Staining of human splenic sinusoids and demonstration of unusual banded structures by monoclonal antisera

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SUMMARY The monoclonal antiserum OKT8 is useful as a marker of splenic sinusoids. Banded and fibrillar structures are also delineated by this antiserum; these appear to be different from the ring fibres of the spleen and the tentative suggestion is made that these represent basal plates or endothelial striations. The reactivity of sinus lining cells with a number of antisera is noted and the conclusion is drawn that splenic sinal endothelium differs substantially from the endothelium of blood vessels.

During an investigation into the immunohistology of the human spleen we noted that the splenic sinusoids were clearly marked by I2 and OKT8 antisera. We wondered if this was due to an affinity of the antisera for endothelial cells, macrophages, ring fibres or basal plates. This paper describes our technical methods and illustrates the staining patterns found.

Material and methods

TISSUE PREPARATION

Tissue from seven fresh splenectomy specimens (four normal histology, two Hodgkin's disease, and one chronic lymphatic leukaemia) was used. For monoclonal antibody studies pieces of tissue 10 mm × 10 mm × 3 mm were surrounded with cytoprotectant, snap frozen in isopentane precooled in liquid nitrogen and stored at −70°C. Blocks for paraffin wax embedding were fixed for 24 h in neutral buffered formaldehyde pH 7-0 then overnight in formol sublimate before routine processing.

LILLIE'S ALLOCHROME METHOD

After removal of mercury and formalin pigments, paraffin sections were treated with the periodic acid-Schiff (PAS) reaction and counterstained with Sirius supra blue GL in aqueous picric acid.1 Basement membranes and ring fibres around the splenic sinuses are coloured purplish red and collagen fibres are stained blue.

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FIXATION

An acetone-chloroform-water fixative was prepared by adding four drops of distilled water to 2 ml of a mixture of equal volumes of the solvents. This fixative preserves the morphology of tissue better than acetone alone. It is not suitable for B1 and I2 preparations which were fixed in acetone only.

MONOCLONAL METHOD

The source specificities and dilution of antisera used is recorded in Table 1. Freshly cut 5 μm cryostat sections were treated as follows:

1 Air dry sections for 30 min.
2 Fix in acetone for 20 min (I2 and B1 preparations) or acetone-chloroform-water to evaporation (OKT3, 4, 6, 8, OKM1 and CLAT 200 preparations).
3 Wash twice in Tris-buffered saline pH 7-6 (TBS).
4 Flood sections with 20% normal rabbit serum in TBS for 10 min.
5 Without rinsing apply mouse monoclonal antisera for 30 min.
6 Wash twice in TBS.
7 Flood sections with 20% normal rabbit serum in TBS for 10 min.
8 Without rinsing apply rabbit antimouse peroxidase conjugated antisera for 30 min.
9 Wash twice in TBS.
10 Demonstrate peroxidase activity using diaminobenzidine.
11 Wash in running tap water.
12 Counterstain in toluidine blue or progressively
Staining of human splenic sinusoids

Table 1  Source, specificities and dilutions of antisera

<table>
<thead>
<tr>
<th>Code</th>
<th>Source</th>
<th>Specificity</th>
<th>Working dilution</th>
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<tr>
<td>Mouse monoclonal antibodies</td>
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<td></td>
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<tr>
<td>OKT3</td>
<td>Ortho Diagnostics</td>
<td>Peripheral T lymphocytes</td>
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</tr>
<tr>
<td>OKT4</td>
<td></td>
<td>Helper T lymphocytes</td>
<td>1/20</td>
</tr>
<tr>
<td>OKT8</td>
<td></td>
<td>Suppressor/cytotoxic T lymphocytes</td>
<td>1/40</td>
</tr>
<tr>
<td>OKT6</td>
<td></td>
<td>“Common thymocytes” and Interdigitating reticulum cells</td>
<td>1/30</td>
</tr>
<tr>
<td>OKM1</td>
<td></td>
<td>Monocytes, null cells, granulocytes</td>
<td>1/20</td>
</tr>
<tr>
<td>I2</td>
<td>Coulter Electronics</td>
<td>HLA D/DR related Ia-like antigen</td>
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</tr>
<tr>
<td>B1</td>
<td>Hybritech</td>
<td>B lymphocytes</td>
<td>1/200</td>
</tr>
<tr>
<td>Common leucocytes</td>
<td></td>
<td>All haemopoietic cells</td>
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<tr>
<td>Antigen T200 (CLAT200)</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>P161</td>
<td>DAKO immunoglobulins a/s</td>
<td>Rabbit antimouse immunoglobulins</td>
<td>1/20</td>
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</table>

*This antiserum was absorbed with human IgG (Koch light 285501) in the ratio 2 mg/100 μl neat antiserum immediately before use.

in Mayer’s haematoxylin.
13 Dehydrate clear and mount.

REDUCTION OF BACKGROUND STAINING
Sections were incubated in normal rabbit serum (Gibco Biocult 200—6120) before the antisera were applied.

DIAMINOBENZIDENE VISUALISATION OF PEROXIDASE ACTIVITY
A modified Graham and Karnovsky solution was used. Like Borowitz et al we have shown that incorporation of imidazole in the diaminobenzidine-Tris buffer significantly improves antigen localisation. The working solution was prepared by adding 4 mg diaminobenzidine tetrahydrochloride (Sigma) to 5 ml of TBS containing 0.01 M imidazole. When this had fully dissolved 0.1 ml of a 1% hydrogen peroxide solution was added. This solution was filtered before use.

ALKALINE PHOSPHATASE
The activity of this enzyme was demonstrated in 5 μm cryostat sections using the “Neufuchsins” method of Stutte. Sections were fixed in 10% formol-methanol at 4°C for 30 s before incubation in the substrate solution, naphthol-ASTR-phosphate.

Results
Under the low power of the microscope the brown deposit of the immunoperoxidase reaction obtained with OKT8 delineated clearly the sinusoids. Little or

Fig. 1  Sinusoidal wall staining by OKT8. Immunoperoxidase × 300 (approx).
no staining was visible in the Malpighian bodies or cords of Billroth. Occasional mononuclear cells situated within the sinusoids or cords were stained but this did not detract from the overall impression of specific sinusoidal wall staining (Fig. 1).

A more diffuse and much less discriminating picture was seen with anti-I2 where the majority of splenic cells gave a positive reaction. The pattern of sinusoidal staining was similar to that obtained with OKT8.

At a higher magnification the staining had a banded appearance. Cross sections of sinusoids showed darkly staining bands arranged at right angles to the luminal surface and extending outwards to the periphery. The bands are separated by a gap of approximately 2–3 μm width. It is difficult to say if these bands are located between endothelial cells or are truly intracytoplasmic structures (Fig. 2). Longitudinal sections through sinusoids showed fine linear streaks or fibres running parallel to the long axis (Fig. 3). The cytoplasm of the endothelial cells was also stained.

Table 2  Reactivity of sinus lining cells with monoclonal antisera

<table>
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<tr>
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<th>5</th>
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<td>±</td>
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<td>ND</td>
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</tbody>
</table>

+ = staining; - = no staining; ± = equivocal staining; ND = not done.

Cases 1–4 Normal histology.
Cases 5–6 Hodgkin's disease.
Cases 7 Chronic lymphatic leukaemia.

Fig. 2  Banded appearance in transverse section of sinusoids. OKT8 and immunoperoxidase × 600 (approx).

Fig. 3  Longitudinal fibrillar structures (top). OKT8 immunoperoxidase × 600 (approx).

No delineation of sinuses or sinal cells was observed with the monoclonals OKT3, OKT6, OKM1, and B1. OKT4 gave equivocal results. Monoclonal CLAT200 stained most cells of the Malpighian body and sinal endothelial cells but did not produce the banded staining of grid structures seen with OKT8.

The reactivity of sinal endothelium with all markers is shown in Table 2.

Alkaline phosphatase activity was noted in the endothelium of trabecular vessels and collecting
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Trabecular vascular endothelium failed to react with the OKT8 serum. Endothelial cells isolated from umbilical veins do not react with OKT8 serum but give positive reactions with la-like antisera.6 The nature of splenic endothelium has been an unsettled question for many years and these marker studies provide additional evidence that vascular and splenic sinus endothelium differ substantially.

It first seemed to us that the banded and fibrillary structures corresponded to the annular or ring fibres which have been well described by King, Puchtler and Sweat.7 These are stained by Lillie's allochrome method and although these fibres are often branched, the majority are annular, surrounding the sinusoids like the hoops of a barrel. However, the fibrils stained by OKT8 run longitudinally and are arranged at right angles to the ring fibres. Furthermore, cross sections of sinusoids reveal a banded venules; sinus lining cells gave a negative reaction.

Classical ring fibres were demonstrated by Lillie's allochrome method in all cases (Fig. 4). These fibres coursed at right angles to the fibres outlined by OKT8 serum.

Discussion

The clear delineation of sinuses by the monoclonal antiserum OKT8 might be due to staining of endothelial cells or sinus associated cells such as the processes of macrophages as described by Burke and Simon8 or Schridde's delicate protoplasmic network composed of small spider-like cells which lie just without the endothelium and extend over its surface. The antimonocyte monoclonal OKM1 stains resident perifollicular macrophages as well as some histiocytes, and thus might reasonably be thought to react with sinus associated macrophages but no such staining pattern was observed. On the other hand the majority of sinus lining cells gave a distinct and positive reaction with OKT8 and it seems inescapable that these cells contain an antigen in common with suppressor or cytotoxic lymphocytes. Sinus lining cells gave positive reactions with anti-I2 and anticommon leucocyte antigen sera; the borderline reactions seen in two cases with OKT4 perhaps requires further investigation. The sinus lining cells failed to stain for alkaline phosphatase unlike the endothelium of the trabecular vessels and collecting venules. It is noteworthy that

Fig. 4 Annular transverse hoops or ring fibres. Lillie's allochrome × 600 (approx).

Fig. 5 Drawing of a sinusoid by Nathan Chandler Foot (1927) showing ring fibres and the endothelial cell striations. Reproduced with permission of the Editor, Dr Lachman, of Anatomical Record published by the Wistar Press.
arrangement quite unlike the dot-like pattern seen in sections stained by Lillie's allochrome method. On the contrary, they resemble Foot's description of the basal plates or striations. He reviewed the early literature and showed that splenic endothelial cells were rod shaped rather than fusiform and contained areas staining blue with phosphotungstic acid haematoxylin (Fig. 5). He thought these were different from the argyrophilic reticular annuli. Our preliminary observations support the view that these basal plates or endothelial cell striations contain OKT8 antigen. It is not known if this reaction is adventitious or whether it conceals a purpose. Because these fibrillary structures form a grid through which cells must pass when migrating between cord and sinus, we put forward the view that the antigens associated with them act as biochemical signposts which modulate cellular traffic through the spleen.

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References


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