Post-tuberculous fibrosing mediastinitis: a review of the literature

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ABSTRACT
Fibrosing mediastinitis (FM) is a rare disease where there is thickening of the fibrous tissue in the mediastinum. While histoplasmosis is the most common recognised cause, the link with tuberculosis (TB) has been rarely documented. We review the link between TB and FM, and describe a case of probable TB-related FM. Our case is of a 74-year-old man who presented with breathlessness 3 years after fully treated TB. Scans revealed a calcified soft tissue mass within the mediastinum, and a diagnosis of fibrosing mediastinitis resulting in pulmonary hypertension was made. Tests for histoplasmosis and IgG4 staining were negative. Surgical intervention was not felt to be beneficial, and he was treated with prednisolone and mycophenolate mofetil. In the review, we highlight the common recognised cause, the link with tuberculosis in 2010, presented in March 2013 in Pakistan with a 4-week history of breathlessness.

His examination findings of the chest revealed mild expiratory wheeze and reduced air entry in the left base. A CT scan showed apical scarring bilaterally, with multiple small nodules in the right middle and lower lobes, the largest measuring 4 mm and a left-sided pleural effusion. Pleural aspiration removed 500 mL of transudative fluid, and cultures were negative for TB. Sputum culture was also negative for TB. He was treated with a week’s course of antibiotics, steroids and inhalers with a diagnosis of asthma. After discharge he was weaned off the steroids and inhalers as he did not feel any subjective benefit.

In October 2013, he was admitted again into hospital following a recurrence of the left pleural effusion, this time in the UK. Lung function testing showed a mixed obstructive and restrictive pattern with forced expiratory volume in 1 s 1.11 (50% predicted), forced vital capacity 1.81 (60%) and transfer factor for carbon monoxide 3.69 (50%). Echocardiography showed an estimated left ventricular ejection fraction of 60%. However, even among patients with histoplasmosis infection, only 3 in 100 000 were found to develop fibrosing mediastinitis.2 Other conditions linked to FM include tuberculosis (TB),3 aspergillosis13 and more recently IgG4 related disease.6 An idiopathic form, with non-calcific fibrosis, has also been described and can be associated with extrathoracic fibrotic conditions such as retroperitoneal fibrosis as well as other autoimmune disease.7,8

The literature describes two forms of mediastinitis, namely granulomatous mediastinitis, where fibrous reaction is limited to the mediastinal lymph nodes, and fibrosing mediastinitis, where there is a more widespread reaction in the mediastinal space.9 The distinction between these two forms of disease is explored in this article.

Despite the link between TB and FM, such cases are sparsely reported in the literature. In this paper, we present a new case of TB-related mediastinal fibrosis and provide a literature review of the association between these two conditions.

CASE PRESENTATION
A 74-year-old man, of Asian origin, with a medical history including a myocardial infarction in 2007 and fully sensitive treated tuberculosis in 2010, presented in March 2013 in Pakistan with a 4-week history of breathlessness.

He was, therefore, referred to the regional centre for pulmonary hypertension and...
subsequently underwent right heart catheterisation which showed a mean pulmonary artery pressure of 31 mm Hg, pulmonary wedge pressure of 10 mm Hg and pulmonary vascular resistance of 512 dynes·sec·cm⁻² in keeping with precapillary pulmonary hypertension.

Crucially, on repeat CT (see figure 1), there was a new calcified soft tissue mass within the mediastinum and hilar region, and a diagnosis of FM was made.

An expert opinion was sought regarding surgical intervention, and it was felt that the mediastinal fibrosis was causing extrinsic vascular compression with multifocal proximal pulmonary artery stenoses and narrowing of the pulmonary veins at the bronchial level. However, angioplasty was considered too high risk for arterial rupture as multiple vessels were affected.

A fluorodeoxyglucose-positron emission tomography (FDG-PET) scan (see figure 2A,B) revealed a metabolically active partially calcified hilar and mediastinal soft tissue mass with activity. The patient underwent an endobronchial ultrasound in May 2014 and histology from the lymph nodes (see figure 3A,B) revealed necrotic debris, anthracotic macrophages and occasional lymphoid cells without evidence of granuloma or malignant cells and negative IgG4 staining. TB culture and PCR were both negative, as were tests for histoplasma, antinuclear antibody and HIV. Due to the increased activity seen on PET scan, he was started on 60 mg prednisolone daily in June 2014 with TB chemoprophylaxis.

After 2 months of treatment, he had symptomatically improved, back almost to his baseline in 2007. However, a repeat CT scan did not show any significant difference to the previous scan.

After reducing the prednisolone to 50 mg, his breathing declined and was admitted again in September 2014 for progressive breathlessness. He was given three doses of intravenous methylprednisolone, another therapeutic aspiration of 500 mL of the left-sided effusion and discharged on prednisolone 40 mg and furosemide.

To wean the steroid dose, he was started on mycophenolate mofetil (MMF) 500 mg twice daily in October 2014. By November 2014, he was taking only 10 mg prednisolone daily. During the subsequent months, he remained steroid dependent but had complications with steroid-related side effects including osteoporotic vertebral fractures and myopathy.

Figure 1 CT scan showing a soft tissue mass with multiple areas of calcification, causing narrowing of the right pulmonary artery. A left-sided small pleural effusion is also present.

Figure 2 CT (left of images) showing calcified areas and PET (right of images) scans showing increased FDG-PET metabolic activity in the mediastinal mass (A) and perihilar lymph nodes (B). FDG-PET, fluorodeoxyglucose-positron emission tomography.
In January 2015, he developed marked lethargy with peripheral oedema and was admitted again for intravenous methylprednisolone, diuresis and a further 660 mL of transudative fluid was aspirated from the left-sided effusion. A repeat echocardiogram showed worsening pulmonary hypertension with a pulmonary artery systolic pressure of 78 mm Hg. Nevertheless, his symptoms have remained stable on 5 mg prednisolone and 500 mg twice daily MMF, with a stable effusion and his exercise tolerance is between 300 and 400 m.

Prevalence
The prevalence of TB-related mediastinal fibrosis is unknown, with only a handful of cases having been reported in the literature. In a large case review of 180 cases by Schowengerdt et al in 1969, there were 13 cases of granulomatous mediastinitis and only one case of FM where TB was identified as the culprit (via confirmation of acid-fast bacilli). The authors reviewed cases that, at the time, were not subjected to current testing methods for TB infection such as TB-PCR. However, the discrepancy between the rates of TB found in the granulomatous versus fibrosing disease spectrum may support the concept of the two states existing in a dynamic spectrum with granulomatous mediastinitis being seen in early active infection, where AFBs are more likely to be detected. Only a proportion of granulomatous mediastinitis cases lead to FM, where the scarring invades into the rest of the mediastinum. Therefore, FM could represent the later stages of an aberrant healing response following an active infection. This theory is observed in many subsequent case reports where tissue from fibrotic mediastinum has yielded negative results for AFB and TB cultures. Indeed, this is also observed in Histoplasmosis-related fibrosis, where most cases are also culture negative. Nevertheless, it is important to highlight that due to the rarity of the disease, it is not clear from the literature whether the two forms of mediastinitis represent ends of a spectrum or distinct clinical entities.

A literature search for other case reports documented in English revealed five single case reports and two case series. The first case series, by Mole and colleagues, detailed eight patients with TB-related FM and a further single case with both TB and histoplasmosis. The second case series reported a further nine patients with active pulmonary TB and dysphagia attributed to FM. The basis for the diagnosis of FM in all of these cases was plain chest radiography alongside normal gastroscopy. A more definitive diagnosis using either CT imaging and/or mediastinal tissue biopsy was not obtained.

Clinical presentation
The initial presentation of FM depends on the anatomical structures that are affected by the fibrotic reaction. Schowengerdt et al identified that in early granulomatous disease, only a quarter (26%) of patients had involvement of adjacent structures, compared with the later fibrotic stage where this percentage increased over threefold (83%). The most common structure involved was the superior vena cava (SVC), with over 60% of the patients with FM having caval obstruction. Other regions involved included the oesophagus, trachea, bronchi, pulmonary vein and artery, inferior vena cava, thoracic duct, atrium and recurrent laryngeal nerve. SVC obstruction was also reported by the Mayo Clinic (6 out of 31 cases), and by Mole and colleagues (6 out of 18).

Although SVC obstruction is a feature, the slowly progressive nature of this condition allows collateral vein formation, and patients may often be asymptomatic initially. Progression of disease may lead to shortness of breath, venous distension of the neck and upper chest region, facial oedema and features of cor pulmonale. Should the bronchial structures be involved, there may be progressive dyspnoea, cough and haemoptysis. Other complaints include dysphagia with oesophageal invasion, hoarseness of voice with recurrent laryngeal nerve entrapment, as well as non-specific chest pain and malaise.

Of the single case reports where FM is specifically believed to be related to prior TB infection, presentations were due to superior vena cava syndrome, bronchial obstruction, pulmonary artery compression, as well as impingement of the transverse sinus and its adjacent vascular structures.

Pathogenesis
The pathogenesis of fibrotic tissue formation in FM is not clear. It has been proposed with Histoplasmosis infection, that the organism spreads along the lymphatic system and a cell-mediated response drives the formation of mediastinal granulomas.

Flieder and colleagues examined the histological morphology of tissue taken from 30 cases of idiopathic FM and proposed the condition as a three-stage progression of fibroinflammatory changes. Stage I was characterised by oedematous fibromyxoid tissue with numerous spindle cells, eosinophils, mast cells, lymphocytes and plasma cells. Stage II developed into thick bands of haphazardly arranged collagen. Stage III revealed dense acellular collagen and occasional dystrophic calcification. Interestingly, a more recent study has revealed...
that CD20-positive B lymphocytes are commonly seen on immunophenotyping and may be involved in potentiating disease progression.19

Diagnosis
Several methods have been reported for obtaining the diagnosis of FM. Prior to the use of more advanced imaging techniques, a chest radiograph was a useful way of raising suspicion, with widening of the mediastinum and lymphadenopathy of the hilar and paratracheal regions.20 The mainstay of imaging currently is CT. The majority of cases with FM will have localised fibrotic areas with calcification and there is much better identification of the surrounding vascular and airway compression with this imaging modality.20, 21 In some cases, a diffuse soft tissue mass without calcification can also be seen,22 and tissue sampling for malignancy should be sought. Although mediastinoscopy and surgical exploration are methods to obtain tissue,3 15 we used the less invasive technique of endobronchial ultrasound.

MRI has equal effect in recognising mediastinal structural involvement, but importantly, it is not as specific for identifying calcification, which is the hallmark of FM.23 MRI obviates the need for intravenous contrast for prominent vascular structures, which may be advantageous in those patients who are iodine allergic or who have severe renal impairment.

There have been some reports of using pulmonary scintigraphy24–26 to visualise perfusion defects and angiography to identify vascular compromise, but they do not diagnose FM, rather they are adjuncts to CT or MRI. In addition, CT or MRI angiography can in the modern era demonstrate sufficient detail of vessel encroachment without such tests.14

The role of FDG PET/CT scanning has been unclear. Chong et al demonstrated negative FDG uptake, concluding that PET scanning is useful in differentiating with malignancy.27 However, others have shown that active inflammatory lesions are also PET avid, so it can be used to gauge response to treatment, although biopsy to exclude a neoplastic process is still mandatory.28 29

Treatment and clinical outcomes
The management of FM is divided into surgical and medical therapy. The surgical approach depends on the anatomical structure involved, but is used as a last resort as fibrosis surrounding vital structures makes the procedure extremely challenging.

In the few reported cases of TB-related FM highlighted in the beginning of this review, outcomes have been variable. In the case reported by Ojeifo et al, the patient was treated for latent TB for 6 months on the basis of a positive interferon-γ release assay with no evidence of active disease from microbiological sampling.13 The main intervention was surgical removal of the fibrotic mass and insertion of a branched homograft to relieve severe stenoses of the right main pulmonary artery and SVC.13 Fijolek et al treated their patient who presented with recurrent haemoptysis, with two courses of anti-TB therapy prior to the emergence of the diagnosis of FM which developed over 10 years after the initial culture-positive TB diagnosis.12 There was severe right lower pulmonary vein and left pulmonary artery narrowing. She underwent a trial of 2 months of 60 mg prednisolone with no surgical intervention, but the mass increased in size, and therapy was stopped.12 However, interestingly they reported that her symptoms had not recurred at follow-up at time of publication.

Zhang and colleagues removed a fibrotic mass, which encroached onto the aorta, pulmonary artery and left atrium, followed by anti-TB drugs, with good resolution of symptoms, although the length of follow-up was not specified.13 Despite compression of vascular structures, collateral circulation often develops over time and without acute obstruction, Baikoussis et al decided to monitor rather than intervene.10

In the case series reported by Mole et al, 9 out of 18 cases of FM had previous TB.15 Unfortunately, the authors do not specify outcomes specifically for the TB patients but for the whole cohort. Moreover, data for treatment with steroids or anti-TB medication was unavailable.15 Several patients underwent surgery, including superior vena cava bypass grafting, lobectomy and excision of fibrous tissue. Follow-up of 12 of the patients was for between 5 and 20 years, with 10 of these free from symptoms, including 3 that had undergone surgery for vena cava obstruction.15 The two that died had developed malignancy and did not die of complications of mediastinal fibrosis.

For those patients that have FM, not related to TB, treatment and outcomes are similar. Peikert et al reviewed 80 cases, with most of the patients having had evidence of previous Histoplasmosis infection from the Mayo Clinic between 1998 and 2007.19 A total of 31 patients had antifungal treatment, 5 received steroids and 2 trialled tamoxifen. Radiological intervention with vascular angioplasty/stenting or endobronchial balloon dilatation/stenting was carried out in 15 patients, while 17 required surgical intervention.19 Of the 34 patients followed up from having medical therapy, less than a fifth (18%) had some response in symptoms or radiological findings; only three patients had imaging response to antifungal therapy; and one patient had response to anti-inflammatory treatment.19 However, 21 medically treated patients remained stable. Those that underwent angioplasty or stenting required a repeat procedure within 6 to 12 months. Five out of the 12 patients who were followed up after surgery needed further intervention such as stenting or embolisation for symptom recurrence.19 However, overall survival was similar to age-matched controls, with only two patients dying as a direct result of FM. Surgery should be used as a last resort as demonstrated in one recent study, with complications from carinal pneumonectomy leading to death in 3 patients out of 18 who underwent surgical interventions relating to vital airway or lung structures.30
Steroid therapy has been successful in some reports. Sherrick et al reported that three out of six patients with diffuse radiological change responded to steroids on repeated imaging, while none of the four patients with localised fibrosis improved. They proposed that widespread fibrosis may be indicative of a more generalised active immunological response and therefore have a better response to anti-inflammatory therapy.

Tamoxifen, a selective oestrogen receptor modulator, has been used in combination with steroids based on its antifibrotic effects against transforming growth factor β activity. Savelli et al used steroid and tamoxifen in combination with good effect initially, and when tamoxifen was stopped disease progression occurred, before regressing once again with combination therapy. In contrast, when relapse occurred following weaning doses of steroids as part of combined therapy with tamoxifen in another patient, the team increased steroid doses to induce remission again with good success. In another case report, a unique triple combination of itraconazole, tamoxifen and high-dose steroids was successful in reducing FDG activity on PET scanning.

DISCUSSION

Our patient had several factors contributing to his breathlessness, namely mild pulmonary hypertension, recurrent pleural effusion and parenchymal lung damage post-TB infection. The onset of mediastinal fibrosis was several years after fully treated active TB infection, and was diagnosed on CT scan, the imaging modality of choice. FDG-PET helped to establish activity within the mass, but biopsy was still required to exclude malignancy and recurrence of TB. We performed endobronchial ultrasound-guided biopsy which has the advantage of being less invasive compared with a video-assisted thorascopic surgical (VATS) biopsy or mediastinoscopy, but not all lesions are accessible via this approach. It is important to note that in our case the biopsy showed fibrous tissue without active granulomatous inflammation, and along with FDG-PET avidity, we postulate this was active fibrotic inflammation.

Treatment was initiated with high-dose steroids, and in line with other cases treated in the literature, TB chemoprophylaxis was given. Our patient had marked symptomatic relief after starting steroids, but relapse of severe dyspnoea occurred within 3 months, even though the steroid dose was only reduced from 60 to 50 mg. However, high-dose steroids with intravenous therapy again produced symptomatic relief. We did not perform a repeat FDG-PET to assess if there was a reduction in the activity within the mass, as the size of the lesion was thought to be the main factor in causing compression of local structures rather than the amount of active inflammation.

The use of MMF in our case as adjunct therapy was based on success with this therapy in treating retroperitoneal fibrosis, where Scheel et al showed that 89% of their patients had an over 25% reduction in size of the periaortic mass but also as a steroid-sparing agent given his diabetes. Surgical and radiological intervention was considered too high risk. The decision to operate should be reserved as a last resort, and moreover does not prevent progression of disease, with repeated procedures likely needed within 6–12 months.

The relationship between TB and mediastinal fibrosis is still poorly understood, and further reporting of cases is required. In this article, we have highlighted the two forms of mediastinitis—granulomatous versus fibrous. It remains unclear whether the granulomatous form inevitably leads to the more widespread fibrous reaction. No studies exist that observe the natural course of disease, and notably this case illustrates the possibility that TB antigen may be the cause of ongoing inflammatory changes even post-treatment. The pathogenesis of TB-related FM is still speculative and partly relies on our understanding of histoplasmosis-related FM. However, it is not known if TB-related FM runs a similar clinical course to histoplasmosis. It is reasonable to trial treatments that have been successful in idiopathic or histoplasmosis-related FM in cases of post-TB mediastinal fibrotic disease.

Although our patient did not show improvement with MMF despite positive outcomes in retroperitoneal fibrosis, we have been able to maintain clinical improvement on a combination of MMF and low-dose corticosteroid. Inoue and colleagues demonstrated the first case of IgG4-related FM with high serum IgG4 level correlating to histological findings. A good response to steroids was indeed observed, and the authors suggest serum IgG4 levels as an indicator for inducing steroid therapy in this specific group of patients. In addition, other agents such as tamoxifen or even B-lymphocyte depleting treatment may be considered on a case-to-case basis.

We have reviewed TB-related FM, and this should be considered in cases of FM with prior or active TB.

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