Electroanatomic Mapping to Determine Scar Regions in Patients with Atrial Fibrillation

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Abstract
Left atrial voltage maps are routinely acquired during electroanatomic mapping in patients undergoing catheter ablation for atrial fibrillation (AF). For patients, who have prior catheter ablation when they are in sinus rhythm (SR), the voltage map can be used to identify low voltage areas (LVAs) using a threshold of 0.2 - 0.45 mV. However, such a voltage threshold for maps acquired during AF has not been well established. A prerequisite for defining a voltage threshold is to maximize the topologically matched LVAs between the electroanatomic mapping acquired during AF and SR. This paper demonstrates a new technique to improve the sensitivity and specificity of the matched LVA. This is achieved by computing omni-directional bipolar voltages and applying Gaussian Process Regression based interpolation to derive the AF map. The proposed method is evaluated on a test cohort of 7 male patients, and a total of 46,589 data points were included in analysis. The LVAs in the posterior left atrium and pulmonary vein junction are determined using the standard method and the proposed method. Overall, the proposed method showed patient-specific sensitivity and specificity in matching LVAs of 75.70% and 65.55% for a geometric mean of 70.69%. On average, there was an improvement of 3.00% in the geometric mean, 7.88% improvement in sensitivity, 0.30% improvement in specificity compared to the standard method. The results show that the proposed method is an improvement in matching LVA. This may help develop the voltage threshold to better identify LVA in the left atrium for patients in AF.

Keywords
atrial fibrillation, electroanatomic mapping, voltage map, omni-polar, omni-directional bipolar, Gaussian process regression interpolation

Disciplines
Computer Engineering | Electrical and Computer Engineering

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Electroanatomic Mapping to determine Scar Regions in patients with Atrial Fibrillation

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Abstract—Left atrial voltage maps are routinely acquired during electroanatomic mapping in patients undergoing catheter ablation for atrial fibrillation (AF). For patients, who have prior catheter ablation when they are in sinus rhythm (SR), the voltage map can be used to identify low voltage areas (LVAs) using a threshold of 0.2 - 0.45 mV. However, such a voltage threshold for maps acquired during AF has not been well established. A prerequisite for defining a voltage threshold is to maximize the topologically matched LVAs between the electroanatomic mapping acquired during AF and SR. This paper demonstrates a new technique to improve the sensitivity and specificity of the matched LVA. This is achieved by computing omni-directional bipolar voltages and applying Gaussian Process Regression based interpolation to derive the AF map. The proposed method is evaluated on a test cohort of 7 male patients, and a total of 46,589 data points were included in analysis. The LVAs in the posterior left atrium and pulmonary vein junction are determined using the standard method and the proposed method. Overall, the proposed method showed patient-specific sensitivity and specificity in matching LVAs of 75.70% and 65.55% for a geometric mean of 70.69%. On average, there was an improvement of 3.00% in the geometric mean, 7.88% improvement in sensitivity, 0.30% improvement in specificity compared to the standard method. The results show that the proposed method is an improvement in matching LVA. This may help develop the voltage threshold to better identify LVA in the left atrium for patients in AF.

I. INTRODUCTION

The voltage map is one form of data provided by electroanatomic mapping (EAM) and is often used to demarcate low voltage areas (LVAs) and preserved voltage areas during catheter ablation therapy to treat atrial fibrillation (AF). LVAs correspond to areas of diseased atrium (fibrosis) or dense scars from prior ablations. Identifying LVA can help in planning ablation strategies especially in patients requiring repeat ablation procedures for arrhythmia recurrences, for example, AF and atypical atrial flutter.

The cutoff threshold voltage for a voltage map that determines LVAs has been established for maps obtained while the patient is in sinus rhythm (SR) [1][2]. However, this same threshold is not applicable to maps collected during AF. For example, Fig. 1(a) depicts the SR map with a cutoff threshold of 0.45 mV. The regions below the threshold are clearly delineated from the healthy regions (magenta). However, in (b), applying the same threshold to the AF map for the same patient distorts the LVAs distribution. After adjusting to a lower threshold, as in (c), the LVAs on the AF map are restored to match the SR map.

Several studies have shown that the local atrial signal acquired during AF is lower than in SR. Thus, identifying LVA during EAM in AF should require a lower cutoff voltage [3]. However, determining a consistent threshold that can be applied to all patients remains challenging. A prerequisite is determining the best match of LVAs that can be obtained between the SR map and the AF map and thereby finding the best threshold to be applied on a patient-by-patient basis.

Problem Statement: Given a set of measurements during SR and AF for a patient, maximize the topologically matched LVAs between the derived SR and AF map and determine the best patient-specific cutoff voltage threshold.

In this paper, we demonstrate a method of deriving the AF map which is robust to noise and error in the measurements and improves the patient-specific sensitivity and specificity of matched LVAs in comparison to the standard method through the following the contributions:

• Compute omni-directional bipolar voltages which are invariant to the orientation of the catheter, thus improving signal strength during AF.
• Apply Gaussian process regression (GPR) interpolation which improves the accuracy of LVA detection in regions of the atrium with lower measurement density.

II. BACKGROUND: STANDARD VOLTAGE MAP

Fig. 2 depicts the steps of deriving the current standard left atrial (LA) voltage map during catheter ablation of AF. Initially, as in (1), a 3D anatomical mesh is generated by manipulating a multi-electrode mapping catheter (Lasso or Pentaray) to different parts of the LA [4]. As the mesh is being created, recordings of 2.5 seconds of electrogram are
Fig. 2: Standard method of deriving voltage map. (1) Anatomical mesh is computed from catheter locations. (2) Bipolar voltages are computed from measurements of electrogram at points along the mesh. (3) The bipolar voltages are interpolated to the remainder of the mesh to derive the voltage map.

Fig. 3: Bipolar voltages are computed as the peak-to-peak voltage in a pre-specified window within a 2.5 seconds recording of the endocardium electrogram. (2) shows the locations of all such sampled points on the LA mesh. The voltage value is interpolated to the remaining areas of the mesh to derive the final voltage map as shown in (3). The colors of the voltage map are based on a pre-specified cutoff threshold, where areas above the threshold are marked in magenta, and areas below the threshold are considered LVAs.

Fig. 3 shows that within the 2.5 seconds, a 300 ms time window is defined relative to the QRS peak, and the peak-to-peak voltage is computed within the time window.

III. ROBUST METHOD FOR VOLTAGE MAP DERIVATION

Our proposed robust method deriving the voltage map differs from the standard method in terms of two components: a) omni-directional bipolar voltage and b) GPR-based interpolation.

a) Omni-directional bipolar voltage: Fig. 4 shows a limitation of bipolar recording, the dependency on electrode orientation. If the bipolar placement is parallel to the isoelectric potential line, the bipolar recording will be zero, which does not reflect the local electric activity [5]. To reduce such dependency, we derive the omni-directional bipolar voltage. For each sample point, we select the unipolar electrogram recorded in the vicinity of the sample and compute all possible bipolar electrogram from this set. We approximate the omni-directional bipolar voltage as the largest bipolar amplitude from this set. Fig. 4 (1) Blue depicts the histogram of the voltages for a patient. Red shows how the corresponding voltages are amplified. In the tail portion, the voltages of some LVA have increased above the threshold to be classified as healthy tissue. (2) and (3) exemplifies how the original red areas of voltage ~0.06 mV are enhanced into yellow and green areas of voltage ~0.17 mV.

b) Gaussian process regression based interpolation: If we model the endocardium as a surface and define the entire set of samples as, \( D = \{ x_n, y_n \}_{n=1}^N \), where inputs \( X = \{ x_n \}_{n=1}^N \) correspond to the locations on the mesh and \( y = \{ y_n \}_{n=1}^N \) are the voltage value at that location. Interpolating from these measured samples to the remainder of the mesh can be thought of as determining the estimates of the voltages at locations \( X^* \). The two major sources of interpolation error are the low measurement density and measurement noise. Both of these can be accounted for by modeling them using a Gaussian process \( GP(m(x), k(x, x')) \) [6], which is characterized by the mean \( m(x) \) and the covariance \( k(x, x') \) kernel functions. We assume the common zero mean function and use the squared exponential function \( k(x, x') = \exp(-||x - x'||^2/(2 \cdot l^2)) \). Fig. 5(1) shows how the standard interpolation can result in regions that are classified as LVAs, due to interpolation error. (2) shows GPR-based interpolation can improve the boundaries of LVAs by considering the surrounding measurements.

For determining the optimum threshold, a search in the range of 0-0.45 mV is performed to maximize the product of sensitivity and specificity. Here, sensitivity = \( \frac{TP}{TP + FN} \)
Fig. 6: Experimental setup. For each patient: From the SR map (1), LVA are determined (2) as well as the region of interest (3). Both (2) and (3) are topologically transferred to the AF map (4) and (5). (4) (5) are intersected to obtain the true LVAs on the AF map (6). The optimal patient-specific threshold is determined by maximizing the product of sensitivity and specificity according to (8).

and specificity \( = \frac{TN}{TN + FP} \), where a true positive (TP) indicates that the corresponding face on the anatomical mesh is detected as a LVA and the true label is a LVA.

IV. EXPERIMENTAL SETUP

The proposed method of voltage map derivation was evaluated on a test cohort of 7 patients that underwent repeat catheter ablation for AF at the Hospital of the University of Pennsylvania. Patient demographics are: age 66 ± 0.7 years, height 5’10” ± 4”, weight 238.9 ± 50 lb and ejection fraction 49.4%. Details are in Table I. For each patient, voltage maps were sequentially obtained using Carto3 (Biosense Webster) during SR and AF or vice-versa. Fill threshold was 5 mm, and filters were set at 2 to 240 Hz for unipolar electrograms (EGMs), 16-500 Hz for bipolar EGMs, and 0.5-200 Hz for surface electrogram recordings. We assumed that the LV A observed based on the SR voltage map is the ground truth.

The overall evaluation process is depicted in Fig. 6. From the original data (1), we apply the standard 0.45 mV cutoff threshold on the SR map to identify regions of LVA (2). We select a region of interest (ROI) on the SR map (3), which consists of the posterior LA and pulmonary vein (PV) junctions. Both the LVA and the ROI is transferred to the AF mesh and intersected to form the final LVA on the AF map (4),(5),(6). Then result (7) is determined as shown in (8) in terms of sensitivity and specificity.

V. RESULTS AND DISCUSSION

A total of 46,589 data points were included in analysis, that was on average 6,656 data points for each of the 7 patients. Table II summarizes the results of evaluation for each patient. On average, our proposed method showed a sensitivity and specificity of 75.70% and 66.55%, respectively. This was a 3.00% improvement in the geometric mean compared to the standard method. Moreover, our proposed method exhibited a 7.88% improvement in sensitivity and a 0.30% improvement in specificity. ROC curves were obtained for each of the methods and the area under the curve was computed as shown in Fig. 7. Our proposed method showed an average of 3.91% improvement in terms of the area under the curve (AUC).

Fig. 7: ROC curve comparing, baseline method vs proposed method using omni-directional bipolar voltages and GPR-based interpolation. The proposed method shows improved or similar performance across various thresholds.

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underestimates the interpolated voltage when spurious low-voltage measurement exist. In this case the combination of omni-directional bipolar voltages enhances the signal, and GPR-based interpolation filters such noise, preventing classification as LVAs.

In patient 7, our proposed method did not improve performance. Upon further inspection, we discovered that 90% of the ROI was LVA as shown in Fig. 9. This patient had undergone prior extensive surgical ablation and so had extensive areas of dense scar in the ROI, making discrimination difficult. Optimizing the threshold with a different criterion which accounts for this bias may result in a better outcome.

**Limitations:** The results of our study are limited in scope, mostly in part by the small cohort size. Bias due to the particular demographics of the cohort may have affected the results and different results may be obtained with a larger cohort. Selection of different hyperparameters for the method, such as a different kernel for GPR-based regression may affect the performance. Finally, establishing a standard protocol for obtaining data may lead to improvement.

**VI. CONCLUSION**

In this work, we have presented a method for deriving the voltage map during AF and comparing it to voltage maps acquired during SR. Our method computes omni-directional bipolar voltages from the measurements and utilizes GPR-based interpolation to derive the voltage map. Evaluation on the test cohort showed that, in general, the method improved the patient-specific sensitivity and specificity in determining LVAs of the AF map compared to the standard method, though some exceptions exist. This improvement in matched areas between the maps is significant and has important practical implications as clinicians interpret voltage maps according to the areas and not by the individual point measurements. More accurate information about LVA distribution is helpful to clinicians in planning ablation strategies for patients who require repeat catheter ablation for arrhythmia recurrences. Immediate future work is to apply the method over a larger cohort. In a practical clinical setting, patient-to-patient variability may need to be accounted for in the criterion. Overall, the results provide evidence that the proposed method improves the detection of LVAs in AF maps. Because of the robustness to measurement noise and interpolation error, the proposed method could lead to a more consistent criterion.

**REFERENCES**


