Electron microscopy of hepatitis B virus components in chronic active liver disease

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SUMMARY  Eighteen liver biopsy specimens from patients with hepatitis B surface antigen (HBsAg) positive chronic aggressive hepatitis were studied by electron microscopy. All cases were selected on the basis of positive liver cell membrane fluorescence for HBsAg on immunohistochemical investigation. Striking changes in the morphology of the liver cell membrane were observed in nearly all cases. Furthermore, a dual aspect of hepatitis B core antigen (HBCAg) is described. HBCAg particles may occur as either 'naked' or 'cloudy' particles surrounded by semi electron dense material. The nature of the 'cloud' remains to be identified.

Hepatitis B viral components represent characteristic structures which can be revealed by electron microscopy. The hepatitis B core component has been described as 21-25 nm sized particles, located predominantly in the nuclei of the hepatocytes and to a lesser extent in their cytoplasm. The hepatitis B surface component was found to correspond to long filamentous or tubular structures located in the cisternae of the smooth endoplasmic reticulum (Huang, 1971; Stein et al., 1972; Huang et al., 1974).

Hepatitis B core antigen (HBCAg) as well as hepatitis B surface antigen (HBsAg) can also be demonstrated by immunohistochemical techniques (immunofluorescence, immunoperoxidase) on frozen and paraffin-embedded liver specimens (Gerber et al., 1974; Huang, 1975; Shikata, 1973; Yamada and Nakane, 1977).

Previous studies in this laboratory (Ray et al., 1976a, 1976b) and elsewhere (Gudat et al., 1975) have shown a differential distribution of HBsAg and HBCAg in various forms of hepatitis B. With these techniques, chronic aggressive (or active) hepatitis and active cirrhosis are characterised by an almost equal proportion of nuclear localisation of HBCAg together with cytoplasmic positivity for HBsAg. Furthermore, expression of HBsAg in the liver cell membrane is a striking feature in these conditions (Ray et al., 1976b).

The present study was undertaken in order to obtain more detail on the ultrastructural counterpart of the immunohistochemical findings in chronic aggressive hepatitis with or without cirrhosis.

Material and methods

Eighteen needle biopsies of liver were performed on 18 patients clinically and histologically diagnosed as HBsAg positive chronic aggressive hepatitis. The sera of all these patients were positive for anti-HBCAg, and three of the nine determined samples were also positive for anti-HBsAg. The titre of DNA-polymerase was determined in only two serum samples and was rather high. The antigen was determined by the Ouchterlony technique in 11 cases; seven samples were negative and four were positive.

The cases were classified on the basis of previously described hepatic histology (De Groote et al., 1968; Bianchi et al., 1971) and immunofluorescence findings (Gudat et al., 1975; Ray et al., 1976b) as chronic aggressive hepatitis (14 cases) and active cirrhosis (4 cases).

The localisation of HBsAg and HBCAg, as studied by immunofluorescence, showed a characteristic pattern: all biopsy specimens had a large number of positive nuclei for HBCAg while the HBsAg expression was mostly prominent in the liver cell membranes: 13 specimens showed a moderate to strong membrane staining in a variable percentage (10-90%) of their hepatocytes; seven of these 13 biopsies showed a moderate to strong membrane staining in 90% of the hepatocytes; only five specimens revealed a rather faint membrane staining.
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Fig. 1  *Hepatocytic nucleus with naked HBcAg particles in the nucleus and in the nuclear pore (× 70,000).*

All the cytoplasm stained positively in a small number of hepatocytes in 15 specimens. In all these biopsy specimens *in vitro* complement fixation was positive as studied with immunofluorescent techniques. In 12 cases, such *in vitro* complement fixation was found in the hepatocyte
nuclei and in the cytoplasm, whereas in six cases it was restricted to the nuclei. On the other hand, 15 specimens showed nuclear fluorescence for immunoglobulin G (IgG) (Ray, 1978).

One part of the biopsy tissue was prepared for light microscopic diagnosis, a second part was prepared for immunofluorescent study, and a third part was prepared for electron microscopy. Thin
blocks (1 x 1 x 1 mm) were immediately fixed by immersion for 1 hour in cold (4°C) 2.5% glutaraldehyde phosphate buffer, 0.1 M, pH 7.2, followed by a phosphate buffer rinse overnight. The specimens were postfixed in 1% OsO4 in phosphate buffer, 0.1 M, pH 7.2, for 1 hour at 4°C, dehydrated in graded alcohols, and embedded in Epon. Ultrathin sections were cut and stained with uranyl acetate followed by lead citrate and studied in a Zeiss EM10 electron microscope.

Results

In all 18 biopsy specimens a varying number of hepatocytes contained non-coated virus-like particles, 21 to 25 nm in size, localised in the nuclei; their number varied from nucleus to nucleus. As demonstrated in Fig. 1, such uncoated or naked core particles could occasionally be demonstrated in the nuclear pores.

These uncoated core particles were also found in the hyaloplasm in 16 specimens; in this location they were observed in any place, even near the cell periphery and in the pericanalicular ectoplasm.

In nine specimens core particles were found lying outside the hepatocytes; they appeared as single particles or in groups and were localised in the intercellular space or in the Disse space. These particles were always surrounded by a narrow, dark cloud of semi-dense fluffy material (Figs 2 and 3).

These clouded core particles were occasionally found in the cytoplasm of one or two hepatocytes in three specimens (Fig. 4).

In six biopsy specimens longitudinally cut tubules and cross-sectioned dots of HBsAg were found in the cisternae of the smooth endoplasmic reticulum (Fig. 5). In many positive cells the whole cytoplasm was filled with HBsAg containing SER; others showed these structures only in parts of their cytoplasm.
In seven specimens, mainly in hepatocytes containing HBsAg in their SER cisternae, the peripheral cell membrane was indistinct over a long segment or even seemed to have disappeared (Fig. 6). In other hepatocytes, the peripheral submembraneous area was thickened by the deposition of fluffy, filamentous material resembling the thin actin filaments described in numerous cell types (Allison, 1973; Marx, 1975; Phillips and Oda, 1974).

Furthermore, an amorphous material could be
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detected in some slightly widened intercellular spaces between hepatocytes with increased microvilli in 11 specimens (Fig. 7). This was mainly the case in the biopsy specimens which showed a clearly delineated membrane positivity for HBsAg on immunofluorescence study.

In seven of the 18 specimens a single core particle was observed surrounded by a clear halo and a dark ring, ± 40 nm in size, located in the cisternae of the SER corresponding to the structures described as Dane particles (Dane et al., 1970) (Fig. 8). On two single occasions a similar Dane particle was lying in a sinusoidal space.

With regard to liver cell morphology, the following striking feature was noted: cilia-like structures composed of protoplasmic extensions containing central microtubules were found on a single liver cell in three cases. These cilia-like structures were localised either at the canaliculus or at the sinusoidal pole.

![Fig. 6](image_url) Hepatocyte with indistinct peripheral cell membrane. The submembranous area is thickened by fluffy filamentous material. Two remnants of desmosomes can be seen (→→) (× 48 100).
In spite of all these changes, it has to be mentioned that in each biopsy specimen a varying number of hepatocytes could be found with a normal morphological appearance.

**Discussion**

This investigation concerns a relatively homogeneous group of patients suffering from chronic aggressive
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hepatitis, a minority of them being already in a stage of cirrhosis. Immunofluorescent investigation showed a nearly equal balance between the numbers of hepatocytes with nuclear HBeAg and cytoplasmic HBsAg. Characteristic for this selected group of cases was a strong fluorescence for HBsAg localised in the liver cell membranes. Although variation due to sampling unavoidably remains a problem, the results obtained are on the whole quite homogeneous. In general, they correspond to published reports on the ultrastructural aspects and localisation of HBeAg and HBsAg (Huang, 1971; Stein et al., 1972; Gerber et al., 1974; Huang et al., 1974; Yamada and Nakone, 1977).

Naked core particles were found mainly in the nucleus and to a smaller extent also in the hyaloplasm. They were found as single particles or in groups. Figure 1 demonstrates their passage through the nuclear pores, probably migrating from the nucleus to the cytoplasm.

A new finding is the occurrence of core particles surrounded by a narrow ‘cloud’ of semi electron dense material. Such ‘clouded particles’ are only rarely found in the hyaloplasm, occasionally near the nucleus, more often in the cell periphery. They are most often observed in the intercellular space or in the space of Disse. Actually all extracellular particles appeared to be of the ‘clouded’ type.

Their morphological appearance does not correspond to that of Dane particles. Pure morphological analysis does not allow identification of the nature of the ‘cloud’.

Huang (1975) described aggregates of ‘coated core particles’ in the nucleus of hepatocytes from poorly processed biopsy specimens and from necropsy-derived liver specimens, and interpreted the ‘coat’ as
autologous anti-HBcAg. However, in the present study, the specimens were immediately and ade-
quately fixed, as evidenced by satisfactory tissue preservation. Moreover, the 'clouded particles' in
the present study are never observed as intranuclear aggregates but are found in extracellular sites and—
to a lesser degree—in the hyaloplasm of the liver cells.

The occurrence of HBcAg—anti HBcAg immune complexes in the nuclei and the cytoplasm of
hepatocytes of chronic aggressive hepatitis patients has been proved in immunofluorescence techniques
by localisation of IgG and in vitro complement fixation (Nowoslawski et al., 1972; Sarno et al., 1975;
Gerber et al., 1976; Rizzetto et al., 1976; Gudat et al., 1977). In vitro complement fixation was present in all specimens examined in the present study and nuclear fluorescence for IgG in 15 cases
(Ray, 1978). The current interpretation of these observations is that autologous circulating anti-
HBcAg (IgG) may enter the hepatocyte, possibly due to altered permeability of the HBsAg containing
cell membrane, and form complexes with HBcAg present in the nucleus. In view of this, a possible
interpretation of the 'clouded particles' could be that they represent core particles complexed with
anti-HBcAg. However, in vitro complement fixation and IgG are mainly demonstrable in the nucleus,
precisely the site where in the present study no clouded particles have been found. Although a
difference in tissue processing between immunofluorescent and electron microscopical techniques
could be invoked to explain this apparent controversy, it seems necessary to consider further possi-
bilities.

As an hypothesis, one could speculate whether the cloud could represent e antigen. This third antigenic
component of the HBV was discovered recently (Magnius and Espmark, 1972; Magnius et al.,
1975); it is usually associated with persistent viral infection in chronic hepatitis (Nielsen et al., 1974;
Elefltheriou et al., 1975; El Sheikh et al., 1975), has been localised by immunofluorescent techniques in
the hepatocytic cytoplasm by one author (Trepo et al., 1976), and is usually demonstrated in the blood
of chronic hepatitis B patients. In this study, all patients had chronic aggressive hepatitis (a few of
them complicated by cirrhosis). However, only four of the 11 serum samples tested were positive for
e antigen, although the procedure used for determination of e antigen is known to be of low sensitivity.
The lack of correlation between the presence of 'clouded' core particles in the biopsy specimen and
of e antigen in the serum, together with the more recent demonstration of e antigen in hepatocytic
nuclei (Arnold et al., 1977), speaks against an identity between the cloud and e antigen.

The finding of naked core particles in the nucleus, in the pores of the nuclear membrane, and, to some
extent also, in the perinuclear region of the hyaloplasm suggests that the core particles are assembled
in the nucleus (some components may come from a cytoplasmic synthesis site), leave the nucleus
through the nuclear pores, and acquire their cloudy material in the cytoplasm (at least in some cells) or in
the extracellular space. The finding of clouded core particles in extracellular sites clearly demonstrates
that these structures can leave the liver cell for the bloodstream via an as yet unidentified pathway.

Although careful attention was paid to identified HBsAg in or near the liver cell membranes,
no undeniable identification of HBsAg structural components was achieved in this location. A
striking finding, however, was the impressive alteration of segments of the liver cell membrane in
hepatocytes loaded with HBsAg: in these areas, the cell membrane became indistinct or even disappeared.
These segments were too long and too frequent to be explained by oblique sectioning of the membrane.
Equally intriguing is the occurrence of broad fibrillar areas beneath the cell membrane and the presence
of amorphous and/or filamentous material in widened intercellular spaces. The submembranous
microfilamentous areas are different from the micro-
filamentous hyperplasia described in cholestasis
(Desmet, 1972; Gabbiani et al., 1975), in which case
this phenomenon is restricted more to the pericanali-
cular zone. Pure morphology alone, however,
cannot identify HBsAg in these altered liver cell
membranes. Immuno-electron microscopy is re-
quired definitely to resolve this issue.

As mentioned in the literature, an occasional
Dane particle could be found in an occasional liver
cell, located inside the cisternae of the SER or in an
extracellular site. The location of core particles in
the hyaloplasm, and the presence of HBsAg and Dane
particles in the cisternal lumina of the SER raise the
question how the Dane particle is assembled.

Cilia-like cytoplasmic extensions on the canali-
cular and sinusoidal pole of the liver cell is a rare
phenomenon which was previously described in a
case of intrahepatic cholestasis (Byler disease) by
De Vos et al. (1975). The significance of this liver
cell alteration remains unknown, although it cannot
be regarded any more as pathognomonic for the
disease entity in which it was originally reported.

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References


Sarno, E., Gerber, M. A., and Vernace, S. (1975). Complement fixing immune complexes in the hepatocytic nuclei of patients with hepatitis B antigen...


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