Visualization of Epicardial Cryoablation Lesions Using Endogenous Tissue Fluorescence

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Background—Percutaneous cryoballoon ablation is a commonly used procedure to treat atrial fibrillation. One of the major limitations of the procedure is the inability to directly visualize tissue damage and functional gaps between the lesions. We seek to develop an approach that will enable real-time visualization of tissue necrosis during cryo- or radiofrequency ablation procedures.

Methods and Results—Cryoablation of either blood-perfused or saline-perfused hearts was associated with a marked decrease in nicotinamide adenine dinucleotide (NADH) fluorescence, leading to a 60% to 70% loss of signal intensity at the lesion site. The total lesion area observed on the NADH channel exhibited a strong correlation with the area identified by triphenyl tetrazolium staining (r=0.89, P<0.001). At physiological temperatures, loss of NADH became visually apparent within 26±8 s after detachment of the cryoprobe from the epicardial surface and plateaued within minutes after which the boundaries of the lesions remained stable for several hours. The loss of electrical activity within the cryoablation site exhibited a close spatial correlation with the loss of NADH (r=0.84±0.06, P<0.001). Cryoablation led to a decrease in diffuse reflectance across the entire visible spectrum, which was in stark contrast to radiofrequency ablation that markedly increased the intensity of reflected light at the lesion sites.

Conclusions—We confirmed the feasibility of using endogenous NADH fluorescence for the real-time visualization of cryoablation lesions in blood-perfused cardiac muscle preparations and revealed similarities and differences between imaging cryo- and radiofrequency ablation lesions when using ultraviolet and visible light illumination. (Circ Arrhythm Electrophysiol. 2014;7:929-937.)

Key Words: ablation • atrial fibrillation arrhythmia

Paroxysmal atrial fibrillation (AF) is thought to be trig-
ergged by focal electrical activity originating mainly within muscle sleeves that extend into the pulmonary veins.1 These focal triggers cause atrial tachycardia, driven by reentran
t electrical activity and rotors, which may then fragment into a multitude of electrical wavelets that are characteris-
tic of AF. Prolonged AF can cause functional alterations in membrane channel expression that further perpetuate AF, a phenom
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membrane channel expression that further perpetuate AF, a phenomenon known as AF begets AF.2 The most common treatment of AF consists of placing ablation lesions in a circular fashion around the ostia of the pulmonary veins to isolate ectopic sources from the rest of the atria.1,3,4 More recently, advanced mapping techniques have allowed the targeting of specific localizations of AF ectopic sources and rotors with some degree of success.5 Whether the surgeon’s goal is complete pulmonary vein isolation with no inter-
lesional gaps or site-targeted ablation, it is critical to know the extent of tissue damage at the site of the ablation. The latter is not a simple function of applied energy; it depends on many factors, including contact between the catheter tip and the tissue, thickness of the myocardium, degree of blood flow around the catheter tip, and presence of fatty tissue and collagen, as well as many other factors. Therefore, it is criti
cal to develop new methods to collect real-time information of the functional state of cardiac tissue whenever an ablation is being performed.

Clinical Perspective on p 937

We have previously reported that a loss of endogenous fluorescence of nicotinamide adenine dinucleotide (fNADH) can serve as an immediate marker of myocardial damage caused by radiofrequency ablation (RFA).6 Cryoablation has recently become an increasingly popular alternative option for ablating abnormal AF sources.7 The main hypothesis of this study is that one can use endogenous NADH fluorescence as a functional marker that allows the direct visualization
of cryoablation lesions in cardiac muscle preparations. We examined the validity of this approach for the real-time visualization of cryoinjury and compared it with fNADH-based visualization of RFA lesions.

Methods

Animal Procedures and General Protocols

All animal experiments were conducted in accordance with the guidelines of the Institutional Animal Care and Use Committee at the George Washington University. Adult Sprague–Dawley rats (200–400 g) of mixed sex were injected with sodium heparin (IP, 500 U/kg) and anesthetized with sodium pentobarbital (IP, 45 mg/kg). For ex vivo studies, the heart was rapidly excised, the aorta cannulated, and the heart Langendorff-perfused with Tyrode’s solution (Sigma T2145). Alternatively, for in vivo studies, the heart was ablated and imaged while still in the chest. Both types of studies included the following steps: (1) expose epicardial surface, (2) position illumination sources to evenly illuminate the surface and record preablation image, (3) perform either cryoablation or RFA, (4) record fNADH postablation images for a specified period of time, (5) switch to room light illumination to record postablation image, and (6) perfuse heart with 2,3,5-tri-phenyl-2H-tetrazolium chloride (TTC; details below) and take a record postablation image.

Ablation Protocol

Cryoablation was performed using either a metal cryoprobe dipped in liquid nitrogen or by using a clinical Freezor cryoablation catheter controlled by a CryoConsole (Medtronic, MN). Cryoablation durations varied from 5 to 30 s (see Table in the Data Supplement for details). When used, the cryoablation catheter was placed perpendicular to the epicardial surface. Radiofrequency energy was delivered using a noncooled blazer catheter with a 4-mm tip (EP Technologies, Boston Scientific). The RFA catheter was placed perpendicular to the epicardial surface. Tip temperatures ranged between 50 and 70°C and ablation durations varied from 5 to 30 s with a maximum power of 8 watts (see Table in the Data Supplement for details). This led to consistent RFA lesions without charring or steam pops.8

NADH Fluorescence Imaging

Since NADH is an endogenous fluorophore, no additional dye is required to visualize its presence in cardiac tissue. To record fNADH, the epicardial surface was illuminated using an LED spotlight with a peak wavelength of 365 nm (PLS-0365-030-S; MightyS systems) unless specified otherwise. The emitted fNADH was bandpass-filtered at 475 nm (Chroma HQ475/50) and imaged using a charge-coupled device camera at 1024×1024 resolution (Andor iKon-M DU934N-BV) fitted with a low magnification lens (Nikon Nikkor 50 mm F/1.4). To illustrate that the presence of blood between tissue and detector interferes with fNADH visualization, the excised heart with ablation lesion was immersed into a blood-filled beaker. Images were taken before and after blood was displaced from the heart surface using a transparent film of polyvinylidine chloride. To obtain the percentage decrease in fNADH intensity across the lesion, values from 3 random pairs of regions of interest from ablated and nearby control tissue were averaged.

Dual NADH and Potentiometric Dye Recordings

A dual imaging approach was used to colocализe changes in the amplitude of epicardial action potentials and ablation-induced loss of endogenous NADH fluorescence. Optical mapping was performed using the potentiometric dye RH237 (N-(4-sulfobutyl)-4-(6-(4-(dibutylamino)phenyl)-hexatrienyl)pyridinium; Molecular Probes). A 50 μg aliquot of RH237 was resuspended in 0.1 mL of dimethyl sulfoxide, vigorously vortexed for 10 minutes, and combined with 3 mL of Tyrode’s solution. The latter was injected into a port at the base of the aortic block that contained an additional 2 mL of perfusate, yielding a 20 μmol/L RH237 concentration. To reduce motion artifacts, the motion uncoupler 2,3-butanediol monoxide was added to the perfusate at a final concentration of 20 mmol/L. A dual optical mapping system comprising of 2 high-frame rate 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 and NADH in the same field of view. fNADH was recorded using filter settings as described earlier. To record optical action potentials (OAPs), the epicardial surface was illuminated using 2 light-emitting diodes (LumiLEDs, 530/35 nm). The resulting RH237 fluorescence was long-pass-filtered at 680 nm. RH237 fluorescence signals were smoothed using a median temporal filter (3 sample width), and the average amplitude of OAPs at each pixel was computed to reveal spatial changes in the amount of electrically active tissue.

Reflectance Spectroscopy Recordings

The epicardial surface of the heart was illuminated using a fiber optic guide connected to a spectrofluorometer (HORIBA Jobin Yvon FluoroMax-3). The illumination wavelength was gradually increased from 400 to 700 nm using a 5 nm slit width. The epicardial surface was continuously imaged throughout the experiment using an Andor iKon-M (DU934N-BV) camera without any filter in the optical path. The full image sequence was then imported into Andor Solis to extract the intensity values from specific regions of interest. Mean values were then normalized to the intensity of the illuminating light by placing a white reflectance standard (Thorlabs, Spectralon SRS-99-010) within a field of view.

Effects of Filter Settings on fNADH-Based Image Visualization

Both RFA and cryoablation lesions were placed next to each other on the epicardial surface of an excised, saline-perfused heart. Illumination through a fiber optic light guide connected to a spectrofluorometer (HORIBA Jobin Yvon FluoroMax-3) was tuned to either 365 or 375 nm using 20 nm slit settings. The images were acquired using an Andor iKon-M (DU934N-BV) camera using either a 475/50 nm or 450/70 nm bandpass filters. Three illumination/bandpass filter combinations were then tested: 375/475, 375/450, and 365/475 nm. The spectral profiles of the bandpass filters were provided by the manufacturers. The spectral profiles of the illuminating light was obtained by directing the light guide toward a white standard and recording the reflected light spectrum using the spectrofluorometer.

TTC Staining

TTC staining is a standard procedure for assessing necrosis. It relies on the ability of dehydrogenase enzymes and NADH to react with tetrazolium salts to form a formazan pigment. Immediately after the imaging protocol, the tissue was Langendorff-perfused with Tyrode’s solution containing 40 mmol/L TTC. Then the heart was submerged in liquid nitrogen or by using a clinical Freezor cryoablation catheter controlled by a CryoConsole (Medtronic, MN). Cryoablation was performed using either a metal cryoprobe dipped in liquid nitrogen or by using a clinical Freezor cryoablation catheter controlled by a CryoConsole (Medtronic, MN). Cryoablation durations varied from 5 to 30 s (see Table in the Data Supplement for details). When used, the cryoablation catheter was placed perpendicular to the epicardial surface. Radiofrequency energy was delivered using a noncooled blazer catheter with a 4-mm tip (EP Technologies, Boston Scientific). The RFA catheter was placed perpendicular to the epicardial surface. Tip temperatures ranged between 50 and 70°C and ablation durations varied from 5 to 30 s with a maximum power of 8 watts (see Table in the Data Supplement for details). This led to consistent RFA lesions without charring or steam pops.8

Statistical Analysis

In total, 42 cryoleisions and 12 RFA lesions in 22 rat hearts were analyzed to derive the quantitative and qualitative conclusions detailed in this article. The Table in the Data Supplement lists the details and the total number of each experiments from which qualitative conclusions and quantitative results were drawn. Values are presented as mean±SD, except noted otherwise. The Wilcoxon signed-rank test was used to compare the differences in mean values from ablated and control regions in the same animal. Spearman’s correlation...
Results

Appearance of Cryoablation Lesions in Saline- and Blood-Perfused Rat Hearts

In both saline-perfused and blood-perfused hearts, cryolesions were barely visible under incandescent illumination with only a small drop in signal intensity across the lesion (14%±4% in saline versus 16%±5% in blood). However, by using ultraviolet illumination (350–370 nm) and collecting the emitted fluorescence within 450 to 500 nm range, we observed distinctly demarcated lesions in both blood-free and blood-perfused hearts, including in vivo beating hearts (Figure 1A and 1B, see right panels). The magnitude of the drop in fNADH signal intensity in blood-perfused preparations and blood-free preparations was the same (65%±7% in blood-perfused versus 67%±8% in saline-perfused). Thus, the presence of the blood within the tissue did not interfere with visualization of cryolesions on the fNADH channel. This assumes, of course, that optically dense blood is displaced from the path between the camera and the heart’s surface (Figure 1C).

To show the clinical applicability of fNADH-based cryolesion visualization, we confirmed our findings using a Medtronic Freezor catheter (Figure 2A) to place cryoablation lesions on the endocardial surface of the heart (Figure 2B). In both cases, cryolesions were readily distinguishable on the fNADH channel with the loss of ≈70% signal intensity at the lesion sites (69%±2% and 71%±3% respectively).

Time Course of Lesion Formation

At the cryoinjury site, there was an immediate increase in signal intensity as tissue reflectance increased dramatically because of formation of ice crystals. As the tissue thawed, the reflectance fell below preablation levels and a gradual loss of fNADH took place. Once the lesion was fully developed, it remained stable throughout the duration of the experiment. This is illustrated in Figures 3 and 4. Figure 3 shows individual snapshots at different points in time and fNADH traces across the lesion boundary during

![Figure 1. Visualization of cryolesions in blood-free and blood-perfused preparations. A, Saline-perfused excised rat heart under incandescent room light and under 365 nm illumination/475 nm acquisition settings. Intensity profiles across dotted white lines for both images are shown in the right. Scale bar 2 mm. B, Color snapshot of cryoablated epicardial surface as seen in an open-chest animal under incandescent illumination. The same heart under 365/475 nm settings showing 2 lesions: 1 ventral and 1 apical. Dot plot on the right compares percent change in signal intensity under the 2 acquisition settings (n=7, P<0.05; scale bar 2 mm) C, When the ablated heart was immersed into a blood-filled beaker, the blood blocked the fluorescence of nicotinamide adenine dinucleotide (fNADH) signal. However, displacement of blood using a transparent film enabled fNADH-based lesion visualization, pointing to the need for a transparent inflatable balloon to be included in future designs of fNADH imaging catheter.](http://circep.ahajournals.org/)

![Figure 2. Visualization of cryolesions made with a clinical catheter and on the endocardial surface. A, Color snapshot of Freezor cryoablation catheter controlled by CryoConsole (Medtronic, MN) next to the cryoablated surface of a saline-perfused excised rat heart (Left; incandescent room light). The cryolesions as they seem on both sides of epicardial surface using fluorescence of nicotinamide adenine dinucleotide (fNADH) imaging settings (Right). Scale bar 2 mm. B, fNADH-based visualization of cryolesions on endocardial surfaces of saline-perfused excised rat ventricle. Scale bar 3 mm. LV indicates left ventricle; and RV, right ventricle.](http://circep.ahajournals.org/)
and immediately after cryoablation. Note again the initial whitening of the lesion, followed by a progressive decline in fNADH intensity. As a result, the difference in fNADH intensity values across the border of the lesion became more and more pronounced, yet the border location stayed static (Figure 3B). Figure 4 compares the time course of cryoablation lesion appearance to that of the RFA lesion. During the RFA procedure, the signal started to decline during the ablation itself. In contrast, the site of cryolesion exhibited the abovementioned increase in signal intensity followed by a rapid loss of fNADH. Notably, when the perfusate is maintained at physiological temperatures, cryolesions became visually apparent on fNADH channel within 26±8 s, making fNADH-based visualization applicable to real-time surgical monitoring of ablation efficacy.

**Correlation of fNADH Loss With Necrotic Tissue Identified Using TTC Staining**

After completion of imaging experiments, the cryoablated hearts were perfused with the vital dye TTC (Figure 5).
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Figure 5. Quantitative comparison between the fluorescence of nicotinamide adenine dinucleotide (fNADH) and the respective 2,3,5-triphenyl-2H-tetrazolium chloride (TTC) staining in cryoablation lesions. Representative pair of fNADH and TTC images and correlation analysis for the total lesion area (n=10).

The latter is a standard staining technique used to distinguish between live and necrotic tissue. The surface area of the TTC-negative tissue directly correlated with the cryolesion surface area as seen on the fNADH channel (r=0.89, P<0.05). Moreover, when the cryoablated heart was TTC-stained and sliced transmurally, the cross-sectional profile of the lesion exhibited strong correlation with the depth profile derived from the epicardial fNADH image (r=0.76, P<0.001). Indeed, although visually the lesions appear to be a uniform grey, when the fNADH intensity values are displayed as a 3D wireframe, it enables direct visualization of lesion depth (Figure 6).

Loss of Electrical Activity at the Cryoablation Lesion Site

The primary reason to ablate viable tissue is to prevent the propagation of electrical activity. We used the voltage sensitive dye RH237 to reveal how much electrically active tissue remains at the ablated site and how this correlates with the loss of endogenous NADH. Figure 7 shows 2 examples of hearts with multiple cryolesions for which dual recordings of fNADH and transmembrane voltage were performed. A change in the RH237 signal intensity corresponds to a change in the transmembrane voltage, yielding OAP amplitude image. Representative OAPs from regions of interests corresponding to unablated tissue, lesion sites, and interlesional gaps are shown on the left. Statistical comparison of the spatial profiles of fNADH loss across the lesions to the amplitude of OAP revealed a close correlation between the 2 (P<0.001; Spearman’s correlation coefficient for each lesion is shown above the 2 profiles). The decrease in OAP amplitude is an indication of less viable tissue contributing to the changes in RH237 intensity. Similarly to fNADH loss, it can serve as an indirect measure of lesion depth.
Changes in Tissue Reflectance at the RFA and Cryoablation Lesion Sites

To quantify wavelength-specific changes in photon absorption and scattering, the entire surface of the heart with both types of lesions was illuminated using increasing wavelengths of light, and the images were acquired without any filter in the optical path. The amount of reflected light from cryoablation sites was significantly lower, whereas that from the RFA sites was significantly higher for each tested wavelength within 400 to 700 nm range of visible light (n=5, P<0.05). This is shown in Figure 8A and 8B using representative images and traces from an individual experiment. Figure 8D displays an average fold change in intensity values collected from ablated and unablated tissue. The latter enables a better appreciation of wavelength-specific changes in tissue reflectance across visible spectra.
Effect of Filter Settings on fNADH-Based Visualization of RFA and Cryoablation Lesions

For RFA lesions visualization, the dramatic increase in tissue reflectance may obscure the loss of NADH fluorescence. This can happen when there is an overlap between the wavelengths of the excitation light and the light passing through the emission filter. This is less of a concern for cryolesions, since the amount of both types of photons, fluoresced and scattered, decreases. To illustrate this effect, we conducted an experiment in which we used 3 different combinations of illumination and acquisition settings to record images from the same heart that had both types of lesions on its epicardial surface (Figure 9). The RFA lesion disappeared (4%±7% drop in signal intensity, insignificant from surrounding unablated tissue) when imaged using the filter set that allowed for a small spectral overlap between the illuminating and collected light. In contrast, at the site of the cryoablation lesion, the decrease in fNADH remained substantial under all 3 illumination/filter combinations (69%±5%, 55%±9%, and 71%±5% drop in signal intensity at 375/475, 375/450, and 365/475 nm settings, respectively). The above information can be useful for implementation of fNADH-based lesion visualization in designs of future imaging catheters.

Discussion

Pulmonary vein isolation with catheter ablation is an effective treatment in patients with symptomatic AF that is refractory or intolerant to antiarrhythmic medications. Cryoballoon catheters have been approved for isolating pulmonary veins in patients with paroxysmal AF.10 After thawing, cryoablation causes necrosis without drastic alterations in tissue structure.11 When compared with RFA, it takes less time to accomplish a cryoballoon ablation procedure, and the lesions created are more homogenous with lower incidence of destruction of the surrounding vasculature.12 Cryoablation is also known to be less thrombogenic and lead to more complete fibrosis of the lesion with better demarcated boundaries.12,13

The most common site of failed cryoballoon ablation is the inferior area of the veins where the atrial muscle sleeves extend into the pulmonary vein.14 If occlusion of the vein by the balloon is not complete, a small blood leak around the balloon can lead to a gap in the ablation line. The latter is formed at the site of the contact between the cryoballoon and the orifice of the vein. Interlesional gaps can then lead to AF recurrence because of pulmonary vein reconnections.15–17 Therefore, the ability to visually inspect the integrity of the ablation lesion border and to address remaining gaps while performing the ablation procedure would be highly beneficial.

The main goal of this study was to demonstrate the feasibility of fNADH imaging to reveal gaps between the cryolesions. The second goal was to compare and contrast fNADH-based visualization of cryolesions to that of RFA lesions. Such information can be useful for designing future surgical catheters that incorporate fNADH-based monitoring of cardiac tissue viability.

The similarities between cryoablation and RFA lesions included a high degree of correlation between the shape of the lesion observed on the fNADH channel versus the actual necrotic area. The latter was identified using TTC, a standard marker of tissue necrosis, which turns viable tissue crimson red and leaves the necrotic area white. Notably, a close correlation between the area as seen on the fNADH channel and after TTC staining was observed for both surface and transmural measurements of the lesions (Figures 5 and 6). Another similarity was stability of the lesion border as a function of time (Figure 3). Finally, there was a direct correlation between the immediate loss of electrical activity and the fall in the fNADH intensity (Figure 7), similar to what we observed for RFA lesions.6

Yet, there were 2 important differences. The first was the difference in the initial timeline of fNADH changes (Figure 4). At the RFA site, fluorescence decreased during the ablation itself. In contrast, at the cryoinjury site there was a transitory increase in signal intensity as diffuse reflectance rose dramatically because of the formation of ice crystals. This was followed by thawing and then fNADH loss at the cryoinjury site. The second major difference between the 2 types of ablation lesions is changes in tissue optical properties (Figure 8). RFA leads to a dramatic increase in tissue scattering. In contrast, across the entire visible spectrum the overall amount of light reflected from the cryolesion site is lower than that from surrounding unablated tissue, making cryolesions almost invisible to the naked eye, particularly in blood-perfused preparations. Therefore, the use of ultraviolet illumination to reveal otherwise invisible cryolesions is particularly appealing.

After defrosting, the sites of cryolesions always looked slightly darker in both blood-perfused and blood-free hearts. The later observation is important because most of the darkening seen during cancer cryoresection is ascribed to hemorrhage. But because in our studies darkening was observed in both blood- and saline-perfused hearts, hemorrhage is an unlikely explanation. The main factor is a loss of highly organized myocyte fibrils resulting from freeze/thaw process. The latter has been shown to significantly decrease the scattering coefficient of cardiac muscle.18 Additionally, changes in hue can be ascribed to changes in myoglobin absorption spectra. Myoglobin is 1 of the major myocardial chromophores within the visible range, and its absorption spectra and extinction coefficient depends on whether it is in its oxy-, met-, or carboxymyoglobin form.19,20 In live cells, metmyoglobin reductase converts metmyoglobin (using NADH as 1 of the cofactor) to myoglobin.21 Upon cell necrosis, this process is disrupted, leading to a net accumulation of metmyoglobin that gives tissue a more brown hue. Finally, the tissue pH can also affect myoglobin’s extinction coefficient, and the pH within necrotic tissue is likely to be rapidly changing.

What are the advantages of fNADH imaging as compared with other methods? Indeed imaging modalities, such as MRI,22,23 C-arm computed tomography,24 and contrast echocardiography,25 are excellent tools in detecting
parameters resulting from thermally induced physical changes. However, they all require contrast agents and have relatively poor resolution for the purpose of identifying lesions. Second, although a drop in fNADH occurs within seconds, MRI and C-arm computed tomography could take ≤30 minutes to visualize cell necrosis. Echocardiography is faster but it suffers from low spatial resolution and a limited field of view. In addition, MRI, C-arm computed tomography, and contrast echocardiography also require significant data processing. Recently, direct visualization of endocardial lesions has become possible. Newer catheters from CardioFocus and Voyage Medical displace blood using an inflatable balloon and catheter shroud, respectively, and both use high-intensity white light illumination to collect reflected and scattered light from the tissue. But because the endocardial surface of the atrium in humans has a significant collagen layer and is white in appearance, the contrast between the cryoablated and unablated tissue is expected to be minimal when direct white light illumination is used. In comparison, use of endogenous tissue fluorescence may provide greater contrast between unablated and ablated tissue.

Limitations

Applicability to Large Animals

Our preliminary data from placing cryoablation lesions on the endocardial surface of canine and porcine atria suggest that fNADH can enable visualization of cryolesions in fresh atrial tissue. More systematic studies, beyond the scope of this study, need to be performed to fully determine the applicability of this approach in large animals and humans.

Lesion Depth

The loss of intensity on fNADH images exhibit strong correlation with cryolesion depth, ≤2 mm of tissue thickness based on our preliminary estimates. Since the reported thickness of human atria is ≈1.5 to 2.5 mm, fNADH imaging should provide useful information about lesion depth, although it might not be able to confirm lesion tranmsurality at all the sites.

Summary

We present the similarities and differences of fNADH-based visualization of cryoablation and RFA lesions. NADH is an endogenous molecule predominantly present in mitochondria that fluoresces when illuminated with UVA light. Because of the abundance of mitochondria in cardiac muscle, NADH loss provides sufficient contrast to delineate the lesion and provides functional information about the ablation site. The main goal of this technique is to enable real-time identification of ablation gaps to eliminate them during the ablation procedure. This should enhance procedural success and lead to lower AF recurrence rates. Integration of this imaging approach into a percutaneous catheter can allow in-surgery monitoring of the endocardial surface of the pulmonary vein area, reveal interlesional gaps, estimate the depth of lesions, and provide information about the functional state of the ablation lesions based on endogenous NADH fluorescence.

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Disclosures

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Atrial fibrillation remains the most commonly occurring cardiac arrhythmia. It is associated with a lower quality of life and a higher rate of morbidity and mortality. Radiofrequency ablation via a percutaneous catheter is a mainstream treatment for atrial fibrillation. More recently, cryoablation has become widely accepted in the electrophysiology community as a safe and effective way to create thermal ablation lesions in cardiac tissue. Both methods produce lesions that block the propagation of electrical activity from sites of abnormal activity, including the pulmonary veins. Today, there are limited options for real-time monitoring of tissue injury during cardiac ablation procedures and even more limited options for direct visualization of lesions. To address this need, we studied the fluorescence of endogenous nicotinamide adenine dinucleotide (NADH), present in all metabolically active cells, as a biomarker of tissue necrosis. We confirmed the feasibility of NADH-based visualization for identification of both radiofrequency ablation and cryoablation lesions. We showed that the loss of NADH closely correlates with the decline in electrical activity, allowing detection of interlesional gaps. We also revealed some significant differences between the optical properties of these 2 types of thermal lesions and how they can affect NADH-based visualization. The ultimate goal of our studies is to help create either a standalone visualization catheter or an ablation catheter that incorporates visualization capabilities. New imaging technologies, such as NADH-based lesion visualization, will enable surgeons to see the ablated areas in real time, paving the way for easier, faster, safer, more cost-effective, and more reliable atrial fibrillation ablation procedures.