P Wave Morphology of an Arrhythmogenic Focus in Patients With Atrial Fibrillation Originating From a Pulmonary Vein or the Superior Vena Cava

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Background  It was hypothesized that atrial premature contractions (APCs) originating in the pulmonary veins (PVs) or superior vena cava (SVC) can be localized by evaluating characteristics of the P wave.

Methods and Results  Thirty-eight patients with paroxysmal atrial fibrillation were studied. P wave polarity and morphology of the ECGs during pacing from PVs were analyzed and compared to those of APCs originating from PVs. The P wave angle and notch in lead II during pacing from the right superior (RS) PV and SVC was compared to those of spontaneous APCs originating from those veins. A positive P wave in lead I was helpful in predicting right PV origin. A positive P wave in lead II distinguished superior PV origin. A notched P wave was helpful in predicting left PV origin. P wave polarity in lead II was positive during RSPV and SVC pacing. P waves in lead II during RSPV pacing had notching in 80%, but all P waves were smooth during SVC pacing. A P wave angle of >40° and notching in lead II showed RSPV origin.

Conclusions  These criteria are helpful in selecting which of the 4 PVs should be isolated when APCs cannot be recorded after transseptal puncture. (Circ J 2008; 72: 1650–1657)

Key Words: P wave morphology; Paroxysmal atrial fibrillation; Surface ECG

P aroxysmal atrial fibrillation (AF) can be triggered by atrial premature contractions (APCs) originating from a focal area. Most APCs originate from the orifice of a pulmonary vein (PV) or from the myocardial sleeve inside a PV.1–6 PV isolation by radiofrequency (RF) catheter ablation has been shown to effectively eliminate this type of AF.1–6 In addition, several studies have revealed the importance of non-PV APCs in triggering AF.7–9 Previous studies showed that pacemapping from each of the PVs and superior vena cava (SVC) reveals unique surface ECG characteristics and spontaneous APC from the PVs and SVC.10–14 It follows that the surface ECG characteristics of spontaneous APCs might help to localize the PV and/or non-PV origin of these triggers and guide ablation when APCs triggering AF do not occur spontaneously or cannot be induced during endocavitary exploration. However, no previous reports compared morphology of the paced P wave and spontaneous P wave from the PVs and SVC by simple algorithm.

The purpose of this 2-part study was to systematically apply simple ECG criteria, which was developed from pacemapping from the PVs and SVC based on previous studies, to spontaneously occurring APCs originating from the PVs and the SVC because the SVC and right superior (RS) PV are anatomically close, and thus, P wave morphology of the ectopic beats originating from the SVC or RSPV might be similar. We hypothesized that criteria developed from PV and SVC pacemapping would allow discrimination of right from left PV, superior from inferior PV origin and RSPV from SVC origin of APCs.

Methods

Study 1a

Patient Population  Fifteen patients (11 men and 4 women; mean age, 46±12 years) undergoing RF catheter ablation by transeptal approach for a left-sided accessory pathway (n=14) or left-sided atrial tachycardia (n=1) were included in this study. Informed consent was obtained from all patients before the procedure (study 1a, 1b and 2) according to the protocol approved by the Nihon University Itabashi Hospital’s Human Research Committee.

Study Protocol  After successful ablation of the accessory pathway or atrial tachycardia, a 7F, 4-mm-tip quadripolar steerable ablation catheter (EP Technologies, Sunnyvale, CA, USA) was introduced through a long sheath into the proximal portion (within 2 cm from the PV orifice) of each (RS, right inferior (RI), left superior (LS) and left inferior (LI)) PV, close to the ostium. The position of the ablation catheter within each PV was verified in all patients by its fluoroscopic appearance outside the cardiac silhouette and by PV angiography. Patients with common left PV ostium and with right middle PV were excluded from the study. Of the 60 PVs in the 15 patients, 60 PVs (100%) were success-
Fig 1. (a) P wave amplitude in lead I during pacing from 4 PVs. Note that a P wave amplitude of ≥50 μV differentiates the P wave of right PV origin from that of left PV origin. (b) P wave amplitude in lead II during pacing from 4 PVs. Note that the P wave amplitude of ≥100 μV differentiates a P wave of superior PV origin from that of inferior PV origin. (c) P wave morphology in lead V1 during pacing from 4 PVs. Note that the notched P wave morphology with the second component larger than the first component differentiates APCs of left PV origin from those of right left pulmonary origin. PV, pulmonary vein; APC, atrial premature contraction; LSPV, left superior pulmonary vein; LIPV, left inferior pulmonary vein; RSPV, right superior pulmonary vein; RIPV, right inferior pulmonary vein; OR, odds ratio; CI, confidence interval.
fully cannulated. Bipolar pacing was performed at a cycle length of 600–800 ms with a minimum current strength of 4.4±1.0 V and 2 ms duration.

**P Wave Analysis** P waves were analyzed by 2 blinded observers on 12-lead ECGs recorded at a paper speed of 50 mm/s and amplitude of 0.5 mV/cm. The P wave configuration was classified according to amplitude as ≥100 μV or <100 μV in lead I (Fig 1a), as ≥50 μV or <50 μV in lead II (Fig 1b) based on the crude method of amplitude measurement through square-counting based on a previous study and notch recording with an amplitude of the second component being larger than the first component in lead V1 (Fig 1c). Only P waves that were clearly visible and isolated from the preceding T waves were analyzed. One V1 lead ECG during RSPV pacing could not be analyzed because of the presence of a large noise.

**Study 1b**

**Patient Population** Twenty-five patients with paroxysmal AF (23 men and 2 women; mean age, 59±11 years) undergoing segmental isolation of the 4 PVs by a transseptal approach were included in this study. Two long sheaths (SL1: Daig Corp, Minnetonka, MN, USA) were inserted into the left atrium via transseptal puncture under fluoroscopic guidance. A 7F Trio guide sheath (Cardiac Pathways Corp, Sunnyvale, CA, USA) with 3 ports was introduced into the SL1 sheath, and 3, 2F quadripolar catheters with 5-mm interelectrode spacing (Ensemble, Cardiac Pathways Corp) were positioned; 1 at the LSPV, 1 at the LIPV and 1 at the RSPV. A 7F deflectable quadripolar catheter with 2.5-2 mm spacing (EP Technologies) was positioned in the RIPV through another SL1 sheath, and 3, 2F quadripolar catheters with 5-mm interelectrode spacing (EP Technologies) was positioned in the RIPV through another SL1 sheath. A 20-pole catheter (A 20; Cordis-Webster, Baldwin Park, CA, USA) was placed in the posterior right atrium, and a decapolar catheter with an end-hole was placed in the coronary sinus (Daig Corp) before PV isolation. Patients with common left PV ostium and with a right middle PV were excluded from the study.

**Study Protocol** Intracardiac bipolar electrograms were displayed simultaneously with ECG leads I, II, III, V1 and V6 on a multichannel recorder (CardioLab System, Prucka Engineering Inc, Houston, TX, USA). The filter was set at 0.05–100 Hz for surface ECG and 30–500 Hz for intracardiac recordings. We first attempted to locate the spontaneous onset of APCs triggering AF in the baseline condition or during isoproterenol infusion (up to 4 μg/min). We could localize the origin of APC by identifying the earliest activation within the PV before activation of the other 3 PVs, coronary sinus activation and right atrial activation. If spontaneous AF did not appear, intermittent atrial pacing (8–12 beats) with a cycle length of 200–300 ms from the high right atrium or coronary sinus was used to induce sustained AF. After the pacing-induced AF was sustained for 5–10 min, external cardioversion was attempted to convert AF to sinus rhythm so that we could observe the spontaneous re-initiation of AF. Then we paced the PV(s) (from which APC originated) in the same manner described in study 1a.

**P-Wave Analysis** Morphology of the APCs was analyzed by 2 blinded observers on 12-lead ECGs, recorded at a paper speed of 50 mm/s and at an amplitude of 0.5 mV/cm, and the morphologies of the paced P wave and APCs originating from the PV were compared. Because of the difficulty in measuring the amplitude of the P waves superimposed on the ST-T segment, we qualitatively measured P wave morphology as positive, biphasic and negative.

**Study 2**

**Patient Population** Thirteen patients with paroxysmal AF (11 men and 2 women; mean age, 52±15 years) who were undergoing segmental isolation of 4 PVs by a transseptal approach were included in this study. After transseptal puncture, 2 SL1 sheaths (Daig Co) were inserted into the left atrium via transseptal puncture under fluoroscopic guidance.

**Study Protocol** A 7F, 4-mm-tip quadripolar steerable ablation catheter (EP Technologies) was introduced through the long sheath into the proximal portion (within 2 cm from the orifice) of the RSPV. Another 6F, 4-mm-tip quadripolar steerable catheter was placed in the SVC (within 2 cm from the SVC and right atrial junction). Bipolar pacing from the RSPV and SVC was performed at a cycle length of 600–800 ms with a minimum current strength of 3.0±1.0 V and 2.3±0.9 V, respectively, and 2 ms duration before PV isolation. We confirmed that pacing from SVC did not directly capture RSPV and vice versa by using an activation sequence; that is, the proximal to distal activation sequence of the SVC and RSPV during RSPV and SVC pacing.

**P Wave Analysis** Morphology of the paced P wave was analyzed by 2 blinded observers on 12-lead ECGs, recorded at a paper speed of 50 mm/s and at an amplitude of
P Wave Morphology of an Atrial Focus

0.5 mV/cm, by measuring the angle of the P waves manually using a protractor. The morphologies of the APCs originating from the RSPV and SVC were compared qualitatively.

**Statistics**

The values are expressed as the mean ± SD. The differences in continuous variables were analyzed by the unpaired Student’s t-test, and the differences in the categorical variables were analyzed by the chi-square test. StatView 5.0 software (SAS Institute) was used for data analysis. Confidence intervals of the odds ratio and the receiver-operating characteristic (ROC) curve of the P wave angle was calculated.
lated using StatFlex software. A p value of less than 0.05 was considered statistically significant.

**Results**

**Study 1a**

**Amplitude of Paced P Waves** Significant differences in the amplitude of paced P waves were observed in leads I and II. In lead I, amplitudes of P waves produced by right PV pacing were significantly higher (≥50 µV) than amplitudes of P waves produced by left PV pacing (p<0.001, Fig 1a). In lead II, pacing from the superior PVs produced P waves with significantly higher amplitudes (≥100 µV) than those of P waves produced by pacing from the inferior PVs (p<0.001, Fig 1b).

**Morphology of Paced P Waves** A significant difference in the morphology of paced P waves was observed in lead V1. Left PV pacing produced positive bimodal P waves with the second component larger than the first component, whereas right PV pacing produced positive unimodal or positive bimodal P waves with the first component larger than the second component (p<0.005, Fig 1c).

**Development of an Algorithm** To predict the PV pacing site origin from the analysis of the morphologic features of P waves during selective PV pacing, we developed an algorithm to predict the origin of the pacing site in the PV. The combination of a P wave amplitude of <50 µV in lead I and a positive bimodal P wave with a larger second component in lead V1 is taken to indicate a left-side PV origin; otherwise, the P wave is regarded to have come from a right-sided PV origin. A positive P wave amplitude in lead II of ≥100 µV was selected to differentiate the superior PVs.

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**Table 1  Evaluation of Surface ECG Criteria**

<table>
<thead>
<tr>
<th>Criteria</th>
<th>RSPV (n=15)</th>
<th>RIPV (n=2)</th>
<th>LSPV (n=8)</th>
<th>LIPV (n=5)</th>
<th>SVC (n=1)</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Lead I</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Positive</td>
<td>13</td>
<td>2</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Biphasic</td>
<td>1</td>
<td>0</td>
<td>4</td>
<td>2</td>
<td>0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Negative</td>
<td>1</td>
<td>0</td>
<td>3</td>
<td>3</td>
<td>0</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Lead II</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Positive</td>
<td>13</td>
<td>0</td>
<td>6</td>
<td>1</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Biphasic</td>
<td>1</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>0</td>
<td></td>
<td></td>
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<tr>
<td>Negative</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>0</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em><em>RPVs vs LPVs</em> (positive P wave in lead I)</em>*</td>
<td>89</td>
<td>92</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td><strong>SPVs vs IPVs</strong> ** (positive P wave in lead II)**</td>
<td>83</td>
<td>86</td>
<td></td>
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</table>

RSPV, right superior pulmonary vein; RIPV, right inferior pulmonary vein; LSPV, left superior pulmonary vein; LIPV, left inferior pulmonary vein; SVC, superior vena cava.

* p<0.001, ** p<0.001.
Thirty-one APC foci were identified in the PVs and SVC: 15 in the RSPV (Fig 3a, Left), 2 in the RIPV (Fig 3a, Right), 8 in the LSPV (Fig 3b, Left), 5 in the LIPV (Fig 3b, Right) and 1 in the SVC (Fig 4). These APCs were located on the preceding T wave. The best sensitivity and specificity were achieved by combining the following criteria: (1) A positive P wave in lead I. This was present in 89% of APCs from right-sided PVs and the SVC, yielding a sensitivity of 89% and a specificity of 92% for distinguishing right (including SVC) from left PV origin of spontaneous APCs; (2) A positive P wave in lead II. This was present in 83% of APCs from superior PVs and the SVC, yielding a sensitivity of 83% and a specificity of 86% for distinguishing inferior from superior PV (including SVC) origin of spontaneous APCs (Table 1).

Study 2

Because APCs originating from the RSPV and SVC showed similar polarity, that is, positive P waves in leads I and II, we classified morphology of the P wave in lead II by notching and by the angle of the P wave formed by 2 lines tangential to the up-slope and down-slope of the P wave (Fig 4). Notching of the P wave was present in 77% during RSPV pacing and 7.6% during SVC pacing (p<0.0005), yielding a sensitivity of 78% and a specificity of 92% for distinguishing right (including SVC) from left PV origin of spontaneous APCs; (2) A positive P wave in lead II. This was present in 83% of APCs from superior PVs and the SVC, yielding a sensitivity of 83% and a specificity of 86% for distinguishing inferior from superior PV (including SVC) origin of spontaneous APCs (Table 1).

Table 2  P Wave Morphology and P Wave Angle in Lead II During RSPV and SVC Pacing

<table>
<thead>
<tr>
<th>Case</th>
<th>Age (years old)</th>
<th>Pacing notch</th>
<th>Angle (degree)</th>
<th>APC</th>
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<tr>
<td></td>
<td>gender</td>
<td>SVC</td>
<td>RSPV</td>
<td>SVC</td>
</tr>
<tr>
<td>1</td>
<td>61/M</td>
<td>(-)</td>
<td>(+)</td>
<td>21</td>
</tr>
<tr>
<td>2</td>
<td>35/M</td>
<td>(-)</td>
<td>(+)</td>
<td>27</td>
</tr>
<tr>
<td>3</td>
<td>50/F</td>
<td>(+)</td>
<td>(+)</td>
<td>17</td>
</tr>
<tr>
<td>4</td>
<td>31/F</td>
<td>(-)</td>
<td>(-)</td>
<td>26</td>
</tr>
<tr>
<td>5</td>
<td>55/M</td>
<td>(-)</td>
<td>(+)</td>
<td>21</td>
</tr>
<tr>
<td>6</td>
<td>66/F</td>
<td>(-)</td>
<td>(+)</td>
<td>25</td>
</tr>
<tr>
<td>7</td>
<td>49/F</td>
<td>(-)</td>
<td>(-)</td>
<td>21</td>
</tr>
<tr>
<td>8</td>
<td>69/M</td>
<td>(-)</td>
<td>(+)</td>
<td>27</td>
</tr>
<tr>
<td>9</td>
<td>74/M</td>
<td>(-)</td>
<td>(+)</td>
<td>39</td>
</tr>
<tr>
<td>10</td>
<td>46/M</td>
<td>(-)</td>
<td>(-)</td>
<td>32</td>
</tr>
<tr>
<td>11</td>
<td>46/M</td>
<td>(-)</td>
<td>(+)</td>
<td>38</td>
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<tr>
<td>12</td>
<td>27/M</td>
<td>(-)</td>
<td>(+)</td>
<td>38</td>
</tr>
<tr>
<td>13</td>
<td>67/M</td>
<td>(-)</td>
<td>(+)</td>
<td>47</td>
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Table 3  Discrimination of RSPV vs SVC Pacing Site by P Wave in Lead II

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
<th>OR</th>
<th>95% CI</th>
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<tr>
<td>Notching (+)*</td>
<td>78</td>
<td>92</td>
<td>40.0</td>
<td>3.6 CI ±447.1</td>
</tr>
<tr>
<td>Angle &gt;40°**</td>
<td>78</td>
<td>92</td>
<td>40.0</td>
<td>3.6 CI ±447.1</td>
</tr>
<tr>
<td>Notching (+) and angle &gt;40°***</td>
<td>67</td>
<td>100</td>
<td>20.8</td>
<td>2.0 CI ±211.8</td>
</tr>
</tbody>
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* p<0.0004, ** p<0.0004, *** p<0.0006.

from inferior PVs (Fig 2).

Study 1b

Because APCs originating from the RSPV and SVC showed similar polarity, that is, positive P waves in leads I and II, we classified morphology of the P wave in lead II by notching and by the angle of the P wave formed by 2 lines tangential to the up-slope and down-slope of the P wave (Fig 4). Notching of the P wave was present in 77% during RSPV pacing and 7.6% during SVC pacing (p<0.0005), yielding a sensitivity of 78% and a specificity of 92% for distinguishing right (including SVC) from left PV origin of spontaneous APCs; (2) A positive P wave in lead II. This was present in 83% of APCs from superior PVs and the SVC, yielding a sensitivity of 83% and a specificity of 86% for distinguishing inferior from superior PV (including SVC) origin of spontaneous APCs (Table 1).

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<td>100</td>
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</table>

OR, odds ratio; CI, confidence interval. Other abbreviations see in Table 1.

Fig 5. ECG of spontaneous APC originating from SVC. P wave polarity was positive in leads I and II, and morphology of the P wave in lead II showed a narrow P wave angle and the absence of a notch. Abbreviations see in Figs 1,3,4.
P wave morphology of these APCs showed similar findings for separating SVC origin from RSPV origin (Fig 5).

Discussion

Major Findings

Study 1  Selective pacing at each of the 4 PVs produced distinct P wave characteristics that could be used to identify the pacing site origin in the PV. Left PV pacing produced a low P wave amplitude (flat or negative P wave) in lead I and a notched morphology in lead V1. Conversely, the right PV origin was suggested by a positive P wave of ≥0.50 mV in lead II and unimodal morphology in lead V1. Superior PV origin was suggested by a positive P wave of ≥100 mV.

Study 2  Differentiation of SVC origin from RSPV origin was possible by a sharp P wave angle (≥40°) and the absence of P wave notching. Thus, the combined use of the P wave amplitude of lead I, the P wave amplitude, notching, and angle in lead II, and the P wave notching in lead V1 is useful for predicting the origin of the APC in the PVs and SVC.

We adopted leads I and II for predicting spontaneous APCs and the absence of P wave notching. The study showed that the right PV pacing produced a positive P wave of ≥0.50 mV in lead II and unimodal morphology in lead V1. Superior PV origin was suggested by a positive P wave of ≥100 mV.

Previous Studies

Previous studies showed pacemapping from each of the PVs (P wave amplitude in lead I, P wave duration, amplitude ratio of III/II and sum of the P wave amplitude in all the inferior leads) and SVC (P wave polarity in leads I and aVl). To the best of our knowledge, our study is the first to attempt to correlate the P wave morphology during pacing in the PVs and SVC with spontaneous atrial ectopic beats of PV and SVC origin.

Tse et al reported P wave polarity during pacing from the PVs and right atrium. They found that the combination of a negative or biphasic P wave in lead I and positive P wave in lead V1 was highly specific for the origin of the pacing site from the PVs. A positive P wave in the inferior leads was associated with pacing sites in superior PVs. In their study, P wave polarity had only limited value in distinguishing pacing sites in left and right PVs. Yaman et al reported P wave characteristics during PV pacing. Their study showed that the right PV pacing produced a positive P wave in lead I, flat P wave in lead aVl, and a low lead III/II amplitude ratio. There was a significant difference in the P wave duration between the right and left PVs; however, the ability to distinguish between right and left PV origin was modest. The P wave amplitude in lead II distinguished superior PVs from inferior PVs with a 74% specificity and a 81% sensitivity. A notched P wave in lead II showed a 92% specificity in predicting left PV origin.

Kuo et al reported that the combination of biphasic or isoelectric P wave polarity in lead V1 or biphasic P wave polarity in lead aV1 had moderate sensitivity (71%) and specificity (82%) in predicting an arrhythmogenic focus of AF from the SVC. They studied spontaneous atrial ectopic beats only from the SVC and RSPV. Their study yielded the following criteria for distinguishing right-sided from left-sided PV origin: (1) a P wave duration <120 ms; (2) a P wave amplitude in lead I of >0.05 mV; and (3) a P wave amplitude in leads II/III of >1.25 mV. The criterion for distinguishing superior from inferior PVs was the sum of the P wave amplitudes in all inferior leads >0.3 mV.

Their study results document that surface ECG P wave morphology during pacing might be different, particularly in patients with infrequent ectopy during an ablation procedure. This is particularly useful in the youngest patients, who typically have a single arrhythmogenic PV. The establishment of surface ECG criteria for identifying the origin of spontaneous atrial ectopy in a PV might facilitate the rapid localization of arrhythmogenic PVs and SVC with a single triggering ectopic beat. The described surface ECG criteria can also be used to localize the arrhythmogenic focus to a PV or the SVC during limited lead monitoring prior to the ablation procedure. These data are of critical importance when spontaneous atrial ectopic beats triggering AF originate only from the SVC. In such cases, transseptal puncture can be avoided. Furthermore, in the absence of frequent spontaneous events that afford the opportunity for more detailed mapping, the presence of confirmed spontaneous events originating from the left and right PVs prior to the procedure allows the physician to anticipate the need to isolate all the PVs and also allow more time to evaluate potential reconnection by isolation of the arrhythmogenic PVs first.

Clinical Implications

Identification of a specific arrhythmogenic PV can be difficult, particularly in patients with infrequent ectopy during an ablation procedure. This is particularly useful in the youngest patients, who typically have a single arrhythmogenic PV. The establishment of surface ECG criteria for identifying the origin of spontaneous atrial ectopy in a PV may facilitate the rapid localization of arrhythmogenic PVs and SVC with a single triggering ectopic beat. The described surface ECG criteria can also be used to localize the arrhythmogenic focus to a PV or the SVC during limited lead monitoring prior to the ablation procedure. These data are of critical importance when spontaneous atrial ectopic beats triggering AF originate only from the SVC. In such cases, transseptal puncture can be avoided. Furthermore, in the absence of frequent spontaneous events that afford the opportunity for more detailed mapping, the presence of confirmed spontaneous events originating from the left and right PVs prior to the procedure allows the physician to anticipate the need to isolate all the PVs and also allow more time to evaluate potential reconnection by isolation of the arrhythmogenic PVs first.
detecting a superior origin is much less reliable than the sensitivity. We did not compare the P wave morphology during pacing from different bipoles in the same PV. However, Dixit et al showed that unique intracardiac activation patterns during ostial pacing from individual PVs are not influenced by the circumferential location of the PV pacing site.

Although all 4 PV isolation had become the standard strategy of AF-ablation, recently published papers demonstrated that in patients with clearly documented arrhythmogenic PVs, the segmental PVI of the PV triggering the AF or an ipsilateral PV had a comparable long-term success rate and shorter RF energy delivery and procedure times than the empirical 4 PV isolation method.

Finally, we did not perform activation sequence mapping of the left and right atria during 4 PVs and SVC pacing to elucidate why P wave morphology is different during pacing from 4 PVs and SVC.

Further studies are needed to evaluate the role of specific anatomic structures such as septalpulmonary bundle and Bachmann’s bundle for intra and inter atrial conduction sequence by using a 3-D mapping system.

Conclusions

Analysis of the surface ECG can often provide enough information to help localize the PV or the SVC origin of spontaneous APCs despite their superimposition upon the T wave. Surface ECG criteria based on P wave amplitude in lead I and notching in V1 can help distinguish right-sided from left-sided origin of spontaneous APCs, P wave amplitude in lead II can help distinguish superior from inferior PV origin of spontaneous APCs, and P wave angle and notching in lead II can help distinguish RS from SVC origin of spontaneous APCs.

References