Rhabdomyomatosis of the heart associated with tuberous sclerosis. A case report and review of the literature

Abstract
Congenital heart rhabdomyomatosis is a rare condition that is associated in most cases with tuberous sclerosis, a genetic syndrome inherited with an autosomal dominant pattern. The last is caused by a mutation in tuberous sclerosis 1 or tuberous sclerosis 2 genes. We report a case of heart rhabdomyomatosis, associated with tuberous sclerosis. The lesion was discovered at 21st week of gestation of a 36-year-old woman.

Keywords: prenatal diagnosis, cardiac rhabdomyomatosis, tuberous sclerosis, tuberous sclerosis gene 1, tuberous sclerosis gene 2

Abbreviations
TSC: Tuberculous Sclerosis

Introduction
Rhabdomyomas, whilst are rare, they are the most common cardiac tumor in pediatric patients [1], accounting for 50-75%. In a postnatal study [2] the referring cases where sporadic, associated with other cardiac anomalies or associated with tuberous sclerosis—approximately 50% of cases are associated with the disease [1]. The latter is a genetically inherited disease with an autosomal dominant pattern of inheritance and an incidence approaching 1:6000 births [3]. Mutation in TSC1 or TSC2 gene remains the main cause, with 60% of cases occurring de novo [3]. TS can affect all organs with skin, heart, brain, and kidneys being the most common targets. In this case we report the appearance of fetal heart rhabdomyomatosis associated with tuberous sclerosis and discovered during the routine follow up of the pregnancy.

Case Report
A 36-year-old female with subfertility and previous premature labor in 26th week of gestation that leads in newborn’s death decided to undergo in vitro fertilization. Her past medical history included cyst in the left ovary, leiomyoma of uterus and hypothyroidism treated with thyrohormone (25 mg per day). The embryo transfer was successful and the woman underwent the routine follow up. At the 21st week of gestation ultrasound monitoring showed multiple nodules in the fetus heart without any other congenital abnormalities.

Embyro’s heart ultrasound detected three hyperechogenic intracardial masses (6-13 mm in diameter). The largest one occupied most of the left ventricle FIGURE 1a. The second one (7 mm) was near the tricuspid valve (with partial impairment of leaflet’s opening) and the third one was at the interventricular septum of the aspect of the right ventricle (6 mm in diameter) FIGURE 1a and 1b. Fetal cardiac function and size were normal. No associated cardiac anomaly was detected. The fetal cranial sonographic examination was normal.

After prenatal counseling for genetic disorders and amniocentesis, the result of the genetic analysis confirmed the diagnosis of tuberous sclerosis. The genetic test detected in the male fetus c.1716+2T>G mutation of TSC2 gene in a heterozygotic pattern. The parents decided to terminate the pregnancy at 23rd week of gestation and the embryo was sent for histological examination.

The autopsy revealed a male embryo, without any congenital malformations, weighing 660 gr with 34 cm crown-heel length, 22 cm crown-rump length and foot length 5 cm. After dissection of the body, the extraction of the heart...
was performed. Its dimensions were $4 \times 3 \times 2$ cm and weighted 7 gm. Cardiac sections showed coagulated blood in right atrium, right ventricle and left atrium. Three firms, well-circumscribed whitish nodules, 0.6 cm, 0.7 cm and 1.2 cm in diameter, respectively were observed. The smallest one was in the interventricular septum at the aspect of the right ventricle, while the other two protruded in the cavity of the left ventricle, pushing the interventricular septum towards right ventricle. In the abdomen, there was an insignificant peritoneal effusion, while sections of all other organs did not show any macroscopic lesions.

Series of sections from these cardiac nodules were obtained and examined. Microscopically, typical findings of rhabdomyoma were noticed. The tumor cells, the so-called spider cells Figure 2a, were large with vacuolated cytoplasm full of glycogen (Periodic acid-Schiff reactivity), some of which had a polygonal shape, centrally located nucleus and distinct cytoplasmic extensions (designated with histochemical stain PTAH). Figure 2b Immunohistochemical investigation of the neoplasm showed reactivity for myoglobin Figure 2c and myoD1, as well as focal positivity to desmin. Proliferation index MIB-1 was minimal.

**Discussion**

Rhabdomyoma is a rare benign tumor of striated muscle fibers, even though some support it as being as hamartoma rather than a true neoplasm [3]. Even though its rarity, it is the most common cardiac tumor, accounting for more than 50% of benign cardiac neoplasms [1,4]. The first reported case was in 1862 [5]. It can occur sporadically [6,7] some appear with other congenital heart disease [8], or in association with tuberous sclerosis [1,8,9] in a rate around 50-60% [1,3].

Usually, it is developed in ventricles with the left one being the most common location. There are reported cases in septum [10] and in intrapericardial space.

**FIGURE 1.** (a): Two round homogeneous hyperechogenic masses (6.1 mm and 13.1 mm). The largest protruded in the left ventricle and the small one in the interventricular septum; (b): Cardiac rhabdomyoma 7.2 mm near the tricuspid valve.

**FIGURE 2.** Microscopic examination of the tumor. (a): “spider” cells with vacuolated cytoplasm (haematoxylin-eosin, x20 magnification); (b): positive immunohistochemical reactivity for myoglobin (x20 magnification); (c): distinct cytoplasmic extensions (histochemical stain PTAH, x60 magnification).
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In intrauterine life they cause echocardiograph changes [13], arrhythmias, hydrops [8] or nothing. Postnatal, infants can be asymptomatic or only having a murmur, whilst some of them suffer from congestive heart failure, severe arrhythmias or even sudden death [3,14]. Cardiac rhabdomyomas may continue to grow postpartum, due to maternal estrogen. The later decrease in the size of the tumor may be due to reduced estrogen levels [15]. These tumors are at their largest in fetal life, then shrink with age and may even disappear completely. The prevalence is, therefore, higher in children than in adults. Surgical resection should be performed if cardiac function is adversely affected [16].

Macroscopically, they are well-circumscribed, without capsule surrounding, gray-white, firm, small nodules with an average size from millimeters to several centimeters, embedded in the myocardium or protruding in cardiac cavities [17]. It can reach up to 10 cm in diameter and often are multiple. The term rhabdomyomatosis refer to numerous nodules that are not greater than 1 mm in diameter.

Microscopically [17], the tumor consists of clusters of large polygonal to rounded cells ("spider cells") with vacuolated cytoplasm full of glycogen deposits (PAS-diastase positive histochemical stain) separated by strands of cytoplasm extending between cell membrane and nucleus, without any mitotic activity. Immunohistochemical investigation [17] reveals reactivity to striated muscle cells antigens, such as myoglobin, actin, desmin, vimentin, and variable to HMB45, although S-100 protein is negative.

Early recognition is really important in order to investigate any connection with tuberous sclerosis [18-20] and to plan treatment after birth, which includes surgery if patient's life is threatened [21,22] and drugs, such as everolimus [3,23,24]. It is worth mentioning that some masses regress spontaneously after birth [8,25] while other investigators believe that the younger the age of the diagnosis, the greater the chance for regression [3].

Tuberous sclerosis is an autosomal dominant inherited disease which affects males and females equally. It is due to mutations of TSC1 and TSC2 gene which are located at chromosome 9q34 and 16p13, respectively, with 60% of cases to be due to de novo mutations [3]. The disease targets almost all organs such as brain, lungs, heart, kidneys, bone, eyes and skin. The clinical manifestations include seizures, hydrocephalus, mental retardation, rhabdomyomas [21], renal angiomyolipomas [11], developmental delay, spine malformations and skin angiofibromas [26]. Some authors focus on prenatal diagnosis of tuberous sclerosis [11,20] and especially the recognition of cardiac rhabdomyomas [1,27] since it is the commonest presentation. Concerning the disease's therapy and management, there is huge progress [28], with mTORC1 inhibitors to be in the forefront [29].

In a meta-analysis, Chao et al. [30] reported that most cardiac rhabdomyomas were detected after the 24th gestational week and only 13.7% of the cases were detected before the 24th gestational week. There have been many cases reported of rhabdomyomas/rhabdomyomatosis connected with tuberous sclerosis [31,22] diagnosed prenatally and in the postnatal period, such as a giant rhabdomyoma [24,32,33] neonatal intracardiac rhabdomyomatosis and a case of diffuse rhabdomyomatosis without any distinct mass formation [34]. There are cases with rhabdomyomatosis of lung [35-37] and meninges [38]. To the best of our knowledge there is only one case of tuberous sclerosis and heart rhabdomyoma in a patient with trisomy 21 [19,39]. Rhabdomyomas usually develop in the left ventricle, but there is a case reported in the right ventricle [40], in the right atrium of a patient suffering from Wolff Parkinson White syndrome [41] and a case occurring in both ventricles and right atrium without any connection with tuberous sclerosis [22].

**Conclusion**

Since heart rhabdomyomas/rhabdomyomatosis is one of the most prominent clinical manifestations in patient with tuberous sclerosis, an early diagnose is critical for the patient itself as well as for the family members since it is a congenital disease.
REFERENCES


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