Case Report: Echocardiography in Fallot Tetralogy

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Summary
The tetralogy of fallot is the most frequent cyanotic congenital heart disease. It presents a clinical case of a pediatric patient who was monitored with transesophageal echocardiography and shows the images and changes in handling due to the echocardiographic monitoring.

Keywords: Tetralogy of fallot; Anesthesia; Echocardiography

We present a five years old boy and six kg of weight, with diagnosis of tetralogy of fallot. He was born from term and he doesn’t present any other congenital anomalies. He is under propanolol treatment due to hypoxia and cyanosis crisis. Having reaches an adequate weight, it is decided to have surgery to repair his cardiac pathology. At his arrival in the operating room, non invasive monitoring is installed, then anesthesia inhalated induction with Sevoflurane 8% plus oxygen 100%. 2 peripheral venous catheters 22g were placed. 3mg of Rocuronium and 40 micrograms of fentanyl were given. It is intubated with an orotracheal tube N° 4 without ballon. Anesthesia is maintained with sevoflurane 1.5%, remifentanil 0.5mcg/kg/min and relaxing (rocuronium) as needed. It is placed right femoral artery cathether for invasive blood pressure monitoring and left anterior jugular venous cathether (4Fr-2 lumens) with ultrasound guide. The pediatric multiplanar Philips ETE probe is placed and the images are obtained from of the 15 cm of the dental arch. The diagnosis of Tetralogy of Fallot is confirmed: a 6mm ventricular septal defect (VSD) is observed, aortic valve overriding the ventricular septum (Figure 1), narrowing of the right ventricular (RV) outflow tract (RVOT) (Figure 2) with a gradient of 67mmHg (Figure 3) and a left ventricular ejection fraction (LVEF) of 65%. In additin, the bad function of right ventricle with a TAPSE (tricuspid annular plane systolic excursion) below the normal values (>14mm) (Figure 4).

The patient is admitted to extracorporeal circulation, guiding fluid therapy under echocardiography and it is not necessary to add vasoactive drugs. It is performed a closure of the VSD by means of a right auriculotomy verifying a 29mmHg gradient, indicating...
the presence of a small residual VSD (Figure 5). At the exit of the cardiopulmonary bypass machine (CPB), adrenaline 0.02 mcg/kg min⁻¹ and milrinone 0.5 mcg/kg/min are used, in a planned way due to the images of the bad function of the RV displayed on echocardiography. The RVOT infundibulum was also taken through the pulmonary artery, being considered the procedure as successful since the gradient in the TSVD decreased to 11 mmHg (Figure 6). Because the RV function reflected by TAPSE did not have a clear improvement in the Post CPB echocardiography it was decided to add milrinone (Figure 7). Finally, the success of the surgery is verified with an adequate ejection fraction (EF) (Figure 8).

Figure 2: Narrowing of the right ventricular outflow tract (RVOT).

Figure 3: It is shown a RVOT gradient of 67 mmHg.

Figure 4: TAPSE: 7mm.

Figure 5: VSD closure: a 29 mmHg gradient is observed, indicating that a small residual CIV remains.

Figure 6: RVOT GRADIENT: 11 mmHg.

Figure 7: TAPSE: 6mm.

Figure 8: LVEF: 56%.

Discussion

The most frequent of congenital cyanotic heart disease is the tetralogy of Fallot that means to be 10% of congenital heart
malformations [1,2]. The characteristics of this pathology are due to a partial anterior septal movement pulmonary aortum which causes:

Pulmonary stenosis: which can be subvalvular - valvular - supravalvular.

VSD: subaortic

Aortic valve overriding ventricular septum: it is because septum IV is deflected forward and right. This causes poor alignment with the anterior wall of the aorta that is connected with both ventricles.

RV hypertrophy: by pulmonary stenosis.

Depending on the degree of displacement of the aortic-pulmonary septum, the TOF can be [3,4]:

a) Total: ductus dependent.

b) Minimal: the output of the RV is slightly narrow with small VSD and minimal aortic riding: pink TOF.

c) What marks the degree of severity of TOF is the size of the pulmonary ring and valve and the degree of obstruction of the RVOT [5].

d) Small ring: transannular patch is needed in surgery which would lead to future reoperations.

e) Normal ring: surgery respects the pulmonary valve.

The most favorable for surgical correction and best prognosis is TOF with normal tree lung.

Special tof subgroups:

a) Small pulmonary branches: in normal RN the pulmonary branches have a diameter 3.5-4mm. A size <3mm is problematic for surgery [6,7].

b) Disconnection of the branches: one of the branches is born from the ascending aorta.

c) Absence of a pulmonary branch: it is supplied by collateral branches.

d) Pulmonary valve agenesis: part of the valvar tissue is missing due to functionally non-existent. Pulmonary insufficiency is severe and with each beat it expels more blood which dilates the pulmonary artery. It is associated with tracheomalacia or broncomalacia [8].

e) Coronary abnormalities: anterior descending artery birth from right coronary artery, crossing the infundibulum of the RV which makes more difficult the surgical correction of pulmonary stenosis [9].

TOF physiopathology

The pathophysiology will depend on the degree of obstruction of the RVOT and the size of the VSD and of the Systemic vascular resistance (SVR) and Pulmonary vascular resistance (PVR). There is an obstruction to the flow of blood to the lungs and due to the VSD produces a shunt of desaturated blood from right to left: D-I shunt causing desaturation and cyanosis. The degree of arterial desaturation depends on the amount of desaturated blood derived by the shunt present in the pulmonary circulation. The more severe the obstruction to the RVOT or the more the D-I shunt causes, the greater the shunt of blood from the RV to the aorta causing cyanosis. If the PVR decreases, the pressure in the RV which diminishes the D-I shunt. On the other hand if the SVR decreases, increases the D-I shunt causing cyanosis. The symptoms depend on Pulmonary Estenosis [10,11], if it is severe there is cyanosis due to increased D-I shunt by the VSD; if it is not severe increases the I-D shunt by VSD and from there to pulmonary circulation causing pulmonary hyperflow and heart failure. The squatting improves symptoms because the Blood pressure (BP) increases by increasing abdominal pressure which results in an increase in blood pressure which favors the I-D shunt for the VSD. Crises of hypoxia and cyanosis occur when the SVR decreases, due to, for example during anesthetic induction or what increases the D-I shunt (cry –adrenergic discharge- RVOT spasm) and this vicious circle is favored [12]. In laboratory tests, polycythemia stands out (Hto 60-70%), hemostasis abnormalities (thrombocytopenia, platelet dysfunction, hypofibrinogenemia, increased fibrinolysis, coagulation factor deficit) everything that contributes to increase perioperative bleeding.

Anesthetic Management

A balance must be made between the PVR and the SVR. If the obstruction is minimal, there is I-D Shunt through VSD with pulmonary hyperflow, you should avoid diminishing PVR and increase the SVR. If the obstruction is severe with cyanosis and polycythemia, the SVR should be maintained or increased to minimize the I-D Shunt and abundant fluid therapy. For cyanotic children, is recommended the use of ketamine IM or EV and midazolam orally. In contrast, in less cyanotic children, inhalation induction can be performed. Nitrous oxide is contraindicated, because it increases the PVR in patients with congenital heart diseases. For the treatment of cyanotic crisis, the trigger must be avoided, (pain, anxiety), hyperventilate with 100% FiO2 to decrease PVR and abundant fluid therapy (7-10ml/kg). It is very important to know the acid base state of the patient and if necessary correct acidosis with bicarbonate (0.5meq/kg). B-blockers follow a very important role reducing infundibular spasm. (esmolol 100mcg/kg)

Echocardiography in Tetralogy of Fallot [12,13,14]

Echocardiography plays a fundamental role, since in the operating room it will allow the evaluation of VSD before surgery, its location, the degree of aortic riding. It is also important to know the degree of pulmonary stenosis, the state of coronary trunk and pulmonary artery and other associated abnormalities such as the presence of patent foramen ovale that is observed in up to a
third of patients. The interventricular septum defect must also be carefully evaluated, a 80% is perimembranous and is generally a large subaortic defect and extends from the non-coronary leaflet and right coronary leaflet of the aortic valve to the inferior membranous septum and with color doppler determine size and direction of the shunt. The anterior and cephalic deviation of the conal septum causes a narrowing of the RVOT, which begins in the crista supraventricularis and extends to the pulmonary valve ring. Depending on the degree of obstruction of the RVOT the symptomatology will be more or less severe, if it is severe there is D-I shunt through VSD and is a cyanotic patient; and if the obstruction is slight, there is an I-D shunt with pulmonary flow and is called Pink Fallot. This is also why it is important to evaluate the pulmonary valve and its ring, which in a high percentage may be hypoplastic. In addition the echocardiography allows the evaluation of the anatomy of the coronary arteries since there are 10% of patients with coronary abnormalities being the most common that is the right coronary artery crossing the RVOT and this makes it difficult to surgery requiring the placement of a tube of the RV to the pulmonary artery.

Conclusion

In conclusion, intraoperative TEE is essential in the management of a correction of tetralogy of Fallot, since it allows the surgeon to make appropriate decisions and evaluate in the operating room the result of their procedures. For anesthesiologists it has become a fundamental tool for the selection of vasoactive drugs and for complex management of Systemic and Pulmonary vascular resistance.

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