Clonality of CD3 negative large granular lymphocyte proliferations determined by PCR based X–inactivation studies

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Abstract

Aims—To examine persistent CD3– large granular lymphocytosis (LGL) cases for clonality, both by lineage specific (T cell receptor) and lineage independent (X–inactivation) molecular methods; and to find out whether X–inactivation studies are more appropriate than gene rearrangement studies for this subset of LGL disorders.

Methods—Patients were selected who had LGL of more than six months’ duration and identified as CD3– by immunophenotyping. T cell receptor studies and, where possible, X–inactivation studies of the phosphoglycerate kinase (PGK) gene were carried out. Analysis of subpopulations was carried out on cases heterozygous for PGK by the use of a polymerase chain reaction (PCR) method for X–inactivation.

Results—Of 17 CD3– LGL cases studied, all were found to be germline for β, γ, and δ T cell receptor studies, and immunoglobulin heavy chain genes. However, six of these were analysed by X–inactivation of the PGK gene and two cases gave clonal band patterns but only within the CD3– subpopulation.

Conclusions—Clonal analysis by the lineage independent method of X–inactivation allows clonal expansion undetected by T and B cell specific markers to be identified. It is therefore a more appropriate method for the analysis of CD3– LGL. This has implications for diagnosis in CD3– LGL disorders.

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Abnormally increased numbers of circulating large granular lymphocytes (LGL) are associated with a diverse range of disorders, including both reactive and malignant conditions.1 Transient (acute reactive) LGL expansions occur primarily in response to viral infections and are often characterised by increases in both CD3+ T cell and CD3– natural killer cell LGL.2 In contrast, persistent lymphocytes with granular morphology are generally due to an abnormal expansion of one LGL type.3 Many persistent LGL expansions with clinical and haematological features of “leukaemia” have been reported, and most cases designated as such by the Morphology–Immunology–Chromosome (MIC) Cooperative Study Group are of the CD3+ subtype.4 However, a recent survey by the Yorkshire Leukaemia Group (YLG) indicates that about 25% of persistent LGL expansions are in fact of CD3– type, and a proportion of these are also thought to be clonal in nature.5,6

Monoclonality is a fundamental characteristic of haemopoietic malignancies, and investigations to establish the clonal nature of abnormal proliferations are therefore important in differentiating reactive from malignant states. The most widely used method for assessing clonality of lymphoid disorders involves molecular analysis of immunoglobulin and T cell receptor (TCR) gene rearrangements.7 TCR studies are informative for most cases of CD3+ malignancy,8 but TCR studies of CD3– proliferations have been of little value.9

An alternative method of clonal analysis is provided by the study of differential methylation in certain X chromosome genes in females.10 The inactivation of one X chromosome with concomitant methylation of the 5′ end of genes, such as PGK and hypoxanthine phosphoribosyl transferase (HPRT) during early embryogenesis, provides a stably inherited genetic marker. A normal population of cells in an adult woman will comprise a random mixture of paternally and maternally derived X–inactivated cells. A clonal population of cells, having originated from a single cell, will feature inactivation of the same X chromosome in each cell. The PGK gene can be used for clonal analysis by a polymerase chain reaction (PCR) method where amplification of DNA around the BstXI polymorphic restriction site and a methylation site permits analysis of heterozygous females.11 The BstXI site is polymorphic in about 33% of females.12

To examine the nature of persistent increases in CD3– LGL/natural killer cells, we analysed X–inactivation at the PGK locus in six informative women from 17 female CD3– cases identified in the YLG survey.5

Methods
To date the YLG survey has identified 97 people with persistently increased LGL and/or lymphocytes expressing natural killer cell associated membrane determinants (defined as lasting more than six months).1 Of these, 23 were found with a primary increase in CD3– natural killer cell associated + cells. Seventeen were women and were screened for heterozygosity at the polymorphic BstXI site in the PGK gene (fig 1). LGL lymphocytes were identified by conventional morphologi-
cal examination and benzylcarbonyl-L-lysine thiobenzyll (BLT) esterase cytochemistry. Lymphocyte counts and immunophenotypic data are shown in table 1. Age and clinical details of the six heterozygous cases identified by this screen are shown in table 2.

LYMPHOCYTE FRACTIONATION AND IMMUNOMAGNETIC MODIFICATION PROCEDURES

Mononuclear cells were initially fractionated from 30 ml of EDTA anticoagulated venous blood by density sedimentation with Lymphoprep (Nycomed, UK). Cells isolated from the interface were washed twice in phosphate buffered saline and azide (0.01%) before immunophenotyping and immunomagnetic depletion procedures.

Washed mononuclear cells were pelleted and mixed with the following combination of monoclonal antibodies at saturating concentrations: 63D3 (CD14); HD37 (CD19); and T3 (CD3). After a 30 minute incubation at 4°C, with occasional mixing, the cells were washed free of excess monoclonal antibodies and resuspended in 0.5 ml Hanks' balanced salt solution (HBSS), supplemented with 1% bovine serum albumin (BSA). Washed sheep anti-mouse immunoglobulin coated M450 magnetic particles (Dynabeads; Dynal, UK) were added (50 μl). The leucocyte/Dynabead mixture was incubated for 30 minutes at room temperature with gentle agitation, then resuspended to 1 ml in 1% BSA/HBSS and placed on a magnetic separator. The unbound cells were removed and tested for residual antibody coated cells by indirect immunorosetting.13 The depletion procedure was repeated with further 50 μl aliquots of Dynabeads until the number of residual mononuclear antibody cells

### Table 1: Summary of haematological and immunophenotypic features of six CD3- natural killer cell associated + persistent expansions examined in this study

<table>
<thead>
<tr>
<th>Case No</th>
<th>Lymphocyte count x 10⁹</th>
<th>CD2+CD3-</th>
<th>Primary natural killer cell associated abnormality</th>
<th>Natural killer cell associated phenotype**</th>
</tr>
</thead>
<tbody>
<tr>
<td>AB</td>
<td>1/7</td>
<td>32/61</td>
<td>0-73</td>
<td>CD11b+CD16+CD56+CD57+</td>
</tr>
<tr>
<td>ZB</td>
<td>6/3</td>
<td>06/39</td>
<td>0-95</td>
<td>CD11b+CD16+CD56+CD57v</td>
</tr>
<tr>
<td>WD</td>
<td>4/7</td>
<td>20/52</td>
<td>1-41</td>
<td>CD11b+CD16+CD56vCD57v</td>
</tr>
<tr>
<td>WF</td>
<td>4/4</td>
<td>36/69</td>
<td>0-97</td>
<td>CD11b+CD16+CD56vCD57v</td>
</tr>
<tr>
<td>LG</td>
<td>5/9</td>
<td>30/75</td>
<td>2-42</td>
<td>CD11b+CD16+CD56vCD57v</td>
</tr>
<tr>
<td>AW</td>
<td>6/0</td>
<td>42/nt</td>
<td>1-56</td>
<td>CD11b+CD16+CD56vCD57v</td>
</tr>
</tbody>
</table>

*These cases represent the six (of 17 tested) women who were shown in preliminary studies to be heterozygous for the PGK gene. † Indicates the percentages of lymphocytes with LGL morphology (Romanowsky) and positive BLT esterase staining.

### Table 2: Summary of clinical data and DNA analysis

<table>
<thead>
<tr>
<th>Case No</th>
<th>Age (µ)</th>
<th>Clinical features</th>
<th>TCR study†</th>
<th>X-inactivation‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>AB</td>
<td>1/7</td>
<td>No important medical history</td>
<td>Germline</td>
<td>Poly</td>
</tr>
<tr>
<td>ZB</td>
<td>6/3</td>
<td>No important medical history</td>
<td>Germline</td>
<td>Poly</td>
</tr>
<tr>
<td>WD</td>
<td>4/7</td>
<td>Polymyalgia rheumatic, gall stones, diverticulitis, hypertension</td>
<td>Germline</td>
<td>Poly</td>
</tr>
<tr>
<td>WF</td>
<td>4/4</td>
<td>Hypothyroidism, maturity onset diabetes mellitus, iron deficiency, menorrhagia, hypertension</td>
<td>Germline</td>
<td>Poly(s)</td>
</tr>
<tr>
<td>LG</td>
<td>5/9</td>
<td>Transient headaches—resolved without treatment, diverticulitis</td>
<td>Germline</td>
<td>Mono</td>
</tr>
<tr>
<td>AW</td>
<td>6/0</td>
<td>Clinically documented viral infection—monospot negative. Now well</td>
<td>Germline</td>
<td>Poly</td>
</tr>
</tbody>
</table>

* These cases (six of 17 tested) are heterozygous for the BrXI polymorphism in the PGK gene.
† Germline configuration of TCR gene was also found in 11 of 17 CD3- NKa + cases tested, which were not polymorphic for PGK.
‡ CD3 + denotes a composite population of cells comprising (CD3+/CD19+/CD14+) separated by immunodepletion. Poly, polyclonal; Mono, monoclonal; Poly(s), skewed X-inactivation in this patient.
was less than 2%. In practice, this rarely required more than two depletion steps. The Dynabead bound fraction comprised CD14+ (monocyte), CD19+ (B lymphocyte), and CD3+ (T lymphocyte) cells. For simplicity this fraction is referred to as CD3+ in fig 2.

**DNA EXTRACTION**

For TCR analysis extraction was performed by a standard phenol/chloroform method after 0.86M ammonium chloride lysis of red cells and overnight digestion with proteinase K. A simplified protocol was used for X-inactivation analysis following lymphocyte fractionation. Cells were resuspended in 1 ml of sodium chloride/TRIS/EDTA (pH8) (STE) buffer, lysed with 20 μl sodium dodecyl sulphate (SDS) (20% w/v) and treated with 100 μg proteinase K for two hours, then extracted once with equal volumes of phenol and chloroform, ethanol precipitated, and pelleted. The dried pellet was resuspended in 100–200 μl of TRIS/EDTA (pH7.5) (TE) buffer.

**T CELL RECEPTOR AND IMMUNOGLOBULIN HEAVY CHAIN GENE ARRANGEMENTS**

DNA (10 μg) was digested overnight with restriction enzymes HindIII and BamHI (and EcoRI when results were inconclusive). The DNA fragments were separated using 0.8% agarose gel and transferred to nylon membranes (Hybond N, Amersham International, UK) using vacuum blotting method (Pharmacia, UK). The membranes were fixed by ultraviolet light and prehybridised at 65°C in buffer solution containing 10% dextran sulphate, 3× sodium citrate and sodium chloride (SSC), 1% sodium dodecyl sulphate (SDS), and denatured salmon sperm DNA, 10 mg/ml, for two hours. Denatured 32P labelled cDNA probes were added to the mixture and hybridised overnight. The probes used in this study were: Jur-β2,15 T-γ (pH60),16 T-β (R2IXH),17 and JH (pUC19).18 The membranes were washed twice in 2× SSC and 1% SDS for 30 minutes at 65°C. Further washes were given if the background radioactivity was still high. The membranes were autoradiographed (Hyperfilm, Amersham International, UK) for three to five days.

**X–INACTIVATION CLONAL ANALYSIS BY THE PCR METHOD**

Primers A1/B3 for the 619 base pair product from the PGK gene were made on Applied Biosystems oligonucleotide synthesiser according to the sequence published elsewhere.11 Primer INT3 was also synthesised which produces a fragment of 309 base pairs enabling greater separation of BstXI digested fragments. The sequence of the 20 nucleotide oligomer INT3 is TTGGTGGTTTCTAGC-CGCATT (fig 1).

The PCR was carried out in an LEP Scientific, model Prem, thermal cycler. Temperatures were 95°C for one minute, denaturation, 56°C for one minute, annealing and 72°C for one minute extension. Amplification with primers A1 and B3 was for 25 cycles using 1.5 U Taq polymerase (Promega) per tube in a total volume of 50 μl. Primers INT3 and B3 were used by taking 1 μl of the 619 base pair A1/B3 product and a further 10 cycles of amplification carried out under the same PCR conditions. Digests of 0.5 μg DNA with 20 U HpaII enzyme (Northumbria Biologicals) were carried out in 20 μl Taq buffer (Promega) before amplification. This digest (10 μl) was used for PCR. Control DNA was not digested with HpaII before amplification. Following amplification 20 μl of products from both control and HpaII digested samples were digested overnight with 12 U BstXI enzyme (Northumbria Biologicals). Fragments were separated using 2% agarose gel and visualised with ethidium bromide (fig 2).

**Results**

**IMMUNOPHENOTYPING**

The immunological characteristics of the six patients who were informative for PGK are
shown in Table 1. Absolute lymphocyte counts were only slightly increased (range 4.7–6.3 x 10^9/l) in four of these, the remaining two being within normal limits. Absolute increases in CD2+CD3− lymphocytes (normal range, 0.15–0.49 x 10^9/l), which are primarily associated with natural killer associated cells, were found in all six cases, and multiple colour flow cytometry showed that the primary natural killer cell abnormality was associated with CD2+CD3−CD4−CD8− (n = 4) or CD2+CD3−CD4−CD8dim + (n = 2) fractions. Expression of natural killer cell associated determinants showed some individual variability but the abnormally increased CD2+CD3− cell numbers in all cases expressed all four natural killer cell associated markers examined (CD11b, CD16, CD56, and CD57) on at least 30% of the cell fraction.

**T cell receptor and immunoglobulin heavy chain gene rearrangements**

DNA from the mononuclear cell fractions of 17 patients with persistent expansions of CD3− natural killer associated cells were investigated for the presence of monoclonal TCR and immunoglobulin gene rearrangements. Figure 3 illustrates germline and rearranged results obtained by this method. All of our 17 cases were found to have germline configuration with each probe used, which is illustrated by results from cases WF and AB (fig 3). T and B cell gene rearrangement studies are carried out using a mononuclear cell fraction. Previous experience in the laboratory indicates that other cell types present in the mixture do not obscure the presence of a minor (as low as 1%) population of cells containing a single clonal rearrangement. Germline bands represent unarranged genes from most of the cells, whereas clonally rearranged cells appear as a distinct band of a different size (fig 3A; lanes 2–4).

**X-inactivation analysis of unfractionated mononuclear cells**

Initially, DNA from the mononuclear cell fractions of 17 patients with persistent CD3− lymphocytosis were screened for the presence of a BstXI polymorphism at the PGK locus. Six heterozygotes were identified and X-inactivation clonal analysis was carried out by the PCR method. None of these showed unequivocal evidence of monoclonality, although cases WF, LG, and WD (data not shown) produced highly “skewed” band patterns, with one product being significantly more intense than the other. These observations could either reflect an unusually high ratio of maternal:paternal X-inactivated or suggest the existence of a mixture of clonal and nonclonal cells. Because X-inactivation studies cannot distinguish cell type, cells not involved in the clone will interfere with the analysis. Therefore, the population of interest must be as pure as possible for meaningful analysis.

**X-inactivation analysis of immunologically fractionated cells**

To isolate the abnormally increased CD3− cell numbers, fresh samples of peripheral blood mononuclear cells from each patient were separated into CD3+ and CD3− fractions. Clonal analysis by the PCR method was then carried out on these subfractions. Results from case AB are shown in fig 2A, and show a polyclonal pattern in both CD3+ and CD3− fractions, where both bands are retained after HpaII digestion. Similarly, cases ZB and AW gave polyclonal patterns (table 2). In contrast, results from case WF in fig 2B showed a polyclonal pattern of products in the CD3+ fraction and a monoclonal pattern in the CD3− fraction (by the loss of one product from the CD3− fraction after HpaII digestion). Likewise, patient LG showed a polyclonal pattern for the CD3+ fraction and a monoclonal pattern for the CD3− fraction (table 2). Results for case WD are shown in fig 2C. This case
shows evidence of uneven (skewed) $X$-inactivation in both the CD3+ and CD3− fractions, shown by a reduction of intensity of the smaller PCR product in the $HpaiI$ digested sample compared with the control. The observation was consistent over three experiments (data not shown) and as the pattern was the same for both the CD3+ and CD3− fractions we concluded that each comprised normal polyclonal subpopulations. The skew towards one $X$-inactivation product is assumed to be consistent for constitutional DNA in this patient because of the comparison with CD3+ cells. In this case there was a slight difference in band intensity in control lanes (fig 2C; lanes 10, 12). This was seen occasionally and might have resulted from a PCR artefact due to template preference, or differences in ethidium bromide incorporation within the agarose gel. However, there was a clear difference between control lanes and $HpaiI$ digested samples (fig 2C; lanes 9, 11).

**Discussion**

Evidence for monoclonality in LGL disorders with CD3+ phenotype has been accumulating for several years by the extensive use of TCR gene rearrangement studies.\(^\text{9, 19}\) These studies, however, have served to confirm the consistent observation of germline TCR genes in LGL with CD3− phenotype. This agrees with our own finding of germline TCR genes in the mononuclear cells of 17 CD3− LGL cases, whereas in CD3+ cases rearrangements can be clearly seen (fig 3). Consequently, a T cell specific genetic marker seems to be inappropriate in the assessment of clonality in CD3− LGL cases.

As there is no specific clonal marker available for cells of natural killer cell lineage we applied the lineage independent method of $X$-inactivation analysis. A preliminary study of mononuclear DNA from CD3− LGL cases showed that polyclonal cells were present in the peripheral blood of these patients; but skewing of the bands suggested that in some cases there could be a subpopulation of clonal cells present within the mononuclear fraction. This was confirmed when fresh samples were divided into CD3+ and CD3− subgroups and analysed separately. Of six informative cases examined, two patients clearly showed a monoclonal pattern in the CD3− cell fraction. Polyclonal patterns were found for the CD3+ fraction of both of these patients, therefore effectively serving as tissue specific internal controls, showing that $X$-inactivation is not extreme within the normal cells of these individuals. Extreme skew of $X$-inactivation was, however, illustrated by another of our patients (WD) who showed evidence of a skewed $X$-inactivation pattern within both cell fractions. This highlights the need for caution when interpreting reduced band intensity as a measure of clonality by the $X$-inactivation method, particularly if constitutional DNA is not available.\(^\text{20}\)

Fortunately, CD3+ cells provided suitable constitutional DNA for this study.

The study of minor subpopulations of cells within the peripheral blood has become practicable by the development of a PCR based technique. This has the advantage of rapidity and has a reduced requirement for starting material compared with the standard Southern blotting method.\(^\text{9, 11}\) This limitation also applies to other Southern blotting based markers, such as HPRT and $M27\beta$. $M27\beta$ may not be ideal as an alternative because the test region suffers from hypermethylation in some leukemic blast cells.\(^\text{20}\)

Analysis of subpopulations makes it possible to isolate and identify the clonal origin of proliferating LGL cells and define the limits of lineage involvement. The abnormal monoclonal population in CD3− LGL seems to be restricted to the natural killer cell subpopulation. Thus the target cell is shown to be committed to the natural killer cell lineage, the remaining peripheral blood cells having a polyclonal pattern of $X$-inactivation. Other evidence for monoclonality in CD3− LGL has been found by Taniwaki et al,\(^\text{21}\) whose patients had chromosomal abnormalities, and by Kawas–Ha et al,\(^\text{22}\) who examined their cases for the presence of monoclonal Epstein–Barr virus termini and suggested the virus may have an aetiological role in some cases of LGL. In contrast, a recent report by Nash et al\(^\text{23}\) did not identify clonal CD3− populations within the seven patients studied.

The clinical relevance of a clonal population of natural killer cells is uncertain. Neither of our two patients had any other evidence of an haematological malignancy. One patient (WF) had a strong history of autoimmune disease. The association of clonality with autoimmune disease is well recognised and is seen both in the manifestation of autoantibodies and clonal proliferation of large granular lymphocytes.\(^\text{24–26}\) The other patient (LG) had no clinical problems known to be associated with clonal proliferation. The increasing use of new molecular techniques is likely to permit identification of previously unidentified minor clonal populations. However, the clinical importance of this can only be ascertained by continued clinical observation.

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