Cardiac tumors: a brief commentary

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INTRODUCTION

It is called tumor any abnormal growth, whether it be cancerous or non-cancerous. Tumors originating from the heart are called primary tumors and can occur in any of its tissues. They can be cancerous or non-cancerous and are rare. Secondary tumors originate elsewhere in the body often the lung, breast, or skin and then spread to the heart (1–3).

Secondary tumors are 30–40 times more common than primary, but still are considered unusual. Cardiac tumors may not cause symptoms or may produce a potentially lethal cardiac dysfunction, mimicking other diseases. Examples of such disorders include sudden heart failure, sudden onset of arrhythmias, and sudden drop in blood pressure due to bleeding in the pericardium (4–6).

Cardiac tumors are difficult to diagnose both being relatively uncommon, as its symptoms are similar to those of many other disorders. To arrive at a diagnosis, it is necessary that the physician has evidence of their presence. For example, if a person has cancer elsewhere in the body, but seeks medical attention because of symptoms related to cardiac dysfunction, the professional may suspect the presence of a cardiac tumor (5, 6). However, if a cardiac mass represents a tumor, its etiology can often be determined by considering four factors: (1) the histology-based likelihood; (2) the age of the patient at time of presentation; (3) the tumor location; and (4) non-invasive tissue characterization.

A PubMed, Europubmed search was performed until July 2014 using the terms “cardiac tumors” in combination with the terms “heart tumors,” “myxomas,” or “cardiac myxomas.”

MYXOMAS

Myxoma is a non-cancerous tumor, and usually has an irregular shape and a gelatinous consistency. Half of all primary cardiac tumors are myxomas. Three quarters of myxomas located in the left atrium, the heart chamber that receives oxygen-rich blood from the lungs. Generally, the left atrial myxomas originate from a pedicle and can swing freely with the blood flow, like a ball attached to a wire. When oscillating, myxomas can move into and out of the mitral valve next to the way between the left atrium and left ventricle. Cardiac myxomas are usually solitary and develop in the atria, 75% originating in the left atrium and 15–20% in the right atrium. They characteristically arise from or near the interatrial septum at the border of the fossa ovalis membrane. They occur in all age groups, most frequently between the third and sixth decades. Women are more commonly affected. Cardiac myxomas range in size from 1 to 15 cm in diameter (7–10).

This oscillation can block and unblock the valve continuously, so that the blood flow is stopped and restarted intermittently. In the standing position, the person may faint or episodes of pulmonary congestion and difficulty breathing, as the force of gravity causes the tumor to move downwards until the valve opening the position reduced the symptoms (11).

The tumor can damage the mitral valve and, consequently, a back flow of blood occurs through this opening, with the production of a murmur that the doctor listens with a stethoscope. Based on the characteristics of the murmur, the doctor considers whether the breath is the result of a flow caused by an
injury arising out of the tumor, a rare cause or a more common cause of rheumatic heart disease. Fragments of a myxoma or blood clots that form on the surface of the myxoma may detach, move to other organs, and clog blood vessels in these locations. Symptoms depend on the blocked vessel: obstruction of a cerebral blood vessel can cause a stroke; obstruction of a pulmonary vessel can cause pain and bloody sputum (12). The clinical features of myxomas, like most cardiac tumors, are determined by their location, size, and mobility. Most patients present with one or more features of the triad of embolism, intracardiac obstruction, and constitutional symptoms. Dyspnea may occur secondary to atriocentric valve obstruction. Other symptoms of myxomas include fever, weight loss, fingers and feet cold and sore when exposed to cold (Raynaud’s phenomenon), anemia, low platelet count (because platelets are involved in blood clotting), and symptoms suggestive of severe infection. Additionally, a myxoma can obstruct blood flow in the heart. Often they grow from a pedicle (resembling a mushroom) and are able to swing freely with the flow of the blood. This oscillating motion can obstruct blood flow through the mitral valve, the passageway between the left atrium and left ventricle. Non-cancerous less common cardiac tumors, such as fibroids and rhabdomyomas can grow directly from the cells of the fibrous tissue of the heart and myocardial cells. Rhabdomyomas, the second most common type of primary tumor, develop in childhood or early adolescence, often associated with a rare childhood disease called tuberous sclerosis. Other primary cardiac tumors, such as primary cancer, are extremely rare and, for them, there is no satisfactory treatment. Children with these tumors have a life expectancy of <1 year. Several tests are used in the diagnosis of cardiac tumors. Often, echocardiography (test that uses ultrasound waves to investigate the internal structures of the heart) is used for the delineation of tumors. The ultrasonic waves can pass through the chest wall or the inside of the esophagus transesophageal (10, 11).

A catheter inserted through a vein to the heart can be used to inject substances outlining a heart tumor on radiographs; but this procedure is less frequently needed. Computed tomography (CT) and magnetic resonance imaging (MRI) are also used. If a tumor is detected, a small sample can be removed through a special catheter; the sample will then be used to identify the type of tumor, which will help in selecting the appropriate treatment (11–13).

Echocardiography provides limited but useful information regarding tissue characteristics, including echogenicity of the mass and whether calcification is present. Vascularity can also be assessed using color flow Doppler and echocardiographic contrast. Strain imaging also has potential in identifying the non-contractile nature of masses such as fibromas. CT provides information regarding vascularity by contrast enhancement, presence of calcification, and presence of fat. Magnetic resonance imaging also provides information regarding vascularity, presence of fat, degree of tissue edema, and possibly iron content (14–23).

An isolated non-cancerous primary cardiac tumor can be removed by surgery, which generally cures the patient. The doctors do not treat primary tumors when there are many gifts, or treat tumors that are so great as to preclude removal. Although cardiac tumors are rare, they are being increasingly recognized ante-mortem, permitting earlier diagnosis and treatment. The most likely etiology of a cardiac mass is a thrombus or vegetation. In the current era, for benign cardiac tumors, an early diagnosis and appropriate treatment is not only possible but often curative.

REFERENCES
1. Shackell M, Mitko A, Williams PL, Stonn GC. Angiosarcoma of the heart. Br Heart J (1979) 41:498–503. doi:10.1136/hrt.41.4.498
2. Bjesoms S, Ivert T. Cardiac myxoma: 40 years. Experience in 63 patients. Ann Thorac Surg (1997) 63:697–700. doi:10.1016/S0003-4975(97)01368-9
3. Meira EBS, Ho CL, Santos-Filho MM, Pimenta J. Left atrial myxoma causing acute myocardial infarction. Arq Bras Cardiol (1996) 67:347–9.
4. Lehmann KL, Prozan GB, Ullnyt D. Atrial myxoma presenting as acute myocardial infarction. Am Heart J (1985) 110:1293–5. doi:10.1016/S0002-8703(85)90274-3
5. Colucci WS, Schoen JF, Braunwald E. Primary tumors of the heart. 5th ed. In: Braunwald E, editor. Heart Disease: A Textbook of Cardiovascular Medicine. Philadelphia, PA: WB Saunders (1998). p. 1464–78.
6. Goodwin JF, Symposium on cardiac tumors. The spectrum of cardiac tumors. Am J Cardiol (1968) 21:307–14. doi:10.1016/0002-9149(68)90135-5
7. Roberts WC. Primary and secondary neoplasm of the heart. Am J Cardiol (1997) 80:671–82. doi:10.1016/0002-9149(97)00587-0
8. Almeida EC, Leite MSB, Silva MA, Rassi L. Rhabdomyoma of the right ventricle causing pulmonary stenosis. Arq Bras Cardiol (1993) 60:417–9.
9. Ianni BM, Mady C, Pereira Barretto AC, Gutierrez PS, Lopes EA, Lemos PC, et al. Angiosarcoma of the heart. Reporting case. Arq Bras Cardiol (1986) 47:1–5.
10. Attar S, Lee Y, Singleton R, Scherli S, David R, Melanghells JS. Cardiac myxoma. Ann Thorac Surg (1980) 29:397–405. doi:10.1016/S0003-4975(10)80166-0
11. Maraj S, Pressman GS, Figueredo VM. Primary cardiac tumors. Int J Cardiol (2002) 133:152–6. doi:10.1016/j.ijcard.2008.11.103
12. Byard RW. Cardiac Conditions. Sudden Death in the Young. 3rd ed. Cambridge: Cambridge University Press (2010). p. 243–9.
13. Byard RW, Smith NM, Bourne AJ. Incidental cardiac rhabdomyomas: a significant finding necessitating additional investigation at the time of autopsy. J Forensic Sci (1991) 36:1229–33.
14. Byard RW. The potential significance of occult cardiac rhabdomyomas in traumatic death. Forensic Sci Med Pathol (2011) 7:267–8. doi:10.1007/s12024-010-9206-5
15. Barreiro M, Renilla A, Jimenez JM, Martin M, Al Musa T, Garcia L, et al. Primary cardiac tumors: 32 years of experience from a Spanish tertiary surgical center. Cardiovasc Pathol (2013) 22:424–7. doi:10.1016/j.carpath.2013.04.006
16. Burke A. Primary malignant cardiac tumors. Semin Diagn Pathol (2008) 25:39–46. doi:10.1053/j.sedp.2007.10.006
17. Reynen K, Kocheritz U, Strasser RH. Metastases to the heart. Ann Oncol (2004) 15:375–81. doi:10.1093/annonc/mdh086
18. Rassasi K, De Georgi F, Abbate A, Silvestre F. Cardiac metastases. J Clin Pathol (2007) 60:27–34. doi:10.1136/jcp.2005.035105
19. Makhija Z, Deshpande R, Desai J. Unusual tumours of the heart: diagnostic and prognostic implications. J Cardiovasc Pathol (2009) 14:4. doi:10.1186/1749-8090-4-4
20. Duband S, Paysant F, Scalon V, Forest P, Peoch M. Sudden death due to myocardial metastasis of lingual squamous cell carcinoma. Cardiovasc Pathol (2011) 20:242–3. doi:10.1016/j.carpath.2010.07.002
21. James CL, Byard RW, Knight WR, Rice MS. Metastatic osteogenic sarcoma to the heart presenting as bacterial endocarditis. Pathology (1993) 25:190–2. doi:10.1111/j.97809084798
22. Chen J, Huang TW, Hsu PS, Yang C, Tsai CS. Cardiac tamponade as the initial manifestation of metastatic adenoarcinoma from the colon: a case report. Heart Surg Forum (2007) 10:E339–30. doi:10.1532/HSF98.20071008
23. Beh P, Byard RW. Cardiac tumors and sudden death. Forensic Sci Med Pathol (2014) 10(2):269–71. doi:10.1007/s12024-013-9526-3
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