

Remarkably similar antigen receptors among a subset of patients with chronic lymphocytic leukemia

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Studies of B cell antigen receptors (BCRs) expressed by leukemic lymphocytes from patients with B cell chronic lymphocytic leukemia (B-CLL) suggest that B lymphocytes with some level of BCR structural restriction become transformed. While analyzing rearranged V_HDJ_H and V_LJ_L genes of 25 non–IgM-producing B-CLL cases, we found five IgG+ cases that display strikingly similar BCRs (use of the same H- and L-chain V gene segments with unique, shared heavy chain third complementarity-determining region [HCDR3] and light chain third complementarity-determining region [LCDR3] motifs). These H- and L-chain characteristics were not identified in other B-CLL cases or in normal B lymphocytes whose sequences are available in the public databases. Three-dimensional modeling studies suggest that these BCRs could bind the same antigenic epitope. The structural features of the B-CLL BCRs resemble those of mAb's reactive with carbohydrate determinants of bacterial capsules or viral coats and with certain autoantigens. These findings suggest that the B lymphocytes that gave rise to these IgG^+ B-CLL cells were selected for this unique BCR structure. This selection could have occurred because the precursors of the B-CLL cells were chosen for their antigen-binding capabilities by antigen(s) of restricted nature and structure, or because the precursors derived from a B cell subpopulation with limited BCR heterogeneity, or both.

Introduction

B cell chronic lymphocytic leukemia (B-CLL), a monoclonal expansion of mature CD5-expressing B lymphocytes, is a heterogeneous disease that affects primarily individuals over 50 years of age (1). Even though B-CLL is the most common leukemia in the Western hemisphere (2), the events that select out an individual normal B cell clone and usher it toward leukemic transformation remain unknown. Genetic abnormalities probably exist in these cells and represent important inducers; however, no single unifying molecular genetic defect or combination of defects has yet been identified (3).

Studies of the characteristics of the B cell antigen receptors (BCRs) expressed by B-CLL cells imply that precursor B lymphocyte clones that eventually become leukemic exhibit varying degrees of BCR structural similarity (4). This restriction in BCR structure suggests that either the precursors of the leukemic B lymphocytes were selected by specific antigens that have affinity

Nonstandard abbreviations used: arsonate (Ars); B cell antigen receptor (BCR); B cell chronic lymphocytic leukemia (B-CLL); double-stranded DNA (dsDNA); framework region (FR); germinal center (GC); heavy chain third complementarity-determining region (HCDR3); light chain third complementarity-determining region (LCDR3); single-stranded DNA (ssDNA); web antibody modeling (WAM).

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for these BCRs, or they were garnered from a B cell subpopulation with restricted BCR structural heterogeneity.

In the present study, we analyzed the rearranged $V_H DJ_H$ and $V_L J_L$ genes of a cohort of 25 B-CLL patients whose leukemic cells express isotype-switched Ig. Our results reveal that a substantial subset of IgG+ cases (~20%) display strikingly similar Ig V region gene features. These include the use of the same H- and L-chain V gene segments, which are combined in unique ways and exhibit little somatic diversification despite their Ig class-switched nature. These findings are compelling evidence that selection of a specific BCR structure is an important component promoting the development of B-CLL. Preliminary abbreviated reports of these findings have appeared previously (5, 6).

Methods

CLL patients and samples. The Institutional Review Board of North Shore University Hospital (Manhasset, New York) and Long Island Jewish Medical Center (New Hyde Park, New York) approved these studies. From a cohort of 237 patients with clinical and laboratory features of B-CLL, 25 patients with expansions of CD5+/CD19+ B cells expressing surface membrane IgG or IgA were chosen and analyzed. All of the patients with surface membrane IgM+ cells were obtained randomly; some of the IgG+ cases were provided by others because of their surface membrane phenotype and therefore were not randomly acquired. Some patients and the V gene



sequences of their leukemic cells were described previously (5–9). PBMCs from these patients, obtained from heparinized blood by density gradient centrifugation (Ficoll-Paque; Amersham Biosciences, Piscataway, New Jersey, USA), were used after thawing samples that had been cryopreserved with a programmable cell-freezing machine (CryoMed, Inc., Mt. Clemens, Michigan, USA).

Isolation of DNA. T lymphocytes were purified from PBMCs by negative selection using the Pan T cell isolation kit (Miltenyi Biotec, Auburn, California, USA), and DNA was isolated from these cells with the DNeasy Tissue Kit (QIAGEN Inc., Valencia, California, USA).

Preparation of RNA and synthesis of cDNA. Total RNA was isolated from PBMCs using Ultraspec RNA (Biotecx Laboratories Inc., Houston, Texas, USA) according to the manufacturer's instructions. RNA (1 μg) was reverse-transcribed to cDNA using 200 U of Moloney murine leukemia virus reverse transcriptase (Invitrogen Corp., Carlsbad, California, USA), 1 U of RNase inhibitor (Eppendorf, Hamburg, Germany), and 20 pmol of oligo dT primer (total volume of 20 μl). These reactants were incubated at 42 °C for 1 hour, heated at 65 °C for 10 minutes to stop the reactions, and then diluted to a final volume of 100 μl .

PCR conditions for IgV gene DNA sequencing. To determine the IgV_H gene family used by various B-CLL cells, cDNA (2 μ l) was amplified using sense framework region 1 (FR1) primers specific for the various IgV_H gene families in conjunction with an appropriate antisense IgC_H primer (10). Reactions were carried out in 50 μ l using 20 pmol of each primer and cycled with a 9600 GeneAmp System (Perkin-Elmer Cetus, Emeryville, California, USA).

The DNA sequence of the B-CLL IgV_H gene was determined by reamplifying the original cDNA (2 µl) using the appropriate IgV_H family leader and IgC_H primers defined above (10). PCR products were sequenced directly after purification with Wizard PCR Preps (Promega Corp., Madison, Wisconsin, USA) using an automated sequencer (Applied Biosystems, Foster City, California, USA). In some instances where mutations were detected, an independent PCR product was generated and either sequenced directly or cloned into TA vector (Invitrogen Corp.), processed using Wizard minipreps (Promega Corp.), and then sequenced using M13 forward and reverse primers.

To determine IgV_L gene sequences, cDNA (2 μ l) was amplified using the leader primers listed in Supplemental Table 1 (supplemental material available at http://www.jci.org/cgi/content/full/113/7/1008/DC1). For the V λ families I, III, and IV, a mixture of primers was used. To amplify V λ families II and X, both forward primers were used in a common reaction, since they cross-prime. Reactions were carried out in a 50- μ l volume using 20 pmol of each primer, 200 μ M dNTPs, 1.5 mM MgCl₂, and 1.25 U of Taq Gold (Perkin-Elmer), and cycled with a 9600 GeneAmp System as follows: denaturation at 94°C for 45 seconds, annealing at 62°C for 30 seconds, and extension at 72°C for 45 seconds. After 35 cycles, extension was continued at 72°C for an additional 10 minutes. IgV_L PCR products were sequenced in the same manner as IgV_H PCR products.

To identify potential polymorphisms in the germline V_H4-39 gene, PCR was performed on DNA from autologous T cells of two patients (nos. 057 and 114) using V_H4-39 CDR2-specific gene primers (forward: 5'-GGTGGCGGCTCCCAGATG-3'; reverse: 5'-TCACACTCACCTCCCCTCAC-3'). PCR products were cloned and sequenced using the TOPO TA cloning kit (Invitrogen Corp.).

Analyses of V_H , D, and J_H sequences. DNA sequences were compared with those in the V BASE sequence directory (11) using MacVector software, version 6.0 (Accelrys, San Diego, California, USA) as previ-

ously described (8). Amino acid sequences were compared with those in GenBank by means of a BLAST search using the tblastn algorithm.

Analyses of heavy chain complementarity region 3 and light chain complementarity region 3 rearrangements. Heavy chain third complementarity-determining region (HCDR3) length was determined according to Kabat and Wu (12) by counting the number of amino acids between position H94 at the 3' end of FR3 (usually two amino acids downstream of the conserved cysteine) and position H102 at the beginning of FR4 (a conserved tryptophan in all J_H segments). Light chain third complementarity-determining region (LCDR3) length was determined by counting amino acids beginning at position L89 (preceded by a conserved cysteine) to position L97 (followed by a conserved Phe-Gly pair). Hypervariable loops were defined according to Chothia and Lesk (13); in particular, the third hypervariable loop of the H chain (H3) spans the amino acids 92–104, in the Kabat et al. numbering scheme (14).

Antibody modeling. Three-dimensional models of the V domains of the Ig's were constructed using the canonical structure model (14) as implemented in the web antibody modeling (WAM) algorithm (15). Models were analyzed using the molecular graphics package Insight II (16).

Results

Identification of five IgG-expressing B-CLL cases with remarkably similar BCR. While determining cDNA sequences of $V_H DJ_H$ and $V_L J_L$ rearrangements expressed in 25 isotype-switched B-CLL samples (23 IgG+ and 2 IgA+, see Supplemental Table 2), we identified five IgG-expressing cases with remarkably similar BCRs. These cases (CLL nos. 039, 057, 114, 202, and 209) expressed the same V_H [4–39], D [6–13], and J_H (5b) gene segments (Table 1). For the four cases in which L-chain data were obtained (additional sample on CLL no. 209 was not available because the patient died in an automobile accident), all used the $V_KO12/2$ gene (Table 1). In cases 057, 114, and 202, this V_L gene recombined with the J_KI gene segment; in case no. 039, the $V_KO12/2$ gene was associated with J_K2 . However, the expressed L chains of these four cases, including CLL no. 039, were virtually identical at the amino acid level (see below).

These five patients had several clinical similarities. The subset comprised primarily women with a 1:4 male/female ratio (Table 1), which differs from that in the other isotype-switched cases in this cohort (14:6; see Supplemental Table 2) and in B-CLL cases in general (~2:1 [refs. 1, 2]). Moreover, the patients experienced aggressive clinical courses complicated by severe recurrent infections (no. 039), Richter's transformation (no. 057), or the occurrence of second solid tumors (nos. 114 and 202). There was no familial or consistent ethnic relationship among these cases, and the patients originated from different parts of the world (two from the United States, one from Italy, one from the Caribbean region, and one from Japan).

The majority of the other 20 isotype-switched cases used either a V_H3 family gene (50%) or a V_H4 family gene (35%; see Supplemental Table 2). All of these V_H4-expressing cases used the V_H4-34 gene. One case (CLL no. 097) used the V_L012/2 gene, although this V_K gene was mutated and was paired with a mutated V_H3-73 gene. Thirteen of the remaining 20 cases expressed V_K, and six expressed V_A genes, with no apparent V_L segment biases. All of the V_Y-expressing cases used a V_H3 gene; among these, only J λ 2 was used.

Ig V gene mutation status. Although the leukemic cells of these five cases synthesized IgG, their $V_{\rm H}$ and $V_{\rm L}$ genes showed minimal deviation from the germline gene sequence ($V_{\rm H}$: 0.3–0.7%, and $V_{\rm L}$



Table 1Gene segments expressed by subset of IgG+ B-CLL cases

CLL no.	Gender	V gene family	Most similar germline V gene	Percent V gene difference from germline	D segment	J gene
039	Female	V_H4	4–39	0.7	6-13	5b
		Vĸ1	012/2	0.0	_	κ2
057	Female	V_H4	4–39	0.3 ^A	6-13	5b
		Vĸ1	012/2	0.0	_	κ1
114	Male	V_H4	4–39	0.0 ^A	6-13	5b
		Vĸ1	012/2	0.7	_	κ1
202	Female	V_H4	4–39	0.3	6-13	5b
		Vĸ1	012/2	0.0	_	κ1
209	Female	V_H4	4–39	0.3	6-13	5b
		NA	NA	NA	NA	NA

ABased on comparison with patient's genomic (T lymphocyte) germline V_H4 –39 genes, which differ at one (CLL 057) and three (CLL 114) positions from the canonical germline V_H4 –39 gene in GenBank. NA, not available.

0.0–0.7%; Table 1). In contrast, among the other 20 isotypeswitched patients, only four (nos. 040, 097, 158, and 185) expressed both $V_{\rm H}$ and $V_{\rm L}$ genes that differed by 1.0% or less from the germline counterparts (Supplemental Table 2).

To determine if some of these nucleotide differences represented allelic polymorphisms, we sequenced the V_H segments from the genomic (T cell) DNA of two patients (nos. 057 and 114; see Supplemental Figure 3). The germline V_H4-39 gene sequence for patient no. 057 exhibited one difference from the canonical germline V_H4 –39 deposited in GenBank, and the germline gene of patient 114 differed at three positions from the GenBank sequence. These findings indicate that some of the nucleotide differences we detected were allelic polymorphisms (Supplemental Figure 3), whereas others were real somatic mutations. A recent communication suggests that most of the Ig V_H sequence differences detected in "unmutated" B-CLL cells are somatically attained (17). A lack of somatic mutations in V_H4-39 genes from patients with predominantly IgM+ B-CLL has been reported previously (8, 18). Supplemental Figure 1, A and B, compares the V_H and V_L genes of the leukemic cells with the GenBank germline counterpart.

Protein sequences of IgV_H and IgV_L . Since most of the nucleotide differences in V_H and V_L were silent, the deduced V_H and V_L protein sequences of the five IgG^+ cases were virtually identical to that predicted from their germline genes (Supplemental Figure 2, A and B).

Figure 1

Amino acid sequences of HCDR3 and LCDR3 of IgG^+ B-CLL cases. Amino acid sequences are shown flanked by the 3′ end of FR3 and the 5′ end of FR4. A consensus sequence is shown for each. Color code: identical amino acids are in brown, chemically similar amino acids in blue, unrelated amino acids in black, and differences within areas of identity or similarity in red. (A) HCDR3. The consensus sequence consists of two hydrophilic amino acids (blue), seven amino acids that represent a portion of the germline D6-13 gene segment (brown), a glycine or serine (both with small side chains; brown), one to two variable amino acids, and five amino acids from the 5′ portion of the germline J_H5b segment. (B) LCDR3. The LCDR3 sequences of all cases are identical, except for a difference in the last amino acid in CLL no. 039, due to the use of the $J\kappa2$ segment. In each case, an arginine occurs at the $V\kappa$ - $J\kappa$ junction (underlined and in italics).

For V_H , there were only two amino acid changes among the five cases. CLL no. 057 exhibited a Pro \rightarrow Thr change at position H63, and no. 039 displayed an Arg \rightarrow Ser change at position H96; these isolated changes occurred in FR3. The V_L protein sequences were also virtually identical; only CLL no. 114 showed a Gln \rightarrow Arg change at position L3 in FR1.

HCDR3 structure. Figure 1A aligns the HCDR3 protein sequences for these five IgG-expressing cases. HCDR3 lengths were very similar (16–17 amino acids). These lengths are longer than most normal B cells (19) and many B-CLL cells (8, 20). The HCDR3 sequences of the five cases were very similar, exhibiting a consensus sequence (XXGYSSSWYG/SX(X)NWFDP; Figure 1A) consisting of two N-terminal hydrophilic amino acids, seven invariant amino acids that represent a portion of the D6–13 gene segment read in the hydrophilic reading frame, a glycine or serine that are similar in their small side chains, one to two variable amino acid(s) that lack chemical similarity, and five amino

acids corresponding to the 5' portion of the J_H5b segment.

LCDR3 structure. The LCDR3 sequences of all cases were identical (Figure 1B), except for a difference in the last amino acid in CLL no. 039; this difference results from the use of the $J\kappa 2$ segment in this instance. In each case, an arginine (R) was located at the Vκ-Jκ junction (position L96). In CLL no. 114 (Figure 2A), CGG, one of the six possibilities that code for arginine, resulted from coding end trimming and recombination of the Vκ and Jκ germline segments. For cases no. 057 and 202 (Figure 2B), the codon CGA yielded the arginine via a process similar to that for CLL no. 114. However, for CLL no. 039, which used the $J\kappa 2$ gene segment (Figure 2C), the combinatorial process that led to the arginine at the Vκ-Jκ junction was more complex. This required the trimming of one nucleotide from the 3' end of V_L and the deletion of seven nucleotides from the 5' end of $J\kappa 2$, along with the nontemplated insertion of two G nucleotides, to generate an arginine codon (CGG).

Lack of similar rearranged Ig V genes in other human B cells. We searched GenBank for $V_H DJ_H$ rearrangements with significant amino acid similarity to the consensus protein sequences of these five B-CLL

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Α
HCDR3
CLL 039:
              (FR3)...CAS
                           SRGYSSSWWS SNWFDP
                                                    WG... (FR4)
CLL 057:
              (FR3) ... CAR
                           HLGYSSSWYGAANWFDP
                                                    WG... (FR4)
CLL 114:
              (FR3) ... CAR
                           RFGYSSSWYG LDWFDP
                                                    WG... (FR4)
CLL 202:
                           STGYSSSWYSWRNWFDP
              (FR3) ... CAR
                                                    WG... (FR4)
CLL 209:
              (FR3) ... CAR
                           OAGYSSSWYGPSNWFDP
                                                    WG... (FR4)
Consensus: (FR3)...CAR
                           XXGYSSSWYG/SXXNWFDP WG... (FR4)
LCDR3
CLL 039:
              (FR3) ... YYC
                             QQSYSTPRS
                                            FGQ... (FR4)
CLL 057:
              (FR3)...YYC
                             OOSYSTPRT
                                            FGO... (FR4)
CLL 114:
              (FR3) ...YYC
                                            FGQ... (FR4))
CLL 202:
              (FR3) ... YYC
                              OOSYSTPRT
                                            FGO... (FR4)
Consensus: (FR3) ... YYC
                              QQSYSTPRT
                                            FGQ... (FR4)
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Arginine codons
                               CGC
                                      CGA
                       AGG
                               CGG CGT
                                Delete
                              C GT GG
VK012/2: ... ACC CCT C
                                           ACG ... Jκ1
CLL 114: ... ACC CCT C
                                       GG
                                            ACG ... Jκ1
                          └─ Arginine codon ─
В
VK012/2: ... ACC CCT C
                                  GT GG
                                            ACG
CLL 057: ... ACC CCT C
                                       GA
                                            \texttt{ACG} \ \dots \ \texttt{J} \kappa \texttt{1}
CLL 202: ... ACC CCT C
                                       GA
                                            \texttt{ACG} \ \dots \ \texttt{J} \kappa 1
                          L Arginine codon L
                                    Delete
VK012/2: ... ACC CCT C | C TG GG TAC
                                                   ACT ... Jκ2
CLL 039: ... ACC CCT C
                                                GG AGT ... JK2

    Arginine codon
```

cases. Only one very similar rearrangement was found, and this was from a lymphoplasmacytoid immunocytoma (GenBank entry no. Y09249 [ref. 21]); however, the corresponding L chain in this case was of the λ isotype (GenBank entry no. Y09250 [ref. 22]).

We also searched for rearranged $V\kappa O12/2$ -Jk L chains of the same LCDR3 length that have an arginine at the V_{L} -J_L junction. Because the V region of an L chain comprises only two gene segments and because six codons can yield an arginine, the likelihood of identifying rearrangements with similar structure is higher than that for a rearranged $V_H DJ_H$. BLAST search revealed 15 close matches (100% to 96.5% similarity) to the $V\kappa O12/2$ -Jk consensus amino acid sequence of the five IgG^+ B-CLL cases (Table 2). For 11 of these 15 cases, the companion $V_H DJ_H$ segments were available; none of these was $V_H 4-39$. Of the $V\kappa O12/2$ -Jk cases with known companion $V_H DJ_H$ segments, eight were of defined antigen specificity. Remarkably, of these eight cases, seven were from autoreactive mAb's (anti-IgG, anti-RhD, anti-IgG, anti-RhD, anti-IgG, anti-RhD, anti-IgG, anti-IgG, and antiapoptotic

Figure 2

Different mechanisms generate an arginine at the V_L - J_L junctions. The codons for arginine are listed. Color code: identical nucleotides of the arginine codon are in brown, nucleotides deleted are in black, nucleotides probably resulting from nontemplated nucleotide addition are in dark blue, and differences within areas of identity or similarity in red. (**A** and **B**) An appropriate codon results from coding end trimming and recombination of the two germline $V\kappa$ and $J\kappa$ 1 segments. (**C**) The arginine codon develops via a more complex combinatorial process, requiring the trimming of one nucleotide from the 3' end of V_L , the deletion of seven nucleotides from the 5' end of $J\kappa$ 2, and the nontemplated insertion of two G nucleotides.

cells); the other bound the capsular polysaccharide of *Neisseria meningitidis* (Table 2). One of the $V\kappa O12/2$ -J κ rearrangements of unknown specificity was from a B-CLL clone (GenBank entry no. AF228327); however, this was paired with a V_H4 –34 gene (GenBank entry no. AF196469 [ref. 23]).

We next searched the Kabat, et al. database (14) and identified 2,950 mAb's for which both H and L Ig V genes were known. Among these, we found 199 with an arginine at L96. There was no statistically significant correlation between the presence of an arginine at the V_L -J_L junction and a specific, unique antigenic reactivity. However, there was a striking enrichment (102/199) for antibodies that react with structures that serve as autoantigens in several autoimmune settings; examples are double-stranded (ds) and single-stranded (ss) DNA, IgG, and thyroid constituents. This was consistent with the autoreactivity identified in the $V\kappa O12/2$ -JK V_L -I_L listed in Table 2.

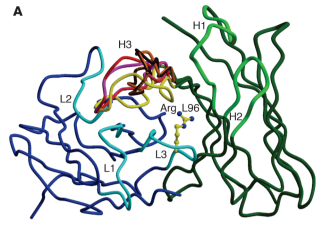
Three-dimensional models of the B-CLL BCR. We used the canonical structure method (13) as implemented in the WAM algorithm to build models for the V domains, and then analyzed these structures to deduce characteristics of the antigen-binding sites (Figure 3). The canonical structure model requires that the main chain con-

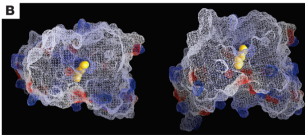
Table 2012/2-Jk Gene rearrangements, with similar LCDR3 lengths and junctional arginine (R) from other human B cells

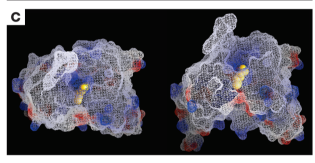
GenBank accession no.	Percent amino acid difference from consensus <i>012/2</i> -Jrc ^A	Jκ	V _H Gene	Source
Consensus	_	1	4–39	IgG+ B-CLL cases
BAC01727	0.0%	1	NA	Antibody library from human B cells (Y. Akahori, et al., unpublished data)
CAA51109	0.0%	1	NA	Phage library using cDNA from human splenic B cells (94)
CAD68160	0.9% (1 AA)	1	NA	Antibody reactive with apoptotic cells (R. Gandhi, et al., unpublished data)
1HEZA	0.9% (1 AA)	1	9.1	κ chain of rheumatoid factor bound by κ -binding protein (95)
AAL65713	0.9% (1 AA)	2	1–46	Hybridoma from human tonsil B lymphocytes (M. Vaisbourd, et al., unpublished data)
CAA85588	0.9% (1 AA)	1	NA	cDNA library from normal blood B cells (22)
CAD43018	0.9% (2 AA)	4	3–21	
CAD43019	1.8% (2 AA)	4	3–21	Human anti-La autoantibodies from phage display library (J.M.H. Raats, unpublished data)
CAD43020	1.8% (2 AA)	4	3–21	
CAC28926	0.9% (2 AA)	2	1–02	Human IgG anti- α (IIb) β 3 antibody from phage-displayed library (96)
AAD19446	1.8% (2 AA)	1	1–24	cDNA library from normal blood B cells (97)
1DEEA	1.8% (2 AA)	1	3–30	Human monoclonal rheumatoid factor (98)
AAN87123	1.8% (2 AA)	4	9.1	Human IgM antibody to capsular polysaccharide of Neisseria meningitidis (63)
AAC13457	2.6% (3 AA)	4	<i>3–33</i>	Phage-displayed human anti-Rh(D) antibody (99)
AAF86916	3.5% (4 AA)	2	4–34	B-CLL cell (23)

[^]Denotes percent and number of amino acid differences from consensus V κO12/2-J κ sequence of IgG+B-CLL cases.









formation of the hypervariable loops depend solely on length and on the nature of a few specific residues. Antibody specificity is therefore determined by the nature of exposed side chains mounted on the main chain of the hypervariable loops, which in turn is determined by their canonical structure.

In the four available B-CLL L chains (case nos. 039, 057, 114, and 202), the L1 loop is six residues long and contains a conserved isoleucine at position 29. In all known antibody structures, sixresidue L1 loops are stabilized by contact of the side chain of this amino acid with residues 2, 25, 33, and 71 (13). These contacts determine the main chain structure of the loop. Each of these five residues is hydrophobic and conserved in the four B-CLL rearranged L chains. Since the exposed amino acids of the loop are also conserved, the contribution of this loop to the antigen-binding site would be virtually identical for all four antibodies. The L2 loop has the same conformation in all known V region structures (13). Since there are no substitutions in the exposed side chains of these cases, the contribution of this loop to the binding site is expected to be the same for all the proteins. Similarly, since the backbone structure of the L3 loop depends on its length and the nature of the residues at positions 90 and 95, which are all identical in the B-CLL sequences, this loop should have the same conformation and solvent-exposed surface in all four cases.

Figure 3

C-alpha trace of the structural models produced by WAM. (A) Overlay of main chains of the five CLL three-dimensional structures. For CLL no. 209, the consensus L-chain amino acid sequence was added for WAM analysis. B-CLL L chains are shown in blue color (hypervariable loops L1, L2, and L3 in light blue), and H chains are shown in green color (hypervariable loops H1, and H2 in light green). The different H3 loop structures are color-coded, as noted by the lime green, red, black, purple, and dark green lines in the H3 area. Figure was prepared with MOLSCRIPT (91), and Raster3D (92). (B and C) Electrostatic and molecular surface representation of the structure of CLL no. 039 as a model representative of the B-CLL cases of nonbulged conformation (nos. 039 and 114; B) and of the CLL no. 057 as a model of the cases of bulged conformation (nos. 057, 202, and 209; C). Negative surface potentials are indicated in red, positive surface potentials in blue, and neutral potentials in white. The arginine L96 side chain is shown in yellow CPK representation. The structures are shown from a top view (left panels) and from a side view, corresponding to an approximately 70° rotation relative to the top view (right panels). Figures were prepared with GRASP (93). Arg L96; arginine L96 side chain.

The H1 loop has a conserved hydrophobic residue in position 29 and a conserved glycine in position 26. The former establishes stabilizing interactions with residues 34, 72, and 77. Each of these five residues is conserved among the $V_{\rm H}$ sequences of the five B-CLL cases. Consistent with these observations, the WAM prediction server produces identical models in this region for the five antibodies. The H2 conformation has been shown to depend only on its length and on the residue at position 71 (24); both of these features are conserved in the five B-CLL sequences.

One cannot infer as clear a sequence–structure relationship for H3, since this loop is the most variable in length and sequence. The conformation of H3 is mainly determined by the presence or absence of a β bulge within its structure, in the region closer to the FR. The presence of a β bulge is in turn determined by the amino acid sequence of the loop (25). Based on our modeling analyses, the H3 loop is predicted not to be bulged in CLL nos. 039 and 114 and bulged at residue H101 in CLL nos. 057, 202, and 209. The creation of the bulge in the latter cases implies that residue H101m of the nonbulged loops is in a position equivalent to that of residue H101 of the bulged loops. Thus, the tip of the loop probably contains the same number of residues (n = 14) in all cases, and the residues in the bulged cases can maintain a relationship in space similar to that of the non-bulged cases.

Discussion

In this study we describe five IgG-expressing B-CLL cases with remarkably similar BCRs. These receptors consist of the same $V_{\rm H},$ D, $J_{\rm H},$ and $V_{\rm L},$ J_L gene segments, except for a different JK segment in one instance. Despite this difference, all BCRs have identical LCDR3s and very similar HCDR3s, each with unique sequence and junctional motifs. This BCR restriction could be the consequence of either random transformation of a subpopulation of B cells with very limited antigen receptor heterogeneity — determined genetically or by antigen selection; or specific transformation of B cells that were selected by antigen from a BCR-restricted or BCR-heterogeneous subpopulation; or both. Whatever the cause, our findings support the concept that B-CLL develops from a limited set of B lymphocytes of defined BCR structure and imply that selection of B cells with such structures represents an important promoting influence in the evolution of the leukemic cells.



Composition, motifs, and three-dimensional structure models. Developing B lymphocytes do not use all germline IgV gene segments with the same frequency, and biases in gene segment rearrangement do occur (26-29). Nevertheless, considering combinatorial diversity, imprecise joining, nucleotide insertion and deletion, and somatic hypermutation, the probability of finding selectively in one disease, by chance alone, five cell clones with such highly similar rearranged H- and L-chain V region pairs is extremely low. The probability that specific V_H, D, and J_H genes would be used in the same V_HDJ_H rearrangement is 1 in 7,128 ($1/44 \times 1/27 \times 1/6$); for a specific V_L and J_L gene pair the probability is $1/230 (1/46 \times 1/5)$ or $1/252 (1/36 \times 1/7)$ for a κ versus a λ rearrangement, respectively. Only 1 in 1,639,440 B cells would be predicted to randomly express the same V_H, D, J_H, Vκ, and Jκ segments in its BCR. These calculations use the number of germline V segments in IgBLAST and assume that κ genes rearrange before λ genes.

Considering these estimations, the frequency at which this BCR occurs in our IgG+ B-CLL cases is extraordinary (~20%). Although this percentage will need confirmation, the frequency of such unique IgG+ cases is very similar to that identified when we first reported on Ig V gene diversification and apparent antigen selection in a smaller number of isotype-switched B-CLL cases (7). This structure was not seen in our IgM+ B-CLL cohort (>175 cases for which both the rearranged $V_H DJ_H$ and $V_L J_L$ are known) or in GenBank, indicating that this BCR is not overexpressed among IgM+ B-CLL cases or in the normal circulating B cell repertoire. We cannot rule out that its frequency increases with age or that it exists among certain distinct noncirculating subpopulations of B cells, although a recent study suggests that the former may be unlikely (30).

Notwithstanding the present limitations of antibody modeling, primarily with respect to the H3 loop, we can reliably conclude that most of the binding site is identical among these IgG+ B-CLL cases. Five of the six H and L loops have the same main chain conformation and differ, in only one case, by a conservative Ser \rightarrow Thr side chain change at position 97 of the L chain. However, this residue does not contact antigen in any known structure (Veronica Morea, personal communication). The H3 loops are also quite similar among the five antibodies, although three are longer by one residue (Figure 1). Since these loops contain a bulge (i.e., they have one residue that is extruded from the regular β structure), the residues at the central region of the loop, which are more relevant for antigen binding, can maintain the same relationship in space in all five cases. Therefore, the antigen-binding surfaces of these antibodies are probably very similar, and they likely bind the same antigenic epitope. Nevertheless since some amino acid differences do exist at the V_H-D and D-J_H junctions, this remains conjecture.

Restricted IgV gene structural features of antibodies with defined antigenic reactivities. Normal and neoplastic B cells of known antigen specificity can display restricted V (D), J segment use, either at both H and L chain loci or individually at either locus. Murine mAb's reactive with β -(1,6)-D-galactan (31), α (1 \rightarrow 6) dextran (32, 33), phosphorylcholine (34), dextrans and fructofurans (35), and phosphatidylcholine (36–38) pair very restricted and characteristic V_HDJ_H and V_LJ_L gene segments. Human antibodies that exhibit individual H- or L-chain restrictions include mAb's specific for the capsular polysaccharides of *Haemophilus influenzae* type b (39–41) and *Streptococcus pneumoniae* (42, 43) (individual V_H3 genes for both), monoclonal cold agglutinins with anti-I specificity (exclusively V_H4–34 [refs. 44–46]), and monoclonal rheumatoid factors with IgG reactivity (V_H1–69 [refs. 47–51]). Murine anti-arsonate (52–54) and

anti-bacterial polysaccharide mAb's (55) frequently use specific individual $V\kappa$ and $J\kappa$ genes.

In addition, mAb's reactive with carbohydrates (30, 32, 40, 56), autoantigens (57, 58), and haptens (52–54) often have characteristic V_L - J_L junctional residues. Human anti–H. influenzae mAb's that use the $V\kappa A2$ gene segment contain an arginine at position L96, although chain recombination experiments suggest that it is not essential for antigen binding (40, 56). Murine mAb's reactive with arsonate (Ars), dsDNA, and β -(1,6)-D-galactan exhibit an arginine or an isoleucine at position L96, respectively. These junctional amino acids are essential for binding to Ars (53), but not to β -(1,6)-D-galactan (59).

Nature of the antigen that could have selected these BCRs. Our findings suggest that the B lymphocytes that gave rise to these IgG+ B-CLL cells were selected for a unique BCR structure. If this selection involved antigen binding and triggering through the BCR, the antigen(s) would most likely have been of restricted nature and structure. Although the identity of such an antigen is unknown, we can infer certain features based on our data.

First, the diverse geographic origins of our patients suggests that the putative structure would probably be distributed worldwide. Second, the epitope would appear to select out and drive B cells to undergo an isotype class switch, since we found this BCR only in non-IgM-expressing B-CLL cells. Third, this selection and drive would not frequently lead to clonal expansion of normal B cells, since we could not identify this BCR in B lymphocytes from normal individuals deposited in GenBank. Fourth, the presence of a positively charged center to the antigen-binding pocket, provided by the side chain of arginine L96, and surrounded by an aromatic area and a ridge of polar residues, would imply that a negatively charged or electron-rich group exists in the epitope interactive with these BCRs (Figure 3). This, however, is not a necessity (41, 53, 59). Of the 199 completely sequenced antibodies with an arginine at L96 (14), we found that about 50% react with autoantigens. This is consistent with the association with autoreactivity in the VKO12/ 2-Jκ rearrangements listed in Table 2. Autoantigens can be negatively charged molecules (e.g., DNA), and their reactivity with positively charged residues in the CDRs of autoantibodies is enhanced as the number of these latter residues increases (58, 60). Finally, in light of the comparisons with antibodies of known specificities, the epitope could be a carbohydrate that is restricted to a unique molecule or shared by several molecules, an autoantigen that is shared by all individuals or polymorphic among populations, or a carbohydrate determinant of an autoantigen. The possibility that two classes of antigens can bind to the same binding site also should not be ruled out (61).

It is of interest that several relationships exist between carbohydrate reactivity and autoreactivity. Autoantibodies can react with carbohydrates (62–65) and can confer protection against infection with encapsulated bacteria (66, 67). In addition, antipneumococal polysaccharide antibodies can convert to anti-dsDNA reactivity after minimal amino acid changes in the Ig V region (68). In this regard, it is known that the BCRs of B-CLL clones can be autoreactive (69, 70) and can express cross-reactive idiotypes of dsDNA antibodies (71).

Importance of B cell precursors with constrained BCR structural diversity in the development and progression of B-CLL. Irrespective of the nature of the antigen(s) possibly involved in these cases, these five IgG+ clones are extraordinary examples of the principle that B-CLL progenitors are selected for certain limited BCR structures (4). Sim-

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ilar evidence for BCR restriction, possibly attained by antigen or superantigen selection, can be found among the IgM-expressing cases that use a $V_H 3-21$ gene with a restricted V λ partner (72, 73), and probably in those cases that use an unmutated $V_H 1-69$ gene with a characteristic HCDR3 (8, 20).

Although the expression of BCRs of class-switched isotype often suggest involvement in T cell-dependent responses, the lack of significant numbers of IgV gene mutations in B lymphocytes is unusual for T cell-mediated differentiation and classical germinal center (GC) passage, unless one invokes the possibilities that the germline sequence is preferred for antigen binding or that secondary rearrangements occurred at both the H- and L-chain loci. However, the lack of *IgV* gene mutations and the occurrence of a limited degree of isotype class switching also occur during B cell clonal expansion in the absence of T cell help and in response to T-independent antigens (74). Previous studies suggest that antigen-stimulated B cells that participate in a GC reaction lead to B cell lymphoproliferative disorders such as follicular cell lymphoma and Burkitt's lymphoma (75). Nevertheless, B cells that follow a different pathway of B cell activation – that is, one not involving a classical GC reaction (76, 77) - could develop into B-CLL cells in certain instances. The marginal zone is considered a possible site for such nonclassical GC reactions (74) and the development of B-CLL from marginal zone B cells has been suggested (4). This suggestion is intriguing because the marginal zone is enriched in B cells that react with carbohydrates (74) and autoantigens (78-80), and its Ig V gene repertoire, at least in rodents, is highly restricted (81, 82). This receptor restriction apparently occurs early in life and is based on BCR composition, specificity, and signal-transducing capacity (81).

Finally, since these and other B-CLL clones express BCRs with restricted antigen-binding sites, antigen drive could promote intraclonal evolution leading to accumulation of deleterious DNA mutations by members of the leukemic clone and subsequently to a more aggressive clinical course. Repetitive engagement of the BCRs by either autoantigens or foreign antigens, possibly from microorganisms that are encountered during intermittent or persistent infections, could elicit such effects. In this regard, the cases that have the most convincing molecular evidence for BCR restriction and antigen or superantigen selection (i.e., these IgG+ cases

and those expressing $V_H 3-21$ and $V_H 1-69$) experience more aggressive clinical courses with shortened mean survival times (72, 83, 84). In addition, ongoing B cell activation and differentiation can occur in B-CLL cells (10, 85-88) and may be initiated or augmented by BCR-mediated signal transduction that is preserved more often in those cases with the worst clinical outcomes (89, 90).

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