Extrahepatic manifestations of hepatitis B virus infection: Addison's disease and myelofibrosis in a patient with persistent hepatitis B surface antigenemia

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F SOMLO, GR BERRY. Extrahepatic manifestations of hepatitis B virus infection: Addison's disease and myelofibrosis in a patient with persistent HBsAg antigenemia. Can J Infect Dis 1993;4(3):139-144. A 60-year-old white male patient was admitted to the hospital with acute abdominal pain, seemingly a self-limited ileus. He was found to be hepatitis B surface antigen (HBsAg)-positive. Previous dental treatment was suspected to be the initial source of the infection with hepatitis B virus. Five months later he was re-admitted with a diagnosis of adrenal insufficiency (Addison's disease) which responded well to steroids. Four years later he developed fever and leucocytosis. A bone marrow biopsy revealed myelofibrosis. He had several episodes of pyrexia during his lifetime. After a 12-year period the patient suffered a fatal myocardial infarction. At autopsy the adrenal glands were reduced to scarred remnants and HBsAg was found to be present in the residual adrenocortical cells by immunofluorescence methods. Bone marrow at autopsy revealed myelosclerosis as well HBsAg (via immunofluorescence). Hepatitis B virus was therefore closely correlated with the development of Addison's disease and myelofibrosis in this case.

Key Words: Addison's disease, Hepatitis B surface antigen, Hepatitis B virus, Myelofibrosis, Myelosclerosis

Manifestations extrahépatiques du virus de l'hépatite B : maladie d'Addison et myélofibrose chez un malade atteint d'antigénémie persistante

RÉSUMÉ: Un mâle caucasien de 60 ans est hospitalisé souffrant de douleurs abdominales aiguës, dûes apparemment à un ileus. Il est HBsAg positif. Un traitement dentaire du passé pourrait être la cause de l'infection initiale au VHB. Cinq mois plus tard présentant une fièvre générale il est hospitalisé de nouveau, la maladie d'Addison est alors diagnostiquée, elle sera par la suite traitée et bien contrôlée par les stéroïdes. Quatre ans plus tard il développe de la fièvre et une leucocytose. Une biopsie de la moëlle osseuse montre une myélofibrose. Il a ensuite plusieurs épisodes de pyrexies. A la fin de la 12ème année de sa maladie, il subit un infarctus fatal. A l'autopsie les surrenales sont réduites à environ 1 cm. HBsAg est détecté dans les cellules résiduelles par la méthode IF. La moëlle osseuse montre une myélosclérose et la présence de HBsAg par IF. On peut supposer une très proche corrélation entre la présence de l'antigène HBs et le développement de la maladie d'Addison et de la myélofibrose.
Acute hepatitis B infection is characterized by a relatively long incubation period, malaise, nausea, vomiting, fever, right upper quadrant tenderness and jaundice. Manifestations may range from asymptomatic to rapidly fulminant disease. The majority of cases are self-limited, with further immunity conferred by the development of anti-HBs antibodies. A certain percentage of patients, however, go on to become chronic carriers, or have variable end-organ damage, both hepatic and extrahepatic (5-7). We report two new extrahepatic manifestations of chronic hepatitis B antigenemia.

**CASE PRESENTATION**

A 60-year-old white male was admitted to St Mary’s Hospital Centre in Montreal, Quebec in March 1978 because of acute abdominal pain, which subsided rapidly and was thought to represent a self-limited ileus. Hemoglobin was 16.2 g/dL and white blood cell count was 18.2x10^9/L, with 80% neutrophils, and platelets 543x10^9/L. Bilirubin was 2.4 mg/dL. The patient was suspected to be the initial source of the hepatitis B virus (HBV) infection.

Five months later, the patient was readmitted because of slowly progressive severe generalized weakness. He had a brownish-grey discoloration on sun-exposed areas and in the palmar creases. Blood pressure was 70/40 mmHg. Adrenal insufficiency (Addison’s disease) was confirmed with the appropriate tests. The patient responded well to treatment with corticosteroids. Hemoglobin was 11.4 g/dL, and white blood cell count was 9.6x10^9/L. Antiadrenal antibody was reported to be negative; HBsAg remained positive. In 1982, he developed bouts of fever with a rising leukocytosis. In 1984, he was found to have newly diagnosed diabetes, as well as a complete blood count as follows: red blood cells, 13.8g/dL; white blood cells, 16.6x10^9/L; and platelets 348x10^9/L. Ovalocytes, tear drops, and nucleated red blood cells were interpreted as a leuko-erythroblastic picture consistent with a diagnosis of myelofibrosis, confirmed with a bone marrow biopsy. The patient was treated with hydroxyurea and remained reasonably well until the fall of 1987, when he again developed severe weakness and dehydration. The spleen at this time was palpable. Hemoglobin was 10 g/dL, and white blood cell count was 30x10^9/L. Splenectomy was performed.

The spleen weighed 2000 g, and showed extramedullary hematopoiesis. Chromosome studies failed to reveal the presence of Philadelphia chromosomes or rearrangement of the 3’BCR focus.

The patient continued to have recurrent bouts of unexplained fever until his death on February 3, 1990 due to an acute myocardial infarction.

Autopsy confirmed myocardial infarction, liver disease, adrenal pathology and myelofibrosis. The liver weighed 2200 g and had signs of extramedullary hematopoiesis and pigmentary fibrosis. The adrenal glands were each only a few centimetres in size. The parenchymal cells were almost completely destroyed and replaced by loose fibro-areolar tissue. Cholesterol-rich accumulation of necrotic debris with focal calcification was present. Mild inflammatory cell infiltration of the remnant of the gland and the periadrenal tissue was found along with some focal necrosis. Stains for mycobacteria and fungi were negative (Figure 1). The bone marrow was replaced by fibrous tissue and increased cancellous bone with only focal remnants of hemopoietic components containing numerous iron-laden macrophages and fibrosis (Figure 2).

**METHODS**

Routine paraffin sections of formalin fixed adrenals and Zenker fixed bone marrows from biopsies as well as from autopsy were placed on alcohol cleaned slides coated with glue (Lepage), and incubated in a ventilated dry air chamber at 56°C overnight. They were deparaffinized in xylene, rehydrated through graded ethanol and rinsed in calcium- and magnesium-free phosphate buffered saline. Treated sections were used in the following procedures:

- Tissues were explored using hematoxilin-eosin, Ziel-Nielsen and Gomory stains;
- Fluorescent antibody method was used for the search and localization of HBsAg.

**HBsAg localization, indirect method:**

1. Pepsin (0.4%) was applied for 1 h, followed by a wash in a calcium- and magnesium-free buffer.
2. Trypsin (0.25%) in calcium chloride (0.1%) was applied for 90 mins, followed by wash as above.
3. Hepatitis B human immunoglobulin (Hyper-Hep, Cutter Biological, California) was applied for 30 mins at 37°C, followed by two washings.
4. Fluorescent-tagged anti-human immunoglobulin G was applied for 30 mins at 37°C, followed by two washings with the same buffer.
Negative controls: Replacement of hepatitis B human immunoglobulin. Slides were submitted to the same procedures, except for step 3, in which hepatitis B human immunoglobulin was replaced by phosphate buffered saline.

Absorption of hepatitis B human immunoglobulin: Hepatitis B human immunoglobulin was absorbed by incubating it at 40°C for 1 h with hepatitis B vaccine (Recombivax, Merck, Sharp & Dohme, New Jersey). This mixture was applied to one slide as a negative control replacing hepatitis B human immunoglobulin in step 3. The hepatitis B human immunoglobulin attachment to the HBsAg in the cells could be eliminated in this control.

RESULTS

Positive immunofluorescent staining indicated the presence of HBsAg in the cells. Such staining was found in the remaining cells of the adrenal glands and the persisting hemopoietic cells from the bone marrow (Figures 3, 4).

The control slides treated with antibody absorbed by hepatitis vaccine gave negative staining results. Those with phosphate-buffered saline instead of hepatitis B human immunoglobulin also gave negative staining results. Thus, immunofluorescent staining could only be elicited if both HBsAg and specific antibodies were present in the cells (Figures 5-10).

In addition, various pathological sections were stained with immunofluorescent antibody to search for the presence of HBsAg. Those organs were removed during the autopsy and the microscopic findings with hematoxylin and eosin stain were described earlier. No HBsAg was found in any of the sections from the heart, lungs, liver, pancreas, kidneys, thyroid gland or lymph nodes.

DISCUSSION

Current epidemiological surveys estimate approximately 300,000,000 individuals to be carriers of HBsAg worldwide, and some 40,000,000 deaths annually are attributed to chronic HBV infection. Six million of these deaths are due to hepatocellular carcinoma (11-13).
In adults, 5 to 10% of HBV infections lead to a chronic carrier state with persistent HBsAg positivity (14). In some parts of the world, HBV is frequently transmitted from infected mothers to babies at the time of birth or by intimate contact in infancy or early childhood, playing a major role in perpetuating the infection. In general, carriers who are positive with hepatitis B early antigen (HBeAg) are particularly infective. An infant born to an HBeAg-positive mother has a very high chance of being infected and of becoming a chronic carrier (15,16).

The chronic carrier state has been demonstrated to be an important factor in the development of several serious diseases. Chronic active hepatitis, hepatic cirrhosis (17,18), hepatocellular carcinoma (19), a variety of renal diseases (20,21), arthritis (22,23), essential mixed cryoglobulinemia (24), aplastic anemia (25), testicular carcinoma (26), and acrodermatitis (27) have all been associated with chronic HBsAg carriers.

It was originally believed that HBV flourished only in the liver parenchymal tissue. The demonstration of HBsAg and HBV DNA in extrahepatic tissues (skin [28], pancreas and biliary secretion [29], leukocytes [30] and mononuclear cells [31]) has raised doubts about the so-called restricted tissue tropism of this virus.

The present patient, with known HBsAg-positivity for 12 years, developed Addison's disease and a myeloproliferative disorder, which progressed over six years from myelofibrosis of the marrow to myelosclerosis. Immunofluorescent staining showed the adenral cortical cells and the persisting hemopoietic cells to contain HBsAg. These findings suggest that the pathological changes occurring in these organs were related to the persistent HBV viremia with seeding of the virus into the cells of those tissues. Addison's disease in this case was of the primary insufficiency type, caused by the destruction of the functioning adrenal cortical tissue. The adenral glands are known to be affected during the course of a variety of acute viral infections: varicella (32), herpes simplex (33,34), cytomegalovirus (35), Coxackie virus (36) and rabies (37).

Recent studies of patients dying of acquired immunodeficiency virus infection have demonstrated destruction of the adenral glands by cytomegalovirus (38-40). Several publications have appeared in the world literature reporting Addison's disease in families, but none of them has documented an associated viral infection, except one which reported Addison's disease in two brothers, with serum hepatitis in one (41).

The exact etiology of myelofibrosis remains unknown. It has been documented to have developed following exposure to toxins, ionizing radiation and chemotherapeutic agents, as well as occurring secondary to metastases to the bone marrow (42,43). Two theories have been advanced to explain the pathogenesis of myelofibrosis: that there is primary damage to hemopoietic cells with secondary marrow fibrosis; and the reverse, that the primary event is in the connective
absorption of the hepatitis B human immunoglobulin by hepatitis B vaccine (Recombivax) no antibody attachment seen in the cells of the bone marrow. Magnification x264

tissue, resulting in a secondary effect on the hemopoietic cells (44,45).

We know of no previous demonstration of any virus related to myelofibrosis in humans, however, transmissible murine virus has been demonstrated to induce myelofibrosis in mice after a long latent period. Neoplastic transformation of fibroblastoid marrow cells have been shown to result from infection of mouse marrow culture by a myeloproliferative sarcoma virus (46).

The mechanisms of the various extrahepatic manifestations of HBV are unknown. In the case of the present patient, there appears to have been necrosis and replacement of the adrenal cortical tissue, but whether this was due to direct viral damage or to an autoimmune reaction, in spite of an early negative report on antiadrenal antibody, remains unanswered. In the case of the myelofibrosis, the presence of HBV in the hemopoietic cells may also lead one to entertain a causative link between infection and pathogenesis of the disease.

CONCLUSION

The occurrence of these unusual manifestations raises the possibility that diseases occurring in association with persistent HBV infection may be the result of end-organ damage directly or indirectly caused by HBV. Possible treatment with antiviral agents such as interferon-alpha might be tried in an effort to arrest the progress. Good clinical studies are needed to develop better treatment modalities. Prevention of the disease by more widespread immunization must be emphasized.

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