Use of endogenous NADH fluorescence for real-time in situ visualization of epicardial radiofrequency ablation lesions and gaps

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Mercader M, Swift L, Sood S, Asfour H, Kay M, Sarvazyan N. Use of endogenous NADH fluorescence for real-time in situ visualization of epicardial radiofrequency ablation lesions and gaps. Am J Physiol Heart Circ Physiol 302: H2131–H2138, 2012. First published March 9, 2012; doi:10.1152/ajpheart.01141.2011.—Radiofrequency ablation (RFA) aims to produce lesions that interrupt reentrant circuits or block the spread of electrical activation from sites of abnormal activity. Today, there are limited means for real-time visualization of cardiac muscle tissue injury during RFA procedures. We hypothesized that the fluorescence of endogenous NADH could be used as a marker of cardiac muscle injury during epicardial RFA procedures. Studies were conducted in blood-free and blood-perfused hearts from healthy adult Sprague-Dawley rats and New Zealand rabbits. Radiofrequency was applied to the epicardial surface of the heart using a 4-mm standard blazer ablation catheter. A dual camera optical mapping system was used to monitor NADH fluorescence upon ultraviolet illumination of the epicardial surface and to record optical action potentials using the voltage-sensitive probe RH237. Epicardial lesions were seen as areas of low NADH fluorescence. The lesions appeared immediately after ablation and remained for several hours. Real-time monitoring of NADH fluorescence allowed visualization of viable tissue between the RFA lesions. Dual recordings of NADH and epicardial electrical activity linked the gaps between lesions to postablation reentries. We found that the fluorescence of endogenous NADH aids the visualization of injured epicardial tissue caused by RFA. This was true for both blood-free and blood-perfused preparations. Gaps between NADH-negative regions revealed unablated tissue, which may promote postablation reentry or provide pathways for the conduction of abnormal electrical activity.

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Radiofrequency ablation (RFA) is an effective therapy for treating atrial and ventricular rhythm disturbances (16). Nearly 100,000 RFA procedures are performed annually in the United States to treat cardiac arrhythmias (29). RFA targets the key elements of reentrant pathways and/or abnormal ectopic loci without damaging a significant amount of adjacent healthy myocardium and coronary vessels. RFA lesion size is not simply a function of delivered energy but depends on many factors, including the contact between the catheter tip and the tissue, the thickness of the myocardium, the amount of blood flow, and the presence of fat (16, 33). Conventional electroanatomical mapping systems map mainly the location of the catheter tip but not the extent of tissue injury caused by ablation. Therefore, as of today, RFA ablation lesions are created with minimal information regarding the physiological condition of the affected tissue.

Gaps of excitable tissue between ablation lesions are directly related to arrhythmia recurrences. Two main strategies have been proposed to reduce the incidence of gaps. The first is to improve ablation devices, which includes the development of multipolar and linear catheters (3, 15), balloon-based technologies using lasers, focused ultrasound and cryoinjury (9, 26), as well as pressure sensor-equipped catheters (34). The second strategy is to visualize RFA lesions during the ablation procedure (1, 7, 10–12, 20, 31). Such visualization could be based upon acute changes in the chemical and/or physical properties of the damaged tissue. The approach described in this article is based upon imaging the fluorescence of endogenous NADH (fNADH) using low-intensity ultraviolet (UV) light illumination.

NADH is a coenzyme that is present within all intact cells. Once NADH is released from the mitochondria of damaged cells and/or converted to its oxidized form, its fluorescence markedly declines. Recent advances in tumor surgery have used endogenous fNADH as a means to monitor tissue necrosis during RFA surgery and provided general proof of the method’s feasibility (21, 22). To the best of our knowledge, this is the first imaging of NADH as a means to visualize viability of cardiac tissue during RFA procedures. Our results show that reductions in epicardial fNADH caused by RFA are so substantial that the method reveals lesions not only in blood-free but also in blood-perfused cardiac preparations.

METHODS

Animal procedures. Ex vivo experiments were conducted using Sprague-Dawley (200–300 g) and New Zealand White rabbits (2.5–3.5 kg). Animals were heparinized and anesthetized using standard procedures (2, 17). Hearts were excised, and the aorta was cannulated and Langendorff-perfused at constant pressure (50 mmHg) with oxygenated, buffered Tyrode solution at room temperature. During the ablation period, hearts were placed on top of a grounding pad and submerged in 37°C Tyrode solution.

In situ experiments were performed using anesthetized open-chest rats (200–300 g Sprague-Dawley). After an intraperitoneal injection of Telazol (40 mg/kg), the hair on the chest and back was shaved, the animal was immobilized on a heated platform, and an ablation pad was placed beneath the animal. Immediately after opening the chest cavity, ablations were carried out as the exposed epicardial surface was imaged. All anesthesia and euthanasia procedures were approved by the George Washington University Medical Center Institutional Animal Care and Use Committee.

Ablation protocols and NADH recordings. Radiofrequency energy was delivered using a noncooled blazer catheter with a 4-mm tip (EP Technologies, Boston Scientific). Tip temperatures ranged between 50 to 70°C. The catheter was placed perpendicular to the epicardial
surface. Ablation durations varied from 15 to 60 s with a maximum power of 50 watts. The epicardial surface was illuminated with UV light (350/25 nm) using a 100-watt mercury lamp (Zeiss HBO100 W/2). To record the epicardial fluorescence of NADH, the emitted light was filtered (460/25 nm) and imaged using a charge-coupled device camera (Andor Ixon DV860) that has high quantum efficiency for wavelengths corresponding to NADH fluorescence (80% quantum efficiency at 460 nm).

**Optical mapping experiments.** Hearts were stained with the potentiometric dye RH237 (10 μM final concentration; Molecular Probes). To reduce motion artifact, blebbistatin was added to the perfusate at a final concentration of 10 μM. A dual optical mapping system comprised of two cameras (Andor Ixon DV860s) fitted with a dual port adapter (Andor CSU Adapter Dual Cam) and a dichroic mirror (610 nm) was used to image the epicardial fluorescence of RH237 (250–500 frames/s (fps)) and NADH (2 fps) from the same field of view (32). To record optical action potentials, the epicardium was illuminated using two light-emitting diodes (LumiLEDs, 530/35 nm). The resulting RH237 fluorescence was long pass filtered at 680 nm. NADH fluorescence was recorded with the other camera as described above. The fluorescence of RH237 was processed to subtract background fluorescence from each image, and signals for each pixel were normalized. RH237 fluorescence signals were smoothed using a median temporal filter (3 sample width). Isochronal maps of activation times were generated to show wave front propagation. The average amplitude of optical action potentials at each pixel was computed to reveal spatial changes in the amount of electrically active tissue.

**Triphenyltetrazolium chloride staining.** Triphenyltetrazolium chloride (TTC) vital staining is a standard procedure for assessing acute necrosis. It relies on the ability of dehydrogenase enzymes and NADH to react with tetrailum salts to form a formazan pigment. Immediately after the imaging protocol, the tissue was retrograde-perfused with Tyrode solution containing 1.0% TTC. Afterward, the heart was submerged in the TTC solution for an additional 8 min. Metabolically active tissue appeared crimson. Necrotic tissue appeared white.

**RESULTS**

**Appearance of RFA lesions in excised, blood-free rat and rabbit hearts.** In total, 16 lesions in 8 rat hearts and 8 lesions in 3 rabbit hearts were studied. When observed on the NADH-sensitive channel, ablated areas appeared markedly dark when compared with the surrounding myocardium (Fig. 1A, center). TTC staining revealed areas of irreversible injury consistent with the shape and the size of RFA lesions (Fig. 1A, right). During the 1- to 2-h experiments, fNADH levels in ablated tissue did not return to their prevablation values, and the size of the lesions did not change significantly (n = 3; Fig. 1B, row on top). Over time, lesion borders became more homogenous (n = 3; Fig. 1C). A systematic comparison of fNADH lesions with TTC-negative areas showed that lesion size and shape matched with better than 95% accuracy (Fig. 2).

**Identification of functional gaps between RFA lesions.** To study propagation through interlesion isthmuses, we optically mapped wavefronts of electrical activity between two closely placed RFA lesions (n = 5). Hearts were stained with RH237 and then ablated at two adjacent sites on the left ventricular epicardium. A bipolar pacing electrode was placed on the

![Fig. 1](http://ajpheart.physiology.org/)

**Fig. 1.** Radiofrequency ablation (RFA) lesions in blood-free excised rat hearts. **A:** a radiofrequency (RF) catheter in the position to deliver energy onto the epicardial surface of a rat heart, followed by appearance of two dark RFA lesions as revealed by fluorescent NADH (fNADH) imaging. The image on the right illustrates appearance of the same two RFA lesions after triphenyltetrazolium chloride (TTC) staining. Metabolically active tissue appears red, and irreversibly damaged tissue shows as white. **B:** three sequential snapshots of epicardial fNADH at different time points after ablation using low-magnification settings. **C:** as time progressed, fNADH lesion borders became more homogenous. This is illustrated by high-magnification snapshots of the lesion shown in B at intermediate time points.
epicardium above the RFA lesions, and current was applied at two times the diastolic threshold. An example of spontaneous reentrant circuits around the lesions when a functional isthmus is present is illustrated in Fig. 3. It shows 1) an overlay of epicardial fNADH with an isochronal activation map, 2) optical action potentials, and 3) sequential RH237 images. The full sequence can be seen as an online supplemental movie (The online version of this article contains supplemental material). To create the isochronal maps shown in Fig. 3A, optical action potentials were normalized to show propagating wavefronts in an all-or-none fashion. Normalization is useful for illustrating propagation but obscures the true optical action potential amplitudes. To better represent true optical action potential amplitudes, the RH237 signal at each pixel was scaled as a percentage of the maximum optical action potential amplitude for all the pixels. The interlesion profile of the optical action potential amplitude is shown in Fig. 4A as an x-t plot for five sequential beats, with the x-axis being the distance between the centers of the two lesions. The interlesion profile of action potential amplitude was then compared with the interlesion profile of fNADH intensity (Fig. 4B). The two were highly correlated (Fig. 4, B and C). These findings suggest that fNADH loss can serve as an endogenous live marker for the diminished functional state of the tissue at the ablation site.

Appearance of RFA lesions on RH237-sensitive channel. The epicardial fluorescence of RH237 faded due to washout of the dye (Fig. 5A). However, in ablated tissue, the fluorescence of RH237 declined much slower (Fig. 5B) than in unablated tissue (Fig. 5B). Over time, the background RH237 fluorescence of the lesions was brighter than that of unablated tissue (Fig. 5A, right). The size of these “RH237 lesions” was significantly smaller (by 19 ± 5%, P < 0.001, n = 7) than corresponding dark “fNADH lesions” (Fig. 5C). This matched with the internal ring-like structures sometimes seen on both fNADH and TTC images (Fig. 2A). We believe that RH237 lesions result from acid damage to epicardial capillaries. This may occur at the site of direct resistive heating immediately beneath the electrode and impede the washout of RH237. We further suggest that, by comparing two types of lesions (fNADH vs. RH237), one can distinguish areas of resistive heating from conductive heat transfer (Fig. 5D).

Visualization of RFA lesions in blood-perfused rat hearts. To show the feasibility of fNADH-based imaging in blood-perfused animals, ablation was performed immediately after opening the chest. fNADH images were acquired in the same way as in the excised heart experiments. Major blood vessels appeared as dark tracks within these images. Even so, RFA lesions were clearly evident, indicating that cardiac muscle with its dense supply of mitochondria provides enough fNADH to reveal the unablated tissue that borders a lesion (Fig. 6A). fNADH images became dark when the epicardial surface of the heart was submerged in blood (Fig. 6B). When blood was displaced from the epicardial surface using a sheet of transparent polyvinylidene chloride, RFA lesions were clearly seen. The data shown in Fig. 6 are representative of four experiments.

DISCUSSION

As of today, RFAs are performed with minimal real-time information regarding the physiological state of the ablated tissue. Limited assessments include electrogram amplitude, the
ability to pace and capture ablated tissue, and maneuvers to pace and detect gaps in linear lesions created by multipoint ablations (23). Yet, these assessments provide only limited information regarding the extent and physiological state of the lesion. Second, it is very difficult to determine whether electrical isolation results from acute irreversible muscle damage, functional changes in reversibly injured cells, or from temporary edema. In the case of edema, it may subside after a few weeks, potentially restoring abnormal electrical conduction. The advantage of fNADH imaging is that it reveals irreversible muscle damage without contrast agents, tracers, or dyes. Because NADH is predominantly localized to the mitochondria, fNADH imaging should be insensitive to edema because the interstitial fluid concentration of NADH is negligible. From a clinical perspective, the fact that fNADH lesions are stable for >1 h (Fig. 1B) suggests that lesions could be examined or reexamined at almost any time during an ablation procedure.

The final size of an RF lesion is dependent upon the magnitude and duration of the applied energy. Our ablation parameters produced lesions having an average diameter of 5 mm, similar to those created in the clinic. Our imaging system provided images that revealed interlesion gaps as small as 0.5 mm (see Fig. 2). Smaller lesions could be revealed using higher-magnification lenses or a camera with higher spatial resolution.

Abundant mitochondria make cardiac myocytes particularly suitable for fNADH imaging. Reduced fNADH at the site of the RFA lesions indicates a loss of myocyte membrane integrity, since cell and mitochondrial membranes are rapidly damaged by thermal stress. Notably, cardiac muscle cell necrosis within the ablation site does not necessarily mean that the integrity of all underlying structures, such as coronary vessels, is destroyed. In our experiments, we did not observe the disruption of major coronary vessel structure. This is because, if vessels were disrupted, then tissue downstream of damaged vessels would become ischemic, causing fNADH to increase (17). Yet, fNADH levels near the lesions did not change significantly before and after ablation. Other evidence of intact coronary structure was the homogeneity of postablation TTC staining: any major vessel damage would have been indicated as areas of unstained tissue outside the RFA lesion. However, all RFA lesions stained with TTC were localized strictly to the RF lesion site (Figs. 1 and 2). Finally, observation of intact vessels on the epicardial surface did not indicate severe damage to major vessels at the ablation sites.

One may perceive a contradiction between the reported increase in fNADH during ischemic injury (17) and fNADH decrease upon thermal damage. This apparent contradiction is explained as follows. About 30% of cardiomyocyte volume is comprised of mitochondria that contain a large amount of NADH. Because of this, changes in the level of fNADH from myocytes can be measured with relative ease. When the sarcolemma and mitochondrial membranes are disrupted by heat, NADH is lost, and fNADH levels immediately fall. During
hypoxia and/or ischemia, cellular integrity is preserved, but oxygen availability is reduced. Oxygen serves as a final electron acceptor in the mitochondrial electron chain, and its decline leads to NADH accumulation (5). Thus, ischemia causes an increase in fNADH in a time-dependent manner (24, 32). For example, if coronary perfusion is disrupted before the ablation, patches of ischemic tissue with elevated fNADH levels may be observed adjacent to the darker circular fNADH lesions. Therefore, one concludes that acute tissue ischemia is, in fact, helpful in visualizing RFA lesions via fNADH due to increased contrast between the lesion and the surrounding tissue. A situation involving a healed myocardial infarction is different. Scar contains a significant amount of collagen and as such can contribute to background fluorescence in a spectral range similar to fNADH. Choosing more selective filters and/or the use of ratiometric approaches should be able to distinguish the two, but additional experiments are required to validate these predictions.

As mentioned earlier, one of the main advantages of monitoring endogenous fNADH is that it can be done without additional probes or contrast agents. Because changes in fluorescence reflect acute biochemical changes, lesions are seen almost immediately. Imaging modalities such as magnetic resonance imaging (MRI) (8, 20), C-arm computed tomography (CT) (12), and contrast echocardiography (18) are excellent tools in detecting parameters resulting from heat-induced biophysical changes. However, contrast agents are required to visualize changes in real time. Although MRI and C-arm CT provide high spatial resolution, it could take up to 30 min to visualize cell necrosis. Echocardiography is faster but suffers from limited spatial resolution and limited field of view. Other modalities based on physical tissue changes, including alterations in tissue elasticity, impedance, or absorption, have also been explored (1, 7, 10, 31). While such strategies provide real-time feedback and may predict lesion size and depth, they also require significant data processing and do not provide direct visualization of the ablated region.

Today a majority of RFA procedures are endocardial, but ~10–20% are epicardial (4, 6, 13, 30). Epicardial substrates are frequently observed for VT, including >20% of postinfarct VTs and >30% of VTs from nonischemic cardiomyopathy, particularly for Chagas disease (14, 30). RFA of these epicardial substrates may use a percutaneous approach that involves the subxiphoid placement of sheaths into an intact, closed pericardial space (30). fNADH imaging seems to be particularly feasible for these procedures. Conventional endoscopes equipped with UV-compatible optics and image capture devices would be suitable for this purpose. Air insufflation
through the endoscope could be used to expand the pericardial space for adequate visualization of ablation sites. In a clinical setting, insufflation with carbon dioxide rather than air would likely reduce the risk of air embolization (25). Most importantly, our findings shown in Fig. 6 show that fNADH imaging is feasible in endocardial procedures. The latter is possible if blood is displaced in front of an endoscope, either by inflatable balloons (9, 19) or via carbon dioxide insufflation mentioned earlier.

Additional studies are required to determine whether fNADH imaging can be used to visualize RFA lesions in the atria, i.e., for the treatment of atrial fibrillation. Trabeculation, the intricate geometry of pulmonary veins, the presence of collagen layers, and other structural heterogeneities make fNADH-based visualization of atrial RFA lesions more complex compared with epicardial lesions.

The method described in this manuscript is a simple, straightforward way to monitor acute myocardial damage while performing RFA. If implemented clinically, it has the potential to shorten the time and improve the efficiency of ablations, minimize unnecessary tissue injury that may cause post-RFA complications, and decrease postablation recurrence of arrhythmias. fNADH imaging may also be useful for mechanistic studies of tissue injury near the RFA sites (28) and for assessment of drugs that may alter electrical propagation between interlesion gaps (27). We also believe that NADH fluorescence could be potentially very useful in identifying the ablation gaps seen after pulmonary vein isolation procedures. Last, loss of NADH fluorescence should occur regardless of energy source that leads to tissue necrosis, making this approach applicable to visualization of cryo or laser-based catheter lesions.

This study has several limitations. The feasibility of the technique was demonstrated using open-chest small animal models and not in more clinically relevant large animal models using percutaneous epicardial access; the feasibility of the technique was demonstrated only for epicardial lesions and not for endocardial lesions; and the method is limited to the
visualization of the surface injury. Electrically active gaps present below the epicardial surface can be missed by this technique.

In conclusion 1) RFA lesions in both blood-free and blood-perfused rat and rabbit hearts can be visualized by imaging endogenous NADH fluorescence, 2) optical action potentials can be imaged simultaneously with the endogenous fluorescence of NADH to study changes in electrical activity and tissue viability around ablation lesions and interlesion gaps, and 3) the data suggest that fNADH imaging could be employed during clinical RFA procedures via dual UV excitation/ emission fiber-optic waveguide located inside a balloon catheter. Such a waveguide system could interface with a three-dimensional mapping system to provide a detailed map of cardiac muscle viability near the catheter.

**GRANTS**

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**DISCLOSURES**

A provisional patent describing the use of NADH imaging for cardiac RF catheters was filed by the George Washington University Technology Transfer Office (#61/537,798).

**AUTHOR CONTRIBUTIONS**


**REFERENCES**


