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*Int J Spine Surg* 2017, 11 (1)
doi: [https://doi.org/10.14444/4005](https://doi.org/10.14444/4005)
http://ijssurgery.com/content/11/1/5

This information is current as of November 27, 2023.

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Surgical Revision after Sacroiliac Joint Fixation or Fusion

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Abstract

Background
Minimally invasive sacroiliac joint (SIJ) fusion has been shown to be safe and effective for the treatment of SIJ dysfunction. Multiple devices are available to perform SIJ fixation or fusion. Surgical revision rates after these procedures have not been directly compared.

Methods
We retrospectively identified all patients in our practice who underwent SIJ fixation or fusion between 2003 and 2015. Using both chart review and focused contact with individual patients, we determined the likelihood of surgical revision. Revision rates were compared using Kaplan-Meier survival analysis.

Results
Thirty-eight patients underwent SIJ fixation with screws and 274 patients underwent SIJ fusion using triangular titanium implants. Four-year cumulative revision rates were 30.8% for fixation and 5.7% for fusion.

Conclusions
In our study, SIJ fixation with screws had a much higher revision rate compared to SIJ fusion with triangular titanium implants designed for bone adherence.

Background

The sacroiliac joint (SIJ) is increasingly recognized as a common cause of chronic low back pain. In patients presenting for evaluation of low back pain, the SIJ may play a role in 15-30% of cases.1-5 The SIJ is even more commonly suspected (up to 40%) as a source of low back pain in patients who have undergone prior lumbar fusion, possibly due to adjacent segment degeneration. Currently available treatment options for SIJ dysfunction include physical therapy,8 SIJ steroid injections,9,10 RF ablation of the lateral branches of the sacral nerve roots,11,12 and open13 or minimally invasive14-18 SIJ surgery. Two recently published surgery vs. non-surgery randomized clinical trials of SIJ fusion using triangular titanium implants (TTI, iFuse Implant System, SI-BONE, Inc., San Jose, CA) showed that subjects undergoing SIJ fusion had large and statistically superior improvements in SIJ pain, pain-related disability and quality of life compared to those randomized to non-surgical care.19,20

Several approaches and implants are available to perform minimally invasive SIJ fixation or fusion, including dorsal and lateral transfixing approaches. Moreover, strategies can include fixation only or fixation plus fusion. Fusion strategies include devices with coatings that have previously been shown to promote bone ongrowth21-23 or joint disruption with placement of graft materials.

Published literature on the dorsal approach is sparse24-26 and some studies report very little improvement with this approach.24 In contrast, the published literature on lateral transfixing approaches is larger, with a small number of reports of hollow modular anchor screws27-29 and a larger number of reports of TTI, including retrospective case series,15,16,30-34 a combined multicenter analysis,35 a prospective multicenter study36 and two prospective randomized trials.19,20

The need to revise a surgically fused joint is a topic of interest to surgeons and their patients. As report-
ed by the manufacturer, the revision rate after SIJ fusion using TTI is low (3.6% at 4 years)\textsuperscript{37} and clinical trials have shown similar revision rates. TTI are specifically designed for SIJ fusion, with the triangular shape preventing rotational motion and the porous coating promoting biological fixation in bone. The extent to which this design improves over SIJ fixation with screws is not known. Herein we report a comparison of surgical revision rates between these two approaches.

**Methods**

Patients at a single spine surgeon practice who were at least 19 years old at the time of surgery and underwent SIJ fixation or fusion were identified through manual review and querying office billing databases. The fixation procedure was performed using cannulated 7.2 mm diameter stainless steel screws (Synthes, Figure 1. Postoperative AP pelvic x-ray showing typical results from SIJ fixation (left, A) and SIJ fusion (right, B). Figure 1a) as follows. With the patient in the prone position on a Jackson table, 2 or 3 guidewires (depending on patient body habitus) were inserted under fluoroscopic guidance across the SIJ. Following confirmation of pin placement, a 5.5 mm drill was used to drill across the joint and about 2.5 mm into the sacrum. Screws were placed and final imaging was done in the AP, outlet, and inlet views. Live fluoroscopy was also utilized in a 180-degree arc to ensure purchase across the joint. The entire procedure was performed through small (2-5 mm) punctures thru the skin in a percutaneous fashion. SIJ fusion with TTI (Figure 1. Postoperative AP pelvic x-ray showing typical results from SIJ fixation (left, A) and SIJ fusion (right, B). Figure 1b) was performed as previously described.\textsuperscript{30} All procedures were performed under general anesthesia and using fluoroscopic guidance. Through chart review, demographic information and index surgery date were collected and recorded in a study database. Patients known to be dead were reviewed but did not otherwise participate in telephone calls (see below). Patients who underwent revision by the practice’s surgeon (TH) were identified, and the circumstances of revision were noted through review of operative notes and clinic charts. Each patient’s date of last clinic visit was identified.

Patients who had not been seen within 120 days of data collection were contacted by telephone. If contacted, verbal consent was obtained for further study participation. Each participating patient was reminded of his/her SIJ fixation or fusion by the study surgeon. Using lay language, the patient was asked whether the index side had undergone surgical revision by a physician other than the study author. If a revision took place unknown to the authors, an operative note was obtained (no such cases were observed). If the patient could not be contacted or refused to speak with study authors, this was noted in the study database. The study was approved by a regional institutional review board prior to initiation. The study was sponsored by SI-BONE, Inc. (which manufactures the TTI used for SIJ fusion in our cohort).

Baseline characteristics were compared across cohorts using t tests or Wilcoxon tests for continuous variables and chi-squared tests for nominal variables. The cumulative likelihood of SIJ revision surgery...
was calculated using Kaplan-Meier analysis.\textsuperscript{38} Analysis was done at the “side” level (to account for a small proportion of patients who underwent bilateral SIJ fusion.) For patients who underwent revision, time was calculated as the number of days from surgery to first revision. For patients who did not undergo revision surgery, follow-up time was calculated as surgery date to either last clinic visit or the date of telephone contact. In a small number of patients, the chart could not be located or the index surgery date could not be identified; these patients were removed from the analysis. The log-rank statistic was calculated. Demographic factors possibly predicting the likelihood of surgical revision were explored using a similar approach. All statistical analysis was performed using R.\textsuperscript{39}

Results

According to available records, 274 patients underwent index SIJ fusion using triangular titanium implants (TTI) and 38 patients underwent SIJ fixation using screws. Records from 20 patients (11 TTI and 9 screw) could not be located (Figure 2). Demographic and clinical characteristics of included patients are shown in Table 1. Age was higher by about 8 years in the SIJ fusion group but gender and body mass index were similar. Patients undergoing SIJ fixation underwent surgery approximately 6 years before those undergoing SIJ fusion with TTI.

All patients whose charts were located and who were seen more than 120 days prior to study start were telephoned. Of the patients not known to have undergone revision who were last seen in clinic more than 120 days prior to study start, 91.1% were successfully contacted, 7.6% could not be contacted, and in 1.3% contact was established with a family member but not the patient himself/herself. No patient contacted by phone had undergone SIJ revision by another surgeon. Mean (median) follow-up time was 2.8 (3.2) years in the SIJ fusion group and 4.6 (4.9) years in the SIJ fixation group.

Based on review of clinic charts, a total of 31 patients underwent SIJ revision (19 in the fixation group and 12 in the SIJ fusion group). Revision was typically for pain recurrence, and some patients showed evidence of radiolucencies around implants (Figure 3). Using survival analysis techniques, the cumulative probability of revision was lower in the fusion group compared to the fixation group (Figure 4, p<.0001); the 4-year cumulative probability of revision in the fusion group was 5.7% in the fusion group and 30.8% in the fixation group. As of the longest follow-up time point, the cumulative probability of revision in the screw group was 79.8%. Subgroup analysis showed no predictors of revision other than implant used.

Discussion

SIJ fusion is an increasingly accepted surgical treatment option for chronic SIJ dysfunction related to degeneration or disruption of the SIJ. Both SIJ fixation and SIJ fusion work by acutely stabilizing the

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>SIJ Fusion (TTI*) (n=263)</th>
<th>SIJ Fixation (Screw) (n=29)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, mean (range)</td>
<td>54.3 (24.0-85.0)</td>
<td>46.6 (27.0-61.0)</td>
<td>&lt;.0001</td>
</tr>
<tr>
<td>BMI, mean (range)</td>
<td>32.7 (16.3-65.5)</td>
<td>33.8 (22.0-47.3)</td>
<td>0.4703</td>
</tr>
<tr>
<td>N (%) female</td>
<td>63.1% (166)</td>
<td>55.2% (16)</td>
<td>0.5247</td>
</tr>
<tr>
<td>Primary underlying cause, N (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Osteoarthritis</td>
<td>260 (98.9%)</td>
<td>29 (100.0%)</td>
<td>1.0</td>
</tr>
<tr>
<td>Trauma</td>
<td>3 (1.1%)</td>
<td>0 (0.0%)</td>
<td></td>
</tr>
</tbody>
</table>

*TTI = triangular titanium implant

Fig. 2. Patient flow diagram.
painful joint. SIJ fusion allows long-term immobility as a result of adherence of the bone to the lateral transfixing implant as well as bone growing in the SIJ itself.\textsuperscript{33}

Surgical approaches and strategies (and associated devices) for the treatment of SIJ dysfunction vary. Several devices are available for a lateral transfixing approach, including screw-based devices and triangular titanium implants (TTI, iFuse Implant System). The literature for TTI includes 2 prospective randomized controlled trials, a large prospective study and several case series. TTI are designed both to resist rotational motion after insertion and to promote biological fixation in bone.

In our practice, we switched from screws to TTI in early 2012. We noticed that a much smaller proportion of patients returned to clinic with complaints of continued SIJ pain. Surgical revision was required in a large proportion of patients who underwent screw-based fixation. In contrast, the surgical revision rate when using TTI was low, and similar to that reported in the literature.\textsuperscript{37}

The failure mode in patients undergoing screw fixation was primarily loosening and recurrence of pain. The failure mode for patients undergoing TTI was traumatic fracture of the iliac wing secondary to a fall (1 case), malposition of the implant and loosening of the implant(s). It is our belief that standard surgical screws do not have a component that promotes biological fixation in bone, and screw loosening in other applications is common. Whether the modifications in screws incorporated into other FDA-cleared devices for SIJ fusion improve upon screw fixation is not known, and few published data are available to support improved effectiveness.

**Limitations**

Our study has several limitations. First, follow-up in the TTI group was shorter than in the SIJ fixation group, primarily because the TTI group was operated on more recently. Not all TTI patients have 4-year follow-up, so the 4-year revision rate is potentially subject to change. Second, we could not contact locate charts for some patients, potentially introducing a bias. Third, we could not contact a small number of patients. It is possible that patients who could not be contacted underwent revision surgery; however, no other surgeon local to our practice performs such procedures. Third, the time periods for the two groups differ, suggesting that temporal factors, such as surgeon learning curve, changes in OR policies, or postoperative care regimens, could play a role. However, the surgical technique used to perform the SIJ fixation and fusion procedures has not evolved in any important way. Differences are therefore mostly likely due to underlying device design. The number of patients in each group was markedly different, with a smaller number of cases undergoing SIJ fixation. The difference in number of cases is due to the availability of TTI devices that appeared to provide better pain relief, at least in the short-term. Observing better results, the desirability of performing the procedure increased, resulting in more treated patients. Finally, we did not assess pain relief or disability improvement, which have been addressed by other studies.
Conclusions
Surgical revision after SIJ fixation using screws was required far more commonly compared to SIJ fusion using TTI. The device and surgical approach may make an important difference in the overall success of surgical treatments for SIJ dysfunction.

References
19. Polly DW, Cher DJ, Wine KD, et al. Randomized Controlled Trial of Minimally Invasive Sacroiliac...


**Disclosures & COI**
The study described herein was sponsored by SI-BONE. SI-BONE staff helped with statistical analysis. Dr. Tim Holt is a consultant to SI-BONE. Ms. Katie Spain has no financial conflict.

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Published 19 January 2017.
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