**Electron microscopic observations and speculations on Australia antigen**

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**Summary**

The use of the electron microscope has simplified methods of detection of Australia antigen in carriers of serum hepatitis, in patients suffering from the disease and those with persistent hepatitis.

Basic studies on the nature and properties of Australia antigen, using the electron microscope, are described.

The correlation between the Australia antigen and serum hepatitis is now so good (Le Bouvier & McCollum, 1970) that a study of the one is automatically a study of the other. This paper deals with one aspect of these studies, the morphology as seen by the electron microscope technique of negative staining. It will consider the practical use to which this morphological information can be put and will go on to speculate on the insight that such studies may give on the nature of the virus of serum hepatitis.

Although the Australia antigen was discovered in the early 1960s it was not until 1968 that it was first examined by negative staining in the electron microscope. This was done from gradient purified preparations and revealed mainly a small spherical component approximately 200 Å in diameter (Bayer, Blumberg & Werner, 1968). This appearance was quickly confirmed by other workers both in purified preparations and also in immune aggregates (Prince, 1968; Millman et al., 1969). Also in 1969, Almeida et al., showed that by simple centrifugation it was possible to visualize the particles of the Australia antigen within a short time of receiving a serum specimen. These negatively stained preparations of the antigen revealed that it exists in two main forms, the spherical component previously mentioned and a tubular one again having a diameter of 200 Å but having variable lengths up to several 1000 Å units (Fig. 1). No regular substructure can be visualized on the spheres and indeed in overall diameter they show an irregularity of size ranging from 160 Å to 250 Å. The tubular component, on the other hand, revealed a cross striation with a periodicity of 30 Å in well oriented examples (Fig. 2). The proof that these structures do indeed identify with the Australia antigen demonstrated by other techniques is that precipitin lines cut out from agar gel are built up from the described particles and in addition specific antisera can be added to preparations containing the antigen and specific complexing observed (Almeida et al., 1969).

Because of the overall simplicity of visualizing these distinctive structures of the Australia antigen by negative staining the electron microscope is now used extensively for this purpose and it appears to be useful on two separate levels. The first level is completely practical and entails the use of the microscope to diagnose the presence of the antigen. The second use is the more basic one of identifying the causative agent of serum hepatitis and, in a disease where there is as yet no biological means of recognizing the presence of the virus, is the most available means of characterizing it. This paper will now deal with these two aspects separately starting first with the practical one.

**Practical uses of the electron microscope in dealing with serum hepatitis**

As has already been stated simple centrifugation of a serum specimen followed by negative staining
of the pellet so obtained allows visualization of the antigen (Almeida et al., 1969). The time-lapse between receiving a specimen and examination in the microscope is a little over 2 hr, a time that compares favourably with other methods of examination. In addition, in our hands, electron microscope examination has been the most sensitive means of detecting the presence of the antigen. Also, in addition to sensitivity, the distinctive structures visualized and the permanent micrograph record mean that it is possible to reach a definite diagnosis with certainty and in some instances, e.g. a potential patient in a haemodialysis unit: it is very important that this can be done.

This then is probably the most frequently employed use of the electron microscope in the serum hepatitis field, simple diagnosis of the presence of Australia antigen (Au-Ag). But in addition to showing the presence of the antigen it was realized that different patterns of distribution could be seen on the microscope grid. Some specimens contain randomly distributed antigen but in other cases it appears clumped, with all the appearances of an antigen–antibody complex, although no antiserum had been added. This effect has been investigated and it now appears that the type of distribution of the antigen can be associated with different states. These are:

(1) **Carriers of serum hepatitis**

These are people who have been recognized by the blood transfusion service as having transmitted serum hepatitis but on investigation are found to have normal liver function. The sera from four such carriers have now been examined and in each case the antigen was found to be randomly distributed (Fig. 3).

(2) **Patients suffering from serum hepatitis**

The haemodialysis unit at the Hammersmith Hospital experienced an outbreak of serum hepatitis from October 1969 to February 1970 and during this time regular specimens were taken from all patients and staff on the unit and stored at −20°C. This allowed us to follow in serial fashion any patients or staff members who contracted the disease and in one case to examine serum obtained at a point before the illness had begun. These findings will be discussed in greater degree elsewhere (Trowell et al., 1971) but in summary it can be stated that in three cases followed during the course of an attack of serum hepatitis the Au-Ag changed from an initially random distribution to an aggregated one and finally these aggregates disappeared from the serum. In the one case where serum was available before the onset of the disease the antigen was found to have been present in the routinely stored specimen a week before the disease became manifest. To sum up this section it appears from our studies that in the normal course of serum hepatitis there is first a stage of non-complexed antigen leading to one where the antigen appears in aggregates which have the appearance of antigen–antibody complexes and that these are finally cleared from the system (Figs. 4–6).
(3) Persistent viral hepatitis

It has been possible to examine the serum from a small number of patients with persistent viral hepatitis and, in those found positive for the antigen, electron microscope examination has revealed the presence of the complex form of the Australia antigen for periods of up to one year. (Fig. 7)

The relevance of these different patterns of distribution is not yet established although there is corroborative evidence from other techniques substantiating the presence of immune complexes in the sera of patients with serum hepatitis (Shulman & Barker, 1969). This has led us to suggest (Almeida & Waterson, 1969) that serum hepatitis may have some aspects in common with serum sickness and that the effect of the Au-Ag may indeed be through its activity as an antigen. This speculation, however, belongs rather in the section dealing with basic uses of the microscope than in that dealing with practical aspects.

To conclude this section on the practical use of the electron microscope for the study of Australia antigen, it can be stated that the negative staining technique allows the recognition not just of the Australia antigen but also gives a possible insight into the status of the patient depending on the distribution of the antigen.

Basic studies on serum hepatitis using the electron microscope

The last section attempted to show that visualization of the Au-Ag can be of considerable diagnostic help but it must be recognized that the exact nature of the antigen is still far from certain.

Numerous attempts have been made to culture it but all of them have been unsuccessful. In addition Gerin et al. (1969) carried out biochemical assays on gradient-purified Australia antigen and were unable to detect the presence of nucleic acid. These two pieces of information, lack of biological activity and absence of nucleic acid, make it very unlikely that the Au-Ag is itself a virus. In addition, the lack of biological activity limits severely the studies that can be carried out on the antigen. This in turn means that the electron microscope is, at the present, the most fruitful approach to the problem of the Au-Ag and serum hepatitis and we shall now consider the various morphological findings that have been made.

The negatively stained appearance of the Au-Ag has already been described, but in addition to the approximately 200 Å spheres and tubules, Dane, Cameron & Biggs, (1970) described a third type of particle present in Australia antigen preparations. This particle has a double-shelled structure, the outer shell having a diameter of 420 Å and the inner, one of 280 Å (Fig. 8). In contrast to the smaller, approximately 200 Å particle, the 420 Å structure is more or less constant in size. The situation was further confounded when typical virus-like particles of 270 Å diameter (Fig. 9) were found in homogenates of two post-mortem livers from patients dying in the acute stage of serum hepatitis (Almeida et al., 1970).

All of these results have been obtained by the negative staining technique but other workers have used the thin-sectioning technique on liver biopsy material. Nowoslawski et al. in 1970 described 200 Å particles in the nuclei of hepatic cells in biopsy material from patients positive for the Australia antigen. Shortly after, Nelson, Barker & Danovitch (1970) although finding fewer infected nuclei confirmed this finding and showed 200 Å particles in liver biopsy material. These intranuclear particles have a regular diameter and have the appearance acceptable for viruses as seen by the thin-sectioning technique.

We have then four different morphological groups: the spheres and tubules of the Australia antigen (Fig. 1); the particles described by Dane et al. (1970) which we shall describe as Dane particles (Fig. 8); the 200 Å particles seen by thin sectioning; and the 270 Å particles present in liver homogenates (Fig. 9). This completes the factual information yielded by the electron microscope in serum hepatitis but it seems worthwhile to speculate on the possible nature and inter-relationship of these
four distinct groups. At first sight it might appear that a 200 Å intranuclear particle in thin-sectioned material correlates well with the 200 Å particle of Australia antigen as seen by negative staining, but this is not true. The Au-Ag particles are distinctly pleomorphic (Almeida et al., 1969) while the intranuclear particles have a reasonably constant diameter of 200 Å. In addition a particle of 200 Å in thin section would be expected to have a diameter between 270 and 300 Å in negatively stained preparations, as considerable shrinkage occurs during processing for thin sectioning. This leads to a consideration of whether the 200 Å intranuclear particle by thin sectioning does not correlate with the 270 Å particle seen by negative staining in liver homogenates. Turning now to the Dane particle the inner component of this has an established size of 280 Å and is described as having a more delicate appearance than that of the Australia antigen. This again could correlate with the 270 Å liver homogenate particle. Antibody to the Australia antigen cross links with the outer shell of the Dane particle and this would mean we are dealing with the same protein (Fig. 8). This speculation would then leave us with the suggestion that the Australia antigen is excess protein of the same type as the outer covering of the Dane particle while the particles described in liver represent the internal component of the same particle. This speculation is strengthened by the fact that Cossart & Field (1970) pointed out that there was a structural similarity between the negatively stained appearance of the Australia antigen and that of the protein component of some plant viruses (Bancroft, Mills & Markham, 1967; Bancroft, Bracker & Wagner, 1969; Bancroft, Hiebert & Bracker, 1969). We are left then with a whole series of speculations based on the multiplicity of electron microscope findings that can be summarized as follows: The most likely candidate for the virus of serum hepatitis is the Dane particle which could have as its internal component the virus-like 270 Å particles seen in liver homogenates while these in turn could identify with the 200 Å particles seen intranuclearily in thin-section preparations. On the basis of an analogous situation in plant virology there is good reason to suspect that the Au-Ag represents viral protein and has its use in diagnosis of serum hepatitis because it has antigenic identity with the outer layer of the Dane particle. However, as must obviously be clear the non-biological information obtained from the electron microscope cannot never stand by itself in characterising a virus and until some method is obtained of handling the serum hepatitis virus in vitro the electron microscope remains as a ready and simple means of visualizing the antigen and perhaps offering a suggestion as to the course future research could take.

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**References**


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