Levels of expression of CD19 and CD20 in chronic B cell leukaemias

Lia Ginaldi, Massimo De Martinis, Estella Matutes, Nahla Farahat, Ricardo Morilla, Daniel Catovsky

Abstract

Aims—To investigate whether the antigen levels of the B cell lineage markers CD19 and CD20 can distinguish between normal and neoplastic B cells or characterise distinct expression patterns among the chronic B cell leukaemias.

Methods—Peripheral blood cells from 70 patients with B cell disorders and 17 healthy donors were analysed by quantitative flow cytometry. Direct immunofluorescence staining was performed with phycoerythrin conjugated CD19 and CD20 monoclonal antibodies. Standard microbeads with different capacities to bind mouse immunoglobulins were used to convert the mean fluorescence intensity (MFI) values into number of antigen molecules/cell, expressed as antibody binding capacity (ABC).

Results—CD19 and CD20 ABC values in leukaemic B cells differed from those of normal blood B lymphocytes. The results identified distinct profiles of CD19 and CD20 expression in the various types of B cell leukaemias. In all leukaemias studied except hairy cell leukaemia (HCL), CD19 expression was significantly lower than the mean (SD) value in normal B cells (22 (7) × 10⁵ molecules/cell), as follows: chronic lymphocytic leukaemia (CLL), 13 (7) × 10⁵; B prolymphocytic leukaemia (B-PLL), 16 (9) × 10⁵; splenic lymphoma with villous lymphocytes (SLVL), 15 (11) × 10⁵; mantle cell lymphoma (MCL), 10 (7) × 10⁵. In HCL there was strong CD19 expression (38 (16) × 10⁵). In contrast, the level of expression of membrane CD20 was higher than the mean (SD) value in normal B cells (94 (16) × 10⁵ molecules/cell) in MCL (123 (51) × 10⁵); B-PLL (129 (47) × 10⁵); SLVL (167 (72) × 10⁵); and HCL (312 (110) × 10⁵); while it was significantly lower (65 (11) × 10⁵) in CLL compared with normal B cells and the other B cell leukaemias.

Conclusions—Quantitative determination of CD19 and CD20 may provide useful diagnostic information for the study of B lymphoproliferative disorders.

Keywords: quantitative flow cytometry; B lymphocytes; B cell leukaemias; CD19 and CD20 antigens

It is well established that immunophenotyping is a powerful tool in the classification of lymphoid disorders, by confirming a suspected diagnosis and demonstrating a characteristic phenotypic profile. However, there is sometimes an overlap of antigen expression in the various B cell disorders, such as CD5 positivity in mantle cell lymphoma (MCL), and chronic lymphocytic leukaemia (CLL). It is also known that some antigens are expressed at different densities in leukaemias, as compared with normal B cells. As a result, quantitative flow cytometry can be employed to distinguish between leukaemic and normal cells, and to some extent to differentiate between the various B cell disorders. We have previously shown that quantification by flow cytometric techniques can disclose differences in CD3 and CD7 expression between normal and leukaemic T cells and among the various T cell leukaemias.

In the context of immature B cell malignancies, quantitative flow cytometry has been used successfully to distinguish between leukaemic blasts and normal B cell precursors. Most previous studies on B lymphoproliferative disorders have focused on the presence or absence of surface antigens on the leukaemic cells, a feature designated “asynchronous phenotype.” Few have considered antigen density as an important biological characteristic of the neoplastic cells, for example low membrane immunoglobulins and CD20 in CLL, or high CD5 and low CD19 density in CLL cells compared with CD5 positive normal B lymphocytes.

We have applied a quantitative flow cytometry method to evaluate the expression of two B cell antigens, CD19 and CD20, in a variety of B cell malignancies in order (1) to compare the expression of these antigens in normal and leukaemic B lymphocytes; (2) to identify any abnormal patterns of antigen expression resulting from the malignant status; and (3) to investigate whether these changes may be useful for differential diagnosis.

Methods

Peripheral blood samples from 70 patients with a B cell disorder were obtained at diagnosis. The leukaemias included: CLL, 23 cases; B cell prolymphocytic leukaemia (B-PLL), 12 cases; hairy cell leukaemia (HCL), 13 cases; splenic lymphoma with villous lymphocytes (SLVL), 13 cases; and mantle cell lymphoma (MCL), nine cases. The diagnosis was based on clinical features, cell morphology, and immunophenotyping. All leukaemias expressed the B cell markers CD19 and CD20; CD5 was expressed in all CLL and MCL cases, whereas CD25 was consistently positive only in HCL.
Peripheral blood lymphocytes from 17 healthy donors were used as controls to establish reference ranges.

The mononuclear cells tested were derived from cryopreserved samples isolated from heparinised peripheral blood by 1.077 density gradient centrifugation with Lymphoprep (Nycomed, Oslo, Norway) from both patients and controls. Cell viability was assessed by trypan blue staining, which was more than 80% in all samples.

**IMMUNOSTAINING, FLOW CYTOMETRY ANALYSIS, AND QUANTIFICATION**

Direct immunofluorescence staining was performed with phycoerythrin conjugated CD19 (B4-RD1) and CD20 (B1-RD1) monoclonal antibodies (Coulter Immunology, Hialeah, Florida, USA). The mononuclear cells were prepared for flow cytometry according to standard methods. Briefly, after washing three times with RPMI 1640 with 20% fetal calf serum (FCS), $1 \times 10^6$ cells were used per test. To each tube, 50 µl of 2% AB serum were added, followed by the saturating amount (10 µl) of monoclonal antibodies. Appropriate directly conjugated isotypic antibodies (MsIgG-RD1; Coulter Immunology) were used as non-specific staining control in all experiments.

The cells were vortexed and incubated for 15 minutes at room temperature, washed three times with phosphate buffered saline (PBS)/azide, and resuspended in 0.5 ml Isoton (Coulter Immunology). All studies were performed using a FACScan flow cytometer (Becton-Dickinson, Mountain View, California, USA) as previously described. The fluorescence intensity was measured with detectors and amplifiers set on a logarithmic scale. For each sample 10 000 mononuclear cells were acquired for list mode

**Table 1 Mean ABC (antibody binding capacity) values $\times 10^3$ in normal peripheral blood B lymphocytes and B lineage leukemias**

<table>
<thead>
<tr>
<th>Antigen</th>
<th>Normal B cells</th>
<th>CLL</th>
<th>PLL</th>
<th>MCL</th>
<th>SLVL</th>
<th>HCL</th>
</tr>
</thead>
<tbody>
<tr>
<td>CD19</td>
<td>22 (7)</td>
<td>13</td>
<td>16</td>
<td>10</td>
<td>15</td>
<td>38</td>
</tr>
<tr>
<td>(p value)*</td>
<td>&lt;0.001</td>
<td>&lt;0.05</td>
<td>&lt;0.001</td>
<td>&lt;0.05</td>
<td>&lt;0.001</td>
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</tr>
<tr>
<td>CD20</td>
<td>94 (16)</td>
<td>65</td>
<td>129 (47)</td>
<td>123 (51)</td>
<td>167 (72)</td>
<td>312 (110)</td>
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<tr>
<td>(p value)*</td>
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Values are mean (SD); *comparison with normal peripheral blood B lymphocytes. CLL, chronic lymphatic leukemia; HCL, hairy cell leukemia; MCL, mantle cell lymphoma; PLL, prolymphocytic leukemia; SLVL, splenic lymphoma with villous lymphocytes.

![Figure 1](http://jgp.bmj.com)  
**Figure 1** Overlays of single parameter histograms showing the difference in CD19 expression between normal peripheral blood B cells (displayed in black) and B cell leukemias. CD19 is lower than in normal B lymphocytes in all B lineage leukemias except HCL.
analysis by setting a live gate around the lymphocyte area in the forward scatter versus side scatter dot plot. Analysis was performed using Lysys II software (Becton Dickinson).

The mean fluorescence intensity (MFI) values of the positive cells were converted into number of molecules of antigen per cell or antibody binding capacity (ABC) by using the Quantum Simply Cellular (QSC) microbeads kit (Sigma, St Louis, Missouri, USA). This is a mixture of four microbead populations which differ by their incremental capacities to bind directly labelled mouse immunoglobulins. The accompanying software regresses the binding capacities of the microbeads against their corresponding peak channels. The regression curve permits quantitative estimates of the monoclonal antibody molecules bound to the target cells, that is, the number of molecules of antigen expressed per cell. Ten microlitres of each monoclonal antibody were added to 50 µl QSC beads. After one hour of incubation, the mixtures were washed and analysed according to the method used for lymphocytes and using the same instrument's settings.

STATISTICS
The QUICKCAL software (FCSC; Sigma) was used for estimation of ABCs. The ABC values of CD19 and CD20 antigens in normal B lymphocytes and leukaemic cells were compared using Student’s t test for independent variables.

Results
The mean (SD) numbers of CD19 and CD20 molecules/cell in normal peripheral blood B lymphocytes were 22 (7) and 94 (16) × 10^3 respectively.

In cells from all the B lineage leukaemias except HCL, the mean number of CD19 molecules/cell was significantly lower than it was in normal B lymphocytes; in HCL, however, the CD19 ABC values were greater than in normal B cells or cells from the other B cell malignancies (table 1, figs 1 and 2).

The lowest CD19 density was found in MCL, followed by CLL, SLVL, and B-PLL, although no statistically significant differences were found between the mean values among these disorders. With a cut off point of 10 × 10^3 molecules/cell (the lowest normal value), five of nine MCL, seven of 13 SLVL, 10 of 23 CLL, and three of 12 B-PLL samples had values < 10, whereas only three cases had ABC values slightly over the normal range (fig 2). There were no differences between the 15 cases of typical CLL (12 (5) × 10^3 and eight cases of CLL with atypical morphology (16 (8) × 10^3). In contrast, a significantly larger mean number of CD19 molecules/cell was found in HCL than in normal B lymphocytes and the other B cell malignancies: eight of 13 HCL cases had ABC values greater than 31 × 10^3 (the highest normal value) and none of them fell below the normal range (fig 2).

A significantly higher mean number of CD20 molecules/cell was found in all B cell malignancies compared with normal B lymphocytes, with the exception of CLL, in which the mean ABC values was lower than in the other leukaemias and normal controls (table 1, figs 3 and 4). Only one CLL had a CD20 value within the normal range, whereas all other cases had CD20 ABCs below the normal range (fig 4). The number of CD20 molecules was lower in typical CLL (15 cases; 60 (5) × 10^3) than in CLL with atypical morphology (eight cases; 74 (13) × 10^3; p < 0.01). The levels in atypical CLL were still lower than in normal B cells and the other B cell leukaemias (p < 0.05). B-PLL, SLVL, and MCL showed comparable CD20 antigen densities, although the values in SLVL were slightly higher (table 1, figs 3 and 4). The highest CD20 density was observed in HCL, in which the mean ABC value was significantly greater than in all other B cell leukaemias, with all HCL cases showing more than 31 × 10^3 CD20 molecules/cell (the maximum normal value) (fig 4).

We have examined whether there was a correlation between the levels of CD19 and CD20 by the Spearman rank correlation test, but none was found. In addition, we have combined the results of CD19 and CD20 to see whether it has greater discriminatory power in distinguishing HCL from other B cell disorders, in particular SLVL, but no additional advantage was detected.

Discussion
CD19 and CD20 are widely expressed on normal and malignant B lymphocytes and thus they are not considered as discriminatory markers to distinguish among the different B cell malignancies. The development of techniques which allow the quantification of antigens by flow cytometry allowed us to quantify the levels of CD19 and CD20
antigens in a range of chronic B leukaemias. Comparing both CD19 and CD20 density in normal and malignant B cell populations, it is possible to characterise these cells more precisely.

The two molecules investigated here have a functional role on B cells. CD19 serves as a coreceptor which modulates B cell growth and differentiation by the induction of cell cycle arrest or programmed cell death. The expression of CD19 on B cells is highly conserved and, together with CD79a, is a feature of early B cell progenitors. CD19 expression increases with differentiation through pre-B and mature B cells, until it is downregulated at the later stages of B cell maturation, for example plasma cells. The CD19 ABC values of peripheral blood B lymphocytes obtained in our study are comparable to those previously reported in bone marrow mature B cells.

CD20 is a transmembrane phosphoprotein that functions as a calcium channel, and it has been shown to play an important role in B cell activation and differentiation. CD20 expression appears later than other B cell markers during normal B lymphocyte development and its membrane density progressively increases during differentiation. B cell precursors characteristically express a lower CD20 density than mature B cells. The lack or very low expression of CD20 is a known feature of B cell lineage ALL. In contrast, membrane CD20 increases either in vitro or in vivo during polyclonal B cell activation.

We have shown here that cells from most chronic B cell malignancies, except HCL, show low CD19 expression. In contrast, a weaker CD20 expression compared with normal B cells was seen only on CLL lymphocytes, whereas strong CD20 was detected in the other B cell leukaemias, the highest value also being found in HCL. Therefore, our analysis has shown that, in combination with other antigenic markers, CD19 and CD20 might be useful to characterise B lymphoproliferative disorders with similar cytology. For example, the distinction between HCL and SLVL can sometimes be difficult based on the cytology of peripheral blood films, especially when the neoplastic cells make up a minor proportion of total leucocytes. Our findings suggest that these conditions can be distinguished by the
CD19 and CD20 antigen density—both significantly higher in HCL (table 1)—together with other HCL associated markers.24 CLL and MCL are consistently positive for CD5 and may share some morphological findings in atypical cases.12 Here we have shown that the levels of CD20 are consistently lower in CLL compared to normal B and MCL cells. Thus bright staining with CD20 in a CD5 expressing B cell leukaemia would suggest the possibility of MCL rather than CLL. It is interesting that cases of CLL with atypical morphological features have higher values of CD20, as shown here and in another study.12 The differential diagnosis between MCL and atypical CLL may present a problem in some cases. However, the levels of CD20 in atypical CLL (74 (13) × 10⁷) were in general lower than in MCL (123 (51) × 10⁷).

The low CD19 ABC values and the faint intensity of CD20 in CLL, also observed by others,25 probably reflects an earlier stage of maturation of the B-CLL lymphocytes compared with normal blood B lymphocytes and other malignant B cells. Robbins et al described bright fluorescence for CD20 as a characteristic of HCL.27 Since HCL cells are considered very late and activated B cells which consistently express IL-2 receptor (CD25),25 26 it is possible that CD20 is upregulated in HCL cells after activation. Alternatively, the strong CD19 and CD20 expression in this disease could result from the expansion of a cell subset with high CD19 and CD20 antigen expression that is rarely represented in normal tissues. However, it is not certain whether the CD19 and CD20 ABC values in leukaemia reflect different stages of B cell development or represent abnormal phenotypes owing to neoplastic transformation.

This study shows that quantitative flow cytometry can detect differences in the levels of expression of the B cell markers CD19 and CD20 in B cell malignancies compared with normal B lymphocytes. Our results suggest that this approach may improve the discrimination between different disease entities and, in combination with morphology and other markers, enhance the accuracy in the diagnosis of B cell disorders.

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