COMPLEX SHOULDER INSTABILITY/ DISLOCATION
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SUMMARY
Traumatic shoulder instability can be complicated by the presence of cartilage injury, glenoid and/or humeral bone defects, rotator cuff injuries and nerve lesions. A high index of suspicion is required in the diagnosis of complex shoulder instability. Patients presenting with continued pain and dysfunction two to three weeks after the initial event should be investigated further. Older patients have a higher risk of associated injuries, but presentation in younger patients is frequent and may lead to devastating outcomes when missed. Correct recognition and treatment of the concomitant injuries is imperative in order to adequately stabilize the glenohumeral joint and avoid long-term dysfunction and degenerative changes. Shoulder instability can also be complicated by prior failed stabilization procedures. Failures are mostly caused by renewed traumatic events, misdiagnosis of the initial pathology or technical errors during the surgery. Type of previous surgical treatment and type of failure will influence the subsequent therapeutic strategy. Surgical history needs to be considered along with patient characteristics, anatomical lesions and functional demands. Clear guidelines in the setting of revision stabilization surgery are not available and treatment should be selected after a thorough case-by-case analysis.

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Introduction
The shoulder offers the greatest range of motion of any joint in the human body. Unfortunately, this generous mobility comes at the cost of an increased risk of instability. Static and dynamic mechanisms interact to stabilize the humeral head in the glenoid socket throughout the range of motion. Nevertheless, the application of sufficient external force can disrupt the stabilizers and result in traumatic subluxation or dislocation. An avulsion or tearing of the anterior capsulolabral complex is typically found in traumatic anterior instability. This, so-called “Bankart” lesion may require surgical repair to prevent further instability. Complex shoulder instability differs from this classical lesion due to the presence of cartilaginous or bony defects on the glenoid or humeral side or both.

Aim
This research report will address glenohumeral instability complicated by cartilaginous or bony defects, associated cuff and nerve lesions or previous stabilization surgery.

Cartilaginous Injury
Glenoid sided cartilaginous injury in the setting of shoulder instability most often presents as a gleno-labral-articular-disruption or GLAD lesion (Figure 1). In these cases, the labral injury is continuous with an articular cartilage component (Neviaser, 1993). Patients with this type of injury will report persistent pain and/or instability after shoulder dislocation. The GLAD lesion is best visualized via advanced imaging techniques such as arthro-CT or arthro-MRI. Arthroscopic evaluation and treatment by labroplasty and, if necessary, microfracture has shown good clinical results. However, only limited evidence is available on long-term outcomes and prevention of early degenerative disease (Page, 2008). Similarly, important chondral lesions can be seen on the humeral head, with variable location and size depending on the type of causative trauma (Figure 2). There are no guidelines for the treatment of central humeral-sided chondral lesions after shoulder instability. Treatment options include conservative, debridement, microfracture, autologous cartilage transplantation, osteochondral allograft or metallic resurfacing (DePalma and Gruson, 2012).

Figure 1. GLAD lesion visualized in a right shoulder in lateral decubitus as seen from the posterior portal. G=Glenoid, L=Labrum, GLAD=Gleno-Labral Articular Disruption.
Associated Bone Lesions

Glenoid bone lesions or “bony Bankart lesions”, are present in up to 90% of recurrent dislocations (Taylor et al., 1997). Humeral bone defects or “Hill-Sachs lesions”, are seen in up to 100% of recurrent dislocations (Wijdaja et al., 2006). The literature regarding bipolar lesions, with bone loss on the glenoid and humeral side, is limited. Reported incidences vary between 67% and 87% (Wijdaja et al., 2006; Griffith et al., 2008). Bone loss disrupts the glenohumeral equilibrium between mobility and stability by altering joint’s congruency and constraint. A diminished resistance against translational forces and a reduced coverage arc lead to increased soft-tissue stresses, resulting in recurrent instability. Large-scale studies have pointed to the importance of addressing bony lesions in the prevention of chronic instability and the onset of osteo-arthritis (Hovelius et al., 2009).

Glenoid Bone Lesions

Anterior glenoid defects are subdivided in ‘fracture’ or ‘erosion’ type depending on the presence of a bony rim fragment. However, since fragments can resorb within the first year, a distinction between both types isn’t always possible. It has been suggested that large fragment defects are caused by high-energy impactions, while smaller bone ‘chip’ avulsions arise from forced rotational trauma (Burkhart and De Beer, 2000). Glenoid defects are typically found at the 3 o’clock position and oriented parallel to the superior-inferior axis of the glenoid (Figure 3A). Three dimensional computed tomography (3D-CT) reconstruction has become the current imaging technique of choice. In en “en face” view, the defect is measured by comparing the actual glenoid width at the inferior glenoid circle, perpendicular to the supero-inferior axis, to the expected or contralateral inferior glenoid circle diameter (Baudi et al., 2013). Traditionally, 20–25% bone loss has been the biomechanically relevant threshold (Itoi et al., 2001), yet more recent publications have shown the clinical relevance of smaller amounts of glenoid bone loss (Shaha et al., 2015).

Humeral Bone Lesions

Impaction fractures of the posterolateral humeral head caused by impression of the soft cancellous bone on the dense anterior cortical genoid rim are referred to as Hill-Sachs lesions (Figure 3B). It is as yet unclear if these lesions arise during the dislocative trauma or as a secondary injury resulting from muscle pull. A recent study has shown that humeral bone loss is more extensive when time until reduction exceeds 5 hours,
supporting the latter theory (Denard et al., 2015). Hill-Sachs bone lesions can be described in detail according to their location, depth, orientation, size and volume. Lesions larger than 4 cm x 0.5 cm, with horizontal positioning, covering 20–25% of articular surface or amounting to 250 mm³ are associated with a higher rate of recurrence (Itoi et al., 2013). This clinical data is support by biomechanical studies showing higher rates of instability with increased humeral bone loss (Kaar et al., 2010). However, recent research has focussed more on the dynamic role of the humeral lesion interacting with the glenoid rim and causing further instability, and less on the static properties of the bony defect.

**Bipolar Bone Lesions**

Burkhart and De Beer first reported on the distinction between engaging and non-engaging humeral lesions (Burkart and De Beer, 2000). The authors found that Hill-Sachs lesions that ran parallel to the glenoid rim in abduction-external rotation presented a high risk of “engaging” the glenoid and resulting in instability. The authors advocated peroperative testing of joint stability after Bankart repair to ascertain if the Hill-Sachs lesion would engage. However, there are several difficulties pertinent to this strategy. First, such a manoeuvre seriously risks injuring the freshly repaired labrum and capsule. Second, peroperative testing in lateral decubitus and under traction may not be possible. Third, if enough energy is applied, all types of instability can be reproduced and rendered “engaging”. Fourth, a post-hoc peroperative diagnosis may be hard to incorporate in the surgical treatment plan. Therefore, an alternative approach was presented by Yamamoto et al. (2007). The authors mapped the path of the glenoid articular surface on the humeral head during abduction and external rotation. They proposed that if the Hill-Sachs lesion was contained within the contact zone of the humeral head with the glenoid, which averages 85% of the glenoid width as measured from the cuff footprint, the humeral head would offer enough bony support to stabilize the glenohumeral joint. This is a so-called “on-track” lesion (= non-engaging). If on the other hand, the lesion is wider or more medial than the glenoid track, the Hill-Sachs lesion will “engage” the glenoid rim and cause instability (“off-track” lesion). Moreover, because of concomitant anterior or glenoid bone loss, the glenoid track can be diminished and a previously stable on-track Hill-Sachs lesion may become unstable (off-track) in the presence of a glenoid rim defect (Di Giacomo et al., 2016).

### Material, methods and results

#### TREATMENT

**Glenoid**

Failure of soft-tissue procedures in the presence of significant gleno-humeral bone loss is well documented (Burkart and De Beer, 2000; Boileau et al., 2006). Historically, glenoid bone block procedures were introduced in order to “block” anterior translation of the humeral head over the glenoid rim. Eden and Hybinette described placing autologous tibial or iliac crest bone grafts as a mechanical barrier on the anterior glenoid rim. These procedures fell out of favour due to high incidences of postoperative degenerative arthritis. Although it is impossible to verify the historic data, it is generally believed that these degenerative changes were either the result of the chronic recurrent instability before the stabilization procedure or due to an excessive lateral graft positioning leading to graft-humeral head hyperpressure, posterior subluxation and rapid chondrolysis. An alternative technique, described by Latarjet, involved transplantation of the coracoid graft to the anterior glenoid neck. This variation theoretically improved stability by the additional stabilizing effect of the conjoint sling and the coraco-acromial ligament anterior reinforcement. Long-term clinical results have shown a dramatic reduction in recurrences after Latarjet-type
procedures compared to the standard capsulolabral repairs (Young et al., 1998). Conversely, complications rates have significantly increased after the popularization of the coracoid transfer procedures. Such complications are mostly benign and treatable, yet can be devastating in cases of hardware or graft malpositioning and neurovascular injuries (Griesser et al., 2013). As an alternative, less invasive and hardware-free techniques have been devised, such as an arthroscopic version of the Latarjet procedure with good clinical outcomes (Lafosse and Boyle, 2010). More recently, further improvements have been suggested by replacing the screws with suture-based fixation methods in order to minimize hardware related issues (Boileau et al., 2016). Nevertheless, the complexity of the procedure and the steep learning curve have prohibited widespread acceptance of this technique. Resch et al. published promising results of the J-bone procedure. This procedure relies on the impaction grafting of a J-shaped iliac crest autograft after creation an anterior glenoid wedge osteotomy. The advantages are the absence of hardware and the option of performing the procedure arthroscopically (Anderl et al., 2016). North-American authors have published extensively on the use of fresh donor osteochondral allografts. These grafts offer the advantage of a cartilage surface, matching radii of curvature and absence of donor-site comorbidity. Availability, cost and risk of disease transmission have to be considered when using fresh frozen allografts (Provencher et al., 2009).

**Humeral Head**

Symptomatic humeral defects can be treated by bony augmentation of the humeral head using autograft, allograft or metallic implants (Armitage et al., 2010). The evidence concerning these procedures is scarce, yet biomechanical data has indicated that significant restoration of stability is feasible using bony humeral reconstruction (Giles et al., 2012). Alternatively, bone can be added to the anterior glenoid with the aim of widening the glenoid track, and thereby reducing the risk of “engagement”. Although biomechanically feasible, bone grafting which exceeds the native glenoid surface area has shown high rates of resorption in clinical series (Di Giacomo et al., 2011). Therefore, we believe that adding an anterior glenoid bone graft in the absence of glenoid bone loss in order to stabilize a large Hill-Sachs lesion is relatively contra-indicated. An alternative is the so-called “remplissage” procedure. As the name suggests (from the French “to fill”), the aim of the procedure is to fill the defect with rotator cuff tissue. This converts the lesion to an extra-articular lesion and acts as a checkrein for anterior translation. Clinical studies have shown good outcomes using this technique in small to moderate sized Hill-Sachs lesions, yet the risk of loss of range of motion is present (Armitage et al., 2010). Humeroplasty, rotational humeral osteotomy and total shoulder replacement have been described for the treatment of large Hill-Sachs lesions but are rarely indicated.

**DECISION MAKING**

The therapeutic decision tree for shoulder instability in the presence of glenoid and humeral bone loss is multifactorial. Age, sex, sportive ambitions, hyperlaxity, and anatomical lesions need to be considered. Glenoid bone defects of 20–25% are traditionally considered the threshold for reconstructive anterior glenoid bone grafting procedures. However, the advent of the glenoid track concept has extended the applicability of glenoid bone augmentation procedures to cases with less pronounced glenoid bone loss (10–20%) when combined with an off-track Hill-Sachs lesions and in selected cases with an increased risk of recurrence such as young age, male sex, failed previous soft-tissue procedures, competitive athletic activity, and hyperlaxity. The correct indication for isolated humeral, isolated glenoid or combined surgical procedures is unknown.
Similarly, the role of anatomic reconstructions such as the Eden-Hybinette, J-bone and allograft reconstructions in relation to the classic non-anatomical Latarjet procedures has not yet been defined. Traditionally soft-tissue capsulo-labral repair should be reserved for low-risk patients with minimal bone loss, while non-operative treatment should be considered in the sedentary population.

ASSOCIATED INJURIES
Cuff Tears
The rotator cuff plays an important role in stabilizing the shoulder joint at both the mid-range of motion, through the concavity-compression mechanism, and at the end-range of motion by muscle action limiting the range and thereby protecting capsulo-labral structures. Rotator cuff structures may be injured due to dislocation, and dislocation may be facilitated by the presence of rotator injuries (Figure 4). Rotator cuff tears are present in 7–32% of dislocations and the incidence increases with age (Berbig et al., 1999; Simank et al., 2006). This is believed to be due to pre-existent thinning and weakness of the rotator cuff in the older population and repetitive micro-injury in the throwing athlete. The surgeon should be suspicious of such injuries in cases with persistent pain and dysfunction 2 to 3 weeks after dislocation. The clinical findings are often attributed to nerve palsy, but may result from underlying cuff pathology. Not infrequently, both arise together, constituting a “terrible triad” of the shoulder (Pevny et al., 1998; Porcellini, 2012). Further investigation by advanced imaging techniques should be ordered in these circumstances to define the type and extent of rotator cuff injury.

Treatment
The presence of a rotator cuff lesion should be considered in conjunction with the traditional factors of decision making in shoulder instability such as enumerated above; patient age, functional demands, recurrence, hyperlaxity and bony defects. Literature has amply shown the importance of surgically addressing the rotator cuff lesions. Lahteenmaki et al. (2006) showed better outcomes after early surgical repair compared to conservative or delayed surgical treatment. A combination of anterior capsulolabral repair and rotator cuff repair is indicated in patients younger than 40–60 years (Hawkins et al., 1999; Porcellini et al., 2012), whereas in older patients, depending on functional demands, solitary cuff repair may be sufficient to stabilize the shoulder. The habitual postoperative capsulitis and resulting stiffness in older patients after arthroscopic or open shoulder surgery, has a protective effect in the cases of instability, and specific anterior capsulolabral repair is rarely required in this population.

NERVE INJURY
Neurologic structures can be injured as a result of anterior shoulder instability. De Laat et al. (1993) recorded an overall 13.8% incidence of EMG abnormalities in patients with persistent pain and loss of function after primary shoulder dislocation. The axillary nerve was most frequently injured, followed by the suprascapular nerve and radial nerve in a mixed population of instability and proximal humeral fractures. Isolated axillary nerve injuries frequently result from closed traction mechanisms. The nerve is tethered proximally by the infraclavicular plexus and distally by the terminal deltoid branches and stretched over the humeral head during glenohumeral dislocation. Patients older than 65 and patients presenting with an associated cuff lesion of greater tuberosity fracture have a higher risk of incurring a concomitant nerve lesion.

Treatment
Spontaneous recovery of axillary is expected in 85–100% of cases after closed shoulder trauma (Gumina et al., 1997; Atef et al., 2016). Patients are generally observed clinically and neurophysiologically for 3–6 months. If signs
of recovery are not seen during this period, surgical intervention is advised before loss of motor end-plate function develops. Early distinction between neurapractic-axonotmetic and neurotmetic lesions is useful in predicting which patients will benefit from surgical intervention. Spontaneous recovery is expected in the former, but not the latter group. A new approach to nerve lesion diagnostics, combining electrodiagnostics and ultrasound imaging, termed “3D assessment”, is currently under development. The combined exam increases the sensitivity and specificity of early detection for lesions requiring operative treatment (Galvin et al., 2016; Terzis et al., 2010). If a lesion is found within a continuous nerve, serial clinical examinations and neurophysiologic testing can be ordered every 6–8 weeks until 3–6 months after injury. If a discontinuity or structural cause of nerve injury is seen, surgical exploration and intraoperative assessment is indicated. Surgical outcomes or surgical interventions have been shown to decline when lead-time exceeds 12–18 months from the injury. In these cases, procedures which do not rely on functional deltoid muscle motor end-plates such as tendon transfers, and salvage procedures such as glenohumeral arthrodesis are indicated.

**REVISION SURGERY IN ANTERIOR SHOULDER INSTABILITY**

Trauma, misdiagnosis, and technical errors are the major causes of surgical failure regardless of the index procedure. Overall, the best results are obtained in revision cases after renewed high-energy trauma, after only one prior attempt at stabilization, and in the absence of a voluntary component to the instability (Boone et al., 2010). Fundamental to successful revision surgery is selecting the correct procedure. Typically, the surgeon’s choice will be either a soft-tissue based repair or a bony procedure.

**REVISION SOFT TISSUE REPAIR**

Although some still consider open Bankart repair to be the gold standard procedure in cases of recurrence, several studies support the equivalence of arthroscopic revision for the appropriate indications (Arce et al.,...
However, revision capsulolabral repair should only be used in patients with minimal risk factors for recurrence, and must not be used in the presence of significant glenoid or humeral bone loss. Essential to the repair is the recreation of an anteroinferior labral bumper. Attachment of the plicated capsule to the glenoid face may serve to replicate a labral bumper in cases of labral deficiency. A rotator interval closure can be considered in cases of persistent laxity after these traditional repairs (Boileau et al., 2009).

BONY PROCEDURES

The coracoid transfer procedures possess numerous advantages in a revision setting and will be the only option left in many cases. The stabilizing effect of the transfer is multifactorial and obviates the need for a robust labral or capsular repair that is often difficult to achieve with a revision Bankart repair. The grafted coracoid process restores the glenoid articular surface and normalizes the glenohumeral contact pressure (Ghodadra et al., 2010). Hill-Sachs engagement is avoided by widening of the glenoid track and the transferred conjoint tendon provides dynamic stability in the main position of apprehension (Wellmann et al., 2011). The postoperative rehabilitation can be accelerated with no limitations on external rotation due to the rigid fixation of the bone graft to the glenoid neck.

In revision cases of failed coracoid transfer procedures, glenoid bone loss may need to be addressed by structural bone grafting, such as in the Eden-Hybinette technique of autogenous tricortical iliac crest bone grafting (Lunn et al. 2008). The use of distal tibial allografts or even glenoid allografts have also been suggested because of the anatomic similarity to the glenoid articular surface, osteochondral nature of the graft, and capacity for secure fixation and incorporation (Provencher et al. 2009).

Most Hill-Sachs lesions can be rendered inconsequential with an adequate anterior repair and capsular plication. In large lesions that threaten stability, engagement may be eliminated with restoration of the articular arc through coracoid transfer or bone grafting as mentioned earlier. Some investigators have recommended filling the defect with an osteochondral allograft to correct the humeral head deformity (Chapovsky et al., 2001). The defect may also be filled with the posterior capsule and the adjacent infraspinatus tendon (“Remplissage procedure”) to prevent engagement and limit anterior translation of the humeral head, yet the indications and long-term outcomes of these procedures are not well understood, and their role in revision surgery has not yet been delineated.

Discussion and conclusions

Specific attention is warranted in cases of traumatic anterior shoulder instability complicated by cartilage injury, glenoid and/or humeral bone defects, rotator cuff injuries, nerve lesions or previous stabilizing surgery. Patients with prolonged pain and dysfunction after an instability event require further investigation. Especially older patients are at risk of associated injuries, but presentation in younger patients is possible and should not be missed. Full clinical and imaging work-up is necessary before decision-making in these complex cases. Clear guidelines are lacking in the literature due to the relative scarcity of the problem. A careful analysis of patient risk factors, anatomical lesions and functional demands is mandatory before proceeding with any therapeutic plans.
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