Fatal Strongyloides Hyperinfection in Post-Deceased Kidney Transplant Presented With Respiratory Failure and Septic Shock


Abstract

Immunocompromised patients are a vulnerable group to develop opportunistic infections. Transplant patients are always immunocompromised either because of their illness or immune-modulatory drugs that they receive. Strongyloidiasis is one of the opportunistic infections that increasingly attack transplant patients. We report this case of strongyloides hyperinfection in post-kidney transplant patient with respiratory failure and Gram-negative sepsis. Clinician should put strongyloidiasis in the differential diagnosis list while dealing with critically ill post-transplant patients and add it to the pre-transplant checkup.

Keywords: Strongyloides; Septic shock; Kidney transplant

Introduction

Infections due to strongyloides stercoralis are unusual in Saudi Arabia and are usually diagnosed in immigrants from endemic areas [1]. Strongyloidiasis can be now diagnosed in non-endemic countries due to the migration flows and travel, being the infection much more common in migrants than in travelers [2].

Hyperinfection describes the syndrome of accelerated autoinfection provided the organism is confined to the organs normally involved in the pulmonary autoinfection cycle [2, 3].

Hyperinfection of this organism sometimes leads to fatal outcome because hyperinfection often occurs in patients with impaired cellular immunity. We report this case of strongyloides hyperinfection presented with respiratory failure and septic shock in post-deceased kidney transplant patients.

Case Report

The patient was a 60-year-old Saudi male known to have hypertension and diabetes mellitus for 23 years complicated by end-stage renal disease for the last 2 years and required hemodialysis.

The patient was enlisted for kidney transplantation and pre-transplant checkup was all within normal. The patient received a post-deceased kidney transplant from a brain dead Pakistani gentleman. His early post-transplant course was complicated by delayed graft function and biopsy from the graft revealed acute cellular rejection. He received a pulse methylprednisolone therapy followed by tapering protocol of oral prednisolone starting with 160 mg daily.

Over next few days, there was slight improvement in the graft function, so the patient was discharged from the hospital with follow-up appointment after 4 weeks. Medication on discharge included prednisolone, mycophenolate and tacrolimus. Laboratory tests showed serum urea 21 mmol/L, creatinine 274 µmol/L, sodium 133 mmol/L, potassium 4 mmol/L and bicarbonate 23 mmol/L.

Three weeks later, he was admitted to the hospital because of severe nausea, vomiting and abdominal pain. His condition deteriorated and became more tachypneic with increasing bilateral chest infiltrates in chest X-ray (Fig. 1), so the patient was intubated and mechanically ventilated.

He developed ARDS with septic shock and Gram-negative (E. coli) bacteremia, and was started on Imipenem, linezolid, levofoxacin, ganciclovir and cotrimoxazol. The laboratory findings showed leucocytic count of 21.1 cc/L (91% neutrophils and 0.1% eosinophils), hemoglobin 7.3 g/dL, mild hyperbilirubinemia and normal liver function tests, bicarbonate 12 mmol/L, urea 22.5 mmol/L and creatinine 249 µmol/L.

Oesophage-gastro-duodenoscopy was done, revealing esophagitis and signs of delayed gastric emptying. Bronchoscopy was performed and bronchoalveolar lavage was sent for analysis which showed parasitic larva consistent with strongyloides stercoralis associated with acute inflam-
matory process mainly macrophages and lymphocytes. The patient was started on albendazol 400 mg BID. With those interventions the patient’s hemodynamics and respiratory status marginally improved. However, his neurological status did not improve, despite daily sedation interruption.

Over the next few days, the patient developed refractory septic shock with acute kidney injury requiring continuous renal replacement therapy. Despite maximizing vasopressor infusions and aggressive other supportive measures, the patient condition deteriorated and he expired.

Discussion

Renal transplantation has become the treatment of choice for most patients with end-stage renal disease [4]. Immunosuppression is the mainstay long-term management for post-transplant patients [5].

A group of known organisms could be reactivated or de novo infect the transplant patients, e.g. HIV, HCV and tuberculosis.

Strongyloides infections are a helminthic intestinal parasite which is endemic in tropical, sub-tropical regions affecting 30 up to 100 million worldwide.

The life cycle of strongyloides is basically comprised of two parts: a free living cycle outside of the host as rhabditiform larvae and a parasitic life cycle as infective filariform larvae (filariae) [6].

The life cycle of strongyloides infection starts by penetration of filariform larvae through skin on exposure to contaminated soil, then reaches the lung and moves to the gastrointestinal tract where they mature into female and male which mate in the intestine and produce eggs shed in stool or hatch into larvae which may re-infect the host penetrating the mucosa circulation disseminating to all organs resulting in disseminated syndrome or hyperinfection.

The source of infection in these patients seems to be either preexisting chronic infections in the recipient or from the allograft itself as in the case presented by Brugemann et al [7].

Strongyloides infection is rare in Saudi Arabia especially in good socioeconomic status like our patient so we assume that infection may be transmitted through the transplanted kidney.

Eosinophilia in strongyloidiasis might be more frequent compared to other chronic intestinal parasitic infections because the adult female worms live within the submucosa, not in the lumen of the gut [8, 9].

Low eosinophilic count and steroid use are risk factors to develop severe strongyloidiasis with high mortality rate [10].

Patients with impaired cellular immunity are also at increased risk of developing hyperinfection syndrome and disseminated infection [11]. From 1991 to 2013, more than 400 deaths due to strongyloides have been reported and about 20 reported cases strongyloides hyperinfection with mortality rate about 69%. In transplant patients, strongyloidiasis has been reported in kidney, liver, heart, intestine, pancreas and hematopoietic recipient [12].

In disseminated strongyloidiasis, the filaria spread all over body organs other than respiratory and gastrointestinal systems, for example brain [13, 14].

It may explain the encephalopathy of our patient which did not improve despite cessation of sedation, normal brain CT and normal liver function tests.

Conclusion

Strongyloides hyperinfection can happen any time after transplantation of different organs with high mortality rates.
However, it seems to have predilection to strike within the first 3 months of transplantation during times of increased Immunosuppression [15]. We recommend testing for strongyloides as pre-transplant workup list in high-risk recipients and donors. Monthly prophylaxis in high-risk patients is a plausible alternative but is difficult to advocate, due to the low frequency of this disease, which makes cost-effectiveness studies challenging [16].

Further studies are needed to define the benefits of routine prophylaxis in high-risk patients. Strongyloides hyperinfection should be considered if the recipient condition deteriorated rapidly despite proper antimicrobial prophylaxis and treatment of critically ill.

Empirical antimicrobial for such critically ill should consider ivermectin and/or albendazole along with a reduction in immunosuppression.

**Conflict of Interests**

The authors declared no conflicts of interest with respect to the authorship and/or publication of this article.

**Grant Support**

Not available.

**References**


