Cardiac tumours simulating collagen vascular disease

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SUMMARY Cardiac tumours can mimic collagen vascular disease and they are often accompanied by profound systemic upset. Both benign and malignant tumours may present in this way. Three cases of cardiac tumour, two malignant and one benign, are reported with just such a presentation. A review of fifteen similar case reports showed that a spectrum of different collagen vascular diseases was diagnosed and treated before the true diagnosis emerged. In half of these cases the cardiac tumour was only diagnosed at necropsy. The diagnosis of collagen vascular disease should not be made in the absence of corroborative laboratory data. In cases of malignant cardiac tumour, and less commonly with atrial myxoma, M mode and cross sectional echocardiography may not exclude the diagnosis. There may be a good response to steroid treatment in cases of suspected but not confirmed collagen vascular disease in which the true diagnosis is cardiac tumour.

The early diagnosis of cardiac tumours is important because most are benign and may be amenable to curative resection. Fifty per cent are myxomas, commonly in the left atrium, and the rest are a wide variety of different histological types, benign and malignant. Initial misdiagnosis is common, and it was only as recently as 1951 that the first myxoma was correctly diagnosed and successfully treated.1

Cardiac tumours may present in three ways. Symptoms may be due to haemodynamic effects, embolic events, or systemic upset.2 Left atrial myxomas are often attached by a pedicle to the atrial wall in the region of the fossa ovalis and may interfere with the function of the mitral valve, and occasionally they have been discovered at operation for mitral stenosis. Myxomas are friable and gelatinous and readily fragment giving rise to microemboli and macroemboli; indeed cardiac myxomas have been diagnosed on several occasions after arterial embolectomy.3 Finally, in a number of cases constitutional upset dominates the clinical presentation. In some of these patients a collagen vascular disease is diagnosed and they are treated with corticosteroids. We describe three such cases; in each case inappropriate treatment was started and correct diagnosis delayed. We have reviewed fifteen similar patients and discuss some principles in the diagnosis and management.

Case reports

CASE 1

A 32 year old woman presented in the twentieth week of pregnancy with right sided chest pain and exertional dyspnoea. She gave a history of transient erythematous rashes, fleeting arthralgia, and Raynaud’s phenomenon. Initial examination showed no abnormality but symptoms persisted and 12 weeks later she developed ankle oedema and tender hepatomegaly. Cardiac auscultation was normal but the jugular venous pulse was increased by 4 cm. There was normochromic, normocytic anaemia, a normal white blood cell count, and thrombocytopenia of $80 \times 10^9$. The erythrocyte sedimentation rate was 93 mm/h. The serum albumin concentration was 25 g/l and globulin fraction 31 g/l. Proteinuria was noted. An electrocardiogram showed small voltage QRS complexes and flattened T waves in the precordial leads. An echocardiogram showed a pericardial effusion but no other abnormality.

A diagnosis of systemic lupus erythematosus was made on the basis of her history and the pericarditis, anaemia, thrombocytopenia, and presumed nephritis. Her condition deteriorated and she was treated with intravenous methylprednisolone, followed by oral prednisolone. Percardiocentesis removed 2.5 l of blood stained fluid, and culture and cytological examination were negative. Steroid treatment was continued and her erythrocyte sedimentation rate fell to 30 mm/h and the haemoglobin concentration rose from 9.8 g/l to 12 g/l. The raised jugular venous

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Cardiac tumours simulating collagen vascular disease

pressure persisted despite pericardiocentesis and an isotope lung scan was performed. This suggested multiple pulmonary emboli and the patient was anticoagulated.

A search for antibodies to nuclear antigens (ANA), extractable nuclear antigens (ENA), and DNA proved negative, raising doubts about the diagnosis. Moreover, 24 hour urinary protein excretion was only 150 mg. A total of six M mode and cross sectional echocardiograms were performed, but only the last of these showed the presence of a mass in the right atrium. A computed tomographic scan of the heart and mediastinum confirmed this finding.

An exploratory thoracotomy showed an extensive infiltrating mass within the pericardium. No attempt was made to remove the tumour. Two days after the operation the patient delivered a 28 week old fetus which survived, but the patient died shortly afterwards. Necropsy showed that the right atrium and part of the great vessels were encased in solid tumour. Histological examination of the tumour indicated that it was a pleomorphic rhabdomyosarcoma. There was no evidence of metastatic spread.

CASE 2

A 23 year old woman presented with a left hemiparesis and a history of five previous episodes of transient left sided weakness associated with headache. Computed tomography suggested cerebral infarction in the region of the basal ganglia. A month later she developed recurrent haemoptyses and pleuritic chest pain. Clinical examination and chest radiography showed a left pleural effusion. A ventilation-perfusion isotope lung scan indicated a pulmonary embolus and she was anticoagulated.

Despite treatment she continued to have headaches and episodes of transient loss of vision. She complained of migratory joint pains and weight loss with persistent fever, and normochromic normocytic anaemia developed. Tests of renal and hepatic function remained normal but tender hepatomegaly and proteinuria were detected. Carotid angiography showed occlusion of the right internal carotid artery. Pulmonary angiography was normal. M mode and cross sectional echocardiography failed to show any abnormality. Systemic vasculitis was diagnosed and she was treated with prednisolone, azathioprine and cyclophosphamide for six weeks.

Ten months after her initial presentation a further echocardiogram was obtained because she had developed a soft third heart sound and a short diastolic murmur. This showed a left atrial mass. Thoracotomy was undertaken and an invasive left atrial angiosarcoma was discovered. The patient died during the operation.

CASE 3

A 40 year old woman presented with malaise, recurrent headache, and visual disturbance. Migraine was diagnosed but both propranolol and clonidine were ineffective as prophylaxis. She developed pain and stiffness in the legs which were accentuated by exercise. There were episodes of regular palpitation but no other cardiovascular symptoms.

General physical examination showed no abnormality. The erythrocyte sedimentation rate was 65 mm/h, and plasma creatine kinase concentration was grossly raised (1000 units/l). Electromyography showed a slight excess of polyphasic action potentials. A skeletal muscle biopsy specimen showed some type II fibre atrophy but no evidence of inflammation. Nevertheless, the clinical picture was thought to support a diagnosis of polymyositis and treatment with oral corticosteroids was started. Despite this, her symptoms persisted and when she was followed up six months later there was an unexplained sinus tachycardia. A cross sectional echocardiogram was performed which showed a tumour in the left atrium. A benign myxoma was subsequently resected and the patient made a complete recovery.

Discussion

Patients with cardiac tumours may have no cardiovascular symptoms or signs but there can be prominent systemic disturbance. The clinical picture may include fever, weight loss, and anaemia, and often there will be a raised erythrocyte sedimentation rate and serum globulin concentrations. Constitutional features often accompany cardiovascular disturbances. Sometimes the absence of the latter may lead to an initial misdiagnosis of collagen vascular disease. There are two reasons for this. Firstly, the clinical diagnosis of systemic vasculitis is not frequently based on non-specific criteria. After the exclusion of other causes of multisystem disease, for example infection and sarcoidosis, it may sometimes be necessary to institute treatment for vasculitis in the severely ill patient with an undiagnosed disorder, before histological confirmation can be obtained. Vasculitis can be difficult to prove, however, if the tissues affected are not amenable to biopsy.

Other clinical features that lead to the mistaken diagnosis of connective tissue disease or vasculitis in patients with cardiac tumours include Raynaud's phenomenon, rash, pericarditis, and arthritis. In the context of a multisystem illness such features strongly suggest a collagen vascular disease. Table 1 lists clinical features and their prominence in fifteen cases that were similar to the three we have reported above.
When clinical features in patients with occult cardiac neoplasms suggest collagen vascular disease, laboratory findings seldom support the diagnosis. Table 1 lists some of these. Non-specific abnormalities, such as a raised erythrocyte sedimentation rate, anaemia, and raised serum globulin concentrations, are common.

Only one patient had a positive antinuclear antibody test (with a titre of 1/64), however, and tests for lupus erythematosus cells, rheumatoid factor, and complement depletion were consistently normal. The absence of supporting laboratory evidence should always cast doubt on the diagnosis of connective tissue disease or vasculitis. There has been considerable interest in the clinical entity “antineuclear antibody-negative lupus” but in such suspected cases antibodies to Ro and La antigens (“SS-A and SS-B”—antibodies commonly present in Sjögren’s syndrome) should always be sought and may help to establish the diagnosis.17

Table 2 summarises fifteen similar cases of cardiac tumour simulating collagen vascular disease. With the exception of a single case of lymphoma all were benign myxomas. Cases 1 and 2 reported above, however, indicate that despite quite distinct histopathological features, malignant primary cardiac neoplasms may present in a similar way. Myxomas tend to be friable, gelatinous tumours and some of their constitutional effects may be explained by micro-embolic events. Nevertheless, if solid, sessile, malignant tumours have similar clinical presentations, some other pathogenesis must exist, presumably immunological.

Immunological mechanisms have been suggested, and antibodies to cardiac muscle have been sought and found in a single case,13 although the significance of this in isolation is unknown.

Detection of these tumours at an early state is important. Table 2 shows that of fourteen previously reported cases of myxoma, initially diagnosed as collagen vascular disease, eight died and the correct diagnosis only came to light at necropsy.

With malignant cardiac tumours early detection is important. Despite their proximity to the circulation such tumours tend to be confined to the heart until relatively late in their course. There have been cases
Cardiac tumours simulating collagen vascular disease

of successful cardiac transplantation for primary cardiac tumours.1

In patients with a non-infective multisystem disease in which serological and autoimmune testing are negative, a cardiac tumour should be considered. Cross sectional echocardiography will usually detect myxomas cheaply, reliably, and quickly. The absence of an obvious myxoma, however, does not rule out the possibility of a myocardial neoplasm. As our report shows, repeated false negative results may occur with malignant infiltrating tumours. If the diagnosis is seriously entertained, the tumour may need to be pursued with other investigations, such as computed tomography or nuclear magnetic resonance, despite a normal echocardiogram.

These cases illustrate the importance of considering cardiac neoplasms in the differential diagnosis of multisystem disease. Moreover, they emphasise the need for an intensive search for supporting histological and laboratory data in patients diagnosed as having connective tissue disease or vasculitis.

References

Cardiac tumours simulating collagen vascular disease.

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