Characteristics of Complex Fractionated Atrial Electrogram in the Electroanatomically Remodeled Left Atrium of Patients With Atrial Fibrillation

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Background: Complex fractionated atrial electrogram (CFAE) guided ablation is effective in some patients with persistent atrial fibrillation (PeAF), but the pattern of CFAE may be different in the remodeled left atrium (LA).

Methods and Results: In 100 AF patients (83 males, 55.0±10.6 years old) with AF (51 paroxysmal AF (PAF), 49 PeAF) who underwent catheter ablation, CFAE cycle length (CL) and distribution (NavX 3D map) were compared according to the LA volume (3D-CT) and endocardial voltage (during high right atrial pacing 500-ms (VolPACE) and AF (VolAF; NavX). The mean CFAE-CL was longer (P=0.003) and the % area CFAE was smaller (P=0.006) in patients with LA ≥125ml than those with <125ml. The mean CFAE-CL was longer in patients with VolPACE <1.7mV than those with ≥1.7mV (P=0.002) and in VolAF <0.7mV than ≥0.7mV (P<0.001). The % area CFAE was smaller in patients with VolPACE <1.7mV than those with ≥1.7mV (P=0.006). The incidence of septal CFAE was consistently high, regardless of the degree of LA remodeling.

Conclusions: In the AF patients with an electroanatomically remodeled LA, the % area of CFAE was smaller and mean CFAE-CL was longer than in those with a less remodeled LA. However, the majority of CFAE are consistently positioned on the septum in the remodeled LA. (Circ J 2010; 74: 1557–1563)

Key Words: Atrial fibrillation; Computed tomography; Left atrium; Morphological remodeling; Voltage mapping

Although radiofrequency catheter ablation (RFCA) of atrial fibrillation (AF) is an effective treatment modality for controlling AF,1 circumferential pulmonary vein (PV) ablation is not enough in some patients, especially those with persistent AF (PeAF).2,3 Nademanee et al reported that complex fractionated atrial electrogram (CFAE)-guided left atrial (LA) ablation terminated sustained AF in 95% of patients, resulting in excellent clinical outcomes (1 year AF-free rate, 91% with combined ibutilide medication in 28% of patients).4 They also reported that maintenance of sinus rhythm (SR) after CFAE-guided LA ablation improved mortality rates.5 However, clinical outcomes of CFAE-guided ablation studies have been quite variable6–10 and the electrophysiologic mechanism of CFAE has not yet been clearly elucidated. We previously characterized CFAE as being predominantly located at the septum, roof, and LA appendage (LAA), with low voltage surrounded by a high voltage area and low conduction velocity in patients with AF.11 We also demonstrated electroanatomical remodeling of the LA in patients with PeAF, compared with paroxysmal AF (PAF), by analyzing LA volume and LA voltage.12,13 However, CFAE in patients with an electroanatomically remodeled LA has not been clearly characterized yet. Therefore, we hypothesized that the CFAE pattern (ie, CFAE-cycle length (CL) or distribution) will be different in patients with a remodeled atrium. In the present study we compared the CFAE-CL and % area of CFAE in patients with an enlarged LA and those without LA enlargement by analyzing 3D-computed tomographic (CT) images and NavX voltage mapping. We also compared the distribution of CFAE in patients with low and high LA voltages.

Methods

Patient Selection
The study protocol was approved by the Institutional Review Board and all patients provided written informed consent. We enrolled 100 patients with AF (M/F=83/17, mean age 55.0±10.6 years) who underwent RFCA. Among them, 51 patients had PAF, and 49 had PeAF. The exclusion criteria

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were: (1) permanent AF refractory to electrical cardioversion; (2) LA >50 mm measured on echocardiogram; (3) AF with rheumatic valvular disease; (4) associated structural heart disease other than left ventricular hypertrophy; (5) prior AF ablation; and (6) SR not maintained for LA voltage mapping before RFCA. Patients with LA thrombus were excluded by transesophageal echocardiography. We imaged all patients with 3D spiral CT (64 Channel, Light Speed Volume CT, Philips, Brilliance 63, the Netherlands) to visually define the anatomy of the LA and PVs.

**Electrophysiological Mapping Procedure**

Intracardiac electrograms were recorded using the Prucka CardioLab™ Electrophysiology system (General Electric Medical Systems) and the catheter ablation procedure was performed on all patients using 3-D electroanatomical mapping (NavX, St Jude Medical) merged with 3-D spiral CT. Before performing catheter ablation, we generated a 3-D electroanatomical map of the LA, a CFAE-CL map, and a voltage map by obtaining contact bipolar electrograms filtered from 32 to 300 Hz. If the initial rhythm was sustained AF, the CFAE-CL map was acquired first. To generate a CFAE-CL map, contact bipolar electrograms of AF were obtained for longer than 6 s at each site approximately equally distributed within the LA. The CFAE-CL was calculated by averaging the peak-to-peak intervals of 6-s recordings of bipolar electrograms. The detection of the CFAE-CL also possessed a downstroke morphology in which the leading local-maximum and the trailing local-minimum amplitudes occurred within a time duration that was set to avoid detection of broad, far-field events, and to exceed a refractory period from the previous detection that was set to avoid multiple detections on a single deflection. CFAE map settings were a refractory period of 49 ms, P-P sensitivity >0.1 mV, and duration of 30 ms. After obtaining the CFAE-CL map, AF was terminated by internal cardioversion (2–10 J, biphasic shocks with R wave synchronization, anodal decapolar catheter in the high right atrium to cathodal duo-decapolar catheter inside the coronary sinus; Lifepak12, Physiocontrol Ltd), and then a voltage map was generated. If the initial rhythm was SR, we acquired the LA voltage map first. Voltage maps were generated by recording bipolar electrograms and measuring peak-to-peak voltage ($\text{Vol}_{\text{PACE}}$) during high right atrial pacing with a CL of 500 ms. We did not obtain a LA voltage map if frequent re-initiation of AF required electrical cardioversion more than 3 times. As a result, LA voltage maps were obtained from 31 patients. The CFAE-CL maps and $\text{Vol}_{\text{PACE}}$ maps (Figure 1A) were color-coded, and the CFAE-CL maps were converted

![Figure 1](image-url)
CFAE in Remodeled LA

into AF voltage (VolAF) maps (Figure 1B) by software. VolAF was calculated by the maximal peak-to-peak voltage of the last 1-s bipolar electrogram recorded during the acquisition of the 6-s CFAE map (Figures 1C, D).

Off-Line Analyses of Color-Coded 3D Maps of LA

We analyzed 3 color-coded maps: CFAE-CL, VolPACE, and VolAF. Both anterior–posterior (AP) and posterior–anterior (PA) views of each map were captured and converted into image files. Each of the 6 color map images per patient (AP and PA views of the 3 color maps) were divided into 4 quadrants, and the percentages of the colored areas were measured and calculated.12 The proportions of the color-coded areas in each quadrant (8 quadrants per patient) of the 3 different maps were analyzed by customized software (Image Pro), and were referenced to the color scale bars. We defined the CFAE area as white, and pink as an area with a CFAE-CL ≤120 ms.4 The low-voltage area was defined as an area with an LA voltage <0.2 mV, and was color-coded gray; the high-voltage areas (>5.0 mV) were coded as purple, as described before.11

Volumetric and Curvilinear Lengths Analyses of 3D Spiral CT Imaging

The 3D spiral CT images of the LA were analyzed on an imaging processing workstation (Aquarius, Terarecon, USA) as described in our previous study.12 The curvilinear lengths of the LA were measured at the linear ablation sites: bilateral antral ablation line, roof line, posterior inferior line, left lateral isthmus line, and anterior line. Each LA image was divided into portions by embryological origin as follows: the venous LA (posterior LA including the antrum and posterior wall), anterior LA (excluding the LAA and venous LA), and LAA.16 Although both the LAA and anterior LA are of embryological primordial atrial origin, they differ in geometry, myocardial fiber orientation, and the distribution of autonomic innervation. Therefore, we divided the LAA and anterior LA, and referenced them to points of inflection and separated the venous LA and anterior LA along the anterior border of both antral ablation lines, roof line, and posterior inferior line on the 3D spiral CT image (Figures 2A, D). The absolute and relative volumes of each portion were calculated and compared.

Data Analyses

We compared the mean and regional CFAE-CL with VolPACE, VolAF, absolute or relative volumes of the embryological portions of the LA, and the length of the linear ablation lines on 3D spiral CT images. We also correlated VolPACE and VolAF. To validate VolAF, we compared the automatically measured VolAF and manually calculated mean voltage of AF electrogram within 1 s of the same segment acquired from 100 different sites. For statistical analyses, we decided the cutoffs as the median rounded to zero decimal places for LA volume and decimal 0.1 for LA voltage, and validated them by receiver-operating characteristic curve analyses. Data are expressed as the mean±standard deviation. Statistical significance of the comparisons was assessed using the Kruskal-Wallis test, and Student’s t-test. A P-value <0.05 was considered significant.

Results

Low Voltage in Enlarged LA

Figure 2 shows representative 3D-CT images of the LA, LA voltage maps, and CFAE maps. LA voltage was lower...
In patients with an electroanatomically remodeled LA (ie, enlarged LA with low mean LA voltage) the CFAE-CL was low and % area of CFAE smaller (Figures 2A–C) than in those without a remodeled LA (Figures 2D–F). When we compared the patients with LA volume ≥125 ml and those with <125 ml, both the mean CFAE-CL and the regional CFAE-CL were lower in patients with an enlarged LA than in those with <125 ml. LA volume showed a weak positive correlation with CFAE-CL (R= 0.4509, P<0.001) and weak negative correlation with % area of CFAE (R=−0.3902, P<0.0001).

When we compared the patients with mean LA VolPACE <1.7 mV with those with >1.7 mV, the mean CFAE-CL and LA volume were negatively correlated (R=−0.5644, P<0.001; Figure 3A). To validate the automatically measured VolAF, we calculated the mean values of manually measured peak-to-peak voltages from 1-s AF electrograms in 100 different segments, and compared them with the automatically measured VolAF in the same segments by a single investigator who was blinded to all the data (Figures 1C,D). The automatically measured maximal peak-to-peak VolAF showed good correlation with the manually calculated mean voltage during AF (R=0.8224, P<0.0001; Figure 3B). We also compared the mean LA VolAF with the mean VolPACE of the same patient, and found a correlation between them (R=0.6155, P<0.0001; Figure 3C).

Prolonged CFAE-CL and Shrinking CFAE Area in Electroanatomically Remodeled LA

In patients with an electroanatomically remodeled LA (ie, enlarged LA with low mean LA voltage) the CFAE-CL was low and % area of CFAE smaller (Figures 2A–C) than in those without a remodeled LA (Figures 2D–F). When we

### Table 1. LA Voltage and CFAE Pattern by LA Volume

<table>
<thead>
<tr>
<th>LA volume</th>
<th>CFAE-CL (ms)</th>
<th>VolPACE</th>
<th>% Area of CFAE</th>
</tr>
</thead>
<tbody>
<tr>
<td>≥125 ml</td>
<td>Entire LA</td>
<td>209.6±83.4</td>
<td>28.4±26.8</td>
</tr>
<tr>
<td></td>
<td>Venous LA</td>
<td>188.6±92.7</td>
<td>20.6±27.2</td>
</tr>
<tr>
<td></td>
<td>LAA</td>
<td>173.1±88.5</td>
<td>19.2±30.7</td>
</tr>
<tr>
<td></td>
<td>Anterior LA</td>
<td>206.1±86.2</td>
<td>30.4±28.4</td>
</tr>
<tr>
<td></td>
<td>Left lateral isthmus area</td>
<td>243.2±108.5</td>
<td>19.2±30.7</td>
</tr>
</tbody>
</table>

| VolPACE (mV) | Entire LA | 1.4±0.7 | 2.2±0.8 | 0.06 |
| Venous LA    | 1.4±1.0 | 2.6±1.4 | 0.04 |
| LAA          | 2.5±1.7 | 4.1±1.7 | 0.08 |
| Anterior LA  | 1.3±0.6 | 1.7±0.6 | 0.04 |
| Left lateral isthmus area | 1.5±1.0 | 1.9±0.9 | 0.12 |

| VolAF (mV) | Entire LA | 0.6±0.3 | 0.8±0.3 | 0.03 |
| Venous LA   | 0.6±0.4 | 0.7±0.4 | 0.01 |
| LAA         | 1.3±1.1 | 1.4±0.9 | 0.24 |
| Anterior LA | 0.6±0.3 | 0.7±0.2 | 0.01 |
| Left lateral isthmus area | 0.6±0.4 | 0.9±0.7 | 0.06 |

Abbreviations see in Table 1.

### Table 2. LA Volume and CFAE Pattern by LA Voltage

<table>
<thead>
<tr>
<th>LA volume (ml)</th>
<th>VolPACE &lt;1.7 mV</th>
<th>VolPACE ≥1.7 mV</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Entire LA</td>
<td>Venous LA</td>
<td>155.7±58.0</td>
<td>92.6±34.7</td>
</tr>
<tr>
<td>LAA</td>
<td>47.5±21.4</td>
<td>33.7±12.4</td>
<td>0.016</td>
</tr>
<tr>
<td>Anterior LA</td>
<td>15.3±7.9</td>
<td>7.8±3.2</td>
<td>0.001</td>
</tr>
<tr>
<td>Right antral volume</td>
<td>9.8±4.3</td>
<td>8.7±2.9</td>
<td>0.018</td>
</tr>
<tr>
<td>Left antral volume</td>
<td>11.0±5.5</td>
<td>9.5±4.0</td>
<td>0.194</td>
</tr>
</tbody>
</table>

Proportional LA volumes (%)

| Venous LA | 30.6±8.2 | 36.9±5.0 | 0.006 |
| LAA       | 10.2±4.2 | 8.7±2.5  | 0.114 |
| Anterior LA | 59.3±9.1 | 54.4±5.9 | 0.043 |
| Right antral volume | 7.6±1.7 | 9.9±2.9 | 0.006 |
| Left antral volume | 7.6±3.8 | 10.5±3.6 | 0.020 |

Distribution of CFAE in Electroanatomically Remodeled LA

Figure 3D shows the distribution of CFAE according to the degree of electroanatomical remodeling. We compared patients with LA VolPACE ≤1.7 mV and LA volume ≥125 ml, those with LA VolPACE ≥1.7 mV and LA volume <125 ml, and those with an intermediate degree of remodeling. The incidence of septal CFAE was consistently high, regardless of the degree of LA remodeling, and the incidence of CFAE in other areas was reduced in the electroanatomically remodeled LA compared with a less remodeled LA.

### Discussion

In this study, we have documented different patterns of CFAE...
CFAE in Remodeled LA

Long-lasting AF induces atrial dilatation by changing LA function, increasing interstitial fibrosis, and atrioventricular nodal dysfunction. LA pressure contributes to AF initiation and maintenance, and LA ischemia reduces conduction velocity, which makes it susceptible to reentry maintenance. Atrial structural remodeling accompanies ultrastructural changes (ie, electrical remodeling).

CFAE in AF Ablation

Although the mechanism of CFAE has not yet been clearly elucidated, electrogram-guided RFCA targeting CFAE has been used as a substrate modification strategy in the catheter ablation of AF, especially for patients with PeAF. CFAE was first mentioned by Konings et al as an intraoperative mapping of human AF. Nademanee et al reported excellent clinical outcomes of CFAE-guided LA ablation. It has been demonstrated that 66–95% of cases of sustained AEs were terminated or organized during CFAE-guided AF ablation, which strongly indicates that the CFAE locus plays an important role in the maintenance of AF. In contrast, the clinical outcomes of CFAE ablation in other reports have varied widely, with SR maintenance rates ranging between 54% and 74%. CFAE-guided AF ablation has the limitation of differentiating CFAE with active driver and passive wavebreaker. Takahashi et al reported that only 17% of the CFAE areas were related to the termination of AF. Therefore, CFAE can be affected by temporal phenomenon caused by autonomic activity or electrical remodeling. In this study, the CFAE-CL was shorter and % area of CFAE was larger in patients with a normal-sized, high-voltage LA as compared with the electroanatomically remodeled atrium. The distribution of CFAE was also different. Most cases of PAF with a normal-sized LA without electrical remodeling have an excellent clinical outcome after circumferential PV isolation without additional CFAE-electrogram-guided AF ablation. Therefore, CFAE alone may not be a parameter sufficient to explain the pathophysiology of AF, at least in patients without electroanatomical remodeling of the LA.

CFAE in the Remodeled LA

Long-lasting AF induces atrial dilatation by changing LA function, increasing interstitial fibrosis, and atrioventricular nodal dysfunction. LA pressure contributes to AF initiation and maintenance, and LA ischemia reduces conduction velocity, which makes it susceptible to reentry maintenance. Atrial structural remodeling accompanies ultrastructural changes (ie, electrical remodeling).

Figure 3. (A) Correlation curve between left atrial (LA) voltage map (VolPACE) and LA volume. (B) Correlation curve between automatically calculated LA voltage map during atrial fibrillation (VolAF) and manually measured VolAF. (C) Correlation curve between VolAF and VolPACE in the same patient. (D) Different distributions of complex fractionated atrial electrogram (CFAE) according to the degree of electroanatomical remodeling of the LA.

in patients with an electroanatomically remodeled LA. CFAE-CL was longer and the % area of CFAE was lower in patients with an enlarged LA or low-voltage LA. Although the actual location of the CFAE differed among patients with a remodeled atrium, it was consistently positioned on the septum. These findings suggest that the patterns of CFAE are closely related to the degree of electroanatomical remodeling of the LA. We also documented a moderate correlation between VolPACE and VolAF.
remodeling), such as downregulation of the L-type calcium channel (\(\text{ICa-L}\)) and delayed rectification of the potassium channel (\(\text{IK}\)), leading to reduced conduction velocity.\textsuperscript{26} Neurohormonal activations, such as atrial natriuretic peptide, brain natriuretic peptide or the renin–angiotensin–aldosterone axis, result in inflammation and fibrosis.\textsuperscript{31} The AF-related reduction in endocardial voltage is associated with local conduction slowdown, which plays a major role in promoting AF.\textsuperscript{28,32} In the present study, the CFAE-CL was longer and % area of CFAE lower in patients with an electroanatomically remodeled LA than in those without it. In contrast, Sanders et al reported higher dominant frequency (DF) in the atria of patients with PeAF and electroanatomical remodeling compared with PAF patients.\textsuperscript{33} These results seem to contradict each other. However, the site of the CFAE does not match exactly the location of the DF,\textsuperscript{15} and the actual site exists at the boundaries of the maximal DF domain.\textsuperscript{34} Although both DF and CFAE reflect the frequency of the local electrogram, refractoriness, conduction velocity, and tissue anisotropy at the location of the electrogram, the former is a simple mathematical number of the fibrillation waveform (Fast Fourier transformation) and the latter is affected by the morphology and amplitude of the electrogram. Differences in recording systems or filter settings may also contribute to this difference. Therefore, we found that CFAE-CL was reduced in patients with AF and an electroanatomically remodeled LA with lowered conduction velocity, in contrast to DF.

Study Limitations

The patients included in this study were a highly selective group referred for RFCA. Although we detected CFAE with an adaptive peak-to-peak sensitivity threshold \(\geq 0.1\) mV, some CFAEs with low voltage might have been overlooked in patients with PeAF. We detected CFAE using a specific algorithm in the time domain, primarily based on the CL, so the findings of this study may not be directly comparable with other studies that used different parameters to identify CFAE. Because we acquired CFAE-CLs by point-by-point contact mapping, the CFAE-CL maps did not reflect a spatiotemporally homogeneous distribution of CFAE. We analyzed 3D maps using 2D measurements.

Conclusion

We documented different patterns of CFAE in patients with an electroanatomically remodeled LA. CFAE-CL was longer and % area of CFAE was lower in patients with an enlarged LA or low-voltage LA. The actual location of the CFAE and % area of CFAE was lower in patients with an electroanatomically remodeled LA. CFAE-CL was longer compared with PAF patients.\textsuperscript{33} These results seem to contradict each other. However, the site of the CFAE does not match exactly the location of the DF,\textsuperscript{15} and the actual site exists at the boundaries of the maximal DF domain.\textsuperscript{34} Although both DF and CFAE reflect the frequency of the local electrogram, refractoriness, conduction velocity, and tissue anisotropy at the location of the electrogram, the former is a simple mathematical number of the fibrillation waveform (Fast Fourier transformation) and the latter is affected by the morphology and amplitude of the electrogram. Differences in recording systems or filter settings may also contribute to this difference. Therefore, we found that CFAE-CL was reduced in patients with AF and an electroanatomically remodeled LA with lowered conduction velocity, in contrast to DF.

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Disclosure

No conflicts to disclose.

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