Gouty Tenosynovitis of the Distal Biceps Tendon Insertion Complicated by Partial Rupture: First Case and Review of the Literature

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Abstract
Background: Given the appreciable prevalence of gout, gout-induced tendon ruptures in the upper extremity are extremely rare. Although these events have been reported only 5 times in the literature, all in patients with a risk factor for or history of gout, they have conspicuously never been diagnosed in the shoulder or elbow. Methods: A 45-year-old, right-hand-dominant man with a history of gout presented with pain in his right anterior elbow and weakness in his forearm after a trivial injury. Results: Here, we report the first case of gouty tenosynovitis of the distal biceps tendon insertion complicated by partial rupture, a composite diagnosis supported by both intraoperative and histological observations. Conclusions: In patients who are clinically diagnosed with biceps tendon rupture and have a history of gout, it is important to consider the possibility of a gout-related pathological manifestation causing or simulating tendon rupture.

Keywords: biceps, tendon, rupture, gout, tenosynovitis

Introduction
Complete and partial tears of the distal biceps tendon are common occurrences, usually due to an eccentric load on the elbow and forearm with the elbow flexed and supinated. Although complete ruptures usually require surgical repair, partial ruptures can be treated conservatively at first but may eventually need operative repair. Such ruptures occur more commonly in males above 30 years of age, smokers lacking tendon nutrition, and individuals taking corticosteroids. In general, however, no obvious predisposing cause for the rupture can be discerned. Distal biceps tendon ruptures are rarely caused by underlying medical conditions.

Gout is a metabolic abnormality caused by the deposition of uric acid, which can form needle-like crystal deposits in soft tissues and joints, and is usually—but not always—seen in individuals with elevated plasma urate levels. It is a systemic disease that may present as an inflammatory deposition arthropathy, soft tissue masses, or both. Kidney involvement includes uric acid lithiasis. Tophaceous deposits can form in any soft tissue or visceral organ, and deposition is favored by biochemical and physical conditions including urate concentration and solubility, pH, temperature, connective tissue structure and ground substance (ie, mucopolysaccharide) matrix material, and crystal metabolic turnover. The most prevalent sites for tophi are the first metatarsal-phalangeal joint (great toe) and the pinna of the ear, due to their lower body temperature and connective tissue ground substance.

The deposition of urates in tissue induces a prominent inflammatory response that may simulate acute septic arthritis. Established tophi show a histiocytic and foreign body giant cell response in a granulomatous-like pattern surrounding the deposited crystals, and the acute gouty attack shows a neutrophil exudate. Tophi may also be present in ligaments and tendons. The complication of gout-induced tendon rupture occurs mostly in the lower extremities, typically affecting the great toe and ankle. Gouty manifestations in the upper extremity are much less common and can include subcutaneous tophi, arthritis, tenosynovitis, and nerve entrapment. Gout-induced tendon rupture in the upper extremity is rare, being reported only 5 times in the hand or wrist but...
never in the elbow or shoulder. Here, we report the first case of gouty tenosynovitis causing rupture of the distal biceps tendon.

**Case Report**

A 45-year-old, right-hand-dominant man with a history of gout, treated with allopurinol, presented with pain in his right elbow and weakness in his forearm after feeling a sudden “pop” while lifting a partition. The pain increased with attempted elbow flexion and forearm supination. He had ecchymosis and swelling extending to the hand. He reported no numbness, tingling, fevers, or chills. Significant medical history included surgical reconstruction of a gouty-infiltrated anterior cruciate ligament several years prior. There was no history of renal disease, smoking, or alcoholism, and no family history of gout. Blood tests revealed no hyperuricemia, evidence of renal disease, or bacteremia.

On physical examination, he had olecranon bursitis with marked posterior elbow tenderness and erythema. There was weakness and pain with elbow flexion and supination. Although swollen, his hand exhibited full range of motion without pain. “Hook” test was positive for pain, but the biceps tendon appeared to be in continuity. The x-ray film of the elbow showed calcifications in the posterior elbow. Magnetic resonance imaging (MRI) of the elbow (Figure 1) showed partial tear of a thickened biceps tendon adjacent to its radial tuberosity insertion and a fluid collection around the insertion. At this time, the patient was tentatively diagnosed with a partial biceps tendon rupture.

After 6 weeks of elbow brace immobilization, his arm remained painful with very limited use. Thus, the patient underwent an olecranon bursectomy and an anterior distal biceps tendon exploration. Intraoperative inspection revealed a gouty biceps tenosynovitis at the tendon insertion site, bathed in a clear exudate with 70% tendon rupture of the biceps head (Figure 2). The tendon was completely detached from its insertion at the radius and substantially infiltrated with gouty tophus (Figure 3). The tendon end was debrided and reinserted into the bicipital tuberosity of the radius using an “endobutton” technique. Intraoperative cultures were negative. Postoperatively, the patient was placed into an IROM™ (incremental range of motion) elbow brace fixed at 90°, and after 3 weeks, a progressive range-of-motion program was started. At 7 weeks, a strengthening program was started.

Microscopic examination of the tenosynovium showed diffuse replacement of normal tendon architecture with mass-like nodules of tophi (Figure 4). The tophi were composed of a palisading inflammatory pattern of mononuclear cells, histiocytes, and multinucleated foreign body giant cells around a crystalline precipitation. The chronicity of the process was appreciated by the generation of a thick
fibrous capsule around some of the tophi. The tophaceous deposits tended to favor the location of degenerative changes within the tenosynovium that were marked with altered matrix mucopolysaccharide changes, increased interstitial edema, and an attempt at reparative cellular proliferation (Figure 5). Some areas of the degenerative changes were accompanied by lytic and cystic changes.

The pathology of our case also demonstrated active tophaceous deposits with an acute fibrinous exudate and connective tissue lysis. This process eroded through normal tendinous connective tissue and caused rupture of the tendon, best appreciated with the trichrome stain (Figure 6).

The destruction of the connective tissue by the inflammatory cells of the gouty tophus was identified as lysis of collagen and reticulin by the inflammatory process. In summary, the pathology sections illustrated tophaceous deposition within tendinous degenerative changes, accompanied by alteration of matrix material. They also showed lytic and cystic degradation of connective tissue, with fibrinous exudation within active tophaceous deposits leading to tendon rupture.

**Discussion**

Here, we report the first case of gouty tenosynovitis leading to distal biceps tendon rupture. Only 5 similar cases where gout is the underlying condition causing or simulating tendon rupture in the upper extremity have been reported in the literature (Table 1). Moore and Weiland\(^6\) presented a 43-year-old man with a long history of gout involving the foot, wrist, and ear who experienced severe burning of his wrist especially at night. Internal neurolysis was performed and tophaceous deposits were observed within the carpal tunnel. Four years later, total infiltration and partial rupture of the extensor pollicis longus (EPL) tendon were observed. Wurapa and Zelouf\(^9\) described a 57-year-old man with a complicated medical history, including gout, who experienced spontaneous loss of index finger flexion. Surgical exploration revealed gouty degradation and rupture of the flexor digitorum profundus (FDP) and the flexor digitorum superficialis tendons. Hung et al\(^4\) reported a 65-year-old man with an extensive gout history and spontaneous painful swelling of the radial dorsal wrist. Surgical examination found rupture of the EPL involving gouty infiltration, with white urate crystals within and around the tendon. Iwamoto et al\(^5\) presented a
A 54-year-old woman with a 3-year history of a painless subcutaneous mass in the dorsum of her hand, an inability to extend her ring finger, and weakness in extending her index finger. Surgical exploration of the wrist exposed hypertrophic crystalline deposits surrounding the extensor digitorum communis and extensor indicis proprius tendons, with rupture of both. Finally, Hankin et al\(^3\) reported a 28-year-old hyperuricemic man presenting with “apparent” rupture of the FDP tendon after sustaining a crush injury. On operative exploration, however, only gouty infiltration of the tendon’s fibrous synovial sheath and adhesions between contiguous structures were observed.

The pathologic mechanism of injury and rupture appears to be due to the deposition of urate crystals within tendinous connective tissue, associated with age-related and possibly trauma-related degenerative changes of tissue structure and matrix material.\(^2\) Deposition and progression of crystal precipitation may be related to nucleation factors, including the fragmentation of collagen and/or matrix material (eg, proteoglycans and chondroitin sulfates). The inflammatory response to urate deposits is associated with connective tissue damage accompanied by collagen lysis and cystic changes. Inflammatory compromise of connective tissue integrity, loss of tendon tensile strength, and fibrinous exudation of active tophi may lead to tendon tears and rupture.

**Conclusion**

Considering the prevalence of gout, it is intriguing that so few cases of gouty tendon infiltration have been reported. Interestingly, gouty infiltration of the distal biceps tendon has never been reported in the literature. In this case, we believe that gouty infiltration was the proximate cause of this patient’s partial rupture and that this etiology should be considered in the differential diagnosis of biceps tendinitis and rupture in patients with a history of gout.

**Ethical Approval**

This study was approved by our institutional review board.

**Statement of Human and Animal Rights**

This article does not contain any studies with human or animal subjects.

**Statement of Informed Consent**

Informed consent was obtained when necessary.

**Declaration of Conflicting Interests**

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