Macrophage origin of Reed-Sternberg cells: an immunohistochemical study

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SUMMARY In an immunohistochemical study of 26 biopsies from 24 patients with Hodgkin's disease a granular staining pattern for alpha-1-antitrypsin (a,AT) and alpha-1-antichymotrypsin (a,ACT) was seen in Reed-Sternberg (RS) cells and mononuclear Hodgkin's (H) cells in over half the cases. The pattern of staining for these antiproteases seen in RS and H cells has previously only been observed in normal and malignant cells of the monocyte/macrophage lineage within the lymphoreticular system. A faintly granular evenly distributed staining for IgG was found in viable RS and H cells. This staining was associated with a similar distribution of both light chains but not J chain, suggesting that the immunoglobulin had not been synthesised by these cells but had been taken up from the extracellular environment. It is suggested that this uptake is an active process occurring in viable RS and H cells, possibly via Fcγ receptors and further supports an origin from cells of the monocyte/macrophage lineage. IgA, IgD, albumin, fibrinogen, C1q, C4 and C3 were present in some cells, IgM was more rarely found and lysozyme was absent. The fact that cells staining for these serum proteins generally showed signs of degeneration and that the extent of staining correlated with the molecular weight, but not serum concentration, of the protein suggests that they are passively acquired by dead or dying cells and thus represent a separate phenomenon from IgG uptake. The function of IgG uptake and accumulation by RS cells and the a,AT and a,ACT markers may prove of use in identifying the macrophage subtype from which these cells are derived.

An understanding of the origin of Reed-Sternberg (RS) cells is central to our understanding of the interaction of these tumour cells with the accompanying reactive lymphoid and histiocytic elements and of the relevance of these interactions to the pathogenesis of Hodgkin's disease. Ultrastructural evidence has been inconclusive since it has been interpreted as indicating both a histiocytic1 2 and lymphocytic origin.3 4 The presence of non-specific esterase and acid phosphatase favours a histiocytic origin5 6 although staining for peroxidase, lysozyme and with antimacrophage serum has proved negative.6 7 The establishment of cell lines with macrophage-like features from Hodgkin's tissue8 9 and the demonstration that in vitro RS cells have phagocytic potential10 suggests that these cells have more in common with macrophages than lymphocytes. More specifically a dendritic pattern seen with Marshall's metalophil stain has been interpreted as showing identity with dendritic reticulum cells.11

Attention has focused on the identification of surface and cytoplasmic IgG on RS cells12-16 taken by some as indicative of a derivation from B lymphocytes.7 However, since this IgG is associated with both light chains within individual cells it seems more likely that rather than having been synthesised it has been acquired from an exogenous source.10 17 Ultrastructural studies have not confirmed this however, since electron microscope immunoperoxidase preparations have shown a cytoplasmic granular product not related to endocytic structures. This distribution has been interpreted as indicating both specific ribosomal synthesis18 19 and nonspecific leakage into the cell.17 Other serum proteins have been found in RS cells but less commonly than IgG.15-17

In the light of the conflicting interpretations as to the extent and significance of IgG and other proteins in RS cells we have performed an immunoperoxidase study of 12 different serum proteins including IgG, on a series of biopsies from patients with Hodgkin's disease. We have stained for J chain although associated extracellularly only with dimeric IgA and

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19 S IgM is synthesised in cells producing immunoglobulin of all classes. The association of J chain with intracellular IgG is therefore indicative of immunoglobulin synthesis, lack of such an association may reflect uptake of serum protein. A characteristic staining pattern for alpha-1-antitrypsin (α1AT) has proved to be a useful marker for cells of the macrophage series. We have, therefore, looked critically at the staining pattern in RS cells for this antiprotease and for the related inhibitor, alpha-1-antichymotrypsin (α1ACT).

Material and methods

Twenty-six biopsies from 24 Hodgkin’s patients were studied. Twenty-one biopsies were of the nodular sclerosing type, one was lymphocyte and histiocyte predominant and two showed mixed cellularity. Sections were cut from routine formalin-fixed paraffin embedded material, deparaffinised in xylol and absolute alcohol and treated with 0·5% hydrogen peroxide in methanol to block endogenous peroxidase activity. Sections were then treated according to the trypsin/peroxidase-antiperoxidase (PAP) method. Briefly, they were incubated with 0·1% trypsin (Sigma Chemical Company, Poole) in 0·1% CaCl2, pH 7·8, at 37°C for 15 min followed by washing in distilled water and TRIS-NaCl. They were then stained for a variety of proteins by first incubating with specific rabbit antisera (Behringwerke AG, Marburg-Lahn) followed by swine anti-rabbit Ig and PAP complexes (Dakopatts AS, Copenhagen). Rabbit antisera specific for the following proteins were used as first layers: γ, μ, δ, α, κ and λ Ig chains, J chain and α1AT, α1ACT, C3, C4, Clq and lysozyme. Suitable dilutions of these antisera were established by staining control tissues. Sites of peroxidase activity were then developed with diaminobenzidine and sections counterstained with haematoxylin. Control sections were treated with normal rabbit serum in place of specific antiserum.

The specificity of the rabbit antisera was established by radial immunodiffusion and immunoelectrophoresis. The antihuman Ig sera were further tested against defined antigen substrate spheres provided by Professor G Stevenson of the Tenvous Research Laboratory, Southampton. The specificity of the anti-J chain, anti-lysozyme and anti-α1AT sera was confirmed by absorption of activity with purified J chain, lysozyme and α1ACT, respectively. Competitive blocking of staining for Clq, C3, C4, albumin and fibrinogen by goat antiserum (Nordic Immunology, Tilburg) established the specificity of these antisera.

Results

IgG was present in RS cells and mononuclear H cells in 26/26 biopsies and both light chains were detected in similar numbers of cells in 25/26 cases (Table). The proportion of IgG-positive RS and H cells varied considerably between biopsies; the numbers of biopsies in which <5%, 5-30% and >30% of RS cells were positive being 31%, 42% and 27% respectively. Hence in the majority of biopsies less than 30% of RS and H cells contained IgG. The staining varied in intensity between neighbouring RS cells, was usually less intense than that of surrounding plasma cells and was in two basic patterns. The majority of positive cells were of type I in which the reaction product was faintly granular and distributed evenly throughout the cytoplasm.

### Extent of staining for 12 serum proteins, κ, λ, and J chains in Reed-Sternberg cells and mononuclear Hodgkin’s cells

<table>
<thead>
<tr>
<th>Protein</th>
<th>Molecular weight (× 10^4)</th>
<th>Serum concentration (mg/100 ml)</th>
<th>Cases positive/cases studied</th>
<th>% Cases positive</th>
</tr>
</thead>
<tbody>
<tr>
<td>(a) Type I granular staining of viable cells and type II marginated staining of degenerate cells</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IgG</td>
<td>150</td>
<td>1200</td>
<td>26/26</td>
<td>100</td>
</tr>
<tr>
<td>IgA</td>
<td>160</td>
<td>210</td>
<td>12/26</td>
<td>46-2</td>
</tr>
<tr>
<td>IgM</td>
<td>900</td>
<td>100</td>
<td>2/26</td>
<td>7-7</td>
</tr>
<tr>
<td>IgD</td>
<td>170</td>
<td>3</td>
<td>5/26</td>
<td>22-7</td>
</tr>
<tr>
<td>α chain</td>
<td></td>
<td></td>
<td>25/26</td>
<td>96-2</td>
</tr>
<tr>
<td>λ chain</td>
<td></td>
<td></td>
<td>3/23</td>
<td>13-0</td>
</tr>
<tr>
<td>J chain</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Clq</td>
<td>400</td>
<td>15</td>
<td>3/19</td>
<td>15-8</td>
</tr>
<tr>
<td>C3</td>
<td>185</td>
<td>82</td>
<td>11/22</td>
<td>50-0</td>
</tr>
<tr>
<td>C4</td>
<td>230</td>
<td>40</td>
<td>4/21</td>
<td>19-0</td>
</tr>
<tr>
<td>Albumin</td>
<td>68</td>
<td>4000</td>
<td>16/25</td>
<td>64-0</td>
</tr>
<tr>
<td>Fibrinogen</td>
<td>54</td>
<td>300</td>
<td>6/21</td>
<td>28-6</td>
</tr>
<tr>
<td>α1AT</td>
<td>68</td>
<td>45</td>
<td>12/17</td>
<td>70-6</td>
</tr>
<tr>
<td>α1ACT</td>
<td>15</td>
<td></td>
<td>0/21</td>
<td>0</td>
</tr>
<tr>
<td>(b) Staining as an accumulation of granules adjacent to the nucleus</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>α1AT</td>
<td></td>
<td></td>
<td>13/24</td>
<td>54-2</td>
</tr>
<tr>
<td>α1ACT</td>
<td></td>
<td></td>
<td>11/17</td>
<td>64-7</td>
</tr>
</tbody>
</table>

*Very weak staining in a few cells.*
with no localisation to the perinuclear space or cell margin. The cytoplasm of these cells extended up to and in between neighbouring cells and the nuclei were intact (Figs. 1 & 2). In cells of type II the cytoplasm had usually retracted away from the neighbouring cells, the nucleus was dark and shrunken and the peroxidase stain was diffuse, darker than in type I and more concentrated at the retracted margin of the cell (Fig. 4).

IgA-positive RS and H cells were seen in 12/26 biopsies and IgD-positive cells in 5/22 (Table). IgA and IgD was present in only a minority of RS and H cells and the staining in these cells was usually of type II although occasionally of type I. Staining for IgM and J chain was negative with the exception of a few shrunken cells positive for IgM in two cases and the occasional cell weakly positive for J chain in three cases (Fig. 3). In 2/3 J chain-positive cases IgM- or IgA-positive RS cells, or both, were also detected. Staining for lysozyme was consistently negative in RS cells but RS cells staining for all other proteins tested, namely Clq, C3, C4, albumin, fibrinogen, α1AT and α1ACT were present in some biopsies (Table). The staining for these proteins was similar to that of IgA and IgD in that it was present in a minority of cells and was predominantly of type II. With the exception of IgG, which was present in all cases, and lysozyme, which was present in none, there was no positive correlation between the proportion of cases containing RS and H cells staining for each protein and the molecular weight of the protein (Table); the smaller proteins being more commonly found. There was no such correlation between the extent of staining and serum concentration.

In addition to the diffuse staining for α1AT in degenerating cells in 79% of cases a distinctive staining pattern, unlike that of the other proteins, was found in RS and H cells in over half the cases studied (Table). The staining in these cases was localised as an accumulation of granules adjacent to the nucleus or tucked into a nuclear indentation (Fig. 5). This pattern of staining was also seen in reactive histiocytes and in normal peripheral blood monocytes (Fig. 6). The proportion of RS and H cells that stained in this pattern varied considerably between biopsies. In a few cases more than 50% RS and H cells were positive but in most cases only a minority were stained. A similar staining pattern for α1ACT was seen in RS and H cells in 11/17 cases (Fig. 5). Reactive histiocytes and monocytes stained for α1ACT in an identical pattern to α1AT.

Plasma cells were common in Hodgkin’s tissue; the frequency of heavy chain classes being γ < α < μ < δ. There was no positive correlation between plasma cell numbers and immunoglobulin positive RS cells, an inverse relation between these two cells being more commonly observed.

Discussion

Many RS cells contain polytypic IgG,7, 10–17 Ultrastructural studies have produced conflicting evidence as to whether this immunoglobulin is associated with ribosomes17–19 and since it does not appear to be associated with endocytotic structures it is not clear whether it has been synthesised or acquired by these cells. Other immunoglobulins, albumin, α1AT and α1ACT, have also been found within RS cells10–17 but reports of their incidence vary widely and little attention has been paid to their staining pattern or cytoplasmic distribution. In this study we have shown that 11 serum proteins other than IgG can be demonstrated in RS cells but that, contrary to other reports,17 their incidence and staining pattern differ from IgG and they are thus thought to represent a different phenomenon.

The proteins detected, in their order of frequency, were α1AT, α1ACT, albumin, C3, IgA, fibrinogen,
IgD, C4, C1q and IgM. This is in agreement with other reports that these first three proteins are quite commonly detected in RS cells, but differs in finding IgM the least commonly detected protein. Poppema et al. reported that albumin and α1AT were present in the same cells that contained IgG. In this study these proteins were found less frequently than IgG and their staining pattern differed in being non-granular and concentrated at the margin of the cell which usually showed morphological signs of degeneration. Together with the observation that the extent of staining for these proteins correlated with their molecular size it is suggested that they are passively acquired by dead or dying cells perhaps as a consequence of membrane damage. It is significant that viable RS cells in vitro do not contain albumin or fibrinogen.

In all but three cases J chain was not present in RS cells. This polypeptide is associated with extra-

Fig. 2 Staining for (a) κ and (b) λ light chains within Reed-Sternberg and mononuclear Hodgkin’s cells. The staining is similar to that of IgG being variable, granular and within cells showing no evidence of degeneration × 640.

Fig. 3 Reed-Sternberg cells and mononuclear Hodgkin’s cells fail to stain for J chain × 640.
Macrophage origin of Reed-Sternberg cells

Fig. 4 Reed-Sternberg cells showing signs of degeneration staining for (a) IgG and (b) albumin. Staining is non-granular and more concentrated at the cell periphery. (a) × 640; (b) × 2500.

Fig. 5 Staining for (a) α,AT and (b) α,ACT in Reed-Sternberg and mononuclear Hodgkin's cells. (c) High power shows focal, granular nature of staining situated close to the nucleus or within a nuclear indentation (arrow) (a) and (b) × 640; (c) × 1600.
cellular dimeric IgA and 19 S IgM but not with IgG. Its synthesis, however, is not confined to IgA and IgM producing cells and may be found in normal and malignant cells producing immunoglobulin of all classes.20–22 The absence of J chain from γ chain-positive RS cells suggests that IgG has not been synthesised by these cells but has been taken up from the extracellular environment. Very weak staining for J chain was seen in a few RS cells in three cases but it was accompanied by IgA- and IgM-positive cells in two of these.

In addition to being more common the staining pattern for IgG differed from that of the other proteins in that it was granular, distributed evenly throughout the cytoplasm and was present in cells showing no morphological evidence of degeneration. It is suggested that it is acquired by a different mechanism from the other proteins and in view of the demonstration that RS cells in vitro can phagocytose immune complexes and internalise aggregated IgG,10 it is likely that a similar process, possibly involving Fcγ receptors, occurs in vivo. In support of an active uptake of serum IgG by RS cells is the observation that IgG is lost or reduced in RS cells during in vitro culture in the absence of human serum.27

The source of the IgG in RS cells in tissue sections is uncertain but the presence of circulating immune complexes in Hodgkin’s patients may be relevant.28 29 The basic reason why RS cells should take up and accumulate IgG remains unclear unless it is considered as the deranged function of a particular macrophage subset that is involved in the uptake or handling of immune complexes. This is in keeping with the suggestion that RS cells are related to dendritic reticulum cells.11 It is interesting that a derivation from the interdigitating reticulum cell of the T cell areas, has also been proposed30 a suggestion that is supported by the close association of autologous T cells with the RS cell surface.31

In addition to the diffuse staining for α1AT and α1ACT seen in degenerating RS cells, an additional staining pattern for these protease inhibitors was found in viable RS cells in over half the cases studied. In these cases a variable number of RS cells showed staining for α1AT or α1ACT, or both, as an accumulation of granules tucked into a nuclear indentation or situated close to the nucleus. Alpha-1-antitrypsin is present in polymorphonuclear leucocytes, platelets, monocytes and macrophages,24 32–35 pancreatic islet and acinar cells36 37 and in lymph node venule endothelial cells and kidney tubular epithelium (unpublished observations). It is not present within cells of the lymphocyte series although it may be associated with the surface of mitogen-activated lymphoblasts.38 The main site of synthesis of the serum protein is the liver but it has been suggested that platelets, pancreatic islet cells

Fig. 6 Staining for α1AT in (a) reactive histiocytes in a Hodgkin’s lymph node and (b) normal peripheral blood monocytes. Note focal, granular nature of stain. (a) × 640; (b) × 1040.
and macrophages may be capable of independent synthesis.\textsuperscript{34} Whatever the origin of $\alpha_1$AT in normal monocytes and macrophages the nuclear-orientated granular staining in immunoperoxidase preparations is characteristic of this lineage. The monocyte/macroage origin of RS cells is supported by several lines of recent evidence including phagocytic activity and tissue culture properties.\textsuperscript{5–10} The expression of protease inhibitors in RS cells is variable as it is among normal monocytes and histiocytes. This variability may arise because they are the products of particular macrophage subsets, and it may be through studies of such macrophage subset markers that a clearer understanding of the nature of RS cells will emerge.

References


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