation exercises were started at 4-6 weeks. RTSA precautions were used for all patients during the first six weeks after surgery.

**Statistical Analysis**

Standard descriptive summaries (means, standard deviations, percentages) were utilized for baseline demographics and radiographic characteristics. For clinical comparison of the two cohorts, chi-squared test and student t-test were utilized for categorical and numerical data. A p-value of < 0.05 was considered statistically significant. All statistical analyses were done using Graph Pad Prism 5® (Graph Pad, LaJolla, CA, USA) and stored using Excel software (Microsoft Corporation, Richmond, WA, USA).

**Results**

**Demographics**

From 2015 to 2020, 87 arthroscopic SCR reconstructions were performed at our institution. 2015 was the first year identified that a patient underwent a RTSA following a failed arthroscopic SCR procedure. One patient (1.1%) had two revision arthroscopic SCR procedures prior to the RTSA operation. Thirteen patients (~15% of SCR cohort) subsequently underwent RTSA on average 14.6 months (5.8–32.4 months) following the SCR procedure and were...
followed up for an average of 17.9 months (1.6–44.6). The average number of shoulder surgeries prior to RTSA was 2.8 (1–7), with the last surgery being SCR. During the same period, we screened all the RTSA done at our institution and identified 31 patients who underwent an RTSA for failed RCR. Of these 31 cases, only 15 patients had a history of failed rotator cuff repair and Hamada grading <3 and were included in the control cohort. The control cohort patients underwent RTSA on average 12.8 months (1.5–39.5 months) following the last shoulder surgery, and patients were followed up for an average of 27.7 months (2.8–53.9). The average number of shoulder surgeries before the RTSA in the control cohort was 1.4 (1–3). Table 1 shows the demographic and radiographic information comparing both cohorts. There were no differences between baseline demographics or radiographic characteristics (P>0.05), except for the number of prior arthroscopic rotator cuff repair surgeries, which were significantly higher in the SCR Cohort (p=0.03).

Clinical Outcomes

Table 2 compares the clinical outcomes between the SCR and control cohorts for PROMIS PFUE score, pain (VAS), and shoulder ROM (FF and ER). Compared to the control cohort, the SCR cohort had lower preoperative PROMIS scores (30.3 vs. 24.5, P=0.08) and higher preoperative VAS scores (4.1 vs. 7.8, P=0.01). Postoperatively, patients in the SCR cohort demonstrated considerable significant improvements in the pain scores (VAS improved from 7.8 preoperatively to 4.3 postoperatively, [Δ=−3.5], P=0.02) and forward flexion (from 76° preoperatively to 119° postoperatively, [Δ=+43°], P=0.02), but there were modest improvements in the PROMIS score (from 24.5 preoperatively to 29.1 postoperatively, P=0.12).
Overall, compared to the SCR cohort, patients in the control cohort had better function (PROMIS), lower VAS score, and greater ROM (FF and ER) both preoperatively and at last postoperative follow-up. However, preoperative to postoperative changes in PROMIS scores ($\Delta=+4.6$ for both cohorts) and FF ($\Delta=+43^\circ[$SCR cohort] vs. $+44^\circ[$control cohort]) were similar in both groups (Table 2).

Complications

There were no 90-day complications (superficial and deep infections, hematomas, and wound drainage) for either cohort. There were 3 (~23%) complications in the SCR cohort and 4 complications (~27%) in the control cohort. The complications in the SCR cohort included periprosthetic joint infection (PJI), acromial stress fracture, and ulnar neuritis. The PJI was treated with two-stage revision, the acromial stress fracture was treated non-surgically, and the ulnar neuritis symptoms resolved without surgical intervention. Overall complications for the control cohort included one patient with acromial stress fracture (treated non-surgically), one patient with peri-prosthetic joint infection (treated with a two-stage revision), one patient with peri-prosthetic humerus shaft fracture (treated non-surgically), and one postoperative stiffness treated with an arthroscopic capsular release. (Table 3) There were no differences in the LOS, ED admission, and revision surgery in the SCR and control cohorts (P>0.05; Table 3).

Discussion

SCR is a treatment option in patients with a symptomatic irreparable posterior-superior RCT with minimal or no glenohumeral arthritis. Although outcomes after SCR have been
recently reported, there are no reports on the outcomes of RTSA after a failed SCR. In this study, we report short-term outcomes and early complications of RTSA after failed SCR. We found that RTSA after failed SCR improves pain and forward flexion but is associated with modest functional improvements and high complication rates.

Considering that SCR is a final attempt at shoulder preservation following multiple failed rotator cuff repairs and commonly involves the use of an allograft and multiple anchors, it was our hypothesis that the complication rates, especially of infection, will be high when RTSA is performed as a salvage operation for failed SCR. In this study, RTSA following a failed SCR improved pain and forward flexion of the shoulder, but there was only modest overall functional improvement and a high complications rate (23%). In order to evaluate SCR as an independent risk factor for poor outcome after RTSA, in this study, we compared the patients who underwent RTSA after failed SCR to patients who had RTSA after failed arthroscopic rotator cuff repair(s). This control group was essential because all patients who underwent SCR had previous arthroscopic repair(s), and comparing the outcomes of the SCR cohort to this control group will allow us to isolate the role of SCR as an independent risk factor. However, the control group was not age/gender matched and the number of patients in both groups was not large enough to refute or accept our hypothesis. However, patients in the SCR cohort had poor shoulder function as determined by higher VAS score, and lower ROM (FF and ER) preoperatively and at last follow-up compared to the control group. As the number of RTSAs for failed SCR increases in the future, studies with large patient numbers will be feasible to test this hypothesis.

To our knowledge, outcomes of RTSA following SCR have not been reported before. However, previous literature has reported up to 55% of SCR surgeries fail, re-tear, or have non-healing of the graft, and 5% undergo a revision SCR procedure, and 20% of patients undergoing
this operation are unsatisfied\textsuperscript{2,5}. In this study at our institution, 14\% of the patients undergoing SCR subsequently underwent RTSA on average 14.6 months following the SCR procedure.\textsuperscript{5} More recent reports describing SCR clinical outcomes and risks factors for failure, show 7.1\% (1/14) and 5.6\% (3/54) conversion from a failed arthroscopic SCR procedure to RTSA but these rates are lower compared to those reported in this study.\textsuperscript{7,12} Gilat et al also reported that RTSA was performed 6 to 12 months following the failed SCR procedure, which is much shorter than what is reported in our study.\textsuperscript{7}

Prior shoulder surgery, including rotator cuff repair, is a risk factor that can negatively affect outcomes after RTSA. The patients in the SCR cohort had an average of 3 arthroscopic procedures (excluding the SCR), and their shoulders were painful and stiff prior to the RTSA, which was reflected in their preoperative scores (VAS, PROMIS, and ROM). Intraoperatively, extensive subdeltoid and subacromial scarring were noted for these patients requiring more surgical dissection, therefore, it was not a surprise that these patients had residual pain and poor function despite significant improvements in the pain scores and forward flexion of the shoulder after RTSA. These findings are of clinical importance and should be discussed during patients’ preoperative counseling.

There are certain weaknesses in this study, including the inherent bias due to the retrospective design. This would include the recall bias for patient-reported outcomes and data collection. Second, the follow-up in both cohorts is short-term and does not capture the long-term performance of RTSA in these clinical scenarios. Third, the control group was not age and gender matched. Finally, the number of patients in the SCR group is relatively small (13). However, we believe that as more RTSAs will be done for failed SCRs studies with higher patient numbers and longer follow-up will be feasible.
Conclusion

RTSA as a salvage operation following failed SCR improves pain and forward elevation, but improvements in overall function are modest with a high complication rate. Whether failed SCR is an independent risk factor for having poor long-term outcomes after RTSA require future investigation.

References


Figure Legend and Tables

Figure I: Humeral-Sided Anchors and Hardware From Previous Arthroscopy
Table I. Demographic and Radiographic Comparison Between SCR and Control Cohort
Table II. Clinical Outcomes Comparison Between SCR and Control Cohort
Table III. Postoperative Complications in SCR and Control Cohort
Table I. Demographic and Radiographic Comparison Between SCR and Control Cohort

<table>
<thead>
<tr>
<th></th>
<th>SCR Cohort</th>
<th>Control Cohort</th>
<th>p-value</th>
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<tbody>
<tr>
<td><strong>N</strong></td>
<td>13</td>
<td>15</td>
<td></td>
</tr>
<tr>
<td><strong>Age (years)</strong></td>
<td>62.5 (43 – 78)</td>
<td>67.7 (51 – 84)</td>
<td>0.22</td>
</tr>
<tr>
<td><strong>Gender (F/M; % F)</strong></td>
<td>6/7; 46.2 %</td>
<td>6/9; 40 %</td>
<td>0.71</td>
</tr>
<tr>
<td><strong>Arm Dominance</strong></td>
<td>6 (46.2 %)</td>
<td>7 (46.7 %)</td>
<td>1.00</td>
</tr>
<tr>
<td><strong>Pre-Operative Hamada Class</strong></td>
<td>1.9</td>
<td>1.7</td>
<td>0.60</td>
</tr>
<tr>
<td><strong>Number of Prior Surgeries</strong></td>
<td>2.7 (1 – 7)</td>
<td>1.4 (1 – 3)</td>
<td>0.03</td>
</tr>
<tr>
<td><strong>Days to RTSA</strong></td>
<td>436.7 (174 – 971)</td>
<td>383.9 (44 – 1,185)</td>
<td>0.71</td>
</tr>
<tr>
<td><strong>Average Follow Up (months)</strong></td>
<td>17.9 (1.6 – 44.6)</td>
<td>27.7 (2.8 – 53.9)</td>
<td>NA</td>
</tr>
</tbody>
</table>

SCR-reverse total shoulder replacement after failed superior capsular reconstruction; Control-reverse total shoulder replacement after failed rotator cuff repair; RTSA-reverse total shoulder arthroplasty.
<table>
<thead>
<tr>
<th></th>
<th>Preoperative p-value</th>
<th>Postoperative p-value</th>
<th>Δ (preoperative to postoperative change)</th>
<th>p-value</th>
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<tbody>
<tr>
<td><strong>PROMIS PFUE</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SCR (N=12)</td>
<td>24.55 (21.6–29.8)</td>
<td>29.11 (24.2–41.4)</td>
<td>+4.66</td>
<td>0.122</td>
</tr>
<tr>
<td>Control (N=15)</td>
<td>30.33 (24.6–42.5)</td>
<td>34.99 (20.8–43.4)</td>
<td>+4.66</td>
<td>0.113</td>
</tr>
<tr>
<td><strong>VAS</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SCR (N=12)</td>
<td>7.8 (3–10)</td>
<td>4.3 (0–7)</td>
<td>-3.5</td>
<td>0.022</td>
</tr>
<tr>
<td>Control (N=15)</td>
<td>4.1 (0–10)</td>
<td>2.9 (0–10)</td>
<td>-1.2</td>
<td>0.155</td>
</tr>
<tr>
<td><strong>Forward Flexion</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SCR (N=12)</td>
<td>76 (40–130)</td>
<td>119 (90–165)</td>
<td>+43</td>
<td>0.022</td>
</tr>
<tr>
<td>Control (N=15)</td>
<td>94 (30–170)</td>
<td>138 (70–170)</td>
<td>+44</td>
<td>0.011</td>
</tr>
<tr>
<td><strong>External Rotation</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SCR (N=12)</td>
<td>22 (0–60)</td>
<td>29 (0–50)</td>
<td>+7</td>
<td>0.233</td>
</tr>
<tr>
<td>Control (N=15)</td>
<td>32 (0–50)</td>
<td>45 (20–50)</td>
<td>+13</td>
<td>0.099</td>
</tr>
</tbody>
</table>

SCR cohort - reverse total shoulder replacement after failed superior capsular reconstruction.
Control cohort - reverse total shoulder replacement after failed rotator cuff repair.
VAS - visual analog scale for pain.
PROMIS PFUE - patient reported outcomes measurement information system, physical function, upper extremity.
Table III. Postoperative Complications in SCR and Control Cohort

<table>
<thead>
<tr>
<th></th>
<th>SCR Cohort (N = 13)</th>
<th>Control Cohort (N = 15)</th>
<th>p-value</th>
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</thead>
<tbody>
<tr>
<td><strong>90 Day Complications</strong></td>
<td>0 (0 %)</td>
<td>0 (0 %)</td>
<td>N/A</td>
</tr>
<tr>
<td><em><em>90 Day ED</em> Visits</em>*</td>
<td>2 (15.4 %)</td>
<td>4 (26.7 %)</td>
<td>0.48</td>
</tr>
<tr>
<td>Hernia Incarceration, Constipation</td>
<td></td>
<td>Dizziness, PE, Seizure, GI Intolerance</td>
<td></td>
</tr>
<tr>
<td><strong>90 Day Readmissions</strong></td>
<td>1 (7.7 %)</td>
<td>2 (13.3 %)</td>
<td>0.64</td>
</tr>
<tr>
<td>Hernia Incarceration (Required Surgery)</td>
<td></td>
<td>PE, Seizure</td>
<td></td>
</tr>
<tr>
<td><strong>Overall Complications</strong></td>
<td>3 (23 %)</td>
<td>4 (26.7 %)</td>
<td>0.48</td>
</tr>
<tr>
<td>Acromial Stress Fracture (type 3), Ulnar Neuropathy, Deep Infection</td>
<td></td>
<td>Acromial Stress Fracture (type 3), Deep Infection, Stiffness peri-prosthetic fracture</td>
<td></td>
</tr>
<tr>
<td><strong>Length of Stay (days)</strong></td>
<td>1.6 (1 – 5)</td>
<td>1.5 (1 – 3)</td>
<td>0.68</td>
</tr>
<tr>
<td><strong>Revision Surgery</strong></td>
<td>1 (7.7 %)</td>
<td>2 (13.3 %)</td>
<td>0.92</td>
</tr>
<tr>
<td>Infection</td>
<td></td>
<td>Infection</td>
<td></td>
</tr>
</tbody>
</table>

*ED: Emergency Department; GI-gastrointestinal; PE-pulmonary embolism; SCR-reverse total shoulder replacement after failed superior capsular reconstruction; Control-reverse total shoulder replacement after failed rotator cuff repair