Temporary iatrogenic fetal tricuspid valve atresia in a case of twin to twin transfusion syndrome

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Abstract
This report describes the sequence of events in the development and subsequent spontaneous resolution of functional tricuspid valve atresia in the donor fetus in a case of twin to twin transfusion syndrome. Fetoscopic laser coagulation of the placental anastomoses was performed at 20 weeks' gestation. Subsequently, there was evidence of increased placental vascular resistance in the donor twin and major impairment of right ventricular function with no forward flow through the tricuspid valve. During the next four weeks, however, there was spontaneous and complete recovery of ventricular function and resolution of the functional tricuspid valve atresia. These findings suggest that alterations in fetal haemodynamics may result in structural cardiac abnormality and may be the precursors of some forms of congenital heart disease.

Almost all monochorionic twin pregnancies have communicating placental vessels between the umbilical cord insertions. Where there are artery to vein anastomoses a chronic shunt of blood from one fetus to the other can cause twin to twin transfusion syndrome. The donor fetus becomes hypovolaemic and oliguric, whereas the hypervolaemic recipient twin develops cardiac insufficiency despite compensatory polyuria. In severe cases presenting in the second trimester perinatal mortality is 80–100% owing to spontaneous abortion, intrauterine death, or very premature delivery of a growth-retarded donor twin and hydroptic recipient twin. The prognosis is improved by repeated amniocentesis for drainage of large volumes of amniotic fluid. This reduces the risk of spontaneous abortion or very premature delivery related to polyhydramnios; approximately 40% of the babies survive.1 A new method of treating this condition is fetoscopically directed laser coagulation of the communicating placental blood vessels.2 In this case report, which is part of a large prospective study, we describe the sequence of events in the development of functional tricuspid atresia in one of the fetuses after laser treatment and its spontaneous resolution.

Case report
Acute polyhydramnios developed at 20 weeks' gestation in a twin pregnancy. Ultrasound showed discordant fetal size and polyhydramnios, and a distended bladder in the bigger fetus (recipient). The smaller fetus (donor) appeared stuck to the common placenta owing to severe oligohydramnios. Doppler studies (Acuson 128, Mountain View, California, USA) of the umbilical arteries showed high placental resistance in the donor, with a pulsatility index (PI) of 1.65, whereas the PI was normal (0.99) in the recipient. Both fetuses had normal waveforms in the descending thoracic aorta. The recipient showed mild tricuspid regurgitation. Under local anaesthesia the communicating vessels on the chorionic placental surface were coagulated by fetoscopically directed (through a 2.7 mm diameter cannula) neodymium yttrium aluminium garnet laser.2

At 21 weeks' gestation, ultrasound examination showed a normal amount of amniotic fluid in both sacs. However, the donor, still identifiable by its smallness, had a pericardial effusion. Doppler studies showed increase in umbilical artery PI in both fetuses (recipient 1.44; donor 2.08).

At 22 weeks' gestation, echocardiography of the donor fetus showed biventricular enlargement and a hyperechogenic tricuspid valve that did not open during diastole (fig 1).
There was no blood flow from the right atrium (RA) to the right ventricle (RV) during diastole. Colour Doppler showed no tricuspid atrioventricular diastolic inflow or systolic regurgitation (fig 2). The right ventricle was filled by retrograde diastolic blood flow in the ductus arteriosus and pulmonary trunk through an incompetent pulmonary valve (fig 3). In both the descending thoracic aorta (PI = 3.34) and umbilical artery (PI = 3.13) there was reversed diastolic flow indicating very high placental resistance. Venous blood flow studies showed reversed velocities in the ductus venosus during atrial contraction, which is never seen in normal fetuses (fig 4). In the former recipient fetus, echocardiography showed normal findings and the PI s in the umbilical artery (1.16) and thoracic aorta (2.11) were within the normal ranges for gestation.

At 24 weeks’ gestation there was intermittent tricuspid regurgitation (fig 5) and at 25 weeks there was transtricuspid right ventricular filling combined with pansystolic regurgitation (fig 6). There was still reversed diastolic flow in the umbilical artery and no end diastolic flow in the thoracic aorta.

At 31 weeks’ gestation the cardiac abnormalities had completely resolved. There was normal opening of the tricuspid valve and right ventricular filling and no systolic regurgitation. End diastolic flow in the umbilical artery had changed from reversed to absent and in the thoracic aorta it was positive.

At 33 weeks’ gestation there was spontaneous rupture of the membranes and two girls were delivered by caesarean section. The birthweights of the former recipient and donor twins were 2100 g and 1500 g respectively and they both had an uneventful postnatal course. When they were a week old, echocardiography of the smaller baby showed mild hypoplasia of the right ventricular cavity but the tricuspid valve was morphologically normal. There was normal ventricular filling without tricuspid regurgitation or pulmonary incompetence. When they were two months old right ventricular growth was normal and the tricuspid annulus was 10 mm in the smaller baby. However, there was a small atrial septal defect with pure left to right shunt. The bigger baby (former recipient) had a small atrial septal defect and mild pulmonary stenosis.

Figure 2 Colour Doppler imaging of left ventricular inflow. There was no blood flow from the right atrium (RA) to the right ventricle (RV) during diastole.

Figure 3 Pulsed Doppler investigation of blood flow at the level of the pulmonary valve showing pulmonary incompetence with retrograde filling of the right ventricle.
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Discussion

Laser coagulation of the placental anastomoses was associated with a major increase in resistance to flow from the donor fetus into the placenta, as shown by reversal of diastolic blood flow velocities in the umbilical arteries and the descending thoracic aorta. Consequently, there was severe impairment of right ventricular function, presumably because the right ventricular afterload was increased. Right ventricular function was compromised to such an extent that there was no forward flow through the tricuspid valve and no discernible right ventricular myocardial contraction. The tricuspid valve was functionally atretic and stationary in a closed position. The abnormal venous waveforms, with marked reversed flow during atrial contraction, are the consequence of the cardiac findings.

During the subsequent weeks there was improvement in the flow velocity waveforms in the umbilical artery, from reversal to absence of diastolic velocities, and this was associated with recovery of cardiac function. The postnatal finding of a smaller than usual right ventricular cavity is consistent with a period of prenatal arrested growth with subsequent recovery. The apparently complete recovery of ventricular function shows that even severely compromised function may be reversible if the abnormality in cardiac loading resolves.

The mild pulmonary valve stenosis present postnatally in the recipient twin is the only apparent residual cardiac abnormality after the prenatal compromise of right ventricular performance that was part of the natural history of the syndrome. The cardiac abnormalities in the donor twin were an unintended consequence of treatment.

The sequence of events suggests that alterations in fetal haemodynamics may result in structural cardiac abnormality and may be the
precursor of some forms of congenital heart disease. Tricuspid regurgitation is a common prenatal abnormality, whereas isolated tricuspid regurgitation is uncommon postnatally, because most cases recognised prenatally are lethal. It seems likely that the most common cause is increased right ventricular afterload. It has long been suggested that abnormalities in fetal flow patterns may be implicated in some forms of structural congenital heart disease, and this case report provides objective support for this hypothesis.

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