Observations on copper associated protein in childhood liver disease

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SUMMARY Hepatic copper concentrations were compared with staining grades of copper associated protein (CAP) and histochemical copper in liver sections from 44 patients (one fetus, one pre-term infant, four term infants, eight normal children, 16 children with various liver diseases, and 14 patients with intrahepatic cholestasis of childhood (IHCC)). A similar comparative study of hepatic copper concentration with CAP and histochemical copper was performed in 21 patients with Wilson’s disease. CAP occurred in the fetus, pre-term infant, and term infants without liver disease. This suggests that CAP is a normal constituent of the hepatocyte and is not a consequence of liver disease or biliary obstruction. CAP was not seen when hepatic copper concentration was normal; it was absent in eight children with no evidence of liver disease, eight children with non-cirrhotic liver disease, and seven of eight children with cirrhosis. When hepatic copper concentration exceeded 4.0 μmol/g dry liver weight grade 2 or grade 3 staining for CAP and histochemical copper was found in the fetus, pre-term infant, infants, and IHCC. CAP was found in IHCC only in the presence of raised hepatic copper levels, supporting evidence of a relationship between copper and CAP. In 17 of 21 patients with Wilson’s disease hepatic copper concentrations exceeded 4 μmol/g. Positive staining for CAP was seen in seven of these patients being usually grade 1. CAP is a normal associated protein, present when hepatic copper concentrations are increased in normal liver cells. It is usually absent in hepatocytes from Wilson’s disease despite similar hepatic copper levels. CAP may represent material which protects the hepatocyte from the toxic effects of copper.

Sipponen et al.1 noted a new material in liver sections from patients when stained with the orcein stain of Shikata.2 This material was present in 83 % of patients with primary biliary cirrhosis (PBC), 20% of patients with active chronic hepatitis, and a few patients with long-standing cirrhosis. It appeared as dark brown granules distinguishable from HBsAg and elastic tissue by staining pattern and location, and occurred in the same cellular and subcellular location as copper. Other materials such as lipofuscin and globules of α-antitrypsin are not stained by orcein.3 The protein nature of this new material was suggested by histochemical stains and it was named copper associated protein (CAP).5 CAP has been considered an abnormal copper binding protein.4 Measured copper concentration in liver sections from patients with PBC has been shown to correlate with grades of staining of CAP and histochemical copper.5 However, this relationship has not previously been systematically studied and reported in children. Hepatic copper concentration is physiologically raised in fetuses and newborn infants6 and remains raised until 6 to 12 months of age.7 Hence, liver sections from a neonatal group have been examined for CAP and histochemical copper in the present study.

In Wilson’s disease, copper accumulation in the liver (usually >4 μmol/g of dry liver) and other organs is associated with tissue damage.8 The relationship between the measured hepatic copper concentration and grades of staining for CAP, and histochemical copper was assessed in patients with Wilson’s disease and the findings compared with other groups.

Methods

STUDY GROUPS
Sixty-five liver sections were assessed from (1) neonatal group (six) (2) childhood group (38), and (3) Wilson’s disease (21).
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Neonatal group
All six patients died from non-hepatic causes and included one fetus of 27 weeks’ gestation, the result of an antepartum haemorrhage, and one pre-term infant of 35 weeks’ gestation with congenital heart disease (CHD) who died on the third post-partum day from heart failure. The four term infants included one child with Down’s syndrome who died on the first day post-partum from a tear in the faél cerebi, one infant with a tracheo-oesophageal fistula who died from respiratory failure at 3 weeks of age, and two infants with CHD who died from heart failure at 7 days and 2 months respectively. Wedge samples were obtained at necropsy in all cases.

Childhood group
The 38 children were at least 18 months old, and each group was age-matched. No child had received copper chelation therapy.

Eight children were ultimately proved to have no evidence of liver disease. Percutaneous liver biopsies were obtained in six and wedge samples were obtained at necropsy in two.

Fourteen children suffered with intrahepatic cholestasis of childhood (IHCC). Of the 14 patients in this group, 13 developed jaundice in the first 3 months of life and one presented with pruritis at 3 years of age. Their ages at the time of study ranged from 18 months to 22 years. Liver specimens were obtained by percutaneous biopsy in 13 and at necropsy in one. Diagnosis was based on clinical, biochemical, and biopsy evidence. Extrahepatic biliary obstruction was excluded in all cases: in nine by laparotomy and in five by percutaneous transhepatic cholangiography and ultrasound examination. A family history of liver disease was obtained only in one case—a brother with IHCC had died after haemorrhage from oesophageal varices. Pulmonary flow murmurs were present in two children and one child also had congenital glaucoma. In these two children there was no evidence of pulmonary valves stenosis as assessed clinically, with ECG, radiograph of the chest, or echocardiography. Arteriography was not performed. All cases were negative for hepatitis B surface antibody and hepatitis B core antibody. One patient, homozygous for α1-antitrypsin deficiency, had cholestasis from the neonatal period. Children with characteristic facies and cardiac lesions, Byler’s syndrome, or arteriohepatic dysplasia were not present in this series. The patients were of various nationalities: six English, three Italians, three Arabs, two Greeks, and one Portuguese. None had had portal systemic shunt surgery.

Eight children suffered from cirrhosis—no cause was identified in five. The test for HBsAg was positive in one child; one child was homozygous for α1-antitrypsin deficiency; one child had fatty cirrhosis, possibly due to type III glycogen storage disorder.

Eight children suffered from various liver diseases excluding cirrhosis, biliary cirrhosis, and Wilson’s disease.

Wilson’s disease
Among the 21 patients in this group there were four sibling pairs (nos. 1, 5; 6, 10; 8, 17; 13, 21). The diagnostic pre-penicillamine biopsy was studied in 11.

Techniques
Tissue copper concentration was determined by neutron activation analysis. The instruments used for taking liver samples were washed with 1% solution of ethylenediaminetetraacetic acid (EDTA) rinsed with ion-free water, and sterilised in a similarly prepared glass container. Copper contamination of necropsy wedge samples was specifically avoided. Hepatic copper concentration was expressed as μmol/g dry liver weight. The coefficient of variation for duplicate measurements of hepatic copper concentration measured by neutron activation analysis is 7% to 15%.

A portion of each liver sample was used for estimation of copper by neutron activation analysis (NAA). The remainder was processed for paraffin sectioning and stained for copper associated protein and tissue copper by the methods of Shikata et al. and Lindquist respectively. If positive staining occurred, the degree was semi-quantitatively assessed at 1, 2, or 3 for mild, moderate, or heavy staining. Absent staining was grade 0.

Histological semi-quantitative grading was performed without prior knowledge of liver copper concentration.

The Mann-Whitney test was used for statistical analysis of hepatic copper levels in the six groups of patients.

Results

Childhood controls
In eight histologically normal liver biopsies the hepatic copper concentration was 0.5 (0.2-1.5) μmol/g dry liver (median and range). This value is similar to figures quoted for normal adults. Liver sections did not stain positively with orcein or rhodanine.

Neonatal group
Hepatic copper concentration was significantly increased in six, being 5.5 (2.2-10.7) μmol/g
Copper associated protein. Typical granules of CAP observed with orcein staining in liver sections from pre-term infants of 35 weeks' gestation. Hepatic copper concentration was 6·9 μmol/g × 300.

![Image: Diagram showing neonatal and childhood groups, excluding Wilson's disease: grades of copper associated protein (orcein) staining and hepatic copper concentration.](image)

**Fig. 2** Neonatal and childhood groups, excluding Wilson's disease: grades of copper associated protein (orcein) staining and hepatic copper concentration. Each group is represented by a different symbol. The hatched area represents normal hepatic copper concentration. A liver copper concentration exceeding 4 μmol/g is considered to be one of the diagnostic features of Wilson's disease.

Fig. 1 Copper associated protein. Typical granules of CAP observed with orcein staining in liver sections from pre-term infants of 35 weeks' gestation. Hepatic copper concentration was 6·9 μmol/g × 300.

Liver histology revealed no evidence of acute or chronic liver disease. Positive staining with orcein and rhodanine was seen in the fetus, pre-term infant, and the four term infants (Fig. 1). CAP seen in these cases appeared to be identical with that described in adults.

**INTRAHEPATIC CHOLESTASIS OF CHILDHOOD (IHCC)**

Hepatic copper concentration was significantly increased in 12 of 14 children being 5·1 (0·9-16·2) μmol/g (p < 0·001). Typical orcein positive granules of CAP were seen in the 12 with raised hepatic copper levels (Fig. 2). Positive staining with rhodanine occurred in 11 of 12 patients with raised hepatic copper levels. Liver sections from one patient with slightly increased hepatic copper concentration did not stain with rhodanine and sections from one patient with normal hepatic copper concentration showed grade 1 staining (Fig. 3).

**LIVER DISEASE**

Liver copper levels in eight children with non-cirrhotic liver disease and eight children with cirrhosis were 0·4 (0·2-1·2) μmol/g and 0·7 (0·3-1·1) μmol/g respectively. These values are in the normal range. Liver sections did not stain positively with orcein or rhodanine in children with non-cirrhotic liver disease.

Sections from one patient with cirrhosis (homozygous α1-antitrypsin deficiency) revealed grade 2 orcein staining with normal hepatic copper concen-
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Fig. 3 Neonatal and childhood groups, excluding Wilson’s disease: grades of histochemical copper (rhodanine) staining and hepatic copper concentration. Each group is represented by a different symbol. The hatched area represents normal hepatic copper concentration. A liver copper concentration exceeding 4 μmol/g is considered to be one of the diagnostic features of Wilson’s disease.

WILSON’S DISEASE

Positive staining patterns of CAP in Wilson’s disease differed qualitatively and semi-quantitatively from the preceding groups (Table). If CAP staining occurred, it was distributed irregularly throughout the hepatic parenchyma.

Hepatic copper concentration was significantly increased being 8.69 (1.91–28.94) μmol/g (p<0.001) and in 17 patients exceeded 4 μmol/g. However, CAP was seen in only seven patients. Grade 1 staining was most common, being present in four patients, and grade 3 staining occurred in one patient (hepatic copper concentration 8.69 μmol/g).

In two patients CAP was absent when liver sections appeared normal and hepatic copper levels were high (nos. 11, 12). CAP was frequently associated with cirrhosis, being present in seven of 11 patients with cirrhosis. Hepatic copper concentrations in the group with cirrhosis was 6.94 (1.91–22.52) μmol/g, which was not significantly different from cases without cirrhosis 9.22 (2.63–28.94) μmol/g. CAP appeared more common in the pre-treatment group. The hepatic copper concentration in the pre-penicillamine group was 11.15 (1.91–28.94) μmol/g, which was not significantly different from patients with treated Wilson’s disease 5.65 (2.63–15.43) μmol/g or patients with IHCC 5.1 (0.9–16.2) μmol/g.

Four siblings were studied. Similar staining patterns for CAP were seen in siblings in all families despite large differences of hepatic copper levels and variable histological appearances. Thus CAP was not seen in both siblings from two families (nos. 6, 10 and 8, 17), grade 1 CAP staining occurred in both siblings from one family (nos. 13, 21) and moderate staining was detected in both siblings (nos. 1, 5) of the fourth family.

There was no correlation between grade of histochemical staining for copper and hepatic copper concentration measured by neutron activation analysis. Ten of the 21 patients had positive staining for copper, being usually grade 1. Grade 3 staining for copper occurred in only two patients and hepatic copper concentrations were 6.14 μmol/g and 8.69 μmol/g.

Discussion

Measurement of liver copper concentration is not widely available.15–18 Histochemical assessment...
of hepatic copper concentration, if reliable, would provide clinically useful information more readily. The orcein stain of Shikata and rhodanine stains appear to demonstrate different components of a copper-protein complex in liver sections. Similar staining grades for copper and copper associated protein (CAP) were usually observed in each liver section in the study of neonatal and childhood groups. When hepatic copper levels were normal positive staining for copper and CAP was usually absent. However, staining for CAP appeared more discriminatory, being positive in one instance only when hepatic copper concentrations were normal. This patient suffered from cirrhosis with homozygous α1-antitrypsin deficiency. α1-antitrypsin globules are not detected by the orcein stain and the discrepancy may be related to different sections of the biopsy being assessed histologically and by neutron activation analysis. There was a relationship between measured hepatic copper concentrations and grades of staining for tissue copper and CAP. Usually, slight rises of hepatic copper concentration (<4 μmol/g) were associated with grade 1 staining and large increases of hepatic copper levels (>4 μmol/g) were associated with grade 2 or grade 3 staining. A wide range of hepatic copper concentrations occurred with grade 2 or grade 3 staining, thus limiting useful interpretation of grades of histological staining in individual patients.

All patients in the neonatal group were less than 3 months of age. The hepatic copper concentrations were physiologically increased and liver sections appeared normal. The study of this group revealed typical granules of CAP in all cases, suggesting that it is also physiological and represents a normal hepatocyte protein rather than an abnormal tissue copper protein (metallothionein) as previously suggested. CAP has been described in primary biliary cirrhosis and chronic liver disease. Histologically, all liver sections in the neonatal group were normal, so demonstrating that CAP is not an artefact of chronic biliary obstruction or liver disease.

CAP was seen in one fetus of 27 weeks gestation demonstrating that it develops in utero, possibly at an early age stage, but more studies will be required to confirm this observation. The biochemical composition of CAP, which is located in lysosomes, is not established. It is unlikely to be caeruloplasmin, which has been localised to the endoplasmic reticulum, but may be similar to a copper-rich form of metallothionein isolated from lysosomes of normal newborn bovine liver.

The study of copper and CAP in the neonatal and childhood groups revealed additional evidence of an association. The changes in hepatic concentration were associated with similar changes in positive orcein staining. Thus CAP was seen in the neonatal group with physiologically raised liver copper levels and was absent in the childhood control group and 15 of 16 children with liver disease and normal hepatic copper concentrations. CAP was found in 12 children with IHCC and raised hepatic copper concentrations, but was not seen in two patients with IHCC and normal hepatic copper concentrations. The appearance of CAP in cases of IHCC with raised hepatic copper concentration might suggest that copper has induced the appearance of a normal copper protein (CAP) (similar effects have been noted experimentally). Similar sequence of events may occur in primary biliary cirrhosis. It is possible that the amount of induced protein is proportional to the copper concentration. Supporting this suggestion is the trend of increasing hepatic copper levels to be associated with increasing grades of CAP staining in these groups.

Hepatic copper concentrations in Wilson’s disease were similar to IHCC and neonates, but positive histochemical staining for copper was infrequent, suggesting that the distribution of copper accumulation in Wilson’s disease may be different. This observation is similar to previous studies. Also, positive staining for CAP in Wilson’s disease was infrequent. In contradistinction with neonates and other forms of childhood liver disease a relationship between CAP and measured hepatic copper content was not found in Wilson’s disease. From data obtained in fetuses, neonates, and children it might be expected that CAP would be present in Wilson’s disease when hepatic copper levels were raised, particularly when the concentration exceeded 4 μmol/g. However, CAP was usually absent in Wilson’s disease, even when the hepatic copper concentration was as high as 28-94 μmol/g. CAP was more commonly associated with cirrhosis in Wilson’s disease. There may be several possible explanations: (1) there is a sampling error in liver biopsies. (2) Wilson’s disease is a pleomorphic disorder and the appearance of CAP may represent a subgroup. Possible support for this suggestion may be the observation that each sibship of the four families revealed similar staining patterns for CAP despite wide variations in hepatic copper concentration, treatment with d-penicillamine or histological appearances. (3) Alternatively, appearance of CAP in Wilson’s disease may be delayed until cirrhosis develops despite raised hepatic copper levels. Cirrhosis per se is not responsible as CAP occurred infrequently in children with cirrhosis of other aetiologies.

Hepatic copper concentrations are increased in fetuses, neonates, IHCC, and PBC. The
reason for the physiological increase of hepatic copper concentration in fetuses and neonates is unknown, while in IHCC and PBC it is probably secondary to chronic cholestasis. The increased hepatic copper levels in fetuses and neonates appear to be non-toxic. The hepatotoxicity of copper is not established in IHCC. Liver cells are normal in fetuses and neonates and raised hepatic copper levels are associated with CAP. In IHCC increased hepatic copper concentrations are always associated with CAP, a normal copper associated protein which is diffusely distributed through hepatic parenchyma. The primary disturbance of copper metabolism in the liver cells of Wilson's disease leads to increased hepatic copper concentrations, usually exceeding 4 μmol/g and liver damage. CAP was usually absent in Wilson's disease even when hepatic copper levels exceeded 4 μmol/g. Thus an absence or relative deficiency of CAP may contribute to the hepatotoxic effects of copper in Wilson's disease.

This study suggests that CAP represents a physiological copper associated protein which is present in normal hepatocytes (fetuses and neonates) and reappears in response to raised concentrations of copper, which may occur with chronic cholestasis thus minimising the toxic effects of an increased copper concentration within the hepatocyte. This response is impaired in liver cells of Wilson's disease and may explain the hepatotoxic effects of copper in this condition.

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