CASE REPORT

Pericarditis with anaemia as a herald syndrome in a fatal presentation of cardiac lymphoma

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SUMMARY
Primary cardiac lymphoma (PCL) is rare, accounting for 2% of all primary cardiac malignancies. Diagnosis is sometimes slow due to the non-specific nature of symptoms, causing a delay to treatment with potentially curative anthracycline chemotherapy. We report an unusual presentation of primary cardiac lymphoma in an immunocompetent man presenting with subacute isolated right-sided heart failure with pericarditis on a background of chronic anaemia and constitutional upset. Echocardiography demonstrated a pericardial mass invading the right atrium and compressing the tricuspid annulus. Diffuse large B-cell lymphoma was diagnosed after biopsy. This case highlights the importance of early imaging and hospitalisation in pericarditis with high-risk features such as high inflammatory markers, myocardial involvement (with troponin elevation), fever, immunosuppression or evidence of heart failure. The differential and diagnostic pathway of an intracardiac mass, and the treatment and prognosis of PCL, are discussed.

BACKGROUND
Pericarditis is a common clinical presentation. It can be idiopathic, or secondary to trauma, autoimmune aetiology, infection or uraemia. Similarly, iron deficiency anaemia is also a common finding in clinical practice and should prompt investigations to exclude malignancy in the older population. Such investigations are often limited to colonoscopy and gastroscopy. A complex constellation of clinical findings, such as pericarditis, iron deficiency anaemia and systemic symptoms, not resolving with initial management, should prompt the clinician to think of other, more unusual, pathologies. Primary cardiac lymphoma (PCL) is a rare form of cardiac neoplasm, accounting for only 2% of all cardiac tumours.1 Although usually presenting inconspicuously, with non-specific symptoms of fever and malaise, making it difficult to diagnose, symptoms due to compression by a tumour mass and catastrophic systemic embolisation can also occur.

An intracardiac mass identified at initial imaging (usually echocardiography or CT) is often the first clue to a diagnosis of PCL. Subsequent histological diagnosis via tissue biopsy allows treatment (which may include chemotherapy and/or surgical resection) to be planned. Timely intervention improves the outcome of PCL, which otherwise carries a poor prognosis.

CASE PRESENTATION
A 65-year-old man presented with a 5-month history of general malaise, dyspnoea, fever and anorexia. These symptoms were thought to be secondary to a new iron deficiency anaemia (iron 8.5 µmol/L, transferrin 1.88 g/L, transferrin saturation 18%), which had prompted gastroscopy and colonoscopy 2 months previously. These investigations were normal, therefore ruling out common gastrointestinal pathology. The patient now described a three-day history of sharp central chest pain with relief sitting upright, bilateral leg oedema, episodes of presyncope relating to standing and dyspnoea graded at New York Heart Association class III prompting cardiology referral. On examination, his blood pressure was 90/60, his jugular venous pressure was raised and he had ankle oedema bilaterally. In addition, he had a pericardial rub. ECG demonstrated atrial fibrillation with ventricular rate 100 bpm and chest radiograph was unremarkable. Blood work demonstrated an inflammatory response with elevated C reactive protein (185 mg/L) and an erythrocyte sedimentation rate of 73 mm/h. Troponin I was significantly raised at 0.85 μg/L. Autoantibodies and immunoglobulins were within normal limits, and lactate dehydrogenase was mildly elevated at 313 U/L.

Transthoracic echocardiography (TTE) demonstrated an intrapericardial mass compressing the tricuspid annulus, with invasion of the right atrium (figure 1). Right ventricular function was severely impaired, while left ventricular function was preserved. MRI characterised the intrapericardial mass as a large cardiac tumour (10×7 cm; figure 2).

Figure 1 Echocardiogram (subcostal four-chamber view) demonstrating a large pericardial mass (1); compressing the right atrium and tricuspid valve annulus (2); with evidence of invasion of the right atrium (3). A small pericardial effusion is also noted (4).
encasing the right ventricle up to the aortic arch and down to the inferior vena cava (video 1), with pedunculated masses seen in the right atrium and severe compression of the tricuspid valve (figure 2). The TTE (figure 1) and MRI (figure 2) both demonstrated a small pericardial effusion. Interestingly, the tumour extension also caused encasement of the right coronary artery (figure 3 and video 2). The patient was started on dexamethasone, with clinical improvement. Endomyocardial biopsy yielded high-grade lymphoid cells (figure 4), and immunohistochemical staining showed these cells were positive for CD20 and bcl-2 (figure 5), consistent with a diagnosis of diffuse large B-cell lymphoma. Unfortunately, the patient deteriorated due to right heart failure developing shock refractory to inotropic support, which proved fatal.

DISCUSSION

PCL is rare, accounting for 2% of all primary cardiac neoplasms.1 The commonest affected sites are the right atrium, right ventricle and the pericardium.2 Symptoms relate to tumour extension and/or embolism, with dyspnoea, chest pain

Figure 2  Steady-state free precession (SSFP) MRI scan demonstrating the large pericardial mass (1); compressing the right atrium and tricuspid valve annulus (2). The small pericardial effusion is also noted (3).

Figure 3  Single frame from short axis video at the atrioventricular valves showing encasement of the right coronary artery (green arrows).

Figure 4

Figure 5

Video 1 Short axis mid-left ventricle video showing tumour encasement of the right ventricle with diastolic dysfunction and delayed right ventricular filling.

Video 2 Short axis video at the level of the atrioventricular valves showing tumour encasement of the right coronary artery and invasion of the right atrium.
Unusual association of diseases/symptoms

Figure 4  Section from the tumour mass demonstrating infiltration by sheets of medium to large sized lymphoid cells featuring round nuclei with small nucleoli and scanty amounts of cytoplasm. The H&E (×40) stain confirms a high-grade lymphoma with high rates of mitosis and apoptosis.

Figure 5  The sample stains positively for CD20 (×20), confirming the lymphoma to be a large B-cell lymphoma.

and constitutional symptoms being the commonest presenting symptoms, occurring in 66%, 26% and 24% of patients, respectively.2 3 Anaemia is a common finding in lymphoproliferative disorders4 such as PCL, and the pathophysiology is thought to be related to abnormal iron metabolism.5 Our patient had progressive anaemia with no evidence of gastrointestinal blood loss; however, diagnosis was only made after chest pain developed. PCL usually presents after the fifth decade of life.6 B-cell lymphoma is the commonest type of PCL and is rapidly progressive, with median survival of less than 1 month without chemotherapy.7 Poor prognostic markers of PCL include left ventricular involvement, immunocompromise and the presence of extracardiac disease.7 In addition to these, anaemia itself is a poor prognostic marker of lymphoma for both overall and progression-free survival,4 therefore its early recognition alone may play a powerful role in reducing mortality.

Making a diagnosis of PCL is often slow, which is to some extent unavoidable given that symptoms such as fevers, anaemia and breathlessness will clearly not prompt echocardiography or thoracic CT prior to more basic investigations. However, with better access to advanced cardiac imaging when requested, the diagnosis is being made earlier with more patients being deemed suitable for receiving chemotherapy with associated survival extending to 30 months.7 Surgical excision and radiotherapy offer less clear benefit, although selection bias may be at play, given that those presenting moribund are less likely to be offered chemotherapy. Our case is testament to the aggressive nature of this disease; despite timely histological diagnosis, the patient passed away within days.

Nearly all the published cases are first identified as right-sided masses at TTE, it thus seems sensible to postulate that early echocardiography affords the patient the best chance of being diagnosed when still a candidate for treatment. The difficulty here is that the denominator is huge. As discussed, cases commonly present with non-specific symptoms of breathlessness, fever, weight loss and lethergy—symptoms that are often benign and investigated routinely as an outpatient. How does the physician pick out the needle from the haystack in deciding who needs early access to an echocardiogram? Rapidly progressive symptoms not responding to initial treatments, or, more critically, a constellation of symptoms (seen in this case as anaemia, chest pain, breathlessness and peripheral oedema) occurring sequentially, should prompt echocardiography to rule out a mass or vegetation.

The finding of an intracardiac mass has extensive differential diagnoses, including the more common primary cardiac neoplasms of myxoma (up to 50% of primary cardiac tumours), angiosarcoma, metastatic deposit and thrombus.8 Therefore, diagnosis of PCL requires further imaging with transoesophageal echocardiography, cardiac MRI, CT and positron emission tomography scanning, leading to endomyocardial biopsy.9 Only when this histological (or, more rarely, cytological via pericardial fluid) diagnosis is obtained, can the patient be considered for chemotherapy or surgery. Right-sided tumours are particularly difficult to debulk, however, and the role of surgery is limited, with the mainstay of treatment being an anthracycline-based chemotherapy regimen of cyclophosphamide, doxorubicin, vincristine and prednisolone (R-CHOP).10

We present an unusual case of PCL in an immunocompetent patient, who initially reported of a constellation of common clinical symptoms ascribed to a new iron deficiency anaemia. The case report highlights the importance of revisiting the extended differential diagnosis when regular diagnostic workup for common clinical presentations (in this case iron deficiency anaemia) yields no diagnosis. In addition, we highlight the importance of early imaging and hospitalisation in pericarditis with high-risk features, such as high inflammatory markers, myocardial involvement, fever and evidence of heart failure.11

Learning points

- Pericarditis is idiopathic in 80–90% of cases, and can be safely managed as an outpatient. Features such as fever, high inflammatory markers, a subacute onset, immunosuppression, myocardial involvement (with troponin elevation) and heart failure, point to an increased risk of complications and warrant hospitalisation for early echocardiography and a full aetiological search.11
- The differential diagnoses of an intracardiac mass are extensive and include treatable causes—anticoagulation for thrombus, excision for myxoma and chemotherapy for lymphoma. The key is early diagnosis, requiring complex imaging and, often, endocardial biopsy.
- Primary cardiac lymphoma is rare and frequently fatal; however, if diagnosed early enough, it is a treatable and sometimes even curable disease.
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