Lymphoid Interstitial Pneumonia in a Renal Transplant Recipient

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Abstract

Objectives: A case report of lymphoid interstitial pneumonia (LIP) in a renal transplant patient who is immunosuppressed with tacrolimus.

Background: LIP is a rare lymphoproliferative disorder affecting the lungs. It is described in the literature as: diffuse infiltration of the alveolar septa by dense collections of lymphocytes and a mixture of plasma cells and other elements. LIP is associated with several conditions, most of which affect the immune system. The development of this condition following immunosuppression with sirolimus is mentioned in the literature but there are currently no documented cases on the development of LIP following treatment with tacrolimus.

Methods: Case report and literature review.

Results: A 44 year old lady with rapidly progressing renal failure secondary to IgA nephropathy developed progressive dyspnoea nine years after a cadaveric renal transplantation. She was immunosuppressed on tacrolimus. Plain chest film revealed bilateral interstitial infiltrates. Following open lung biopsy she was diagnosed with lymphoid interstitial pneumonia and made a good recovery after treatment with Prednisolone.

Conclusion: LIP is a rare complication of treatment with tacrolimus, to our knowledge this is the first reported case of this condition in a patient maintained on tacrolimus. A lung biopsy may be required as the differential diagnosis is vast and treatment with prednisolone can lead to good outcome.

Keywords
Lymphoid; Pneumonia; Renal Transplant; Immunosuppression

Introduction

Here we report a case of a patient with IgA nephropathy who developed lymphoid interstitial pneumonia (LIP) 11 years after her renal transplant. Initially identified by Liebow and Carrington, LIP, a rare disorder, was described as a "benign lymphoproliferative disorder found in the lungs" [1]. They described it as: "diffuse infiltration of the alveolar septa by dense collections of lymphocytes and a mixture of plasma cells and other elements". This description further evolved into "a hypoplastic disorder of the mucosa associated lymphoid tissue, affecting the lungs". LIP is associated with several conditions, notably common variable immune deficiency (CVID), Sjögren's syndrome, post bone marrow transplantation as graft versus host disease, AID's, Epstein-Barr virus infection and related to hypogammaglobulinaemia [2-5] In renal transplant recipients, LIP has been associated with sirolimus immunosuppression but there have been no reports of association with calcineurin inhibitors (CNI) [6,7]. The patient presented to our transplant centre where we proceeded to do a high-resolution computed tomography (HRCT) scan (Figure 1). This showed bilateral hilar lymphadenopathy (BHL), mediastinal lymphadenopathy, multiple nodules in left lung and right base, interlobular septal thickening in both the middle and lower right lobe and ground glass change in right middle lobe.

Case Report

This 44 year old retired air stewardess developed rapidly progressive renal failure secondary to IgA nephropathy in 2005, and underwent a successful deceased donor renal transplant in 2006. She has no other past medical history, and was immunosuppressed on single agent of tacrolimus. She remained fit and well until late 2014, when she presented to her primary care physician with a six week history of a dry cough and progressive shortness of breath. She was uncomplaining of other symptoms. Her symptoms were initially treated as a lower respiratory tract infection with co-trimoxazole and later doxycycline, but this failed to improve her condition. Her inflammatory markers remained within the normal parameters (CRP<5 mg/dl) but a plain radiograph of her chest (Figure 2) revealed bilateral interstitial infiltrates.

Figure 1: Plain chest X ray.

Figure 2: HRCT of lung.
An urgent bronchoalveolar lavage (BAL) was performed, to exclude pneumocystis jiroveci pneumonia (PJP) among other opportunistic pathogens. Interestingly the PJP polymerase chain reaction (PCR), was negative from her BAL but her serum EBV PCR was positive at moderate titre, she was previously negative for serum EBV PCR at pre-transplant stage. After consulting our local microbiology and respiratory physicians for advice, the EBV viremia was thought to be an incidental finding and unlikely to explain her radiological appearances and the clinical features.

The patient’s lack of clinical improvement with antibiotics and lack of clear etiology for the extensive changes in the HRCT combined with concerns about possible post-transplantation lymphoproliferative disorder (PTLD) led us to perform an open lung biopsy. Histology of this biopsy was described as "widespread granulomatous non-necrotising lesions". Our differential included sarcoidosis and LIP. Prednisolone was initiated and patient made good recovery and was discharged home with a plan to taper prednisolone. On review of the HRCT the appearances are typical of LIP.

Discussion

LIP is a rare lung disorder that is usually associated with connective tissue disorders and infections. It is rarer in transplant patients but has been reported in patients maintained on sirolimus immunosuppression. To our knowledge this is the first case of LIP related to tacrolimus, in difficult to diagnose cases an open lung biopsy may be needed to diagnose the condition.

References