Decreasing the Pressure Gradient of the Left Ventricular Outflow Tract by Single-lead VDD Pacing in a Patient with Hypertrophic Obstructive Cardiomyopathy

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A 59-year-old woman with hypertrophic cardiomyopathy of 8 years duration, who had been taking β-blocker, was admitted to our hospital for exertional dyspnea and previous syncope. Cardiac catheterization showed a prominent left-ventricular outflow tract (LVOT) pressure gradient, and hypertrophic obstructive cardiomyopathy (HOCM) was diagnosed. To reduce LVOT obstruction, we implanted a single-lead VDD-mode pacemaker. Cardiac catheterization after the implantation revealed a remarkable decrease in the LVOT pressure gradient with short atrioventricular delay, 80 msec, and her symptoms disappeared. A single-lead VDD pacemaker is also a useful treatment for an HOCM patient due to the relative ease with which it can be implanted.

Key words: single-lead VDD pacing, hypertrophic obstructive cardiomyopathy (HOCM), left-ventricular outflow tract obstruction

Introduction

The degree of obstruction of the left ventricular outflow tract (LVOT) is an important determining factor in the clinical course of patients with hypertrophic cardiomyopathy (HCM). To relieve the LVOT obstruction, the initial therapeutic approach is pharmacotherapy such as β-blocker and verapamil. However, many patients develop a resistance to such therapies. The next approach is either surgery involving left ventricular septal myectomy or mitral valve replacement, or atrioventricular sequential pacing. But surgery frequently fails to reduce LVOT obstruction and leads to a 2-11% possibility of early mortality. Dual-chamber pacing has been shown to induce the reduction of the LVOT pressure gradient and to lead to improvement in symptoms of hypertrophic obstructive cardiomyopathy (HOCM). Recently single-lead VDD pacing has been used widely to patients with advanced atrioventricular block. We report a decrease in the LVOT pressure gradient by single-lead VDD pacing in an HOCM patient.

Case Report

On 25 December 1998, a 59-year-old woman was admitted to our hospital for further examinations due to exertional dyspnea, chest pain and a history of syncope. She had been diagnosed as having HCM in another hospital about 8 years before and had received medicinal treatment, β-blocker, and antiplatelet therapy. On admission, electrocardiography (ECG) showed T inversion in leads I, II, aVL, aVF, and V4-V6; ST elevation in leads V1 and V2; and high voltage (SV1 + RV5 = 8.59 mV). Cardiomegaly (cardiothoracic ratio: 63.7%) was observed by thoracic radiography (Fig. 1, left). Concentric left-ventricular hypertrophy with 17- to 19-mm wall thickness and LVOT narrowing were shown by echocardiography (Fig. 2). Hematology and biochemistry results showed iron deficiency anemia and high LDH (679 IU/l). We performed cardiac catheterization on the day of admission. Pressure study revealed a left ventricle (LV)-aorta (Ao) pressure gradient of 112 mmHg, and left ventriculography of right-anterior, oblique view showed a spade-like shape, which was compatible with HCM, during systolic phase. Coronary angiography showed no significant stenosis.
Fig. 1. Chest X-ray. Left, before implantation of pacemaker (CTR: 63.7%); right, after single-lead VDD pacemaker implantation (CTR: 61.0%). CTR: cardiothoracic ratio.

Fig. 2. Echocardiography. A, two-dimensional apical view during systolic phase; B, short axis view of LV at end-diastolic phase; C, systolic anterior movement (SAM) of mitral valve on M-mode. LV: left ventricle; LVOT: left ventricular out flow tract; Ao: Aorta; LA: left atrium.
Holter ECG recording showed occasional premature atrial contractions, including atrial tachycardia, but no long pause and no ventricular tachycardia that caused syncope, and no significant ST-T change. We thought her symptoms were caused by HOCM and the pharmacotherapy was not enough to decrease the high degree of pressure gradient in LV. Therefore it was necessary for her to undergo further treatment, either atrioventricular (AV) sequential pacing or surgery. She hoped the treatment of the pacing, and we implanted her a single-lead VDD-mode pacemaker (Unity 292-07, Intermedics) on 13 January (Fig. 1, right). Before this procedure, we did not identify the acute useful effects of this treatment with temporary pacing. One reason was that she hoped early treatment and short hospitalization. Another reason was that Fananaparzir et al reported the acute study is not always necessary because chronic effects are more beneficial than acute effects. One week after the implantation, the second cardiac catheterization was performed. We measured the LVOT pressure gradient (Fig. 3) while changing the AV delay intervals (AV delay). As AV delay was shortened, the LV systolic pressure and the LVOT pressure gradient decreased (Table 1). There were, however, no remarkable changes under 80 msec AV delay, and so we programmed the AV delay at 80 msec. Cardiac output was not measured with the thermo-dilution method, because the manipulation involved in such measurements might displace the pacing lead from the endocardium. Echocardiography revealed no differences in LV function between before and after implantation of the pacemaker. After the implantation of pacemaker, her symptoms were not occurred during the hospitalization.

**Table 1. Changes of pressure in LV and Ao at various AV delay**

<table>
<thead>
<tr>
<th>AV delay (msec)</th>
<th>LVSP (mmHg)</th>
<th>AoSP (mmHg)</th>
<th>LV-AoPG (mmHg)</th>
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<tr>
<td>250</td>
<td>250</td>
<td>152</td>
<td>98</td>
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<td>120</td>
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<td>60</td>
<td>198</td>
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LV: left-ventricle. Ao: Aorta. LVSP: left-ventricular systolic pressure, mmHg. AoSP: Aortic systolic pressure, mmHg. LV-AoPG: Pressure gradient between left-ventricle and aorta, mmHg.

**Discussion**

Dual-chamber pacing for patients with HOCM has been shown to be an effective treatment for reduction of...
LVOT obstruction and improvement of symptoms\(^9\)

Jeanrenaud et al.\(^1\) reported that decrease of the LVOT pressure gradient without decrease of arterial pressure and cardiac output was observed immediately after implantation of a DDD pacemaker, and after 3 to 30 months, the symptoms such as dyspnea, angina, and syncope, were also reduced. In another study that included a 1.5 to 3 months follow-up, the nearly identical hemodynamic changes to those reported by Jeanrenaud et al. were apparent, even when DDD pacing was discontinued.\(^1\) A long term study (2.3 years) revealed that the LVOT pressure gradient and LV systolic pressure continued to decrease overtime, indicating an excellent prognosis.\(^1\) It has been considered that these beneficial effects of DDD pacing to HOCM patients were due to the right ventricular apical preexcitation, which occurred asynchronous ventricular contraction and consequently decreased LVOT pressure gradient. Furthermore, in the long term study, chronic DDD pacing was reported to reduce LV wall thickness, especially of the basal anterior and distal anterior septal segments. Jeanrenaud and Kappenberger\(^1\) also demonstrated by echocardiography that DDD pacing reduces ventricular septal wall motion, which contributes to relieving LVOT obstruction. Regarding VDD pacing, Pak et al.\(^9\) reported that pacing shifts the end-systolic pressure-volume relation to rightward and increases end-systolic volume in LV, which reduces intracavity pressure gradient and lowers total chamber workload in HCM. Short AV delay VDD pacing is apparently superior to DDD pacing because it does not influence mean arterial pressure and stroke volume of the LV in patients with poor left ventricular function who have undergone coronary bypass surgery.\(^1\) Single-lead VDD pacing was established about 10 years ago, and its use has spread. It is easier to implant a VDD pacemaker than that a DDD pacemaker because the VDD uses only one lead. Furthermore, a VDD pacemaker preserves the patients' own sinus beat better than a DDD pacemaker does. Therefore we chose VDD pacing for our patient.

The optimal AV delay is an important factor when using atrioventricular sequential pacing for HOCM patients. The pacing must always excite the right ventricular apex before normal cardiac conduction and must induce atrial contraction with sufficient ventricular volume. Reports of the optimal AV delay have varied: 75-125 msec, 76 msec on average, and 75 msec.\(^1\) Jeanrenaud et al.\(^1\) suggested that optimal AV delay should be programmed individually. We determined the most beneficial AV delay to reduce LVOT obstruction in our particular patient and programmed in the delay of 80 msec. At this delay, echocardiography showed no remarkable change in her LV function before and after implantation of the pacemaker.

The implantation of a single-lead VDD pacemaker improved hemodynamics and symptoms in our patient. However, her intracavity pressure has remained high, about 200 mmHg, and further follow-up examination, therefore, will be required.

References


