A Rare Case of Asymptomatic Giant Right Atrial Aneurysm: Case Report

Asemptomatik Dev Sağ Atriyal Anevrizmalı Nadir Bir Olgu

ABSTRACT: Right atrial aneurysm is a rare abnormality of the heart. This defect is seen in all age groups and is usually diagnosed incidentally. The aneurysm can be asymptomatic or present with symptoms related to supraventricular arrhythmias or thromboembolic phenomenon. We present the case report of a 74-year-old male with right atrial aneurysm which was incidentally detected during echocardiographic evaluation.

Keywords: Heart atria; aneurysm; echocardiography; tomography, x-ray computed

ÖZET Sağ atriyal anevrizma kalbin nadir ve sıklıkla konjenital orjinli bir anomalisidir. Fetal dönemden geçişerik döneme kadar, tüm yaş gruplarında görülen bu defekt genellikle rastlantısal saptanmaktadır. Asemptomatik olabileceği gibi supraventriküler aritmiler veya tromboembolik olaylar ile de ortaya çıkabilir. Bu yazida ileri yaşa kadar tanı almayan, asemptomatik 74 yaşında bir erkek olgu ekokardiografi sırasında tesadüfen saptanması nedeniyle sunulacaktır.

Anahtar Kelimeler: Kalp atriumları; anevrizma; ekokardiografi; tomografi, bilgisayarlı x-ray

The right atrial aneurysm is a cardiac malformation of unknown etiology, rarely found in cardiological practice. It can be detected at any time between fetal and adult life. Approximately half of patients show no symptoms. If no treatment is carried out, the risk of complications such as arrhythmia and thromboembolism will increase; therefore, medical or surgical treatment is recommended. In our article we present a rare and asymptomatic patient who was diagnosed with right atrial aneurysm at 74 years of age by echocardiography and computed tomography (CT) angiography.

CASE REPORT

A 74-year-old male patient was admitted to our department after referral from another hospital because of hypertension, and an abnormality was detected incidentally via echocardiography. His and his family’s past history showed non specific findings. In the physical examination there
weren’t any pathologic findings. An electrocardiogram (ECG) showed sinus rhythm with QRS axis of -30°. Surprisingly, the P wave was normal (Figure 1). A chest radiograph showed marked cardiomegaly with a cardiothoracic ratio of 60%. Laboratory findings were normal. On an echocardiogram there was a large cystic structure arising from the anterior freewall of the right atrium and extended across the atrioventricular groove over the anterior aspect of the right ventricle (Figure 2). The right atrial dimensions were 8.7x9.7 cm. No abnormality was found in the structure and function of the valves, and abnormal wall motion was not observed. Transesophageal echocardiogram (TEE) showed a cystic mass arising from the anterior freewall of right atrium, but no thrombus was found. The lesion was confirmed by CT angiogram, which depicted a giant aneurysm of the right atrial free wall and causing slight compression of the right ventricle, otherwise two coronary systems were intact (Figure 3). It was considered that his signs were associated with the presumably congenital right atrial aneurysm. The patient was asked to undergo right atrial aneurysm resection due to the risk of complications such as arrhythmia and thromboembolism, but he denied the procedure and instead started anticoagulant therapy. At a recent 1-year follow-up visit, this patient was asymptomatic and doing well clinically.

**DISCUSSION**

An atrial aneurysm is defined as the dilatation of the atrium involving all layers of the atrial wall. Aneurysms of the right atrial are very rare and its origin is not clear. Although most right atrial

**FIGURE 1:** ECG showing sinus rhythm.

**FIGURE 2:** Transthoracic echocardiography from the apical four-chamber a) and modified b) view demonstrating an aneurysmal structure related to the right atrium. LA; Left atrium, LV; Left ventricle, RA; Right atrium, RV; Right ventricle, TV; Tricuspid valve, RAA; Right atrial aneurysm.

**FIGURE 3:** Computed tomography angiography scan demonstrating a right atrial aneurysm. LA; Left atrium, LV; Left ventricle, RA; Right atrium, RV; Right ventricle, RAA; Right atrial aneurysm.
aneurysms have a congenital origin, acquired cases due to trauma have also been reported. The entity was first described in 1955 and performed the first surgical resection of a right atrial aneurysm in 1968.

In a review of 103 sporadic cases with congenital malformations of the right atrium and the coronary sinus published between 1955 and 1998 of the 105 cases in this series, 60 were cases of congenital enlargement of the right atrium. The age at presentation varies from the neonatal period to late adulthood. The most common symptoms are palpitations due to atrial tachycardia, including atrial flutter and fibrillation. Other rhythm abnormalities include pre-excitation, junctional rhythm, atrioventricular block and incessant supraventricular tachycardia. In addition, potential complications include pulmonary thrombo-embolism, paradoxical systemic embolization and atrial rupture.

The majority of patients are asymptomatic and diagnosed with giant right atrial aneurysm after an incidental finding of cardiomegaly on chest radiograph. The diagnosis can be difficult as it is easily mistaken for more common anomalies that involve right atrial enlargement. Alternative diagnoses that can mimic giant right atrial aneurysm include Ebstein disease, pericardial effusion, pericardial cysts, and tumors. It is distinguishable from Ebstein’s anomaly with normal placement and function of tricuspid valve. Correct diagnosis is necessary for planning therapy. The diagnosis of right atrial aneurysm can be established with echocardiography, angiography, CT, or MRI. In our case, right atrial aneurysm was diagnosed by echocardiography, TEE, and confirmed by CT angiography.

Treatment options depend on clinical presentation, and include anticoagulation, catheter ablation, and surgery. Surgical treatment can be considered in symptomatic cases and in patients with symptoms due to compression of large aneurysms. Follow-up with conservative treatment may be appropriate in most cases. Asymptomatic patients should be managed conservatively with treatment directed toward prevention of thromboembolic complications.

In any event, asymptomatic patients should be evaluated individually for treatment. We recommended surgical treatment due to the risk of death from complications. However, he denied the treatment for the reasons of old age and the absence of symptoms. Currently, his progress is being observed with medication.

**Informed Consent**

Written informed consent was obtained from patient who participated in this case.

**Conflict of Interest**

Authors declared no conflict of interest or financial support.

**Authorship Contributions**

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