Diagnosis of tuberculous aetiology in pericardial effusions

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The diagnosis of tuberculous aetiology in pericardial effusions is important since the prognosis is excellent with specific treatment. The clinical features may not be distinctive and the diagnosis could be missed particularly with tamponade. With the spread of HIV infection the incidence has increased. The diagnosis largely depends on histopathology of the pericardial tissue or culture of Mycobacterium tuberculosis from this tissue or fluid, but patients without haemodynamic compromise do not require pericardiocentesis. Histopathology may, however, show non-specific findings in a significant number. This review is an update on the diagnostic difficulties, current research, and criteria for diagnosis.

Tuberculous pericardial effusion (TPE) is still common in African and Asian countries, but with migrant populations and the spread of AIDS there are now more frequent reports from other parts of the world like Europe and the Western hemisphere. TPE usually presents as a slowly progressive febrile illness. When it presents as an acute pericarditis, which is uncommon, or as cardiac tamponade, which is frequent, the diagnosis is more likely to be delayed or missed. The delay from hospital admission to diagnosis was 5.2 weeks in a report from Spain and in another from the USA the diagnosis was first made only at necropsy in 17% of patients. Malignant, viral, and chronic idiopathic effusions are some of the other causes of large effusions. Effusions associated with malignancy are usually but not always apparent at presentation. Chronic idiopathic effusions in which no aetiology could be established are a common cause of tamponade even in countries like the UK, France, and the USA with a reported incidence varying from 11%–32%.

The diagnosis of tuberculosis as the aetiological cause is important. Without specific treatment the average survival was 3.7 months in a report from Africa and only 4/20 (20%) were alive at six months. On the other hand the prognosis with appropriate medical treatment is excellent as seen in recent reports. The confirmation of tuberculous aetiology currently depends on histopathological study of the pericardium, the demonstration of tubercle bacilli in the pericardial tissue or pericardial fluid, the presence of proved tuberculosis elsewhere in the body, or the response to specific treatment. Histopathological examination, however, can at times give non-specific findings even when tubercle bacilli have been cultured from the pericardial fluid.

PATHOGENESIS OF TUBERCULOUS PERICARDITIS

There are many ways in which the pericardium may be involved in tuberculosis. In a rare case there may be direct spread from tuberculous pneumonia. The pericardium can be seeded in miliary tuberculosis and in such instances other organ systems dominate the presentation. Direct extension from an infected visceral pleura or rib is rare. Most often the spread is from the breakdown of infection in mediastinal nodes directly into the pericardium and particularly those at the tracheobronchial bifurcation. The spread is over lymph channels that merge at points where the parietal pericardium and the pleura separate.

LYMPHATIC DRAINAGE OF THE PERICARDIUM

Studies in human cadavers, in the macaque monkey, and in the dog show that the lymphatic drainage of the pericardium is mainly to the anterior mediastinal, tracheobronchial, hilar, and posterior mediastinal (juxta oesophageal) lymph nodes and not into the hilar nodes. This has an important bearing on the routes of mediastinal lymph nodes that are enlarged in TPE. The mediastinal node enlargement in TPE does not show up on routine chest radiographs (fig 1) but can be seen only on chest computed tomography or magnetic resonance imaging (MRI) studies. In other conditions associated with pericardial effusion and mediastinal adenopathy like lymphomas, malignancy, and sarcoid hilar node involvement is prominent even though in sarcoidosis the pericardium is involved in fewer than 3% of patients.

MEDIASTINAL LYMPH NODES IN TPE

Sir William Osler from his review of consecutive necropsies found 275 with tuberculosis of whom seven had pericarditis. He concluded that caseous mediastinal lymph nodes were the usual focus of pericardial involvement. In another report, at necropsy all patients with TPE had mediastinal adenopathy. The groups of glands involved were not described. In another study, right scalene node biopsy was performed in six patients with TPE and even though the node was not enlarged, all six had evidence of tuberculosis.

Abbreviations: MRI, magnetic resonance imaging; PCR, polymerase chain reaction; TPE, tuberculous pericardial effusion
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The adenopathy clears or regresses on specific therapy. 12 24 the computed tomogram as coalescing of adjacent glands. 29

CD4 + there is a broad clonal heterogeneity of antigen specific will be widely applied. A more recent report suggests that microscopy for acid fast bacilli. A sensitivity of 61% (at 96% was used. All but one of the patients had negative sputum of PCR approached the results of conventional methods, PCR was often faster. The sensitivity for pericardial fluid of PCR approached the results of conventional methods, although PCR was faster. The sensitivity for pericardial fluid was poor and false positive results with PCR remain a concern.27

**Pleural effusion and peripheral lymphadenitis**

Pleural effusion occur as a result of tuberculous effusion. Tuberculous lymphadenopathy is characterised by caseation, which shows on computed tomographic studies as hypodense central areas,27 and “matting” on palpation, which is seen on the computed tomograms as coalising of adjacent glands. The adenopathy clears or regresses on specific therapy.13 24

**RECENT DEVELOPMENTS IN THE DIAGNOSIS OF TPE**

**Polymerase chain reaction (PCR)**

PCR technology has been used for nucleic acid amplification in the diagnosis of tuberculosis.25 Cegielski et al compared PCR, culture, and histopathology in the diagnosis of TPE.26 PCR was performed with both pericardial fluid and tissue with IS6110 based primers specific for the *Mycobacterium tuberculosis* complex. They concluded that the overall accuracy of PCR approached the results of conventional methods, although PCR was faster. The sensitivity for pericardial fluid was poor and false positive results with PCR remain a concern.27

**Serodiagnosis**

The potential for serodiagnosis of TPE has been reported by Ng et al who tested a solid phase antibody competition sandwich ELISA (SACT-CE) method.28 A monoclonal antibody (CDC/WHO reference number IT39) which was raised against a specific epitope on the *M tuberculosis* 30 kDa antigen was used. All but one of the patients had negative sputum microscopy for acid fast bacilli. A sensitivity of 61% (at 96% specificity) was achieved. It is unlikely that this technology will be widely applied. A more recent report suggests that there is a broad clonal heterogeneity of antigen specific CD4+T-cells localising at the site of disease during tuberculosis.29

**Adenosine deaminase and other markers**

Adenosine deaminase levels are believed to reflect T-cell activity. The levels with TPE have varied from 10–303 U/l in one report,10 and with a cut off level of 30 U/l the sensitivity was 94% and specificity 68% with a positive predictive accuracy of 80%. In this study the median interferon-gamma concentration in TPE was ≥1000 pg/l and significantly higher than malignancy or non-tuberculous effusions (p<0.0005). A cut off value of 200 pg/l for interferon-gamma resulted in a sensitivity and specificity of 100% for the diagnosis of TPE. In another report the adenosine deaminase values were a mean (SD) of 126 (16.68) U/l and significantly higher than neoplastic, idiopathic, purulent bacterial, and radiotherapy groups.11 On the other hand it has been suggested adenosine deaminase and lysozome levels have to be taken into account in attempting an early diagnosis of TPE.12

**Nuclear imaging**

Gallium-67 and indium-111 scintigraphy have been used in the diagnosis of TPE. The results are, however, non-specific and there are other cardiac causes of gallium-67 uptake.33 34

**Chest computed tomography**

Recent observations on chest computed tomography regarding the detection of mediastinal lymph nodes in TPE and their value in diagnosis of TPE are discussed later.12 Computed tomography also allows study of the pericardium and the pleural changes (fig 2).

**EVALUATION OF TUBERCULOUS PERICARDIAL EFFUSION**

**Clinical features**

Asymptomatic presentation or small effusions are infrequent unless picked up during screening during the course of an illness like AIDS. The onset is most often insidious with loss of weight and fatigue,1 7 but could be more acute and occasionally explosive with tamponade.12 The presenting symptoms appear to be similar all over the world. Fever (73%–97%) and dyspnoea (80%–88%) are the most frequent symptoms while precordial pain (39%–59%) is seen less often.1 7 12 15 Pericardial rub is present in (37%–84%)12 and rarely can even be triphasic.12 Raised jugular venous pressure is frequent and pulsatil paradoxus is present in (23%–71%) with tamponade (box 1).15

Peripheral lymphadenopathy with matting which most often affects the cervical glands has been reported in 13%–28% with TPE1 and biopsy of the gland is diagnostic. This is unlikely to be seen in Western countries as it is a late manifestation.

**Cardiac tamponade**

Cardiac tamponade is a common complication of TPE. Jain et al found that 60% of patients with tamponade had tuberculosis.7 In another study of 44 children with TPE, 90% had tamponade.15 In a recent report from the USA both patients reported had tamponade16 and in an earlier report 6/11 deaths were due to tamponade.7 Since tamponade may

![Figure 1](http://pmj.bmj.com/)

**Figure 1** Chest radiograph after aspiration of same patient as in fig 4 taken at time of chest computed tomography. Nodes not seen on chest radiograph but seen only on chest computed tomogram.

![Figure 2](http://pmj.bmj.com/)

**Figure 2** Pre-treatment chest computed tomogram after aspiration showing irregular pericardial thickening (P) and some fluid (F). Right (RPE) and left (LPE) pleural effusion with strands and loculation (LS) in pleural space.
in part be due to late presentation, the incidence is likely to be higher in Afro-Asian countries. 12

HIV infection
Tuberculosis has been reported as the predominant cause of large pericardial effusions in HIV infected patients in Tanzania.37 These patients were in an early stage of HIV infection and responded well to treatment. In an inner city hospital in the USA, out of 122 patients with pericardial effusion of varying sizes, 40 were HIV related and accounted for 20% with tamponade. 3

Chest radiography
Apart from the signs of pericardial effusion, radiological evidence of pulmonary tuberculosis may be present.17 Pleural effusions are present in about 50% because of tuberculous pleuritis,7 and a tuberculous granuloma on pleural biopsy can help in a specific diagnosis.

Chest computed tomography
Chest computed tomography can yield valuable information in addition to the pleural changes (fig 2). The pericardium is irregular, not calcified, and is thickened measuring a mean (SD) of 4.4 (1.9) mm.12 Normal mediastinal lymph nodes are <10 mm and do not change in size on follow up.39 Enlarged mediastinal lymph nodes >10 mm detected on chest computed tomography have been reported recently (figs 3 and 4) and for the first time in virtually 100% of patients with TPE.12 They were found in all 22 patients with TPE and none of a control group with large viral/idiopathic or postoperative pericardial effusion (fig 5). The gland size varied from 10–50 mm, mean (SD) 19.5 (8.6), and the number of anatomical gland areas from 1–5, mean (SD) 2.5 (1.22), with varying combinations. The aortopulmonary glands were most frequently enlarged (63%) followed by the paratracheal (51.8%), carinal (40.7%), pretracheal (25.9%), and hilar (14.8%). The glands showed a hypodense centre in 52%. Matting of adjacent glands was seen in 57.7%. The hilar glands were least frequently enlarged which can be explained from the lymphatic drainage of the pericardium. The enlarged nodes could not be seen on the plain chest radiograph (fig 1) and only on chest computed tomography and disappeared or regressed with specific anti tuberculous therapy.12 Features of mediastinal lymph nodes in TPE are shown in box 2.
Echocardiography
The pericardial exudate is thick and fibrinous with a tendency to form adhesions and in some instances constriction. On echocardiography there are patchy deposits 4–8 mm in thickness with “fibrinous” strands criss crossing the pericardial space. The appearance is quite characteristic.

Tuberculin skin test
The tuberculin skin test is done using the purified protein derivative. A positive skin reaction is an induration ≥10 mm and a strongly positive response is defined as one ≥15 mm with or without excoriation of the skin. A positive tuberculin skin test has been found in all patients with TPE in one report and in 239/240 in another. In the latter it was >15 mm in 78%. In a recent publication the induration measured 16.4 (3) mm. A strongly positive tuberculin skin test is of value when associated with tissue granuloma without acid fast bacilli or when typical non-hilar mediastinal adenopathy is detected on chest computed tomography.

Pericardial fluid
The greatest value of the pericardial fluid is when M tuberculosis can be cultured as it is rare to find acid fast bacilli in a spun smear. The raised protein and lactic dehydrogenase values speak for an exudate. Lymphocytosis has been found in some studies but not in others. All these non-specific findings can be found in chronic idiopathic pericardial effusion as well. The value of adenosine deaminase and other markers has been discussed earlier.

Culture of mycobacterium tuberculosis
M tuberculosis may be cultured from the sputum, tissue like the pericardium, pleura, scalene node, or other accessible enlarged nodes but such information is patchy. Recovery from the pericardial fluid has varied from 30%–100%. Strange et al reporting on patients with TPE found that out of 49 patients in whom M tuberculosis was recovered from the pericardial fluid the pericardial histological changes “characteristic of tuberculosis” was present only in 8 of 19 patients with tuberculous pericarditis. In another study 2/19 biopsies showed non-specific findings. This difference may be due to the number of samples obtained at biopsy.

Scalene node biopsy
The report on scalene node biopsy has been discussed earlier. This would be a safe and relatively easy route to a specific diagnosis but so far there has been only one report.

Histological evidence
Like recovery of M tuberculosis from the pericardial tissue or fluid, histological evidence of a tuberculous granuloma with the demonstration of acid fast bacilli would be a definite diagnostic criterion. The typical granuloma is however not always found and the pericardial biopsy may show non-specific findings even when M tuberculosis is found in the pericardial fluid. Strange et al reporting on patients with TPE found that out of 49 patients in whom M tuberculosis was recovered from the pericardial fluid the pericardial histological changes “characteristic of tuberculosis” was present only in 35/49 and 14/49 (29%) had non-specific findings. In another study 2/19 biopsies showed non specific findings. This difference may be due to the number of samples obtained at biopsy.

Box 3: Criteria for diagnosis of tuberculous aetiology

Any of the following:

Invasive
1. Culture of M tuberculosis from pericardial fluid or tissue.
2. Pericardial tuberculous granuloma with acid fast bacilli.
3. Pericardial tuberculous granuloma + positive tuberculin skin test.
4. Pleural tuberculous granuloma with acid fast bacilli.
5. Pleural tuberculous granuloma + positive tuberculin skin test.
6. Tuberculous granuloma in scalene node or peripheral lymph node.

Non-invasive
1. Active tuberculous elsewhere in the body.
2. Mediastinal (non-hilar) lymph nodes on chest computed tomography with hypodense centre and matting + positive tuberculin skin test (matting = coalescing of adjacent lymph nodes).
3. Response to specific antituberculous therapy.

Box 4: Questions (answers at end of references)

1. What are three common causes of large pericardial effusions which can pose difficulties in differential diagnosis?
2. What is the most frequent route by which the pericardium is affected in tuberculosis?
3. Which group of mediastinal nodes are least often enlarged in tuberculous pericardial effusion?
4. Can mediastinal lymph nodes be detected both on a plain chest radiograph as well as a chest computed tomogram?
5. Non-specific findings on histopathology of the pericardium may be found in around 30% of patients with proven tuberculous pericarditis. True or false?
6. What is the prognosis of tuberculous pericardial effusion with specific treatment?
7. Cardiac tamponade is a frequent mode of presentation with tuberculous pericardial effusion. True or false?

CONCLUSIONS
Tuberculosis is an important cause of pericardial effusion in Afro-Asian countries and with the spread of HIV infection there is an increased incidence all over the world. Significant recent developments have been the use of PCR technology, improved techniques for the recovery of M tuberculosis, observations on mediastinal lymph nodes on chest computed tomography, more clearly defined observations on echocardiography. The diagnosis of tuberculous aetiology in pericardial effusions can be based on the criteria shown in Box 3.

REFERENCES
ANSWERS

(1) Tuberculosis, viral, and chronic idiopathic.  
(2) From infected mediastinal lymph nodes.  
(3) Hilar group.  
(4) No. Not seen on plain chest radiograph.  
(5) True.  
(6) Excellent.  
(7) True.


