The liver in siblings of patients with Indian childhood cirrhosis: a light and electron microscopic study

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SUMMARY Liver biopsies from 29 siblings of patients with Indian childhood cirrhosis (ICC) and from two age-matched controls were examined by routine light and transmission electron microscopy. Histochemical stainings for copper and copper-binding protein were also carried out. The mild and non-specific structural alterations that were observed did not differ from those seen in control livers, even though a slight to moderate excess of copper and copper-binding protein was demonstrated in the majority. Aggregates of microtubules seen in some siblings, as well as in control livers, may indicate the preconditions for development of Mallory hyaline. It is possible that these features suggest a susceptibility for the development of ICC but not early disease.

Among several interesting features in the clinical and epidemiological spectrum of Indian childhood cirrhosis (ICC), there is one enigmatic one: the peculiar familial distribution of the disease. Even though no clear-cut pattern of inheritance is apparent, affliction of more than one child, twins, and other collateral members in the family has been described.1-4 It has been suggested that an inherited susceptibility in children of the affected families may develop into frank disease under the precipitating influence of some unknown agent or agents.7,8 In the established stage of ICC, both clinical and pathological features appear to be characteristic.5 Hallmarks of earlier phases, however, remain to be clearly identified, though a precirrhotic symptom complex believed to herald the disease has been described.5 In a follow-up study of asymptomatic siblings, progression of early, non-specific hepatic alterations to classical changes of ICC was observed.7 Recently, considerable interest has centred around the finding of large excesses of copper and copper binding protein in the livers of patients with ICC.5-11 The precise pathological effects of these materials in the development of hepatic lesions in this disease remain to be clarified. When we investigated a group of ICC patients and siblings of such patients to ascertain the part played by excessive hepatic copper and copper-binding protein in the evolution of the disease (Marwaha et al., submitted for publication), we took particular note of the structural alterations in the sibling liver, which are described in the present communication. It was considered important from the point of view of the affected families to determine if there was any specific abnormality in the sibling livers and, because of ethical considerations, permission was obtained before performing the biopsies.

Methods

Percutaneous needle biopsy of the liver was obtained from 29 siblings of patients with Indian childhood cirrhosis (ICC); in 15 of the latter the diagnosis had been histologically confirmed. The age distribution was as follows: ICC patients—age range 10 months to 3 years (mean age 1.77 years), siblings of ICC patients—age range 2 months to 6 years (mean age 1.6 years). The major part of the biopsy specimen was fixed in formol saline and processed routinely for paraffin embedding followed by serial sectioning. Deparaffinised sections were stained with haematoxylin and eosin, modified Shikata's orcein stain12 for copper binding protein, p-dimethyl aminobenzylidene rhodanine stain for copper, and the periodic acid schiff (PAS) stain. A small portion of the liver biopsy specimen from 19 siblings was fixed in 3% glutaraldehyde, post-fixed in 1% osmium tetroxide, and processed in the conventional manner for transmission electron microscopy. Sections were examined with a Philips 300 electron microscope for specific organelle damage and for the presence of any abnormal structures within the hepatocyte. Wedge biopsy of the liver from two children who were operated on for congenital megacolon, but who had

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Fig. 1  Sibling liver showing moderate amounts of orcein positive deposits (arrow heads) in the periportal hepatocytes. P: Portal tract. × 380.

Fig. 2  Sibling liver showing irregular configuration of the nuclear membrane (N) and numerous vesicles of smooth endoplasmic reticulum (ser) in the cytoplasm. × 6000.

Fig. 3  Abundant glycogen rosettes of alpha particles, dilated sacs of smooth endoplasmic reticulum, and a few profiles of rough endoplasmic reticulum close to the nucleus. Lipofuscin pigment granules and two autophagic vacuoles are also seen. × 16400.
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no clinical or biochemical evidence of liver disorder, served as control material. These were also processed for light and electron microscopy.

Results

Light microscopic structure

Histological examination of the sibling livers revealed non-specific morphological features in the form of cytoplasmic and/or nuclear vacuolation in hepatocytes, and focal fatty change and portal fibrosis with or without the presence of a few mononuclear inflammatory cells in 16 out of 29 cases; the propositi of eight of these cases had been histologically proven. There appeared to be no difference in these minor abnormalities between siblings in whom the propositi had proved to have Indian childhood cirrhosis or those who had not. In the remaining cases the liver appeared to be normal by light microscopy. Copper was seen as orange red rhodanine positive intracytoplasmic granules, 0.5–2 μ in diameter, localised to the periportal hepatocytes in mild to moderate amounts in 18 out of 29 livers. Copper-binding protein, appearing as orcein-

Fig. 4 Three adjacent hepatocytes (H) show excess of lipofuscin pigment and numerous autophagic vacuoles. × 6600.

Fig. 5 An aggregate of microtubules (mt arrows) in proximity to polysomes and profiles of rough endoplasmic reticulum. × 20000.
positive, caramel coloured, coarse granules (Fig. 1),
was present in 23 livers, its distribution and amount
.correlating well with those of copper. Non-specific
histological alterations were seen in 12 out of 18
(66.6%) of copper positive cases and in four out of
11 (36.4%) of copper negative cases, the difference
not being statistically significant. Also, no consistent
histological abnormality could be observed in
association with excess of tissue copper. The livers
of two control cases were histologically unremark-
able, except for mild excess of lipofuscin pigment
in one. No copper or copper-binding protein was
detected.

ULTRASTRUCTURE
The livers revealed a variable picture. In two biopsies
the nuclear membranes of hepatocytes appeared to
be irregular (Fig. 2), leading to loss of the normal
rounded contour of the nucleus, but the chromatin
and nucleolus seemed unremarkable. Moderate to
abundant glycogen deposits were observed in the
cytoplasm of hepatocytes in 14 siblings, displacing
the mitochondria to the perinuclear area or to the
cell membrane. Glycogen appeared as rosette-like
aggregations of electron dense alpha particles (Fig.
3). In two biopsies relatively electron-lucent particles
resembling the beta form of glycogen were scattered
in some areas of the cytoplasm. Smooth endoplasmic
reticulum (SER) was seen as groups of somewhat
widened vesicles, diffusely scattered or interspersed
among glycogen rosettes (Figs. 2 and 3). Rough
endoplasmic reticulum (RER) was mainly located in
the perimitochondrial and perinuclear regions and
occasionally appeared to be dilated. In some hepatocytes focal aggregates of polysomes were seen in
abundance. Mitochondria varied markedly in size
and somewhat in shape, but their finer structure
seemed to be undisturbed. Lipofuscin granules were
conspicuous in seven sibling livers and, in two of
these, prominent autophagic vacuoles were seen (Fig.
4). No zonal distribution of lipofuscin could be
discerned. In seven biopsies occasional aggregates of
microtubules were observed (Figs 5 and 6). These
were random collections of haphazardly oriented
microtubules ranging in diameter from 20-41 nm.
Some were in continuity with, while others were
in proximity to, the RER profiles. An inter-
mingled granular material was present in some of
the microtubule aggregates.

Livers of eight siblings showed numerous irregu-
lar clear spaces devoid of any organelles or
glycogen. Myelin figures and lipid vacuoles were
prominent in a few biopsies and occasionally a
widened intercellular junction or a dilated bile
canalculus was noted.

Hepatic ultrastructure in control livers was similar

Fig. 6 Details of microtubules (arrow) in
an aggregate in the cytoplasm of a
hepatocyte. × 46 000.
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to that described in sibling livers including the
presence of beta particles of glycogen in one liver.
In one case moderately large collections of micro-
tubules, morphologically identical to those seen in
some of the sibling livers, were observed (Fig. 7).

Discussion

Hepatic morphology in siblings of ICC patients
reveals no significant or consistent abnormality and
the non-specific alterations seen in some do not
correlate with the presence of mild to moderate
excess of copper and copper-binding protein. Ultra-
structural features of these livers do not differ from
those of the two controls who had no apparent liver
disease. In fact, although, for obvious reasons,
information on hepatic ultrastructure in the healthy
human is lacking, observations made in our subjects
fall well within the spectrum of organelle morpho-
logy described by Ma and Biempica\(^12\) in biopsies
from apparently normal livers obtained from patients
with nonhepatic diseases who were subjected to
different surgical procedures. We, therefore, believe
that, in the asymptomatic siblings of those with
Indian childhood cirrhosis, hepatic morphology at
best presents mild and non-specific alterations within
the range of the so-called 'normal'. Moreover, a
follow-up of 200 siblings over a period of six months
to three years revealed not only that none progressed
to ICC but that the hepatomegaly present in some
regressed (Marwaha \etal., submitted for publication).

An interesting finding in seven out of 19 sibling
livers and in one of the two control livers was
random aggregates of microtubules in the cytoplasm
of liver cells—a feature not mentioned by Ma and
Biempica\(^12\) in their report on the ultrastructure of
the normal human liver cell. In the human liver
occasional microtubules have been observed in the
aggregates of microfilaments representing Mallory
hyaline in various disorders including ICC.\(^14\)\(^15\)
Random aggregates of microtubules have also been
described in hepatocytes of alcoholics even in the
absence of characteristic Mallory bodies.\(^15\) Micro-
tubules are formed by circumferential alignment of
protofilaments constituted by tubulin—a protein
synthesised by ribosomes.\(^16\) Tubulin monomers
undergo polymerisation in the presence of a micro-
tubule-associated protein to form dimers and oligo-

Fig. 7  Hepatocyte from a control liver showing a
well defined aggregate of microtubules (mt) along
with scattered beta particles of glycogen.
\(\times 12\) 000.
mers which assume globular configurations and link up to form protofilaments. The exact mode of degradation of microtubules is not known, but this possibly occurs through formation of intermediate-sized filaments which are 6–11 nm in diameter. Studies on experimental animals indicate that these intermediate filaments are likely precursors of Mallory hyaline, which is composed of aggregates of 20–27 nm diameter filaments. Griseofulvin administration to mice for a period of four to six weeks induces the formation of classical Mallory hyaline showing all three ultrastructural variants described in alcoholic liver disease, and ICC. In vitro cell culture systems low doses of griseofulvin applied for a few hours give rise to random ‘broken’ microtubule aggregates, while with higher doses given for a somewhat longer period of time collections of intermediate filaments are observed. Thus microtubule disintegration may be an important step for Mallory hyaline induction, though the precise steps involved in its evolution are not clear. Priming and precipitating agents have been suggested, the nature of the former being unknown, while the latter are possibly anti-microtubule agents.

Development of Mallory hyaline is an important structural landmark in ICC. The finding of aggregates of microtubules in some of the sibling livers, therefore, assumes significance, even though no intermediate filaments or Mallory hyaline filaments were seen. Our studies on hepatic copper and copper-binding protein in children with ICC and their healthy siblings show that, in almost all of these subjects, a transient maturation defect of hepatocytes manifests itself in excess accumulation of these materials (Marwaha et al., submitted for publication). Copper in mild to moderate excess does not appear to be significantly injurious, but, once liver damage is initiated by other—as yet unknown—causes, copper may precipitate Mallory hyaline in an already ‘primed’ cell. Recent studies have shown that copper, by binding to sulphydryl groups, can inhibit polymerisation of tubulin. The finding of collections of microtubules in some of the sibling as well as control livers may indicate some type of a ‘priming’ phenomenon. These changes probably resolve spontaneously unless a super-added injury results in progressive damage and Mallory hyaline formation.

References