Primary Cardiac Fibroma and Cardiac Conduction System Alterations in a Case of Sudden Death of a 4-month-old Infant

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**Abstract:** A 4-month-old female infant considered to be in good health died suddenly and unexpectedly. Post-mortem examination was requested, with clinical diagnosis of sudden infant death syndrome. At autopsy the infant was described in good health. Histological examination of the heart found a cardiac fibroma compressing the atrio-ventricular node and the examination of the cardiac conduction system showed an accessory fiber of Mahaim (nodo-ventricular) and cartilaginous metaplasia of the cardiac fibrous body. Probably the concomitant presence of cardiac conduction system abnormalities and a septal fibroma, compressing the atrio-ventricular node, could have an important role in causing the sudden death.

**Keywords:** SIDS, Primary cardiac fibroma, cardiac conduction system.

**INTRODUCTION**

Primary cardiac tumors are very rare. In infancy and childhood they are even more infrequent, with a prevalence of 0.0017 to 0.28 in autopsy series. In contrast, the incidence during fetal life is approximately 0.14% [1-3].

Cardiac fibroma is the second most common childhood cardiac tumor [4, 5]. It typically originates from the ventricular septum or ventricular free wall. Although cardiac fibroma is histologically benign and does not metastasize or infiltrate, it can cause hemodynamic and conductive abnormalities resulting in sudden death [1].

This report will focus on the histological examination of the heart and conduction system in a case of sudden death in a 4-month-old infant.

**CASE REPORT**

The infant was a female 4-months-old, born at 37 weeks of gestation, with a birth weight of 2270 g (low birthweight, at the 5th percentile). She was in apparent good health until the day she died, suddenly and unexpectedly. Her parents said that they knew of no illness or complaint of any nature, except for two episodes of vomit two days before. There was no significant family history relevant to the case. In particular, there was no familiarity for tuberous sclerosis and Gorlin’s syndrome (multiple nevoid basal carcinomas, jaw cysts, and bifid ribs), both associated with cardiac fibromas [1].

A complete autopsy was performed, according to the autopsy protocol usually followed by our center in cases of sudden death [6]. Multiple samples of all organs were fixed in 10% formalin buffer, processed, and embedded in paraffin. Five-micrometer sections were stained with hematoxylin and eosin. The heart had been regularly examined for pathological changes in the atria, septa, ventricules, pericardium, endocardium, and coronary arteries. Samples of the myocardium were stained with hematoxylin and eosin and trichromic Heidenhain (azan). Histological observations were focused on the cardiac conduction system, which was removed in two blocks, according to our guidelines [6]: the first included the sino-atrial node (SAN) and the “crista terminalis,” the second contained the atrioventricular node (AVN), His bundle, and bundle branches. The samples were stained alternately with hematoxylin and eosin and trichromic Heidenhain (azan).

At autopsy, the infant was described in good health, with body length and weight at the 5th percentile. There were no marks of violence. The lungs were congested and slightly edematous. The heart, arrived without the tip, weighed 25 g; the cardiac diameters were as follows: transverse 3.5 cm and anteroposterior 2.9 cm. We could not measure the longitudinal diameter due to the absence of the tip. The myocardium was brownish and homogeneous in appearance, without any evident alteration. The foramen of Botalio was open for about 3 mm. The coronaries appeared normal.

Histological examination of the atrio-ventricular node showed an accessory fiber of Mahaim (nodo-ventricular), cartilaginous metaplasia of the cardiac fibrous body and atherosclerotic lesions of the artery of the atrio-ventricular node (Fig. 1). It was also present a non-encapsulated proliferation of fibroblasts with regular nuclei and an abundant network of collagen and elastic fibers that dissociated and entrapped the myocytes (Fig. 2). These morphological fea-
tures led to the diagnosis of primary cardiac fibroma compressing the atrio-ventricular node [5]. Mitotic figures and cytologic atypia were not observed. No foci of dystrophic calcification were present. The SAN, adjacent nerve plexuses, and internodal right atrial myocardium were free of alterations.

DISCUSSION AND CONCLUSIONS

Sudden infant death syndrome affects about 0.7-1 per 1000 live births [7]. The most accredited pathogenetic hypothesis of SIDS (cardiac-arrhythmogenic, respiratory-apnea and/or suffocation and dyskinetic-visceral) seem to give preference, in most cases, to a dysfunction of the autonomic nervous system which control the cardio-respiratory activity, the first upper digestive tract and the awakening [8-10].

Although the diagnosis of SIDS and sudden unexplained perinatal death is by exclusion, changes of the cardiac conduction system and brainstem are of great value in establishing and characterizing these forms of death [11-13]. Although the majority of these fatalities are related to sudden infant death syndrome (SIDS), a very small percentage is due to primary cardiac neoplasm [11]. This is the second case of fibroma of the fibrous trigone found in a series of 69 cases of death examined.

Cardiac fibroma is a rare tumor in pediatric practice but, after rhabdomyoma, it is the second most frequent tumor of childhood in autopsy series [3-5]. Fibromas are usually large and intramural, occupying the septum and/or the anterior wall of the left ventricle. The tumor is composed of fibroblasts, dense connective tissue, and a small admixture of myocardial cells [5]. Intramural cardiac tumors are frequently associated with disorders of the rhythm, particularly in children [8]. In our case the neoplasm was adjacent and compressing the atrio-ventricular node. In the cases described in literature were reported significant alterations of the rhythm [1, 5, 14]. In our case the tumor was clinically silent.

Sudden infant death syndrome (SIDS) “gray zone” or borderline cases are defined as those cases in which it is difficult to establish whether the pathological findings are really cause of death [15,16]. Krous et al. described as “unclassified sudden infant deaths” those cases that do not meet the criteria for a diagnosis of SIDS [17]. In fact, according to the original definition, SIDS is “… the sudden death of any infant or young child, which is unexpected by history, and in which a thorough post-mortem examination fails to demonstrate an adequate cause for death” [18]. In 1991 the National Institute of Child Health and Human Development (NICHHD) described more precisely inclusion and exclusion criteria for SIDS [19], but it is still not present a clear classification of these deaths.

In our case, the concomitant presence of cardiac conduction system abnormalities (an accessory fiber of Mahaim and cartilaginous metaplasia of the cardiac fibrous body) and a septal fibroma, compressing the atrio-ventricular node, could have an important role in causing the sudden death.

Infact, the circumstance that the patient died suddenly and unexpectedly and that a fibroma was found compressing...
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the atrioventricular pathway suggest that the two facts could be causally related.

The anatomic abnormalities observed in the present case concur with a diagnosis of SIDS/sudden unexplained death gray zone, without any proven sequential occurrence.

CONFLICT OF INTEREST

The authors confirm that this article content has no conflicts of interest.

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