Atrial Myxoma Saddle Aortic Embolism: An Overview

Huthayfa Ghanem*, Sami Asfar2, Marzouk Albader3, Jassim Al-Ali3 and Ibrahim Hanbal4

1Senior Vascular Fellow, Bedford Hospital NHS Trust, UK
2Professor of Surgery and Senior Consultant Vascular and Hepatic Surgeon, Kuwait University Health Sciences Center, Kuwait
3Consultant Vascular Surgeon, Mubarak Al-Kabeer Hospital, Kuwait
4Professor of Vascular Surgery, Al-Azhar Faculty of Medicine, Egypt

*Corresponding author: Huthayfa Ghanem, Senior Vascular Fellow, Bedford Hospital NHS Trust, UK.

Abstract

Although they are rare, atrial myxomas are the most common primary cardiac tumours, to be followed by sarcomas. These tumours are benign and don't metastasize; however, they do embolize. Myxoma emboli usually go into the systemic circulation because of their usual location in the left atrium. Early diagnosis is usually challenging because of the non-specific symptoms. Therefore, it is not uncommon to have them presented with embolism, congestive heart failure or sudden cardiac death. Vascular surgeons are usually exposed to these tumours when the affected patients come to A/E department with acutely ischaemic limb(s) and / or strokes. Only after initial imaging and/or embolectomy, myxoma might be considered as a possible culprit. Once suspected, further detailed cardiac imaging must be done urgently to confirm the diagnosis in order to proceed with tumour resection at the earliest, considering myxoma tumour propensity for recurrent and multiple embolization with the subsequent high risk of morbidity and mortality. Hence, a high index of suspicion, early diagnosis and excision of these masses are vital to avoid the adverse clinical outcome, taking into account the excellent prognosis after resection [1-4].

Keywords: Left Atrium; Myxoma; Myxomectomy; Saddle Embolus; Acute Limb Ischaemia

Introduction

In 1952, myxoma was first diagnosed by Goldberg [5]. These tumours account for up to 50% of primary cardiac tumours. Around 80% of them occur in the left atrium, usually at the border of fossa ovalis in the interatrial septum. The estimated incidence is 0.5 per million population per year; however, the incidence at autopsy is 0.02%. Most of these tumours are sporadic. Nonetheless, 10% of myxomas have an autosomal dominant inheritance such in the case of Carney syndrome.

Myxomas can occur at any age; however, they are most commonly seen between the third and sixth decade, and the mean age of presentation was 56 in sporadic cases, 25 in familial cases and 37.1 overall. There is female preponderance of 2-3:1 [6-9].

The most common symptom is dyspnoea (75%). Constitutional symptoms due to IL-6 production by the tumour such as arthralgia, fatigue, cyanosis, clubbing, weight loss, rashes and Raynaud’s phenomenon occur in 50%, dizziness and syncope occur in 20%, and haemoptysis in up to 15% of patients. Only in around 20% of cases, myxomas can be asymptomatic or discovered incidentally [7, 8].

More than 50% of the patients with left atrial myxoma have clinical evidence of mitral valve dysfunction, with systolic and diastolic murmurs on auscultation. Large left atrial tumours produce diastolic sound “tumor plop” because of obstruction to ventricular in-flow (mechanical interference). In some patients, these tumours can cause arrhythmias [10, 11].

Features of embolism can be seen in any vascular bed in up to 40% of myxomas, commonly affecting arteries of CNS (50% of embolic events) and retinae. Furthermore, 10% of coronary embolic events are thought to be due to myxomas. Right sided myxomas can embolize to pulmonary arteries or cause trans-septal paradoxical embolism [12,13].

The embolus can be: [14]

a) A fragment of the tumour
b) The whole tumour
c) An associated thrombus attached to the tumour (difficult to diagnose)
The following features are associated with higher embolism potentials: [15-18]

1. Size: 4.5 cm or less (more aggressive tumours with IL-6 production)
2. Shape: polypoid multilobed villous tumours
3. Consistency: 35% of myxomas are soft and friable
4. Location: left atrial and aortic valve tumours
5. Mobility: intermittently prolapsing masses
6. Growth rate: higher growth rates are associated with greater embolic likelihood (The reported growth rates vary from no growth, to between 1.3 to 6.9 mm/month). Growth rates are usually faster in recurrent cases.
7. Embolized tumours will usually embolize again concurrently.
8. Function: IL-6 production
9. Symptoms: less symptomatology is associated with higher risk
10. Associated atrial tachyarrhythmias
11. Mitral valve destruction
12. High platelets count and mean platelets volume
13. Associated thrombophilia

Morphologically, myxomas appear as yellowish, glistening, gelatinous, polypoid, smooth, lobulated or villiform, sessile or pedunculated masses usually 5 – 6 cm in diameter and, with a short stalk. Sometimes, the tumour can be large and mobile enough to cause valve obstruction [14] Figure 1. Microscopically, they show loose myxoid stroma containing proteoglycans. Polygonal stellate cells are found within the matrix, singly or in small clusters [14].

Five Cases with Cardiac Myxoma Saddle Aortic Embolism

Over a period of 10 years (2007-2017), 5 of the patients with acute bilateral lower limb ischaemia presented to Mubarak Al-Kabeer Hospital found to have left atrial myxoma saddle aortic embolism. All were otherwise healthy females and the mean age was 41.4 years (37 – 59). The average duration of symptoms to presentation in A/E department was 5.2 hours (4 – 11). Atrial myxomas were diagnosed by post-embolectomy echo scan in 3 and by histology in all patients. The left atrial myxoma residual average diameter was 3.7 cm. All the 3 ladies with left atrial myxomas had excision after an average duration of 9 hours post embolectomy (6 – 15). Histology confirmed myxoma cardiac residuals in all examined masses. Post-operative period was uneventful in all patients. No recurrences detected with at least 2-year follow up duration.

During the same period, a total of 104 saddle aortic embolisms operations were performed, and the average duration of symptoms was 7.1 hours (2 – 19).

Therefore, cardiac myxoma saddle aortic embolism accounted for 4.8% of all saddle aortic embolism (less than 0.3% in peripheral embolism). In addition, those patients tend to present earlier than other saddle aortic embolism patients.

According to our experience, the typical patient is:

a. Young female
b. Previously healthy (no arteriopathy, no atrial fibrillation)

Often, presents with multiple organ embolism. Less frequently, symptomatic myxomas presents with embolic events.

Acute ischaemia due to cardiac myxomatosa saddle embolism is usually severe (Rutherford stage II: limb threatening ischaemia). Embolus volume is usually higher in myxomas than other forms of embolism. The typical clinical scenario is a patient with the above-mentioned characteristics comes to A/E department with features...
of acute bilateral lower limb ischaemia after only a few hours of onset of symptoms (earlier than with other causes of acute limb ischaemia) due to profound symptoms and massive embolism. Classically, those patients will have bilateral femoral embolectomy after an intravenous bolus of heparin directly without having pre-operative scans. Intra-operatively, the retrieval of yellowish jelly like material should be alarming enough to justify arrangements for an echocardiography (preferably trans-oesophageal) in the immediate post-operative period (bedside). Once confirmed, prompt cardiac surgery input is mandatory to clear the cardiac residuals as soon as possible. Otherwise, the new embolism risk is exceedingly high and is usually fatal. It is not uncommon to have unremarkable echo scans after embolectomy due to embolization of the whole tumour; hence, tissue diagnosis will be the only way to confirm the diagnosis. Moreover, in such cases it is highly advisable to arrange for regular echo scans (possibly biannually) as the risk of having tumour cells left at myxoma bed in fossa ovalis cannot be omitted.

Cardiac myxomectomy for sporadic tumours is usually curative. Tumour manipulation should be done only after cardioplegia as the risk of fragmentation and embolism is usually there. Tumour excision along with its stalk and septal bed is recommended with pericardial patch septoplasty. Inspection of all champer is omitted.

Surgical risks:

a. Early mortality: 2%
b. Post-operative AF: 30%
c. Neurologic complications: 3%
d. Re-exploration: 5% and usually due to bleeding [21].

Recurrences (up to 5% in sporadic tumours and more than 20% in familial cases and is uncommon after 5 years) are usually occurred due to:

a. Incomplete excision
b. Other foci
c. Implantation of tumour cells (intra- or extracardiac) [22].

There is no current guidelines or recommendations exist regarding the treatment of embolic manifestations; however, anticoagulation is recommended if associated with:

- AF
- Some cases of dysrhythmia or valve destruction [23].

Conclusion

The incidence of atrial myxoma in saddle aortic embolism cases seems to be higher than its incidence in peripheral embolism in general. In other words, atrial myxoma embolism to the lower limbs is usually of the saddle type. This condition has to be considered in “typical patients” presented to A/E department with acute bilateral lower limb ischaemia, as the prompt diagnosis and timely intervention will significantly improve the clinical outcome.

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Conflict of Interest

No conflict of interest.

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