Chronic performance of a novel radiofrequency ablation device on the beating heart: Limitations of conduction delay to assess transmurality

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Objective: The creation of consistently transmural lesions with epicardial ablation on the beating heart has represented a significant challenge for current technology. This study examined the chronic performance of the AtriCure Coolrail device (AtriCure Inc, West Chester, Ohio), an internally cooled, bipolar radiofrequency ablation device designed for off-pump epicardial ablation. The study also examined the reliability of using acute intraoperative conduction delay to evaluate lesion integrity.

Methods: Seven swine underwent median sternotomy. The right atrial appendage and inferior vena cava were isolated with a bipolar radiofrequency clamp. Linear ablation lines were created between these structures with the AtriCure Coolrail. Paced activation maps were recorded with epicardial patch electrodes acutely before and after ablation and after keeping the animals alive for 4 weeks. The conduction time across the linear ablation was calculated from these maps. The lesions were histologically evaluated with trichrome staining.

Results: Only 76% of cross-sections of Coolrail lesions were transmural, and only 1 of 12 ablation lines was transmural in every cross-section examined. Mapping data were available in 5 of the animals. Significant conduction delay was present after the creation of each line of ablation acutely; however, after 4 weeks, conduction time returned to preablation values, demonstrating lack of transmurality.

Conclusions: The AtriCure Coolrail failed to reliably create transmural lesions. Although the Coolrail was able to create acute conduction delay, its failure to transmurally ablate the atrial myocardium left gaps along the length of the lesion, which resulted in neither chronic conduction block nor delay across any line of ablation.

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assess whether a lesion has resulted in a transmural scar in the operating room. This study tested the efficacy of this technique to predict chronic conduction block in a porcine model.

**MATERIALS AND METHODS**

Seven Hanford miniature swine weighing 50 to 70 kg were used in this study. All animals received humane care in compliance with the Guide for the Care and Use of Laboratory Animals (National Academy Press, Washington, DC). Each animal was premedicated with tiletamine HCl (Telazol; Fort Dodge Animal Health, New York, NY), ketamine, and xylazine; intubated; anesthetized with isoflurane; and monitored continuously throughout the procedure with electrocardiography and invasive arterial pressure recordings. Perioperative antibiotics were given during the initial surgery and consisted of cefitofur (2 mg/kg). Antibiotics were continued for 3 days postoperatively.

Each pig had a set of right and left atrial ablations performed through a median sternotomy. After creation of a pericardial sling, a set of 3 molded silicone plaques with a total of 252 unipolar electrodes were placed onto the epicardial surface to obtain epicardial activation maps. The electrode templates were constructed from a form-fitting silicone elastomer (Specialty Silicone Fabricators, Paso Robles, Calif) that fit snugly on the entire atrial epicardium and contained 0.5-mm-diameter silver electrodes (Pacific Wire & Cable, Inc, Santa Ana, Calif). The interelectrode distance was 5 mm. These plaques were secured with Rommel tourniquets to allow for consistent placement before and after the lesion set creation. After a baseline activated clotting time was measured, a heparin bolus (100 units/kg) was given intravenously and activated clotting time was maintained at more than 350 seconds until after all ablations were completed.

Pacing thresholds were obtained on either side of the planned lines of ablation as shown in Figure 1. Activation maps during normal sinus rhythm and paced activation (twice threshold, 120 beats/min) were recorded from identical sites before and immediately after completion of the lesion set. The lesion set is shown in Figure 1. The atrial appendages were isolated with a single application of the AtriCure Synergy device, a bipolar RF clamp device with dual electrodes on each jaw of the clamp. The device was used per the manufacturer’s instructions. Ablations around the inferior vena cava and the left pulmonary veins were also created with single applications of the Synergy device. On the right atrium, a line of ablation was created with multiple applications of the AtriCure Coolrail device to connect the ablation at the base of the right atrial appendage to the ablation around the inferior vena cava. This epicardial RF ablation device has been described in another publication from our laboratory (Figure 2). Briefly, the device is a handheld pen-like device that has 2 hollow, 3-cm-long electrodes through which sterile water is circulated during ablation to reduce tissue-electrode temperature. An analogous line on the left atrium was created with multiple applications of the Coolrail device between the base of the left atrial appendage and the left pulmonary veins. Initially, application of the Coolrail was planned for 50 seconds with an overlap of 0.5 cm per application based on dosimetry data previously published. After perforation and subsequent death due to intraoperative hemorrhage in the first pig with this ablation time, the protocol was changed to a 40-second ablation with only minimal overlap of prior ablation lines. After completion of the ablation protocol and acquisition of the acute postablation activation maps, hemostasis was secured. The pericardium was closed, and the sternum was reapproximated. After the operation, the animal was closely monitored for at least 48 hours and survived for 28 days. Each animal received 325 mg of aspirin daily beginning on the first postoperative day. The animals received buprenorphine (0.005–0.01 mg/kg intramuscularly) for postoperative analgesia.

At 28 days, the animals underwent a reoperative sternotomy using the same anesthesia protocol used for the initial operation. The pericardium was reopened, and the atria were carefully dissected free of adhesions. The epicardial electrode plaques were repositioned, and activation maps were obtained during normal sinus rhythm and paced rhythm from the same sites as during the initial operation.

Once the chronic postablation maps were obtained, the animals were euthanized humanely with a concentrated potassium chloride solution. The aorta was crossclamped, and the hearts were perfused with 60 mL of 1% 2,3,5-triphenyl-tetrazolium chloride solution through the coronary circulation via infusion into the root of the aorta. The hearts were removed en bloc and examined grossly for any evidence of intra-atrial thrombus formation. They were then placed in 1% 2,3,5-triphenyl-tetrazolium chloride solution and incubated at room temperature for 45 minutes to aid in visualization of the ablations. Each lesion was sectioned at 5-mm intervals perpendicular to the long axis of the ablation. The sections were fixed in formalin, molded in paraffin, and stained with Gomori trichrome. The sections were microscopically examined to assess transmurality and measure lesion depth and tissue thickness.

Analysis of mapping data was done offline using custom-designed software as described previously. Unipolar electrogram data were recorded at a gain of 250 and a frequency response of 0.5 to 500 Hz. The data were digitized at 1000 Hz. Activation times were picked automatically by selecting the peak derivative of voltage over time from the electrogram after paced activation. Automatic activation time was manually reviewed and edited for accuracy. Only the activation maps on the right side of the atria were analyzed because the left atrium in the Hanford miniature swine was too narrow in relation to the Coolrail ablation line, and the spatial resolution of the mapping system was not fine enough to map over the remaining viable tissue of the left atrium.

**Statistical Analysis**

All data are expressed as mean ± standard deviation. Comparison of conduction time across the line of ablation was done with a Student t test.

**RESULTS**

**Operative Results**

The first animal died intraoperatively during the initial procedure from a perforation caused by the Coolrail device. In the initial protocol, applications of the device were for 50 seconds, and there was at least 0.5 cm of overlap between applications to ensure a continuous linear ablation. Given the perforation, the protocol was modified as noted in the previous section, and subsequently, the remaining 6 animals survived the operation. There were no further perforations caused by the device after adaptation of the new protocol. There were no postoperative complications related to ablation. None of the animals showed any signs of cerebrovascular accidents or other signs of thromboembolic events.

**Histology**

Gross visual inspection of ablations revealed no intra-atrial thrombi. The device performance for the Coolrail is summarized in Figure 3. Sectioning of the 12 connecting lesions created by the Coolrail yielded 72 total cross-sections.
Of these, 76% (55/72) were transmural. Only 1 of the 12 connecting lesions created in the study had transmural ablations at every cross-section examined. This was one of the lines connecting the base of the right atrial appendage to the inferior vena cava. At least 1 (range, 1–4) cross-section in each of the other 11 lines demonstrated failure to achieve transmural ablation. In 2 cases, there was an outright gap in the ablation line at the point of overlap between applications of the Coolrail device. These gaps were excluded from analysis because they represented tissue that was not ablated, rather than failed device performance.

Device performance varied with tissue depth (Figure 3). In tissue less than 4 mm thick, 97% (33/34) of all cross-sections examined were transmurally ablated. In tissue 4 mm thick or greater, 58% (22/38) of cross-sections were transmurally ablated. Average lesion depth for the study was 3.7 ± 1.5 mm, and tissue thickness was 4.4 ± 2.2 mm. Average lesion width was 8.6 ± 2.5 mm.

Electrophysiology

Complete mapping data, including preablation, acute postablation, and chronic postablation activation maps, were available for analysis in 5 animals. The pericardium of the sixth animal was too tightly adhered to the atrial epicardium to allow for safe dissection and subsequent mapping.
A combination of conduction delay and change in activation sequence was used to evaluate for conduction block on acute and chronic maps (Figure 4). Conduction block was defined by a change in the activation sequence around the observed line of block, and conduction delay was defined as delay in conduction without a change in the sequence of activation. Four of five activation maps demonstrated acute conduction block after the ablation line was performed. The other showed only conduction delay without block (from 13 ms preablation to 38 ms postablation) through the line of ablation. Of note, this map correlated with the sole line of ablation with completely transmural cross-sections in histologic analysis. Conduction time in the chronic setting for this right atrium decreased to 28 ms.

Conduction time across the Coolrail line of ablation was significantly increased after ablation in the initial operation. Conduction time before ablation averaged 14.4 ± 5.6 ms. This increased significantly to 36.8 ± 15.1 ms acutely postablation (P = .007). Four weeks after ablation, epicardial mapping demonstrated a return to baseline conduction times. The chronic postablation conduction time averaged 19.4 ± 6.8 ms and was not statistically different from preablation values (P = .242). The conduction time for all 5 animals with complete maps is plotted in Figure 5. In each case, conduction delay across the Coolrail line was increased in the acute postablation map but decreased toward preablation values in the chronic epicardial map at 4 weeks.

DISCUSSION

In the last decade, the Cox-Maze procedure has been modified to make it more accessible and less invasive. Devices have been used to replace the incisions of the

FIGURE 4. Representative sequence of activation maps demonstrating normal paced activation preablation, acute conduction block with a change in activation sequence around the ablation line (dotted line) acutely, and recovery of conduction through nontransmural gaps chronically (4 weeks). Conduction times were measured between the 2 points marked by asterisks.

FIGURE 5. Conduction time across the Coolrail lesion line (see Figure 4) is plotted for the preablation, acute postablation, and chronic postablation periods. There was no statistical difference between conduction times preablation and chronic postablation 4 weeks after the initial operation (P = .242).
Cox-Maze III with lines of ablation to make it easier and faster to perform. At Barnes-Jewish Hospital, the use of bipolar RF clamps has decreased the operative time by more than half. Moreover, these bipolar devices have been shown experimentally to create reliable transmural lesions. The bipolar RF-based Cox-Maze IV had achieved more than 80% freedom from AF off antiarrhythmic drugs at 6 months. However, these devices still require cardiopulmonary bypass and small incisions in the atria to perform the entire Cox-Maze lesion set.

Less-extensive lesion sets have been introduced to decrease the invasiveness of these procedures so they can be performed without cardiopulmonary bypass. The most common are limited to pulmonary vein isolation alone and occasionally involve surgical ablation of the ganglionated plexi. These operations are attractive because they can be done with minimal access techniques and without cardiopulmonary bypass, avoiding potential morbidity. The results of these new procedures have been less than satisfactory, particularly in patients with long-standing, persistent AF, in whom the success rates can be less than 50%. This is not surprising, and similar results have been demonstrated in the electrophysiology laboratory with pulmonary vein isolation alone in patients with long-standing persistent AF.

To improve results, more extensive left atrial lesion sets have been proposed. These have required creating linear lesions on the beating heart. To answer this need, devices have been recently introduced to perform epicardial beating heart ablation on the atria. However, experience in our laboratory and others has shown that it is difficult to perform consistently transmural lesions on the beating heart. We have shown that even small gaps (<1 mm) in a linear lesion can result in a lack of conduction block. Therefore, it is vital that any new device be tested experimentally before widespread clinical use. The AtriCure Coolrail device was introduced specifically to make linear lesions on the beating heart. The device was not independently tested experimentally before clinical use. The first goal of this study was to examine whether the device was capable of creating consistent transmural ablation in a chronic animal model. This study simulated the proposed clinical use of the device by making connecting lesions to create lines of conduction block.

The Coolrail was not capable of creating a long linear lesion in this chronic study. The performance of the Coolrail has been described by this laboratory in an acute animal study. None of the 12 lines created with the device created a pathologically or electrically transmural lesion. The only ablation line that had all histologic sections transmural had no evidence of conduction block by epicardial mapping. Although every cross-section examined in the line of ablation was transmural, the pattern of conduction in the chronic activation map did not demonstrate acute conduction block, indicating that there must have been a gap or gaps along the ablation line.

These chronic results confirm the acute results found after staining with 1% 2,3,5-triphenyl-tetrazolium chloride solution. Although this study was not designed specifically to compare acute histology with chronic histology, comparing the chronic histologic data with previously acquired acute data, there was no statistically significant difference in depth of tissue penetration (P = .147) or likelihood of transmural ablation (P = .621) as determined by acute staining with 1% 2,3,5-triphenyl-tetrazolium chloride solution in our prior study versus chronic histologic evaluation with trichrome staining.

The inability of this device and other unidirectional devices we have tested to create reliably transmural lesions emphasizes the importance of developing an intraoperative method to test for lesion integrity. Various groups have proposed using conduction block or delay as a way to verify lesion transmurality. This study also demonstrated that acute conduction delay or block did not predict chronic transmurality when used to assess epicardial RF ablation in the porcine atria. Histologic analysis demonstrated that the majority of lines created in this study with the Coolrail contained at least 1 nontransmural gap. These gaps were not readily identifiable on the basis of activation sequence mapping. In each case where a gap was seen clearly in histology, the activation map demonstrated a pattern consistent with acute conduction block. The wavefront of electrical activation traveled from the right atrial free wall around the anterior surface of the atrium to activate the atrial tissue on the opposite side of the line of ablation. The Coolrail contained at least 1 nontransmural gap. These gaps were clearly identified, and conduction times returned to baseline.

A shortcoming of acute intraoperative activation mapping is demonstrated in this study. When gaps were present histologically, no evidence of this was found acutely with activation mapping. The phenomenon of chronic recovery of conduction is well described in the electrophysiology literature. Catheter-based pulmonary vein isolation for AF is based on identifying and ablating conduction gaps until the pulmonary veins demonstrate entrance and exit block. Although every effort is made at the initial ablation to isolate the pulmonary veins, patients can present with recurrent AF, and repeat electrophysiologic mapping often reveals recovery of conduction from the majority of pulmonary veins. This recovery seen clinically is similar to the recovery of conduction in this study. There is evidence of acute injury and recovery at the tissue level. One echocardiography-based study found swelling in the ablated crista terminalis that resolved over time.
the course of 4 weeks. This observation likely explains the acute conduction block, because localized tissue swelling and inflammation may lead to acute stunning and electrical block. However, as the tissue recovered, electrical activity was restored. Further evidence in the electrophysiology literature suggests that this recovery of activation may not take an extended period of time. Observing the results of pulmonary vein isolation over the course of an hour revealed recovery of conduction in a significant number of patients.

This study underscores the importance of using a reliable, rigorously investigated energy source when performing surgical ablation until a reliable method of intraoperative lesion evaluation is developed. The practice at Barnes-Jewish Hospital is to use devices that have been shown to reliably create transmural ablation, such as bipolar RF clamps. In addition, exit block from the pulmonary veins reliably create transmural ablation, such as bipolar RF lesions. Although the Coolrail was able to create acute lesions. The animal model used in this study may not perfectly simulate conditions encountered clinically. The average thickness of atrial tissue encountered in this study was 4.4 ± 2.2 mm. In some areas of the diseased human atrial myocardium left gaps along the length of the lesion, which resulted in neither chronic conduction block nor delay across any line of ablation.

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Study Limitations

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Conclusions

The AtriCure Coolrail failed to reliably create transmural lesions. Although the Coolrail was able to create acute lesions, it failed to transmurally ablate the atrial myocardium left gaps along the length of the lesion, which resulted in neither chronic conduction block nor delay across any line of ablation.

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References


