of the hepatic anlage (Fraser, 1952) or by further branching of the foregut diverticulum (Ashby, 1969).

Accessory lobes may be attached to the liver by a pedicle of the liver tissue or by a mesentery (Johnstone, 1965). For the accessory lobe to be viable it should have the hepatic artery, hepatic vein, portal vein and bile duct, which usually communicate through the liver, but occasionally may have extrahepatic connection as in the present case.

All reported cases of this phenomenon, and their symptoms, are summarized in Table 1.

Late presentation is due to gradual worsening of the circulation in the lobe. All the six cases reported were diagnosed at operation. Pre-operative diagnosis can be made by using one or more techniques of pneumoperitoneum, percutaneous hepatic cholangiography, splenic portography, regional arteriography, coelioscopy and scintillation scanning (Omanik and Jablonsky, 1972; Llorente and Dardik, 1971).

References


Chronic hepatitis in polymyalgia rheumatica

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Summary

A review of the literature relating hepatic abnormalities with the giant cell arteritis, polymyalgia rheumatica syndrome is presented. A case of polymyalgia rheumatica with biochemical and histological abnormalities of the liver is described. It is suggested that a more detailed investigation of liver function should be made in patients with this syndrome where reasonably indicated.

Introduction

Recently, attention has been focused on the association between polymyalgia rheumatica, giant cell arteritis and an elevated serum alkaline phosphatase with, or without, other evidence of hepatic dysfunction (Hall and Hargreaves, 1972; Glick, 1972; Dickson et al., 1973; Long and James, 1974). These findings have raised the possibility of an association between liver disease and these two disorders. Long and James (1974) reported finding granulomatous hepatitis in a single case of polymyalgia rheumatica. A case with different histopathological changes in the liver is now reported.

Case report

A 77-year-old woman with a long history of psoriasis, presented 8 years ago with malaise and severe pain in the muscles of the neck, back and proximal limb muscles. On examination, these muscles were tender to palpation and there was no evidence of psoriatic arthropathy. X-rays of the joints and chest were normal, the haemoglobin was 11 g/100 ml and the ESR 100 mm in 1 hr. Serological tests for gout and rheumatoid arthritis were negative. A diagnosis of polymyalgia rheumatica was made. The symptoms abated with oral prednisone therapy but the ESR remained elevated. Over the next 6 years she remained well apart from exacerbations of the disease when steroid withdrawal was

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attempted. Two years ago she began to develop anorexia and lassitude and abdominal palpation revealed an enlarged liver. Liver function tests were: alkaline phosphatase 44 KA \(\text{u}/100\;\text{ml}\) (normal range 4–14), aspartate transaminase (AST) 42 i.u./l (normal range 0–27), alanine transaminase (ALT) 63 i.u./l (normal range 4–24). Over the next 6 months, the liver function tests, having initially returned to normal values, again became abnormal. The ESR continued to be elevated at 69 mm/hr. At the end of the 6 months, prednisone therapy was stopped. The patient was then lost to follow-up until June, 1974, when she presented with depression, anorexia and recurrence of muscle pain.

On examination, she was jaundiced and had hepatomegaly and so she was admitted to hospital for investigation. She had had no injections in the previous 6 months and was receiving nitrazepam only. There was no history of excess alcohol ingestion. She had not been taking oral therapy for her psoriasis. The laboratory results were: bilirubin 1·9 mg/100 ml (direct 1·4 mg); alkaline phosphatase 80·5 KA \(\text{u}/100\;\text{ml}\); AST 75 i.u./l; ALT 54 i.u./l; total protein 8·8 g/100 ml (normal 5·7–7·9); albumin 3·7 g/100 ml (normal 2·5–5·0); globulin 5·1 g/100 ml (normal 2·3–4·0); lactate dehydrogenase 760 i.u./l (normal 250–650); serum calcium and phosphate normal; urinary bilirubin and urobilinogen elevated; Wasserman reaction negative; Australia antigen negative; haemoglobin normal; ESR 93 mm/hr; bleeding time and coagulation studies normal; there were no LE cells and no antibodies to nuclei, mitochondria, smooth muscle, parietal cells, thyroid microsomes and thyroid colloid. A liver biopsy showed increased fibrosis in the portal tract with irregularity of the limiting plate (Fig. 1) with a lymphocytic and occasional polymorph and plasma cell exudate (Fig. 2). There was no bile duct proliferation and no granulomata were seen. Kupffer's cells were prominent and showed ceroid phagocytosis, indicating recent liver cell damage. Bile stasis was not a feature and there was no active cellular necrosis. There was no evidence of cirrhosis or fatty change.

The appearances were considered to represent either a chronic persistent hepatitis or a stage in the development of primary biliary cirrhosis. In view of these findings and the patient's general symptomatic improvement, she was discharged from hospital but was not started on prednisone therapy. Two months later she was re-admitted to hospital with a fractured femur and the most recent liver function tests were: bilirubin 1·9 mg/100 ml (direct 1·3); AST 50 i.u./l; ALT 34 i.u./l; alkaline phosphatase 101 KA \(\text{u}/100\;\text{ml}\).

**Discussion**

Undoubtedly, this is a case of polymyalgia rheumatica with chronic liver disease. Laboratory tests in previous cases (Dickson et al., 1973; Long and James, 1974) particularly with isoenzyme studies of alkaline phosphatase, suggest the presence of hepatic dysfunction with involvement of the biliary excretory system. In this case, although no mitochondrial antibodies were present, the histological changes were more in favour of a disease process such as primary biliary cirrhosis rather than large duct obstruction or a hypersensitivity cholestatic reaction. In a recent review of liver diseases in

![Fig. 1. The portal tract is enlarged owing to irregular fibrosis and there is an inflammatory exudate extending into the parenchyma. Haematoxylin and eosin \(\times\) 90.](http://pmj.bmj.com/ on June 20, 2017 - Published by group.bmj.com)
patients with joint symptoms (Hilton et al., 1974), attention was drawn to the importance of drugs in the disturbance of hepatic function, particularly salicylates, indomethacin, phenylbutazone, gold and azathioprine, all anti-rheumatic agents. The one case of giant cell arteritis included in the review showed hepatic dysfunction and this was believed to be due to azathioprine therapy. No liver biopsy was performed. Previous reports of histopathological changes in the liver have included two with normal appearances and two with fatty change (Dickson et al., 1973) and one with a granulomatous hepatitis (Long and James, 1974). There now seems to be a case for linking hepatic dysfunction with the polymyalgia rheumatica, giant cell arteritis syndrome. It is only by following up such cases with raised alkaline phosphatase levels, with liver biopsies where reasonably indicated in these elderly patients, that a full assessment of this complication can be made.

References
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*Postgrad Med J* 1976 52: 236-238
doi: 10.1136/pgmj.52.606.236

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