The feasibility of complete anatomical correction in the setting of discordant atrioventricular connections

C Alva, E Horowitz, S Y Ho, M L Rigby, R H Anderson

Abstract

Objective—To evaluate the feasibility of complete anatomical correction based on morphological and echocardiographic findings in patients and preserved hearts with discordant atrioventricular connections.

Design—A retrospective study with clinicomorphological correlations to assess potential contraindications for anatomical correction in the setting of discordant atrioventricular connections.

Setting—A tertiary referral centre for congenital heart disease.

Material—25 specimens and 53 patients unified by presence of discordant atrioventricular connections.

Methods—The potential contraindications for anatomical correction were first evaluated on the basis of morphological findings in all 25 specimens with discordant atrioventricular connections collected in the department museum, including study of the major coronary arterial patterns in 20. These contraindications were then sought in a population of 53 patients examined echocardiographically between January 1992 and October 1997.

Results—At least one lesion was discovered that might have contraindicated anatomical correction in 14 of the specimens and in 16 of the patients. The most common lesions that might mitigate against the anatomical approach were severe Ebstein’s malformation or straddling of the tricuspid valve, each when combined with hypoplasia of the morphologically right ventricle. Other potential contraindications were atrioventricular septal defect with common atrioventricular junction, and obstruction of the left ventricular outlet combined with a restrictive ventricular septal defect, although these may be overcome with increasing experience and expertise.

Conclusions—According to the morphological and echocardiographic findings, at least 10 hearts and 37 patients would have produced no anatomical problems for the type of surgical correction in which the morphologically left ventricle is restored its rightful role as the systemic pumping chamber.

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Keywords: atrioventricular discordant connections; congenitally corrected transposition; double switch operation

It is now more than 100 years since Von Rokitansky1 coined the term “corrected transposition.” Since then, the characteristic morphology of this fascinating lesion, with discordant atrioventricular connections as its primary feature, has been well described.2–5 Attention has appropriately been focused on complicating factors, such as obstruction in the ventricular outflow tracts,6–7 straddling of the atrioventricular valves,8 and the abnormal disposition of the atrioventricular conduction tissues.9–10 These studies have established that the potential “physiological” correction of the circulation produced by the presence of discordant connections at both atrioventricular and ventriculo-arterial junctions is usually “uncorrected” by the associated malformations.

In the present era, it would seem a simple matter to correct these malformations surgically, and thus restore the physiologically corrected circulation. Despite much experience, however, the results of simple repair of the intracardiac lesions, when viewed over the longer term, have been less than ideal. The rates of operative mortality, the incidence of tricuspid valvar replacement, and the iatrogenic production of atrioventricular block all remain alarmingly high. Furthermore, failure of the morphologically right ventricle is by no means uncommon in the long term.11–12 Thus, although extended survival into adult life would be expected,13 Connolly and coworkers14 found that, in those undergoing “classical” surgery in Toronto, one quarter had died by the end of their fourth decade. Furthermore, many of the survivors showed progressive atrioventricular block, arrhythmias, and left atrioventricular valve regurgitation. These findings lend support to the long held suspicion that, in time, the morphologically right ventricle will fail when called upon to support the systemic circulation.

Considerations of this kind prompted Ilbawi et al to seek an alternative surgical approach.15 They combined an atrial switch with the Rastelli procedure, thus recruiting the morphologically left ventricle as the systemic ventricle. This approach, producing both physiological and anatomical correction, has now evolved to include a further combination of procedures. In patients with a normal pulmonary valve, the atrial switch is combined with an arterial switch. In this setting, if the morphologically left ventricle is not capable of supporting the systemic circulation immediately, banding of the pulmonary trunk is first performed to train...
the left ventricle. These two combinations of surgical procedure have now been expanded by incorporating surgical repair of various associated malformations.\textsuperscript{16–19}

These experiences, all of which restore the morphologically left ventricle to its systemic role and thus produce anatomical correction, have now encouraged most centres to reassess their populations of patients with discordant atrioventricular connections. The purpose of our study, therefore, was to evaluate the feasibility of such operative procedures in our own patients with discordant atrioventricular connections, assessing echocardiographic findings in the light of morphological observations.

As yet, we do not have sufficient surgical experience to confirm the validity of our prognostications.

**Methods**

We analysed first the morphological features of all the 25 specimens with discordant atrioventricular connections stored in the cardiopathological museum at the Royal Brompton Campus of the National Heart and Lung Institute, Imperial College School of Medicine, London. These had been collected over a period of 25 years. Each specimen was studied to establish any potential morphological contraindications to anatomical correction. We were able to determine the patterns of the major coronary arteries in 20 of these hearts, describing the findings as suggested by McKay et al.\textsuperscript{20} Thus we evaluated the mutual relation of the aortic and pulmonary valvar orifices, the alignment of the facing arterial sinuses, the location of the coronary arterial orifices within the sinuses, and the origin and epicardial course of the coronary arteries.

Those features discovered in the necropsied hearts which might have militated against surgical success were then evaluated in a population of 53 patients with discordant atrioventricular connections examined echocardiographically at the Royal Brompton Hospital between January 1992 and October 1997 (33 males and 20 females, ratio 1.6). The patients had a mean (SD) age of 4.6 (3.5) years. Trans-thoracic echocardiography had been performed in all, and additional transoesophageal studies

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**Table 1** Lesions observed in the necropsied hearts that could have militated against surgical success

<table>
<thead>
<tr>
<th>Contraindications for atrial and arterial switch operations</th>
<th>n</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Severe Ebstein’s malformation with very small trabecular component of the right ventricle</td>
<td>2</td>
<td>8</td>
</tr>
<tr>
<td>Atrioventricular septal defect with common atrioventricular junction</td>
<td>2</td>
<td>8</td>
</tr>
<tr>
<td>Straddling and overriding of the tricuspid valve with hypoplasia of the right ventricle</td>
<td>4</td>
<td>16</td>
</tr>
<tr>
<td>Ebstein’s malformation combined with mitral stenosis</td>
<td>1</td>
<td>4</td>
</tr>
<tr>
<td>Subtotal</td>
<td>9</td>
<td>36</td>
</tr>
</tbody>
</table>

**Contraindications for atrial switch and Rastelli procedure**

<table>
<thead>
<tr>
<th>Contraindications for atrial switch and Rastelli procedure</th>
<th>n</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Severe subpulmonary diaphragmatic fibrous shelf, and restrictive* perimembranous VSD</td>
<td>2</td>
<td>8</td>
</tr>
<tr>
<td>Mixed subvalvar and valvar pulmonary stenosis and restrictive* perimembranous VSD</td>
<td>1</td>
<td>4</td>
</tr>
<tr>
<td>Severe valvar pulmonary stenosis and restrictive* perimembranous VSD</td>
<td>2</td>
<td>8</td>
</tr>
<tr>
<td>Subtotal</td>
<td>5</td>
<td>20</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>14</td>
<td>56</td>
</tr>
</tbody>
</table>

*Septal defect diameter less than the aortic valve.

VSD, ventricular septal defect.
were performed in seven. Echocardiography was performed using Hewlett-Packard Sonos 1500 and 2000 machines (Hewlett-Packard Inc, Andover, Massachusetts, USA). All images were recorded on tape. The examination included the inferential diagnosis of atrial arrangement (situs) based on the relations of the great vessels relative to the spine within the abdomen. In those undergoing transoesophageal studies, it proved possible to determine the morphology of the atrial appendages directly. The subcostal views in long and short axis, along with the four chamber projection, proved best for establishing the segmental connections. In the interest of brevity, when discussing our findings we shall describe the morphologically left ventricle simply as the left ventricle, and the morphologically right ventricle as the right ventricle, irrespective of their locations in space. We catalogued all the associated anomalies, noting also the position of the heart within the thorax. Some patients were studied on more than one occasion.

Results

CARDIOPATHOLOGICAL FINDINGS

The 25 specimens were obtained from patients in whom the heart had been left sided in 18 (72%), and right sided in seven (28%), with the apex pointing to the side of the heart in all. The atrial appendages were of usual arrangement in 23 specimens (92%), and showed mirror imagery in two (8%). In 23 hearts the ventriculo-arterial connections were discordant, with double outlet right ventricle in one and double outlet left ventricle in the other.

We considered that, on basis of the associated lesions, significant impediments to anatomical correction were present in 15 specimens (60%) (table 1).

Severe Ebstein’s malformation was found in combination with a very small apical trabecular component of the right ventricle in two specimens (8%). The ventricular size was only one third of the expected dimensions.

Straddling and overriding of the tricuspid valve (fig 1), again when combined with significant hypoplasia of the right ventricle, was also considered to produce a situation in which anatomical correction would have proven exceedingly difficult. This was found in four hearts (16%).

Mitral stenosis, observed in one specimen (4%), was also considered a contraindication, taking account of the fact that the left ventricle must drive the systemic circulation.

Atrioventricular septal defect with common atrioventricular junction, seen in two hearts (8%) (fig 2), was considered a potential contraindication in the present era because of the added complexity in achieving anatomical correction.

A ventricular septal defect was judged to be restrictive (fig 3) when its diameter was significantly less than the diameter of the aortic valve. This finding, seen in five hearts (20%) would have precluded the construction of an intraventricular tunnel as part of the Rastelli procedure. It is nearly impossible to enlarge such defects in the setting of the discordant atrioventricular connections without damaging the atrioventricular conduction tissues (fig 3), although
some might choose to accept the complication of surgically induced heart block as a lesser evil than the alternatives.

The coronary arterial patterns were studied in 20 hearts (fig 4). Of these, 18 had the usual (solitus) arrangement of the atrial appendages, and two had mirror imagery (inversus). The aorta was leftward and anterior to the pulmonary trunk in all the specimens with the usual atrial arrangement, and rightward and anterior in the specimens with mirror imagery. Abnormal commissural alignment of the valvar leaflets was seen in 11 hearts (55%), all having the zone of apposition between the valvar leaflets supporting the facing aortic sinuses deviated toward the right hand sinus. In 18 hearts with the usual atrial arrangement, the artery arising from the right hand sinus (as viewed from the non-facing aortic sinus) supplied the morphologically right ventricle. The anterior descending and circumflex coronary arteries supplying the morphologically left ventricle arose from the left hand sinus. One heart had a solitary orifice in the right hand sinus (fig 4C). In this case, the anterior descending and circumflex coronary arteries originated as a branch of the artery which supplied the morphologically right ventricle (fig 4A). In one heart, the morphologically right coronary artery continued in the atroventricular groove to end as an anterior diagonal branch, crossing a short circumflex artery (fig 4D). Most of the arterial orifices arose within the aortic sinuses. Four orifices were at the sinutubular junction, and only one orifice was located above the sinutubular junction. None of these patterns was considered to offer any contraindications for anatomical correction, nor were any intramural arterial segments discovered.

On the basis of the abnormalities observed, therefore, we considered that anatomical correction would have proved feasible in 11 specimens (44%) by means of combining an atrial switch with either an arterial switch or the Rastelli procedure. Of the four specimens suitable for combined atrial and arterial switching, three would not have required previous banding of the pulmonary trunk. Banding would have been necessary in one specimen with an intact ventricular septum. The remaining seven specimens, all having large ventricular septal defects associated with severe pulmonary stenosis, were considered suitable for correction using the atrial switch combined with a Rastelli procedure.

CLINICAL FINDINGS

The heart was left sided in 29 patients (55%), right sided in 13 (25%), and centrally located in 11 (21%). The atrial chambers were arranged in usual fashion in 51 patients (96%).
**Table 2** Lesions observed echocardiographically that could have mitigated against surgical success

<table>
<thead>
<tr>
<th>Lesions observed echocardiographically that could have mitigated against surgical success</th>
<th>n</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Militating against the combined atrial and arterial switch operations</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Severe Ebstein’s malformation with very small trabecular component of the right ventricle</td>
<td>2</td>
<td>3.7</td>
</tr>
<tr>
<td>Straddling and overriding of the tricuspid valve with severe hypoplasia of the right ventricle</td>
<td>6</td>
<td>11.3</td>
</tr>
<tr>
<td>Mitral stenosis with hypoplasia of the left ventricle</td>
<td>1</td>
<td>1.8</td>
</tr>
<tr>
<td>Atrioventricular septal defect with common atrioventricular junction</td>
<td>1</td>
<td>1.8</td>
</tr>
<tr>
<td><strong>Subtotal</strong></td>
<td>10</td>
<td>18.9</td>
</tr>
<tr>
<td><strong>Producing problems for the atrial switch combined with the Rastelli procedure</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Valvar pulmonary stenosis and restrictive perimembranous VSD*</td>
<td>3</td>
<td>5.6</td>
</tr>
<tr>
<td>Severe subpulmonary diaphragmatic fibrous shelf and perimembranous VSD*</td>
<td>3</td>
<td>5.6</td>
</tr>
<tr>
<td><strong>Subtotal</strong></td>
<td>6</td>
<td>11.3</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>16</td>
<td>30.2</td>
</tr>
</tbody>
</table>

*Septal defect diameter less than the aortic valve, except in pulmonary atresia.

VSD, ventricular septal defect.

and were mirror imaged in two (4%). The ventriculo-arterial connections were discordant in 47 cases (89%), single outlet from the right ventricle through the aorta with pulmonary atresia in five (9%), and double outlet from the right ventricle in one (2%).

In 16 patients (30%), we discovered at least one of the lesions encountered in our necropsy study which might have produced significant, and potentially insurmountable, problems during any attempted anatomical repair (table 2).

Of the remaining 37 patients (70%), 22 (42%) could readily have been corrected by combining the atrial and arterial switch procedures, and 15 (28%) by combining the atrial switch with the Rastelli procedure. Of the 22 patients considered suitable for the combined atrial and arterial switch operations, seven would have needed preparation of the left ventricle. These patients had no potential for intracardiac shunting, but most had tricuspid regurgitation. The other patients would not have required banding of the pulmonary trunk, since all had ventricular septal defects.

Discussion

After “classical” repair of the associated lesions in patients with discordant atrioventricular connections, which will correct the pattern of the circulations physiologically, the major concern in the long term is the fate of the morphologically right ventricle. Indeed, it is precisely because the morphologically right ventricle must function as the systemic ventricle that so-called “classical” repair does not provide anatomical correction. From neither morphological nor physiological points of view was the right ventricle designed to work as a systemic ventricle. Rather, the morphologically right ventricle has the structure to function best at low pressure. It should be no surprise, therefore, to discover that right ventricular failure occurs after a substantial period of pumping against high pressure. Such systemic right ventricular failure, particularly when associated with tricuspid regurgitation, has been well documented as one of the major deficiencies of the Senning or Mustard operations when used for surgical correction of complete transposition (concordant atrioventricular and discordant ventriculo-arterial connections).21 In this setting, recent experience has shown that, in properly selected cases, the failing right ventricle can be rescued by the arterial switch procedure, restoring it to its role as the subpulmonary ventricle.22

In the setting of congenitally corrected transposition, deterioration of the systemic morphologically right ventricle is seen most often when associated with tricuspid valvar regurgitation. Systemic ventricular dysfunction, however, is common in the overall population of patients with corrected transposition after childhood, reflecting the inability of the right ventricle to support the systemic circulation over a lifetime.23 Such right ventricular dysfunction is increased when there is the need for multiple reoperations for tricuspid valvar replacement, and these procedures themselves carry significant mortality.24 Indeed, heart failure owing to right ventricular dysfunction is the main cause of late death in these patients.11 25 26 Even patients with uncomplicated congenitally corrected transposition, when followed over the long term, develop tricuspid regurgitation, dysfunction of the right ventricle, and complete atrioventricular block. Only a few patients have been reported to survive beyond the fourth decade.25

In this equation, when the few patients living normal lives are contrasted with the much larger proportion destined in time to develop failure of the morphologically right ventricle, it is not unreasonable to support the trend towards anatomical rather than physiological surgical correction, accepting that other options, such as the Fontan procedure,28 are also applicable to many, if not most, of the patients.

As far as we can establish, it was Subirana et al who first performed a double switch operation for surgical repair of corrected transposition,29 although their attempt was unsuccessful. It was Ilbawi and his colleagues who were the first to report success with the combination of an atrial switch and intracardiac rerouting of the outlet of the left ventricle, coupled with placement of a conduit from the right ventricle to the pulmonary trunk.30 Other reports followed rapidly, describing anatomical correction by combining the atrial switch with an arterial switch when the left ventricular outflow tract was unobstructed, or with the Rastelli procedure when there was pulmonary stenosis or atresia. In the latter setting, of course, a large ventricular septal defect is needed to permit construction of an adequate intraventricular tunnel.16-18 22 30 This is because enlargement of a perimembranous ventricular septal defect in the setting of discordant atrioventricular connections is extremely hazardous owing to the abnormal position of the atrioventricular conduction tissues.9

It seems, therefore, that anatomical correction will increasingly become the surgical option of choice for patients with double discordance. In this light, it is important to identify properly those patients in whom the operation can be safely undertaken. In our present study, we considered that while conditions such as a common atrioventricular junction and straddling of the tricuspid valve or severe Ebstein’s malformation, when
combined with significant hypoplasia of the apical component of the right ventricle, might not necessarily be absolute contraindications for anatomical correction, they would have required technically hazardous and extremely long surgical procedures, hampering the chances of successful repair. We recognise that, with continuing improvements, the skilful surgeon will now almost certainly be able to overcome such purely anatomical problems. It is imperative in the early stages of surgical experience, nonetheless, to select patients with adequate anatomy so as to ensure appropriate survival. With increasing surgical experience, therefore, we recognise that our predictions may prove unduly pessimistic.

Fortunately, as far we could judge, the anatomy of the coronary arteries would not have precluded an arterial switch in any of our necropsied hearts, despite the discovery of abnormal commissural alignment in more than half. An intramural origin of the coronary arteries is now the major complicating factor recognised in repair of complete transposition, although it can be circumvented in the best hands. This abnormality was not observed among our specimens, but it must surely be anticipated. Other patterns, such as origin from the non-left sinus, if it exists, would probably produce greater problems.

We acknowledge that a major deficiency of our investigation is its retrospective nature, along with the fact that, as yet, the number of patients undergoing anatomical correction in our own centre is too small to permit validation of our predictions. It is now possible, nonetheless, to recognise all possible contraindications by cross sectional echocardiography, particularly when combined with colour Doppler imaging and transesophageal imaging. We also anticipate that, with experience gained from patients with complete transposition, it will prove possible to identify any abnormal origins of the coronary arteries, including their intramural location, should this be present. As we have already discussed, it is also likely to prove that, with increasing experience, many of the problems we presently consider as contraindications can be neutralised. Such clinical experience will also enable us to document accurately features such as hypoplasia of the right ventricle and the size of restrictive ventricular septal defects which, in our study, have been assessed subjectively rather than objectively. In the final analysis, only experience gained in the clinical arena will determine the proportion of patients with discordant atrioventricular connections who will become realistic candidates for anatomical repair, restoring the morphologically left ventricle to its rightful role as the systemic pumping chamber.

We are grateful to Karen McCarthy, Zarlasht Najem, and Vi-Hue Tran for their excellent technical assistance, and to Mrs Christine Anderson for expert secretarial help. We are also indebted to Professor A N Redington and Dr E A Shinebourne, who were involved in the clinical care of many of the patients studied, and Dr Leon Gerlis, who collected many of the specimens. RHA and SVH are supported by the British Heart Foundation and the Joseph Levy Foundation. During the course of this investigation CA was a visiting research fellow from Hospit al de Cardiologia Centro Medico Nacional Siglo XXI, Mexico City, Mexico.


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