Immunoglobulin negative follicle centre cell lymphoma

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Summary

Immunoglobulin (Ig) could not be detected on the surface or in the cytoplasm of neoplastic cells from five cases of follicle centre cell lymphoma with centroblastic/centrocytic follicular histology when examined by immunohistology of frozen or wax embedded sections. Examination by fluorescein labelled antibodies of cells in suspensions prepared from the biopsies revealed a monotypic surface Ig positive population in one case and a surface or cytoplasmic Ig k:λ light chain imbalance in a further two cases consistent with neoplastic B cell involvement: in all cases the proportion of cells failing to express Ig or T cell markers ranged from 24 to 75%. The monoclonal antibodies B1 (Pan B cell), FMC4 (HLA class II) and J5 (cALL antigen) stained the majority of cells in suspension with residual cells stained with UCHT1 or OKT11 (T cell monoclonal antibodies). In frozen sections, neoplastic follicular cells did not stain with UCHT1. However, in the one case tested these cells stained with the antibodies B1 and FMC4. In paraffin sections J chain could be demonstrated in the cytoplasm of three out of five cases. Cells from four cases were cultured \textit{in vitro} for Ig production: two failed to produce Ig and monotypic light chains were the sole Ig product of the remaining two cases.

The failure to express Ig by the majority of the neoplastic cells from the cases described in this report is at variance with the follicular histology of these neoplasms. Mechanisms responsible for this failure are discussed with reference to current models of B cell differentiation.

Normal lymphocyte development is accompanied by sequential changes in cellular enzyme content and surface antigen expression. In the B lymphocytic lineage the best characterized of these are changes in immunoglobulin (Ig) synthesis, expression and secretion. The earliest event which distinguishes B cells is the rearrangement of Ig heavy chain (HC) genes and subsequent expression of cytoplasmic μ (\textit{cy}) chains (Raff \textit{et al.}, 1976; Burrows \textit{et al.}, 1979). This is followed by rearrangement of Ig light chain (LC) genes and the expression of surface membrane Ig (sIg), initially sIg M followed by sIg D (Osmand & Nossal, 1974). Further development is usually accompanied by heavy chain class switching and an increase in Ig secretion, both of these effects being antigen and/or factor dependent (Melchers \textit{et al.}, 1982).

Current classifications of non-Hodgkin's lymphoma (NHL) relate the neoplastic cell type to simplified schemes of normal lymphocyte maturation. The Kiel classification of B cell NHL (Gerard-Marchant \textit{et al.}, 1974) distinguishes two predominant cell types in lymphoid follicles, large and small cleaved centrocytes (cc) and large and small non-cleaved centroblasts (cb) The neoplastic analogues of these cell types are found as mixtures forming neoplastic follicles or in diffuse sheets effacing the normal architecture of the lymph node.

The diagnosis and classification of NHL has been facilitated by the demonstration of cell markers in sections and on isolated cells prepared from biopsy tissue. One of the most valuable markers for neoplastic lymphocytes of the B cell lineage is the demonstration of monotypic sIg and/or clg, although many monoclonal antibodies to non Ig determinants have also proved to be of value. The assay of Ig production by cells in culture can provide additional information where the sIg isotype is either difficult to assess or absent (Hannam–Harris \textit{et al.}, 1980).

In this paper we report the findings in five cases of follicular cb/cc NHL, in which the majority of the neoplastic cells failed to express sIg clg or produce Ig \textit{in vitro}.

Materials and methods

Biopsies

Lymph node biopsy material was collected fresh and divided for routine histology, frozen sections and preparation of cell suspensions as described by Stevenson \textit{et al.} (1983).

Paraffin sections

Sections of routine formalin fixed paraffin embedded material were deparaffinized in xylol and taken to alcohol. Endogeneous peroxidase activity was inhibited with 0.5% hydrogen peroxide in

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Received 13 May 1984; accepted 19 September 1984.

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methanol and the sections were washed in water and treated for 15 min with a solution of 0.1% trypsin in 0.1% calcium chloride pH 7.8 (Mepham et al., 1979). The sections were washed with agitation in cold distilled water and further rinsed in tris buffered saline (TBS) pH 7.8, followed by a 30 min incubation with rabbit antiserum to immunoglobulin determinants or J chain (see below). Sections were given a further three washes in TBS before incubation with swine anti-rabbit antibody followed by a thirty minute incubation with rabbit peroxidase-anti-peroxidase (PAP, Dakopatts) complexes at appropriate dilutions. Specific staining was demonstrated by the development of brown coloration on incubation with diaminobenzidine (DAB). The sections were counterstained with Mayer's haematoxylin and differentiated in 1% acid alcohol. Appropriate controls were performed for each layer of the immunostaining technique and for the specificity of the antibody.

Frozen sections

Fresh biopsy material was snap frozen and stored in liquid nitrogen. Six μm cryostat sections were air dried for 30 min at room temperature and stored over silica gel at −20°C in an airtight container until stained. Frozen sections were allowed to reach room temperature and fixed in dry acetone for 20 min. The sections were then immediately transferred to TBS, drained and incubated for 30 min with rabbit antiserum to human Ig or with monoclonal antibodies directed against other cell surface determinants (see below). This was followed with 3 washes in TBS and incubation with peroxidase conjugated rabbit anti-mouse Ig or with the two stages of PAP reagent as appropriate. After a further 3 washes in PBS sections were incubated with DAB, and counterstained in Harris's haematoxylin (Stein et al., 1980).

Cell suspensions

Fresh biopsy material was minced through sterile wire mesh and a cell suspension prepared by density gradient centrifugation over ficoll-triosil as described previously (Payne et al., 1977). Cells collected at the interface were washed 3 times in PBS. In all cases viability was >90% by trypan blue exclusion. Cells were incubated for 30 min at 37°C and washed once in minimal essential medium (MEM, Flow Laboratories) containing 10% foetal calf serum (FCS, Sera Laboratories), also at 37°C, to remove cytophilic antibody. The cell suspensions were investigated for T and B markers and Ig synthesis in culture.

The rosette test for the indentification of T cells with receptors for sheep erythrocytes (E) has been described previously (Payne et al., 1977). Surface Ig was characterized by staining cell suspensions with fluorescein conjugated rabbit antiser to Ig heavy and light chain determinants. Mouse monoclonal antibodies direct against T and B cell determinants were used when sufficient cells were recovered. These antibodies were used in an indirect fluorescent antibody assay with fluorescein conjugated rabbit antibody to mouse Ig as second antibody. Details of antibodies are given separately. Cytospin preparations fixed in methanol and washed with PBS were stained with fluorescein conjugated antiser for the detection of clg heavy or light chains. Appropriate controls were included in each batch of stains. The fluorescein labelled preparations were examined using a Leitz labolux 12 microscope fitted with an HBO 50w mercury vapour floempak fluorescence vertical illuminator.

Diagnostic antibodies

Polyclonal antibodies used in this study were: fluorescein conjugated and unconjugated rabbit antibodies to Ig heavy and light chains (κ, λ, γ, μ, or σ chain specific, Dakopatts), rabbit antibody to J chain (Dakopatts), horseradish peroxidase and fluorescein conjugated rabbit antibody to mouse Ig (Dakopatts) and swine antibody to rabbit Ig (Dakopatts).

Mouse monoclonal antibodies to T cells (Pan T markers: UCHT1, (Beverley & Callard, 1981) University College Hospital, London and OKT11 (Verbi et al., 1982) Orthodiagnostics), B cells (Pan B marker, B1 (Stashenko et al., 1980) Orthodiagnostics and a marker for a B cell subset FMC7, (Catovsky et al., 1981) Sera Laboratories), HLA-DR (FMC4, (Beckman et al., 1980) Sera Laboratories), the CR1 (C3b) receptor, (E11, which delineates dendritic reticulum cells, macrophages and B cell subsets (E11, (Hogg et al., 1984)) and to the common acute lymphoblastic (cALL) antigen (J5 (Ritz et al., 1980) Coulter Electronics). The mouse monoclonal antibody, RF-A1, (gift from Professor G. Janossy) reactive to T cells and a B cell subset was also used in this study (Caligaris-Cappis et al., 1982; Martin et al., 1981).

Cell culture

Cells were resuspended at 2 × 10^7 ml^-1 in Eagle's MEM (Flow Laboratories) containing 10% heat inactivated FCS, 1% non-essential amino acids, 2mM L-glutamine and 100IU ml^-1 of benzylpenicillin and streptomycin. The cells were incubated at 37°C with gentle swirling for 6 h. Aliquots were removed at 0, 3 and 6 h and viable cell numbers assessed by trypan blue exclusion. The viable cell numbers remained within 2% of the starting value throughout the culture period. The cells were
pelleted by centrifugation at 250 g for 10 min and the supernatants retained for determination of Ig production by enzyme linked immunosorbent assay (ELISA) or radioimmunoassay (RIA).

Radioimmunoassay

The RIA system has been previously described by one of us (Stevenson et al., 1980, 1981). Briefly sheep antibodies to human Fd γ, μ, δ, Fabγκ or Fabγλ were prepared and characterized as described previously and were coupled to Sephadex G-25 superfine beads (Pharmacia, Sweden) and used as solid phase antibodies. Radiolabelled Fab γ, Fab μ, Fab δ and κ or λ light chains were used as antigens. All were labelled with I125 (Amersham) by the lactoperoxidase technique (Morrison et al., 1971). Solid phase antibodies were incubated for 48 h at room temperature with labelled antigen and unknown or standards in the presence of 0.5% nonidet P40 (BDH) 1 mg ml⁻¹ bovine haemoglobin (BDH) and 20 μg ml⁻¹ soya bean trypsin inhibitor (BDH). After washing, counts associated with the solid phase were determined on an LKB rackgamma counter. These assays were sensitive to the respective immunoglobulin components in the range 1–200 ng ml⁻¹.

Enzyme linked immunoabsorbent assay

The method is adapted from Engvall & Perlman (1972). A triple layer sandwich technique was used. Sheep antibodies to human Ig κ, λ, γ, δ and µ chains or Fd δ were prepared and characterized (Stevenson et al., 1980). Antibodies diluted in Na₂CO₃/NaHCO₃ buffer (pH 9.6) were coated onto a 96 well flat bottomed, Nunc Immunoplate (Gibco Europe Ltd, Uxbridge) by incubation for 60 min at 37°C followed by 16 h at 4°C. Uncoated plastic was blocked by incubation with 1% BSA in PBS for 60 min at 37°C. Plates were washed 4 times with 0.1% Tween 20 (Koch Light Laboratories) in PBS. Cell culture supernatants were added to individual wells and the plate was incubated for 90 min at 37°C. The plates were again washed 4 times before incubation for 90 min at 37°C with horseradish peroxidase (HRP) conjugated antibody of similar specificity to the coating antibody. (HRP rabbit anti-human κ and λ chains, Dakopatts; HRP goat anti-human γ, μ, α chains, Sigma) and HRP sheep anti-human Fd δ, (Stevenson et al., 1980). All conjugated antibodies not raised in sheep were absorbed with normal sheep IgG prior to use and were negative in control incubations. After a further four washes colour was developed by the addition of freshly prepared o-phenylenediamine substrate in a phosphate: citric acid buffer (pH 5.0), catalysed by the addition of H₂O₂. The reaction was stopped by the addition of 2.5 M sulphuric acid and the absorbance at 490 nm determined using a Microelisa Auto Reader (Dynatech MR580).

These assays were sensitive to Ig products in the range 1–25 ng ml⁻¹ and were performed in triplicate with positive and negative standards on each plate. For each assay system antibodies were titrated to give optimum sensitivity and these optimal values were always used subsequently. Each new batch of antibody was re-titrated.

Serum and urine Ig assay

Serum and urine were assayed by standard methods for Ig levels, paraproteins and monoclonal urinary light chains by routine methods previously described (Stevenson et al., 1983).

Results

Marker studies

The results of marker studies carried out on both frozen and paraffin embedded biopsies are given in Table I. In frozen sections the neoplastic cells failed to stain for Ig or with the T cell monoclonal antibody (UCHT1). Cells staining with UCHT1 were scattered within neoplastic follicles and as a rim around the follicle (Figure 1a). Polytigic Ig staining cells were found in the interfollicular areas and in the mantle zone (Figures 1b and c) surrounding the follicles. The antibody E11 stained dendritic reticulum cells within the follicles but did not stain the neoplastic cells (Figure 1d). In J K the sections were stained with B1 and FMC4; these antibodies stained both neoplastic cells within the follicles and normal B cells in the interfollicular areas.

No cytoplasmic Ig could be detected within neoplastic cells embedded in paraffin sections, however cytoplasmic J chain was detectable in 3 cases (LV, AT, SH) (Table I).

The results of marker studies carried out on cells dispersed from biopsy material are given in Table II. The slg staining of AT was consistent with a predominant population of monotypic slg κ staining cells. In two other cases the number of cells staining for slg κ and λ differed from the normal ratio of 2:1, with a predominance of slg λ cells (SH and SA). In SH a predominant population of cytoplasmic Ig M λ positive cells was detectable. T cells detectable by monoclonal antibody or E rosetting were present in all preparations. The total percentage of cells detectable by slg κ and λ staining and T cell monoclonal antibody markers was less than 100% in all cases examined and ranged from 76% (SA) to 25% (JK). In LV, 87% of the cells could be accounted for by E rosetting and slg κ and λ staining.
Table I  Immunohistochemical investigation of frozen and paraffin embedded material

| Patient | Frozen sectiona | Paraffin sectionsb |
|---------|-----------------|--------------------|
|         | Surface Ig      | UCH-T1 E11c        | Cytoplasmic Igb | Cytoplasmic J chain |
| JK      | Neoplastic follicles |                | neg             | neg |
|         | Interfollicular areas | (+)            | +               | +   |
|         |                   | M,D             | k,λ             |     |
| LV      | Neoplastic follicles |                | neg             | +   |
|         | Interfollicular areas | (+)            | n.d.            | +   |
|         |                   | M,D             | k,λ             |     |
| AT      | Neoplastic follicles |                | neg             | +   |
|         | Interfollicular areas | (+)            | +               | +   |
|         |                   | M,D             | k,λ             |     |
| SH      | Neoplastic follicles |                | neg             | +   |
|         | Interfollicular areas | (+)            | +               | +   |
|         |                   | M,D             | k,λ             |     |
| SA      | Neoplastic follicles |                | neg             | +   |
|         | Interfollicular areas | (+)            | +               | +   |
|         |                   | M,D             | k,λ             |     |

*aFor details see methods.
bCytoplasmic staining was negative for all classes of Ig heavy and light chains in neoplastic follicles and polyclonal in interfollicular areas.
c'dendritic cell staining.
neg=no staining of cells seen.
n.d.=not done.
(+)=scattered positive cells within follicles see Figure 1.

Table II  Markers on dispersed cells

| % Cells expressing | Surface Ig type | Cytoplasmic Ig and light chains | Surface antigens detected by monoclonal antibodies |
|-------------------|----------------|-------------------------------|-----------------------------------------------|
| Patient           | G  A  M  D  k | λ                             | B1   FMC7   FMC4   J5   RFAI   T*   E rosettes |
| JK                | 0  0  8  3  10 | 5  2 (MGl)                   | 85   5     95    72   40   10*   4 |
| SH                | 3  0  20  4  13 | 15  12 (10λ, 2k, 12M)        | 63   5     50    67   55   35*   30 |
| LV                | 4  0  11  6  6  | 1  0                         | nd   nd    nd    nd   nd   80 |
| AT                | 17  7  30  21  43| 6  0                         | 69   3     65    61   91   30*   18 |
| SA                | 4  1  27  2  12 | 20  1 (Mλ)                   | 79   18    65    53   82   44*   38 |

*bResults from ten cases of reactive lymphadenopathy: mean (range).

The monoclonal antibodies B1, J5 and FMC4 detected a predominant population of cells in the four cases in which they were used (JK, SH, AT, SA) (Table II). The total percentage of cells detected by B1 and the T cell markers accounted for more than 95% of the cells in suspension from all the cases tested, with a small overlap (23%) between the numbers of cells detected by B1 and OKT11 in cell suspensions from SA. The proportion of cells stained by the monoclonal antibody FMC4 ranged from 50% (SH) to 95% (JK), similarly J5 stained 61% (AT) to 72% (JK) of the cells in suspension. More than 82% of the cells in JK, SH, AT and SA could be accounted for by
Figure 1 Immunoperoxidase staining of frozen sections from case SA. The staining patterns are typical of the other cases in this study. (a) Section stained with UCH T1 showing a rim of T cells around a neoplastic follicle with scattered T cells within the follicle (Immunoperoxidase × 300). (b) Section stained to show IgD. A narrow mantle of IgD positive cells is visible around the neoplastic follicle (Immunoperoxidase × 120). (c) Section stained to show IgM. A narrow mantle of IgM positive cells, similar to IgD, is visible around the neoplastic follicle (Immunoperoxidase × 120). (d) Section stained with E11. The neoplastic follicles contain a network of positively staining dendritic cells (Immunoperoxidase × 120).
staining with FMC4 or J5 and the T cell monoclonal antibodies (Table II). Material was not available from LV for monoclonal antibody studies.

The results of cell suspensions, prepared from reactive lymph nodes, stained for sIg k and l and the T cell monoclonal antibody, UCHT1, are given in Table II for comparison.

**Ig production**

Cells from four of the five cases were available for cell culture and assessment of Ig production (Table III). Cell culture supernatants were assessed for k and l light chains and for \( \gamma, \alpha, \mu \) and \( \delta \) heavy chains as described in the methods. In two cases (SH, AT) monotypic Ig light chain secretion was found with no detectable whole Ig secretion. No production of Ig was observed by cells from JK or SA. Results from five typical cases of cb/cc follicular lymphoma and from nine reactive lymph nodes are given for comparison (Table III).

**Table III** Ig production by neoplastic cells (ng per 2 x 10⁷ cells at 6h)

| Patient | Heavy chain | Light chain k | Light chain l |
|---------|-------------|---------------|---------------|
| JK      | 0           | 0             | 0             |
| SH      | 0           | 0             | 25            |
| AT      | 0           | 19            | 0             |
| SA      | 0           | 0             | 0             |

*Results from five typical cases of cb/cc follicular lymphoma: mean (range). Monotypic Ig light chain secretion was found in all cases (3k:2l).

bResults from nine reactive lymph nodes: mean (range). Polyclonal Ig light chain secretion was found in all cases; heavy chain class was not determined (nd).

**Discussion**

Cells of the follicle centre represent antigenically stimulated B lymphocytes at an intermediate stage of differentiation thought to be actively expressing and secreting Ig (Wakefield & Thorbecke, 1968; Grobler et al., 1974; Cooper et al., 1973). Studies from this and other laboratories using similar immunoperoxidase techniques for the detection of Ig in tissue sections have shown that follicle centre cells from normal reactive and neoplastic tonsils, lymph nodes and spleen are either surface and/or cytoplasmic Ig positive (Isaacson et al., 1980; Stein et al., 1980, 1982). These data contrast with a recent report from Hsu et al. (1983) who failed to detect Ig expression by the majority of follicle centre cells in frozen sections from normal reactive tissue. Clearly methods need to be compared and standardized for the detection of Ig in sections. Recently Cordell et al. (1984) have drawn attention to differences in sensitivity between techniques.

The large majority of cells in suspensions prepared from reactive lymph nodes can be labelled with the T cell monoclonal antibody, UCHT1, and antibodies to Ig k and l chains (Table II). Data from this and other laboratories have shown that the majority of neoplastic cells in suspensions prepared from cb/cc follicular NHL express sIg and occasionally clg predominantly of IgM isotype (Payne et al., 1977; Leech et al., 1975; Godal & Funderund, 1982).

Cell suspensions, prepared from biopsy tissue, offer optimal conditions for the detection of sIg, but may selectively enrich for normal or neoplastic subpopulations. Consequently results obtained by tissue section staining techniques or by staining of dispersed cells in suspension will occasionally show discrepancy. This discrepancy was clearly observed in one case in this study (AT) where monotypic sIg was demonstrable on cells in suspension but not in tissue section. A further two cases (SA & SH) also negative for sIg and clg in sections exhibited sIg and/or clg \( \kappa: \lambda \) light chain imbalance when cell suspensions were examined suggestive of IgM \( \lambda \) neoplastic B cell involvement. Despite these findings, sIg was not detected on 24–75% of the neoplastic cells, which failed to stain with monoclonal antibodies to T cells, in JK, SH, AT and SA.

In two cases (JK & LV) cell suspension analysis and examination of tissue sections both failed to demonstrate an Ig phenotype. E rosetting cells were recovered in high numbers in suspensions prepared from LV. In this particular case it was not possible to decide whether we had failed to recover the neoplastic population or if the E positive cells represented a neoplastic B cell population with

**Serum and urine**

Serum IgA, G, and M levels for all cases were within normal range and there was no evidence for a paraprotein. A search for monoclonal urinary Ig light chain in concentrated urine revealed a trace \( \kappa \) light chain in JK. The remaining four patients did not have detectable urinary light chains. None of these patients had proteinuria.
affinity for sheep erythrocytes (Prieur & Brouet, 1974).

The B cell nature of the neoplastic cells was sought by using antibody probes for J chain and B cell related antigens. J chain was demonstrable in the cytoplasm of neoplastic cells in paraffin embedded sections of LV, AT and SA, but was not found in cells from JK & SH. The monoclonal antibody B1, which recognises a determinant widely expressed throughout the B cell lineage (Stasheniko et al., 1980) and the monoclonal antibody FCM4, which is specific for class II antigens (Beckman et al., 1980), stained the majority of cells in frozen sections from JK. The other cases were not tested. In our experience both these antibodies reliably stain normal and neoplastic B cells in lymphomas of follicular histology. B1 stained the majority of cells in suspensions prepared from JK, SH, AT and SA, and together with the T cell monoclonal antibody accounted for more than 95% of the cells. A small population of cells expressing antigen determinants recognised by B1 and OKT11 was observed in SA. This population of cells is analogous to previously reported neoplastic B cell populations from a small number of malignant lymphomas, which express the T cell antigen recognised by OKT11, (Aisenberg et al., 1981). FMC4 stained the majority of cells in suspension from JK, SH, AT and SA and together with the T cell monoclonal antibody accounted for more than 85% of the cells. Neoplastic cell populations from all five cases expressed one or more determinants associated with B cells, consistent with these tumours being of B cell origin.

The monoclonal antibody J5, which reacts with cALL cells and some B and T lymphomas including neoplastic and normal B cells of follicle centres (Ritz et al., 1980, 1981; Habeshaw et al., 1983; Stein et al., 1982), stained 53–72% of cells in suspensions from JK, SH, AT and SA. This antibody does not appear to react with T cells in follicular lymphomas (Stein et al., 1982) and this finding is a further demonstration of the similarity of the neoplasms described in this report to other cases of follicular lymphoma.

The majority of cells in suspension (82–97%) from JK, SH, AT and SA were negative for the B cell subset antibody FMC 7, which reacts preferentially with neoplastic cells in prolymphocytic leukaemia and variably with neoplastic cells in lymphoma (Catovsky et al., 1981; Collins et al., 1983). The monoclonal antibody RF-A1 which recognises T cells and a shared determinant present on a normal B cell subset and neoplastic B cells of CLL (Martin et al., 1981) reacted with more cells than expected when tested with cell suspensions prepared from the four cases.

A similar observation for cb/cc lymphoma of follicular histology has been reported by Habeshaw et al. (1983).

All of the lymphomas in this study showed histology typical of follicle centre cell lymphoma, cb/cc follicular. Lymphomas with this histology in which the neoplastic cells do not express slg are rare and represent less than 3% of follicular tumours in our series. In a recent immunohistochemical study of NHL Tubbs et al. (1984) found one Ig negative lymphoma with a follicular histology in a group of 53 such cases examined, a similar incidence to that in our own series.

Human “null” lymphocytes defined by a lack of T lymphocyte markers and Ig expression have been identified among mononuclear cells from normal peripheral blood, lymph nodes and spleen. A brief review of the earlier literature relating to this subject can be found in Haegert & Coombs (1979).

In this article and a subsequent paper (Haegert, 1981) it is claimed that the majority of “null” lymphocytes are of B cell lineage by virtue of surface Ig demonstrable by the mixed antiglobulin technique. Our earlier studies (Payne et al., 1977) and the data relating to normal lymph node populations presented in this paper support this conclusion. Nevertheless whether all “null” cells detected in these tissues bear slg and are therefore of B cell lineage remains to be resolved. Technical improvements such as those described by Haegert (1981) and Cordell et al. (1984) together with the application of T and B cell specific monoclonal antibodies will help to resolve this problem.

Neoplastic “null” cells occur in ALL, CLL and NHL. “Null” cells in CLL can be shown to be capable of synthesizing Ig in vitro (Gordon et al., 1978) thus providing evidence for their B cell origin (Gordon, 1984). We have also shown that isolated neoplastic cells from cases of follicular NHL secrete whole Ig and/or free Ig light chains when cultured in vitro (Hannam-Harris et al., 1982; Stevenson et al., 1984). Neoplastic cells from four cases in this study were investigated for their ability to secrete Ig. Neoplastic cells from two of these, AT and SH secreted monotypic Ig light chain but did not secrete whole Ig in vitro. This pattern of Ig secretion is unusual in follicular NHL and has been associated with immature normal and neoplastic B lymphocyte populations in CLL (Hannam-Harris et al., 1980; Gordon et al., 1983).

In ALL the majority of “null” cells can be shown to be of B cell lineage by the presence of cytoplasmic µ chain (Vogler et al., 1978) or by gene rearrangement (Korsmeyer et al., 1981). Studies of Ig gene rearrangement have been helpful in determining the monoclonality of lymphoid tumours, including those of follicular histology,
where this cannot be established by Ig phenotypic analysis (Arnold et al., 1983). Similarly this technique has been used to establish the monoclonality of lymphomas in transplant patients (Cleary et al., 1984). A JH probe has been used to demonstrate Ig gene rearrangement in the neoplastic cells isolated from one case included in this study (SA), strongly supporting a neoplastic B cell origin, (Dr Nigel O’Connor, personal communication).

We have shown by a variety of techniques that the neoplastic cells from the lymphomas described in this report are of B cell origin. However we are not able to decide on the basis of the present investigations, whether these cells are at an early stage of B cell maturation. These tumours may represent a separate developmental lineage of B lymphocytes within follicle centres or alternatively asynchrony between Ig expression and homing to sites of B cell development, analogous to that reported in myeloid neoplasms for the development of cytoplasmic enzymes and surface antigen expression (Scott et al., 1982). Alternatively failure to express Ig by the majority of the neoplastic cells may arise as a consequence of an acquired defect in production by mechanisms similar to that observed in myeloma (Williams et al., 1966), by aberrant Ig gene rearrangement (McIntosh et al., 1983) or by factors unknown.

This report serves to demonstrate that a coordinated approach is required for the investigation of B cell neoplasms. If routine immunohistology and immunological analysis of cell suspensions fails to demonstrate the cell of origin, functional studies as described in this report may be useful. However, in a small number of cases where the B cell origin remains undetermined the ultimate investigation would appear to be Ig gene rearrangement.

We acknowledge the Leukaemia Research Fund for financial support.

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