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# Catheter ablation lesion visualization with intracardiac strain imaging in canines and humans

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2 Abstract—Catheter ablation is a common treatment for 3 arrhythmia, but can fail if lesion lines are non-contiguous. 4 Identification of gaps and non-transmural lesions can reduce the 5 likelihood of treatment failure and recurrent arrhythmia. 6 Intracardiac myocardial elastograph is a strain imaging technique 7 that provides visualization of the lesion line. Lesion size estimation 8 and gap resolution was evaluated in an open chest canine model 9 (n=3), and clinical feasibility was investigated in patients 10 undergoing ablation to treat typical cavotricuspid isthmus atrial 11 flutter (n=5). A lesion line consisting of three lesions and two gaps 12 was generated in each canine left ventricle via epicardial ablation. 13 One lesion was generated in one canine right ventricle. Average 14 lesion and gap areas were measured with high agreement  $(33 \pm 14)$ 15  $mm^2$  and  $30 \pm 15 mm^2$ , respectively) when compared against gross 16 pathology (34  $\pm$  19 mm<sup>2</sup> and 26  $\pm$  11 mm<sup>2</sup>, respectively). Gaps as 17 small as 11 mm<sup>2</sup> (3.6 mm on epicardial surface) were identifiable. 18 Absolute error and relative error in estimated lesion area were 9.3 19  $\pm$  8.4 mm<sup>2</sup> and 31  $\pm$  34 %; error in estimated gap area was 11  $\pm$  9.0 20  $mm^2$  and 40 ± 29 %. Flutter patients were imaged throughout the 21 procedure. Strain was shown to be capable of differentiating 22 between baseline and after ablation completion as confirmed by conduction block. In all patients, strain decreased in the 23 24 cavotricuspid isthmus after ablation (mean paired difference of -25  $17 \pm 11$  %, p < 0.05). IME could potentially become a useful 26 ablation monitoring tool in the clinic.

Index Terms—ablation, electrophysiology, lesion monitoring,
 strain imaging, intracardiac echocardiography.

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#### I. INTRODUCTION

ATHETER ablation is one of the most effective treatments for atrial and ventricular arrhythmias [1]–[3]. An ablation catheter deposits energy in specific areas of the heart to form a barrier of non-conductive tissue that interrupts the electrical circuit responsible for the arrhythmia. The lesion line must be contiguous; gaps between lesions or non-transmural lesions can serve as a pathway of conductive tissue through which the arrhythmia circuit can recover, leading to treatment failure [4]–

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40 [7].

41 Currently, lesion geometry and location is estimated by using 42 surrogate parameters (e.g. force-time integral or FTI, and 43 contact force) that have been found to correlate with lesion size, 44 in conjunction with an electroanatomic mapping system [8]-45 [10]. These parameters are indirect measurements of lesion 46 formation, which limits their accuracy. Furthermore, the 47 inherent variance in atrial volume, tissue deformation from 48 application of the ablation catheter to the endocardial surface, 49 and unexpected procedural complications can cause significant 50 errors in the integration of the electroanatomic map with the CT 51 and MRI images, leading to inaccurate lesion localization [11], 52 [12]. Given that the single-procedure success of catheter 53 ablation of persistent atrial fibrillation is estimated to be as low 54 as 25% [13], there is an urgent need for an ablation monitoring 55 technique capable of accurately and robustly depicting 56 myocardial anatomy and lesion formation that can easily be integrated into the clinical routine. 57

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58 Ultrasound-based lesion mapping methods are a promising 59 means of obtaining real-time feedback during ablations. Unlike 60 MRI-based lesion mapping techniques-which would require extensive adjustments in the current procedure workflow due to 61 the required additional hardware and technicians-ultrasound 62 63 (particularly intracardiac echocardiography or ICE) is already a 64 common imaging modality during ablations[12], [14]. A novel 65 ablation catheter with near field ultrasound imaging capabilities has been shown capable of providing realtime feedback about 66 lesion formation [15]. While an improvement over the current 67 68 state-of-the art catheters which relay contact force or FTI, the 69 catheter cannot provide information about the position of 70 lesions relative to one another. Photoacoustic imaging of 71 ablation has been investigated, but thus far reports have been limited to benchtop experiments with ex-vivo tissue [16]–[19]. 72 73 Elastography-based methods employing shear waves or 74 acoustic radiation force impulse (ARFI) to identify lesions based on its mechanical moduli are an area of active research. 75

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76 Shear wave approaches have yet to investigate lesion gap 77 resolution [20], [21], and ARFI has stringent requirements 78 regarding the imaging configuration in terms of depth and 79 catheter orientation relative the myocardial wall [22].

80 Intracardiac Myocardial Elastography (IME) is a strain-81 based lesion mapping technique developed by our group. An 82 ICE catheter employs a high-frame rate acquisition sequence to estimate strain in the atrial or ventricular walls. IME has 83 84 minimal hardware requirements, and theoretically can be 85 programmed onto any ultrasound system capable of high-frame rate acquisitions. Grondin et al demonstrated feasibility by 86 capturing strain images in the canine LA, and by observing that 87 88 ablation lead to reduced local strain in the ablated region of the 89 human atria in atrial fibrillation patients [23]. Bunting et al used 90 IME in the canine LV to detect lesion gaps of 15 mm and 45 91 mm in vivo [24].

92 This paper presents the next stage of development in the 93 IME. Its objectives are twofold: 1) investigate lesion area and 94 lesion gap resolution in an open-chest canine model, and 2) 95 explore the potential diagnostic utility of IME strain in a clinical 96 feasibility study. Lesion gap resolution was investigated by 97 creating lesion lines comprised of three epicardial lesions and 98 two gaps in three canine left ventricles (LV). In patients, IME 99 was used to monitor ablation in the cavotricuspid isthmus (CTI) throughout an atrial flutter procedure in five subjects. 100

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#### II. METHODS

# 104 A. Animal model experimental protocol

Due to its cellular, functional, and physiological similarities with the human heart, the canine heart is one of the most popular large animal models in cardiac research [25]. Furthermore, since mongrel canines are genetically diverse, they are close representations to the non-homogenous genetic background of humans [26].

The study protocol was approved by the Institutional Animal 111 112 Care and Use Committee (IACUC) at Columbia University, and 113 was compliant with the Public Health Service Policy on 114 Humane Care and Use of Laboratory Animals. Lateral 115 thoracotomy was performed on anesthesized mongrel canines (n = 3, 100% male,  $26 \pm 2.1$  kg) to expose the myocardium for 116 117 epicardial ablation. The intracardiac ultrasound catheter (Carto Soundstar, Biosense Webster, Irvine, CA, USA) was introduced 118 119 via the external jugular vein and advanced through the superior 120 vena cava. Positioning the probe in the right ventricle (RV) 121 provided images of the anterior and antero-lateral segments of 122 the left ventricle (LV). Imaging of the RV was performed by 123 positioning the probe in the LV via apical puncture.

Epicardial lesions were created in the LV and RV by catheter ablation (Carto 3 System, Biosense Webster, Irvine, CA, USA). In the LV, a lesion line consisting of three lesions with two gaps were generated in three animals. In the RV, one lesion was created in one animal. Images were acquired prior to and after each lesion (SoundStar 10 F Catheter, Biosense Webster, Ivrine, CA, USA). For the LV lesion lines, images were acquired at baseline, and then after each lesion for a total of fourtime points. For the RV lesion, images were acquired beforeand after the ablation.

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134 Lesions were imaged with one of two ultrasound platforms 135 (the same catheter, Soundstar, was employed). Two canines 136 (two LV lesions lines and one RV lesion) were imaged with an 137 Acuson SC2000 in research mode (Siemens, Munich, 138 Germany). The transmit sequence consisted of 24 steered plane 139 waves (virtual source located >300 mm behind the transducer) 140 at a frame rate of 200-250 fps and depth of 80 mm. The angular 141 aperture of the field-of-view was 70°. One canine (one LV 142 lesion line) was imaged with a Verasonics Vantage (WA, USA) 143 and а modified Acuson Swiftlink Connector 144 (TransducerWorks, PA, USA). A high-frame rate compounded 145 sequence was employed (15 virtual sources, focus located 21 146 mm behind the transducer, 460 fps, depth 80 mm, angular 147 aperture 90°) [27].

148 At the conclusion of the procedure, the myocardium was 149 excised. The lesion line was segmented and placed in the 150 freezer (-18° C) for 40-60 minutes. The sample was sliced 151 transmurally along the axis of the lesion line. Sections were submerged in 1% tetrazolium chloride (TTC) and placed in an 152 153 incubator (37°) for at least 40 minutes. TTC stained the lesions 154 white. Photos of the sections with scale bar for reference were 155 obtained (Nikon EOS Rebel T3i, Tokyo, Japan).

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#### 157 B. Human experimental protocol

The study protocol was approved by the Institutional Review
Board (IRB) of Columbia University. Patients were informed
of the study's risk prior to obtaining consent. The transmit
sequence complied with the U.S. Food and Drug
Administration (FDA) limits on acoustic output.

163 Patients with typical cavotricuspid isthmus atrial flutter (n = 164 5, men = 60%, age =  $67 \pm 16$  years old) undergoing RF ablation 165 of the cavotricuspid isthmus of the right atrium (RA) were 166 imaged with an ICE clinical machine in research mode 167 (ViewMate ICE Catheter and Viewmate Z, Abbott, Chicago, IL). The ICE catheter was positioned in the RA. A custom 168 transmit sequence was implemented: 1.5s of conventionally 169 170 focused imaging at 30 fps, followed by 1 s of a single diverging 171 wave sequence (-6.5 mm virtual source, 600 fps frame rate). 172 The latter acquisition was used to estimate myocardial 173 displacement and strain; the former was used to provide a B-174 mode reference frame of end-systole over which to overlay the 175 strain. The ICE ultrasound field-of-view was set to the CTI region proximal to the tricuspid valve. Images were acquired 176 177 prior, during, and after the CTI ablation procedure. The 178 ultrasound view was updated throughout the procedure to 179 ensure that the ablation catheter was in view before, during, and 180 after each lesion. The ablation procedure was considered 181 complete once achievement of block was confirmed via 182 coronary sinus pacing.

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# 184 C. Intracardiac Myocardial Elastography (IME) imaging

185 Three different ultrasound platforms were used in this study: 186 Siemens Acuson, Verasonics Vantage, and Abbott Viewmate Z. Each hardware platform possessed different strengths. The 187 188 Acuson provides a means to coregister the lesion line and the 189 ICE view through the CARTOSOUND software (Carto 3 190 System, Biosense Webster, Irvine, CA, USA), which can mark 191 the position of the ablation catheter in real time when it is in 192 plane with the ICE catheter [28]. The Vantage's open 193 programmability allows for the implementation of optimal high 194 frame rate strategies. The Viewmate Z is a clinically available 195 FDA-approved scanner electrophysiologists have experience 196 operating. Beamforming, displacement estimation, and strain 197 estimation parameters slightly differed between the three 198 platforms.

Beamforming was performed internally with the Acuson. Raw RF data obtained with the Vantage and Viewmate Z was beamformed by delay-and-sum [27].

202 For data obtained with the Viewmate Z and Vantage, axial 203 displacements were calculated from beamformed RF signals 204 using a 1-D normalized cross-correlation kernel [29]. Due to 205 its lower achievable acquisition frame rate, axial displacements 206 were estimated from the beamformed envelope (as opposed to 207 RF) signals on data obtained from the Acuson. For the canine 208 imaging protocol, the displacements observed in the LV and 209 RV were accumulated throughout LV systole, and RV systole, 210 respectively. In the human protocol, the displacements were 211 accumulated during atrial filling, a segment of the cardiac cycle 212 during which the CTI lengthens. Different phrases of the 213 cardiac cycle were imaged in the animal and human models to 214 preserve the directionality of the strain being observed, i.e. 215 positive strain was estimated in both. Axial strains were derived 216 from cumulative axial displacements with a 1-D least-squares estimator (LSQSE) [30]. Strain images were smoothed using a 217 218 2-D median filter. The specific processing parameters used with each hardware platform—displacement kernel, strain (LSQSE) 219 kernel, and median filter kernel-are summarized in a table in 220 221 the Appendix (Table A.1).

D. Canine study – image analysis and statistics

manual segmentation (Fig. 1).

Canine strain images were validated against gross pathology.

The images of the TTC-stained, excised lesion line were

converted to grayscale, scaled, aligned with ICE, and overlaid

onto the B-mode image based on anatomical landmarks (Fig.

1). TTC stains the lesions white; in grayscale, the lesions are

brighter compared to non-ablated myocardium. The lesions

were masked by a combination of brightness thresholding and

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233 234 Fig. 1. Coregistration procedure to validate IME lesion maps with gross 235 pathology in canines. The lesion line was excised from the myocardium after 236 sacrifice and sliced transmurally along the axis of the lesion line to provide a 237 cross-sectional view of the lesion and gap area (A). Using anatomical markers 238 (e.g. papillary muscles and epicardial surface), the intensity image was 239 manually rotated and translated (B) to coregister with the B-mode image (C). 240Intensity thresholding and manual segmentation were employed to create a 241 binary mask that indicated the lesion area by gross pathology (indicated in light 242 blue) (D).

243

244 Due to translation and deformation of the myocardium 245 during the cardiac cycle, the lesion line may move out of the 246 field-of-view at certain time points during systole. This drop-247 out is evident upon examination of the strain movie through the 248 entirety of systole. Thus, the number of displacement frames 249 accumulated varied for each acquisition. The strain magnitude 250 is dependent on the number of displacement frames 251 accumulated. The strain image dynamic range was adjusted 252 accordingly in order to maintain high image contrast between 253 unablated and ablated tissue. The upper bound of the dynamic 254 range  $(DR_{upper})$  was empirically chosen to be half of the 255 median strain at baseline at the number of frames accumulated: 256

$$DR_{upper} = \frac{median(\varepsilon_{baseline}(n))}{2} \tag{1}$$

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258 where  $\varepsilon_{baseline}(n)$  is the masked strain values in the  $n^{th}$ 259 accumulated frame at baseline. The lower bound of the dynamic 260 range was set to 0%.

Lesion area as estimated by IME was calculated as follows:

$$A_{strain}(mm) = A_{pixel} *$$

$$\sum_{i=1}^{N} \begin{cases} 0 & if \ \varepsilon(i) \ge \varepsilon_{thresh} \\ 1 & if \ \varepsilon(i) < \varepsilon_{thresh} \end{cases},$$
(2)

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264 where  $\varepsilon(i)$  represents strain at a given pixel *i* within a masked 265 region consisting of *N* pixels,  $\varepsilon_{thresh}$  is the strain threshold, and 266  $A_{pixel}$  is the area of each pixel in mm. Masks were manually 267 delineated to isolate lesions (n=10) and gaps (n=6). The 268 boundary of the gap masks was set by a vector spanning the 269 apex of the two lesions, at the points closest to the endocardial 270 wall. The other borders consisted of the lesion perimeters and

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271 the epicardial wall. The absolute ( $\delta A$ ) and relative difference 272 between  $A_{strain}$  and the areas reported by gross pathology 273 ( $A_{gross}$ ) were calculated, given

$$A_{gross}(mm) = A_{pixel} * \sum_{i=1}^{N} \begin{cases} 0 & if \ pixel(i) \ge pixel_{thresh} \\ 1 & if \ pixel(i) < pixel_{thresh} \end{cases}$$
(3)

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276 The brightness threshold  $pixel_{thresh}$  was determined 277 empirically based on qualitative assessment of the gross 278 pathology images, and  $A_{pixel}$  is the area of each pixel in mm. 279 Manual segmentation was performed to isolate individual 280 lesion areas.

281 The threshold under which a point would be considered a 282 lesion,  $\varepsilon_{thresh}$ , was empirically derived. The Dice similarity coefficient (DSC) ranges from 0 to 1, and measures the 283 284 similarity between two binary sets. A DSC of 1 indicates that 285 the binary masks are identical, while a score of 0 indicates that 286 there is no intersection between the positive values in the binary 287 masks. The DSC has frequently been employed in medical imaging analysis (e.g. to compare manual segmentation against 288 289 an automated method) [31], [32]. In terms of true positives 290 (TP), false positives (FP), and false negatives (FN), the DSC 291 between two binary sets A and B is given by,

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$$dice(A,B) = \frac{2TP}{2TP + FP + FN}.$$
 (4)

In this study, TP corresponds to tissue that was ablated and
correctly identified as such by IME, FP corresponds to regions
of unablated tissue that was incorrectly identified as ablated,
and FN corresponds to ablated tissue that was incorrectly
identified as unablated.

299 The DSC of the lesion maps produced by strain imaging and 300 gross pathology (the ground truth) were compared for a range 301 of  $\varepsilon_{thresh}$ , defined as,

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$$\varepsilon_{thresh} = \alpha * median(\varepsilon_{baseline}(n)),$$
 (5)

304 with  $\alpha$  evaluated within the range of [0, 1.5]. The  $\alpha$  value 305 that yields the optimal  $\varepsilon_{thresh}$  was determined by calculating 306 the mean DSC curve across the four lesion lines. The  $\alpha$  value 307 corresponding to the maximum of the mean DSC was chosen to 308 calculate  $\varepsilon_{thresh}$  in the canine model.

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#### 310 E. Human study – image analysis and statistics

The CTI was manually segmented. Median axial strain was 311 312 calculated within the CTI for the five patients. Median strain  $(\varepsilon_{median})$  at the CTI at baseline and at the conclusion of the 313 procedure (once block was achieved) was statistically 314 315 compared via the Student's paired t test. In contrast to the canine protocol, no thresholding was performed to isolate 316 317 individual lesions, and the dynamic range was set to [-40%, 318 40%] for all cases.

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III. RESULTS

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322 A. Open chest canine ablation

323 ME was capable of accurately capturing the formation of the 324 LV lesion line throughout the ablation procedure (Fig. 2). At baseline, LV strain was homogenously positive and high 325 magnitude ( $\varepsilon \ge DR_{upper}$ ) throughout (Fig. 2A). In contrast, 326 lesions manifested as regions of low strain ( $\varepsilon < \frac{DR_{upper}}{2}$ ). The 327 328 strain images demonstrate the progression of lesion line 329 formation, lesion-by-lesion (Fig. 2b-d). At the conclusion of the 330 ablation experiment, three distinct lesions and two distinct gaps 331 are visible (Fig. 2d).

The contours of the lesion areas as indicated by the TTCstained tissue sections (shown in green) were overlaid onto the post-ablation strain-based lesion maps (Fig. 3). Qualitatively, there is good agreement between the lesions detected by ME and the gross pathology. In each of the three LV lesion lines, ME correctly identified three lesions and two gaps. The singular lesion in the RV was also correctly identified.

339 Calculating the DSC across a range of  $\alpha$  values (Eq. 3) 340 yielded the plot summarized in Fig. 4. The maximum of the 341 mean DSC curve was found at  $\alpha = 0.27$ , wherein a DSC value 342 of 0.62 was found.

The lesion and gap areas as determined by IME versus gross
pathology are summarized in Fig. 5. The lesion areas are
designated in turquoise, with the gaps indicated in yellow.
Qualitatively, the thresholded strain lesion maps compare well
against the gross pathology. The lesion areas found by strain
and gross pathology are summarized and compared in Table I.
Gap area assessment is summarized in Table II.

By gross pathology, the lesion and gap areas  $(A_{gross})$ measured 34 ± 19 mm<sup>2</sup> and 26 ± 11 mm<sup>2</sup> on average, respectively. By IME strain imaging, the lesion areas and gaps  $(A_{strain})$  were estimated to be 33 ± 14 mm<sup>2</sup> and 30 ± 15 mm<sup>2</sup> on average, respectively.

The individual difference in lesion area between strain and gross pathology ( $\delta A$ ) ranged 0.82-24 mm<sup>2</sup>, with a mean difference of 9.3 ± 8.4 mm<sup>2</sup>. In terms of relative error, the difference ranged 3.9 - 120 %, with a mean relative difference of 31 ± 34 %.

The individual difference in gap areas between strain and gross pathology ranged  $0.63 - 23 \text{ mm}^2$ , with a mean difference of  $11 \pm 9.0 \text{ mm}^2$ . In relative error, the difference ranged 5.5 - 78%, with a mean relative difference of  $40 \pm 29$  %.

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#### 365 B. Atrial flutter CTI ablation

366 Lesion mapping in a patient receiving CTI ablation to relieve 367 atrial flutter is demonstrated in Fig. 6. Images were acquired at 368 baseline (Fig. 6A), during (Fig. 6B), and after (Fig. 6C) CTI 369 ablation. The lesion line was initiated proximal to the tricuspid 370 valve, and progressed towards the direction distal the tricuspid 371 valve during the ablation. At baseline, the CTI exhibits 372 homogenously positive, high-magnitude strain (i.e. strain 373 greater than about 20%) (Fig. 6a). During the procedure, strain

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in the region proximal the tricuspid valve is low magnitude (i.e.
strain less than about 20%), while the region distal the valve
(thus far in the procedure unablated) still possesses highmagnitude positive strain (Fig. 6b). Finally, the entire CTI is
observed to have low-magnitude strain at the conclusion of the
ablation procedure (Fig. 6c).

180 In all five patients imaged,  $\varepsilon_{median}$  in the CTI decreased after 181 ablation compared to baseline. The mean paired difference in 182 CTI strain was -17 ± 11 %. Employing a two-sided paired t-test, 183 the difference was determined to be statistically significant (p < 184 0.05).



385 386 Fig. 2. Strain-based lesion maps obtained with ICE imaging. IME imaged the 387 development of a lesion line generated in an open-chest canine via epicardial 388 ablation. Unablated myocardium exhibits high-magnitude positive strain over 389 systole, indicated in red. At baseline, prior to any ablations, high-magnitude 390 strain is evident throughout the LV wall (A). Ablated tissue is stiff and non-391 contractile, manifesting as low-magnitude regions, indicated in blue. IME 392 accurately tracked the ablation lesion-by-lesion: the first lesion is indicated in 393 (B), followed by a second lesion and the first gap in (C), and finally all three 394 lesions and both gaps in (D). 395

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В line Line 1 20 С D Line 2 20 mm 10 mm Е 10 Line 3 -20 -10 0 10 20 G н 20 10 -10 0 5 mm -40 -30 -20 -10 0 10 20 30

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406  $\alpha$  coefficient (Eq. 3) 407 Fig. 4. Dice similarity coefficient (DSC) for each lesion line. The metric 408 measures the similarity between two binary images, with higher values 409 indicating more similarity. The three lesion lines and one RV lesion were 410 evaluated against gross pathology. A range of  $\alpha$  values [-0.5, 1.5] were 411 evaluated to determine the optimal strain threshold,  $\varepsilon_{thresh}$  (Eq. 3). The 412 maximum DSC of the mean Dice curve occurs at  $\alpha = 0.27$ .

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414 415 Fig. 5. Thresholded strain-based lesion maps versus gross pathology. 416  $A_{strain}(\alpha = 0.27)$  for the lesion and gaps detected by IME were assessed against the ground truth areas according to gross pathology  $(A_{gross})$  for LV 417 418 lesion line 1 (A, B), LV lesion line 2 (C, D), LV lesion line 3 (E, F), and the 419 RV lesion (G, H). Lesion areas are indicated in turquoise, with gap areas in 420 yellow. 421

TABLE I

LESION AREA BY IME STRAIN AND GROSS PATHOLOGY IN THE CANINE
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Lesion	Agross	$A_{strain}$	$\delta A$	Relative error
	(mm <sup>2</sup> )	(mm <sup>-</sup> )	(mm <sup>-</sup> )	(%)
Line 1, Lesion A	28	15	13	46
Line 1, Lesion B	27	34	7.9	30
Line 1, Lesion C	18	23	4.8	26
Line 2, Lesion A	30	28	2.1	7.0
Line 2, Lesion B	20	45	24	120
Line 2, Lesion C	21	22	0.82	3.9
Line 3, Lesion A	29	28	1.1	3.9
Line 3, Lesion B	26	32	6.8	26
Line 3, Lesion C	65	42	23	35
RV, Lesion A	72	62	9.5	13
Mean	34	33	9.3	31
STD	19	14	8.4	34

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		TABLE II		
GAP AREA BY IN	AE STRAIN A	ND GROSS P	ATHOLOGY I	N THE CANINE
Gap	A <sub>aross</sub>	A <sub>strain</sub>	δΑ	Relative error
	(mm <sup>2</sup> )	(mm <sup>2</sup> )	(mm <sup>2</sup> )	(%)
Line 1, Gap AB	19	32	13	66
Line 1, Gap BC	11	11	0.63	5.5
Line 2, Gap AB	43	25	18	43
Line 2, Gap BC	25	22	3.2	13
Line 3, Gap AB	29	39	10	35
Line 3, Gap BB	30	53	23	78
Mean	26	30	11	40
STD	11	15	9.0	29

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435 436 Fig. 6. Tracking ablation of the cavotricuspid isthmus (CTI) in the RA of a 437 patient with atrial flutter. The catheter is outlined by a yellow dotted line. 438 Unablated myocardium in the CTI exhibits high-magnitude (>20%) positive 439 strain (indicated in red) over the atrial filling phase of the cardiac cycle. Ablated 440 tissue manifests as comparatively low magnitude (<20%) or near-zero 441 magnitude strain (indicated in pink or white). At baseline, the CTI is observed 442 as healthy throughout (A). Lesions were first generated proximal to the 443 tricuspid valve. There is a clear difference in strain magnitude between the 444 ablated tissue proximal the valve and the unablated tissue distal the valve (B). 445 At the end of the ablation procedure, the CTI is observed to have low-magnitude 446 strain throughout (C).

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Fig. 7. Median strain within the CTI decreases after ablation. Patients receiving
ablation of the CTI to relieve atrial flutter were imaged with IME (n=5). Median
strain decreased for all patients, with a statistically significant difference in the
mean paired difference (paired Student's t-test, \* <0.05).</li>

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#### IV. DISCUSSION

The potential of IME for lesion and gap visualization and 456 quantification with intracardiac echocardiography 457 was 458 investigated. Employing an open-chest canine model, IME was 459 capable of resolving all ten lesions and all six lesion gaps generated in three LVs and one RV. IME was then used to track 460 461 atrial flutter ablation in human subjects. There was a reduction 462 in strain in the ablated region (the CTI) in all five patients after 463 the ablation procedure.

464 In both the canine and clinical studies, lesions manifested as 465 areas of low-magnitude or near-zero percent strain, as opposed to areas of negative strain. The mechanics behind non-466 compliant and compliant passive myocardial tissue dictates the 467 directionality of the strain. Following acute ischemia, the 468 affected region of the myocardium becomes passive [33]. 469 470 During systole, the compliant ischemic tissue stretches, exhibiting negative strain. In RF catheter ablation, the 471 472 mechanism is different. The lesion that is generated is non-473 compliant, and becomes stiffer compared to unablated tissue 474 [34]. Since non-compliant, stiff tissue does not deform easily, 475 0% strain is expected over both systole (as shown in the canine 476 model) and diastole (as shown in the patient model).

477 The strain magnitude within the ablated area was 478 significantly reduced compared to unablated tissue in the 479 canine. The dynamic range as defined in Eqn. 1 was chosen to 480 increase the visual contrast between scar and unablated tissue; healthy tissue manifests as regions of high magnitude positive 481 strain (in red) while scarred tissue manifests are regions of low 482 483 magnitude strain (in blue) (Fig. 2). These low-magnitude strain 484 regions were indicative of non-contractile scar tissue, verified 485 by overlaying the ground truth lesion contours as defined by gross pathology (Fig. 3). Qualitatively, there was excellent 486 487 agreement between the IME lesion maps and the gross 488 pathology: IME correctly represented the three lesions in each 489 canine LV as a non-contiguous linear line (Fig. 3A-F).

490 Thresholding was employed to allow for quantitative

491 comparison of lesion and gap area between strain imaging and 492 gross pathology (Tables I and II). The strain threshold under 493 which a region of tissue should be classified as scar was chosen 494 based on the peak mean Dice curve (Fig. 4). Thresholding can 495 also be a useful tool for visualization, simplifying interpretation 496 of the strain images. The optimal hard threshold will likely be 497 variable among patients, imaging conditions, and the heart 498 chamber being imaged. However, given that lesions are stiff 499 and non-compliant, it can be hypothesized that lesion strain 500 magnitude should always be close to zero. Future clinical 501 studies should be conducted to determine the degree of 502 variation in the strain threshold between patients. If the 503 variability is small, a general threshold may be implemented 504 instead of adjusting the value for each patient.

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505 Gap resolution was improved compared to previous 506 implementations of IME. In earlier work by our group, the 507 smallest detectable gap measured from lesion edges at the 508 epicardial level was 15 mm [24]. This study demonstrated that 509 IME was capable of resolving gaps as small as 11 mm<sup>2</sup>, or 3.6 510 mm measured edge-to-edge at the epicardial level (Table II, 511 Line 1 Gap BC, Fig. 5A-B). The improved lesion mapping is 512 due to the implementation of superior high frame-rate transmit 513 strategies. Instead of a single-diverging wave sequence, a 15-514 source compounding (Verasonics) and 23-source composite 515 plane wave sequence (Siemens) were applied in the canine 516 model. Increased gap resolution improvement is likely 517 attributable to improved SNR and lateral resolution over single-518 source diverging wave imaging [35], [36].

519 Great care was taken to best align the lesion line and 520 ultrasound plane in the canine model. In conjunction with 521 SOUNDSTAR, the Acuson was able to graphically mark the 522 location of the ablation catheter when it was in-plane. In the 523 acquisitions taken with the Vantage, each potential location was 524 manually palpated prior to ablation; if the location was in-plane, 525 the resulting tissue deformation would be evident in the B-526 mode. Nonetheless, an inherent source of error in the canine 527 study design is that a 2-D strain image of a live tissue target 528 featuring 3-D translation and deformation was compared 529 against a 2-D section of gross pathology. A lower degree of 530 agreement in the lesion and gap areas are attributable to the 531 imperfect coregistration of the two 2-D representations of the 532 lesion line. This imperfect coregistration likely inflated the 533 error of the lesion and gap area estimation— $31 \pm 34$  % and 40 534  $\pm$  29 %, respectively (Table I and II)—and led to a relatively 535 low maximum DSC of 0.62. In future large animal studies, the 536 addition of Gadolinium-enhanced Magnetic Resonance 537 Imaging could mitigate this source of error [37]. The combination of MRI 3-D visualization of lesion line and gross 538 539 pathology would allow for more reliable assessment of lesion 540 and gap area accuracy, and provide insight on how much error is attributable to poor coregistration versus the resolution 541 542 limitations of IME.

543 The feasibility of IME in intracardiac echocardiography as a 544 useful diagnostic tool was investigated via a pilot study 545 comparing the strain in the CTI before and after atrial flutter 546 ablation. Strain was calculated during atrial filling of the RV, 547 during which the CTI exhibits extension (i.e. stretches). Scar

tissue is stiffer and less compliant [38]; ablated regions of the 548 549 atrial wall exhibit significantly reduced extension during atrial 550 filling, manifesting as regions of low-magnitude strain (<20%) in IME lesion maps. The progression of a CTI ablation is 551 552 summarized in Fig. 6. The ablation catheter begins proximal the 553 valve and progresses along the CTI in the direction distal the 554 valve, the location of previous lesions clearly indicated by the 555 strain drop compared to the baseline. IME was able to 556 differentiate between the CTI before and after ablation (mean 557  $\Delta \varepsilon = -17 \pm 11$  %), with every patient recording a decrease in  $\varepsilon_{median}$  at the end of the procedure (Fig. 7). 558

The clinical pilot study demonstrated that IME could be 559 560 integrated into the current ablation workflow with minimal adaptations, especially since the use of ICE is standard of care 561 in many cardiac ablation procedures. Many of the proposed 562 563 approaches to imaging the lesion line require additional 564 hardware, such as MRI [12], [14], or an additional probe to 565 induce a push beam [21], [38], [39]. In contrast, IME was 566 integrated into an ICE platform that was already being used in 567 ablation procedures at the EP clinic. Furthermore, IME can 568 provide a large and deep field-of-view of the lesion line in contrast to photoacoustic [16]–[18] or ARFI [38], [39] methods, 569 570 allowing for quicker assessment of an ablation procedure's 571 progress.

572 In contrast to the canine study, the IME was unable to 573 observe the development of the lesion line lesion-by-lesion. Due to constraints on the programmability of the Abbott ICE 574 platform's research mode, a single diverging wave transmit 575 576 sequence was employed, as opposed to the compounding or 577 composite plane wave sequences used in the canine protocol. 578 The reduced imaging SNR, as well as the lower thickness of the human RA compared to the canine LV and RV, lead to poorer 579 580 strain resolution and less precise lesion maps.

581 As stated in the Methods section, the displacement estimation algorithm employed depended on the hardware platform that 582 583 was used to collect the acquisitions. Similarly to previous 584 iterations of IME [23], [24], displacement estimation was 585 performed using 1-D cross-correlation on the RF data derived 586 from the Verasonics Vantage and Abbott Viewmate Z. In 587 contrast, 1-D cross-correlation was performed on the envelope 588 of the RF signal on data derived from the Siemens Acuson 589 system. RF-based motion estimators are generally considered 590 more accurate than envelope-based estimators at high frame 591 rates, since the former contains phase information [40]-[42]. 592 However, RF-based estimators can perform poorly if the 593 acquisition frame rate is too low, or a large window size is 594 employed, due to decorrelation from false peak or jitter errors [41], [43]. The Siemens Acuson transmit sequence frame rate 595 596 used herein ( $\leq 250$  Hz) was substantially lower than that of the Verasonics (460 Hz) or Abbott (600 Hz) systems. Envelope-597 598 based displacement estimation was necessary as limitations in 599 the programmability of the Acuson prevented imaging at higher frame rates. Preliminary analysis confirmed that the envelope-600 601 based estimator would be preferred for the Siemens data, given the frame rate limitations. While displacement estimation on RF 602 versus beamformed envelope signals is comparable in the 603 canine study, the latter approach is less optimal because it 604

requires a larger displacement window size (Table A.1). When 605 606 imaging thin tissue such as the atria, a large displacement kernel 607 precludes high resolution imaging. Finally, 1-D crosscorrelation of the envelope signal was performed instead of 608 609 using a 2-D kernel, which is more common [41], [44]. It was 610 hypothesized that a 1-D kernel would be more accurate due to 611 the edge mismatch between adjacent plane waves in the 612 transmit sequence, visible in the B-mode (Fig 1C, 1D, for 613 example). While this mismatch in the lateral direction would lead to substantial decorrelation when employing a 2-D kernel, 614 615 this artifact is not relevant when using a uniaxial estimator.

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616 There are some limitations to consider and future work that 617 must be performed before IME is integrated into the clinic. The 618 implementation of IME described here only estimated 1-D 619 (axial) motion. In both the canine and human study, ultrasound 620 views were chosen such that the predominate direction of 621 myocardial motion was in the axial direction. However, prior to 622 implementation in the clinic, integration of 2-D or 3-D motion 623 estimation is necessary to circumvent angle dependence in the 624 strain calculation.

625 IME requires that the ultrasound plane is well-aligned with 626 the lesion line. Due to the myocardial translation and rotation 627 inherent in the cardiac cycle, this can be a challenging demand 628 for the ICE operator. Furthermore, this study evaluated IME's 629 performance in monitoring ablation of the CTI, a region of the 630 myocardium that is particularly amenable to ICE imaging. 631 Imaging regions of the myocardium that are frequently ablated, 632 such as the pulmonary veins in atrial fibrillation ablation, is 633 more challenging. A future clinical implementation of IME 634 could be integrated with an electroanatomic system. The 635 clinical model could be improved by using the Acuson platform 636 and SOUNDSTAR software, which is capable of visually 637 tagging the ablation catheter when it is view, leading to better 638 alignment with the lesion line. Linking IME to an 639 electroanatomic system could theoretically allow for the 640 generation of stain lesion maps that are registered to specific 641 positions in the myocardium, allowing for a pseudo-3D 642 visualization. Alternatively, a 3D ICE catheter could be 643 employed.

This study investigated gap resolution in the canine LV. 644 645 Clinically, ablation in the atria are more common than ablation 646 in the ventricles. Unfortunately, the smaller size of the canine 647 atrial chamber compared to the human equivalent complicated 648 navigation of the ICE catheter, and good coregistration of the 649 lesion line and ultrasound windows was unattainable. 650 Furthermore, lesions are typically generated on the endocardial 651 wall, as opposed to the epicardial wall as performed in the canine protocol. Further studies are needed to investigate the 652 653 gap resolution of IME in thinner myocardial tissue and in 654 endocardial catheter ablation.

Although three different ultrasound platforms were used in this study, determining the optimal hardware platform for IME is outside the scope of this paper. We hypothesize that any platform capable to achieving high frame rate imaging and integration with electroanatomic mapping systems would enable IME to be a diagnostic tool of high clinical value.

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# V. CONCLUSION

663 The potential of IME to visualize the lesion line and better inform ablation procedures was investigated. Gap resolution of 664 IME lesion mapping was validated in an open-chest canine 665 666 model that tracked epicardial ablations in the ventricles, with the smallest gap tested being 11 mm<sup>2</sup> (3.6 mm on epicardial 667 surface). A clinical feasibility study was also performed to 668 demonstrate the diagnostic utility of strain, and to show that 669 IME could be integrated into ablation procedures with minimal 670 modifications to the current workflow. Additional feasibility in 671 672 animals and humans are warranted to prove that IME is a viable 673 ablation monitoring approach for atrial arrhythmias. 674

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#### VI. APPENDIX

TABLE A. I           Ultrasound platform processing parameters								
Ultrasound platform	Probe center freq. (MHz)	Disp. kernel (mm)	Disp. kernel overlap (%)	Strain (LSQSE) kernel (mm)	2D median filter kernel (mm, °)			
Verasonics Vantage	5.2	3.9	90	4.3	(2.1, 2.5)			
Siemens SC2000 Acuson	6.0	4.8	90	3.0	(4.6, 2.9)			
Abbott Viewmate Z	6.0	1.0	90	2.3	(1.4, 2.3)			

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