Angiocardiographic quantitative evaluation of double-outlet right ventricle: special reference to the morphology of the common outflow tract.

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Abstract

Geometrical measurements of angiocardiograms of the common outflow tract (COT) of 13 patients were made to determine in which cases internal conduit repair was feasible, and under which conditions a patch enlargement of the COT was indicated. In the pulmonary stenosis (PS) group, the area of the narrowest cross-section of the COT was significantly smaller than that in the pulmonary hypertension (PH) group (p less than 0.025). In the PS group, the area was rarely sufficient to be shared by systemic and pulmonary circulation. Therefore, stenosis in the outflow tract to the pulmonary artery will occur if the intraventricular tunnel technique is applied, without patch enlargement of this portion, to patients with PS. On the contrary, the cross-sectional areas of the COT and pulmonary arteries were significantly larger in the PH group than in the PS group. Accordingly, the intracardiac conduit operation may be possible in such patients without a patch enlargement, even in young patients if other intracardiac conditions allow. Preoperative angiocardioangiographic evaluation of the COT is helpful in preoperatively selecting the proper operative procedure for this anomaly.

KEYWORDS: double-outlet right ventricle, angiocardiography, common outflow tract, intraventricular tunnel technique, patch enlargement

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Key words: double-outlet right ventricle, angiocardiography, common outflow tract, intraventricular tunnel technique, patch enlargement

Witham coined the phrase double-outlet right ventricle (DORV) in 1957 (1). Since then, much research has been done to define the anomaly as a distinct entity (2). Lev (3) and Kawashima (4) indicated that the various conotruncal malformations formed a continuous spectrum, rather than being distinct and separate entities. Therefore, preoperative evaluation of each case of DORV is necessary to select the best operative procedure. Many procedures have been developed since the successful repair by Kirklin et al. (5). Intraventricular rerouting through a baffle is now the most common operative method, and results with this procedure are particularly satisfactory in patients with subaortic ventricular septal defect (VSD) (6). It has not been clearly demonstrated, however, under what anatomical conditions intraventricular tunnel repair is possible. We undertook this study to clarify what geometrical dimensions allowed the intraventricular rerouting technique. We also studied how to avoid postoperative

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stenosis in the pulmonary part of the right ventricular outflow tract.

Methods

The definition of DORV used in this study was that of Lev (3) and Pacifico et al.(7). The classification referring to pulmonary stenosis (PS) and/or pulmonary hypertension (PH) was based on the relationship of VSD to the great arteries.

Patients. During the past eleven years, we experienced a total of thirteen patients with DORV (Table 1). The mean age at the time of the angiocardiographic study was four years and ten months, ranging from 0.1 to 21.0 years. Of the patients with PS, three had undergone one or two palliative operations. None of the patients with PH received pulmonary artery banding. Corrective surgery was performed for five of the patients with PS, and three of those with PH.

Surgical Methods. The basic surgical procedure was the creation of an internal conduit (IC) from VSD to the aortic ostium (Table 1). PS was concomitantly relieved by infundibular cardiomyectomy in four cases and by pulmonary valvulotomy in three cases. Enlargement with an outflow patch was used in only one case (Case 6). The Damus method (9) was applied to one case (Case 12), and a modification of the Fontan operation (10) was applied to another case (Case 11).

Angiocardiographic evaluation method. For the evaluation of the COT, anteroposterior and lateral biplane projections of biventriculograms were obtained preoperatively. The external diameter of the catheter used was measured at the diaphragmatic level to calculate the magnification rate. The diameter of the pulmonary arterial outflow tract (DPAOT), the diameter of the pulmonary annulus, and that of the aortic annulus (DAoa) were measured in accordance with the report by Blackstone et al.(11). The narrowest point of the main pulmonary artery was measured at the

<p>| Table 1 Characteristics of patients, operative procedures, and results |
|---|---|---|---|---|---|---|---|</p>
<table>
<thead>
<tr>
<th>Cases</th>
<th>Age (years)</th>
<th>Sex</th>
<th>Type</th>
<th>VSD</th>
<th>PAP</th>
<th>Previous op.</th>
<th>Operative procedure</th>
<th>Operative results</th>
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<tr>
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<td>SA</td>
<td>PH</td>
<td>-</td>
<td>IC</td>
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<tr>
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<td>1.8</td>
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<td>SDD</td>
<td>SA</td>
<td>PS</td>
<td>-</td>
<td>IC, IM, PVT</td>
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<td>-</td>
<td>M</td>
<td>SLL</td>
<td>NC</td>
<td>PS</td>
<td>BT, PVT</td>
<td>-</td>
</tr>
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<td>-</td>
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<td>NDD</td>
<td>NC</td>
<td>PS</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
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<td>-</td>
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<td>DC</td>
<td>PS</td>
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<td>-</td>
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<td>9.5</td>
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<td>PS</td>
<td>BT</td>
<td>IC, IM, OTP, PVT</td>
</tr>
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<td>0.6</td>
<td>-</td>
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<td>NC</td>
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<td>-</td>
<td>-</td>
</tr>
<tr>
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<td>DC</td>
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<td>IC</td>
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<td>-</td>
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<tr>
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<td>PS</td>
<td>BT</td>
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<td>SDD</td>
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*Cases 3, 4, 5, 7, and 13 did not undergo corrective surgery.*

*Van Praagh's terminology (8).*

*Relationship of VSD to the great arteries.*

*Pulmonary arterial pressure (PAP) is expressed as PS or PH.*

Abbreviations: angio, angiocardiography; op., operation; VSD, ventricular septal defect; F, female; SA, subaortic; PH, pulmonary hypertension; IC, internal conduit; M, male; PS, pulmonary stenosis; IM, infundibular myectomy; PVT, pulmonary valvulotomy; NC, non-committed; BT, Blalock-Taussig shunt; DC, doubly committed; OTP, outflow tract patch; SP, subpulmonary.*

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systolic phase, as was the DPAOT. These diameters were used to calculated the cross-sectional area of the pulmonary arterial outflow tract (PAOTa) and that of the aortic annulus (AoAa) under the assumption that the transsectional shape of these portions were truely circular. The anteroposterior projection was used whenever possible, but occasionally the lateral projection was also used if the details were clearly shown on it. From diameters obtained, the cross-sectional areas of the right pulmonary artery (RPAa), the left pulmonary artery, and the main pulmonary artery (PAa) were calculated. These values were compared with the respective areas of the normal heart obtained from equations below (12-14). The mean value (PAa) of the cross-sectional areas of the right and left pulmonary arteries was also calculated, and was compared with the RPAa of the normal heart.

The cross-sectional area of the right pulmonary artery of the normal heart (N-RPAA) was calculated by:

$$N\text{-}RPAA = -0.3546 + 2.7989 \cdot (BSA),$$

where BSA is the body surface area.

The cross-sectional area of the main pulmonary artery of the normal heart (N-PAA) was calculated by:

$$N\text{-}PAA = -0.536 + 4.035 \cdot (BSA).$$

The diameter of the pulmonary arterial outflow tract of the normal heart (N-DPAOT) was calculated by:

$$N\text{-}DPAOT = 15.29 + 11.42 \cdot \log_{10}(BSA).$$

The diameter of the aortic annulus of the normal heart (N-DAA) was calculated by:

$$N\text{-}DAA = 0.967 + 3.51 \cdot \log_{10}(BSA).$$

Fig. 1 shows the method of angiocardiographic evaluation of the COT. The line A is drawn from the right margin of the aortic annulus to the left margin of the pulmonary annulus. The aortic annular diameter (DAa) is indicated as 2R, and the pulmonary annular diameter as 2r. The distance between the two annuli is shown as a; this does not represent the conal septal thickness. The cross-sectional area of the COT (S) at the
level of the line A was geometrically calculated. The area of the residual orifice to the pulmonary artery is indicated by X (S minus πR^4, the area covered by oblique lines in the upper right figure) under the supposition that the internal tunnel has the same cross-sectional area as the AoAa, that is πR^2.

The line B indicates the narrowest part of the COT, where the smallest diameter (a) of the COT is found at the end-systole in the anteroposterior projections. The line B in the lateral projection is at the same level as the line B, and is where the sagittal diameter of the COT at the end-systole (β) is found. The cross-sectional area of the COT at the level of the line B, indicated as COTa, was also calculated geometrically, under the assumption that the transsectional shape of this portion was elliptical with the longest diameter of a and the shortest one of β, as shown in the lower right figure. The cross-sectional area of the internal tunnel, which is created so as to have the same cross-sectional area as the AoAa (πR^2), is labeled IC and covered with oblique lines slanting down to the left. The residual area of the COTa (labeled Y and covered with oblique lines slanting down to the right) after tunnel formation is calculated by the COTa minus AoAa. S and COTa were standardized using the body surface area of each case, and were then expressed as SI and COTaSI in cm^2/m^2.

Variables were evaluated by dividing the subjects into two groups, the PS and PH groups. The data were comparatively analyzed with Student’s unpaired t test, and a p < 0.05 was considered to be statistically significant. All data are presented as means ± standard deviation (SD).

**Results**

Some values in Case 1 were not available because of poor angiocardiograms. The distal pulmonary arterial size was evaluated by the ratios RPAa to N-RPAa (RPAa/N-RPAa), and PAa to the N-RPAa (PAa/N-RPAa). RPAa/N-RPAa in the PH group was 1.87 ± 0.31, which was significantly larger than 0.88 ± 0.50 of the PS group (p < 0.01). PAa/N-RPAa in the PH group was also statistically larger than that of the PS group (1.71 ± 0.29 vs. 0.85 ± 0.38, p < 0.05; Fig. 2). The size of the proximal part of the pulmonary artery was evaluated by the ratios PAa to N-PAa (PAa/N-PAa), and DPAa to N-DPAa (DPAa/N-DPAa) (Fig. 3). PAa/N-PAa was significantly smaller in the PS group than in the PH group (1.00 ± 0.62 vs. 3.14 ± 1.01, p < 0.01), and DPAa/N-DPAa was markedly larger in the PH group. The aortic size was evaluated by the ratio DAAo to N-DAAo (DAAo/N-DAAo), which was 1.20 ± 0.08 in the PH group and 1.44 ± 0.26 in the PS group, there being no sig-

![Fig. 2](http://escholarship.lib.okayama-u.ac.jp/amo/vol43/iss1/8)
significant difference (Fig. 4). The ratio $\beta$ to D AoA was calculated to be $1.45 \pm 0.34$ in the PH group and $0.80 \pm 0.10$ in the PS group; the difference was significant ($p < 0.005$).

The cross-sectional area of the COT at the level of the line A, i.e., SI, was $9.42 \pm 0.29 \, \text{cm}^2/\text{m}^2$ in the PH group and $7.04 \, \text{cm}^2/\text{m}^2$ in the PS group; the difference was not significant. The cross-sectional area at the level of the line B, i.e., COTaL, was $8.23 \pm 1.96 \, \text{cm}^2/\text{m}^2$ in the PH group and $4.16 \pm 1.83 \, \text{cm}^2/\text{m}^2$ in the PS group. This difference was statistically significant ($p < 0.025$) (Fig. 5).

The ratio X to N PAOTa (X/N PAOTa) was $3.37 \pm 0.38$ and $1.31 \pm 0.47$ in the PH and PS groups, respectively. This difference was significant ($p < 0.005$) (Fig. 6). However, the residual area at this level (line A) was sufficiently large in most

![Fig. 4](image)

**Fig. 4** The aortic annular diameter and comparison with the sagittal diameter of the common outflow tract ($\beta$). Symbols are the same as in Fig. 2. Abbreviations: D AoA, diameter of the aortic annulus; N D AoA, D AoA of the normal heart (14); $\beta$, sagittal diameter of the common outflow tract (Fig. 1); NS, not significant. ****, $p < 0.005$.

![Fig. 5](image)

**Fig. 5** Evaluation of the cross-sectional area of the right ventricular outflow tract. Symbols are the same as in Fig. 2. Abbreviations: SI, S standardized according to the body surface area (See Fig. 1); COTaL, standardized value of COTa (See Fig. 1); NS, not significant. **, $p < 0.025$. 

Fig. 3 Evaluation of the size of the proximal part of pulmonary arteries. Symbols are the same as in Fig. 2. Abbreviations: PAa, cross-sectional area of the main pulmonary artery; N PAa, the PAa of the normal heart (13); D PAOT, diameter of the pulmonary arterial outflow tract; N D PAOT, D PAOT of the normal heart (14). ***, $p < 0.01$; ****, $p < 0.005$. 

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cases of the PS group not to cause an outflow tract stenosis to the pulmonary artery, since X/N-PAOTa was larger than 1.0. On the contrary, Y/N-PAOTa in the PH group (2.62±0.80) was significantly different from that in the PS group (−0.10±0.70) (p < 0.005). It was revealed that the residual area to the pulmonary circulation at this level (line B) was very restricted in the PS group, since Y/N-PAOTa was much smaller than 1.0 (Fig. 6).

Discussion

This study was undertaken to clarify the preoperative criteria for deciding whether a complete or palliative operation should be selected, and to determine under what geometrical circumstances a patch enlargement of the COT, pulmonary annulus, or both were indicated. The ultimate purpose of this study was to clarify what intracardiac, geometrical dimensions allowed or prohibited intraventricular tunnel repair.

It is generally considered that the fundamental policy for surgical managements of PS associated with various cardiac anomalies is based on that for PS in the tetralogy of Fallot. From our experience with the tetralogy of Fallot, we have considered that a palliative operation is indicated when RPAa/N-RPAa is less than 0.36, or when PAA/N-RPAa is less than 0.32 (15, 16). Accordingly, with only the size of distal pulmonary arteries as a criterion, complete repair of the DORV was contraindicated in few cases of this series. The size of the main pulmonary artery and its orifice were sufficient in most of the cases. However, PAA/N-PAAa was 0.12 and 0.19 in Cases 4 and 11, and DPAOT/N-DPAOT was 0.36 and 0.64 in Cases 3 and 11, respectively. In Cases 3 and 4, no surgery was done, and in Case 11 a modification of the Fontan operation was applied. As in this way, when the case does not meet with the criteria indicating either corrective surgery on the tetralogy of Fallot or the Fontan procedure (17), a palliative operation should be preferred to a corrective one. In the PH group, on the contrary, pulmonary annulus and the distal and main pulmonary arteries were dilated, and there was no problem concerning the size of pulmonary arteries.

It is necessary to determine where a stenosis in the outflow tract to the pulmonary artery could develop in cases where tunnel repair is applied. The sagittal diameter of the COT (β) was measured and compared with the DAAoA to give the ratio β/DAoA. The result of the analysis with this ratio indicates that the cross-sectional shape of the tunnel has to be elliptic in order to give an internal tunnel the same cross-sectional

Fig. 6 Evaluation of residual cross-sectional area of the right ventricular outflow tract to the pulmonary circulation after the internal conduit operation. Symbols are the same as in Fig. 1. Abbreviations: N-PAOTa, transsectional area of the pulmonary arterial outflow tract of the normal heart (14); OTP, outflow tract patch. ****, p < 0.005.
area as AoAa. That is, the tunnel will laterally expand and will be anteroposteriorly compressed as in Fig. 1, even if the patch roof of the tunnel fully expands anteriorly to the inner surface of the free wall of the outflow tract, since \( \beta \) is always smaller than the aortic annular diameter, i.e., \( \beta / \text{DAOA} < 1.0 \). Accordingly, the conduit wall must protrude into the pulmonary part of the COT. In the present series, the COTa was statistically smaller in the PS group than in the PH group, so that Y was significantly smaller than the N-PAOTA in the PS group.

Although X/N-PAOTA was statistically smaller in the PS group than in the PH group, all patients in the former group were observed to have sufficient cross-sectional area of the remaining outflow tract to the pulmonary circulation after the internal tunnel operation, except Case 3 in which no operation has been performed. This observation indicates that the pulmonary outflow tract stenosis would not develop at the level of the line A after tunnel repair. However, the residual cross-sectional area at the level of the line B in the COT, represented by Y/N-PAOTA, was found to be significantly smaller in the PS group, and consequently postoperative stenosis in the pulmonary outflow tract could develop here unless this portion were enlarged using a patch or some other procedure. Among the patients with PS upon whom we operated, Y/N-PAOTA was the lowest in Case 6, at −1.14, and the highest in Case 2, at 0.78. By our own criteria for the tetralogy of Fallot (15, 16), all the patients in the PS group should have received a patch enlargement of the outflow tract, although the procedure was used only in Case 6. However, in spite of the patch enlargement, the patient died from low cardiac output. It was thought that the low cardiac output was caused by the residual stenosis of newly created outflow tract to the pulmonary artery (Y/N-PAOTA of −1.14 in this case) being inadequately narrow even after the patch enlargement. Other operative procedures such as extracardiac conduit operation or a modification of the Fontan operation should have been performed in this case. Stewart et al. (2, 19) also recommended a patch enlargement or an external conduit operation even when PS was absent because simple infundibular dissection and pulmonary valvulotomy could not prevent postoperative stenosis of the pulmonary outflow tract after tunnel repair. Angiographic evaluation of the outflow tract as described in the present study should help surgeons to decide preoperatively whether to patch or not to patch, and whether the patch enlargement technique is feasible or not.

In our cases of PH, the SI, COTa, X/N-PAOTA, and Y/N-PAOTA were all large, and none of the unfavorable geometries was present, indicating that there was no risk of postoperative stenosis in the pulmonary outflow tract developing after the intraventricular tunnel operation, although there remained a problem concerning the location of VSD (6). Nevertheless, the operative results in the PH group of our series were poor, with high pulmonary arterial pressure early after corrective surgery. Pulmonary artery banding to prevent the development of pulmonary vascular obstructive disease was not performed in these cases, and that might account for the unsatisfactory results in the PH group. There also may be some question as to whether intraventricular tunnel repair is feasible in very young patients. However, our observation was that the COT in the PH group had sufficient area to be shared by both the systemic and pulmonary circulation even during the early time of life, so that an intracardiac tunnel operation should be possible.

It was concluded that preoperative evalu-
ation of the morphology of the COT as described herein might help physicians to select the proper operative procedure for DORV. It was recommended that the COT should be enlarged using a patch in most cases of PS when intraventricular tunnel repair is to be applied.

References


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