Clozapine-induced fatal fulminant hepatic failure: A case report

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Fulminant hepatic failure (FHF) refers to the rapid development of severe acute liver injury with impaired synthetic function and encephalopathy in a person who previously had a normal liver or had well-compensated liver disease. The potential causes of FHF are numerous, but viral or toxin-induced hepatitis are the most common. Clozapine-induced hepatotoxicity has rarely been reported in the literature, occurs via an unknown mechanism and results in liver biochemical abnormalities that are usually of no clinical significance. In approximately 30% to 50% of patients treated with clozapine, there is an asymptomatic rise in serum aminotransaminase levels; however, there are no current guidelines for routine monitoring of liver function tests and liver enzymes during its use. Fatal fulminant hepatitis has only been reported in three patients receiving clozapine. A case of fatal FHF that occurred in a schizophrenic woman who began clozapine therapy shortly before her illness developed is described.

Key Words: Clozapine; Fulminant hepatic failure; Schizophrenia

Many medications are associated with hepatic toxicity. The severity of drug-induced hepatic injury can range from transient asymptomatic liver enzyme elevation (transaminis) to fulminant hepatic failure (FHF). The exact incidence of FHF remains uncertain, but studies suggest that the leading causes in North America are drug overdose and idiosyncratic drug reactions (1). Clozapine (Clozaril, Sandoz Pharmaceuticals, Germany) is an atypical antipsychotic used in the treatment of schizophrenia. It was introduced in the early 1970s, and subsequent clinical trials demonstrating fatal agranulocytosis resulted in a reduction in the frequency of its use. Clozapine has also been associated with numerous gastrointestinal side effects including constipation, bowel ischemia and hepatitis (2-4). There is little reported in the literature regarding acute clozapine-induced FHF apart from one case report (2). The present article describes a case of clozapine-induced fatal FHF.

CASE PRESENTATION

A woman in her fifth decade was examined by the gastroenterology service because of jaundice. She was admitted to hospital under the care of the psychiatry service due to increased delusions and a decreased ability to function independently. She had a history of paranoid schizophrenia with catatonic features and passive-aggressive behaviour. She also had type 2 diabetes and a gynecological malignancy that had been treated surgically one year earlier. Her medications before hospitalization included levothyroxine, risperidone and olanzapine, but she had a longstanding history of poor compliance. She denied having a history of alcohol use, smoking or intravenous drug use. She had no known history of liver disease.

Her regular antipsychotic medications were not continued after hospital admission and she was started on clozapine with the dose titrated from 150 mg daily to 300 mg daily. Baseline laboratory results collected on admission included a normal platelet count, normal liver enzymes and liver function tests. Her preadmission laboratory investigations were also normal, including an alanine aminotransferase (ALT) level of 60 U/L and an aspartate aminotransferase (AST) level of 26 U/L (ALT normal at less than 38 U/L). Six weeks after initiation of clozapine, the patient complained of mild nausea and right upper quadrant pain, and was noted to have developed jaundice. She subsequently developed nausea, emesis and anorexia. Her physical examination revealed severe drowsiness with easy rousability and no asterixis. She had a mildly distended soft abdomen with tenderness in the right upper quadrant. There were no stigmas of chronic liver disease. Laboratory results obtained 8.5 weeks after clozapine initiation revealed a white blood cell count of of 7.5×10^9/L without eosinophilia, hemoglobin 122 g/L, platelet count of 198×10^9/L, AST 890 U/L, ALT 1668 U/L, direct bilirubin 212 μmol/L, total bilirubin 321 μmol/L, alkaline phosphatase 273 U/L, albumin 27 g/L, international normalized ratio 2.9 and creatinine 54 mmol/L. Her clozapine was immediately discontinued. An abdominal ultrasound with Doppler assessment of
hepatic vessel flow revealed a small collapsed liver with no focal lesions.

Laboratory results for infectious (viral) serology, including hepatitis A, B and C, Epstein-Barr virus and cytomegalovirus, were negative. An autoimmune analysis was negative for antimitochondrial antibody, antinuclear antibody, antismooth muscle antibody and tissue transglutaminase antibody. Her urine toxicology screen, acetaminophen and alcohol levels were negative. Her serum ceruloplasmin and serum iron studies showed no evidence of copper or iron overload. Her serum alpha 1 antitrypsin level, serum immunoglobulin (Ig) G, IgA and IgM levels were normal. A full assessment was performed by the liver transplant service and she was deemed not to be a suitable transplant candidate. This decision was based on the patient’s history of poor compliance with her psychiatric medications, poorly controlled psychiatric disease, limited social support and recent gynecological malignancy, factors that significantly compromised the likelihood of a reasonable post-transplant outcome.

She subsequently developed periods of hypothermia, marked ascites and peripheral edema. Paracentesis revealed an ascitic fluid white blood cell count of $600 \times 10^9$\text{/L} with 26% neutrophils. Her blood and urine were taken several times; the resulting cultures were negative. Her platelet count progressively decreased and her renal function began to fail. She received intravenous albumin infusions daily for five days. She then developed mild encephalopathy, which was treated with lactulose.

Eleven-and-a-half weeks after starting clozapine, and three weeks after its discontinuation, she underwent a transjugular liver biopsy. Because she had persistent coagulopathy with thrombocytopenia, she received four units of fresh frozen plasma before the procedure. Middle hepatic wedge pressure was 44 mmHg and the central venous pressure was 10 mmHg. A liver needle core biopsy revealed massive zonal necrosis with approximately two-thirds of the tissue being necrotic (Figure 1). Periportal sparing, collapse of the reticulin framework, mild chronic inflammation, marked bile stasis and immature fibrosis was also demonstrated (Figure 1). Steatosis was not seen and there was no demonstrable excess of iron or copper.

The patient subsequently developed acute respiratory distress syndrome and multiorgan failure. Despite admission to the intensive care unit, mechanical ventilation and intense critical care support, she died 12.5 weeks after clozapine was started.

**DISCUSSION**

Clozapine-induced FHF is an uncommon, but recognized, complication of clozapine therapy (2). Well-documented side effects limiting clozapine use include agranulocytosis, seizures, orthostatic hypotension, hypertension and drowsiness. From a gastrointestinal perspective, documented side effects include abnormal liver function tests, constipation, bowel ischemia and ascites (3,4). There have been five cases of clozapine-associated hepatitis reported in the literature (4-7) and, to the best of our knowledge, the present case is only the second documented report of a patient with FHF leading to death as a result of treatment with clozapine (2).

The mechanism of clozapine-induced liver injury is unclear. The drug is metabolized in the liver via the cytochrome P450 pathway. It has been suggested that patients who experience drug-induced hepatitis with any medication may be vulnerable to clozapine-induced acute hepatitis (8). Icteric hepatitis with accompanying nausea has been documented four times and zone 3 necrosis was demonstrated on liver biopsy in one case (7,9). Fulminant hepatitis with encephalopathy and coagulopathy has rarely been reported (2).

It has been suggested that regular monitoring of liver enzymes is not necessary, given that the incidence and risk of serious clozapine-induced hepatotoxicity is low. Approximately 40% of subjects receiving clozapine have ALT or alkaline phosphatase levels twice the upper limit of normal (10,11). Recovery from clozapine-induced hepatitis appears to occur in the majority of patients when the medication is stopped (4). In retrospect, documentation of abnormalities in liver enzymes along with a worsening clinical picture and earlier cessation of clozapine may have prevented the present patient from developing fatal FHF. Therefore, our experience should be of interest to clinicians prescribing clozapine.

**IMPLICATIONS FOR CLINICAL CARE**

Asymptomatic transaminitis is commonly associated with clozapine therapy. However, caution is advised for patients with elevated liver enzymes who are receiving clozapine because the transaminitis may be the earliest manifestation in the development of FHF. Unfortunately, it is not possible to reliably predict which patients will progress to hepatic failure. Nevertheless, it may be prudent to diligently follow biochemical liver tests in patients receiving clozapine to monitor for early evidence of the development of FHF and possibly death.

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**Figure 1** A Liver biopsy stained with hematoxylin and eosin demonstrating hepatocyte necrosis with some periportal sparing, mild chronic inflammation, marked bile stasis and relative preservation of portal to central vein relationships; B High-powered view of a portal triad; C Reticulin stain demonstrating immature fibrosis and collapse of the reticulin framework.
REFERENCES