Morphological characterisation of ventricular septal defects associated with coarctation of aorta by cross-sectional echocardiography

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SUMMARY Cross-sectional echocardiograms of 18 neonates and infants with coarctation of the aorta and ventricular septal defect were retrospectively assessed. With a combination of subcostal and precordial cuts the site and anatomical relations of the defects were determined. In one case there was a muscular trabecular ventricular septal defect. Three had a malalignment defect with associated left ventricular outflow tract obstruction. In two there was a doubly committed subarterial defect with associated malalignment of the point of continuity between the aortic and pulmonary valves and the crest of the trabecular septum. Twelve cases had a perimembranous defect, with varying degrees of extension into the inlet, trabecular, or outlet septum. In this group 10 had associated aortic override, with varying degrees of left ventricular outflow tract narrowing. Abnormal insertion of the tricuspid valve was observed in 10 cases, such that it partly obscured the ventricular septal defect.

Thus in most cases of coarctation with ventricular septal defect, the morphology of the ventricular septal defect and ventricular outflow tracts is such that left ventricular ejection is directed towards the pulmonary artery rather than the aorta.

In a recent study on cross-sectional echocardiography in aortic interruption it was confirmed, as has previously been described from anatomical studies, that the ventricular septal defect is situated such that left ventricular blood preferentially ejects into the pulmonary artery. In the majority of cases this was the result of posterior deviation of the outlet (infundibular) septum into the left ventricular outflow tract.

In a further study on the cross-sectional echocardiographic features of coarctation of the aorta in the sick neonate and infant, it was also observed that in some with a ventricular septal defect there was associated aortic override or posterior infundibular displacement. A recent morphological study by Anderson, Lenox, and Zuberbuhler of the specimens in the cardiopathological collection of The Children's Hospital of Pittsburgh, with ventricular septal defect associated with aortic coarctation, showed that aortic override or posterior displacement of the outlet septum was a common association, occurring in 24 out of 25 hearts studied. As it is likely that a necropsy study does not present a true picture of the spectrum seen in clinical practice, we investigated in more detail the cross-sectional echocardiographic findings of those patients who had presented over the past two years in The Hospital for Sick Children, London, with a diagnosis of coarctation of the aorta and ventricular septal defect.

Subjects and methods

The cross-sectional echocardiograms were examined retrospectively and the results correlated with available angiographic or necropsy material. Only those cases with atroventricular and ventriculoarterial concordance and “normal relations” were included in the study. During the examination the level of atroventricular valves and the status of the muscular septum were assessed in a subcostal and apical long axis cut at right angles to the inlet septum (“four chamber”). Then with clockwise rotation of the

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transducer the left ventricular outflow tract and perimembranous region were inspected. With further clockwise rotation the morphology of the outlet septum was assessed. Next a precordial long axis cut was performed at right angles to the outlet septum and with clockwise rotation of the transducer the right ventricular outflow tract could be seen. The presence of an overriding aortic valve was only assessed in the precordial long axis cut. Finally a precordial short axis cut at the level of the papillary muscles was obtained, and the transducer then directed so as to scan the left ventricular outflow tract. With these combined manoeuvres the presence and site of a ventricular septal defect could be determined, provided it fell within the range of axial and lateral resolution.

In one patient who died the specimen was examined and tomographic sections obtained.

### Results

Eighteen cases were judged to have a ventricular septal defect by cross-sectional echocardiography. The defect was considered large in 15, medium sized in one, and small in two. Angiocardiographic or necropsy confirmation was available in all of those with a significant ventricular septal defect. A summary of the echocardiographic findings is shown in the Table. One case had posterior muscular trabecular ventricular septal defect which was plugged by tricuspid valve.
tissue during systole. Three had a malalignment ventricular septal defect, with posterior displacement of the outlet septum in the left ventricular outflow tract (Fig. 1). Two cases had a doubly committed subarterial ventricular septal defect without extension back towards the membranous septum (Fig. 2). In both of these the point of continuity between the aortic and pulmonary valves was displaced relative to the crest of the trabecular septum, again resulting in malalignment and the substrate for preferential flow from left ventricle to pulmonary artery (Fig. 2).

Four cases had a perimembranous trabecular ventricular septal defect, two of whom had associated aortic override (Fig. 3). In one the left ventricular outflow tract appeared narrow (Fig. 3). Four cases had a perimembranous defect which excavated both the trabecular and outlet septum (Fig. 4). In one there was an associated low trabecular ventricular septal
defect. Aortic override was present in all cases, with the left ventricular outflow tract appearing narrow in three. In the other four cases the defect was perimembranous and extended into the inlet, trabecular, and outlet septum (Fig. 4). Aortic override was present in all cases and the left ventricular outflow tract appeared narrow in one (Fig. 4).

In no cases did the mitral valve obstruct the left ventricular outflow tract. Similarly there was also no evidence of a subaortic ridge. In nine cases with a perimembranous defect and one with a muscular trabecular defect, the tricuspid valve was inserted abnormally high into the infundibular septum (Fig. 3). The medial papillary muscle was also situated more anteriorly in the trabecular septum (Fig. 5). The result of this was that in the precordial long axis cut at right angles to the outlet septum and the four chamber aortic root cut the defect appeared to be obscured by the tricuspid valve (Fig. 5). In four cases the tricuspid valve tissue entered the left ventricular outflow tract through the ventricular septal defect during systole (Fig. 4 and 5). This appearance could be seen both in the precordial short axis cut as the transducer was scanned towards the left ventricular outflow tract (Fig. 6) and in the precordial long axis cut. Necropsy confirmation of the defect was available in one case with a perimembranous trabecular ventricular septal defect (Fig. 7).
Fig. 4  These images are from a patient with a coarctation of the aorta and perimembranous defect with extension into the inlet septum, trabecular septum, and outlet septum. The upper left picture is a precordial long axis cut showing the narrow left ventricular outflow tract seen during diastole. The lower left picture is taken during systole and shows the tricuspid valve bulging into left ventricular outflow tract. Also note the aortic override. The upper right picture is a precordial cut with clockwise rotation showing excavation of the infundibular septum. Note that the infundibular septum is not displaced anteriorly. The lower right picture is a precordial four chamber cut with the atrioventricular valves at the same level. Note the ventricular septal defect which is indicated by the small arrows. For abbreviations see Fig. 1, 2, and 3.
Fig. 5. The upper left picture is a precordial long axis cut in a patient with coarctation and ventricular septal defect. Note that the tricuspid valve is seen crossing the right ventricular cavity in this cut. Though there is aortic override the left ventricular outflow tract appears to be adequate. The lower left picture from the same patient is taken during systole and shows the tricuspid valve protruding into the left ventricular outflow tract. The right picture is a precordial four chamber cut with superior angulation of the transducer. Note the large ventricular septal defect indicated by the arrows and the tricuspid valve which obscures the defect. PM, papillary muscle. For rest of abbreviations see Fig. 1, 2, and 3.
Fig. 6  The lower left picture is a precordial short axis cut in coarctation of the aorta, ventricular septal defect, and left ventricular outflow tract obstruction. Note the narrow left ventricular outflow tract seen in this cut. The lower right picture from the same case is taken during early diastole and demonstrates that the abnormal anterior echo is connected to the tricuspid valve. The upper picture from the same case is during maximum diastole. PV, pulmonary veins. For rest of abbreviations see Fig. 1, 2, and 3.
Fig. 7  The upper picture is a precordial long axis cut in coarctation of the aorta, ventricular septal defect, and abnormal insertion of the tricuspid valve. Note during systole the tricuspid valve protrudes into the left ventricular outflow tract. The tricuspid valve tissue obscures the ventricular septal defect. In addition, note the aortic override. The lower picture is a specimen from the same patient cut to simulate the echocardiographic appearance. For abbreviations see Fig. 1 and 3.

Discussion

Cross-sectional echocardiography has proved to be a reliable method for both identifying and accurately locating the position of a ventricular septal defect, provided the defect falls within the range of axial and lateral resolution of the equipment. Visualisation of the left ventricular outflow tract is readily achieved by the above technique, which, in the presence of an adjacent ventricular septal defect, frequently showed severe narrowing in the absence of a demonstrable pressure gradient at the site.

The range of defects seen in this series differs slightly from that reported after anatomical examination of the necropsy population of The Children's Hospital of Pittsburgh. It is likely that these findings represent a truer spectrum of the frequency seen in clinical practice of the types of ventricular septal defect associated with aortic coarctation. In the necropsy study four cases had a muscular outlet defect
Coarctation, ventricular septal defect

Fig. 8  The upper picture is a precordial long axis cut in coarctation of the aorta without a ventricular septal defect. Note that the aortic root appears to override the interventricular septum, despite the lack of ventricular component. The lower picture is also from a patient with coarctation and intact ventricular septum. Note that the tricuspid valve is inserted abnormally. For abbreviations see Fig. 1, 3, and 5.

associated with posterior displacement of the outlet septum, while 20 out of the remaining 21 had a large perimembranous defect with extension into all components of the septum together with associated aortic override.

Even so, in this clinical series aortic override associated with a large perimembranous defect was seen in 10 cases (58%), while left ventricular outflow tract narrowing caused by posterior displacement of the outlet septum was seen in five (28%). In all cases with aortic override the outlet septum was normally positioned (Fig. 3 and 4), unlike patients with Fallot's
tetralogy in whom the override is a consequence of anterior displacement of the outlet septum.

The abnormal insertion of the tricuspid valve, such that it crosses the ventricular septal defect before inserting into the underside of the outlet septum provides a pocket of tissue which obscures the defect. This feature was also noted in the necropsy series. The appearance of the tricuspid leaflet tissue crossing the place of the transducer in the long axis has only in our experience been seen previously in patients with double outlet right ventricle and subpulmonary ventricular septal defect, or complete transposition associated with a ventricular septal defect.\(^9\)\(^10\)

Right ventricular hypertension is common in infants with coarctation of the aorta even when there is no ventricular septal defect or a small one.\(^11\) Thus in many cases with coarctation and ventricular septal defect, and in all cases with associated complete transposition or double outlet right ventricle, the combination of a high right ventricular pressure and a high insertion of the tricuspid valve allows the tricuspid tissue tag to be forced through the defect, this being readily demonstrable echocardiographically. Whether this tissue eventually closes the defect is open to speculation and can only be determined by a prospective study. Interestingly, similar tricuspid valve anatomy (Fig. 7) was seen in some patients presenting to our institution with a diagnosis of coarctation of the aorta but with an intact ventricular septum. Aortic override was also observed in some of these patients and it was evident that a pre-existing ventricular septal defect had been completely closed either \textit{in utero} or during the first weeks of life (Fig. 8). Thus, as in the necropsy series, the most common type of ventricular septal defect seen in aortic coarctation in patients who survive the critical period of early infancy is one whose morphology, in combination with that of the ventricular outflow tracts, might be expected significantly to reduce the amount of left ventricular blood reaching the aortic isthmus. This entire complex, and the association of tricuspid valve abnormalities in those with aortic override, are readily demonstrated by cross-sectional echocardiography.

**References**


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