Cytoplasmic vacuolation of peripheral blood cells in acute alcoholism

RJ DAVIDSON AND JL McPHIE

From the Haematology Unit and Department of Pathology, University of Aberdeen, Foresterhill, Aberdeen AB9 2ZD, UK

SUMMARY Cytoplasmic vacuolation of peripheral blood leucocytes is reported in three patients with acute alcoholism. Apparently this is the first report of vacuolation of peripheral blood leucocytes associated with acute alcoholism. Apart from any direct cytotoxic effect of alcohol the metabolic acidosis and hypoglycaemia associated with acute alcoholic poisoning may be important factors in the pathogenesis of the morphological changes.

The association of alcoholism with reversible vacuolation of red cell and, less consistently, white cell marrow precursors is well recognised.1 When experimentally induced by ethanol administration to human volunteers such vacuolation appears to be dose-related, and its development in the presence of folate supplementation and normoblastic haemopoiesis seems to indicate that it is due to a direct toxic effect of alcohol.2

We report, apparently for the first time, vacuolation in granulocytes and lymphocytes in the peripheral blood in three alcoholics, two of whom succumbed to acute alcoholic poisoning.

Case reports

Case 1
A 38-year-old man with a long history of alcoholism was admitted to hospital after being found unconscious in his home after a bout of drinking thought to have lasted for six days. On admission he was comatose and his blood pressure unrecordable. The blood alcohol concentration measured some 36 hours later was still 1200 mg/l. Examination of the cerebrospinal fluid (CSF) was normal. Clinically he was diagnosed as having acute liver failure with hypoglycaemia and metabolic acidosis. Confirmatory biochemical results included arterial blood pH 7·4; serum bilirubin 66 µmol/l; aspartate amino transferase (AAT) > 100 IU/l; alkaline phosphatase 115 IU/l; plasma glucose < 1·2 mmol/l; blood lactate 65·3 mg/l and pyruvate 22·5 mg/l. The Coulter S peripheral blood profile showed WBC 7·0 × 10⁹/l, RBC 3·3 × 10¹²/l, Hb 10·8 g/dl, PCV 0·37, MCV 111 fl, MCH 33 pg, and MCHC 29 g/dl. Platelets were 79 × 10⁹/l and reticulocytes 4·2%. The differential WBC comprised neutrophils 74%, lymphocytes 22%, monocytes 4%. On microscopy the red cells were generally normochromic but included some round macrocytes. Polychromatic forms were increased and of the occasional late normoblasts detected some contained multiple cytoplasmic vacuoles (Fig. d). Neutrophils showed prominent cytoplasmic vacuolation but the usual accompanying features of toxic or reactive change—namely, a nuclear shift to the left and toxic granulation—were absent. Vacuolation was also present in a significant number of monocytes and lymphocytes (Table).

After intravenous electrolyte and fluid replacement his systolic BP rose to 100 mm Hg and his depth of unconsciousness lightened. But he remained anuric and died some 24 hours after admission. Necropsy 20 hours after death revealed subcutaneous crepitus of neck, shoulder, and scrotal tissues. Both lungs were grossly oedematous. The liver (2100 g) was pale and fatty and its cut surface showed a micronodular cirrhotic pattern. Both kidneys displayed cortical pallor and the terminal ileum showed a 5-cm long segment of congestion with subserosal gas bubbles. Histological examination confirmed the presence of pulmonary oedema, an active micronodular hepatic cirrhosis, and bilateral renal cortical necrosis. Staining by the Oil-Red-O technique showed severe fatty change in hepatocytes, focal fatty vacuolation of subendocardial muscle fibres,
Vacuolation of peripheral blood cells in (a) neutrophil, (b) monocyte, (c) monocyte and lymphocyte, and (d) nucleated red cell adjacent to non-vacuolated 'band' neutrophil (magnification × 1000).

and fat accumulation in proximal renal tubular cells. Swabs taken from the spleen yielded a growth of Clostridium welchii.

**CASE 2**

A 32-year-old married woman with a long and tragic history of alcohol abuse had had several previous hospital admissions because of episodes of delirium tremens or liver failure. Her final admission was after she was found unconscious in her home after a bout of drinking of uncertain duration. She was moribund, BP 70/50 mm Hg, deeply jaundiced, and her blood alcohol concentration was 990 mg/l. She was diagnosed as suffering from acute-on-chronic alcohol-induced liver damage with encephalopathy, metabolic acidosis, and hypoglycaemia. Biochemical findings included standard bicarbonate 17 mmol/l; urea 8-4 mmol/l; plasma glucose < 1-0 mmol/l; serum bilirubin 123 µmol/l; AAT 470 IU/l, and alkaline phosphatase 188 IU/l. The Coulter S peripheral blood profile was WBC 11-7 x 10⁹/l, RBC 3-1 x 10¹²/l, Hb 10-8 g/dl, PCV 0-33, MCV 106 fl, MCH 36 pg, MCHC 33 g/dl. Platelets were 96 x 10⁹/l and reticulocytes 4%. The differential WBC comprised neutrophils 75%, lymphocytes 18%, monocytes 5%, and eosinophils 2%. On microscopy the red cells were normochromic but included some round macrocytes and stomatocytes. Neutrophils showed prominent cytoplasmic vacuolation with toxic granular change but no accompanying nuclear shift to the left. Vacuolation was again noted in many monocytes and lymphocytes (Table). Despite intensive resuscitative measures the patient died within 18 hours of admission. Necropsy 39 hours after death revealed patchy haemorrhages in both upper lung lobes, oesophageal varices, splenic enlargement (280 g) with congestion, and a fatty liver (1350 g) with a pattern of micronodular cirrhosis. Histological examination confirmed pulmonary oedema and focal haemorrhages consistent with aspiration of blood, splenic changes of portal hypertension, a micronodular cirrhosis with marked

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<th>Case No.</th>
<th>Vacuolated cells (%)</th>
<th>Neutrophils</th>
<th>Monocytes</th>
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<tr>
<td>1</td>
<td>56</td>
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cholestasis, and alcoholic hepatitis. Lipid staining again revealed gross fatty change within hepatocytes, focal fatty degeneration of myocardial fibres, and fatty vacuolation of proximal renal tubular cells.

CASE 3

A 53-year-old woman office cleaner was admitted as an emergency after being found at work in a collapsed and confused state. She had a history of alcoholic abuse for one year. When she arrived for work on the day of her admission to hospital she was seen to be 'shaky and trembling'. On arrival in the casualty department she was confused and aggressive with delusional features consistent with an alcohol withdrawal syndrome despite an absence of alcohol in the blood. An empty whisky bottle was found in her bag. After treatment and surveillance in a medical ward for 48 hours her behaviour necessitated her transfer to a psychiatric unit for further treatment of her withdrawal symptoms.

The biochemical profile included normal plasma electrolytes and urea; blood glucose 4.8 mmol/l; serum bilirubin 11 μmol/l; AAT 84 IU/l; alkaline phosphatase 80 IU/l, and gamma-glutamyl-transferase 28 U/l. Skull and chest radiographs, brain scan, and blood cultures were all normal. The Coulter S peripheral blood profile revealed WBC 5.7 × 10⁹/l, RBC 4.0 × 10¹²/l, Hb 12.5 g/dl, PCV 0.4, MCV 101 fl, MCH 31 pg, MCHC 31 g/dl. Platelets were 139 × 10⁹/l and reticulocytes 2%. The differential WBC comprised neutrophils 77%, lymphocytes 17%, monocytes 5%, and eosinophils 1%. On microscopy the red cells were normochromic but included some round macrocytes and stomatocytes with very occasional target cells. Neutrophils again showed prominent cytoplasmic vacuolation without a nuclear shift to the left or toxic granular change. A smaller number of monocytes and lymphocytes displayed vacuolar change (Table).

Discussion

The vacuolation of bone-marrow cells in acute alcoholism is reversible, occurs predominantly in red cell precursors, affects early rather than late forms (mainly pronormoblasts and promyelocytes), and the cytoplasmic rather than nuclear component. These changes have also been noted in marrow cells of premature infants born to mothers who had intravenous alcohol administered to delay the onset of labour. The vacuoles do not stain positively for fat, mucopolysaccharide, DNA, RNA, peroxidase, or acid and alkaline phosphatase. Similar marrow vacuoles have been reported in association with chloramphenicol administration, in experimentally induced riboflavin deficiency and phenylalanine deficiency, and in the course of treatment of phenylketonuria. Riboflavin or phenylalanine deficiency is not thought to be involved in the formation of vacuolation in acute alcoholism. Cytoplasmic and nuclear vacuolation of only red cell precursors may be prominent in erythraemic myelosis, while their presence in only the primitive white cells is usual in bone marrow involvement in Burkitt's lymphoma and occasional in acute leukaemias.

Unfortunately we were unable to ascertain the marrow cytology in our patients and hence to compare any changes with those in peripheral blood. In cases 1 and 2 the marrow obtained at necropsy was too autolysed for valid study, while marrow aspiration was not attempted in case 3 because of the patient's disturbed mental state. In the peripheral blood degenerative vacuolation of neutrophils, monocytes, and lymphocytes may be discernible in the stained film when prepared from blood which has been allowed to stand, particularly at room temperature, for more than three hours after collection. Vacuolation of neutrophils and monocytes is most commonly associated with infective, especially septicaemic, states and toxic conditions including metabolic disturbances such as diabetic ketoacidosis. Similar vacuolar changes have also been reported in progressive muscular dystrophy. In contrast, vacuolation of peripheral blood lymphocytes is unusual and apart from its occasional presence in viral infections, including infectious mononucleosis, it is, as described, virtually limited to a number of rare hereditary disorders—sphingomyelin lipidosis (Niemann-Pick disease), ganglioside lipidosis (Tay-Sach and Batten-Spielmeyer-Vogt diseases), type II glycogen storage disease (Pompe's disease), and genetic mucopolysaccharidosis (Hurler-Hunter syndrome).

Both quantitative and functional granulocytic changes with predisposition to infection are well documented in alcoholics and in case 1 we cannot exclude that clostridial infection contributed to the leucocytosis and red cell changes, although its detection only at necropsy 20 hours after death favours an alogical event. In cases 2 and 3, however, there was no clinical or laboratory evidence of infection and we could not therefore attribute the leucocyte changes to any of the aforementioned causes other than direct toxicity or the metabolic disturbances induced by the acute alcoholism. This conclusion is supported by recent reviews which have not only incriminated alcohol as a direct haematoxinin but suggested that the pathogenesis of its effects on haemopoietic cells may not be different from that on any other body cell or tissue. Our present findings not only support this concept but extend the
spectrum of cytotoxic change to include peripheral blood leucocytes and in severe cases even the mature red cell.

We would further postulate that, apart from the 'direct' toxic effect of alcohol and its metabolites, the metabolic acidosis and hypoglycaemia21 induced by acute alcoholism may be important factors contributing to the cell damage by impairing and, in severe cases, largely inhibiting glycolysis and other metabolic processes essential to normal cell function and membrane integrity. Such a sequence of events is analogous to that in diabetic ketoacidosis where a similar spectrum of morphological and functional change may be manifest in different tissues or cells ranging from cloudy swelling to fatty vacuolation in hepatocytes and toxic granulation to vacuolation in peripheral blood neutrophils.14

Finally, our observations of vacuolar change in peripheral blood leucocytes is supported by experimental studies in dogs22 fed varying amounts of alcohol and where marked cytoplasmic vacuolation and nuclear pyknosis were not only noted in both peripheral blood lymphocytes and granulocytes but the frequency and degree of these morphological changes were clearly dose-related.

We thank Dr AM Lessells for the necropsy report in case 1 and several of our clinical colleagues for permission to publish details of patients under their care.

References

5 McCurdy PR. Chloramphenicol bone marrow toxicity. JAMA 1961;176:588-93.

Requests for reprints to: Dr RJ Davidson, Haematology Unit, Department of Pathology, University Medical Buildings, Foresterhill, Aberdeen AB9 2ZD.