High temperature antigen retrieval and loss of nuclear morphology: a comparison of microwave and autoclave techniques

N C A Hunt, R Attanoos, B Jasani

Abstract
The use of high temperature antigen retrieval methods has been of major importance in increasing the diagnostic utility of immunocytochemistry. However, these techniques are not without their problems and in this report attention is drawn to a loss of nuclear morphological detail, including mitotic figures, following microwave antigen retrieval. This was not seen with an equivalent autoclave technique. This phenomenon was quantified using image analysis in a group of B cell lymphomas stained with the antibody L26. Loss of nuclear morphological detail may lead to difficulty in identifying cells accurately, which is important in the diagnostic setting—for example, when trying to distinguish a malignant lymphoid infiltrate within a mixed cell popu-
lation. In such cases it would clearly be wise to consider the use of alternative high temperature retrieval methods and accept their slightly lower staining enhancement capability compared with the microwave technique.


Keywords: antigen retrieval, artefacts, immunocytochemistry.

The masking of tissue antigens in routinely processed histological specimens has been a major obstacle to the application of immunocytochemistry to archival material. A variety of techniques have been successfully developed to reveal such antigens, including enzymatic and high temperature methods. These techniques have been important in overcoming the obvious limitations imposed by the dependence on frozen tissue and their development has permitted the diagnostic utility of immunocytochemistry to increase dramatically. However, these methods are not without disadvantages. Here, we present histomorphometric data from a study comparing our standard autoclave and microwave pretreatment protocols in a group of B cell lymphomas stained with the L26 antibody (anti-CD20 B cell lineage marker). Our results demonstrate a potentially significant difference between these methods with implications for diagnostic work.

**Methods**

Six cases of non-Hodgkin’s B cell lymphoma were identified retrospectively from the archives of the Department of Histopathology, University Hospital of Wales. All cases were of diffuse, centroblastic/centrocytic type. Sections, 4–5 μm thick, were cut from each tumour and were untreated or pretreated in an autoclave (0.01 M sodium citrate buffer, pH 6.5; 15 minutes at 120°C; 15 psi), or pretreated in a microwave oven (0.01 M sodium citrate buffer, pH 6.5; 15 minutes at 800 W (Matsui TC, Harrow, UK)).

Following pretreatment, all sections were stained with L26 (Dako, High Wycombe, UK) (diluted 1 in 40) and incubated at 4°C overnight. Sections were then stained using a simple, indirect immunoperoxidase method. All sections then underwent histomorphometric analysis of mean nuclear area and mitotic index. All samples were examined blind.

To assess the apparent difference in nuclear size observed between the control, autoclave and microwave treated sections, the cross-sectional area of 300 randomly selected nuclei was measured from each section using an IBAS image analyser (Kontron, Eching, Germany), with a Videoplan software package, coupled to an Olympus (London, UK) BH2 microscope. The data were gathered by interactively defining the nuclear circumference using computer generated graphics superimposed over a live, colour television image. From these data the mean cross-sectional nuclear area for each section was calculated. Mitotic indices were then determined for each section using a slight modification of the methodology described by van Diest et al. In each case 50 high power fields were counted using an Olympus BH2 microscope (×40 objective; field diameter 500 μm).

The histomorphometric data were analysed by comparison of means using standard errors. The mitotic indices were analysed for significant differences using a Wilcoxon rank sum test for non-parametric data.

**Results**

The results of the histomorphometric analysis of nuclear area are presented in table 1. There seems to be a constant difference between the sections pretreated with microwaves compared with the control and the autoclave pretreated sections in terms of nuclear area, which was consistently higher in the former. This reaches statistical significance in each case with the null hypothesis being rejected at p < 0.001 (table 1). Analysis of the mitotic indices for sections from each case showed a consistent difference between the sections pretreated with microwaves compared with both control and autoclave pretreated sections. In all cases the mitotic index was lower in the microwave treated sections.

In qualitative terms a number of differences were noted between the variously treated sections. The microwave pretreated sections showed a clear increase in intensity of staining when compared with controls. The autoclave pretreated sections also showed an increase in staining intensity compared with the control sections, although overall enhancement was slightly less than in the microwave pretreated group. However, there was also a clear, qualitative difference between the sections in terms of cellular morphology with the control sections having the best morphology and those pretreated with microwaves the worst. The autoclave pretreated group had lost much less morphological detail than the microwave pretreated group (fig 1) and compared favourably with the control group. Specifically, the cells in the microwave pretreated specimens appeared rather swollen with associated loss of nuclear detail and often complete blurring of mitotic figures. These observations are, we believe, the qualitative correlates of the quantitative data presented above.

**Discussion**

Immunocytochemistry has become an invaluable tool in routine diagnostic histopathology as well as an important research technique. The development of antigen retrieval tech-

---

**Table 1** Comparison of mean nuclear area of centrocytic/centroblastic cells and mitotic index with and without pretreatment by autoclave or microwave antigen retrieval methods. Results expressed as mean (SD) for area

<table>
<thead>
<tr>
<th>Control</th>
<th>Autoclave</th>
<th>Microwave</th>
</tr>
</thead>
<tbody>
<tr>
<td>Case number</td>
<td>Area (μm²)</td>
<td>Mitotic count</td>
</tr>
<tr>
<td>1</td>
<td>19.03 (4.42)</td>
<td>11</td>
</tr>
<tr>
<td>2</td>
<td>18.25 (3.29)</td>
<td>8</td>
</tr>
<tr>
<td>3</td>
<td>20.56 (4.65)</td>
<td>30</td>
</tr>
<tr>
<td>4</td>
<td>17.29 (3.49)</td>
<td>37</td>
</tr>
<tr>
<td>5</td>
<td>16.43 (3.65)</td>
<td>33</td>
</tr>
<tr>
<td>6</td>
<td>17.36 (3.65)</td>
<td>17</td>
</tr>
</tbody>
</table>
niques has been instrumental in this process as it has permitted the application of antibodies to routinely processed tissue samples and has, in many cases, abrogated the need for frozen material. Such antigen retrieval can be achieved using both enzymatic and, more recently, high temperature techniques. The former techniques, whilst offering the potential for using formalin fixed, paraffin wax embedded material for immunocytochemistry, were difficult to standardise and do not offer significant staining enhancement with many antibodies. The latter techniques involve superheating tissue sections with either a microwave oven, autoclave or pressure cooker. There are a number of possible mechanisms likely to be responsible for antigen retrieval using these techniques, including breakage of formaldehyde induced protein cross linkages, denaturation of proteins to reveal previously masked epitopes and unmasking of epitopes by removal of calcium ions.

To date, most experience has been gained with microwave techniques, perhaps because workers already have experience with this technology. However, microwaves have certain drawbacks including the fact that they work best with a small number of cases, problems with evaporation of buffer, inconsistent staining due to local superheating, and their relatively frequent mechanical failures due to overuse. In addition, we have also noted in our own work on B cell lymphomas that microwave pretreatment seems to impair cellular morphology significantly. A diminution in morphology following microwave pretreatment has also been noted by other workers, although they do not seem to have assessed this phenomenon in any detail. Because of the implications of this in diagnostic work, especially when only limited material is available and enhancement is needed, we decided to undertake a formal study of this phenomenon as presented in this short report.

The cellular distortion with microwave pretreatment of tissue sections is reflected in the increase in mean nuclear area and the apparent decrease in the mitotic index. While it is possible to speculate on the underlying changes responsible for these observations, it would be rather more difficult to delineate these in exact detail. However, one possibility is that microwaves cause some structural damage to intracellular macromolecules which, in turn, leads to an increase in the number of osmotically active moieties within the nuclear compartment, thus attracting water and causing nuclear swelling. An intrinsic property of microwaves is proposed as superheating alone cannot be the answer as evinced by the much better morphological preservation using an autoclave. Such changes would certainly explain the increase in mean nuclear area observed and furthermore, this swelling would have the effect of blurring mitotic figures and making them less easy to distinguish accurately from apoptotic bodies or lymphocytes and thus less likely to be counted. Given that this study was undertaken using the optimised protocols, used in our laboratory for diagnostic work, these observed changes may have important consequences for selection of retrieval methods in the diagnostic setting when it becomes important to identify accurately the cells which have been stained. Both mitotic figures and morphology may be of importance in achieving this—for example, when trying to distinguish a malignant lymphoid infiltrate within a mixed cell population. In such cases it would clearly be wise to consider the use of alternative high temperature retrieval methods and accept their slightly lower staining enhancement capability compared with the microwave technique.

Figure 1 Photomicrographs showing (A) an autoclave pretreated section and (B) a microwave pretreated section of lymphoma (original magnification ×200).

Abstract

A unique case of primary myxoid liposarcoma of the thigh, in which focal pleomorphic areas were present containing rhabdomyoblasts, is described. Focal rhabdomyosarcoma in liposarcoma has only rarely been reported previously and only in dedifferentiated liposarcomas of the retroperitoneum. All but one have been recurrences with rhabdomyoblasts being absent in the primary liposarcoma. As rhabdomyoblasts were only focally present, the present case is regarded as liposarcoma with focal divergent rhabdomyoblastic differentiation rather than malignant mesenchymoma.

Keywords: liposarcoma, rhabdomyosarcoma, malignant mesenchymoma.

Liposarcoma may contain benign or malignant heterologous mesenchymal elements. Benign cartilage,1 smooth muscle,2 or rarely bone3 have been described in well differentiated liposarcoma/atypical lipomatous tumours or in myxoid liposarcoma. Until now, malignant heterologous elements have been found only in dedifferentiated liposarcoma.1,5,6

Malignant myogenic elements found in liposarcoma have included leiomyosarcoma, rhabdomyosarcoma, or both. Other heterologous elements including angiosarcoma or osteoid have also been described rarely.1 Divergent rhabdomyosarcomatous differentiation has been reported previously in only seven cases of liposarcoma.4,5,6 All were dedifferentiated liposarcomas arising in the retroperitoneum and in all but one case rhabdomyosarcoma was found only in tumour recurrences. There is only a single previous case of rhabdomyosarcomatous differentiation in de novo liposarcoma, a retroperitoneal dedifferentiated tumour.3

Here, we describe a unique case of divergent rhabdomyosarcomatous differentiation in a de novo liposarcoma, which was of combined myxoid and pleomorphic subtypes arising in the thigh. Immunohistochemical and ultrastructural evidence is provided to support our diagnosis.

Case report

A 75 year old man presented with a six week history of a swelling in the right thigh. This was associated with oedema of the right leg, but there were no systemic symptoms. There was no evidence of any abdominal mass. On exploration, a mass was found to be displacing, but not invading, the femoral vessels and nerves. The tumour was easily mobilised except on the posterior surface where it was adherent to the pectineus muscle.

Macroscopically, the tumour consisted of a partly encapsulated fatty mass measuring 12 cm at its maximum dimension with a small amount of surrounding soft tissue. It was partly necrotic. Histologically, the tumour was a liposarcoma which was predominantly of myxoid type with a typical delicate branching capillary network and myxoid stroma (fig 1). In places, the tumour had more cellular pleomorphic areas. The latter were composed of spindle cells with notable nuclear pleomorphism and bizarre lipoblasts containing typical scalloped nuclei and cytoplasmic lipid vacuoles (fig 1, left upper inset). Mitotic activity was frequent (21/10 high power fields) within the pleomorphic areas and large foci of necrosis were seen. The tumour grade was 3 based on the method by Trojani et al.8

Cells within the pleomorphic areas had abundant eosinophilic cytoplasm and noticeably hyperchromatic nuclei (fig 1, right lower inset), many showing notable nuclear enlargement and irregular nuclear contours. Pleomorphic multinucleate tumour giant cells were also seen. No cross striations were visible by light microscopy. Immunohistochemistry was performed using the streptavidin-peroxidase complex technique with diaminobenzidine chromogen substrate. The pleomorphic cells showed intense cytoplasmic positivity using immunostains for desmin (diluted 1 in 25; Dako, High Wycombe, UK). Myoglobin (diluted 1 in 1000; Dako) and muscle specific actin (HHF 35) (diluted 1 in 40; Biomen, Finchampstead, UK) were also positive in these cells. A proportion of them were positive for fast myosin (diluted 1 in 600; Sigma, Poole, Dorset, UK) and sarcomeric actin (diluted 1 in 100; Sigma). No tumour cells stained for

Department of Histopathology, Christie Hospital NHS Trust, Wilmslow Road, Withington, Manchester M20 4BX

Correspondence to:
Dr SS Banerjee.

Accepted for publication
7 May 1996