Light and electron microscopic demonstration of extracellular immunoglobulin deposition in renal tissue

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SUMMARY Extracellular immunoglobulin (IgG) deposits were shown by both light and electron microscopy in renal biopsy material using immunogold labelling. After fixation of tissue in 4% paraformaldehyde and embedding in Lowicryl K4M, semithin sections were cut and stained using the immunogold silver stain. The sections were then viewed and areas of interest were noted; ultrathin sections were cut from the same block of tissue, then stained using immunogold. Good localisation was achieved at both optical and ultrastructural levels allowing direct correlation to be made in the same area of tissue.

The demonstration of extracellular immunoglobulin deposition in renal tissue is a very important aspect in the diagnosis of renal disease in man. At the light microscopical level antibodies labelled with fluorescein or with peroxidase are the most widely used. A previously developed peroxidase technique has been used to demonstrate immunoglobulin deposition both at the light and ultrastructural level in human disease. This pre-embedding technique and its variants have some major disadvantages, the most important of which is reliably securing adequate penetration of tissue. There is also the difficulty of defining the extent of the peroxidase reaction product and the possibility of diffusion and spurious localisation. The recent development of hydrophilic resins for use in electron microscopy has meant that immunolabelling after embedding can be done and avoids the potentially damaging effects of the “etching” needed with most hydrophobic resins. These methods avoid the problems of inadequate penetration of tissue by large antibody and tracer molecules because most tissue barriers are disrupted during the sectioning of the tissue, allowing direct access of immunological reagents to the interior of the cells.

Another boost to post-embedding immunostaining has been the introduction of gold labelled antibodies. Colloidal gold offers many advantages as a label: the colloidal gold particle is electron dense and has a characteristic appearance, making it easily distinguishable from biological structures. Colloidal gold particles can be produced in virtually any size between 1–150 nm. Particles between the sizes of 5–40 nm are the most widely used: they give more efficient labelling than larger particles because they produce less steric hindrance. Gold particles of this size are not visible by light microscopy, but may be rendered so by the use of silver enhancement. These silver enhancement techniques were technically demanding and the results have been unreliable. The recent development of silver enhancement kits which are insensitive to light and are less critically dependent on ionic concentration has, in our hands, facilitated the consistent production of good light microscopic results.

The method we describe here makes use of all of the technical advances mentioned and allows direct correlation between light and electron microscopy on the same block of tissue to be made. The survival of the antigens in the resin enables post-embedding immunocytochemistry to be carried out and this has many advantages over pre-embedding techniques.

Material and methods

Fresh renal tissue was obtained from routine needle biopsies. The specimens were fixed and processed into Lowicryl K4M.

Tissue was immersion-fixed in 4% paraformaldehyde for two hours and then processed into Lowicryl K4M as follows. After fixation the tissue was dehydrated in a graded series of alcohol (methanol):
1 50% methanol at −20°C for 30 minutes;
2 80% methanol at −20°C for 60 minutes;
3 90% methanol at −20°C for 60 minutes.
The tissue was then infiltrated with the resin as follows:
1. pure methanol/pure Lowicryl in a 1:1 ratio at 
   \(-20\)°C for 30 minutes;
2. pure methanol/pure Lowicryl in a 1:2 ratio at 
   \(-20\)°C for 60 minutes;
3. pure Lowicryl at \(-20\)°C for 60 minutes;
4. pure Lowicryl at \(-20\)°C overnight.

The tissue was then embedded in fresh resin and put in a desiccator for one hour before staining was started.

** IMMUNOGOLD STAINING (IGS) **

Ultra-thin sections were cut from the same blocks of tissue used for the IGSS method using a diamond knife. Gold coloured sections (about 2 nm thick) were picked up on 200 mesh nickel grids. They were left in a desiccator for 20 minutes. Finally, the sections were counterstained with toluidine blue for two minutes, dried, and mounted.

** IMMUNOGOLD-SILVER STAIN (IGSS) **

Semithin sections (about 2 \(\mu\)m in thickness) were cut from the polymerised blocks using glass knives on an LKB ultramicrotome. They were applied to multispot glass slides coated with 0-05% poly-l-lysine (Sigma) to ensure that the sections would not separate during staining and washing of the slides. The slides were air dried at room temperature for one hour before further treatment.

The sections were covered in phosphate buffered saline (pH 7-4) for two minutes before applying dilute normal goat serum (1/1 dilution) to block non-specific binding of antibodies to the tissue. This was left for 20 minutes each in a wet chamber. The goat serum was then tipped off, but not washed off, and the sides of the slide wiped without touching the section.

Varying dilutions of rabbit anti-human IgG (heavy chain) were applied to find the optimal dilution that allowed the best staining of the sections. The diluent was PBS (pH 7-4) containing 1% cold fish gelatin. The dilutions initially used were between 1/10 to 1/8000. This primary incubation was carried out for one hour in a wet chamber followed by washing of the slides in PBS (pH 7-4), twice, for seven and a half minutes each with agitation. This was then followed by a 60 minute incubation with the secondary antibody (10 nm gold labelled antibody of anti-rabbit IgG antibody) (Jansen) in the wet chamber. The slides were removed from the wet chamber and immersed in PBS in a Coplin jar for five minutes with agitation, followed by a further wash in double distilled water twice for five minutes. The slide was dried (back and sides) with a clean drying cloth. Silver stain amplification was begun using the Intense M silver enhancement kit (Jansen). The reaction was allowed to continue for nine minutes before it was terminated by washing the slides in double distilled water for five minutes. Finally, the sections were counterstained with toluidine blue for two minutes, dried, and mounted.

** CONTROLS **

Negative and positive control tests were carried out on renal tissue, where absence or presence of human IgG had been shown previously by conventional immunofluorescence studies on frozen sections. For absorption tests, the antigen (purified human IgG, about 95% pure) was purchased from Sigma, and not from the company that supplied the antibody (Dako). Characterisation of the antigen was carried out by determining the molecular weight and isoelectric point.
of the antigen. For convenience, the absorption was measured by the quantity of antigen added in nmol/ml of optimally diluted antibody. The antigen and antibody were placed in a sealed phial and were shaken vigorously to distribute the antibody over the entire surface to make sure all the antigen was taken up. The antigen and antibody were incubated for 24 hours at 4°C before the solution was used for staining. As a further control, the antibody was also absorbed with inappropriate purified antigen, under the same conditions.

Results

Semithin sections from renal biopsy material embedded in Lowicryl K4M, incubated primarily in various concentrations of rabbit and anti-human IgG antibody, followed by a secondary incubation with goat anti-rabbit gold labelled antibody (GAR 10 nm LM grade, Jansen), showed no visible staining. When the sections were further subjected to silver amplification using the Intense M silver enhancement kit (Jansen), however, all concentrations of the anti-human IgG antibody except 1/8000 showed a positive dark, brownish-black staining on glomerular deposits. In cases of lupus nephritis there was granular staining of the glomerular capillary walls and also of the mesangium (fig 1). Staining was also recorded in the tubular epithelial cells. In cases of idiopathic membranous glomerulonephritis staining was restricted to the glomerular capillary wall but no staining was seen in the mesangial matrix (fig 2). There was also staining of resorption granules in some tubular epithelial cells.

Fig 2  Idiopathic membranous glomerulonephritis. Global, granular, staining in the glomerular basement membrane without mesangial staining. (Indirect IGSS.)

Fig 3  Electron micrograph of lupus nephritis showing labelling of a subendothelial deposit (Indirect IGS, 20 nm gold label.)
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These staining patterns were also seen on frozen sections from the same cases using immunofluorescence. Although positive staining was observed at low dilutions, the level of background staining was unacceptable and the best signal:noise ratio was obtained at a dilution of 1/2600. Both negative and positive controls confirmed the specificity and sensitivity of the method. The absorption tests showed that immunoreactivity was abolished by addition of any concentration above 0.001 nmol/ml of specific antigen.

After viewing the semithin sections, areas of interest were noted and ultra-thin sections were cut from the tissue blocks. In the cases of lupus nephritis, subepithelial, subendothelial, and mesangial electron dense deposits were intensely labelled with gold particles (fig 3). In the cases of idiopathic membranous glomerulonephritis subepithelial electron dense deposits were intensely labelled but no subendothelial labelling could be seen (fig 4). The best immunolabelling was obtained at a dilution of 1/2600, the same dilution that was considered to be optimal for staining at the light microscopical level.

Discussion

A disadvantage found in most immunohistochemical techniques is the absence of direct correlation between light and electron microscopical images, although some techniques have been reported to achieve direct correlation between the optical and ultrastructural levels. The combination of techniques we have described here allows light microscopical immunohistochemistry on one section to be compared with ultrastructural immunohistochemistry of an adjacent section from the same block. There is also the added advantage of the possibility of staining ultra-thin sections for different antigens.

Roth et al described the use of lectin-gold complexes in various animal tissues in two papers published four years apart. The first paper described localisation of...
lectin binding sites at the ultrastructural level, while the other described the localisation at the light microscopical level of those same sites. In both papers the tissues were embedded in Lowicryl K4M. In the second paper the authors reported that for cytochemical labelling with lectin-gold complexes, the same concentrations of labelling reagents applied for the incubation of ultra-thin sections were used for light microscopical examination. Our observation that the optimal dilution of primary antibody used for staining at the light microscopical level was also the optimal dilution for immunostaining at the electron microscopical level was very time saving in determining the optimal dilution for immunostaining at the ultrastructural level. Further studies are necessary with different antigens before this can be made a general rule.

The method described here has been shown to be very sensitive in localising IgG in human glomerulonephritis. We have also successfully localised IgM, IgA, and C3 in human renal tissue using the same methods (data not shown). The light microscopical demonstration of the antigen is permanent, so that the sections can be referred to at a later date and can be viewed using an ordinary light microscope. The tissue blocks can be stored at room temperature, with no loss of antigenicity in the tissue for at least eight months, or as reported by other researchers, for many years. The most important advantage is the ability to correlate the light microscopical findings with the electron microscopical level on the same block of tissue. These virtues make the method potentially valuable in histopathology.

References