PAEDIATRIC CARDIOLOGY - CASE PRESENTATIONS

Sudden cardiac arrest as first symptom of a benign cardiac tumor growth

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KEYWORDS
Tumors; Pediatrics; Cardiac arrest

Abstract Primary cardiac tumors are rare, especially in the pediatric age. Most of them are benign in the sense they are not invasive. However, benign tumors maintain the potential for serious illness related to significant hemodynamic compromise or life-threatening dysrhythmias. We present the case of an infant with an initial diagnosis of cardiac rhabdomyoma who suffered ventricular arrhythmia and cardiac arrest. He suffered irreversible severe neurologic sequelae, due to his prolonged cardiopulmonary arrest and was finally diagnosed of cardiac fibroma. Good arrhythmia control was obtained after an extensive partial surgical resection of the tumor. This case highlights the importance of arrhythmia burden in this condition. A correct diagnosis based essentially in different imaging modalities and closer clinical and rhythm follow up could have avoided this ominous event.

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PALABRAS CLAVE
Tumores; Pediatría; Parada cardiaca

Muerte súbita como primer síntoma de crecimiento de tumor cardiaco benigno

Resumen Los tumores cardíacos primarios son raros, especialmente en la edad pediátrica. La mayoría de ellos son benignos, en el sentido de que no son invasivos. Sin embargo, los tumores benignos tienen el potencial para producir enfermedades graves que pueden causar compromiso hemodinámico significativo o arritmias potencialmente letales. Se presenta el caso de un niño con un diagnóstico inicial de rhabdomyoma cardíaco, quien sufrió una arritmia ventricular y una parada cardíaca. Tuvo secuelas neurológicas severas irreversibles debido al tiempo prolongado en parada cardiorrespiratoria y se le diagnosticó finalmente fibroma cardíaco. Se obtuvo un...
Primary cardiac tumors are rare, especially in the pediatric age. The incidence in this age group is approximately 0.17% in the echocardiographic database review at Boston Children’s Hospital, which is similar to the incidence of 0.2% reported at the Hospital for Sick Children, Toronto.

Most primary pediatric cardiac tumors are benign in the sense they are not invasive. Fewer than 10% of these primary tumors are malignant. However, benign tumors maintain the potential for serious illness related to significant hemodynamic compromise or life-threatening dysrhythmias.

Is presented a case of a patient diagnosed of a benign primary cardiac tumor who developed a serious complication and poor outcome in follow up.

Case

A two year-old infant with prenatal diagnosis of cardiac mass, who collapsed at home while playing. He was being followed since birth in a cardiology outpatient clinic with the diagnosis of benign cardiac rhabdomyoma. He suffered a cardiopulmonary arrest and was taken to the emergency department where advanced resuscitation maneuvers were practiced for 40 minutes. First documented cardiac rhythm was ventricular fibrillation. After initial stabilization and treatment, he was transferred to our centre for multidisciplinary treatment.

Chest x-ray demonstrated an increased cardiothoracic index (0.70) with pulmonary congestion. 12-lead electrocardiogram showed unremarkable sinus rhythm with an average heart rate of 130 bpm, normal PR and QRS intervals, and QRS axis in 0°. There were symmetrical and deep T-wave inversions in the lateral precordial leads. Transthoracic echocardiography demonstrated a homogeneous intramural left ventricular mass in the free wall and apex, which involved the anterior papillary muscle, with no valvar incompetence nor mechanical left ventricular outflow or inflow obstruction. The image suggested a fibroma as the first diagnostic possibility (fig. 1). Cardiac magnetic resonance images revealed a 4.5 × 4.8 × 4.1 cm, firm, well-demarcated intramural mass related to the anterior papillary muscle and the subendocardial free wall region of the middle and apical segments of the left ventricle. Late contrast-enhanced cardiac magnetic resonance images were acquired after gadolinium injection, showing a homogeneous and intense bright mass, suggesting the nature of fibroma (fig. 2). A slight global left ventricular systolic dysfunction was observed, with an ejection fraction of 55%.

The patient had a severe irreversible neurological dysfunction due to a severe hypoxic ischemic encephalopathy secondary to the prolonged cardiac arrest. Given this situation, he was unable to swallow and a percutaneous gastrostomy was placed.

During admission, he presented episodes of non-sustained ventricular tachycardia as well as multiple episodes of different monomorphic sustained ventricular tachycardia. Ventricular tachycardia morphology suggested its origin from left ventricular apex, coinciding with cardiac tumor’s location. Pharmacologic treatment was initiated with amiodarone and propranolol with poor control of the arrhythmia. The decision was taken to perform a complete surgical resection of the tumor (fig. 3). Finally, only an extensive partial tumor resection was undertaken due to technical

Figure 1  Two-dimensional echocardiogram in a 2-year-old child with a large left ventricular fibroma. A. Subcostal sagittal sweeps showing homogenous intramural mass extending from the left ventricular free wall into the cavity. B. Subcostal coronal sweeps revealing no left ventricular outflow obstruction. C. Parasternal short-axis shows tumor involving the anterior papillary muscle.
Sudden cardiac arrest as first symptom of a benign cardiac tumor growth

Figure 2  Cardiac magnetic resonance imaging. A. Axial T1-weighted spin-echo image; B. DP-weighted spin-echo image; C. 4 chamber view gradient-echo cine; D. Short axis gradient-echo cine. An isointense solid mass is observed in T1, slightly hyperintense relative to muscle on T2 and gradient-echo sequences, located in left ventricular lateral wall. E, F. Late contrast-enhanced cardiac magnetic resonance images after gadolinium injection in short axis (E) and four chamber view (F), showing a homogeneous mass. Lateral wall and apex myocardium can be appreciated, closely to the mass on its lateral, top and bottom margins and papillary muscle (arrow).

Discussion
Cardiac fibroma could be considered a benign condition, but slow and continual growth has been reported and may cause significant morbidity related to obstruction, infiltration, thromboembolism, arrhythmias or even death. Clinical significant arrhythmia can occur in up to 24% of pediatric patients with cardiac tumors, being fibromas the subtype with the highest risk, with risk of ventricular tachycardia exceeding 50%. Special attention must
be taken then to arrhythmia in this condition. A rapid and precise tumor diagnosis and close follow up could be essential to prevent an ominous outcome as in the case presented.

Diagnosis is based in different imaging modalities. Echocardiography has been and remains an invaluable tool for cardiac evaluation. It permits a rapid and cost-effective evaluation of cardiac masses. The typical echocardiographic appearance of a fibroma is a discrete often obstructive, echogenic, noncontractile mass ranging in size from 1 to 10 cm in diameter in a ventricular wall. They are single, bright, intramural masses within the interventricular septum or the left ventricular free wall. In appearance it may mimic hypertrophic cardiomyopathy or ventricular septal hypertrophy. Macroscopically, they are characteristically solitary, white, firm, nonencapsulated, intramural tumors that involve the left ventricular free wall, apex or interventricular septum. Less frequently, they are multiple and invade the right ventricular free wall, atrial septum, or atrial free wall. Extensive intramural fibromas can encroach and obliterate the intracavitary space. Although rare, intracavitary fibromas have been reported, occurring either as a single pedunculated mass or attached by a broad base to the endocardium. They may be echocardiographically differentiated from rhabdomyomas by the presence of calcification and cystic degeneration; also with strain imaging, fibromas are shown not to be compressed during the cardiac cycle.

Computed tomography and cardiac magnetic resonance allow for better tissue characterization. At computed tomography, fibromas appear as homogenous masses with soft tissue attenuation that may be either infiltrative or sharply marginated. Calcification is often seen. Cardiac Magnetic Resonance can help to assess tumor location, size, morphology, borders and most importantly, provide definite tumor characterization while avoiding radiation. Cardiac magnetic resonance has been shown to correctly classify tumors as benign or malignant in 95% of cases, when comparing with histology as the gold standard. On cardiac magnetic resonance, fibromas are strong hyperenhancement on myocardial delayed enhancement imaging with or without a hypoenhancing (dark) core. They have a heterogeneous appearance on T1- and T2-weighted turbo (fast) spin echo sequences with variable areas of slightly hypo- or hyperintense areas. It also evaluates the extent of myocardial infiltration as well as the relationship of the tumor mass to coronary artery anatomy.

Regular frequent clinical assessments and Holter monitoring could be necessary in order to determine arrhythmia burden. Clinically significant ventricular arrhythmias are common and should be promptly evaluated. The baseline electrocardiogram may show QRS axis deviation, atrial enlargement, or ventricular hypertrophy, and invariably present ST-T wave abnormalities consistent with strain or ischemia typically in left lateral leads. If high degree ventricular ectopy or non-sustained ventricular tachycardia is present, treatment should be preferably encountered.

Optimal management strategies for pediatric patients with cardiac tumors remain unclear, particularly when serious arrhythmias are present. Surgical excision of such tumors has been reported to be a highly effective management option, being curative and lifesaving therapy. Surgery for cardiac tumors in pediatric age carries an acceptable mortality risk and appears to be the optimal treatment in patients with symptomatic resectable tumors. In patients with severe tumor involvement, subtotal excision has also shown good long term survival. The role of surgery in patients with asymptomatic tumors that could remain dormant for many years and even regress is less clear. However, because of fatal arrhythmias, surgery is often recommended despite absence of symptoms. Even transplantation is considered for large symptomatic unresectable tumors.

In conclusion, this case report shows how a benign cardiac tumor could have devastating consequences. Clinicians should be aware of arrhythmia risks in cardiac tumors in children, especially in fibromas. A correct tumor type diagnosis based essentially in different imaging modalities, and closer clinical and rhythm follow up could have avoided the event and severe neurologic sequelae.

Ethical disclosures

Protection of human and animal subjects. The authors declare that no experiments were performed on humans or animals for this study.

Confidentiality of data. The authors declare that no patient data appear in this article.

Right to privacy and informed consent. The authors declare that no patient data appear in this article.

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Conflict of interest

Any.
References


