Endothelin receptor antagonists are commonly used in the treatment of pulmonary hypertension. Sitaxsentan, a selective endothelin A receptor blocker, induces a mild transaminis in approximately 3% to 5% of patients, but rarely an acute severe hepatitis. A case involving a 61-year-old female with sitaxsentan-induced acute severe liver failure is presented. Despite withdrawal of therapy, her liver tests failed to improve. After six weeks of monitoring, the patient was administered high-dose corticosteroids, with a good clinical and biochemical response. While endothelin receptor antagonists are postulated to cause hepatitis by inhibition of a bile salt transporter pump, an immune-mediated or idiosyncratic mechanism should be considered.

**Key Words:** Endothelin receptor antagonists; Hepatitis; Sitaxsentan

**CASE PRESENTATION**

The patient, a 61-year-old female with pulmonary arterial hypertension on a background of an atrial septal defect repair 15 years earlier, atrial flutter, atrioventricular node ablation and pacemaker insertion was commenced on sitaxsentan (100 mg/day for 16 weeks). Her liver enzyme levels were near normal at baseline (alanine aminotransferase 24 U/L, alkaline phosphatase 112 U/L, bilirubin 11.6 μmol/L), and she had no known history of liver disease. Concomitant medications included warfarin 2 mg once/day, diltiazem 100 mg four times/day, furosemide 20 mg four times/day orally, vitamin D and calcium. Liver enzyme levels were checked on a regular basis. After eight weeks of therapy, her liver enzyme levels became abnormal. Due to a continued deterioration in liver enzymes, sitaxsentan was discontinued six weeks later.

Despite cessation of therapy, there was continued worsening of all liver parameters over the next six weeks, and the patient was admitted to hospital for monitoring and investigation. On examination, she was clearly jaundiced, but had no stigmata of chronic liver disease nor encephalopathy. Investigations performed included a normal abdominal ultrasound and autoimmune screen (antinuclear antibody, antineutrophil cytoplasmic antibody, antismooth muscle antibody, antiliver kidney microsomal antibody, antimitochondrial antibody), negative hepatitis A, B and C, and Epstein-Barr virus and cytomegalovirus serology, and negative toxicology screen. The patient underwent a liver biopsy, which revealed a mixed inflammatory infiltrate with piecemeal necrosis of hepatocytes, without evidence of Mallory's hyaline, alpha-1 antitrypsin bodies nor iron deposition.

With the clinical scenario and investigations suggesting drug-induced hepatitis, and the bilirubin failing to improve despite cessation of sitaxsentan six weeks previously, the patient was commenced on high-dose corticosteroids, with good effect. Following four weeks of tapering oral corticosteroid therapy, her liver enzymes normalized (Figures 1 and 2).

**DISCUSSION**

The mechanism of ETRA-induced hepatitis is not clearly known. Bosentan-induced hepatitis is postulated to occur by inhibition of a bile-salt transporter pump (6). Sitaxsentan is also postulated to have a direct hepatotoxic effect on hepatocytes (4,7). Ambisentan, a newer ETRA, is not believed to have an effect on the bile-salt transporter pump, and remains an alternative therapy for patients intolerant to bosentan/sitaxsentan for reasons of hepatotoxicity (7).

The classical hallmarks of a direct hepatotoxin are that they are dose related, with rapid improvement of liver function on drug withdrawal. In our case, there was no resolution of jaundice despite a long observation period of six weeks until corticosteroids were administered.
Of the five published cases of sitaxsentan-induced hepatitis, one was treated with steroids after an observation period following cessation of therapy, and did not result in improvement of liver function (5). There was a rapid resolution of the hepatitis following steroid induction. Of the four other cases that were treated supportively after withdrawal of sitaxsentan, two died. While the delayed recovery may occur as a result of the nonlinear pharmacokinetics of sitaxsentan (4), an immune-mediated mechanism should be considered.

CONCLUSION
Nonclinically significant elevation of transaminase levels with sitaxsentan therapy occur in 3% to 5% of patients. To date, there are five cases of severe sitaxsentan-induced hepatitis in the literature. We presented a case of severe sitaxsentan-induced hepatitis that failed to resolve despite prolonged cessation of therapy, which required rescue glucocorticoid therapy. While it is currently postulated that sitaxsentan is directly hepatotoxic, an idiosyncratic or immune-mediated mechanism needs to be considered.

REFERENCES
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