Combined interventional and surgical treatment in pediatric patients with tetralogy of Fallot

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\section*{ARTICLE INFORMATION}
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\section*{Competing interests}
The authors declare no competing interests

\section*{Acronyms}
VSD: ventricular septal defect

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\section*{ABSTRACT}
In tetralogy of Fallot with hypoplastic pulmonary arteries, an alternative to avoid palliative surgeries is percutaneous pulmonary valvuloplasty using a balloon catheter. When there are major arteriovenous collaterals producing volume overload of the left chambers, their previous occlusion improves surgical outcomes. Two patients with aorto-pulmonary collaterals, closed in the laboratory of hemodynamics 24 hours before surgery, and two children who underwent pulmonary valve dilation and then surgery, seven and nine months afterwards, respectively, are reported. Mean follow-up was five years without complications. Interventional catheterization techniques before surgery for tetralogy of Fallot are feasible and can help reduce the number of palliations and improve the results of surgical correction of the disease in selected cases.

Key words: Cardiac catheterization, Cardiovascular surgery, Tetralogy of Fallot

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\section*{Tratamiento combinado intervencionista y quirúrgico en pacientes pediátricos con tetralogía de Fallot}

RESUMEN
En la tetralogía de Fallot con ramas pulmonares hipoplásicas, una alternativa para evitar cirugías paliativas es la valvuloplastia percutánea pulmonar con catéter de globo. Cuando existen colaterales arteriovenosas mayores que producen sobrecarga de volumen de las cavidades izquierdas, ocluirlas previamente mejora los resultados quirúrgicos. Se presentan dos pacientes con colaterales aorto-pulmonares, cerradas en el laboratorio de hemodinámica 24 horas antes de la cirugía, y dos niños a quienes se les dilató la válvula pulmonar y luego recibieron cirugía, siete y nueve meses después, respectivamente. El seguimiento medio ha sido de cinco años sin complicaciones. Las técnicas de cateterismo intervencionista previas a la cirugía de la tetralogía de Fallot, son factibles y pueden contribuir a disminuir el número de paliaciones y a mejorar los resultados de la corrección quirúrgica de la enfermedad en casos seleccionados.

Palabras clave: Cateterismo cardíaco, Cirugía cardiovascular, Tetralogía de Fallot
INTRODUCTION

Diameter of pulmonary arteries is a determining factor for the correction of tetralogy of Fallot. Sometimes palliative surgeries are needed to stimulate the growth of the central pulmonary and intrapulmonary arteries. However, such interventions are not without high surgical risks\(^1\)-\(^3\).

A less invasive alternative is percutaneous transluminal pulmonary valvuloplasty with balloon catheter, in those who have not developed a significant infundibular stenosis\(^2\)-\(^4\). This technique was described in 1986 by McCredie\(^5\) and allows a better distributed antegrade blood flow to both pulmonary artery branches, resulting in better anatomical conditions of patients at the time of surgical correction. Stents in the right ventricle outflow tract have also been used\(^1\)-\(^6\).

In other cases there are major arteriovenous collaterals producing volume overload of the left cardiac chambers and pulmonary hypertension, seriously complicating intraoperative and postoperative recovery. Their surgical closure is difficult, so that occluding them previously, via catheterization, could improve the results of corrective surgery\(^7\)-\(^10\).

The aim of this case report is to describe the combined use of interventional catheterization techniques and corrective surgery in four patients with tetralogy of Fallot.

CASE REPORT

Under intravenous anesthesia with ketamine and midazolam a central venous access via the femoral artery was performed in the catheterization laboratory and in cases with multiple aorto-pulmonary collaterals, the femoral artery was also inserted. Anatomy was precisely defined by angiography.

Aorto-pulmonary collaterals of significant diameter were closed with controlled release devices, 24 hours before surgery, after verifying the dual irrigation of lung segments by true pulmonary arteries.

The pulmonary valve dilation was performed with balloon catheters that exceeded 1.4 times the diameter of the pulmonary annulus (Figure).

Surgical and interventional details are reflected in tables 1 and 2. There were no hypoxia crises or serious arrhythmias during interventional procedures. In patients who received pulmonary valve dilation an improvement of the overall condition was achieved as well as improvement of arterial oxygen saturation and of the pulmonary arteries size at the time of surgical correction.

In patients to whom collaterals were closed, cardiopulmonary bypass was conducted with adequate pulmonary venous return and a good visualization of the surgical field, without having to lower the temperature and perfusion flow. The mean follow-up of cases has been of five years without complications. In all patients informed consent signed by the parents was obtained and procedures are part of protocols approved by the Scientific Council and the Ethics Committee of the research.

A. Anteroposterior view of angiocardiogram with contrast injection into the right ventricle in a newborn of 11 days and 1.9 kg in weight, where the tiny branches of the pulmonary artery (2 mm) are observed. B. Side view which shows the balloon catheter in the right ventricular outflow and the notch at the pulmonary annulus level is indicated.
Table 1. Data of interventional treatment.

<table>
<thead>
<tr>
<th>Age</th>
<th>Weight</th>
<th>Anatomy</th>
<th>Indication</th>
<th>Technique</th>
<th>Evolution</th>
</tr>
</thead>
<tbody>
<tr>
<td>11 days</td>
<td>1.9 kg</td>
<td>Pulmonary branches of 2 mm</td>
<td>Arterial oxygen saturation: 40 %</td>
<td>Pulmonary valve dilation</td>
<td>No complications. Arterial oxygen saturation: 85 %.</td>
</tr>
<tr>
<td>9 months</td>
<td>5 kg</td>
<td>Valve stenosis and pulmonary branches of 3 mm</td>
<td>Arterial oxygen saturation: 55 %</td>
<td>Pulmonary valve dilation</td>
<td>No complications. Arterial oxygen saturation: 88 %.</td>
</tr>
<tr>
<td>2 years</td>
<td>14 kg</td>
<td>Thick direct collateral arteries of the aorta</td>
<td>Volume overload of left cardiac cavities</td>
<td>Closure of three collaterals of the thoracic aorta and one infradiafragmatic</td>
<td>No complications. No drop in arterial oxygen saturation.</td>
</tr>
<tr>
<td>10 years</td>
<td>32 kg</td>
<td>Thick direct collateral arteries of the aorta</td>
<td>Volume overload of left cardiac cavities</td>
<td>Closure of two collaterals of the thoracic aorta</td>
<td>No complications. No drop in arterial oxygen saturation.</td>
</tr>
</tbody>
</table>

Table 2. Data of surgical treatment.

<table>
<thead>
<tr>
<th>Edad</th>
<th>Peso</th>
<th>Características anatómicas</th>
<th>Técnica quirúrgica</th>
<th>Evolución y seguimiento</th>
</tr>
</thead>
<tbody>
<tr>
<td>9 months</td>
<td>8 kg</td>
<td>Significant improvement of pulmonary branches. Right 5 mm and left 5.7 mm</td>
<td>Correction. No pulmonary annulus section.</td>
<td>No complications at 5 years</td>
</tr>
<tr>
<td>16 months</td>
<td>10 kg</td>
<td>Mixed infundibular and valvular stenosis. Appropriate branches</td>
<td>Correction with a section of the pulmonary annulus and patch enlargement with monocuspid valve of autologous pericardium</td>
<td>No complications at 3 years</td>
</tr>
<tr>
<td>2 years</td>
<td>14 kg</td>
<td>Mixed supravalvular, infundibular and valvular stenosis.</td>
<td>Correction with a section of the pulmonary annulus and patch enlargement with monocuspid valve of autologous pericardium</td>
<td>No complications at 7 years</td>
</tr>
<tr>
<td>10 years</td>
<td>32 kg</td>
<td>Mixed supravalvular, infundibular and valvular stenosis.</td>
<td>Correction with a section of the pulmonary annulus and patch enlargement with monocuspid valve of autologous pericardium</td>
<td>No complications at 5 years</td>
</tr>
</tbody>
</table>

COMMENT
In tetralogy of Fallot, although the correction at a time is preferred, sometimes the anatomical characteristics make us choose palliative measures. Systemic pulmonary Blalock-Taussig shunt is the usual measure in such cases. However, its morbidity is significant. The rate of mortality in preterm or low weight at birth is 10 to 18%2,3.

The opening of the outflow tract of the right ventricle, without closing the ventricular septal defect
(VSD) is another option that increases the effective transpulmonary antegrade flow and may promote the growth of the pulmonary arteries. However, this must be done under cardiopulmonary bypass and is very difficult to predict the magnitude of the flow through the VSD, so congestive heart failure, prolonged mechanical ventilation and long term pulmonary hypertension occur frequently.2

Percutaneous transluminal pulmonary valvuloplasty with balloon catheter is an alternative in which valvular obstruction predominates and hypoplasia at the pulmonary arterial branches is also present2-5. The accepted indications are: arterial oxygen saturation below 80 %, hypoxia crisis and lung perfusion dependent of ductus arteriosus2,5.

In these circumstances, an improvement in the growth of the main pulmonary artery, in the size of its branches and in oxygen arterial saturation has been reported2,3,5. For some, although there is an immediate increase in the annulus, its growth remains abnormal and a transannular patch may be required at the surgical moment4.

This method of treatment that could palliate about half of symptomatic newborns and young infants with tetralogy of Fallot, is less effective in those who have had recurrent hypoxia crisis. The growth achieved in the pulmonary arteries is comparable to that of the systemic pulmonary shunt2,3.

The complications reported are mainly in older children in whom the infundibular component is predominant and thus the incidence of serious arrhythmias, or in newborns with less than 2,500 grams, in whom crossing the valve is more difficult3.

Benefits with the use of stents in the right ventricular outflow are identified when compared with systemic pulmonary shunts, as they prevent complications and technical difficulties of placing them in so tiny branches. Potential limitations of these interventional procedures include obstruction due to endothelial proliferation, device fracture and cardiac perforations1,4,6.

Surgical ligation or section of undesirable vascular connections between the aorta and the pulmonary vasculature, which is established in some patients, can considerably increase surgical time, extracorporeal circulation, and besides being technically difficult due to its position, can predispose to profuse bleeding. Pulmonary venous return during cardiopulmonary bypass can be of great magnitude, and the perfusion flow is diverted from such important organs such as the brain and kidneys7,8.

The usefulness and effectiveness of closing them before surgery has currently been demonstrated, with a level of evidence B, as long as there is native pulmonary circulation in the affected segments9. It was recently proposed to carry it out as hybrid procedures during surgery10. Valvular balloon dilation and the closing of major collaterals in a single patient was also reported4.

Interventional catheterization techniques before surgery for tetralogy of Fallot are feasible and can help reduce the number of palliations and improve the results of surgical correction of the disease in selected cases.

REFERENCES
