Angiographic diagnosis of anomalous coronary artery in tetralogy of Fallot

Julene S Carvalho, Celia M C Silva, Michael L Rigby, Elliot A Shinebourne

Abstract

**Objective**—To obtain angiographic views in tetralogy of Fallot that can show whether or not an anomalous coronary artery passes anterior to the right ventricular outflow tract.

**Design**—(a) A 10 year retrospective review of all patients who underwent repair of tetralogy of Fallot up to December 1990; (b) a prospective study of 30 children undergoing routine cardiac catheterisation.

**Patients and methods**—295 cases in whom standard angiographic views had been used were reviewed retrospectively. Thirty non-consecutive children with tetralogy of Fallot were studied prospectively, including one child previously studied in whom diagnosis of an unsuspected anomalous coronary artery was made only at operation. The aortogram was performed with >45° caudocranial and 20°-30° left anterior oblique angles.

**Setting**—Tertiary referral centre.

**Results**—Ten of the 295 cases reviewed were shown to have a coronary vessel traversing the right ventricular outflow tract. In one case the diagnosis was suspected before operation but it was missed in the others. Even in retrospect we could not be certain of the precise anatomy with the use of standard angiographic views. In the prospective study the caudocranial aortogram showed the aortic valve face on in all the patients. The right ventricular outflow tract lay in a left and anterior (seen as superior) position in relation to the aortic root. Thus any vessel crossing the outflow tract could be identified. Identification of the aortic cusps allowed precise definition of the origin of the coronary arteries. All but four had normal origin and course of the coronary arteries. Four had paired left anterior descending arteries (including the restudied patient), in all cases with a large vessel originating from the right coronary artery passing across the right ventricular outflow tract.

**Conclusions**—Important anomalies of the coronary arteries in tetralogy of Fallot may remain undiagnosed if standard angiographic projections are used. Aortography with >45° caudocranial and 20°-30° left anterior oblique angles allows precise definition of the anatomy and certainty as to whether any major vessel crosses the right ventricular outflow tract. Interpretation, however, can only be correct if the projection is technically adequate with a view of the aortic valve face on. Furthermore, a normal bifurcation of the left main stem does not exclude a second left anterior descending artery crossing the pulmonary outflow tract.

(Tetralogy of Fallot is one of the conditions for which cardiac catheterisation is still performed before operation in most centres. This is because non-invasive techniques may not always permit identification of abnormalities of pulmonary arteries or of coronary anatomy. The sensitivity and specificity of angiography for diagnosing coronary anomalies, however, remains uncertain. Surgical and postmortem series suggest that important coronary anomalies occur in 2% to 10% of patients. The most common anomaly, the left anterior descending artery arising from the right coronary artery and crossing the right ventricular outflow tract, is important in young infants undergoing complete repair. It may influence surgical mortality and morbidity and has implications regarding technique and age of repair. Though age in itself may not represent an incremental risk factor for surgery, preoperative diagnosis of a vessel crossing the right ventricular outflow tract remains a contraindication for early repair if the use of an extracardiac conduit is to be avoided. We have found it difficult to make the diagnosis with certainty based on standard angiographic views and decided to review our 10 year experience. The use of a caudocranial aortogram has been described previously in patients with aortic valve disease. More recently its use in children with transposition of the great arteries has been reported. This prompted us and others to assess its use in tetralogy of Fallot. A prospective study was then carried out with this purpose.

**Patients and methods**

To obtain retrospective data records of patients who underwent complete repair of tetralogy of Fallot at the Royal Brompton Hospital from January 1980 to December 1990 were reviewed. To obtain prospective data we studied 30 non-consecutive patients with tetralogy of
Fallot undergoing routine cardiac catheterisation. This included one child with an anomalous left anterior descending coronary artery missed at previous cardiac catheterisation but identified at surgery.

Twenty had a left and 10 a right sided aortic arch. Age varied from one month to 11·8 years (median 17 months). Either a retrograde (n = 25) or anterograde approach (n = 5) was used for the biplane aortogram, which was performed with a minimal 45\(^\circ\) caudocranial and 20\(^\circ\)-30\(^\circ\) left anterior oblique angulation as well as a straight lateral projection. In four of the cases studied by a retrograde approach aortography was performed with a pigtail catheter but in general a standard National Institute of Health catheter gave better opacification of the coronary arteries. When possible, however, an anterograde approach with a balloon tipped catheter with the balloon inflated in the ascending aorta, was preferable.

Results

RETROSPECTIVE STUDY

Records for 295 patients were reviewed. Ten (five males and five females) had an associated anomalous coronary artery. Age at time of repair ranged from 10 months to 7·6 years. All had undergone preoperative echocardiography and cardiac catheterisation. Aortography and left ventriculography had been performed in one or two of the following views: anteroposterior, lateral, or left anterior oblique for the aortogram and lateral, long axis, or right oblique for the left ventriculogram. In only one case was there a high degree of doubt that an abnormality of coronary artery anatomy was present. Six had had previous palliative surgery. One of these had first stage correction (relief of the right ventricular outflow tract but no closure of the ventricular septal defect) and an anomalous vessel was identified at surgery. Subsequently, but before definitive repair, angiography was carried out with selective coronary artery injections. In this case, all six standard views, anteroposterior, lateral, right and left anterior oblique with and without craniocaudal tilt, were used but still there was no certainty as to the course of the vessel passing in front of the right ventricular outflow tract. In all cases it was necessary to modify the technique of repair as the usual incision to the right ventricular outflow tract could not be made. A combination of modified right ventriculotomy, right atriotomy, and pulmonary arteriotomy was used. Good relief of right sided obstruction was achieved in all patients but four required interposition of a conduit between the right ventricle and the pulmonary artery.

PROSPECTIVE STUDY

Selective coronary arteriography (fig 1) in one case and placement of a second catheter in the right ventricular outflow tract (fig 2) in another facilitated understanding and confirmed how the anatomy is displayed in the caudocranial angiographic projection with the left oblique angle. In a third case the aorta was reached by an anterograde route with the venous catheter and spillage of contrast into the right ventricle and the right ventricular outflow tract by accident contributed further to the understanding of the relation of the coronary arteries to the right ventricular outflow tract (fig 3). The course of the left anterior descending artery was seen by looking at the interventricular septum with the same angles during left ventriculography in two cases. Four patients (including the patient with a known abnormality) were shown to have a large left anterior descending artery from the right coronary artery as well as a second left descending artery from the left coronary cusp, so two vessels were supplying the area of the left anterior descending artery (figs 4 and 5). In the remaining 26 patients the coronary distribution was normal and none had an abnormal branch across the area.

Figure 1  Selective right (A) and left (B) coronary artery injections displayed in the caudocranial left anterior oblique view in a patient with tetralogy of Fallot and normal distribution of coronary arteries. Note the course of the vessels as they arise from the right (A) and left (B) coronary sinuses. Cs, circumflex artery; LAD, left anterior descending artery; LMS, left main stem; RCA, right coronary artery.
Angiographic diagnosis of anomalous coronary artery in tetralogy of Fallot

of the right ventricular outflow tract. In three cases the conus branch, arising from the right coronary artery, was enlarged but did not run across the outflow tract area. In 21 cases the anatomy was known or was confirmed at time of surgery, including three of four with abnormality. In one, the vessel originating from the right coronary artery and supplying the left descending territory was considered smaller than the artery coming from the left coronary sinus. Relief of the right ventricular outflow tract obstruction was through the pulmonary artery and right atrium. Three patients have undergone palliative procedures, including one with abnormality. The rest await surgical treatment.

The lateral aortogram performed simultaneously was unhelpful in assessing the coronary artery anatomy with certainty but was used to show the presence or absence of an arterial duct.

Discussion
Successful display of coronary artery anatomy in tetralogy of Fallot, reported by means of a ventriculogram or flush aortogram or with selective coronary artery injections has been reported. This has not been our experience with aortography and we do not carry out selective coronary arteriography routinely. Cross sectional echocardiography may also be used. Although it is possible to identify abnormal vessels across the right ventricular outflow tract, there may not be an adequate view of the coronary anatomy in a significant proportion of patients. Berry et al correctly and blindly distinguished normal and abnor-
ormal patterns in the coronary circulation of some patients with tetralogy of Fallot but it was not possible to see the coronary arteries well enough in over 20% of their patients. In only two cases was this due to large patient size.  

To show a branch of a major coronary artery passing anterior to the right ventricular outflow tract in tetralogy of Fallot it is necessary to show the coronary arteries clearly and also to know where the right ventricular outflow tract lies in the projection chosen for the aortogram. Overlapping of the coronary branches is the rule with most standard views. This leads to confusion and failure to recognize abnormal patterns when interpreting the angiograms. In our retrospective series the prevalence of surgically important anomalies of the coronary artery was 3-4% but the diagnosis was not made with certainty in any of the cases although suspected in one. Conversely, in our prospective series of patients in whom a caudocranial aortogram was performed, the diagnosis was made with certainty in four cases and, equally important, we could be certain that no major coronary artery passed anterior to the right ventricular outflow tract in the rest.

With the use of a caudocranial angle for the aortogram the aortic valve is seen face on. Its three cusps are clearly seen and the origin and course of the coronary arteries are easily followed by the absence of overlapping as pointed out by Mandel et al. Moreover, and of crucial importance in patients with this condition, knowing that the position of the right ventricular outflow tract is to the left and slightly anterior to the aortic valve allows identification of any abnormal branch crossing the area. To be certain, however, a correct projection of the base of the heart is required (as if the observer were looking at the heart from below and slightly from the left). The aortic arch itself is foreshortened and therefore the aortic valve must be seen face on or nearly so. This view is thus analogous to the echocardiographic parasternal short axis section at the level of the aortic valve. In both, the normal left mainstem can be seen passing behind (beneath) the right ventricular outflow tract and dividing into left anterior descending and circumflex arteries, but we have already emphasized the potential deficiency of echocardiography in the individual patient.

The caudocranial aortogram, when performed correctly, allows distinction between right and left, posterior and anterior (but not inferior and superior) relations (fig 3). Thus it should be possible to detect an abnormal origin or course of a coronary artery and to see its anterior or posterior relation to the pulmonary outflow tract. Any major coronary branch passing anterior to the right ventricular outflow tract is likely to be of importance to the surgeon, and management can then be directed accordingly. It is also important to see both coronary arteries as identification of one normal left anterior descending artery does not exclude a paired artery originating from the right coronary artery and crossing the pulmonary outflow tract. Our current practice is to use 45° caudocranial and 30° left anterior angulation in all patients. We have shown that by using this angiographic projection it is possible to detect these abnormalities before operation.

Angiographic diagnosis of anomalous coronary artery in tetralogy of Fallot.

J S Carvalho, C M Silva, M L Rigby and E A Shinebourne

Br Heart J 1993 70: 75-78
doi: 10.1136/hrt.70.1.75

Updated information and services can be found at:
http://heart.bmj.com/content/70/1/75

These include:

Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/