Mitral valve replacement with a pulmonary autograft: long-term follow-up in an infant

Anne Moreau de Bellaing, Amel Mathiron, Yves Lecompte and Pascal Vouhé

INTRODUCTION

A 7-month-old baby was referred for massive left atrioventricular valve regurgitation 4 months after complete repair. As the patient’s clinical conditions did not improve with medication, a surgical correction was indicated. Since the beginning of the operation, the repair of the valve was deemed impossible because of lack of the valvar tissue and severe malformation of the chordae. The LAV annulus was too small to accommodate the smallest mechanical valve, and the left atrium was not dilated enough to allow supra-annular implantation. We decided to use the modified ‘top hat’ mitral Ross procedure [Ross 2]. The pulmonary autograft was harvested. A cuff of the glutaraldehyde-treated autologous pericardium was sewn onto the proximal (infundibular) end of the autograft. The distal (pulmonary) end of the autograft was secured to the mitral annulus, whereas the native mitral apparatus was conserved to preserve the left ventricular function. At the level of the left atrial free wall, the pericardial cuff was progressively tailored and sewn onto the atrial wall to remain away from the ostia of the pulmonary veins and to maintain normal morphology of the autograft. At the level of the atrial septum, the autograft itself was used to close the os- tum primum defect avoiding distortion of the autograft. There was no complication during the postoperative period. The echocardiography showed normal functioning of the autograft.

The course of the child was uneventful for more than 11 years. However, mitral stenosis and regurgitation progressively developed without significant pulmonary hypertension (Fig. 1).

Reoperation was performed at the age of 11.75 years. The autograft was severely degenerated but not calcified; this and the native, still identifiable LAV were resected. A 21 mm-mechanical prosthesis was implanted in the supra-annular position. The pul- monary heterograft was removed as well. Four years later, the patient’s clinical conditions continue to be excellent.

Figure 1: Echocardiographic follow-up after the Ross II procedure.
COMMENTS

Whatever the mitral valve defect requiring surgery during childhood, conservative procedures should remain the first choice since they are the most compatible with regular child growth. Otherwise, when mitral valve repairs fail or are not technically feasible, surgical options for replacing the mitral valve in children are limited. Because mechanical mitral valve replacement (MVR) was challenging for infants or in developing countries, Ross and Kabbani [1] updated, in 1997, the procedure using the autologous pulmonary valve as an inverted autograft. This avoids anticoagulation treatment and size mismatch between the prosthesis and the LAV annulus.

The long-term course of 92 patients, all older than 4 years, was reported with good results in terms of freedom from degeneration, reoperation and death [2]. In a review of the literature, Athanasiou et al. [3] reported similar results, again without including infants. So far, there is only 1 medium-term follow-up reported in an 11-month-old infant [4]. An MVR was required 7 years after the Ross II procedure.

The conventional technique involves the implantation of the pulmonary autograft into a prosthetic conduit [1]. The latter reinforces the external wall of the autograft avoiding longitudinal deformation or constriction, but it prevents any growth potential of the valve. Slit opening of the prosthetic conduit, along 1 or both of its sides, as suggested by Kabbani et al. [2], has not been shown to solve the growth issue.

We used a technical modification without prosthetic support. Recently, Jeong and Yun [5] reported a similar procedure, but the long-term follow-up of their patient is still unknown. The autologous pericardial cuff allowed to secure the valve on the left atrial free wall and the interatrial septum (Fig. 2). Precise tailoring and suturing of the cuff are of utmost importance to avoid any distortion and dysfunction of the autograft.

CONCLUSION

In conclusion, we report the favourable long-term follow-up of a child having undergone MVR using a pulmonary graft without a prosthetic material. In case of irreparable congenital mitral malformation, this procedure could be an alternative substitute, especially for infants in whom mechanical MVR remains a concern. Further experience is obviously necessary before recommending its widespread use. However, this alternative should be kept in mind when the MVR is required in infants.

Conflict of interest: none declared.

REFERENCES