Assessment of Atrial Conduction Abnormalities in Patients with Hypertrophic Cardiomyopathy Before and One Year after Alcohol Septal Ablation

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Key Words
Hypertrophic cardiomyopathy · Atrial fibrillation · Atrial electromechanical delay · Percutaneous transluminal septal myocardial ablation

Abstract
Objectives: Intra- and interatrial electromechanical delay (AEMD) can be used to evaluate the development of atrial fibrillation (AF). Percutaneous transluminal septal myocardial ablation (PTSMA) is an alternative therapy for patients with hypertrophic obstructive cardiomyopathy (HOCM) that results in sustained improvements in atrial structure and function. We investigated the effects of PTSMA on the intra- and inter-AEMD of HOCM patients using tissue Doppler imaging. Methods: Conventional echocardiographic and AEMD parameters were obtained in 25 healthy controls and 31 HOCM patients before and 1 year after septal ablation procedures. Results: Compared with the healthy controls, the left atrial volumes indexed by body surface area (LAVI) and the intra- and inter-AEMD were significantly higher in the HOCM patients. At 1 year after PTSMA, the LAVI was decreased (37.2 ± 11.4 to 27.0 ± 8.5 ml/m\textsuperscript{2}, p < 0.001). The intra- and inter-AEMD were also significantly decreased (22.7 ± 9.2 to 16.6 ± 7.7 ms, p < 0.001 and 37.0 ± 8.4 to 26.6 ± 8.0 ms, p < 0.001, respectively). These changes correlated well with the reductions in LAVI (r = 0.83, p < 0.001; r = 0.66, p < 0.001). Conclusions: Both the intra- and inter-AEMD were significantly prolonged in the HOCM patients. PTSMA can improve the prolonged and inhomogeneous propagation of sinus impulses in atria.

Introduction

In patients with hypertrophic cardiomyopathy (HCM), atrial fibrillation (AF) is the most common clinical complication, with a prevalence exceeding 20% in recent studies [1–3]. For HCM patients, AF has been associated with substantial risks of heart failure-related mortalities, strokes and severe functional disabilities [4]. Left ventricular (LV) diastolic dysfunction is an important pathophysiological feature of HCM that contributes to elevated left atrial (LA) and pulmonary vascular pressure [5]. The thin-walled left atrium is sensitive to volume and pressure changes. Sustained elevated pressure on the left atrium leads to its remodeling, which is closely associated with the occurrence of AF [6–8].

S.C. and J.Y. contributed equally to this study and share first authorship.
Percutaneous transluminal septal myocardial ablation (PTSMA) is an alternative therapy for patients with hypertrophic obstructive cardiomyopathy (HCM) that results in long-term improvement in diastolic dysfunction and also partially reverses LA remodeling [9–11]. However, it is still uncertain whether PTSMA can reduce the risk of AF in HCM patients.

The echocardiographic evaluation of electrical events, especially atrial electromechanical delay (AEMD), is novel in cardiac ultrasound practice. With recent developments in tissue Doppler imaging (TDI), it is possible to precisely assess atrial mechanical events from different regions with high temporal resolutions. AEMD can be measured from the onset of the P-wave on the electrocardiogram (ECG) to the onset of atrial contraction as determined by TDI [12, 13]. Previous studies have proven that TDI-derived intra- and inter-AEMD values are predictors of AF [14, 15].

To our knowledge, AEMD has not been evaluated in HCM patients. In this study, we assessed AEMD using TDI in HCM patients before and 1 year after septal ablation to investigate the severities of their atrial electrophysiological abnormalities and the effects of PTSMA on the intra- and inter-AEMD.

**Methods**

**Study Population**

Between January and December 2010, 45 consecutive patients with symptomatic HCM were referred for PTSMA to our center. The diagnoses of HCM were obtained by means of 2-dimensional echocardiography in patients with interventricular septal thicknesses of ≥1.5 cm who had no other causes attributed to their LV hypertrophy. The selection criteria for PTSMA were as follows: the persistence of symptoms despite being administered the maximum tolerated dosage of medication, LV outflow tract (LVOT) gradient >50 mm Hg at rest or >100 mm Hg after provocation, accessible septal branches, typically of the left anterior descending coronary artery, the absence of a significant intrinsic abnormality of the mitral valve and other conditions for which cardiac surgery was indicated. Finally, 37 patients underwent septal ablation and 8 underwent surgical myectomy and mitral valve replacement because of serious mitral regurgitation. We excluded 6 patients due to the inability to deliver ethanol (3 cases) or incomplete echocardiographic parameters (3 cases) from the 37 patients who underwent PTSMA. The final cohort included 31 patients for whom the echocardiographic parameters were available at the baseline and 1-year follow-up evaluation. Twenty-five age- and gender-matched healthy controls were included from the subjects who visited our hospital for annual routine medical examinations. This study was approved by the ethics committee of the Fuwai Hospital, China.

**PTSMA Procedure**

A 6-French transfemoral temporary pacemaker was positioned in the right ventricle. After initial angiography to localize the origin of the septal perforating arteries, a 9-mm-long, oversized, over-the-wire balloon (1.5–2.5 mm) catheter was introduced over a 0.014-inch guidewire into the septal perforator artery and inflated. Contrast medium (SonoVue, Bracco Imaging BV, Geneva, Switzerland) was then injected through the balloon catheter lumen under continuous echocardiographic imaging. Once it was confirmed that the territory that was supplied by that septal branch corresponded to the basal septum causing the LVOT obstruction and that no other myocardial territory was involved, 1–4 ml of ethanol was injected into the septal branch. Repeated evaluations of the presence and extent of LVOT obstruction at rest and after amyl nitrite inhalation were performed using catheters and Doppler echocardiography to ensure the success of the intervention. A successful procedure was defined as a reduction in the LVOT pressure gradient of 50% of the baseline. For patients with <50% reduction in either the resting or provoked LVOT gradient, other septal arteries were targeted and treated in a similar fashion.

**Echocardiography**

Each subject underwent an echocardiographic evaluation using an ie33 echocardiographic system (Philips Medical Systems, Best, The Netherlands) that was equipped with an SS-1 transducer (frequency transmitted, 1.7 MHz; frequency received, 3.4 MHz) before and 1 year after the PTSMA procedure. One-lead ECG was recorded continuously. Two-dimensionally guided M-mode echocardiography was performed to measure the thicknesses of the interventricular septum and LV posterior wall (LVPW) and the diameter of the left atrium by standard methods. The LV end-diastolic and end-systolic volumes (LVEDV and LVESV), ejection fraction and LA end-systolic volume (LAESV) were calculated using a modified Simpson's biplane method in the apical 4- and 2-chamber views. The maximal early and late diastolic inflow velocities (E and A waves), E/A ratio, and deceleration time (DT) of the E wave were obtained using pulsed-wave Doppler. The sample volume was placed just below the level of the mitral leaflet tips in the apical 4-chamber view. The LVOT gradient was measured using continuous-wave Doppler in the apical 5-chamber view. The TDI of the mitral annulus movement was performed using the apical 4-chamber view. A 1.5-mm sample volume was placed at the lateral mitral annular sites. Analyses of the early and late diastolic peak velocities (Ea and Aa) and isovolumic relaxation time (IVRT) were performed. The AEMD was defined as the time interval from the onset of the P wave on the surface ECG to the beginning of the late diastolic wave (Aa wave) (fig. 1). It was measured from the lateral mitral annulus (mitral AEMD), inferoseptal mitral annulus (septum AEMD) and tricuspid annulus (tricuspid AEMD). All AEMD intervals were averaged over 3 consecutive beats. The difference between the mitral AEMD and tricuspid AEMD intervals was defined as the inter-AEMD, and the difference between the septum AEMD and tricuspid AEMD intervals was defined as the intra-AEMD.

**Electrocardiography**

All subjects underwent a 12-lead ECG recording after a 20-min resting period in the supine position at a paper speed of 50 mm/s and 2 mV/cm before and 1 year after PTSMA. The P-wave
durations were measured manually in all of the simultaneously recorded 12 leads of the surface ECG by 1 observer who was unaware of the study hypothesis. The mean P-wave duration of at least 3 complexes was calculated for each lead. The onset of the P wave was defined as the point of the first visible upward departure from the baseline for the positive waveforms and as the point of the first downward departure from the baseline for the negative waveforms. The return to baseline was considered to be the end of the P wave. The $P_{\text{max}}$ that was measured in from any of the 12 leads of the surface ECG was used as the longest atrial conduction time. The difference between $P_{\text{max}}$ and $P_{\text{min}}$ was calculated and defined as $P_d$.

Statistical Analyses

The data are presented as the means ± SD for the continuous variables or as percentages for the categorical variables. The clinical characteristics were compared using the t test for the continuous variables and the $\chi^2$ test for the categorical variables. The correlations between changes in the intra- and inter-AEMD and the improvement of the LA volume indexed to the body surface area (LAVI) were examined using the Spearman’s test. All of the probability values that were used were for 2-tailed tests. A value of $p < 0.05$ was considered to be indicative of a statistically significant result. The data processing and statistical analyses were performed using the SPSS 17.0 software (SPSS, Chicago, Ill., USA).

Results

The basic clinical characteristics of the 31 HOCM patients and 25 normal subjects are listed in table 1. There were no significant differences between patients and healthy controls in terms of age, gender, height, weight, heart rate, systolic blood pressure and diastolic blood pressure. In patients with HOCM, the mean NYHA class was $2.6 \pm 0.7$, despite optimal medical therapy that consisted of beta-blockers in 24 (77.4%) patients and calcium-channel blockers in 8 (25.8%).

The conventional echocardiographic characteristics are shown in table 2. Compared with healthy controls, HOCM patients prior to the PTSMA procedure exhibited higher septal and LV PW thicknesses, LAESV, LAVI and LV ejection fraction, but lower LVEDV and LVESV. The transmitral E-wave velocities, A-wave velocities and E/A ratios were similar between the 2 groups, whereas the DT, IVRT and E/Ea ratios were higher in patients with HOCM. The lateral Ea was significantly lower in the HOCM patients. The mean baseline resting LVOT gradient for the HOCM patients was $91.7 \pm 22.4 \text{ mm Hg}$.

P-wave indices are shown in table 3. The $P_{\text{max}}$ was significantly greater in the HOCM patients prior to PTSMA compared with the healthy controls. The $P_{\text{min}}$ did not differ significantly between the groups. The $P_d$ was significantly higher in the patients with HOCM. Table 4 shows the TDI-derived AEMD parameters. Compared with the controls, the AEMD values that were measured at the lateral mitral annulus, inferoseptal mitral annulus, and tricuspid annulus were significantly prolonged in the HOCM patients prior to PTSMA and both the intra- and inter-AEMD were significantly higher ($22.7 \pm 9.2$ vs.
branch blocks occurred at a rate of 51.6% (16 patients). Transitory trifascicular blocks occurred at a rate of 41.9% (13 patients). No patient underwent permanent pacemaker implantation following the procedure. There was no peri-interventional mortality during the observational period.

Changes in the conventional echocardiographic parameters in the patients with HOCM at the 1-year follow-up are shown in Table 2. The LVOT gradients decreased from 91.7 ± 22.4 to 27.1 ± 15.4 mm Hg following PTSMA. Both the LAESV (68.4 ± 21.3 to 49.2 ± 14.8 ml, p < 0.001) and LAVI (37.2 ± 11.4 to 27.0 ± 8.5 ml/m², p < 0.001) were significantly reduced at 1 year after PTSMA. There were no significant changes in E-wave velocity, A-wave velocity or the E/A ratio. Both the DT and IVRT were significantly shorter at the 1-year follow-up after septal ablation. Moreover, the lateral Ea and E/Ea ratio significantly improved following PTSMA. However, the diastolic function indexes including LAVI, E/A ratio, DT, IVRT and E/Ea ratio were still abnormal compared with the control group.

Although P-wave indices and TDI-derived AEMD were still longer than those in healthy controls, they were significantly improved after septal ablation. Table 3 and Table 4 list detail parameters. There was no significant change in $P_{min}$, but $P_{max}$ and $P_d$ decreased significantly (116.9 ± 7.0 to 105.0 ± 6.4 ms, p < 0.001 and 51.1 ± 6.9 to 40.5 ± 7.0 ms, p < 0.001, respectively) at 1 year after PTSMA. PTSMA significantly decreased all of the AEMD parameters in the patients (mitral AEMD 76.9 ± 11.5 to 64.0 ± 8.9 ms, p < 0.001; septum AEMD 54.1 ± 5.2 to 47.4 ± 3.5 ms, p < 0.001; tricuspid AEMD 39.8 ± 7.6 to 37.4 ± 5.3 ms, p = 0.046). Both the intra- and inter-AEMD were significantly shorter at 1 year after septal ablation (22.7 ± 9.2 to 16.6 ± 7.7 ms, p < 0.001 and 37.0 ± 8.4 to 26.6 ± 8.0 ms, p < 0.001, respectively). Moreover, the changes in the intra- and inter-AEMD correlated well with the reductions in LAVI (r = 0.83, p < 0.001, fig. 2a; r = 0.66, p < 0.001, fig. 2b).

**Discussion**

Our study provides unprecedented data regarding the AEMD parameters in HOCM patients before and 1 year after the PTSMA procedure. We found that compared with healthy controls, the AEMD measurements of different sites by TDI were significantly prolonged in the HOCM patients. Moreover, both the intra- and inter-AEMD were significantly higher. At 1 year after septal

| Table 2. Conventional echocardiographic characteristics of the healthy controls and HOCM patients at baseline and 1 year after PTSMA |
|------------------------|---------------------|---------------------|
| Characteristics        | Healthy controls (n = 25) | HOCM patients at baseline (n = 31) | HOCM patients 1 year after PTSMA (n = 31) |
| Septum thickness, mm   | 8.4 ± 1.3            | 19.6 ± 4.7*          | 16.5 ± 4.3* |
| LV PW thickness, mm    | 8.5 ± 1.5            | 12.6 ± 3.4*          | 11.7 ± 2.4* |
| LA diameters, mm       | 31.3 ± 3.2           | 40.8 ± 9.2*          | 36.9 ± 6.2* |
| LA ESV, ml             | 34.6 ± 8.6           | 68.4 ± 21.3*         | 49.2 ± 14.8* |
| LAV index, ml/m²       | 19.5 ± 4.2           | 37.2 ± 11.4*         | 27.0 ± 8.5* |
| LVEDV, ml              | 88.0 ± 20.3          | 76.9 ± 17.7*         | 82.5 ± 12.1* |
| LVE SV, ml             | 32.7 ± 12.6          | 18.4 ± 6.9*          | 24.6 ± 5.0h* |
| LVEF, %                | 64.6 ± 5.0           | 77.4 ± 6.6*          | 77.5 ± 6.0* |
| LVOT gradient, mm Hg   | 0                   | 91.7 ± 22.4*         | 27.1 ± 15.4* |
| Mitral E velocity, cm/s | 83.1 ± 15.2     | 83.8 ± 24.7           | 72.9 ± 18.4* |
| Mitral A velocity, cm/s | 65.1 ± 20.1     | 81.0 ± 34.5*          | 81.2 ± 32.3* |
| E/A ratio              | 1.4 ± 0.6            | 1.2 ± 0.6             | 1.0 ± 0.5*  |
| E deceleration time, ms | 174.6 ± 8.9 | 226.6 ± 11.7*         | 192.5 ± 10.1* |
| IVRT, ms               | 76.6 ± 6.8           | 122.8 ± 8.6*         | 96.2 ± 8.0*  |
| Lateral Ea, cm/s       | 11.9 ± 3.8           | 6.0 ± 1.9*            | 7.2 ± 1.8*  |
| Lateral Aa, cm/s       | 8.4 ± 2.4            | 7.5 ± 2.6             | 9.6 ± 3.0*  |
| E/Ea ratio             | 7.6 ± 2.5            | 15.1 ± 5.4*           | 10.9 ± 4.4* |

* p < 0.001 vs. healthy controls; $^b$ p < 0.01 vs. control subjects; $^c$ p < 0.001 vs. control subjects; $^d$ p < 0.005 vs. HOCM patients at baseline; $^e$ p < 0.01 vs. HOCM patients at baseline; $^f$ p < 0.001 vs. HOCM patients at baseline.

| Table 3. Electrocardiographic variables of the healthy controls and HOCM patients at baseline and 1 year after PTSMA |
|------------------------|---------------------|---------------------|
| Characteristics        | Healthy controls (n = 25) | HOCM patients at baseline (n = 31) | HOCM patients 1 year after PTSMA (n = 31) |
| $P_{max}$, ms           | 94.4 ± 7.5           | 116.9 ± 7.0*         | 105.0 ± 6.4* |
| $P_{min}$, s            | 63.6 ± 4.0           | 65.8 ± 5.5           | 64.5 ± 6.8 |
| $P_d$, ms               | 30.8 ± 5.6           | 51.1 ± 6.9*          | 40.5 ± 7.0* |

* p < 0.001 vs. healthy controls; $^d$ p < 0.001 vs. HOCM patients at baseline.

12.3 ± 6.0, p < 0.001 and 37.0 ± 8.4 vs. 21.1 ± 5.9, p < 0.001, respectively.

During the ablation procedure, the mean amount of alcohol that was injected was 2.49 ± 0.78 ml. In 24 patients, only 1 septal branch was treated and in the other 7 patients, 2 septal branches were treated. Right-bundle
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ablation, all of the AEMD parameters, including the intra- and inter-AEMD, had improved significantly.

AF is the most common sustained arrhythmia in patients with HCM. The recently reported prevalence of AF in several HCM populations ranged from 21 to 28% [1–3]. A previous study reported that HCM patients have a 4- to 6-fold increased risk of developing AF compared to the general population [2]. AF is an important cause of symptoms, morbidity and even mortality in patients with HCM. For these patients, the loss of the atrial systolic contribution to ventricular filling during AF may lead to clinical deterioration, involving progressive cardiac failure and systemic thromboembolism. Once AF has developed, the risk of ischemic stroke may increase 8-fold in HCM patients relative to those with sinus rhythm, presumably via an increased risk of cardioembolic stroke [4].

The main pathophysiological feature of HCM is diastolic dysfunction due to abnormal relaxation and decreased chamber compliance. Abnormal diastolic functioning results in elevated LV filling pressures that are transmitted back to the left atrium [5]. The thin-walled atrium is seriously affected by increased load, and the elevated LA pressures result in an increase in LA wall tension and enlargement of the LA [6]. Mild LA enlargement is common in HCM. Consistent with previous studies, in our study, the mean LA in the HOCM patients was much higher than that in the healthy controls. LA enlargement has been regarded as a sensitive and specific predictor of the occurrence of AF in HCM patients [7, 8]; it can affect atrial conduction time and may be closely related to atrial electrophysiological abnormalities [16, 17]. A previous study reported that not only structure remodeling but also electrical remodeling developed in the atria of patients with HCM [18]. Prolonged intra- and inter-AEMD and inhomogeneous propagation of sinus impulses are the major electrophysiological findings that predispose a patient to developing AF [19, 20]. TDI-derived AEMD is a new parameter for the assessment of atrial conduction abnormalities. AEMD has been evaluated by this method in some cardiac disorders. Ozer et al. [14] evaluated the intra- and inter-AEMD in patients with mitral stenosis and reported that the inter-AEMD was longer and correlated with

### Table 4. AEMD of the healthy controls and HOCM patients at baseline and 1 year after PTSMA

<table>
<thead>
<tr>
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<th>Healthy controls (n = 25)</th>
<th>HOCM patients at baseline (n = 31)</th>
<th>HOCM patients 1 year after PTSMA (n = 31)</th>
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<tbody>
<tr>
<td>Mitral AEMD, ms</td>
<td>51.7 ± 7.4</td>
<td>76.9 ± 11.5^c</td>
<td>64.0 ± 8.9^c</td>
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<tr>
<td>Septum AEMD, ms</td>
<td>39.4 ± 6.3</td>
<td>54.1 ± 5.2^c</td>
<td>47.4 ± 3.5^c</td>
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<tr>
<td>Tricuspid AEMD, ms</td>
<td>30.6 ± 5.4</td>
<td>39.8 ± 7.6^c</td>
<td>37.4 ± 5.3^c</td>
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<tr>
<td>Intra-AEMD, ms</td>
<td>12.3 ± 6.0</td>
<td>22.7 ± 9.2^c</td>
<td>16.6 ± 7.7^c</td>
</tr>
<tr>
<td>Inter-AEMD, ms</td>
<td>21.1 ± 5.9</td>
<td>37.0 ± 8.4^c</td>
<td>26.6 ± 8.0^c</td>
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^a p < 0.05 vs. healthy controls; ^b p < 0.01 vs. healthy controls; ^c p < 0.001 vs. healthy controls; ^d p < 0.01 vs. HOCM patients at baseline; ^e p < 0.001 vs. HOCM patients at baseline.

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**Fig. 2.** The relationship between the reductions in the LAVI and the changes in the AEMD parameters in an HOCM patient at 1 year after successful septal ablation. 

a The correlation of the changes in the intra-AEMD with the changes in the LAVI. 

b The correlation of the changes in the inter-AEMD with the changes in the LAVI.
P-wave dispersion, which is known to be a marker of AF. Cui et al. [15] proved that the prolongation of the intra- and inter-AEMD with TDI is an electrophysiological characteristic of atria that are prone to fibrillation. In this study, we demonstrated that both the intra- and inter-AEMD were significantly longer in HOCM patients than in healthy controls, indicating the existence of atrial electrophysiological remodeling and possibly predicting the development of new-onset AF in HOCM patients.

LVOT obstruction is present in up to two thirds of patients with HCM. Over the past 17 years, PTSMA has become an alternative to surgical myectomy to relieve LVOT obstruction and has been the preferred therapy since it was first reported in 1995 [21]. LV global diastolic function has been shown to improve immediately following a successful septal ablation. In addition, a long-term effect of PTSMA, involving the persistent improvement of global LV diastolic functioning for up to several years following the procedure, has been demonstrated [9, 10]. In this study, the DT, IVRT and E/Ea ratios significantly improved in the HOCM patients following PTSMA, again reflecting the improved diastolic function resulting from the procedure. Septal ablation has been previously shown to result in the decrement of LA dimensions using 2-dimensional echocardiography. Hage et al. [11] assessed the effects of PTSMA on LA volume and the LA ejection fraction using 3D echocardiography. They found that the LA volume was significantly reduced and the LA ejection fraction significantly increased in HOCM patients following septal ablation. Our data were consistent with the results of previous studies. In the HOCM patients, the LAVI was significantly reduced and the LA ejection fraction significantly increased at 1 year after PTSMA. These data confirm that PTSMA can partially reverse atrial structure remodeling and improve atrial function. John et al. [22] studied 21 patients with mitral stenosis undergoing mitral commissurotomy. Immediately following these procedures, LA volumes were reduced and the atrial electrophysiological and electroanatomical abnormalities improved consistently. This observation suggests that relieving the atrial stretch may reverse the AF substrate. However, it is unclear whether the improvement in atrial structure and function following PTSMA can reduce the risk of AF or the prevalence of paroxysmal AF in HOCM patients. Recently, Hosokawa et al. [23] reported on 3 HOCM patients with refractory paroxysmal AF. Following PTSMA, the incidence of episodes of paroxysmal AF became significantly less frequent. Because there were only 3 participants in their study, we could not obtain a validated conclusion. In our study, we found that both the intra- and inter-AEMD were significantly shorter in HOCM patients following PTSMA. Our results may indicate improved impulse conduction and less electrical heterogeneity in the atria following the septal ablation. This study indirectly supports the idea that PTSMA may reduce the incidence of AF in patients with HOCM. Moreover, we found that the decrements of the intra- and inter-AEMD correlated well with the reduced LAVI values. Therefore, we can speculate that LA structural reverse remodeling may reduce the arrhythmogenic substrate, ultimately resulting in atrial electrophysiological reverse remodeling in HOCM patients following PTSMA.

Our study has several limitations. First, the study population size was small. Second, changes in atrial conduction time were assessed before and 1 year after septal ablation, but patients were not followed up for arrhythmic episodes. We could not compare the incidence of AF in HOCM patients who underwent PTSMA with that in HOCM patients who do not receive septal ablation. Therefore, our findings should be interpreted with caution. Third, in this study, we used a modified Simpson’s biplane method based on 2-dimensional but not 3-dimensional echocardiography to measure the LA volume. Therefore, a large, prospective study of extended duration is necessary to confirm the effects of PTSMA on the primary prevention of AF in HOCM patients.

In conclusion, compared to the healthy controls, the intra- and inter-AEMD were significantly prolonged in the HOCM patients prior to septal ablation. PTSMA can improve upon LV diastolic dysfunction and partially reverse LA remodeling. Both the intra- and inter-AEMD were significantly reduced in the HOCM patients following PTSMA. The reduced intra- and inter-AEMD values correlated well with the changes in LAVI. Whether these reductions in the intra- and inter-AEMD reflect the decreased incidence of AF in the long-term requires further study.

Conflict of Interest

All authors have no conflicts of interest to declare.
References


