An Experimental Study of Transvenous Defibrillation Using a Coronary Sinus Catheter

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KADISH, A.H., ET AL.: An Experimental Study of Transvenous Defibrillation Using a Coronary Sinus Catheter. The efficacy of a transvenous defibrillating system, utilizing bipolar right ventricular and coronary sinus catheters was evaluated in 14 normal mongrel dogs. Two groups of seven animals each were studied. During all shocks, the right ventricular apex electrode served as the anode. In both groups, defibrillation was performed using the proximal pole of the right ventricular catheter (superior vena cava), as the cathode served as a control (configuration A). In group 1, a coronary sinus cathode (configuration B) was compared to control. The mean energy at which 50% or more of the shocks were successful was similar for configuration B (20.7 ± 7.9 joules) and for configuration A (18.8 ± 9.4 joules). In group 2, the superior vena cava and coronary sinus electrodes served as a common cathode (configuration C). Mean defibrillation energy at which 50% or more of the shocks was successful was 21.4 ± 9.0 joules for configuration C and 27.1 ± 9.5 joules for configuration A (P < 0.01). Leading edge voltage was similar for all three configurations, but shock duration was longer for configuration A (11.3 ± 2.8 msec) than configuration B (6.6 ± 1.8 msec) or C (6.1 ± 1.5; P < 0.05). Nonsustained ventricular tachycardia and transient heart block were common, but no damage to the coronary sinus was noted despite the delivery of up to 38 shocks. Conclusions: (1) With the catheter system used, coronary sinus to right ventricular apex defibrillation system offered no advantages over a superior vena cava to right ventricular apex system; (2) A three-electrode system with the high right atrium and coronary sinus serving as the common cathode reduced defibrillation thresholds significantly without any severe short-term adverse consequences; and (3) Improvements in catheter design may make a coronary sinus catheter part of a feasible transvenous defibrillating system. (J Electrophysiol 3:253–260, 1989)

defibrillation, AICD, catheter defibrillation, coronary sinus, ventricular fibrillation

The use of the automatic implantable cardioverter defibrillator (AICD) has resulted in a decrease in mortality in patients suffering life-threatening cardiac arrhythmias. However, significant morbidity is caused by the current implantation procedure involving a thoracotomy, subxyphoid or intercostal incision, and implantation of one or two epicardial patch electrodes. Recently, a nonthoracotomy device involving a subcutaneous patch placed in the left chest has undergone experimental and preliminary clinical testing. Defibrillation thresholds have been somewhat variable using this configuration.

Transvenous defibrillation was investigated early in the development of the AICD using a bipolar electrode placed in the right ventricular apex with the proximal pole being located in the high right atrium. A recent experimental study has investigated defibrillation thresholds using this configuration and found a mean threshold of 26 joules. Since successful defibrillation appears to be related to the volume of the left ventricle that is subjected to
an adequate current density during the shock, we hypothesize that placing a catheter in the coronary sinus in addition to the right ventricular apex might interpose a larger portion of myocardium between the defibrillating electrodes and thus lead to a lowering of the transvenous defibrillation threshold.

Methods

Surgical Procedure

Adult mongrel dogs with a mean weight of 21.6 kg were anesthetized intravenously with sodium pentobarbital, 25 mg/kg and ventilated with a volume ventilator (Harvard Apparatus). The right femoral vein was cannulated for intravenous access, both jugular veins were isolated, and a carotid artery cannulated for blood pressure measurement. The chest was opened via a median sternotomy and pericardial cradle was created.

Two custom-made tripolar catheters (CPI, St. Paul, MN, USA) were used for defibrillation. The catheters had a small tip electrode and two large proximal defibrillating electrodes. The tip electrode was used (along with the distal large electrode) for the induction of fibrillation and was 6 mm in length and had a surface area of 9.0 mm². The proximal defibrillating electrode was 73 mm in length, the distal electrode was 35 mm in length, and the interelectrode distance was 62 mm. Catheter diameter was 3 mm, distal electrode surface area was 277 mm², and proximal electrode surface area was 578 mm². One catheter was placed in the coronary sinus (CS) via the left jugular vein and the second in the right ventricular apex through the right jugular vein. The proximal pole of the right ventricular catheter was in the distal superior vena cava (SVC). Blood pressure was recorded using a fluid-filled catheter. Electrocardiograms and blood pressure were amplified and displayed on an Electronics for Medicine Oscilloscope (Electronics for Medicine, Pleasantville, NY, USA) and recorded on heat sensitive paper.

Ventricular fibrillation was induced through the right ventricular catheter using alternating current. An external cardioverter defibrillator (CPI) connected to the transvenous leads was used for defibrillation. If the transvenous shocks were unsuccessful, dogs were rescued with internal paddles connected to a standard defibrillator. Bipolar Teflon-coated silver wire pacing electrodes were threaded with a 21-gauge needle into the subepicardium. A Bloom DTU-110 Stimulator (Bloom Assoc., Norbeth PA, USA) was used to pace the heart at a cycle length of 500 msec if temporary heart block lasting more than 3 seconds developed following fibrillation and defibrillation.

Experimental Protocol

Two groups of seven animals each were studied to examine the use of a coronary sinus catheter in a bipolar (group 1) or tripolar (group 2) configuration. In both groups, a right ventricular apex anode and a superior vena cathode (configuration A) both of which were on the same catheter served as the control. In each group, a test configuration was compared to configuration A. In group 1, defibrillation using the distal coronary sinus cathode and right ventricular apex anode (configuration B) was compared to control. In group 2, configuration C which consisted of the right ventricular apex as the anode and the superior vena cava (proximal right ventricular catheter) and distal coronary sinus as a common cathode was compared to control.

A truncated exponential waveform was delivered at a pulse energy varying between 5-40 joules. Defibrillating attempts began 12 seconds after the induction of fibrillation. At least 5 minutes elapsed between defibrillation attempts. In order to account for the variability in defibrillation success at a given energy, a crude defibrillation threshold (DFT) was first identified and then multiple shocks were delivered around that energy to perform probability calculations. To determine the crude DFT, the first shock was delivered at a 40-joules energy level. If defibrillation was successful, energy output was decreased in 5-joules steps to 10 joules and 2-joules steps below that. At least 5 minutes elapsed between defibrillation attempts. After the crude DFT was identified,
up to five shocks were delivered using each configuration at three energy levels adjacent to the crude DFT to allow probability calculations. At each energy level, shocks using one of the configurations was available for comparison with the other configuration. Animals were included for analysis if at least three pairs of the shocks could be delivered at each of these three adjacent energy levels. Ventricular tachycardia was defined as that having a uniform beat to beat morphology and lasting more than 6 seconds following the delivery of a defibrillation shock.

Voltage waveforms were monitored during defibrillation shocks using a standard 10x voltage probe (Tektronix, model P6122, Tektronix, Beaverton, OR, USA) with an input impedance of 11 Mohms. Voltage waveforms were displayed on an oscilloscope and photographed on Polaroid film (Polaroid, Boston, MA, USA).

Data Analysis

Data are expressed as mean ± SD unless otherwise indicated. The successful energy mean was computed from the successful total energy at each configuration divided by the number of successful shocks. Defibrillation efficacies were compared by defining the energy at which more than 50% of shocks were successful in a given configuration ($E_{50}$). Variability in heart rate and blood pressure were analyzed using a coefficient of variation. The incidence of ventricular tachycardia and heart block were compared by Chi-square analysis with Yates continuity correction. Voltage measurements in the different configurations were compared using repeated measures analysis of variance using Scheffe's F-test.

Results

The animals remained physiologically stable through the multiple shock procedure. Mean systolic blood pressure was 116 ± 17 mmHg, mean diastolic blood pressure was 91 ± 21 mmHg, and mean pulse was 156 ± 22 beats/min. Groups A and B had similar pressure and pulse measurements during the protocol. There was no significant change in blood pressure or pulse with time (Fig. 1).

Defibrillation

Mean weight was 19.6 ± 6.6 kg in group 1. A total of 196 shocks were analyzed in group 1 (mean 28 per animal per configuration). One hundred and ten (56%) were successful. There was no significant difference in the number of successful shocks between the two configurations. Fifty-four shocks (55%) were successful using configuration B (RV apex to distal coronary sinus) and 56 shocks using configuration A (57%; $P = NS$). Mean defibrillation energy at which 50% or more of the shocks were
successful ($E_{50}$) was $20.7 \pm 7.9$ joules for configuration B and $18.8 \pm 9.4$ joules for configuration A ($P = \text{NS}$). Values for $E_{50}$ in individual animals are shown in Figure 2. There was also no difference in the mean defibrillation energy per kilogram between the two configurations ($1.17 \pm 0.63$ joules/kg for configuration B and $1.06 \pm 0.23$ joules/kg for configuration A; $P = \text{NS}$).

Mean weight was $24.4 \pm 2.8$ kg in group 2 ($P = 0.010$ vs group 1). One hundred and fourteen of 214 shocks were successful at defibrillation (53%). Forty-three shocks (40%) were successful using configuration A (RV apex to common SVC distal CS) and 71 (66%) were successful using configuration C (RV apex to superior vena cava). This difference was significant at the $P < 0.001$ level. Mean defibrillation energy at which 50% or more of shocks were successful (Fig. 3) was significantly lower using configuration C ($21.4 \pm 9.0$) than configuration A ($27.1 \pm 9.5; P < 0.01$). Mean $E_{50}$ was $0.89 \pm 0.4$ joules/kg for configuration C and $1.12 \pm 3.8$ joules/kg ($P < 0.01$) for configuration A.

Voltage Measurements

Shock amplitude and duration were measured during test pulses using each of the configurations. Mean leading edge voltage was similar in the three configurations ($622 \pm 137$ volts for configuration A, $610 \pm 131$ volts for configuration B, and $617 \pm 151$ volts for configuration C). However, pulse duration was significantly shorter when using a catheter configuration including the coronary sinus catheter (configurations B and C). Mean pulse duration was $11.3 \pm 2.8$ msec for configuration A, $6.6 \pm 1.8$ msec for configuration B, and $6.1 \pm 1.5$ msec for configuration C ($P < 0.01$ configurations B and C vs configuration A). To determine the effects of catheter location within the coronary sinus, voltage measurements were made with a catheter tip in the proximal coronary sinus and compared to those with a catheter tip in the posterior coronary sinus. Mean pulse duration was $10.5 \pm 2$ msec in the proximal coronary sinus and $6.0 \pm 2.2$ msec in the lateral coronary sinus ($P < 0.05$).

Adverse Effects

Despite the delivery of up to 38 shocks (including rescue shocks) through the coronary sinus catheter, no instances of hemorrhage, per-
foration or visible damage to the coronary sinus occurred. Data on arrhythmias was not available for two group 1 animals.

Nonsustained episodes of ventricular tachycardia and transient complete heart block were seen frequently. A total of 162 episodes of nonsustained ventricular tachycardia with a mean rate of 211 ± 31 beats/min and mean duration of 15 ± 10 seconds were seen. The occurrence of ventricular tachycardia did not appear to be related to energy level but because of the study design, only a narrow range of energy levels (near the defibrillation threshold) were frequently tested in each animal. In group 1, nonsustained ventricular tachycardia occurred in 40 of 68 (59%) defibrillation attempts using configuration B and 14 of 68 (21%) attempts using configuration A (P < 0.001). In group 2, nonsustained ventricular tachycardia was equally frequent in configuration A (50 of 107; 48%) and configuration C (58 of 107; 54%).

Transient complete heart block was observed frequently, but the duration could not be precisely determined because back-up pacing was begun at a rate of 117 beats/min for asystole. Except in one group 1 animal, heart block was always resolved by the time of delivery of the next shock. The incidence of heart block was not different between the two configurations in either group. In group 1, 11 of 53 shocks (21%) delivered between the distal CS and right ventricular (RV) apex (configuration A) were associated with complete heart block versus 6 of 53 (11%) with configuration B (P = NS). Four of five of the animals in which arrhythmia data was available developed at least one episode of complete heart block. In group 2, 22 of 107 (21%) shocks delivered with configuration A developed complete heart block versus 15 of 197 with configuration C (P = NS). All but one of seven animals developed at least one episode of transient complete heart block.

Discussion

The major finding of this study is that an electrode configuration including a transvenous catheter in the coronary sinus reduces defibrillation thresholds in comparison to a bipolar superior vena cava to right ventricular apex configuration. Although bipolar defibrillation from the coronary sinus to right ventricular apex did not significantly reduce defibrillation threshold, the use of the superior vena cava and coronary sinus as a common cathode did. Despite up to 38 shocks in individual animals, no damage to the coronary sinus was seen. Thus, the use of a catheter in the coronary sinus may prove practical in order to reduce transvenous defibrillation thresholds.

Defibrillation Thresholds

Successful defibrillation depends on the delivery energy and electrode location. Several studies have addressed the energy dependence of defibrillation. Defibrillation is a statistical phenomenon without a single clear cutoff energy between successful and unsuccessful defibrillation. Gold et al.11 suggested in 1979 that a single cutoff energy was not appropriate to characterize defibrillation efficacy and devised a contour graph relating percent success of defibrillation to the energy characteristics of the device. More recently, Davey et al.12 showed that in dogs subjected to internal defibrillation there was no clear cutoff energy between successful and unsuccessful energies. They constructed dose response curves for defibrillation versus energy, and suggested analyzing defibrillation success in individual animals to improve the sensitivity in detecting differences between defibrillation methods. Although other investigators have supported the concept of a defibrillation threshold, data in the current study was analyzed using the level of energy at which at least 50% of shocks of E50 produced successful defibrillation. A total of up to five shocks were delivered at each energy level using each configuration.

Determinants of Defibrillation

Differing shock waveforms, energy levels, electrode positions, body and heart weight, cardioactive drugs and ischemia have been shown to affect energy requirements for defibrillation.13-16 The location of defibrillating elec-
trodes also has a major effect on defibrillation efficacy. Although it had previously been hypothesized that a specific location of defibrillation electrodes was necessary for adequately depolarizing a critical mass of myocardium, it has recently been shown that even unsuccessful shocks can lead to depolarization over the entire heart. However, the isoelectric interval before resumption of fibrillation was shorter in unsuccessful shocks and ventricular fibrillation regenerated at sites distal from the defibrillating electrodes. Thus, although simple depolarization of distal myocardial sites is not adequate to insure successful defibrillation, the current density around such sites and thus defibrillating electrode location, can affect defibrillation efficacy. In the current study, catheters were positioned in the lateral coronary sinus and right ventricle in an attempt to interpose large amounts of myocardium between the electrodes.

Defibrillation Lead Systems

Clinical defibrillators use two large patches placed directly on the epicardial surface of the heart or a high right atrial superior vena cava lead combined with an epicardial patch. This approach has resulted in defibrillation thresholds that have generally been < 30 joules, but requires epicardial implantation of at least one lead. Experimental approaches to other lead systems have included the use of a subcutaneous patch to right-sided electrodes in an attempt to reduce current pathways across large portions of the left ventricle during defibrillating shocks efficacy. Recent clinical trials have demonstrated some success with this approach. Several studies have examined three-electrode configurations for defibrillation. Many of these have used sequential shocks using two separate anode cathode configurations have previously examined a coronary sinus electrode as part of a sequential shock system, but they did not systematically test the effect of this electrode on defibrillation thresholds.

A three-electrode configuration was also evaluated in this study. Although pulse width was also approximately 6 msec in duration (suggesting some current shunting), energy requirements for defibrillation were lower than control using a common high right atrial coronary sinus cathode. Thus, this configuration could potentially contribute to an effective transvenous defibrillating system.

Limitations

There are several limitations to the current study. Defibrillation was tested in an open-chested model and current pathways could potentially be different in such a situation. However, this should be less of a problem in testing a transvenous system than an epicardial one and a recent study showed that even with epicardial patches there was no difference in defibrillation threshold between open- and
closed-chested animals. In addition, testing was performed on mongrel dogs without heart disease and there may be limitations in applying these findings to patients with organic heart disease.

Only a single polarity for the right ventricular apex electrode (positive) was evaluated in this study. Official recommendations have suggested that the left ventricular patch electrode always has negative polarity. However, a recent study by Bardy et al. showed that defibrillation thresholds were actually lower when the left ventricular patch was negative. It is possible that if the coronary sinus electrode had served as the anode, even lower defibrillation thresholds would have been found since this catheter was in closest proximity to the left ventricle. However, the fairest comparison to the control (RV apex to SVC) configuration involved using the right ventricular apex as the anode since it was the only electrode in proximity to the left ventricular myocardium in both configurations.

The absolute value of defibrillation threshold for the control configuration was substantially higher in group 2 than in group 1. However, weight which may be a determinant of defibrillation threshold was higher in group 2 and the mean defibrillation threshold per kg was similar in the two groups.

No gross damage to the coronary sinus was observed in this study. However, histologic analysis after chronic lead implantation was not performed and will be required in future studies before the technique can be deemed safe.

**Clinical Implications**

The results of this study suggest that a catheter in the coronary sinus could potentially form a part of an effective transvenous defibrillating system. Although the mean defibrillation energy using this configuration was still relatively high, the use of bipolar waveforms and improvements in catheter design including shortening the electrodes could potentially reduce energy requirements to the point where they could be clinically useful. Further work including chronic studies on damage to the coronary sinus and incorporating changes in catheter design and waveform pulses will be necessary to confirm the feasibility of this approach.

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**References**


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