**CASE REPORT**

**Fibrosing cholestatic hepatitis following cytotoxic chemotherapy for small-cell lung cancer**

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**Abstract**

Fibrosing cholestatic hepatitis (FCH) is a variant of viral hepatitis reported in hepatitis B virus or hepatitis C virus infected liver, renal or bone transplantation recipients and in leukemia and lymphoma patients after conventional cytotoxic chemotherapy. FCH constitutes a well-described form of fulminant hepatitis having extensive fibrosis and severe cholestasis as its most characteristic pathological findings. Here, we report a case of a 49-year-old male patient with a 38 pack-year smoking history and a past medical history of hepatitis C virus infection who developed this condition following conventional chemotherapy. The patient had a smoking history and a past medical history of hepatitis C virus infection for decades. He was diagnosed with small-cell lung cancer who developed this condition following conventional chemotherapy. The patient was responsible for writing of the paper and its supervision; Gil-Bazo I was responsible for writing of the paper and its supervision; Sola JJ was responsible for the pathology assessment and graphic material editing; Aisa G contributed to the pathology assessment and graphic material editing; Gil-Bazo I was responsible for data gathering and paper writing; López-Picazo JM was responsible for the patient’s treatment, writing of the paper and its supervision; Pérez-Gracia JL performed writing of the paper and its supervision; Sola JJ was responsible for the pathology assessment and graphic material editing; Aisa G contributed to the pathology assessment and graphic material editing; Gil-Bazo I was responsible for writing of the paper and its supervision.

**INTRODUCTION**

Hepatitis B virus (HBV) is a well-known pathogen which can cause fulminant hepatitis in some patients undergoing cytotoxic chemotherapy. Patients who have completely recovered from acute hepatitis can harbor a latent HBV infection for decades. Fibrosing cholestatic hepatitis (FCH) is a recognized unique variant of viral hepatitis which was originally reported in 1991 by Davies et al. in HBV-infected recipients of liver allografts. Other authors have also reported FCH cases in HCV-infected recipients after liver transplantation and in renal, cardiac or bone marrow transplanted patients as well as in patients with acquired immunodeficiency syndrome. FCH has additionally been reported secondary to hepatitis C allograft reinfection.

However, to our knowledge only 3 cases of FCH have been reported after conventional cytotoxic chemotherapy. Nonetheless, all of them were diagnosed with non-solid tumors (acute myelogenous leukemia, acute lymphoblastic leukemia and acute lymphoblastic leukemia and low grade non-Hodgkin’s lymphoma). FCH is associated with extremely high mortality.

We report a case of a patient diagnosed with small-cell lung cancer who developed FCH under chemotherapy-induced immunosuppression.

**CASE REPORT**

This is a case of a 49-year-old male with a 38 pack-year smoking history and a past medical history of hepatitis B. He had not received previous blood transfusions. At the time of admission, the patient presented with pain in the right hemithorax. Initial investigations included thoracic X-ray, thoracic computed tomography (CT) and positron emission tomography (PET)-CT imaging demonstrating right pleural effusion, right upper lobe...
atelectasia, mediastinal adenopathies and liver and bone metastases. After a bronchoscopy-guided biopsy, he was diagnosed with extensive-disease small-cell lung cancer. The patient accepted chemotherapy and a combination regimen of cyclophosphamide 400 mg/m$^2$ (days 1-3), adriamycin 40 mg/m$^2$ (day 1), cisplatin 100 mg/m$^2$ (day 2), vincristine 2 mg (day 3) and etoposide 100 mg/m$^2$ (days 1-3) was administered. After the first cycle, the patient presented with grade IV febrile neutropenia in spite of pegfilgrastim prophylaxis. For this reason, a 20% dose reduction was applied on the second cycle. Doses were escalated to the original levels in the third cycle, during which the patient presented with elevation of glutamic oxalacetic transaminase (GOT) (232 UI/mL) and glutamic pyruvic transaminase (GPT) (639 UI/mL).

Following the third cycle, a PET-CT scan showed a radiological complete response. In contrast, the patient showed persistent elevation of transaminases (GOT: 198 UI/mL; GPT: 400 UI/mL).

After administration of the fourth cycle, he developed grade 4 thrombocytopenia and neutropenia. A complete serological study was performed demonstrating hepatitis B reactivation (HBsAg, HBeAb and HBcAb IgG positive). Despite treatment with the nucleoside analogue entecavir and methylprednisolone, the hepatitis progressed towards acute hepatic failure (GOT: 1879 UI/mL; GPT: 2663 UI/mL; total bilirubin: 42 mg/dL; direct bilirubin: 37 mg/dL; indirect bilirubin: 5 mg/dL; prothrombin time: 10%). A liver biopsy was not performed, due to thrombocytopenia and clotting time elevation.

The patient was transferred to the Intensive Care Unit and received plasma exchange, vasoactive drugs and wide spectrum antibiotics (piperacillin-tazobactam). In spite of these therapies, he died a month after the last chemotherapy cycle from acute liver failure.

The autopsy showed residual tumor in the mediastinum and peribronchial nodes, a hepatic reaction and fibrosing cholestatic hepatitis. The microscopic study revealed marked hepatocyte degeneration, whereas the immunohistochemical analysis demonstrated hepatocytes diffusely stained for HBsAg following both intracytoplasmic and cytoplasmic membranous patterns and nuclear and intracytoplasmic HBcAg staining patterns (Figure 1). Serum levels of HBV DNA polymerase were 17,857,100 UI/mL.

**DISCUSSION**

To our knowledge, this is the first reported case of FCH after conventional cytotoxic chemotherapy treatment in a patient diagnosed with a solid tumor. There have only been 3 case reports published of FCH patients who had received conventional chemotherapy for hematological malignancies. FCH is a subtype of viral hepatitis in HVB-infected patients which is associated with extremely high mortality.

Degeneration of hepatocytes with minimal infiltration of inflammatory cells (which is clearly distinguishable from other fulminant hepatitis), extensive fibrosis and severe cholestasis are the most characteristic pathological findings. Liver parenchymatous changes include hepatocyte swelling and cholestasis, with only mild or no portal or lobular inflammation, combined with acute cholangiolitis and fibrosis surrounding the cholangiomes.

Although the ultimate physiopathological mechanism in this condition remains elusive\[14\], the extremely high levels of viral replication, the massive HBcAg and HBsAg expression in the liver and the non-significant inflammatory component suggest a direct HBV cytopathologic effect. The accumulation of viral antigens in the endoplasmic reticulum damages vital cell functions leading to cell death\[13\]. Although specific treatment protocols are lacking, a few case reports
have described a clinical benefit after administration of ganciclovir/foscarnet antiviral therapy[15] in these patients. Additionally, other reports describe the efficacy of lamivudine, a nucleoside analog reverse transcriptase inhibitor, in the treatment of chronic hepatitis[12] and in prophylaxis of chronic hepatitis and FCH[16]. Moreover, lamivudine has been recommended by some authors for the prophylaxis of HBV hepatitis in carrier subjects undergoing cytotoxic chemotherapy for lymphoid malignancies[17], although no data from clinical trials are available.

This case report suggests that viral analysis might be indicated in patients presenting with solid tumors before initiating intensive chemotherapy regimens. In addition, prophylactic lamivudine should be considered in HVB chronic infection carrier patients (HBsAg positive and/or HBcAb IgG positive) in this setting.

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