Heatstroke and rhabdomyolysis presenting as fulminant hepatic failure

Sarah Fidler,1 Elizabeth Fagan,1 Roger Williams,1 I. Dewhurst2 and C.E. Cory3

1The Liver Unit, King's College Hospital, Denmark Hill, London SE5 8RX, 2RAF Cranwell, Cranwell, Lincs NG34 8RN and 3Grantham & Kesteven Hospital, Grantham, Lincs NG31 8DG, UK.

Summary: Heatstroke following vigorous exercise is reported in a young male. Severe hepatic and renal failure resulted but the patient completely recovered following treatment by active measures to reduce his core temperature.

Introduction

Heatstroke associated with rigorous exercise is well documented especially during initial training1 and biochemical evidence of renal impairment and hepatic dysfunction including abnormal levels of blood urea, serum aspartate aminotransferase (AST) and hyperbilirubinaemia are commonly detected.1, 2 The occurrence of established renal failure is rare, being reported in 4% of severe cases and is usually found with accompanying rhabdomyolysis.1–4 Only a few instances of severe hepatic necrosis and liver failure have been reported, usually in association with renal failure, and have a very poor prognosis.4

In the present report we describe a patient with heatstroke following vigorous exercise. He, in addition, developed massive hepatic necrosis and fulminant hepatic failure as well as renal impairment and muscle injury. Active measures to reduce his core temperature were followed by complete recovery.

Case report

A 23 year old male Royal Air Force cadet suddenly collapsed whilst he was on his first training run which involved running for four miles in physical training kit in high summer. On admission to hospital his core temperature was 40.6°C, rising to 41.6°C one hour later. He was disorientated and mildly jaundiced with evidence of marked bruising, muscle tenderness and pitting oedema of all limbs and abdominal wall. The initial blood pressure was 180/70 mmHg, pulse 160/min. An electrocardiogram showed a sinus tachycardia. On admission his haemoglobin was 13.5 g/dl, packed cell volume (PCV) 38.4, total white blood cell count (WBC) 11.7 x 10⁹/l, platelets 43 x 10⁹/l, fibrinogen degradation products (FDPs) >1:128. Serum K⁺ 4.1 mmol/l, Na⁺ 145 mmol/l, HCO₃, 20 mmol/l, urea 6.3 mmol/l, creatinine 173 μmol/l, glucose 3.5 mmol/l, pH 7.4, Pco₂ 3.6 kPa, Po₂ 12.8 kPa. AST 380 IU/l, bilirubin 23 μmol/l, alkaline phosphatase 166 IU/l, prothrombin 5 seconds prolonged, creatinine kinase >10,000, urinalysis: positive for haemoglobin, myoglobin, urobilinogen and bilirubin. Attempts to lower his temperature with tepid sponging, electric fans and packing in ice were unsuccessful. He was therefore treated with intravenous chlorpromazine 12.5 mg every 15 min, fentanyl and pancuronium prior to ventilation which reduced the core temperature to 38.8°C, 6 hours later. Over the ensuing 4 days he remained pyrexial (37.7–38.9°C core), hypertensive with a wide pulse pressure (190–200 mm Hg, systolic/70–90 mm Hg diastolic pressure) and remained confused despite treatment with chlorpromazine 5 mg/h, maintained ventilation, continuous cooling and fluid replacement. The jaundice deepened (Figure 1) with further elevation of serum AST and alkaline phosphatase. The prothrombin time became 40 seconds prolonged by day 2 and peaked at 67 seconds prolonged by day 3. On transfer to King's College Hospital (day 4), his hypertension and disorientation, attributed to cerebral oedema, responded to bolus doses (2 x 100 ml) of 20% mannitol over 24 hours and a subsequent diuresis. The chlorpromazine was discontinued on day 4 after the prothrombin time had begun to fall and by day 8, the platelet count had risen to

Correspondence: Roger Williams M.D., F.R.C.P.
Accepted: 3 September 1987

© The Fellowship of Postgraduate Medicine, 1988
157 x 10^9/l. His clinical condition improved rapidly and he was discharged from hospital on the 14th day of illness with complete clinical and biochemical resolution of liver and renal damage.

**Discussion**

Survival from heatstroke after this degree of hepatic and renal failure must be very uncommon, particularly in association with marked rhabdomyolysis.1-4 The rise in serum AST and creatinine could be attributed partly to a skeletal muscle component but the progressive elevation in prothrombin time was indicative of severe hepatic dysfunction. Some degree of disseminated intravascular coagulation was apparent from the fall in platelet count and elevation in FDPs although these changes are also common in fulminant hepatic failure.5 The confusion may have been related, at least early on, to the heatstroke. Later, his disorientation was probably confounded by the hepatic encephalopathy since his mental state rapidly improved with treatment for cerebral oedema. Sustained systolic hypertension is not a feature of heatstroke per se but, in the context of this case, was more typical of cerebral oedema in association with fulminant hepatic failure6 and characteristically responded to administration of mannitol.

Rapid, effective cooling in heatstroke is mandatory but the optimal method remains controversial. Packing in ice may have been detrimental by effectively insulating the patient’s core and hence delaying the temperature drop. Control of muscle hyperactivity to limit further generation of heat by paralysis with non-depolarising agents and chlorpromazine was almost certainly life-saving but required 4–5 days to be effective.
References


