Late diagnosis of total anomalous connection of pulmonary veins during the puerperium

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Abstract

The total anomalous pulmonary venous connection (TAPVC) is a rare congenital anomaly. We present a case of a 36-year-old woman, in the postpartum period during the physical examination there was a heart murmur and beating in hands and feet. Cardiac magnetic resonance reported supracardiac TAPVC and large atrial septal defect. The survival of these patients is very uncommon in adults and even worse during pregnancy.

Keywords

Anomalous Systemic Venous Drainage; Postpartum Period; Adult; Phase-Contrast MRI

Introduction

The total anomalous pulmonary venous connection (TAPVC) is a rare congenital anomaly, which corresponds to approximately 2% of all congenital heart defects [1,2]. The majority of patients present in the neonatal period with decompensated heart failure due to severe pulmonary hypertension or obstruction of pulmonary venous return [3,4]. The natural history of the disease shows that 20% survive until the first year of life without treatment [5]. Exceptionally, the symptoms may appear later in life or progressively, depending on some factors, which will be discussed later [6], the case we present is of a woman who is diagnosed with congenital heart disease during the puerperium. In some series, maternal mortality of patients with pulmonary hypertension, during pregnancy and the puerperium is very high.

Presentation of the Case

A 36-year-old female patient, from 13 years of age, began with dyspnea at great exertion and cyanosis of the lips. At 35 years of age, during his first pregnancy, he presented a slight increase in cyanosis in his lips, hands, and dyspnea at medium and large efforts, so he limited his physical activities [7]. She went to a health care center for the first time to attend her delivery, which was eutocic, without hemodynamic complications. The product was a newborn, full term, with adequate weight for gestational age, healthy. The hospital discharge was usually the second postpartum day. The findings in the physical examination were cyanosis, digital clubbing, without signs of respiratory distress (oxygen saturation 85%, FiO2 24%), the hyperdynamic precordium, cardiac auscultation revealed a regular heart rhythm, a wide and fixed division of the second heart sound with accentuation of the pulmonary component and holosystolic murmur 4/6 in the lung area irradiated to the neck. Liver palpable 4cm below the costal margin. The peripheral pulses were symmetrical and there was no peripheral edema. The chest x-ray revealed...
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cardiomegaly and cardiac silhouette in the shape of a “snowman” (Fig-1A). Echocardiography was not optimal, found dilation of right cavities, atrial septal defect (ASD) of 15 mm and systolic pulmonary pressure of 70 mmHg. Twenty days postpartum, the patient comes to our institution, where the magnetic resonance imaging (MRI) shows the presence of total anomalous connection of pulmonary veins supracardiac variety. The confluence of the four pulmonary veins was wide (diameter 36x25 mm) and was connected to a dilated vertical vein, which in turn drained into a dilated innominate vein (at the mouth its diameter measured 32 x 20 mm), without evidence of obstruction (Fig-1B). Right atrium

Fig-1: Chest X-ray
RPA (Right Pulmonary Artery)
A: Configuration of the heart and the superior mediastinal borders resembling a “snowman”.
B: Echo gradient acquisition shows the pulmonary veins (PV) drain into an enlarged vertical vein (VV) which in turn drains into the innominate (IV) vein and then to the superior vena cava (arrow).

Fig-2: Cardiac Magnetic Resonance (CMR)
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measured 23 cm²/m², left atrium 7.5 cm²/m². The ASD was superior venous sinus type of 25 x 20 mm, the short circuit from right to left and high pulmonary flow (Qp / Qs 2.9). Dilated right ventricle, systolic septal flattening, ejection fraction 48% and moderate tricuspid regurgitation; left ventricle with preserved ejection fraction. Trunk and branches of the pulmonary artery dilated. Presence of late gadolinium enhancement intramyocardial (Fig-2B).

Cardiac catheterization (Fig-3) reported pulmonary arterial hypertension, due to volume overload, even without the involvement of the pulmonary microvasculature (mean of the pulmonary pressure 60mmHg, pulmonary vascular resistance (PVR) 2.5 U Wood/m², PVR: SVR 0.1). With these hemodynamic parameters, it was decided to perform the respective total surgical repair.

Due to the large variations in anatomy and hemodynamics, TAPVC demonstrates a broad spectrum of clinical presentations ranging from the absence of symptoms to severe hypoxemia [8-11]. To survive to adulthood, there should be no obstruction between the pulmonary venous catheter, the systemic vein and the right atrium, an adequate size of the ASD, adequate oxygenation of the arterial blood and exclude the presence of severe pulmonary hypertension. Jian et al [11] in their study describes the average diameter of the pulmonary venous confluence (25mm), the average size of the atrial septal defect (3mm) and the non-restrictive drainage site can provide conditions compatible with long-term survival term. In the present case, similar conditions were observed.

The thorough evaluation of these patients must include right cardiac catheterization, for the
measurement of pulmonary pressure, PVR. Criteria for the closure of the short circuit have been proposed based on the baseline RVP < 4 U Wood / m^2ASC. Additional criteria are the type of defect, the age, the ratio PVR: SVR and the quotient Qp: Qs. There are no prospective data on the usefulness of the vasoreactivity tests, the closure tests or the lung biopsy for the evaluation of operability, however, its use will depend on the treating doctor [12].

In patients with PH, characterized by having fixed and elevated RVP, there is great difficulty in managing high cardiac output during pregnancy and its increase during delivery, which leads to higher mortality. Clinical deterioration appears more frequently in the second trimester of pregnancy, corresponding to a 40% increase in cardiac output [12]. The diagnosis of this case is very rare in the adult, due to the high infant mortality when the corrective treatment is not carried out. The presence of a broad atrial septal defect, which allowed blood flow from right to left, the absence of obstruction of the venous system and the preservation of the pulmonary microvasculature prevented a fatal event during the development of this patient’s life, in this way could be a candidate for corrective treatment.

Conclusions
The diagnosis of TAPVC is infrequent in adulthood. The case presented is exceptional, as it is an adult woman and in the postpartum period. Survival in these cases will depend on the absence of pulmonary venous obstruction, presence of large intracardiac septal defects that allow the right to left shunts. However, the presence of severe pulmonary hypertension and damage to the pulmonary vasculature is common and may limit the corrective treatment. MRI and catheterization are suitable diagnostic methods, for use in preoperative evaluation.

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