An analysis of functional outcome following tendon augmentation surgeries in patients presenting with steroid-induced tendo achilles rupture and spontaneous tendo achilles ruptures

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Abstract

Introduction: Achilles tendon is the strongest tendon in the body, paradoxically is also the commonest one to undergo rupture. Corticosteroids are one of the most commonly used drugs in tendon disorders; also it is blamed for early and late ruptures. We wanted to assess whether there is any difference in functional outcome in steroid-induced TA rupture group versus spontaneous (no steroid injection) TA rupture.

Materials and Methods: A total of 12 patients were included in the study as per inclusion-exclusion criteria and preoperatively grouped based on previous history of intralesional steroid injection. Both the group underwent same tendon augmentation surgeries and similar physiotherapy protocols. Functional outcome was assessed using AOFAS and Leppilahti Scores.

Results: Better functional outcome was seen in the group with no previous exposure of intralesional steroid injection which was statistically significant (p <0.05) which was confirmed in both scoring systems.

Conclusion: Uses of intralesional steroid correlates negatively with outcome.

Keywords: Achilles tendon disorders, Tendo Achilles ruptures, Peroneus Brevis, Intralesional steroid injection, Tendon augmentation surgeries.

Introduction

Achilles tendon is the strongest tendon in the body, paradoxically is also the commonest one to undergo rupture.1 Disorders of Tendo Achilles (TA) are quite common in sporting as well as in general population.

Repetitive micro trauma and decreased tendon vascularity associated with increasing age have been definitely shown to correlate to Achilles tendon disorders.2 Several mechanical factors like improper footwear with rigid soles, and running on irregular grounds play an important role in causing TA ruptures.2

Management options for TA tendinosis include rest, ice therapy, non-steroidal anti-inflammatory medications, physiotherapy and footwear modifications with custom insoles.3 Eccentric loading exercises have been proven to be effective to the tune of 60- 90%.3,5

Fredberg et al. found good improvement after peritendinous injection of corticosteroid only in short term.6,7 Oral as well as intralesional usage of steroid has been associated with increased incidence of TA ruptures. Corticosteroids are one of the most commonly used drugs in tendon disorders.8 Often misused; the potential risks of adverse effects outweigh benefits. Histological studies in tendinopathy have shown the presence of degeneration and absence of inflammatory markers.9 Hence the role of steroids as an anti-inflammatory agent in tendinopathies is doubtful and debatable. Overall incidence of complications after corticosteroid injection is difficult to measure because of its widespread use and its complications are not commonly reported. Spontaneous complete ruptures have also been reported after steroid injections.10,11

There is paucity of literature detailing long-term effects of Corticosteroid injections in TA tendinopathies. We wanted to assess whether there is any association in patients presenting concerning Steroid induced TA ruptures as against spontaneous TA ruptures(patients have not received intralesional steroid injections).

With this background, we conducted a study in our hospital to assess the functional outcome after surgical treatment for TA rupture. We wanted to evaluate whether there is any difference in functional outcome in steroid-induced TA rupture group versus spontaneous (no steroid injection) TA rupture.

Materials and Methods

This study was a prospective, intention to treat study done at PES Institute of medical sciences and research, Kuppam during March 2015 to Sept 2017. The study subjects included patients who presented with TA ruptures. Patients were allocated into two groups. Group A (Steroid group) included all patients who had received at least one intralesional steroid injection in the past one year prior to rupture of Achilles tendon. Group B included all other patients who had presented with tendo Achillies ruptures without previous history of local steroid injections. Selection of patients was based on inclusion and exclusion criteria as given below.

Inclusion Criteria

1. All patients aged above 18yrs with insertional tendinosis and TA ruptures

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Exclusion Criterion
1. Age less than 18yrs with TA ruptures
2. Peripheral vascular disease
3. Patients refusing to be a part of the study
4. Patients with TA ruptures associated with neurovascular injuries
5. Patients with TA ruptures and Fractures of ipsilateral lower limb

The American Orthopaedic Foot and Ankle Society (AOFAS) Ankle-Hindfoot scale and The Leppihiati Scores were employed for outcome assessment following Tendon augmentation surgeries.

The AOFAS Ankle-Hindfoot scale developed in 1994 is universally accepted outcome measure for comparing different conditions of hind foot\(^{12}\) (Table 1). This is a clinician-based outcome scale collects both subjective and objective factors into numerical scales and has a maximum total Score of 100 points. The subjective portion has been shown to have satisfactory reliability and responsiveness.\(^{13,14}\)

The Leppilahti Score, described in 1998 by Leppilahti et al (Table 2) is a specific evaluation tool for assessing outcome after Achilles tendon ruptures.\(^{15}\) This scoring system combines both subjective assessments of symptoms and objective measures, such as ankle range of motion (ROM) and isokinetic calf strength. It has a total of seven items giving a sum of 100 points as the best possible score.

A detailed history was taken and clinical examination performed for all the subjects included in the study. Preoperative AOFAS Ankle-Hindfoot scale and The Leppihiati scores were tabulated. After appropriate lab investigations and preanesthetic evaluation, patients underwent surgery under spinal anesthesia and tourniquet control. Postero lateral skin incision was used. Ruptured ends of TA were freshened, intratendinous calcification was excised. Peroneus Brevis tendon was used for tendon augmentation. Interferential screw fixation technique was employed for securing the tendons after drilling a tunnel in calcaneum (Fig. 1-8). Layered tension free suturing was done. Operated Limb was immobilized in plaster slab in resting equines position. Suture removal was done at 2 weeks and later limb was immobilized in plaster cast in plantigrade position. At 6 weeks passive range of motion (ROM) exercises were initiated (Fig. 11-12). Graduated weight-bearing was allowed on the operated limb from 12 weeks onwards (Fig. 13).

Patients were assessed before surgery and during follow-up at six weeks, 3\(^{rd}\) month and at 6\(^{th}\) month using AOFAS Ankle-Hindfoot scale and The Leppihiati Scores. Standard physical rehabilitation was given to all patients in the post operative period.
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Fig. 5: Augmentation of TA; Pulvertaft method

Fig. 6: Osseous tunnel in calcaneum

Fig. 7: Anchoring of peroneus brevis with interference screw

Fig. 8: Final repair of paratenon

Fig. 9: Post operative x-ray

Fig. 10: Post operative scar at 6 weeks follow-up

Fig. 11: Thompson test-part 1- before squeezing calf muscle
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Fig. 12: Thompson test- Plantar flexion noted after squeezing calf muscle confirming integrity of gastroc soleus complex.

Fig. 13: Patient bearing weight on operated limb

Fig. 14: Delayed wound healing noticed in Patient 6- settled with conservative management

Statistical Analysis
Comparison was done between Group A (Steroid group) and Group B (spontaneous rupture- no steroid injection) using data obtained from AOFAS Ankle-Hindfoot scale and The Leppihiilati Score. Statistical analysis was done by using SPSS 20 software. Independent t-test was used to compare the mean difference between two groups. Tests for statistical significance were assessed.

Results
During study period, a total of 15 patients with TA rupture were admitted. Only 12 patients were included in the study after applying inclusion and exclusion criteria. 6 patients presented with previous history of local steroid injections in the last one year. One patient had received 2 local steroid injections. All patients were available for follow up for a minimum period of 9 months. Average age at presentation was 58.16 years and 51yrs respectively in Group A and Group B. Both the groups had equal male and female patients. 2 Patients in Group A had well-controlled type II diabetes mellitus. One patient had wound healing problem in Group A (Fig. 14) which settled with regular dressings. There was no re-ruptures of the augmented TA in both the groups. One patient in Group A had sensory problem. (Table 5)

Average Pre operative AOFAS Ankle-Hindfoot scale was 35.50 and 41.17 respectively in Group A and Group B. In Group A Ankle-Hindfoot score improved to 46.83, 75.67 and 81.00 at 6 weeks, 3rd month and 6th months respectively. In Group B the scores were 50.67, 82.00 and 90.17 at similar 6th week, 3rd month and 6th month. Although improvement in scores in group B was noted at 6th week and 3rd month, the difference was not statistically significant. At 6th month follow up however the mean differences in group A and Group B was statistically significant (p-value 0.009) implying better functional outcome in Group B.

The Mean pre operative Leppiihilati Score was 24.17 and 21.67 in Group A and B respectively. The mean difference at six weeks and 3rd month between scores between Group A and Group B were not statistically significant (p-value 0.45 and 0.64 respectively). At 6th month however, the mean difference was statistically significant (p-value 0.01<0.05).
Table 1: The AOFAS Ankle - hindfoot scale

|                      | Pain (40 points) |
|----------------------|------------------|
| None                 | 40               |
| Mild, occasional     | 30               |
| Moderate, daily      | 20               |
| Severe, almost always present | 0               |

**Function (50 points)**

Activity limitations, support requirement
- No limitations, no support: 10
- No limitation of daily activities, limitation of recreational activities, no support: 7
- Limited daily and recreational activities, cane: 4
- Severe limitation of daily and recreational activities, walker, crutches, wheelchair, brace: 0

Maximum walking distance, blocks
- Greater than 6: 5
- 4–6: 4
- 1–3: 2
- Less than 1: 0

Walking surfaces
- No difficulty on any surface: 5
- Some difficulty on uneven terrain, stairs, inclines, ladders: 3
- Severe difficulty on uneven terrain, stairs, inclines, ladders: 0

Gait abnormality
- None, slight: 8
- Obvious: 4
- Marked: 0

Sagittal motion (flexion plus extension)
- Normal or mild restriction (30° or more): 8
- Moderate restriction (15°–29°): 4
- Severe restriction (less than 150): 0

Hindfoot motion (inversion plus eversion)
- Normal or mild restriction (75%–100% normal): 6
- Moderate restriction (25%–74% normal): 3
- Marked restriction (less than 25% normal): 0

Ankle-hindfoot stability (anteroposterior, varus-valgus)
- Stable: 8
- Definitely unstable: 0

Alignment (10 points)
- Good, plantigrade foot, midfoot well aligned: 10
- Fair, plantigrade foot, some degree of midfoot malalignment observed, no symptoms: 5
- Poor, nonplantigrade foot, severe malalignment, symptoms: 0

Table 2: The leppilahti score

|                      | Pain |
|----------------------|------|
| None                 | 15   |
| Mild, no limitations on recreational activities | 10   |
| Moderate, limitations on recreational and daily activities | 5    |
| Severe, limitations on recreational and daily activities | 0    |

**Stiffness**

|                      | Pain |
|----------------------|------|
| None                 | 15   |
| Mild, occasional, no limitations on recreational activities | 10   |
| Moderate, limitations on recreational, but not daily activities | 5    |
| Severe, limitations on recreational and daily activities | 0    |

**Calf muscle weakness (subjective)**

|                      | Pain |
|----------------------|------|
| None                 | 15   |
| Mild, no limitations on recreational activities | 10   |
| Moderate, limitations on recreational, but not daily activities | 5    |
Severe, limitations on recreational and daily activities 0

**Footwear restrictions**

None 10
Mild, most shoes tolerated 5
Moderate, unable to tolerate fashionable shoes, modified shoes tolerated 0

**Active range of motion (ROM) difference between ankles**

Normal (<6°) 15
Mild (6°–10°) 0
Moderate (11°–15°) 5
Severe (>15°) 0

**Subjective result**

Very satisfied 15
Satisfied with minor reservations 10
Satisfied with major reservations 5
dissatisfied 0

**Isokinetic muscle strength (score)**

Excellent 15
Good 10
Fair 5
Poor 0

| Table 3: Group statistics- functional outcome with respect to previous steroid injections |
|-----------------------------------------------|-----------------------------------------------|
| Previous Steroid injections | Total Number (N) | Mean | Standard Deviation | Significance |
| Preoperative AOFAS score | Group A (yes) | 6 | 35.50 | 9.894 | 0.34 |
| | Group B (no) | 6 | 41.17 | 10.722 |
| Preoperative leppilahti score | Group A (yes) | 6 | 24.17 | 10.685 | 0.68 |
| | Group B (no) | 6 | 21.67 | 9.832 |
| 6 week postoperative AOFAS score | Group A (yes) | 6 | 46.83 | 8.232 | 0.38 |
| | Group B (no) | 6 | 50.67 | 6.314 |
| 6 week postoperative leppilahti score | Group A (yes) | 6 | 31.67 | 10.801 | 0.45 |
| | Group B (no) | 6 | 35.83 | 7.360 |
| 3 month postoperative AOFAS score | Group A (yes) | 6 | 75.67 | 5.989 | 0.11 |
| | Group B (no) | 6 | 82.00 | 6.512 |
| 3 months postoperative leppilahti score | Group A (yes) | 6 | 77.50 | 7.360 | 0.64 |
| | Group B (no) | 6 | 79.17 | 3.764 |
| 6 months postoperative AOFAS score | Group A (yes) | 6 | 81.00 | 5.177 | 0.009 |
| | Group B (no) | 6 | 90.17 | 4.579 |
| 6 Months postoperative leppilahti score | Group A (yes) | 6 | 85.83 | 3.764 | 0.01 |
| | Group B (no) | 6 | 91.67 | 2.582 |

| Table 4: Functional outcome with different scoring systems |
|-------------------------------------------------------|-------------------------------------------------------|
| **Group A(Steroid group) (n=6)** | **Group B (N=6)** |
|---------------------------------------------|---------------------------------------------|
| **Group A(Steroid group) (n=6)** | **Group B (N=6)** |
| AOFAS Scale | Leppilahti Score | AOFAS Scale | Leppilahti Score |
| Excellent results (>85) | 33% (n=2) | 5 | 100 % (n=6) | 6 |
| Good (70-85) | 67% (n=4) | 1 | nil | nil |
| Fair (50-75) | nil | nil | nil | nil |
| poor(<50) | nil | nil | nil | nil |

| Table 5: Complications |
|-------------------------|-------------------------|
| **Complications** | **Group A(Steroid group) (n=6)** | **Group B (N=6)** |
| Wound healing problems | 1 | 0 |
| Infections | 0 | 0 |
| Rerupture | 0 | 0 |
Discussion

It was a beginning of a new era when the discovery of cortisone and its effects fetched a group of scientists The Nobel Prize. Use of corticosteroid in select indications has been well documented in the literature. Hench et al and coworkers showed beneficial effects of cortisone in rheumatoid arthritis for which they received Nobel prize in 1950.16 Hollander et al proved that deleterious systemic side effects of steroids could be avoided by local injections.17 In the last 60 years there has been debate about beneficial and deleterious effects of local corticosteroids.

Tendo Achilles tendinosis is a spectrum of disorders ranging from paratenonitis where there is inflammation of peritendinous structure on one end and insertional tendinitis with rupture on the other end.18 Few Studies have shown short-term improvement following local steroid injections.5,7 Frequent relapses were seen in long-term. Intratendinous injection of steroid causes tendon degeneration at site of injection and has been associated with more incidence of ruptures.19 In a study done by Ferland et al for evaluating the effect of local corticosteroid injection in adult albino rabbit’s Achilles tendon the following results were observed. Tendon necrosis seen in intratendinous injection group was an astonishing 100% whereas in peritendinous injection group, tendon necrosis was a mere 5%.20

In a 10 year follow-up study done by Johansen et al sonographic assessment of TA thickness was measured in 3 study groups 0cs-no steroid injection group, 1cs-1 steroid injection group and 2cs-2 steroid injection group. The groups receiving steroid injection showed statistically significant decrease in tendon size with a tendency to worsening. Also in the steroid group late ruptures were noted at 5-8 years post steroid injection.21 A meta-analysis done in 2010 by Coombes et al also showed poor outcomes with steroid injection.22

In the present study (N=12), all of the patients (100%) had previous complaints of retrocalcaneal pain lasting for months prior to rupture and (N=6) 50% of the patients had received intratendinous steroid injections in the last one year prior to rupture. With all of the patients (N=12) presenting with insertional ruptures, tendon transfer was done for all patients. Peroneus Brevis was the tendon of choice in our institute because of its advantages like (a) no residual eversion weakness; (b) PB is an In-phase transfer (c) No weakness of great toe flexion associated with FHL.23

Raghunandan et al in 2014 in a study on neglected tendon ruptures (n=48) observed 29 patients had received intratendinos injection.54 Effect of pre operative steroid on outcome was not studied in this study. In their study author preferred FHL as the preferred choice for tendon augmentation and claimed higher post operative complications with Peroneus Brevis like wound infection (14.28%), delayed wound healing (28.57%) and superficial sensory problems. In our study done with Peroneus Brevis, complications like wound healing (8%) and sensory problem (8%) were seen. Both these complications were seen in Group A (steroid group). There were no complications in Group B.

In the present study, 100% good to excellent outcomes were seen both Group A and Group B, however the percentage of excellent outcome was better in Group B (Table 4). Where as in Raghunandan et al. good to excellent outcome was seen in 91% (44 of48). There is no mention as to whether 9% fair to poor outcome was associated with prior steroid injections.

The present study there was an improvement in Mean AOFAS scores in both Group A (35.5 to 81) and Group B (41.16 to 90.16). The results in Group B was more in line with a study done by Wegryn J25 in 2010 where there was an improvement in AOFAS score from 64 to 98 at 79 months follow-up

In the present study, it was observed that there was improvement after surgery in both group A and B. However there was better improvement in Group B with no previous history of steroid injections as evidenced by both the scoring systems (Table 3 and 4). There was statistically significant improvement in Group B (No steroid) at 6 months postoperatively. The difference in outcomes in steroid group could be due to tendon degeneration, significant decrease in tensile strength and disruption of collagen bundles. In our study all the patients had insertional tendinitis and rupture. Microscopic tendon degeneration at a more proximal area because of steroid injection could also have resulted in lower outcome.

Limitations

This was a single center study with smaller sample size. Hence projection of the results for a general population may not be accurate. A multicentre study with a larger sample and longer-term follow-up is needed for confirming the results obtained in the present study.

Conclusion

Better Functional outcome following augmentation surgeries is seen in patients with tendo Achilles rupture with no previous history of steroid injection than in patients with steroid-induced TA rupture. Uses of intralesional steroid correlates negatively with outcome.

Conflict of Interest: None

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