Defibrillation has been used to treat disturbances of the cardiac rhythm since the late 1800s. In order to appreciate the mechanisms of defibrillation, one must also understand the basic electrophysiological principles of fibrillation.

**ELECTROPHYSIOLOGICAL BASIS OF FIBRILLATION**

Atrial and ventricular fibrillation are both based upon a re-entrant mechanism, a phenomenon that was first recognised almost 100 years ago and initially described by Mines and Garrey. Mines also suggested that conduction block was necessary to facilitate re-entry. It was not until the 1950s that Moe and colleagues advanced the circus movement theory of re-entry when they suggested the multiple wavelet hypothesis of atrial fibrillation (AF). This proposed that AF occurred in the presence of inhomogeneous atrial tissue. It was postulated that multiple wavelets propagated randomly through the atria. These waves of excitation were presumed capable of spreading through the atrial myocardium at a time when some of its components had recovered while others remained partially or fully refractory as a result of a preceding activation. Thus AF could be maintained as a turbulent arrhythmia in a stable state for long periods of time. By 1985, after the development of high resolution electrode mapping systems, Allessie and colleagues provided evidence of multiple propagating wavelets that created turbulent atrial activity in the canine heart. Subsequent experiments have proven Moe’s idea that multiple wavelets distributed randomly throughout the atria gave rise to the chaotic activation patterns observed on the ECG.

More recent studies have further expanded on some of the experimental findings of Allessie and colleagues, suggesting that some forms of AF may actually be the result of high frequency activation by a single re-entrant source in some patients. The drivers of AF may be relatively stationary “rotors” of electrical activity that are anchored to an unexcited (anatomical) core in the posterior left atrium. The atrial anatomy may facilitate this re-entry process by providing areas of conduction block (for example, pulmonary vein ostia). It is also increasingly recognised that ectopic foci originating from the pulmonary veins may be responsible for initiation of AF in patients, and that electrical isolation of these foci may provide long term correction of AF in some patients. Thus, AF is based upon re-entry. Multiple wavelets of electrical activity propagate randomly in the atrial tissue producing a chaotic but continuous activation of the atrial tissue. Defibrillation (or direct current cardioversion) in this setting aims to bring an abrupt halt to this process and restore sinus rhythm.

**MECHANISMS OF ATRIAL DEFIBRILLATION**

There are only limited data that examine the interaction between an electrical shock and atrial tissue. Research into the mechanisms of defibrillation has occurred almost exclusively in the setting of ventricular fibrillation. While there may be distinct differences between the precise mechanisms of atrial and ventricular defibrillation, none has been described. There are several theories of the mechanisms of defibrillation that are derived by ventricular defibrillation.

It is thought that shocks defibrillate by altering the potential difference across the cell membrane (that is, the transmembrane potential). However, the situation in the heart is extremely complex with the intracellular space and extracellular space both possessing different electrical properties than the cell membrane that divides them. Unfortunately, there is no current consensus on how the extracellular potential gradient relates to the transmembrane potential gradient. Thus, the exact mechanism of electrical defibrillation remains incompletely understood.

The critical mass theory proposes that defibrillation can be successfully achieved by the depolarisation of a sufficient or “critical” mass of tissue. Zipes and colleagues demonstrated that chemical depolarisation of a critical mass of ventricular myocardium (~75%), using selective infusion of potassium chloride into the left coronary artery, would successfully abolish ventricular...
Therefore, if a sufficient voltage potential gradient could be generated by a shock to halt the fibrillation activation fronts in the myocardium, fibrillation could be abolished.

When an electrical field with strength of 5 V/cm or less is introduced to ventricular myocardium, it exhibits an all or nothing response. Therefore, recovered myocardium produces a new action potential, while the stimulus has no effect on even partially refractory cells. Above 5 V/cm a graded response can be produced in refractory cells. The result of this graded response is a prolongation of both refractoriness and the action potential, although of shorter duration than the production of a new action potential. These observations form the basis of the “extension of refractoriness hypothesis”. This hypothesis states “the defibrillating shock interacts primarily with cells during their refractory period and prolongs refractoriness in most of the myocardium so that fibrillation wavefronts cannot propagate and fibrillation ceases”.

A lower limit of vulnerability also exists for ventricular myocardium. This is the minimum voltage required by an electrical stimulus to induce fibrillation during the vulnerable period. It was noted in the 1960s that there was an upper limit to the strengths of shocks delivered during the vulnerable period that induce ventricular fibrillation. It was also observed that the strengths of these shocks at “the upper limit of vulnerability” were approximately equivalent to the shocks at defibrillation threshold. The upper limit of vulnerability hypothesis for defibrillation states “to defibrillate, a shock not only must halt the activation fronts of fibrillation, but it also must not reinitiate fibrillation by the same mechanism that a shock of the same strength during the vulnerable period of sinus or paced rhythm initiates fibrillation”.

These theories are supported by the Fundamental Law of Electrostimulation, which was adapted by Irnich and applied to ventricular defibrillation. If the concept of nearfield and farfield rheobase (lowest current strength which successfully stimulates myocardium) are introduced, then attempted defibrillation with shocks of too low voltage may be effective close to the electrode (nearfield) but fail to stimulate cardiac tissue at a remote site (farfield). In order to gain a more accurate picture of the cellular response to electrical stimuli, optical mapping techniques were developed. This involves the use of potentiometric fluorescent dyes that bind to cell membranes in proportion to the polarisation of the membrane caused by a shock. Using this technique, Kwaku and Dillon reported that activation fronts in the ventricles arose from the border of shock-depolarised areas and that these fronts propagated unidirectionally away from these regions. They also found that increasing refractoriness decreased the likelihood of a reactivation wavefront, and concluded that for successful defibrillation a shock must depolarise myocardial tissue, even during its refractory period.

Transmembrane potentials have been recorded from the atria during AF and after biphasic shocks. The shocks have been shown to produce four types of responses: immediate cessation of epicardial activity, single post-shock activation, organised activation for 0.8–1.5 seconds followed by termination, and organised activity followed by degeneration back into AF. These experiments led the authors to conclude, “These results are consistent with both the critical mass hypothesis and the upper limit of vulnerability hypothesis”. Therefore, given the similarity between atrial and ventricular myocytes, it seems reasonable to assume that the effects of an electric shock are similar in both chambers.

DEFIBRILLATION WAVEFORMS

Early devices employed both alternating and direct current for defibrillation. However, by the 1960s it was apparent that

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**Figure 1** Examples of some monophasic, biphasic, and triphasic waveforms (3–20 ms in duration).
alternating current was more detrimental to the heart and
direct current has been used in defibrillators since. There
are a variety of different waveforms in use today (fig 1).

The critically damped monophasic waveforms have been
the mainstream of external defibrillation for over three decades.
However, these are produced with inductors used in
conjunction with capacitors. Inductors are electrical
components that are too large for implantable devices.
Interest in more recent years has therefore focused on
straightforward capacitor discharges, as capacitors were
suitable for use in implantable defibrillators. Consider fig 2.
When the switch is in position X, the capacitor C will charge
until the potential difference across it is equivalent to the
potential difference across the battery. When the switch
is moved to position Y, exponential discharge occurs through
the resistor R (as shown in the graph of voltage versus time).

The rate of capacitor discharge is described by the equation
\[ Q = Q_0 e^{-t/CR} \]
where \( Q \) represents the charge on the capacitor at a
given time, \( Q_0 \) is the charge at time zero, \( t \) is time and \( e \) is the
exponential function. The time constant of the circuit
(amount of time required for charge to reduce to 37% of its
initial value) is therefore proportional to \( CR \) (the capacitance
value and the resistance through which the discharge
occurs). Adjusting capacitance allows the slope or “tilt” of
the resultant waveform to be controlled. Duration can be
truncated at the desired point. Switching the polarity of the
circuit during the discharge produces biphasic and triphasic
waveforms. The energy produced by a waveform is calculated
from the product of voltage and current (power) multiplied
by time.

There has been a gradual switch to biphasic waveforms in
more recent years due to improved performance in various
clinical trials over their monophasic counterparts. It is
generally agreed that the first phase of a biphasic waveform
should have a positive polarity and duration of equal to or
greater than the second phase of the waveform. The actual
electrode polarity (placing the anode in the apical or
alternative pad position) does not influence the efficacy of
biphasic waveforms. It has been shown that biphasic shocks
produced less dispersion of refractory period extension and
suggested that a more homogeneous dispersion of refractory
period extension was a more important factor in explaining
why biphasic shocks are more efficacious in ventricular
defibrillation. In effect a biphasic shock produces less
heterogeneous repolarisation and is therefore less likely to
re-induce fibrillation.

**PRACTICE OF ATRIAL DEFIBRILLATION (DIRECT
CURRENT CARDIOVERSION)**

**Waveforms**

Several studies of epicardial and endocardial atrial defibrilla-
tion have demonstrated biphasic waveforms to be superior to
their monophasic counterparts of equivalent overall duration.
The biphasic waveforms offer superior efficacy in terms of
peak voltage, current and total energy delivered. This has
been shown in animal models of AF as well as in patients.

The development of devices that deliver impedance
compensated biphasic (ICB) waveforms for use in cardiac
arrest has stimulated interest in their use for direct current
cardioversion. The evidence to date shows that the biphasic
deVICES offer successful cardioversion rates comparable to or
superior to standard monophasic defibrillators, but at
significantly lower energies (table 1). Some physicians may argue that biphasic devices do not offer a
clear advantage over monophasic defibrillators. Arguments
include: there is no large increase in overall success of the
procedure (when both protocols are completed to the highest
energy); there does not appear to be any functional
advantage for the patient from the use of biphasic wave-
forms; biphasic devices have not been shown to influence the
duration of sinus rhythm; and exposure to high cumulative
energies does not lead to troponin release.

However, skin burns are less common with biphasic
waveforms with the likelihood of a skin burn increasing
proportionally with the total energy delivered for both monophasic and biphasic waveforms. Biphasic
CARDIOVERSION is also associated with less skeletal muscle
damage. In addition, fewer shocks are needed with a
biphasic device, potentially leading to shorter cardioversion
procedures. Biphasic waveforms have been shown to be
 efficacious for patients with previously failed attempted
cardioversion with monophasic waveforms. Finally, given
the clear advantages of impedance compensated biphasic
waveforms, monophasic devices in hospital are likely to be
phased out and replaced by their biphasic counterparts. This
will result in an inevitable shift in practice toward biphasic
direct current cardioversion of atrial arrhythmias.

**Electrode positions**

Historically, anteroposterior (AP) configurations have been con-
sidered to provide a superior “shock vector” through the atria
compared with the antero-apical (AA) configuration.
However, the actual path taken by current through human tissue is complex, with a small percentage (~4%) of current delivered by the transthoracic route reaching the heart in the AA configuration. The route taken by current during AP defibrillation has not been described. Two studies have suggested that an AP electrode configuration is superior to an AA electrode configuration when monophasic waveforms are used for direct current cardioversion of AF. However, neither study reported whether there was a difference in transthoracic impedance for the configurations studied. Transthoracic impedance (TTI) has been shown to be an important determinant of atrial and ventricular defibrillation and is now widely accepted as an important predictor of success. We have recently shown in a large series of patients (using ICB waveforms) that the AP pad position is associated with lower transthoracic impedance than the AA configuration. It is likely that this explains the findings of the older monophasic studies, and it is therefore reasonable to recommend the AP electrode positions for monophasic direct current cardioversion.

Several AP pad positions have been described. Some authors feel that a right anterior to left posterior configuration is better where the underlying pathology involves both atria, while a left anterior to posterior configuration is better when the left atrium is primarily affected. The right anterior to left posterior configuration is described, as are the left anterior to right posterior and AP midline configurations. No data have been published showing superiority of one AP configuration over the other. Finally, our comparison of AP and AA electrode configurations using ICB waveforms demonstrated no difference between these approaches with a modern defibrillator.

### Internal atrial defibrillation

Internal atrial defibrillation can also be reliably achieved using electrodes placed in the right atrium and distal coronary sinus. This approach has been widely used in patients and subsequently led to the development of a single chamber atrial defibrillator. This device has been shown to be sensitive and specific in the detection of AF and wave synchronised shocks were delivered appropriately. Follow up of initial implants revealed treatment to be highly efficacious—96% of 227 AF episodes were converted to sinus rhythm. Unfortunately, the frequency of short lived non-treated episodes does not appear to be decreased by this device, and the thromboembolic risk from paroxysmal AF is therefore likely to remain. In addition, a significant number of patients can expect to have the device explanted because of uncontrolled AF.

The single chamber atrial defibrillator is now superseded by dual chamber alternatives. Several of these are currently available. They also offer AF preventative pacing, atrial antitachycardia pacing, and atrial defibrillation functions in addition to ventricular defibrillation. However, the efficacy of these devices for the prevention of AF is unproven and long term clinical benefit has not been demonstrated in large numbers of patients. The role of implantable devices for the treatment of AF alone is undefined at present. The place of transvenous atrial defibrillation as a stand alone procedure has not been established in the era of ICB waveforms. However, there may be a use for low energy internal cardioversion for sustained arrhythmia in the electrophysiology laboratory or in selected patient groups (such as the very obese, after cardiac surgery, or in those where external cardioversion fails).

### Energy selection for direct current cardioversion

There are two approaches to energy selection for direct current cardioversion. After developing the technique of cardioversion, Low began the escalating energy approach. The rationale for this was to minimise post-shock arrhythmia. This approach also allows cardioversion at the lowest energy for each individual patient and may prevent high cumulative doses in some. Practitioners prefer to begin the procedure at the highest energy in order to minimise the total number of shocks delivered and the duration of the procedure (including exposure to sedation or anaesthesia) for the majority of patients, while accepting that a proportion will be relatively “overdosed”. The most recent guidelines for AF suggest a starting energy of at least 200 J for attempted cardioversion of AF using monophasic waveforms. This recommendation is made on the basis that energies lower than this are unlikely to be successful. However, AF of very short duration (for example, after cardiac surgery) may be amenable to lower energy shocks.

Variable energies have been recommended for defibrillators that deliver impedance compensated biphasic waveforms in the correction of AF. The recommended energy depends on the manufacturer of the device. Published evidence suggests that a shock of 150 J will result in success for ~80% of patients with these new devices and there does not appear to be a significant additional benefit in the selection of energies above 200 J (table 2). Indeed, this manoeuvre may result in an excess of skin burns. Using an ICB defibrillator, we favour an initial energy of 150 J followed by a 200 J shock.

### ROLE OF DIRECT CURRENT CARDIOVERSION FOR AF IN THE ERA OF THE RATE CONTROL STRATEGY

The high likelihood of relapse into AF after a successful cardioversion has been recognised for some time. AF has been shown to recur in 88% of patients over 19 months of follow up on “optimal” antiarrhythmic treatment. Thus, withdrawal of anticoagulation during periods of sinus rhythm is likely to carry a risk of thromboembolism in a number of patients. Two large clinical studies have recently...
challenged the rationale of recurrent cardioversion and pharmacological prophylaxis for the maintenance of sinus rhythm in AF patients (table 3). A total of 4060 patients were enrolled in the AFFIRM study. In the rhythm control group (2033 patients), the antiarrhythmic drugs used were chosen by the treating physician. Attempts to maintain sinus rhythm could include cardioversion as necessary. In the rate control group (2027 patients), drugs that were acceptable in the protocol for this purpose were β blockers, verapamil, diltiazem, digoxin, and combinations of these drugs. The goal for antiocoagulation with warfarin was an international normalised ratio (INR) of 2.0–3.0.

In the rhythm control group, continuous antiocoagulation was encouraged but could be stopped at the physician’s discretion if sinus rhythm had apparently been maintained for at least four, and preferably 12, consecutive weeks with antiarrhythmic drug treatment. In the rate control group, continuous antiocoagulation was mandated by the protocol. For patients in the rate control group, 35% were in sinus rhythm at five years and over 80% of those in AF had adequate heart rate control. The prevalence of sinus rhythm in the rhythm control group at follow up was 82%, 73%, and 63% at one, three, and five years, respectively. Electrical cardioversion was attempted once during follow up in 368 patients, twice in 214 patients, and three or more times in 187 patients in this group.

There were 356 deaths among the patients assigned to rhythm control treatment and 310 deaths among those assigned to rate control treatment (mortality at five years, 24% and 21%, respectively; hazard ratio 1.15, 95% confidence interval 0.99 to 1.34; p = 0.08). More patients in the rhythm control group than in the rate control group were hospitalised, and there were more adverse drug effects in the rhythm control group. In both groups, the majority of strokes occurred after warfarin had been stopped or when the INR was subtherapeutic. The authors concluded that “management of AF with the rhythm-control strategy offers no survival advantage over the rate-control strategy, and there are potential advantages, such as a lower risk of adverse drug effects, with the rate-control strategy”. Additionally, anticoagulation should be continued in this group of high risk patients.

The RACE study was smaller, with 522 patients enrolled, but of similar design. Patients in the rate control group received oral antiocoagulant drugs and rate slowing medication. The rhythm control group had a more rigorous management protocol. These patients initially underwent electrical cardioversion without previous treatment with antiarrhythmic drugs. Thereafter, patients received sotalol. If there was a recurrence within six months, electrical cardioversion was repeated and sotalol was replaced by flecainide or propafenone. If there was a recurrence within six months after the start of this regimen, a loading dose of amiodarone was given, and electrical cardioversion was repeated. The dose of amiodarone was then lowered to 200 mg daily. If sinus rhythm was maintained beyond one month, warfarin could be stopped or substituted by aspirin. A total of 35 thromboembolic complications occurred (14 rate control group and 21 rhythm control group). These occurred more commonly during periods of no or subtherapeutic antiocoagulation. The conclusion of the authors was strikingly similar to the AFFIRM study: “rate control is not inferior to rhythm control for the prevention of death and morbidity from cardiovascular causes and may be appropriate therapy in patients with a recurrence of persistent AF after electrical cardioversion”. The STAF pilot study provides further support for a rate control strategy.

Despite these studies, direct current cardioversion is still an important treatment option for patients with symptomatic AF. Many of our patients will complain of lethargy or poor exercise capacity when in AF. It is still reasonable to cardiovert these patients and attempt to maintain sinus rhythm for symptom control. An important lesson from AFFIRM and RACE is that antiocoagulation should be continued in those patients who are at risk from thromboembolism. Cardioversion will also be useful where AF is secondary to another condition (for example, postoperative, pneumonia or an exacerbation of thyrotoxicosis, etc.), where
restoration of sinus rhythm will often be “permanent” after the precipitating event has resolved. Aggressive pursuit of sinus rhythm can no longer be justified in asymptomatic AF. Where the arrhythmia is symptomatic but refractory, alternative strategies such as atrioventricular nodal ablation and pacing should be considered.

FUTURE DEVELOPMENTS IN DIRECT CURRENT CARDIOVERSION

Biphasic defibrillators are replacing their monophasic counterparts for this procedure (monophasic devices are no longer available from most manufacturers). In the next few years warfarin is likely to be superseded by alternative anticoagulants. New antiarrhythmic agents may revitalise the rhythm control strategy in the future if these drugs have comparable efficacy to amiodarone with safer side effect profiles. The role of implantable devices is still under investigation. Hybrid devices that maintain sinus rhythm by multisite atrial pacing and facilitate recurrent internal direct current cardioversion by the delivery of radiofrequency energy\(^6\) have potential for some patients with AF.

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13. A rate control strategy (pharmacological including anti-coagulant treatment) is satisfactory in those where AF has recurred or in those with chronic AF.

Additional references appear on the Heart website—http://heart.bmj.com/supplemental
Theory and practice of defibrillation: (1) Atrial fibrillation and DC conversion

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