Axillary Artery Injury Following Closed Reduction of an Age-Indeterminate Anterior Glenohumeral Dislocation

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Abstract

Case study: This report details an axillary artery injury associated with an anterior glenohumeral dislocation. The patient was initially evaluated at an outside hospital where the vascular injury was not immediately identified, and then was subsequently transferred to our institution where he underwent bypass grafting without significant sequela.

Conclusion: Although rare, clinicians should actively rule out vascular injuries when evaluating shoulder dislocations, especially in the elderly patient with a known history of atherosclerotic disease or those with evidence of chronic joint instability. Hard signs of vascular injury including diminished distal pulses are the hallmark of this complication, and should always prompt vascular consultation.

Keywords: Axillary artery, Shoulder dislocation, Bypass repair

Introduction

Given the shallow geometry of the glenoid fossa and large range of motion permitted, glenohumeral joint dislocation is a relatively common injury with an incidence of 24/100,000 persons per year. Frequent complications of shoulder dislocation include deltoid weakness, rotator cuff tear, nerve injury, bony deformities, persistent joint laxity, and in rare instances axillary artery damage [1]. Vascular injury is identified by “hard signs” including: active pulsatile hemorrhage, expanding hematoma, palpable/audible bruit, overt limb ischemia, or diminished pulses. Further, “soft signs” of vascular injury include: hypotension or shock, neurologic deficits, stable non-pulsatile hematoma, or proximity of injury to major vascular structures [2,3]. When a vascular injury is suspected urgent evaluation and treatment should be initiated in order to prevent limb ischemia and possible irreversible damage. Here we present and discuss a unique case of an anterior shoulder dislocation with associated axillary artery injury treated with reduction and prosthetic bypass grafting.

Case Report

A 51-year-old intoxicated, homeless man with a history of alcohol abuse initially presented to an outside hospital after being found unconscious at a gas station. Initial survey revealed a right shoulder deformity, and X-ray demonstrated an anterior glenohumeral dislocation (Figure 1). There was no evidence of bony deformities seen on X-ray. Following successful reduction under sedation with etomidate, the patient became increasingly apneic and desaturatated. He was subsequently intubated in order to protect his airway given his level of intoxication coupled with the sedative used for reduction. At this time, the decision was made to transfer the patient urgently via helicopter to a tertiary care center for definitive care.

Upon presentation to our Level 1 Trauma Center as a mid-level trauma, the man remained intubated and sedated. Initial assessment was significant for diminished, but still present, distal pulses in the right upper extremity. Additionally, the capillary refill was delayed to five seconds and the extremity was cooler compared to the left. Due to concerns for vascular involvement, a computerized tomography (CT) angiogram of the right upper extremity was performed and revealed an occlusion of the axillary artery just distal to the takeoff of the subscapular artery (Figure 2). Reconstitution of the brachial artery was found 11.5 cm proximal to the antebrachial fossa. Presence of enhanced extraluminal contrast at the point of occlusion on delayed images was consistent with active extravasation. In addition, an expanding hematoma, which measured 7.3 x 7.7 x 12 centimeters, was noted in the proximal arm. Incidentally, CT imaging also showed evidence of multiple corticated and incompletely corticated ossuous fragments along the superolateral humeral head, and proximal humeral shaft. Additionally, a Hill Sachs deformity was noted in the humeral head. At this time, approximately 7 h after initial presentation to the outside hospital, vascular surgery was notified, and it was decided that immediate surgery was necessary in order to establish hemostasis based on the active extravasation seen on vascular imaging. Given...
that distal pulses were still present, albeit diminished, and that there was reconstitution distally on imaging, it was thought tissue ischemia would be limited. Still, the risk of reperfusion injury was high given the anticipated relative change in vascular flow following bypass, so orthopaedic surgery was consulted in order to perform prophylactic fasciotomies.

Vascular surgery performed a right common carotid artery to mid brachial artery bypass with a 40 cm vascular allograft (CryoArtery, CryoLife- Kennesaw, GA). The decision to use a Cryo Artery allograft rather than saphenous vein autograft was made intraoperatively due to the small size of the patient’s saphenous vein after exposure. Given the extensive amount of hemarthrosis in the shoulder the vascular surgery team chose to tunnel the bypass superficially. For similar reasons, the common carotid was used as an inflow source, as opposed to a more distal segment closer to the apparent vascular injury. Following successful bypass, arm and forearm fasciotomies as well as carpal tunnel release were performed by the orthopaedic surgery team. Upon inspection, the forearm muscles and surrounding tissue appeared viable and not necrotic. Fasciotomy was extended to include carpal tunnel release due to swelling that was noted more distally in the hand. The patient was then brought to the intensive care unit for monitoring post-operatively. The patient returned to the operating room on post-op days 2, 4, and 5 from the index procedure for irrigation, debridement, and delayed primary closure of all wounds. Throughout his hospital course, vascular examination was performed routinely, and showed no evidence of stenosis, thrombosis, or bypass failure. The right upper extremity remained well perfused with palpable distal pulses and brisk capillary refill. In addition, distal neurologic and distal pulses and brisk capillary refill. In addition, distal neurologic and distal revascularization, as denoted by arrows.

![Figure 2](image.png)

**Figure 2:** A) Right axial upper extremity CT angiogram with corresponding coronal view. The arrow denotes the damaged axillary artery. B) Enhancement on delayed imaging is consistent with active extravasation. C) Coronal Maximum Intensity Projection (MIP) showing the proximal transection with distal revascularization, as denoted by arrows.

more rare, a major complication of shoulder dislocation is axillary artery transection, and is associated with a high morbidity if not properly diagnosed. The axillary artery begins as a direct continuation of the subclavian artery as it passes lateral to the first rib. It is further subdivided into three parts, proximal, posterior, and distal, which are defined by the location relative to the pectoralis minor muscle. Major branches of the third portion include the subscapular and circumflex humeral arteries [4]. The distal portion is most commonly injured given its fixed position, which subjects the artery to significant shear forces during traumatic events such as glenohumeral dislocation or subsequent reduction. Clinically, axillary artery injury after an anterior shoulder dislocation is rare [5-7]. Most commonly, this injury is seen in elderly populations over the age of 50 and in the setting of atherosclerotic disease [3]. Atherosclerotic disease leads to decreased arterial elasticity and laxity, which reduces the threshold of shear force required for injury to occur during dislocation or even during dislocation reduction [8]. Axillary artery injury has also been reported in the setting of recurrent shoulder dislocations [5]. Specifically in the setting of relocation, one study showed that close to 70% of patients who underwent closed reduction for a chronically dislocated shoulder experienced a vascular injury [5,6,9,10]. Mechanistically, chronic subluxation and spontaneous relocations lead to increased fibrous tissue deposition over time [9]. As the surrounding soft tissue becomes increasingly stiff so too do the surrounding vascular structures. Via a similar mechanism to that of atherosclerosis, this lowers the shear force threshold needed for vascular injury. CT findings of a Hill Sachs deformity and osseous fragments in this case suggest this patient was subject to chronic joint pathology lending this presentation to previously reported cases. While not known at the time of initial presentation and reduction, care should be taken in the select populations where chronic joint pathology or atherosclerosis is a concern. This could potentially include using more aggressive anesthesia to allow for greater muscle relaxation and easier reduction. Closed reduction under general anesthesia could also be employed especially in patients with an already tenuous cardiopulmonary status as in the case here.

Every patient with a shoulder dislocation requires thorough neurovascular examination, since failure to recognize an underlying arterial injury could lead to limb loss and possibly death. Prompt diagnosis and revascularization is critical, ideally within 2-3 h [10,11]. Clinically, the triad of expanding hematoma, anterior dislocation, and diminished distal pulses has been reported [12,13]. However, given the extensive collateral circulation of the upper extremity, axillary injury has been reported without evidence of decreased pulses [8]. Collateral circulation could propose a mechanism whereby irreversible tissue ischemia can be prevented during extended periods of ischemia [14]. As in this case, CT angiogram showed revascularization distal to the arterial injury consistent with an extensive collateral circulation. This presumably played a significant role in the prevention of tissue necrosis and overall return of limb function post operatively.

If clinical suspicion is high for vascular injury, advanced angiographic imaging should be employed to better quantify the injury. If an injury is confirmed, vascular surgery should be consulted urgently. Most commonly, axillary artery injuries leading to near or complete occlusion are treated surgically by vein or prosthetic bypass grafting. In the case here, surgery was primarily performed in order to establish hemostasis and prevent further active extravasation [15]. In addition, prophylactic forearm fasciotomies should be considered to prevent reperfusion injury associated compartment syndrome. One retrospective study of 62 patients with a variety of upper and lower extremity injuries found the incidence of compartment syndrome to
be 60% (37 patients) following revascularization [16,17]. The majority (31/37) of these patients had a prophylactic fasciotomy at the time of revascularization [18]. Of the remaining patients (6/37), a second surgery was required after revascularization. In these six patients rhabdomyolysis was found in five patients. Three patients required amputation. Additionally, in a recent report of 378 patients with varying vascular injuries of the extremities, six patients required amputation following revascularization as a result of compartment syndrome and subsequent tissue necrosis. These studies highlight the importance and potentially devastating sequelae of reperfusion injury, specifically in the setting of delayed revascularization [19]. Regardless of the warm ischemic time or the presence of collateral circulation, fasciotomies should always be performed if there is concern for reperfusion injury as failure to do so could lead to potentially irreversible limb necrosis.

While this arterial injury is an uncommon complication associated with shoulder dislocation, appropriate identification and treatment in this case resulted in recovery without permanent sequelae.

References