Clinical Case: Place of Cardiac Stimulation in Asymptomatic Significant Chronic Aortic Insufficiency Associated with Sinus Dysfunction

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Abstract

The association between aortic insufficiency and sinus dysfunction is rare. We reported a case of significant chronic degenerative aortic insufficiency associated with sinus dysfunction. This was a 66-year-old patient on follow-up for asymptomatic chronic aortic insufficiency who was referred to us for a preoperative invasive hemodynamic assessment. The indication for surgery was based on the left ventricular dilation criteria. The echocardiography and cardiac catheterization showed a disappearance between the aortic insufficiency and the left ventricular repercussion. Upon discovery of the sinus bradycardia, the patient received a single chamber pacemaker set at a base rate of 70 seconds in AAIR at first and then the patient was measured remotely. At five months, cardiac pacing resulted in a significant reduction in the left ventricular diameter and volume on the echocardiography and magnetic resonance imaging; therefore surgery was postponed.

Keywords

Aortic Insufficiency, Sinus Dysfunction, Cardiac Stimulation, Sainte Clotilde Clinic, Reunion Island

1. Introduction

Dystrophic aortic insufficiency (AI) is common in western countries [1] [2], especially in the presence of aortic bicuspid disease. The association of a significant AI and sinus dysfunction is a rare entity. Bradycardia aggravates AI by increasing the duration of diastole, where the treatment of this bradycardia can delay the evolution of AI. We reported a case of significant chronic degenerative...
2. Observation

Mr. B.J, 66 years old, was hospitalized in the cardiology department at Clinique Sainte Clotilde (Reunion Island) from 01/11/2016 to 07/11/2016 for preoperative invasive evaluation of a leaking aortic valve. His cardiovascular risk factors were: arterial hypertension treated with Aprovel 150 mg/day and dyslipidemia with Lipanthyl 145 mg/day. In August 2016, a polysomnography revealed a permanent bradycardia of less than 40 cycles/min; there was no sign of sleep apnea syndrome. The patient was followed for several years for a degenerative AI on bicuspid valve without dilatation of the aortic ring. He was asymptomatic when running and jogging three times a week (around 240 minutes). On the physical examination, he presented a good general condition. His clinical parameters were: body mass index at 24 kg/m², temperature at 36.5 °C, blood pressure at 135/65 mm Hg and heart rate at 45 beats per minute. On cardiac auscultation the cardiac rhythm was regular with a diastolic murmur 2/6th along the sternum; there were signs of vascular hyperpulsatility with no signs of heart failure. Pulmonary auscultation was normal. The examination of the other apparatus was insignificant.

The biological blood tests were as follows: hemoglobinemia at 12.9 g/dl, prothrombin level at 90%, C reactive protein at 4 mg/l, glycaemia at 1.10 g/l, serum creatinine at 91 μmol/l with a clearance of 76 ml/min/m², normal hypersensitive troponin < 0.010 ng/l, Brain Natriuretic Peptide (BNP) at 70 μg/ml (normal < 100), normal TSH at 3 mU/l, total cholesterol at 1.44 g/ml, HDL at 0.37 g/l, LDL at 0.92 g/l, and triglycerides at 0.75 g/l.

The electrocardiogram showed a sinus bradycardia at 48 cycles per minute, a PR gap duration of 221 ms, an incomplete left block, and nonspecific abnormalities of repolarization.

The 24-hour rhythmic Holter confirmed a polysomnography data with a permanent sinus bradycardia, a mean night time frequency of less than 40/min and mean day time frequency of around 50/min. There was no prolonged pause or significant supraventricular or ventricular arrhythmia.

The stress test was conducted at 90% of the theoretical maximum frequency (MTF) which did not show a chronotropic deficit. The tension profile was normal.

In the transthoracic echocardiography (Figure 1) the left ventricle (LV) was hypertrophied (eccentric) and dilated with an end diastolic diameter (DTD) of 73.4 mm and a telencephalic diameter (SDR) of 47.6 mm (25.6 mm/m²), without a segmental kinetic disorder and a 60% systolic ejection fraction (EF). The filling pressures were normal. The left atrium was not dilated, there was an average AI on an anteroposterior bicuspid type 0. The ascending AO was not expanded measured at the Valsalva level at 42 mm and at the isthmus at 33 mm. The right cavities were not dilated, however, the inferior vena cava was dilated. Systolic pulmonary arterial pressures were normal and the pericardium was dry.
Figure 1. Cut-time motion (TM) on echocardiography showing LV and LVEF diameters.

On coronary angiography (Figure 2), the coronary arteries were atheromatous without significant focal lesions. On ventriculography, the left ventricle was dilated with an end-diastolic volume (EDV) of 149.5 ml/m^2 and a telesystolic volume (TSV) of 53 ml/m^2 with no segmental kinetic disturbance and a 64% left ventricular ejection fraction (LVEF). The sigmoidal aortogram revealed a medium to large AI.

The patient underwent implantation of a single-chamber pacemaker (AAI with placement of a Boston Accolade MRI SR compatible with thoracic magnetic resonance imaging (MRI) set at a base rate of 70 seconds in AAIR).

Five months later, the patient remained clinically stable. On echocardiography (Figure 3) and MRI (Figure 4) there was a coordinating and significant reduction in DTD (<65 mm), DTS (<45 mm) and LV volumes. The indication for surgery has been deferred.

On 05/07/2019 (32 months later), the patient was still asymptomatic with stable echocardiographic data (61 mm DTD and 60% LVEF).

3. Discussion

The size of the leak depends on the size of the regurgitant orifice, the duration of the diastole, and the VG-aortic pressure gradient. In chronic IA, there is a decrease in aortic diastolic pressure, a vascular erethism responsible for peripheral signs of AI, an increase in residual LV volume resulting in LV dilatation by volumetric overload followed by eccentric LV hypertrophy, at least in the early phase [3]. The bradycardia prolongs the duration of the diastole, therefore the volume of the regurgitation is very important for the long diastole. The optimal heart rate is probably around 80 - 90 beats/minute [4]. A transthoracic echocardiography, possibly supplemented by transoesophageal examination, is the key diagnostic examination. It allows a reliable and reproducible quantification of regurgitation, the evaluation of its etiology, its mechanism, its left ventricular repercussion and the associated lesions, in particular the ascending aorta [5]-[11].
MRI is complementary to or even an alternative in the evaluation of valvulopathies in both anatomical imaging and functional imaging [7] [12] [13]. However,
there is a considerable intra- and interobserver variability [14]. A long chronic AI can remain asymptomatic for long periods of time, possibly exceeding 10 years without impairment of systolic function, especially in rheumatic diseases. The risk of sudden death in the absence of symptoms is low (<0.2%/year) [15]. In asymptomatic AI, surgery is indicated [2] if there is impairment of the LVEF (≤50) or significant dilatation of LV (DTD > 70 mm or DTSVG > 50 mm) or (>25 mm/m²). Surgery has a risk of serious complications, hence the interest of waiting for the ideal moment before carrying out the surgery. The optimal date of surgery in severe asymptomatic AI is determined by a follow-up and an existence of a significant LV resonance according to the criteria described above. The surgical treatment usually consists of a valve replacement by mechanical or biological prosthesis. Medical treatment is appropriate for delaying surgery in asymptomatic patients with large AI without surgical criteria or in symptomatic and/or LV dysfunction patients who can’t be operated on due to major comorbidities. To our knowledge only one similar case has been published [16]. Our patient had a significant chronic AI, asymptomatic with an operative indication validated on echocardiographic criteria. According to current recommendations [2] [17], the surgical indication initially selected was based solely on VG dilatation criteria (normal LVEF, and normal BNP level). LV dilatation seemed disproportionate to the aortic insufficiency that wasn’t major. The presence of permanent diurnal and especially nocturnal sinus bradycardia confirmed by the rhythmic holter of 24 hours suggested that LV dilatation was related to aortic leakage but was worsened by long diastole associated with bradycardia. The international recommendations and the level of evidence in the management of a severe asymptomatic AI left us with the opportunity to reflect and discuss the operative indications and to consider another therapeutic alternative without altering the patient’s prognosis. The indication of stimulation in this case does not appear in the recommendations of cardiac pacemakers [18]. On the basis of physiopathology, our team has opted for an alternative stimulation by cardiac stimulation. The evolution was marked after five months by a clinical stability and a significant reduction of the diameters and volumes of the VG at the echocardiography and the MRI to exclude the patient of the surgical indication criteria. Thirty-two (32) months later the patient remained clinically stable and echocardiographically stable.

4. Conclusion

The association of AI and sinus dysfunction is not frequent. Significant sinus bradycardia should be sought in cases of severe asymptomatic AI while cardiac pacing helps to delay surgery.

Conflicts of Interest

The authors declare no conflicts of interest regarding the publication of this paper.
References


**Abbreviation**

AI: aortic insufficiency;
LV: left ventricular;
MTF: theoretical maximum frequency;
TDD: telediastolic diameter;
TSD: systolic diameter;
EF: ejection fraction;
TDV: telediastolic volume;
TSV: telesystolic volume;
MRI: magnetic resonance imaging.