High-intensity ultrasound catheter ablation achieves deep mid-myocardial lesions in vivo

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BACKGROUND Radiofrequency ablation of epicardial and mid-myocardial ventricular arrhythmias is limited by lesion depth.

OBJECTIVE The purpose of this study was to generate deep mid-interventricular septal (IVS) lesions using high-intensity ultrasound (US) from an endocardial catheter-based approach.

METHODS Irrigated US catheters (12 F) were fabricated with 3 × 5 mm transducers of 5.0, 6.5, and 8.0 MHz frequencies and compared in an ex vivo perfused myocardial ablation model. In vivo septal ablation in swine (n = 12) was performed via femoral venous access to the right ventricle. Lesions were characterized by echocardiography, cardiac magnetic resonance imaging, and electroanatomic voltage mapping pre- and post-ablation, and at 30 days. Four animals were euthanized immediately post-ablation to compare acute and chronic lesion histology and gross pathology.

RESULTS In ex vivo models, maximal lesion depth and volume was achieved by 6.5 MHz catheters, which were used in vivo. Lesion depth by gross pathology was similar post-ablation (10.8 mm; 95% confidence interval [CI] 9.9–12.4 mm) and at 30 days (11.2 mm; 95% CI 10.6–12.4 mm) (P = .56). Lesion volume decreased post-ablation to 30 days (from 255 [95% CI 198–440] to 162 [95% CI 133–234] mm³; P = .05), yet transmurality increased from 58% (95% CI 50%–76%) to 81% (95% CI 74%–93%), attributable to a reduction in IVS thickness (from 16.0 ± 1.7 to 10.6 ± 2.4 mm; P = .007). Magnetic resonance imaging confirmed dense septal ablation by delayed enhancement, with increased T1 time post-ablation and at 30 days and increased T2 time only post-ablation. Voltage mapping of both sides of IVS demonstrated reduced unipolar (but not bipolar) voltage along the IVS.

CONCLUSION High-intensity US catheter ablation may be an effective treatment of mid-myocardial or epicardial ventricular arrhythmias from an endocardial approach.

KEYWORDS Catheter ablation; Mid-myocardial; Ultrasound; Ventricular arrhythmia; Ventricular tachycardia

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to the His-Purkinje system when performing ablation of the IVS from an RV approach.

In this study, we designed, fabricated, acoustically characterized, optimized, and tested HIU ablation catheters suitable for ablation of the IVS from a femoral venous approach to the RV. We hypothesized that this approach could produce deep mid-myocardial lesions without His-Purkinje injury and iatrogenic atrioventricular (AV) block.

Methods

Catheter assembly and acoustic testing

Side-facing HIU ablation catheters were fabricated from piezoelectric crystals, electrical cable, modified stainless steel tubing, nylon tubing, and electrical connectors (Figure 1A). Each catheter contained a planar 5.0, 6.5, and 8.0 MHz single-element PZT-4 piezoelectric crystal 3 × 5 mm in dimension (EBL Products Inc., East Hartford, CT), which was soldered to 34 AWG micro coaxial electrical cable (AlphaWire, Elizabeth, NJ) and a BNC connector (Amphenol, Wallingford, CT). Edges of the US crystal were bonded to a partially cut stainless steel hypodermic tubing that served as a rigid transducer platform with air backing. The transducer platform assembly was then bonded to a 12-F flexible nylon catheter tubing (Frellein-Wade Inc., McMinnville, OR). The transducer assemblies were encapsulated within a thin-walled polyester balloon through which closed-loop internal irrigation was performed with deionized and degassed water at 10°C with a flow rate of 50 mL/min supplied by a peristaltic pump. Finally, impedance matching networks were built to achieve >99% electrical power transmission.

Catheters were operated in the continuous wave mode using signals produced by a waveform generator (Agilent 33250A, Santa Clara, CA), amplified by a power amplifier (Amplifier Research 600A225, Souderton, PA), and monitored using a high-voltage passive detection US probe (LeCroy PPE 1.2 kV, Chestnut Ridge, NY) connected to an oscilloscope (LeCroy Waverunner 44 MXi-A, Chestnut Ridge, NY). After fabrication, the US pressure field for each catheter was characterized in filtered, deionized, and degassed water at room temperature using a needle hydrophone (75 μm, Precision Acoustics, Dorchester, UK) to map peak negative acoustic pressures (Figure 1B-C).

Ex vivo myocardial ablation protocol

Ovine hearts were obtained from animals receiving intravenous heparin immediately before killing. Hearts were studied within 30 minutes of explantation. A 5-F perfusion catheter was sutured in the proximal left anterior descending coronary artery (LAD) to perfuse the anterior myocardium. The heart was suspended in a degassed water bath at room temperature. A solution of 1% bovine serum albumin in normal saline was perfused through the LAD at 30 mL/min using a non-peristaltic syringe pump. Epicardial HIU lesions were applied at a power of 30 W for 60 seconds. Eight separate lesions were made in the LAD perfusion territory: one anteroseptal line of 4 extending from the basal to the apical anteroseptum 1 cm lateral and parallel to the LAD and a second anterior line of 4 just 1 cm lateral to the first line. US frequency (n = 32 sonifications for each frequency) was randomized on a per-heart basis.

In vivo myocardial ablation protocol

This study was approved and monitored by the Oregon Health & Science University Institutional Animal Care and Use Committee under guidelines set forth by the Association for Assessment and Accreditation of Laboratory Animal Care International and consistent with the Guide for the Care and Use of Laboratory Animals (National Research Council). Female farm swine (Sus domesticus) (40–45 kg) were anesthetized with tiletamine and zolazepam and intubated. Anesthesia was maintained with inhaled isoflurane (1.5%–3.5% via 2 L/min O2). Isothermia was maintained with water-heated blankets. The HIU catheter was advanced via femoral venous access through a 14-F steerable sheath to the RV and guided to the RV septum using fluoroscopic and transthoracic echocardiography (TTE) guidance. Eight soninations at a power of 30 W were performed for 60 seconds, approximately 1 cm apart in a × 2 pattern with 4 lesions extending from the basal to mid-anterosesum and a second row from the basal to mid-inferoseptum. Continuous electrocardiographic monitoring was performed during the procedure, with attention to AV conduction, QRS width, and morphology. Four swine were euthanized immediately post-ablation per protocol for the assessment of acute lesions. The remainder were survived for 30 days.

Imaging and electroanatomic mapping

TTE (Epiq CV, Philips Ultrasound, Andover, MA) was performed with a phased array probe and tissue harmonic filtering pre-ablation, immediately post-ablation, and, when applicable, at 30 days. Parasternal short-axis views at the basal and mid-ventricular planes and apical 4-chamber views were obtained at a transmit frequency of 1.6 MHz. A subset of swine underwent cardiac magnetic resonance imaging (MRI) preablation (n = 4), immediately post-ablation (n = 6), and at 30 days (n = 6). MRI was performed on a 3.0 T Siemens Prisma system (Erlangen, Germany) using the body coil for RF transmitting as well as spine and body matrix coils for signal receiving and with electrocardiographic gating. After standard T1- (spin-lattice relaxation time) and T2- (spin-spin relaxation time) weighted scans, 0.1 mmol/kg of ProHance gadolinium contrast agent (Bracco, Milan, Italy) was administered intravenously. First-pass perfusion as well as T1-weighted delayed enhanced images were acquired ~20 minutes postinjection with phase-sensitive inversion recovery reconstruction. MRI analysis was performed using dedicated MRI software (Circle Cardiovascular Imaging, Calgary, Canada). The number of lesions and their locations were first assessed on short-axis slices of delayed enhancement images. The corresponding regions on the native (precontrast) T1 and T2 maps were sampled for T1 and T2 times at lesion locations. We used MODified Look Locker Inversion recovery (MOLLI) sequences to measure T1. These measurements were also...
normalized to their respective T1 and T2 times directly opposite in the non-ablated, inferolateral LV myocardium to account for variations between swine. T1 and T2 times were made in the middle third of the ventricular myocardium to avoid partial volume effects.

Electroanatomic voltage mapping of the RV and LV septum was also performed in a subset of 4 swine using the PentaRay catheter (Biosense Webster International, Diamond Bar, CA) pre- and post-ablation and at 30 days. Bipolar voltage mapping was assessed at the standard threshold of 0.5 and 1.5 mV and unipolar voltage mapping at 1.0 and 5.5 mV for the RV septum and 1.0 and 8.3 mV for the LV septum.

Gross pathology and histology
For both ex vivo and in vivo studies, the heart was perfused with 10 g of 2,3,5-triphenyl-2H-tetrazolium chloride in 50 mL of saline (through the perfusion catheter for ex vivo studies and via central venous access for in vivo studies) to aid visual inspection of myocardial necrosis. Hearts were fixed in 10% formalin at 4°C for 1 week and then sectioned in 3-mm-thick short-axis segments. Segments containing lesions were further sectioned to measure individual lesion sizes in 3 dimensions and then further fixed and stained with Masson’s trichrome and hematoxylin and eosin.

Lesion depth was measured starting from and orthogonal to the RV endocardium and ending at the distal (toward the LV) end of the lesion (Figure 2B). RV subendocardial sparing measurement also started at the RV endocardium, but extended toward the proximal (toward the RV) end of the lesion (Figure 2B). Lesion volume was calculated using the visual observation of lesion boundaries as input to the revolved ellipsoid volume formula (4/3πL/2W/2T/2), where L is the lesion length (measured in the axis going away from the transducer from the RV to the LV endocardium), W is the width (from the antero- to the inferoseptum), and T is the thickness (along the basal-to-apical axis). This equation has been validated for the assessment of ablation lesions.10 Lesion transmurality (defined as percentage of the IVS thickness bounded by the lesion at its cross-section) and near-field (RV side of the mid-septum) vs far-field (LV side of the mid-septum) deposition of necrosis was quantified by determining the IVS midline with a custom image processing algorithm using spline analysis (Online Supplement).

For histological analysis of necrosis, inflammatory infiltrate, hemorrhage, and fibrosis, lesion blocks were further sectioned at a thickness of 10 μm and stained with hematoxylin and eosin and Masson’s trichrome.

Statistical analysis
Data were analyzed using GraphPad Prism software (version 8.4.1; San Diego, CA) and SAS (version 9.4; Cary, NC). Groupwise differences were assessed using the
Mann-Whitney \( U \) test for data that were determined to be nonnormally distributed. For data with normal distribution, groupwise differences were assessed using the Student \( t \) test (paired for comparisons at different time points) or analysis of variance (ANOVA). These data are presented as mean \( \pm \) SD. A 1-way analysis of variance test was used when comparing multiple groups per time point. One survival swine died at 14 days. Gross pathological data from this swine were included with the 30-day survival swine. When comparing repeated measures (multiple ablation lesions per animal) among multiple swine, fixed effects regression models were fit and SEs adjusted to address the correlation among measurements from the same animal. Means with bootstrapped 95% confidence intervals (CIs) based on 500 replications are reported from these models. All statistical tests were 2-sided, with \( P \leq .05 \) considered significant.

**Results**

**Comparison of US frequencies**

The acoustic fields of HIU catheters with fundamental frequencies of 5.0, 6.5, and 8.0 MHz were mapped (Figure 1) and predicted that 6.5 MHz would generate the largest and deepest acoustic field. Ablation at 5 MHz did not reproducibly result in necrotic lesions (32 sonications yielded only 4 identifiable small lesions on gross pathology), likely because of the reduced thermal effects of US at lower frequencies having fewer harmonics formed during nonlinear acoustic propagation.\(^{11,12}\) Ablation at 6.5 MHz generated lesions that were deeper (\( 10.7 \pm 3.2 \) mm vs \( 8.3 \pm 2.6 \) mm from the RV endocardium; \( P = .0015 \)) and larger in volume (\( 223 \pm 171 \) mm\(^3 \) vs \( 137 \pm 95 \) mm\(^3 \) in volume; \( P = .028 \)) than at 8 MHz (Online Supplemental Figure 1). Sparing of the RV subendocardial surface tissue immediately adjacent to the transducer was similar between both frequencies (1.6 \( \pm \) 1.2 mm vs 1.5 \( \pm \) 0.6 mm distance from the RV endocardial surface to ablation lesion; \( P = .797 \)). To increase the study efficiency, in vivo studies were performed only with 6.5 MHz HIU catheters.

**In vivo ablation: Pathological analysis**

In the 4 animals euthanized after immediate post-ablation imaging, histology detected 24 necrotic lesions out of 32 HIU ablation attempts (75% efficiency); while 42 necrotic lesions were found out of 64 HIU ablation attempts (66% efficiency) in survival animals. Sonication locations that were more apically located were least likely to result in visible lesions on gross pathology, largely because of the inability of the custom steerable sheath to reach the apical septum. HIU lesions were typically ellipsoid in shape (Figure 2), similar to prior descriptions of HIU epicardial LV lesions.\(^9\) The mean lesion depth was similar immediately post-ablation (10.8 mm; 95% CI 9.9–12.4 mm) and at 30 days (11.2 mm; 95% CI 10.6–12.4 mm) \( (P = .56) \), although lesion volume decreased over time (from 255 [95% CI 198–440] to 162 [95% CI 133–234] mm\(^3 \); \( P = .05 \)) (Figure 2). The RV subendocardial surface was similarly spared at both time points (2.2 mm; 95% CI 1.7–3.3 mm vs 2.7 mm; 95% CI 2.3–3.4 mm; \( P = .3 \)). Similar to lesion depth, lesion length (L, going away from the transducer from the RV to the LV endocardium and excluding RV subendocardial sparing) was similar at both

\begin{figure}
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\includegraphics[width=\textwidth]{figure2.png}
\caption{In vivo myocardial lesion characteristics. A: Short-axis gross pathology of 2 adjacent lesions in the interventricular septum (IVS). B: Magnification of IVS demonstrating lesion measurements (D, depth; S, subendocardial surface sparing; L, length; W, width; T, thickness is not shown, but was measured by sectioning serial short-axis cross-sections). C: Quantification of gross pathological lesion characteristics. Graph bars represent means, and error bars represent SD. RV = right ventricular.}
\end{figure}
time points. However, lesion width (W) decreased from 6.4 (95% CI 5.9–7.0) to 5.5 (95% CI 5.0–6.1) mm ($P = .006$) and lesion thickness (T) decreased from 7.8 (95% CI 7.2–8.4) to 6.2 (95% CI 5.6–6.8) mm at 30 days ($P < .0001$).

Based on spline analysis image processing, a similar portion of the lesion surface area was deposited on the LV side of the IVS (30%; 95% CI 22%–46% post-ablation vs 39%; 95% CI 33%–51% at 30 days; $P = .29$) (Figure 3). Spline analysis showed that lesion transmurality increased from 58% (95% CI 50%–76%) to 81% (95% CI 74%–93%) over 30 days ($P = .002$). Increased transmurality despite decreased lesion volume at 30 days compared with immediately post-ablation was attributable to IVS remodeling resulting in reduced thickness: IVS thickness at the level of the lesions decreased from 16.0 ± 1.7 mm post-ablation to 10.6 ± 2.4 mm at 30 days ($P = .0074$) (Figure 3).

Histology demonstrated myocyte disruption without fibrosis or inflammation immediately post-ablation. At 30 days, dense midseptal fibrosis was noted without a surrounding inflammatory infiltrate (Online Supplemental Figure 2).

### TTE, MRI, and electroanatomic mapping

TTE confirmed gross pathological findings: diastolic IVS thickness by TTE increased from 11.8 ± 1.3 mm at baseline to 13.4 ± 1.6 mm ($P = 0.002$) immediately post-ablation (presumably because of edema) but decreased to 10.0 ± 1.3 mm at 30 days ($P = .0003$ by ANOVA for between-group differences; $P = .004$ for comparison between pre-ablation and 30 days post-ablation) despite swine growth. The LV ejection fraction at baseline was 58% ± 6%, decreased to 49% ± 4% immediately post-ablation (due to hypokinesis of the basal IVS), and recovered to 58% ± 5% at 30 days ($P = .0002$) by ANOVA for between-group differences; $P = .92$ for comparison between pre-ablation and 30 days post-ablation).

MRI also confirmed IVS remodeling: IVS thickness increased from 11.0 ± 0.8 mm at baseline to 13.8 ± 1.7 mm at 30 days.

**Table 1.** Summary of lesion parameters at baseline and at 30 days post-ablation.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Immediately post-ablation (n=24)</th>
<th>30 day survival (n=42)</th>
</tr>
</thead>
<tbody>
<tr>
<td>% transmurality</td>
<td>58 ± 19%</td>
<td>81 ± 20%</td>
</tr>
<tr>
<td>% left of mid-septum</td>
<td>30 ± 23%</td>
<td>39 ± 28%</td>
</tr>
</tbody>
</table>

**Figure 3.** Gross pathology images of *in vivo* lesions immediately post-ablation (left) and at 30 days (right). Bottom row images demonstrate image processing with spline analysis defining the portion of the lesion surface area to the right (RV; red) and left (LV; green) ventricular side of the septum, with the mid-septum defined as the white dotted line.
mm immediately postablation (presumably because of edema) but decreased to 10.6 ± 1.1 mm at 30 days (P = .003 by ANOVA) despite swine growth. First-pass MRI contrast perfusion imaging demonstrated dense perfusion defects in the basal to mid-IVS immediately post-ablation (Online Supplemental Videos 1 and 2). Late gadolinium enhancement (LGE) detected areas of HIU ablation in the IVS, but resolution was insufficient to quantify individual lesion size on a per-lesion basis: 3.6 ± 1.1 regions of LGE were present post-ablation and 3.4 ± 1.1 regions at 30 days (Figure 4; Online Supplemental Figure 3), likely because of the coalescence of multiple HIU lesions into single LGE regions. T1 time at the basal IVS HIU ablation sites increased from 1176 ± 18 ms preablation to 1408 ± 87 ms postablation and remained elevated at 30 days (1319 ± 128 ms; P = .008 by ANOVA). T2 time increased from 40.4 ± 5.1 to 64.2 ± 6.0 ms postablation but then returned to near baseline 47.1 ± 5.8 ms at 30 days (P = .1169 for comparison between pre-ablation and 30 days; P < .0001 by ANOVA for groupwise comparison). Data were similar when normalizing septal ablation site T1 and T2 values with basal inferolateral (non-ablated) T1 and T2 values.

Electroanatomic voltage mapping did not reveal any bipolar voltage abnormalities at standard voltage thresholds, but demonstrated new areas of reduced unipolar voltage along the basal and mid-RV septum immediately post-ablation (10.6 ± 6.1 cm²) and at 30 days (7.0 ± 3.5 cm²) (P = .35). Similar unipolar voltage findings were present in the LV septum (7.6 ± 0.5 cm² vs 6.0 ± 1.8 cm²; P = .31) confirming deep lesions (Figure 5). No fractionated or late electrograms were noted, suggesting fairly homogeneous, deep ablation.

**Procedural complications**

No AV block, QRS widening, pericardial effusions, or ventricular septal defects were noted intra- or post-procedure. All survival swine recovered uneventfully from anesthesia after the procedure. One survival swine died suddenly at 14 days postprocedure, with no structural abnormalities determined at autopsy.

**Discussion**

There are spatial challenges related to catheter ablation of VAs. Mid-myocardial sites of VA origin and epicardial sites of origin in patients unsuitable for percutaneous subxiphoid pericardial access pose significant challenges for RF ablation. Several novel strategies have arisen to improve upon shallow lesion depth of standard RF ablation, including an RF needle catheter, stereotactic body radiation therapy, use of hypotonic irrigated RF ablation, bipolar RF ablation, and...
alcohol venous ablation, although access to these therapies is limited, and recurrence and complication rates remain high.

There are several potential advantages of using HIU over RF for targeting mid-myocardial and epicardial sites from the endocardium. Lower US absorption in tissue allows deeper penetration. Moreover, US transducers can be geometrically focused, or the natural focus of the US element may permit an "offset" that, with catheter cooling by irrigation, can spare tissue immediately adjacent to the transducer, thereby avoiding the subendocardial His-Purkinje system with septal ablation. Finally, US-based ablation may permit the real-time monitoring of the region of thermal injury that could help guide lesion placement or size.

In our study, HIU ablation from the RV side of the IVS resulted in lesions of ≈ 11 mm depth, initially spanning 58% of the IVS acutely increasing to 81% transmural at 30 days. This increase in transmurality was, in part, explained by remodeling of the IVS, which was noted to markedly decrease in thickness in response to septal ablation. Indeed, 38% of the lesion surface area was deposited on the far-field, LV side of the IVS, with the first 2–3 mm of the RV septal subendocardium spared. No ORS widening, bundle branch block, or AV block was noted, suggesting protection of the subendocardial His-Purkinje system. However, it should be noted that while extensive ablation of the basal-mid RV anteroseptum was performed, sonications were not performed in a dedicated manner over the His and right bundle potentials, but simply in their general region. Furthermore, we did not histopathologically define the His-Purkinje system after euthanasia and thus were unable to define the proximity of our ablation lesions to the conduction system. No acute complications were noted, but there was 1 autopsy-negative sudden death event at 14 days that we presume was due to VA secondary to heterogeneous necrosis in the IVS with lesions spaced 1 cm apart, although late AV block cannot be excluded.

HIU lesions were successfully identified by cardiac MRI, which may serve as a means of assessing lesion extent clinically using either LGE or noncontrast T1 imaging, as has previously been shown for RF. In our study, T1 and T2 times at the IVS were both elevated immediately post-ablation whereas only T1 time remained elevated at 30 days, suggesting an acute edematous and/or inflammatory reaction (although significant inflammation not noted histologically post-ablation), resolving to dense, mid-septal fibrosis without edema or inflammation at 30 days (which was confirmed by histology). It should be noted that our MRI lacked the resolution to confidently measure individual lesion dimensions for correlation with gross pathology.

Limitations of our study include lack of an RF control group, although the shallow depth of RF ablation has been well-documented, including in our prior study where it was 4.7 ± 4.0 mm and significantly less deep than epicardial HIU. Swine were not monitored with telemetry or implantable loop recorder monitoring over the 30-day survival period; we cannot exclude delayed transient AV block. We also did not test a large range of frequencies, transducer sizes (restricted to 3 × 5 mm transducers that would accommodate a 12-F catheter), or differences in acoustic power or lesion duration time. Instead, we used HIU frequencies within a range known to promote thermal necrosis and our previous experience to narrow our range of HIU variables, compared 5, 6.5, and 8 MHz in ex vivo models, which identified 6.5 MHz as the optimal frequency for the transducer size in the in vivo study.

Another limitation is the large variability in lesion sizes. While TTE was used to confirm placement against the RV side of the IVS, this did not correlate with histology findings. Additionally, we did not perform histology as a means of confirming lesion extent, as this was thought to be less likely to be correlated with lesion size in the ex vivo setting. Further work is required to determine the optimal range of HIU variables for mid-myocardial ablation, as well as the impact of transducer size, frequency, and power on lesion size and transmurality. Finally, we did not test a large range of transducer sizes, and further work is required to determine the optimal transducer size for mid-myocardial ablation.

Figure 5  Electroanatomic unipolar voltage mapping of the right (RV) and left (LV) ventricular septum. New unipolar voltage abnormalities noted postablation and at 30 days, confirming the presence of deep mid-myocardial lesions.
endocardial surface, we cannot exclude the periodic noncontact, which could explain the variability as well as why lesion efficiency was not 100% (ie, fewer histological lesions than ablation procedures performed; most evident for sonication made in the most apical portions of the RV) and also why lesion size varied. Furthermore, our side-facing transducer may not have always been exactly parallel to the IVS. Future iterations of the catheter should use a contact- and orientation-sensing mechanism. Integration of the catheter into electroanatomic mapping may also aid in accessing contact as well as allow us to guide HIU ablation directly over the His-Purkinje system to formally demonstrate sparing of the conduction system with surface-sparing lesions. Finally, our studies were not performed in a model of arrhythmia or cardiomyopathy. It has been shown that RF applications within the myocardial scar are less likely to generate reliable and large necrotic lesions compared with applications performed in healthy myocardium because of heterogeneous electrical impedance of scar tissue.22 HIU thermal necrosis relies on tissue acoustic impedance, which is more homogeneous across tissue types than electrical impedance.23 Thus, HIU lesions may perform more efficiently in scar, although this has yet to be tested.

Future investigations of HIU should assess lesions in large animal cardiomyopathy models, use long-term ambulatory rhythm monitoring for proarrhythmia, and assess endocardial to epicardial lesions with ablation over the non-septal LV endocardium. While we previously demonstrated a dose-response relationship between HIU powers of 15–30 W and lesion depth and volume9 (and thus chose 30 W for this study as it is the maximal acoustic power that the transducer and circuitry can tolerate in the continuous wave mode), further control of lesion depth and size may also be achieved with adjusting transducer size or ablation time and should be investigated. Our finding of reduction in IVS thickness is intriguing and warrants further investigation with a consolidated lesion set over the basal septum to determine whether HIU may be a suitable approach for septal reduction therapy in hypertrophic cardiomyopathy.

**Conclusion**

This first study of HIU endocardial ventricular ablation demonstrates deep, nearly transmural mid-myocardial lesion formation with sparing of the subendocardium and His-Purkinje system.

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**Appendix**

**Supplementary data**

Supplementary data associated with this article can be found in the online version at https://doi.org/10.1016/j.hrthm.2020.01.027.

**References**
