one patient with tricuspid incompetence due to a ruptured papillary muscle after myocardial infarction. The higher incidence of left ventricular papillary muscle dysfunction reported (Heikkilä, 1967) must be largely due to the higher incidence of left ventricular infarction observed by various workers (Wartman & Hellerstein, 1948; Zaus & Kearns, 1952).

This patient had minimal dilatation of the tricuspid valve ring and the valves at post-mortem appeared competent. The necropsy impression is occasionally misleading as the valve ring may be larger during life, when the pressure in the right ventricle is increased. However, in this patient tricuspid incompetence persisted, although the clinical signs of heart failure were mild, although such failure as was present improved with treatment, and despite a decrease in heart size on the chest X-ray. This clinical course in conjunction with the autopsy findings make papillary muscle dysfunction the only tenable explanation for this patient’s tricuspid incompetence.

The mechanism of papillary muscle dysfunction in the left ventricle was reviewed by Burch, De Pasquale & Phillips (1963) and more recently by Raftery, Oakley & Goodwin (1966). The very prominent ‘v’ waves and ‘y’ descent in the jugular veins of this patient accord well with the left atrial pressure tracings found in patients with left ventricular papillary muscle dysfunction (Raftery et al., 1966). The right atrium in this patient was presumably normal before the sudden development of tricuspid incompetence and the venous pressure excursions were not damped because the atrium was still relatively uncompliant.

Diagnosis of the cause of tricuspid incompetence is important. When due to papillary muscle dysfunction it should be amenable to surgical treatment as is mitral incompetence (Fluck et al., 1966). If this patient had survived with persistent tricuspid incompetence the question of prosthetic valve replacement would have been seriously considered.

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References


Intrahepatic granulomatous arteriopathy

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Summary

The case history is detailed of a 55-year-old woman who was investigated for persistent pyrexia and anaemia. A liver biopsy specimen showed an unusual lesion of the small intrahepatic arteries. Many of these vessels showed circumferential replacement of their adventitial coat by a non-caseating granuloma whilst others showed localized granulomata focally interrupting the adventitial coat. The medial and intimal coats of the affected arteries were normal.

This was an intrahepatic granulomatous arteriopathy of unknown origin. The patient responded promptly and completely to steroid therapy.

Introduction

It is the purpose of this report to describe an unusual form of arteriopathy involving the intrahepatic vessels.

Case report

A woman, aged 55, was seen at Manchester
Royal Infirmary in August 1965. She had a previous history of inguinal herniorrhapathy in 1951, hysterectomy for uterine fibroids in 1958, left tympanotomy for otosclerosis in 1962 and appendicectomy for acute appendicitis in 1964. She now gave a 2-month history of headaches, vomiting, nausea for sweet and fatty foods, anorexia, weight loss, fatigue and depression.

On examination, she was pale and sallow but, apart from some tenderness in the right hypochondrium, there was no other abnormality. Her haemoglobin was 9·8 g/100 ml and her white cell count was 11,400/mm³ with a normal differential count. A barium meal and a chest X-ray showed no abnormality.

The patient was treated with oral iron, but there was no symptomatic improvement, and when seen in September 1965 her haemoglobin had fallen to 6·9 g/100 ml and her ESR was 66 mm in 1 hr. She was therefore admitted to hospital for more detailed study.

Re-examination revealed no further abnormality, though there was still some tenderness in the right hypochondrium and it was thought by some observers that her liver was just palpable. She had a persistent pyrexia, her temperature usually being about 100°F, but rising on occasions to 102–103°F. Numerous investigations were performed, the majority of which gave negative results.

The positive findings were an iron-deficiency anaemia, a persistently elevated erythrocyte sedimentation rate and a leucocytosis. Liver function tests showed a serum albumen of 2·5 g/100 ml, globulin 4·0 g/100 ml, alkaline phosphatase 22 K.A. units/100 ml, serum bilirubin 0·3 mg/100 ml and serum glutamic pyruvate transaminase 35 u/100 ml. The bromsulphalein retention was 33% at 45 min on one occasion and 23% when repeated. Multiple radiological studies failed to reveal any abnormality in the chest, gastrointestinal tract or urinary system, and tests for LE cells, antinuclear factor and rheumatoid factor were consistently negative. There was no serological evidence of syphilis. The routine investigations for a pyrexia of unknown origin proved unrewarding except for the finding of a positive brucellin skin test. However, on no occasion, either before or after this test, did the patient have a positive brucella antibody titre.

A diagnostic impasse having been reached, a laparotomy was performed in November 1965. At operation, all abdominal organs appeared normal and no masses were present. Biopsy specimens were taken from jejunum and liver.

**Biopsy findings**

The jejunal biopsy specimen showed no histological abnormality. The specimen of liver was a wedge of tissue measuring 1 X 1 X 0·5 cm. Histologically, the hepatic parenchyma appeared normal; there was no evidence of liver cell necrosis or fibrosis and no granulomatous lesions were present in the parenchyma. The hepatic sinuses were normal and there was no hyperplasia of the Kupfer cells. Most, but not all, of the medium-sized branches of the hepatic artery in the portal tracts of the specimen showed an unusual form of granulomatous inflammation of their adventitial coats. In many of the smaller of these arteries the adventitia was completely replaced by a non-casing granuloma involving the full circumference of the arterial wall (Fig. 1), while in slightly larger vessels only a portion of the adventitial coat was replaced by a granuloma, the remaining portion of the vessel wall appearing normal. In larger arterial branches discrete granulomata were seen in the adventitia (Fig. 2), there producing sharply localized destruction of the adventitial fibro-muscular tissue, with the intervening portions of the adventitia
presence of an unusual granulomatous condition affecting the arterioles it was impossible to make a definitive diagnosis. On clinical grounds the only investigation pointing to a possible aetiological agent was the positive brucellin skin test. The patient was therefore given 1 g of tetracycline daily for 4 months. This failed to produce any symptomatic improvement and at the end of this period the patient’s ESR was still 44 mm in 1 hr and her liver function tests were still abnormal. This treatment was therefore abandoned and the patient was started on prednisone, 30 mg daily. This produced an immediate alleviation of her symptoms, and when seen 3 months later in August 1966 she was well; her ESR had fallen to 3 mm in 1 hr. She has continued on a gradually reducing dose of prednisone and has remained well.

**Discussion**

The clinical features of this patient were non-specific and diagnosis was dependent on the findings in the liver biopsy specimen. These were difficult to interpret for the histological findings were not typical of any of the recognized arterial diseases. Thus, the absence of foreign-body giant-cells, medial damage, intimal thickening, elastic damage and fibrinoid necrosis was considered sufficient to exclude temporal arteritis, Takayasu’s arteritis, hypersensitivity angiitis and the vascular lesions of polyarteritis nodosa, lupus erythematosus, Wegener’s granulomatosis, systemic sclerosis, rheumatoid disease and dermatomyositis.

There was similarly no clinical or histological evidence that the patient was suffering from a generalized granulomatous disease in which the arteries were secondarily involved. The lesions in this case do indeed bear some resemblance to those occasionally seen in sarcoidosis (Erickson, Osom & Stern, 1948; Meyer, Foley & Campagna-Pinto, 1953). In the reported cases of sarcoid vasculitis, however, the granulomata have been basically parenchymal and vascular involvement has been an accidental result of enlargement of the granulomata, either arteries or veins being involved in an indiscriminate manner. In the present case in which granulomata were confined to the arterial walls and were totally absent from the hepatic parenchyma the diagnosis of sarcoidosis appeared unlikely.

The absence of caseation and of parenchymatous lesions also argued against tuberculosis and there were no findings to suggest syphilis, tuberaemia, histoplasmosis or erythema nodosum. The patient had not been exposed to beryllium. Despite the positive brucellin test it was difficult

![Fig. 2. A localized granuloma, which is focally interrupting the adventitia of an intrahepatic artery. H & E, × 190.](image-url)
to support the diagnosis of brucellosis for the agglutination test was persistently negative and the patient, though improving promptly with steroid therapy, failed to improve with prolonged administration of tetracycline.

It was therefore concluded that the patient was suffering from an intrahepatic granulomatous arteriopathy of unknown aetiology. Biopsies of other organs were not performed and hence it was not known whether this was an arteriopathy confined to the hepatic vessels or whether the liver biopsy allowed the opportunity to observe a local manifestation of a generalized disease. There were no clinical features to indicate specific involvement of any other organ but the persistent and unexplained anaemia indicates that the lesion, even if localized, was accompanied by systemic effects.

The prompt response to steroid treatment in this patient suggests, in the absence of any other obvious aetiological factors, a hypersensitivity reaction, though the arterial lesions did not resemble those described as typical of hypersensitivity states (Chure & Strauss, 1951).

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References


Sideropenic anaemia with reticulo-endothelial siderosis in a case of hypernephroma

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Summary

A case of sideropenic anaemia with reticulo-endothelial siderosis is described. At the time of admission the anaemia was severe and the cause was not apparent. Extensive search had to be made to reveal the neoplasm, a hypernephroma.

Introduction

Of the various types of anaemias encountered in patients with neoplastic disease, sideropenic anaemia with reticulo-endothelial siderosis is the most interesting and puzzling. According to Cartwright (1966), it is usually mild in degree and not progressive in severity, but when the anaemia is moderately severe the causal neoplasm is almost always obvious. The case reported here presented with severe anaemia long before the underlying neoplasm could be detected.

Case report

The patient, a man aged 58, a caterer by profession, was admitted with a 3 months' history of increasing tiredness, generalized weakness and shortness of breath on exertion. He had lost approximately 1 stone of weight in the 6 months preceding admission. He had never had indigestion and his appetite had always been good. The only relevant past history was that he had contracted syphilis about 30 years before.

On examination, he appeared very anaemic, but not jaundiced. The tongue, buccal mucous membranes, nails, palms and soles of the feet appeared pale but otherwise normal. There were no palpable glands or sternal tenderness. The skin and joints were normal. The liver was just palpable, one finger breadth below the right costal margin, smooth and not tender. All the features of a hyperkinetic circulation were present, with moderate tachycardia, slightly raised JVP, hyperdynamic cardiac impulse, flow murmurs in all cardiac areas, wide pulse-pressure, warm hands and digital pulsion. The only other abnormal physical finding was the presence of Argyll Robertson pupils—evidence of neurosyphilis. Capillary fragility test, rectal examination and proctoscopy were normal.
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