Truncus arteriosus: successful surgical correction without the use of a valved conduit

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SUMMARY

A new operation for the restoration of continuity between the right ventricle and pulmonary circulation in truncus arteriosus in infancy without the use of a tube graft is described. This was achieved by anastomosing the inferior margin of the detached confluent branch pulmonary arteries to the apex of a vertical infundibular ventriculotomy to form the posterior wall of the reconstructed right ventricular outflow tract. A patch of pericardium was used to form the anterior wall and complete the primary cardiac repair.

The surgical management of the infant with truncus arteriosus remains a challenge, although recent reports have indicated satisfactory results of corrective surgery in patients under one year of age. Major centres, however, still report high operative risks in small infants. The principles of repair and of intraoperative and postoperative management are becoming increasingly clear in this complex condition but invariably reports advise the placement of an appropriately sized Rastelli conduit, valved or otherwise, between the right ventricle and pulmonary arteries and accept the need for reoperation and further reconstruction at a later date. All conduits are liable to progressive stenosis, endocarditis, and valve failure if a valved conduit is used; reoperation carries its own important risk. Even if a second operation for conduit replacement is successful, it remains likely that a considerable number of children will require further operation especially after ten years with such a conduit.

The preliminary work of Bailey et al has led us to devise an operative technique that obviates the need for conduit placement in the common types of truncus arteriosus, allows for potential growth, and seems to have important advantages over conventional techniques while still retaining the advantages of anatomical correction. We have used this technique to correct truncus arteriosus in an infant weighing 3 kg.

Case report

A 10 week old infant was referred to the Royal Hospital for Sick Children, Edinburgh with a history of failure to thrive and recurrent chest infections. She was tachypnoeic with a respiratory rate of 75 per minute, there was intercostal indrawing, the peripheral pulses were all of full volume with a collapsing quality, and the blood pressure in the right arm was 100/30 mm Hg. On auscultation both heart sounds were single and there was a long, harsh grade 4/6 systolic murmur and a grade 3/6 early diastolic murmur that were both loudest at the lower left sternal edge. A chest x-ray showed cardiomegaly with a cardiothoracic ratio of 65% and increased pulmonary vascularity. Electrocardiographic examination showed biventricular hypertrophy with ST-T changes in the left lateral precordial leads.

The clinical features suggested a diagnosis of truncus arteriosus and this was confirmed on cardiac ultrasound examination which was interpreted as being in keeping with a type 1 truncus arteriosus and truncal valve regurgitation. After 48 hours of medical support and treatment with sedation, digoxin, frusemide, and ventilation she was taken to the operating theatre for repair of the defect.

OPERATIVE PROCEDURE

Figure 1a shows the anatomy of the vessels. There was no main pulmonary artery, both the right and left pulmonary arteries arose from the left lateral margin of the aortic root, the right artery being inferior in position and passing below the left vessel. Both arteries were extensively mobilised into both hila. The infant was cooled to 20–21°C while on
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Fig 1  Diagrams showing restoration of continuity between the right ventricle and pulmonary circulation. (For details see text.)

cardiopulmonary bypass; the pulmonary vessels were gently clamped to avoid bypass perfusion of the lungs and they were resected from the aorta. Then a cannula was manipulated into the left coronary ostium and cardioplegia was continued. A superior right ventriculotomy was made and through it the ventricular septal defect was closed. The aortotomy was repaired. The inferior margin of the right pulmonary artery was sewn to the superior margin of the ventriculotomy by means of a continuous to and fro Prolene suture. This suture was carried around one half of the inferior circumference of the right pulmonary artery opening, bringing it down on to the upper one quarter to one third of the ventriculotomy margin (fig 1b). The anterior defect thus created was closed with a carefully tailored patch of xenograft bovine pericardium (Shiley Corporation). The procedure established generous egress from the right ventricle.

POSTOPERATIVE COURSE
The infant's postoperative course was uneventful. She was extubated 48 hours after operation and required no inotropic support. Postoperatively there was a residual grade 2/6 to 3/6 early diastolic murmur. Fifteen days after operation cardiac catheterisation was performed. The right ventricular systolic pressure was 55 mm Hg and the pulmonary artery systolic pressure was 35 mm Hg, and a left ventricular peak systolic pressure was 90 mm Hg. Oximetry demonstrated a small residual left to right shunt of 1:5:1. There was no pressure difference across the left ventricular outflow tract. A selective right ventricular angiogram demonstrated free egress of contrast into the pulmonary arteries with no important narrowing or obstruction (fig 2). Left ventricular angiography showed a small residual ventricular septal defect at the lower margin of the

Fig 2  Postoperative right ventricular angiogram (anteroposterior and lateral views) demonstrating free egress of contrast into the pulmonary arteries with no important narrowing or obstruction.
ventricular septal patch that had been inserted at operation. On aortography there was a mild to moderate degree of aortic valvar regurgitation.

Discussion

Orthodox right ventricular outflow tract reconstruction for other types of congenital heart disease is accomplished by transannular patching with pericardium or preclotted Dacron and follow up studies show an apparently normal propensity for growth. What is less certain is the efficacy of the valve leaflets and whether they are necessary for medium and long term success since adult cases can be symptom free in the face of chronic pulmonary valvar incompetence which, while not necessarily benign, is well tolerated. Our experience with this case agrees with that of others who have used non-valved conduits in this condition; we cannot agree with Bailey’s conclusion that a competent pulmonary valve is required for short term survival. Moreover, the development in Bailey’s case of a right ventricular outflow tract aneurysm within 40 days is strongly suggestive of important residual right ventricular hypertension that could have contributed to early death.

There are practical and theoretical advantages in applying this technique of right ventricular outflow tract reconstruction to types I and II truncus arteriosus. These include (a) ease of operation, (b) growth potential of the right ventricular/pulmonary artery junction, and (c) avoidance of the acknowledged severe early and late complications of conduit placement in very young infants. We acknowledge the persistence of the native truncal valve regurgitation and the possibility that this could worsen with time.

References

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