Outcomes of Nitinol Compression Staples in Tarsometatarsal Fusion

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Abstract

Background: Tarsometatarsal (TMT) arthrodesis is commonly performed in the management of midfoot arthritis, trauma, or deformity. The purpose of this study was to collect aggregate data (demographic, surgical, and perioperative outcomes) on patients who previously had a TMT fusion with BME compression staples.

Methods: Sixty-six patients underwent TMT fusion with BME compression staples. Outcomes included demographics, surgical information, the Veterans Rand VR-12 Health Survey, Foot and Ankle Ability Measure (FAAM), visual analog scale (VAS), Revised-Foot Function Index (FFI-R), Ankle Osteoarthritis Scale (AOS), patient satisfaction survey scores, radiographic fusion rate, level of pain reduction, and complications. Sixty-six patients (68 feet) were analyzed (59 females) with an average age of 64 years (range, 18-83). The mean latest follow-up was 35.9 (range, 6-56.6 months).

Results: The average surgical time was 38.1 ± 14.3 minutes (range, 11-75). All outcomes improved significantly (P < .001) from preoperative to latest follow-up except for the VR-12 Mental and Physical score. The average time to fusion determined by radiographs was 8.4 weeks (range, 6.1-46.1 weeks). Wound complications were not seen. Indications for subsequent surgeries (26.5%, 18/68 feet) in this current study included pain (n = 14), broken staples, and nonunion (n = 3).

Conclusions: The fusion rate in this study, 89.7%, was similar to values reported in the literature. The patient satisfaction score of 81.9 at latest follow-up is consistent with patient satisfaction for other methods of fusion.

Level of Evidence: Level IV, retrospective case series.

Keywords: tarsometatarsal fusion, staples

Introduction

Tarsometatarsal (TMT) arthrodesis is commonly performed in the management of midfoot arthritis, trauma, or deformity. The vast majority of patients experience substantial improvement in both pain and function with a successful fusion. A variety of conventional fixation devices have been used in TMT fusion, including Kirschner wires, lag screws, staples, compression plate devices, external fixators, and their combinations. Conventional screw placement across the TMT joints is difficult because of the acute angle of screw insertion. Recently, there has been an increase in popularity in the use of shape memory compression staples in orthopedic surgical procedures, including TMT fusions. Purported benefits of staple fixation include ease of insertion, faster time to union, low-profile design, and maximization of joint coaptation. Older-generation nitinol staples required refrigeration prior to implantation and subsequent heating after implantation to achieve their dynamic compression state. In contrast, new generation of nitinol staples, including BME ELITE (Synthes GmbH, Oberdorf, Switzerland), have the ability to elastically recover from deformations, which may occur in vivo, imparting a dynamic compressive capability not possible in conventional fusion methods. This has been demonstrated in numerous in vitro biomechanical studies. This feature of recovering a prior shape, enables the specific staple used...
in this series to be implanted after reduction and alignment of the fusion site with the expectation that the staple would begin to impart compression at the intended fusion site after it was released from its “inserter.” It is reported that compression staples generate a significantly greater compression force across a stimulated osteotomy compared to mechanical staples, and resist permanent deformation, fully recovering their shape following loading.1,3,5,9,14

Numerous studies have demonstrated the safety and efficacy of the use of nitinol compression staples in TMT fusion3,9,13,14,16,23 and generally have fusion rates near 90%, comparable to literature rated for conventional fixation.19 Although some clinical data exist regarding union rate using BME ELITE compression staples in TMT fusion, the data lack objective preoperative and postoperative data and analysis. In this report, validated clinical outcome scores were used to assess the subjective efficacy of midfoot arthrodesis using the new generation of nitinol staples. The purpose of this study was to collect aggregate data (demographic, surgical, and perioperative outcomes) on patients who previously had a TMT fusion with BME compression staples. Primary endpoints include arthrodesis rate and level of pain reduction. Whereas secondary endpoints include Veterans Rand 12-Item Health Survey (VR-12), Foot and Ankle Ability Measure (FAAM), visual analog scale (VAS) for pain, Revised-Foot Function Index (FFI-R), Ankle Osteoarthritis Scale (AOS), and patient satisfaction.

**Methods**

After institutional review board approval was obtained, a retrospective chart review of prospectively collected outcome data was conducted to investigate the surgical and perioperative outcomes on TMT fusion with BME compression staples (Synthes USA, LLC, Monument, CO; or Bio-Medical Enterprises, Inc, San Antonio, TX). All procedures were performed by a single, fellowship-trained foot and ankle surgeon (J.C.C.) between March 2014 and April 2018.

Inclusion criteria were patients between the age of 18 and 85 years, a prior single or multi-TMT joint primary or revision (previously failed TMT fusion procedure) fusion with the use of BME ELITE compression staples (first, second, or third TMT joint, with or without naviculocuneiform joint); and willingness to participate in external research via their clinic admitting form. Patients were excluded if they were younger than 18 years, had diabetes, less than 6 months of follow-up, and/or had an associated talonavicular or calcaneocuboid fusion.

Data collection included patient demographics; medical and surgical history; complications; and pre- and postoperative patient-reported outcomes. Outcome scores included the VR-12,12 AOS,8 VAS,20 FAAM,17 and a patient satisfaction survey. The VR-12 evaluated 8 domains, the scores are tabulated into a summary physical score (PCS) and a summary mental score (MCS) and it follows patient-reported changes in physical and emotional health over time.12 AOS8 is a validated and reliable outcome measure derived from the Foot Function Index. The patient satisfaction survey consists of 6 questions: 5 multiple-choice questions ask the patient to describe their pain relief, ability to perform daily tasks, ability to perform heavy work or recreational activities, meeting expectations (answers range from excellent to poor), and if they would have the operation again. The last question on the patient satisfaction asked, “How satisfied are you with your medical care?” and is a standard 0-100 numeric rating scale, with 0 denoting “least satisfied” and 100 being “most satisfied.”

Eighty-four patients were originally screened, 18 patients were excluded. Sixty-six patients (68 feet) were analyzed (59 women), with an average age of 64 years (range, 18-83). The mean follow-up was 35.9 months (range, 6-56.6). The majority (48/66) of patients were nonsmokers whereas 25.8% (17/66) of patients were former smokers. The average body mass index was 29.6 (range, 20.7-44.3). Primary TMT fusions accounted for 92.4% of the cohort whereas 5 patients/5 feet had a previously failed TMT fusion procedure. The fusions performed included 32 single-TMT fusions, 27 multiple-TMT fusions, 4 naviculocuneiform and single-TMT fusions, and 5 naviculocuneiform and multiple TMT fusions (Table 1). The average surgical time was 38.1 ± 14.4 minutes (range, 11-75).

**Surgical Procedure**

Surgery was performed as an outpatient procedure. The decision to use 1 or 2 incisions was dependent on the number of TMT joints involved in the injury or arthritic process. Second- and third-TMT joint fusions were performed using 1 incision; however, if the first TMT was also involved, 2 dorsal longitudinal incision were made.

The first incision was made over the first TMT joint, just lateral to the extensor hallucis longus. This allowed access to the first and most of the second TMT joints. Pathology involving only the medial 2 TMT joints could be corrected with this single incision. Accessing the entire second TMT joint through this incision carried a risk of injury to the dorsalis pedis artery and deep peroneal nerve. If there was any concern about the reduction accuracy of the second TMT joint or if a surgery included a third TMT joint fusion, a second more lateral incision was used to facilitate exposure and visualization of the second and third TMT joint. The second, lateral incision was in line with the third dorsal

| Table 1. Fusion Procedures. |
|-----------------------------|
|                            | First TMT | Second TMT | Third TMT |
| Single TMT                  | 4         | 25          | 3          |
| Multiple TMT                | 2         | 27          | 25         |
| NC and single TMT           | 0         | 4           | 0          |
| NC and multiple TMT         | 0         | 5           | 5          |

Abbreviations: NC, naviculocuneiform; TMT, tarsometatarsal.
webspace, which was further lateral than what is typically appreciated.

Dorsal bone spurs were removed to expose the joints. If the joints were well aligned, small osteotomes and curettes were used to remove articular cartilage remnants and expose subchondral bone. A saw was used only when significant angular correction was needed to correct the alignment. The opposing surfaces of the joints were perforated with a 2 mm diameter drill to enter subchondral bone. A small curved osteotome was used to microfracture the subchondral bone. If there are small gaps it was filled with local bone graft. As a general rule, the second and third TMT joints are immobilized with a single staple. With the first TMT joint, there is enough room to use two staples for added stability and strength. It can either be placed next to each other over the dorsum, or one over the dorsum and a second medial, or dorso-medial.

The patients were immobilized in a short leg cast splint for 2 weeks, followed by a controlled ankle movement boot for 4 weeks. During that time, patients were encouraged to do active range of motion and did not have to sleep with the boot. Patients were advised to be heel touch weightbearing for the first 6 weeks and could then progress to weightbearing as tolerated if the radiographs looked fine.

Statistical Analysis

Paired-sample t tests were used to determine significant differences in outcome variables from preoperative to latest follow-up. Statistical analyses were performed with SPSS, version 24.0 (IBM Corp, Armonk, NY), and significance was set at $P < .05$.

### Table 2. Clinical Outcome Scores (Mean ± SD).

| Clinical Outcome Measure | Preoperative | Latest Follow-up | P Value |
|--------------------------|--------------|------------------|---------|
| VR-12 Physical           | 35.9 ± 10.1  | 40.1 ± 12.8      | .007    |
| VR-12 Mental             | 54.8 ± 10.1  | 55.0 ± 9.7       | .881    |
| AOS Pain                 | 55.4 ± 22.6  | 27.6 ± 23.5      | <.001   |
| AOS Disability           | 59.8 ± 23.5  | 34.2 ± 26.6      | <.001   |
| FFI-R                    | 69.7 ± 21.3  | 50.3 ± 19.0      | <.001   |
| FAAM ADL                 | 53.3 ± 18.6  | 81.3 ± 19.9      | <.001   |
| FAAM Sports              | 33.4 ± 26.5  | 63.4 ± 33.1      | <.001   |
| VAS                      | 6.0 ± 2.2    | 2.5 ± 2.3        | <.001   |
| Patient satisfaction     | 81.9 ± 22.3  |                  |         |

Abbreviations: ADL, Activities of Daily Living; AOS, Ankle Osteoarthritis Scale; FFI-R, Revised-Foot Function Index; FAAM, Foot and Ankle Ability Measure; VAS, visual analog scale for pain; VR-12, Veterans Rand 12-Item Health Survey.

### Results

Implant size selections were made by the surgeon. Of the 6 first-TMT fusions, 4 two-prong staples and 1 four-prong staples were used. One patient had both two-prong and four-prong staples implanted in the first TMT (Figure 1). For the second TMT, 41 two-prong staples and 20 four-prong staples were used. For the third TMT, 32 two-prong staples and 1 four-prong staple were used.

Clinical outcome scores are summarized in Table 2. Good to excellent relief of their pain after surgery was reported by 63.6% of patients whereas 59.1% patients reported that the surgery met their expectations. Eighty-six percent of the patients stated they would definitely or probably have the operation again.

To determine time to fusion and fusion rate, a single orthopedic surgeon (K.L.F.) not involved in the surgical or clinical care of the patient reviewed sequential postoperative radiographs. Fusion rate was reported for each joint fused as well as the presence of any hardware complications. The radiographic end point fusion was defined as at least 50% osseous bridging across each joint. The average time to fusion was 8.4 weeks (range, 6.1-46.1 weeks), with the longest time of 12.7 weeks for the first TMT compared to the second (7.0 weeks) and third TMT (8.3 weeks) joints, respectively.

Indications for subsequent surgeries (26.5%, 18/68 feet) in this current study included discomfort over the hardware (n = 14), shortening osteotomies (n = 1), and revision surgery for nonunion of the joint (n = 3). The average time to subsequent surgery was 16.8 months (6-48 months). Of the 14 feet with discomfort over the hardware, 5 surgeries were performed because of pain over a single staple and 9 because of pain over multiple staples. Four patients had broken staples; 3 were broken 2-prong staples of the third TMT joint (Figure 2A) and occurred at 6, 15, and 48 months postoperatively. One patient had broken staples of both the second (4-prong) and third (2-prong) TMT joints at 14 months.
postoperatively. Hardware removal performed because of pain resulted in resolution of symptoms in all patients (Figure 2B). One other surgery included distal metatarsal shortening osteotomies due to overload. There were no wound complications.

There were 8 nonunions. Nonunions were defined as no radiologic sign of healing on plain radiographs or computed tomographic (CT) scan. Five of the 8 nonunions were obvious on plain radiographs, whereas 3 were questionable and then confirmed with CT scan. Four of the 8 nonunions had broken hardware, whereas in the other 4 the staples were intact. Four patients had a single first-TMT joint nonunion whereas the other 4 patients were multi-TMT joint. Of the single-joint nonunions, 1 presented in the first TMT, 1 in the second TMT joint, and 2 in the third TMT joint. Four patients had nonunions of both the second- and third-TMT joints. Three of the 8 nonunions had a subsequent revision surgery. The average time to the revision surgery was 20.6 months (range, 6-48). In the 3 patients who underwent nonunion revision surgery, BME staples were removed (1 patient had a broken staple; Figure 3), and different fixation methods were employed (Infuse, fusion with 2 compression screws; CrossRoads, Memphis, TN).

Discussion
Most contemporary studies on TMT fusions report fusion rates of greater than 90% and satisfaction scores surpassing 85%. The occasional discord between lower patient satisfaction despite successful fusion is typically the result of mild residual pain, sesamoid discomfort, or persistent functional limitation. The arthrodesis rate, 89.7% (61/68 feet), and the overall patient satisfaction score, 81.9, in this study are in accordance with literature values.

Smoking has been found to significantly increase the nonunion rate in patients undergoing conventional TMT fusions, with rates ranging from 18.6% to 27%. In the current study, subsequent surgeries in the population of former smokers and current smokers consisted of 5.9% of this cohort (4/68). None of these were nonunions. The current smoker had hardware related pain and removal of both staples at 10.8 months. The other 3 patients needing surgery were former smokers who also had continued pain; in 1 patient, the hardware was removed and shortening osteotomies to unload the metatarsals was performed, and for the other 2 patients' painful hardware was removed.

Symptomatic hardware with conventional methods is common and has been reported to require subsequent hardware removal in 9% to 25% of patients; fortunately, patients predictably experience pain relief with hardware removal. In the current study, subsequent surgeries due to symptomatic hardware was 20.6% (14/68). Eight patients reported improvement in pain and less swelling, but 6 had ongoing pain issues without any identifiable reason for pain.

Recent studies have found nonunion rates with conventional fixation methods to range between 0% and 10% for isolated TMT fusions, with the naviculocuneiform and talonavicular joints generally having the highest rates of nonunion. In this current study, the focus was on the tarsometatarsal joints only. With time, this can be expanded to the larger joints as well.

One of the concerns of using nitinol staples was that the hardware failure rate would be unacceptably high. This did not prove to be the case in this study.

Reduction and fixation can be challenging with midfoot fusion, especially the second- and third-TMT joints. Staple
fixation appeared to be simple and predictable in this patient cohort.

This study is not without limitations. There was no control group or comparative intervention cohort in this study because of the number of surgical options (plates, screws, other staples) for this pathology. Also, because of the retrospective case-series design of the study, a nonresponder bias exists because of incomplete patient data or inability to contact patients for follow-up for outcome data and complication variables. Furthermore, CT scans would have been more reliable assessing the extent and accuracy of fusion.

Conclusion

Numerous studies have demonstrated the safety and efficacy of nitinol compression staples used for in TMT fusion procedures. TMT fusions performed using nitinol compression staples have fusion rates near 90%, which is comparable to the reported values for conventional fixation. The fusion rate in this study, 89.7%, is in agreement with the current available evidence. The patient satisfaction score of 81.9 at latest follow-up is consistent with the reported patient satisfaction for conventional methods of fusion.

Ethical Approval

Ethical approval for this study was obtained from IntegReview IRB (approved 11062017).

Declaration of Conflicting Interests

The author(s) declared the following potential conflicts of interest with respect to the research, authorship, and/or publication of this article: J. Chris Coetzee, MD, reports other from DePuy, during the conduct of the study. ICMJE forms for all authors are available online.

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