Two-dimensional sector scanner echocardiography in cyanotic congenital heart disease

A. B. HOUSTON, N. L. GREGORY, AND E. N. COLEMAN

From the Department of Cardiology and the University Department of Child Health, Royal Hospital for Sick Children, and the Department of Clinical Physics and Bio-Engineering, Glasgow

A wide angle (60°) mechanical sector scanner producing a real-time two-dimensional echocardiogram has been used to assess the anatomy of the great arteries in 30 infants and children with cyanotic congenital heart disease. Longitudinal scans show the relations of the mitral valve and ventricular septum to the posterior great artery, and transverse scans show the spatial relations of the great arteries.

In 19 patients the echocardiographic examination was preceded by angiocardiographic diagnosis. Echocardiographic examination in 10 with complete transposition of the great arteries showed parallel great arteries; in 4 with Fallot's tetralogy, normally related great arteries with a narrow right ventricular outflow and an overriding aorta; in 4 with double outlet right ventricle, parallel great arteries, but in only 2 of these an unambiguous right ventricular origin of both great arteries; in 1 with persistent truncus arteriosus, a single artery with abnormal cusp echoes.

In 9 infants and 2 children, aged 9 and 16 years, echocardiographic examination was followed by angiocardiography. In the 9 infants the assessment of great artery size and relation by echocardiography closely correlated with that determined by angiocardiography. In the older child whose previous cardiac catheterisation without angiocardiography had suggested transposition of the great arteries, echocardiography showed normally related great arteries which were subsequently confirmed by angiocardiography. In the younger child with corrected transposition of the great arteries the echocardiographic appearance was initially misinterpreted as normally related great arteries.

Two-dimensional echocardiography, because of its capacity to display the anatomy of the ventricular outflow tracts, permits better planning of the subsequent cardiac catheterisation and angiocardiography in children with cyanotic congenital heart disease.

The findings of time-position echocardiography in transposition of the great arteries (Dillon et al., 1973; Gramiak et al., 1973), double outlet right ventricle (Chesler et al., 1971), persistent truncus arteriosus (Chung et al., 1973a), Fallot's tetralogy (Chung et al., 1973b), total anomalous pulmonary venous drainage (Paquet and Gutgesell, 1975), and in cyanotic newborn infants (Godman et al., 1974) have been reported. To increase the information available from echocardiography two-dimensional real-time scanning systems have recently been introduced and have proved of value in cyanotic congenital heart disease (Sahn et al., 1974; Henry et al., 1975; Maron et al., 1975).

Subjects and methods

Thirty patients with cyanotic congenital heart disease were studied. In 19 with known anomalies of the origin of the great arteries the diagnosis of
complete transposition of the great arteries (10),
double outlet right ventricle (4), Fallot's tetralogy
(4), or persistent truncus arteriosus (1) had been
made by angiocardiography; the age range was from
4 weeks to 16 years.
In 11 patients echocardiographic assessment of
great artery size and position was made before elucida-
tion of the abnormality by angiocardio-
graphy; the age range was 3 days to 16 years.
The ultrasonic equipment used was a mechanical
60° sector scanner (Shaw et al., 1976) incorporating
later developments (Houston et al., 1977). In the
initial part of the study a 2-5 MHz 12-5 mm diam-
ter transducer was used but when a 5-0 MHz
6 mm diameter transducer became available it was
used where appropriate. All illustrations in this
report were obtained as still frame photographs
from the television monitor. The convention adopted
in transverse views is that the patient’s left is to the
right of the illustration, anterior being upwards.
The scanning planes employed were longitudinal
and transverse as previously described (Houston
et al., 1977). The initial approach was, where pos-
sible, to obtain a longitudinal section through the
great artery continuous with the anterior mitral
leaflet and to determine its relation to the ventricu-
lar septum. A transverse scan was then obtained
through this great artery and an attempt made to display
a normal right ventricular outflow and main pul-
monary artery crossing in front. In the absence of
this normal relation the scanning plane was ad-
justed in an attempt to show the adjacent circular
spaces characteristic of the parallel vessels in
transposition of the great arteries (King et al., 1973;
Henry et al., 1975); to confirm that such a circular
space represented a great artery the semilunar valve
echo was sought. In complex abnormalities where
there was no mitral leaflet continuity with the
posterior artery, the technique was modified with
the intention of locating the ventricular septum and
great arteries.

Results

COMPLETE TRANPOSITION OF
THE GREAT ARTERIES
In all 10 patients with complete transposition of the
great arteries a high transverse scan showed two
circular spaces (Fig. 1) which were confirmed as
representing great arteries by the demonstration
of central linear cusp echoes in diastole. It was not
possible to display both great arteries at valve
level in a single transverse scan in 3 of the 10
patients (Fig. 2). These illustrations were obtained
with the scanner directly transverse and tilted to
carry out a headwards sweep as with time-position
echocardiography. In Fig. 2a a small posterior
vessel, the pulmonary artery, is apparent. As the
scanning plane passes upwards (Fig. 2b) the pul-
monary cusp echoes are lost and a larger anterior
vessel, with central cusp echoes, appears to the
right; this indicates that the valve of the anterior
vessel lies to the right and superior to the posterior
one. The posterior vessel appears to give rise to an
elongated transverse echo-free space passing pos-
terior to the anterior artery (Fig. 2b and c). We
consider that this represents the bifurcation of the
pulmonary artery and confirms the identity of the
posterior vessel. Definite evidence of branching of
the posterior great artery was noted in 4 of the 10
patients. In all 10 the echocardiographic spatial
relations were found to correspond to those deter-
mined by angiocardiology.

DOUBLE OUTLET RIGHT VENTRICLE
The 4 patients with double outlet right ventricle
had echocardiographic evidence of two parallel
vessels in the transverse scan. When the posterior
great artery and the ventricular septum were
viewed simultaneously their relation was seen to
vary throughout the heart cycle, the septum moving
posteriorly with respect to the great artery root dur-
ing systole. The diastolic position was considered to
give the best indication of the relation of the great
arteries to the ventricles. This confirmed that both arteries arose from the same ventricle in 2 cases. In the other 2, less than one-and-a-half vessels appeared to arise from the right ventricle. Longitudinal scans showed an abnormal relation between the anterior mitral leaflet and the posterior artery in 1 patient from each of these pairs; mitral-semilunar valve continuity was interrupted by intervening conal tissue.

**PERSISTENT TRUNCUS ARTERIOSUS**
The one patient with persistent truncus arteriosus had a single large anterior vessel (Fig. 3) overriding the ventricular septum. Within this vessel a double central echo and several eccentric echoes were seen, representing the abnormal truncal valve cusps.

**FALLOT'S TETRALOGY**
In the 4 patients with tetralogy a high transverse scan confirmed normally related great arteries. Three of these showed overriding of the aorta on longitudinal scan (Fig. 4). The fourth showed no overriding in diastole but the aorta and ventricular septum moved asynchronously, the septum being posterior to the anterior wall of the aorta in systole.

**ECHOCARDIOGRAPHY BEFORE ANGIOCARDIOGRAPHY**
The Table shows the data on the 11 cyanotic patients (including 9 infants) in whom an echocardiographic assessment of great artery anatomy was made before the elucidation of the anomaly by

---

**Fig. 2** High transverse scans from a patient with complete transposition of the great arteries and pulmonary artery hypoplasia obtained as the scanning plane is tilted superiorly from (a) the level of the pulmonary valve (PV), to (b) the aortic valve, and to (c) a supravalvar level. The light in the upper left of the picture indicates the onset of the QRS complex, i.e. end-diastole.

**Fig. 3** Transverse scan from the patient with persistent truncus arteriosus at the level of the truncal valve cusps.
angiocardiology. In all 9 infants the echocardiographic assessment of great artery size and relation showed close correlation with the angiographic findings. In 2 of the 4 patients with complete transposition of the great arteries, branching of the posterior artery confirmed that it was the pulmonary artery. The 16-year-old patient appeared to have normally related great arteries with a narrow right ventricular outflow and main pulmonary artery. The diagnosis of transposition of the great arteries with tricuspid atresia had been made at a previous cardiac catheterisation but because of arrhythmia no angiogram had been obtained. Angiocardiography performed subsequent to the echocardiographic examination confirmed that he had normally related great arteries. An incorrect diagnosis of normally related great arteries with a narrow outflow was made in the 9-year-old child with corrected transposition. Review of the videotape recording showed that the anterior wall of the anterior left vessel had been misinterpreted as a thick immobile pulmonary valve; failure to recognise the semilunar cusp echo posterior to this caused the diagnostic error.

**Discussion**

Time-position echocardiography allows exact measurement of the distances between structures along the axis of the ultrasound beam. The determination of the relations of different structures not lying along a single beam is dependent on the operator's assessment of small changes he makes in transducer angulation; the technique is difficult, particularly when it is applied to the rapidly beating heart of a restless infant. Two-dimensional echocardiography allows exact determination of the relation of structures, thus facilitating the assessment of great artery anatomy. Other workers who have reported on the use of real-time two-dimensional echocardiography in the assessment of great artery anomalies have concluded that the method is of diagnostic value (Sahn et al., 1974; Henry et al., 1975; Maron et al., 1975). There were some differences between their techniques and their findings in normal high transverse scans; their experience and ours have been discussed elsewhere (Houston et al., 1977). We have found that a high transverse scan can show the aortic origin as a circular space with a central diastolic cusp echo, and the pulmonary artery either in the same manner when the great arteries are parallel, or as an anterior crescentic space representing the right ventricular outflow and main pulmonary artery crossing in the normal way.

In the tetralogy of Fallot or other conditions in which there are normally related great arteries the circular aorta and crescentic right ventricular outflow and main pulmonary outflow are recognised. In complete transposition of the great arteries the diagnosis is suggested by the abnormal relation of the great arteries shown by echocardiography as

---

**Table  Echocardiographic assessment of the anatomy of the great arteries compared with subsequent angiographic findings**

<table>
<thead>
<tr>
<th>Age</th>
<th>Transducer (MHz)</th>
<th>Echocardiogram</th>
<th>Angiocardiogram</th>
</tr>
</thead>
<tbody>
<tr>
<td>3 d</td>
<td>5 o</td>
<td>CTGA</td>
<td>CTGA</td>
</tr>
<tr>
<td>8 d</td>
<td>5 o</td>
<td>CTGA</td>
<td>CTGA</td>
</tr>
<tr>
<td>10 d</td>
<td>5 o</td>
<td>Normal relations; narrow RVO and MPA</td>
<td>CTGA</td>
</tr>
<tr>
<td>6 w</td>
<td>2/5</td>
<td>CTGA; narrow MPA</td>
<td>CTGA; hypoplastic MPA</td>
</tr>
<tr>
<td>2 m</td>
<td>2/5</td>
<td>Normal relations; wide RVO and MPA</td>
<td>TAPVD</td>
</tr>
<tr>
<td>4 m</td>
<td>2/5</td>
<td>Normal relations; narrow RVO and MPA</td>
<td>Primitive ventricle with outflow chamber</td>
</tr>
<tr>
<td>6 m</td>
<td>2/5</td>
<td>Overriding aorta; narrow RVO</td>
<td>Fallot's tetralogy</td>
</tr>
<tr>
<td>10 m</td>
<td>2/5</td>
<td>Overriding aorta; narrow RVO</td>
<td>Fallot's tetralogy</td>
</tr>
<tr>
<td>9 y</td>
<td>2/5</td>
<td>Normal relations; narrow RVO</td>
<td>Corrected TGA</td>
</tr>
<tr>
<td>16 y</td>
<td>2/5</td>
<td>Normal relations; narrow RVO and MPA</td>
<td>Primitive ventricle with outflow chamber; normally related arteries</td>
</tr>
</tbody>
</table>

CTGA, complete transposition of the great arteries; RVO, right ventricular outflow; MPA, main pulmonary artery; RV, right ventricle; TAPVD, total anomalous pulmonary venous drainage.
two echo-free circles which represent the vessels lying parallel at their origin; it is not dependent upon the separate identification of the aorta and the pulmonary artery. If, in some of the rarer types of transposition of the great arteries (Van Praagh, 1973), there should be any diagnostic difficulty it may be important to show branching of an artery which can thus be positively identified as the pulmonary artery. A problem is likely to arise when only one artery can be viewed in transverse section and no right ventricular outflow or other artery is visible, making a differential diagnosis between persistent truncus arteriosus and pulmonary atresia difficult. The abnormal eccentric cusp echoes reported by Sahn et al. (1974) and confirmed in our patient would be an important feature in the positive diagnosis of persistent truncus. For the diagnosis of double outflow right ventricle as defined by Lev et al. (1972) more than one-and-a-half great arteries should arise from the right ventricle. Echocardiographic examination of our patients showed that the relation of the great arteries to the ventricular septum varied throughout the cardiac cycle. We considered that the position in diastole was likely to show the closest agreement with findings at necropsy or operation and we used this in our assessment of their relation. In 2 of our patients more than half of one great artery appeared to be posterior to the septum, the inferior margin of the ventricular septal defect being used in this assessment. Subsequent review of the material has led to the suggestion that the superior and lateral margins of the septal defect may have been posterior to both great arteries. Chesler et al. (1971) with echocardiography showed mitral-semilunar valve discontinuity in double outflow right ventricle, whereas Cameron et al. (1976) found continuity in a minority of cases at necropsy. Two of our patients showed mitral-semilunar valve continuity and one of these on echocardiography also failed to satisfy the criterion of Lev et al. (1972). Thus the recognition that the ventricular septum is posterior to both arteries or that there is definite mitral-semilunar valve discontinuity is likely to be diagnostic of double outflow right ventricle, but our findings suggest that failure to recognize either by echocardiography does not exclude the diagnosis.

Correct assessment of the size of the great arteries and relation was made in 10 of the 11 patients examined before angiocardiography. In the 9-year-old child with corrected transposition an incorrect echocardiographic assessment was made while scanning and without viewing the videotape recording. This has stressed the importance of reviewing the recording before giving a diagnostic opinion. Newborn infants were not included until the 5 MHz transducer became available because the 2-5 MHz transducer could not consistently provide scans of diagnostic quality.

In conclusion we believe that the results obtained from this wide angle sector scanner in the study of the size and relation of the great arteries have shown a satisfactory correlation with the angiographic diagnosis. The extensive planar display which it provides makes it particularly suited to the exploratory role which the diagnosis of complex congenital malformations demands, and in this respect more powerful than the linear display to which time-position echocardiography is confined. When it is used in the planning of the cardiac catheterisation and angiography of infants with cyanotic congenital heart disease we are convinced that these investigations can be performed more expeditiously.

We are indebted to Mr. A. Shaw, B.Sc., who directed the development of the scanning equipment, and to Mr. D. J. Wheatly, F.R.C.S., who initiated its clinical application. We acknowledge the help we have received from Dr. E. M. Sweet, Dr. M. McNair, and Miss J. L. McLardy of the Radiology Department of the Royal Hospital for Sick Children, Glasgow.

References


Sector scanner echocardiography in cyanotic congenital heart disease


Requests for reprints to Dr. A. B. Houston, Royal Hospital for Sick Children, Yorkhill, Glasgow G3 8SJ.