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Core decompression for early-stage avascular necrosis of the humeral head: current concepts and techniques

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Avascular necrosis (AVN) of the humeral head is a rare, yet detrimental complication. Left untreated, humeral head AVN frequently progresses to subchondral fracturing and articular collapse. Cases of late-stage humeral head AVN commonly require invasive procedures including humeral head resurfacing, hemiarthroplasty, and total shoulder arthroplasty (TSA) to improve clinical outcomes. However, in cases of early-stage AVN, core decompression of the humeral head is a viable and efficacious short-term treatment option for patients with pre-collapse AVN of the humeral head to improve clinical outcomes and prevent disease progression. Several techniques have been described, however, a percutaneous, arthroscopic-assisted technique may allow for accurate staging and concomitant treatment of intraarticular pathology during surgery, although further long-term clinical studies are necessary to assess its overall outcomes compared with standard techniques. Biologic adjunctive treatments, including synthetic bone grafting, autologous mesenchymal stem cell/bone marrow grafts, and bone allografts are viable options for reducing the progression of AVN to further collapse in the short term, although long-term follow-up with sufficient study power is lacking in current clinical studies. Further long-term outcome studies are required to determine the longevity of core decompression as a conservative measure for early-stage AVN of the humeral head.

Keywords: Core decompression; Avascular necrosis of bone; Humeral head; Review

INTRODUCTION

Avascular necrosis (AVN) of the humeral head is a rare complication. Atraumatic AVN of the humeral head is infrequently isolated, coinciding with multifocal osteonecrosis including the hip [1,2]. Following AVN, inflammation, fibrosis, and sclerosis of the humeral head occurs as the body attempts to heal the damaged bone. However, if left untreated, AVN frequently progresses to subchondral fracturing and articular collapse [3,4]. AVN symptoms of the humeral head are commonly unrecognized and non-specific in early stages prior to collapse of the humeral head, often leading to misdiagnosis and inaccurate treatment. Particu-

larly, in a prior study, up to 71% of patients in one cohort had progression of their humeral head AVN in the setting of nonoperative management, highlighting a devastating natural history of the condition [5]. In patients with late-stage AVN, including Cruess stage III–V AVN [6], treatment often requires extensive intervention, including humeral head resurfacing [7,8], hemiarthroplasty [9,10], and total shoulder arthroplasty (TSA) [5,10], because management of the osteonecrosis is unamendable to more conservative measures. In patients with early-stage AVN, including Cruess stages I and II, several measures have been described, most notably core decompression [11-14]. Although core decompression has been well described as a viable option

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for management of early-stage AVN in the femoral head [15-21], outcome studies related to the humeral head have been relatively limited without complete consensus on superior techniques to optimize patient outcomes and reduce overall AVN progression.

In this article, the etiology, clinical presentation, and classification of AVN of the humeral head, the indications and current techniques for core decompression of the humeral head, and a complete summary of all current clinical outcomes for each core decompression technique, including the use of adjunctive biologics for the humeral head, are presented.

PATHOGENESIS AND ETIOLOGY

Trauma

Traumatic injury of the proximal humerus is the leading cause of AVN of the humeral head. Proximal humerus fractures increase the risk of disrupting both anterior circumflex and posterior circumflex arteries, the primary vascular supply of the proximal humerus. Significant proximal humerus fracture patterns of humeral head AVN have been reported, including 0%–25% risk for three-part fractures [22-24] and 0%–77% risk for four-part fractures [22-25]. The risk of traumatic AVN has a strong correlation with initial medial hinge integrity and length of the metaphyseal head extension [26], and likely associated with displacement-induced stripping of the periosteum and vessels [27]. However, despite the fracture pattern severity, younger age has been associated with greater resistance to AVN [28].

Atraumatic AVN

Although consensus is lacking on the pathogenesis of atraumatic AVN of the humeral head, several theories have been presented. As previously described, AVN develops due to vascular compromise, commonly due to traumatic etiology. However, certain risk factors and genetic predisposition may increase the risk of vascular disruption, increasing the risk for osteonecrosis through increased intraosseous pressure and compromised blood supply [6,29-32]. Although cases can be idiopathic, several risk factors, both direct and indirect, have been proposed, most notably corticosteroid use [33], sickle cell anemia [31,34,35], systemic lupus erythematosus [36], alcohol consumption [37,38], human immunodeficiency virus (HIV) [39-41], and dysbaric osteonecrosis [42,43].

CLINICAL PRESENTATION

AVN of the humeral head is characterized by insidious onset of

shoulder pain, often without evidence of an inciting event. Patients can experience significant glenohumeral pain, either at rest or with shoulder motion. Some patients may experience pain at night, limiting sleep. In addition to pain, patients may experience shoulder stiffness or weakness of the rotator cuff. In cases of advanced AVN, there may be clicking/clunking of the shoulder with movement due to either an arthritic or collapsed humeral head. Progressive AVN can lead to intraarticular osteochondral fragments, causing locking or popping. Tenderness with shoulder motion is often delayed to later stages of the disease, with most discomfort occurring with maximum glenohumeral loading at > 90° of arm abduction [23].

RADIOLOGIC ASSESSMENT

Initial assessment in suspected AVN of the humeral head should include an extensive clinical and radiographic examination to assess for osteosclerosis, subchondral lucency, and frank joint collapse of the humeral head. A history and physical examination for suspected AVN of the humeral head should first be assessed with plain radiography. On plain films, early osteonecrosis of the humeral head is often not detected (Fig. 1). However, signs of sclerosis or cystic changes may be suggestive of underlying AVN (Fig. 2) [44]. With progression of AVN, the articular surface of the humeral head may collapse due to subchondral fracturing in the area of bone necrosis and identified on plain radiographs by the "crescent sign," an area of subchondral lucency and incongruency of the articular surface. Later stages of AVN progress to flattening, collapse, and progressive degenerative changes [6].

After plain radiography, magnetic resonance imaging (MRI) is considered the modality of choice for the diagnosis of early-stage AVN. The benefits of MRI are associated with sensitive identification of early changes of water and fat content of the bone marrow not evident on plain radiographs [23]. Thus, in cases of suspected osteonecrosis with negative results on plain radiography, MRI of the shoulder should be performed. However, in late stages of AVN, plain radiography and occasionally computed topography (CT) imaging are recommended for further evaluation of subchondral fracturing [23].

CLASSIFICATION

AVN of the humeral head is most commonly classified according to the Cruess staging for osteonecrosis of the humeral head [6] as modified from the Ficat and Arlet classification of the hip [45]. In brief, the Cruess classification describes the severity of osteo-

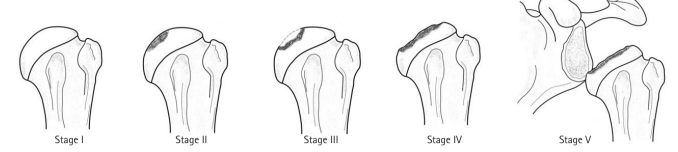


Fig. 1. Illustration depicting the Cruess staging of osteonecrosis of the humeral head. Stage I, no evident changes on radiographic imaging with bone marrow changes only observed on magnetic resonance imaging; stage II, plain radiographs showing focal subchondral osteolysis and/or mottled or wedged sclerosis of the superior humeral head without articular collapse; stage III, subchondral collapse and loss of humeral head sphericity, commonly associated with the "crescent sign"; stage IV, extensive subchondral bone collapse and loss of humeral head sphericity and secondary arthritic changes; stage V, progressive osteonecrosis with arthritic changes extending onto the glenoid articular surface.

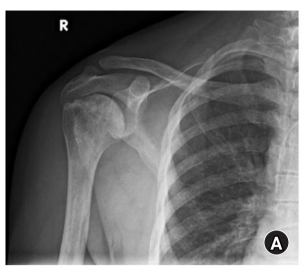


Fig. 2. Preoperative plain anterior-posterior with internal rotation and T2-weighted coronal magnetic resonance imaging (MRI) of the shoulder showing stage I avascular necrosis of the humeral head. (A) The plain radiographic view of the shoulder showing absent osteosclerosis or articular abnormality. (B) T2-weighted MRI showing a T2 signal intensity of the medial humeral head, consistent with osteonecrosis.

necrosis of the humeral head as follows: stage I, negative findings on plain radiographs and bone marrow signal changes on MRI; stage II, plain radiographs showing focal subchondral osteolysis and/or mottled or wedged sclerosis of the superior humeral head without articular collapse; stage III, subchondral collapse and loss of humeral head sphericity, commonly associated with the "crescent sign"; stage IV, extensive subchondral bone collapse and loss of humeral head sphericity and secondary arthritic changes; stage V, progressive osteonecrosis with arthritic changes extending onto the glenoid articular surface (Fig. 3) [11]. In general, the criteria for early (stage I or II) versus late (stage III–V) stage AVN of the humeral head depends on the radiographic and clinical presence of humeral head subchondral collapse in late-stage AVN [11].

NONOPERATIVE MANAGEMENT

In the setting of early-stage osteonecrosis of the humeral head, including stages I and II osteonecrosis where subchondral collapse of the humeral head has not yet occurred, nonoperative management can be explored to address shoulder pain and mechanical symptoms. Prior to considering core decompression or other operative measures, management can include shoulder physical therapy for strengthening and improving range of motion, activity modification, and medical management of any underlying medical conditions that may have contributed to the development of the osteonecrosis as aforementioned. Anti-inflammatory medications may be useful for pain control. However, as previously mentioned, a high rate of progression of humeral head osteonecrosis has been emphasized in prior studies, with more



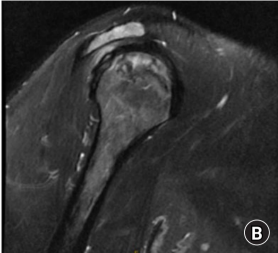


Fig. 3. Preoperative plain anterior-posterior radiograph (A) and sagittal T2 magnetic resonance imaging (MRI; B) view of the shoulder showing stage II avascular necrosis (AVN) of the humeral head. Plain radiographic views of the right shoulder and right humerus show diffuse osteosclerosis consistent with AVN. (A) A subtle subchondral linear lucency at the humeral head without frank collapse is present. (B) T2-weighted MRI showing a focal region of subchondral osteonecrosis of the humeral head with areas of increased T2 signal and a subchondral fracture line without humeral head collapse.

than 71% of patients in mid to late stages having disease progression as early as the 2-year follow-up [5]. This reaffirms the necessity for early diagnosis and intervention to prevent further progression to later-stage AVN, including progression to subchondral collapse and further development of AVN-associated glenohumeral arthritis.

CORE DECOMPRESSION INDICATIONS

Since the introduction of core decompression by Mont et al. [13] for the treatment of the humeral head osteonecrosis, the method has been considered a viable conservative option to prevent further progression of AVN and prolong the time before extensive treatment measures, including humeral head resurfacing, hemiarthroplasty, and TSA, are performed. Although initially described for all stages of humeral head AVN, poor clinical outcomes for patients with late-stage AVN of the humeral head, primarily in Cruess stages IV and V, have been shown in clinical studies. In particular, excellent outcomes in \geq 88% of patients with Cruess stage I or II AVN of the humeral head were reported in early studies, however, in studies by LaPorte et al. [12] and Mont et al. [36], excellent outcomes in stage III osteonecrosis were observed in only 70% of patients. Additional decline in success rates was observed in stage IV AVN, with excellent outcomes as low as 14% [12]. Furthermore, studies by L'Insalata et al. [5] described five shoulders with stage III AVN that progressed to TSA (80%) or stage IV AVN (20%). Thus, based on early evidence, whether core decompression in stage III AVN produces long-term clinical benefit is unclear because subchondral collapse has already occurred. Nevertheless, core decompression is indicated in symptomatic patients with Cruess stage I or II AVN, however, alternatives to core decompression may be considered in stages III-V depending on patient age, functional status, and consideration of arthroplasty longevity. To date, there have not been sufficient clinical studies to further clarify outcomes of core decompression of the humeral head based on other patient characteristics or preoperative shoulder patient reported outcome scores, limiting selection based on clinically symptomatic patients with radiographic signs of early-stage AVN. However, in general, core decompression should be performed in patients with early-stage AVN of the humeral head without articular collapse (Cruess stages I and II), and in patients in whom nonoperative measures including analgesics, physical therapy, and corticosteroid injections failed [44]. Several techniques have been developed for core decompression of the humeral head, and are described in detail in the following section.

CORE DECOMPRESSION TECHNIQUES AND OUTCOMES

A summary of available core decompression techniques and clinical outcome studies for the humeral head is presented in Table 1.

Anesthesia and Patient Positioning

Each technique for core decompression is performed under gen-

Table 1. Clinical outcome studies for core decompression of the humeral head

Complication/	5/6 Stage IV progressed to TSA		4/5 Progressed to arthroplasty at intervals of 1 month, 1 year, 1 year, and 3 years; fifth shoulder with progression to stage IV	4/5 Progressed to arthroplasty at intervals of 1 month, 1 year, 1 year, and 3 years, fifth shoulder with progression to stage IV 16 Shoulders (1 stage I, 2 stage II, 7 stage III, 6 stage IV) required shoulder arthroplasty	4/5 Progressed to arthroplasty at intervals of 1 month, 1 year, 1 year, and 3 years; fifth shoulder with progression to stage IV 16 Shoulders (1 stage I, 2 stage II, 7 stage III, 6 stage IV) required shoulder arthroplasty houlder arthroplasty 8 stage II progressed to stage III progressed to stage III progressed to stage III progressed to stage III progressed to stage IV	4/5 Progressed to arthroplasty at intervals of 1 month, 1 year, 1 year, and 3 years; fifth shoulder with progression to stage IV 16 Shoulders (1 stage I, 2 stage II, 7 stage III, 6 stage IV) required shoulder arthroplasty houlder arthroplasty arge III, 12 stage III progressed to stage III progressed to stage III progressed to stage III progressed to stage IV 19 attent with spondy-loarthropathy had poor outcome (11>14 postop) although with multijoint involvement
Outcome	ng 20	good or excellent; at time of follow-up, 11/14 stage I or II deteriorated radiologically; of 10 stage III—7 with excellent, 3 with arthroplasty; of stage IV, 1 with good and 5 requiring arthroplasty	4,	ellent; at time of foldes tage I or II deterioogically; of 10 stage excellent, 3 with arbitage IV, 1 with good ing arthroplasty 27, stage 1, 17–27; 27; stage 3, 14–25; 20	4 2 Z	1
Disease stage	Six shoulders had UC stage I changes, e. e. 8 stage II, 10 stage III, and 6 stage ii IIV, and 6 stage ii IV	(ELT)	Stage III ^{b)} No	N ". stage A ye II, 13, S group: group: stage e III, 5;	N stage A ge II, 13, 18, 5 group: group: stage e III, 5; 2 stage II, 0 stage II, 13, 14, 8b)	N: stage A 3e II, 13; 18; 5 6 III, 13; 2 stage e III, 5; 2 stage II, U III, 34; 8 ^{b)} U fied by
Graft	No graff		No graff		r.	r er
Procedure	5-mm diameter coring device (single large trephine)		Humeral head drilling	Humeral head drilling drilling Single large trephine, percutaneous fluoroscopic guided core decompression	Humeral head drilling Single large trephine, percutaneous fluoroscopic guided core decompression 5-mm diameter coring device (single large trephine)	Humeral head drilling Single large trephine, percutaneous fluoroscopic guided core decompression coring device (single large trephine) Percutaneous small-diameter perforations technique under fluoroscopy guidance
Follow-up	5.6 yr		Range, 1 mo–4 Humeral head yr drilling	Range, 1 mo-4 yr 10 yr (2-20)		
Age (range, yr)	/sex 38 (22–76)/ 11 E 9 M	Not indicated	ואסן ווותורפוכת	CSI group: 37 (22–60)/ 20 F 14 M; Non-CSI group: 41 (23–78)/ 4 F 5 M	CSI group: 37 (22–60)/ 20 F 14 M; Non-CSI group: 41 (23–78)/ 4 F 5 M Total project: 41 (20–60)/ 47 F 26 M; Core decompression specific not listed	CSI group: 37 (22–60)/ 20 F 14 M; Non-CSI group: 41 (23–78)/ 4 F 5 M Total project: 41 (20–60)/ 47 F 26 M; Core decompression specific not listed 37 (15–50)/ 7 F, 8 M
No. of patients	20 (30)	4 (5)		43 (63)	43 (63)	43 (63) 95 Shoulders 15 (26)
Study type	Retrospective review	Retrospective	Icview	review Retrospective review	Retrospective 43 (63) CSI group: review 20 F 14 M; Non-CSI group: 41 (23–78)/4 F 5 M review 95 Shoulders Total project: review 47 F26 M; decompressi	Retrospective review review review Retrospective review review review
Study	Mont et al. (1993) ^{al} [13]	L'Insalata et al. (1996)	<u>\(\tag{\alpha} \)</u>	rre et al. 98) ^{a,,d)}]		al. 36]

(Continued to the next page)

Table 1. Continued

Complication/ advancement of disease	, Patient 2: persistent pain at 9 month FU, resolved with injection of Decadron and xylocaine	None 1	Stage IV shoulder requiring TSA at 20 months post-index procedure	3/30 Mesenchymal treated progressed to collapse vs. 25/34 in untreated group (P < 0.0001); 2/30 mesenchymal treated progressed to arthroplasty vs. 25 in untreated group	2 Stage II and 1 stage IV progressed to hemiar- throplasty
Outcome	Final FU: patient 1, FF 165°; Ext rot, Patient 2: persistent pain 85°; IR T12, no pain medication for shoulder; patient 2, IR T12 final follow-up; patient 3, no pain medication at final follow-up, and xylocaine range of motion was same as prior to surgery	Asymptomatic with full range of motion (shoulder 1); mild residual stiffness (shoulder 2)	Stage III: average change of VAS from 33 mm preoperative to 5 mm postoperative; stage IV: failed joint sparing requiring TSA at 20 months post-index procedure	Average improvements in constant scores of $40 (35-56)$ to $71 (50-88)$ in cell therapy group and $43 (31-60)$ to $57 (43-70)$ in untreated (P=0.01); average improvement in VAS scores of 45.8 ± 7.2 to 16 ± 3.2 in cell therapy group and 43.2 ± 8.5 to 29 ± 4.1 in untreated group (P=0.01)	Precollapse SPADI scores ¹ : 1 poor, 4 good, 2 excellent; postcollapse SPADI scores: one poor, 3 excellent, 4 missing outcome scores
Disease stage	Stage II	Extensive subchondral osteonecrosis without joint collapse - bilateral; staging not indicated	Stage III, 4; stage IV, 1°)	Percutaneous Core Mesenchymal Mesenchymal cell decompression cell injection treated 1: stage II, (30/64 8; stage II, 10; patients); stage III, 7; stage untreated IV, 5 simple coredecompression (34/64)	8 Precollapse ^{e)} (stage I and stage II); 6 postcollapse (2 stage III)
Graft	No graft	Synthetic bone Extensive grafting subchon (Pro-Dense, osteonec Wright without Medical) collapse bilateral; not indic	Autologous concentrated bone mar- row	Mesenchymal cell injection (30/64 patients); untreated simple coredecompression (34/64)	Fibular strut Allograft
Procedure	Arthroscopic- assisted core decompression with ACL tibial drill guide	Arthroscopic- assisted core decompression	Percutaneous fluoroscopic guided core decompression	Percutaneous Core decompression	Arthroscopic- assisted core decompression
Follow-up (range)	Patient 1: 7 mo; patient 2: 9 mo; patient 3: 11 mo	8 mo (shoulder 1); 4 mo (shoulder 2)	49.4 mo (24–73)	7 yr (5–10)	Precollapse: 7 yr (3–18); postcollapse: 2.4 yr (1.5–3)
Age (range, yr) /sex	37, 49, 36 /3 F	51/F	48 (38–63)/ 3 E 1 M	24 F, 37 M	Precollapse: 44 (33–61); postcollapse: 30 (19–40)
No. of patients (shoulders)	3 (3)	1 (2)	4 (5)	30 (30) Mesenchymal cell treated; 34 (34) matched pair control group	14 (17)
Study type	Case report	Case report	Case report	Retrospective 30 (30) Mes- 46.5 (28-60)/ review enchymal 24 F; 37 M cell treated; 34 (34) matched pair control group	Retrospective review
Study	Dines et al. (2007) [46]	Steffensmeier Case report et al. (2016) [47]	Makihara et al. (2017) [48]	Hemigou et al. (2020) [23]	Galloway et al. (2013) [49]

^{a)}Contain same patient population; ^{b)}Staging based upon Ficat and Arlet Classification; ^{c)}UCLA shoulder rating system. Ten points assigned for each of three categories; pain, function and active movement with max of 30; excellent 27 or more, good is 24–26, fair 21–23, poor <21; ^{a)}20 (30) from Mont et al. (1993) [13]; ^{e)}Staging based upon Cruess classification; ^{b)}SPADI; scores 0–25 deemed excellent; 26–50 are good; 51–75 are fair; 76–100 are poor. UCLA: University of California Los Angeles, TSA: total shoulder arthroplasty, CSI: corticosteroid injection, FU: follow-up, SCD: sickle cell disease, RTSA: reverse total shoulder arthroplasty, FF: forward flexion, Ext. external rotation, IR: internal rotation, VAS: visual analog scale, SPADI: Shoulder Disability and Shoulder Pain and Disability Index.

eral anesthesia with the patient either in the supine or beach chair position on a standard operating table.

Open Technique

Mont et al. [13] were the first to present core decompression for AVN of the humeral head using a standard technique with a 5-mm coring device (single large trephine). The standard open technique involves utilizing the deltopectoral interval. To perform the open technique, a 2-cm incision is made superior to the pectoralis major tendon anterior to the axillary fold. The deltopectoral interval is developed by blunt dissection until the proximal humerus is reached. Care should be taken to not disrupt the common neurovascular structures within the interval [50]. A single coring device is then utilized and drilled into the proximal humeral metaphysis slightly lateral to the bicipital groove. Accurate decompression should be confirmed with intraoperative fluoroscopy corresponding to the lesion position observed on preoperative imaging. Although originally described with a 5 mm trephine, several other techniques have been developed with single trephines ranging from 6 mm- 10 mm in size [12,13,36].

Outcomes of Open Technique

Mont et al. [13] reported good to excellent University of California Los Angeles (UCLA) shoulder score outcomes for core decompression of Ficat and Arlet [45] stage I or stage II humeral head AVN with no progression to TSA. Conversely, 3 of 10 stage III and 5 of 6 stage IV shoulders progressed to TSA, with the 1 stage IV shoulder not progressing to TSA having an outcome of only good. Similar results were observed in follow-up studies by LaPorte et al. [12] and Mont et al. [36], with excellent outcomes of early-stage humeral head AVN and high failure rates in shoulders with stage III–V AVN. This was consistent with studies by L'Insalata et al. [5] in which all five shoulders with stage III AVN progressed to TSA (80%) or stage IV AVN (20%).

Percutaneous Technique

Although the traditional technique for core decompression has been shown an effective technique for both the hip and humeral head, a small-diameter percutaneous technique showed excellent results for core decompression of the hip [16,51]. The inception of this technique was based on the theory that multiple passes into the necrotic lesion will increase the coverage of the necrotic lesion, and the percutaneous technique reduces surgical morbidity in comparison with the larger surgical incisions required for a large trephine. More recently, the technique has been adapted for the humeral head [11]. In brief, the technique by Harreld et al. [11] utilizes multiple small diameter percutaneous perforations using a 3.2-mm Steinmann pin advanced twice, for smaller lesions, or three times for larger lesions, to the epiphyseal lesion under fluoroscopic guidance (Fig. 4). Insertion of the Steinmann pin should be localized lateral to the bicipital groove, reducing risk of damage to the ascending branch of the anterior humeral circumflex artery [52]. Kennon et al. [10] also presented this technique utilizing a standard 2.7-mm drill bit passed several times into the necrotic lesion under fluoroscopic guidance. Although a small-diameter drill is more commonly used, a single large trephine may also be utilized for the percutaneous technique albeit with a larger initial incision to accommodate the larger drill (Fig. 5). The authors recommend a lateral approach when a larger drill, reamer, or trephine is used. Under fluoroscopy, a lateral point of entry inferior to the level of the axillary nerve is localized using a spinal needle. An incision is then made and a blunt dissection performed down to the lateral cortex of the proximal humerus. A blunt trocar can then be used for dilation of the incision and to reduce risk of iatrogenic nerve injury. Next, a standard core decompression with a large trephine is performed as previously described.

Outcomes of Percutaneous Technique

Harreld et al. [11] assessed the outcomes of patients with Cruess

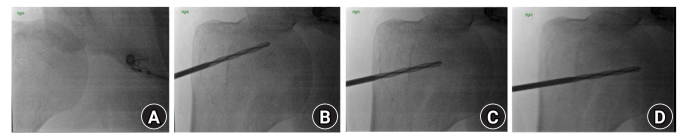


Fig. 4. Right shoulder, beach chair position. Illustration of small-diameter percutaneous core decompression of the humeral head using a 3.5-mm drill bit. (A) Under fluoroscopic guidance, the area of necrosis is identified. (B-D) Several passes with the drill are then made into the necrotic lesion, creating 2–4 decompression tracts.

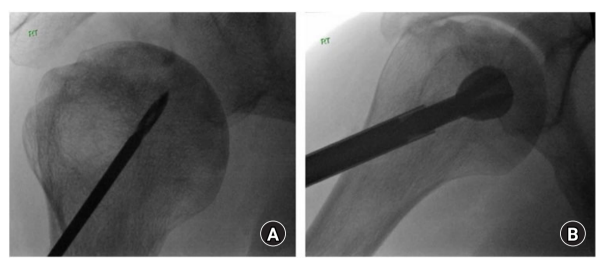


Fig. 5. Right shoulder, beach chair position. Illustration of core decompression using a 9-mm reamer. (A) Placement of the guide pin within the humeral head. The guide pin is directionally placed from a lateral approach, directed under fluoroscopic guidance to the necrotic lesion. (B) The avascular necrosis lesion is then over-reamed using a 9-mm diameter reamer with a single pass.

stage I or stage II AVN of the humeral head treated with percutaneous core decompression. In a cohort of 15 patients (26 shoulders), the mean UCLA shoulder scores increased from 14 (range, 10–22) preoperatively to 27 (range, 14–30) postoperatively at the mean follow-up of 32 months (range, 24–41 months). Only 1 patient had poor outcome with a UCLA score of 11 preoperatively and 14 postoperatively, however, this patient was complicated by spondyloarthropathy and multiple joint involvement of the AVN.

Although good outcomes for core decompression were observed in previous studies, mixed results were reported in a case series by Kennon et al. [10] in which 11 patients (11 shoulders) with stage I or stage II humeral head AVN were assessed for outcomes following core decompression. The procedure was performed using a standard 2.7-mm drill bit with multiple fluoroscopically guided passes with concurrent intraoperative ultrasound bone stimulation. Among the 11 patients, 3 were lost to 1-year follow-up. However, two of three patients continued to have persistent pain postoperatively. At the mean follow-up of 17.4 months, seven of the eight remaining patients progressed to further collapse and five required additional resurfacing procedures due to continued postoperative symptoms. Functional outcomes in that study were only assessed following resurfacing, TSA, or reverse TSA, however, the high failure rate and progression within 2 years postoperatively raises concerns for long-term durability of the procedure.

Arthroscopic-Assisted Technique

The use of arthroscopy in core decompression of the humeral head was first introduced by Chapman et al. [53] as an adjunctive measure for accurate articular assessment and placement of the guide pin. The main premise of this technique is to reduce the risk of iatrogenic perforation of the articular surface while additionally allowing the surgeon to treat coinciding shoulder pathology, including synovitis and chondral pathology, which may limit optimal postoperative recovery. Furthermore, arthroscopy provides the benefit of direct observation of articular surface collapse without the need of an open incision. In brief, arthroscopic-assisted core decompression is performed using two standard portals, including posterior and anterior portals. After the arthroscope is placed in the posterior portal, a standard diagnostic arthroscopy is performed with full assessment of the intraarticular cartilage, labrum, rotator cuff, biceps, and subacromial bursa. During the arthroscopy, any intraarticular pathology should be appropriately managed. In addition, arthroscopy allows for the absence of humeral head collapse, which would contraindicate core decompression, to be confirmed. Once diagnostic arthroscopy is complete, a lateral portal is established and a standard percutaneous core decompression is completed. Intraoperatively, the arthroscope is maintained within the posterior portal to confirm the drill has not disrupted the articular surface of the humeral head.

In addition, the use of an anterior cruciate ligament (ACL) tibial drill guide has been described an effective tool for accurate triangulation of the humeral head AVN lesion while reducing the risk of unnecessary drilling [46,49]. Dines et al. [46] described the technique in terms of a standard arthroscopic-assisted core decompression with use of only the anterior and posterior arthroscopic portals, however, the authors prefer a laterally directed mini-open incision for placement of the ACL targeting guide (Fig. 6). Due to the proximity of the axillary nerve in this ana-

tomic area, the incision should be carefully extended down to the lateral cortex of the humerus. Fluoroscopic imaging is used to localize the necrotic lesion and the ACL tibial drill guide is positioned to triangulate the area of necrosis. Next, a soft tissue protector is placed within the incision and a guide pin is drilled into the lesion. Direct visualization using arthroscopy and fluoroscopy is performed to confirm the lesion has been reached and the pin has not penetrated through the subchondral bone into the cartilage. Then, using a 4-mm cannulated drill, a decompression channel is created by aligning with the ACL guide (Fig. 6). The guide is then moved two to three more times with concurrent drilling as described to achieve appropriate spread within the lesion. Then, the humeral head cartilage should be reinspected to ensure iatrogenic penetration has not occurred.

Outcomes of Arthroscopic-Assisted Technique

Dines et al. [46] reported the outcomes of three patients (3 shoulders) that underwent arthroscopic-assisted core decompression as first described by Chapman et al. [53]. Their technique included the use of an ACL tibial drill guide to reduce the risk of articular perforation. All three patients presented with Cruess stage II humeral head AVN based on preoperative radiographs and MRI. At an average follow-up of 9 months (range, 7–11 months), two of three patients had complete resolution of their preoperative pain, and shoulder range of motion was completely restored in three of three patients. One patient presented with persistent pain at the 9-month follow-up, although this was completely resolved with a corticosteroid injection. Long-term outcomes for this technique with and without the use of an ACL tibial drill guide are not currently available for the humeral head.

Adjunctive Biologic Treatment

More recently, significant focus has been on the concurrent use of growth and differentiation-inducing agents following core decompression to promote native bone regeneration of the decompressed tract. In particular, methods including autologous bone grafting [15], tantalum rod implants [54-56], autologous bone marrow cell implants [57], and synthetic calcium sulfate/calcium phosphate bone cementing [58-64] have been described for core decompression in the femoral head. Although current literature on biologics for core decompression of the humeral head is limited, several techniques have been proposed.

Synthetic Bone Grafting Technique

As presented by Steffensmeier et al. [47], the use of a calcium sulfate and calcium phosphate synthetic bone graft (Pro-Dense, Wright Medical Group, Memphis, TN, USA) is a viable option for adjunctive management with core decompression. Although not sufficiently studied for the humeral head, synthetic bone graft has been proven to promote bone regeneration following femoral head AVN [65,66]. The technique is performed following a standard percutaneous core decompression. Through the original percutaneous incision, an extension cannula is placed at the decompression tract and synthetic bone graft is injected into the proximal humerus (Fig. 7). Fluoroscopy is utilized to visualize complete cementation of the decompression tract. Arthroscopy with irrigation is recommended to ensure no cement is present within the subacromial space.

Outcomes of Synthetic Bone Grafting Technique

Steffensmeier et al. [47] reported on one patient (two shoulders)

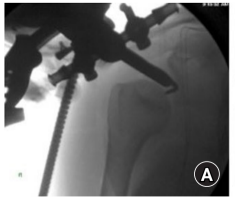






Fig. 6. Right shoulder, beach chair position. Intraoperative fluoroscopic imaging depicting percutaneous core decompression of the humeral head using an anterior cruciate ligament (ACL) tibial guide. (A) An ACL tibial drill guide is positioned through a lateral portal to triangulate the area of necrosis on the humeral head. (B) A soft tissue protector is then placed within the incision and a guide pin is drilled into the lesion. (C) With a 4-mm cannulated drill, a decompression channel is created by aligning with the ACL guide. The guide is then moved two to three more times with concurrent drilling as described to achieve appropriate spread within the lesion.

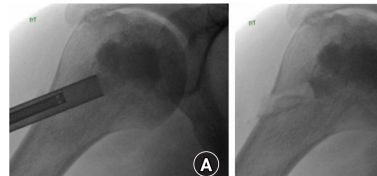






Fig. 7. Right shoulder, beach chair position. Intraoperative fluoroscopic imaging demonstrating core decompression with synthetic bone grafting. (A) Through the original percutaneous incision, an extension cannula is place at the decompression tract and synthetic bone graft is injected into the proximal humerus. (B, C) Fluoroscopy is used to visualize complete cementation of the decompression tract. Arthroscopy with irrigation is recommended to ensure no cement is present within the subacromial space.

with early-stage AVN of the humeral head who underwent core decompression with synthetic bone graft cement (Pro-Dense, Wright Medical Group). Improvement in pain and range of motion was observed at the final follow-up of 8 months for 1 shoulder and mild residual stiffness was observed at the 4-month follow-up. Neither shoulder progressed to collapse and pain in both shoulders was resolved.

Autologous Concentrated Bone Marrow Cell Technique

The use of autologous mesenchymal cell grafts for core decompression of the femoral head has been well described. In a meta-analysis by Li et al. [57], improved outcomes were observed in patients receiving congruent mesenchymal injections compared with core decompression alone. Makihara et al. [48] recently showed this technique was a viable adjunctive treatment for core decompression of the humeral head. Bone marrow aspiration, centrifugation, and transplantation were performed according to the technique by Yoshioka et al. [67] for the femoral head. In brief, under general anesthesia, bone marrow aspirate is harvested using a bone marrow biopsy needle with an acid citrate dextrose containing syringe. Centrifugation is performed twice. The first centrifugation is performed at 1,200 ×g for 10 minutes to r emove the red blood cell layer. Following a second centrifugation at 3,870 $\times g$ for 7 minutes, the blood plasma is r emoved, leaving buffy coats containing the mesenchymal cell grafts. After standard percutaneous core decompression, the autologous concentrated bone marrow graft is grafted with a cylindrical rod smaller in diameter than the diameter of the drill to ease implantation. The graft is then implanted within the decompression tract. The procedure is completed by closing the incision using standard technique.

Outcomes of Autologous Concentrated Bone Marrow Cells Technique

Makihara et al. [48] reported four patients (five shoulders) who underwent core decompression for four shoulders with Cruess stage III and 1 shoulder with stage IV AVN of the humeral head using autologous concentrated bone marrow grafts. At an average final follow-up of 49.4 months (range, 24-73 months), the average visual analog scale changed from 33 mm preoperatively to 5 mm postoperatively, and progression to further collapse or TSA did not occur. However, the patient with stage IV shoulder failed to improve following core decompression, requiring TSA prior to the final follow-up. Similarly, Hernigou et al. [68] reported the outcomes of 30 patients (30 shoulders) with adjunctive mesenchymal stem cell injection and 34 patients (34 shoulders) with no adjunctive injection who underwent core decompression for AVN of the humeral head. In the mesenchymal group, 18 patients were pre-collapse and 12 patients were post-collapse. At an average of 7-year follow-up (range, 5-10 years), 3 of 30 patients in the mesenchymal group progressed to humeral head collapse compared with 25 of 34 patients in the untreated group (P<0.0001). In addition, only 2 of 30 patients in the mesenchymal group progressed to shoulder arthroplasty compared with 25 of 34 in the untreated group (p < 0.0001).

Bone Allograft Technique

Galloway et al. [49] described bone allograft in conjunction with core decompression of the humeral head with use of a fibular allograft strut. Bone allograft is used to support the subchondral bone and decrease the risk of articular collapse following AVN lesion decompression rather than bony revascularization [49]. Following a standard arthroscopic-assisted percutaneous core decompression, a bone allograft is prepared with the same diameter of the drilled decompression tract. Introduction of the bone allograft is conducted over the previously placed guide pin and the pin is removed once the graft is secured. The remaining graft is then cut flush with the outer cortex with all edges smoothed using a burr. Galloway et al. [49] recommend under-contouring the graft progressing from proximal to distal to aid with interference fit of the graft within the tract.

Outcomes of Bone Allograft Technique

In a study including 14 pre-collapse shoulders (stage II) and 6 post-collapse shoulders (2 stage III and 4 stage IV), Galloway et al. [49] reported good outcomes for core decompression with adjunctive fibular strut allograft. Prior to the final follow-up, 3 shoulders, including 2 stage II and 1 stage IV, underwent shoulder arthroplasty due to pain. In the pre-collapse group, Shoulder Disability and Shoulder Pain and Disability Index (SPADI) scores were excellent in two shoulders, good in four shoulders, and poor in one shoulder. In contrast, SPADI scores in the post-collapse group included three excellent and one poor result. Limited improvement was observed in range of motion for both groups postoperatively. The better outcome scores in the post-collapse group were attributed to the younger patient age compared with pre-collapse group.

Postoperative Protocol

In the literature searched, formal consensus on postoperative restrictions following core decompression was not found. In a few articles, a short period of several days of sling immobilization with early progressive range of motion and limited heavy lifting for 8-12 weeks was recommended [11-13]. Conversely, Steffensmeier et al. [47] recommended no formal postoperative restrictions aside from tolerated activity and Makihara et al. [48] only recommended limiting carrying loads following core decompression. Although postoperative protocols differ, formal evidence indicating the gold standard for postoperative rehabilitation does not exist. However, the concern for postoperative stiffness and functional limitations should be considered with any postoperative protocol. Thus, the authors recommend patients begin a postoperative protocol with early range of motion, including shoulder pendulums, elbow, wrist, and hand exercises, to reduce stiffness. However, restrictions should be guided based on concomitant procedures. If the operation is an isolated core decompression, the authors recommend no formal restrictions and patients allowed to return to their normal activity level as tolerated.

CONCLUSIONS

Core decompression of the humeral head is an excellent treatment modality for shoulder preservation in patients with early-stage AVN of the humeral head without articular collapse. Several techniques have been developed, with good to excellent outcomes for each modality. In theory, a percutaneous, arthroscopic-assisted technique may allow accurate staging and concomitant treatment of intraarticular pathology during surgery, although further long-term clinical studies are necessary to assess the overall outcomes compared with standard techniques. In addition, although use of adjunctive biologics has been well described for core decompression of the hip, further high-quality studies are required in which their benefit in core decompression of the humeral head is further assessed.

NOTES

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Author contributions

Conceptualization: MDS, SA, DS, NG. Visualization: MDS. Writing – original draft: MDS, SA, DS, NG. Writing – review & editing: MDS, SA, DS, NG.

Conflict of interest

Dane Salazar reports personal fees from Tornier Wright Medical, Personal fees from Zimmer Biomet, outside the submitted work. Nickolas Garbis reports personal fees from DJO and Arthrex, outside the submitted work. No other potential conflicts of interest relevant to this article were reported.

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