ORTHOTOPIC LIVER TRANSPLANTATION FOR WILSON'S DISEASE

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Summary
An 11-year-old boy with terminal hepatic failure due to Wilson's disease was treated 18 months ago with orthotopic liver transplantation. Postoperatively, there has been evidence of clearance of body copper stores but without accumulation of copper in biopsy specimens of the transplanted liver after 6 and 17 months. Further follow-up will be necessary before deciding whether the disorder has been cured by liver replacement and in turn whether this constitutes proof that Wilson's disease is an inborn error of hepatic metabolism. The observations so far are consistent with these conclusions.

Introduction
D-Penicillamine is recognised as the treatment of choice in Wilson's disease. However, even with therapy some patients are left with post-necrotic cirrhosis and portal hypertension. In addition, a small group of patients has been labelled as having "acute Wilson's disease"—an entity which has the distinctive hepatic histopathological feature of Mallory bodies, and an extremely high mortality rate despite treatment with specific copper-chelating agents. Moreover, even the more chronic form of Wilson's disease may lead to severe hepatic insufficiency, particularly in children, before the appearance of any diagnostic neurological or ocular signs.

We report here a case of Wilson's disease which was treated successfully by total hepatectomy and orthotopic homotransplantation of a cadaveric liver. There has been no clinical or biochemical evidence of recurrence of Wilson's disease in the one and a half years of postoperative follow-up.

Case-report
An 11-year-old boy was admitted on July 14, 1969, in hepatic pre-coma. He was said to have had a protuberant abdomen since birth. Although he had developed generalised pruritus at 8 months of age, the diagnosis of a liver disorder was not definitely established until admission to another hospital in April, 1966. At that time the cephalin flocculation was 4+; bromsulphalein retention was 30% at 45 minutes, and serum-alkaline-phosphatase was 15 Bodansky units (normal less than 5). An open liver biopsy was reported as "post-necrotic cirrhosis". In April, 1969, he was admitted to a second hospital with obvious advanced liver disease. There were no Kayser-Fleischer rings on slit-lamp examination. A second liver biopsy again showed cirrhosis, with the additional finding of the alcoholic hyaline of Mallory. This biopsy was interpreted as being consistent with Wilson's disease. The diagnosis of Wilson's disease was supported by the eventual histopathological study of the excised liver by members of our staffs and several consulting pathologists, including Dr. Hans Popper (New York) and Prof. K. A. Porter (London).

The patient was transferred to the University of Colorado Medical Center in hepatic pre-coma. He had anasarca and massive ascites. There were prominent venous collaterals on the abdomen, spider angiomas on the chest and upper extremities, and liver palms. Gynecomastia was not present. He was somnolent, disoriented, and incontinent with fetor hepaticus. He had marked clonus and was hyper-reflexic, but the Babinski responses were equivocal. He had no evidence of asterixis. The serum-bilirubin was 4.5 mg. per 100 ml., the serum-alkaline-phosphatase 35 Bodansky units (normal less than 5), S.G.O.T. 49 i.u. per litre (normal 3-27), S.G.P.T. 14 i.u. per litre (normal 1-30), serum-total-protein 4.5 g. per 100 ml., serum-albumin 1.5 g. per 100 ml., y-globulin 2.2 g. per 100 ml., prothrombin-time 31-5 seconds (control 13 seconds) or 13%. The haematocrit was 27%; white blood-cells 15,000 per c.mm.; platelets 30,000 per c.mm. The blood-ammonia was 432 mg. nitrogen per 100 ml. and the electroencephalogram was markedly abnormal, showing diffuse slow waves, compatible with a metabolic encephalopathy.

Within a few hours of his arrival the patient developed upper-gastrointestinal haemorrhage and became agonal. Although the preoperative diagnosis was Wilson's disease, no consideration was given to the possibility of d-penicillamine treatment, since death was imminent.

Transplantation
When the abdomen had been entered through a bilateral subcostal incision, several litres of ascitic fluid were aspirated. The liver was dark brown, firm, and the site of finely nodular cirrhosis. The organ was removed in its entirety and replaced with an orthotopic hepatic homograft from a cadaveric donor of approximately the same age as the recipient. The donor had no stigmata or family history suggestive of Wilson's disease and the serum copper and ceruloplasmin were within normal limits.

The excised native liver weighed 540 g., compared to a predicted 840 g. for a child of this age and weight. Bilary drainage was with choledochoduodenostomy. The recipient's greatly enlarged spleen was also removed. Grossly and microscopically there were multiple splenic infarcts.
Because of extensive venous collaterals, the operation was a difficult one, necessitating transfusion of 12 litres blood.

Post-transplantation Course

There was an immediate clearing of the sensorium, a fall in the serum-bilirubin and blood-ammonia levels, an increase in the total-serum-protein, and a reduction in the prothrombin-time. The patient was treated with a triple-drug immunosuppressive regimen that included azathioprine, prednisone, and heterologous antilymphocyte globulin (A.L.G.). Two severe rejection episodes occurred—one after a few days and the other after 3 weeks. The most prominent biochemical findings at these times were changes suggestive of biliary-tract obstruction, with an extreme degree of hyperbilirubinaemia (maximum 43 mg. per 100 ml.), of which most was glucuronide, and raised alkaline phosphatase (maximum 990 I.U. per litre) (fig. 1).

With reversal of the second rejection episode, these and the other adverse findings of rejection receded completely. The patient has had essentially normal hepatic function since 6 months post-transplantation (fig. 1).

Open biopsies of the homograft, obtained after 6 and 17 months, were studied for evidence of rejection by Prof. K. A. Porter, of St. Mary's Hospital and Medical School, London. The initial biopsy, apart from some cellular infiltration of the portal tracts and prominence of the Kupffer cells, was essentially normal. The second biopsy showed some excess reticulin and collagen in and around the portal tracts and also around some of the liver lobules. A few of the small hepatic-artery branches now showed some intimal thickening.

The patient is receiving 37.5 mg. azathioprine and 15 mg. of prednisone daily. He now has no demonstrable neurological impairment or evidence of Kayser-Fleischer rings on slit-lamp examination. No dietary limitations have

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**Fig. 1—Course after liver replacement.**
A.L.G. = horse antilymphocyte globulin.
been imposed, and d-penicillamine or potassium sulphide have never been administered except during the tests to be described below.

Studies of Copper Metabolism

The diagnosis of Wilson’s disease was confirmed by determining the copper content of the frozen native liver. As measured in the Denver laboratory, the excised organ had 216 µg. copper per gramme wet weight. In the New York laboratory, the copper per gramme dry weight was 1313 µg. Except for Wilson’s disease, hepatic copper values approaching these levels have been reported only in primary biliary cirrhosis or biliary atresia. As controls in the Colorado laboratory, the copper concentration was determined in 11 livers that were affected by biliary atresia (2 examples), hepatoma (2 examples), chronic active hepatitis (5 examples), and cirrhosis (2 examples). The values averaged 17.4 µg. of copper per gramme wet liver (range 9–45), the highest content being in the liver of an 11-year-old girl with intrahepatic biliary atresia.

During both the previous admissions to other hospitals, several other relevant studies were obtained. In April, 1966, the 24-hour urinary copper excretion was 50 µg. on two occasions, a normal value in that laboratory. The serum-copper was 178 µg. per 100 ml. (normal stated to be 80–280). In April, 1969, during the second admission to hospital elsewhere, serum-ceruloplasmin was reported as 26 mg. per 100 ml. (normal 20–40). Preoperative ceruloplasmin values in the Denver laboratories in three different preoperative specimens after freeze storage for 6 months were 26, 27, and 31 mg. per 100 ml., using an enzymatic assay. The concentration in the first sample was found to be 28 mg. per 100 ml. by Dr. George Cartwright, of the University of Utah, using a similar analytical method. However, in the New York laboratory, ceruloplasmin concentration of this specimen was 15 mg. per 100 ml. by enzymatic assay as well as with a calibrated immunochromatographic method.

Following liver transplantation, serial 24-hour urinary copper excretion was determined repeatedly. There was a massive cupriuresis which declined to normal (less than 30 µg. per 24 hours) after several months (fig. 2). At the end of 7 months a 3-day course of D-penicillamine (1.5 g. per day) was given and there was a 20-fold increase in urine copper excretion, which then rapidly returned to normal. Copper excretion remained normal until 15 months post-transplantation, at which time it became slightly elevated (fig. 2). During the 18 months of follow-up, 15 serial serum-ceruloplasmins varied from 29.5 to 55 mg. per 100 ml. (normal 22–43) and coincidentally measured serum-coppers ranged from 80 to 171 µg. per 100 ml. (normal 70–118).

The homograft biopsies obtained at 6 months and 17 months after transplantation contained 15 and 13-6 µg. copper per gramme wet weight in the Denver laboratory (normal less than 20). The dry weight per gramme liver in the New York laboratories at 6 and 17 months was 54 and 21.3 µg.

As controls, cases were studied in which orthotopic hepatic transplantation was carried out for indications other than Wilson’s disease. 24-hour urine-copper excretion was estimated in three patients 40–665 days after liver replacement for biliary atresia; this ranged from 18 to 26 µg. per 24 hours. In addition, copper was measured in twelve livers that had been transplanted 35–400 days previously to patients with biliary atresia or hepatoma. The grafts contained an average of 10.5 µg. copper per gramme wet weight liver tissue (range 7–19).

Genetic Studies

Both parents and all three siblings of the patient had 24-hour urinary copper excretion and serum copper and ceruloplasmin estimations. In addition, these family members had ⁶⁷⁷Cu incorporation studies performed (see table). There was no clear evidence that either parent or any of the other children were heterozygous. The mother was not taking contraceptive pills at the time of testing, nor has she ever done so in the past.

Renal Function

Before operation the blood-urea-nitrogen was 17 mg. per 100 ml. and the serum-creatinine 1.0 mg. per 100 ml. Unfortunately, no other evaluation of renal function could be obtained. Afterwards, serial phosphorus and creatinine clearances were performed. Amino acids were measured by ion-exchange chromatography on a Beckman-Spinco amino acid analyser and phenylalanine, tyrosine, alanine, glycine, glutamine, and serine clearances were determined. The measures were within normal limits at all times post-transplantation, thereby ruling out the possibility that the renal injury sometimes associated with Wilson’s disease had developed.

Discussion

The characteristic findings in Wilson’s disease can include a low serum-ceruloplasmin, an increased urinary excretion of copper, an elevation of non-ceruloplasmin copper in the serum, Kayser-Fleischer rings, and an elevated copper content of the liver and other organs. Any or all of these diagnostic criteria, except the last one, may be absent; and this may lead to the tragedy of not providing treatment with the chelating agent, penicillamine. Moreover, since ceruloplasmin values are depressed in only about 10% of heterozygous carriers of the abnormal gene, and the ⁶⁷⁷Cu incorporation test is of only fair reliability, the findings in the siblings and parents of our
patient cannot be regarded as proof that some of them do not possess a single "Wilson's disease gene".

In our patient, Wilson's disease was suspected intermittently in the 3 years preceding transplantation, but the results of the biochemical tests of serum and/or urine were either normal or equivocal. For example, all the preoperative ceruloplasmin determinations were within normal limits except one result from the New York laboratories. These generally negative findings plus the absence of the Kayser-Fleischer rings delayed recognition of the disease until there was no hope of conservative medicinal treatment. Nevertheless, the firm diagnosis of Wilson's disease was ultimately established from the high copper concentration in the native liver as well as from the presence of Mallory bodies in this specimen. The first of these abnormalities can be found in primary biliary cirrhosis or atresia, but the second may be present in alcoholic cirrhosis or Indian childhood cirrhosis, but obviously these diagnoses did not pertain to this 11-year-old white American boy.

Despite the absence of several of the common criteria for the diagnosis of Wilson's disease, evidence of extensive copper deposition in extrapathic tissues was provided indirectly by the massive cupriuresis which followed operation and which continued for several months before the urinary copper level returned to normal. Even after 7 months, the magnitude of urinary copper excretion in response to a 3-day penicillamine course was somewhat greater than is usually observed in normal people. At all times in the past year, urine excretion of copper has been slightly elevated, either at high normal or above.

It has been suggested by Walshe and Sternlieb and Scheinberg and others that in Wilson's disease the primary defect is hepatic. This hypothesis is based upon the fact that in all clinical permutations of the disease the highest concentrations of copper are in the liver. The extrapathic organ systems are presumably not affected until the buffering capacity of the liver for copper is exceeded. Thereafter toxic levels presumably develop in other parts of the body.

If the essential defect in Wilson's disease were hepatic, provision of a new liver would be expected to be curative, since liver homografts and the proteins they synthesize retain their donor specificity in the new host for a long time and probably permanently. Consequently, the studies in our case are consistent with the hypothesis that Wilson's disease is a liver-based inborn error of metabolism, although they do not prove it. In the 18 months since the liver was transplanted the total body-copper has declined steadily, while the copper concentration in the new liver has remained normal. In so far as can be determined within the limits of the follow-up, this patient now has no clinical or biochemical evidence of Wilson's disease.

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