Limited-Incision Knotless Achilles Tendon Repair

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The incidence of midsubstance Achilles tendon ruptures is increasing in patients 30 years to 50 years of age, and more than 50% of these injuries occur during recreational basketball.1,2 Achilles ruptures occur more in deconditioned individuals engaged in explosive push-off and jumping activities. Management of these injuries has been controversial over the past decade; there is no consensus on nonoperative treatment, surgical repair, or optimal repair technique.1,3-7 According to American Academy of Orthopaedic Surgeons (AAOS) clinical practice guidelines, limited-incision approaches have fewer overall complications relative to traditional open repair.3,4 Modern repair techniques, such as the Percutaneous Achilles Repair System (PARS; Arthrex), combine limited soft-tissue dissection with percutaneous suture insertion and knot tying.1,8 This limited-incision technique, employed since 2010, uses a 2-cm transverse incision and nondisposable metal jig with divergent needle passes and locking suture fixation options to secure and fix both tendon ends with minimal dissection of skin, subcutaneous tissue, and paratenon. A review of 270 surgically treated Achilles tendon ruptures (101 PARS, 169 traditional open repair) found that, compared with the open repair group, the PARS group had significantly shorter operative times and more patients returning to baseline physical activities within 5 months after surgery.1 Although the difference was not statistically significant, the overall postoperative complication rate was 5% for the PARS group and 11% for the open repair group. The PARS group had no cases of sural neuritis or deep infection requiring reoperation.

Although the PARS technique has had good outcomes with few complications, care must be taken during surgery to prevent sutures from pulling through the tendon near the rupture site, which can result from overtensioning and from suture knot irritation against superficial soft tissues. Given these potential issues, the PARS procedure was modified (Achilles Midsubstance SpeedBridge; Arthrex) to provide knotless restoration of musculotendinous length in a reliable, reproducible fashion and direct fixation of tendon to bone for early mobilization.9 This new procedure bypasses suture fixation in the compromised tendon ends adjacent to the rupture site, thereby reducing suture slippage and allowing for potential early range of motion and weight-bearing relative to previous techniques. Preliminary results from a cohort of 34 patients treated with this technique are promising: Average return to baseline activities was 18.2 weeks (range, 9-26 weeks), and there were no wound complications, nerve injuries, or reruptures.9 Indications are overall health and an acute midsubstance Achilles rupture that presents within 3 weeks after injury (the time limit is used to ensure that both tendon ends can be mobilized and repaired to appropriate length). A relative contraindication is delayed presentation (≥4 weeks), which may require open reconstruction in combination with V-Y lengthening or other adjuvant procedures. Other relative contraindications are insertional rupture, Achilles tendinopathy, and a significant medical comorbidity that prohibits surgical intervention.
Surgical Technique

Operating Room Setup and Approach

The patient is positioned prone with chest rolls and kneepads and with arms at <90° of abduction (Figures 1A-1E).

A thigh tourniquet is placed on the operative extremity, and the feet are placed slightly hanging off the end of the bed with a small bump underneath to adjust the degree of ankle plantarflexion and Achilles tension during the case. It is important that the operative leg be in neutral rotation to allow for central positioning of the PARS jig. After sterile preparation and draping, the extremity is exsanguinated and the tourniquet inflated. The defect within the Achilles tendon is palpated and marked out, and a 2-cm transverse skin incision is made along the proximal aspect of the rupture site.

A “no-touch” technique is used without pickups, and soft tissues are carefully dissected with small scissors down to the paratenon. The sural nerve typically is not visible in the operative field, but, if it is, it can be dissected out and retracted out of the way. A transverse incision is made through the paratenon, and expression of rupture hematoma often follows. Paratenon preservation is key in minimizing disruption of the native vascular supply of the tendon and allowing for repair at the end of the case. A freer can be placed within the wound to confirm that the center of the rupture has been identified.

An Allis clamp is inserted into the wound, and the proximal tendon stump is secured and then pulled about 1 cm through the wound. A freer is circumferentially run along the sides of the proximal tendon to release any potential adhesions that may limit distal excursion.

PARS Jig Insertion and Suture Passing

The PARS jig is inserted into the wound with the inner prongs in the narrowest position possible. The curved jig is inserted proximally, and the center turn wheel is used to widen the inner prongs so they can slide along the sides of the tendon in the paratenon. Proper jig placement should be smooth and encounter little resistance. The proximal tendon is in a superficial location and can be palpated within the prongs of the jig to double-check that the tendon is centered within the jig. A frequent error is to insert the jig too deep, which subsequently causes needles and sutures to miss the tendon and pull through.

Keeping the jig centralized in neutral rotation minimizes improper suture passing and avoids iatrogenic injury to the medial and lateral neurovascular structures. During suture passing, all needles (1.6 mm) with nitinol loops are first used unloaded without suture. The first 2 needles are inserted into their respective, numbered holes, through
the tendon, and then through the opposite side of the jig. Each needle is checked to make sure that it does not pass outside the jig. Having 2 needles within the jig and tendon at all times during suture passing helps stabilize the jig and avoids adjacent suture piercing with the subsequent needle.

A No. 2 FiberWire suture (Arthrex) is then passed through the first hole using the needle suture passer and made even in length on both sides. The specific colors of the suture are not important, but the order of the sutures placed is. An assistant can write down the colors and order of the sutures passed. Before the second suture is passed, the first needle is inserted back through the jig and tendon into the third hole. The third and fourth sutures (green-striped) differ from the other sutures in that one end has a loop and the other has a tail, and they are passed in an oblique, crossing pattern. These sutures later help create a locking suture on either side of the tendon.

After these sutures are passed, the final result should be 1 green-striped loop and 1 green-striped tail on either side of the tendon. The fifth suture is passed straight across the tendon in a trajectory similar to that of the first suture. In large laborers, obese patients, and elite athletes, 2 additional green-striped sutures can be passed through the optional sixth and seventh holes to create an additional locking suture.

**PARS Jig Removal and Suture Management**

After all sutures are passed, the turn wheel is used to narrow the inner prongs while gentle, controlled tension is applied to the jig to remove it from the wound (*Figures 2A-2C*).

All sutures from both sides of the tendon should emerge from the wound. Before the jig is completely pulled out, a hemostat is used through each loop of sutures to guide them out of the wound and reduce tangling. Both pairs of sutures are pulled distally to ensure adequate proximal fixation.

Pullout of any suture from the tendon indicates that the tendon was not centered in the jig or was not proximal enough along the tendon during suture passing. If a suture pulls out, it is removed, and the previous steps are repeated with close attention paid to tendon positioning within the jig. It is not advised to extend the incision longitudinally on either end of the transverse incision, as doing so can lead to potential wound-healing complications. After proximal fixation is achieved, all sutures on each side of the tendon are neatly spread apart in the following order from proximal to distal: first suture, second suture, looped green-striped (third) suture, tail green-striped (fourth) suture, fifth suture. The second suture on both sides is then looped around the 2 green-striped sutures and back proximally through the looped end of the green-striped suture.

The green-striped suture tail is pulled through the tendon to the opposite side to create a locking suture on both sides of the tendon. In the end, there are 2 nonlocking sutures and 1 locking suture on either side of the tendon. Each pair of sutures is pulled distally to confirm fixation and remove any initial suture creep from the system. A
hemostat is placed on each group of 3 sutures to keep them out of the way during distal anchor preparation.

**Distal Anchor Preparation and Banana SutureLasso Passing**

Two longitudinal 5-mm incisions are made along the posterior aspect of the heel just distal to the area of maximal heel convexity. Incisions are spaced 1.5 cm apart along the sides of the Achilles tendon insertion. A 3.5-mm drill and a drill guide are used through each incision and placed flush against bone (Figures 3A-3E).

The drill is inserted into bone oriented slightly proximally and toward midline until it bottoms out against the guide. Each drill hole is then tapped to receive a 4.75-mm SwiveLock anchor (Arthrex).

A Banana SutureLasso (Arthrex) with inner nitinol wire is passed through the center of the distal Achilles tendon stump and out the proximal incision to retrieve one side of the proximal sutures. SutureLasso passage through tendon can be facilitated with tactile feedback. The surgeon’s nondominant thumb is placed directly against the distal tendon while the dominant hand grasps the SutureLasso with the thumb near the tip. As the SutureLasso is advanced proximally through the tendon, the surgeon can feel its tip meeting mild resistance. Confirm that the tip of the SutureLasso is in the center of the distal tendon by direct visual inspection through the wound.

The inner nitinol wire is advanced 2 cm to 3 cm out of the tip of the SutureLasso, and sutures are passed through the distal Achilles tendon. During suture passing, the nitinol wire is drawn back to the tip of the SutureLasso, and then the entire SutureLasso is removed from the distal incision. Trying to pass the sutures only through the inner nitinol wire can result in suture tangling and increased resistance. The process is then repeated for the sutures on the opposite side. Suture pairs are placed under maximal tension and cycled multiple times (5-10) to remove any residual proximal suture creep.10

**Achilles Tensioning and Anchor Insertion**

The ankle is plantar flexed to tension the Achilles tendon relative to the contralateral limb and is held in place by an assistant (Figures 4A-4E).
At the same time, the assistant holds tension on the opposite pair of sutures to ensure that Achilles length and tension do not change before initial anchor insertion. The rupture site can be palpated to confirm there is no residual gap or excessive overlap of the tendon ends. Sutures are passed through the eyelet of the SwiveLock anchor, and then the anchor is gently malleted into the calcaneal drill hole and hand-tightened until flush with bone. Often, squeaking can be heard as the anchor reaches its final depth in bone.

Position of the drill holes can be rechecked with a Kirschner wire before anchor insertion, as their relative position changes with ankle plantar flexion. It is not necessary to premeasure and adjust suture length at the tip of the anchor as in other blind tunnel anchor insertion techniques (eg, InternalBrace; Arthrex). Once the anchor tip is malleted into bone, the free suture ends are released to avoid overtensioning the tendon. Before the anchor insertion handle is completely removed, the tip of a mosquito clamp can be used to feel the bony surface and confirm the anchor is completely seated.

With the ankle still held in the appropriate amount of plantarflexion, the process is repeated and the other SwiveLock anchor inserted. Sutures are cut flush with the anchor, and the surgeon performs wound irrigation and layered closure, with absorbable suture, of the paratenon and subcutaneous tissues. After skin closure with nylon suture, resting ankle plantarflexion is assessed and the Thompson test performed. The patient is placed in a well-padded non-weight-bearing plantarflexion splint for incision and initial tendon healing during the first 2 weeks after surgery.

**Discussion**

A key aspect of recovery is the balance achieved between skin and tendon healing and early mobilization, as outcomes of surgical repair of Achilles ruptures are improved with early weight-bearing and functional rehabilitation. Some surgeons recommend weight-bearing immediately after surgery, given the direct tendon-to-bone fixation achieved with repair. I prefer 2 weeks of non-weight-bearing, which allows the skin to heal adequately and the initial soft-tissue inflammation to subside. If the incision is healed at 2 weeks, sutures are removed, and the patient is transitioned to a tall, non-weight-bearing CAM (controlled ankle motion) boot, worn for 1 to 2 weeks with initiation of gentle ankle range-of-motion exercises. If there is any concern about wound healing, sutures are maintained for another 1 to 2 weeks.

Between 3 and 8 weeks after surgery, progressive weight-bearing is initiated with a peel-away heel lift (~2 cm thick total, 3 layers). Each lift layer is removed as pain allows, every 2 to 3 days. The goal is full weight-bearing with the foot flat 5 to 6 weeks after surgery. Physical therapy focusing on ankle motion and gentle Achilles stretching and strengthening is started 5 to 6 weeks after surgery, depending on progression and functional needs. Between 8 and 12 weeks after surgery, the patient is transitioned to normal shoe wear with increased activities. Running and jumping are allowed, as pain and swelling allow, starting at 12 weeks.

Although preliminary outcomes and experience with the Achilles Midsubstance SpeedBridge have been favorable, long-term clinical and functional studies are needed to determine the specific advantages and disadvantages of this new technique relative to other repairs. The main benefits observed thus far are reduced subjective knot tying and tensioning, decreased reliance on suture fixation in compromised tissue at the rupture site, reduced risk of FiberWire knot irritation of superficial soft tissues, lower risk of distal suture pullout, and earlier mobilization owing to bony fixation of the tendon. Potential complications include anchor-site heel pain caused by prominent anchors or by the bone edema that occurs when a patient increases physical activity by a significant amount at 12 weeks. Heel pain caused by bone edema resolves by 20 weeks without intervention.
Stress shielding of the distal Achilles tendon is a theoretical concern given the tendon–bone construct, but there have been no reports of tendon atrophy or repair failure caused by stress shielding. The original PARS technique was often used to create Achilles tension with the ankle maximally plantar flexed—the idea being that the tendon would gradually stretch over time.\(^1\) Overtensioning the Achilles repair is a potential complication with the SpeedBridge, as the distal anchors provide a more rigid point of distal fixation. Surgeons can avoid this complication by cycling the sutures to remove any residual creep and then tensioning the Achilles according to the contralateral limb and/or palpating tendon opposition at the rupture site.

Overall, this new limited-incision knotless Achilles tendon repair technique allows for minimal soft-tissue dissection, restoration of Achilles musculotendinous length, and direct tendon-to-bone fixation. Early results are promising, but long-term clinical outcomes and comparative analysis are needed. In addition, many details of this technique must be clarified—including incidence of short- and long-term complications in larger cohorts, optimal suture material and configuration, and risks and benefits of immediate (<2 weeks) and delayed (2-4 weeks) weight-bearing.

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**Key Info**

**Figures/Tables**

**References**

**References**


**Multimedia**

**Product Guide**

- STRATAFIX™ Symmetric PDS™ Plus Knotless Tissue Control Device
- STRATAFIX™ Spiral Knotless Tissue Control Device
- BioComposite SwiveLock Anchor
- BioComposite SwiveLock C, with White/Black TigerTape™ Loop
Citation


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