Liver disease in infancy: a 20 year perspective

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In Northern Europe and North America the majority of children with chronic or life threatening liver disease first develop features of liver disease in early infancy. They usually have a hepatitis syndrome characterised by conjugated hyperbilirubinaemia, abnormal biochemical tests of liver function, hepatomegaly with or without splenomegaly and partial or complete cholestasis. There may be features caused by malabsorption, particularly of fat soluble vitamin K. It is from this cohort that the majority of children requiring transplantation are drawn. Such liver disease may occur in as many as 1 in 250 newborn infants.¹²

During the development of the paediatric liver service at King's, in fruitful cooperation with the adult Liver Unit, we have identified important differences in some aspects of the clinical and pathobiological features of chronic liver disease which occur in children and adults, such as autoimmune chronic active hepatitis and primary sclerosing cholangitis.³⁴ Our major efforts, however, have been to identify and improve the treatment of those with progressive liver disease starting in infancy. These account for over 50% of patients referred, the numbers in the main categories being given in the Table. This review provides an opportunity to consider some developments in which we have participated in the last 20 years.

We highlight four advances which have had a major impact on the management of these disorders. Most important has been the observation that a large proportion of patients with biliary atresia could have prolonged survival with a good quality of life after successful porto-enterostomy. The second has been identifying the role of alpha-1-antitrypsin deficiency (PIZZ) as a genetic factor associated with a particularly severe form of the syndrome frequently leading to cirrhosis. The third has been the developments in molecular biology which have led to the identification of rare genetic or congenital disorders presenting as neonatal hepatitis, gradually reducing the percentage of benign cryptogenic cases. Three will be considered: Niemann-Pick type 2, Zellweger's syndrome and Alagille's syndrome. Lastly, there has been the increased awareness of the importance of liver disease in children and the provision of improved treatment and research facilities which has come from the activities of the Children's Liver Disease Foundation.

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Biliary atresia

In the last 30 years it has been confirmed that the underlying pathological process in this disorder is usually a destructive sclerosing inflammatory process. It initially causes atresia of all or part of the extrahepatic biliary system which in uncorrected cases extends into the major intrahepatic ducts. A biliary cirrhosis rapidly develops with a mean age of death of 11 months with less than 5% surviving beyond two years.⁵ Up to 25% of patients have congenital abnormalities in cardiovascular, gastrointestinal and genitourinary systems with up to 7% having a distinct constellation of abnormalities forming the polysplenia syndrome which may include *situs inversus abdominis*, preduodenal portal vein and intestinal malrotation as well as multiple or absent spleen.⁶ The aetiology of biliary atresia remains unknown. Those with other abnormalities may be a separate aetiological subgroup.

The vast majority of these infants are entirely well during the first four to eight weeks of life apart from their jaundice. Their wellbeing often causes paediatricians and other health workers to dismiss consideration of this disorder until successful surgery is less likely, the process having destroyed the major intrahepatic bile ducts. Kasai and his coworkers in the late 1950's pioneered the operation of portoenterostomy.8 In this procedure an anastomosis is fashioned between the area of the porta hepatis from which the inflamed bile duct remnants have been resected and a 30-40 cm Roux loop. This allows bile to drain from patent major intrahepatic bile ducts directly into the bowel. Before Kasai's work less than 15% of patients with biliary atresia were operated on at the stage in the development of the process in which the surgeon could identify a bile-containing duct at the porta hepatis and the surgeon was able to fashion a direct bile duct to bowel anastomosis. With this procedure a few patients cleared their jaundice and achieved longterm survival.

The portoenterostomy procedure was introduced only slowly into the surgical practice of Europe and North America.⁵ The first 32 patients seen between 1970 and 1973 in the newly established children's liver service at King's College Hospital all died of liver failure.⁵ Surgical experience with an experimental model of this condition in the pig, however, led to improvements in technique and the first longterm survivor (Fig 1) was treated in 1973.⁵ Since then it has been possible to obtain good bile flow with normal serum bilirubin values within one to six months in between 80% and 90% of infants if they are operated on by 60 days of age. In infants operated on later the percentage becoming jaundice free is between 20% and 35%. Results are less satisfactory in centres with less experience.¹⁰

Biochemical tests of liver function remain abnormal for many years after surgery. Hepatic fibrosis is usually well established at the time of surgery and is aggravated if ascending cholangitis occurs. Nevertheless in infants becoming jaundice free the 15 years survival with a good quality of life is now almost 90%.¹¹ In infants in whom the bilirubin is not reduced the rate of progression to cirrhosis is not slowed. If bile drainage is

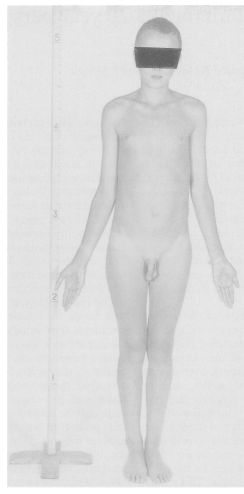


Figure 1: Fifteen year old boy with extra hepatic biliary atresia corrected at the age of 10 weeks.

partially effective death may be delayed up to the second decade. $^{\rm u}$

Cholangitis is a major and unresolved problem after portoenterostomy. It occurs in over 50% of cases in the first two years after surgery. It is characterised by fever, recurrence or aggravation of jaundice and frequently features of septicaemia. Blood culture, ascitic aspirate or liver biopsy may identify the organism responsible. Broad spectrum antibiotic therapy is given pending *in vitro* sensitivity results. Neither prophylactic antibiotics nor complicated surgical techniques of cutaneous diversion of bile influence the frequency or severity of cholangitis.¹²

With successful surgery portal pressure falls, but approximately 50% of those aged five years have oesophageal varices. Ten per cent to 15% have alimentary bleeding. We have shown that injection sclerotherapy is a very satisfactory method of management of this complication and devascularisation or shunt procedures are rarely needed.¹³

All require dietary supplements because of malabsorption in the first year of life. Thereafter normal growth can be expected in these patients even though there may be severe intrahepatic fibrosis or even cirrhosis.

In approximately 10% of cases in whom the serum bilirubin returns to normal cirrhosis progresses either because of the ascending cholangitis or possibly the continuation of the original cholangiopathy. For these patients and for those in whom surgery has not been effective in returning the serum bilirubin to normal liver transplantation is now possible. The morbidity and mortality of portoenterostomy (and the cost) is minor to that of transplantation, making portoenterostomy the initial procedure of choice in all except infants who have decompensated cirrhosis when they first present.^{14 15} Transplantation should be considered for the infant in whom portoenterostomy fails completely or for the longterm survivor with deteriorating liver function or life threatening portal hypertension. Sadly, biliary atresia remains the most frequent reason for transplantation in children in the United Kingdom because of delay in initial recognition of the possibility of serious hepatobiliary disease in infants remaining jaundiced after two weeks of age.7

Alpha-1-antitrypsin deficiency and liver disease

Alpha-1-antitrypsin was isolated by Schultz and coworkers in 1955.16 An association with liver disease and alpha-1-antitrypsin deficiency was first identified in two brothers with cirrhosis.17 Since then genetic deficiency of alpha-1antitrypsin has had a major impact in the clinical practice of paediatric hepatology.¹⁸⁻²⁰ The deficiency state PIZZ (PI=protease inhibitor, phenotype ZZ) which is inherited in an autosomal fashion is the second most common single diagnosis after biliary atresia in infants with the hepatitis syndrome in populations of European descent. Alpha-1-antitrypsin phenotyping, preferably by isoelectric focusing, is required for diagnosis. Serum concentrations, measured by immunological techniques, may be increased or decreased by associated diseases or drugs and therefore unreliable in making a diagnosis.¹⁸

The deficient infant has an increased susceptibility to liver disease with 15% developing clinical features of a hepatitis.^{21 22} What initiates the liver damage is controversial.^{20 23} Alpha-1antitrypsin is thought to inhibit tissue-damaging proteases. It inhibits a wide range of serine proteases, particularly neutrophil elastase. This enzyme functions as an extracellular protease. Its prime substrate is elastin but it also attacks many other proteins including a variety of proteins in the coagulation and complement cascades, E coli cell wall components and all major components of extracellular matrix.^{18 24} In the deficient individual uninhibited action of proteases may cause progressive liver disease. The HLA status may contribute to the severity of liver injury.25

Liver disease is most commonly identified in early infancy as a conjugated hyperbilirubinaemia with hepatitis in 11% or a bleeding state caused by vitamin K malabsorption in 2%. Up to 70% have abnormal biochemical tests of liver function in early infancy. The mean age at recognition of icteric hepatitis is between two and three weeks. Jaundice lasts on average for three months but may persist for as long as a year. The infants commonly have slow weight gain, some may show irritability or lethargy. They are at risk of septicaemia which can cause a devastating deterioration in liver function with marked pro-

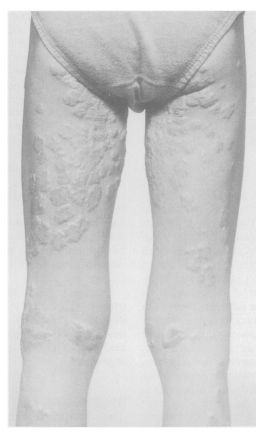


Figure 2: Xanthelasma in a 10 year old boy with Alagille's syndrome.

longation of the prothrombin time. All have hepatomegaly and approximately 50% have splenomegaly. Those presenting with a severe bleeding episode do so at two to six weeks of age. The prothrombin time (prothrombin ratio) is greatly prolonged. It reverts to normal within six hours with intravenous vitamin K. Rarely the presentation is with ascites in the newborn period. The course of the liver disease is independent of the mode of presentation in infancy.

About 5% remain jaundiced, progress to decompensated cirrhosis and die in the first year of life. The remainder recover from the acute hepatitis and in approximately 25% the clinical and biochemical abnormalities gradually improve and results come within the normal range at ages ranging from three to 10 years. Survival into the third decade without features of cirrhosis has been recorded in such patients.

Approximately 25% have died from complications of cirrhosis at ages ranging from six months to 17 years. Death from liver disease occurs within two months to four years of the onset of complications. Haematuria and/or albuminuria as a result of glomerular lesions is a late complication which may predispose to severe systemic hypertension after transplantation.²⁸

 TABLE
 Infants with conjugated hyperbilirubinaemia,

 referred to King's College Hospital between 1970–90

Biliary atresia	377
Idiopathic hepatitis of infancy	331
Alpha-1-Antitrypsin deficiency	189
Other hepatitis of infancy	94
Alagille's syndrome	61
Choledochal cvst	34

Approximately 25% survive through the first decade although they have histologically confirmed cirrhosis. A further 25% without liver biopsy evidence of cirrhosis have persistently abnormal liver function tests with or without clinical features of portal hypertension. In some of these without clinical abnormality, liver function test may eventually become normal but the outcome for those with clinical or biochemical evidence of liver disease is guarded.^{18 19 26-28}

The management is that of chronic cholestasis and of cirrhosis.² There is no specific treatment for liver disease associated with alpha-1antitrypsin deficiency short of liver transplantation. It corrects the serum phenotype to that of the donor.²⁹ The longest follow up is only 16 years making it too early to determine whether it will prevent emphysema.³⁰ In up to 80% of children within a sibship the liver disease is of the same severity as the proband.26 Antenatal diagnosis of the PIZZ state initially possible only with fetal blood sampling³¹ is now possible by examining the DNA of chorionic villus samples using synthetic oligonucleotide probes specific for the M and Z gene or by restriction fragment length polymorphism. With the polymerase chain reaction results can be made available within a few days of sampling at 11 weeks gestation. Preliminary studies confirm the validity of such techniques.32

In adults with emphysema it has been possible to bring plasma alpha-1-antitrypsin levels up to normal values with infusions of plasma derived alpha-1-antitrypsin. This has not been shown to modify the emphysema. Nor did it decrease to normal levels the raised serum transaminases values found in a few patients who were treated in this fashion.³³

Direct gene targeting to the liver in vitro is another possibility having been used successfully stimulate albumin production to in analbuminaemic rats.³⁴ The gene was contained in a plasmid which is targeted to the hepatocytespecific asialoglycoprotein receptor and carried into the hepatocyte by pinocytosis. In this way the PIMZ state would be created. Another theoretical possibility is the correction of the hepatic secretory problem by insertion of another gene which makes a complementary change in the polypeptide allowing it to assume normal tertiary configuration which may facilitate secretion.^{33 35} Perhaps more immediately practicable and attractive, to paediatric hepatologist if not to pulmonary physicians, would be a trial of exogenous serum-derived alpha-1-antitrypsin, given iv at one to four weekly intervals, as is being used in emphysema but commencing as soon as significant liver damage is identified, perhaps in conjunction with a neonatal screening programme. It should be possible to conduct a trial of therapy lasting six to 12 months which in the course of at most a decade, and perhaps much shorter, would indicate whether such therapy was efficacious.

Niemann-Pick type 2

The Niemann-Pick group of diseases are sphingomyelin-cholesterol lipidosis.³⁶ Failure of lipid degradation causes abnormal lipid storage with hepatosplenomegaly, the formation of foam cells in the bone marrow and in some forms progressive central nervous system involvement. In type 1 disease sphingomyelinase activity is <10% of normal. In type 2 disease sphingomyelinase activity is normal except in cultured fibroblasts in which activity may be reduced by 80%. A defect in cholesterol esterification has recently been demonstrated *in vitro*.³⁷ Within both types there is considerable clinical and biochemical variability.

Niemann-Pick type 2 is the second most common genetic cause of liver disease in infancy in the United Kingdom, after alpha-1antitrypsin deficiency. It has previously been designated type C, D, III, IV and juvenile Niemann-Pick with sea blue histiocytes, DAF syndrome and neurovisceral storage disease with supranuclear ophthalmoplegia. Over 50% of cases present with hepatitis in infancy or intrauterine ascites.^{37 38} A third of these die by six months of age while in the survivors features of liver disease regress in infancy leaving an asymptomatic cirrhosis. Unfortunately they, like those in whom liver involvement is asymptomatic, develop features of progressive, ultimately fatal, neurological involvement starting between two years of age and the sixth decade. Diagnosis is suspected by finding storage material in bone marrow and within neurones in a rectal biopsy and is confirmed by observing defective cholesterol esterification in cultured fibroblasts. There is no treatment. Liver transplantation does not arrest the disease. Diagnosis is essential for genetic counselling. Prenatal diagnosis by amniocentesis is possible.

Zellweger's syndrome

Zellweger's syndrome, an autosomal recessive disorder, is characterised by absence of peroxisomes and defective function of peroxisomal enzymes.^{39 40} It is a multisystem disorder with profound abnormalities of neuronal migration and brain structure, hepatic malfunction, renal cortical cysts, abnormal calcification, retinal degeneration and multiple congenital anomalies. The defect is associated with a sometimes diagnostic facial appearance, severe mental retardation, multiple congenital anomalies, including renal cortical cysts, retinal degeneration and deafness.^{37 38} It is the most severe of the disorders of peroxisome biosynthesis. Hepatomegaly and raised transaminases with jaundice in approxmately 60% of cases are the initial hepatic features. In the first weeks of life there is mild periportal fibrosis with an irregular relationship between portal tracts and central veins but by eight weeks of age fibrosis is more marked and frequently a micronodular cirrhosis is established by 20 weeks of age. Eighty per cent of patients die by six months of age, with few surviving beyond two years, but less severe incomplete forms of the condition, with more prolonged survival are increasingly described. There is no treatment for this condition but prenatal diagnosis is possible. The main scientific interest in the condition has been to identify the key metabolic role of peroxisomes. They account for an estimated 20% of the oxygen consumption

of the liver being engaged in the oxidation of a large number of endogenous and exogenous substrates. The latter include ethanol, methanol, and nitrites. Peroxisomes also have an important role in the catabolism, by beta oxidation, of fatty acids of chain length greater than 12 (producing much energy), long chain dicarboxylic acids, prostaglandins, xenobiotics and the side chain of cholesterol. They have an essential role in the synthesis of the plasmalogens (phospholipids) and bile acids. In this syndrome beta oxidative cleavage of the side chain does not occur. Therefore 5 β -cholestane-3 α , 7 α , 12 α -triol is oxidised to 3a, 7a, 12a-trihydroxy-5ß-cholestanoic acid, 5α cholestane- 3α , 7α -diol to 3α , 7αdihydroxy-5ß-cholestanoic acid. These are not metabolised to cholic acid or desoxycholic acid but accumulate and undergo either side chain elongation to form C₂₉ bile acids or nuclear changes.

Diagnostic biochemical abnormalities include greatly diminished levels of plasmalogens, defects in bile acid synthesis, raised serum concentrations of long chain fatty acids, phytanic acid and pipecolic acid (a product of lysine degradation).

Alagille's syndrome

This disorder is characterised by paucity of interlobular bile ducts (intrahepatic bile duct hypoplasia) occurring in association with a range of characteristic cardiovascular, skeletal and ocular anomalies (arteriohepatic dysplasia, syndromic paucity of the interlobular bile ducts).^{41,42} The syndrome is probably inherited in an autosomal dominant fashion with variable expression. The family history is positive in 15% of cases, although full expression of the syndrome may not be present in earlier generations.

Essential to the diagnosis is a decrease in the number of interlobular bile ducts seen in portal tracts (ratio <0.06). Such paucity may be found also in association with genetic disorders such as alpha-1-antitrypsin deficiency or failure of primary bile salt synthesis, in chromosomal abnormalities and rarely with intrauterine infections. It is frequently present in biliary atresia after nine months of age. Diagnosis, based on histological finding, is supported by the finding of the typical facies, deep set eyes, mild hypertelorism, overhanging forehead, a straight nose which in profile is in the same plane as the forehead and a small pointed chin, posterior embryotoxon, vertebral body defects on spinal radiographs and the presence of a cardiac murmur. Peripheral pulmonary stenosis is the most common abnormality but a wide range of other forms of cyanotic and acyanotic heart disease may occur. As none of these abnormalities are unique to this syndrome if the biopsy findings are not typical (see below) and cholestasis is complete ERCP may be required to distinguish this disorder from biliary atresia.43

There is long standing cholestasis varying in severity from mild pruritus to life long jaundice. Severe cases start with jaundice in the neonatal period, developing pruritus at five months by which time there is hypercholesterolaemia (frequently >15 mmol/l) with xanthelasma appear-

ing six to 12 months later. Xanthelasma usually clear some years before the jaundice. Less severe cases have no xanthelasma. Jaundice may persist, but in others clears in late childhood or early adult life. Renal impairment and renal tubular acidosis may occur in association with 'mesangiolipidosis' or juvenile nephronophthisis. The longterm prognosis is uncertain but some 15% may go on to develop cirrhosis and 5%–10% die from liver disease. In one series 25% died from cardiac involvement, classically a peripheral pulmonary stenosis, or infection.41 The pathogenesis is intriguing in that during the first three to nine months of age it may be impossible to make the histological diagnosis in up to 25% of cases because bile ductular proliferation with features suggestive of bile duct obstruction may be the main finding. If the jaundice clears all features of liver disease may eventually remit and where liver biopsies have been performed the bile duct to portal tract ratio has been normal.

The treatment is that of chronic cholestasis with particular emphasis on adequacy of fat soluble vitamin replacement, particularly vitamin E. After vitamin E deficiency for periods ranging from six months to four years these patients develop a neuromuscular degenerative syndrome. This can be prevented if the serum vitamin E concentration is maintained within the normal range. Growth failure and pruritus are major problems. It is not clear whether growth failure can be reversed by liver transplantation.

The Children's Liver Disease Foundation

In the last decade paediatric hepatology in the United Kingdom has received a considerable boost from the activities of the Children's Liver Disease Foundation. This organisation has as its aims the improved management of liver disease in children by provision of better facilities, promotion of research and teaching and by giving emotional support for the families of children affected, Parents and friends of children with liver disease have been largely responsible for collecting the magnificent sum of £1.2 million pounds which has been spent in furthering these aims. To them and to the trustees we owe a large debt of gratitude. At last there is increasing professional and public awareness of the problem of liver disease in children. We hope that the British Liver Foundation can achieve as much for adult hepatology.

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