Electrophysiological End Point for Catheter Ablation of Atrial Fibrillation Initiated From Multiple Pulmonary Venous Foci

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Background—The end point for catheter ablation of pulmonary vein (PV) foci initiating atrial fibrillation (AF) has not been determined.

Methods and Results—Ninety patients underwent mapping during spontaneous or induced ectopy and/or AF initiation. Ostial PV ablation was performed by use of angiograms to precisely define targeted sites. Success defined by elimination of AF without drugs was correlated with the procedural end point of the abolition of distal PV potentials. A total of 197 arrhythmogenic PV foci (97%)—single in 31% and multiple in 69%—and 6 atrial foci were identified. A discrete radiofrequency (RF) application eliminated the PV potentials in 9 PV foci, whereas 2 foci from the same PV required RF applications at separate sites in 19 cases. In others, a wider region was targeted with progressive elimination of ectopy. In 49 patients, multiple sessions were necessary owing to recurrent or new ectopy. The clinical success rates were 93%, 73%, and 55% in patients with 1, 2, and ≥3 arrhythmogenic PV foci. Recovery of local PV potential and the inability to abolish it were significantly associated with AF recurrences (90% success rate with versus 55% without PV potential abolition). PV stenosis was noted acutely in 5 of 6 cases, remained unchanged at restudy, and was associated with RF power >45 W.

Conclusions—Multiple PV foci are involved in initiation of AF, and elimination of PV muscle conduction is associated with clinical success. (Circulation. 2000;101:1409-1417.)

Key Words: atrial fibrillation • catheter ablation • electrophysiology

The pulmonary veins (PVs) have been shown to trigger paroxysms of atrial fibrillation (AF), and radiofrequency (RF) ablation of these foci can eliminate AF.1-5 However, the efficacy of ablation is limited by a high percentage of recurrences and the unknown frequency of venous stenosis, and the end point of successful ablation, particularly in patients with scant ectopy, has not been defined. The present study was prospectively performed to assess these issues.

Methods

Patient Characteristics
The study population consisted of 90 consecutive patients (Table 1). Thirty-seven had >700 ectopies per day (Figure 1).

Electrophysiological Study
The study was performed as described previously with the use of 2 or 3 electrode catheters and after confirmation of the absence of intracardiac thrombi by transesophageal echocardiography.5 The left atrium (LA) and PVs were explored through either a patent foramen ovale (19 patients) or a transseptal catheterization. Selective PV angiography was performed by hand injection of 5 to 10 mL of contrast with an NIH catheter and displayed during the procedure.

Heparin was titrated to maintain a partial thromboplastin time of 60 to 90 seconds (control, 30 seconds).

In patients with few ectopies or sustained AF necessitating cardioversion, 2 steerable ablation catheters (with different curves) were used to map the PVs simultaneously. When few ectopies were observed, the following provocative maneuvers were performed: vagal maneuvers, including carotid sinus massage, deep breathing, and Valsalva; slow-rate atrial pacing (bursts of 3 to 10 stimuli at 100 to 200 bpm for postpause ectopy); isoproterenol infusion (2 to 4 μg/min); high-rate pacing; and their combinations. If long-lasting AF was induced, internal (n=6) or external electrical cardioversions were performed, sometimes followed by AF reinitiation. ATP injection (20 to 80 mg) or saline infusion (500 mL in 15 minutes) was tried in 4 patients each. A swallowing maneuver was a reproducible mode of induction in 1 patient.

Mapping was performed in ≥2 sites in the superior PV(s) either sequentially or simultaneously and thereafter in the inferior veins (Figures 2 and 3). If the earliest activity could not be traced from any PV, both atria were mapped.

Definition of an Arrhythmogenic PV
An arrhythmogenic PV4,5 was defined on the basis of documented ectopy—single or multiple and with or without conduction to the LA. During sinus rhythm, double or multiple potentials were...
recorded in a progressively later temporal sequence, synchronous with the first (right PVs) or second (left PVs) half of the P wave (Figure 3). The first low-frequency potential reflected activation of the adjacent atrium. The latest high-frequency potentials indicated PV muscle potentials (PVPs). The ostial PVPs were sometimes not obvious in the left PVs because of the superimposed LA potential during sinus rhythm; however, pacing of the distal coronary sinus (or LA appendage) allowed their separation and thus easy recognition of PVP (Figure 4).

During ectopy from an arrhythmogenic PV, there was a reversal in activation sequence, from the distal PV trunk (source) to the ostium and LA (exit), with the PVP preceding the LA potential. Ectopic discharges with the shortest coupling interval were not conducted to the LA producing isolated PV confined within the PV (Figures 2 and 3). They were recognized as a PVP coincident with or just after the ventricular electrogram and could be distinguished from a potential of ventricular origin by their spontaneous disappearance (intermittent PVP) or suppression during atrial pacing.

Conversely, if the explored PV was not the origin of ectopy, it was "passively" activated as in sinus rhythm with a proximal-to-distal sequence and a PVP after or fusing with the LA potential. Multiple foci in the same PV were defined by the presence of ≥2 different sources and exits at opposite locations of the PV perimeter (eg, roof and bottom) demonstrated by selective PV angiography.

**Ablation Procedure**

RF ablation of atrial foci was performed at the site of earliest activation. RF ablation of arrhythmogenic PV was performed within 15 mm of the PV ostium; the exact location depended on catheter stability. This was thought to be safer (vis-à-vis the risk of PV stenosis) than more distal applications in smaller veins. The perimeter of the PV ostium was mapped to localize the ostial sector showing the earliest activity during ectopy, and the highest and sharpest PV potential was first targeted for ablation. Subsequent RF applications were performed if needed at contiguous sites showing PVP. The end point was elimination of ectopy, spontaneous or induced by provocative maneuvers, and elimination of PV muscle conduction (during sinus rhythm) distally to the ablation site(s) on the basis of either abolition or dissociation of distal PVP. To assess capture of the LA, local pacing was performed from these sites.

RF energy was delivered at the distal electrode of the thermocouple-equipped catheter (target, 50°C) with a power limit of...
45 to 50 W in the first 35 patients. Because of the occurrence of several acute PV stenoses (see later), a maximal power limit of 25 to 30 W was subsequently set for the last 55 patients. If 1 PV was arrhythmogenic, the PV producing the most repetitive ectopy and/or initiating AF was first targeted; then, RF energy was delivered in a second PV after angiographic verification of the unchanged patency of the first one. PV angiography was performed at the end of ablation in all but 4 patients (technical problems). PV stenosis was defined as a diameter reduction of >50%. Repeated PV angiography was performed 1 month later in 58 patients.

Patients were discharged beyond the third day under oral anticoagulant. In the event of recurrent AF, all patients were advised to undergo a new study, and ablation was performed if there was no PV stenosis. The procedure was considered a success when the AF episodes were totally eliminated without antiarrhythmic drugs both clinically and on Holter recordings. Any suggestive but undocumented symptoms were attributed to AF recurrence. Anticoagulants were interrupted 3 months after successful elimination of AF unless there were other risk factors.

Statistical Analysis
Continuous variables are expressed as group mean±SD and compared with the use of a Kruskal-Wallis or Student’s t test. Noncontinuous variables were compared by use of the χ² or Fisher’s exact test. Statistical significance was selected at P<0.05.

Results
In 4 patients, initiation and maintenance of AF were due to a rapidly firing focus with a persistent distal-to-proximal PV activation sequence during AF (focal AF). The remaining patients had focally initiated AF with a short burst of focal discharges initiating typical intra-atrial fibrillation, which continued independently.

Mapping of Triggering Ectopic Beats
In 58 patients, provocative maneuvers were needed to elicit ectopy (Table 2). Sustained AF required electrical external or internal cardioversions in 38 patients.

Ectopy originating from a single PV was found in 27 patients, 2 in 22, 3 in 29, and 4 in 11, for a total of 197 arrhythmogenic PVs in 89 patients, and from the atrial muscle in 6 patients (right atrium in 1, atrial septum in 1, and posterior LA in 4). A single arrhythmogenic PV was associated with female sex, younger age, less frequent and shorter AF paroxysms, and smaller atrial dimensions (Table 3).

RF Ablation of Foci Initiating AF

Atrial Foci
One patient (patient 11) had a single atrial focus and underwent 3 ablation sessions because of 2 recurrences after ablation. Five patients had atrial foci in addition to arrhythmogenic PV; ablation was successful in 3 and failed in 2 because of insufficient ectopy.

Frequently Discharging PV Foci
Mapping of ectopy from 9 PVs (4 left inferior PVs [LIPVs], 3 right inferior PVs [RIPVs], 1 right superior PV [RSPV], and 1 left superior PV [LSPV]) showed a discrete source from the ostium of a venous branch and a discrete exit at the ostium of the venous trunk reflected by a localized (which disappeared with small catheter movements on either side) continuum of the PVP-LA potential. Ectopy was abruptly eliminated by a

![Figure 3](image1.png)

**Figure 3.** In sinus rhythm (first beat), fractionated electrograms are recorded from 2 veins with initial far-field atrial component followed by late sharper PV component, electrograms being early (with respect to P wave) in RSPV and late in LIPV. Activation during ectopic (second beat) is relatively late and passive in both veins; ie, local PV component does not precede atrial component, indicating that this ectopic discharge originates from third site, in this case, LSPV. After second sinus beat, sharp-spike discharges that are not followed by atrial deflection (concealed ectopy) are noted in both PVs.

![Figure 4](image2.png)

**Figure 4.** Local left PV potentials are not always obvious during sinus rhythm and may require pacing from within distal coronary sinus (CS) to be revealed. Cine frame shows catheters placed in LSPV, LIPV, and coronary sinus. Bipolar tracings show 1 sinus beat followed by paced beat from distal coronary sinus. In each case, although electrograms during sinus rhythm do not show evident local PV potentials, characteristic sharp potential (•) distinct from initial far-field potential is recorded during pacing. S indicates stimulation artefact.
discrete RF ablation, which abolished the ostial and corresponding distal (site activated later into the venous branch) PVP during sinus rhythm (Figure 5).

In 19 PVs (9 LIPVs, 6 LSPVs, and 4 RSPVs), there were 2 sources of ectopy from different PV branches associated with 2 distinct ostial exits necessitating ablation at separate points of the ostium (Figure 6). This produced abrupt disappearance of targeted ectopies and sequentially eliminated the ostial and distal PVPs in each corresponding venous branch.

In the remaining PVs, mapping of the ostium during ectopy showed synchronous local activation in a large sector of the venous perimeter (Figure 7). Correspondingly, during sinus rhythm, PVPs were recorded in a large part of the ostium ranging from one quarter to the full circumference. In 14, differing activation times during ectopy suggested various ectopy sources or activation courses. Sequential RF applications at the ostium produced a progressive decrease in ectopies or an initial disappearance of repetitive followed by that of isolated ectopy; the first applications commonly produced exacerbation of ectopies. Ablation eliminated progressively the local PVPs, usually requiring 2 or 3 series of RF applications along the entire sector (Figure 8). A progressive prolongation of the LA-PVP conduction time (both locally and distally) was observed during sinus rhythm, and ablation at the remaining site(s) without atrial-PVP delay (gap like continuum electrogram) resulted in abrupt disappearance of local and distal PVPs, indicating disconnection. An abrupt reappearance of local and distal PVPs was noted in 19 patients, and reablation as described above was successfully performed.

In 9 patients, after ablation, residual isolated ectopies were found to originate at the atrial edge of the PV ostium (instead of inside the PV), indicating a modification of the source of earliest activity as a result of RF delivery and necessitating additional RF applications.

**Patients With Insufficient PV Ectopy**

In 12 patients, only 1 to 5 ectopies were noted despite multiple provocative maneuvers; however, this was sufficient to identify the arrhythmogenic PV. RF applications were performed during sinus rhythm or distal coronary sinus pacing at the sites exhibiting PVPs, resulting in total elimination of the distal PVPs (Figure 9).

In 8 other patients, PV ectopy immediately produced sustained AF, which recurred after repeated cardioversions (5±2). In 5 patients, a circumferential PV ostial ablation was performed during AF, and progressive abolition of the PVP (sharpest potentials) could be seen distally inside the PV; after a new cardioversion, AF did not reappear, and additional RF applications abolished the residual PVP during sinus rhythm. In the last 3 patients, amiodarone (300 to 600 mg) was infused and ablation was performed after conversion in sinus rhythm.

**Reablation and Final Mapping Results**

A total of 155 ablation sessions were performed: 1 in 41 patients, 2 in 36, 3 in 10, and 4 in 3, for total procedure duration and fluoroscopy times of 278±154 and 82±44 minutes, respectively (149±71 and 45±21 minutes per session).

At the end of the first session, ectopy was persistent in 9 patients, and all had early recurrences of AF. Of the remaining 81, 40 (50%) had recurrent AF. These 49 patients underwent 1 to 3 (total, 65) reablation sessions. During these
sessions, mapping showed that the ectopy was related to the persistence or recovery of a previously targeted PVP in 24, originated from another part of a previously targeted PV in 12, originated from a different PV in 21, and was related to combinations of the above in 8. The recovered PVP was delayed and thus obvious during sinus rhythm; it was present (ostially and distally) at a discrete area in 9 patients and abolished with a single RF application (Figure 10). In 4, it disappeared intermittently by simple mechanical catheter pressure. Sequential RF applications were needed in others to eliminate PVPs.

After all ablation, mapping in 96 PVs showed preservation of the distal PVP. Local PV pacing from the area with PVP, even when it was very discrete, could capture the LA. All PVPs were eliminated in 86 PVs or dissociated in 15 (superior) PVs; pacing could not capture the LA.

Safety
Air embolism during catheter exchange was responsible for transient ECG changes in 5 patients. A transient bradycardia occurred in 3 patients during LSPV ablation. Patient 11 developed a hemopericardium that was percutaneously drained during the first ablation session. One patient had an episode of binocular blurred vision for 2 minutes without thrombus visible at transesophageal echocardiography. One patient developed a reversible ischemic neurological deficit 36 hours after ablation.

Angiograms performed just after ablation in 86 patients revealed a narrowing of the ostium in 5 ablated PVs; in an additional patient who received a single RF application at 30 W, the angiogram performed 5 minutes later showed stenosis of a 6-mm-diameter branch. PV angiograms (n = 115, including all those not angiographically studied just after ablation) 1 month after ablation (mean, 5 ± 4 months) demonstrated a persistent similar stenosis in these PVs plus an additional case that had not had an angiography just after ablation. Thus, a total of 6 ablated PVs (3%) had stenosis of the PV trunk: 1 patient had a double PV stenosis (RSPV and LIPV) and 4 had a single PV stenosis (3 LIPV and 1 LSPV). All stenoses were located at the most distal ablation site in the PV trunk. The mean gradient across the stenosis was 4 ± 2 mm Hg (range, 2 to 7 mm Hg), but no patient had pulmonary hypertension. However, 2 had symptoms: 1 described dyspnea during strong effort, and 1 patient developed hemoptysis and pleural effusion 1 week after interruption of warfarin (3 months after ablation). PV thrombosis demonstrated by spiral CT scan subsided after 2 weeks of heparin; a repeated scan showed only PV stenosis. Stress tests in 4 of 5 patients showed an unchanged performance relative to their peak effort before ablation. Analysis showed that the prevalence of PV trunk stenosis was the highest for LIPV (7% of LIPV versus 1.5% of all other PVs, P < 0.05) and when the maximal delivered RF power reached 45 W (5 of 37 versus 1 of 160, P < 0.01).

### TABLE 3. Characteristics of Patients With Single Versus Multiple Arrhythmogenic PVs

<table>
<thead>
<tr>
<th></th>
<th>Single PV (n=27)</th>
<th>Multiple PV (n=62)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinical characteristics</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sex, M/F</td>
<td>16/11</td>
<td>56/6</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Age, y</td>
<td>44±11</td>
<td>53±11</td>
<td>0.008</td>
</tr>
<tr>
<td>Duration of AF, y</td>
<td>4±4</td>
<td>6±5</td>
<td>NS (0.07)</td>
</tr>
<tr>
<td>AF maximal duration within 8 days before ablation, min</td>
<td>356±389</td>
<td>911±1029</td>
<td>0.006</td>
</tr>
<tr>
<td>Frequency of episodes, daily/every 2 days/weekly, n</td>
<td>12/6/9</td>
<td>46/7/9</td>
<td>0.04</td>
</tr>
<tr>
<td>Preprocedural AF duration on Holter, min/24 h</td>
<td>210±271</td>
<td>567±477</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Structural heart disease, n (%)</td>
<td>5 (19)</td>
<td>12 (19)</td>
<td>NS</td>
</tr>
<tr>
<td>Echocardiographic parameters, mm</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anterior-posterior LA dimension</td>
<td>36±6</td>
<td>40±7</td>
<td>0.008</td>
</tr>
<tr>
<td>Inferior-superior LA dimension</td>
<td>49±7</td>
<td>54±8</td>
<td>0.009</td>
</tr>
<tr>
<td>Medial-lateral LA dimension</td>
<td>36±6</td>
<td>38±6</td>
<td>NS</td>
</tr>
<tr>
<td>Inferior-superior RA dimension</td>
<td>43±6</td>
<td>51±8</td>
<td>0.007</td>
</tr>
<tr>
<td>Medial-lateral RA dimension</td>
<td>34±6</td>
<td>37±7</td>
<td>NS</td>
</tr>
<tr>
<td>Procedural data</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PV diameter, mm</td>
<td>16±3</td>
<td>16±4</td>
<td>NS</td>
</tr>
<tr>
<td>PV distribution, LS/RS/LI/RI, n</td>
<td>8/11/7/1</td>
<td>54/52/51/14</td>
<td>NS</td>
</tr>
<tr>
<td>Multiple procedures, n (%)</td>
<td>8 (30)</td>
<td>41 (66)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Cardioversion required, n (%)</td>
<td>7 (26)</td>
<td>31 (50)</td>
<td>0.02</td>
</tr>
<tr>
<td>Total duration, min</td>
<td>169±86</td>
<td>325±154</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Fluoroscopy time, min</td>
<td>49±26</td>
<td>97±43</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>RF duration, min</td>
<td>12±9</td>
<td>23±12</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

RA indicates right atrial.

*Patient 11 with a single atrial focus is not included.
No ostial PV stenosis was observed with a RF power limit of 30 W, including after circumferential ablation.

**Final Outcome**

With a mean follow-up of 8±5 months after discharge, AF was completely eliminated in 64 patients (71%) without antiarrhythmic drugs (success group), and anticoagulants were interrupted in 52. The other 26 patients were prescribed a drug that was ineffective before ablation, resulting in total elimination of AF in 12.

Table 4 compares the variables according to outcome. The presence of a single arrhythmogenic PV and successful elimination of PVPs were significant predictors of success. The success rate was 90% (36 of 40) when PVPs were

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**Figure 5.** Example of discrete arrhythmogenic triggering substrate within LIPV. Tracing (top) shows AF initiation by rapid train of discharges (•) from distal LIPV. Mapping during sinus rhythm revealed local PVP at roof extending deep into vein (arrows). Ablation at roof eliminated arrhythmia and spike both at ostium and distally, signifying discrete nature of this tissue.

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**Figure 6.** LIPV angiogram and electrograms during sinus rhythm showing late potentials distally within both branches. Sharp potential could be followed up to ostium; it is visible at roof and can be discerned at bottom (fusing with LA potential). Ablation at roof of ostium eliminated PVP only in superior branch. Further ablation at bottom of ostium eliminated activity extending into inferior branch. Thus, there appeared to be dual-input connection to LIPV.
Figure 7. Sustained regular atrial tachycardia from LSPV with various degrees of exit block, mainly 2:1. First tracing shows characteristic sequence of activation during sinus rhythm, with atrial activation preceding PVP (arrowhead) followed by 3 discharges (+) from PV, 2 of which are conducted to atria. During sustained tachycardia (second tracing), there is 2:1 exit block from within LSPV. Ablation limited to bottom sector of PV ostial (OS) perimeter (angiogram frame) terminated arrhythmia, and postablation mapping showed that local PV activity was dissociated distally (+) at slow rate.

Figure 8. Two examples of elimination of PV activity by RF delivery. In both examples, within 3 to 4 beats after onset of RF delivery (downward-pointing arrowhead), PV activity (upward-pointing arrowhead) is seen to disappear.
eliminated in all arrhythmogenic PVs and 55% (27 of 49) when PVPs were still persistent in ≥1 arrhythmogenic PVs.

**Discussion**

This study describes the results of electrophysiologically directed curative treatment for paroxysmal AF targeting the initiating triggers. It demonstrates plurifocal sources within the PV, a significant extent of local arrhythmogenic tissue, and better clinical outcome with the interruption of all conduction from the PV.

**Multiple Sources of Ectopy**

The study population included consecutively referred patients with drug-resistant paroxysmal AF without selection criteria, thus covering a broad range of situations encountered clinically. Spontaneous ectopy was absent in most patients, and the average AF duration was variable, ranging from 0 to 24 h/d. Spontaneous or provoked AF paroxysms were initiated.
Electrophysiologically Guided Ablation

Ablation guided by mapping ectopy was limited by its unpredictability, its inconsistent inducibility, and the risk of inducing AF requiring cardioversion. In patients with few ectopies, the arrhythmogenic PV could still be readily identified, and then all PVPs were ablated in sinus rhythm at the ostium, possibly eliminating multiple potential foci in the same trunk. LA or distal coronary sinus pacing separated the LA and PVPs, facilitating PVP identification notably in the left PV.

The results indicate that elimination of the PVPs correlated better with clinical success than the acute suppression of arrhythmias. However, it should be noted that a successful outcome could be observed without complete PVP elimination and that some “unsuccessfully” treated patients were remarkably improved with a previously ineffective drug. The PVP reflects activation of muscular LA bands extending into the PV with longitudinal, oblique, or complex courses and ending in a cul-de-sac or even looping back in the LA.1–3 The prevalence of multiple arrhythmogenic PVPs was high (69%), perhaps because of wide inclusion criteria, greater mapping experience, the use of provocative maneuvers with simultaneous exploration of different PV, and repeated electrophysiological studies in cases of recurrence. Moreover, multiple sources of AF in the same PV or even a focal source “wandering” to the atrial edge of the PV were observed. Multiple arrhythmogenic PVPs were associated with older age, longer AF duration, and larger atrial dimensions.

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PV stenosis occurred in 5 patients; however, all had no pulmonary hypertension, so no treatment was required. With the limitation of RF power to <30 W, acute and late PV stenosis was not observed at the ostium (although it occurred in a branch), but a longer follow-up is needed for confirmation. Given this uncertainty, a curative AF procedure based on PV ablation that uses current technologies should be reserved for patients with frequent drug-resistant paroxysmal AF who are exposed to the thromboembolic and hemodynamic risks of persistent arrhythmia and its inherent increased morbidity and mortality.

**References**

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