The form factor of alpha-naphthyl acetate esterase-positive cells in non-Hodgkin’s lymphomas and reactive lymph nodes

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SUMMARY The shape of alpha-naphthyl acetate esterase (ANAE)-positive cells (other than T lymphocytes) has been measured in 40 lymph nodes. The specimens comprised 15 high-grade lymphomas, 15 low-grade lymphomas and 10 reactive lymph nodes. The parameter used for the measurement of shape was form factor (FF), which is readily calculated by the Reichert-Jung (Kontron) MOP-AMO, user-controlled image analyzer. Perfectly round cells have an FF value of 1·0, whereas the FF of irregularly-shaped cells diverges from unity. It has been demonstrated that the ANAE-positive cells in high-grade lymphomas have mean values for FF of 0·8–0·9, whereas in low-grade lymphomas and reactive nodes the mean value is 0·4–0·5. Thus, high-grade lymphomas contain many more rounded ANAE-positive macrophage type cells than do low-grade lymphomas and reactive nodes. In the latter two types of specimen there is an excess of branching and spindle-shaped ANAE-positive cells.

During a study of the numbers of alpha-naphthyl acetate esterase (ANAE)-positive cells in non-Hodgkin’s lymphomas (NHL)1 it was subjectively noted, in passing, that these cells were, in high-grade NHL, more often of the rounded, tingible-body type. Conversely, the impression was gained that branching forms of ANAE-positive cells outnumbered the rounded forms in low-grade NHL and in reactive nodes. To test this impression objectively, 40 specimens were examined using the Reichert-Jung (Kontron) MOP-AMO, user-controlled (“interactive”) image analyzer. One of the parameters measurable by means of the MOP-AMO, is that of form factor (FF).2 The FF of a structure is a measurement whose value is given by the simple formula:

\[ FF = \frac{4 \pi A}{P^2} \]

A = area of structure.
\[ P = \text{perimeter of structure.} \]

For a perfect circle, FF = 1·0. Deviations from a circle, whether elliptical or, for example, dendritic or spindle-shaped, result in an FF value of less than 1·0. Highly complex dendritic structures have values approaching 0·2 or even 0·1. Therefore, FF appears to be a useful means of determining the degree of roundness of structures such as ANAE-positive cells.

Material and methods

LYMPH NODES

Forty lymph nodes were examined, from the same number of patients. These were histologically diagnosed and typed according to the Kiel classification3 prior to measurement of the FF of ANAE-positive cells. All measurements were performed “blind,” the specimens being coded by numbers. There were six centrocytic, nine centrocytic-centroblastic, two centroblastic, nine immunoblastic, two lymphoblastic and two unclassifiable high-grade malignancy lymphomas. Ten ”reactive” lymph nodes were also examined, showing follicular hyperplasia with or without sinus histiocytosis (Table).

FIXATION AND STAINING

These were performed as described before4; lymph nodes were collected immediately after surgical removal, cut into 2 mm thick slices with a degreased

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Specimens examined

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>No of specimens</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reactive lymph nodes</td>
<td>10</td>
</tr>
<tr>
<td>Centrocytic lymphoma</td>
<td>6</td>
</tr>
<tr>
<td>Centrocytic-centroblastic lymphoma</td>
<td>9</td>
</tr>
<tr>
<td>Centroblastic lymphoma</td>
<td>2</td>
</tr>
<tr>
<td>Immunoblastic lymphoma</td>
<td>9</td>
</tr>
<tr>
<td>Lymphoblastic lymphoma</td>
<td>2</td>
</tr>
<tr>
<td>Unclassifiable high-grade lymphoma</td>
<td>2</td>
</tr>
</tbody>
</table>

drazor blade and fixed in cold (4°C) formol-calcium for 24 h. The slices were then transferred to gum-sucrose solution for a further 24 h, then sectioned in a cryostat at 6 μm thickness and stained for ANAE by the simultaneous coupling azo-dye technique. Slices were also fixed in 10% formol saline at room temperature and dehydrated and embedded in paraffin wax. Sections were cut at 2–3 μm and stained by Harris's haematoxylin and eosin for diagnostic histology.

MEASUREMENT OF FORM FACTOR

The use of the MOP-AMO₃ has previously been described in detail by the authors. The microprocessor was programmed to measure FF in steps of 0.05 units, from 0.05 to 1.0 and to give the data as a linear histogram, with their mean value. One hundred ANAE-positive cells' images were outlined (under a × 100 oil immersion objective) using the graphic tablet and sensitive pen. Fields were selected randomly and all ANAE-positive cells in each field were outlined (excluding lymphocytes bearing punctate enzyme activity, which include some T lymphocytes). In addition to linear histograms, the data for each specimen were plotted as cumulative percentages on logarithmic probability plot paper. The latter technique is useful as a means of demonstrating different populations of cells.

Results

Cell outlines were clearly defined by the ANAE method and their images were easily and accurately outlined on the graphic tablet using the × 100 oil immersion objective lens. Typical FF values for cells of differing shape are shown in Figs. 1 and 2; clearly these have values which correspond to their roundness or degree of dendritic shape.

Figure 3 shows the mean FF values for the 15 high-grade and 15 low-grade lymphomas; the 10 reactive lymph nodes' values are also shown. The mean values for FF in high-grade lesions tend to be in the range of 0.8-0.9; this is much greater than the values for low-grade lymphomas, whose FF values are usually 0.5 (one having a value of 0.4). The ANAE-positive cells in reactive lymph nodes also have an FF of 0.5 (one being 0.4). Thus the mean FF for low-grade lymphomas is very similar to that of reactive nodes.

Figure 4 shows the data plotted on cumulative probability paper. All specimens have non-linear plots and it is possible to project two lines through
The form factor of alpha-naphthyl acetate esterase-positive cells

Fig. 3  The mean values for FF of ten reactive nodes, 15 specimens of low-grade and 15 specimens of high-grade lymphoma.

Fig. 4  Cumulative probability plots for the 40 specimens measured. The FF values lie along the x-axis. Plots for high-grade lymphomas lie more to the right of the x-axis, in a cluster, separate from the large mixed cluster of values for low-grade lymphomas and reactive nodes, whose values for FF lie nearer to the mid-range.

Discussion

Form factor is readily and accurately measured by the MOP-AMO₃, but is a parameter that has been used only rarely in past studies. However, there is a recent description of the usefulness of FF measurement (with the MOP-1 system) in a study of cellular size and shape in normal and diseased skeletal muscle.²

In the field of lymphoma research, another parameter of shape has been used to distinguish early lesions of mycosis fungoides from chronic benign skin lesions. This parameter is the nuclear contour index (NCI) which is given by the relation:

\[ \text{NCI} = \frac{\text{perimeter}}{\text{area}} \]

The NCI gives values which reflect the degree of nuclear indentation (a feature of, for example, Sézary cells). The NCI of isolated lymph node cells has also been measured in specimens of mycosis fungoides and Sézary’s syndrome.¹¹¹² In a subsequent study,¹³ quantitative electron microscopy was used to measure NCI in skin biopsies from 109 patients, 77 of whom had chronic benign inflammatory disease and 16 had mycosis fungoides. A further 16 patients had infiltrates, the nature of which was uncertain. A “false-negative” rate of 50% was found for early mycosis fungoides lesions.

It appears, therefore, that measurements of shape as well as size are potentially useful in histopathology. FF is readily measurable by the MOP-AMO₃ and is well suited to measurements of shape of structures such as ANAE-positive cells in lymphoid tissue. On subjective grounds the shape of these cells varies, although cells (such as tingible-body macrophages) which appear rounded in haematoxylin and eosin-stained sections often have small cytoplasmic processes when stained for ANAE. Nonetheless, this sort of cell is morphologically quite different from the highly branching ANAE-positive macrophage (or, of course, the very weakly ANAE-positive dendritic reticulum cell). A striking “starry-sky” appearance with mainly round macrophages is seen, for example,
in Burkitt-type lymphoblastic lymphoma which is a high-grade variety of NHL. Such rounded macrophages have a value of FF approaching unity.

The results of this study show that, on the basis of FF, the ANAE-positive macrophage-type cells in reactive nodes and low-grade NHL tend to be more often of the branching type than in high-grade lymphomas, where more rounded esterase-positive cells predominate. The probability plots show that, as might be expected, all specimens possess two populations of such cells, of both branching and rounded type. The plots for high-grade lymphomas are shifted towards a FF of 1·0 to a greater extent than those for reactive specimens and low-grade lymphomas, with many more cells in the range of 0·8 to 1·0.

The significance of these findings is uncertain, but it is clear that FF values of ANAE-positive cells can usefully discriminate between high- and low-grade lymphomas, when correlated with morphological diagnosis using the Kiel classification. However, there is no discrimination between low-grade lymphomas and reactive nodes. It may be, in some instances, that rounded macrophages are in a highly "activated" state and are mounting an attack on high-grade lymphoma cells, where there is cell death and necrosis. In low-grade lymphomas, where growth is less rapid, the branching forms seen in normal and reactive nodes are predominant. It is possible that in the high-grade lymphomas there is an in-pouring of rounded, "reactive" macrophages, their numbers being greatly in excess of those present in normal or reactive lymphoid tissue. These rounded macrophages may be much less "specialised" or "differentiated" than their branching counterparts.

The ANAE technique offers a major advantage over the use of immunohistochemical methods (such as staining for muramidase or alpha-1-antitrypsin) for the study of size or shape of macrophages. This is because the frozen sections used for ANAE-staining are thicker than those used in immuno-histochemistry; thus the extent of branching processes is more often evident with the former than the latter method.

It is possible that factors such as mitotic activity of lymphoid cells adjacent to the ANAE-positive cells may affect the shape of the latter. Certainly, in two studies of reactive lymphoid follicles using metalophil staining methods, it was noticed that rounded macrophage-type cells were predominant in the lower centroblast rich part of the follicle, where mitoses are frequent. Conversely, in the upper parts of the reactive follicle, where fewer mitoses are seen and centrocytes predominate, metalophil branching macrophages and dendritic reticulum cells were observed. Thus, local microenvironmental factors may, in some unknown fashion, influence macrophage shape in lymphoid follicles.

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References

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