Investigating the efficacy of chest pressure for direct current cardioversion in atrial fibrillation: a randomised control trial protocol (Pressure-AF)

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ABSTRACT

Introduction Atrial fibrillation (AF) is the most common sustained arrhythmia worldwide. Direct current cardioversion is commonly used to restore sinus rhythm in patients with AF. Chest pressure may improve cardioversion success through decreasing transthoracic impedance and increasing cardiac energy delivery. We aim to assess the efficacy and safety of routine chest pressure with direct current cardioversion for AF.

Methods and analysis Multicentre, double blind (patient and outcome assessment), randomised clinical trial based in New South Wales, Australia. Patients will be randomised 1:1 to control and interventional arms. The control group will receive four sequential biphasic shocks of 150 J, 200 J, 360 J and 360 J with chest pressure on the last shock, until cardioversion success. The intervention group will receive the same shocks with chest pressure from the first defibrillation. Pads will be placed in an anteroposterior position. Success of cardioversion will be defined as sinus rhythm at 1 min after shock. The primary outcome will be total energy provided. Secondary outcomes will be success of first shock to achieve cardioversion, transthoracic impedance and sinus rhythm at post cardioversion ECG.

Ethics and dissemination Ethics approval has been confirmed at all participating sites via the Research Ethics Governance Information System. The trial has been registered on the Australia New Zealand Clinical Trials Registry (ACTRN12620001028998). De-identified patient level data will be available to reputable researchers who provide sound analysis proposals.

INTRODUCTION

Atrial fibrillation (AF) is the most common persistent arrhythmia worldwide and a significant contributor to poor quality of life.1,2 Currently, there are an estimated 37 million individuals with AF globally. Projections to the year 2050 indicate a 60% absolute increase in AF prevalence.3 Given the huge burden of disease, measures to prevent and optimally treat AF are of great importance.

Recently, there has been renewed interest in the benefit of rhythm control in those with AF.4 Direct current cardioversion is an efficacious procedure commonly used to terminate arrhythmias and plays an important role in rhythm control for AF.

Rhythm control for AF

Initial trials comparing rate and rhythm control for AF showed no difference in outcomes, and perhaps increased hospitalisations with rhythm control.5 6 Despite this, recent trials have shown benefit to rhythm control in select populations. In those with heart failure with reduced ejection fraction and persistent ventricular dysfunction despite medical therapy, mortality reduction is seen with catheter ablation.7 8 Moreover, in younger patients with early AF, there may be small improvements in cardiovascular endpoints with rhythm control.7 Clearly, a tailored approach to AF management is needed. Therapeutic strategies are dependent on patient preference, symptoms, the presence of structural heart disease and risks associated with interventions.

In those whom rhythm control is felt to be appropriate, electrical cardioversion offers a promising means of restoring sinus rhythm. The limitation of electrical cardioversion is that initial restoration of sinus rhythm does not ensure maintenance of sinus rhythm. Patients generally also require longer term antiarrhythmic therapy and lifestyle modification. Early success of cardioversion for AF was low, around 75%.9 Over the past decades, numerous techniques have been implemented to improve restoration of sinus rhythm. With these methods, sinus rhythm is achieved in over 90% of cases.10
Cardioversion technique to improve success

Chance of successful cardioversion is directly proportional to cardiac energy delivery. Increased energy delivery heightens the chance of simultaneous depolarisation of all cardiac tissue. There are four major clinician-controlled modifiers that can increase cardiac energy delivery (see Table 1). The first is the intensity of energy provided during cardioversion. While other arrhythmias require less energy, 50–100 J for atrial flutter, for example, the electrical disorganisation of AF requires higher energies. International guidelines recommend initiating treatment at 150–200 J. The second variable is defibrillator pad position. Anteroposterior pad placement seems to improve success compared with anterolateral positioning. The mechanism may be through increased atrial energy delivery. The third variable is the energy form delivered. Biphasic energy is more effective than monophasic energy and has been consistently shown to improve cardioversion success. The final modifier is the provision of chest pressure during cardioversion. Chest pressure appears efficacious through reducing transthoracic impedance which is inversely proportional to cardiac energy delivery.

Risks of electrical cardioversion

While increasing energy will improve cardioversion success, this is also associated with risk. Moderate to severe post-procedural chest pain is reported to occur in 23% of cases and is directly proportional to energy delivery. Cutaneous burns are another significant complication of cardioversion and often associated with high energy shocks. Measures to reduce energy requirements, while maintaining cardioversion efficacy, have the potential benefits of decreasing patient discomfort and reducing burn incidence.

Chest pressure with cardioversion

While chest pressure during cardioversion is reproducible, simple and associated with minimal cost, there is little evidence for its use. Most clinicians still regard close patient contact during cardioversion to be unsafe to the bystander. A systematic review investigating bystander adverse events during defibrillation showed that events were uncommon, always minor and often associated with inappropriate defibrillator use. Moreover, a recent single centre randomised trial involving only 100 patients demonstrated no bystander adverse events with chest pressure during cardioversion. This trial used starting energies of 50 J and found improved cardioversion success, reduced shocks and reduced energy requirements with active compression. There is a clear need for external validity in an adequately powered multicentre randomised trial to examine the utility of routine chest pressure.

Hypothesis and aims

In this trial, we hypothesise that upfront manual chest pressure will reduce energy required for cardioversion and be safe for both patient and proceduralist. Moreover, we hypothesise that upfront chest pressure will increase the rate of first shock success. We aim to assess the efficacy and safety of manual chest pressure in AF with direct current cardioversion.

METHODS AND ANALYSIS

Trial design

This will be a prospective multicentre randomised control trial located in Australia. Patients and statistical analysis will be blinded. Inclusion criteria are age over 18 years, AF referred for cardioversion and 3 weeks of therapeutic anticoagulation or transoesophageal echocardiography excluding left atrial appendage thrombus (see box 1). Exclusion criteria are other atrial arrhythmias (atrial flutter or atrial tachycardia), those who are pregnant or breast feeding and medical comorbidity where anticoagulation is contraindicated. Definition of successful cardioversion will be sinus rhythm at 1 min after shock to maintain consistency with other studies. Patients will be randomised via an online tool in a 1:1 ratio to control and interventional arms.

Cardioversion technique

Anteroposterior placement of self-adhesive pads with biphasic energy will be provided to all patients. Sedation will be performed by an anaesthetist with a recommended anaesthetic regime of weight-based dosing of propofolmidazolam titrated to effect. Direct chest pressure will be provided on the anterior defibrillator pad with plastic gloves and a folded towel (see figure 1). Estimated chest pressure provision was assessed among cardiology advanced trainees at the centres involved. Four trainees provided 36 applications of blinded simulated chest pressure on weight scales with mean pressure of 25.5±2.6 kg.

The control arm will receive four sequential shocks until cardioversion success with energies of 150, 200, 360 and 360 J. Chest pressure will be provided with the final shock if the arrhythmia is refractory to the first three defibrillations. The interventional arm will receive chest pressure from the first shock with the same energies. Both arms will receive maximal energy with chest pressure (figure 2).
Arrhythmias and sudden death

Primary and secondary endpoints
The primary efficacy endpoint will be total joules provided during the cardioversion encounter. Secondary efficacy endpoints will be success of first shock for reversion to sinus rhythm, transthoracic impedance at the time of shock and sinus rhythm at post cardioversion ECG (30 min after). Safety and tolerability endpoints will be patient chest pain post cardioversion (Ordinal Scale 0–10) and incidence of shock delivered to proceduralist.

Sample size and power calculation
Power calculation was performed based on a retrospective analysis of direct current cardioversions over the prior year. The mean energy provided per encounter for cardioversion in AF was 280±188 J. Assuming a mean reduction of 60 J in energy in the intervention group (approximately one-third of the SD), a total of 308 patients would be required for an alpha of 0.05 and power of 80%. An interim analysis will be performed at 12 months to ensure adequate recruitment (at least 30% of the target) and no safety signals. Trial recruitment has begun, and 11 patients have been enrolled.

Statistical analysis
Continuous variables will be reported as mean±SD or as median and percentiles. Normally distributed variables will be compared using the paired Student’s t-test. Otherwise, comparisons between both the groups will be performed using the Mann-Whitney U test. Categorical variables will be stated as absolute and relative frequencies and compared using the χ² test. All tests are two-tailed. A p value of <0.05 will be considered statistically significant. Pre-specified post hoc analyses will be performed based on body mass index, left atrial size, age, antiarrhythmic therapy and duration of AF (online supplemental file 1).

ETHICS AND DISSEMINATION
Ethics approval has been confirmed at all participating sites via the Research Ethics Governance Information System. The trial has been registered on the Australia New Zealand Clinical Trials Registry (ACTRN12620001028998). In the spirit of integrity and openness, de-identified patient level data will be available to reputable researchers who provide methodologically sound proposals for analysis.

DISCUSSION
Chest pressure is a simple intervention which may be effective in reducing cardioversion energy requirements and improving cardioversion success. A small randomised trial investigated chest pressure in cardioversion for AF and found reduced energy requirements and improved cardioversion success. This trial had a sample size of 100 patients and defibrillation energy started at 50 J. In our study, we aim to perform this multicentre study using guideline-directed energies (150 J) with adequately large sample size powered to definitively answer this important clinical question.

Consideration was made to quantify and standardise chest pressure provision during cardioversion in this trial. There were two main reasons why this was not pursued. First, one major benefit of chest pressure is its simplicity. External validity would be diminished if complex measures were taken to standardise chest pressure. Second, the degree of chest pressure is highly variable and depends on the body habitus of the patient. Having a ‘one size fits all’ approach to chest pressure provision is unlikely to be advantageous to the individual.

A theoretical concern is that of a shock being sustained by the proceduralist performing manual pressure. Despite this, a systematic review investigating bystander adverse events during defibrillation showed that events were uncommon and minor. At our centres, under controlled circumstances with trained staff, chest pressure has not been associated with adverse events to the proceduralist. This trial will provide quality data around the safety of manual chest pressure during defibrillation.

Box 1 Eligibility criteria

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<th>Inclusion criteria</th>
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<td>► Age over 18 years of age.</td>
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<td>► Ability to provide written informed consent.</td>
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<td>► Atrial fibrillation referred for cardioversion.</td>
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<td>► Three weeks of therapeutic anticoagulation.</td>
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<td>► Transesophageal echocardiography excluding left atrial appendage thrombus.</td>
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<th>Exclusion criteria</th>
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<tr>
<td>► Other atrial arrhythmias (atrial tachycardia, atrial flutter).</td>
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<td>► Pregnant or breast feeding.</td>
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<td>► Medical comorbidity where anticoagulation is contraindicated.</td>
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Figure 1 Diagram of cardioversion technique.
Improving success of direct current cardioversion in a reproducible, cost effective and safe manner has significant implications. First, it may assist in improving symptoms in patients who suffer with symptomatic arrhythmias. Second, we may be able to provide an energy threshold by which a large proportion of patients will be successfully cardioverted with a single shock and chest pressure. Third, there are indirect implications for defibrillation during advanced life support. Manual compression when providing defibrillation to refractory shockable rhythms may improve success of resuscitation.23

In conclusion, Pressure-AF is an important clinical trial evaluating the simple intervention of chest pressure with potential to improve cardioversion success for AF.

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**Data availability statement** Data are available upon reasonable request. De-identified patient level data will be available to reputable researchers who provide sound analysis proposals.

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**REFERENCES**


**Figure 2** Day of cardioversion: study flow diagram.
Arrhythmias and sudden death


