

# Multimodal Examination of Atrial Fibrillation Substrate

## Correlation of Left Atrial Bipolar Voltage Using Multi-Electrode Fast Automated Mapping, Point-by-Point Mapping, and Magnetic Resonance Image Intensity Ratio

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### ABSTRACT

**OBJECTIVES** The aim of this study was to examine atrial fibrillation (AF) substrate using different modalities (point-by-point [PBP], fast anatomic mapping [FAM], and late gadolinium enhancement [LGE] magnetic resonance imaging [MRI] mapping) in patients presenting for AF ablation.

**BACKGROUND** Bipolar voltage mapping, as part of AF ablation, is traditionally performed in a PBP approach using single-tip ablation catheters. Alternative techniques for fibrosis delineation include FAM with multi-electrode circular catheters and LGE MRI. The correlation between PBP, FAM, and LGE-MRI fibrosis assessment is unknown.

**METHODS** LGE MRI was performed pre-ablation in 26 patients (73% men, mean age  $63 \pm 8$  years). Local image intensity ratio (IIR) was used to normalize myocardial intensities. PBP and FAM voltage maps were acquired, in sinus rhythm, prior to ablation and coregistered with LGE MRI.

**RESULTS** The mean bipolar voltage for all 19,087 FAM voltage points was  $0.88 \pm 1.27$  mV, and the average IIR was  $1.08 \pm 0.18$ . In an adjusted mixed-effects model, each unit increase in local IIR was associated with a 57% decrease in bipolar voltage ( $p < 0.0001$ ). An IIR  $>0.74$  corresponded to a bipolar voltage  $<0.5$  mV. A total of 1,554 PBP mapping points were matched to the nearest FAM point. In an adjusted mixed-effects model, log FAM bipolar voltage was significantly associated with log PBP bipolar voltage ( $\beta = 0.36$ ,  $p < 0.0001$ ). At low voltages, FAM distribution was shifted to the left compared with PBP mapping; at intermediate voltages, FAM and PBP voltages were overlapping; and at high voltages, FAM voltages exceeded PBP voltages.

**CONCLUSIONS** LGE-MRI, FAM, and PBP mapping showed good correlation in delineating electroanatomic AF substrate. Each approach has fundamental technical characteristics, the awareness of which allows proper assessment of atrial fibrosis. (J Am Coll Cardiol EP 2017; ■:■-■) © 2017 by the American College of Cardiology Foundation.

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**ABBREVIATIONS  
AND ACRONYMS****AF** = atrial fibrillation**EAM** = electroanatomic mapping**FAM** = fast anatomic mapping**GEE** = generalized estimating equation**IIR** = image intensity ratio**LA** = left atrial**LGE** = late gadolinium enhancement**MRI** = magnetic resonance imaging**PBP** = point-by-point**PV** = pulmonary vein**3D** = 3-dimensional

**P**ulmonary vein (PV) isolation by radiofrequency ablation has emerged as an effective treatment for atrial fibrillation (AF) by interrupting signal conduction arising from ectopic triggers in the PVs to the left atrial (LA) myocardium (1). Atrial myocardial substrate including areas with low bipolar voltage, shortened effective refractory periods, and conduction heterogeneity all likely play a role in AF perpetuation (2-4). These regional disparities in myocardial properties are thought to support re-entry and triggered activity in the atrium (5). Assessment of atrial fibrosis burden is increasingly used for prognostication and to inform ablation strategies (6-8).

Initial studies examining low-voltage substrate and its correlation with late gadolinium enhancement (LGE) magnetic resonance imaging (MRI) used a point-by-point (PBP) mapping approach using ablation catheters (6,7). Although this approach offers the advantage of confirming tissue contact using force-sensing catheters, it is time-consuming. An alternative is fast anatomic mapping (FAM) using decapolar circular mapping catheters (CARTO LASSO 2515 Variable Loop Eco Nav, Biosense Webster, Diamond Bar, California) with a smaller electrode size that can acquire highly detailed (>1,000 points) voltage maps in significantly less time. However, the correlation of LA fibrosis detected with LGE MRI, PBP mapping, and FAM is unknown. Voltage correlation between these techniques has not been established despite the use, in clinical practice, of common voltage thresholds (0.2 to 0.5 mV) to define diseased myocardium irrespective of the mapping technique. In this study, we sought to examine and compare AF substrate characteristics using different modalities (LGE MRI, FAM, and PBP mapping) in patients presenting for AF ablation.

**METHODS**

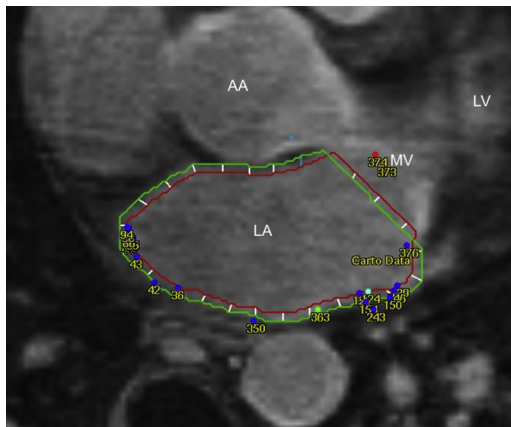
**PATIENT POPULATION.** Twenty-six patients with symptomatic, drug-refractory paroxysmal or persistent AF referred for AF ablation underwent preprocedural MRI between June 2016 and December 2016. The Johns Hopkins Institutional Review Board approved the study, and all patients provided written informed consent for both the ablation procedure and inclusion in medical research at the time of the procedure.

**MRI ACQUISITION.** Images were obtained using a 1.5-T MRI scanner (Avanto, Siemens, Erlangen,

Germany) and a 6-channel phased-array body coil in combination with a 6-channel spine matrix coil. Contrast enhanced 3-dimensional (3D) fast low-angle shot magnetic resonance angiographic sequences, obtained immediately following intravenous administration of 0.1 mmol/kg of contrast, were used to define LA and PV anatomy (echo time 0.8 ms, repetition time 2.2 ms, in-plane resolution  $1.4 \times 1.4$  mm, slice thickness 1.4 mm). To optimize image quality, patients with persistent AF were kept on antiarrhythmic medications and/or referred for cardioversion prior to MRI. LGE MRI scans were acquired within a range of 15 to 25 minutes (mean  $18.8 \pm 2.4$  min) following 0.2 mmol/kg gadolinium injection (gadopentetate dimeglumine, Bayer Healthcare Pharmaceuticals, Montville, New Jersey) using a fat-saturated 3D inversion recovery-prepared fast spoiled gradient-recalled echo sequence with respiratory navigation and electrocardiographic gating, echo time of 1.52 ms, repetition time of 3.8 ms, in-plane resolution of  $1.3 \times 1.3$ , slice thickness of 2.0 mm, and flip angle of  $10^\circ$ . Trigger time for 3D LGE magnetic resonance images was optimized to acquire data during LA diastole. The optimal inversion time was identified with a scout scan (median 270 ms; range: 240 to 290 ms) to maximize nulling of LA myocardium. A parallel imaging technique, generalized autocalibrating partially parallel acquisition (reduction factor 2), was used.

**IMAGE ANALYSIS.** Pre-ablation LGE magnetic resonance images were processed off-line using QMass MR version 7.4 (Medis, Leiden, the Netherlands). Epicardial and endocardial contours were manually drawn around LA myocardium. The reference point was placed at the anterior base of the LA septum, and the LA myocardium in each axial plane was divided into 25 sectors, as seen in Figure 1. The image intensity ratio (IIR), a previously described LGE MRI technique that normalizes mean myocardial intensities by the mean intensity of the entire blood pool, was calculated for each sector. The methodology for calculation of the IIR and its association with regional voltage by electroanatomic mapping (EAM) were previously validated (6,8). Interobserver and intraobserver variability of our LA analysis was previously assessed and found to be excellent (6).

**MAPPING AND ABLATION PROCEDURE.** A double transatrial septal puncture was performed under fluoroscopic guidance. Intravenous heparin was administered to achieve an activated clotting time >350 s. With hemodynamic and electrocardiographic monitoring, a 3.5-mm irrigated tip with 2-mm inter-electrode spacing ablation catheter (ThermoCool

**FIGURE 1** Left Atrial Segmentation and Voltage Correlation

Endocardial and epicardial contours are manually drawn on axial late gadolinium enhancement (LGE) magnetic resonance images. Fast anatomic mapping points are coregistered and projected to LGE magnetic resonance images. Bipolar voltage of each point is correlated to the image intensity ratio of the nearest segment. AA = ascending aorta; LA = left atrium; LV = left ventricle; MV = mitral valve.

SmartTouch, Biosense Webster) and a decapolar circular mapping catheter, with 25-15 loop diameter, 8-mm interelectrode spacing (LASSO 2515 Variable Loop Eco Nav) were advanced under fluoroscopic guidance to the left atrium. In patients presenting in AF on the day of the procedure, cardioversion was performed prior to image registration and EAM to minimize patient movement and registration errors. If sinus rhythm could not be restored, patients were excluded from analysis. Mapping was performed with an equal distribution of points using a fill threshold of 15 mm, excluding PV ostia. A target of >100 PBP mapping points and >1,000 FAM points was set by consensus at study onset, but the extent of mapping was ultimately left to the operator's discretion. In 19 patients (73%), PBP mapping was performed initially with the ThermoCool SmartTouch ablation catheter and the CARTO3 mapping system (Biosense Webster). In those patients, target contact force >10 g was adopted when obtaining points to ensure adequate tissue-catheter contact. The decision to undergo PBP mapping was left to the operator's discretion. LA magnetic resonance angiography was registered to the voltage map by using standard landmark image registration techniques. A detailed fast anatomic map of the entire left atrium was also created with the LASSO Nav catheter and superimposed on pre-existing magnetic resonance images. An internal

point filter was used to limit data acquisition to within 5 mm from the anatomic shell to reduce mapping in the blood pool. The ConfiDENSE module, which uses impedance measurement to determine tissue proximity for each electrode, was used to assess electrode-tissue contact during FAM. Intra-cardiac echocardiography, orthogonal fluoroscopy, and electrogram characteristics were also used to monitor for adequate contact.

After voltage mapping was complete, circumferential lesions were applied surrounding the PV. If conduction from PV to left atrium persisted despite PV isolation, additional lesions were delivered along the original ablation line at sites of earliest activation on the circular mapping catheter. Entrance block into the PVs was confirmed in all patients as the primary procedural endpoint. To prevent short-term recurrences of AF, pre-procedural antiarrhythmic medications were continued for at least 3 months. No immediate post-operative complications were reported.

**ABLATION MAP REGISTRATION.** FAM-derived voltage maps were registered off-line to pre-ablation LGE images using MASS Research Software (Leiden University Medical Center, Leiden, the Netherlands). Mapping points were extracted from the CARTO workstation and imported into LGE magnetic resonance images using the registration matrix defined during the procedure (Figure 1). An intermediate registration step, performed on ParaView (Kitware, Clifton Park, New York), was needed to reconcile the CARTO coordinate system to the magnetic resonance coordinate system. The mean IIR of the neighboring sector to each mapping point was recorded to evaluate the correlation between bipolar voltage and signal intensity.

**FAM AND PBP VOLTAGE COMPARISON.** Each of the points acquired during FAM and PBP mapping have coordinates (x, y, and z). For each PBP point, the distances between this point and all FAM points were calculated using a custom-written Python script. PBP points were matched in a 1:1 ratio with FAM points using the shortest orthogonal distance.

**STATISTICAL ANALYSIS.** Continuous variables are expressed as mean  $\pm$  SD. Categorical variables are expressed as number (percentage). The unadjusted relationship between IIR and bipolar voltage was initially explored using scatterplots. Because of the skewed nature of bipolar voltage measures, log transformation was used to accommodate modeling within a linear normal framework. A generalized estimating equation (GEE) marginal model with exchangeable

**TABLE 1** Baseline Patient Characteristics

	Total (N = 26)	Paroxysmal AF (n = 14)	Persistent AF (n = 12)	p Value (Paroxysmal vs. Persistent)
Age, yrs	63.1 ± 7.9	62.3 ± 7.3	64.1 ± 8.8	0.575
Male	19 (73)	10 (71)	9 (75)	0.838
Caucasian	22 (85)	12 (86)	10 (83)	0.867
BMI, kg/m <sup>2</sup>	27.2 ± 3.9	26.3 ± 4.9	28.3 ± 2.0	0.199
Hypertension	17 (65%)	10 (71%)	7 (58)	0.484
Diabetes mellitus	1 (4)	0 (0)	1 (8)	0.271
Coronary/vascular disease	4 (15)	2 (14)	2 (17)	0.867
Congestive heart failure	1 (4)	0 (0)	1 (8)	0.271
Prior thromboembolism	2 (8)	2 (14)	0 (0)	0.173
Obstructive sleep apnea	3 (12)	2 (14)	1 (8)	0.636
CHA <sub>2</sub> DS <sub>2</sub> -VASc score	1.8 ± 1.1	1.9 ± 1.0	1.8 ± 1.2	0.957
0	3 (8)	1 (7)	1 (8)	0.918
1	6 (8)	4 (29)	2 (17)	
2	11 (8)	6 (43)	5 (42)	
3	4 (8)	2 (14)	2 (16)	
4	2 (8)	1 (7)	1 (8)	
5	0 (0)	0 (0)	0 (0)	
6	0 (0)	0 (0)	0 (0)	
AF duration, yrs	4.8 ± 4.3	4.6 ± 3.8	5.0 ± 4.9	0.813
Prior ablations	0.7 ± 0.6	0.6 ± 0.5	0.8 ± 0.7	0.434
0	9 (35)	5 (36)	4 (33)	0.276
1	15 (58)	9 (75)	6 (50)	
2	2 (7)	0 (0)	2 (17)	
Prior ablation strategy				0.565
Radiofrequency ablation	11 (42)	7 (50)	4 (33)	
Cryoballoon ablation	6 (23)	2 (14)	4 (33)	
Smoking				0.532
Never	12 (46)	7 (50)	5 (42)	
Prior	13 (50)	7 (50)	6 (50)	
Current	1 (4)	0 (0)	1 (8)	
DCV pre-mapping	7 (27)	0 (0)	7 (58)	0.0008
LVEF, %	61.0 ± 9.4	65.9 ± 6.1	54.8 ± 9.4	0.002
LA end-systolic area, cm <sup>2</sup>	26.8 ± 8.4	24.5 ± 5.9	29.3 ± 10.1	0.157

Values are mean ± SD or n (%).  
AF = atrial fibrillation; BMI = body mass index; DCV = direct-current cardioversion; LA = left atrial; LVEF = left ventricular ejection fraction.

working correlation structure was used to examine the association between bipolar voltage and IIR. The model was clustered by patient and adjusted for age, LA end-diastolic area, sex, body mass index, CHA<sub>2</sub>DS<sub>2</sub>-VASc score, AF type, AF duration, left ventricular ejection fraction, and history of ablation. Similarly, the correlation between log-transformed PBP voltage and FAM voltage was assessed using an adjusted, patient-clustered GEE marginal model with exchangeable working correlation structure. Receiver-operator characteristic analysis was used to assess the sensitivity and specificity of FAM voltage with PBP-derived fibrosis. Two-sided p values < 0.05 were considered to indicate statistical significance. Statistical analysis was performed using STATA version 13 (StataCorp, College Station, Texas).

## RESULTS

**PATIENT CHARACTERISTICS.** We prospectively enrolled 26 patients (19 men [73%], mean age 63.1 ± 7.9 years). **Table 1** summarizes the baseline characteristics of patients. Fourteen patients (54%) had paroxysmal AF, and 9 (35%) were undergoing their first ablation procedure. The mean body mass index was 27.2 ± 3.9 kg/m<sup>2</sup>, and the median CHA<sub>2</sub>DS<sub>2</sub>-VASc score was 2 (interquartile range: 1 to 2). Patients with persistent AF had lower left ventricular ejection fractions and higher rates of pre-mapping cardioversion than those with paroxysmal AF; there were no other statistically significant differences between the 2 patient groups.

**IMAGE ANALYSIS.** MRI studies were performed a median of 1 day (interquartile range: 0 to 6 days) prior to the procedure. Image intensity in arbitrary units and IIR were measured for each sector. The relationship of signal intensity versus IIR and the distribution of each measure are plotted in **Online Figure 1**. Although image intensity and IIR measures had a linear relationship within each patient, the slope of this association varied across patients, with a range of 7.79 to 75.62 arbitrary units. In contrast to the right-skewed distribution of image intensity, IIR displayed a normal distribution (**Figure 2**).

**CORRELATION OF FAM VOLTAGE AND IIR.** Seven of the 26 patients (26%) presented in AF at the beginning of the procedure and required cardioversion prior to mapping. A mean of 734 FAM points (range: 397 to 1,089 points) were acquired per patient, and a total of 19,087 points were registered to LGE magnetic resonance images. The average bipolar voltage was 0.88 ± 1.27 mV, and the average IIR was 1.08 ± 0.18. Local IIR was strongly associated with bipolar voltage; however, the relationship was more linear with log transformation of bipolar voltage measures (**Figure 2**). Using a GEE model, clustered by patient and adjusted for age, sex, CHA<sub>2</sub>DS<sub>2</sub>-VASc score, AF type, AF duration, previous ablations, LA end-diastolic area, and left ventricular ejection fraction, each unit increase in local IIR was associated with 57% decrease in LA bipolar voltage (p < 0.0001) (**Table 2**). On the basis of unadjusted analyses, local IIR thresholds of >0.74 corresponded to bipolar voltage <0.5 mV.

We performed a stratified analysis in patients with prior ablations versus first-time ablation. The findings are summarized in **Online Tables 1 and 2**. In brief, mean bipolar voltage was 1.20 mV in ablation-naive patients and 0.78 mV in previously ablated patients. The adjusted relationship between IIR and FAM

**FIGURE 2** Histograms of Distribution of MRI-Signal Intensity and Bipolar Voltage

**(Top)** Histogram of signal intensity and image intensity ratio as well as fast-anatomical-mapping and point-by-point bipolar voltage and its log transformation **(middle and bottom panels)**. FAM = fast anatomic mapping; PBP = point-by-point.

voltage was significantly negative in both groups. We also performed a subset analysis of 4 patients with the least percentage of low-voltage mapping points (characterized by voltage  $\leq 0.5$  mV). All 4 patients had paroxysmal AF, with a mean bipolar voltage of 1.84 mV and an average proportion of low-voltage points of  $8.7 \pm 4.1\%$ . As seen in [Online Table 3](#), the adjusted relationship between IIR and

FAM-measured bipolar voltage was significantly negative. It was similarly negative upon stratifying mapping points into “normal” voltage ( $>0.5$  mV) and “low” voltage ( $\leq 0.5$  mV) ([Online Tables 4 and 5](#)).

**FAM AND PBP VOLTAGE COMPARISON.** Serial PBP mapping and FAM were performed in 19 patients. A mean of 82 points per patient were acquired using

**TABLE 2 Results of the Generalized Estimating Equation Model**

Log FAM Bipolar Voltage (mV)	Univariate		Multivariate	
	Coefficient	p Value	Coefficient	p Value
IIR (per U)	-0.90	<0.0001	-0.85	<0.0001
Age (per yr)	-0.03	0.054	-0.01	0.661
Male	-0.60	0.019	-0.53	0.014
BMI (per kg/m <sup>2</sup> )	0.06	0.036	0.07	0.004
CHA <sub>2</sub> DS <sub>2</sub> -VASc score (per U)	-0.02	0.162	-0.13	0.187
Persistent AF	-0.14	0.565	0.02	0.911
Repeat ablation	-0.53	0.027	-0.43	0.026
AF duration (per yr)	-0.05	0.076	-0.06	0.012
LA area (per cm <sup>2</sup> )	-0.02	0.162	-0.001	0.952
LVEF (per U %)	0.01	0.415	0.01	0.308

FAM = fast anatomic mapping; IIR = image intensity ratio; other abbreviations as in Table 1.

the PBP technique, and a total of 1,554 points were matched in a 1:1 ratio to the nearest FAM-EAM point. In the matched 1,554 voltage-point pairs, mean PBP voltage was  $0.91 \pm 0.99$  mV, and corresponding mean FAM voltage was  $1.20 \pm 1.36$  mV. Distribution of PBP voltage and its log transformation are displayed in Figure 2. A scatterplot of voltage measurements correlating PBP-EAM and FAM-EAM is displayed in Online Figure 2. Using an adjusted, patient-clustered GEE model, log FAM bipolar voltage was significantly associated with log PBP bipolar voltage ( $\beta = 0.36$ ,  $p < 0.0001$ ), as shown in Table 3. Individual log PBP versus log FAM voltage linear relationships are demonstrated in Online Figure 3. At low voltages (<40th percentile), the FAM distribution was shifted to the left compared with PBP mapping (Table 4). On FAM, the 20th and 40th percentiles were 0.08 and 0.25 mV, respectively, compared with 0.18 and 0.42 mV on PBP mapping. At intermediate voltages (40th to 80th percentiles), FAM and PBP voltages were

**TABLE 3 Results of the Generalized Estimating Equation Model**

Log Bipolar PBP Voltage (mV)	Univariate		Multivariate	
	Coefficient	p Value	Coefficient	p Value
Log bipolar FAM voltage (mV)	0.38	<0.0001	0.36	<0.0001
Age (per yr)	0.02	0.382	0.02	0.366
Male	-0.29	0.426	-0.45	0.042
BMI (per kg/m <sup>2</sup> )	0.04	0.356	-0.01	0.661
CHA <sub>2</sub> DS <sub>2</sub> -VASc score (per U)	-0.12	0.428	-0.17	0.244
Persistent AF	-0.25	0.422	-0.08	0.767
Repeat ablation	-0.72	0.004	-0.62	0.033
AF duration (per yr)	-0.01	0.935	0.02	0.518
LA area (per cm <sup>2</sup> )	-0.01	0.797	0.01	0.490
LVEF (per U %)	0.02	0.170	0.01	0.510

PBP = point-by-point; other abbreviations as in Table 2.

overlapping (the 60th and 80th percentiles on FAM were 0.69 and 1.52 mV, respectively, vs. 0.73 and 1.54 mV on PBP mapping). At high voltages (>80th percentile), FAM voltage exceeded PBP voltage (the 99th percentile was 6.26 mV on FAM vs. 4.65 mV on PBP mapping). Receiver-operating characteristic analysis of FAM voltage predicting low voltage (<0.5 mV) on PBP mapping showed an area under the curve of 0.75 (Online Figure 4). The FAM voltage with the highest sensitivity and specificity was 0.58 mV (63% sensitivity, 77% specificity).

Figure 3 compares IIR maps and FAM and PBP voltage maps in an ablation-naive patient with minimal baseline fibrosis, an ablation-naive patient with posterior wall fibrosis, and a patient with a history of radiofrequency AF ablation. The figure demonstrates a good correlation of LA fibrotic substrate between the 3 modalities. Similarly, Online Figure 5 demonstrates an anterior view of IIR and FAM voltage maps in 2 patients.

## DISCUSSION

In this study, we sought to compare LA substrate using different modalities (LGE MRI, FAM, and PBP mapping) in patients presenting for AF ablation. The major findings are as follows: 1) voltage acquired by FAM using a decapolar circular (LASSO 2515 Variable Loop Eco Nav) mapping catheter with a small electrode size correlated with IIR on LGE MRI; and 2) voltage acquired by FAM correlated with voltage on PBP mapping. These findings might have a significant impact on the interpretation of substrate mapping in ablation strategies targeting extra-PV LA remodeling.

**FAM VERSUS IIR CORRELATION.** A variety of novel AF substrate-mapping approaches have been introduced in recent years, including dominant-frequency analysis (9,10), Shannon entropy (11), complex fractionated electrogram activity (3), focal impulse and rotor modulation (12), bipolar electrogram amplitude (13), and LGE MRI (6-8). Prior studies have shown poor correlation among various techniques (14-17), with the exception of bipolar voltage, which correlates well with LGE MRI (6,7). In a landmark study, Oakes et al. (7) found a positive correlation ( $r^2 = 0.61$ ) between the extent of LGE on MRI with low-voltage zones. Kapa et al. (18) projected EAM points, acquired in sinus rhythm, onto LGE magnetic resonance images and found that mean bipolar voltage in areas of MRI fibrosis was significantly lower than in non-fibrosis areas. In a study by Khurram et al. (6), IIR was used to normalize signal intensities, and intracardiac



**TABLE 4** Correlations of Voltage Thresholds Between Voltage-Mapping Modalities

	Percentile					
	5th (95% CI)	20th (95% CI)	40th (95% CI)	60th (95% CI)	80th (95% CI)	99th (95% CI)
FAM voltage	0.03 (0.02-0.03)	0.09 (0.09-0.10)	0.28 (0.27-0.29)	0.69 (0.67-0.71)	1.52 (1.50-1.54)	6.26 (5.99-6.45)
PBP voltage	0.04 (0.03-0.04)	0.18 (0.16-0.20)	0.42 (0.39-0.45)	0.73 (0.68-0.79)	1.54 (1.41-1.67)	4.65 (4.27-4.79)

CI = confidence interval; other abbreviations as in Tables 2 and 3.

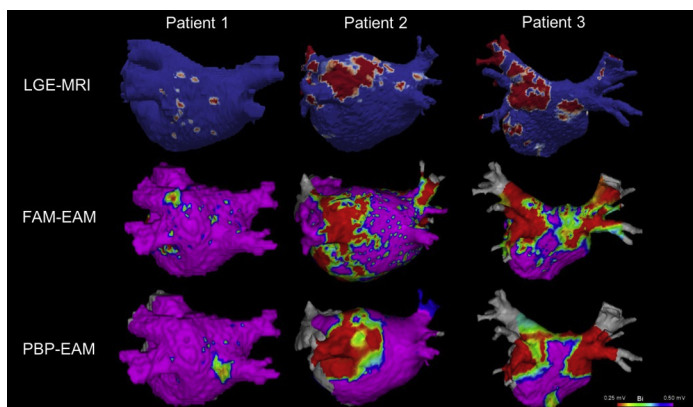
mapping points were registered to LGE images. In a multivariate model, each unit increase in IIR was associated with 91.3% decrease in bipolar LA voltage ( $p < 0.0001$ ). In the aforementioned studies, voltage maps were created using PBP acquisition (with or without force sensing). In our investigation, we used a multipolar circular mapping catheter with small electrodes (~1 mm) to acquire a detailed (>1,000 points) voltage maps of the left atrium. In a semi-automated fashion, we were able to merge mapping points onto LGE images and correlate bipolar voltage with neighboring IIR on MRI. Using an adjusted patient-clustered GEE model, each unit increase in local IIR was associated with a 57% decrease in local bipolar LA voltage, demonstrating a good correlation between FAM voltage and IIR in mapping AF substrate.

The distinction between ablation-induced scar and AF-related interstitial fibrosis is an important one to make. On the microscopic level, these represent different pathological processes: necrosis and replacement fibrosis in ablation-related scar and interstitial, diffuse collagen deposition in native AF-related fibrosis. Our group has recently established the distinct characteristics of these processes on LGE MRI (19). Regions with ablation-induced scar exhibit increased contrast uptake (higher IIR), likely signifying higher scar density, compared with regions with native interstitial fibrosis. For these reasons, we performed a stratified analysis in patients with prior ablations versus first-time ablation and found an inverse relationship between IIR and FAM-measured bipolar voltage in both groups.

These findings further demonstrate the utility of MRI in mapping abnormal myocardium as well as guiding AF ablation. In particular, long-term procedural outcomes of patients with persistent AF remain dismal (~50%) (20) and are attributed to extensive atrial fibrotic remodeling not targeted by PV isolation (7,21-23). This is supported by the DECAAF (Delayed-Enhancement MRI Determinant of Successful Radiofrequency Catheter Ablation of Atrial Fibrillation) study, which showed that pre-ablation fibrotic tissue is a major predictor of ablation outcome (21). The

efficacy of targeting atrial fibrosis tissue during persistent AF ablation will be assessed in the much awaited DECAAF II study (NCT02529319).

**FAM VERSUS PBP CORRELATION.** Several studies have shown smaller low-voltage areas and higher mean regional bipolar voltages on FAM compared with PBP mapping (24-26). In line with previous studies, the mean PBP bipolar voltage in our study was  $0.91 \pm 0.99$  mV and mean FAM voltage was  $1.20 \pm 1.36$  mV. Log FAM bipolar voltage was significantly associated with log PBP bipolar voltage ( $\beta = 0.36$ ,  $p < 0.0001$ ). Several factors might influence bipolar voltage amplitude, such as catheter characteristics (electrode size, interelectrode spacing), wave front of activation, electrode-wall contact force, and angle (25,27,28). In fact, our linear ablation catheters had

**FIGURE 3** Illustration of Image Intensity Ratio, FAM-Voltage and PBP-Voltage Maps

This illustration compares IIR maps (Top Panel), FAM voltage maps (Middle Panel), and PBP voltage maps (Bottom Panel) in an ablation-naïve patient with minimal baseline fibrosis (Patient 1), an ablation-naïve patient with posterior wall fibrosis (Patient 2), and a patient with a history of prior radiofrequency ablation for AF (Patient 3). The figure demonstrates good correlation between the 3 mapping modalities. Of note, no voltage mapping was done in PV ostia and this area was excluded from any multimodal correlation analysis. Indeed, the thin myocardial sleeves in that area render voltage and magnetic resonance imaging less suited for myocardial substrate mapping. EAM = electroanatomic mapping; FAM = fast anatomic mapping; LGE = late gadolinium enhancement; PBP = point-by-point.

3.5-mm distal electrodes with interelectrode spacing of about 5 mm, whereas multipolar circular catheters had 20 poles, 1-mm electrode size, and 2-6-2-mm interelectrode spacing. Smaller electrode sizes allow increased mapping resolution and a greater ability to discriminate between surviving myocardial fibers and fibrosis; at the same time, smaller electrode size may increase susceptibility of the electrodes to far-field sensing. This was seen on the voltage distribution of the 2 modalities. At low voltages (<40th percentile), the FAM distribution was shifted to the left compared with PBP mapping, which may reflect increased capacity of FAM to detect fibrotic fibers interspaced with normal myocytes, whereas at high voltages (>80th percentile), FAM voltages exceeded PBP voltages (the 99th percentile was 6.26 mV on FAM vs. 4.65 mV on PBP mapping), which may be due to far-field sensing. Moreover, although adequate tissue contact during mapping with the PBP catheter was confirmed by contact force measurement, such data were not available for FAM catheters, which could account for some of the voltage differences.

Voltage mapping in the left atrium is increasingly used to guide AF ablation, the accuracy of which relies heavily on the accurate identification of LA fibrosis (3,29-32). Our work raises awareness of the differences in substrate delineation among various mapping techniques and catheters available that would influence the interpretation of mapped substrate. Based in part on our observations in the present study, our current clinical practice is to use 0.25 to 0.5 mV as a voltage range indicative of abnormal fibrotic tissue and <0.25 mV as consistent with scar when performing either FAM or PBP mapping; in cases in which LGE MRI data are used for scar assessment, we use an IIR value of >1.22 on the basis of equations derived from our work. An important caveat, of course, is that data are lacking regarding how these parameters correlate with scar assessment on the basis of actual tissue evaluation.

**STUDY LIMITATIONS.** There were a number of limitations to this study. The gold standard in assessing atrial substrate in AF is tissue evaluation. In patients with AF, the muscle bundles were surrounded by thick connective tissue fibers. These thick fibers were also interspaced between the single muscle cells (33). The resolution of PBP or fast anatomic maps using bipolar electrodes is very poor compared with this gold standard, regardless of the mapping density. The fill threshold used by the mapping systems compounds the problem by interpolating areas that are not directly touched by the catheter. Although

LGE MRI characterizes the entirety of atrial tissue within each axial slice, the thickness of the slices adds error and interpolation as well. Given the limitations of both EAM and IIR methodologies, in addition to the fact that IIR has not been validated against tissue samples as of yet, the results of this study should be interpreted as 2 methods in relative agreement rather than against a gold standard. This study includes a relatively small cohort of patients who were heterogeneous with regard to prior ablations. However, such heterogeneity did not modify the association between IIR and voltage and demonstrated the wide applicability of LGE MRI in detecting abnormal LA myocardium. Voltage can be affected by tissue contact. Although it cannot be completely ruled out that diminished bipolar voltage on FAM was measured as a consequence of reduced catheter contact (because FAM catheters are not capable of force sensing), it is unlikely to account for the differences between catheters: 1) every attempt was made to ensure adequate contact by means of splaying of the electrodes in contact with the myocardium on fluoroscopy; 2) points were accepted only if they were within 5 mm of the anatomic shell; and 3) patient-clustered association of IIR with voltage suggests that on average, the measures represent LA myocardial tissue characteristics. Our results may also be affected by positional errors when registering ablation points to the magnetic resonance-based segmentation; however, the extent of positional errors appears to be very low on the basis of prior validation studies of our technique (34). For the PBP versus FAM correlation, distances between PBP and FAM points were calculated using the coordinates extracted from CARTO. As such, our results are contingent on having an accurate 3D registration of PBP and FAM-EAM and would be affected by patient movement between the 2 mapping modalities (as seen in individual graphs in Figure 3).

## CONCLUSIONS

LGE MRI, FAM, and PBP mapping show a good degree of correlation for delineating fibrosis-based AF substrate. Each approach has fundamental technical characteristics, the awareness of which allows proper assessment of electroanatomic AF substrate and potentially avoidance of unnecessary ablation.

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## PERSPECTIVES

**COMPETENCY IN MEDICAL KNOWLEDGE:** Substrate modification, targeting low-voltage areas and/or enhancing areas on LGE MRI, has been proposed as an individualized ablation approach. As seen in our study, differences in catheters used for LA substrate mapping may explain the variable results of fibrosis-based ablation strategies. Our findings reflect the importance of adjusting voltage and/or IIR thresholds when examining AF substrate using different modalities.

**TRANSLATIONAL OUTLOOK:** Intracardiac electrograms and LGE MRI can delineate fibrosis in the left atria of patients with AF, and this study has shown good correlation between the 2 modalities. However, histological confirmation remains of paramount necessity to establish the sensitivity and specificity of these less invasive tools compared with a gold standard. Moreover, a study on healthy volunteers without AF is needed to validate the associations among the various modalities in structurally normal atria.

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**KEY WORDS** atrial fibrillation, fibrosis, magnetic resonance imaging, substrate, voltage mapping

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**APPENDIX** For supplemental figures and tables, please see the online version of this paper.