Aneurysm of the Right Atrial Appendage

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Aneurysms involving the free wall or atrial appendage are rare entities in cardiology practice and may be associated with atrial arrhythmias or embolic phenomena. We review the literature and report a case of aneurysm of the right atrial appendage in a young adult, whose diagnosis was established with echocardiography after an episode of paroxysmal atrial flutter.

Atrial aneurysms are extremely rare entities in cardiology practice. In the literature, we found 49 reported cases in the left atrium and only 7 in the right atrium. Left atrial aneurysms may be congenital and intrapericardial or secondary to the partial absence of the pericardium (due to herniation of the left atrial auricula). They most frequently manifest as recurring or incessant atrial arrhythmias, which may be refractory to medicamentous treatment and require surgical resection of the aneurysm. In addition, systemic embolization may occur as a severe complication; therefore, long-term anticoagulation is indicated.

Right atrial aneurysms may be congenital and intrapericardial involving the free wall or they may result from trauma. No report of aneurysm of the right atrial appendage exists in the literature. Patients may be asymptomatic or have atrial arrhythmias and repetitive pulmonary embolism.

Case report

The patient is a 23-year-old male who sought emergency treatment complaining of rhythmic tachycardic palpitations of sudden onset. He reported a similar episode 4 years before, which was treated in the emergency department with intravenous medication. The physical examination was within the normal range, except for a heart rate of 150 bpm and deviation of the ictus cordis to the left. The chest X-ray showed global enlargement of the cardiac area. The electrocardiogram diagnosed regular atrial flutter (type I), which was successfully treated with chemical cardioversion with amiodarone. The following complementary examinations (thoracic echocardiogram, transesophageal echocardiogram and chest tomography) revealed a giant intrapericardial aneurysm of the right atrial appendage. On thoracic echocardiogram, the aneurysm measured 15x8 cm, caused compression of the middle and basal regions of the right ventricle and deviation of the cardiac structures to the left (fig. 1). Both ventricles were of normal size and function. The left atrium was normal. On Doppler, significant diastolic restriction to the filling flows of both ventricles was not found. During transesophageal echocardiography, intense spontaneous contrast (stasis) was detected inside the aneurysm, but with no thrombi (figs. 2, 3, and 4). No thrombi could be seen in the left atrium. Patent oval foramen was also diagnosed, with no hemodynamic repercussions.

The patient refused to undergo surgery, therefore, being kept on clinical medicamentous treatment with amiodarone and oral anticoagulant. Currently, the patient is asymptomatic and free from morbid events in the 9th month of ambulatory follow-up.

Discussion

The case we report differs from those in the consulted literature in regard to anatomical features and clinical evolution. In regard to the location of the aneurysm, all those reported aneurysms were located in the trabecular portion of the right atrial free wall, anterior to the right ventricle. Our patient is the first reported with an aneurysm located specifically in the right atrial appendage, similar to the cases reported for the left atrial appendage (this location is more common for aneurysms of the left side). In regard to clinical evolution, even though the patient may be asymptomatic and the diagnosis established as a surgical or complementary examination finding, the most common occurrence was the arrhythmic manifestation. The resulting atrial arrhythmias evolve in an incessant or recurring way, and the potential risk of systemic or pulmonary embolic phenomena occurs. In the literature, we found reports of 2 asymptomatic patients. One was only diagnosed in the surgical suite during myocardial revascularization, when the aneurysm was resected and the patient evolved uneventfully; the other
patient, whose diagnosis was an examination finding \(^1\), chose clinical management. One patient had a recurring pulmonary embolism \(^10\), was treated with oral anticoagulation, and remained asymptomatic for more than 4 years, when the case was published. One patient was diagnosed in the prenatal period, and gestation was interrupted \(^12\). The remaining 3 cases had incessant atrial arrhythmias (1 fibrillation, 1 flutter and 1 atrial tachycardia). Due to refractoriness to clinical treatment, the surgical treatment was indicated and resolved the arrhythmias found in 2 patients \(^8-9\). The unsuccessful case \(^1\), a patient with atrial fibrillation, also had 1 complication on the 4\(^{th}\) postoperative day; embolism to the anterior descending artery, which had not been recanalized by angioplasty, and new surgery was required. This event showed the need for surgical exploration of the left atrium in surgical resection of right atrial aneurysms, even when complementary examinations do not reveal thrombi. No death related to right atrial aneurysm has been reported in the literature. Our patient had only 2 episodes of arrhythmia during his 23 years of life; he evolved asymptomatically and not medicated until the 2\(^{nd}\) episode. No manifestation of embolic phenomena could be detected through current or previous histories or on physical or complementary examinations. Because of the good clinical evolution, absence of incessant arrhythmia, and the patient’s desire not to undergo surgery on that occasion, we chose medicamentous treatment with oral anticoagulants and antiarrhythmic agents, with follow-up with a complementary imaging examination every 6 months.

Despite the tendency found in the literature reports towards surgical indication, especially in symptomatic cases, the rarity of the affliction does not allow conclusions about the efficacy of surgery in curing arrhythmia and improving the prognosis of embolic events. We believe that, when facing such a rare affliction, individualization of the treatment according to clinical features (valuing the presence of incessant atrial arrhythmia and embolism) and complementary examinations (valuing the presence of thrombi, compression of adjacent structures, and associated diseases) is required.

In conclusion, aneurysm of the right atrium or atrial auricula is a rare malformation, which may evolve with high morbidity, and, therefore, should be remembered as a potential anatomic cause of atrial arrhythmias or embolic phenomena, or both. The diagnosis may be easily established through noninvasive complementary techniques, such as echocardiography.
References