Hepatocellular carcinoma complicating chronic granulomatous hepatitis

WM MELIA, H CALVEY, B PORTMANN, ROGER WILLIAMS

From the Liver Unit, King’s College Hospital and Medical School, Denmark Hill, London SE5

SUMMARY The development of hepatocellular carcinoma, is reported in a patient with chronic granulomatous hepatitis after a seven year interval in which clinical and biochemical improvement had occurred on corticosteroid therapy and in whom, the development of cirrhosis was excluded by liver biopsy.

Hepatocellular carcinoma is a well recognised complication of cirrhosis, particularly those types that occur in males and in association with hepatitis B virus infection. It has also been seen very rarely in long-standing cases of hepatic fibrosis consequent on various chronic liver disorders—for example, HBsAg positive chronic active hepatitis, in which cirrhosis is more usually found. To our knowledge, however, there have been no previous reports of tumour development in chronic granulomatous hepatitis. We describe here a patient in whom hepatocellular carcinoma occurred seven years after the diagnosis of this condition and in whom serial liver biopsies over the years have shown little progression of the lesion and no development of cirrhosis.

Case report

A 48-year-old man was admitted to the Royal Infirmary, Doncaster in December 1973 following an episode of haematemesis and melaena. Clinical examination revealed hepatosplenomegaly, and oesophageal varices were demonstrated on barium swallow. Subsequently he was transferred to the Liver Unit where a history of gradually increasing abdominal distension during the previous year was obtained. On abdominal examination, the spleen was palpable 16 cm below the left costal margin, the liver edge 6 cm below the xiphisternum, and there was some ascites.

Laboratory investigations were as follows (Fig. 1): Erythrocyte sedimentation rate (ESR) 4 mm/h; haemoglobin concentration 11·6 g/dl; white cell count 3·7 × 10⁹/l with a normal differential count; serum bilirubin 24 μmol/l; alkaline phosphatase (ALP) 96 IU/l; aspartate aminotransferase (AST) 33 IU/l; serum calcium 2·40 mmol/l and other electrolytes normal. The prothrombin time was six seconds prolonged.

Serum was negative for HBsAg (RIA, Travenol Laboratories Ltd) and anti-HBs (Ausab, Abbott Laboratories) and alpha fetoprotein (AFP) was in the normal range (RIA, Radiochemicals Ltd). Autoantibody screening was negative apart from positive antinuclear factor in a titre of 1/20.

A radionuclide liver scan (99mTc Technetium sulphur colloid) showed that isotope uptake by the liver was reduced and that the right lobe had decreased in size. The spleen was markedly enlarged with an uptake of isotope greater than the liver. Mantoux and Kveim tests were negative; chest x-ray and slit lamp examination of the eyes were normal. Splenic venography showed an intrasplenic pressure of 32 mm Hg with patent splenic and portal veins.

Splenectomy with an end-to-side lienorenal shunt was performed on 6 March 1974, after which the oesophageal varices, assessed radiologically, became smaller. Histological examination of the spleen revealed multiple granulomata as well as congestive changes and a liver biopsy performed at that time showed widespread granuloma formation. After discharge from hospital he was able to recommence work and remained well until March 1975 when serum AST was found to have risen to 770 IU/l, the ALP to 333 IU/l and bilirubin to 23 μmol/l. After treatment with prednisolone (10 mg daily) there was a steady improvement in liver function tests, and by October, serum AST had fallen to 29 IU/l, ALP 67 IU/l and bilirubin to 12 μmol/l. Over the next two years he remained well and the liver became impalpable. From October 1977, prednisolone was gradually withdrawn and stopped altogether in June.

Accepted for publication 21 April 1983
Hepatocellular carcinoma complicating chronic granulomatous hepatitis

1978. However, a further liver biopsy in December revealed more florid changes and prednisolone (10 mg daily) was restarted. When he developed mild exertional dyspnoea in February 1979 the dosage was increased to 15 mg daily.

In February 1981 he complained that he became easily fatigued, and on examination was noticed to be jaundiced. The liver edge was palpable 6 cm below the right costal margin. Liver function tests showed a serum AST of 176 IU/l, ALP 517 IU/l and bilirubin 30 μmol/l. The dosage of prednisolone was increased to 20 mg daily, but he continued to lose weight and complained of abdominal discomfort, together with increased breathlessness on exertion. In August the liver had enlarged further (10 cm below the costal margin) and liver function tests showed deterioration with a serum AST of 590 IU/l, ALP 825 IU/l and bilirubin 44 μmol/l. Serum AFP was slightly raised at 35 ng/ml (normal range 0–10 ng/ml) and liver ultrasound revealed multifocal solid filling defects in the right lobe. Percutaneous liver biopsy from this area confirmed hepatocellular carcinoma. Investigations into the cause of his breathlessness included a chest x-ray which revealed a metastasis in the right lower lobe. Lung function tests showed a vital capacity of 3-4 l (predicted value 4-6 l), one-second forced expiratory volume of 2-6 l (predicted 3-4 l), CO transfer factor 22-4 mmol/kPa/min (predicted 30 mmol/kPa/min) and permeability constant 3-8/min (predicted 4-4/min), changes indicative of a transfer defect.

Chemotherapy with adriamycin was commenced but serum AFP concentrations continued to rise and there was an increase in tumour size on serial ultrasound examinations. There was a gradual deterioration and he died at home in April 1982.

SERIAL LIVER HISTOLOGY

Eight needle liver biopsies obtained at approximately yearly intervals since 1974 were available for review.

The initial biopsy showed the portal tracts to be expanded due to fibrous tissue deposition with conspicuous mixed cell infiltrates and discrete non-necrotising granuloma formation. A mild and patchy lobular inflammation was also observed. Similar changes were seen on the 1975 specimen, except that the granulomata were more numerous, in places confluent, and the portal cell infiltrate denser and often aggressive towards the nearby limiting plates (Fig. 2). There was a substantial improvement in this activity on the 1976 specimen (Fig. 3) when the patient had been on a daily dose of 10 mg of prednisolone for one year, and the 1977 biopsy showed

![Fig. 1 Serum aspartate aminotransferase (AST) and alkaline phosphatase (ALP) activities and bilirubin concentrations in a 48-year-old man with granulomatous hepatitis first diagnosed in January 1974, who developed hepatocellular carcinoma in August 1981 (LB = liver biopsy).]
the features of chronic persistent hepatitis with mild residual lobular inflammation and was the only non-tumoral specimen devoid of granulomatous infiltration. A further biopsy performed after steroid withdrawal in 1978 revealed florid portal and periportal activity identical to that seen on the 1975 sample. In July 1980, a year after steroid treatment had been resumed, activity was again markedly reduced, although giant cell granulomata were still present within slightly expanded portal tracts. Over the years there had been no significant increase in extent or degree of fibrosis, which was moderate on all specimens. The lobular pattern was well preserved in five and slightly distorted in one (1975). None of the biopsies showed evidence of nodular regeneration. The Ziehl Nielsen method for acid-fast bacilli, Grocott's method for fungi, and periodic acid-Schiff reaction after diastase treatment did not reveal any aetiological agent in the six biopsy specimens with granulomata. The biopsy performed in August 1981 revealed a very different picture, with a severely collapsed liver architecture and a mixed pattern of venous outflow block and cholangitis, strongly suggestive of compression of the liver by an expanding process. A repeat biopsy under

Fig. 2 Florid giant cell granulomata in and around a markedly enlarged portal area. Haematoxylin and eosin × 100.

Fig. 3 Mild residual inflammation with one granuloma and a well-preserved lobular architecture. Haematoxylin and eosin × 100.
Hepatocellular carcinoma complicating chronic granulomatous hepatitis

The development of hepatocellular carcinoma in such a patient in the absence of cirrhosis is thus of considerable interest. An increased incidence of neoplasia (malignant lymphoma and lung cancer) has been reported by Brucker and Wilder in patients with respiratory sarcoidosis although this has not been confirmed, and there is one report of malignancy at another site (malignant melanoma) in a patient with long-standing granulomatous hepatitis. However, we have been unable to find any previous report of hepatocellular carcinoma either in respiratory or other clinical variants of sarcoidosis; or in chronic granulomatous hepatitis from other causes. Other factors to be considered in relation to the development of malignancy in this patient are splenectomy and the prolonged use of corticosteroids. Robinette and Fraumeni did not find an increased incidence of malignancy in World War II veterans who had undergone splenectomy following trauma.

The present patient was on prednisolone therapy for six years and a convincing relation has been established between immunosuppressive drug therapy and an increased incidence of malignant tumours in patients after organ transplantation. Although the tumours which are found are most frequently mesenchymal (lymphoma) or epithelial (particularly carcinomas of skin, lip and uterine cervix) origin, there is one reported case of hepatocellular carcinoma developing in a child after renal transplantation. There is also some suggestion that cancers occur in non-transplant patients being treated with corticosteroids and the question of whether patients with autoimmune liver disease on immunosuppressive drugs have an increased risk of hepatocellular carcinoma, has been raised by several workers. However, with the exception of carcinoma of the skin, the results of the Oxford collaborative study performed between 1970 and 1978 failed to provide any clear evidence of an increased risk of the more common cancers in non-transplant patients treated with immunosuppressive drugs.

**Discussion**

The most likely cause of the granulomatous hepatitis in this patient was sarcoidosis. Although a Kveim test was negative and there was no radiological evidence of pulmonary involvement, epithelioid granulomata were demonstrated in both liver and spleen tissue, and dyspnoea had been present for over two years before lung function tests, compatible with fibrosing alveolitis, were demonstrated.

The definite clinical and biochemical response to corticosteroids is also consistent with hepatic sarcoidosis. Indeed, the Kveim test is often negative in patients with hepatic sarcoidosis. In a review of 30 patients with granulomatous hepatitis, Israel and Goldstein describe nine with definite, and another seven with probable, sarcoidosis of whom only two had positive Kveim tests. Similarly, in six of 13 patients with hepatic sarcoidosis described by Maddrey the Kveim test was negative.

Progression of granulomatous hepatitis to cirrhosis has been reported in occasional cases, but in the present patient the eight serial liver biopsies obtained over a seven-year period showed no such evidence. The portal hypertension was due to extensive granulomata and fibrosis in the portal tracts and is described in hepatic granulomatosis in the absence of cirrhosis. The ultrasound direction two weeks later confirmed the presence of a moderately differentiated hepatocellular carcinoma (Fig. 4).

**Fig. 4** Hepatocellular carcinoma with relatively compact growth of clear cells of which two are in mitosis. Haematoxylin and eosin × 180.

References


Requests for reprints to: Dr R Williams, The Liver Unit, King’s College Hospital and Medical School, Denmark Hill, London SE5 8RX, England.
Hepatocellular carcinoma complicating chronic granulomatous hepatitis.

W M Melia, H Calvey, B Portmann and R Williams

doi: 10.1136/jcp.36.9.1062

Updated information and services can be found at:
http://jcp.bmj.com/content/36/9/1062

**Email alerting service**

Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

**Notes**

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/