Severe symptomatic aortic stenosis at high surgical risk: the new „classic” treatment

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Abstract: During the past 10 years, transcatheter aortic valve replacement (TAVR) has emerged as a valuable alternative treatment option for patients with severe symptomatic aortic stenosis at high surgical risk. We present the case of a 69 year-old man with symptomatic severe aortic stenosis deemed to be inoperable because of severe biventricular systolic dysfunction. TAVR was successfully performed as a last-resort option within a recently launched national programme covering such procedures in the public healthcare system.

Keywords: severe aortic stenosis, transfemoral TAVR.

INTRODUCTION

When patients with severe calcific aortic stenosis (AS) develop symptoms, survival at 2 years is less than 50%, and by five years less than 10% of these patients are alive1. Surgical aortic valve replacement (SAVR) improves not only symptoms, but also survival2. Transcatheter aortic valve replacement (TAVR) is recommended in patients with severe symptomatic AS who are not suitable for surgery and with an estimated life expectancy >1 year (class I, level B), or should be considered when there is a high surgical risk as assessed by a heart team (class IIa, level B)3.

CASE REPORT

We present the case of a 69 year-old patient who was diagnosed 8 months previously with symptomatic calcific aortic valve disease (severe stenosis, mild regurgitation) and biventricular dysfunction. Since the diagnosis he has suffered two episodes of acute decompensated congestive heart failure, the last one requiring high doses of intravenous furosemide with progressive loss of 10 kg of accumulated fluid. Associated comorbidities were type 2 diabetes mellitus and chronic kidney disease stage II.

On the current admission the patient described NYHA class III heart failure symptoms. The clinical examination revealed systemic congestion (jugular vein distention, hepatomegaly), and an ejection-type systolic murmur (grade 3/6) best heard in the aortic area that irradiated on both carotid arteries. He had no peripheral oedema or pulmonary rales. Electrocardiogram showed sinus rhythm and left ventricular hypertrophy with secondary ST-T changes. The BNP level was severely increased (4040 pg/ml).

Transthoracic echocardiography (TTE) revealed a severely stenotic, calcified trileaflet aortic valve (functional aortic valve area, AVA, of 0.6 cm², mean transaortic gradient of 50 mmHg) and mild aortic and mitral regurgitation (Figure 1 A and B). The left ventricle (LV) was hypertrophied, with normal cavity size and severely reduced global systolic function (LV ejection fraction, LVEF; of 25%, as assessed by biplane Simpson’s method). TTE also showed moderate secondary tricuspid regurgitation with an estimated systolic pulmo-

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nary artery pressure (PAPs) of 60 mmHg. The right ventricle (RV) was dilated (RV basal diameter 43 mm) with significant global systolic dysfunction (RV fractional area change of 25%, TAPSE of 12 mm).

Coronary angiography showed normal coronary arteries and right heart catheterization confirmed the isolated post-capillary pulmonary hypertension (PAP 69/24 mmHg, mean PAP 41 mmHg, mean pulmonary capillary wedge pressure 31 mmHg, pulmonary vascular resistance 3.03 WU). According to the current European Society of Cardiology guidelines on the management of valvular heart disease, the patient has a class I indication for aortic valve replacement. The surgical risk score was assessed by logistic EUROSCORE (16%) and by the Society of Thoracic Surgeons (STS) criteria (2.64%). Due to the severe biventricular systolic dysfunction, the 'heart team' considered this patient at high-surgical risk and TAVR was recommended.

Therefore, the patient was evaluated for the feasibility of TAVR by transesophageal echocardiography (TEE) and multi-slice angio-CT reconstruction for evaluation of both the aortic root and arch and also for the ilio-femoral axes. TAVR was considered feasible and given the results of TEE and CT tests, a 26 mm Edwards Sapien valve was chosen.

The procedure was performed under general anesthesia using transfemoral access, under 2D and real-time 3D TEE monitoring. After balloon predilatation of the stenotic valve (Figure 2), the prosthesis was positioned under fluoroscopic and TEE control and the valve was released by dilating the balloon during an induced episode of ventricular tachycardia. At the end of the procedure the prosthetic cusps were moving well (Figure 1C), the valve stent had a circular configuration, there was no significant residual transprosthetic gradient (Figure 1D), no significant transprosthetic regurgitation and only mild paraprosthetic regurgitation with a posterior location, at the level of a severe calcification of the aortic annulus (Figure 3).

The patient’s in-hospital course was good, with swift clinical and echocardiographic improvement. At discharge he was in NYHA class II with a global LVEF of 45% (Figure 4). PAPs decreased from 60 to 35 mmHg as assessed by CW Doppler echocardiography and the RV systolic function improved (FAC 35%, TAPSE 22 mm) (Figure 5). No neurological or haemorrhagic complications were observed. He was discharged under treatment with clopidogrel (for 6 months) asso-

Figure 1. TEE long-axis view and CW Doppler through the aortic valve before and after TAVR. A. Severely stenotic and calcified native aortic valve with a severely reduced systolic opening. B. CW Doppler through the native stenotic aortic valve showing high transvalvular gradient (50 mmHg mean gradient). C. Edwards Sapien aortic valve prosthesis with normal opening of the cusps. D. CW Doppler through the prosthesis: 11 mmHg mean gradient. Ao = aorta, LA = left atrium, LV = left ventricle.

Figure 2. Intraprocedural TEE, mid-esophageal long-axis view. Balloon predilatation (red arrow) of the stenotic calcific aortic valve (blue arrow). Ao = aorta, LV = left ventricle.

Figure 3. Post procedural TEE long axis view showing mild posterior paravalvular regurgitation (red arrow). Ao = aorta, LA = left atrium, LV = left ventricle.
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RE >20%. However, a number of comorbid conditions associated with adverse surgical outcomes are not included in these risk calculation scores, including right ventricular dysfunction/failure, such as in this case. A retrospective analysis performed by Cavalcante et al. showed that the presence of RV dysfunction in patients with AS provides important prognostic value and risk stratification adjunctive to STS score and it should be incorporated into the decision-making process.

It is without doubt that RV dysfunction in patients undergoing SAVR leads to adverse outcomes. On the other hand, Poliacikova et al showed no significant effect of RV dysfunction on mortality outcomes with TAVR. Furthermore, TAVR provided significant improvement of RV dysfunction at the 1 month and 6 months follow-up in another prospective echocardiographic study. However, one cannot make the assumption that all patients with RV dysfunction would be better treated by TAVR. Future studies to develop risk models for TAVR outcomes that include RV dysfunction as a variable are needed.

The guidelines recognize the limitations of the current risk scores and recommend that the indication for TAVR should be decided by a multidisciplinary heart valve team consisting of clinical cardiologists, interventional cardiologists, cardiovascular surgeons, imaging specialists, and anesthetists.

There is now a clear trend towards performing TAVR even in intermediate-risk patients. Offering TAVR to patients at lower risk for surgical morbidity and mortality is intuitively attractive, but remains controversial. The Nordic Aortic Valve Intervention (NOTION) trial was the first randomized trial addressing low- and intermediate-risk patients. The primary outcome of
stroke, myocardial infarction, and all-cause mortality at 1 year was 13.1% for TAVR and 16.3% for SAVR (p = 0.43), but these results are limited by the relatively small population included (280 patients). The results of larger randomized trials are awaited (SURTAVI trial, PARTNER II trial – intermediate risk cohort).

Perioperative complications (stroke, paravalvular regurgitation, renal failure, access site vascular complications and advanced atrioventricular block requiring permanent pacemaker implantation), as well as lack of data on long-term durability of the percutaneous valves, are among the key concerns for expanding indications for TAVR to lower-risk patients.

In summary, we presented the case of a 69 year-old man with symptomatic severe AS considered by the heart team to be at high surgical risk because of severe biventricular systolic dysfunction. TAVR was successfully performed as a last-resort option within a recently launched national programme covering such procedures in the public healthcare system.

**Conflict of interests:** none declared.

**References**