BMJ Open

Biomechanical repositioning techniques in anterior shoulder dislocation: a randomised multicentre clinical trial—the BRASD-trial protocol

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ABSTRACT

Introduction Glenohumeral (shoulder) dislocations are the most common large joint dislocations seen in the emergency department (ED). They cause pain, often severe, and require timely interventions to minimise discomfort and tissue damage. Commonly used reposition or relocation techniques often involve traction and/or leverage. These techniques have high success rates but may be painful and time consuming. They may also cause complications. Recently, other techniques—the biomechanical reposition techniques (BRTs)—have become more popular since they may cause less pain, require less time and cause fewer complications. To our knowledge, no research exists comparing the various BRTs. Our objective is to establish which BRT or BRT combination is fastest, least painful and associated with the lowest complication rate for adult ED patients with anterior glenohumeral dislocations (AGDs).

Methods and analysis Adults presenting to the participating EDs with isolated AGDs, as determined by radiographs, will be randomised to one of three BRTs: Cunningham, modified Milch or scapular manipulation. Main study parameters/endpoints are ED length of stay and patients’ self-report of pain. Secondary study parameters/endpoints are procedure times, need for analgesic and/or sedative medications, iatrogenic complications and rates of successful reduction.

Ethics and dissemination Non-biomechanical AGD repositioning techniques based on traction and/or leverage are inherently painful and potentially harmful. We believe that the three BRTs used in this study are more physiological, more patient friendly, less likely to cause pain, more time efficient and less likely to produce complications. By comparing these three techniques, we hope to improve the care provided to adults with acute AGDs by reducing their ED length of stay and minimising pain and procedure-related complications. We also hope to define which of the three BRTs is quickest, most likely to be successful and least likely to require sedative or analgesic medications to achieve reduction.

Trial registration number NTR5839.
Reduction techniques

More than 50 glenohumeral dislocation repositioning techniques exist. They are described unambiguously in the literature but often inconsistently performed in clinical practice. The wide range of techniques can be divided into three groups based on their major mode of action: traction, leverage or biomechanical. The most commonly used techniques in Dutch EDs are the traction-based Hippocratic method and the leverage-based Kocher method.

Traction-based techniques

Traction-based techniques—such as the Hippocratic method and its variants—rely on force to overcome muscle spasm. The idea being that, by applying traction, muscles will tire and relocation will occur. The amount of traction the operator can apply can be increased by means of countertraction. Many modifications of the pure Hippocratic method exist, some already suggested by Hippocrates himself, including the application of countertraction with a sheet, the operator’s shoulder, the operator’s knee, the patient’s bodyweight (Eskimo technique), a bed, a chair and a ladder. Since applying traction will increase muscle spasm and pain, traction techniques often require analgesia and/or sedation, resulting in prolonged ED lengths of stay (LOSs). Traction–countertraction techniques may result in neurovascular damage in the axillary region, although incidence is unknown.

Leverage-based techniques

Kocher’s method, originally described in 1870, is the best-known leverage technique for AGD reduction. The technique has been altered by clinicians since, and often includes traction, which is commonly associated with increased pain. This combined technique achieves good results, but some force is still needed to manipulate the humeral head over the glenoid. Additionally, iatrogenically induced humeral fractures and axillary vessel ruptures are seen with the technique and were, in fact, described by Kocher in his original article. Two other studies describe the risk of postreduction humeral neck fractures during leverage techniques in patients over 40 years of age.

Biomechanical techniques

More recently, several techniques with a biomechanical basis have been described. These biomechanical repositioning techniques (BRTs) depend on muscular relaxation without force and often start with the patient’s arm in an analgesic position, thus eliminating or minimising the need for sedatives or analgesics. They do require patient cooperation, making it essential that patients receive accurate instruction about the procedure. BRTs can be separated into three approaches: positioning and relaxation, zero position and scapular manipulation.

The Cunningham technique involves positioning and relaxation. The patient’s arm must be fully adducted for the technique to succeed. This reduces spasm in the stretched rotator cuff muscles. By massaging the trapezius, deltoïd and especially the biceps brachii muscles, tension in the ‘bowstringed’ biceps brachii will decrease and relocation will occur. No traction is applied.

The original Milch technique first described in 1938 and the modified Milch technique described in 1992 involve positioning the patient’s arm such that all the muscles acting on the shoulder joint align with the humerus (the so-called ‘zero position’). No traction is applied. The scapular manipulation technique (SMT) was developed in the late 1970s and published in 1982. As the name implies, SMT involves scapular movement with the patient prone so that the glenoid fossa re-engages the humeral head, achieving reduction. In a sense, arm traction is involved as well, but only to stabilise the humeral head, not to fatigue muscles. Patient pain is thereby limited. The classically described SMT is often modified to a sitting or supine position.

Pain relief

Many methods exist to address the pain associated with AGD, ranging from intra-articular anaesthesia to nitrous oxide, nerve blocks and various procedural sedation and analgesia regimens. Not one method is clearly superior in every regard, and all involve time to gather medications, consent the patient, administer (and possibly re-administer) medications, wait for effects and observe the patient postprocedure as he/she recovers. In addition to these delays and the possibility of inadequate pain relief, there is the real risk of complications associated with procedural sedation: nausea and vomiting, hypotension, hypoxaemia, prolonged drowsiness, headache, aspiration, respiratory depression and untoward medication reactions, among others. Many authors have advocated that the best relief for AGD pain is reduction.

Conclusion

A variety of traction-based or leverage-based techniques are often successful in repositioning AGDs, with success rates ranging from 60% to 100% in generally small studies. However, since pain is increased by traction, countertraction and leverage, these techniques often require the administration of analgesics and sedations, which may be associated with complications. Additionally, the techniques themselves may not be quick, painless or complication free and do not pay heed to patient satisfaction or ED throughput. Consequently, total ED time can be 3 hours or more for a procedure with a performance time of less than 10 min.

In contrast, BRTs do meet the requirements for optimal repositioning. The ideal method should be simple, easy, quick, effective, atraumatic and pain
free; require little assistance or medication; and cause no additional injury to the shoulder joint or to the musculoskeletal or neurovascular structures.’

Data on the BRTs are scarce, but the reported minimal inflicted pain, high success rates and the avoidance or reduced need for sedation or analgesia seem promising for a shorter ED stay, lower resource utilisation and a better patient experience. However, which BRT or BRT combination is fastest, least painful and least likely to cause complications is unaddressed in the current medical literature.

Methods and analysis

Primary research question
Which BRT or BRT combination is fastest and least painful for adult ED patients with AGDs?

Secondary research questions
- Are complications caused by BRTs or BRT combinations? If so, what are those complications?
- What are the reposition success rates of the BRTs or BRT combinations?
- What are the ED LOSs associated with the BRTs or BRT combinations?

Study design
A randomised controlled trial (RCT) will be conducted in two Dutch hospital-based EDs comparing the three BRTs: modified Milch, Cunningham and SMT.

To optimise technique execution from study outset, we intend to enrol 222 patients. We calculated the sample size on ED LOS per combination of techniques as shown in figure 1 flowchart. We assumed a probability of type 1 (alpha) error of 0.05 and a type 2 (beta) error probability of 0.20. One hundred and eighty-five inclusions are therefore needed in total. Based on other studies done at one of our hospitals, we are anticipating a 20% data loss, so we intend to enrol 222 patients.

Investigational treatment
Patients able to adduct (‘can adduct’ path) will be randomised to BRT using Cunningham, modified Milch or SMT. Those unable to adduct (‘cannot adduct’ path) will be randomised to BRT using either modified Milch or SMT (see figure 1 flowchart).

AGD reduction will be defined as the re-establishment of a normal glenohumeral relationship on postintervention radiographs. After reduction, an internal rotation sling will be applied and follow-up arranged in the outpatient clinic.

Data collection
Baseline demographics, medical history and study-specific data will be collected. ED LOS will be defined as the time in minutes from patient arrival in the ED until discharge. The well-validated Numeric Rating Scale of 0–10 will be used to assess patients’ pain, before, during and after reduction attempts.

Other data to be collected:
- reduction time (in minutes, from start to end of procedure)
- number of reduction techniques used
- sedatives and analgesics used (types, dosages, prehospital and/or in-hospital administration)
- preintervention and postintervention radiograph interpretation
- physical examination (with particular attention to neurovascular status of the affected arm)
- iatrogenic complications (caused by the interventions)
- patient age
- patient gender
- time of last oral intake
- dislocation number (first or recurrence number)
- dislocation mechanism (sports, seizures, falls, traffic accidents, other).

Statistical methodology
We calculated the sample size on ED LOS per combination of techniques as shown in figure 1. A 15-minute difference between the combinations of techniques is considered clinically relevant. We assumed a probability of type 1 (alpha) error of 0.05 and a type 2 (beta) error probability of 0.20.

In the can adduct group, we will compare two combinations of techniques. Assuming non-normality and using the Mann-Whitney U test, power calculations lead to a sample size per combination of 31, with a total of 62 inclusions.

In the cannot adduct group, we will compare three combinations of techniques. Similar to the calculation of the cannot adduct group, assuming non-normality and using the Kruskal-Wallis test, power calculations lead to a sample size per combination of 41, with a total of 123 participants required.

One hundred and eighty-five inclusions are therefore needed in total. Based on other studies done at one of our hospitals, we are anticipating a 20% data loss, so we intend to enrol 222 patients.

Box 1 Inclusion/exclusion criteria

Inclusion
- All adult patients (≥18 years) with an isolated anterior glenohumeral dislocation of less than 24 hours and able to understand and sign consent

Exclusion criteria
A potential subject who meets any of the following criteria will be excluded from participation in this study:
- subcapital humeral fractures—major multi-trauma
- subclavicular, intrathoracic, inferior or posterior dislocations
- dislocations presenting after 24 hours.
Enrolment will continue until the required sample size in each arm is reached.

The techniques will be randomised in advance per centre by creating a stratified block randomisation list.

Nominal variables related to subgroups will be analysed with the chi-square test. Ordinal variables related to subgroups will be analysed using the Mann-Whitney U test or the Kruskal-Wallis test. Variables will be compared with each other (depending on the scaling level) with the Wilcoxon test, the Friedman test and the correlation coefficient of Spearman. Nominal and ordinal variables will be described using frequency tables, mode and median. A value of $p<0.05$ will be accepted as statistically significant.

In cases of missing data, the treating physician will make inquiries. If more than 30% of the data are still missing postinquiry, the patient will be excluded from the study. SPSS V.22 will be used for data processing.

**ETHICS AND DISSEMINATION**

This RCT will compare three BRTs described in the medical literature, the modified Milch, Cunningham and the SMT, and will be one of the first comparative studies on BRT outcomes. Its aim is to establish whether different BRTs produce different ED LOS and patient discomfort during and after reduction. This trial will add valuable information to the presently limited knowledge about these techniques. Results of the study will be made publicly available by submitting the results to a peer-reviewed medical journal. No veto or disclosures are made with the sponsors.

AGDs are painful and require timely intervention to relieve and/or minimise discomfort and potential tissue damage. Non-BRTs are based on traction or leverage and therefore inherently painful and potentially harmful. We posit that the BRTs used in our study

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**Figure 1** Flowchart showing the randomisation of the biomechanical reposition techniques in anterior shoulder dislocation trial. AP, anteroposterior view; SMT, scapular manipulation technique.
are more physiological, likely less pain producing and will lead to a decreased ED LOS while being just as successful as older techniques at repositioning the acute AGD. To date, no adverse events have been described for these techniques. Our study results may help define a more standardised, less risky and improved treatment regimen for patients with AGD by minimising pain and shortening ED throughput times. This may not only benefit individual patients but also healthcare systems.

LIMITATIONS
Since it is impossible to blind physicians and patients to the technique used for shoulder reduction, this may introduce a bias toward techniques more favoured by some physicians. We are also aware that practitioner learning will occur over the course of the study and individual physicians may gravitate toward or become increasingly adept at certain techniques.

We will attempt to minimise the bias introduced by the absence of blinding, learning effect and optimise technique execution by training the participating doctors, NPs and nurses before the study starts. After the study start, we plan to revisit the participating centres to train and answer questions about the techniques used. Also, visual and written instructions will be provided at the start of the study and learning material is also be available on our YouTube channel.

Videos
1. Cunningham: https://youtu.be/6TF3h3RN80M?si=10
2. Modified Milch: https://youtu.be/yOm1bF-U9Q8
3. SMT: https://youtu.be/Cig7XRH8cZs

Acknowledgements We thank Tjeerd van der Ploeg for statistical guidance and Sylvia van Rossum for her endless support.

Contributors DNB is the corresponding author. DNB and MHR were responsible for conception and study design and, as coordinators of project, coordination and study refinement/focus. DNB was responsible for manuscript writing, editing and study refinement/focus.MDB and patient recruitment. SP was responsible for guidance in protocol design. MDB for conception and study design and, as coordinators of project, coordination and study refinement/focus. DNB was responsible for manuscript writing, editing and study refinement/focus. DNB and MHR contributed equally to this work and are both first authors.

Competing interests None declared.

Ethics approval Approval for the BRASD-trial has been obtained from the ethics committee (MREC) Northern Holland. The BRASD-trial is conducted in accordance with the principles of Good Clinical Practice as defined by the International Council on Harmonisation (ICH-E6, 17/07/96) as well as with specific laws and regulations that apply in the Netherlands.

Provenance and peer review Not commissioned; externally peer reviewed.

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BMJ Open 2017 7:
doi: 10.1136/bmjopen-2016-013676

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